

(12) STANDARD PATENT
(19) AUSTRALIAN PATENT OFFICE

(11) Application No. **AU 2014259755 B2**

(54) Title
Compositions and methods for modulating apolipoprotein (a) expression

(51) International Patent Classification(s)
C07H 21/00 (2006.01) **C07K 14/775** (2006.01)
A61K 48/00 (2006.01) **C12N 15/113** (2010.01)

(21) Application No: **2014259755** (22) Date of Filing: **2014.05.01**

(87) WIPO No: **WO14/179625**

(30) Priority Data

(31) Number	(32) Date	(33) Country
61/871,673	2013.08.29	US
61/986,867	2014.04.30	US
61/843,887	2013.07.08	US
61/818,442	2013.05.01	US
61/823,826	2013.05.15	US
61/976,991	2014.04.08	US
61/880,790	2013.09.20	US

(43) Publication Date: **2014.11.06**

(44) Accepted Journal Date: **2018.08.30**

(71) Applicant(s)
Ionis Pharmaceuticals, Inc.

(72) Inventor(s)
Prakash, Thazha P.;Seth, Punit P.;Swayze, Eric E.;Graham, Mark J.

(74) Agent / Attorney
Pizzeys Patent and Trade Mark Attorneys Pty Ltd, PO Box 291, WODEN, ACT, 2606, AU

(56) Related Art
WO 2009/073809 A2



(51) International Patent Classification:

C07K 14/775 (2006.01) **A61K 48/00** (2006.01)
C12N 15/113 (2010.01)

(21) International Application Number:

PCT/US2014/036460

(22) International Filing Date:

1 May 2014 (01.05.2014)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

61/818,442	1 May 2013 (01.05.2013)	US
61/823,826	15 May 2013 (15.05.2013)	US
61/843,887	8 July 2013 (08.07.2013)	US
61/871,673	29 August 2013 (29.08.2013)	US
61/880,790	20 September 2013 (20.09.2013)	US
61/976,991	8 April 2014 (08.04.2014)	US
61/986,867	30 April 2014 (30.04.2014)	US

(71) **Applicant: ISIS PHARMACEUTICALS, INC.** [US/US];
2855 Gazelle Court, Carlsbad, CA 92010 (US).

(72) **Inventors: PRAKASH, Thazha, P.**; 2855 Gazelle Court,
Carlsbad, CA 92010 (US). **SETH, Punit, P.**; 2855 Gazelle
Court, Carlsbad, CA 92010 (US). **SWAYZE, Eric, E.**;
2855 Gazelle Court, Carlsbad, CA 92010 (US). **GRA-**
HAM, Mark, J.; 2855 Gazelle Court, Carlsbad, CA 92010
(US).

(74) **Agents: LIANG, Teresa, Y.** et al.; Isis Pharmaceuticals,
Inc., 2855 Gazelle Court, Carlsbad, CA 92010 (US).

(81) **Designated States** (unless otherwise indicated, for every
kind of national protection available): AE, AG, AL, AM,
AO, AT, AU, AZ, BA, BB, BG, BH, BN, BR, BW, BY,
BZ, CA, CH, CL, CN, CO, CR, CU, CZ, DE, DK, DM,
DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT,
HN, HR, HU, ID, IL, IN, IR, IS, JP, KE, KG, KN, KP, KR,
KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME,
MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ,
OM, PA, PE, PG, PH, PL, PT, QA, RO, RS, RU, RW, SA,
SC, SD, SE, SG, SK, SL, SM, ST, SV, SY, TH, TJ, TM,
TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM,
ZW.

(84) **Designated States** (unless otherwise indicated, for every
kind of regional protection available): ARIPO (BW, GH,
GM, KE, LR, LS, MW, MZ, NA, RW, SD, SL, SZ, TZ,
UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, RU, TJ,
TM), European (AL, AT, BE, BG, CH, CY, CZ, DE, DK,
EE, ES, FI, FR, GB, GR, HR, HU, IE, IS, IT, LT, LU, LV,
MC, MK, MT, NL, NO, PL, PT, RO, RS, SE, SI, SK, SM,
TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW,
KM, ML, MR, NE, SN, TD, TG).

Published:

- with international search report (Art. 21(3))
- before the expiration of the time limit for amending the
claims and to be republished in the event of receipt of
amendments (Rule 48.2(h))
- with sequence listing part of description (Rule 5.2(a))

(54) **Title:** COMPOSITIONS AND METHODS FOR MODULATING APOLIPOPROTEIN (a) EXPRESSION

(57) **Abstract:** Provided herein are oligomeric compounds with conjugate groups targeting apolipoprotein (a) [apo(a)]. In certain embodiments, the apo(a) targeting oligomeric compounds are conjugated to *N*-Acetylgalactosamine. Also disclosed herein are conjugated oligomeric compounds targeting apo(a) for use in decreasing apo(a) to treat, prevent, or ameliorate diseases, disorders or conditions related to apo(a) and/or Lp(a). Certain diseases, disorders or conditions related to apo(a) and/or Lp(a) include inflammatory, cardiovascular and/or metabolic diseases, disorders or conditions. The conjugated oligomeric compounds disclosed herein can be used to treat such diseases, disorders or conditions in an individual in need thereof.



COMPOSITIONS AND METHODS FOR MODULATING APOLIPOPROTEIN (a) EXPRESSION**SEQUENCE LISTING**

The present application is being filed along with a Sequence Listing in electronic format. The Sequence Listing is provided as a file entitled BIOL0250WOSEQ_ST25.txt, created on May 1, 2014, which is 432 Kb in size. The information in the electronic format of the sequence listing is incorporated herein by reference in its entirety.

BACKGROUND OF THE INVENTION

The principle behind antisense technology is that an antisense compound hybridizes to a target nucleic acid and modulates the amount, activity, and/or function of the target nucleic acid. For example in certain instances, antisense compounds result in altered transcription or translation of a target. Such modulation of expression can be achieved by, for example, target mRNA degradation or occupancy-based inhibition. An example of modulation of RNA target function by degradation is RNase H-based degradation of the target RNA upon hybridization with a DNA-like antisense compound. Another example of modulation of gene expression by target degradation is RNA interference (RNAi). RNAi refers to antisense-mediated gene silencing through a mechanism that utilizes the RNA-induced silencing complex (RISC). An additional example of modulation of RNA target function is by an occupancy-based mechanism such as is employed naturally by microRNA. MicroRNAs are small non-coding RNAs that regulate the expression of protein-coding RNAs. The binding of an antisense compound to a microRNA prevents that microRNA from binding to its messenger RNA targets, and thus interferes with the function of the microRNA. MicroRNA mimics can enhance native microRNA function. Certain antisense compounds alter splicing of pre-mRNA. Regardless of the specific mechanism, sequence-specificity makes antisense compounds attractive as tools for target validation and gene functionalization, as well as therapeutics to selectively modulate the expression of genes involved in the pathogenesis of diseases.

Antisense technology is an effective means for modulating the expression of one or more specific gene products and can therefore prove to be uniquely useful in a number of therapeutic, diagnostic, and research applications. Chemically modified nucleosides may be incorporated into antisense compounds to enhance one or more properties, such as nuclease resistance, pharmacokinetics or affinity for a target nucleic acid. In 1998, the antisense compound, Vitravene® (fomivirsen; developed by Isis Pharmaceuticals Inc., Carlsbad, CA) was the first antisense drug to achieve marketing clearance from the U.S. Food and Drug Administration (FDA), and is currently a treatment of cytomegalovirus (CMV)-induced retinitis in AIDS patients.

New chemical modifications have improved the potency and efficacy of antisense compounds, uncovering the potential for oral delivery as well as enhancing subcutaneous administration, decreasing

potential for side effects, and leading to improvements in patient convenience. Chemical modifications increasing potency of antisense compounds allow administration of lower doses, which reduces the potential for toxicity, as well as decreasing overall cost of therapy. Modifications increasing the resistance to degradation result in slower clearance from the body, allowing for less frequent dosing. Different types of chemical modifications can be combined in one compound to further optimize the compound's efficacy.

Lipoproteins are globular, micelle-like particles that consist of a non-polar core of acylglycerols and cholesteryl esters surrounded by an amphiphilic coating of protein, phospholipid and cholesterol. Lipoproteins have been classified into five broad categories on the basis of their functional and physical properties: chylomicrons, very low density lipoproteins (VLDL), intermediate density lipoproteins (IDL), low density lipoproteins (LDL), and high density lipoproteins (HDL). Chylomicrons transport dietary lipids from intestine to tissues. VLDLs, IDLs and LDLs all transport triacylglycerols and cholesterol from the liver to tissues. HDLs transport endogenous cholesterol from tissues to the liver

Lipoprotein particles undergo continuous metabolic processing and have variable properties and compositions. Lipoprotein densities increase without increasing particle diameter because the density of their outer coatings is less than that of the inner core. The protein components of lipoproteins are known as apolipoproteins. At least nine apolipoproteins are distributed in significant amounts among the various human lipoproteins.

The lipoprotein(a) [Lp(a)] particle was identified nearly 50 years ago and is comprised of a highly unique LDL particle in which one apolipoprotein B (apoB) protein is linked via a disulfide bond to a single apolipoprotein(a) [apo(a)] protein. The apo(a) protein shares a high degree of homology with plasminogen particularly within the kringle IV type 2 repetitive domain. Levels of circulating Lp(a) are inversely proportional to the number of kringle IV type 2 variable repeats present in the molecule and, as both alleles are co-expressed within individuals, can display heterozygous plasma isoform profiles (Kraft et al., Eur J Hum Genet, 1996; 4(2): 74-87). It is thought that this kringle repeat domain in apo(a) may be responsible for its pro-thrombotic and anti-fibrinolytic properties, potentially enhancing atherosclerotic progression.

Apo(a) is transcriptionally regulated by IL-6 and in studies in rheumatoid arthritis patients treated with an IL-6 inhibitor (tocilizumab), plasma levels were reduced by 30% after 3 month treatment (Schultz et al., PLoS One 2010; 5:e14328).

Apo(a) has been shown to preferentially bind oxidized phospholipids and potentiate vascular inflammation (Bergmark et al., J Lipid Res 2008; 49:2230–2239; Tsimikas et al., Circulation. 2009; 119(13):1711–1719).

Further, studies suggest that the Lp(a) particle may also stimulate endothelial permeability, induce plasminogen activator inhibitor type-1 expression and activate macrophage interleukin-8 secretion (Koschinsky and Marcovina, Curr Opin Lipidol 2004; 15:167–174). Importantly, recent genetic association studies revealed that Lp(a) was an independent risk factor for myocardial infarction, stroke, peripheral vascular disease and abdominal aortic aneurysm (Rifai et al., Clin Chem 2004; 50:1364–71; Erqou et al.,

JAMA 2009;302:412–23; Kamstrup et al., Circulation 2008;117:176–84). Further, in the recent Precocious Coronary Artery Disease (PROCARDIS) study, Clarke *et al.* (Clarke et al., NEJM (2009)361; 2518-2528) described robust and independent associations between coronary heart disease and plasma Lp(a) concentrations. Additionally, Solfrizzi et al., suggested that increased serum Lp(a) may be linked to an increased risk for Alzheimer's Disease (AD) (Solfrizzi et al., J Neurol Neurosurg Psychiatry 2002, 72:732-736. Currently, in the clinic setting, examples of indirect apo(a) inhibitors for treating cardiovascular disease include aspirin, Niaspan, Mipomersen, Anacetrapib, Epirotirome and Lomitapide which reduce plasma Lp(a) levels by 18%, 39%, 32%, 36%, 43% and 17%, respectively. Additionally, Lp(a) apheresis has been used in the clinic to reduce apo(a) containing Lp(a) particles.

To date, therapeutic strategies to treat cardiovascular disease by directly targeting apo(a) levels have been limited. Ribozyme oligonucleotides (U.S. Patent 5,877,022) and antisense oligonucleotides (WO 2005/000201; WO 2003/014397; WO2013/177468; US20040242516; U.S. Patent Nos. 8,138,328, 8,673,632 and 7,259,150; Merki et al., J Am Coll Cardiol 2011; 57:1611–1621; each publication incorporated by reference in its entirety) have been developed but none have been approved for commercial use.

Thus, there remains a clear unmet medical need for novel agents which can potently and selectively reduce apo(a) levels in patients at enhanced risk for cardiovascular events due to chronically elevated plasma Lp(a) levels.

SUMMARY OF THE INVENTION

Provided herein are compositions and methods for modulating expression of apo(a) mRNA and protein. In certain embodiments, the apo(a) specific inhibitor decreases expression of apo(a) mRNA and protein. Provided herein are compositions and methods for modulating expression of Lp(a) levels.

In certain embodiments, the composition is an apo(a) specific inhibitor. In certain embodiments, the apo(a) specific inhibitor is a nucleic acid, protein, or small molecule. In certain embodiments, the apo(a) specific inhibitor is an antisense oligonucleotide targeting apo(a) with a conjugate. In certain embodiments, the apo(a) specific inhibitor is a modified oligonucleotide and a conjugate, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and comprises a nucleobase sequence comprising a portion of at least 8 contiguous nucleobases complementary to an equal length portion of nucleobases 3901 to 3920 of SEQ ID NO: 1, wherein the nucleobase sequence of the modified oligonucleotide is at least 80% complementary to SEQ ID NO: 1. In certain embodiments, the apo(a) specific inhibitor is a modified oligonucleotide and a conjugate, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and has a nucleobase sequence comprising at least 8, least 9, least 10, least 11, at least 12, least 13, at least 14, at least 15, at least 16, least 17, least 18, least 19, or 20 contiguous nucleobases of the nucleobase sequence of SEQ ID NO: 1-130, 133, 134. In certain embodiments, the apo(a) specific inhibitor is a modified oligonucleotide and a conjugate, wherein the modified oligonucleotide consists of 20 linked nucleosides and

has a nucleobase sequence comprising at least 8 contiguous nucleobases of any of SEQ ID NO: 58, wherein the modified oligonucleotide comprises: (a) a gap segment consisting of ten linked deoxynucleosides; (b) a 5' wing segment consisting of five linked nucleosides; (c) a 3' wing segment consisting of five linked nucleosides; and wherein the gap segment is positioned between the 5' wing segment and the 3' wing segment, wherein each nucleoside of each wing segment comprises a 2'-O-methoxyethyl sugar, wherein at least one internucleoside linkage is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

Certain embodiments provide a composition comprising a conjugated antisense compound described herein, or a salt thereof, and a pharmaceutically acceptable carrier or diluent.

In certain embodiments, the modulation of apo(a) expression occurs in a cell or tissue. In certain embodiments, the modulations occur in a cell or tissue in an animal. In certain embodiments, the animal is a human. In certain embodiments, the modulation is a reduction in apo(a) mRNA level. In certain embodiments, the modulation is a reduction in apo(a) protein level. In certain embodiments, both apo(a) mRNA and protein levels are reduced. In certain embodiments, the modulation is a reduction in Lp(a) level. Such reduction may occur in a time-dependent or in a dose-dependent manner.

Certain embodiments provide conjugated antisense compositions and methods for use in therapy. Certain embodiments provide compositions and methods for preventing, treating, delaying, slowing the progression and/or ameliorating apo(a) related diseases, disorders, and conditions. Certain embodiments provide compositions and methods for preventing, treating, delaying, slowing the progression and/or ameliorating Lp(a) related diseases, disorders, and conditions. In certain embodiments, such diseases, disorders, and conditions are inflammatory, cardiovascular and/or metabolic diseases, disorders, and conditions. In certain embodiments, the compositions and methods for therapy include administering an apo(a) specific inhibitor to an individual in need thereof. In certain embodiments, the apo(a) specific inhibitor is a nucleic acid. In certain embodiments, the nucleic acid is an antisense compound. In certain embodiments, the antisense compound is a modified oligonucleotide. In certain embodiments, the antisense compound is a modified oligonucleotide with a conjugate.

In certain embodiments, the present disclosure provides conjugated antisense compounds. In certain embodiments, the present disclosure provides conjugated antisense compounds comprising an antisense oligonucleotide complementary to a nucleic acid transcript. In certain embodiments, the present disclosure provides methods comprising contacting a cell with a conjugated antisense compound comprising an antisense oligonucleotide complementary to a nucleic acid transcript. In certain embodiments, the present disclosure provides methods comprising contacting a cell with a conjugated antisense compound comprising an antisense oligonucleotide and reducing the amount or activity of a nucleic acid transcript in a cell.

The asialoglycoprotein receptor (ASGP-R) has been described previously. See e.g., Park et al., PNAS vol. 102, No. 47, pp 17125-17129 (2005). Such receptors are expressed on liver cells, particularly hepatocytes. Further, it has been shown that compounds comprising clusters of three N-

acetylgalactosamine (GalNAc) ligands are capable of binding to the ASGP-R, resulting in uptake of the compound into the cell. See e.g., Khorev et al., *Bioorganic and Medicinal Chemistry*, 16, 9, pp 5216-5231 (May 2008). Accordingly, conjugates comprising such GalNAc clusters have been used to facilitate uptake of certain compounds into liver cells, specifically hepatocytes. For example it has been shown that certain GalNAc-containing conjugates increase activity of duplex siRNA compounds in liver cells in vivo. In such instances, the GalNAc-containing conjugate is typically attached to the sense strand of the siRNA duplex. Since the sense strand is discarded before the antisense strand ultimately hybridizes with the target nucleic acid, there is little concern that the conjugate will interfere with activity. Typically, the conjugate is attached to the 3' end of the sense strand of the siRNA. See e.g., U.S. Patent 8,106,022. Certain conjugate groups described herein are more active and/or easier to synthesize than conjugate groups previously described.

In certain embodiments of the present invention, conjugates are attached to single-stranded antisense compounds, including, but not limited to RNase H based antisense compounds and antisense compounds that alter splicing of a pre-mRNA target nucleic acid. In such embodiments, the conjugate should remain attached to the antisense compound long enough to provide benefit (improved uptake into cells) but then should either be cleaved, or otherwise not interfere with the subsequent steps necessary for activity, such as hybridization to a target nucleic acid and interaction with RNase H or enzymes associated with splicing or splice modulation. This balance of properties is more important in the setting of single-stranded antisense compounds than in siRNA compounds, where the conjugate may simply be attached to the sense strand. Disclosed herein are conjugated single-stranded antisense compounds having improved potency in liver cells in vivo compared with the same antisense compound lacking the conjugate. Given the required balance of properties for these compounds such improved potency is surprising.

In certain embodiments, conjugate groups herein comprise a cleavable moiety. As noted, without wishing to be bound by mechanism, it is logical that the conjugate should remain on the compound long enough to provide enhancement in uptake, but after that, it is desirable for some portion or, ideally, all of the conjugate to be cleaved, releasing the parent compound (e.g., antisense compound) in its most active form. In certain embodiments, the cleavable moiety is a cleavable nucleoside. Such embodiments take advantage of endogenous nucleases in the cell by attaching the rest of the conjugate (the cluster) to the antisense oligonucleotide through a nucleoside via one or more cleavable bonds, such as those of a phosphodiester linkage. In certain embodiments, the cluster is bound to the cleavable nucleoside through a phosphodiester linkage. In certain embodiments, the cleavable nucleoside is attached to the antisense oligonucleotide (antisense compound) by a phosphodiester linkage. In certain embodiments, the conjugate group may comprise two or three cleavable nucleosides. In such embodiments, such cleavable nucleosides are linked to one another, to the antisense compound and/or to the cluster via cleavable bonds (such as those of a phosphodiester linkage). Certain conjugates herein do not comprise a cleavable nucleoside and instead comprise a cleavable bond. It is shown that that sufficient cleavage of the conjugate from the oligonucleotide is provided by at least one bond that is vulnerable to cleavage in the cell (a cleavable bond).

In certain embodiments, conjugated antisense compounds are prodrugs. Such prodrugs are administered to an animal and are ultimately metabolized to a more active form. For example, conjugated antisense compounds are cleaved to remove all or part of the conjugate resulting in the active (or more active) form of the antisense compound lacking all or some of the conjugate.

5 In certain embodiments, conjugates are attached at the 5' end of an oligonucleotide. Certain such 5'-conjugates are cleaved more efficiently than counterparts having a similar conjugate group attached at the 3' end. In certain embodiments, improved activity may correlate with improved cleavage. In certain embodiments, oligonucleotides comprising a conjugate at the 5' end have greater efficacy than oligonucleotides comprising a conjugate at the 3' end (see, for example, Examples 56, 81, 83, and 84).
10 Further, 5'-attachment allows simpler oligonucleotide synthesis. Typically, oligonucleotides are synthesized on a solid support in the 3' to 5' direction. To make a 3'-conjugated oligonucleotide, typically one attaches a pre-conjugated 3' nucleoside to the solid support and then builds the oligonucleotide as usual. However, attaching that conjugated nucleoside to the solid support adds complication to the synthesis. Further, using that approach, the conjugate is then present throughout the synthesis of the oligonucleotide and can become
15 degraded during subsequent steps or may limit the sorts of reactions and reagents that can be used. Using the structures and techniques described herein for 5'-conjugated oligonucleotides, one can synthesize the oligonucleotide using standard automated techniques and introduce the conjugate with the final (5'-most) nucleoside or after the oligonucleotide has been cleaved from the solid support.

In view of the art and the present disclosure, one of ordinary skill can easily make any of the
20 conjugates and conjugated oligonucleotides herein. Moreover, synthesis of certain such conjugates and conjugated oligonucleotides disclosed herein is easier and/or requires few steps, and is therefore less expensive than that of conjugates previously disclosed, providing advantages in manufacturing. For example, the synthesis of certain conjugate groups consists of fewer synthetic steps, resulting in increased yield, relative to conjugate groups previously described. Conjugate groups such as GalNAc3-10 in Example 46 and
25 GalNAc3-7 in Example 48 are much simpler than previously described conjugates such as those described in U.S. 8,106,022 or U.S. 7,262,177 that require assembly of more chemical intermediates. Accordingly, these and other conjugates described herein have advantages over previously described compounds for use with any oligonucleotide, including single-stranded oligonucleotides and either strand of double-stranded oligonucleotides (e.g., siRNA).

30 Similarly, disclosed herein are conjugate groups having only one or two GalNAc ligands. As shown, such conjugates groups improve activity of antisense compounds. Such compounds are much easier to prepare than conjugates comprising three GalNAc ligands. Conjugate groups comprising one or two GalNAc ligands may be attached to any antisense compounds, including single-stranded oligonucleotides and either strand of double-stranded oligonucleotides (e.g., siRNA).

35 In certain embodiments, the conjugates herein do not substantially alter certain measures of tolerability. For example, it is shown herein that conjugated antisense compounds are not more immunogenic

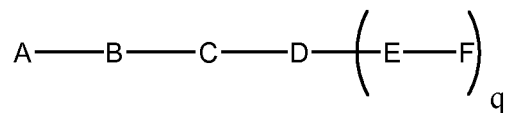
than unconjugated parent compounds. Since potency is improved, embodiments in which tolerability remains the same (or indeed even if tolerability worsens only slightly compared to the gains in potency) have improved properties for therapy.

5 In certain embodiments, conjugation allows one to alter antisense compounds in ways that have less attractive consequences in the absence of conjugation. For example, in certain embodiments, replacing one or more phosphorothioate linkages of a fully phosphorothioate antisense compound with phosphodiester linkages results in improvement in some measures of tolerability. For example, in certain instances, such antisense compounds having one or more phosphodiester are less immunogenic than the same compound in which each linkage is a phosphorothioate. However, in certain instances, as shown in Example 26, that same
10 replacement of one or more phosphorothioate linkages with phosphodiester linkages also results in reduced cellular uptake and/or loss in potency. In certain embodiments, conjugated antisense compounds described herein tolerate such change in linkages with little or no loss in uptake and potency when compared to the conjugated full-phosphorothioate counterpart. In fact, in certain embodiments, for example, in Examples 44, 57, 59, and 86, oligonucleotides comprising a conjugate and at least one phosphodiester internucleoside
15 linkage actually exhibit increased potency in vivo even relative to a full phosphorothioate counterpart also comprising the same conjugate. Moreover, since conjugation results in substantial increases in uptake/potency a small loss in that substantial gain may be acceptable to achieve improved tolerability. Accordingly, in certain embodiments, conjugated antisense compounds comprise at least one phosphodiester linkage.

20 In certain embodiments, conjugation of antisense compounds herein results in increased delivery, uptake and activity in hepatocytes. Thus, more compound is delivered to liver tissue. However, in certain embodiments, that increased delivery alone does not explain the entire increase in activity. In certain such embodiments, more compound enters hepatocytes. In certain embodiments, even that increased hepatocyte uptake does not explain the entire increase in activity. In such embodiments, productive uptake of the
25 conjugated compound is increased. For example, as shown in Example 102, certain embodiments of GalNAc-containing conjugates increase enrichment of antisense oligonucleotides in hepatocytes versus non-parenchymal cells. This enrichment is beneficial for oligonucleotides that target genes that are expressed in hepatocytes.

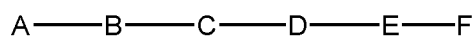
In certain embodiments, conjugated antisense compounds herein result in reduced kidney exposure.
30 For example, as shown in Example 20, the concentrations of antisense oligonucleotides comprising certain embodiments of GalNAc-containing conjugates are lower in the kidney than that of antisense oligonucleotides lacking a GalNAc-containing conjugate. This has several beneficial therapeutic implications. For therapeutic indications where activity in the kidney is not sought, exposure to kidney risks kidney toxicity without corresponding benefit. Moreover, high concentration in kidney typically results in
35 loss of compound to the urine resulting in faster clearance. Accordingly for non-kidney targets, kidney accumulation is undesired.

In certain embodiments, the present disclosure provides conjugated antisense compounds represented by the formula:

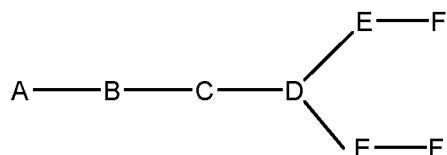


- 5 wherein
 A is the antisense oligonucleotide;
 B is the cleavable moiety
 C is the conjugate linker
 D is the branching group
 10 each E is a tether;
 each F is a ligand; and
 q is an integer between 1 and 5.

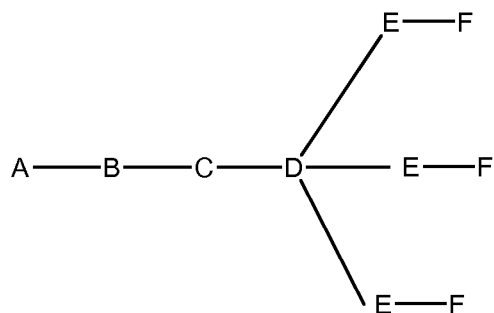
In the above diagram and in similar diagrams herein, the branching group “D” branches as many times as is necessary to accommodate the number of (E-F) groups as indicated by “q”. Thus, where q = 1,
 15 the formula is:



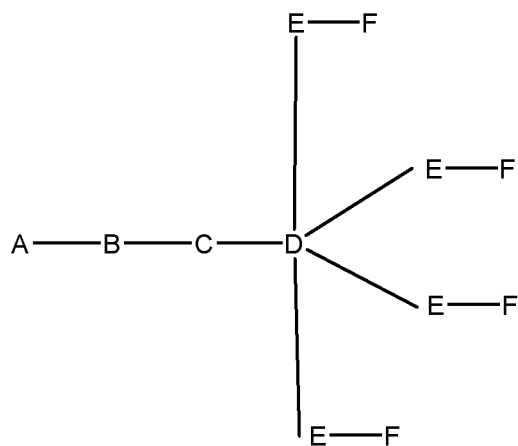
where q = 2, the formula is:



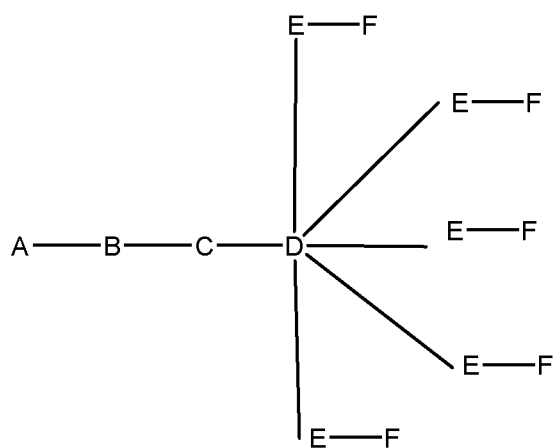
where q = 3, the formula is:



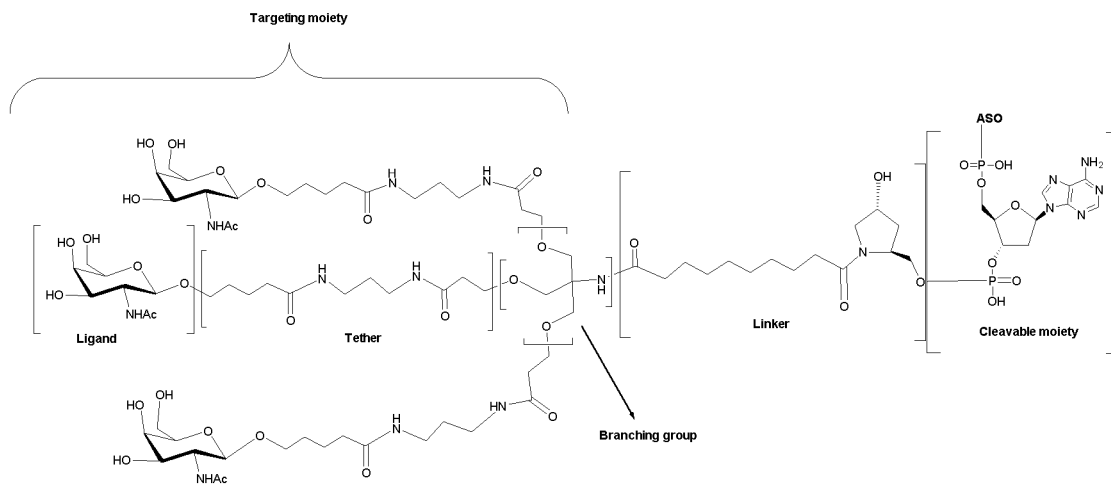
where q = 4, the formula is:



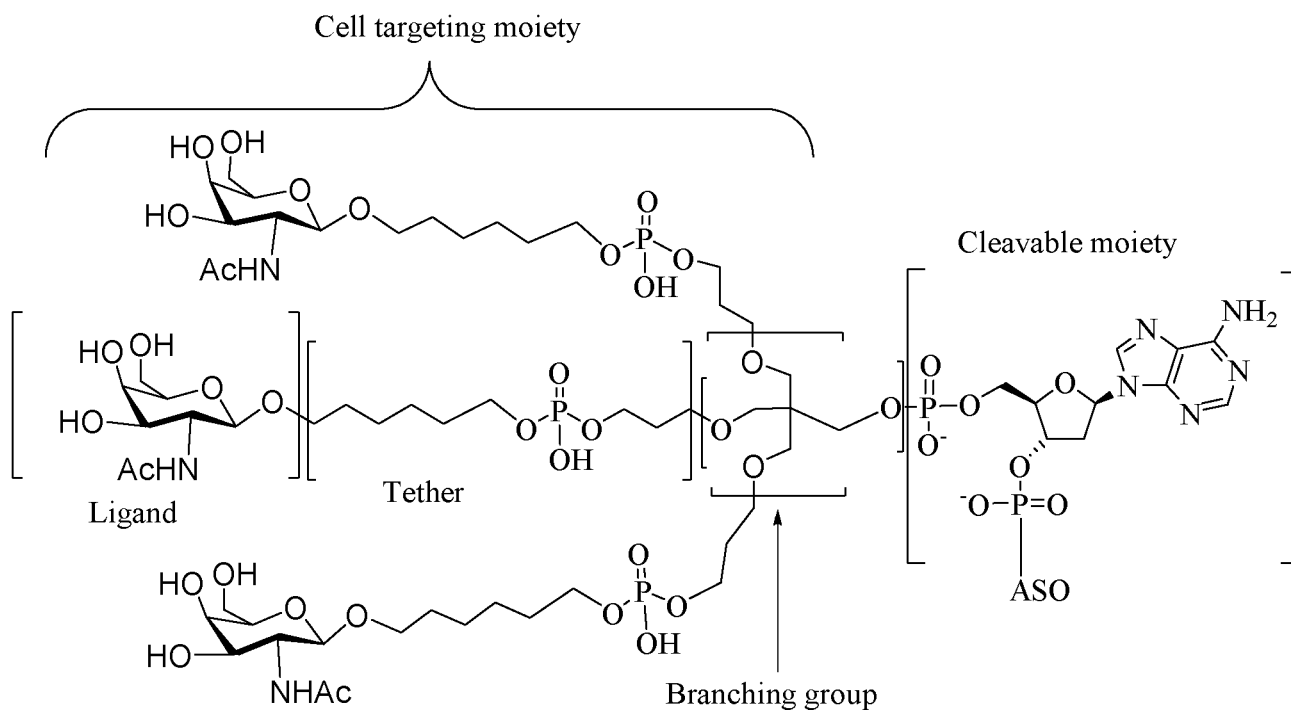
where $q = 5$, the formula is:



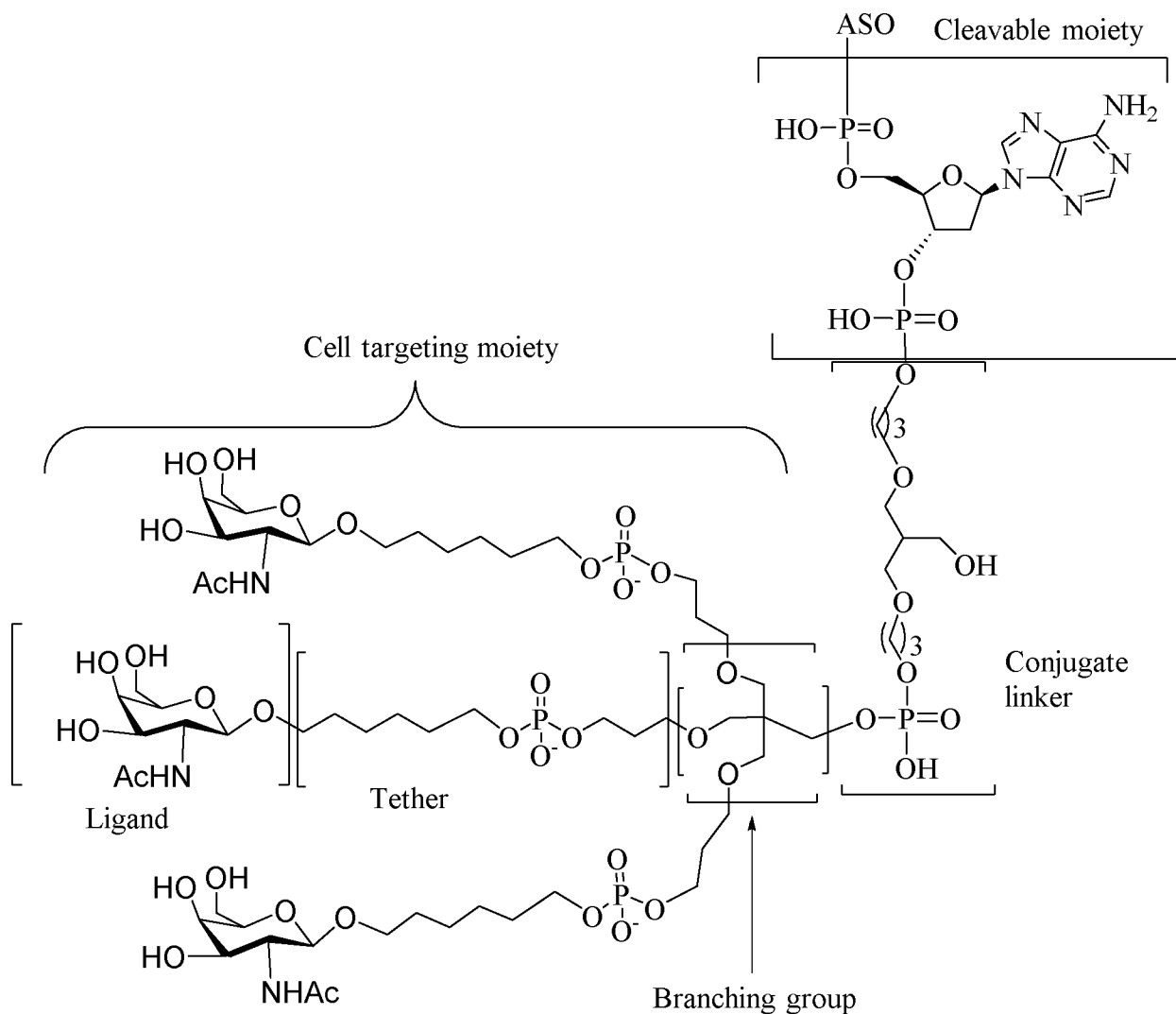
In certain embodiments, conjugated antisense compounds are provided having the structure:



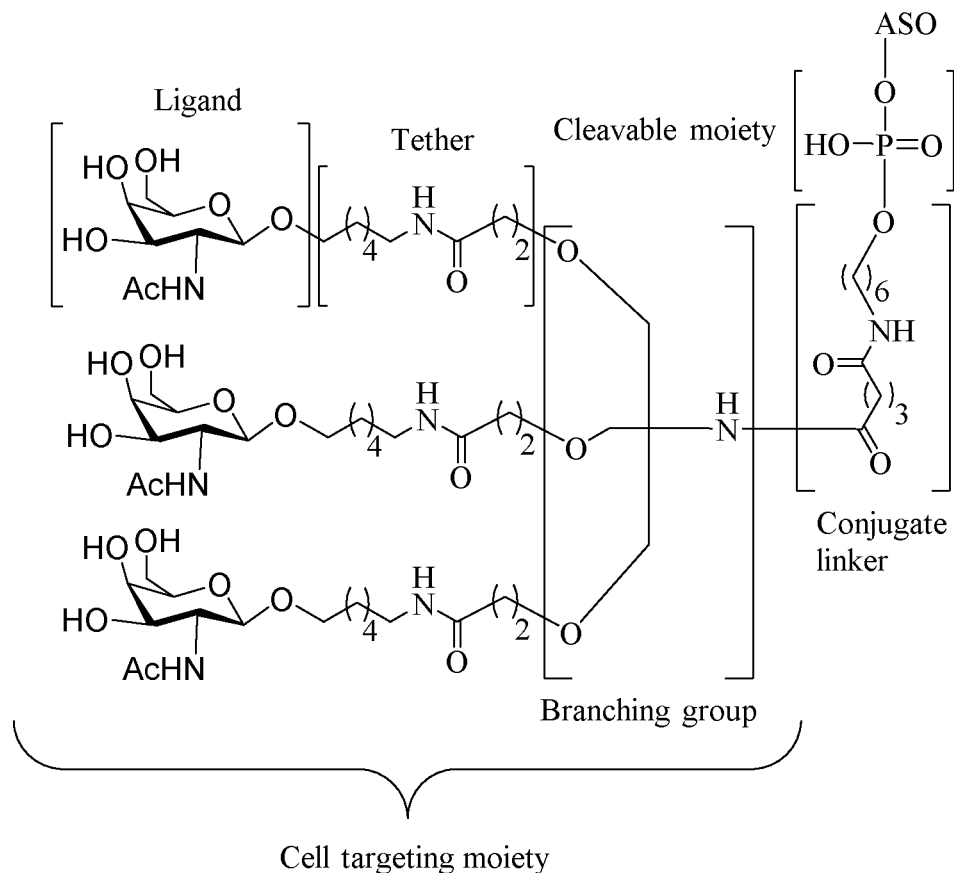
In certain embodiments, conjugated antisense compounds are provided having the structure:



In certain embodiments, conjugated antisense compounds are provided having the structure:



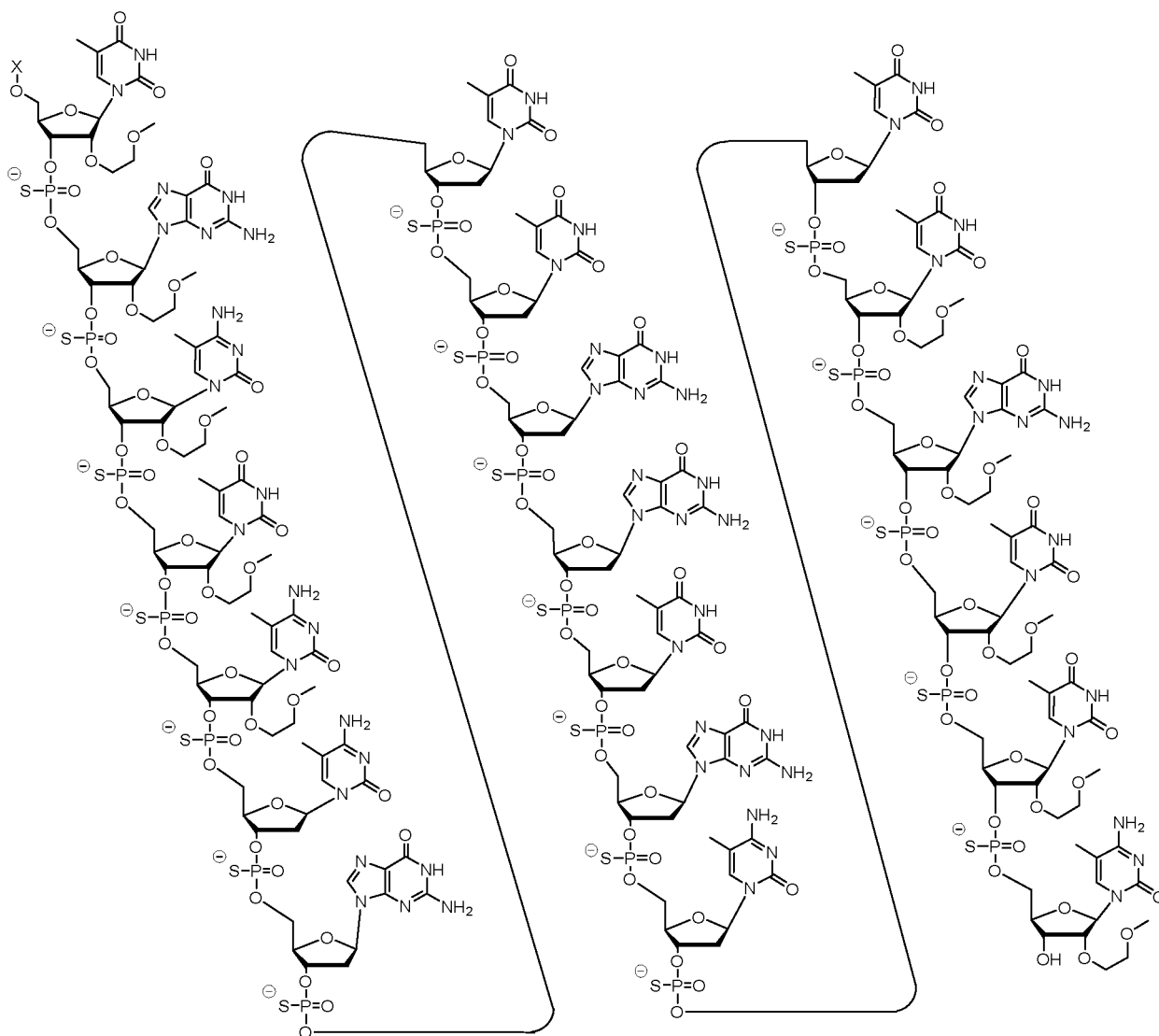
In certain embodiments, conjugated antisense compounds are provided having the structure:



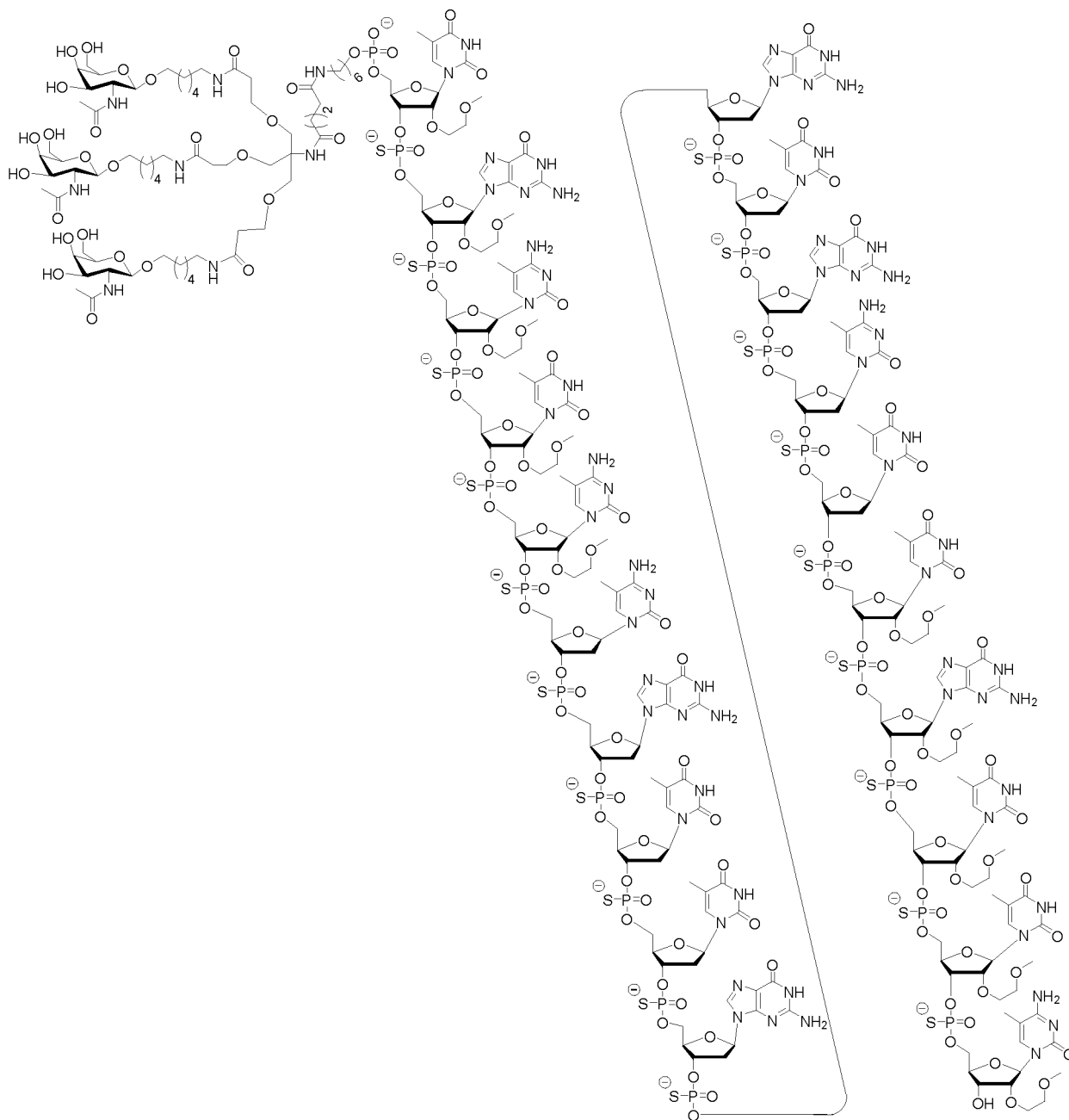
The present disclosure provides the following non-limiting numbered embodiments:

In embodiments having more than one of a particular variable (e.g., more than one “m” or “n”),
 5 unless otherwise indicated, each such particular variable is selected independently. Thus, for a structure having more than one n, each n is selected independently, so they may or may not be the same as one another.

In certain embodiments, the present disclosure provides conjugated antisense compounds represented by the following structure. In certain embodiments, the antisense compound comprises modified
 oligonucleotide ISIS 494372 with a 5'-X, wherein X is a conjugate group comprising GalNAc. In certain
 10 embodiments, the antisense compound consists of modified oligonucleotide ISIS 494372 with a 5'-X, wherein X is a conjugate group comprising GalNAc.

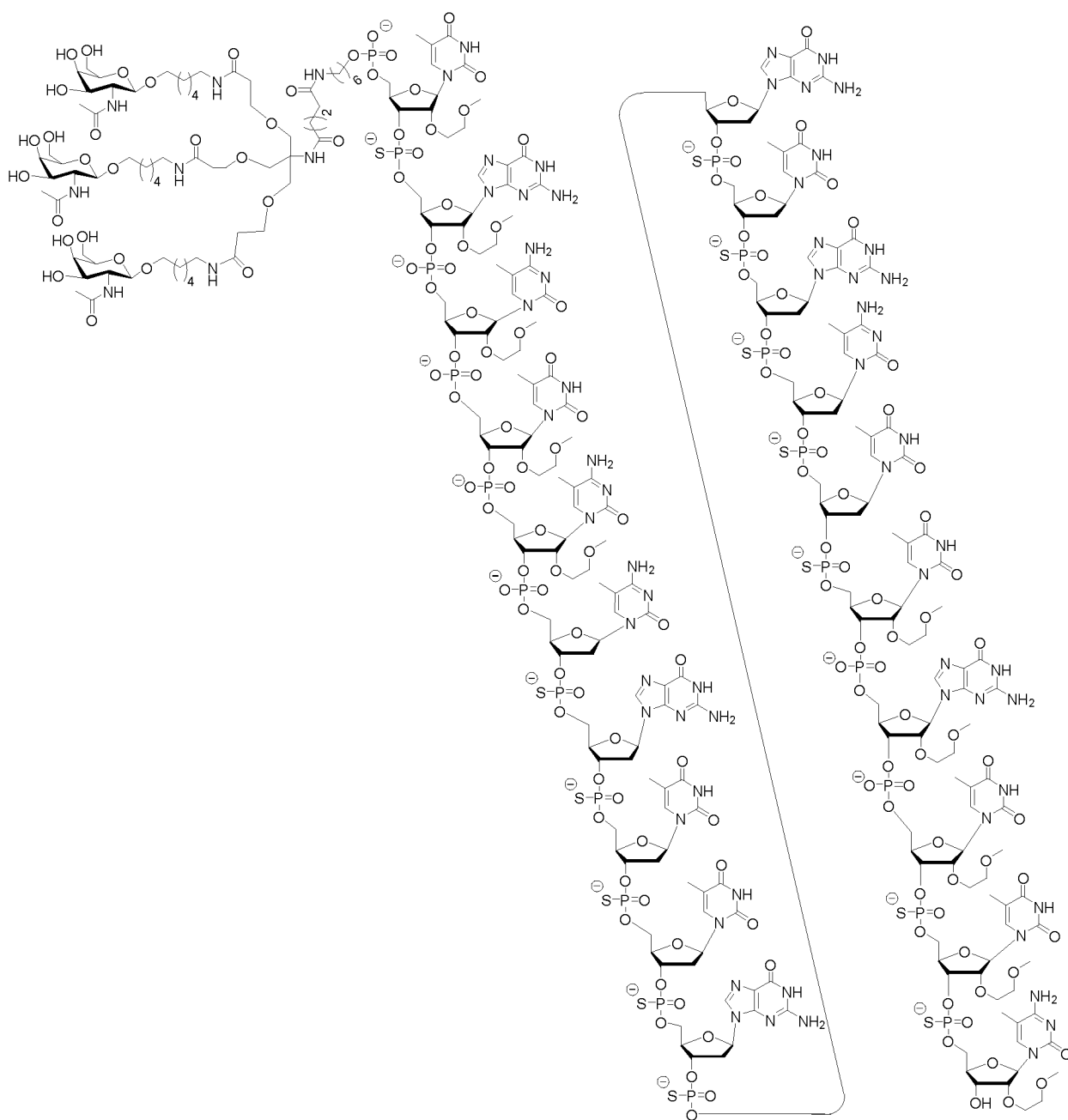


In certain embodiments, the present disclosure provides conjugated antisense compounds represented by the following structure. In certain embodiments, the antisense compound comprises the conjugated modified oligonucleotide ISIS 681251. In certain embodiments, the antisense compound consists of the conjugated modified oligonucleotide ISIS 681251.

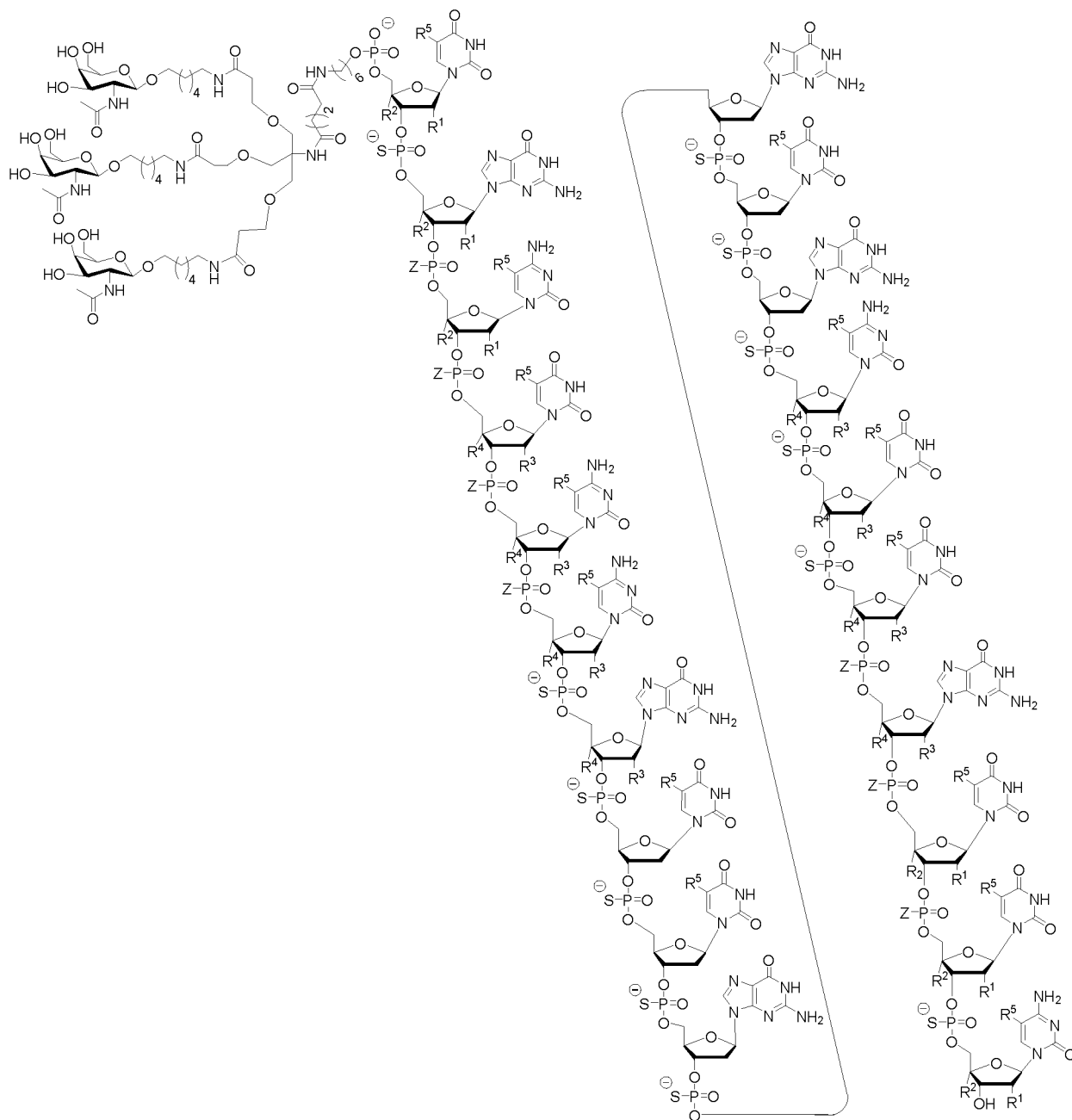


In certain embodiments, the present disclosure provides conjugated antisense compounds represented by the following structure. In certain embodiments, the antisense compound comprises the conjugated modified oligonucleotide ISIS 681257. In certain embodiments, the antisense compound consists of the conjugated modified oligonucleotide ISIS 681257.

5



In certain embodiments, the present disclosure provides conjugated antisense compounds represented by the following structure. In certain embodiments, the antisense compound comprises a modified oligonucleotide with SEQ ID NO: 58 with a 5'-GalNAc with variability in the sugar mods of the wings. In certain embodiments, the antisense compound consists of a modified oligonucleotide with SEQ ID NO: 58 with a 5'-GalNAc with variability in the sugar mods of the wings.



Wherein either R^1 is $-\text{OCH}_2\text{CH}_2\text{OCH}_3$ (MOE) and R^2 is H; or R^1 and R^2 together form a bridge, wherein R^1 is $-\text{O}-$ and R^2 is $-\text{CH}_2-$, $-\text{CH}(\text{CH}_3)-$, or $-\text{CH}_2\text{CH}_2-$, and R^1 and R^2 are directly connected such that the resulting bridge is selected from: $-\text{O}-\text{CH}_2-$, $-\text{O}-\text{CH}(\text{CH}_3)-$, and $-\text{O}-\text{CH}_2\text{CH}_2-$;

And for each pair of R^3 and R^4 on the same ring, independently for each ring: either R^3 is selected from H and $-OCH_2CH_2OCH_3$ and R^4 is H; or R^3 and R^4 together form a bridge, wherein R^3 is $-O-$, and R^4 is $-CH_2-$, $-CH(CH_3)-$, or $-CH_2CH_2-$ and R^3 and R^4 are directly connected such that the resulting bridge is selected from: $-O-CH_2-$, $-O-CH(CH_3)-$, and $-O-CH_2CH_2-$;

And R^5 is selected from H and $-CH_3$;

And Z is selected from S^- and O^- .

The present disclosure provides the following non-limiting numbered embodiments:

DETAILED DESCRIPTION

It is to be understood that both the foregoing general description and the following detailed description are exemplary and explanatory only and are not restrictive of the disclosure. Herein, the use of the singular includes the plural unless specifically stated otherwise. As used herein, the use of "or" means "and/or" unless stated otherwise. Furthermore, the use of the term "including" as well as other forms, such as "includes" and "included", is not limiting. Also, terms such as "element" or "component" encompass both elements and components comprising one unit and elements and components that comprise more than one subunit, unless specifically stated otherwise.

The section headings used herein are for organizational purposes only and are not to be construed as limiting the subject matter described. All documents, or portions of documents, cited in this application, including, but not limited to, patents, patent applications, articles, books, and treatises, are hereby expressly incorporated by reference in their entirety for any purpose.

A. Definitions

Unless specific definitions are provided, the nomenclature used in connection with, and the procedures and techniques of, analytical chemistry, synthetic organic chemistry, and medicinal and pharmaceutical chemistry described herein are those well known and commonly used in the art. Standard techniques may be used for chemical synthesis, and chemical analysis. Certain such techniques and procedures may be found for example in "Carbohydrate Modifications in Antisense Research" Edited by Sangvi and Cook, American Chemical Society, Washington D.C., 1994; "Remington's Pharmaceutical Sciences," Mack Publishing Co., Easton, Pa., 21st edition, 2005; and "Antisense Drug Technology, Principles, Strategies, and Applications" Edited by Stanley T. Crooke, CRC Press, Boca Raton, Florida; and Sambrook et al., "Molecular Cloning, A laboratory Manual," 2nd Edition, Cold Spring Harbor Laboratory Press, 1989, which are hereby incorporated by reference for any purpose. Where permitted, all patents, applications, published applications and other publications and other data referred to throughout in the disclosure are incorporated by reference herein in their entirety.

Unless otherwise indicated, the following terms have the following meanings:

As used herein, “nucleoside” means a compound comprising a nucleobase moiety and a sugar moiety. Nucleosides include, but are not limited to, naturally occurring nucleosides (as found in DNA and RNA) and modified nucleosides. Nucleosides may be linked to a phosphate moiety.

As used herein, “chemical modification” means a chemical difference in a compound when compared to a naturally occurring counterpart. Chemical modifications of oligonucleotides include nucleoside modifications (including sugar moiety modifications and nucleobase modifications) and internucleoside linkage modifications. In reference to an oligonucleotide, chemical modification does not include differences only in nucleobase sequence.

As used herein, “furanosyl” means a structure comprising a 5-membered ring comprising four carbon atoms and one oxygen atom.

As used herein, “naturally occurring sugar moiety” means a ribofuranosyl as found in naturally occurring RNA or a deoxyribofuranosyl as found in naturally occurring DNA.

As used herein, “sugar moiety” means a naturally occurring sugar moiety or a modified sugar moiety of a nucleoside.

As used herein, “modified sugar moiety” means a substituted sugar moiety or a sugar surrogate.

As used herein, “substituted sugar moiety” means a furanosyl that is not a naturally occurring sugar moiety. Substituted sugar moieties include, but are not limited to furanosyls comprising substituents at the 2'-position, the 3'-position, the 5'-position and/or the 4'-position. Certain substituted sugar moieties are bicyclic sugar moieties.

As used herein, “2'-substituted sugar moiety” means a furanosyl comprising a substituent at the 2'-position other than H or OH. Unless otherwise indicated, a 2'-substituted sugar moiety is not a bicyclic sugar moiety (i.e., the 2'-substituent of a 2'-substituted sugar moiety does not form a bridge to another atom of the furanosyl ring).

As used herein, “MOE” means $-OCH_2CH_2OCH_3$.

As used herein, “2'-F nucleoside” refers to a nucleoside comprising a sugar comprising fluorine at the 2' position. Unless otherwise indicated, the fluorine in a 2'-F nucleoside is in the ribo position (replacing the OH of a natural ribose).

As used herein the term “sugar surrogate” means a structure that does not comprise a furanosyl and that is capable of replacing the naturally occurring sugar moiety of a nucleoside, such that the resulting nucleoside sub-units are capable of linking together and/or linking to other nucleosides to form an oligomeric compound which is capable of hybridizing to a complementary oligomeric compound. Such structures include rings comprising a different number of atoms than furanosyl (e.g., 4, 6, or 7-membered rings); replacement of the oxygen of a furanosyl with a non-oxygen atom (e.g., carbon, sulfur, or nitrogen); or both a change in the number of atoms and a replacement of the oxygen. Such structures may also comprise substitutions corresponding to those described for substituted sugar moieties (e.g., 6-membered carbocyclic bicyclic sugar surrogates optionally comprising additional substituents). Sugar surrogates also include more

complex sugar replacements (e.g., the non-ring systems of peptide nucleic acid). Sugar surrogates include without limitation morpholinos, cyclohexenyls and cyclohexitols.

As used herein, “bicyclic sugar moiety” means a modified sugar moiety comprising a 4 to 7 membered ring (including but not limited to a furanosyl) comprising a bridge connecting two atoms of the 4 to 7 membered ring to form a second ring, resulting in a bicyclic structure. In certain embodiments, the 4 to 7 membered ring is a sugar ring. In certain embodiments the 4 to 7 membered ring is a furanosyl. In certain such embodiments, the bridge connects the 2'-carbon and the 4'-carbon of the furanosyl.

As used herein, “nucleic acid” refers to molecules composed of monomeric nucleotides. A nucleic acid includes ribonucleic acids (RNA), deoxyribonucleic acids (DNA), single-stranded nucleic acids (ssDNA), double-stranded nucleic acids (dsDNA), small interfering ribonucleic acids (siRNA), and microRNAs (miRNA). A nucleic acid may also comprise any combination of these elements in a single molecule.

As used herein, “nucleotide” means a nucleoside further comprising a phosphate linking group. As used herein, “linked nucleosides” may or may not be linked by phosphate linkages and thus includes, but is not limited to “linked nucleotides.” As used herein, “linked nucleosides” are nucleosides that are connected in a continuous sequence (i.e. no additional nucleosides are present between those that are linked).

As used herein, “nucleobase” means a group of atoms that can be linked to a sugar moiety to create a nucleoside that is capable of incorporation into an oligonucleotide, and wherein the group of atoms is capable of bonding with a complementary naturally occurring nucleobase of another oligonucleotide or nucleic acid. Nucleobases may be naturally occurring or may be modified. As used herein, “nucleobase sequence” means the order of contiguous nucleobases independent of any sugar, linkage, or nucleobase modification.

As used herein the terms, “unmodified nucleobase” or “naturally occurring nucleobase” means the naturally occurring heterocyclic nucleobases of RNA or DNA: the purine bases adenine (A) and guanine (G), and the pyrimidine bases thymine (T), cytosine (C) (including 5-methyl C), and uracil (U).

As used herein, “modified nucleobase” means any nucleobase that is not a naturally occurring nucleobase.

As used herein, “modified nucleoside” means a nucleoside comprising at least one chemical modification compared to naturally occurring RNA or DNA nucleosides. Modified nucleosides comprise a modified sugar moiety and/or a modified nucleobase.

As used herein, “bicyclic nucleoside” or “BNA” means a nucleoside comprising a bicyclic sugar moiety.

As used herein, “constrained ethyl nucleoside” or “cEt” means a nucleoside comprising a bicyclic sugar moiety comprising a 4'-CH(CH₃)-O-2' bridge.

As used herein, “locked nucleic acid nucleoside” or “LNA” means a nucleoside comprising a bicyclic sugar moiety comprising a 4'-CH₂-O-2' bridge.

As used herein, “2'-substituted nucleoside” means a nucleoside comprising a substituent at the 2'-

position other than H or OH. Unless otherwise indicated, a 2'-substituted nucleoside is not a bicyclic nucleoside.

As used herein, "deoxynucleoside" means a nucleoside comprising 2'-H furanosyl sugar moiety, as found in naturally occurring deoxyribonucleosides (DNA). In certain embodiments, a 2'-deoxynucleoside may comprise a modified nucleobase or may comprise an RNA nucleobase (e.g., uracil).

As used herein, "oligonucleotide" means a compound comprising a plurality of linked nucleosides. In certain embodiments, an oligonucleotide comprises one or more unmodified ribonucleosides (RNA) and/or unmodified deoxyribonucleosides (DNA) and/or one or more modified nucleosides.

As used herein "oligonucleoside" means an oligonucleotide in which none of the internucleoside linkages contains a phosphorus atom. As used herein, oligonucleotides include oligonucleosides.

As used herein, "modified oligonucleotide" means an oligonucleotide comprising at least one modified nucleoside and/or at least one modified internucleoside linkage.

As used herein, "linkage" or "linking group" means a group of atoms that link together two or more other groups of atoms.

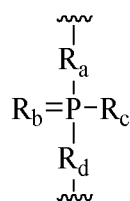
As used herein "internucleoside linkage" means a covalent linkage between adjacent nucleosides in an oligonucleotide.

As used herein "naturally occurring internucleoside linkage" means a 3' to 5' phosphodiester linkage.

As used herein, "modified internucleoside linkage" means any internucleoside linkage other than a naturally occurring internucleoside linkage.

As used herein, "terminal internucleoside linkage" means the linkage between the last two nucleosides of an oligonucleotide or defined region thereof.

As used herein, "phosphorus linking group" means a linking group comprising a phosphorus atom. Phosphorus linking groups include without limitation groups having the formula:



wherein:

R_a and R_d are each, independently, O, S, CH_2 , NH, or NJ_1 wherein J_1 is C_1 - C_6 alkyl or substituted C_1 - C_6 alkyl;

R_b is O or S;

R_c is OH, SH, C_1 - C_6 alkyl, substituted C_1 - C_6 alkyl, C_1 - C_6 alkoxy, substituted C_1 - C_6 alkoxy, amino or substituted amino; and

J_1 is R_b is O or S.

Phosphorus linking groups include without limitation, phosphodiester, phosphorothioate, phosphorodithioate,

phosphonate, phosphoramidate, phosphorothioamidate, thionoalkylphosphonate, phosphotriesters, thionoalkylphosphotriester and boranophosphate.

As used herein, "internucleoside phosphorus linking group" means a phosphorus linking group that directly links two nucleosides.

5 As used herein, "non-internucleoside phosphorus linking group" means a phosphorus linking group that does not directly link two nucleosides. In certain embodiments, a non-internucleoside phosphorus linking group links a nucleoside to a group other than a nucleoside. In certain embodiments, a non-internucleoside phosphorus linking group links two groups, neither of which is a nucleoside.

As used herein, "neutral linking group" means a linking group that is not charged. Neutral linking
10 groups include without limitation phosphotriesters, methylphosphonates, MMI (-CH₂-N(CH₃)-O-), amide-3 (-CH₂-C(=O)-N(H)-), amide-4 (-CH₂-N(H)-C(=O)-), formacetal (-O-CH₂-O-), and thioformacetal (-S-CH₂-O-). Further neutral linking groups include nonionic linkages comprising siloxane (dialkylsiloxane), carboxylate ester, carboxamide, sulfide, sulfonate ester and amides (See for example: Carbohydrate Modifications in Antisense Research; Y.S. Sanghvi and P.D. Cook Eds. ACS Symposium Series 580; Chapters 3 and 4, (pp.
15 40-65)). Further neutral linking groups include nonionic linkages comprising mixed N, O, S and CH₂ component parts.

As used herein, "internucleoside neutral linking group" means a neutral linking group that directly links two nucleosides.

As used herein, "non-internucleoside neutral linking group" means a neutral linking group that does
20 not directly link two nucleosides. In certain embodiments, a non-internucleoside neutral linking group links a nucleoside to a group other than a nucleoside. In certain embodiments, a non-internucleoside neutral linking group links two groups, neither of which is a nucleoside.

As used herein, "oligomeric compound" means a polymeric structure comprising two or more sub-
structures. In certain embodiments, an oligomeric compound comprises an oligonucleotide. In certain
25 embodiments, an oligomeric compound comprises one or more conjugate groups and/or terminal groups. In certain embodiments, an oligomeric compound consists of an oligonucleotide. Oligomeric compounds also include naturally occurring nucleic acids. In certain embodiments, an oligomeric compound comprises a backbone of one or more linked monomeric subunits where each linked monomeric subunit is directly or indirectly attached to a heterocyclic base moiety. In certain embodiments, oligomeric compounds may also
30 include monomeric subunits that are not linked to a heterocyclic base moiety, thereby providing abasic sites. In certain embodiments, the linkages joining the monomeric subunits, the sugar moieties or surrogates and the heterocyclic base moieties can be independently modified. In certain embodiments, the linkage-sugar unit, which may or may not include a heterocyclic base, may be substituted with a mimetic such as the monomers in peptide nucleic acids.

35 As used herein, "terminal group" means one or more atom attached to either, or both, the 3' end or the 5' end of an oligonucleotide. In certain embodiments a terminal group is a conjugate group. In certain

embodiments, a terminal group comprises one or more terminal group nucleosides.

As used herein, “conjugate” or “conjugate group” means an atom or group of atoms bound to an oligonucleotide or oligomeric compound. In general, conjugate groups modify one or more properties of the compound to which they are attached, including, but not limited to pharmacodynamic, pharmacokinetic, binding, absorption, cellular distribution, cellular uptake, charge and/or clearance properties.

As used herein, “conjugate linker” or “linker” in the context of a conjugate group means a portion of a conjugate group comprising any atom or group of atoms and which covalently link (1) an oligonucleotide to another portion of the conjugate group or (2) two or more portions of the conjugate group.

Conjugate groups are shown herein as radicals, providing a bond for forming covalent attachment to an oligomeric compound such as an antisense oligonucleotide. In certain embodiments, the point of attachment on the oligomeric compound is the 3'-oxygen atom of the 3'-hydroxyl group of the 3' terminal nucleoside of the oligomeric compound. In certain embodiments the point of attachment on the oligomeric compound is the 5'-oxygen atom of the 5'-hydroxyl group of the 5' terminal nucleoside of the oligomeric compound. In certain embodiments, the bond for forming attachment to the oligomeric compound is a cleavable bond. In certain such embodiments, such cleavable bond constitutes all or part of a cleavable moiety.

In certain embodiments, conjugate groups comprise a cleavable moiety (e.g., a cleavable bond or cleavable nucleoside) and a carbohydrate cluster portion, such as a GalNAc cluster portion. Such carbohydrate cluster portion comprises: a targeting moiety and, optionally, a conjugate linker. In certain embodiments, the carbohydrate cluster portion is identified by the number and identity of the ligand. For example, in certain embodiments, the carbohydrate cluster portion comprises 3 GalNAc groups and is designated “GalNAc₃”. In certain embodiments, the carbohydrate cluster portion comprises 4 GalNAc groups and is designated “GalNAc₄”. Specific carbohydrate cluster portions (having specific tether, branching and conjugate linker groups) are described herein and designated by Roman numeral followed by subscript “a”. Accordingly “GalNAc3-1_a” refers to a specific carbohydrate cluster portion of a conjugate group having 3 GalNAc groups and specifically identified tether, branching and linking groups. Such carbohydrate cluster fragment is attached to an oligomeric compound via a cleavable moiety, such as a cleavable bond or cleavable nucleoside.

As used herein, “cleavable moiety” means a bond or group that is capable of being split under physiological conditions. In certain embodiments, a cleavable moiety is cleaved inside a cell or sub-cellular compartments, such as a lysosome. In certain embodiments, a cleavable moiety is cleaved by endogenous enzymes, such as nucleases. In certain embodiments, a cleavable moiety comprises a group of atoms having one, two, three, four, or more than four cleavable bonds.

As used herein, “cleavable bond” means any chemical bond capable of being split. In certain embodiments, a cleavable bond is selected from among: an amide, a polyamide, an ester, an ether, one or both esters of a phosphodiester, a phosphate ester, a carbamate, a di-sulfide, or a peptide.

As used herein, "carbohydrate cluster" means a compound having one or more carbohydrate residues attached to a scaffold or linker group. (see, e.g., Maier et al., "Synthesis of Antisense Oligonucleotides Conjugated to a Multivalent Carbohydrate Cluster for Cellular Targeting," *Bioconjugate Chemistry*, 2003, (14): 18-29, which is incorporated herein by reference in its entirety, or Rensen et al., "Design and Synthesis of Novel *N*-Acetylgalactosamine-Terminated Glycolipids for Targeting of Lipoproteins to the Hepatic Asialoglycoprotein Receptor," *J. Med. Chem.* 2004, (47): 5798-5808, for examples of carbohydrate conjugate clusters).

As used herein, "modified carbohydrate" means any carbohydrate having one or more chemical modifications relative to naturally occurring carbohydrates.

As used herein, "carbohydrate derivative" means any compound which may be synthesized using a carbohydrate as a starting material or intermediate.

As used herein, "carbohydrate" means a naturally occurring carbohydrate, a modified carbohydrate, or a carbohydrate derivative.

As used herein "protecting group" means any compound or protecting group known to those having skill in the art. Non-limiting examples of protecting groups may be found in "Protective Groups in Organic Chemistry", T. W. Greene, P. G. M. Wuts, ISBN 0-471-62301-6, John Wiley & Sons, Inc, New York, which is incorporated herein by reference in its entirety.

As used herein, "single-stranded" means an oligomeric compound that is not hybridized to its complement and which lacks sufficient self-complementarity to form a stable self-duplex.

As used herein, "double stranded" means a pair of oligomeric compounds that are hybridized to one another or a single self-complementary oligomeric compound that forms a hairpin structure. In certain embodiments, a double-stranded oligomeric compound comprises a first and a second oligomeric compound.

As used herein, "antisense compound" means a compound comprising or consisting of an oligonucleotide at least a portion of which is complementary to a target nucleic acid to which it is capable of hybridizing, resulting in at least one antisense activity.

As used herein, "antisense activity" means any detectable and/or measurable change attributable to the hybridization of an antisense compound to its target nucleic acid. In certain embodiments, antisense activity includes modulation of the amount or activity of a target nucleic acid transcript (e.g. mRNA). In certain embodiments, antisense activity includes modulation of the splicing of pre-mRNA.

As used herein, "RNase H based antisense compound" means an antisense compound wherein at least some of the antisense activity of the antisense compound is attributable to hybridization of the antisense compound to a target nucleic acid and subsequent cleavage of the target nucleic acid by RNase H.

As used herein, "RISC based antisense compound" means an antisense compound wherein at least some of the antisense activity of the antisense compound is attributable to the RNA Induced Silencing Complex (RISC).

As used herein, "detecting" or "measuring" means that a test or assay for detecting or measuring is

performed. Such detection and/or measuring may result in a value of zero. Thus, if a test for detection or measuring results in a finding of no activity (activity of zero), the step of detecting or measuring the activity has nevertheless been performed.

As used herein, “detectable and/or measureable activity” means a statistically significant activity that is not zero.

As used herein, “essentially unchanged” means little or no change in a particular parameter, particularly relative to another parameter which changes much more. In certain embodiments, a parameter is essentially unchanged when it changes less than 5%. In certain embodiments, a parameter is essentially unchanged if it changes less than two-fold while another parameter changes at least ten-fold. For example, in certain embodiments, an antisense activity is a change in the amount of a target nucleic acid. In certain such embodiments, the amount of a non-target nucleic acid is essentially unchanged if it changes much less than the target nucleic acid does, but the change need not be zero.

As used herein, “expression” means the process by which a gene ultimately results in a protein. Expression includes, but is not limited to, transcription, post-transcriptional modification (e.g., splicing, polyadenylation, addition of 5'-cap), and translation.

As used herein, “target nucleic acid” means a nucleic acid molecule to which an antisense compound is intended to hybridize to result in a desired antisense activity. Antisense oligonucleotides have sufficient complementarity to their target nucleic acids to allow hybridization under physiological conditions.

As used herein, “nucleobase complementarity” or “complementarity” when in reference to nucleobases means a nucleobase that is capable of base pairing with another nucleobase. For example, in DNA, adenine (A) is complementary to thymine (T). For example, in RNA, adenine (A) is complementary to uracil (U). In certain embodiments, complementary nucleobase means a nucleobase of an antisense compound that is capable of base pairing with a nucleobase of its target nucleic acid. For example, if a nucleobase at a certain position of an antisense compound is capable of hydrogen bonding with a nucleobase at a certain position of a target nucleic acid, then the position of hydrogen bonding between the oligonucleotide and the target nucleic acid is considered to be complementary at that nucleobase pair. Nucleobases comprising certain modifications may maintain the ability to pair with a counterpart nucleobase and thus, are still capable of nucleobase complementarity.

As used herein, “non-complementary” in reference to nucleobases means a pair of nucleobases that do not form hydrogen bonds with one another.

As used herein, “complementary” in reference to oligomeric compounds (e.g., linked nucleosides, oligonucleotides, or nucleic acids) means the capacity of such oligomeric compounds or regions thereof to hybridize to another oligomeric compound or region thereof through nucleobase complementarity. Complementary oligomeric compounds need not have nucleobase complementarity at each nucleoside. Rather, some mismatches are tolerated. In certain embodiments, complementary oligomeric compounds or regions are complementary at 70% of the nucleobases (70% complementary). In certain embodiments,

complementary oligomeric compounds or regions are 80% complementary. In certain embodiments, complementary oligomeric compounds or regions are 90% complementary. In certain embodiments, complementary oligomeric compounds or regions are 95% complementary. In certain embodiments, complementary oligomeric compounds or regions are 100% complementary.

5 As used herein, “mismatch” means a nucleobase of a first oligomeric compound that is not capable of pairing with a nucleobase at a corresponding position of a second oligomeric compound, when the first and second oligomeric compound are aligned. Either or both of the first and second oligomeric compounds may be oligonucleotides.

10 As used herein, “hybridization” means the pairing of complementary oligomeric compounds (e.g., an antisense compound and its target nucleic acid). While not limited to a particular mechanism, the most common mechanism of pairing involves hydrogen bonding, which may be Watson-Crick, Hoogsteen or reversed Hoogsteen hydrogen bonding, between complementary nucleobases.

As used herein, “specifically hybridizes” means the ability of an oligomeric compound to hybridize to one nucleic acid site with greater affinity than it hybridizes to another nucleic acid site.

15 As used herein, “fully complementary” in reference to an oligonucleotide or portion thereof means that each nucleobase of the oligonucleotide or portion thereof is capable of pairing with a nucleobase of a complementary nucleic acid or contiguous portion thereof. Thus, a fully complementary region comprises no mismatches or unhybridized nucleobases in either strand.

20 As used herein, “percent complementarity” means the percentage of nucleobases of an oligomeric compound that are complementary to an equal-length portion of a target nucleic acid. Percent complementarity is calculated by dividing the number of nucleobases of the oligomeric compound that are complementary to nucleobases at corresponding positions in the target nucleic acid by the total length of the oligomeric compound.

25 As used herein, “percent identity” means the number of nucleobases in a first nucleic acid that are the same type (independent of chemical modification) as nucleobases at corresponding positions in a second nucleic acid, divided by the total number of nucleobases in the first nucleic acid.

30 As used herein, “modulation” means a change of amount or quality of a molecule, function, or activity when compared to the amount or quality of a molecule, function, or activity prior to modulation. For example, modulation includes the change, either an increase (stimulation or induction) or a decrease (inhibition or reduction) in gene expression. As a further example, modulation of expression can include a change in splice site selection of pre-mRNA processing, resulting in a change in the absolute or relative amount of a particular splice-variant compared to the amount in the absence of modulation.

35 As used herein, “chemical motif” means a pattern of chemical modifications in an oligonucleotide or a region thereof. Motifs may be defined by modifications at certain nucleosides and/or at certain linking groups of an oligonucleotide.

As used herein, “nucleoside motif” means a pattern of nucleoside modifications in an oligonucleotide or a region thereof. The linkages of such an oligonucleotide may be modified or unmodified. Unless otherwise indicated, motifs herein describing only nucleosides are intended to be nucleoside motifs. Thus, in such instances, the linkages are not limited.

5 As used herein, “sugar motif” means a pattern of sugar modifications in an oligonucleotide or a region thereof.

As used herein, “linkage motif” means a pattern of linkage modifications in an oligonucleotide or region thereof. The nucleosides of such an oligonucleotide may be modified or unmodified. Unless otherwise indicated, motifs herein describing only linkages are intended to be linkage motifs. Thus, in such
10 instances, the nucleosides are not limited.

As used herein, “nucleobase modification motif” means a pattern of modifications to nucleobases along an oligonucleotide. Unless otherwise indicated, a nucleobase modification motif is independent of the nucleobase sequence.

As used herein, “sequence motif” means a pattern of nucleobases arranged along an oligonucleotide
15 or portion thereof. Unless otherwise indicated, a sequence motif is independent of chemical modifications and thus may have any combination of chemical modifications, including no chemical modifications.

As used herein, “type of modification” in reference to a nucleoside or a nucleoside of a “type” means the chemical modification of a nucleoside and includes modified and unmodified nucleosides. Accordingly, unless otherwise indicated, a “nucleoside having a modification of a first type” may be an unmodified
20 nucleoside.

As used herein, “differently modified” mean chemical modifications or chemical substituents that are different from one another, including absence of modifications. Thus, for example, a MOE nucleoside and an unmodified DNA nucleoside are “differently modified,” even though the DNA nucleoside is unmodified. Likewise, DNA and RNA are “differently modified,” even though both are naturally-occurring unmodified
25 nucleosides. Nucleosides that are the same but for comprising different nucleobases are not differently modified. For example, a nucleoside comprising a 2'-OMe modified sugar and an unmodified adenine nucleobase and a nucleoside comprising a 2'-OMe modified sugar and an unmodified thymine nucleobase are not differently modified.

As used herein, “the same type of modifications” refers to modifications that are the same as one
30 another, including absence of modifications. Thus, for example, two unmodified DNA nucleosides have “the same type of modification,” even though the DNA nucleoside is unmodified. Such nucleosides having the same type modification may comprise different nucleobases.

As used herein, “separate regions” means portions of an oligonucleotide wherein the chemical modifications or the motif of chemical modifications of any neighboring portions include at least one
35 difference to allow the separate regions to be distinguished from one another.

As used herein, “pharmaceutically acceptable carrier or diluent” means any substance suitable for use

in administering to an animal. In certain embodiments, a pharmaceutically acceptable carrier or diluent is sterile saline. In certain embodiments, such sterile saline is pharmaceutical grade saline.

As used herein the term "metabolic disorder" means a disease or condition principally characterized by dysregulation of metabolism – the complex set of chemical reactions associated with breakdown of food to produce energy.

As used herein, the term "cardiovascular disorder" means a disease or condition principally characterized by impaired function of the heart or blood vessels.

As used herein the term "mono or polycyclic ring system" is meant to include all ring systems selected from single or polycyclic radical ring systems wherein the rings are fused or linked and is meant to be inclusive of single and mixed ring systems individually selected from aliphatic, alicyclic, aryl, heteroaryl, aralkyl, arylalkyl, heterocyclic, heteroaryl, heteroaromatic and heteroarylalkyl. Such mono and poly cyclic structures can contain rings that each have the same level of saturation or each, independently, have varying degrees of saturation including fully saturated, partially saturated or fully unsaturated. Each ring can comprise ring atoms selected from C, N, O and S to give rise to heterocyclic rings as well as rings comprising only C ring atoms which can be present in a mixed motif such as for example benzimidazole wherein one ring has only carbon ring atoms and the fused ring has two nitrogen atoms. The mono or polycyclic ring system can be further substituted with substituent groups such as for example phthalimide which has two =O groups attached to one of the rings. Mono or polycyclic ring systems can be attached to parent molecules using various strategies such as directly through a ring atom, fused through multiple ring atoms, through a substituent group or through a bifunctional linking moiety.

As used herein, "prodrug" means an inactive or less active form of a compound which, when administered to a subject, is metabolized to form the active, or more active, compound (e.g., drug).

As used herein, "substituent" and "substituent group," means an atom or group that replaces the atom or group of a named parent compound. For example a substituent of a modified nucleoside is any atom or group that differs from the atom or group found in a naturally occurring nucleoside (e.g., a modified 2'-substituent is any atom or group at the 2'-position of a nucleoside other than H or OH). Substituent groups can be protected or unprotected. In certain embodiments, compounds of the present disclosure have substituents at one or at more than one position of the parent compound. Substituents may also be further substituted with other substituent groups and may be attached directly or via a linking group such as an alkyl or hydrocarbyl group to a parent compound.

Likewise, as used herein, "substituent" in reference to a chemical functional group means an atom or group of atoms that differs from the atom or a group of atoms normally present in the named functional group. In certain embodiments, a substituent replaces a hydrogen atom of the functional group (e.g., in certain embodiments, the substituent of a substituted methyl group is an atom or group other than hydrogen which replaces one of the hydrogen atoms of an unsubstituted methyl group). Unless otherwise indicated, groups amenable for use as substituents include without limitation, halogen, hydroxyl, alkyl, alkenyl, alkynyl,

acyl ($-C(O)R_{aa}$), carboxyl ($-C(O)O-R_{aa}$), aliphatic groups, alicyclic groups, alkoxy, substituted oxy ($-O-R_{aa}$), aryl, aralkyl, heterocyclic radical, heteroaryl, heteroarylalkyl, amino ($-N(R_{bb})(R_{cc})$), imino ($=NR_{bb}$), amido ($-C(O)N(R_{bb})(R_{cc})$ or $-N(R_{bb})C(O)R_{aa}$), azido ($-N_3$), nitro ($-NO_2$), cyano ($-CN$), carbamido ($-OC(O)N(R_{bb})(R_{cc})$ or $-N(R_{bb})C(O)OR_{aa}$), ureido ($-N(R_{bb})C(O)N(R_{bb})(R_{cc})$), thioureido ($-N(R_{bb})C(S)N(R_{bb})(R_{cc})$), guanidiny ($-N(R_{bb})C(=NR_{bb})N(R_{bb})(R_{cc})$), amidiny ($-C(=NR_{bb})N(R_{bb})(R_{cc})$ or $-N(R_{bb})C(=NR_{bb})(R_{aa})$), thiol ($-SR_{bb}$), sulfinyl ($-S(O)R_{bb}$), sulfonyl ($-S(O)_2R_{bb}$) and sulfonamidyl ($-S(O)_2N(R_{bb})(R_{cc})$ or $-N(R_{bb})S(O)_2R_{bb}$). Wherein each R_{aa} , R_{bb} and R_{cc} is, independently, H, an optionally linked chemical functional group or a further substituent group with a preferred list including without limitation, alkyl, alkenyl, alkynyl, aliphatic, alkoxy, acyl, aryl, aralkyl, heteroaryl, alicyclic, heterocyclic and heteroarylalkyl. Selected substituents within the compounds described herein are present to a recursive degree.

As used herein, "alkyl," as used herein, means a saturated straight or branched hydrocarbon radical containing up to twenty four carbon atoms. Examples of alkyl groups include without limitation, methyl, ethyl, propyl, butyl, isopropyl, n-hexyl, octyl, decyl, dodecyl and the like. Alkyl groups typically include from 1 to about 24 carbon atoms, more typically from 1 to about 12 carbon atoms (C_1 - C_{12} alkyl) with from 1 to about 6 carbon atoms being more preferred.

As used herein, "alkenyl," means a straight or branched hydrocarbon chain radical containing up to twenty four carbon atoms and having at least one carbon-carbon double bond. Examples of alkenyl groups include without limitation, ethenyl, propenyl, butenyl, 1-methyl-2-buten-1-yl, dienes such as 1,3-butadiene and the like. Alkenyl groups typically include from 2 to about 24 carbon atoms, more typically from 2 to about 12 carbon atoms with from 2 to about 6 carbon atoms being more preferred. Alkenyl groups as used herein may optionally include one or more further substituent groups.

As used herein, "alkynyl," means a straight or branched hydrocarbon radical containing up to twenty four carbon atoms and having at least one carbon-carbon triple bond. Examples of alkynyl groups include, without limitation, ethynyl, 1-propynyl, 1-butylnyl, and the like. Alkynyl groups typically include from 2 to about 24 carbon atoms, more typically from 2 to about 12 carbon atoms with from 2 to about 6 carbon atoms being more preferred. Alkynyl groups as used herein may optionally include one or more further substituent groups.

As used herein, "acyl," means a radical formed by removal of a hydroxyl group from an organic acid and has the general Formula $-C(O)-X$ where X is typically aliphatic, alicyclic or aromatic. Examples include aliphatic carbonyls, aromatic carbonyls, aliphatic sulfonyls, aromatic sulfinyls, aliphatic sulfinyls, aromatic phosphates, aliphatic phosphates and the like. Acyl groups as used herein may optionally include further substituent groups.

As used herein, "alicyclic" means a cyclic ring system wherein the ring is aliphatic. The ring system can comprise one or more rings wherein at least one ring is aliphatic. Preferred alicyclics include rings having from about 5 to about 9 carbon atoms in the ring. Alicyclic as used herein may optionally include further substituent groups.

As used herein, "aliphatic" means a straight or branched hydrocarbon radical containing up to twenty four carbon atoms wherein the saturation between any two carbon atoms is a single, double or triple bond. An aliphatic group preferably contains from 1 to about 24 carbon atoms, more typically from 1 to about 12 carbon atoms with from 1 to about 6 carbon atoms being more preferred. The straight or branched chain of an aliphatic group may be interrupted with one or more heteroatoms that include nitrogen, oxygen, sulfur and phosphorus. Such aliphatic groups interrupted by heteroatoms include without limitation, polyalkoxys, such as polyalkylene glycols, polyamines, and polyimines. Aliphatic groups as used herein may optionally include further substituent groups.

As used herein, "alkoxy" means a radical formed between an alkyl group and an oxygen atom wherein the oxygen atom is used to attach the alkoxy group to a parent molecule. Examples of alkoxy groups include without limitation, methoxy, ethoxy, propoxy, isopropoxy, *n*-butoxy, *sec*-butoxy, *tert*-butoxy, *n*-pentoxy, neopentoxy, *n*-hexoxy and the like. Alkoxy groups as used herein may optionally include further substituent groups.

As used herein, "aminoalkyl" means an amino substituted C₁-C₁₂ alkyl radical. The alkyl portion of the radical forms a covalent bond with a parent molecule. The amino group can be located at any position and the aminoalkyl group can be substituted with a further substituent group at the alkyl and/or amino portions.

As used herein, "aralkyl" and "arylalkyl" mean an aromatic group that is covalently linked to a C₁-C₁₂ alkyl radical. The alkyl radical portion of the resulting aralkyl (or arylalkyl) group forms a covalent bond with a parent molecule. Examples include without limitation, benzyl, phenethyl and the like. Aralkyl groups as used herein may optionally include further substituent groups attached to the alkyl, the aryl or both groups that form the radical group.

As used herein, "aryl" and "aromatic" mean a mono- or polycyclic carbocyclic ring system radicals having one or more aromatic rings. Examples of aryl groups include without limitation, phenyl, naphthyl, tetrahydronaphthyl, indanyl, idenyl and the like. Preferred aryl ring systems have from about 5 to about 20 carbon atoms in one or more rings. Aryl groups as used herein may optionally include further substituent groups.

As used herein, "halo" and "halogen," mean an atom selected from fluorine, chlorine, bromine and iodine.

As used herein, "heteroaryl," and "heteroaromatic," mean a radical comprising a mono- or polycyclic aromatic ring, ring system or fused ring system wherein at least one of the rings is aromatic and includes one or more heteroatoms. Heteroaryl is also meant to include fused ring systems including systems where one or more of the fused rings contain no heteroatoms. Heteroaryl groups typically include one ring atom selected from sulfur, nitrogen or oxygen. Examples of heteroaryl groups include without limitation, pyridinyl, pyrazinyl, pyrimidinyl, pyrrolyl, pyrazolyl, imidazolyl, thiazolyl, oxazolyl, isooxazolyl, thiadiazolyl, oxadiazolyl, thiophenyl, furanyl, quinolinyl, isoquinolinyl, benzimidazolyl, benzooxazolyl,

quinoxalinyll and the like. Heteroaryl radicals can be attached to a parent molecule directly or through a linking moiety such as an aliphatic group or hetero atom. Heteroaryl groups as used herein may optionally include further substituent groups.

As used herein, “conjugate compound” means any atoms, group of atoms, or group of linked atoms suitable for use as a conjugate group. In certain embodiments, conjugate compounds may possess or impart one or more properties, including, but not limited to pharmacodynamic, pharmacokinetic, binding, absorption, cellular distribution, cellular uptake, charge and/or clearance properties.

As used herein, unless otherwise indicated or modified, the term “double-stranded” refers to two separate oligomeric compounds that are hybridized to one another. Such double stranded compounds may have one or more or non-hybridizing nucleosides at one or both ends of one or both strands (overhangs) and/or one or more internal non-hybridizing nucleosides (mismatches) provided there is sufficient complementarity to maintain hybridization under physiologically relevant conditions.

As used herein, “5’ target site” refers to the nucleotide of a target nucleic acid which is complementary to the 5’-most nucleotide of a particular antisense compound.

As used herein, “About” means within $\pm 10\%$ of a value. For example, if it is stated, “a marker may be increased by about 50%”, it is implied that the marker may be increased between 45%-55%.

As used herein, “administered concomitantly” refers to the co-administration of two agents in any manner in which the pharmacological effects of both are manifest in the patient at the same time. Concomitant administration does not require that both agents be administered in a single pharmaceutical composition, in the same dosage form, or by the same route of administration. The effects of both agents need not manifest themselves at the same time. The effects need only be overlapping for a period of time and need not be coextensive.

As used herein, “administering” or “administration” means providing a pharmaceutical agent to an individual, and includes, but is not limited to, administering by a medical professional and self-administering. Administration of a pharmaceutical agent to an individual can be continuous, chronic, short or intermittent. Administration can parenteral or non-parenteral.

As used herein, “agent” means an active substance that can provide a therapeutic benefit when administered to an animal. “First agent” means a therapeutic compound of the invention. For example, a first agent can be an antisense oligonucleotide targeting apo(a). “Second agent” means a second therapeutic compound of the invention (e.g. a second antisense oligonucleotide targeting apo(a)) and/or a non-apo(a) therapeutic compound.

As used herein, “amelioration” or “ameliorate” or “ameliorating” refers to a lessening of at least one indicator, sign, or symptom of an associated disease, disorder, or condition. The severity of indicators can be determined by subjective or objective measures, which are known to those skilled in the art.

As used herein, “animal” refers to a human or non-human animal, including, but not limited to, mice, rats, rabbits, dogs, cats, pigs, and non-human primates, including, but not limited to, monkeys and chimpanzees.

As used herein, “apo(a)” means any nucleic acid or protein sequence encoding apo(a). For example, in certain embodiments, apo(a) includes a DNA sequence encoding apo(a), a RNA sequence transcribed from DNA encoding apo(a) (including genomic DNA comprising introns and exons), a mRNA sequence encoding apo(a), or a peptide sequence encoding apo(a).

As used herein, “apo(a) nucleic acid” means any nucleic acid encoding apo(a). For example, in certain embodiments, an apo(a) nucleic acid includes a DNA sequence encoding apo(a), a RNA sequence transcribed from DNA encoding apo(a) (including genomic DNA comprising introns and exons), and a mRNA sequence encoding apo(a).

As used herein, “apo(a) mRNA” means a mRNA encoding an apo(a) protein.

As used herein, “apo(a) protein” means any protein sequence encoding Apo(a).

As used herein, “apo(a) specific inhibitor” refers to any agent capable of specifically inhibiting the expression of an apo(a) nucleic acid and/or apo(a) protein. For example, apo(a) specific inhibitors include nucleic acids (including antisense compounds), peptides, antibodies, small molecules, and other agents capable of inhibiting the expression of apo(a) nucleic acid and/or apo(a) protein. In certain embodiments, by specifically modulating apo(a) nucleic acid expression and/or apo(a) protein expression, apo(a) specific inhibitors can affect other components of the lipid transport system including downstream components. Similarly, in certain embodiments, apo(a) specific inhibitors can affect other molecular processes in an animal.

As used herein, “atherosclerosis” means a hardening of the arteries affecting large and medium-sized arteries and is characterized by the presence of fatty deposits. The fatty deposits are called “atheromas” or “plaques,” which consist mainly of cholesterol and other fats, calcium and scar tissue, and damage the lining of arteries.

As used herein, “coronary heart disease (CHD)” means a narrowing of the small blood vessels that supply blood and oxygen to the heart, which is often a result of atherosclerosis.

As used herein, “diabetes mellitus” or “diabetes” is a syndrome characterized by disordered metabolism and abnormally high blood sugar (hyperglycemia) resulting from insufficient levels of insulin or reduced insulin sensitivity. The characteristic symptoms are excessive urine production (polyuria) due to high blood glucose levels, excessive thirst and increased fluid intake (polydipsia) attempting to compensate for increased urination, blurred vision due to high blood glucose effects on the eye's optics, unexplained weight loss, and lethargy.

As used herein, “diabetic dyslipidemia” or “type 2 diabetes with dyslipidemia” means a condition characterized by Type 2 diabetes, reduced HDL-C, elevated triglycerides (TG), and elevated small, dense LDL particles.

As used herein, “diluent” means an ingredient in a composition that lacks pharmacological activity, but is pharmaceutically necessary or desirable. For example, the diluent in an injected composition can be a liquid, e.g. saline solution.

As used herein, “dyslipidemia” refers to a disorder of lipid and/or lipoprotein metabolism, including lipid and/or lipoprotein overproduction or deficiency. Dyslipidemias can be manifested by elevation of lipids such as chylomicron, cholesterol and triglycerides as well as lipoproteins such as low-density lipoprotein (LDL) cholesterol.

As used herein, “dosage unit” means a form in which a pharmaceutical agent is provided, e.g. pill, tablet, or other dosage unit known in the art. In certain embodiments, a dosage unit is a vial containing lyophilized antisense oligonucleotide. In certain embodiments, a dosage unit is a vial containing reconstituted antisense oligonucleotide.

As used herein, “dose” means a specified quantity of a pharmaceutical agent provided in a single administration, or in a specified time period. In certain embodiments, a dose can be administered in one, two, or more boluses, tablets, or injections. For example, in certain embodiments where subcutaneous administration is desired, the desired dose requires a volume not easily accommodated by a single injection, therefore, two or more injections can be used to achieve the desired dose. In certain embodiments, the pharmaceutical agent is administered by infusion over an extended period of time or continuously. Doses can be stated as the amount of pharmaceutical agent per hour, day, week, or month. Doses can also be stated as mg/kg or g/kg.

As used herein, “effective amount” or “therapeutically effective amount” means the amount of active pharmaceutical agent sufficient to effectuate a desired physiological outcome in an individual in need of the agent. The effective amount can vary among individuals depending on the health and physical condition of the individual to be treated, the taxonomic group of the individuals to be treated, the formulation of the composition, assessment of the individual’s medical condition, and other relevant factors.

As used herein, “fully complementary” or “100% complementary” means each nucleobase of a nucleobase sequence of a first nucleic acid has a complementary nucleobase in a second nucleobase sequence of a second nucleic acid. In certain embodiments, a first nucleic acid is an antisense compound and a second nucleic acid is a target nucleic acid.

As used herein, “glucose” is a monosaccharide used by cells as a source of energy and inflammatory intermediate. “Plasma glucose” refers to glucose present in the plasma.

As used herein, “high density lipoprotein-C” or “HDL-C” means cholesterol associated with high density lipoprotein particles. Concentration of HDL-C in serum (or plasma) is typically quantified in mg/dL or nmol/L. “Serum HDL-C” and “plasma HDL-C” mean HDL-C in serum and plasma, respectively.

As used herein, “HMG-CoA reductase inhibitor” means an agent that acts through the inhibition of the enzyme HMG-CoA reductase, such as atorvastatin, rosuvastatin, fluvastatin, lovastatin, pravastatin, and simvastatin.

As used herein, “hypercholesterolemia” means a condition characterized by elevated cholesterol or circulating (plasma) cholesterol, LDL-cholesterol and VLDL-cholesterol, as per the guidelines of the Expert Panel Report of the National Cholesterol Educational Program (NCEP) of Detection, Evaluation of Treatment of high cholesterol in adults (see, Arch. Int. Med. (1988) 148, 36-39).

5 As used herein, “hyperlipidemia” or “hyperlipemia” is a condition characterized by elevated serum lipids or circulating (plasma) lipids. This condition manifests an abnormally high concentration of fats. The lipid fractions in the circulating blood are cholesterol, low density lipoproteins, very low density lipoproteins, chylomicrons and triglycerides. The Fredrickson classification of hyperlipidemias is based on the pattern of TG and cholesterol-rich lipoprotein particles, as measured by electrophoresis or ultracentrifugation and is
10 commonly used to characterize primary causes of hyperlipidemias such as hypertriglyceridemia (Fredrickson and Lee, Circulation, 1965, 31:321-327; Fredrickson et al., New Eng J Med, 1967, 276 (1): 34-42).

As used herein, “hypertriglyceridemia” means a condition characterized by elevated triglyceride levels. Its etiology includes primary (i.e. genetic causes) and secondary (other underlying causes such as diabetes, metabolic syndrome/insulin resistance, obesity, physical inactivity, cigarette smoking, excess
15 alcohol and a diet very high in carbohydrates) factors or, most often, a combination of both (Yuan *et al.* CMAJ, 2007, 176:1113-1120).

As used herein, “identifying” or “selecting an animal with metabolic or cardiovascular disease” means identifying or selecting a subject prone to or having been diagnosed with a metabolic disease, a cardiovascular disease, or a metabolic syndrome; or, identifying or selecting a subject having any symptom of
20 a metabolic disease, cardiovascular disease, or metabolic syndrome including, but not limited to, hypercholesterolemia, hyperglycemia, hyperlipidemia, hypertriglyceridemia, hypertension increased insulin resistance, decreased insulin sensitivity, above normal body weight, and/or above normal body fat content or any combination thereof. Such identification can be accomplished by any method, including but not limited to, standard clinical tests or assessments, such as measuring serum or circulating (plasma) cholesterol,
25 measuring serum or circulating (plasma) blood-glucose, measuring serum or circulating (plasma) triglycerides, measuring blood-pressure, measuring body fat content, measuring body weight, and the like.

As used herein, “improved cardiovascular outcome” means a reduction in the occurrence of adverse cardiovascular events, or the risk thereof. Examples of adverse cardiovascular events include, without
30 limitation, death, reinfarction, stroke, cardiogenic shock, pulmonary edema, cardiac arrest, and atrial dysrhythmia.

As used herein, “immediately adjacent” means there are no intervening elements between the immediately adjacent elements, for example, between regions, segments, nucleotides and/or nucleosides.

As used herein, “increasing HDL” or “raising HDL” means increasing the level of HDL in an animal after administration of at least one compound of the invention, compared to the HDL level in an animal not
35 administered any compound.

As used herein, “individual” or “subject” or “animal” means a human or non-human animal selected for treatment or therapy.

As used herein, “individual in need thereof” refers to a human or non-human animal selected for treatment or therapy that is in need of such treatment or therapy.

As used herein, “induce”, “inhibit”, “potentiate”, “elevate”, “increase”, “decrease”, “reduce” or the like denote quantitative differences between two states. For example, “an amount effective to inhibit the activity or expression of apo(a)” means that the level of activity or expression of apo(a) in a treated sample will differ from the level of apo(a) activity or expression in an untreated sample. Such terms are applied to, for example, levels of expression, and levels of activity.

As used herein, “inflammatory condition” refers to a disease, disease state, syndrome, or other condition resulting in inflammation. For example, rheumatoid arthritis and liver fibrosis are inflammatory conditions. Other examples of inflammatory conditions include sepsis, myocardial ischemia/reperfusion injury, adult respiratory distress syndrome, nephritis, graft rejection, inflammatory bowel disease, multiple sclerosis, arteriosclerosis, atherosclerosis and vasculitis.

As used herein, “inhibiting the expression or activity” refers to a reduction or blockade of the expression or activity of a RNA or protein and does not necessarily indicate a total elimination of expression or activity.

As used herein, “insulin resistance” is defined as the condition in which normal amounts of insulin are inadequate to produce a normal insulin response from fat, muscle and liver cells. Insulin resistance in fat cells results in hydrolysis of stored triglycerides, which elevates free fatty acids in the blood plasma. Insulin resistance in muscle reduces glucose uptake whereas insulin resistance in liver reduces glucose storage, with both effects serving to elevate blood glucose. High plasma levels of insulin and glucose due to insulin resistance often leads to metabolic syndrome and type 2 diabetes.

As used herein, “insulin sensitivity” is a measure of how effectively an individual processes glucose. An individual having high insulin sensitivity effectively processes glucose whereas an individual with low insulin sensitivity does not effectively process glucose.

As used herein, “lipid-lowering” means a reduction in one or more lipids (e.g., LDL, VLDL) in a subject. “Lipid-raising” means an increase in a lipid (e.g., HDL) in a subject. Lipid-lowering or lipid-raising can occur with one or more doses over time.

As used herein, “lipid-lowering therapy” or “lipid lowering agent” means a therapeutic regimen provided to a subject to reduce one or more lipids in a subject. In certain embodiments, a lipid-lowering therapy is provided to reduce one or more of apo(a), CETP, apoB, total cholesterol, LDL-C, VLDL-C, IDL-C, non-HDL-C, triglycerides, small dense LDL particles, and Lp(a) in a subject. Examples of lipid-lowering therapy include, but are not limited to, apoB inhibitors, statins, fibrates and MTP inhibitors.

As used herein, “lipoprotein”, such as VLDL, LDL and HDL, refers to a group of proteins found in the serum, plasma and lymph and are important for lipid transport. The chemical composition of each

lipoprotein differs, for example, in that the HDL has a higher proportion of protein versus lipid, whereas the VLDL has a lower proportion of protein versus lipid.

As used herein, “Lp(a)” comprises apo(a) and a LDL like particle containing apoB. The apo(a) is linked to the apoB by a disulfide bond.

5 As used herein, “low density lipoprotein-cholesterol (LDL-C)” means cholesterol carried in low density lipoprotein particles. Concentration of LDL-C in serum (or plasma) is typically quantified in mg/dL or nmol/L. “Serum LDL-C” and “plasma LDL-C” mean LDL-C in the serum and plasma, respectively.

As used herein, “major risk factors” refers to factors that contribute to a high risk for a particular disease or condition. In certain embodiments, major risk factors for coronary heart disease include, without
10 limitation, cigarette smoking, hypertension, high LDL, low HDL-C, family history of coronary heart disease, age, and other factors disclosed herein.

As used herein, “metabolic disorder” or “metabolic disease” refers to a condition characterized by an alteration or disturbance in metabolic function. “Metabolic” and “metabolism” are terms well known in the art and generally include the whole range of biochemical processes that occur within a living organism.
15 Metabolic disorders include, but are not limited to, hyperglycemia, prediabetes, diabetes (type 1 and type 2), obesity, insulin resistance, metabolic syndrome and dyslipidemia due to type 2 diabetes.

As used herein, “metabolic syndrome” means a condition characterized by a clustering of lipid and non-lipid cardiovascular risk factors of metabolic origin. In certain embodiments, metabolic syndrome is identified by the presence of any 3 of the following factors: waist circumference of greater than 102 cm in
20 men or greater than 88 cm in women; serum triglyceride of at least 150 mg/dL; HDL-C less than 40 mg/dL in men or less than 50 mg/dL in women; blood pressure of at least 130/85 mmHg; and fasting glucose of at least 110 mg/dL. These determinants can be readily measured in clinical practice (JAMA, 2001, 285: 2486-2497).

“Parenteral administration” means administration through injection or infusion. Parenteral administration includes subcutaneous administration, intravenous administration, intramuscular
25 administration, intraarterial administration, intraperitoneal administration, or intracranial administration, e.g. intrathecal or intracerebroventricular administration. Administration can be continuous, chronic, short or intermittent.

As used herein, “peptide” means a molecule formed by linking at least two amino acids by amide bonds. Peptide refers to polypeptides and proteins.

30 As used herein, “pharmaceutical agent” means a substance that provides a therapeutic benefit when administered to an individual. For example, in certain embodiments, an antisense oligonucleotide targeted to apo(a) is a pharmaceutical agent.

As used herein, “pharmaceutical composition” or “composition” means a mixture of substances suitable for administering to an individual. For example, a pharmaceutical composition can comprise one or
35 more active agents and a pharmaceutical carrier e.g., a sterile aqueous solution.

As used herein, “pharmaceutically acceptable derivative” encompasses derivatives of the compounds described herein such as solvates, hydrates, esters, prodrugs, polymorphs, isomers, isotopically labelled variants, pharmaceutically acceptable salts and other derivatives known in the art.

As used herein, “pharmaceutically acceptable salts” means physiologically and pharmaceutically acceptable salts of antisense compounds, i.e., salts that retain the desired biological activity of the parent compound and do not impart undesired toxicological effects thereto. The term “pharmaceutically acceptable salt” or “salt” includes a salt prepared from pharmaceutically acceptable non-toxic acids or bases, including inorganic or organic acids and bases. “Pharmaceutically acceptable salts” of the compounds described herein may be prepared by methods well-known in the art. For a review of pharmaceutically acceptable salts, see Stahl and Wermuth, Handbook of Pharmaceutical Salts: Properties, Selection and Use (Wiley-VCH, Weinheim, Germany, 2002). Sodium salts of antisense oligonucleotides are useful and are well accepted for therapeutic administration to humans. Accordingly, in one embodiment the compounds described herein are in the form of a sodium salt.

As used herein, “portion” means a defined number of contiguous (i.e. linked) nucleobases of a nucleic acid. In certain embodiments, a portion is a defined number of contiguous nucleobases of a target nucleic acid. In certain embodiments, a portion is a defined number of contiguous nucleobases of an antisense compound.

As used herein, “prevent” or “preventing” refers to delaying or forestalling the onset or development of a disease, disorder, or condition for a period of time from minutes to indefinitely. Prevent also means reducing risk of developing a disease, disorder, or condition.

As used herein, “raise” means to increase in amount. For example, to raise plasma HDL levels means to increase the amount of HDL in the plasma.

As used herein, “reduce” means to bring down to a smaller extent, size, amount, or number. For example, to reduce plasma triglyceride levels means to bring down the amount of triglyceride in the plasma.

As used herein, “region” or “target region” is defined as a portion of the target nucleic acid having at least one identifiable structure, function, or characteristic. For example, a target region may encompass a 3’ UTR, a 5’ UTR, an exon, an intron, an exon/intron junction, a coding region, a translation initiation region, translation termination region, or other defined nucleic acid region. The structurally defined regions for apo(a) can be obtained by accession number from sequence databases such as NCBI and such information is incorporated herein by reference. In certain embodiments, a target region may encompass the sequence from a 5’ target site of one target segment within the target region to a 3’ target site of another target segment within the target region.

As used herein, “second agent” or “second therapeutic agent” means an agent that can be used in combination with a “first agent”. A second therapeutic agent can include, but is not limited to, antisense oligonucleotides targeting apo(a) or apoB. A second agent can also include anti- apo(a) antibodies, apo(a)

peptide inhibitors, cholesterol lowering agents, lipid lowering agents, glucose lowering agents and anti-inflammatory agents.

As used herein, “segments” are defined as smaller, sub-portions of regions within a nucleic acid. For example, a “target segment” means the sequence of nucleotides of a target nucleic acid to which one or more antisense compounds is targeted. “5’ target site” refers to the 5’-most nucleotide of a target segment. “3’ target site” refers to the 3’-most nucleotide of a target segment. Alternatively, a “start site” can refer to the 5’-most nucleotide of a target segment and a “stop site” refers to the 3’-most nucleotide of a target segment. A target segment can also begin at the “start site” of one sequence and end at the “stop site” of another sequence.

As used herein, “statin” means an agent that inhibits the activity of HMG-CoA reductase.

As used herein, “subcutaneous administration” means administration just below the skin.

As used herein, “subject” means a human or non-human animal selected for treatment or therapy.

As used herein, “symptom of cardiovascular disease or disorder” means a phenomenon that arises from and accompanies the cardiovascular disease or disorder and serves as an indication of it. For example, angina; chest pain; shortness of breath; palpitations; weakness; dizziness; nausea; sweating; tachycardia; bradycardia; arrhythmia; atrial fibrillation; swelling in the lower extremities; cyanosis; fatigue; fainting; numbness of the face; numbness of the limbs; claudication or cramping of muscles; bloating of the abdomen; or fever are symptoms of cardiovascular disease or disorder.

As used herein, “targeting” or “targeted” means the process of design and selection of an antisense compound that will specifically hybridize to a target nucleic acid and induce a desired effect.

As used herein, “therapeutically effective amount” means an amount of a pharmaceutical agent that provides a therapeutic benefit to an individual.

As used herein, “therapeutic lifestyle change” means dietary and lifestyle changes intended to lower fat/adipose tissue mass and/or cholesterol. Such change can reduce the risk of developing heart disease, and may includes recommendations for dietary intake of total daily calories, total fat, saturated fat, polyunsaturated fat, monounsaturated fat, carbohydrate, protein, cholesterol, insoluble fiber, as well as recommendations for physical activity.

As used herein, “treat” or “treating” refers to administering a compound described herein to effect an alteration or improvement of a disease, disorder, or condition.

As used herein, “triglyceride” or “TG” means a lipid or neutral fat consisting of glycerol combined with three fatty acid molecules.

As used herein, “type 2 diabetes,” (also known as “type 2 diabetes mellitus”, “diabetes mellitus, type 2”, “non-insulin-dependent diabetes”, “NIDDM”, “obesity related diabetes”, or “adult-onset diabetes”) is a metabolic disorder that is primarily characterized by insulin resistance, relative insulin deficiency, and hyperglycemia.

Certain Embodiments

In certain embodiments, a compound comprises a siRNA or antisense oligonucleotide targeted to apolipoprotein(a) (apo(a)) known in the art and a conjugate group described herein. Examples of antisense oligonucleotides targeted to apo(a) suitable for conjugation include but are not limited to those disclosed in
5 WO 2013/177468; US 8,673,632; US 7,259,150; and US Patent Application Publication No. US 2004/0242516; which are incorporated by reference in their entireties herein. In certain embodiments, a compound comprises an antisense oligonucleotide having a nucleobase sequence of any of SEQ ID NOs 12-130, 133, 134 disclosed in WO 2013/177468 and a conjugate group described herein. In certain
10 embodiments, a compound comprises an antisense oligonucleotide having a nucleobase sequence of any of SEQ ID NOs 11-45 and 85-96 disclosed in US 8,673,632 and a conjugate group described herein. In certain embodiments, a compound comprises an antisense oligonucleotide having a nucleobase sequence of any of SEQ ID NOs 11-45 disclosed in US 7,259,150 and a conjugate group described herein. In certain
15 embodiments, a compound comprises an antisense oligonucleotide having a nucleobase sequence of any of SEQ ID NOs 7-41 disclosed in US Patent Application Publication No. US 2004/0242516 and a conjugate group described herein. The nucleobase sequences of all of the aforementioned referenced SEQ ID NOs are incorporated by reference herein.

Certain embodiments provide a compounds and methods for decreasing apo(a) mRNA and protein expression. In certain embodiments, the compound is an apo(a) specific inhibitor for treating, preventing, or ameliorating an apo(a) associated disease. In certain embodiments, the compound is an antisense
20 oligonucleotide targeting apo(a). In certain embodiments, the compound is an antisense oligonucleotide targeting apo(a) and a conjugate group.

Certain embodiments provide a compounds and methods for decreasing Lp(a) levels. In certain embodiments, the compound is an apo(a) specific inhibitor for treating, preventing, or ameliorating an Lp(a) associated disease. In certain embodiments, the compound is an antisense oligonucleotide targeting apo(a). In
25 certain embodiments, the compound is an antisense oligonucleotide targeting apo(a) and a conjugate group.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides. In certain embodiments, the modified oligonucleotide with the conjugate group consists of 15 to 30, 18 to 24, 19 to 22, 13 to 25, 14 to 25, 15 to 25 linked nucleosides. In certain embodiments, the modified oligonucleotide
30 with the conjugate group comprises at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, at least 27, at least 28, at least 29 or 30 linked nucleosides. In certain embodiments, the modified oligonucleotide with the conjugate group consists of 20 linked nucleosides.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a)
35 and a conjugate group, wherein the modified oligonucleotide comprises at least 8, at least 9, at least 10, at

least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases complementary to an equal length portion of any of SEQ ID NOs: 1-4.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting an apo(a) segment and a conjugate group, wherein the modified oligonucleotide comprises at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases complementary to an equal length portion of any of the target segments shown in, for example, Examples 114 and 117. In the tables, the "Start Site" refers to the 5'-most nucleotide of a target segment and "Stop Site" refers to the 3'-most nucleotide of a target segment. A target segment can range from the start site to the stop site of each sequence listed in the tables. Alternatively, the target segment can range from the start site of one sequence and end at the stop site of another sequence. For example, as shown in Table 125, a target segment can range from 3901-3920, the start site to the stop site of SEQ ID NO: 58. In another example, as shown in Table 125, a target segment can range from 3900-3923, the start site of SEQ ID NO: 57 to the stop site of SEQ ID NO: 61.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the nucleobase sequence of the modified oligonucleotide is at least 80%, at least 85%, at least 90%, at least 95%, or 100% complementary to any of SEQ ID NOs: 1-4. Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the nucleobase sequence of the modified oligonucleotide is at least 80%, at least 85%, at least 90%, at least 95%, or 100% complementary to any of the target segments shown in, for example, Examples 114 and 117.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and comprises a nucleobase sequence comprising a portion of at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases complementary to an equal length portion of nucleobases 3901 to 3920 of SEQ ID NO: 1, wherein the nucleobase sequence of the modified oligonucleotide is at least 80% complementary to SEQ ID NO: 1.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and comprises a nucleobase sequence comprising at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, at least 27, at least 28, at least 29 or 30 contiguous nucleobases complementary to an equal length portion of nucleobases 3900 to 3923 of SEQ ID NO: 1, wherein the nucleobase sequence of the modified oligonucleotide is at least 80% complementary to SEQ ID NO: 1.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and has

a nucleobase sequence comprising at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases of any of the nucleobase sequences of SEQ ID NOs: 12-130, 133, 134. In certain embodiments, the modified oligonucleotide has a nucleobase sequence comprising at least 8 contiguous nucleobases of any one of the nucleobase sequences of SEQ ID NOs: 12-130, 133, 134. In certain embodiments, the compound consists of any one of SEQ ID NOs: 12-130, 133, 134 and a conjugate group.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and has a nucleobase sequence comprising at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases of any of the nucleobase sequences of SEQ ID NOs: 12-20, 22-33, 35-44, 47-50, 51, 53, 57-62, 65-66, 68, 70-79, 81, 85-86, 89-90, 92-94, 97, 105-110, 103-104, 133-134. In certain embodiments, the compound consists of any of the nucleobase sequences of SEQ ID NOs: 12-20, 22-33, 35-44, 47-50, 51, 53, 57-62, 65-66, 68, 70-79, 81, 85-86, 89-90, 92-94, 97, 105-110, 103-104, 133-134 and a conjugate group.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and has a nucleobase sequence comprising at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases of any of the nucleobase sequences of SEQ ID NOs: 12-19, 26-30, 32, 35, 38-44, 46-47, 50, 57-58, 61, 64-66, 68, 72-74, 76-77, 92-94, 103-110. In certain embodiments, the compound consists of any of the nucleobase sequences of SEQ ID NOs: 12-19, 26-30, 32, 35, 38-44, 46-47, 50, 57-58, 61, 64-66, 68, 72-74, 76-77, 92-94, 103-110 and a conjugate group.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and has a nucleobase sequence comprising at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases of any of the nucleobase sequences of SEQ ID NOs: 111, 114-121, 123-129. In certain embodiments, the compound consists of any of the nucleobase sequences of SEQ ID NOs: 111, 114-121, 123-129 and a conjugate group.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and has a nucleobase sequence comprising at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases of any of the nucleobase sequences of SEQ ID NOs: 14, 17, 18, 26-28, 39, 71, 106-107. In certain embodiments, the compound consists of any of the nucleobase sequences of SEQ ID NOs: 14, 17, 18, 26-28, 39, 71, 106-107 and a conjugate group.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and has a nucleobase sequence comprising at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases of any of the nucleobase sequences of SEQ ID NOs: 14, 26-29, 39-40, 82. In certain embodiments, the compound consists of any of the nucleobase sequences of SEQ ID NOs: 14, 26-29, 39-40, 82 and a conjugate group.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and has a nucleobase sequence comprising at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases of any of the nucleobase sequences of SEQ ID NOs: 14, 16-18. In certain embodiments, the compound consists of any of the nucleobase sequences of SEQ ID NOs: 14, 16-18 and a conjugate group.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and has a nucleobase sequence comprising at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases of any of the nucleobase sequences of SEQ ID NOs: 26-27, 107. In certain embodiments, the compound consists of any of the nucleobase sequences of SEQ ID NOs: 26-27, 107 and a conjugate group.

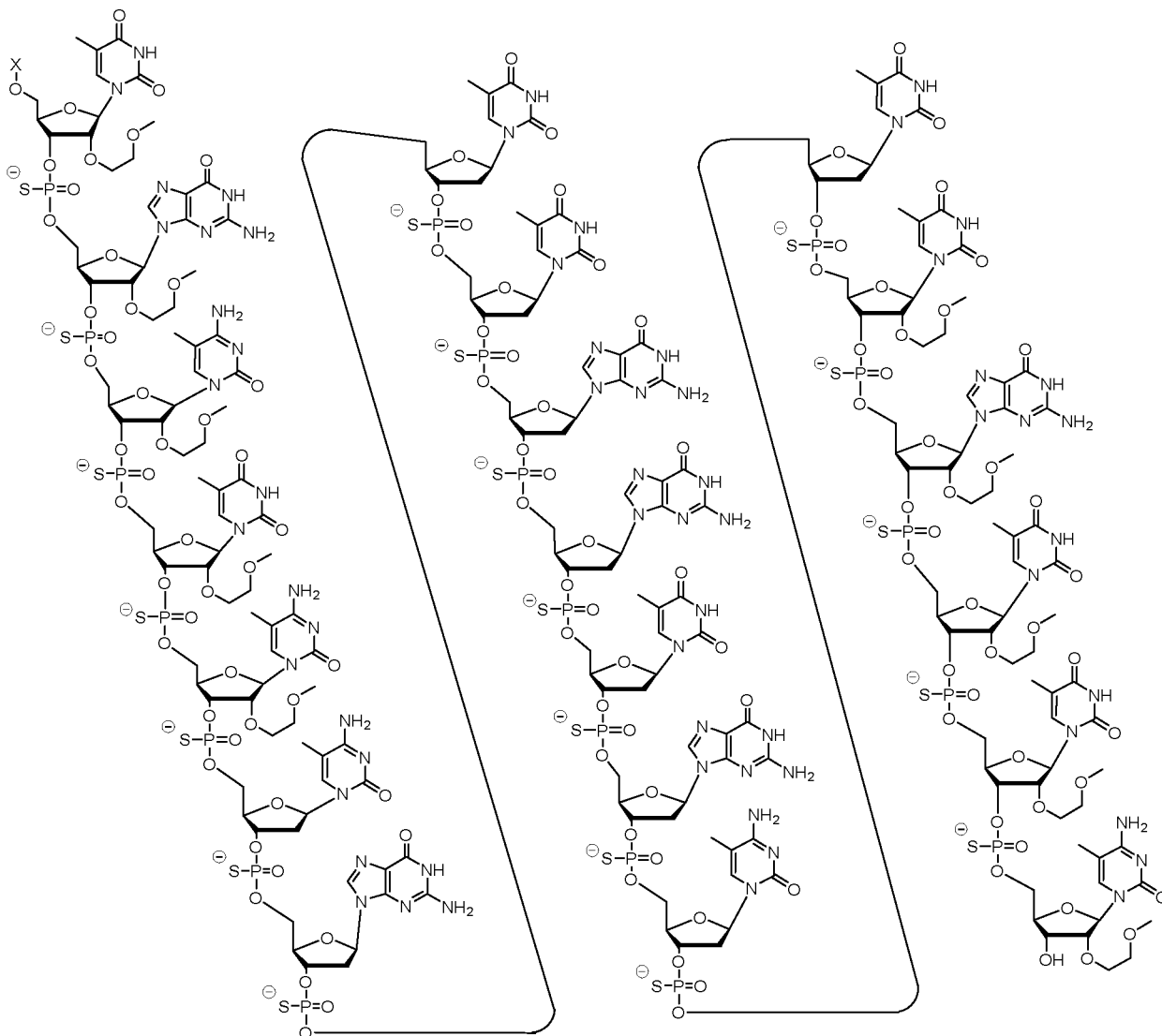
Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and has a nucleobase sequence comprising at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases of any of the nucleobase sequences of SEQ ID NOs: 28-29, 39-40, 47. In certain embodiments, the compound consists of any of the nucleobase sequences of SEQ ID NOs: 28-29, 39-40, 47 and a conjugate group.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and has a nucleobase sequence comprising at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases of any of the nucleobase sequences of SEQ ID NOs: 28, 93, 104, 134. In certain embodiments, the compound consists of any of the nucleobase sequences of SEQ ID NOs: 28, 93, 104, 134 and a conjugate group.

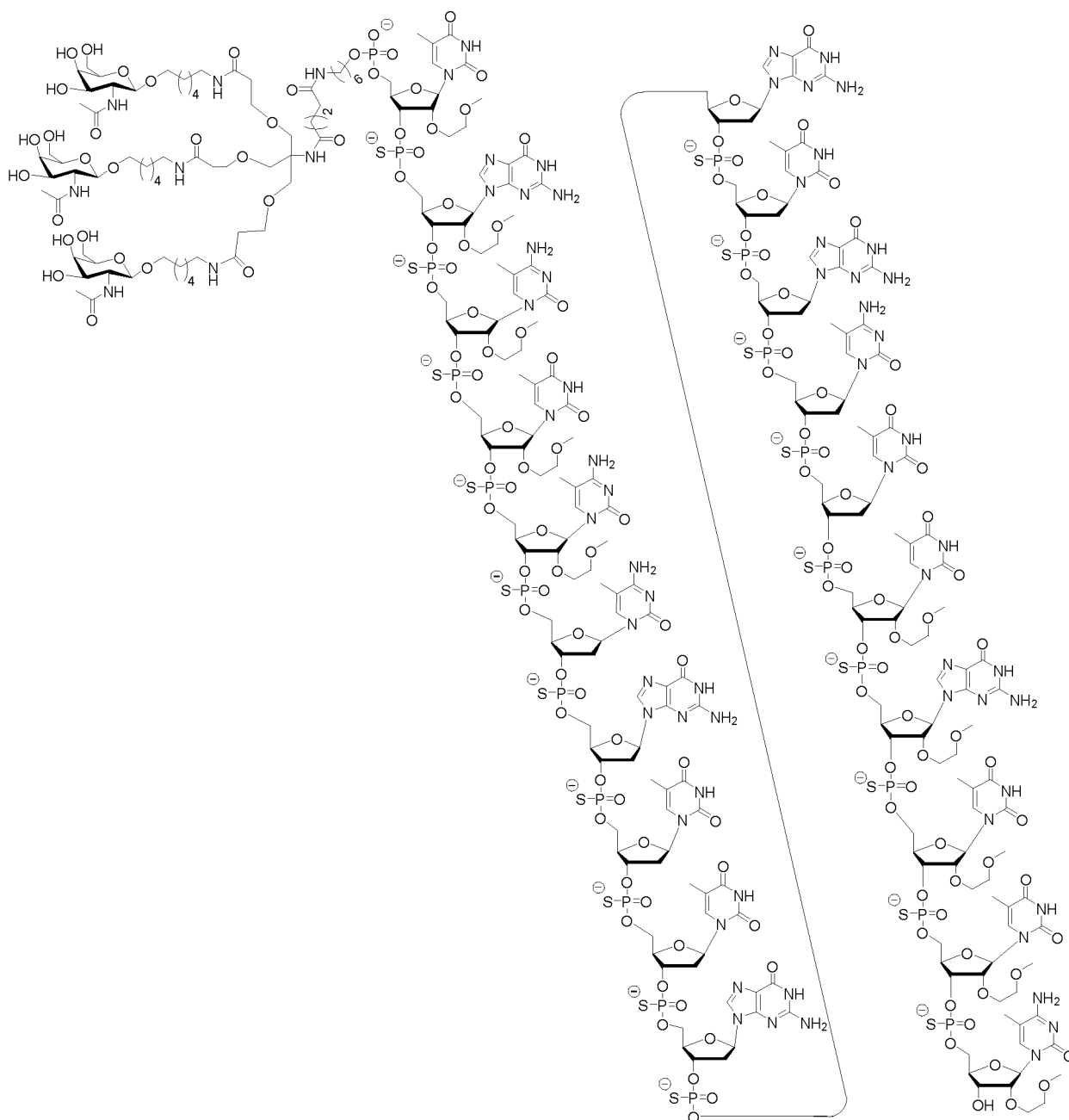
Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and has a nucleobase sequence comprising at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases of the nucleobase sequence of SEQ ID NO: 58. In certain embodiments, the modified oligonucleotide with the conjugate group

has a nucleobase sequence comprising at least 8 contiguous nucleobases of the nucleobase sequence of SEQ ID NO: 58. In certain embodiments, the compound consists of SEQ ID NO: 58 and a conjugate group.

In certain embodiments, the present disclosure provides conjugated antisense compounds represented by the following structure. In certain embodiments, the antisense compound comprises the modified oligonucleotide ISIS 494372 with a 5'-X, wherein X is a conjugate group comprising GalNAc. In certain
5
embodiments, the antisense compound consists of the modified oligonucleotide ISIS 494372 with a 5'-X, wherein X is a conjugate group comprising GalNAc.



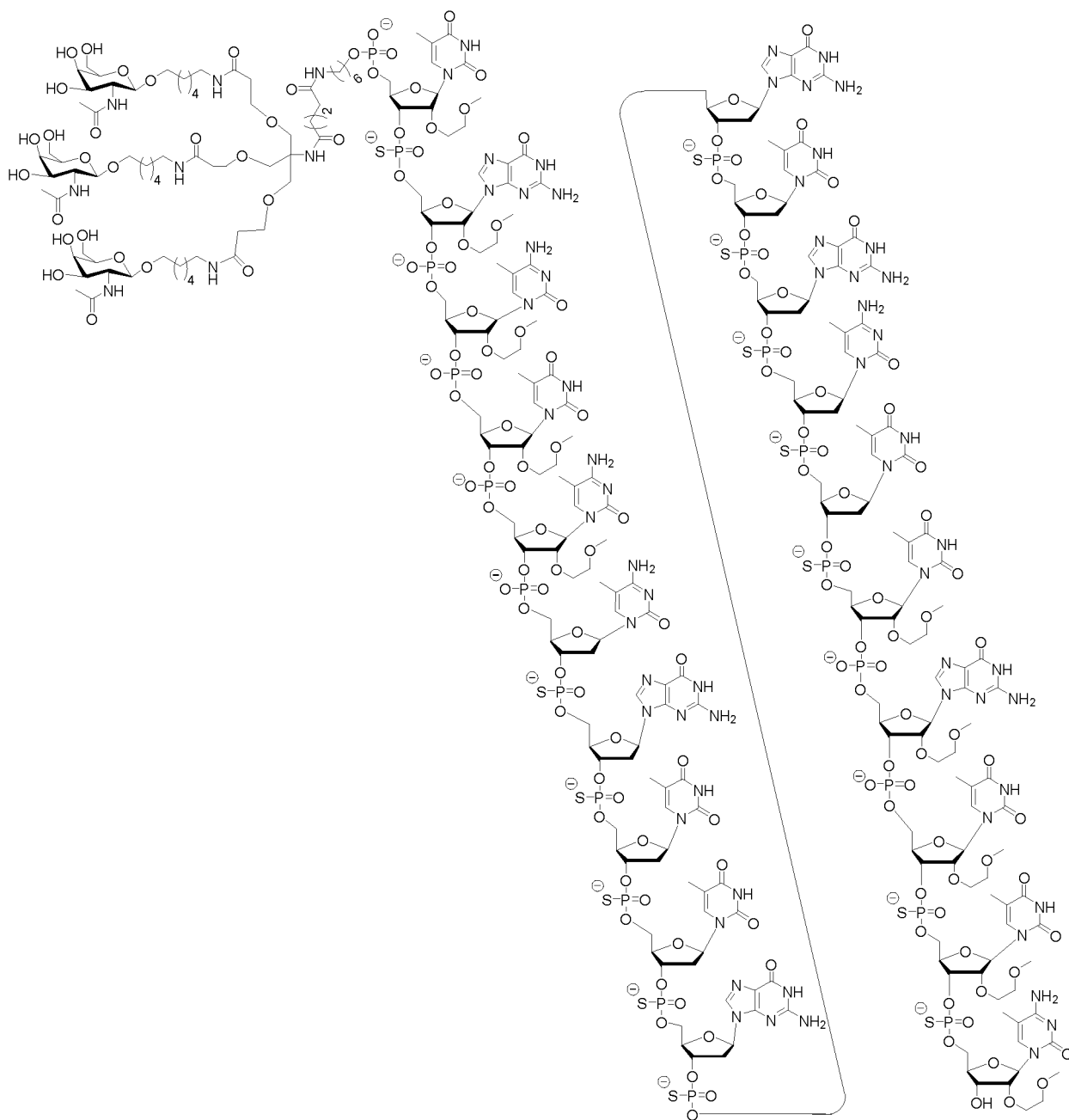
In certain embodiments, the present disclosure provides conjugated antisense compounds represented by the following structure. In certain embodiments, the antisense compound comprises the conjugated modified oligonucleotide ISIS 681251. In certain embodiments, the antisense compound consists of the conjugated modified oligonucleotide ISIS 681251.



5

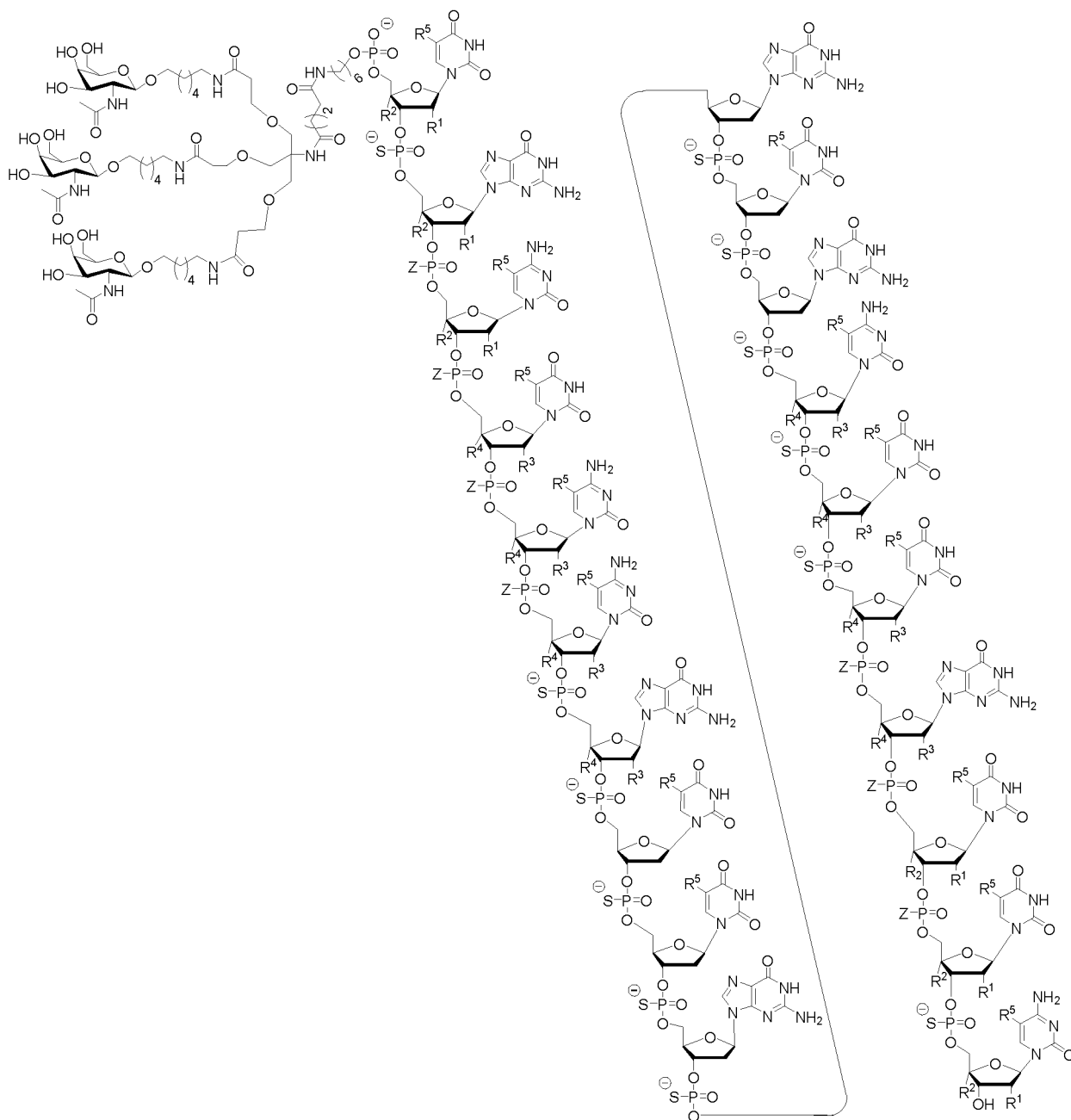
In certain embodiments, the present disclosure provides conjugated antisense compounds represented by the following structure. In certain embodiments, the antisense compound comprises the conjugated modified oligonucleotide ISIS 681257. In certain embodiments, the antisense compound consists of the conjugated modified oligonucleotide ISIS 681257.

5



In certain embodiments, the present disclosure provides conjugated antisense compounds represented by the following structure. In certain embodiments, the antisense compound comprises a modified oligonucleotide with the nucleobase sequence of SEQ ID NO: 58 with a 5'-GalNAc with variability in the sugar mods of the wings.

5 oligonucleotide with the nucleobase sequence of SEQ ID NO: 58 with a 5'-GalNAc with variability in the sugar mods of the wings.



Wherein either R^1 is $-\text{OCH}_2\text{CH}_2\text{OCH}_3$ (MOE) and R^2 is H; or R^1 and R^2 together form a bridge, wherein R^1 is $-\text{O}-$ and R^2 is $-\text{CH}_2-$, $-\text{CH}(\text{CH}_3)-$, or $-\text{CH}_2\text{CH}_2-$, and R^1 and R^2 are directly connected such that the resulting bridge is selected from: $-\text{O}-\text{CH}_2-$, $-\text{O}-\text{CH}(\text{CH}_3)-$, and $-\text{O}-\text{CH}_2\text{CH}_2-$;

10

And for each pair of R^3 and R^4 on the same ring, independently for each ring: either R^3 is selected from H and $-OCH_2CH_2OCH_3$ and R^4 is H; or R^3 and R^4 together form a bridge, wherein R^3 is $-O-$, and R^4 is $-CH_2-$, $-CH(CH_3)-$, or $-CH_2CH_2-$ and R^3 and R^4 are directly connected such that the resulting bridge is selected from: $-O-CH_2-$, $-O-CH(CH_3)-$, and $-O-CH_2CH_2-$;

5 And R^5 is selected from H and $-CH_3$;

And Z is selected from S^- and O^- .

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide is single-stranded.

10 Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein at least one internucleoside linkage is a modified internucleoside linkage. In certain embodiments, the modified internucleoside linkage is a phosphorothioate internucleoside linkage. In certain embodiments, at least 1, at least 2, at least 3, at least 4, at least 5, at least 6, at least 7, at least 8, at least 9 or at least 10 internucleoside linkages of said modified oligonucleotide are phosphorothioate internucleoside linkages. In certain embodiments, each internucleoside linkage is a phosphorothioate
15 internucleoside linkage. In certain embodiments, the modified oligonucleotide comprises at least 1, at least 2, at least 3, at least 4, at least 5, at least 6, at least 7, at least 8, at least 9 or at least 10 phosphodiester internucleoside linkages. In certain embodiments, each internucleoside linkage of the modified oligonucleotide is selected from a phosphodiester internucleoside linkage and a phosphorothioate internucleoside linkage.

20 Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein at least one nucleoside comprises a modified nucleobase. In certain embodiments, the modified nucleobase is a 5-methylcytosine.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide comprises at least one modified sugar. In certain
25 embodiments, the modified sugar is a bicyclic sugar. In certain embodiments, the modified sugar comprises a 2'-O-methoxyethyl, a constrained ethyl, a 3'-fluoro-HNA or a 4'-(CH_2)_n-O-2' bridge, wherein n is 1 or 2.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and comprises: (a) a gap segment consisting of linked deoxynucleosides; (b) a 5' wing segment consisting of
30 linked nucleosides; (c) a 3' wing segment consisting of linked nucleosides; and wherein the gap segment is positioned between the 5' wing segment and the 3' wing segment and wherein each nucleoside of each wing segment comprises a modified sugar.

Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 20 linked nucleosides and
35 comprises: (a) a gap segment consisting of ten linked deoxynucleosides; (b) a 5' wing segment consisting of

five linked nucleosides; (c) a 3' wing segment consisting of five linked nucleosides; and wherein the gap segment is positioned between the 5' wing segment and the 3' wing segment, wherein each nucleoside of each wing segment comprises a 2'-O-methoxyethyl sugar, wherein at least one internucleoside linkage is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

5 Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 20 linked nucleosides and has a nucleobase sequence comprising at least 8 contiguous nucleobases of any of SEQ ID NOs: 12-130, 133, 134, wherein the modified oligonucleotide comprises: (a) a gap segment consisting of ten linked deoxynucleosides; (b) a 5' wing segment consisting of five linked nucleosides; (c) a 3' wing segment
10 consisting of five linked nucleosides; and wherein the gap segment is positioned between the 5' wing segment and the 3' wing segment, wherein each nucleoside of each wing segment comprises a 2'-O-methoxyethyl sugar, wherein at least one internucleoside linkage is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

 Certain embodiments provide a compound comprising a modified oligonucleotide targeting apo(a)
15 and a conjugate group, wherein the modified oligonucleotide consists of 20 linked nucleosides and has a nucleobase sequence comprising at least 8 contiguous nucleobases of SEQ ID NO: 58, wherein the modified oligonucleotide comprises: (a) a gap segment consisting of ten linked deoxynucleosides; (b) a 5' wing segment consisting of five linked nucleosides; (c) a 3' wing segment consisting of five linked nucleosides; and wherein the gap segment is positioned between the 5' wing segment and the 3' wing segment, wherein
20 each nucleoside of each wing segment comprises a 2'-O-methoxyethyl sugar, wherein at least one internucleoside linkage is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

 Certain embodiments provide a modified oligonucleotide targeting apo(a) and a conjugate group, wherein the modified oligonucleotide consists of 20 linked nucleosides with the nucleobase sequence of SEQ
25 ID NO: 58, wherein the modified oligonucleotide comprises: (a) a gap segment consisting of ten linked deoxynucleosides; (b) a 5' wing segment consisting of five linked nucleosides; (c) a 3' wing segment consisting of five linked nucleosides; and wherein the gap segment is positioned between the 5' wing segment and the 3' wing segment, wherein each nucleoside of each wing segment comprises a 2'-O-methoxyethyl sugar, wherein at least one internucleoside linkage is a phosphorothioate linkage and wherein
30 each cytosine residue is a 5-methylcytosine.

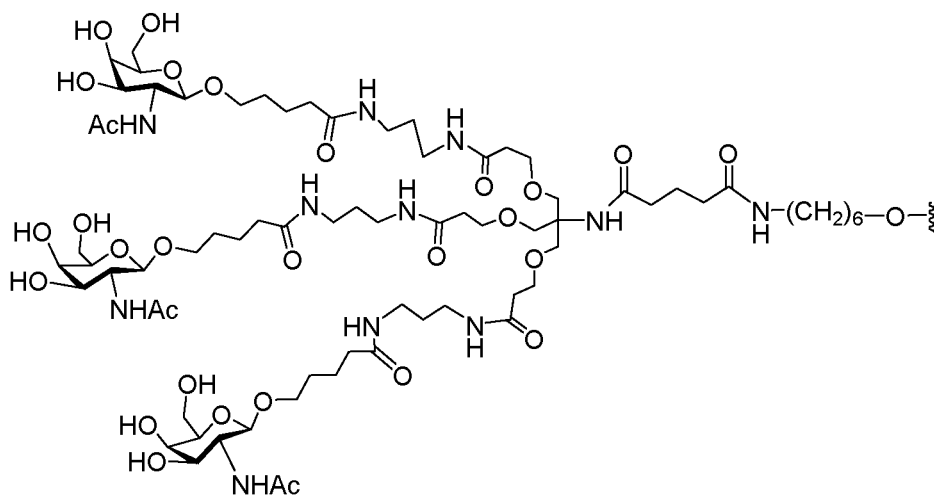
 In certain embodiments, the conjugate group is linked to the modified oligonucleotide at the 5' end of the modified oligonucleotide. In certain embodiments, the conjugate group is linked to the modified oligonucleotide at the 3' end of the modified oligonucleotide.

 In certain embodiments, the conjugate group comprises one or more ligands. In certain embodiments,
35 the conjugate group comprises two or more ligands. In certain embodiments, the conjugate group comprises three or more ligands. In certain embodiments, the conjugate group comprises three ligands. In certain

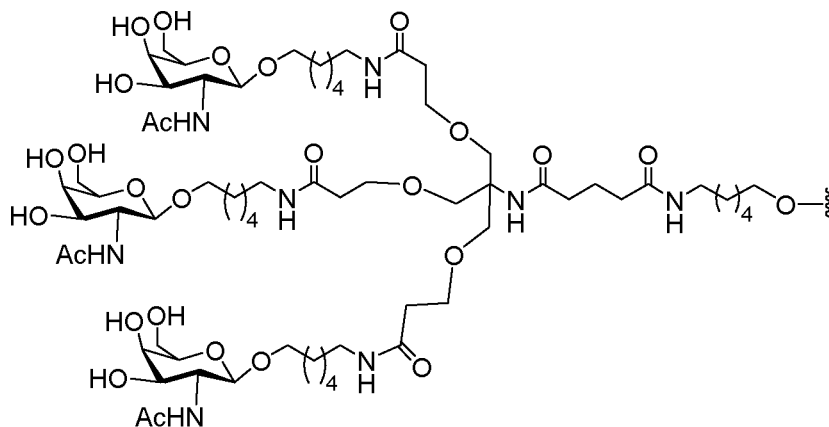
embodiments, each ligand is selected from among: a polysaccharide, modified polysaccharide, mannose, galactose, a mannose derivative, a galactose derivative, D-mannopyranose, L-Mannopyranose, D-Arabinose, L-Galactose, D-xylofuranose, L-xylofuranose, D-glucose, L-glucose, D-Galactose, L-Galactose, α -D-Mannofuranose, β -D-Mannofuranose, α -D-Mannopyranose, β -D-Mannopyranose, α -D-Glucopyranose, β -D-Glucopyranose, α -D-Glucofuranose, β -D-Glucofuranose, α -D-fructofuranose, α -D-fructopyranose, α -D-Galactopyranose, β -D-Galactopyranose, α -D-Galactofuranose, β -D-Galactofuranose, glucosamine, sialic acid, α -D-galactosamine, N-Acetylgalactosamine, 2-Amino-3-*O*-[(*R*)-1-carboxyethyl]-2-deoxy- β -D-glucopyranose, 2-Deoxy-2-methylamino-L-glucopyranose, 4,6-Dideoxy-4-formamido-2,3-di-*O*-methyl-D-mannopyranose, 2-Deoxy-2-sulfoamino-D-glucopyranose, *N*-Glycoloyl- α -neuraminic acid, 5-thio- β -D-glucopyranose, methyl 2,3,4-tri-*O*-acetyl-1-thio-6-*O*-trityl- α -D-glucopyranoside, 4-Thio- β -D-galactopyranose, ethyl 3,4,6,7-tetra-*O*-acetyl-2-deoxy-1,5-dithio- α -D-*gluco*-heptopyranoside, 2,5-Anhydro-D-allonitrile, ribose, D-ribose, D-4-thioribose, L-ribose, L-4-thioribose. In certain embodiments, each ligand is N-acetyl galactosamine.

In certain embodiments, each ligand is N-acetyl galactosamine.

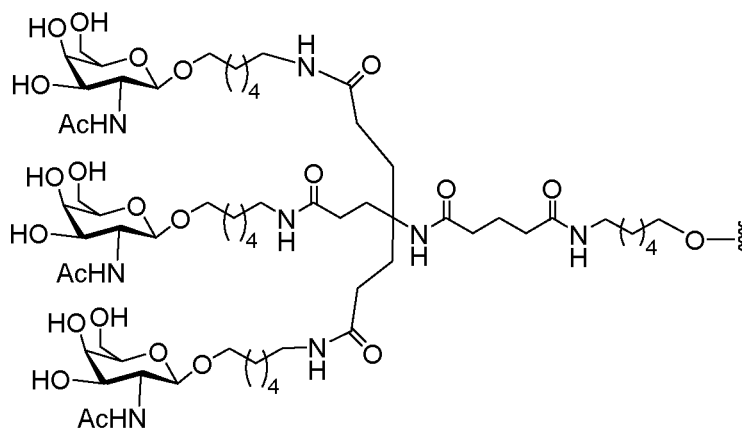
In certain embodiments, the conjugate group comprises:



In certain embodiments, the conjugate group comprises:

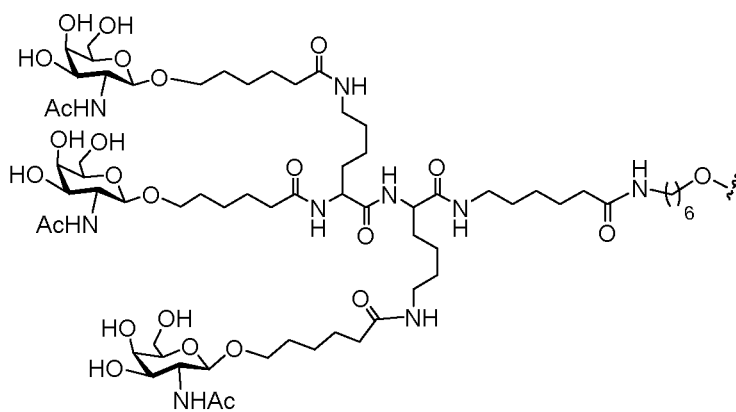


In certain embodiments, the conjugate group comprises:



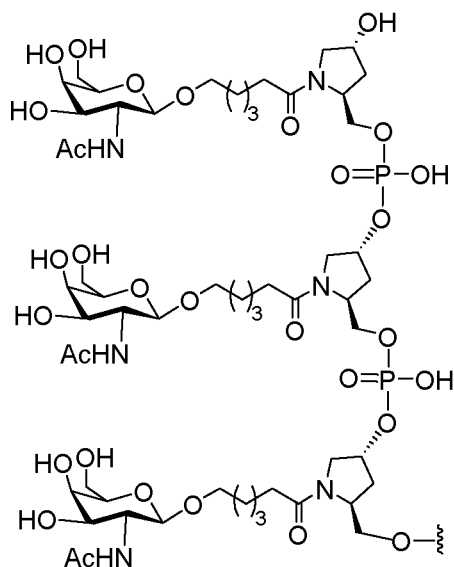
5

In certain embodiments, the conjugate group comprises:



10

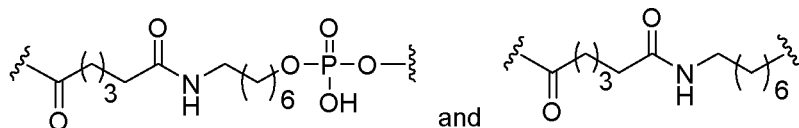
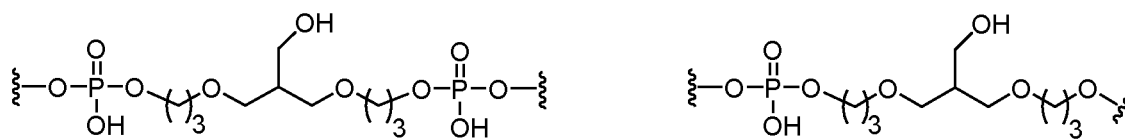
In certain embodiments, the conjugate group comprises:



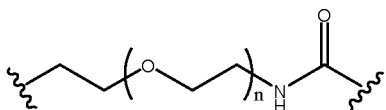
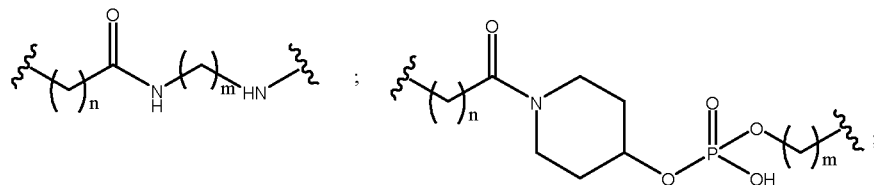
In certain embodiments, the conjugate group comprises at least one phosphorus linking group or neutral linking group.

5

In certain embodiments, the conjugate group comprises a structure selected from among:



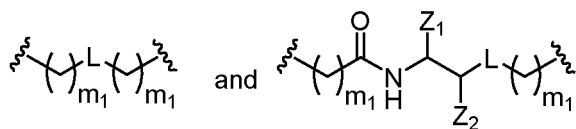
and



wherein n is from 1 to 12; and

wherein m is from 1 to 12.

In certain embodiments, the conjugate group has a tether having a structure selected from among:



wherein L is either a phosphorus linking group or a neutral linking group;

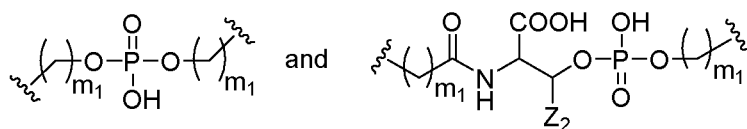
Z1 is C(=O)O-R2;

Z2 is H, C1-C6 alkyl or substituted C1-C6 alkyl;

R2 is H, C1-C6 alkyl or substituted C1-C6 alkyl; and

each m1 is, independently, from 0 to 20 wherein at least one m1 is greater than 0 for each tether.

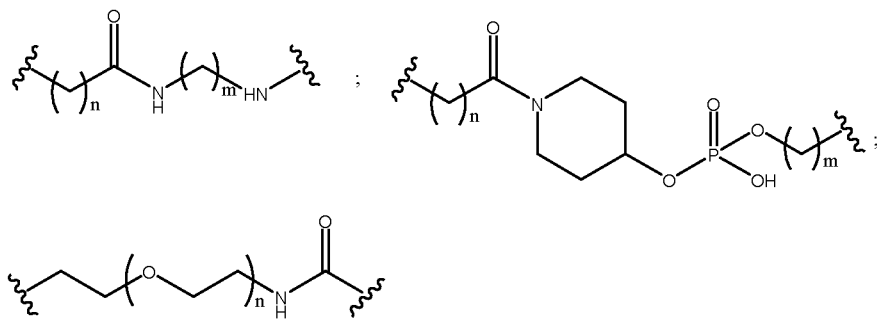
In certain embodiments, conjugate group has a tether having a structure selected from among:



wherein Z2 is H or CH3; and

each m1 is, independently, from 0 to 20 wherein at least one m1 is greater than 0 for each tether.

In certain embodiments, the conjugate group has tether having a structure selected from among:

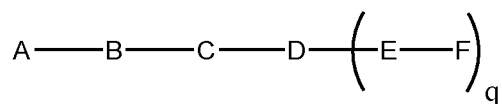


wherein n is from 1 to 12; and

wherein m is from 1 to 12.

In certain embodiments, the conjugate group is covalently attached to the modified oligonucleotide.

In certain embodiments, the compound has a structure represented by the formula:



wherein

A is the modified oligonucleotide;

B is the cleavable moiety

C is the conjugate linker

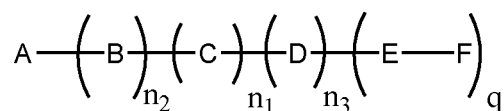
D is the branching group

each E is a tether;

each F is a ligand; and

q is an integer between 1 and 5.

In certain embodiments, the compound has a structure represented by the formula:



wherein:

A is the modified oligonucleotide;

B is the cleavable moiety

C is the conjugate linker

D is the branching group

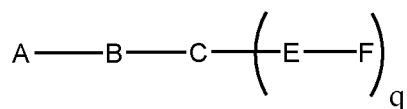
each E is a tether;

each F is a ligand;

each n is independently 0 or 1; and

q is an integer between 1 and 5.

In certain embodiments, the compound has a structure represented by the formula:



wherein

A is the modified oligonucleotide;

B is the cleavable moiety;

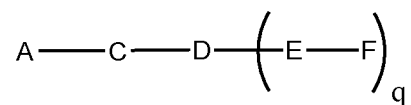
C is the conjugate linker;

each E is a tether;

each F is a ligand; and
q is an integer between 1 and 5.

In certain embodiments, the compound has a structure represented by the formula:

5



wherein

A is the modified oligonucleotide;

C is the conjugate linker;

10

D is the branching group;

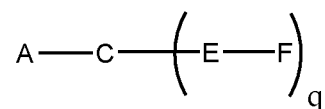
each E is a tether;

each F is a ligand; and

q is an integer between 1 and 5.

15

In certain embodiments, the compound has a structure represented by the formula:



wherein

A is the modified oligonucleotide;

20

C is the conjugate linker;

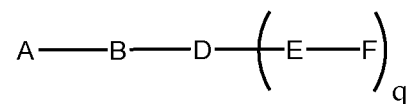
each E is a tether;

each F is a ligand; and

q is an integer between 1 and 5.

25

In certain embodiments, the compound has a structure represented by the formula:



wherein

A is the modified oligonucleotide;

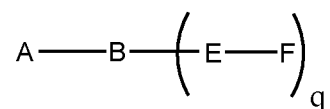
30

B is the cleavable moiety;

D is the branching group;

each E is a tether;
 each F is a ligand; and
 q is an integer between 1 and 5.

- 5 In certain embodiments, the compound has a structure represented by the formula:



wherein

A is the modified oligonucleotide;

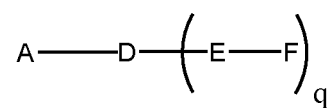
- 10 B is the cleavable moiety;

each E is a tether;

each F is a ligand; and

q is an integer between 1 and 5.

- 15 In certain embodiments, the compound has a structure represented by the formula:



wherein

A is the modified oligonucleotide;

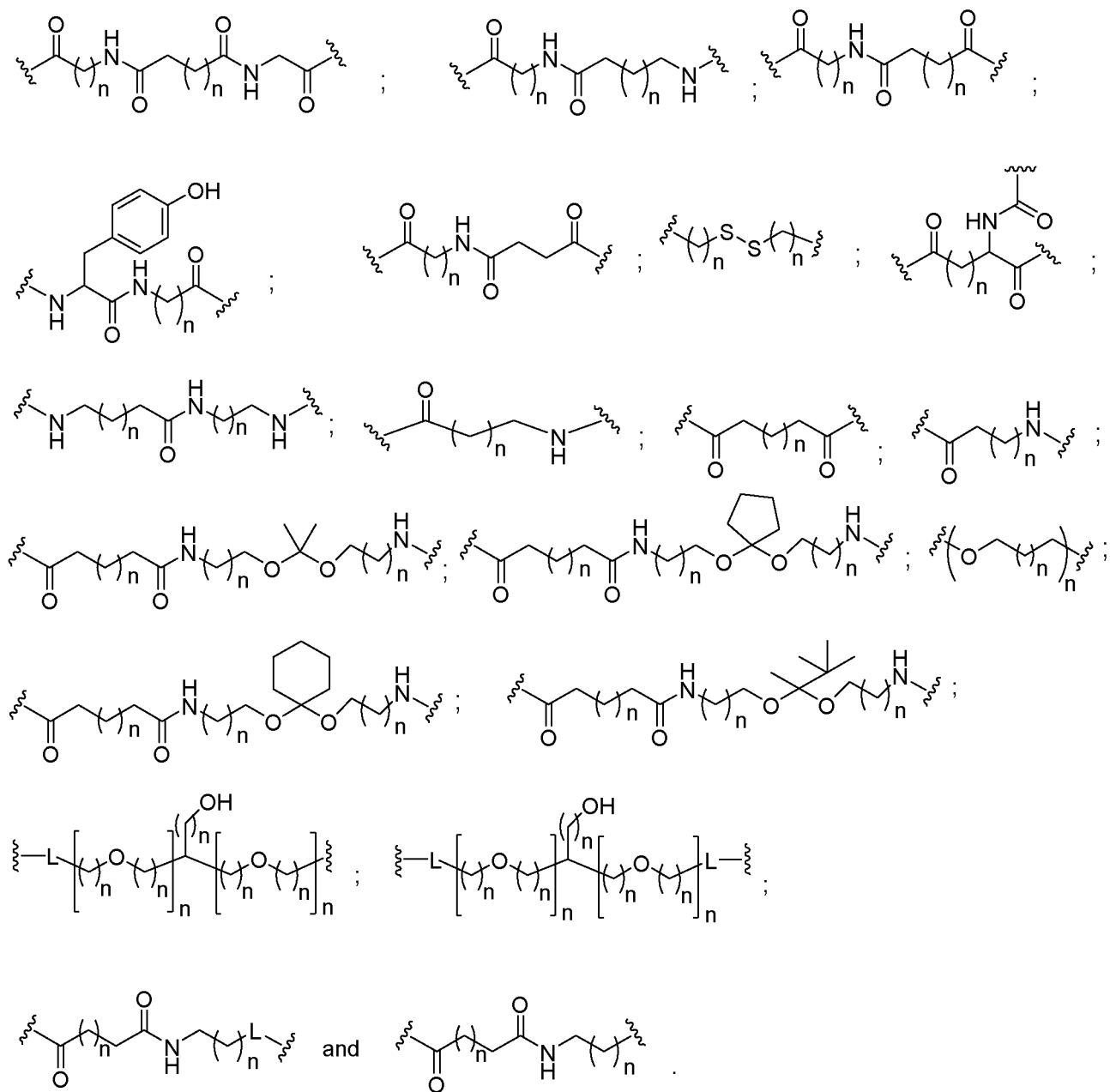
- 20 D is the branching group;

each E is a tether;

each F is a ligand; and

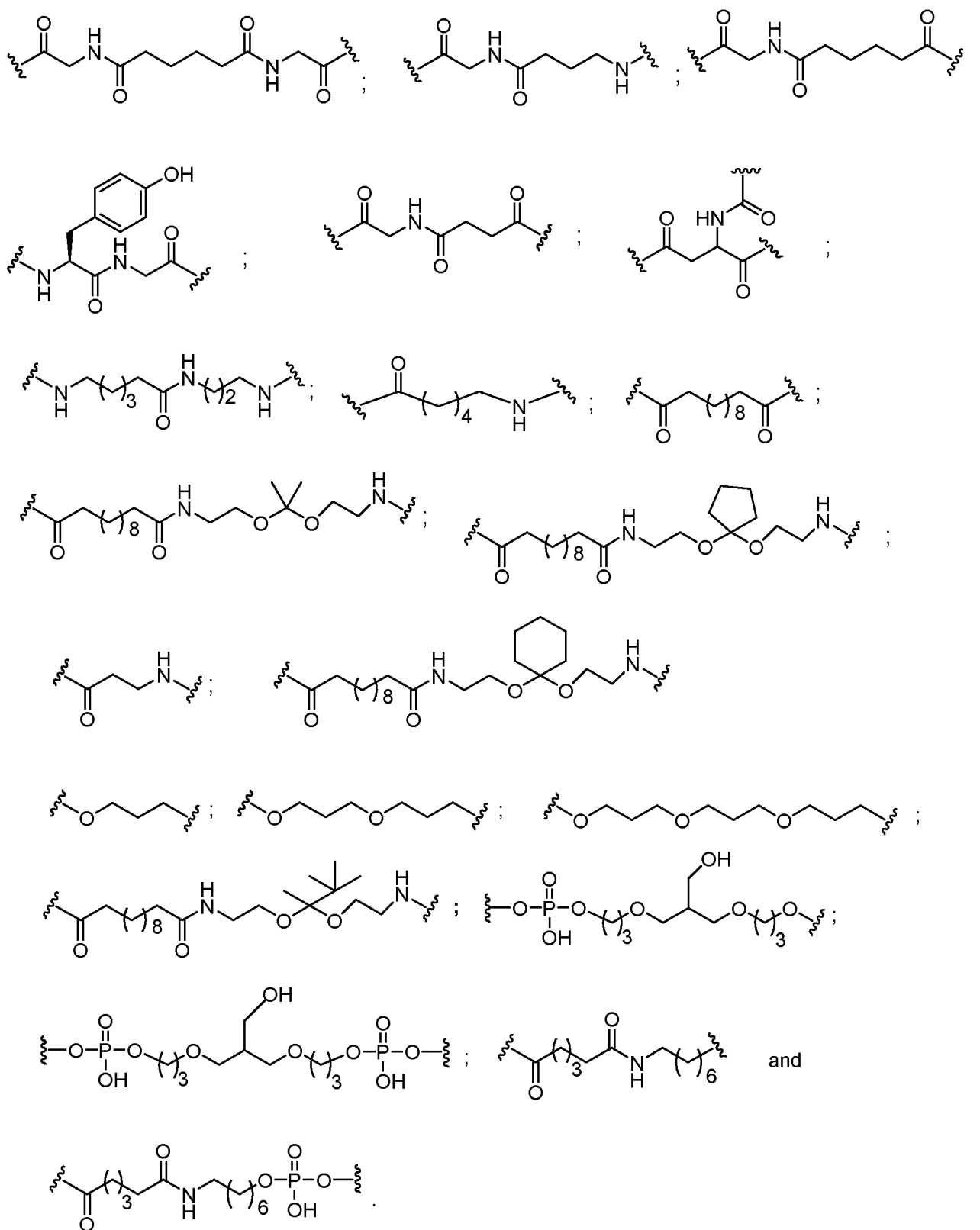
q is an integer between 1 and 5.

In certain embodiments, the conjugate linker has a structure selected from among:

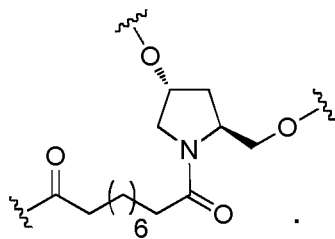


wherein each L is, independently, a phosphorus linking group or a neutral linking group; and
 5 each n is, independently, from 1 to 20.

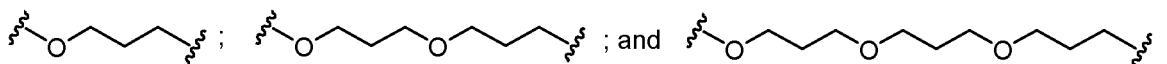
In certain embodiments, the conjugate linker has a structure selected from among:



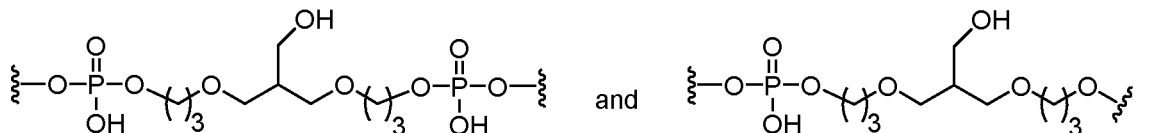
In certain embodiments, the conjugate linker has the following structure:



5 In certain embodiments, the conjugate linker has a structure selected from among:

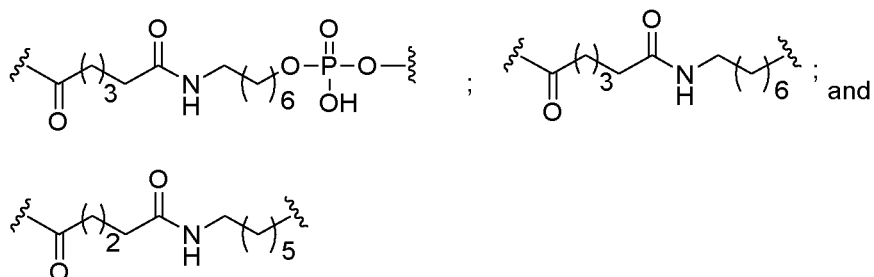


In certain embodiments, the conjugate linker has a structure selected from among:



10

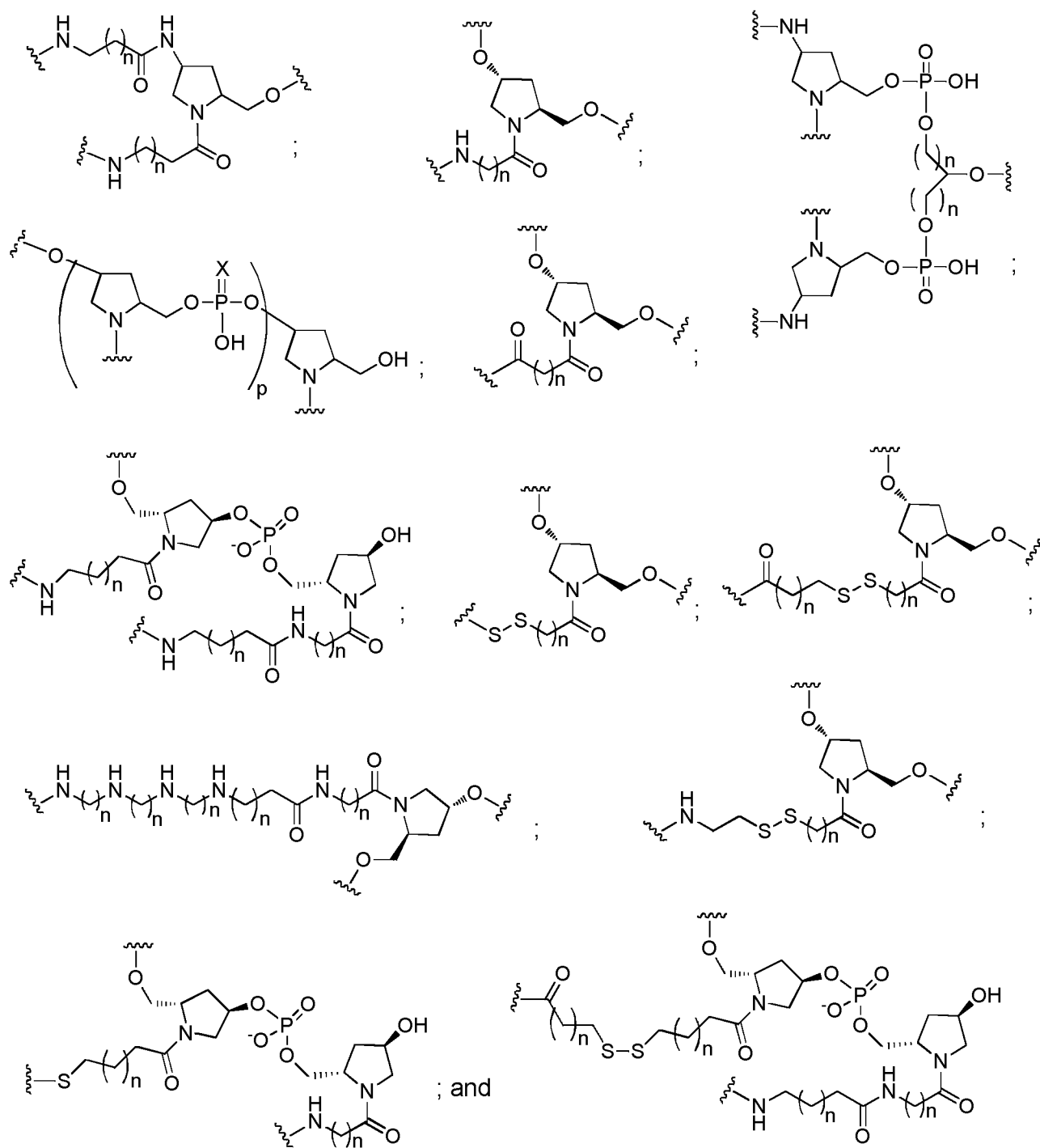
In certain embodiments, the conjugate linker has a structure selected from among:



15 In certain embodiments, the conjugate linker comprises a pyrrolidine. In certain embodiments, the conjugate linker does not comprise a pyrrolidine. In certain embodiments, the conjugate linker comprises PEG. In certain embodiments, the conjugate linker comprises an amide. In certain embodiments, the conjugate linker comprises at least two amides. In certain embodiments, the conjugate linker does not comprise an amide. In certain embodiments, the conjugate linker comprises a polyamide. In certain
20 In certain embodiments, the conjugate linker comprises an amine. In certain embodiments, the conjugate linker comprises one or more disulfide bonds. In certain embodiments, the conjugate linker comprises a protein binding moiety. In certain embodiments, the protein binding moiety comprises a lipid.

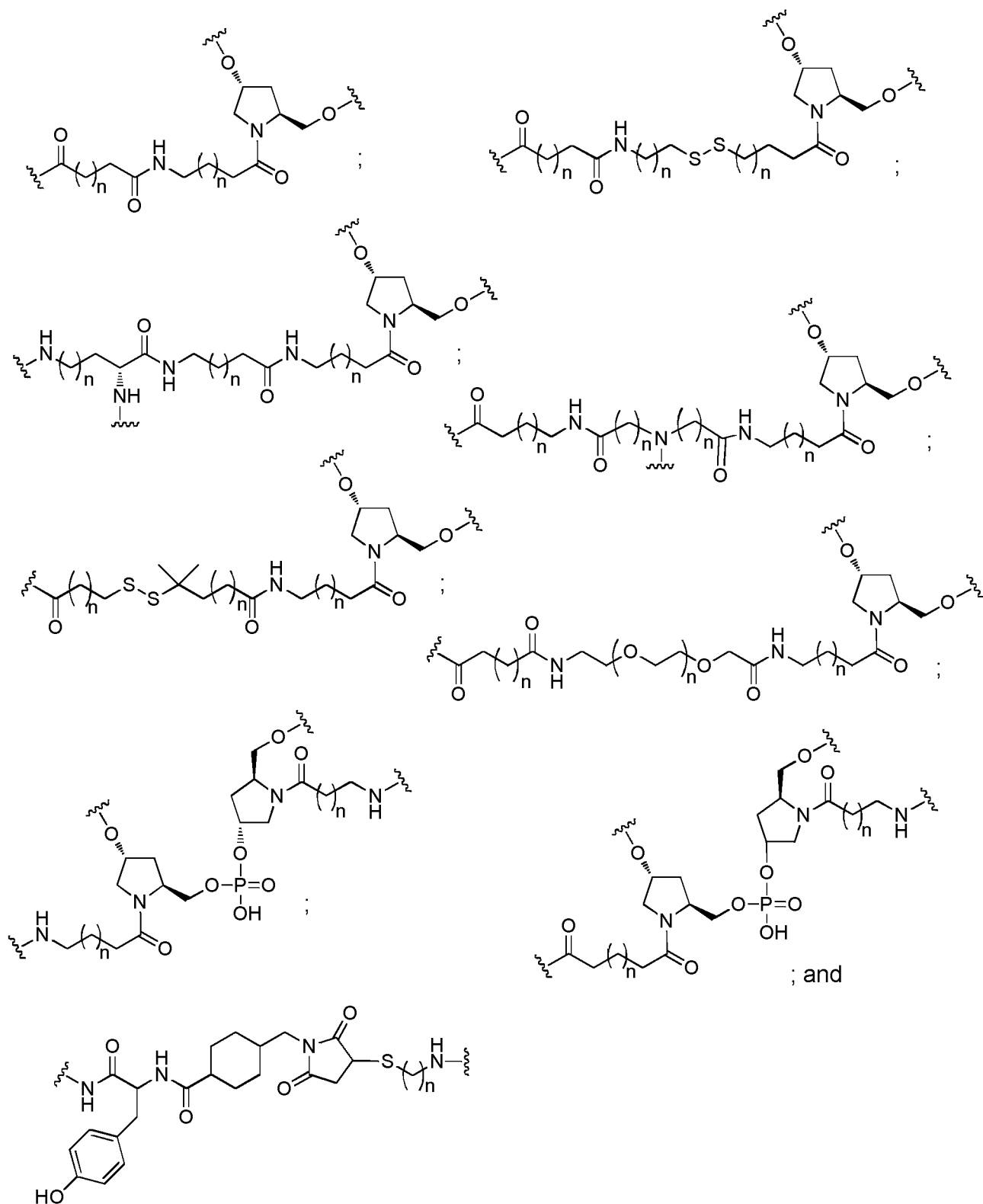
- In certain embodiments, the protein binding moiety is selected from among: cholesterol, cholic acid, adamantane acetic acid, 1-pyrene butyric acid, dihydrotestosterone, 1,3-Bis-O(hexadecyl)glycerol, geranyloxyhexyl group, hexadecylglycerol, borneol, menthol, 1,3-propanediol, heptadecyl group, palmitic acid, myristic acid, O3-(oleoyl)lithocholic acid, O3-(oleoyl)cholenic acid, dimethoxytrityl, or phenoxazine), a
- 5 vitamin (e.g., folate, vitamin A, vitamin E, biotin, pyridoxal), a peptide, a carbohydrate (e.g., monosaccharide, disaccharide, trisaccharide, tetrasaccharide, oligosaccharide, polysaccharide), an endosomolytic component, a steroid (e.g., uvaol, hecigenin, diosgenin), a terpene (e.g., triterpene, e.g., sarsasapogenin, friedelin, epifriedelanol derivatized lithocholic acid), or a cationic lipid.
- 10 In certain embodiments, the protein binding moiety is selected from among: a C16 to C22 long chain saturated or unsaturated fatty acid, cholesterol, cholic acid, vitamin E, adamantane or 1-pentafluoropropyl.

In certain embodiments, the conjugate linker has a structure selected from among:



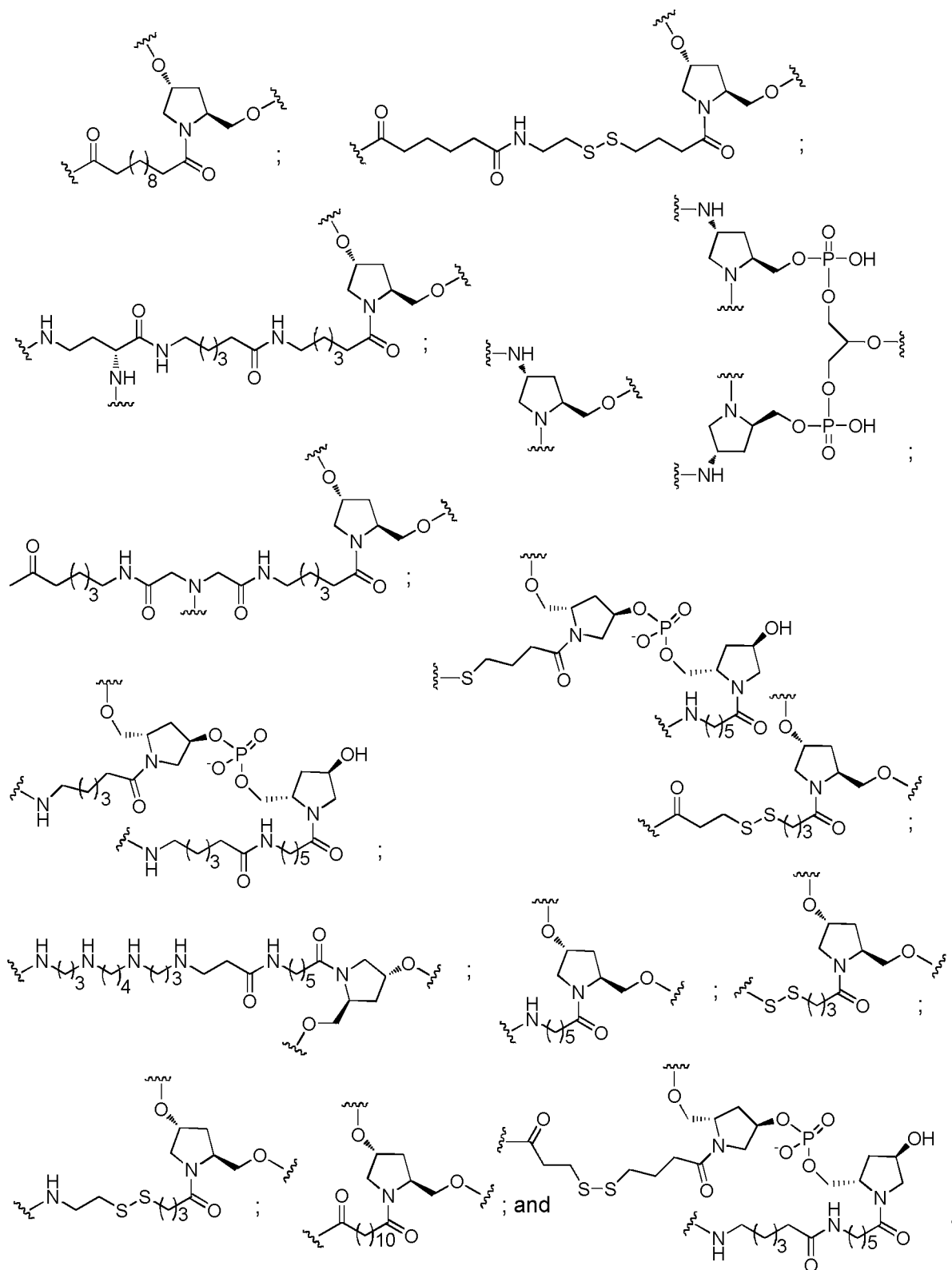
wherein each n is, independently, is from 1 to 20; and p is from 1 to 6.

In certain embodiments, the conjugate linker has a structure selected from among:

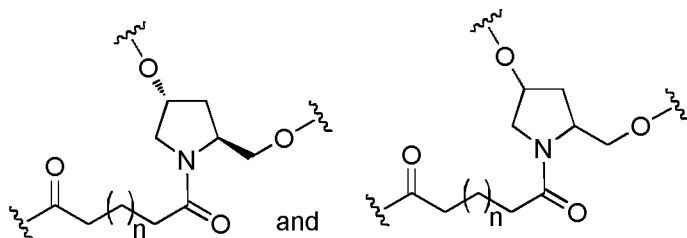


5 wherein each n is, independently, from 1 to 20.

In certain embodiments, the conjugate linker has a structure selected from among:



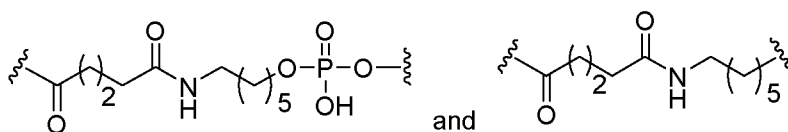
In certain embodiments, the conjugate linker has a structure selected from among:



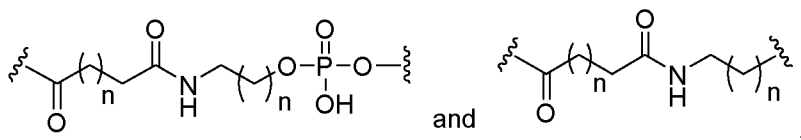
wherein n is from 1 to 20.

5

In certain embodiments, the conjugate linker has a structure selected from among:



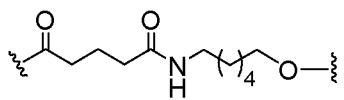
In certain embodiments, the conjugate linker has a structure selected from among:



10

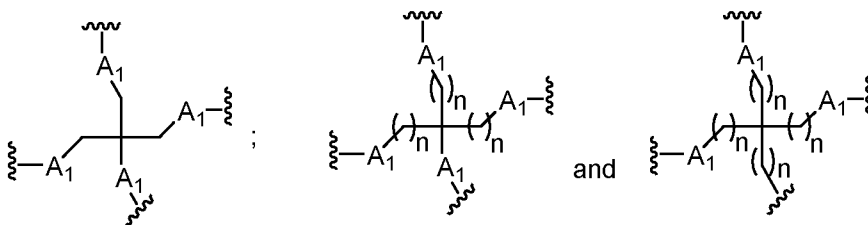
wherein each n is independently, 0, 1, 2, 3, 4, 5, 6, or 7.

In certain embodiments, the conjugate linker has the following structure:



15

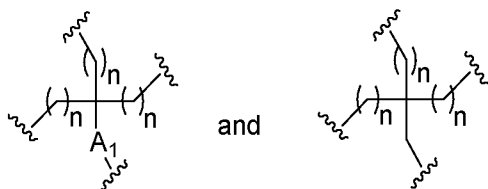
In certain embodiments, the branching group has one of the following structures:



20

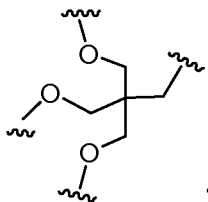
wherein each A₁ is independently, O, S, C=O or NH; and
each n is, independently, from 1 to 20.

In certain embodiments, the branching group has one of the following structures:

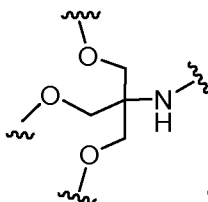


wherein each A1 is independently, O, S, C=O or NH; and
each n is, independently, from 1 to 20.

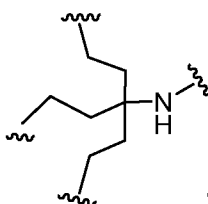
In certain embodiments, the branching group has the following structure:



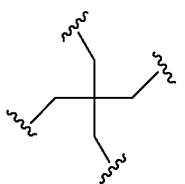
In certain embodiments, the branching group has the following structure:



In certain embodiments, the branching group has the following structure:

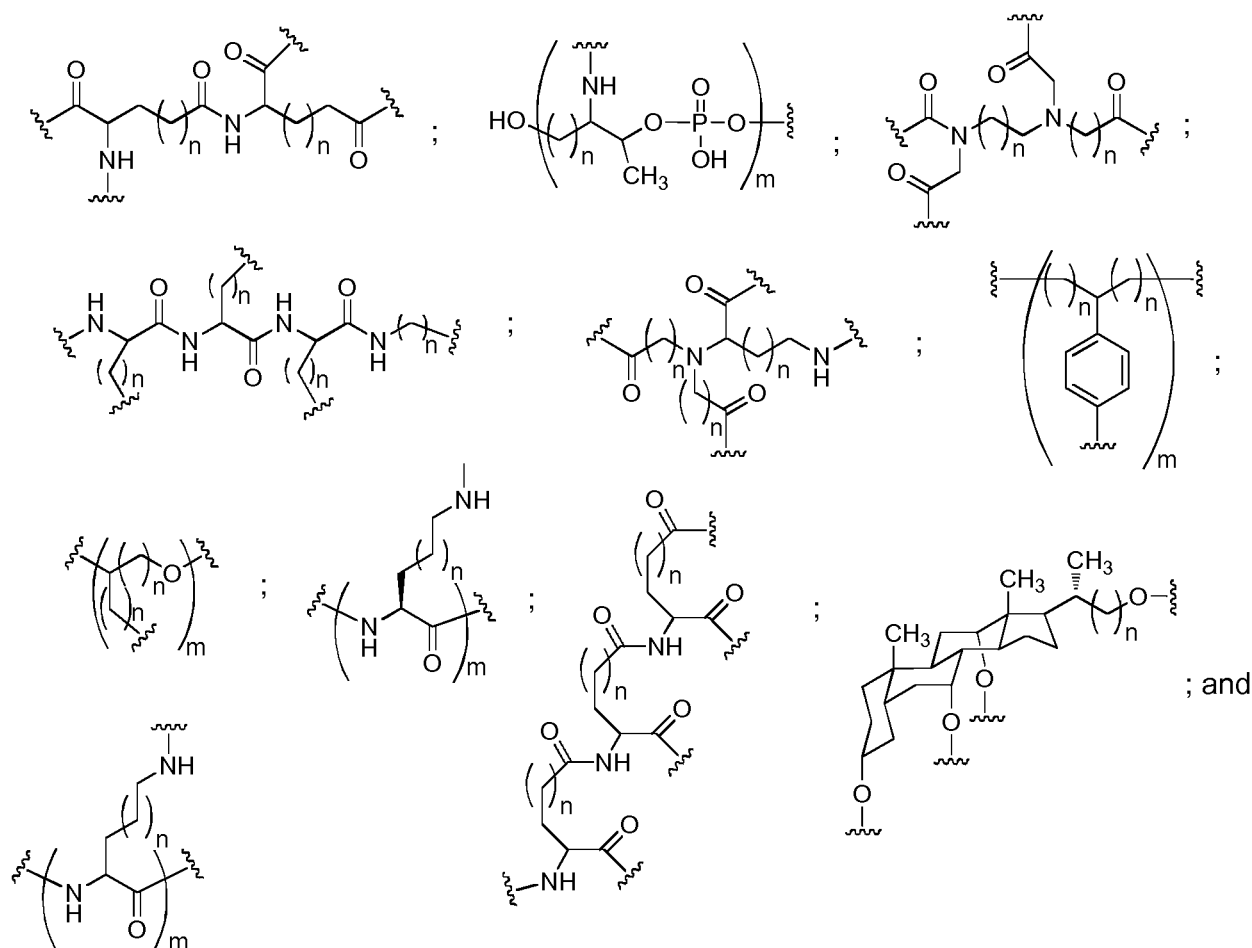


In certain embodiments, the branching group has the following structure:



In certain embodiments, the branching group comprises an ether.

In certain embodiments, the branching group has the following structure:

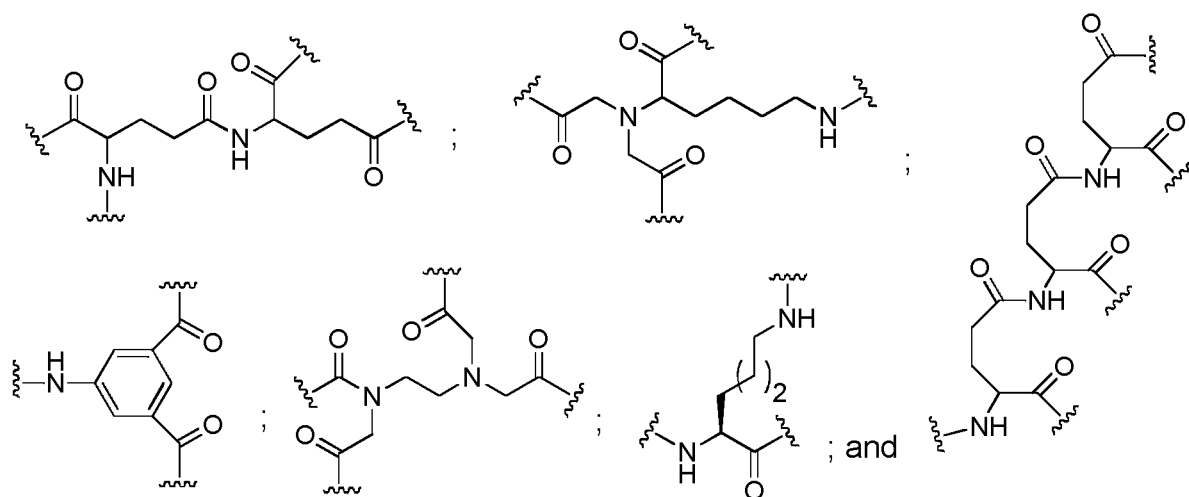


5

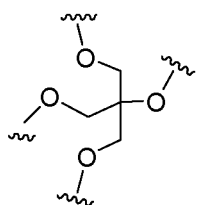
each n is, independently, from 1 to 20; and
 m is from 2 to 6.

10

In certain embodiments, the branching group has the following structure:

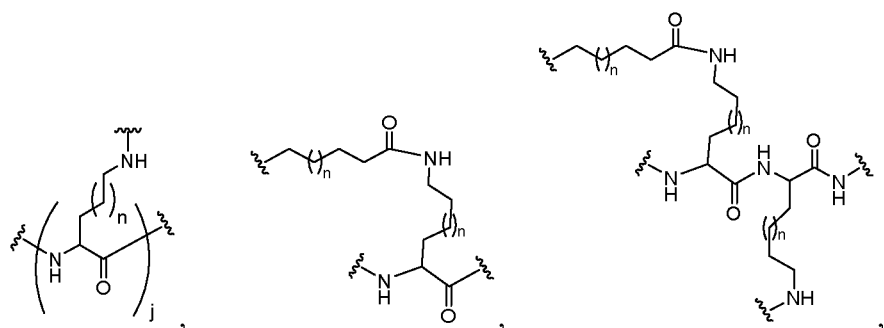


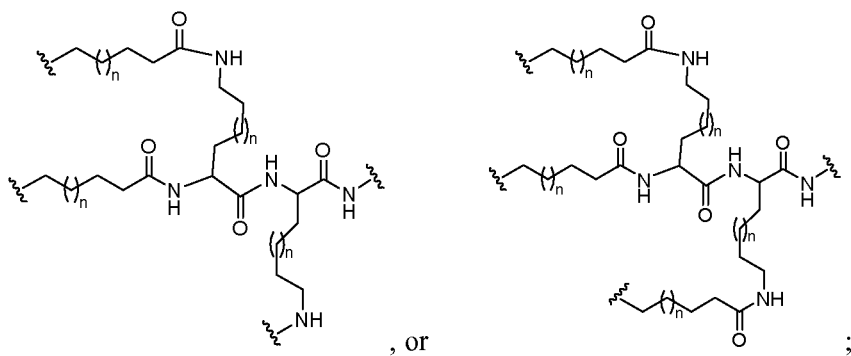
5 In certain embodiments, the branching group has the following structure:



In certain embodiments, the branching group comprises:

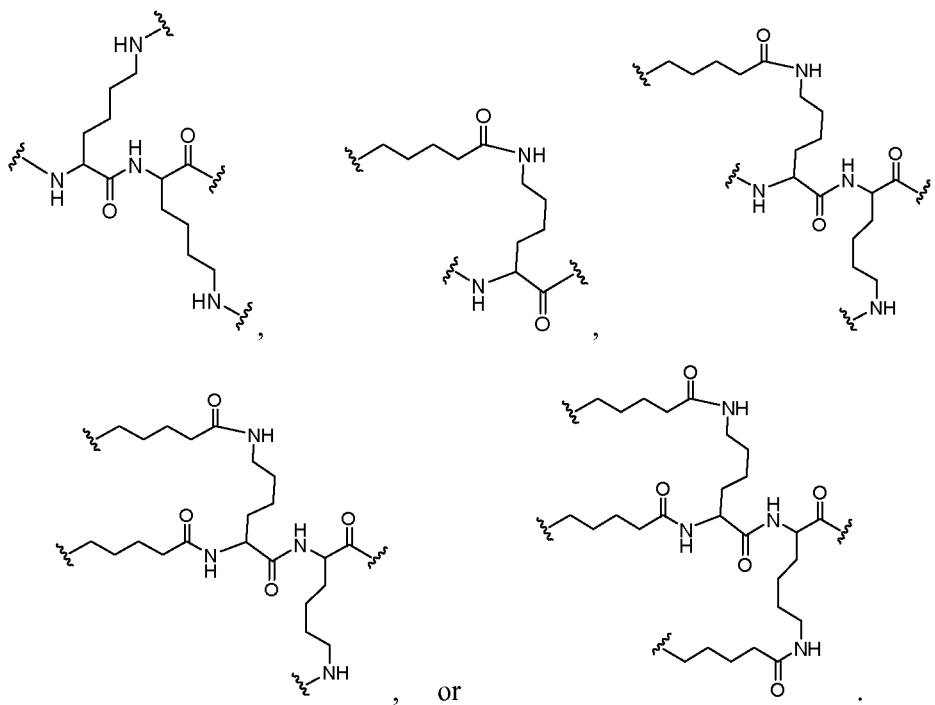
10





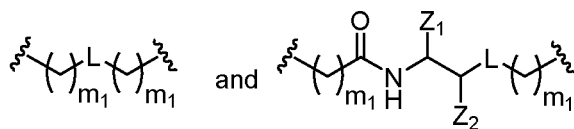
wherein each j is an integer from 1 to 3; and
 wherein each n is an integer from 1 to 20.

5 In certain embodiments, the branching group comprises:



10

In certain embodiments, each tether is selected from among:



15 wherein L is selected from a phosphorus linking group and a neutral linking group;

Z1 is C(=O)O-R2;

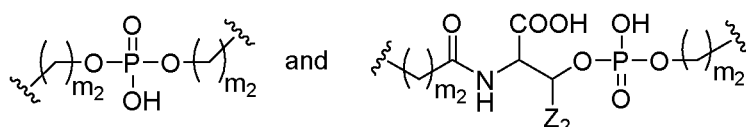
Z2 is H, C1-C6 alkyl or substituted C1-C6 alky;

R2 is H, C1-C6 alkyl or substituted C1-C6 alky; and

each m_i is, independently, from 0 to 20 wherein at least one m_i is greater than 0 for each tether.

5

In certain embodiments, each tether is selected from among:

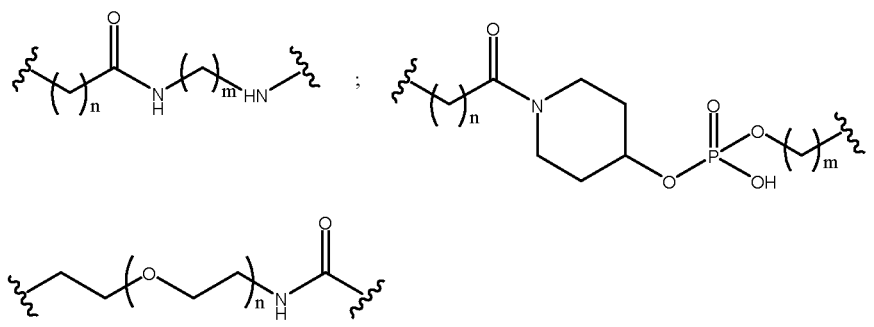


wherein Z2 is H or CH3; and

each m_2 is, independently, from 0 to 20 wherein at least one m_2 is greater than 0 for each tether.

10

In certain embodiments, each tether is selected from among:

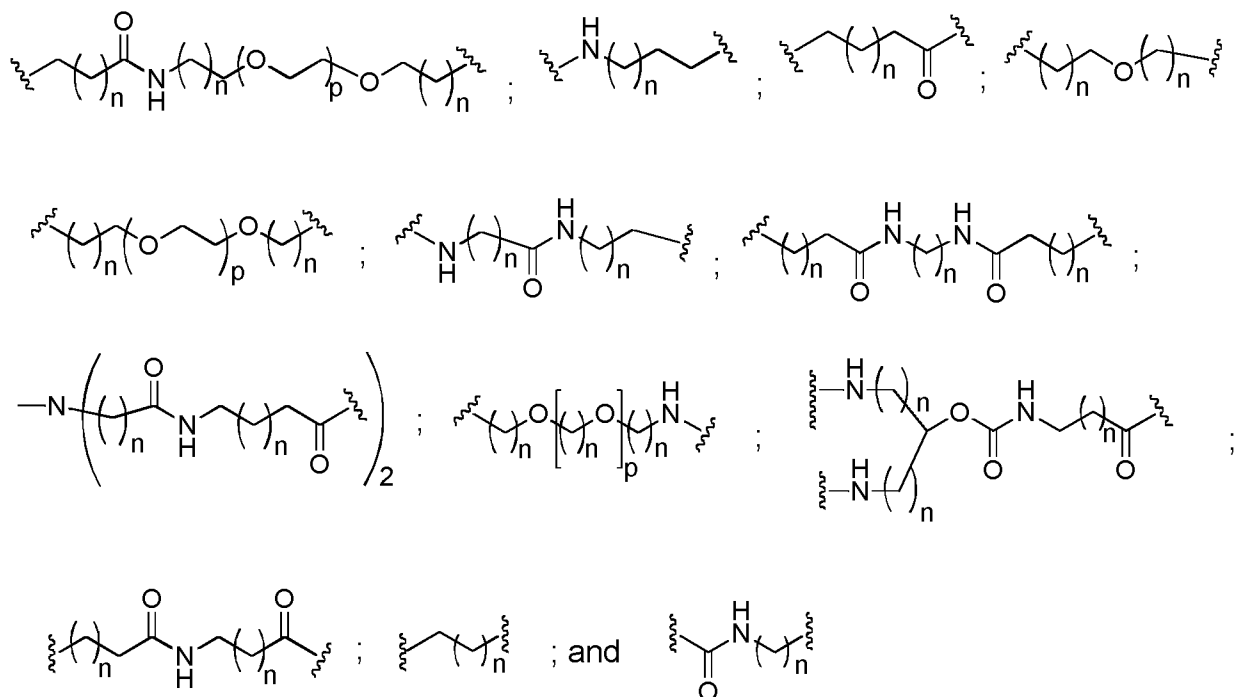


wherein n is from 1 to 12; and

wherein m is from 1 to 12.

15

In certain embodiments, at least one tether comprises ethylene glycol. In certain embodiments, at least one tether comprises an amide. In certain embodiments, at least one tether comprises a polyamide. In certain embodiments, at least one tether comprises an amine. In certain embodiments, at least two tethers are different from one another. In certain embodiments, all of the tethers are the same as one another. In certain embodiments, each tether is selected from among:

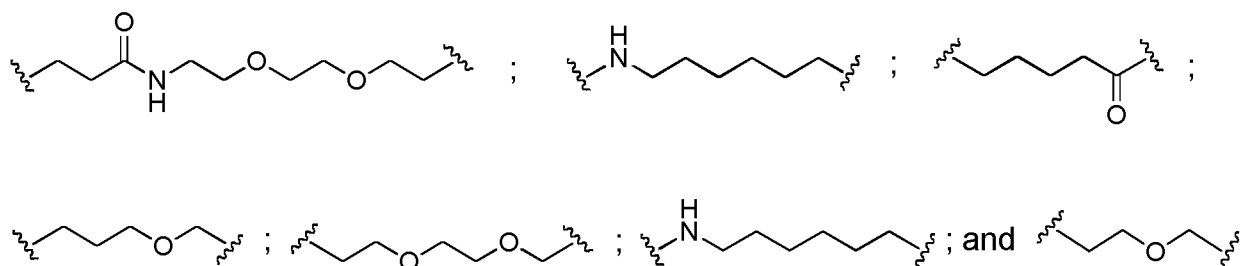


wherein each n is, independently, from 1 to 20; and

each p is from 1 to about 6.

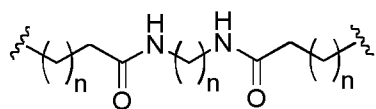
5

In certain embodiments, each tether is selected from among:



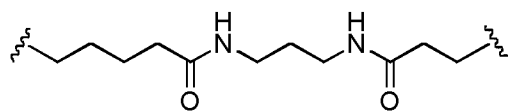
10

In certain embodiments, each tether has the following structure:



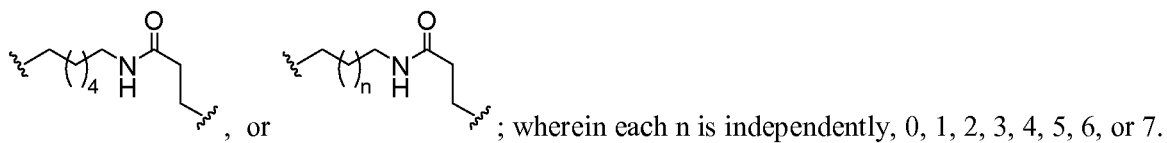
wherein each n is, independently, from 1 to 20.

In certain embodiments, each tether has the following structure:

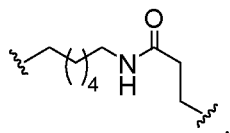


15

- 5 In certain embodiments, the tether has a structure selected from among:



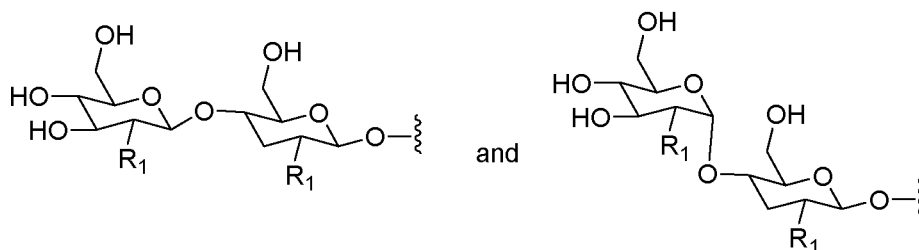
In certain embodiments, the tether has a structure selected from among:



10

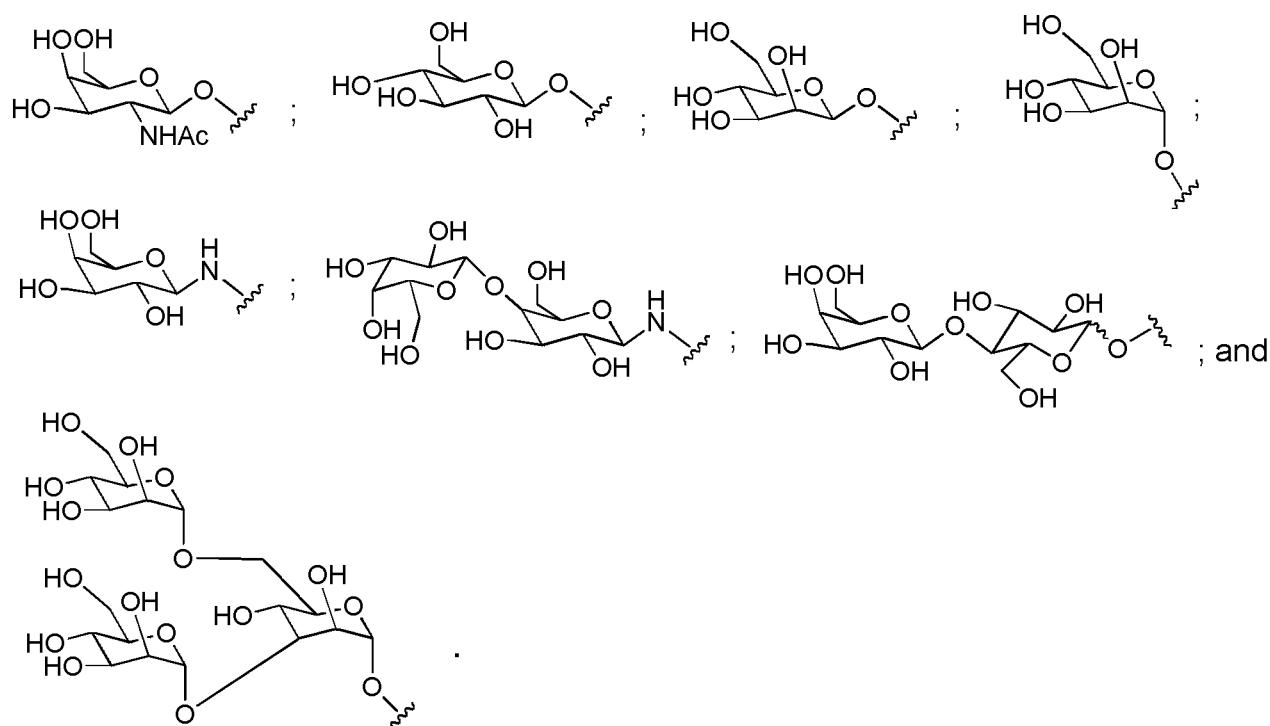
In certain embodiments, the ligand is galactose. In certain embodiments, the ligand is mannose-6-phosphate.

- 15 In certain embodiments, each ligand is selected from among:

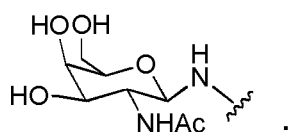


wherein each R1 is selected from OH and NHCOOH.

In certain embodiments, each ligand is selected from among:

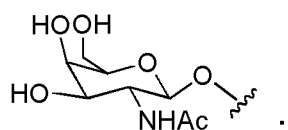


In certain embodiments, each ligand has the following structure:



5

In certain embodiments, each ligand has the following structure:

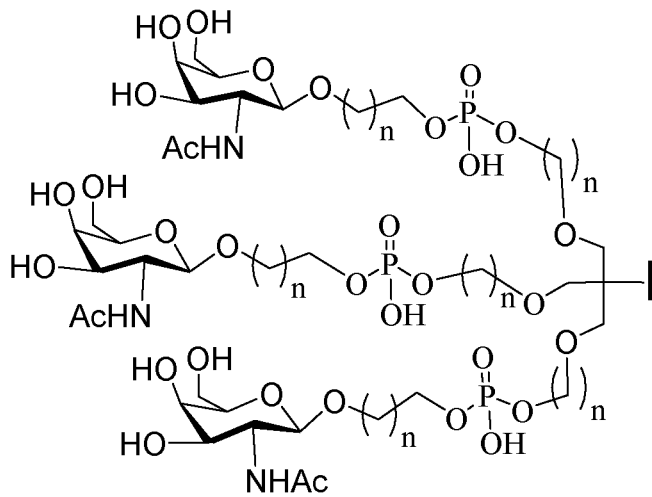


10

In certain embodiments, the conjugate group comprises a cell-targeting moiety.

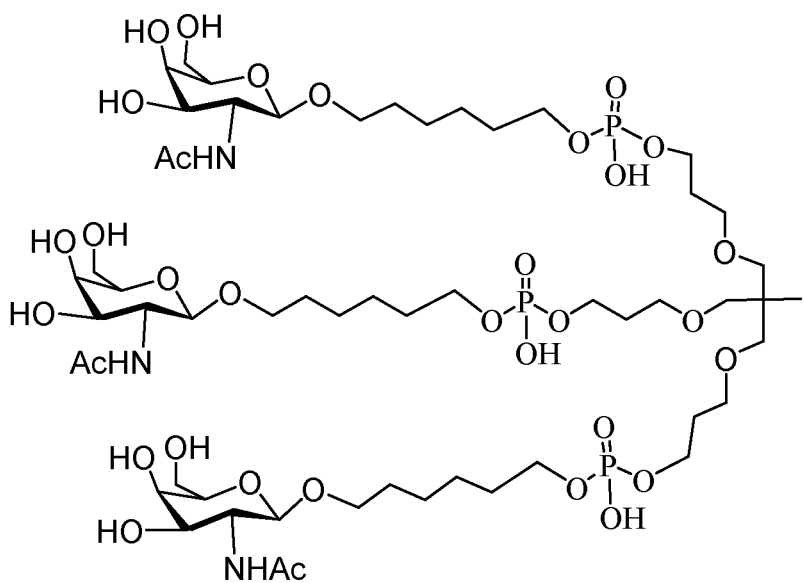
In certain embodiments, the conjugate group comprises a cell-targeting moiety having the following structure:

15



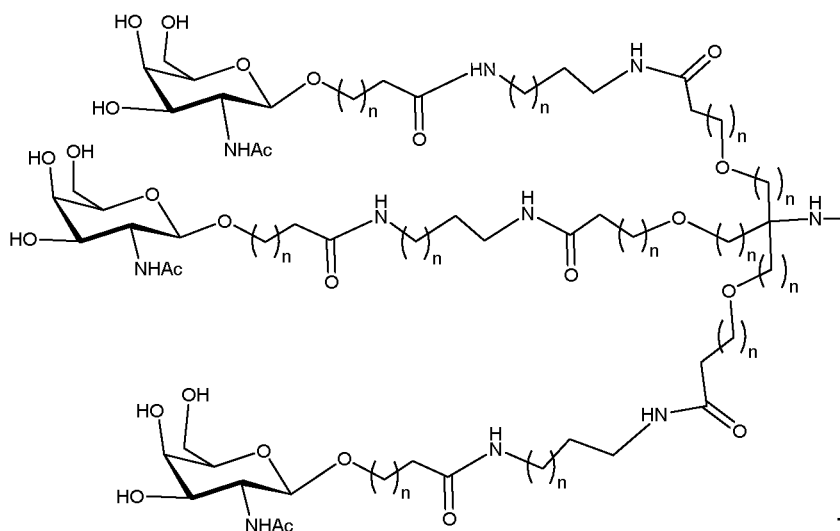
wherein each n is, independently, from 1 to 20.

5 In certain embodiments, the cell-targeting moiety has the following structure:



In certain embodiments, the cell-targeting moiety has the following structure:

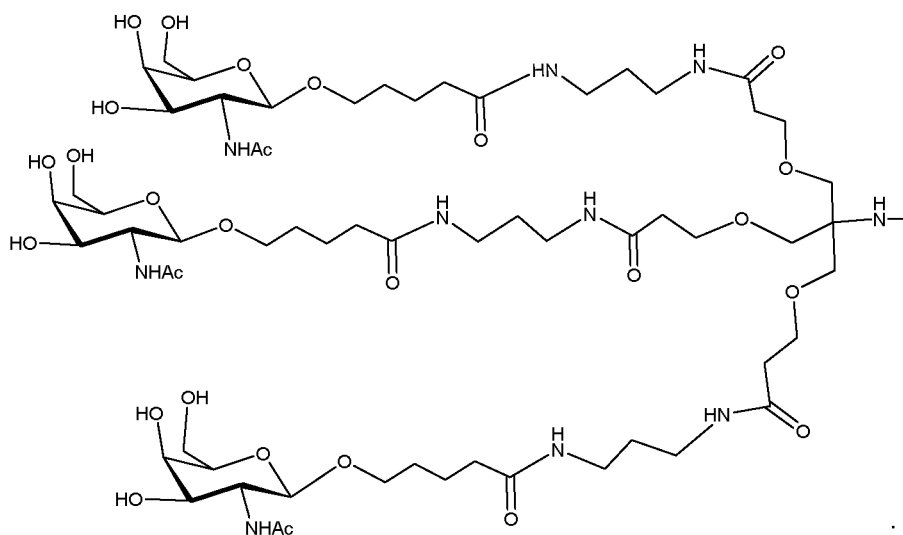
10



wherein each n is, independently, from 1 to 20.

5

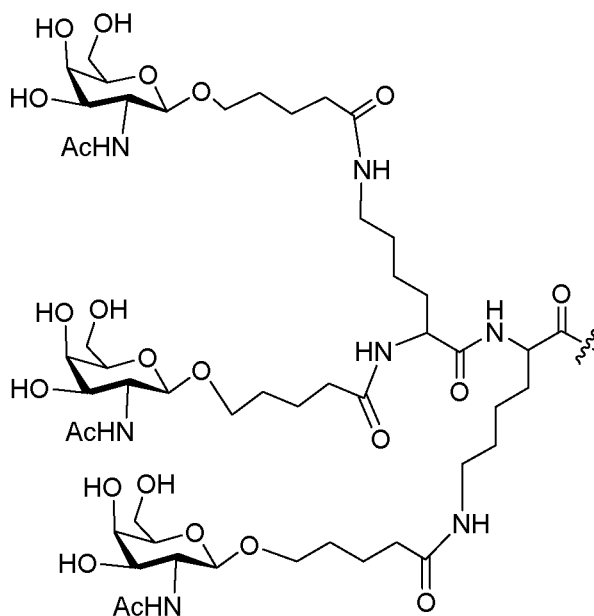
In certain embodiments, the cell-targeting moiety has the following structure:



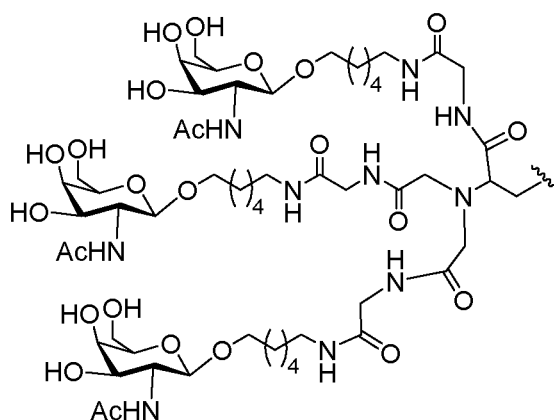
10

15

In certain embodiments, the cell-targeting moiety comprises:

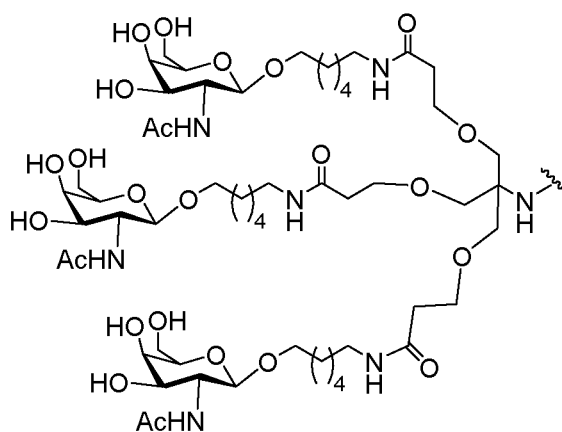


In certain embodiments, the cell-targeting moiety comprises:

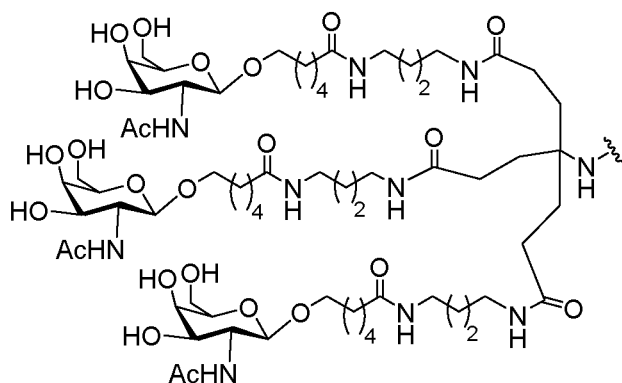


5

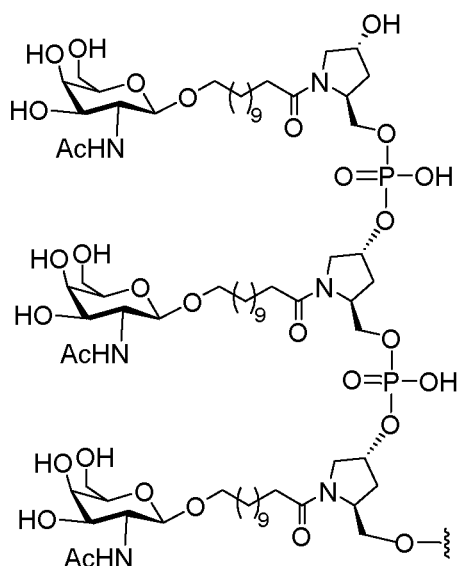
In certain embodiments, the cell-targeting moiety comprises:



In certain embodiments, the cell-targeting moiety comprises:

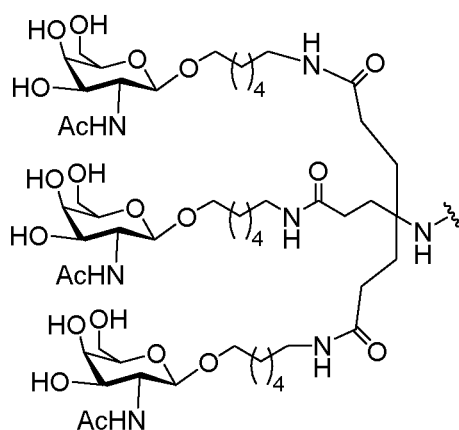


In certain embodiments, the cell-targeting moiety comprises:

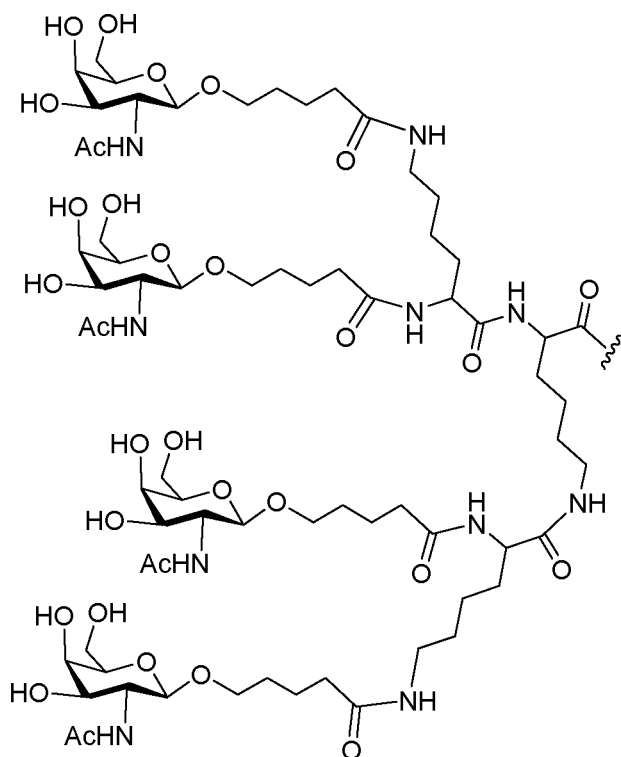


5

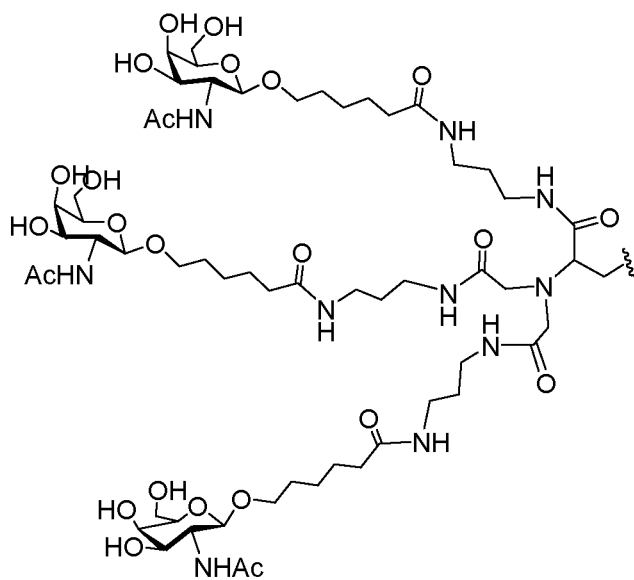
In certain embodiments, the cell-targeting moiety comprises:



In certain embodiments, the cell-targeting moiety comprises:

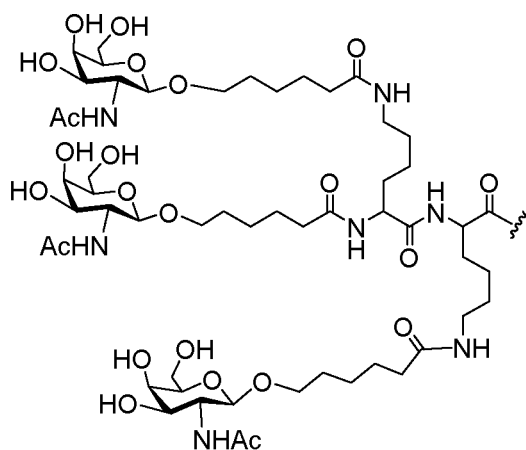


In certain embodiments, the cell-targeting moiety comprises:

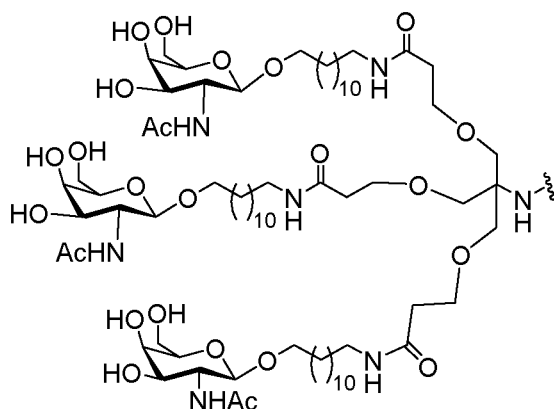


5

In certain embodiments, the cell-targeting moiety comprises:

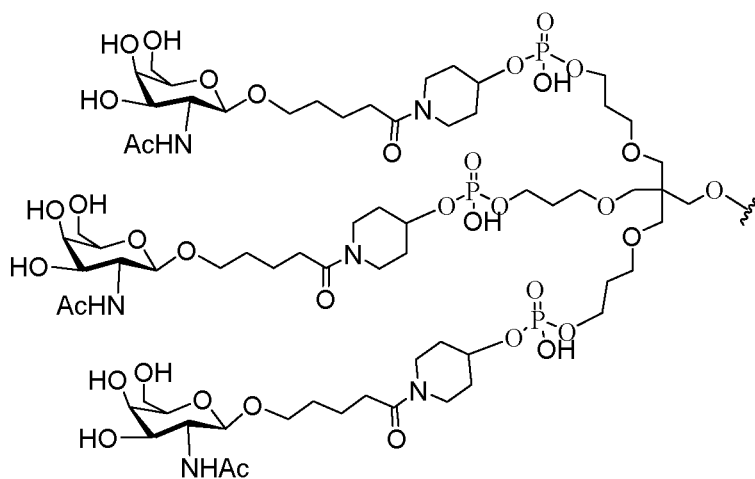


In certain embodiments, the cell-targeting moiety comprises:

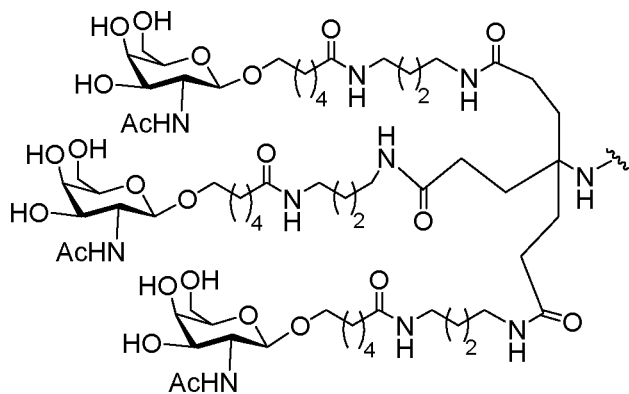


5

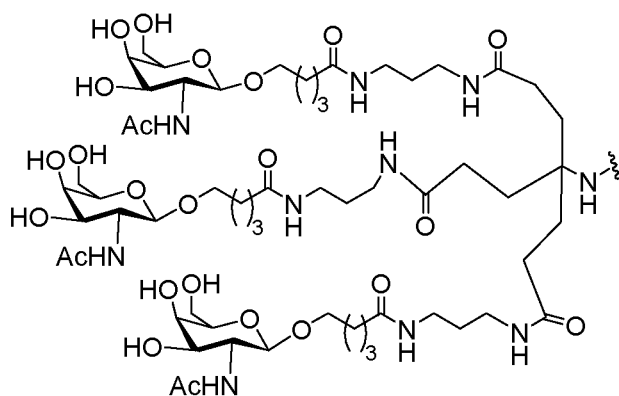
In certain embodiments, the cell-targeting moiety comprises:



In certain embodiments, the cell-targeting moiety comprises:

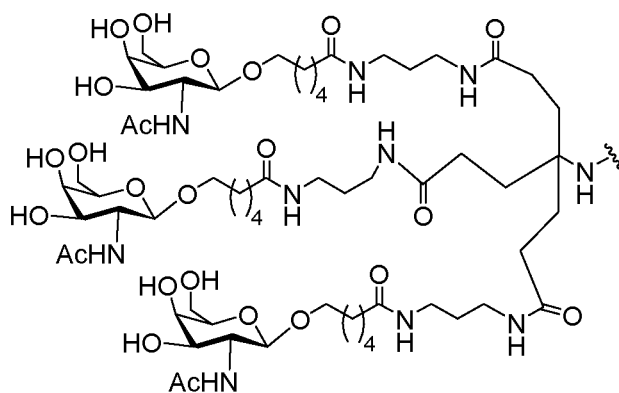


In certain embodiments, the cell-targeting moiety comprises:

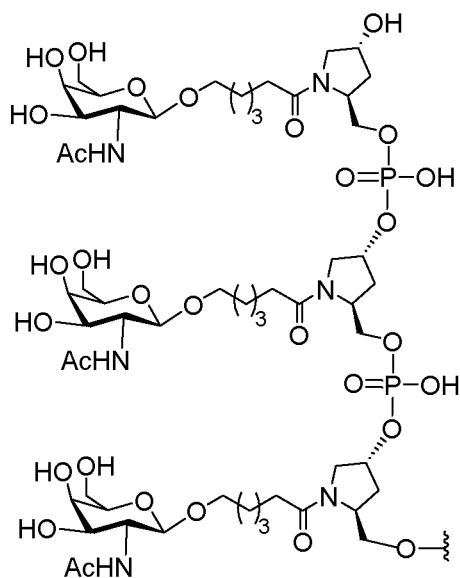


5

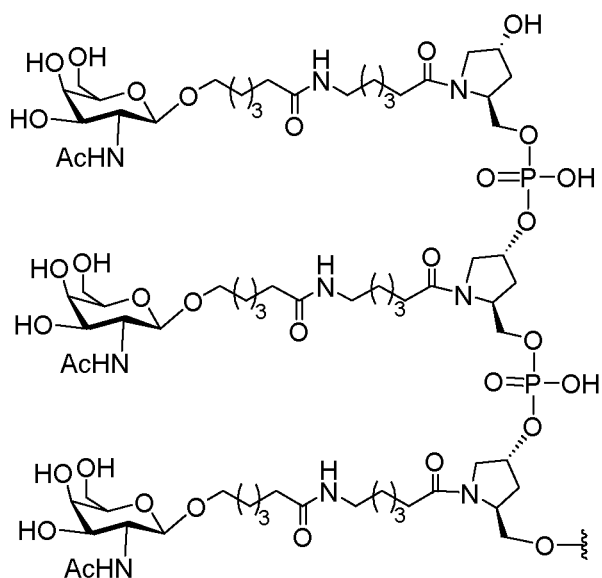
In certain embodiments, the cell-targeting moiety comprises:



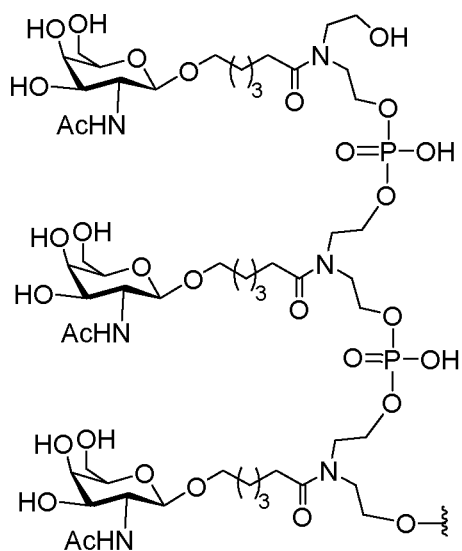
In certain embodiments, the cell-targeting moiety comprises:



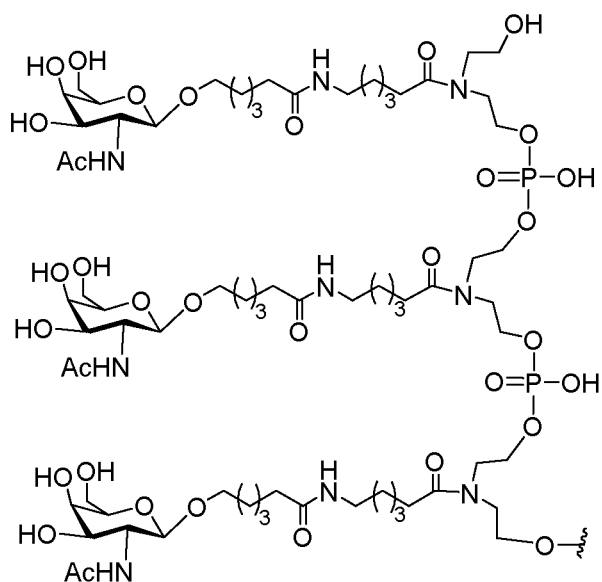
In certain embodiments, the cell-targeting moiety comprises:



In certain embodiments, the cell-targeting moiety comprises:

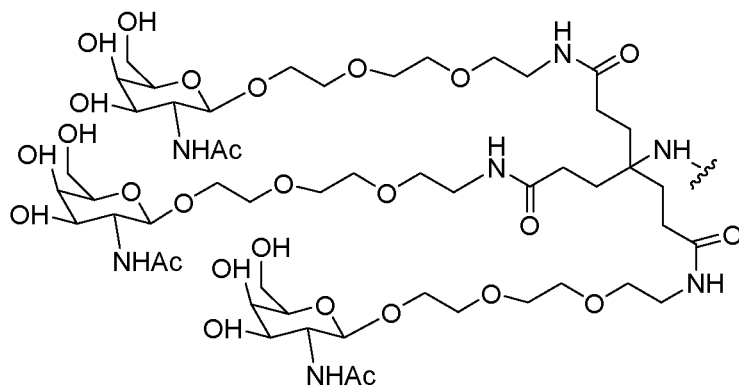


In certain embodiments, the cell-targeting moiety comprises:

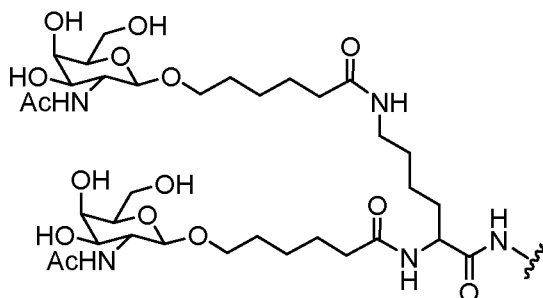


5

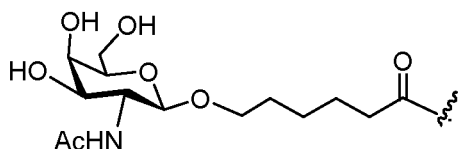
In certain embodiments, the cell-targeting moiety comprises:



In certain embodiments, the cell-targeting moiety comprises:

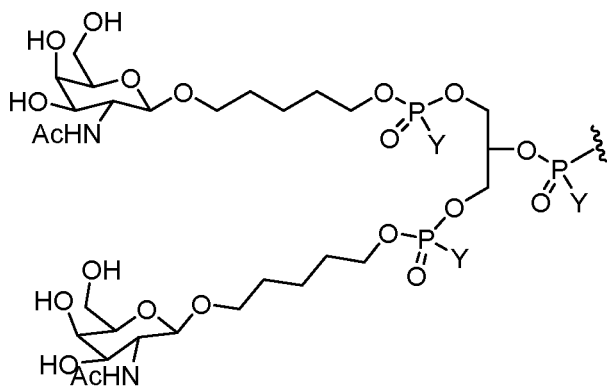


In certain embodiments, the cell-targeting moiety comprises:



5

In certain embodiments, the cell-targeting moiety comprises:

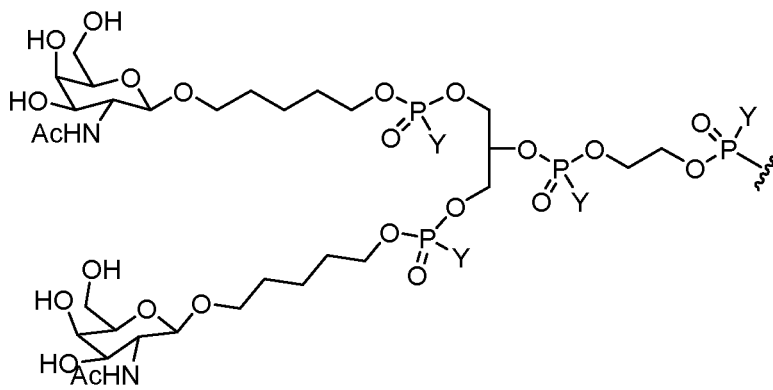


;

wherein each Y is selected from O, S, a substituted or unsubstituted C1-C10 alkyl, amino, substituted amino, azido, alkenyl or alkynyl.

10

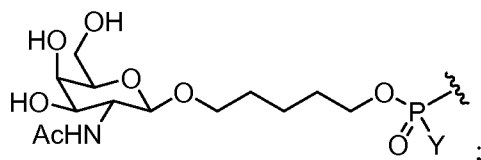
In certain embodiments, the conjugate group comprises:



;

wherein each Y is selected from O, S, a substituted or unsubstituted C1-C10 alkyl, amino, substituted amino, azido, alkenyl or alkynyl.

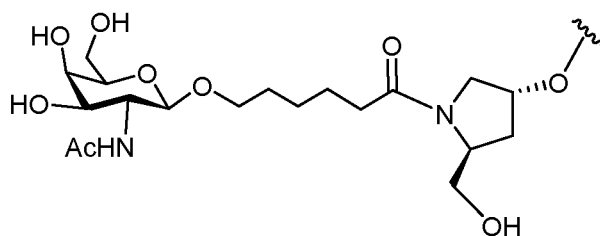
In certain embodiments, the conjugate group comprises:



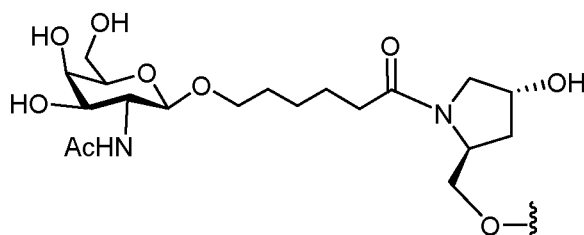
wherein each Y is selected from O, S, a substituted or unsubstituted C1-C10 alkyl, amino, substituted

5 amino, azido, alkenyl or alkynyl.

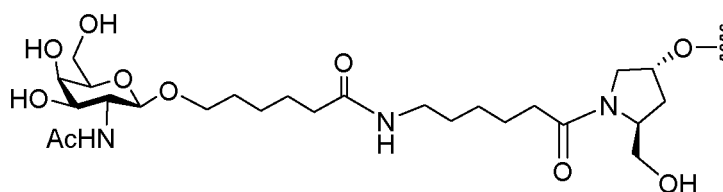
In certain embodiments, the conjugate group comprises:



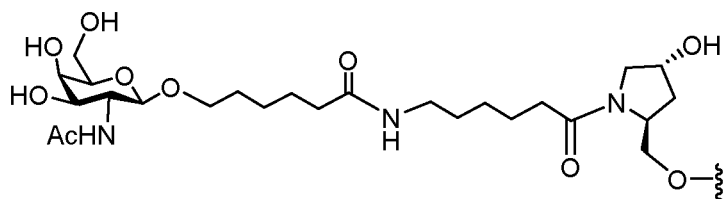
In certain embodiments, the conjugate group comprises:



10 In certain embodiments, the conjugate group comprises:



In certain embodiments, the conjugate group comprises:



15 In certain embodiments, the conjugate group comprises a cleavable moiety selected from among: a phosphodiester, an amide, or an ester.

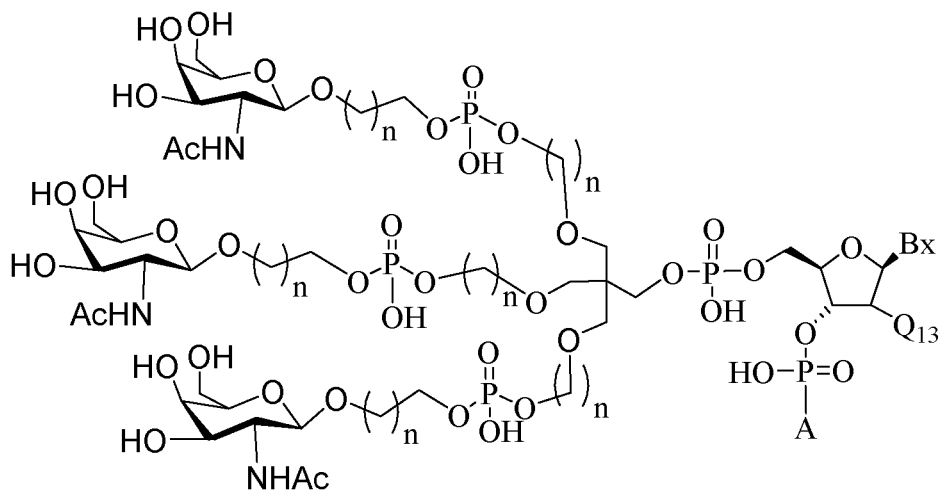
In certain embodiments, the conjugate group comprises a phosphodiester cleavable moiety.

In certain embodiments, the conjugate group does not comprise a cleavable moiety, and wherein the conjugate group comprises a phosphorothioate linkage between the conjugate group and the oligonucleotide.

In certain embodiments, the conjugate group comprises an amide cleavable moiety. In certain embodiments,

5 the conjugate group comprises an ester cleavable moiety.

In certain embodiments, the compound has the following structure:



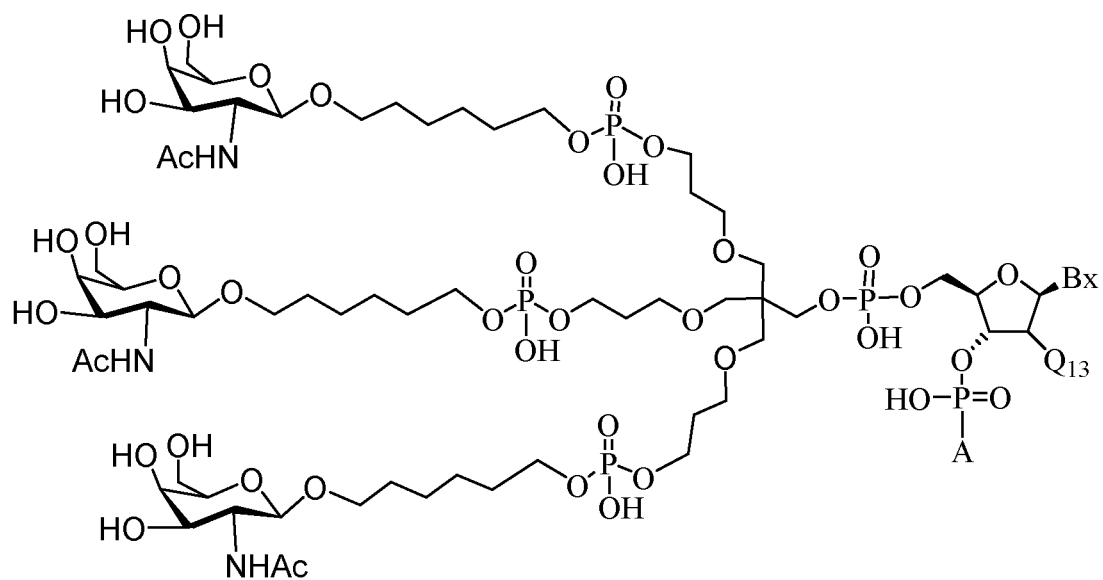
10 wherein each n is, independently, from 1 to 20;

Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

15 In certain embodiments, the compound has the following structure:



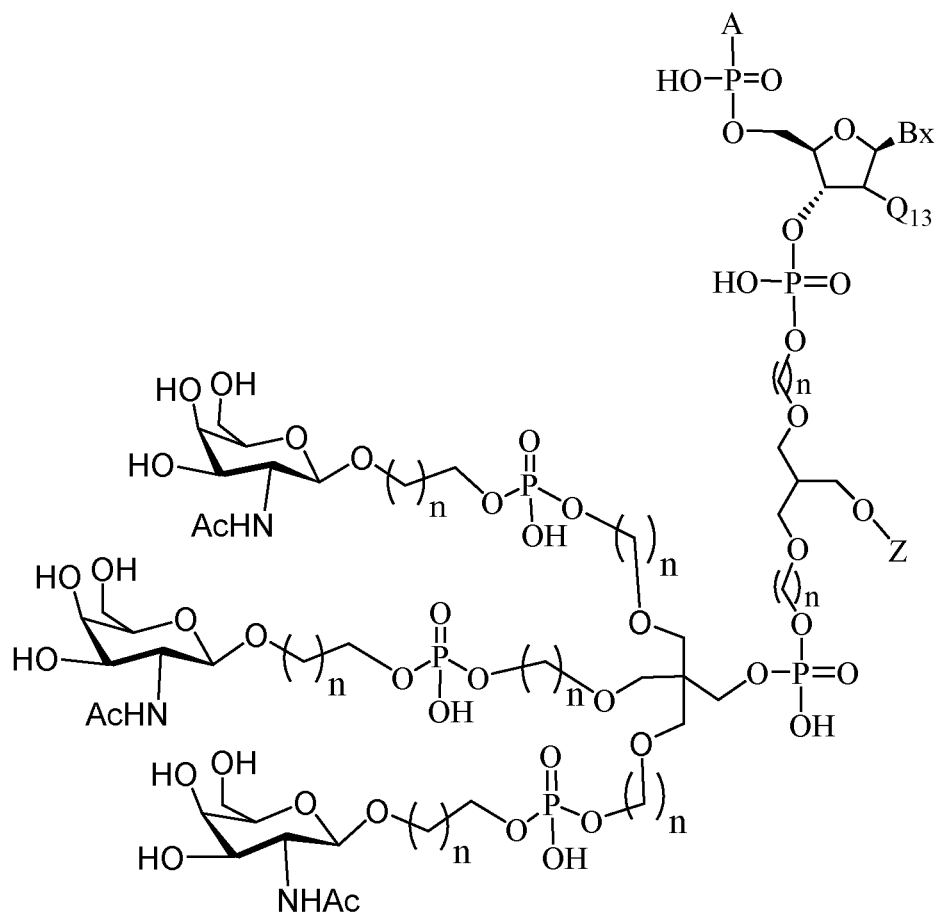
wherein each n is, independently, from 1 to 20;

Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

In certain embodiments, the compound has the following structure:



wherein each n is, independently, from 1 to 20;

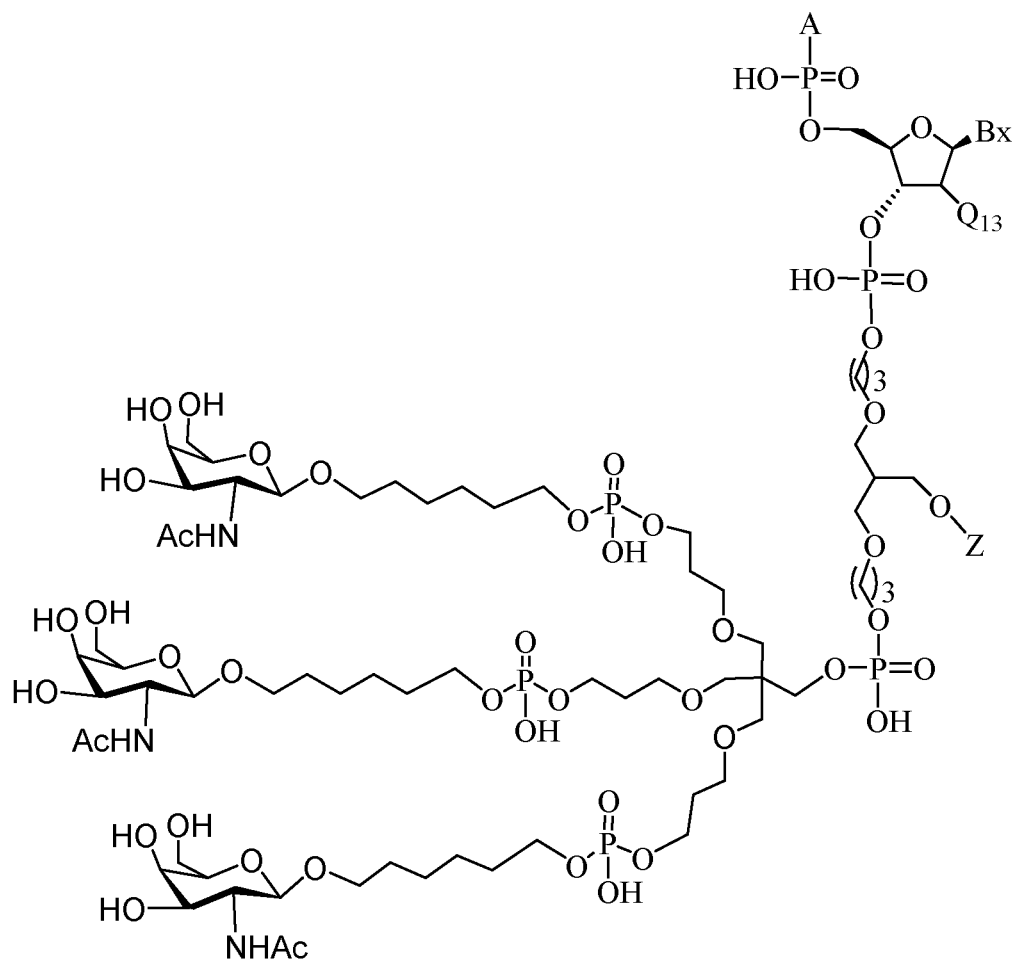
Q_{13} is H or $O(CH_2)_2-OCH_3$;

A is the modified oligonucleotide;

Z is H or a linked solid support; and

Bx is a heterocyclic base moiety.

In certain embodiments, the compound has the following structure:

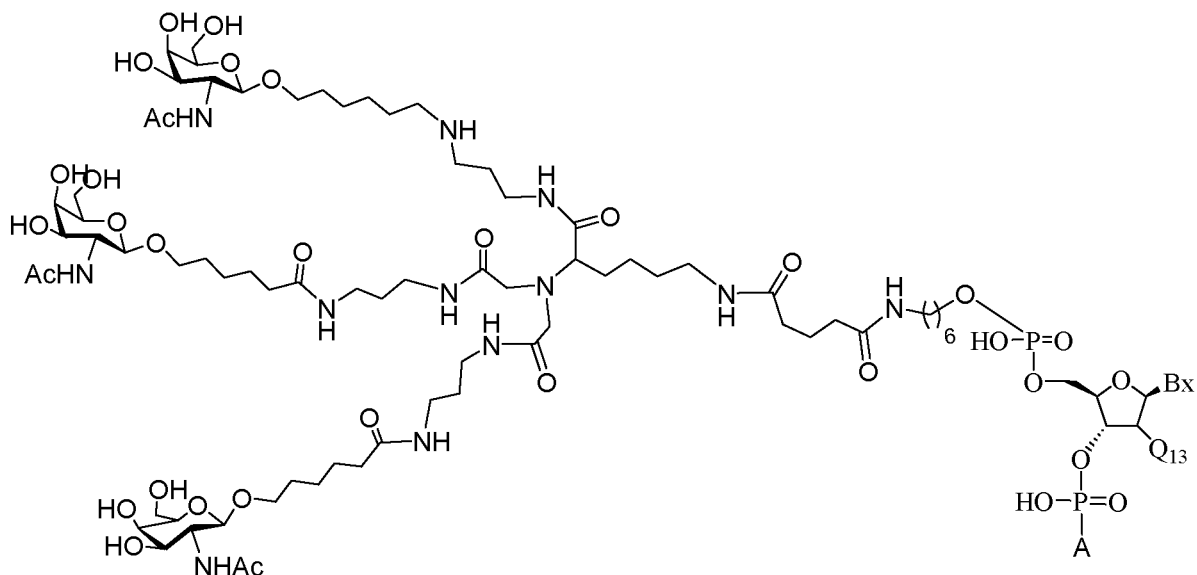


5

A is the modified oligonucleotide;

Bx is a heterocyclic base moiety.

In certain embodiments, the compound has the following structure:



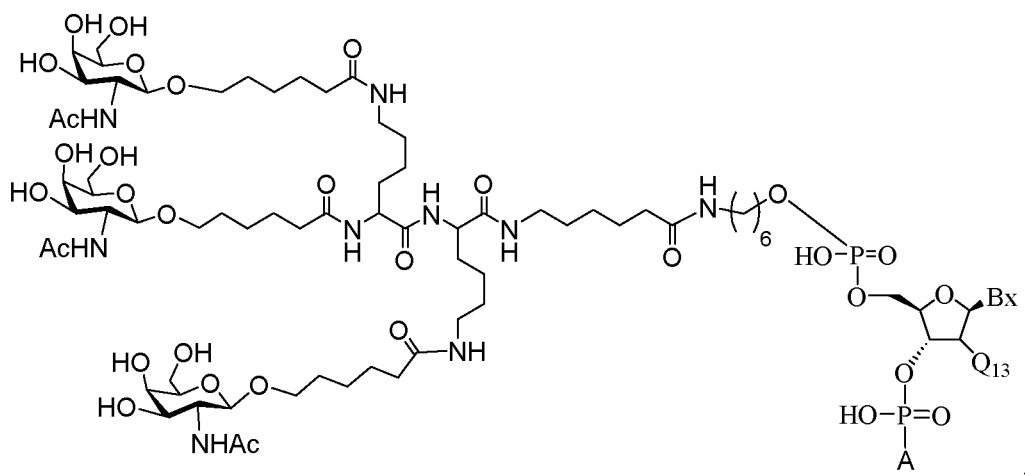
wherein Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

5

In certain embodiments, the compound has the following structure:



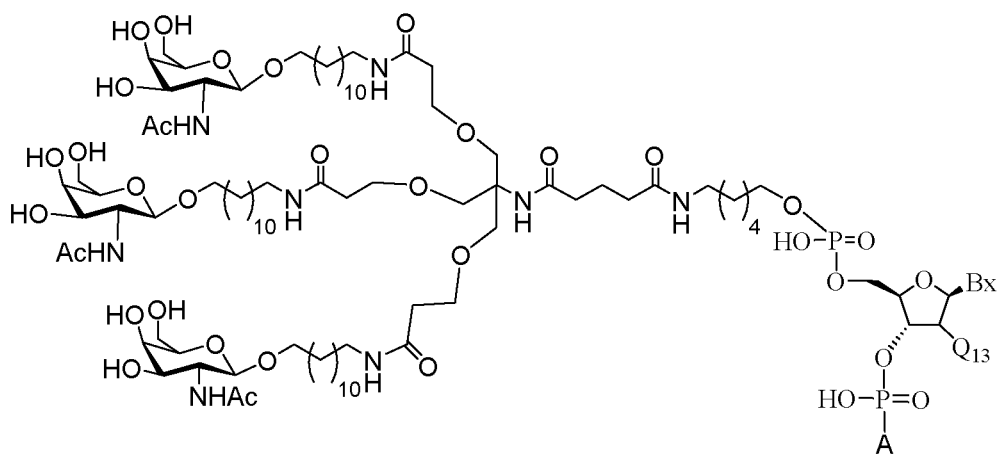
wherein Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

10

In certain embodiments, the compound has the following structure:

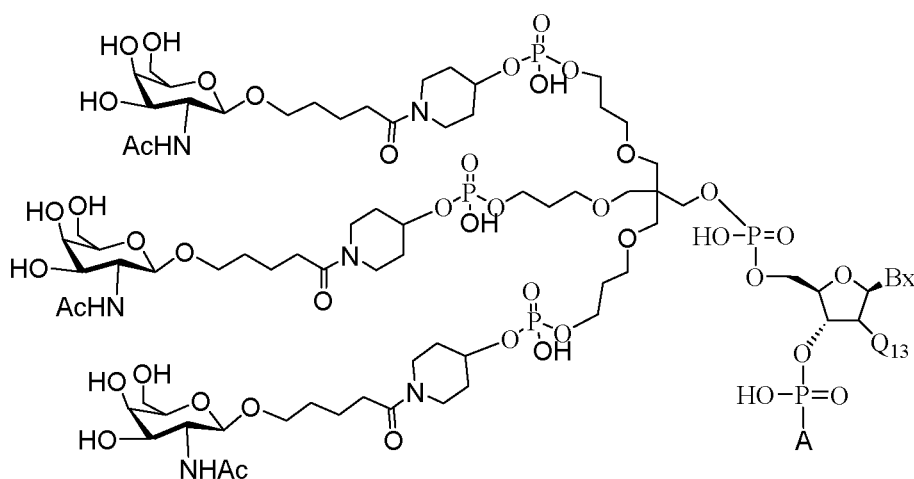


wherein Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

In certain embodiments, the compound has the following structure:

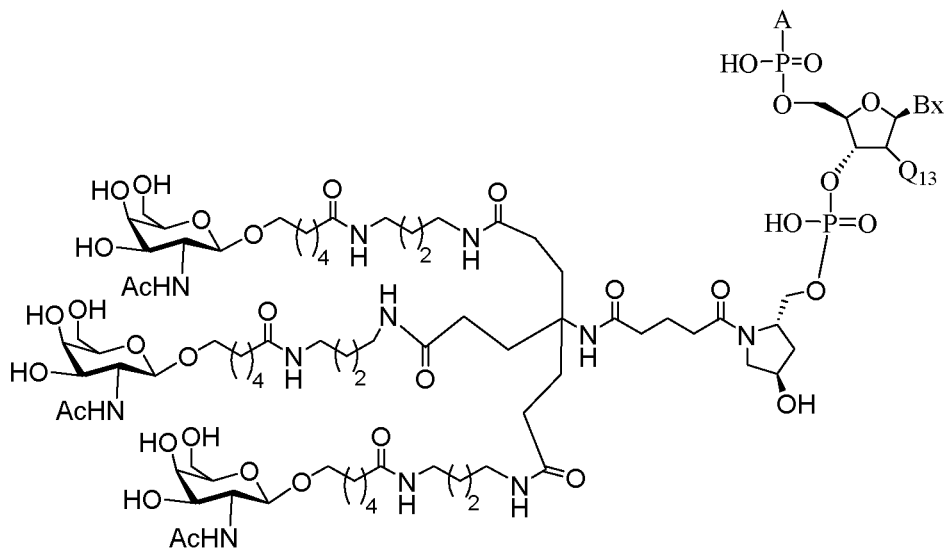


wherein Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

In certain embodiments, the compound has the following structure:



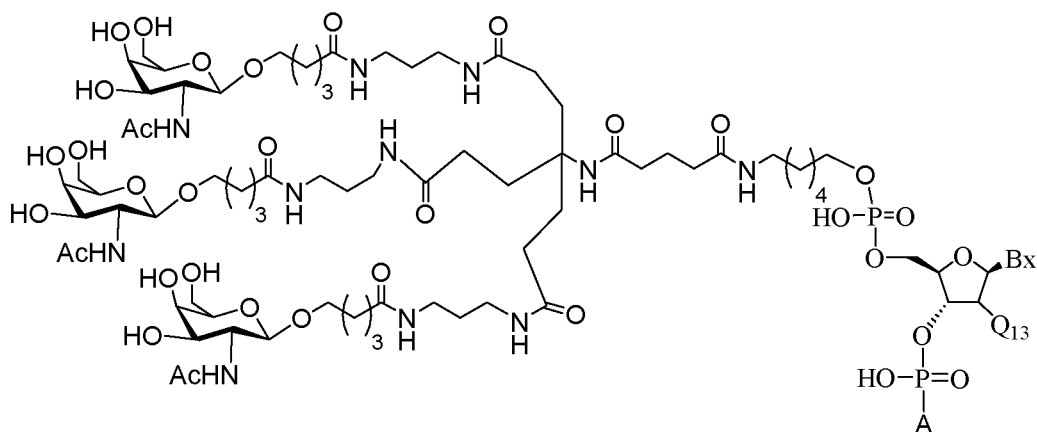
wherein Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

5

In certain embodiments, the compound has the following structure:



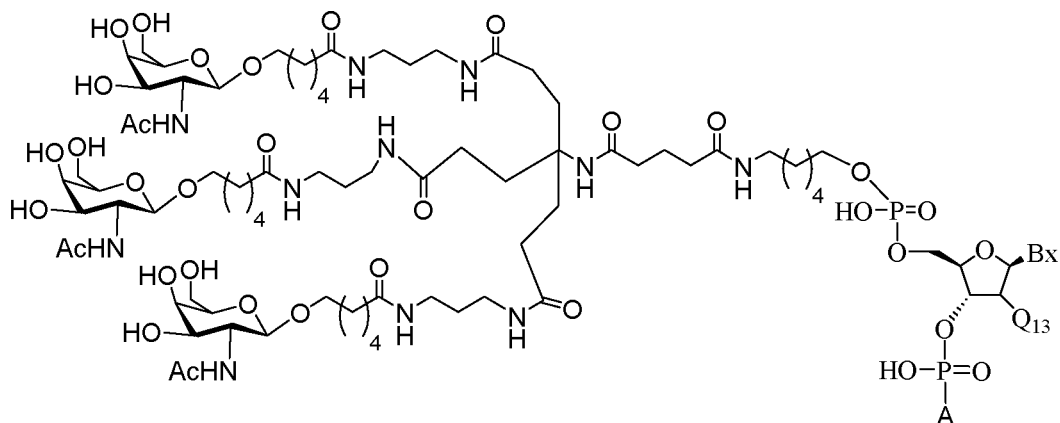
wherein Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

In certain embodiments, the compound has the following structure:

10



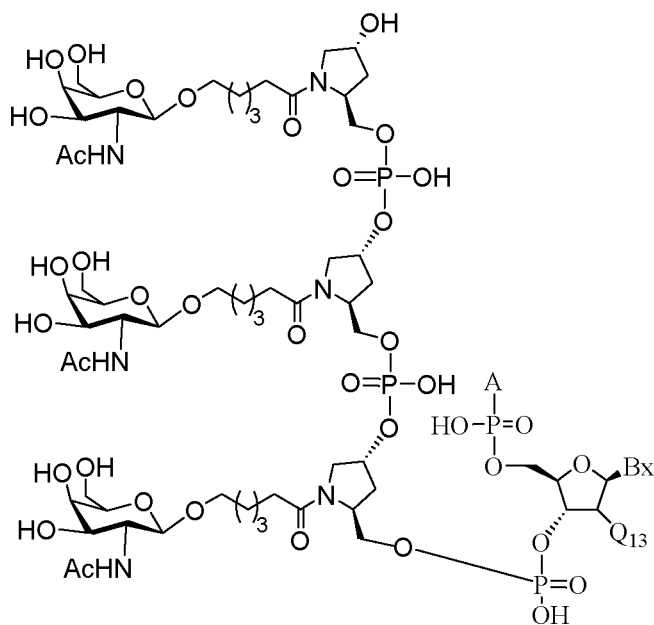
wherein Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

5

In certain embodiments, the compound has the following structure:



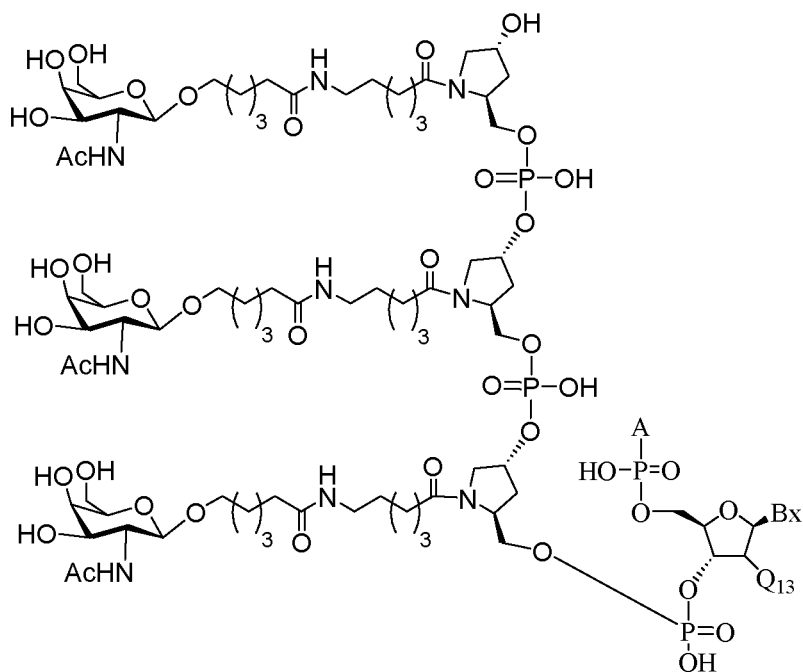
wherein Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

10

In certain embodiments, the compound has the following structure:



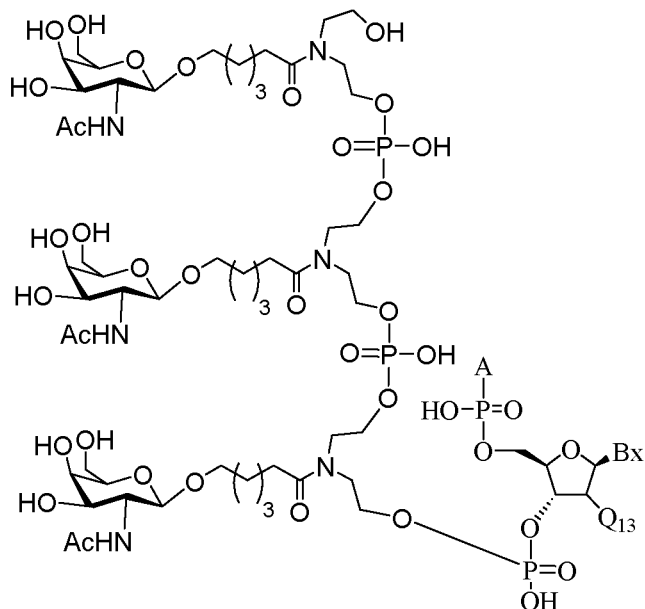
wherein Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

5

In certain embodiments, the compound has the following structure:



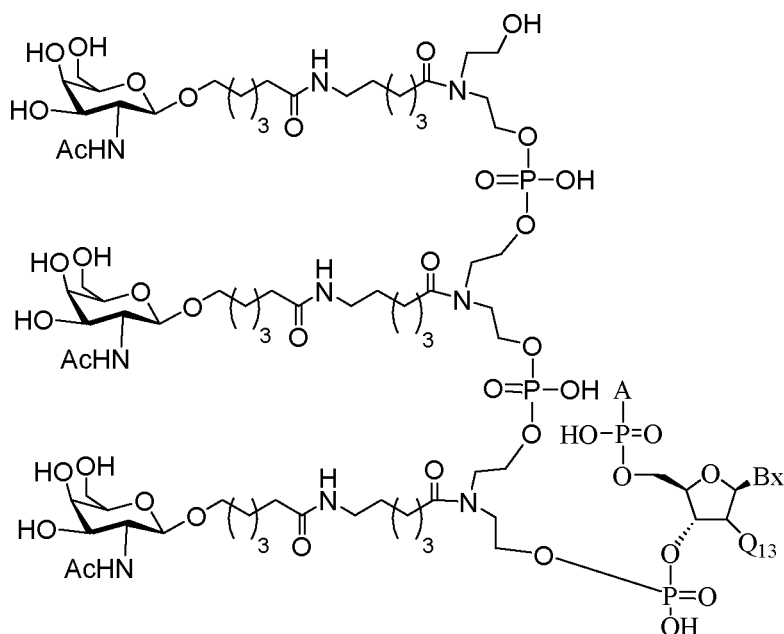
wherein Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

10

In certain embodiments, the compound has the following structure:



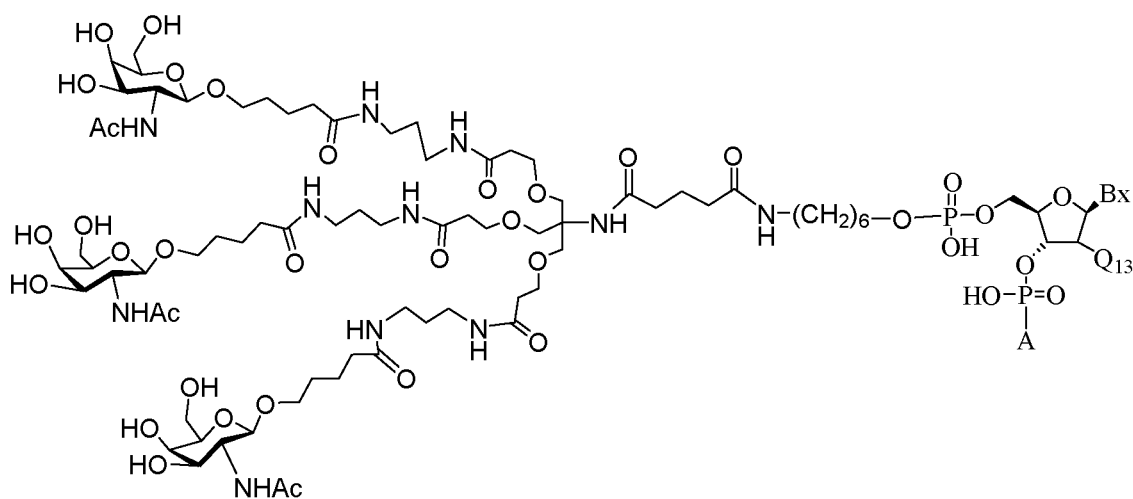
wherein Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

5

In certain embodiments, the conjugate group comprises:



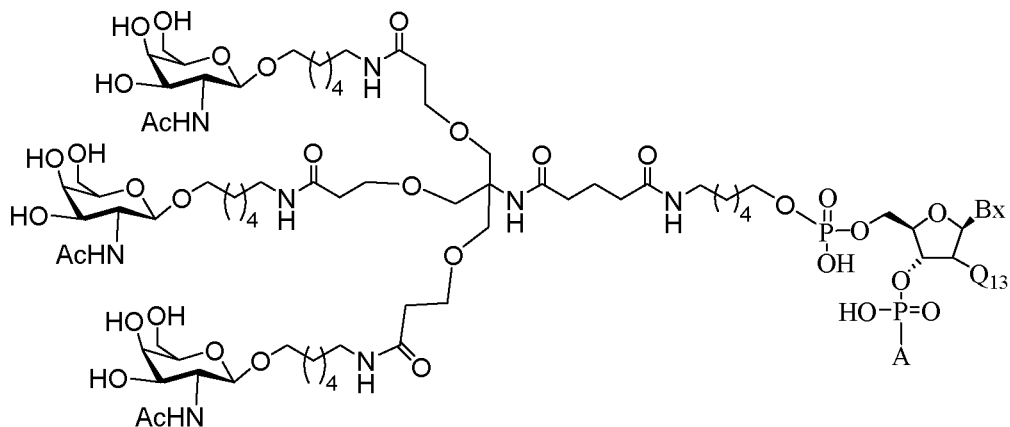
wherein Q13 is H or O(CH₂)₂-OCH₃;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

10

In certain embodiments, the conjugate group comprises:



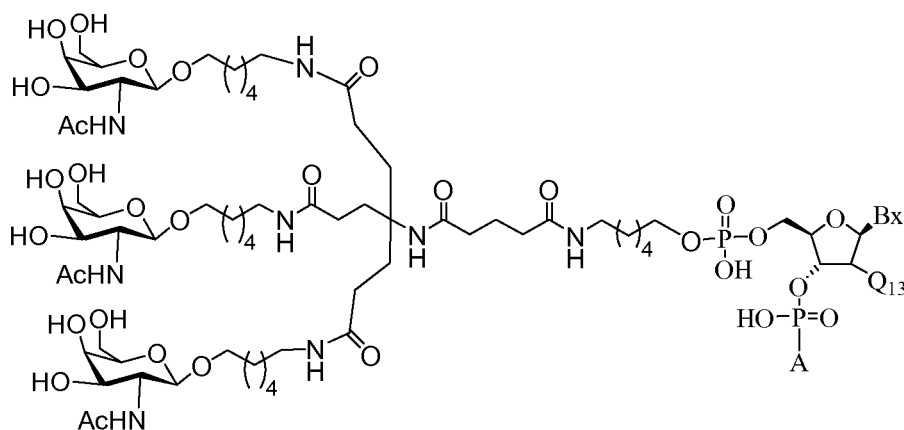
wherein Q13 is H or $\text{O}(\text{CH}_2)_2\text{-OCH}_3$;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

5

In certain embodiments, the conjugate group comprises:



wherein Q13 is H or $\text{O}(\text{CH}_2)_2\text{-OCH}_3$;

A is the modified oligonucleotide; and

Bx is a heterocyclic base moiety.

10

In certain embodiments, Bx is selected from among adenine, guanine, thymine, uracil, or cytosine, or 5-methyl cytosine. In certain embodiments, Bx is adenine. In certain embodiments, Bx is thymine. In certain embodiments, Q13 is $\text{O}(\text{CH}_2)_2\text{-OCH}_3$. In certain embodiments, Q13 is H.

15

In certain embodiments, the compound is in a salt form. In further embodiments, the compound further comprises of a pharmaceutically acceptable carrier or diluent. In certain embodiments, the compound comprises a modified oligonucleotide targeting apo(a) and a conjugate group, or a salt thereof, and a pharmaceutically acceptable carrier or diluent.

Certain embodiments provide a composition comprising a conjugated antisense compound as described herein, wherein the viscosity level of the compound is less than 40 centipoise (cP). In certain embodiments, the conjugated antisense compounds as described herein are efficacious by virtue of having a viscosity of less than 40 cP, less than 35 cP, less than 30 cP, less than 25 cP, less than 20 cP or less than 15 cP when measured by the parameters as described in Example 125.

Certain embodiments provide compositions and methods comprising administering to an animal a conjugated antisense compound or composition disclosed herein. In certain embodiments, administering the conjugated antisense compound prevents, treats, ameliorates, or slows progression of a cardiovascular, metabolic and/or inflammatory disease

Certain embodiments provide compositions and methods for use in therapy to treat an apo(a) related disease, disorder or condition. Certain embodiments provide compositions and methods for use in therapy to treat an Lp(a) related disease, disorder or condition. In certain embodiments, apo(a) and/or Lp(a) levels are elevated in an animal. In certain embodiments, the composition is a compound comprising an apo(a) specific inhibitor. In certain embodiments, the apo(a) specific inhibitor is a nucleic acid. In certain embodiments, the nucleic acid is an antisense compound. In certain embodiments, the antisense compound is a modified oligonucleotide targeting apo(a). In certain embodiments, the antisense compound is a modified oligonucleotide targeting apo(a) and a conjugate group. In certain embodiments, the modified oligonucleotide targeting apo(a) with the conjugate group, is used in treating, preventing, slowing progression, ameliorating a cardiovascular and/or metabolic disease, disorder or condition. In certain embodiments, the compositions and methods for therapy include administering an apo(a) specific inhibitor to an individual in need thereof.

Certain embodiments provide compositions and methods for reducing apo(a) levels. Certain embodiments provide compositions and methods for reducing Lp(a) levels. In certain embodiments, reducing apo(a) levels in a tissue, organ or subject improves the ratio of LDL to HDL or the ratio of TG to HDL. Certain embodiments provide compositions and methods to reduce apo(a) mRNA or protein expression in an animal comprising administering to the animal a conjugated antisense compound or composition disclosed herein to reduce apo(a) mRNA or protein expression in the animal. Certain embodiments provide compositions and methods to reduce Lp(a) levels in an animal comprising administering to the animal a conjugated antisense compound or composition disclosed herein to reduce apo(a) mRNA or protein expression in the animal.

Certain embodiments provide compositions and methods for preventing, treating, delaying, slowing the progression and/or ameliorating apo(a) related diseases, disorders, and conditions in a subject in need thereof. Certain embodiments provide compositions and methods for preventing, treating, delaying, slowing the progression and/or ameliorating Lp(a) related diseases, disorders, and conditions in a subject in need thereof. In certain embodiments, such diseases, disorders, and conditions include inflammatory, cardiovascular and/or metabolic diseases, disorders, and conditions. Certain such cardiovascular diseases, disorders or conditions include, but are not limited to, aortic stenosis, aneurysm (e.g., abdominal aortic

aneurysm), angina, arrhythmia, atherosclerosis, cerebrovascular disease, coronary artery disease, coronary heart disease, dyslipidemia, hypercholesterolemia, hyperlipidemia, hypertension, hypertriglyceridemia, myocardial infarction, peripheral vascular disease (e.g., peripheral artery disease, peripheral artery occlusive disease), retinal vascular occlusion, or stroke. Certain such metabolic diseases, disorders or conditions include, but are not limited to, hyperglycemia, prediabetes, diabetes (type I and type II), obesity, insulin resistance, metabolic syndrome and diabetic dyslipidemia. Certain such inflammatory diseases, disorders or conditions include, but are not limited to, aortic stenosis, coronary artery disease (CAD), Alzheimer's Disease and thromboembolic diseases, disorder or conditions. Certain thromboembolic diseases, disorders or conditions include, but are not limited to, stroke, thrombosis (e.g., venous thromboembolism), myocardial infarction and peripheral vascular disease. Certain embodiments provide compositions and methods for preventing, treating, delaying, slowing the progression and/or ameliorating aortic stenosis.

Certain embodiments provide a method of reducing at least one symptom of a cardiovascular disease, disorder or condition. In certain embodiments, the symptoms include, but are not limited to, angina, chest pain, shortness of breath, palpitations, weakness, dizziness, nausea, sweating, tachycardia, bradycardia, arrhythmia, atrial fibrillation, swelling in the lower extremities, cyanosis, fatigue, fainting, numbness of the face, numbness of the limbs, claudication or cramping of muscles, bloating of the abdomen, and fever. Certain embodiments provide a method of reducing at least one symptom of aortic stenosis.

In certain embodiments, the modulation of apo(a) or Lp(a) expression occurs in a cell, tissue or organ. In certain embodiments, the modulations occur in a cell, tissue or organ in an animal. In certain embodiments, the modulation is a reduction in apo(a) mRNA level. In certain embodiments, the modulation is a reduction in apo(a) protein level. In certain embodiments, both apo(a) mRNA and protein levels are reduced. In certain embodiments, the modulation is a reduction in Lp(a) level. Such reduction may occur in a time-dependent or in a dose-dependent manner.

In certain embodiments, the subject or animal is human.

In certain embodiments, the conjugated antisense compound is parenterally administered. In further embodiments, the parenteral administration is subcutaneous.

In certain embodiments, the conjugated antisense compound or composition is co-administered with a second agent or therapy. In certain embodiments, the conjugated antisense compound or composition and the second agent are administered concomitantly.

In certain embodiments, the second agent is a glucose-lowering agent. In certain embodiments, the second agent is a LDL, TG or cholesterol lowering agent. In certain embodiments, the second agent is an anti-inflammatory agent. In certain embodiments, the second agent is an Alzheimer Disease drug. In certain embodiments, the second agent can be, but is not limited to, a non-steroidal anti-inflammatory drug (NSAID e.g., aspirin), niacin (e.g., Niaspan), nicotinic acid, an apoB inhibitor (e.g., Mipomersen), a CETP inhibitor (e.g., Anacetrapib), an apo(a) inhibitor, a thyroid hormone analog (e.g., Eprotirome), a HMG-CoA reductase inhibitor (e.g., a statin), a fibrate (e.g., Gemfibrozil) and an microsomal triglyceride transfer protein inhibitor

(e.g., Lomitapide). The therapy can be, but is not limited to, Lp(a) apheresis. Agents or therapies can be co-administered or administered concomitantly. Agents or therapies can be sequentially or subsequently administered.

Certain embodiments provide use of a conjugated antisense compound targeted to apo(a) for decreasing apo(a) levels in an animal. Certain embodiments provide use of a conjugated antisense compound targeted to apo(a) for decreasing Lp(a) levels in an animal. Certain embodiments provide use of a conjugated antisense compounds targeted to apo(a) for the treatment, prevention, or amelioration of a disease, disorder, or condition associated with apo(a). Certain embodiments provide use of a conjugated antisense compounds targeted to apo(a) for the treatment, prevention, or amelioration of a disease, disorder, or condition associated with Lp(a).

Certain embodiments provide use of a conjugated antisense compound targeted to apo(a) in the preparation of a medicament for decreasing apo(a) levels in an animal. Certain embodiments provide use of a conjugated antisense compound targeted to apo(a) in the preparation of a medicament for decreasing Lp(a) levels in an animal. Certain embodiments provide use of a conjugated antisense compound for the preparation of a medicament for the treatment, prevention, or amelioration of a disease, disorder, or condition associated with apo(a). Certain embodiments provide use of a conjugated antisense compound for the preparation of a medicament for the treatment, prevention, or amelioration of a disease, disorder, or condition associated with Lp(a).

Certain embodiments provide the use of a conjugated antisense compound as described herein in the manufacture of a medicament for treating, ameliorating, delaying or preventing one or more of a disease related to apo(a) and/or Lp(a).

Certain embodiments provide a kit for treating, preventing, or ameliorating a disease, disorder or condition as described herein wherein the kit comprises: (i) an apo(a) specific inhibitor as described herein; and optionally (ii) a second agent or therapy as described herein.

A kit of the present invention can further include instructions for using the kit to treat, prevent, or ameliorate a disease, disorder or condition as described herein by combination therapy as described herein.

B. Certain Compounds

In certain embodiments, the invention provides conjugated antisense compounds comprising antisense oligonucleotides and a conjugate.

a. Certain Antisense Oligonucleotides

In certain embodiments, the invention provides antisense oligonucleotides. Such antisense oligonucleotides comprise linked nucleosides, each nucleoside comprising a sugar moiety and a nucleobase. The structure of such antisense oligonucleotides may be considered in terms of chemical features (e.g., modifications and patterns of modifications) and nucleobase sequence (e.g., sequence of antisense

oligonucleotide, identity and sequence of target nucleic acid).

i. **Certain Chemistry Features**

In certain embodiments, antisense oligonucleotide comprise one or more modification. In certain such embodiments, antisense oligonucleotides comprise one or more modified nucleosides and/or modified internucleoside linkages. In certain embodiments, modified nucleosides comprise a modified sugar moiety and/or modified nucleobase.

1. **Certain Sugar Moieties**

In certain embodiments, compounds of the disclosure comprise one or more modified nucleosides comprising a modified sugar moiety. Such compounds comprising one or more sugar-modified nucleosides may have desirable properties, such as enhanced nuclease stability or increased binding affinity with a target nucleic acid relative to an oligonucleotide comprising only nucleosides comprising naturally occurring sugar moieties. In certain embodiments, modified sugar moieties are substituted sugar moieties. In certain embodiments, modified sugar moieties are sugar surrogates. Such sugar surrogates may comprise one or more substitutions corresponding to those of substituted sugar moieties.

In certain embodiments, modified sugar moieties are substituted sugar moieties comprising one or more non-bridging sugar substituent, including but not limited to substituents at the 2' and/or 5' positions. Examples of sugar substituents suitable for the 2'-position, include, but are not limited to: 2'-F, 2'-OCH₃ ("OMe" or "O-methyl"), and 2'-O(CH₂)₂OCH₃ ("MOE"). In certain embodiments, sugar substituents at the 2' position is selected from allyl, amino, azido, thio, O-allyl, O-C₁-C₁₀ alkyl, O-C₁-C₁₀ substituted alkyl; OCF₃, O(CH₂)₂SCH₃, O(CH₂)₂-O-N(R_m)(R_n), and O-CH₂-C(=O)-N(R_m)(R_n), where each R_m and R_n is, independently, H or substituted or unsubstituted C₁-C₁₀ alkyl. Examples of sugar substituents at the 5'-position, include, but are not limited to: 5'-methyl (R or S); 5'-vinyl, and 5'-methoxy. In certain embodiments, substituted sugars comprise more than one non-bridging sugar substituent, for example, 2'-F-5'-methyl sugar moieties (*see, e.g.*, PCT International Application WO 2008/101157, for additional 5', 2'-bis substituted sugar moieties and nucleosides).

Nucleosides comprising 2'-substituted sugar moieties are referred to as 2'-substituted nucleosides. In certain embodiments, a 2'-substituted nucleoside comprises a 2'-substituent group selected from halo, allyl, amino, azido, SH, CN, OCN, CF₃, OCF₃, O, S, or N(R_m)-alkyl; O, S, or N(R_m)-alkenyl; O, S or N(R_m)-alkynyl; O-alkylenyl-O-alkyl, alkynyl, alkaryl, aralkyl, O-alkaryl, O-aralkyl, O(CH₂)₂SCH₃, O-(CH₂)₂-O-N(R_m)(R_n) or O-CH₂-C(=O)-N(R_m)(R_n), where each R_m and R_n is, independently, H, an amino protecting group or substituted or unsubstituted C₁-C₁₀ alkyl. These 2'-substituent groups can be further substituted with one or more substituent groups independently selected from hydroxyl, amino, alkoxy, carboxy, benzyl, phenyl, nitro (NO₂), thiol, thioalkoxy (S-alkyl), halogen, alkyl, aryl, alkenyl and alkynyl.

In certain embodiments, a 2'-substituted nucleoside comprises a 2'-substituent group selected from F, NH₂, N₃, OCF₃, O-CH₃, O(CH₂)₃NH₂, CH₂-CH=CH₂, O-CH₂-CH=CH₂, OCH₂CH₂OCH₃, O(CH₂)₂SCH₃,

O-(CH₂)₂-O-N(R_m)(R_n), O(CH₂)₂O(CH₂)₂N(CH₃)₂, and N-substituted acetamide (O-CH₂-C(=O)-N(R_m)(R_n)) where each R_m and R_n is, independently, H, an amino protecting group or substituted or unsubstituted C₁-C₁₀ alkyl.

In certain embodiments, a 2'-substituted nucleoside comprises a sugar moiety comprising a 2'-substituent group selected from F, OCF₃, O-CH₃, OCH₂CH₂OCH₃, O(CH₂)₂SCH₃, O-(CH₂)₂-O-N(CH₃)₂, -O(CH₂)₂O(CH₂)₂N(CH₃)₂, and O-CH₂-C(=O)-N(H)CH₃.

In certain embodiments, a 2'-substituted nucleoside comprises a sugar moiety comprising a 2'-substituent group selected from F, O-CH₃, and OCH₂CH₂OCH₃.

Certain modified sugar moieties comprise a bridging sugar substituent that forms a second ring resulting in a bicyclic sugar moiety. In certain such embodiments, the bicyclic sugar moiety comprises a bridge between the 4' and the 2' furanose ring atoms. Examples of such 4' to 2' sugar substituents, include, but are not limited to: -[C(R_a)(R_b)]_n-, -[C(R_a)(R_b)]_n-O-, -C(R_aR_b)-N(R)-O- or, -C(R_aR_b)-O-N(R)-; 4'-CH₂-2', 4'-(CH₂)₂-2', 4'-(CH₂)₃-2', 4'-(CH₂)-O-2' (LNA); 4'-(CH₂)-S-2'; 4'-(CH₂)₂-O-2' (ENA); 4'-CH(CH₃)-O-2' (cEt) and 4'-CH(CH₂OCH₃)-O-2', and analogs thereof (*see, e.g.*, U.S. Patent 7,399,845, issued on July 15, 2008); 4'-C(CH₃)(CH₃)-O-2' and analogs thereof (*see, e.g.*, WO2009/006478, published January 8, 2009); 4'-CH₂-N(OCH₃)-2' and analogs thereof (*see, e.g.*, WO2008/150729, published December 11, 2008); 4'-CH₂-O-N(CH₃)-2' (*see, e.g.*, US2004/0171570, published September 2, 2004); 4'-CH₂-O-N(R)-2', and 4'-CH₂-N(R)-O-2', wherein each R is, independently, H, a protecting group, or C₁-C₁₂ alkyl; 4'-CH₂-N(R)-O-2', wherein R is H, C₁-C₁₂ alkyl, or a protecting group (*see*, U.S. Patent 7,427,672, issued on September 23, 2008); 4'-CH₂-C(H)(CH₃)-2' (*see, e.g.*, Chattopadhyaya, *et al.*, *J. Org. Chem.*, 2009, 74, 118-134); and 4'-CH₂-C(=CH₂)-2' and analogs thereof (*see*, published PCT International Application WO 2008/154401, published on December 8, 2008).

In certain embodiments, such 4' to 2' bridges independently comprise from 1 to 4 linked groups independently selected from -[C(R_a)(R_b)]_n-, -C(R_a)=C(R_b)-, -C(R_a)=N-, -C(=NR_a)-, -C(=O)-, -C(=S)-, -O-, -Si(R_a)₂-, -S(=O)_x-, and -N(R_a)-;

wherein:

x is 0, 1, or 2;

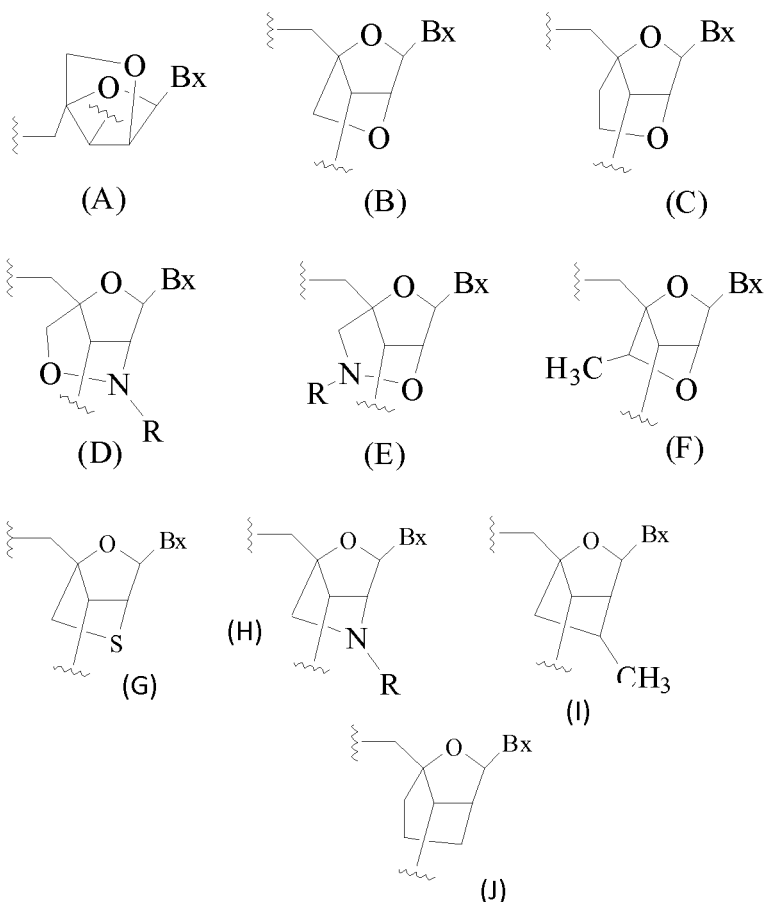
n is 1, 2, 3, or 4;

each R_a and R_b is, independently, H, a protecting group, hydroxyl, C₁-C₁₂ alkyl, substituted C₁-C₁₂ alkyl, C₂-C₁₂ alkenyl, substituted C₂-C₁₂ alkenyl, C₂-C₁₂ alkynyl, substituted C₂-C₁₂ alkynyl, C₅-C₂₀ aryl, substituted C₅-C₂₀ aryl, heterocycle radical, substituted heterocycle radical, heteroaryl, substituted heteroaryl, C₅-C₇ alicyclic radical, substituted C₅-C₇ alicyclic radical, halogen, OJ₁, NJ₁J₂, SJ₁, N₃, COOJ₁, acyl (C(=O)-H), substituted acyl, CN, sulfonyl (S(=O)₂-J₁), or sulfoxyl (S(=O)-J₁); and

each J₁ and J₂ is, independently, H, C₁-C₁₂ alkyl, substituted C₁-C₁₂ alkyl, C₂-C₁₂ alkenyl, substituted C₂-C₁₂ alkenyl, C₂-C₁₂ alkynyl, substituted C₂-C₁₂ alkynyl, C₅-C₂₀ aryl, substituted C₅-C₂₀ aryl, acyl (C(=O)-H), substituted acyl, a heterocycle radical, a substituted heterocycle radical, C₁-C₁₂ aminoalkyl, substituted

C₁-C₁₂ aminoalkyl, or a protecting group.

Nucleosides comprising bicyclic sugar moieties are referred to as bicyclic nucleosides or BNAs. Bicyclic nucleosides include, but are not limited to, (A) α -L-Methyleneoxy (4'-CH₂-O-2') BNA, (B) β -D-Methyleneoxy (4'-CH₂-O-2') BNA (also referred to as locked nucleic acid or LNA), (C) Ethyleneoxy (4'-(CH₂)₂-O-2') BNA, (D) Aminooxy (4'-CH₂-O-N(R)-2') BNA, (E) Oxyamino (4'-CH₂-N(R)-O-2') BNA, (F) Methyl(methyleneoxy) (4'-CH(CH₃)-O-2') BNA (also referred to as constrained ethyl or cEt), (G) methylene-thio (4'-CH₂-S-2') BNA, (H) methylene-amino (4'-CH₂-N(R)-2') BNA, (I) methyl carbocyclic (4'-CH₂-CH(CH₃)-2') BNA, and (J) propylene carbocyclic (4'-(CH₂)₃-2') BNA as depicted below.



wherein Bx is a nucleobase moiety and R is, independently, H, a protecting group, or C₁-C₁₂ alkyl.

Additional bicyclic sugar moieties are known in the art, for example: Singh et al., *Chem. Commun.*, 1998, 4, 455-456; Koshkin et al., *Tetrahedron*, 1998, 54, 3607-3630; Wahlestedt et al., *Proc. Natl. Acad. Sci. U. S. A.*, 2000, 97, 5633-5638; Kumar et al., *Bioorg. Med. Chem. Lett.*, 1998, 8, 2219-2222; Singh et al., *J. Org. Chem.*, 1998, 63, 10035-10039; Srivastava et al., *J. Am. Chem. Soc.*, 129(26) 8362-8379 (Jul. 4, 2007); Elayadi et al., *Curr. Opin. Invers. Drugs*, 2001, 2, 558-561; Braasch et al., *Chem. Biol.*, 2001, 8, 1-7; Orum et al., *Curr. Opin. Mol. Ther.*, 2001, 3, 239-243; U.S. Patent Nos. 7,053,207, 6,268,490, 6,770,748, 6,794,499, 7,034,133, 6,525,191, 6,670,461, and 7,399,845; WO 2004/106356, WO 1994/14226, WO 2005/021570, and WO 2007/134181; U.S. Patent Publication Nos. US2004/0171570, US2007/0287831, and

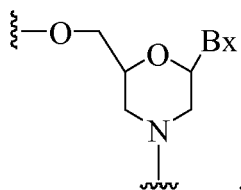
US2008/0039618; U.S. Patent Serial Nos. 12/129,154, 60/989,574, 61/026,995, 61/026,998, 61/056,564, 61/086,231, 61/097,787, and 61/099,844; and PCT International Applications Nos. PCT/US2008/064591, PCT/US2008/066154, and PCT/US2008/068922.

In certain embodiments, bicyclic sugar moieties and nucleosides incorporating such bicyclic sugar moieties are further defined by isomeric configuration. For example, a nucleoside comprising a 4'-2' methylene-oxy bridge, may be in the α -L configuration or in the β -D configuration. Previously, α -L-methyleneoxy (4'-CH₂-O-2') bicyclic nucleosides have been incorporated into antisense oligonucleotides that showed antisense activity (Frieden *et al.*, *Nucleic Acids Research*, 2003, 21, 6365-6372).

In certain embodiments, substituted sugar moieties comprise one or more non-bridging sugar substituent and one or more bridging sugar substituent (e.g., 5'-substituted and 4'-2' bridged sugars). (*see*, PCT International Application WO 2007/134181, published on 11/22/07, wherein LNA is substituted with, for example, a 5'-methyl or a 5'-vinyl group).

In certain embodiments, modified sugar moieties are sugar surrogates. In certain such embodiments, the oxygen atom of the naturally occurring sugar is substituted, e.g., with a sulfur, carbon or nitrogen atom. In certain such embodiments, such modified sugar moiety also comprises bridging and/or non-bridging substituents as described above. For example, certain sugar surrogates comprise a 4'-sulfur atom and a substitution at the 2'-position (*see*, e.g., published U.S. Patent Application US2005/0130923, published on June 16, 2005) and/or the 5' position. By way of additional example, carbocyclic bicyclic nucleosides having a 4'-2' bridge have been described (*see, e.g.*, Freier *et al.*, *Nucleic Acids Research*, 1997, 25(22), 4429-4443 and Albaek *et al.*, *J. Org. Chem.*, 2006, 71, 7731-7740).

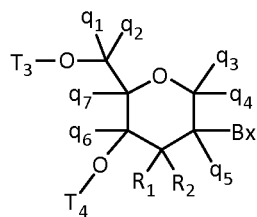
In certain embodiments, sugar surrogates comprise rings having other than 5-atoms. For example, in certain embodiments, a sugar surrogate comprises a morpholino. Morpholino compounds and their use in oligomeric compounds has been reported in numerous patents and published articles (*see* for example: Braasch *et al.*, *Biochemistry*, 2002, 41, 4503-4510; and U.S. Patents 5,698,685; 5,166,315; 5,185,444; and 5,034,506). As used here, the term "morpholino" means a sugar surrogate having the following structure:



In certain embodiments, morpholinos may be modified, for example by adding or altering various substituent groups from the above morpholino structure. Such sugar surrogates are referred to herein as "modified morpholinos."

For another example, in certain embodiments, a sugar surrogate comprises a six-membered tetrahydropyran. Such tetrahydropyrans may be further modified or substituted. Nucleosides comprising

such modified tetrahydropyrans include, but are not limited to, hexitol nucleic acid (HNA), anitol nucleic acid (ANA), manitol nucleic acid (MNA) (*see* Leumann, C.J. *Bioorg. & Med. Chem.* (2002) **10**:841-854), fluoro HNA (F-HNA), and those compounds having Formula VI:



VI

wherein independently for each of said at least one tetrahydropyran nucleoside analog of Formula VI:

Bx is a nucleobase moiety;

5 T₃ and T₄ are each, independently, an internucleoside linking group linking the tetrahydropyran nucleoside analog to the antisense compound or one of T₃ and T₄ is an internucleoside linking group linking the tetrahydropyran nucleoside analog to the antisense compound and the other of T₃ and T₄ is H, a hydroxyl protecting group, a linked conjugate group, or a 5' or 3'-terminal group;

10 q₁, q₂, q₃, q₄, q₅, q₆ and q₇ are each, independently, H, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₂-C₆ alkenyl, substituted C₂-C₆ alkenyl, C₂-C₆ alkynyl, or substituted C₂-C₆ alkynyl; and

each of R₁ and R₂ is independently selected from among: hydrogen, halogen, substituted or unsubstituted alkoxy, NJ₁J₂, SJ₁, N₃, OC(=X)J₁, OC(=X)NJ₁J₂, NJ₃C(=X)NJ₁J₂, and CN, wherein X is O, S or NJ₁, and each J₁, J₂, and J₃ is, independently, H or C₁-C₆ alkyl.

15 In certain embodiments, the modified THP nucleosides of Formula VI are provided wherein q₁, q₂, q₃, q₄, q₅, q₆ and q₇ are each H. In certain embodiments, at least one of q₁, q₂, q₃, q₄, q₅, q₆ and q₇ is other than H. In certain embodiments, at least one of q₁, q₂, q₃, q₄, q₅, q₆ and q₇ is methyl. In certain embodiments, THP nucleosides of Formula VI are provided wherein one of R₁ and R₂ is F. In certain embodiments, R₁ is fluoro and R₂ is H, R₁ is methoxy and R₂ is H, and R₁ is methoxyethoxy and R₂ is H.

20 Many other bicyclo and tricyclo sugar surrogate ring systems are also known in the art that can be used to modify nucleosides for incorporation into antisense compounds (*see, e.g.*, review article: Leumann, J. C, *Bioorganic & Medicinal Chemistry*, **2002**, *10*, 841-854).

25 Combinations of modifications are also provided without limitation, such as 2'-F-5'-methyl substituted nucleosides (see PCT International Application WO 2008/101157 Published on 8/21/08 for other disclosed 5', 2'-bis substituted nucleosides) and replacement of the ribosyl ring oxygen atom with S and further substitution at the 2'-position (see published U.S. Patent Application US2005-0130923, published on June 16, 2005) or alternatively 5'-substitution of a bicyclic nucleic acid (see PCT International Application WO 2007/134181, published on 11/22/07 wherein a 4'-CH₂-O-2' bicyclic nucleoside is further substituted at the 5' position with a 5'-methyl or a 5'-vinyl group). The synthesis and preparation of carbocyclic bicyclic

nucleosides along with their oligomerization and biochemical studies have also been described (*see, e.g.,* Srivastava *et al.*, *J. Am. Chem. Soc.* 2007, 129(26), 8362-8379).

In certain embodiments, the present disclosure provides oligonucleotides comprising modified nucleosides. Those modified nucleotides may include modified sugars, modified nucleobases, and/or modified linkages. The specific modifications are selected such that the resulting oligonucleotides possess desirable characteristics. In certain embodiments, oligonucleotides comprise one or more RNA-like nucleosides. In certain embodiments, oligonucleotides comprise one or more DNA-like nucleotides.

2. Certain Nucleobase Modifications

In certain embodiments, nucleosides of the present disclosure comprise one or more unmodified nucleobases. In certain embodiments, nucleosides of the present disclosure comprise one or more modified nucleobases.

In certain embodiments, modified nucleobases are selected from: universal bases, hydrophobic bases, promiscuous bases, size-expanded bases, and fluorinated bases as defined herein. 5-substituted pyrimidines, 6-azapyrimidines and N-2, N-6 and O-6 substituted purines, including 2-aminopropyladenine, 5-propynyluracil; 5-propynylcytosine; 5-hydroxymethyl cytosine, xanthine, hypoxanthine, 2-aminoadenine, 6-methyl and other alkyl derivatives of adenine and guanine, 2-propyl and other alkyl derivatives of adenine and guanine, 2-thiouracil, 2-thiothymine and 2-thiocytosine, 5-halouracil and cytosine, 5-propynyl ($-C\equiv C-CH_3$) uracil and cytosine and other alkynyl derivatives of pyrimidine bases, 6-azo uracil, cytosine and thymine, 5-uracil (pseudouracil), 4-thiouracil, 8-halo, 8-amino, 8-thiol, 8-thioalkyl, 8-hydroxyl and other 8-substituted adenines and guanines, 5-halo particularly 5-bromo, 5-trifluoromethyl and other 5-substituted uracils and cytosines, 7-methylguanine and 7-methyladenine, 2-F-adenine, 2-amino-adenine, 8-azaguanine and 8-azaadenine, 7-deazaguanine and 7-deazaadenine, 3-deazaguanine and 3-deazaadenine, universal bases, hydrophobic bases, promiscuous bases, size-expanded bases, and fluorinated bases as defined herein. Further modified nucleobases include tricyclic pyrimidines such as phenoxazine cytidine([5,4-b][1,4]benzoxazin-2(3H)-one), phenothiazine cytidine (1H-pyrimido[5,4-b][1,4]benzothiazin-2(3H)-one), G-clamps such as a substituted phenoxazine cytidine (e.g. 9-(2-aminoethoxy)-H-pyrimido[5,4-b][1,4]benzoxazin-2(3H)-one), carbazole cytidine (2H-pyrimido[4,5-b]indol-2-one), pyridoindole cytidine (H-pyrido[3',2':4,5]pyrrolo[2,3-d]pyrimidin-2-one). Modified nucleobases may also include those in which the purine or pyrimidine base is replaced with other heterocycles, for example 7-deaza-adenine, 7-deazaguanosine, 2-aminopyridine and 2-pyridone. Further nucleobases include those disclosed in United States Patent No. 3,687,808, those disclosed in *The Concise Encyclopedia Of Polymer Science And Engineering*, Kroschwitz, J.I., Ed., John Wiley & Sons, 1990, 858-859; those disclosed by Englisch *et al.*, *Angewandte Chemie*, International Edition, 1991, 30, 613; and those disclosed by Sanghvi, Y.S., Chapter 15, *Antisense Research and Applications*, Crooke, S.T. and Lebleu, B., Eds., CRC Press, 1993, 273-288.

Representative United States patents that teach the preparation of certain of the above noted modified nucleobases as well as other modified nucleobases include without limitation, U.S. 3,687,808; 4,845,205;

5,130,302; 5,134,066; 5,175,273; 5,367,066; 5,432,272; 5,457,187; 5,459,255; 5,484,908; 5,502,177; 5,525,711; 5,552,540; 5,587,469; 5,594,121; 5,596,091; 5,614,617; 5,645,985; 5,681,941; 5,750,692; 5,763,588; 5,830,653 and 6,005,096, certain of which are commonly owned with the instant application, and each of which is herein incorporated by reference in its entirety.

3. Certain Internucleoside Linkages

In certain embodiments, the present disclosure provides oligonucleotides comprising linked nucleosides. In such embodiments, nucleosides may be linked together using any internucleoside linkage. The two main classes of internucleoside linking groups are defined by the presence or absence of a phosphorus atom. Representative phosphorus containing internucleoside linkages include, but are not limited to, phosphodiester (PO), phosphotriesters, methylphosphonates, phosphoramidate, and phosphorothioates (PS). Representative non-phosphorus containing internucleoside linking groups include, but are not limited to, methylenemethylimino ($-\text{CH}_2-\text{N}(\text{CH}_3)-\text{O}-\text{CH}_2-$), thiodiester ($-\text{O}-\text{C}(\text{O})-\text{S}-$), thionocarbamate ($-\text{O}-\text{C}(\text{O})(\text{NH})-\text{S}-$); siloxane ($-\text{O}-\text{Si}(\text{H})_2-\text{O}-$); and N,N'-dimethylhydrazine ($-\text{CH}_2-\text{N}(\text{CH}_3)-\text{N}(\text{CH}_3)-$). Modified linkages, compared to natural phosphodiester linkages, can be used to alter, typically increase, nuclease resistance of the oligonucleotide. In certain embodiments, internucleoside linkages having a chiral atom can be prepared as a racemic mixture, or as separate enantiomers. Representative chiral linkages include, but are not limited to, alkylphosphonates and phosphorothioates. Methods of preparation of phosphorous-containing and non-phosphorous-containing internucleoside linkages are well known to those skilled in the art.

The oligonucleotides described herein contain one or more asymmetric centers and thus give rise to enantiomers, diastereomers, and other stereoisomeric configurations that may be defined, in terms of absolute stereochemistry, as (R) or (S), a or b such as for sugar anomers, or as (D) or (L) such as for amino acids etc. Included in the antisense compounds provided herein are all such possible isomers, as well as their racemic and optically pure forms.

Neutral internucleoside linkages include without limitation, phosphotriesters, methylphosphonates, MMI ($3'-\text{CH}_2-\text{N}(\text{CH}_3)-\text{O}-5'$), amide-3 ($3'-\text{CH}_2-\text{C}(=\text{O})-\text{N}(\text{H})-5'$), amide-4 ($3'-\text{CH}_2-\text{N}(\text{H})-\text{C}(=\text{O})-5'$), formacetal ($3'-\text{O}-\text{CH}_2-\text{O}-5'$), and thioformacetal ($3'-\text{S}-\text{CH}_2-\text{O}-5'$). Further neutral internucleoside linkages include nonionic linkages comprising siloxane (dialkylsiloxane), carboxylate ester, carboxamide, sulfide, sulfonate ester and amides (See for example: *Carbohydrate Modifications in Antisense Research*; Y.S. Sanghvi and P.D. Cook, Eds., ACS Symposium Series 580; Chapters 3 and 4, 40-65). Further neutral internucleoside linkages include nonionic linkages comprising mixed N, O, S and CH_2 component parts.

4. Certain Motifs

In certain embodiments, antisense oligonucleotides comprise one or more modified nucleoside (e.g., nucleoside comprising a modified sugar and/or modified nucleobase) and/or one or more modified internucleoside linkage. The pattern of such modifications on an oligonucleotide is referred to herein as a motif. In certain embodiments, sugar, nucleobase, and linkage motifs are independent of one another.

a. Certain sugar motifs

In certain embodiments, oligonucleotides comprise one or more type of modified sugar moieties and/or naturally occurring sugar moieties arranged along an oligonucleotide or region thereof in a defined pattern or sugar modification motif. Such motifs may include any of the sugar modifications discussed herein and/or other known sugar modifications.

5 In certain embodiments, the oligonucleotides comprise or consist of a region having a gapmer sugar motif, which comprises two external regions or “wings” and a central or internal region or “gap.” The three regions of a gapmer sugar motif (the 5'-wing, the gap, and the 3'-wing) form a contiguous sequence of nucleosides wherein at least some of the sugar moieties of the nucleosides of each of the wings differ from at least some of the sugar moieties of the nucleosides of the gap. Specifically, at least the sugar moieties of the
10 nucleosides of each wing that are closest to the gap (the 3'-most nucleoside of the 5'-wing and the 5'-most nucleoside of the 3'-wing) differ from the sugar moiety of the neighboring gap nucleosides, thus defining the boundary between the wings and the gap. In certain embodiments, the sugar moieties within the gap are the same as one another. In certain embodiments, the gap includes one or more nucleoside having a sugar moiety that differs from the sugar moiety of one or more other nucleosides of the gap. In certain embodiments, the
15 sugar motifs of the two wings are the same as one another (symmetric sugar gapmer). In certain embodiments, the sugar motifs of the 5'-wing differs from the sugar motif of the 3'-wing (asymmetric sugar gapmer).

i. Certain 5'-wings

In certain embodiments, the 5'- wing of a gapmer consists of 1 to 8 linked nucleosides. In certain
20 embodiments, the 5'- wing of a gapmer consists of 1 to 7 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 1 to 6 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 1 to 5 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 2 to 5 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 3 to 5 linked nucleosides. In certain
25 embodiments, the 5'- wing of a gapmer consists of 4 or 5 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 1 to 4 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 1 to 3 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 1 or 2 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 2 to 4 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 2 or 3 linked nucleosides. In certain
30 embodiments, the 5'- wing of a gapmer consists of 3 or 4 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 1 nucleoside. In certain embodiments, the 5'- wing of a gapmer consists of 2 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 3 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 4 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 5 linked nucleosides. In certain embodiments, the 5'- wing of a gapmer consists of 6 linked nucleosides.

35 In certain embodiments, the 5'- wing of a gapmer comprises at least one bicyclic nucleoside. In certain embodiments, the 5'- wing of a gapmer comprises at least two bicyclic nucleosides. In certain

embodiments, the 5'- wing of a gapmer comprises at least three bicyclic nucleosides. In certain embodiments, the 5'- wing of a gapmer comprises at least four bicyclic nucleosides. In certain embodiments, the 5'- wing of a gapmer comprises at least one constrained ethyl nucleoside. In certain embodiments, the 5'- wing of a gapmer comprises at least one LNA nucleoside. In certain embodiments, each nucleoside of the 5'- wing of a gapmer is a bicyclic nucleoside. In certain embodiments, each nucleoside of the 5'- wing of a gapmer is a constrained ethyl nucleoside. In certain embodiments, each nucleoside of the 5'- wing of a gapmer is a LNA nucleoside.

In certain embodiments, the 5'- wing of a gapmer comprises at least one non-bicyclic modified nucleoside. In certain embodiments, the 5'- wing of a gapmer comprises at least one 2'-substituted nucleoside. In certain embodiments, the 5'- wing of a gapmer comprises at least one 2'-MOE nucleoside. In certain embodiments, the 5'- wing of a gapmer comprises at least one 2'-OMe nucleoside. In certain embodiments, each nucleoside of the 5'- wing of a gapmer is a non-bicyclic modified nucleoside. In certain embodiments, each nucleoside of the 5'- wing of a gapmer is a 2'-substituted nucleoside. In certain embodiments, each nucleoside of the 5'- wing of a gapmer is a 2'-MOE nucleoside. In certain embodiments, each nucleoside of the 5'- wing of a gapmer is a 2'-OMe nucleoside.

In certain embodiments, the 5'- wing of a gapmer comprises at least one 2'-deoxynucleoside. In certain embodiments, each nucleoside of the 5'- wing of a gapmer is a 2'-deoxynucleoside. In certain embodiments, the 5'- wing of a gapmer comprises at least one ribonucleoside. In certain embodiments, each nucleoside of the 5'- wing of a gapmer is a ribonucleoside. In certain embodiments, one, more than one, or each of the nucleosides of the 5'- wing is an RNA-like nucleoside.

In certain embodiments, the 5'-wing of a gapmer comprises at least one bicyclic nucleoside and at least one non-bicyclic modified nucleoside. In certain embodiments, the 5'-wing of a gapmer comprises at least one bicyclic nucleoside and at least one 2'-substituted nucleoside. In certain embodiments, the 5'-wing of a gapmer comprises at least one bicyclic nucleoside and at least one 2'-MOE nucleoside. In certain embodiments, the 5'-wing of a gapmer comprises at least one bicyclic nucleoside and at least one 2'-OMe nucleoside. In certain embodiments, the 5'-wing of a gapmer comprises at least one bicyclic nucleoside and at least one 2'-deoxynucleoside.

In certain embodiments, the 5'-wing of a gapmer comprises at least one constrained ethyl nucleoside and at least one non-bicyclic modified nucleoside. In certain embodiments, the 5'-wing of a gapmer comprises at least one constrained ethyl nucleoside and at least one 2'-substituted nucleoside. In certain embodiments, the 5'-wing of a gapmer comprises at least one constrained ethyl nucleoside and at least one 2'-MOE nucleoside. In certain embodiments, the 5'-wing of a gapmer comprises at least one constrained ethyl nucleoside and at least one 2'-OMe nucleoside. In certain embodiments, the 5'-wing of a gapmer comprises at least one constrained ethyl nucleoside and at least one 2'-deoxynucleoside.

ii. Certain 3'-wings

In certain embodiments, the 3'- wing of a gapmer consists of 1 to 8 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 1 to 7 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 1 to 6 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 1 to 5 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 2 to 5
5 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 3 to 5 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 4 or 5 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 1 to 4 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 1 to 3 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 1 or 2 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 2 to 4
10 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 2 or 3 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 3 or 4 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 1 nucleoside. In certain embodiments, the 3'- wing of a gapmer consists of 2 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 3 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 4 linked nucleosides. In
15 certain embodiments, the 3'- wing of a gapmer consists of 5 linked nucleosides. In certain embodiments, the 3'- wing of a gapmer consists of 6 linked nucleosides.

In certain embodiments, the 3'- wing of a gapmer comprises at least one bicyclic nucleoside. In certain embodiments, the 3'- wing of a gapmer comprises at least one constrained ethyl nucleoside. In certain embodiments, the 3'- wing of a gapmer comprises at least one LNA nucleoside. In certain embodiments,
20 each nucleoside of the 3'- wing of a gapmer is a bicyclic nucleoside. In certain embodiments, each nucleoside of the 3'- wing of a gapmer is a constrained ethyl nucleoside. In certain embodiments, each nucleoside of the 3'- wing of a gapmer is a LNA nucleoside.

In certain embodiments, the 3'- wing of a gapmer comprises at least one non-bicyclic modified nucleoside. In certain embodiments, the 3'- wing of a gapmer comprises at least two non-bicyclic modified
25 nucleosides. In certain embodiments, the 3'- wing of a gapmer comprises at least three non-bicyclic modified nucleosides. In certain embodiments, the 3'- wing of a gapmer comprises at least four non-bicyclic modified nucleosides. In certain embodiments, the 3'- wing of a gapmer comprises at least one 2'-substituted nucleoside. In certain embodiments, the 3'- wing of a gapmer comprises at least one 2'-MOE nucleoside. In
30 certain embodiments, the 3'- wing of a gapmer comprises at least one 2'-OMe nucleoside. In certain embodiments, each nucleoside of the 3'- wing of a gapmer is a non-bicyclic modified nucleoside. In certain embodiments, each nucleoside of the 3'- wing of a gapmer is a 2'-substituted nucleoside. In certain embodiments, each nucleoside of the 3'- wing of a gapmer is a 2'-MOE nucleoside. In certain embodiments, each nucleoside of the 3'- wing of a gapmer is a 2'-OMe nucleoside.

In certain embodiments, the 3'- wing of a gapmer comprises at least one 2'-deoxynucleoside. In
35 certain embodiments, each nucleoside of the 3'- wing of a gapmer is a 2'-deoxynucleoside. In a certain embodiments, the 3'- wing of a gapmer comprises at least one ribonucleoside. In certain embodiments, each

nucleoside of the 3'-wing of a gapmer is a ribonucleoside. In certain embodiments, one, more than one, or each of the nucleosides of the 5'-wing is an RNA-like nucleoside.

In certain embodiments, the 3'-wing of a gapmer comprises at least one bicyclic nucleoside and at least one non-bicyclic modified nucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one bicyclic nucleoside and at least one 2'-substituted nucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one bicyclic nucleoside and at least one 2'-MOE nucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one bicyclic nucleoside and at least one 2'-OMe nucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one bicyclic nucleoside and at least one 2'-deoxynucleoside.

In certain embodiments, the 3'-wing of a gapmer comprises at least one constrained ethyl nucleoside and at least one non-bicyclic modified nucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one constrained ethyl nucleoside and at least one 2'-substituted nucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one constrained ethyl nucleoside and at least one 2'-MOE nucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one constrained ethyl nucleoside and at least one 2'-OMe nucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one constrained ethyl nucleoside and at least one 2'-deoxynucleoside.

In certain embodiments, the 3'-wing of a gapmer comprises at least one LNA nucleoside and at least one non-bicyclic modified nucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one LNA nucleoside and at least one 2'-substituted nucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one LNA nucleoside and at least one 2'-MOE nucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one LNA nucleoside and at least one 2'-OMe nucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one LNA nucleoside and at least one 2'-deoxynucleoside.

In certain embodiments, the 3'-wing of a gapmer comprises at least one bicyclic nucleoside, at least one non-bicyclic modified nucleoside, and at least one 2'-deoxynucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one constrained ethyl nucleoside, at least one non-bicyclic modified nucleoside, and at least one 2'-deoxynucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one LNA nucleoside, at least one non-bicyclic modified nucleoside, and at least one 2'-deoxynucleoside.

In certain embodiments, the 3'-wing of a gapmer comprises at least one bicyclic nucleoside, at least one 2'-substituted nucleoside, and at least one 2'-deoxynucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one constrained ethyl nucleoside, at least one 2'-substituted nucleoside, and at least one 2'-deoxynucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one LNA nucleoside, at least one 2'-substituted nucleoside, and at least one 2'-deoxynucleoside.

In certain embodiments, the 3'-wing of a gapmer comprises at least one bicyclic nucleoside, at least one 2'-MOE nucleoside, and at least one 2'-deoxynucleoside. In certain embodiments, the 3'-wing of a

gapmer comprises at least one constrained ethyl nucleoside, at least one 2'-MOE nucleoside, and at least one 2'-deoxynucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one LNA nucleoside, at least one 2'-MOE nucleoside, and at least one 2'-deoxynucleoside.

In certain embodiments, the 3'-wing of a gapmer comprises at least one bicyclic nucleoside, at least one 2'-OMe nucleoside, and at least one 2'-deoxynucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one constrained ethyl nucleoside, at least one 2'-OMe nucleoside, and at least one 2'-deoxynucleoside. In certain embodiments, the 3'-wing of a gapmer comprises at least one LNA nucleoside, at least one 2'-OMe nucleoside, and at least one 2'-deoxynucleoside.

iii. Certain Central Regions (gaps)

In certain embodiments, the gap of a gapmer consists of 6 to 20 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 6 to 15 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 6 to 12 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 6 to 10 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 6 to 9 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 6 to 8 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 6 or 7 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 7 to 10 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 7 to 9 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 7 or 8 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 8 to 10 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 8 or 9 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 6 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 7 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 8 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 9 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 10 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 11 linked nucleosides. In certain embodiments, the gap of a gapmer consists of 12 linked nucleosides.

In certain embodiments, each nucleoside of the gap of a gapmer is a 2'-deoxynucleoside. In certain embodiments, the gap comprises one or more modified nucleosides. In certain embodiments, each nucleoside of the gap of a gapmer is a 2'-deoxynucleoside or is a modified nucleoside that is "DNA-like." In such embodiments, "DNA-like" means that the nucleoside has similar characteristics to DNA, such that a duplex comprising the gapmer and an RNA molecule is capable of activating RNase H. For example, under certain conditions, 2'-(ara)-F have been shown to support RNase H activation, and thus is DNA-like. In certain embodiments, one or more nucleosides of the gap of a gapmer is not a 2'-deoxynucleoside and is not DNA-like. In certain such embodiments, the gapmer nonetheless supports RNase H activation (e.g., by virtue of the number or placement of the non-DNA nucleosides).

In certain embodiments, gaps comprise a stretch of unmodified 2'-deoxynucleoside interrupted by one or more modified nucleosides, thus resulting in three sub-regions (two stretches of one or more 2'-deoxynucleosides and a stretch of one or more interrupting modified nucleosides). In certain embodiments,

no stretch of unmodified 2'-deoxynucleosides is longer than 5, 6, or 7 nucleosides. In certain embodiments, such short stretches is achieved by using short gap regions. In certain embodiments, short stretches are achieved by interrupting a longer gap region.

In certain embodiments, the gap comprises one or more modified nucleosides. In certain
5 embodiments, the gap comprises one or more modified nucleosides selected from among cEt, FHNA, LNA, and 2-thio-thymidine. In certain embodiments, the gap comprises one modified nucleoside. In certain
embodiments, the gap comprises a 5'-substituted sugar moiety selected from among 5'-Me, and 5'-(*R*)-Me. In certain embodiments, the gap comprises two modified nucleosides. In certain embodiments, the gap
comprises three modified nucleosides. In certain embodiments, the gap comprises four modified nucleosides.
10 In certain embodiments, the gap comprises two or more modified nucleosides and each modified nucleoside is the same. In certain embodiments, the gap comprises two or more modified nucleosides and each modified
nucleoside is different.

In certain embodiments, the gap comprises one or more modified linkages. In certain embodiments,
the gap comprises one or more methyl phosphonate linkages. In certain embodiments the gap comprises two
15 or more modified linkages. In certain embodiments, the gap comprises one or more modified linkages and
one or more modified nucleosides. In certain embodiments, the gap comprises one modified linkage and one
modified nucleoside. In certain embodiments, the gap comprises two modified linkages and two or more
modified nucleosides.

20 **b. Certain Internucleoside Linkage Motifs**

In certain embodiments, oligonucleotides comprise modified internucleoside linkages arranged along
the oligonucleotide or region thereof in a defined pattern or modified internucleoside linkage motif. In
certain embodiments, oligonucleotides comprise a region having an alternating internucleoside linkage motif.
In certain embodiments, oligonucleotides of the present disclosure comprise a region of uniformly modified
25 internucleoside linkages. In certain such embodiments, the oligonucleotide comprises a region that is
uniformly linked by phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide
is uniformly linked by phosphorothioate internucleoside linkages. In certain embodiments, each
internucleoside linkage of the oligonucleotide is selected from phosphodiester and phosphorothioate. In
certain embodiments, each internucleoside linkage of the oligonucleotide is selected from phosphodiester and
30 phosphorothioate and at least one internucleoside linkage is phosphorothioate.

In certain embodiments, the oligonucleotide comprises at least 6 phosphorothioate internucleoside
linkages. In certain embodiments, the oligonucleotide comprises at least 7 phosphorothioate internucleoside
linkages. In certain embodiments, the oligonucleotide comprises at least 8 phosphorothioate internucleoside
linkages. In certain embodiments, the oligonucleotide comprises at least 9 phosphorothioate internucleoside
35 linkages. In certain embodiments, the oligonucleotide comprises at least 10 phosphorothioate internucleoside
linkages. In certain embodiments, the oligonucleotide comprises at least 11 phosphorothioate internucleoside

linkages. In certain embodiments, the oligonucleotide comprises at least 12 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises at least 13 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises at least 14 phosphorothioate internucleoside linkages.

5 In certain embodiments, the oligonucleotide comprises at least one block of at least 6 consecutive phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises at least one block of at least 7 consecutive phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises at least one block of at least 8 consecutive phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises at least one block of at least 9 consecutive
10 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises at least one block of at least 10 consecutive phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises at least one block of at least one 12 consecutive phosphorothioate internucleoside linkages. In certain such embodiments, at least one such block is located at the 3' end of the oligonucleotide. In certain such embodiments, at least one such block is located within 3 nucleosides of the 3' end of the
15 oligonucleotide. In certain embodiments, the oligonucleotide comprises less than 15 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises less than 14 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises less than 13 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises less than 12 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises less than 11 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises
20 less than 10 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises less than 9 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises less than 8 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises less than 7 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises less than 6 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide
25 comprises less than 5 phosphorothioate internucleoside linkages.

c. Certain Nucleobase Modification Motifs

In certain embodiments, oligonucleotides comprise chemical modifications to nucleobases arranged along the oligonucleotide or region thereof in a defined pattern or nucleobases modification motif. In certain
30 such embodiments, nucleobase modifications are arranged in a gapped motif. In certain embodiments, nucleobase modifications are arranged in an alternating motif. In certain embodiments, each nucleobase is modified. In certain embodiments, none of the nucleobases is chemically modified.

In certain embodiments, oligonucleotides comprise a block of modified nucleobases. In certain such embodiments, the block is at the 3'-end of the oligonucleotide. In certain embodiments the block is within 3
35 nucleotides of the 3'-end of the oligonucleotide. In certain such embodiments, the block is at the 5'-end of

the oligonucleotide. In certain embodiments the block is within 3 nucleotides of the 5'-end of the oligonucleotide.

In certain embodiments, nucleobase modifications are a function of the natural base at a particular position of an oligonucleotide. For example, in certain embodiments each purine or each pyrimidine in an oligonucleotide is modified. In certain embodiments, each adenine is modified. In certain embodiments, each guanine is modified. In certain embodiments, each thymine is modified. In certain embodiments, each cytosine is modified. In certain embodiments, each uracil is modified.

In certain embodiments, some, all, or none of the cytosine moieties in an oligonucleotide are 5-methyl cytosine moieties. Herein, 5-methyl cytosine is not a "modified nucleobase." Accordingly, unless otherwise indicated, unmodified nucleobases include both cytosine residues having a 5-methyl and those lacking a 5 methyl. In certain embodiments, the methylation state of all or some cytosine nucleobases is specified.

In certain embodiments, chemical modifications to nucleobases comprise attachment of certain conjugate groups to nucleobases. In certain embodiments, each purine or each pyrimidine in an oligonucleotide may be optionally modified to comprise a conjugate group.

d. Certain Overall Lengths

In certain embodiments, the present disclosure provides oligonucleotides of any of a variety of ranges of lengths. In certain embodiments, oligonucleotides consist of X to Y linked nucleosides, where X represents the fewest number of nucleosides in the range and Y represents the largest number of nucleosides in the range. In certain such embodiments, X and Y are each independently selected from 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, and 50; provided that $X \leq Y$. For example, in certain embodiments, the oligonucleotide may consist of 8 to 9, 8 to 10, 8 to 11, 8 to 12, 8 to 13, 8 to 14, 8 to 15, 8 to 16, 8 to 17, 8 to 18, 8 to 19, 8 to 20, 8 to 21, 8 to 22, 8 to 23, 8 to 24, 8 to 25, 8 to 26, 8 to 27, 8 to 28, 8 to 29, 8 to 30, 9 to 10, 9 to 11, 9 to 12, 9 to 13, 9 to 14, 9 to 15, 9 to 16, 9 to 17, 9 to 18, 9 to 19, 9 to 20, 9 to 21, 9 to 22, 9 to 23, 9 to 24, 9 to 25, 9 to 26, 9 to 27, 9 to 28, 9 to 29, 9 to 30, 10 to 11, 10 to 12, 10 to 13, 10 to 14, 10 to 15, 10 to 16, 10 to 17, 10 to 18, 10 to 19, 10 to 20, 10 to 21, 10 to 22, 10 to 23, 10 to 24, 10 to 25, 10 to 26, 10 to 27, 10 to 28, 10 to 29, 10 to 30, 11 to 12, 11 to 13, 11 to 14, 11 to 15, 11 to 16, 11 to 17, 11 to 18, 11 to 19, 11 to 20, 11 to 21, 11 to 22, 11 to 23, 11 to 24, 11 to 25, 11 to 26, 11 to 27, 11 to 28, 11 to 29, 11 to 30, 12 to 13, 12 to 14, 12 to 15, 12 to 16, 12 to 17, 12 to 18, 12 to 19, 12 to 20, 12 to 21, 12 to 22, 12 to 23, 12 to 24, 12 to 25, 12 to 26, 12 to 27, 12 to 28, 12 to 29, 12 to 30, 13 to 14, 13 to 15, 13 to 16, 13 to 17, 13 to 18, 13 to 19, 13 to 20, 13 to 21, 13 to 22, 13 to 23, 13 to 24, 13 to 25, 13 to 26, 13 to 27, 13 to 28, 13 to 29, 13 to 30, 14 to 15, 14 to 16, 14 to 17, 14 to 18, 14 to 19, 14 to 20, 14 to 21, 14 to 22, 14 to 23, 14 to 24, 14 to 25, 14 to 26, 14 to 27, 14 to 28, 14 to 29, 14 to 30, 15 to 16, 15 to 17, 15 to 18, 15 to 19, 15 to 20, 15 to 21, 15 to 22, 15 to 23, 15 to 24, 15 to 25, 15 to 26, 15 to 27, 15 to 28, 15 to 29, 15 to 30, 16 to 17, 16 to 18, 16 to 19, 16 to 20, 16 to 21, 16 to 22, 16 to 23, 16 to 24, 16 to 25, 16 to 26, 16 to 27, 16 to 28, 16 to 29, 16 to 30, 17 to 18, 17 to

19, 17 to 20, 17 to 21, 17 to 22, 17 to 23, 17 to 24, 17 to 25, 17 to 26, 17 to 27, 17 to 28, 17 to 29, 17 to 30, 18 to 19, 18 to 20, 18 to 21, 18 to 22, 18 to 23, 18 to 24, 18 to 25, 18 to 26, 18 to 27, 18 to 28, 18 to 29, 18 to 30, 19 to 20, 19 to 21, 19 to 22, 19 to 23, 19 to 24, 19 to 25, 19 to 26, 19 to 27, 19 to 28, 19 to 29, 19 to 30, 20 to 21, 20 to 22, 20 to 23, 20 to 24, 20 to 25, 20 to 26, 20 to 27, 20 to 28, 20 to 29, 20 to 30, 21 to 22, 21 to 23, 21 to 24, 21 to 25, 21 to 26, 21 to 27, 21 to 28, 21 to 29, 21 to 30, 22 to 23, 22 to 24, 22 to 25, 22 to 26, 22 to 27, 22 to 28, 22 to 29, 22 to 30, 23 to 24, 23 to 25, 23 to 26, 23 to 27, 23 to 28, 23 to 29, 23 to 30, 24 to 25, 24 to 26, 24 to 27, 24 to 28, 24 to 29, 24 to 30, 25 to 26, 25 to 27, 25 to 28, 25 to 29, 25 to 30, 26 to 27, 26 to 28, 26 to 29, 26 to 30, 27 to 28, 27 to 29, 27 to 30, 28 to 29, 28 to 30, or 29 to 30 linked nucleosides. In embodiments where the number of nucleosides of an oligonucleotide of a compound is limited, whether to a range or to a specific number, the compound may, nonetheless further comprise additional other substituents. For example, an oligonucleotide comprising 8-30 nucleosides excludes oligonucleotides having 31 nucleosides, but, unless otherwise indicated, such an oligonucleotide may further comprise, for example one or more conjugate groups, terminal groups, or other substituents.

Further, where an oligonucleotide is described by an overall length range and by regions having specified lengths, and where the sum of specified lengths of the regions is less than the upper limit of the overall length range, the oligonucleotide may have additional nucleosides, beyond those of the specified regions, provided that the total number of nucleosides does not exceed the upper limit of the overall length range.

5. Certain Antisense Oligonucleotide Chemistry Motifs

In certain embodiments, the chemical structural features of antisense oligonucleotides are characterized by their sugar motif, internucleoside linkage motif, nucleobase modification motif and overall length. In certain embodiments, such parameters are each independent of one another. Thus, each internucleoside linkage of an oligonucleotide having a gapmer sugar motif may be modified or unmodified and may or may not follow the gapmer modification pattern of the sugar modifications. Thus, the internucleoside linkages within the wing regions of a sugar-gapmer may be the same or different from one another and may be the same or different from the internucleoside linkages of the gap region. Likewise, such sugar-gapmer oligonucleotides may comprise one or more modified nucleobase independent of the gapmer pattern of the sugar modifications. One of skill in the art will appreciate that such motifs may be combined to create a variety of oligonucleotides.

In certain embodiments, the selection of internucleoside linkage and nucleoside modification are not independent of one another.

i. Certain Sequences and Targets

In certain embodiments, the invention provides antisense oligonucleotides having a sequence complementary to a target nucleic acid. Such antisense compounds are capable of hybridizing to a target nucleic acid, resulting in at least one antisense activity. In certain embodiments, antisense compounds

specifically hybridize to one or more target nucleic acid. In certain embodiments, a specifically hybridizing antisense compound has a nucleobase sequence comprising a region having sufficient complementarity to a target nucleic acid to allow hybridization and result in antisense activity and insufficient complementarity to any non-target so as to avoid or reduce non-specific hybridization to non-target nucleic acid sequences under conditions in which specific hybridization is desired (e.g., under physiological conditions for *in vivo* or therapeutic uses, and under conditions in which assays are performed in the case of *in vitro* assays). In certain embodiments, oligonucleotides are selective between a target and non-target, even though both target and non-target comprise the target sequence. In such embodiments, selectivity may result from relative accessibility of the target region of one nucleic acid molecule compared to the other.

In certain embodiments, the present disclosure provides antisense compounds comprising oligonucleotides that are fully complementary to the target nucleic acid over the entire length of the oligonucleotide. In certain embodiments, oligonucleotides are 99% complementary to the target nucleic acid. In certain embodiments, oligonucleotides are 95% complementary to the target nucleic acid. In certain embodiments, such oligonucleotides are 90% complementary to the target nucleic acid.

In certain embodiments, such oligonucleotides are 85% complementary to the target nucleic acid. In certain embodiments, such oligonucleotides are 80% complementary to the target nucleic acid. In certain embodiments, an antisense compound comprises a region that is fully complementary to a target nucleic acid and is at least 80% complementary to the target nucleic acid over the entire length of the oligonucleotide. In certain such embodiments, the region of full complementarity is from 6 to 14 nucleobases in length.

In certain embodiments, oligonucleotides comprise a hybridizing region and a terminal region. In certain such embodiments, the hybridizing region consists of 12-30 linked nucleosides and is fully complementary to the target nucleic acid. In certain embodiments, the hybridizing region includes one mismatch relative to the target nucleic acid. In certain embodiments, the hybridizing region includes two mismatches relative to the target nucleic acid. In certain embodiments, the hybridizing region includes three mismatches relative to the target nucleic acid. In certain embodiments, the terminal region consists of 1-4 terminal nucleosides. In certain embodiments, the terminal nucleosides are at the 3' end. In certain embodiments, one or more of the terminal nucleosides are not complementary to the target nucleic acid.

Antisense mechanisms include any mechanism involving the hybridization of an oligonucleotide with target nucleic acid, wherein the hybridization results in a biological effect. In certain embodiments, such hybridization results in either target nucleic acid degradation or occupancy with concomitant inhibition or stimulation of the cellular machinery involving, for example, translation, transcription, or splicing of the target nucleic acid.

One type of antisense mechanism involving degradation of target RNA is RNase H mediated antisense. RNase H is a cellular endonuclease which cleaves the RNA strand of an RNA:DNA duplex. It is known in the art that single-stranded antisense compounds which are "DNA-like" elicit RNase H activity in

mammalian cells. Activation of RNase H, therefore, results in cleavage of the RNA target, thereby greatly enhancing the efficiency of DNA-like oligonucleotide-mediated inhibition of gene expression.

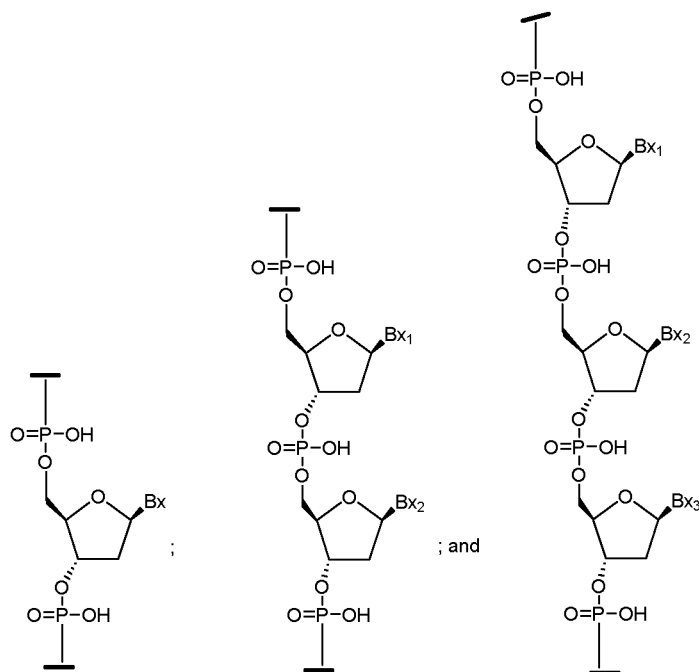
In certain embodiments, a conjugate group comprises a cleavable moiety. In certain embodiments, a conjugate group comprises one or more cleavable bond. In certain embodiments, a conjugate group comprises a linker. In certain embodiments, a linker comprises a protein binding moiety. In certain
 5 embodiments, a conjugate group comprises a cell-targeting moiety (also referred to as a cell-targeting group). In certain embodiments a cell-targeting moiety comprises a branching group. In certain embodiments, a cell-targeting moiety comprises one or more tethers. In certain embodiments, a cell-targeting moiety comprises a carbohydrate or carbohydrate cluster.

10 ii. Certain Cleavable Moieties

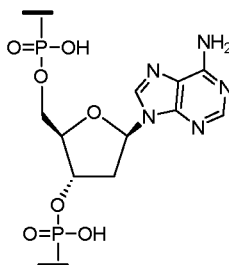
In certain embodiments, a cleavable moiety is a cleavable bond. In certain embodiments, a cleavable moiety comprises a cleavable bond. In certain embodiments, the conjugate group comprises a cleavable moiety. In certain such embodiments, the cleavable moiety attaches to the antisense
 15 oligonucleotide. In certain such embodiments, the cleavable moiety attaches directly to the cell-targeting moiety. In certain such embodiments, the cleavable moiety attaches to the conjugate linker. In certain embodiments, the cleavable moiety comprises a phosphate or phosphodiester. In certain embodiments, the cleavable moiety is a cleavable nucleoside or nucleoside analog. In certain embodiments, the nucleoside or nucleoside analog comprises an optionally protected heterocyclic base selected from a purine, substituted purine, pyrimidine or substituted pyrimidine. In certain embodiments, the cleavable moiety is a nucleoside
 20 comprising an optionally protected heterocyclic base selected from uracil, thymine, cytosine, 4-N-benzoylcytosine, 5-methylcytosine, 4-N-benzoyl-5-methylcytosine, adenine, 6-N-benzoyladenine, guanine and 2-N-isobutyrylguanine. In certain embodiments, the cleavable moiety is 2'-deoxy nucleoside that is attached to the 3' position of the antisense oligonucleotide by a phosphodiester linkage and is attached to the linker by a phosphodiester or phosphorothioate linkage. In certain embodiments, the cleavable moiety is 2'-
 25 deoxy adenosine that is attached to the 3' position of the antisense oligonucleotide by a phosphodiester linkage and is attached to the linker by a phosphodiester or phosphorothioate linkage. In certain embodiments, the cleavable moiety is 2'-deoxy adenosine that is attached to the 3' position of the antisense oligonucleotide by a phosphodiester linkage and is attached to the linker by a phosphodiester linkage.

In certain embodiments, the cleavable moiety is attached to the 3' position of the antisense
 30 oligonucleotide. In certain embodiments, the cleavable moiety is attached to the 5' position of the antisense oligonucleotide. In certain embodiments, the cleavable moiety is attached to a 2' position of the antisense oligonucleotide. In certain embodiments, the cleavable moiety is attached to the antisense oligonucleotide by a phosphodiester linkage. In certain embodiments, the cleavable moiety is attached to the linker by either a phosphodiester or a phosphorothioate linkage. In certain embodiments, the cleavable moiety is attached to
 35 the linker by a phosphodiester linkage. In certain embodiments, the conjugate group does not include a cleavable moiety.

In certain embodiments, the cleavable moiety is cleaved after the complex has been administered to an animal only after being internalized by a targeted cell. Inside the cell the cleavable moiety is cleaved thereby releasing the active antisense oligonucleotide. While not wanting to be bound by theory it is believed that the cleavable moiety is cleaved by one or more nucleases within the cell. In certain embodiments, the one or more nucleases cleave the phosphodiester linkage between the cleavable moiety and the linker. In certain embodiments, the cleavable moiety has a structure selected from among the following:



wherein each of Bx, Bx₁, Bx₂, and Bx₃ is independently a heterocyclic base moiety. In certain embodiments, the cleavable moiety has a structure selected from among the following:



iii. Certain Linkers

In certain embodiments, the conjugate groups comprise a linker. In certain such embodiments, the linker is covalently bound to the cleavable moiety. In certain such embodiments, the linker is covalently bound to the antisense oligonucleotide. In certain embodiments, the linker is covalently bound to a cell-

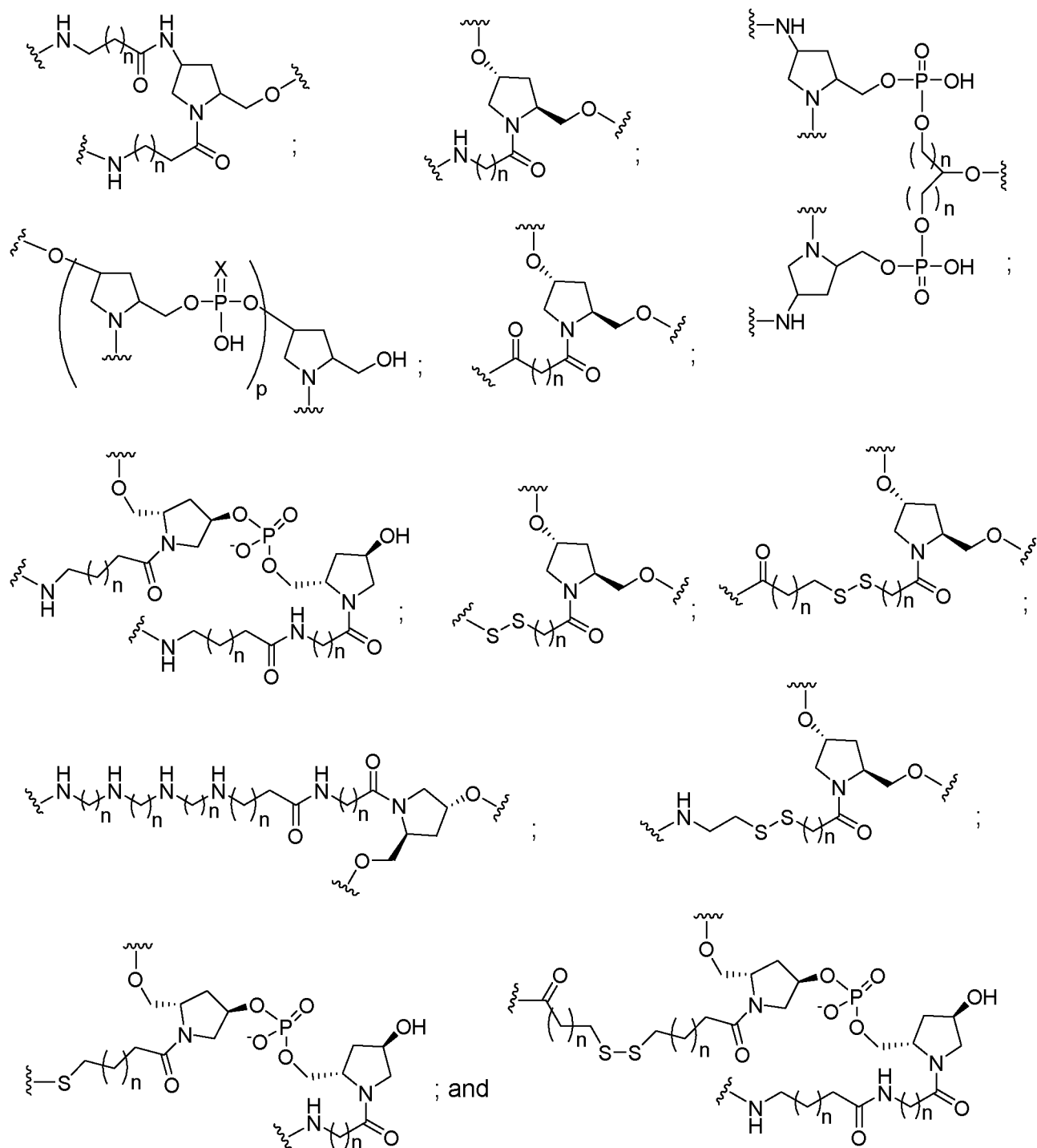
targeting moiety. In certain embodiments, the linker further comprises a covalent attachment to a solid support. In certain embodiments, the linker further comprises a covalent attachment to a protein binding moiety. In certain embodiments, the linker further comprises a covalent attachment to a solid support and further comprises a covalent attachment to a protein binding moiety. In certain embodiments, the linker includes multiple positions for attachment of tethered ligands. In certain embodiments, the linker includes multiple positions for attachment of tethered ligands and is not attached to a branching group. In certain embodiments, the linker further comprises one or more cleavable bond. In certain embodiments, the conjugate group does not include a linker.

In certain embodiments, the linker includes at least a linear group comprising groups selected from alkyl, amide, disulfide, polyethylene glycol, ether, thioether (-S-) and hydroxylamino (-O-N(H)-) groups. In certain embodiments, the linear group comprises groups selected from alkyl, amide and ether groups. In certain embodiments, the linear group comprises groups selected from alkyl and ether groups. In certain embodiments, the linear group comprises at least one phosphorus linking group. In certain embodiments, the linear group comprises at least one phosphodiester group. In certain embodiments, the linear group includes at least one neutral linking group. In certain embodiments, the linear group is covalently attached to the cell-targeting moiety and the cleavable moiety. In certain embodiments, the linear group is covalently attached to the cell-targeting moiety and the antisense oligonucleotide. In certain embodiments, the linear group is covalently attached to the cell-targeting moiety, the cleavable moiety and a solid support. In certain embodiments, the linear group is covalently attached to the cell-targeting moiety, the cleavable moiety, a solid support and a protein binding moiety. In certain embodiments, the linear group includes one or more cleavable bond.

In certain embodiments, the linker includes the linear group covalently attached to a scaffold group. In certain embodiments, the scaffold includes a branched aliphatic group comprising groups selected from alkyl, amide, disulfide, polyethylene glycol, ether, thioether and hydroxylamino groups. In certain embodiments, the scaffold includes a branched aliphatic group comprising groups selected from alkyl, amide and ether groups. In certain embodiments, the scaffold includes at least one mono or polycyclic ring system. In certain embodiments, the scaffold includes at least two mono or polycyclic ring systems. In certain embodiments, the linear group is covalently attached to the scaffold group and the scaffold group is covalently attached to the cleavable moiety and the linker. In certain embodiments, the linear group is covalently attached to the scaffold group and the scaffold group is covalently attached to the cleavable moiety, the linker and a solid support. In certain embodiments, the linear group is covalently attached to the scaffold group and the scaffold group is covalently attached to the cleavable moiety, the linker and a protein binding moiety. In certain embodiments, the linear group is covalently attached to the scaffold group and the scaffold group is covalently attached to the cleavable moiety, the linker, a protein binding moiety and a solid support. In certain embodiments, the scaffold group includes one or more cleavable bond.

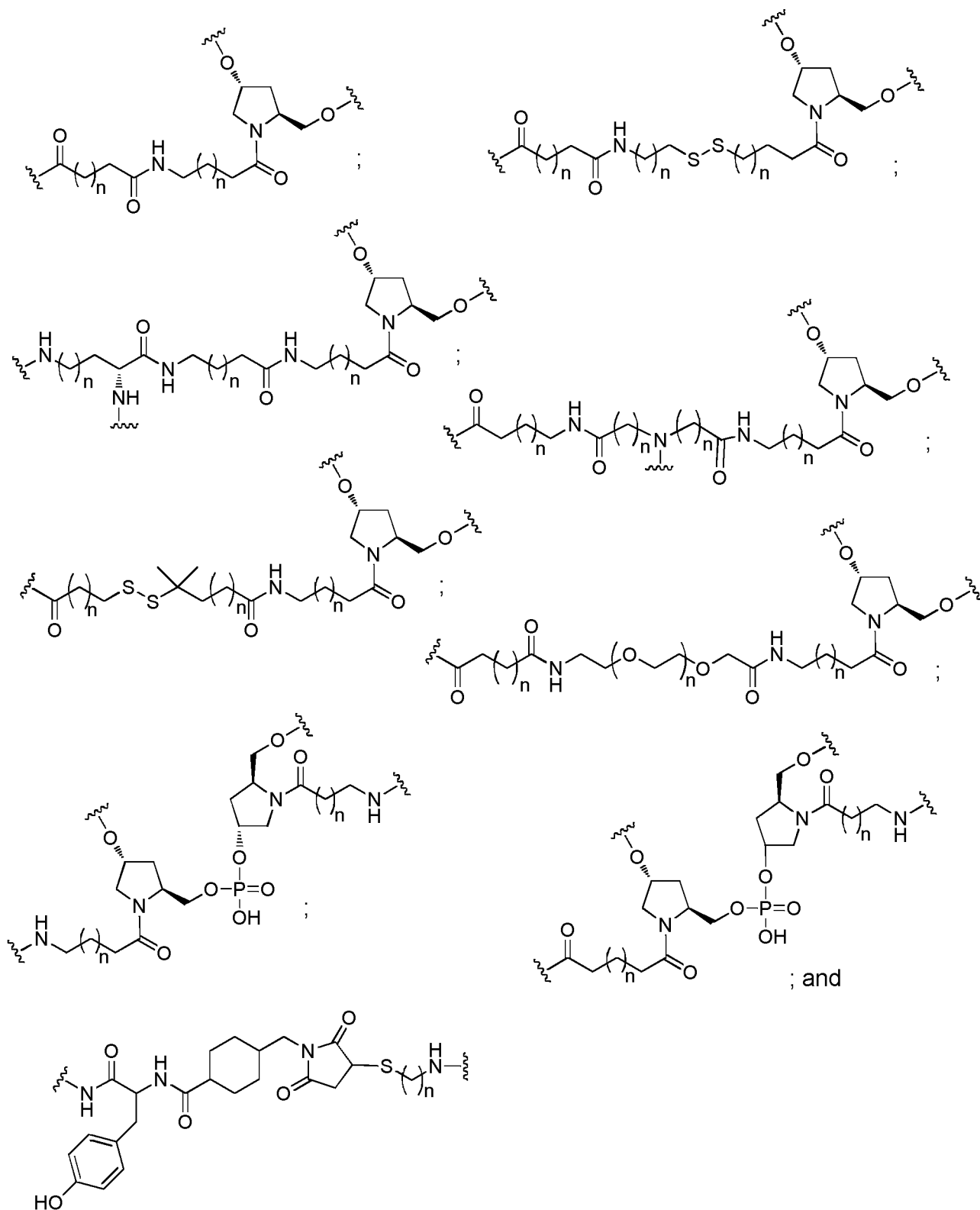
In certain embodiments, the linker includes a protein binding moiety. In certain embodiments, the protein binding moiety is a lipid such as for example including but not limited to cholesterol, cholic acid, adamantane acetic acid, 1-pyrene butyric acid, dihydrotestosterone, 1,3-Bis-O(hexadecyl)glycerol, geranyloxyhexyl group, hexadecylglycerol, borneol, menthol, 1,3-propanediol, heptadecyl group, palmitic acid, myristic acid, O3-(oleoyl)lithocholic acid, O3-(oleoyl)cholenic acid, dimethoxytrityl, or phenoxazine), a vitamin (e.g., folate, vitamin A, vitamin E, biotin, pyridoxal), a peptide, a carbohydrate (e.g., monosaccharide, disaccharide, trisaccharide, tetrasaccharide, oligosaccharide, polysaccharide), an endosomolytic component, a steroid (e.g., uvaol, hecigenin, diosgenin), a terpene (e.g., triterpene, e.g., sarsasapogenin, friedelin, epifriedelanol derivatized lithocholic acid), or a cationic lipid. In certain
5
10
embodiments, the protein binding moiety is a C16 to C22 long chain saturated or unsaturated fatty acid, cholesterol, cholic acid, vitamin E, adamantane or 1-pentafluoropropyl.

In certain embodiments, a linker has a structure selected from among:



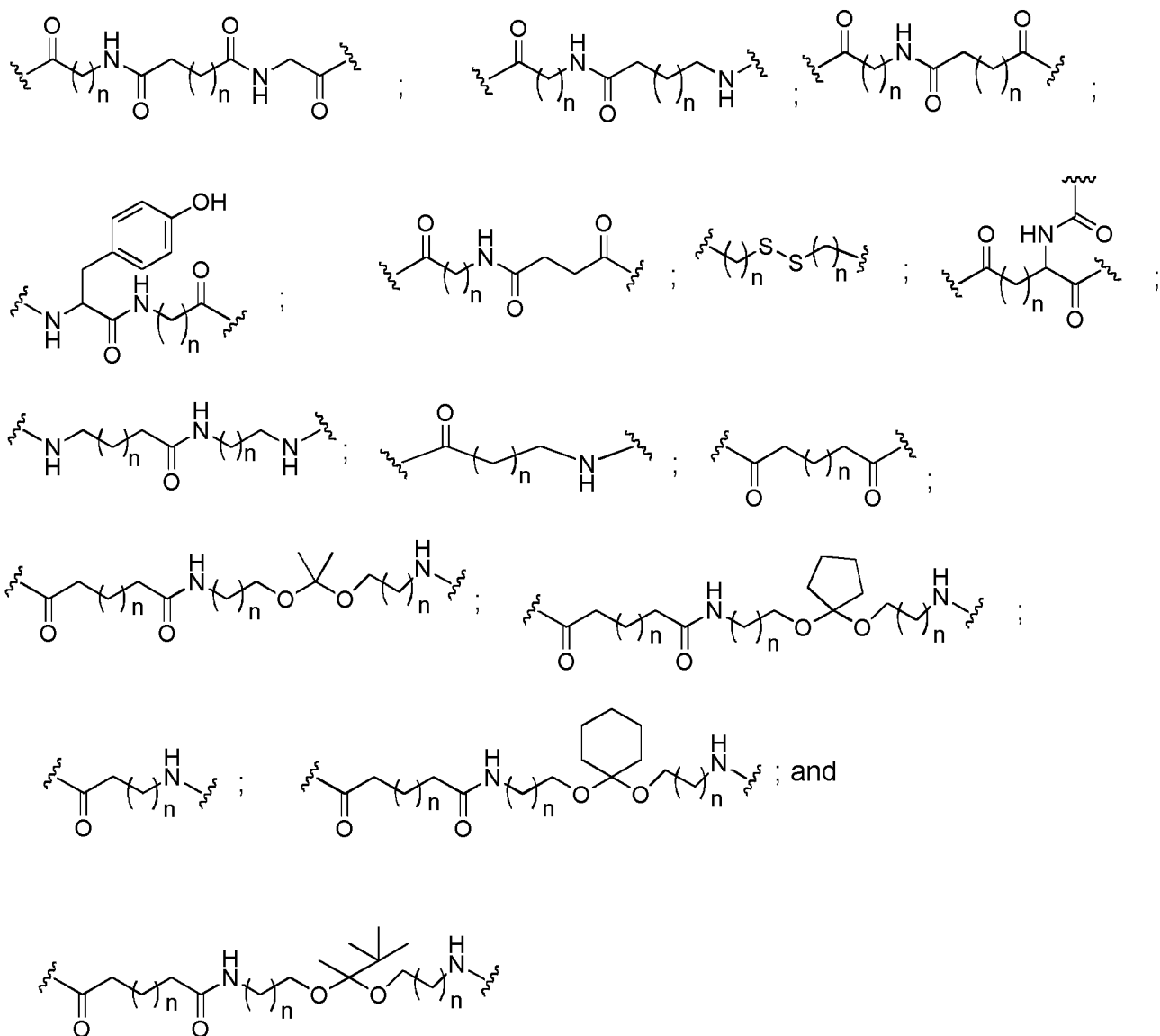
wherein each n is, independently, from 1 to 20; and p is from 1 to 6.

In certain embodiments, a linker has a structure selected from among:



5 wherein each n is, independently, from 1 to 20.

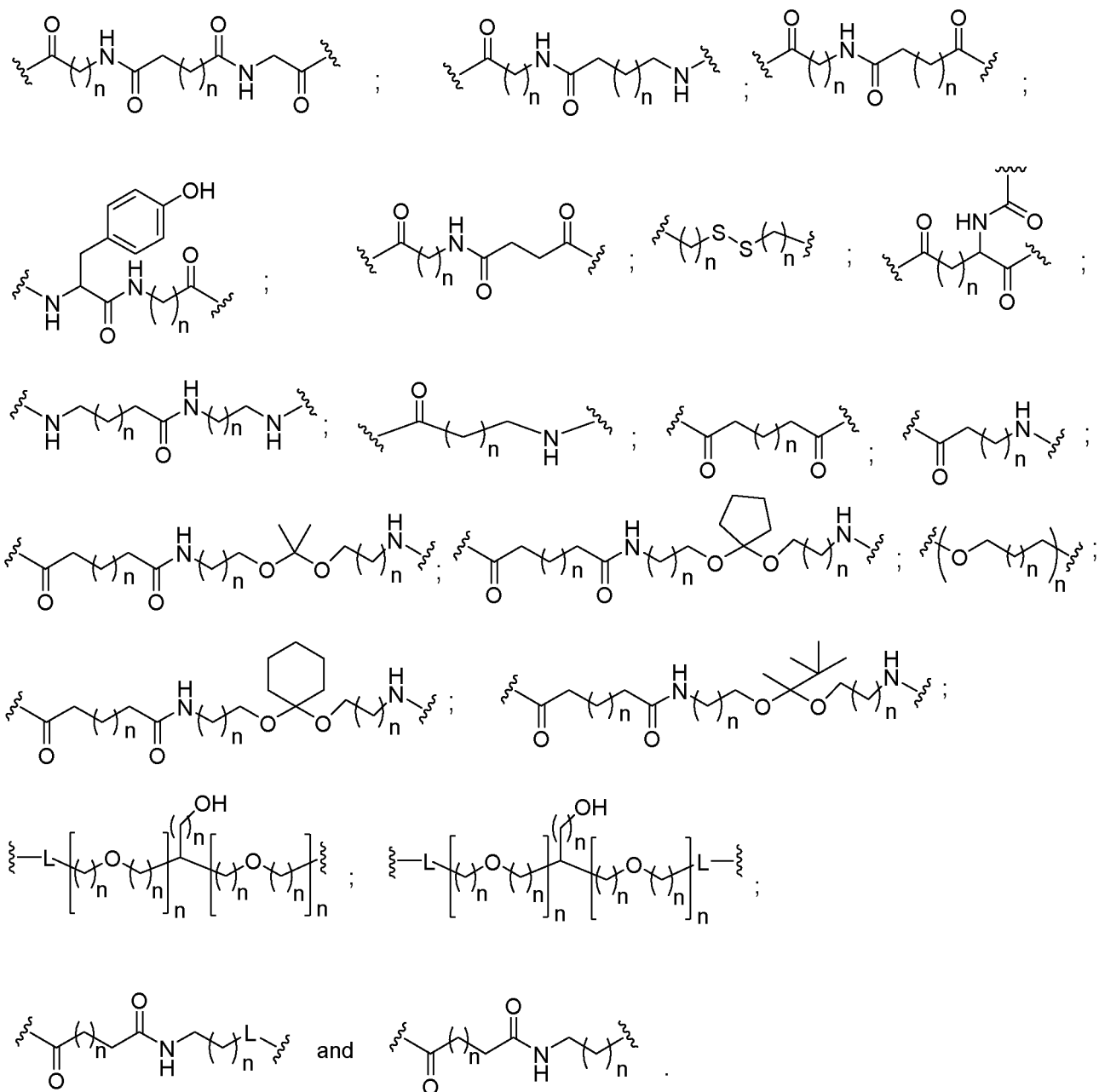
In certain embodiments, a linker has a structure selected from among:



5

wherein n is from 1 to 20.

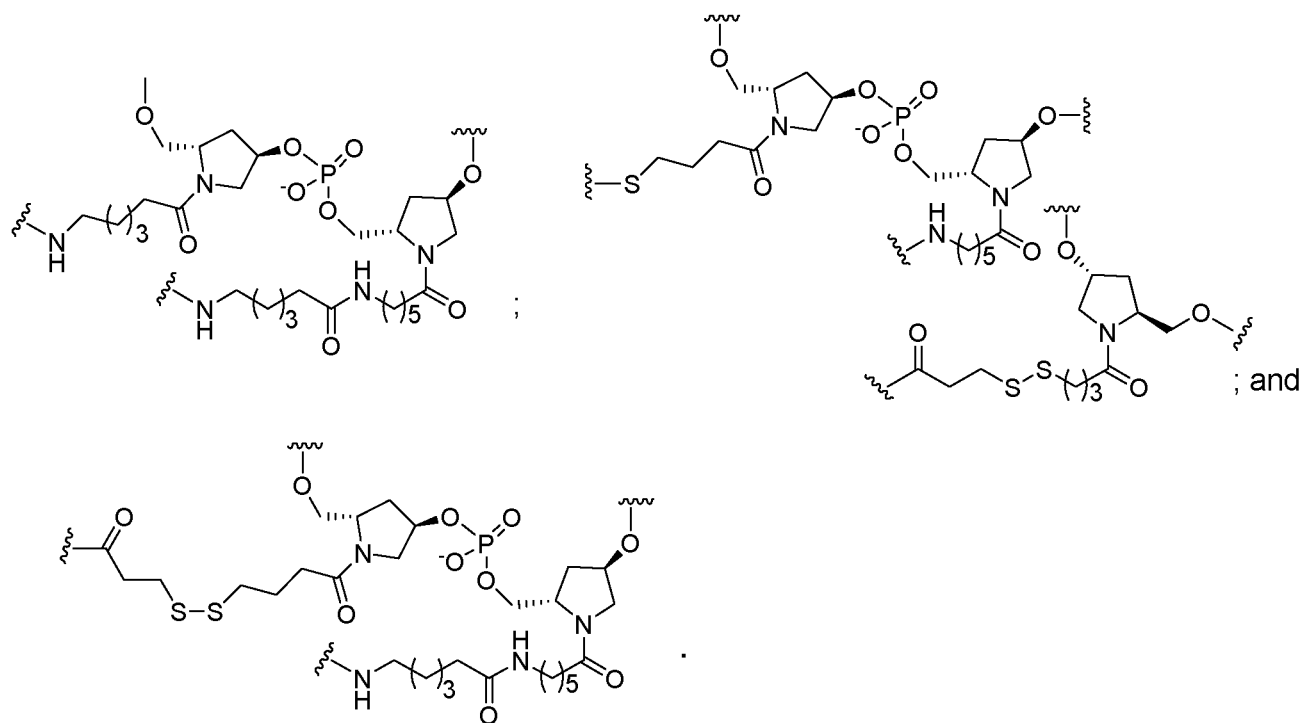
In certain embodiments, a linker has a structure selected from among:



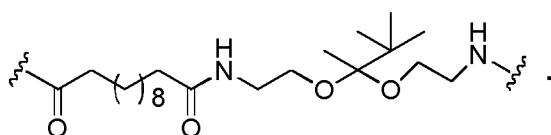
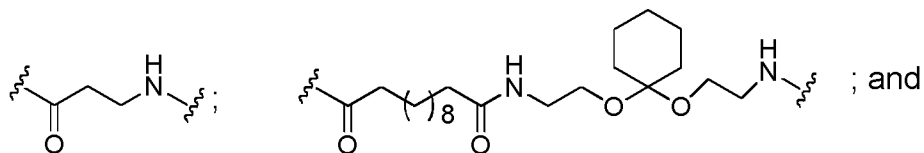
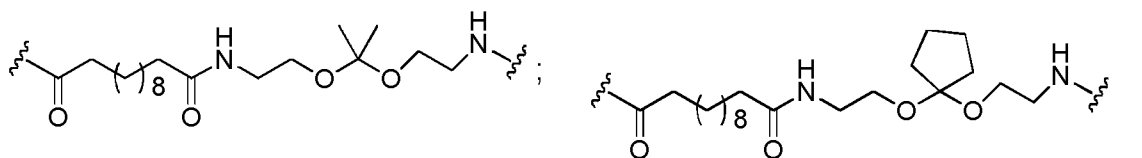
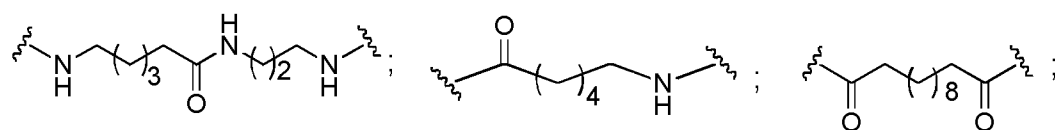
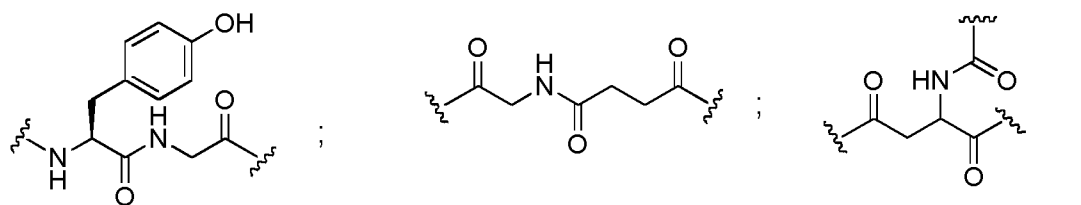
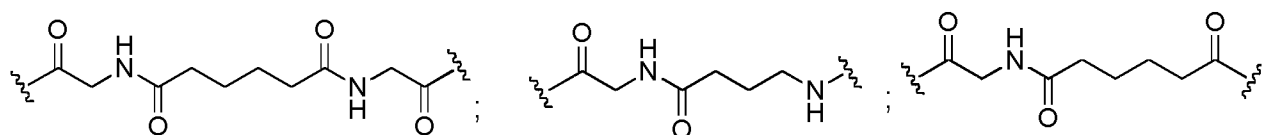
wherein each L is, independently, a phosphorus linking group or a neutral linking group; and
 5 each n is, independently, from 1 to 20.

In certain embodiments, a linker has a structure selected from among:

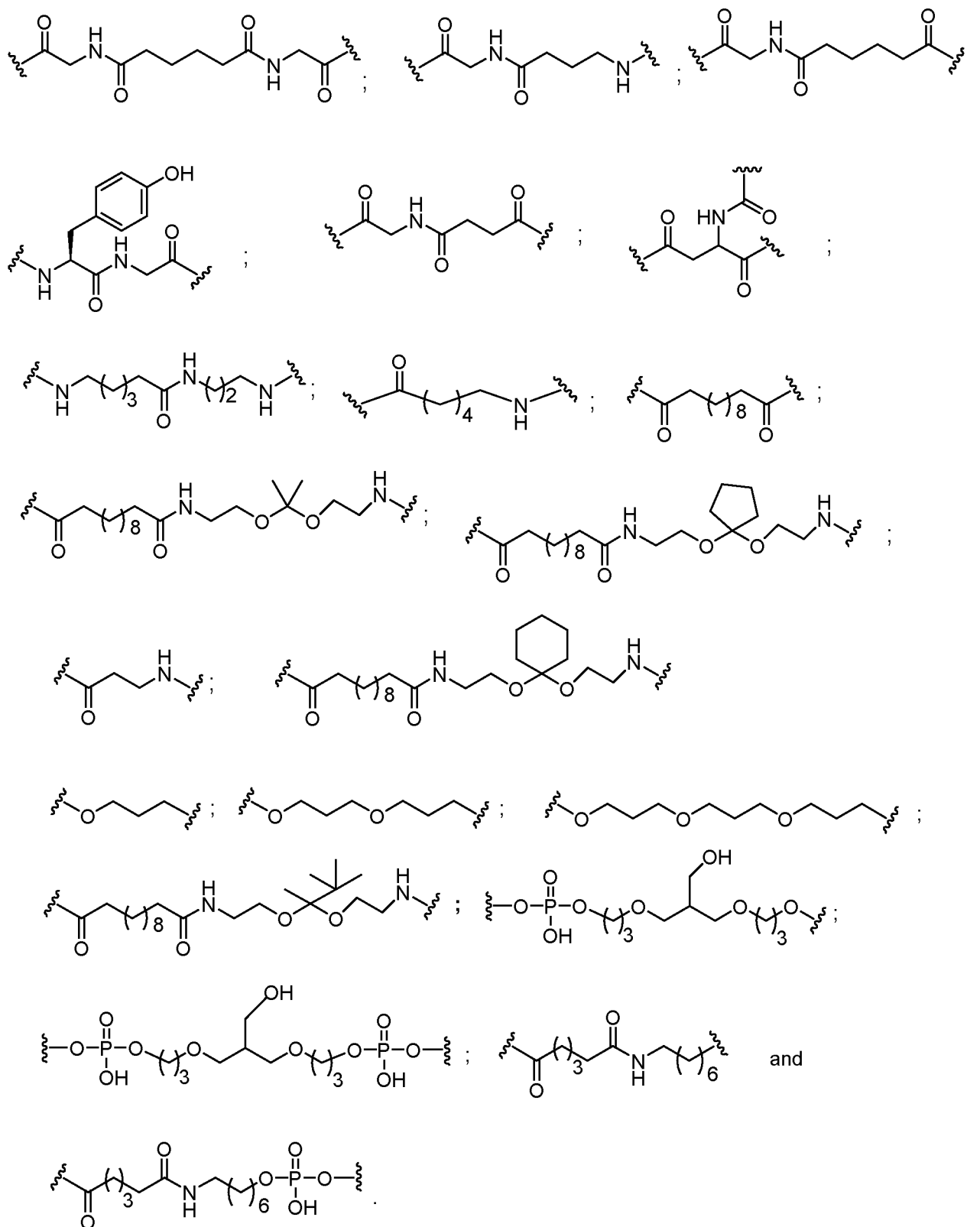




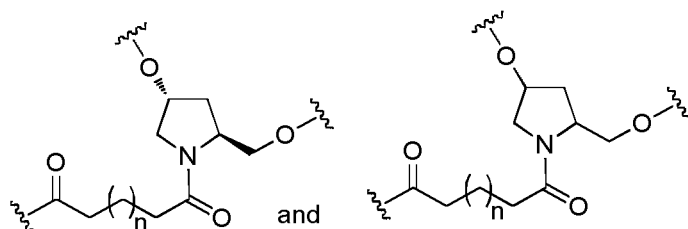
In certain embodiments, a linker has a structure selected from among:



In certain embodiments, a linker has a structure selected from among:



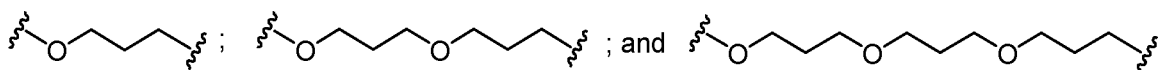
In certain embodiments, a linker has a structure selected from among:



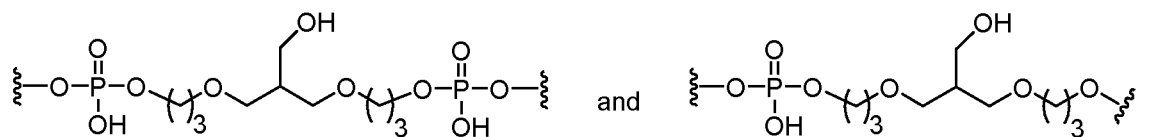
wherein n is from 1 to 20.

5

In certain embodiments, a linker has a structure selected from among:

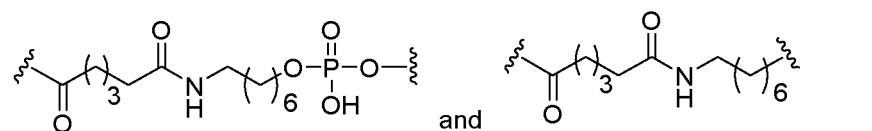


In certain embodiments, a linker has a structure selected from among:



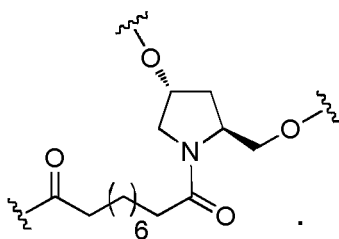
10

In certain embodiments, a linker has a structure selected from among:

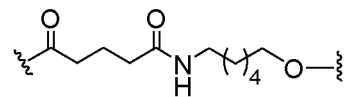


15

In certain embodiments, the conjugate linker has the structure:

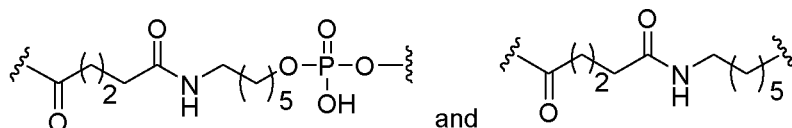


In certain embodiments, the conjugate linker has the structure:

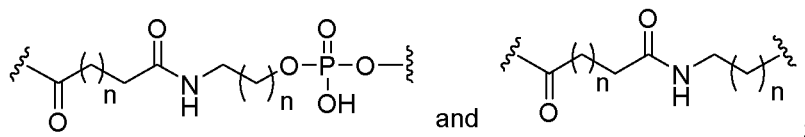


In certain embodiments, a linker has a structure selected from among:

20



In certain embodiments, a linker has a structure selected from among:



wherein each n is independently, 0, 1, 2, 3, 4, 5, 6, or 7.

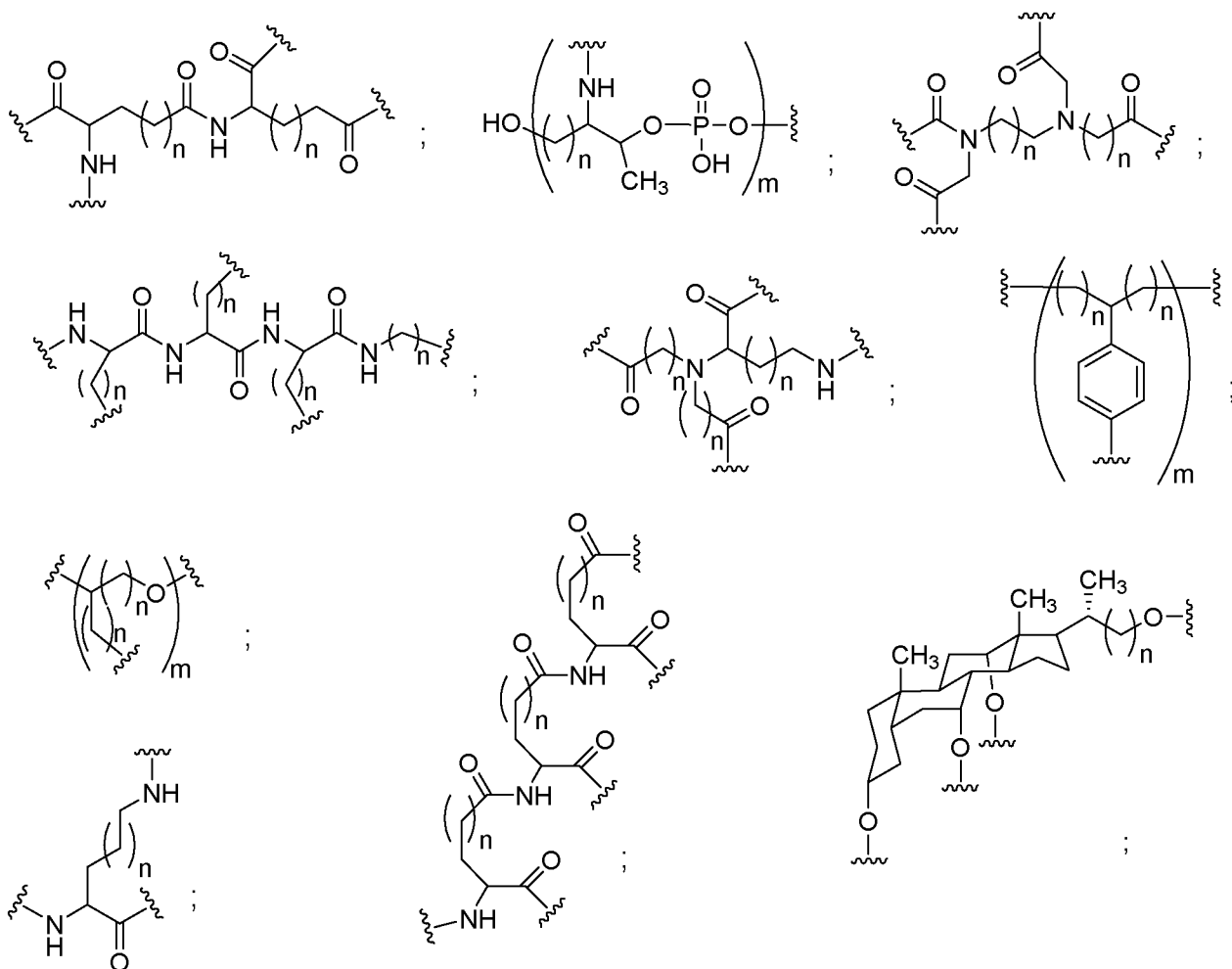
iv. Certain Cell-Targeting Moieties

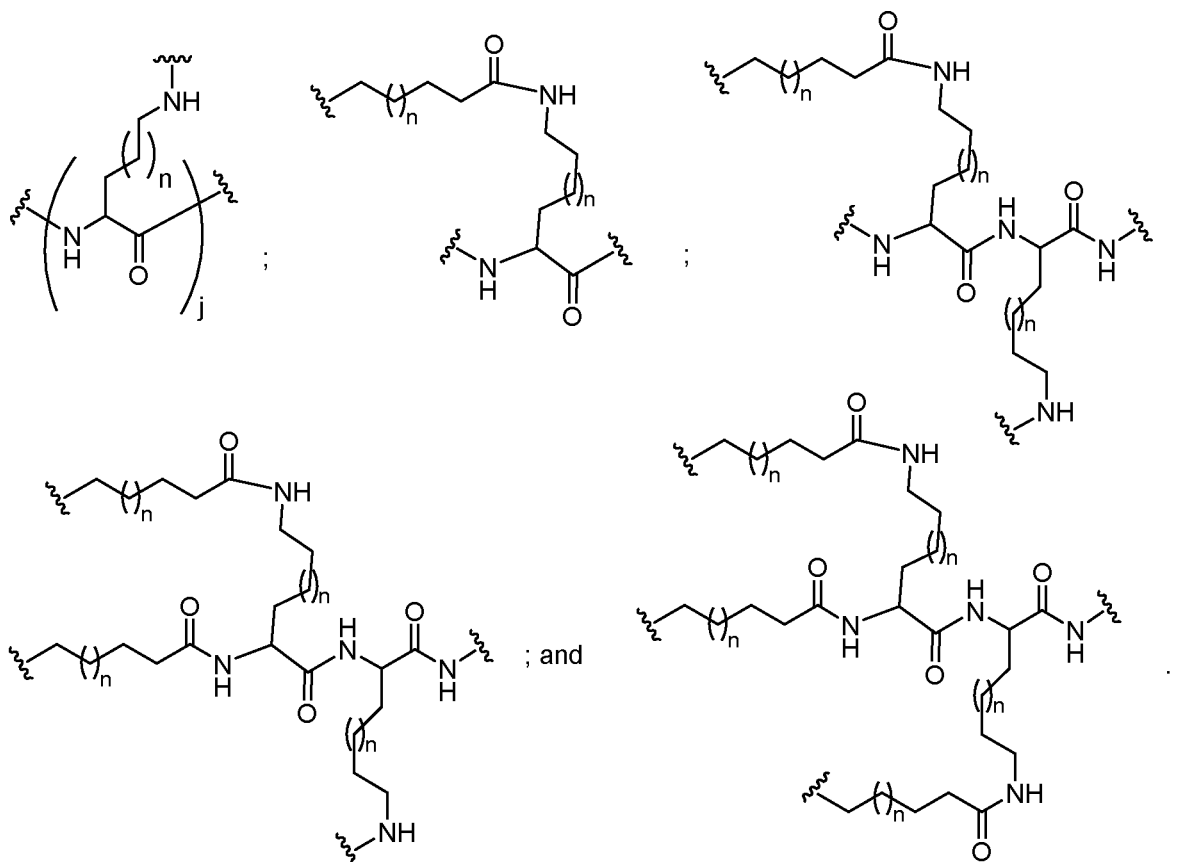
In certain embodiments, conjugate groups comprise cell-targeting moieties. Certain such cell-targeting moieties increase cellular uptake of antisense compounds. In certain embodiments, cell-targeting moieties comprise a branching group, one or more tether, and one or more ligand. In certain embodiments, cell-targeting moieties comprise a branching group, one or more tether, one or more ligand and one or more cleavable bond.

1. Certain Branching Groups

In certain embodiments, the conjugate groups comprise a targeting moiety comprising a branching group and at least two tethered ligands. In certain embodiments, the branching group attaches the conjugate linker. In certain embodiments, the branching group attaches the cleavable moiety. In certain embodiments, the branching group attaches the antisense oligonucleotide. In certain embodiments, the branching group is covalently attached to the linker and each of the tethered ligands. In certain embodiments, the branching group comprises a branched aliphatic group comprising groups selected from alkyl, amide, disulfide, polyethylene glycol, ether, thioether and hydroxylamino groups. In certain embodiments, the branching group comprises groups selected from alkyl, amide and ether groups. In certain embodiments, the branching group comprises groups selected from alkyl and ether groups. In certain embodiments, the branching group comprises a mono or polycyclic ring system. In certain embodiments, the branching group comprises one or more cleavable bond. In certain embodiments, the conjugate group does not include a branching group.

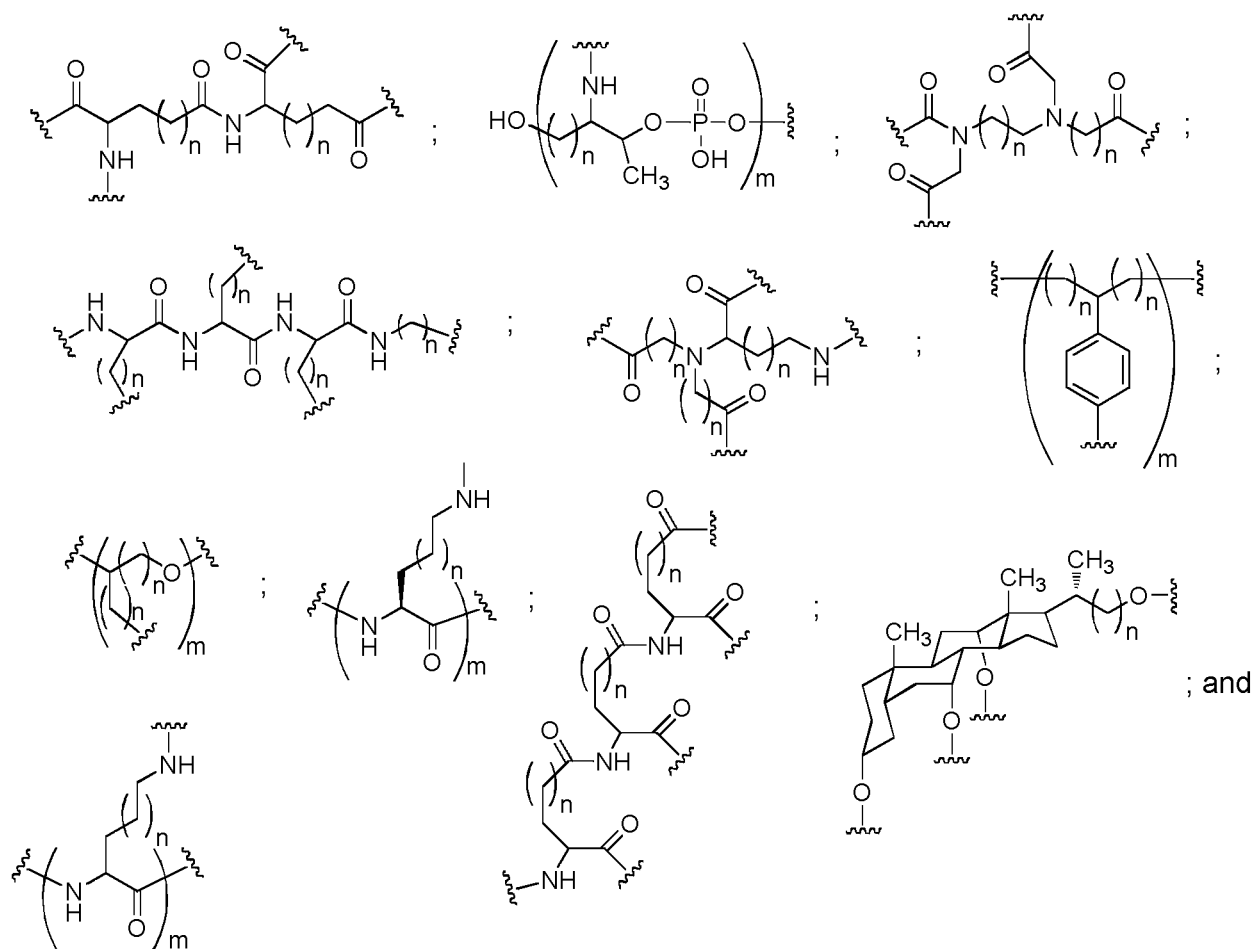
In certain embodiments, a branching group has a structure selected from among:





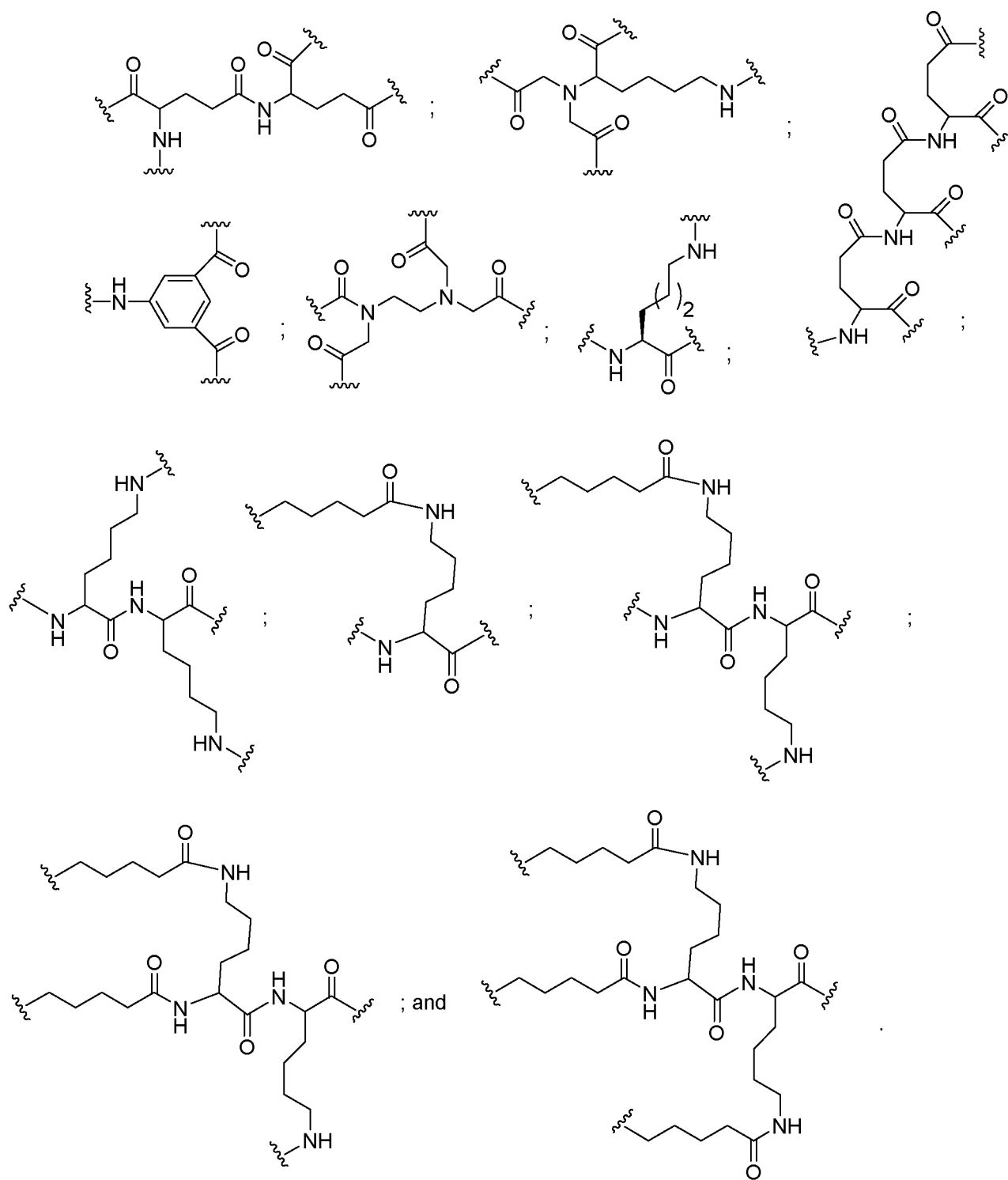
wherein each n is, independently, from 1 to 20;
 j is from 1 to 3; and
 m is from 2 to 6.

In certain embodiments, a branching group has a structure selected from among:

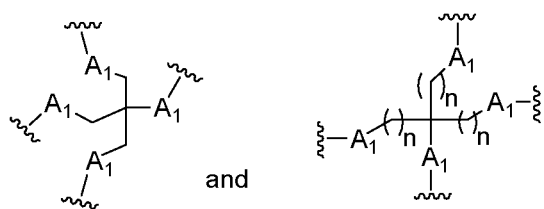


wherein each n is, independently, from 1 to 20; and m is from 2 to 6.

In certain embodiments, a branching group has a structure selected from among:

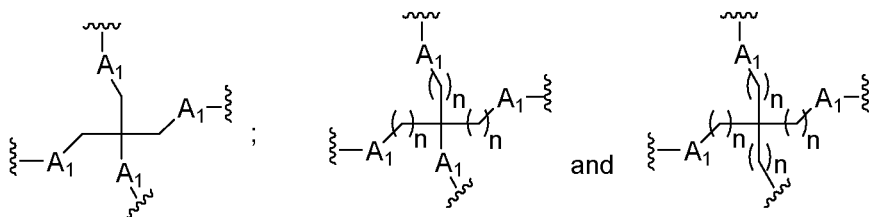


In certain embodiments, a branching group has a structure selected from among:



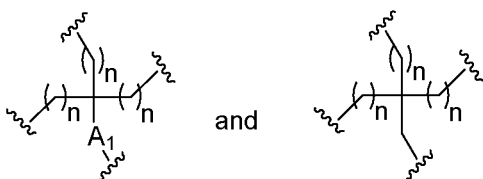
wherein each A_1 is independently, O, S, C=O or NH; and
each n is, independently, from 1 to 20.

- 5 In certain embodiments, a branching group has a structure selected from among:



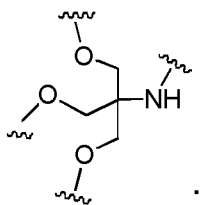
wherein each A_1 is independently, O, S, C=O or NH; and
each n is, independently, from 1 to 20.

- 10 In certain embodiments, a branching group has a structure selected from among:

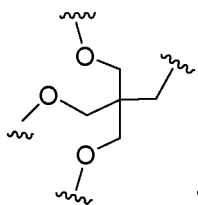


wherein A_1 is O, S, C=O or NH; and
each n is, independently, from 1 to 20.

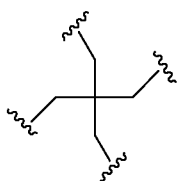
- 15 In certain embodiments, a branching group has a structure selected from among:



In certain embodiments, a branching group has a structure selected from among:



In certain embodiments, a branching group has a structure selected from among:



5

2. Certain Tethers

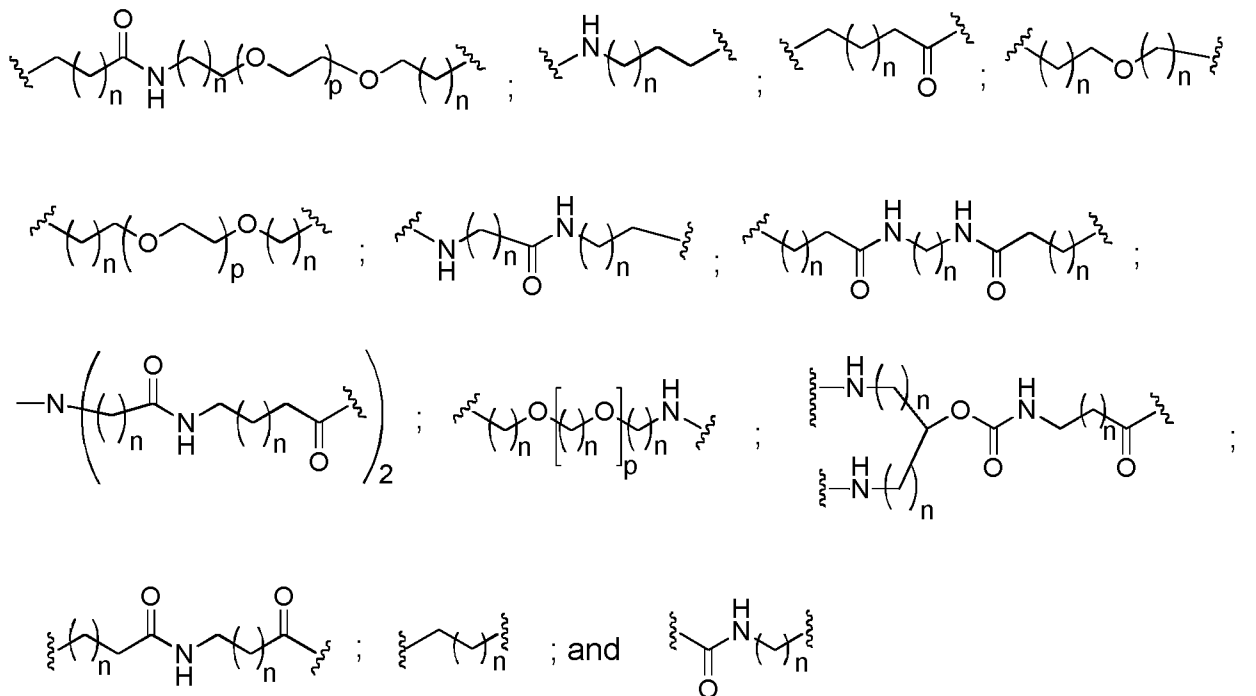
In certain embodiments, conjugate groups comprise one or more tethers covalently attached to the branching group. In certain embodiments, conjugate groups comprise one or more tethers covalently attached to the linking group. In certain embodiments, each tether is a linear aliphatic group comprising one or more groups selected from alkyl, ether, thioether, disulfide, amide and polyethylene glycol groups in any combination. In certain embodiments, each tether is a linear aliphatic group comprising one or more groups selected from alkyl, substituted alkyl, ether, thioether, disulfide, amide, phosphodiester and polyethylene glycol groups in any combination. In certain embodiments, each tether is a linear aliphatic group comprising one or more groups selected from alkyl, ether and amide groups in any combination. In certain embodiments, each tether is a linear aliphatic group comprising one or more groups selected from alkyl, substituted alkyl, phosphodiester, ether and amide groups in any combination. In certain embodiments, each tether is a linear aliphatic group comprising one or more groups selected from alkyl and phosphodiester in any combination. In certain embodiments, each tether comprises at least one phosphorus linking group or neutral linking group.

In certain embodiments, the tether includes one or more cleavable bond. In certain embodiments, the tether is attached to the branching group through either an amide or an ether group. In certain embodiments, the tether is attached to the branching group through a phosphodiester group. In certain embodiments, the tether is attached to the branching group through a phosphorus linking group or neutral linking group. In certain embodiments, the tether is attached to the branching group through an ether group. In certain embodiments, the tether is attached to the ligand through either an amide or an ether group. In certain embodiments, the tether is attached to the ligand through an ether group. In certain embodiments, the tether is attached to the ligand through either an amide or an ether group. In certain embodiments, the tether is attached to the ligand through an ether group.

In certain embodiments, each tether comprises from about 8 to about 20 atoms in chain length between the ligand and the branching group. In certain embodiments, each tether group comprises from

about 10 to about 18 atoms in chain length between the ligand and the branching group. In certain embodiments, each tether group comprises about 13 atoms in chain length.

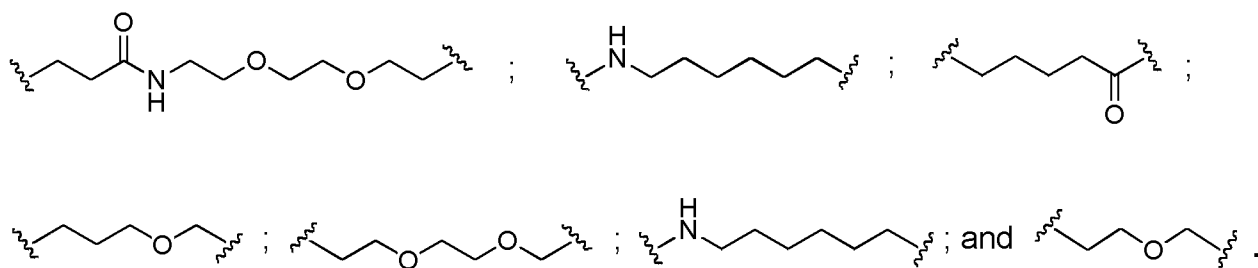
In certain embodiments, a tether has a structure selected from among:



5

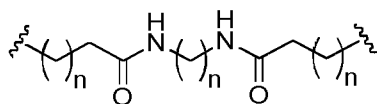
wherein each n is, independently, from 1 to 20; and each p is from 1 to about 6.

In certain embodiments, a tether has a structure selected from among:



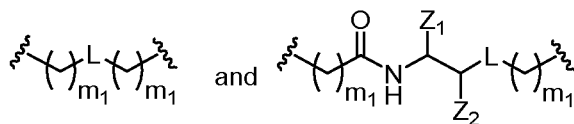
10

In certain embodiments, a tether has a structure selected from among:



wherein each n is, independently, from 1 to 20.

In certain embodiments, a tether has a structure selected from among:



wherein L is either a phosphorus linking group or a neutral linking group;

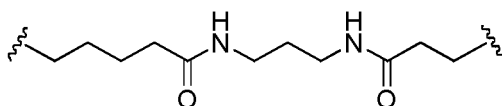
Z_1 is $C(=O)O-R_2$;

Z_2 is H, C_1-C_6 alkyl or substituted C_1-C_6 alkyl;

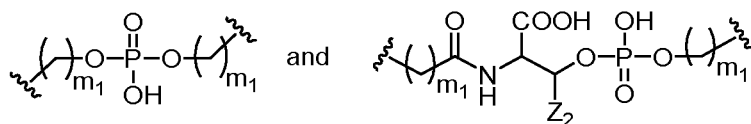
R_2 is H, C_1-C_6 alkyl or substituted C_1-C_6 alkyl; and

each m_1 is, independently, from 0 to 20 wherein at least one m_1 is greater than 0 for each tether.

In certain embodiments, a tether has a structure selected from among:



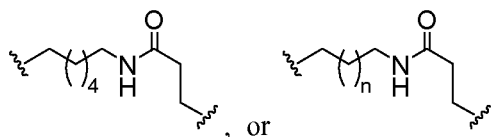
In certain embodiments, a tether has a structure selected from among:



wherein Z_2 is H or CH_3 ; and

each m_1 is, independently, from 0 to 20 wherein at least one m_1 is greater than 0 for each tether.

In certain embodiments, a tether has a structure selected from among:



; wherein each n is independently, 0, 1, 2, 3, 4, 5, 6, or 7.

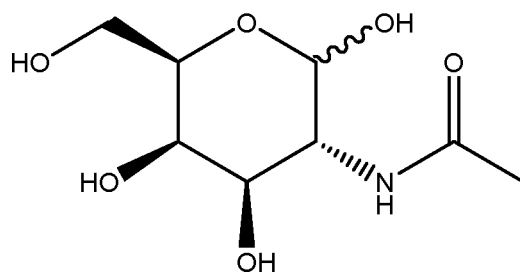
In certain embodiments, a tether comprises a phosphorus linking group. In certain embodiments, a tether does not comprise any amide bonds. In certain embodiments, a tether comprises a phosphorus linking group and does not comprise any amide bonds.

3. Certain Ligands

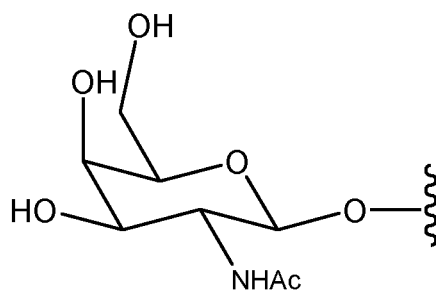
In certain embodiments, the present disclosure provides ligands wherein each ligand is covalently attached to a tether. In certain embodiments, each ligand is selected to have an affinity for at least one type of receptor on a target cell. In certain embodiments, ligands are selected that have an affinity for at least one type of receptor on the surface of a mammalian liver cell. In certain embodiments, ligands are selected that have an affinity for the hepatic asialoglycoprotein receptor (ASGP-R). In certain embodiments, each ligand is a carbohydrate. In certain embodiments, each ligand is, independently selected from galactose, N-acetyl galactoseamine, mannose, glucose, glucosamine and fucose. In certain embodiments, each ligand is N-acetyl galactoseamine (GalNAc). In certain embodiments, the targeting moiety comprises 2 to 6 ligands. In certain embodiments, the targeting moiety comprises 3 ligands. In certain embodiments, the targeting moiety comprises 3 N-acetyl galactoseamine ligands.

In certain embodiments, the ligand is a carbohydrate, carbohydrate derivative, modified carbohydrate, multivalent carbohydrate cluster, polysaccharide, modified polysaccharide, or polysaccharide derivative. In certain embodiments, the ligand is an amino sugar or a thio sugar. For example, amino sugars may be selected from any number of compounds known in the art, for example glucosamine, sialic acid, α -D-galactosamine, N-Acetylgalactosamine, 2-acetamido-2-deoxy-D-galactopyranose (GalNAc), 2-Amino-3-*O*-[(*R*)-1-carboxyethyl]-2-deoxy- β -D-glucopyranose (β -muramic acid), 2-Deoxy-2-methylamino-L-glucopyranose, 4,6-Dideoxy-4-formamido-2,3-di-*O*-methyl-D-mannopyranose, 2-Deoxy-2-sulfoamino-D-glucopyranose and *N*-sulfo-D-glucosamine, and *N*-Glycoloyl- α -neuraminic acid. For example, thio sugars may be selected from the group consisting of 5-Thio- β -D-glucopyranose, Methyl 2,3,4-tri-*O*-acetyl-1-thio-6-*O*-trityl- α -D-glucopyranoside, 4-Thio- β -D-galactopyranose, and ethyl 3,4,6,7-tetra-*O*-acetyl-2-deoxy-1,5-dithio- α -D-*gluco*-heptopyranoside.

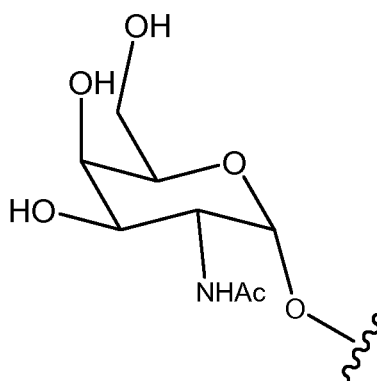
In certain embodiments, “GalNAc” or “Gal-NAc” refers to 2-(Acetylamino)-2-deoxy-D-galactopyranose, commonly referred to in the literature as N-acetyl galactosamine. In certain embodiments, “N-acetyl galactosamine” refers to 2-(Acetylamino)-2-deoxy-D-galactopyranose. In certain embodiments, “GalNAc” or “Gal-NAc” refers to 2-(Acetylamino)-2-deoxy-D-galactopyranose. In certain embodiments, “GalNAc” or “Gal-NAc” refers to 2-(Acetylamino)-2-deoxy-D-galactopyranose, which includes both the β -form: 2-(Acetylamino)-2-deoxy- β -D-galactopyranose and α -form: 2-(Acetylamino)-2-deoxy-D-galactopyranose. In certain embodiments, both the β -form: 2-(Acetylamino)-2-deoxy- β -D-galactopyranose and α -form: 2-(Acetylamino)-2-deoxy-D-galactopyranose may be used interchangeably. Accordingly, in structures in which one form is depicted, these structures are intended to include the other form as well. For example, where the structure for an α -form: 2-(Acetylamino)-2-deoxy-D-galactopyranose is shown, this structure is intended to include the other form as well. In certain embodiments, In certain preferred embodiments, the β -form 2-(Acetylamino)-2-deoxy-D-galactopyranose is the preferred embodiment.



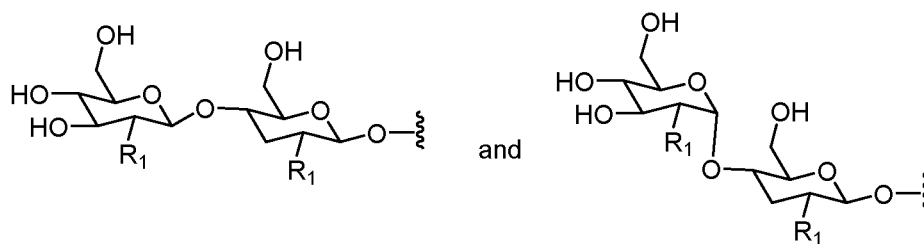
2-(Acetylamino)-2-deoxy-D-galactopyranose

2-(Acetylamino)-2-deoxy- β -D-galactopyranose

5

2-(Acetylamino)-2-deoxy- α -D-galactopyranose

In certain embodiments one or more ligand has a structure selected from among:

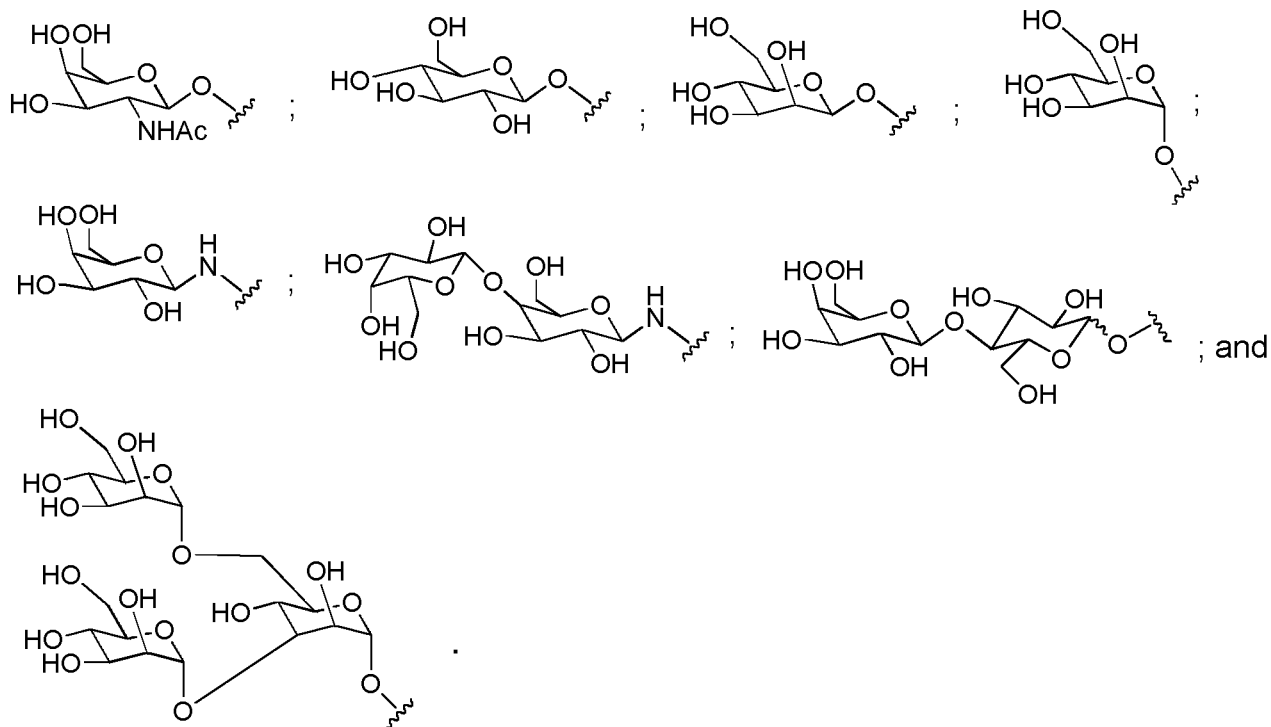


and

10

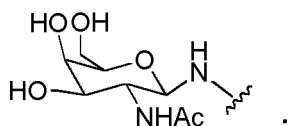
wherein each R_1 is selected from OH and $NHCOOH$.

In certain embodiments one or more ligand has a structure selected from among:



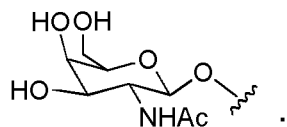
5

In certain embodiments one or more ligand has a structure selected from among:



10

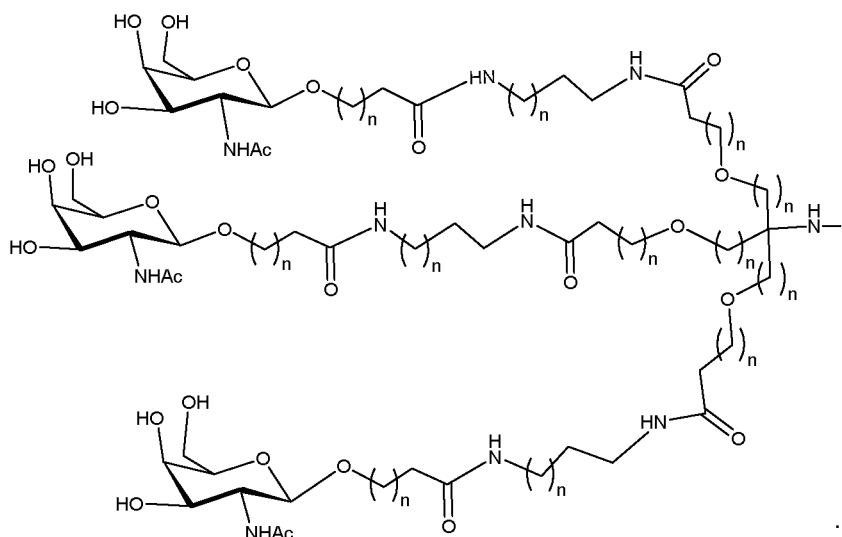
In certain embodiments one or more ligand has a structure selected from among:



i. Certain Conjugates

In certain embodiments, conjugate groups comprise the structural features above. In certain such embodiments, conjugate groups have the following structure:

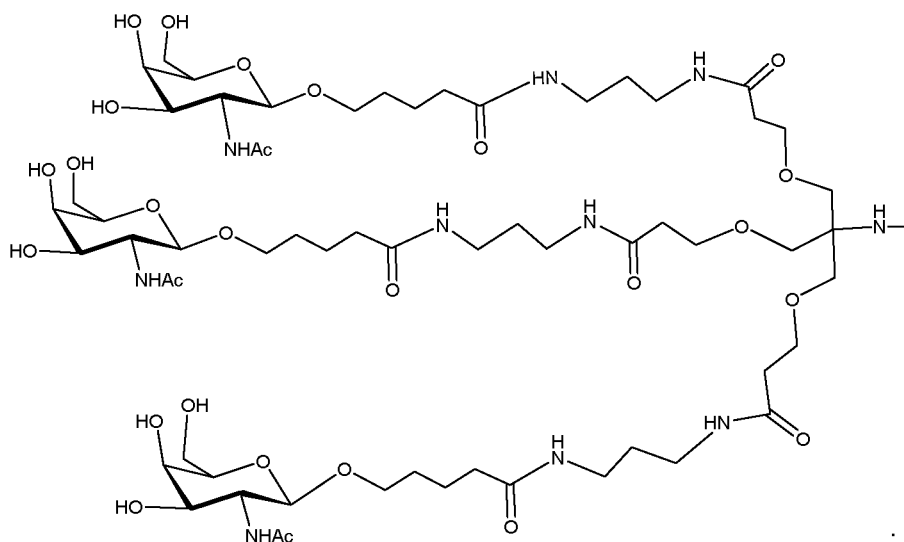
5



wherein each n is, independently, from 1 to 20.

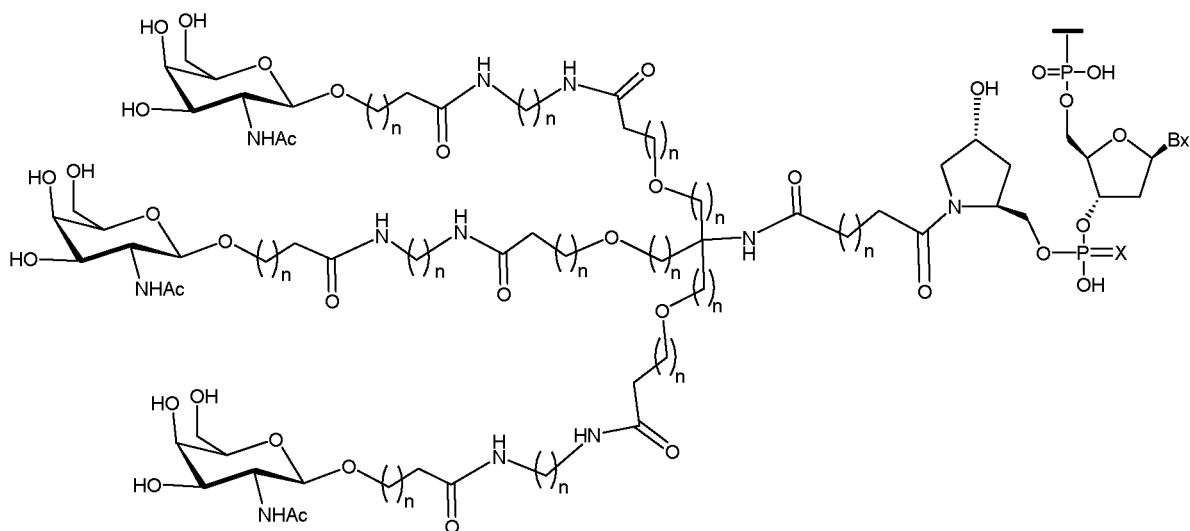
10

In certain such embodiments, conjugate groups have the following structure:



15

In certain such embodiments, conjugate groups have the following structure:



wherein each n is, independently, from 1 to 20;

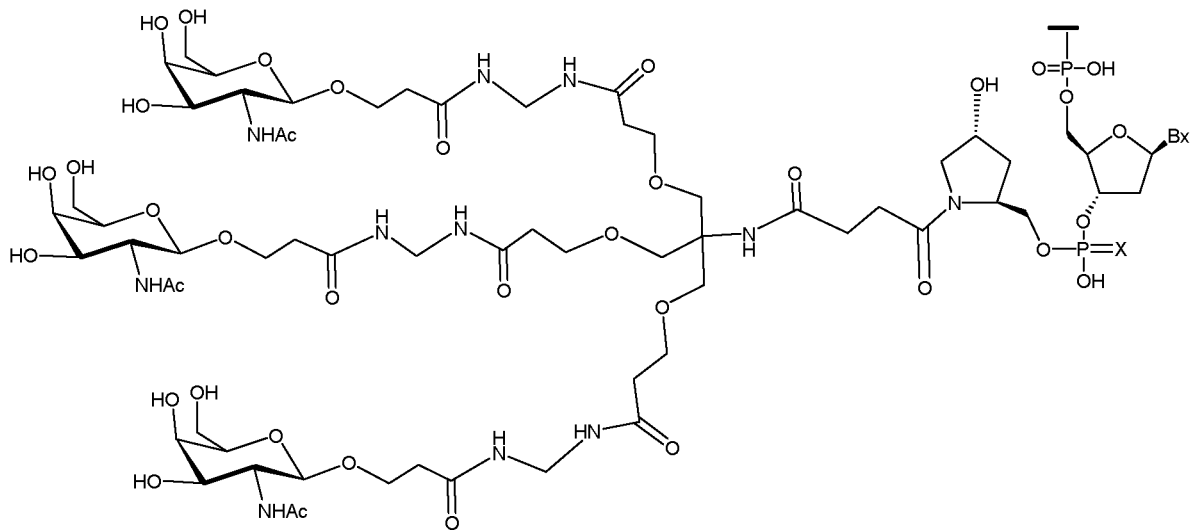
Z is H or a linked solid support;

Q is an antisense compound;

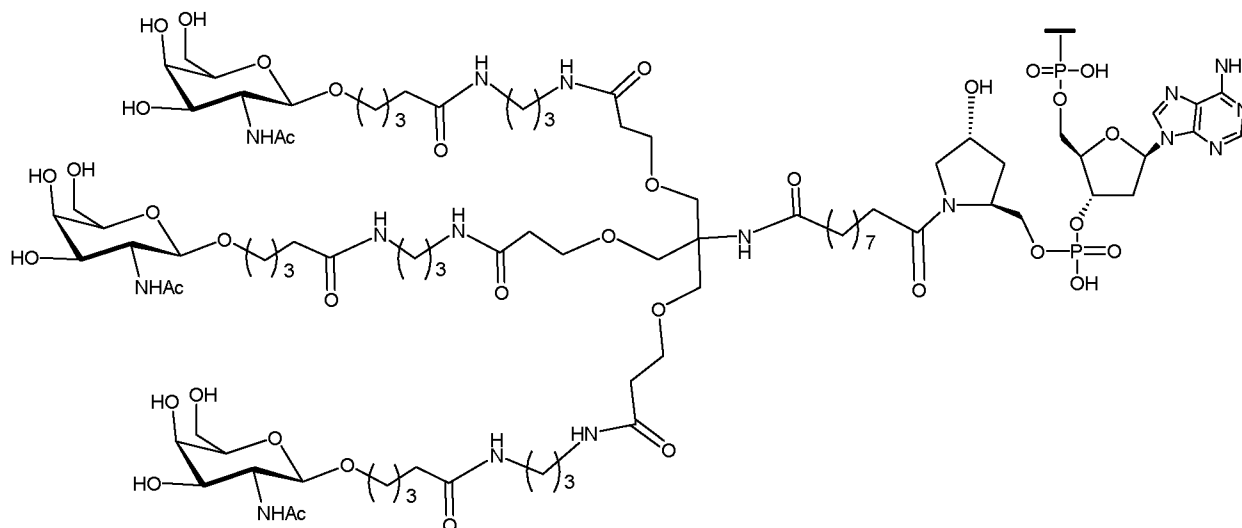
X is O or S; and

Bx is a heterocyclic base moiety.

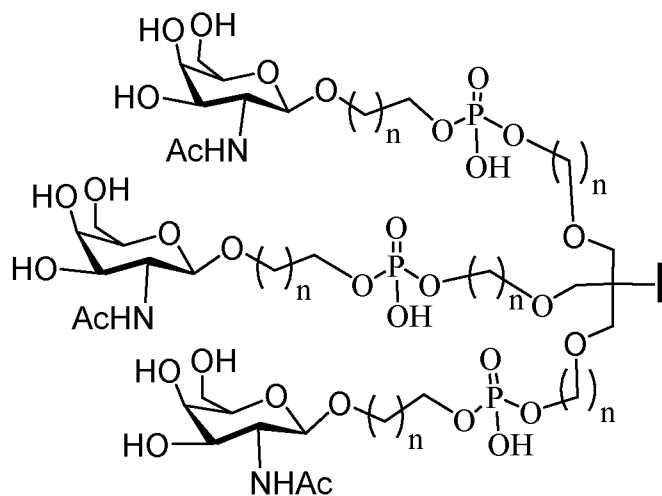
In certain such embodiments, conjugate groups have the following structure:



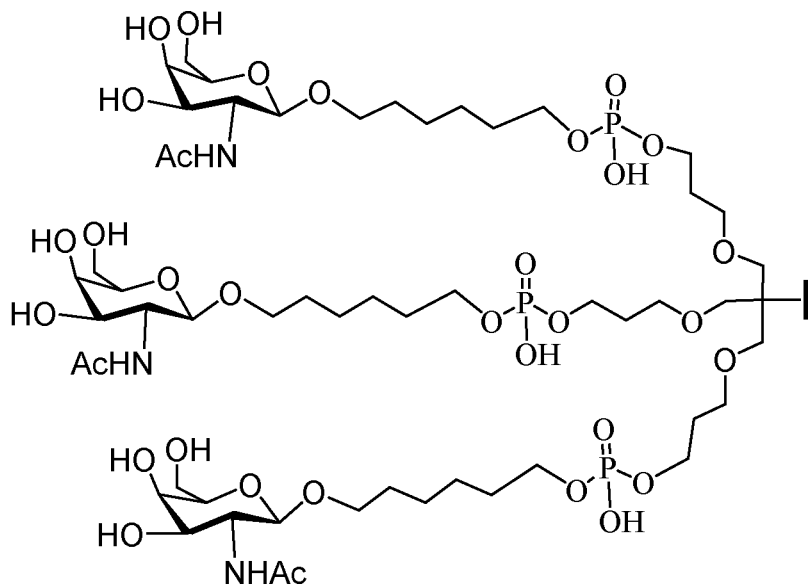
In certain such embodiments, conjugate groups have the following structure:



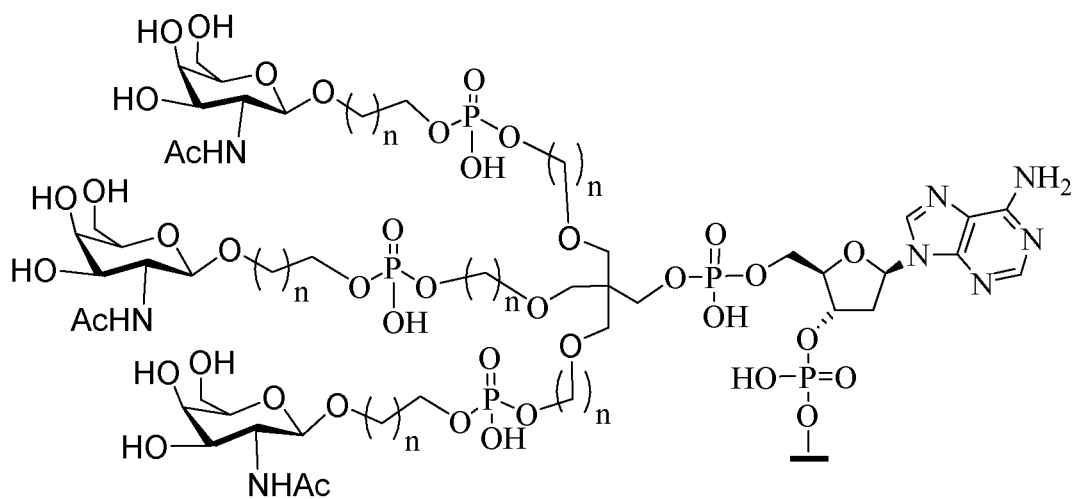
In certain such embodiments, conjugate groups have the following structure:



In certain such embodiments, conjugate groups have the following structure:

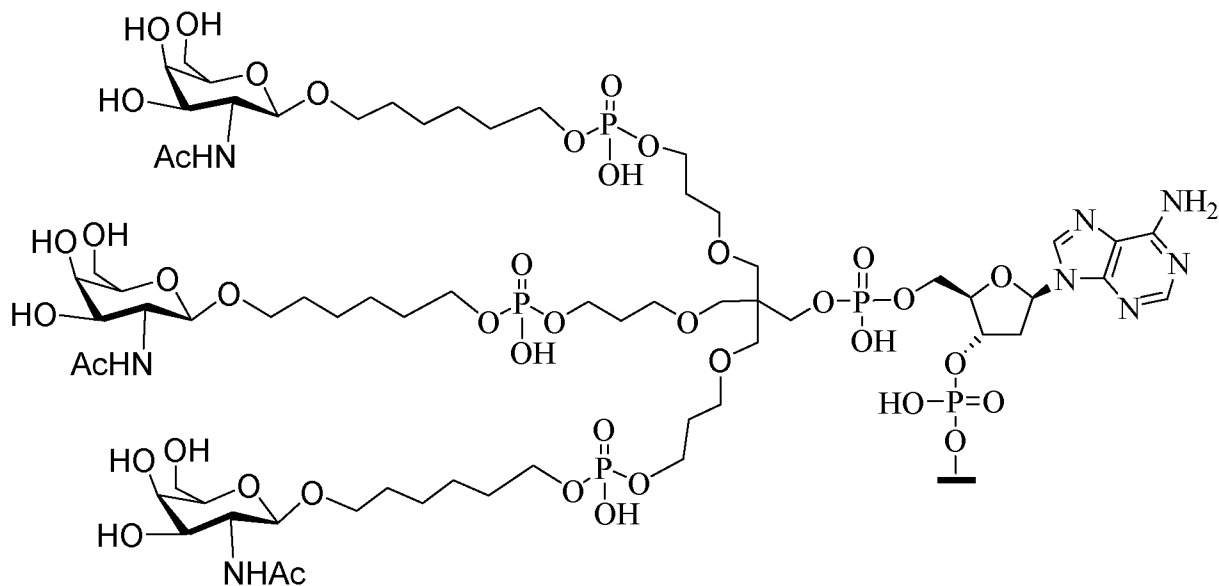


In certain such embodiments, conjugate groups have the following structure:



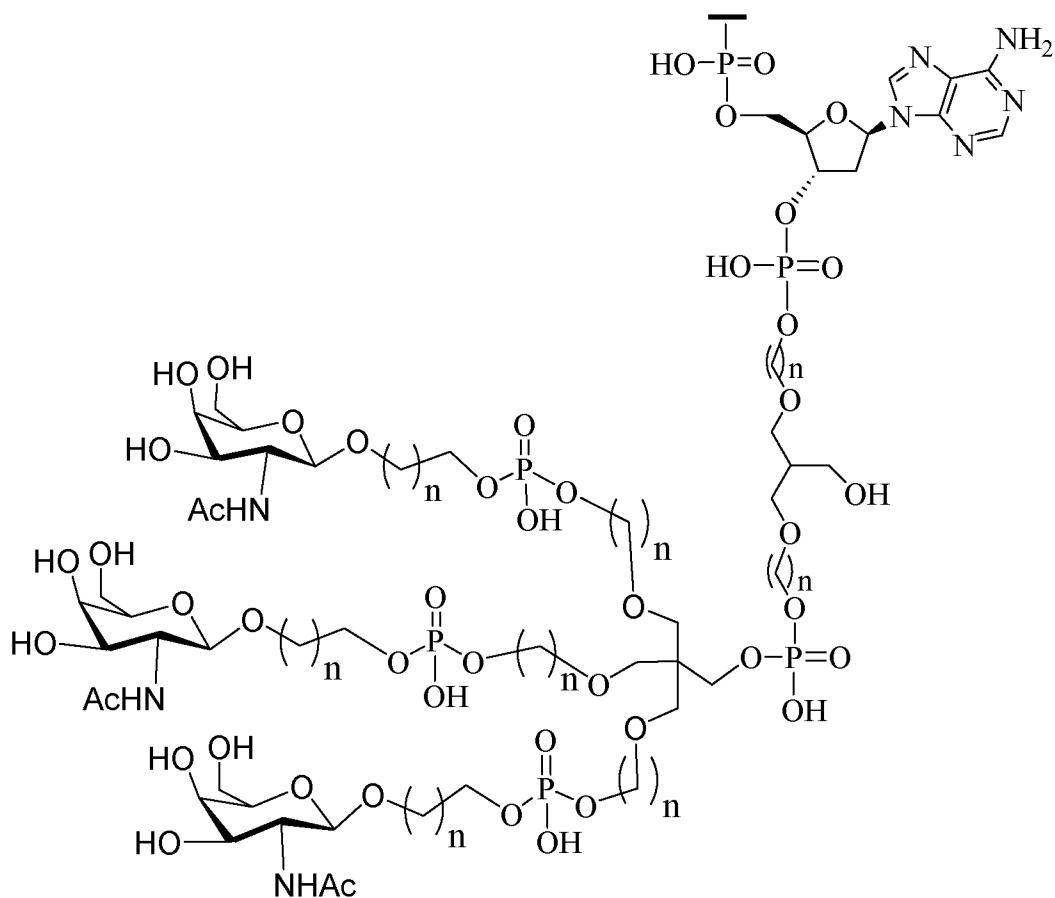
5

In certain such embodiments, conjugate groups have the following structure:

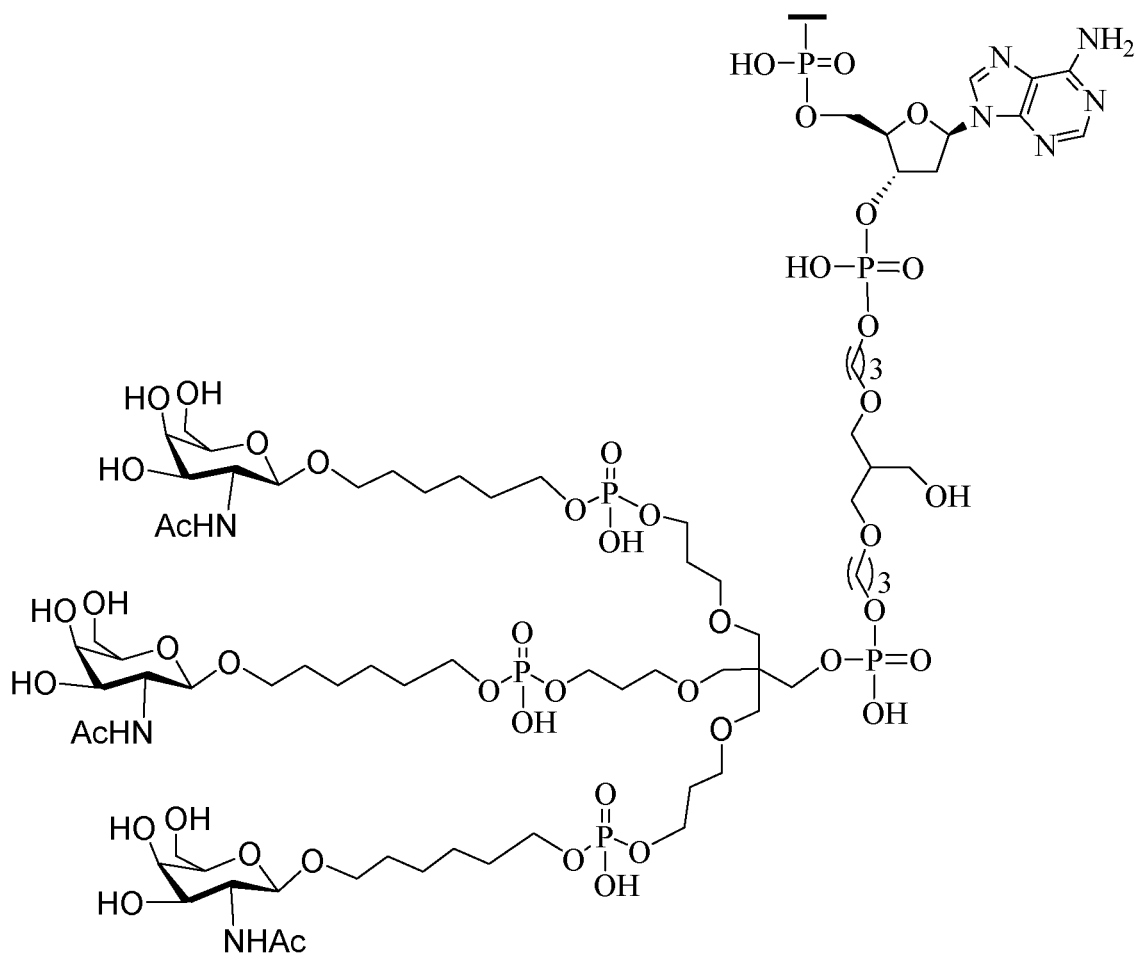


In certain such embodiments, conjugate groups have the following structure:

5

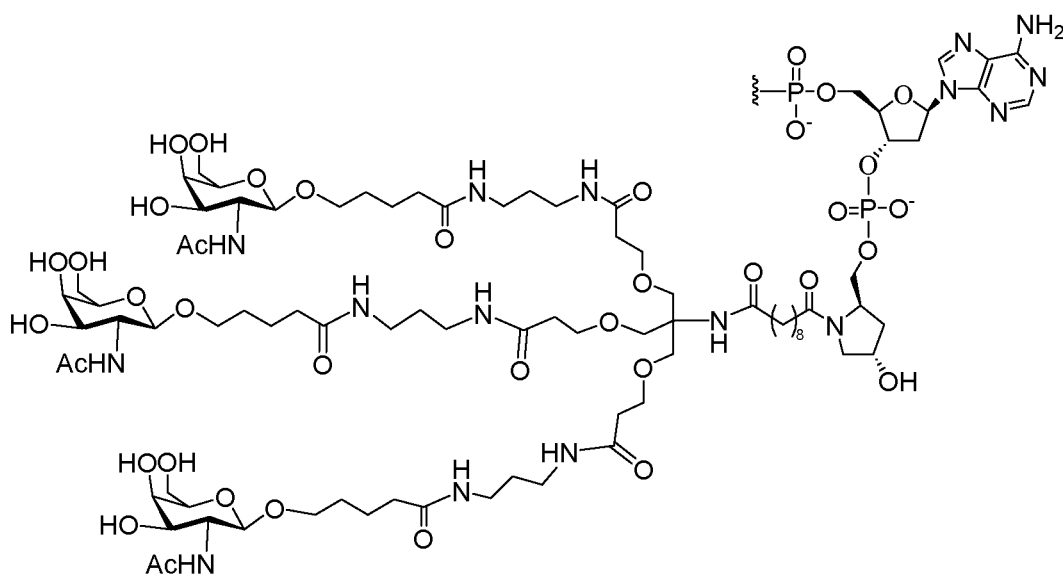


In certain such embodiments, conjugate groups have the following structure:

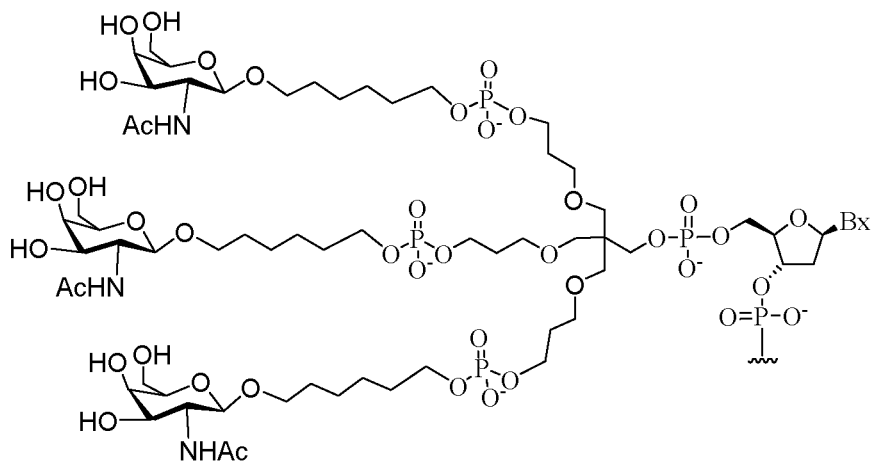


In certain embodiments, conjugates do not comprise a pyrrolidine.

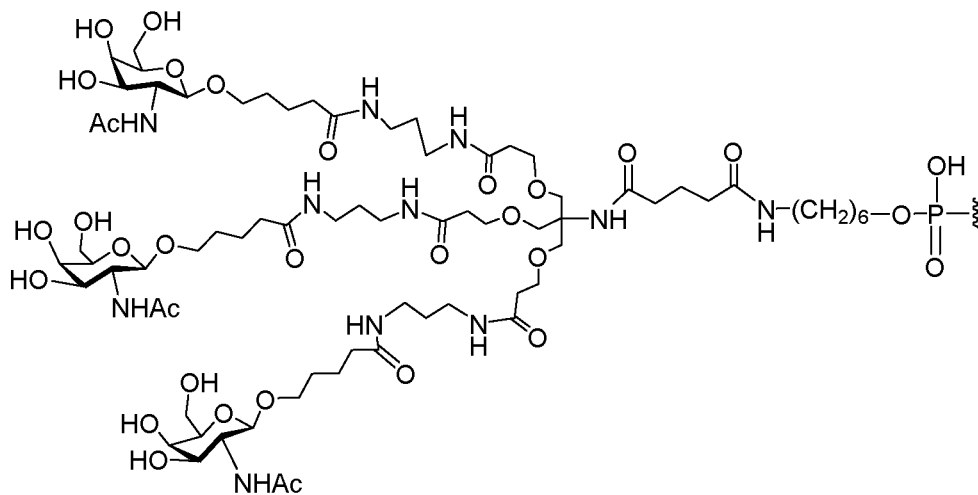
5 In certain such embodiments, conjugate groups have the following structure:



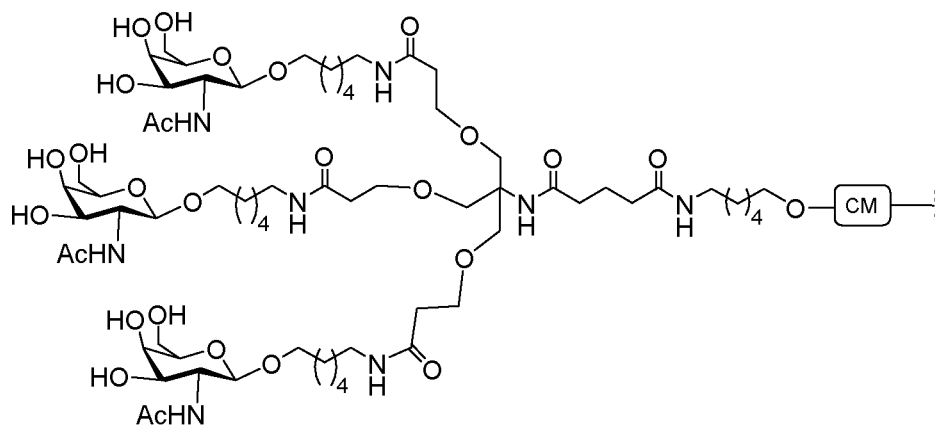
In certain such embodiments, conjugate groups have the following structure:



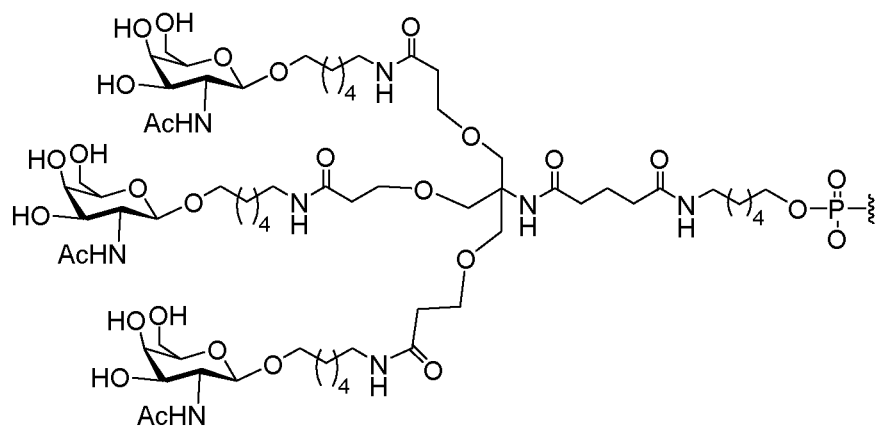
In certain such embodiments, conjugate groups have the following structure:



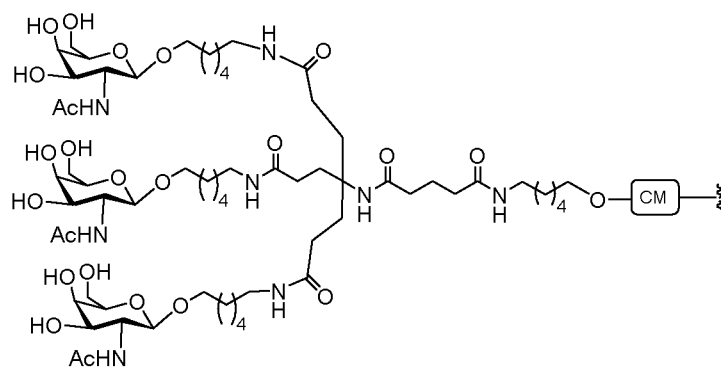
In certain such embodiments, conjugate groups have the following structure:



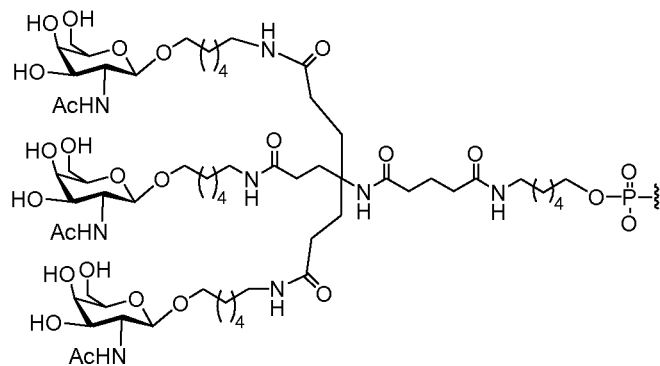
In certain such embodiments, conjugate groups have the following structure:



In certain such embodiments, conjugate groups have the following structure:

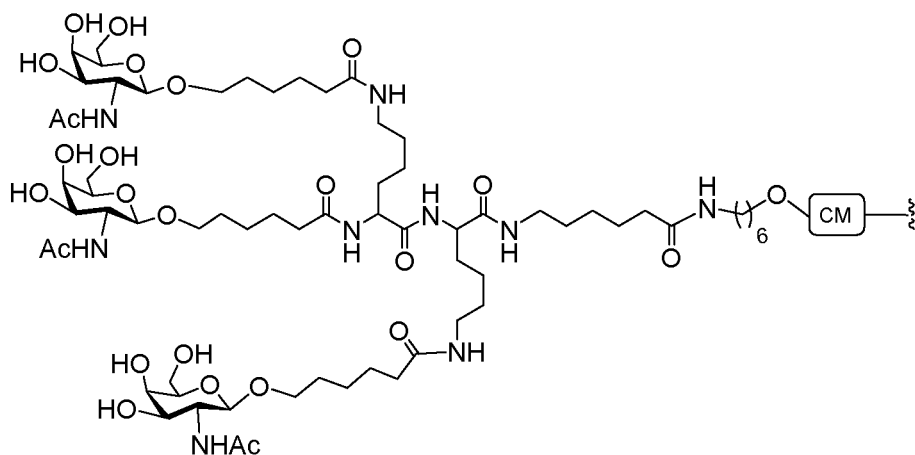


In certain such embodiments, conjugate groups have the following structure:

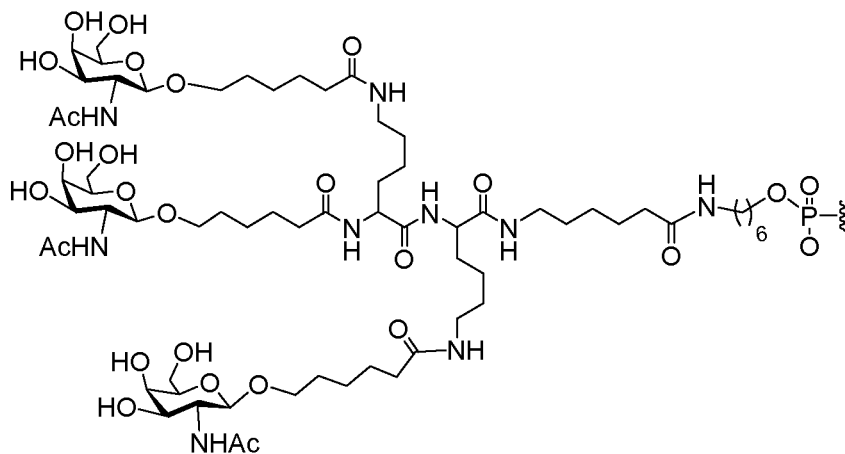


5

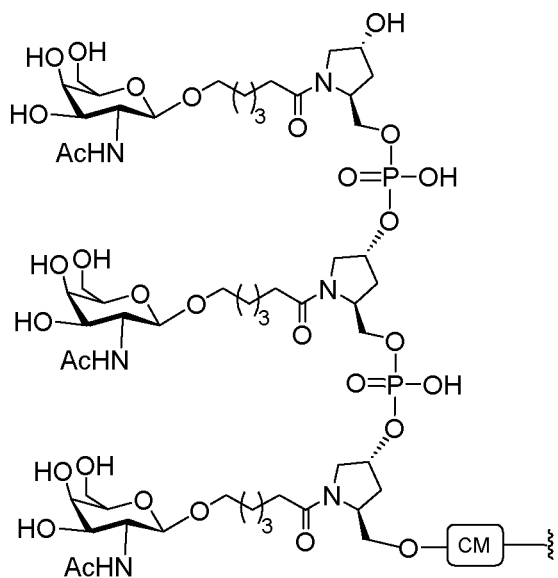
In certain such embodiments, conjugate groups have the following structure:



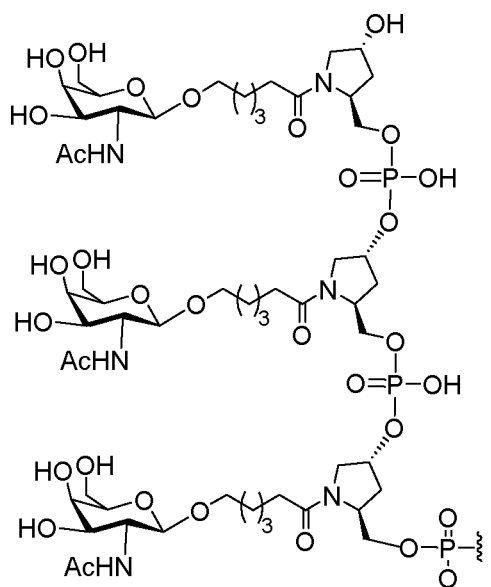
In certain such embodiments, conjugate groups have the following structure:



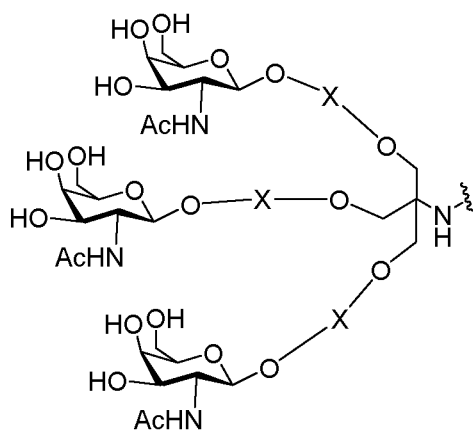
In certain such embodiments, conjugate groups have the following structure:



In certain such embodiments, conjugate groups have the following structure:

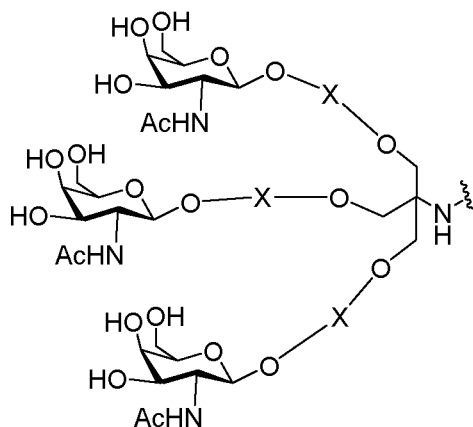


In certain embodiments, the cell-targeting moiety of the conjugate group has the following structure:



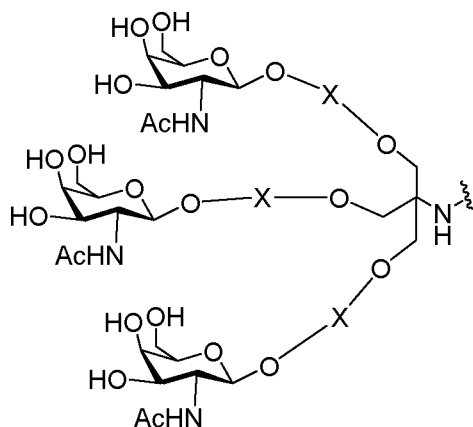
wherein X is a substituted or unsubstituted tether of six to eleven consecutively bonded atoms.

- 5 In certain embodiments, the cell-targeting moiety of the conjugate group has the following structure:



wherein X is a substituted or unsubstituted tether of ten consecutively bonded atoms.

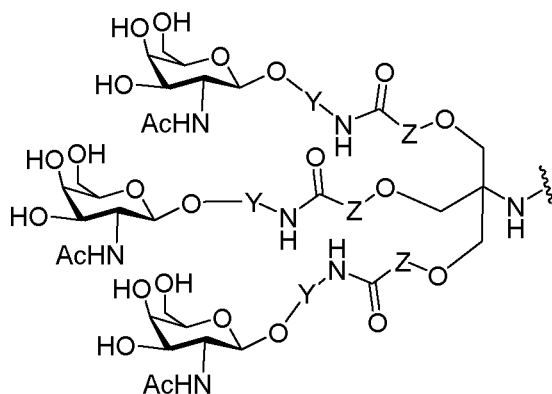
In certain embodiments, the cell-targeting moiety of the conjugate group has the following structure:



wherein X is a substituted or unsubstituted tether of four to eleven consecutively bonded atoms and wherein the tether comprises exactly one amide bond.

5

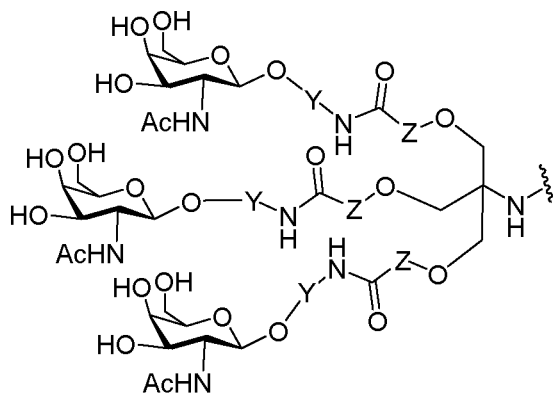
In certain embodiments, the cell-targeting moiety of the conjugate group has the following structure:



wherein Y and Z are independently selected from a C₁-C₁₂ substituted or unsubstituted alkyl, alkenyl, or alkynyl group, or a group comprising an ether, a ketone, an amide, an ester, a carbamate, an amine, a piperidine, a phosphate, a phosphodiester, a phosphorothioate, a triazole, a pyrrolidine, a disulfide, or a thioether.

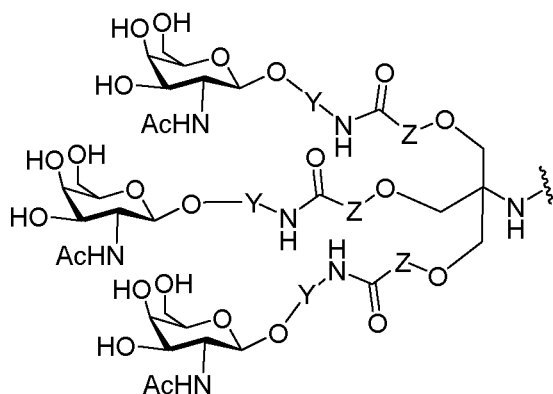
10

In certain such embodiments, the cell-targeting moiety of the conjugate group has the following structure:



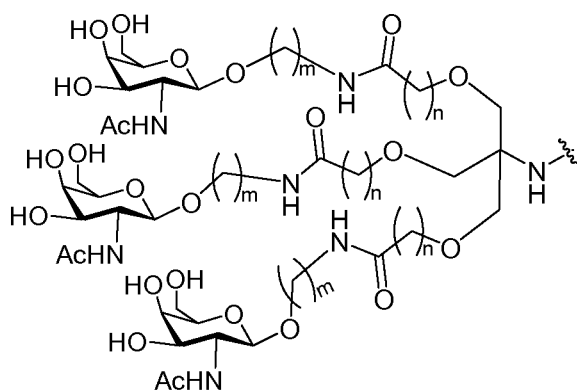
wherein Y and Z are independently selected from a C_1 - C_{12} substituted or unsubstituted alkyl group, or a group comprising exactly one ether or exactly two ethers, an amide, an amine, a piperidine, a phosphate, a phosphodiester, or a phosphorothioate.

- 5 In certain such embodiments, the cell-targeting moiety of the conjugate group has the following structure:



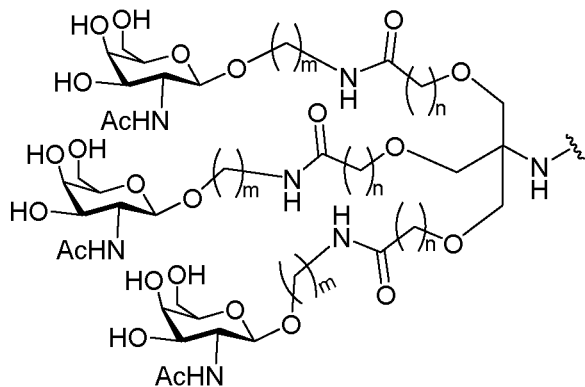
wherein Y and Z are independently selected from a C_1 - C_{12} substituted or unsubstituted alkyl group.

In certain such embodiments, the cell-targeting moiety of the conjugate group has the following structure:



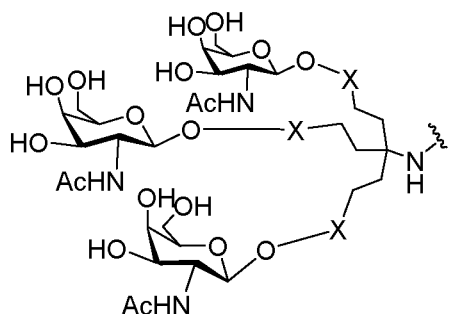
- 10 wherein m and n are independently selected from 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, and 12.

In certain such embodiments, the cell-targeting moiety of the conjugate group has the following structure:



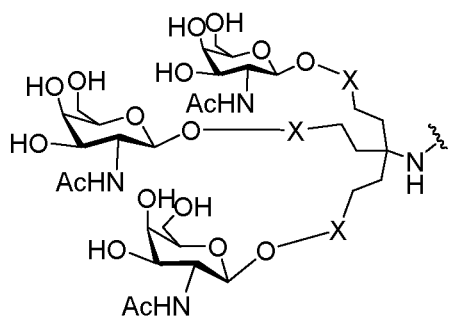
wherein m is 4, 5, 6, 7, or 8, and n is 1, 2, 3, or 4.

In certain embodiments, the cell-targeting moiety of the conjugate group has the following structure:



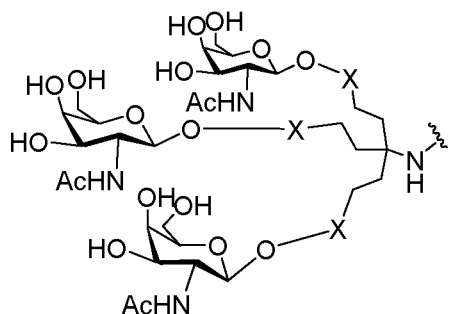
- 5 wherein X is a substituted or unsubstituted tether of four to thirteen consecutively bonded atoms, and wherein X does not comprise an ether group.

In certain embodiments, the cell-targeting moiety of the conjugate group has the following structure:



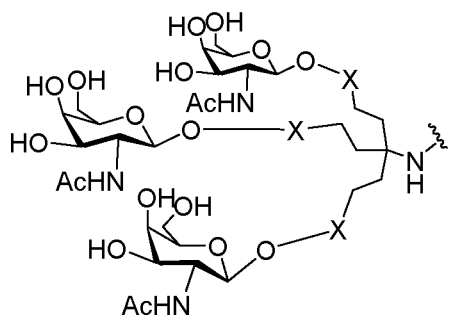
- 10 wherein X is a substituted or unsubstituted tether of eight consecutively bonded atoms, and wherein X does not comprise an ether group.

In certain embodiments, the cell-targeting moiety of the conjugate group has the following structure:



wherein X is a substituted or unsubstituted tether of four to thirteen consecutively bonded atoms, and wherein the tether comprises exactly one amide bond, and wherein X does not comprise an ether group.

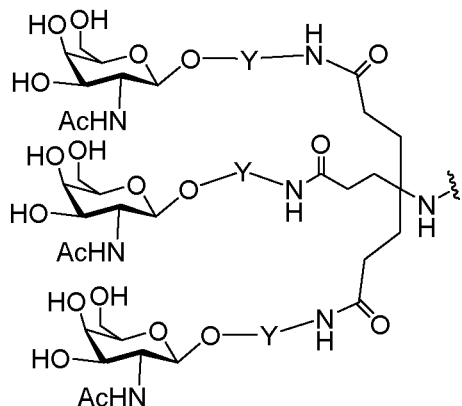
In certain embodiments, the cell-targeting moiety of the conjugate group has the following structure:



5

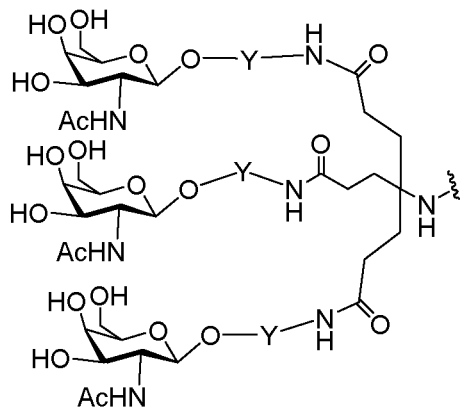
wherein X is a substituted or unsubstituted tether of four to thirteen consecutively bonded atoms and wherein the tether consists of an amide bond and a substituted or unsubstituted C_2 - C_{11} alkyl group.

In certain embodiments, the cell-targeting moiety of the conjugate group has the following structure:



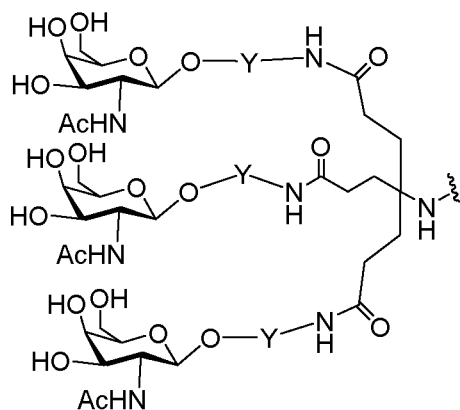
- 10 wherein Y is selected from a C_1 - C_{12} substituted or unsubstituted alkyl, alkenyl, or alkynyl group, or a group comprising an ether, a ketone, an amide, an ester, a carbamate, an amine, a piperidine, a phosphate, a phosphodiester, a phosphorothioate, a triazole, a pyrrolidine, a disulfide, or a thioether.

In certain such embodiments, the cell-targeting moiety of the conjugate group has the following structure:



wherein Y is selected from a C_1 - C_{12} substituted or unsubstituted alkyl group, or a group comprising an ether, an amine, a piperidine, a phosphate, a phosphodiester, or a phosphorothioate.

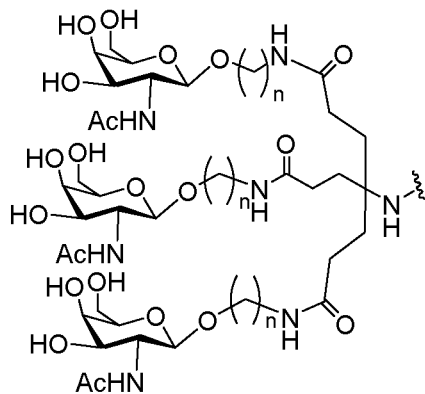
In certain such embodiments, the cell-targeting moiety of the conjugate group has the following structure:



5

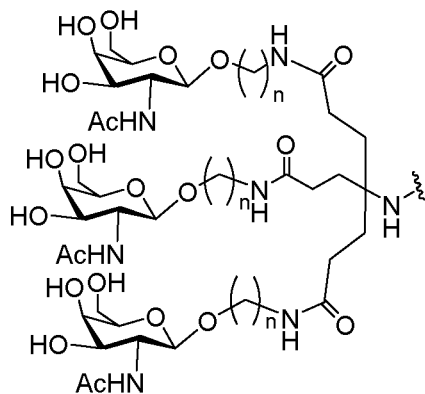
wherein Y is selected from a C_1 - C_{12} substituted or unsubstituted alkyl group.

In certain such embodiments, the cell-targeting moiety of the conjugate group has the following structure:



Wherein n is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12.

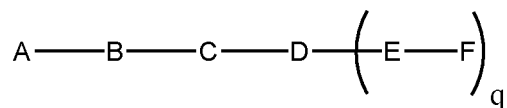
10 In certain such embodiments, the cell-targeting moiety of the conjugate group has the following structure:



wherein n is 4, 5, 6, 7, or 8.

b. Certain conjugated antisense compounds

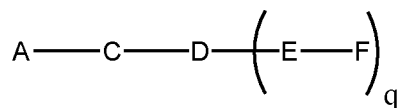
- 5 In certain embodiments, the conjugates are bound to a nucleoside of the antisense oligonucleotide at the 2', 3', or 5' position of the nucleoside. In certain embodiments, a conjugated antisense compound has the following structure:



- 10 wherein
 A is the antisense oligonucleotide;
 B is the cleavable moiety
 C is the conjugate linker
 D is the branching group
 15 each E is a tether;
 each F is a ligand; and
 q is an integer between 1 and 5.

In certain embodiments, a conjugated antisense compound has the following structure:

20



- wherein
 A is the antisense oligonucleotide;
 C is the conjugate linker

D is the branching group

each E is a tether;

each F is a ligand; and

q is an integer between 1 and 5.

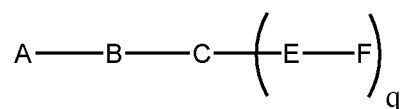
5 In certain such embodiments, the conjugate linker comprises at least one cleavable bond.

In certain such embodiments, the branching group comprises at least one cleavable bond.

In certain embodiments each tether comprises at least one cleavable bond.

10 In certain embodiments, the conjugates are bound to a nucleoside of the antisense oligonucleotide at the 2', 3', of 5' position of the nucleoside.

In certain embodiments, a conjugated antisense compound has the following structure:



wherein

15 A is the antisense oligonucleotide;

B is the cleavable moiety

C is the conjugate linker

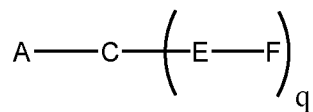
each E is a tether;

each F is a ligand; and

20 q is an integer between 1 and 5.

In certain embodiments, the conjugates are bound to a nucleoside of the antisense oligonucleotide at the 2', 3', of 5' position of the nucleoside. In certain embodiments, a conjugated antisense compound has the following structure:

25



wherein

A is the antisense oligonucleotide;

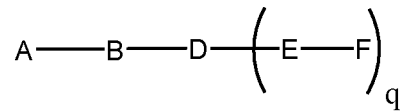
C is the conjugate linker

30 each E is a tether;

each F is a ligand; and

q is an integer between 1 and 5.

In certain embodiments, a conjugated antisense compound has the following structure:



wherein

A is the antisense oligonucleotide;

B is the cleavable moiety

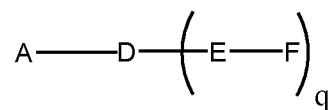
D is the branching group

each E is a tether;

each F is a ligand; and

q is an integer between 1 and 5.

In certain embodiments, a conjugated antisense compound has the following structure:



wherein

A is the antisense oligonucleotide;

D is the branching group

each E is a tether;

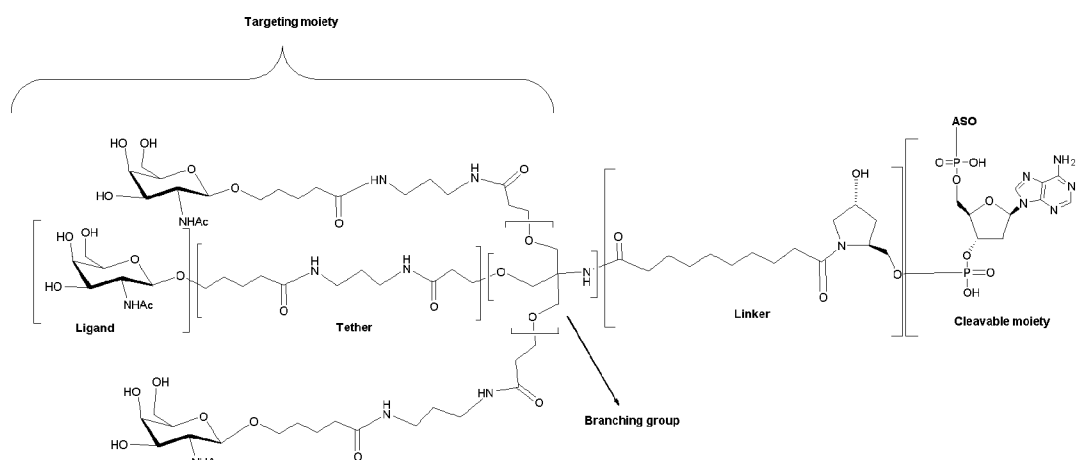
each F is a ligand; and

q is an integer between 1 and 5.

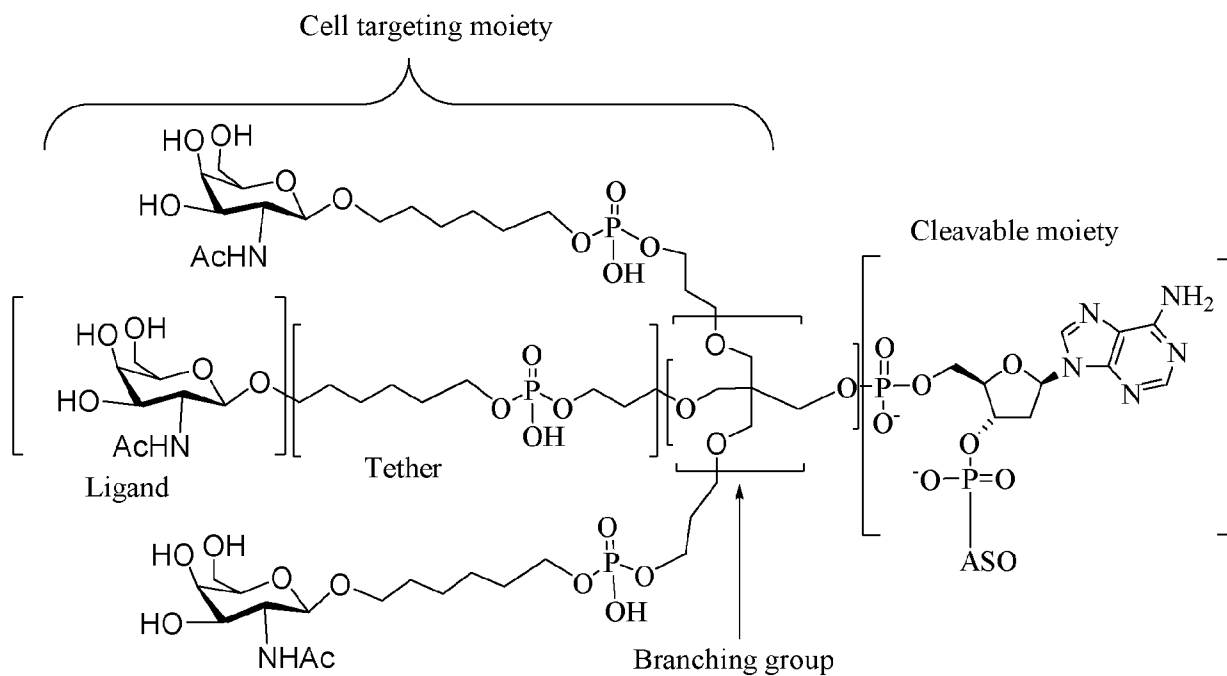
In certain such embodiments, the conjugate linker comprises at least one cleavable bond.

In certain embodiments each tether comprises at least one cleavable bond.

In certain embodiments, a conjugated antisense compound has a structure selected from among the following:

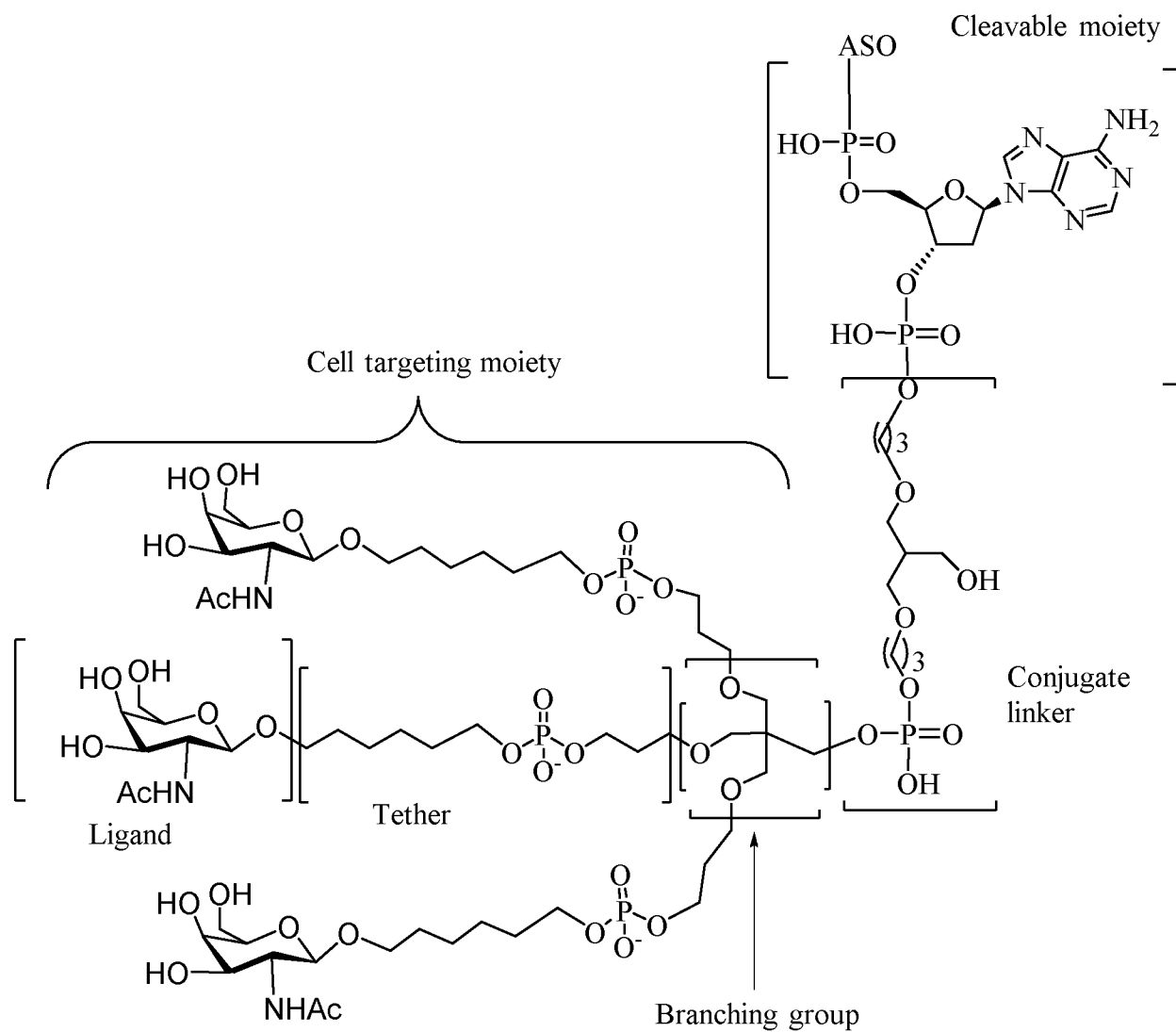


In certain embodiments, a conjugated antisense compound has a structure selected from among the following:

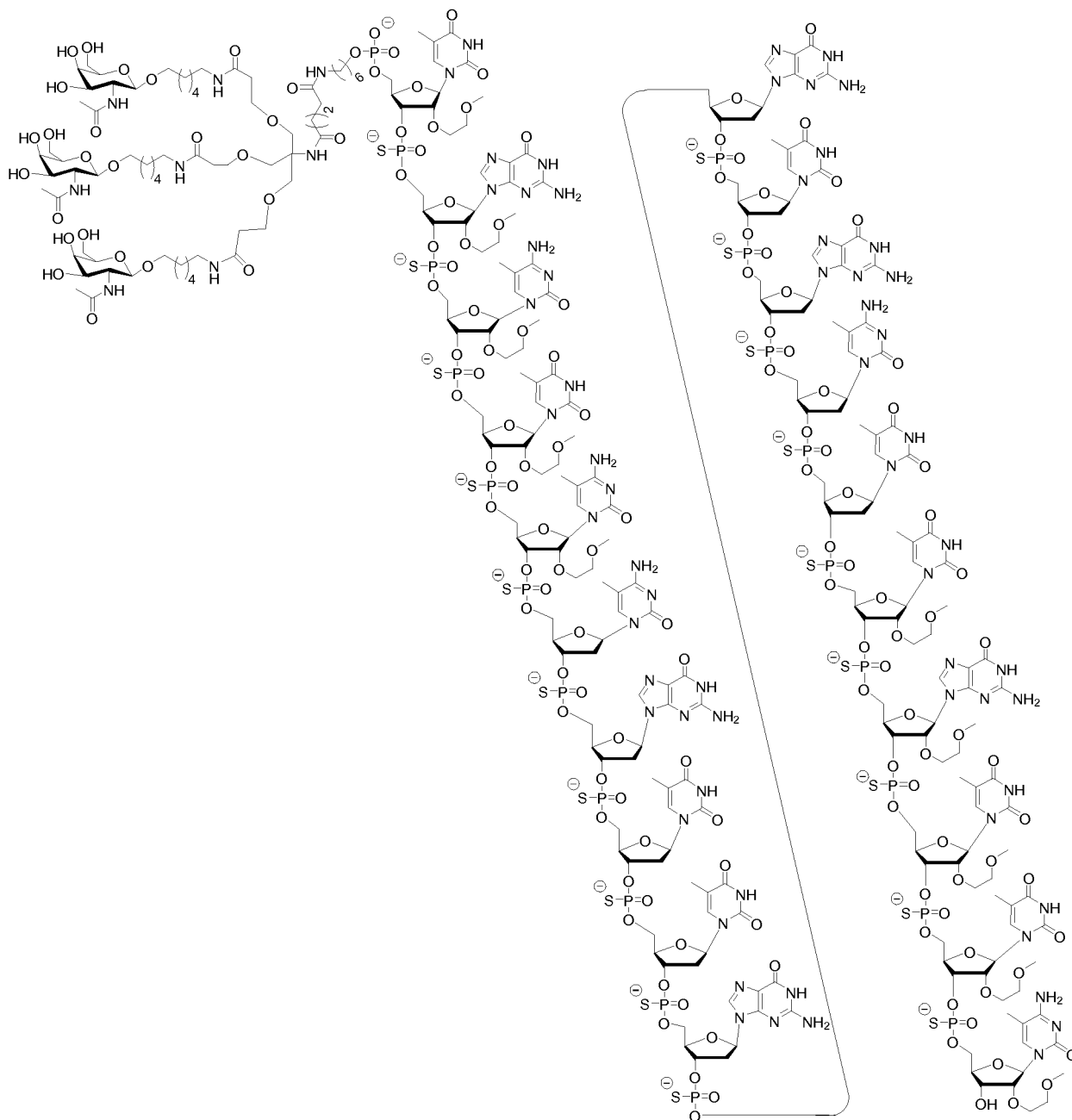


5

In certain embodiments, a conjugated antisense compound has a structure selected from among the following:



In certain embodiments, the conjugated antisense compound has the following structure:



Representative United States patents, United States patent application publications, and international patent application publications that teach the preparation of certain of the above noted conjugates, conjugated antisense compounds, tethers, linkers, branching groups, ligands, cleavable moieties as well as other
 5 modifications include without limitation, US 5,994,517, US 6,300,319, US 6,660,720, US 6,906,182, US 7,262,177, US 7,491,805, US 8,106,022, US 7,723,509, US 2006/0148740, US 2011/0123520, WO 2013/033230 and WO 2012/037254, each of which is incorporated by reference herein in its entirety.

Representative publications that teach the preparation of certain of the above noted conjugates, conjugated antisense compounds, tethers, linkers, branching groups, ligands, cleavable moieties as well as

other modifications include without limitation, BIESSEN et al., "The Cholesterol Derivative of a Triantennary Galactoside with High Affinity for the Hepatic Asialoglycoprotein Receptor: a Potent Cholesterol Lowering Agent" *J. Med. Chem.* (1995) 38:1846-1852, BIESSEN et al., "Synthesis of Cluster Galactosides with High Affinity for the Hepatic Asialoglycoprotein Receptor" *J. Med. Chem.* (1995) 38:1538-1546, LEE et al., "New and more efficient multivalent glyco-ligands for asialoglycoprotein receptor of mammalian hepatocytes" *Bioorganic & Medicinal Chemistry* (2011) 19:2494-2500, RENSEN et al., "Determination of the Upper Size Limit for Uptake and Processing of Ligands by the Asialoglycoprotein Receptor on Hepatocytes in Vitro and in Vivo" *J. Biol. Chem.* (2001) 276(40):37577-37584, RENSEN et al., "Design and Synthesis of Novel N-Acetylgalactosamine-Terminated Glycolipids for Targeting of Lipoproteins to the Hepatic Asialoglycoprotein Receptor" *J. Med. Chem.* (2004) 47:5798-5808, SLIEDREGT et al., "Design and Synthesis of Novel Amphiphilic Dendritic Galactosides for Selective Targeting of Liposomes to the Hepatic Asialoglycoprotein Receptor" *J. Med. Chem.* (1999) 42:609-618, and Valentijn *et al.*, "Solid-phase synthesis of lysine-based cluster galactosides with high affinity for the Asialoglycoprotein Receptor" *Tetrahedron*, 1997, 53(2), 759-770, each of which is incorporated by reference herein in its entirety.

In certain embodiments, conjugated antisense compounds comprise an RNase H based oligonucleotide (such as a gapmer) or a splice modulating oligonucleotide (such as a fully modified oligonucleotide) and any conjugate group comprising at least one, two, or three GalNAc groups. In certain embodiments a conjugated antisense compound comprises any conjugate group found in any of the following references: Lee, *Carbohydr Res*, 1978, 67, 509-514; Connolly et al., *J Biol Chem*, 1982, 257, 939-945; Pavia et al., *Int J Pep Protein Res*, 1983, 22, 539-548; Lee et al., *Biochem*, 1984, 23, 4255-4261; Lee et al., *Glycoconjugate J*, 1987, 4, 317-328; Toyokuni et al., *Tetrahedron Lett*, 1990, 31, 2673-2676; Biessen et al., *J Med Chem*, 1995, 38, 1538-1546; Valentijn et al., *Tetrahedron*, 1997, 53, 759-770; Kim et al., *Tetrahedron Lett*, 1997, 38, 3487-3490; Lee et al., *Bioconjug Chem*, 1997, 8, 762-765; Kato et al., *Glycobiol*, 2001, 11, 821-829; Rensen et al., *J Biol Chem*, 2001, 276, 37577-37584; Lee et al., *Methods Enzymol*, 2003, 362, 38-43; Westerlind et al., *Glycoconj J*, 2004, 21, 227-241; Lee et al., *Bioorg Med Chem Lett*, 2006, 16(19), 5132-5135; Maierhofer et al., *Bioorg Med Chem*, 2007, 15, 7661-7676; Khorev et al., *Bioorg Med Chem*, 2008, 16, 5216-5231; Lee et al., *Bioorg Med Chem*, 2011, 19, 2494-2500; Kornilova et al., *Analyt Biochem*, 2012, 425, 43-46; Pujol et al., *Angew Chemie Int Ed Engl*, 2012, 51, 7445-7448; Biessen et al., *J Med Chem*, 1995, 38, 1846-1852; Slidregt et al., *J Med Chem*, 1999, 42, 609-618; Rensen et al., *J Med Chem*, 2004, 47, 5798-5808; Rensen et al., *Arterioscler Thromb Vasc Biol*, 2006, 26, 169-175; van Rossenberg et al., *Gene Ther*, 2004, 11, 457-464; Sato et al., *J Am Chem Soc*, 2004, 126, 14013-14022; Lee et al., *J Org Chem*, 2012, 77, 7564-7571; Biessen et al., *FASEB J*, 2000, 14, 1784-1792; Rajur et al., *Bioconjug Chem*, 1997, 8, 935-940; Duff et al., *Methods Enzymol*, 2000, 313, 297-321; Maier et al., *Bioconjug Chem*, 2003, 14, 18-29; Jayaprakash et al., *Org Lett*, 2010, 12, 5410-5413; Manoharan, *Antisense Nucleic Acid Drug Dev*, 2002, 12,

103-128; Merwin et al., *Bioconjug Chem*, 1994, 5, 612-620; Tomiya et al., *Bioorg Med Chem*, 2013, 21, 5275-5281; International applications WO1998/013381; WO2011/038356; WO1997/046098; WO2008/098788; WO2004/101619; WO2012/037254; WO2011/120053; WO2011/100131; WO2011/163121; WO2012/177947; WO2013/033230; WO2013/075035; WO2012/083185; 5 WO2012/083046; WO2009/082607; WO2009/134487; WO2010/144740; WO2010/148013; WO1997/020563; WO2010/088537; WO2002/043771; WO2010/129709; WO2012/068187; WO2009/126933; WO2004/024757; WO2010/054406; WO2012/089352; WO2012/089602; WO2013/166121; WO2013/165816; U.S. Patents 4,751,219; 8,552,163; 6,908,903; 7,262,177; 5,994,517; 6,300,319; 8,106,022; 7,491,805; 7,491,805; 7,582,744; 8,137,695; 6,383,812; 6,525,031; 6,660,720; 10 7,723,509; 8,541,548; 8,344,125; 8,313,772; 8,349,308; 8,450,467; 8,501,930; 8,158,601; 7,262,177; 6,906,182; 6,620,916; 8,435,491; 8,404,862; 7,851,615; Published U.S. Patent Application Publications US2011/0097264; US2011/0097265; US2013/0004427; US2005/0164235; US2006/0148740; US2008/0281044; US2010/0240730; US2003/0119724; US2006/0183886; US2008/0206869; US2011/0269814; US2009/0286973; US2011/0207799; US2012/0136042; US2012/0165393; 15 US2008/0281041; US2009/0203135; US2012/0035115; US2012/0095075; US2012/0101148; US2012/0128760; US2012/0157509; US2012/0230938; US2013/0109817; US2013/0121954; US2013/0178512; US2013/0236968; US2011/0123520; US2003/0077829; US2008/0108801; and US2009/0203132; each of which is incorporated by reference in its entirety.

C. Certain Uses and Features

20 In certain embodiments, conjugated antisense compounds exhibit potent target RNA reduction *in vivo*. In certain embodiments, unconjugated antisense compounds accumulate in the kidney. In certain embodiments, conjugated antisense compounds accumulate in the liver. In certain embodiments, conjugated antisense compounds are well tolerated. Such properties render conjugated antisense compounds particularly useful for inhibition of many target RNAs, including, but not limited to those involved in metabolic, 25 cardiovascular and other diseases, disorders or conditions. Thus, provided herein are methods of treating such diseases, disorders or conditions by contacting liver tissues with the conjugated antisense compounds targeted to RNAs associated with such diseases, disorders or conditions. Thus, also provided are methods for ameliorating any of a variety of metabolic, cardiovascular and other diseases, disorders or conditions with the conjugated antisense compounds of the present invention.

30 In certain embodiments, conjugated antisense compounds are more potent than unconjugated counterpart at a particular tissue concentration. Without wishing to be bound by any theory or mechanism, in certain embodiments, the conjugate may allow the conjugated antisense compound to enter the cell more efficiently or to enter the cell more productively. For example, in certain embodiments conjugated antisense compounds may exhibit greater target reduction as compared to its unconjugated counterpart wherein both 35 the conjugated antisense compound and its unconjugated counterpart are present in the tissue at the same

concentrations. For example, in certain embodiments conjugated antisense compounds may exhibit greater target reduction as compared to its unconjugated counterpart wherein both the conjugated antisense compound and its unconjugated counterpart are present in the liver at the same concentrations.

Productive and non-productive uptake of oligonucleotides has been discussed previously (*See e.g.* Geary, R. S., E. Wancewicz, et al. (2009). "Effect of Dose and Plasma Concentration on Liver Uptake and Pharmacologic Activity of a 2'-Methoxyethyl Modified Chimeric Antisense Oligonucleotide Targeting PTEN." *Biochem. Pharmacol.* 78(3): 284-91; & Koller, E., T. M. Vincent, et al. (2011). "Mechanisms of single-stranded phosphorothioate modified antisense oligonucleotide accumulation in hepatocytes." *Nucleic Acids Res.* 39(11): 4795-807). Conjugate groups described herein may improve productive uptake.

In certain embodiments, the conjugate groups described herein may further improve potency by increasing the affinity of the conjugated antisense compound for a particular type of cell or tissue. In certain embodiments, the conjugate groups described herein may further improve potency by increasing recognition of the conjugated antisense compound by one or more cell-surface receptors. . In certain embodiments, the conjugate groups described herein may further improve potency by facilitating endocytosis of the conjugated antisense compound.

In certain embodiments, the cleavable moiety may further improve potency by allowing the conjugate to be cleaved from the antisense oligonucleotide after the conjugated antisense compound has entered the cell. Accordingly, in certain embodiments, conjugated antisense compounds can be administered at doses lower than would be necessary for unconjugated antisense oligonucleotides.

Phosphorothioate linkages have been incorporated into antisense oligonucleotides previously. Such phosphorothioate linkages are resistant to nucleases and so improve stability of the oligonucleotide. Further, phosphorothioate linkages also bind certain proteins, which results in accumulation of antisense oligonucleotide in the liver. Oligonucleotides with fewer phosphorothioate linkages accumulate less in the liver and more in the kidney (see, for example, Geary, R., "Pharmacokinetic Properties of 2'-O-(2-Methoxyethyl)-Modified Oligonucleotide Analogs in Rats," *Journal of Pharmacology and Experimental Therapeutics*, Vol. 296, No. 3, 890-897; & *Pharmacological Properties of 2'-O-Methoxyethyl Modified Oligonucleotides* in *Antisense a Drug Technology*, Chapter 10, Crooke, S.T., ed., 2008) In certain embodiments, oligonucleotides with fewer phosphorothioate internucleoside linkages and more phosphodiester internucleoside linkages accumulate less in the liver and more in the kidney. When treating diseases in the liver, this is undesirable for several reasons (1) less drug is getting to the site of desired action (liver); (2) drug is escaping into the urine; and (3) the kidney is exposed to relatively high concentration of drug which can result in toxicities in the kidney. Thus, for liver diseases, phosphorothioate linkages provide important benefits.

In certain embodiments, however, administration of oligonucleotides uniformly linked by phosphorothioate internucleoside linkages induces one or more proinflammatory reactions. (see for example: *J Lab Clin Med.* 1996 Sep; 128(3):329-38. "Amplification of antibody production by phosphorothioate oligodeoxynucleotides". Branda et al.; and see also for example: *Toxicologic Properties in Antisense a Drug Technology*, Chapter 12, pages 342-351, Crooke, S.T., ed., 2008). In certain embodiments, administration of oligonucleotides wherein most of the internucleoside linkages comprise phosphorothioate internucleoside linkages induces one or more proinflammatory reactions.

In certain embodiments, the degree of proinflammatory effect may depend on several variables (e.g. backbone modification, off-target effects, nucleobase modifications, and/or nucleoside modifications) see for example: *Toxicologic Properties in Antisense a Drug Technology*, Chapter 12, pages 342-351, Crooke, S.T., ed., 2008). In certain embodiments, the degree of proinflammatory effect may be mitigated by adjusting one or more variables. For example the degree of proinflammatory effect of a given oligonucleotide may be mitigated by replacing any number of phosphorothioate internucleoside linkages with phosphodiester internucleoside linkages and thereby reducing the total number of phosphorothioate internucleoside linkages.

In certain embodiments, it would be desirable to reduce the number of phosphorothioate linkages, if doing so could be done without losing stability and without shifting the distribution from liver to kidney. For example, in certain embodiments, the number of phosphorothioate linkages may be reduced by replacing phosphorothioate linkages with phosphodiester linkages. In such an embodiment, the antisense compound having fewer phosphorothioate linkages and more phosphodiester linkages may induce less proinflammatory reactions or no proinflammatory reaction. Although the the antisense compound having fewer phosphorothioate linkages and more phosphodiester linkages may induce fewer proinflammatory reactions, the antisense compound having fewer phosphorothioate linkages and more phosphodiester linkages may not accumulate in the liver and may be less efficacious at the same or similar dose as compared to an antisense compound having more phosphorothioate linkages. In certain embodiments, it is therefore desirable to design an antisense compound that has a plurality of phosphodiester bonds and a plurality of phosphorothioate bonds but which also possesses stability and good distribution to the liver.

In certain embodiments, conjugated antisense compounds accumulate more in the liver and less in the kidney than unconjugated counterparts, even when some of the phosphorothioate linkages are replaced with less proinflammatory phosphodiester internucleoside linkages. In certain embodiments, conjugated antisense compounds accumulate more in the liver and are not excreted as much in the urine compared to its unconjugated counterparts, even when some of the phosphorothioate linkages are replaced with less proinflammatory phosphodiester internucleoside linkages. In certain embodiments, the use of a conjugate allows one to design more potent and better tolerated antisense drugs. Indeed, in certain emobidments, conjugated antisense compounds have larger therapeutic indexes than unconjugated counterparts. This allows the conjugated antisense compound to be administered at a higher absolute dose, because there is less risk of proinflammatory response and less risk of kidney toxicity. This higher dose, allows one to dose less

frequently, since the clearance (metabolism) is expected to be similar. Further, because the compound is more potent, as described above, one can allow the concentration to go lower before the next dose without losing therapeutic activity, allowing for even longer periods between dosing.

5 In certain embodiments, the inclusion of some phosphorothioate linkages remains desirable. For example, the terminal linkages are vulnerable to exonucleases and so in certain embodiments, those linkages are phosphorothioate or other modified linkage. Internucleoside linkages linking two deoxynucleosides are vulnerable to endonucleases and so in certain embodiments those those linkages are phosphorothioate or other modified linkage. Internucleoside linkages between a modified nucleoside and a deoxynucleoside where the deoxynucleoside is on the 5' side of the linkage deoxynucleosides are vulnerable to endonucleases
10 and so in certain embodiments those those linkages are phosphorothioate or other modified linkage. Internucleoside linkages between two modified nucleosides of certain types and between a deoxynucleoside and a modified nucleoside of certain type where the modified nucleoside is at the 5' side of the linkage are sufficiently resistant to nuclease digestion, that the linkage can be phosphodiester.

In certain embodiments, the antisense oligonucleotide of a conjugated antisense compound
15 comprises fewer than 16 phosphorothioate linkages. In certain embodiments, the antisense oligonucleotide of a conjugated antisense compound comprises fewer than 15 phosphorothioate linkages. In certain embodiments, the antisense oligonucleotide of a conjugated antisense compound comprises fewer than 14 phosphorothioate linkages. In certain embodiments, the antisense oligonucleotide of a conjugated antisense compound comprises fewer than 13 phosphorothioate linkages. In certain embodiments, the antisense
20 oligonucleotide of a conjugated antisense compound comprises fewer than 12 phosphorothioate linkages. In certain embodiments, the antisense oligonucleotide of a conjugated antisense compound comprises fewer than 11 phosphorothioate linkages. In certain embodiments, the antisense oligonucleotide of a conjugated antisense compound comprises fewer than 10 phosphorothioate linkages. In certain embodiments, the antisense oligonucleotide of a conjugated antisense compound comprises fewer than 9 phosphorothioate
25 linkages. In certain embodiments, the antisense oligonucleotide of a conjugated antisense compound comprises fewer than 8 phosphorothioate linkages.

In certain embodiments, antisense compounds comprising one or more conjugate group described herein has increased activity and/or potency and/or tolerability compared to a parent antisense compound lacking such one or more conjugate group. Accordingly, in certain embodiments, attachment of such
30 conjugate groups to an oligonucleotide is desirable. Such conjugate groups may be attached at the 5'-, and/or 3'- end of an oligonucleotide. In certain instances, attachment at the 5'-end is synthetically desirable. Typically, oligonucleotides are synthesized by attachment of the 3' terminal nucleoside to a solid support and sequential coupling of nucleosides from 3' to 5' using techniques that are well known in the art. Accordingly if a conjugate group is desired at the 3'-terminus, one may (1) attach the conjugate group to the

3'-terminal nucleoside and attach that conjugated nucleoside to the solid support for subsequent preparation of the oligonucleotide or (2) attach the conjugate group to the 3'-terminal nucleoside of a completed oligonucleotide after synthesis. Neither of these approaches is very efficient and thus both are costly. In particular, attachment of the conjugated nucleoside to the solid support, while demonstrated in the Examples
5 herein, is an inefficient process. In certain embodiments, attaching a conjugate group to the 5'-terminal nucleoside is synthetically easier than attachment at the 3'-end. One may attach a non-conjugated 3' terminal nucleoside to the solid support and prepare the oligonucleotide using standard and well characterized reactions. One then needs only to attach a 5' nucleoside having a conjugate group at the final coupling step. In certain embodiments, this is more efficient than attaching a conjugated nucleoside directly to the solid
10 support as is typically done to prepare a 3'-conjugated oligonucleotide. The Examples herein demonstrate attachment at the 5'-end. In addition, certain conjugate groups have synthetic advantages. For Example, certain conjugate groups comprising phosphorus linkage groups are synthetically simpler and more efficiently prepared than other conjugate groups, including conjugate groups reported previously (e.g., WO/2012/037254).

15 In certain embodiments, conjugated antisense compounds are administered to a subject. In such embodiments, antisense compounds comprising one or more conjugate group described herein has increased activity and/or potency and/or tolerability compared to a parent antisense compound lacking such one or more conjugate group. Without being bound by mechanism, it is believed that the conjugate group helps with distribution, delivery, and/or uptake into a target cell or tissue. In certain embodiments, once inside the target
20 cell or tissue, it is desirable that all or part of the conjugate group to be cleaved to release the active oligonucleotide. In certain embodiments, it is not necessary that the entire conjugate group be cleaved from the oligonucleotide. For example, in Example 20 a conjugated oligonucleotide was administered to mice and a number of different chemical species, each comprising a different portion of the conjugate group remaining on the oligonucleotide, were detected (Table 23a). This conjugated antisense compound demonstrated good
25 potency (Table 23). Thus, in certain embodiments, such metabolite profile of multiple partial cleavage of the conjugate group does not interfere with activity/potency. Nevertheless, in certain embodiments it is desirable that a prodrug (conjugated oligonucleotide) yield a single active compound. In certain instances, if multiple forms of the active compound are found, it may be necessary to determine relative amounts and activities for each one. In certain embodiments where regulatory review is required (e.g., USFDA or counterpart) it is
30 desirable to have a single (or predominantly single) active species. In certain such embodiments, it is desirable that such single active species be the antisense oligonucleotide lacking any portion of the conjugate group. In certain embodiments, conjugate groups at the 5'-end are more likely to result in complete metabolism of the conjugate group. Without being bound by mechanism it may be that endogenous enzymes responsible for metabolism at the 5' end (e.g., 5' nucleases) are more active/efficient than the 3' counterparts.
35 In certain embodiments, the specific conjugate groups are more amenable to metabolism to a single active

species. In certain embodiments, certain conjugate groups are more amenable to metabolism to the oligonucleotide.

D. Antisense

In certain embodiments, oligomeric compounds of the present invention are antisense compounds.

5 In such embodiments, the oligomeric compound is complementary to a target nucleic acid. In certain embodiments, a target nucleic acid is an RNA. In certain embodiments, a target nucleic acid is a non-coding RNA. In certain embodiments, a target nucleic acid encodes a protein. In certain embodiments, a target nucleic acid is selected from a mRNA, a pre-mRNA, a microRNA, a non-coding RNA, including small non-coding RNA, and a promoter-directed RNA. In certain embodiments, oligomeric compounds are at least
10 partially complementary to more than one target nucleic acid. For example, oligomeric compounds of the present invention may be microRNA mimics, which typically bind to multiple targets.

In certain embodiments, antisense compounds comprise a portion having a nucleobase sequence at least 70% complementary to the nucleobase sequence of a target nucleic acid. In certain embodiments, antisense compounds comprise a portion having a nucleobase sequence at least 80% complementary to the
15 nucleobase sequence of a target nucleic acid. In certain embodiments, antisense compounds comprise a portion having a nucleobase sequence at least 90% complementary to the nucleobase sequence of a target nucleic acid. In certain embodiments, antisense compounds comprise a portion having a nucleobase sequence at least 95% complementary to the nucleobase sequence of a target nucleic acid. In certain
20 embodiments, antisense compounds comprise a portion having a nucleobase sequence at least 98% complementary to the nucleobase sequence of a target nucleic acid. In certain embodiments, antisense compounds comprise a portion having a nucleobase sequence that is 100% complementary to the nucleobase sequence of a target nucleic acid. In certain embodiments, antisense compounds are at least 70%, 80%, 90%, 95%, 98%, or 100% complementary to the nucleobase sequence of a target nucleic acid over the entire length of the antisense compound.

25 Antisense mechanisms include any mechanism involving the hybridization of an oligomeric compound with target nucleic acid, wherein the hybridization results in a biological effect. In certain embodiments, such hybridization results in either target nucleic acid degradation or occupancy with concomitant inhibition or stimulation of the cellular machinery involving, for example, translation, transcription, or polyadenylation of the target nucleic acid or of a nucleic acid with which the target nucleic
30 acid may otherwise interact.

One type of antisense mechanism involving degradation of target RNA is RNase H mediated antisense. RNase H is a cellular endonuclease which cleaves the RNA strand of an RNA:DNA duplex. It is known in the art that single-stranded antisense compounds which are "DNA-like" elicit RNase H activity in

mammalian cells. Activation of RNase H, therefore, results in cleavage of the RNA target, thereby greatly enhancing the efficiency of DNA-like oligonucleotide-mediated inhibition of gene expression.

Antisense mechanisms also include, without limitation RNAi mechanisms, which utilize the RISC pathway. Such RNAi mechanisms include, without limitation siRNA, ssRNA and microRNA mechanisms.

5 Such mechanisms include creation of a microRNA mimic and/or an anti-microRNA.

Antisense mechanisms also include, without limitation, mechanisms that hybridize or mimic non-coding RNA other than microRNA or mRNA. Such non-coding RNA includes, but is not limited to promoter-directed RNA and short and long RNA that effects transcription or translation of one or more nucleic acids.

10 In certain embodiments, oligonucleotides comprising conjugates described herein are RNAi compounds. In certain embodiments, oligomeric oligonucleotides comprising conjugates described herein are ssRNA compounds. In certain embodiments, oligonucleotides comprising conjugates described herein are paired with a second oligomeric compound to form an siRNA. In certain such embodiments, the second oligomeric compound also comprises a conjugate. In certain embodiments, the second oligomeric compound
15 is any modified or unmodified nucleic acid. In certain embodiments, the oligonucleotides comprising conjugates described herein is the antisense strand in an siRNA compound. In certain embodiments, the oligonucleotides comprising conjugates described herein is the sense strand in an siRNA compound. In embodiments in which the conjugated oligomeric compound is double-stranded siRNA, the conjugate may be on the sense strand, the antisense strand or both the sense strand and the antisense strand.

20 **C. Apolipoprotein (a) (apo(a))**

In certain embodiments, conjugated antisense compounds target any apo(a) nucleic acid. In certain embodiments, the target nucleic acid encodes an apo(a) target protein that is clinically relevant. In such embodiments, modulation of the target nucleic acid results in clinical benefit.

25 The targeting process usually includes determination of at least one target region, segment, or site within the target nucleic acid for the antisense interaction to occur such that the desired effect will result.

In certain embodiments, a target region is a structurally defined region of the nucleic acid. For example, in certain such embodiments, a target region may encompass a 3' UTR, a 5' UTR, an exon, an intron, a coding region, a translation initiation region, translation termination region, or other defined nucleic acid region or target segment.

30 In certain embodiments, a target segment is at least about an 8-nucleobase portion of a target region to which a conjugated antisense compound is targeted. Target segments can include DNA or RNA sequences that comprise at least 8 consecutive nucleobases from the 5'-terminus of one of the target segments (the

remaining nucleobases being a consecutive stretch of the same DNA or RNA beginning immediately upstream of the 5'-terminus of the target segment and continuing until the DNA or RNA comprises about 8 to about 30 nucleobases). Target segments are also represented by DNA or RNA sequences that comprise at least 8 consecutive nucleobases from the 3'-terminus of one of the target segments (the remaining
5 nucleobases being a consecutive stretch of the same DNA or RNA beginning immediately downstream of the 3'-terminus of the target segment and continuing until the DNA or RNA comprises about 8 to about 30 nucleobases). Target segments can also be represented by DNA or RNA sequences that comprise at least 8 consecutive nucleobases from an internal portion of the sequence of a target segment, and may extend in either or both directions until the conjugated antisense compound comprises about 8 to about 30 nucleobases.

10 In certain embodiments, antisense compounds targeted to an apo(a) nucleic acid can be modified as described herein. In certain embodiments, the antisense compounds can have a modified sugar moiety, an unmodified sugar moiety or a mixture of modified and unmodified sugar moieties as described herein. In certain embodiments, the antisense compounds can have a modified internucleoside linkage, an unmodified internucleoside linkage or a mixture of modified and unmodified internucleoside linkages as described
15 herein. In certain embodiments, the antisense compounds can have a modified nucleobase, an unmodified nucleobase or a mixture of modified and unmodified nucleobases as described herein. In certain embodiments, the antisense compounds can have a motif as described herein.

In certain embodiments, antisense compounds targeted to apo(a) nucleic acids can be conjugated as described herein.

20 One apo(a) protein is linked via a disulfide bond to a single apolipoprotein B (apoB) protein to form a lipoprotein(a) (Lp(a)) particle. The apo(a) protein shares a high degree of homology with plasminogen particularly within the kringle IV type 2 repetitive domain. It is thought that the kringle repeat domain in apo(a) may be responsible for its pro-thrombotic and anti-fibrinolytic properties, potentially enhancing atherosclerotic progression. Apo(a) is transcriptionally regulated by IL-6 and in studies in rheumatoid
25 arthritis patients treated with an IL-6 inhibitor (tocilizumab), plasma levels were reduced by 30% after 3 month treatment. Apo(a) has been shown to preferentially bind oxidized phospholipids and potentiate vascular inflammation. Further, studies suggest that the Lp(a) particle may also stimulate endothelial permeability, induce plasminogen activator inhibitor type-1 expression and activate macrophage interleukin-8 secretion. Importantly, recent genetic association studies revealed that Lp(a) was an independent risk factor
30 for myocardial infarction, stroke, peripheral vascular disease and abdominal aortic aneurysm. Further, in the Precocious Coronary Artery Disease (PROCARDIS) study, Clarke *et al.* described robust and independent associations between coronary heart disease and plasma Lp(a) concentrations. Additionally, Solfrizzi *et al.*, suggested that increased serum Lp(a) may be linked to an increased risk for Alzheimer's Disease (AD). Antisense compounds targeting apo(a) have been previously disclosed in WO2005/000201 and US2010-

0331390, herein incorporated by reference in its entirety. An antisense oligonucleobase targeting Apo(a), ISIS-APOA_{Rxx}, was assessed in a Phase I clinical trial to study its safety profile.

Certain Conjugated Antisense Compounds Targeted to an Apo(a) Nucleic Acid

5 In certain embodiments, conjugated antisense compounds are targeted to an Apo(a) nucleic acid having the sequence of GENBANK® Accession No. NM_005577.2, incorporated herein as SEQ ID NO: 1; GENBANK Accession No. NT_007422.12 truncated from nucleotides 3230000 to 3380000, incorporated herein as SEQ ID NO: 2; GENBANK Accession No. NT_025741.15 truncated from nucleotides 65120000 to 65258000, designated herein as SEQ ID NO: 3; and GENBANK Accession No. NM_005577.1, incorporated
10 herein as SEQ ID NO: 4. In certain such embodiments, a conjugated antisense compound is at least 90%, at least 95%, or 100% complementary to any of the nucleobase sequences of SEQ ID NOs: 1-4.

In certain embodiments, a conjugated antisense compound targeted to any of the nucleobase sequences of SEQ ID NOs: 1-4 comprises an at least 8 consecutive nucleobase sequence selected from the nucleobase sequence of any of SEQ ID NOs: 12-130, 133, 134. In certain embodiments, a conjugated
15 antisense compound targeted to any of SEQ ID NOs: 1-4 comprises a nucleobase sequence selected from the nucleobase sequence of any of SEQ ID NOs: 12-130, 133, 134.

Table A: Antisense Compounds targeted to Apo(a) SEQ ID NO: 1

ISIS No	Target Start Site	Sequence (5'-3')	Motif	SEQ ID NO
494372	3901	TGCTCCGTTGGTGCTTGTC	eeeeeddddddddeeeee	58
494283	584	TCTTCCTGTGACAGTGGTGG	eeeeeddddddddeeeee	26
	926			
	1610			
	1952			
	2294			
494284	3320	TTCTTCCTGTGACAGTGGTG	eeeeeddddddddeeeee	27
	585			
	927			
	1611			
	1953			
494286	2295	GGTTCTTCCTGTGACAGTGG	eeeeeddddddddeeeee	29
	3321			
	587			
	929			
	1613			
494301	1955	CGACTATGCGAGTGTGGTGT	eeeeeddddddddeeeee	38
	2297			
	628			
	970			

	1312			
	1654			
	1996			
	2338			
	2680			
	3022			
494302	629	CCGACTATGCGAGTGTGGTG	eeeeeddddddddeeeee	39
	971			
	1313			
	1655			
	1997			
	2339			
	2681			
	3023			

Apo(a) Therapeutic Indications

In certain embodiments, the invention provides methods for using a conjugated antisense compound targeted to an apo(a) nucleic acid for modulating the expression of apo(a) in a subject. In certain
5 embodiments, the expression of apo(a) is reduced.

In certain embodiments, provided herein are methods of treating a subject comprising administering one or more pharmaceutical compositions as described herein. In certain embodiments, the invention provides methods for using a conjugated antisense compound targeted to an apo(a) nucleic acid in a pharmaceutical composition for treating a subject. In certain embodiments, the individual has an apo(a)
10 related disease. In certain embodiments, the individual has an Lp(a) related disease. In certain embodiments, the individual has an inflammatory, cardiovascular and/or a metabolic disease, disorder or condition.

In certain embodiments, the subject has an inflammatory, cardiovascular and/or metabolic disease, disorder or condition.

In certain embodiments, the cardiovascular diseases, disorders or conditions include, but are not
15 limited to, aortic stenosis, aneurysm (e.g., abdominal aortic aneurysm), angina, arrhythmia, atherosclerosis, cerebrovascular disease, coronary artery disease, coronary heart disease, dyslipidemia, hypercholesterolemia, hyperlipidemia, hypertension, hypertriglyceridemia, myocardial infarction, peripheral vascular disease (e.g., peripheral artery disease), stroke and the like.

In certain embodiments, the compounds targeted to apo(a) described herein modulate physiological
20 markers or phenotypes of the cardiovascular disease, disorder or condition. For example, administration of the compounds to animals can decrease LDL and cholesterol levels in those animals compared to untreated animals. In certain embodiments, the modulation of the physiological markers or phenotypes can be associated with inhibition of apo(a) by the compounds.

In certain embodiments, the physiological markers of the cardiovascular disease, disorder or
25 condition can be quantifiable. For example, LDL or cholesterol levels can be measured and quantified by, for

example, standard lipid tests. For such markers, in certain embodiments, the marker can be decreased by about 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95 or 99%, or a range defined by any two of these values.

Also, provided herein are methods for preventing, treating or ameliorating a symptom associated with the cardiovascular disease, disorder or condition in a subject in need thereof. In certain embodiments, provided is a method for reducing the rate of onset of a symptom associated with the cardiovascular disease, disorder or condition. In certain embodiments, provided is a method for reducing the severity of a symptom associated with the cardiovascular disease, disorder or condition. In such embodiments, the methods comprise administering a therapeutically effective amount of a compound targeted to an apo(a) nucleic acid to an individual in need thereof.

The cardiovascular disease, disorder or condition can be characterized by numerous physical symptoms. Any symptom known to one of skill in the art to be associated with the cardiovascular disease, disorder or condition can be prevented, treated, ameliorated or otherwise modulated with the compounds and methods described herein. In certain embodiments, the symptom can be any of, but not limited to, angina, chest pain, shortness of breath, palpitations, weakness, dizziness, nausea, sweating, tachycardia, bradycardia, arrhythmia, atrial fibrillation, swelling in the lower extremities, cyanosis, fatigue, fainting, numbness of the face, numbness of the limbs, claudication or cramping of muscles, bloating of the abdomen or fever.

In certain embodiments, the metabolic diseases, disorders or conditions include, but are not limited to, hyperglycemia, prediabetes, diabetes (type I and type II), obesity, insulin resistance, metabolic syndrome and diabetic dyslipidemia.

In certain embodiments, compounds targeted to apo(a) as described herein modulate physiological markers or phenotypes of the metabolic disease, disorder or condition. For example, administration of the compounds to animals can decrease glucose and insulin resistance levels in those animals compared to untreated animals. In certain embodiments, the modulation of the physiological markers or phenotypes can be associated with inhibition of apo(a) by the compounds.

In certain embodiments, physiological markers of the metabolic disease, disorder or condition can be quantifiable. For example, glucose levels or insulin resistance can be measured and quantified by standard tests known in the art. For such markers, in certain embodiments, the marker can be decreased by about 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95 or 99%, or a range defined by any two of these values. In another example, insulin sensitivity can be measured and quantified by standard tests known in the art. For such markers, in certain embodiments, the marker can be increase by about 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95 or 99%, or a range defined by any two of these values.

Also, provided herein are methods for preventing, treating or ameliorating a symptom associated with the metabolic disease, disorder or condition in a subject in need thereof. In certain embodiments, provided is a method for reducing the rate of onset of a symptom associated with the metabolic disease, disorder or condition. In certain embodiments, provided is a method for reducing the severity of a symptom associated

with the metabolic disease, disorder or condition. In such embodiments, the methods comprise administering a therapeutically effective amount of a compound targeted to an apo(a) nucleic acid to an individual in need thereof.

The metabolic disease, disorder or condition can be characterized by numerous physical symptoms.

Any symptom known to one of skill in the art to be associated with the metabolic disease, disorder or condition can be prevented, treated, ameliorated or otherwise modulated with the compounds and methods described herein. In certain embodiments, the symptom can be any of, but not limited to, excessive urine production (polyuria), excessive thirst and increased fluid intake (polydipsia), blurred vision, unexplained weight loss and lethargy.

In certain embodiments, the inflammatory diseases, disorders or conditions include, but are not limited to, aortic stenosis, coronary artery disease (CAD), Alzheimer's Disease and thromboembolic diseases, disorder or conditions. Certain thromboembolic diseases, disorders or conditions include, but are not limited to, stroke, thrombosis, myocardial infarction and peripheral vascular disease.

In certain embodiments, the compounds targeted to apo(a) described herein modulate physiological markers or phenotypes of the inflammatory disease, disorder or condition. For example, administration of the compounds to animals can decrease inflammatory cytokine or other inflammatory markers levels in those animals compared to untreated animals. In certain embodiments, the modulation of the physiological markers or phenotypes can be associated with inhibition of apo(a) by the compounds.

In certain embodiments, the physiological markers of the inflammatory disease, disorder or condition can be quantifiable. For example, cytokine levels can be measured and quantified by standard tests known in the art. For such markers, in certain embodiments, the marker can be decreased by at least about 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95% or 99%, or a range defined by any two of these values.

Also, provided herein are methods for preventing, treating or ameliorating a symptom associated with the inflammatory disease, disorder or condition in a subject in need thereof. In certain embodiments, provided is a method for reducing the rate of onset of a symptom associated with the inflammatory disease, disorder or condition. In certain embodiments, provided is a method for reducing the severity of a symptom associated with the inflammatory disease, disorder or condition. In such embodiments, the methods comprise administering a therapeutically effective amount of a compound targeted to an apo(a) nucleic acid to an individual in need thereof.

In certain embodiments, provided are methods of treating an individual with an apo(a) related disease, disorder or condition comprising administering a therapeutically effective amount of one or more pharmaceutical compositions as described herein. In certain embodiments, the individual has elevated apo(a) levels. In certain embodiments, provided are methods of treating an individual with an Lp(a) related disease, disorder or condition comprising administering a therapeutically effective amount of one or more pharmaceutical compositions as described herein. In certain embodiments, the individual has elevated Lp(a)

levels. In certain embodiments, the individual has an inflammatory, cardiovascular and/or metabolic disease, disorder or condition. In certain embodiments, administration of a therapeutically effective amount of an antisense compound targeted to an apo(a) nucleic acid is accompanied by monitoring of apo(a) or Lp(a) levels. In certain embodiments, administration of a therapeutically effective amount of an antisense compound targeted to an apo(a) nucleic acid is accompanied by monitoring of markers of inflammatory, cardiovascular and/or metabolic disease, or other disease process associated with the expression of apo(a), to determine an individual's response to the antisense compound. An individual's response to administration of the antisense compound targeting apo(a) can be used by a physician to determine the amount and duration of therapeutic intervention with the compound.

In certain embodiments, administration of an antisense compound targeted to an apo(a) nucleic acid results in reduction of apo(a) expression by at least about 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95% or 99%, or a range defined by any two of these values. In certain embodiments, apo(a) expression is reduced to at least ≤ 100 mg/dL, ≤ 90 mg/dL, ≤ 80 mg/dL, ≤ 70 mg/dL, ≤ 60 mg/dL, ≤ 50 mg/dL, ≤ 40 mg/dL, ≤ 30 mg/dL, ≤ 20 mg/dL or ≤ 10 mg/dL.

In certain embodiments, administration of an antisense compound targeted to an apo(a) nucleic acid results in reduction of Lp(a) expression by at least about 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95% or 99%, or a range defined by any two of these values. In certain embodiments, Lp(a) expression is reduced to at least ≤ 200 mg/dL, ≤ 190 mg/dL, ≤ 180 mg/dL, ≤ 175 mg/dL, ≤ 170 mg/dL, ≤ 160 mg/dL, ≤ 150 mg/dL, ≤ 140 mg/dL, ≤ 130 mg/dL, ≤ 120 mg/dL, ≤ 110 mg/dL, ≤ 100 mg/dL, ≤ 90 mg/dL, ≤ 80 mg/dL, ≤ 70 mg/dL, ≤ 60 mg/dL, ≤ 55 mg/dL, ≤ 50 mg/dL, ≤ 45 mg/dL, ≤ 40 mg/dL, ≤ 35 mg/dL, ≤ 30 mg/dL, ≤ 25 mg/dL, ≤ 20 mg/dL, ≤ 15 mg/dL, or ≤ 10 mg/dL.

In certain embodiments, the invention provides methods for using a conjugated antisense compound targeted to an apo(a) nucleic acid in the preparation of a medicament. In certain embodiments, pharmaceutical compositions comprising a conjugated antisense compound targeted to apo(a) are used for the preparation of a medicament for treating a patient suffering or susceptible to an inflammatory, cardiovascular and/or a metabolic disease, disorder or condition.

Apo(a) Treatment Populations

Certain subjects with high Lp(a) levels are at a significant risk of various diseases (Lippi et al., Clinica Chimica Acta, 2011, 412:797-801; Solfrizz et al.). In many subjects with high Lp(a) levels, current treatments cannot reduce their Lp(a) levels to safe levels. Apo(a) plays an important role in the formation of Lp(a), hence reducing apo(a) can reduce Lp(a) and prevent, treat or ameliorate a disease associated with Lp(a).

In certain embodiments, treatment with the compounds and methods disclosed herein is indicated for a human animal with elevated apo(a) levels and/or Lp(a) levels. In certain embodiments, the human has

apo(a) levels ≥ 10 mg/dL, ≥ 20 mg/dL, ≥ 30 mg/dL, ≥ 40 mg/dL, ≥ 50 mg/dL, ≥ 60 mg/dL, ≥ 70 mg/dL, ≥ 80 mg/dL, ≥ 90 mg/dL or ≥ 100 mg/dL. In certain embodiments, the human has Lp(a) levels ≥ 10 mg/dL, ≥ 15 mg/dL, ≥ 20 mg/dL, ≥ 25 mg/dL, ≥ 30 mg/dL, ≥ 35 mg/dL, ≥ 40 mg/dL, ≥ 50 mg/dL, ≥ 60 mg/dL, ≥ 70 mg/dL, ≥ 80 mg/dL, ≥ 90 mg/dL, ≥ 100 mg/dL, ≥ 110 mg/dL, ≥ 120 mg/dL, ≥ 130 mg/dL, ≥ 140 mg/dL, ≥ 150 mg/dL, ≥ 160 mg/dL, ≥ 170 mg/dL, ≥ 175 mg/dL, ≥ 180 mg/dL, ≥ 190 mg/dL, ≥ 200 mg/dL.

D. Certain Pharmaceutical Compositions

In certain embodiments, the present disclosure provides pharmaceutical compositions comprising one or more antisense compound. In certain embodiments, such pharmaceutical composition comprises a suitable pharmaceutically acceptable diluent or carrier. In certain embodiments, a pharmaceutical composition comprises a sterile saline solution and one or more antisense compound. In certain embodiments, such pharmaceutical composition consists of a sterile saline solution and one or more antisense compound. In certain embodiments, the sterile saline is pharmaceutical grade saline. In certain embodiments, a pharmaceutical composition comprises one or more antisense compound and sterile water. In certain embodiments, a pharmaceutical composition consists of one or more antisense compound and sterile water. In certain embodiments, the sterile saline is pharmaceutical grade water. In certain embodiments, a pharmaceutical composition comprises one or more antisense compound and phosphate-buffered saline (PBS). In certain embodiments, a pharmaceutical composition consists of one or more antisense compound and sterile phosphate-buffered saline (PBS). In certain embodiments, the sterile saline is pharmaceutical grade PBS.

In certain embodiments, antisense compounds may be admixed with pharmaceutically acceptable active and/or inert substances for the preparation of pharmaceutical compositions or formulations. Compositions and methods for the formulation of pharmaceutical compositions depend on a number of criteria, including, but not limited to, route of administration, extent of disease, or dose to be administered.

Pharmaceutical compositions comprising antisense compounds encompass any pharmaceutically acceptable salts, esters, or salts of such esters. In certain embodiments, pharmaceutical compositions comprising antisense compounds comprise one or more oligonucleotide which, upon administration to an animal, including a human, is capable of providing (directly or indirectly) the biologically active metabolite or residue thereof. Accordingly, for example, the disclosure is also drawn to pharmaceutically acceptable salts of antisense compounds, prodrugs, pharmaceutically acceptable salts of such prodrugs, and other bioequivalents. Suitable pharmaceutically acceptable salts include, but are not limited to, sodium and potassium salts.

A prodrug can include the incorporation of additional nucleosides at one or both ends of an oligonucleotide which are cleaved by endogenous nucleases within the body, to form the active antisense oligonucleotide.

Lipid moieties have been used in nucleic acid therapies in a variety of methods. In certain such methods, the nucleic acid is introduced into preformed liposomes or lipoplexes made of mixtures of cationic lipids and neutral lipids. In certain methods, DNA complexes with mono- or poly-cationic lipids are formed without the presence of a neutral lipid. In certain embodiments, a lipid moiety is selected to increase
5 distribution of a pharmaceutical agent to a particular cell or tissue. In certain embodiments, a lipid moiety is selected to increase distribution of a pharmaceutical agent to fat tissue. In certain embodiments, a lipid moiety is selected to increase distribution of a pharmaceutical agent to muscle tissue.

In certain embodiments, pharmaceutical compositions provided herein comprise one or more modified oligonucleotides and one or more excipients. In certain such embodiments, excipients are selected
10 from water, salt solutions, alcohol, polyethylene glycols, gelatin, lactose, amylase, magnesium stearate, talc, silicic acid, viscous paraffin, hydroxymethylcellulose and polyvinylpyrrolidone.

In certain embodiments, a pharmaceutical composition provided herein comprises a delivery system. Examples of delivery systems include, but are not limited to, liposomes and emulsions. Certain delivery systems are useful for preparing certain pharmaceutical compositions including those comprising
15 hydrophobic compounds. In certain embodiments, certain organic solvents such as dimethylsulfoxide are used.

In certain embodiments, a pharmaceutical composition provided herein comprises one or more tissue-specific delivery molecules designed to deliver the one or more pharmaceutical agents of the present disclosure to specific tissues or cell types. For example, in certain embodiments, pharmaceutical
20 compositions include liposomes coated with a tissue-specific antibody.

In certain embodiments, a pharmaceutical composition provided herein comprises a co-solvent system. Certain of such co-solvent systems comprise, for example, benzyl alcohol, a nonpolar surfactant, a water-miscible organic polymer, and an aqueous phase. In certain embodiments, such co-solvent systems are used for hydrophobic compounds. A non-limiting example of such a co-solvent system is the VPD co-solvent
25 system, which is a solution of absolute ethanol comprising 3% w/v benzyl alcohol, 8% w/v of the nonpolar surfactant Polysorbate 80™ and 65% w/v polyethylene glycol 300. The proportions of such co-solvent systems may be varied considerably without significantly altering their solubility and toxicity characteristics. Furthermore, the identity of co-solvent components may be varied: for example, other surfactants may be used instead of Polysorbate 80™; the fraction size of polyethylene glycol may be varied; other biocompatible
30 polymers may replace polyethylene glycol, e.g., polyvinyl pyrrolidone; and other sugars or polysaccharides may substitute for dextrose.

In certain embodiments, a pharmaceutical composition provided herein is prepared for oral administration. In certain embodiments, pharmaceutical compositions are prepared for buccal administration.

In certain embodiments, a pharmaceutical composition is prepared for administration by injection
35 (e.g., intravenous, subcutaneous, intramuscular, etc.). In certain of such embodiments, a pharmaceutical composition comprises a carrier and is formulated in aqueous solution, such as water or physiologically

compatible buffers such as Hanks's solution, Ringer's solution, or physiological saline buffer. In certain embodiments, other ingredients are included (e.g., ingredients that aid in solubility or serve as preservatives). In certain embodiments, injectable suspensions are prepared using appropriate liquid carriers, suspending agents and the like. Certain pharmaceutical compositions for injection are presented in unit dosage form, e.g.,
5 in ampoules or in multi-dose containers. Certain pharmaceutical compositions for injection are suspensions, solutions or emulsions in oily or aqueous vehicles, and may contain formulatory agents such as suspending, stabilizing and/or dispersing agents. Certain solvents suitable for use in pharmaceutical compositions for injection include, but are not limited to, lipophilic solvents and fatty oils, such as sesame oil, synthetic fatty acid esters, such as ethyl oleate or triglycerides, and liposomes. Aqueous injection suspensions may contain
10 substances that increase the viscosity of the suspension, such as sodium carboxymethyl cellulose, sorbitol, or dextran. Optionally, such suspensions may also contain suitable stabilizers or agents that increase the solubility of the pharmaceutical agents to allow for the preparation of highly concentrated solutions.

In certain embodiments, a pharmaceutical composition is prepared for transmucosal administration. In certain of such embodiments penetrants appropriate to the barrier to be permeated are used in the
15 formulation. Such penetrants are generally known in the art.

In certain embodiments, a pharmaceutical composition provided herein comprises an oligonucleotide in a therapeutically effective amount. In certain embodiments, the therapeutically effective amount is sufficient to prevent, alleviate or ameliorate symptoms of a disease or to prolong the survival of the subject being treated. Determination of a therapeutically effective amount is well within the capability of those
20 skilled in the art.

In certain embodiments, one or more modified oligonucleotide provided herein is formulated as a prodrug. In certain embodiments, upon *in vivo* administration, a prodrug is chemically converted to the biologically, pharmaceutically or therapeutically more active form of an oligonucleotide. In certain
25 embodiments, prodrugs are useful because they are easier to administer than the corresponding active form. For example, in certain instances, a prodrug may be more bioavailable (e.g., through oral administration) than is the corresponding active form. In certain instances, a prodrug may have improved solubility compared to the corresponding active form. In certain embodiments, prodrugs are less water soluble than the corresponding active form. In certain instances, such prodrugs possess superior transmittal across cell
30 membranes, where water solubility is detrimental to mobility. In certain embodiments, a prodrug is an ester. In certain such embodiments, the ester is metabolically hydrolyzed to carboxylic acid upon administration. In certain instances the carboxylic acid containing compound is the corresponding active form. In certain embodiments, a prodrug comprises a short peptide (polyaminoacid) bound to an acid group. In certain of such embodiments, the peptide is cleaved upon administration to form the corresponding active form.

In certain embodiments, the present disclosure provides compositions and methods for reducing the
35 amount or activity of a target nucleic acid in a cell. In certain embodiments, the cell is in an animal. In certain embodiments, the animal is a mammal. In certain embodiments, the animal is a rodent. In certain

embodiments, the animal is a primate. In certain embodiments, the animal is a non-human primate. In certain embodiments, the animal is a human.

In certain embodiments, the present disclosure provides methods of administering a pharmaceutical composition comprising an oligonucleotide of the present disclosure to an animal. Suitable administration routes include, but are not limited to, oral, rectal, transmucosal, intestinal, enteral, topical, suppository, through inhalation, intrathecal, intracerebroventricular, intraperitoneal, intranasal, intraocular, intratumoral, and parenteral (e.g., intravenous, intramuscular, intramedullary, and subcutaneous). In certain embodiments, pharmaceutical intrathecal are administered to achieve local rather than systemic exposures. For example, pharmaceutical compositions may be injected directly in the area of desired effect (e.g., into the liver).

Nonlimiting disclosure and incorporation by reference

While certain compounds, compositions and methods described herein have been described with specificity in accordance with certain embodiments, the following examples serve only to illustrate the compounds described herein and are not intended to limit the same. Each of the references, GenBank accession numbers, and the like recited in the present application is incorporated herein by reference in its entirety.

Although the sequence listing accompanying this filing identifies each sequence as either “RNA” or “DNA” as required, in reality, those sequences may be modified with any combination of chemical modifications. One of skill in the art will readily appreciate that such designation as “RNA” or “DNA” to describe modified oligonucleotides is, in certain instances, arbitrary. For example, an oligonucleotide comprising a nucleoside comprising a 2'-OH sugar moiety and a thymine base could be described as a DNA having a modified sugar (2'-OH for the natural 2'-H of DNA) or as an RNA having a modified base (thymine (methylated uracil) for natural uracil of RNA).

Accordingly, nucleic acid sequences provided herein, including, but not limited to those in the sequence listing, are intended to encompass nucleic acids containing any combination of natural or modified RNA and/or DNA, including, but not limited to such nucleic acids having modified nucleobases. By way of further example and without limitation, an oligonucleotide having the nucleobase sequence “ATCGATCG” encompasses any oligonucleotides having such nucleobase sequence, whether modified or unmodified, including, but not limited to, such compounds comprising RNA bases, such as those having sequence “AUCGAUCG” and those having some DNA bases and some RNA bases such as “AUCGATCG” and oligonucleotides having other modified bases, such as “AT^{me}CGAUCG,” wherein ^{me}C indicates a cytosine base comprising a methyl group at the 5-position.

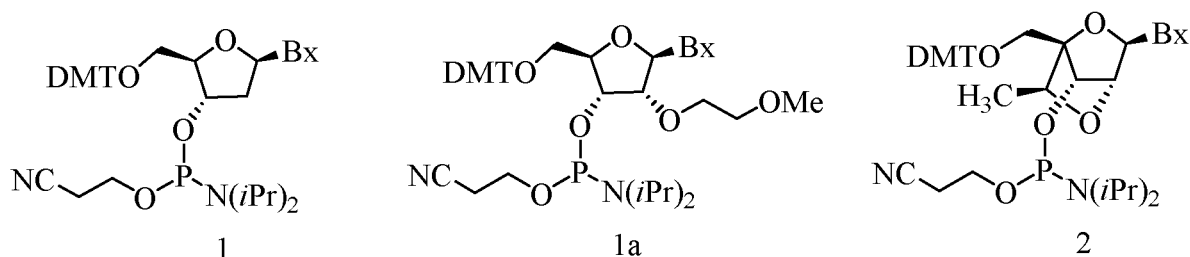
EXAMPLES

The following examples illustrate certain embodiments of the present disclosure and are not limiting. Moreover, where specific embodiments are provided, the inventors have contemplated generic application of

those specific embodiments. For example, disclosure of an oligonucleotide having a particular motif provides reasonable support for additional oligonucleotides having the same or similar motif. And, for example, where a particular high-affinity modification appears at a particular position, other high-affinity modifications at the same position are considered suitable, unless otherwise indicated.

5

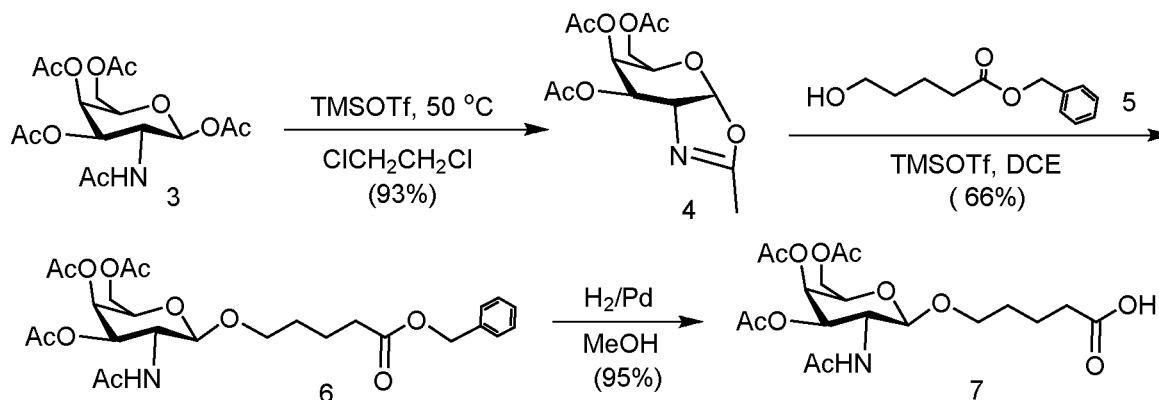
Example 1: General Method for the Preparation of Phosphoramidites, Compounds 1, 1a and 2



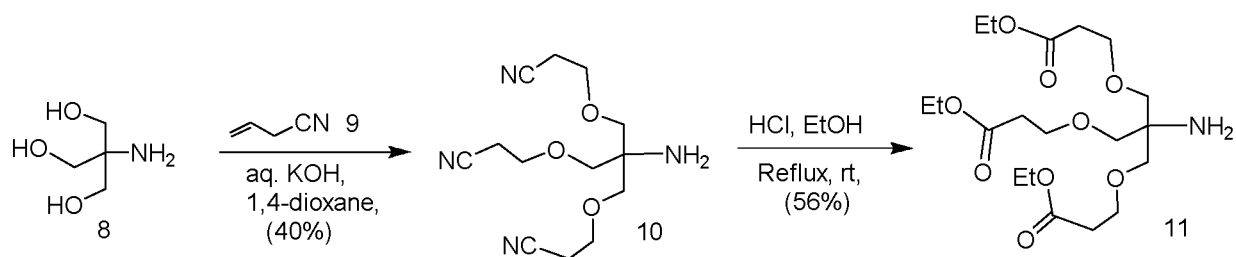
Bx is a heterocyclic base;

Compounds 1, 1a and 2 were prepared as per the procedures well known in the art as described in the specification herein (see Seth et al., *Bioorg. Med. Chem.*, 2011, 21(4), 1122-1125, *J. Org. Chem.*, 2010, 75(5), 1569-1581, *Nucleic Acids Symposium Series*, 2008, 52(1), 553-554); and also see published PCT International Applications (WO 2011/115818, WO 2010/077578, WO2010/036698, WO2009/143369, WO 2009/006478, and WO 2007/090071), and US patent 7,569,686).

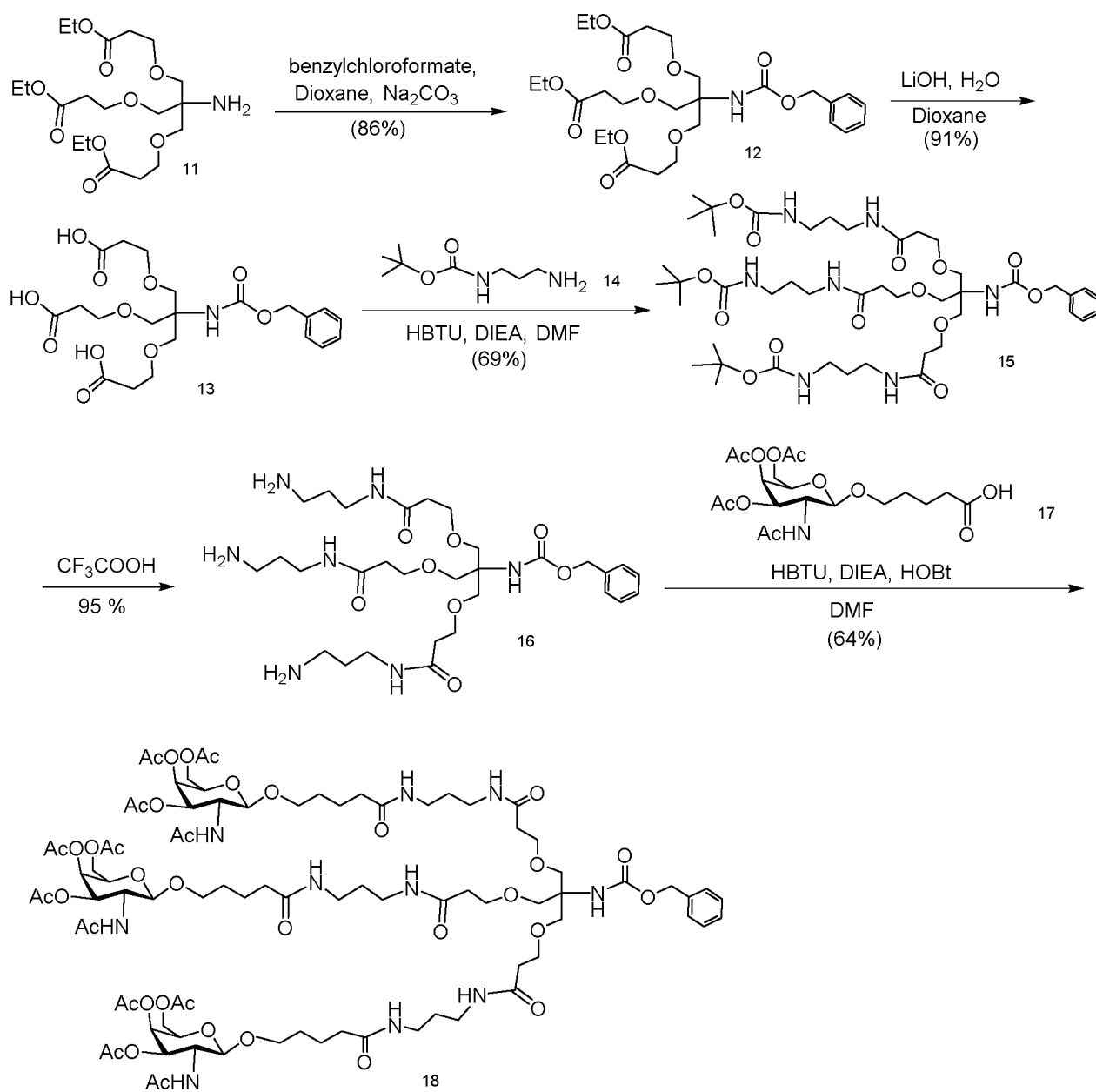
Example 2: Preparation of Compound 7



Compounds 3 (2-acetamido-1,3,4,6-tetra-*O*-acetyl-2-deoxy- β -Dgalactopyranose or galactosamine pentaacetate) is commercially available. Compound 5 was prepared according to published procedures (Weber *et al.*, *J. Med. Chem.*, 1991, 34, 2692).

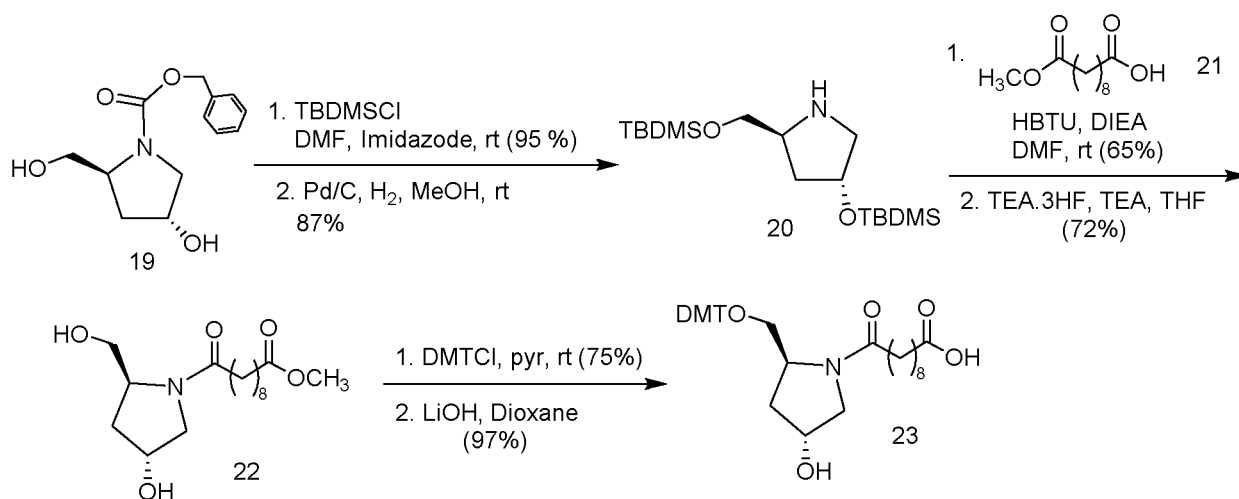
Example 3: Preparation of Compound 11

Compounds 8 and 9 are commercially available.

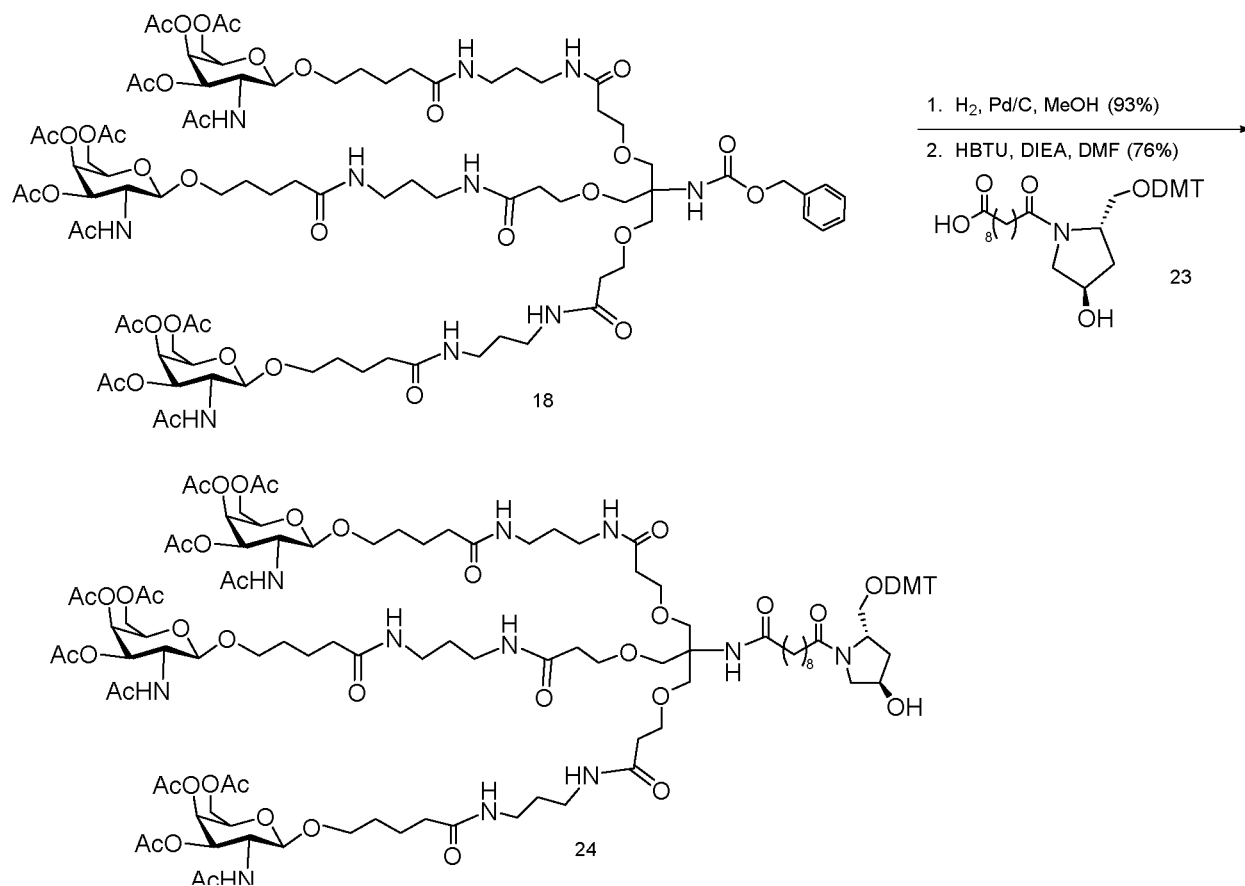
Example 4: Preparation of Compound 18

Compound 11 was prepared as per the procedures illustrated in Example 3. Compound 14 is commercially available. Compound 17 was prepared using similar procedures reported by Rensen *et al.*, *J. Med. Chem.*, 2004, 47, 5798-5808.

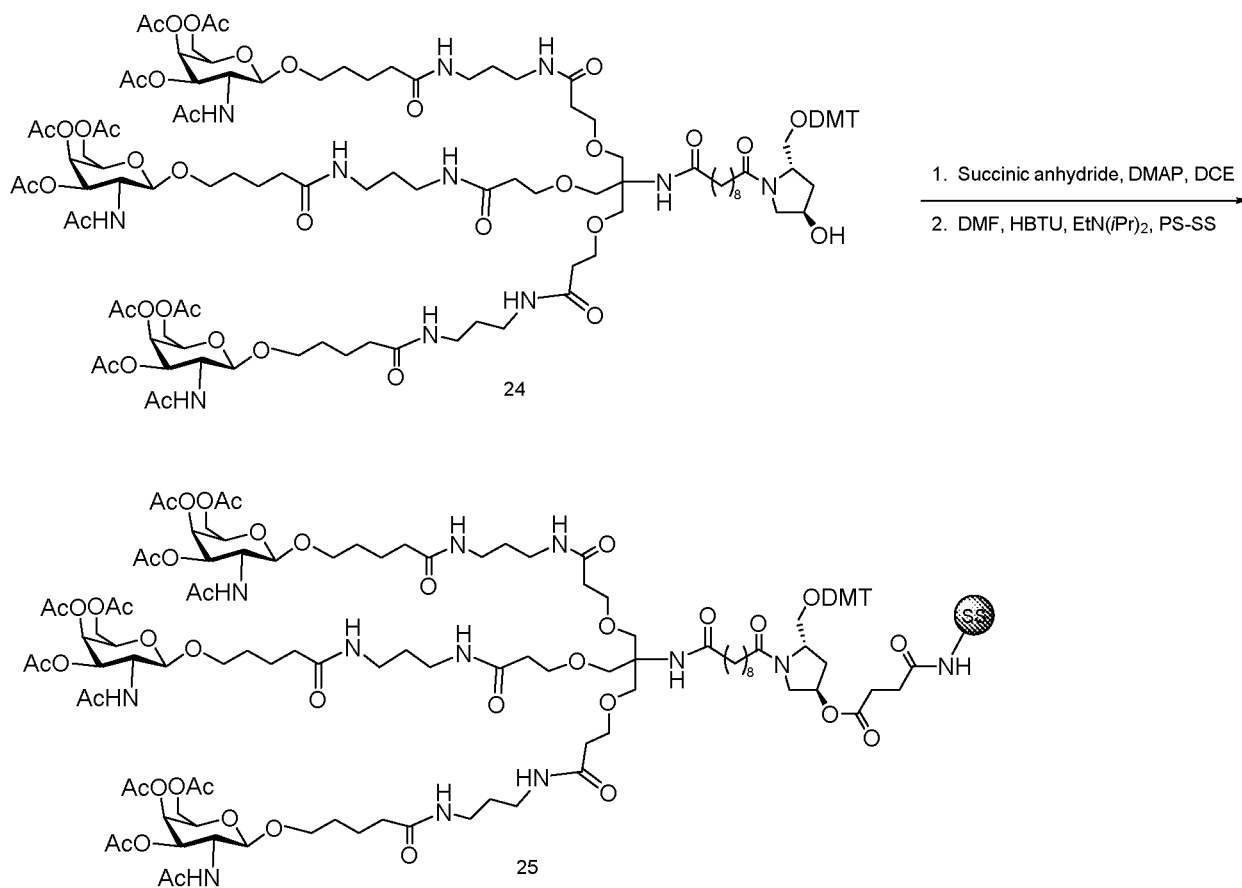
5 Example 5: Preparation of Compound 23



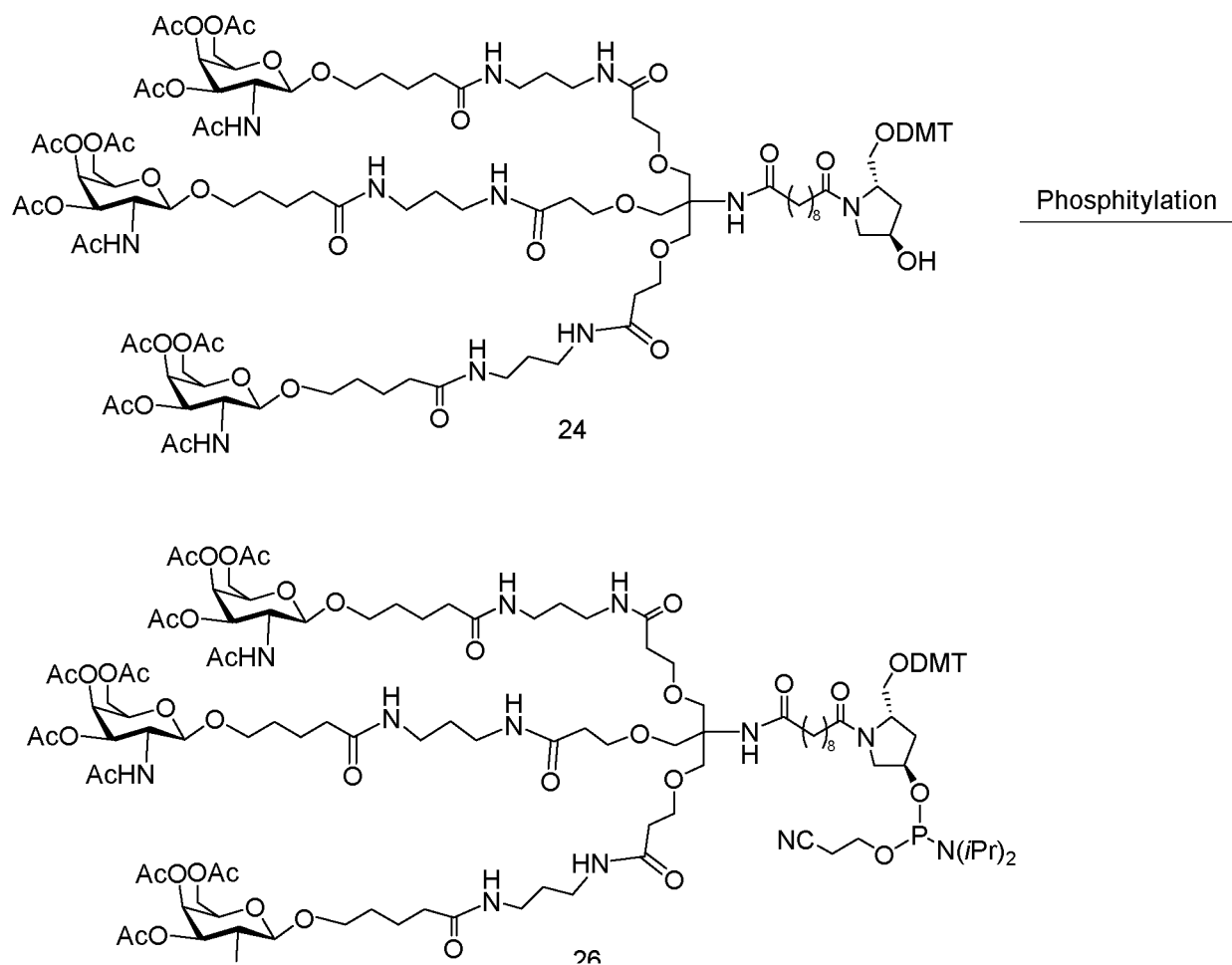
Compounds 19 and 21 are commercially available.

Example 6: Preparation of Compound 24

Compounds 18 and 23 were prepared as per the procedures illustrated in Examples 4 and 5.

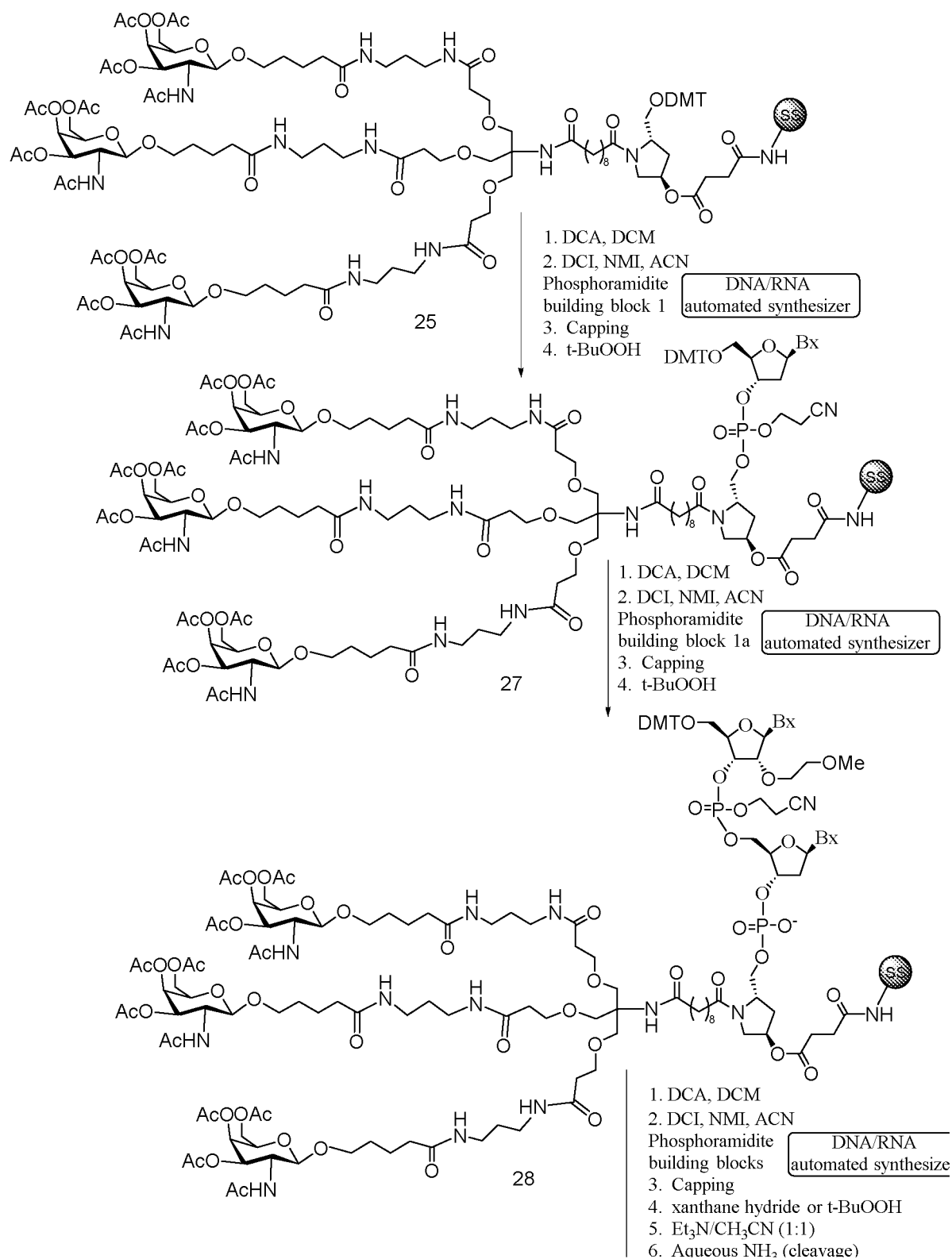
Example 7: Preparation of Compound 25

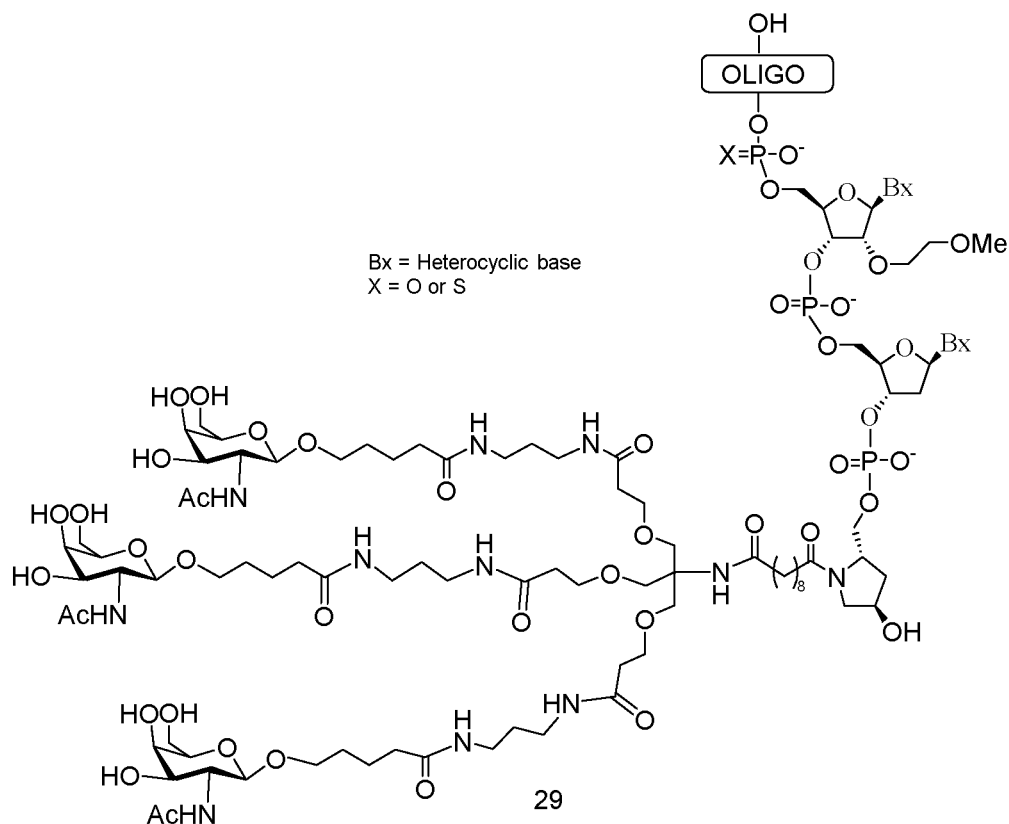
Compound 24 was prepared as per the procedures illustrated in Example 6.

Example 8: Preparation of Compound 26

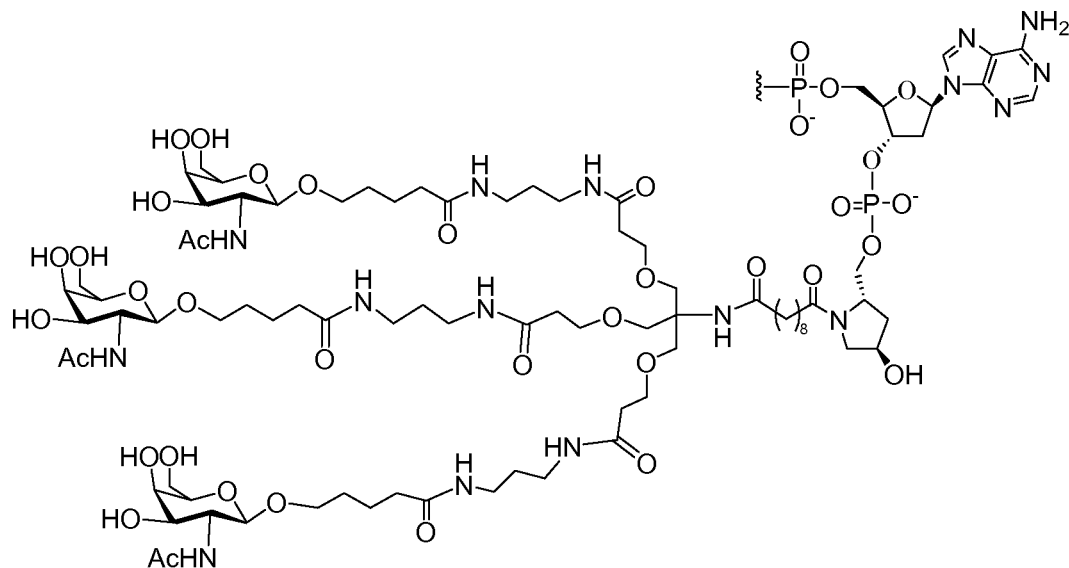
Compound 24 is prepared as per the procedures illustrated in Example 6.

**Example 9: General preparation of conjugated ASOs comprising GalNAc₃-1 at the 3' terminus,
Compound 29**

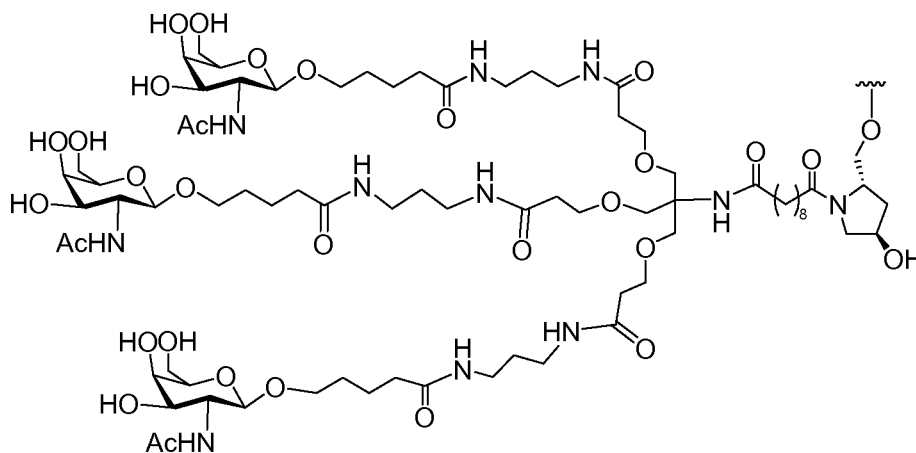




Wherein the protected **GalNAc₃-1** has the structure:

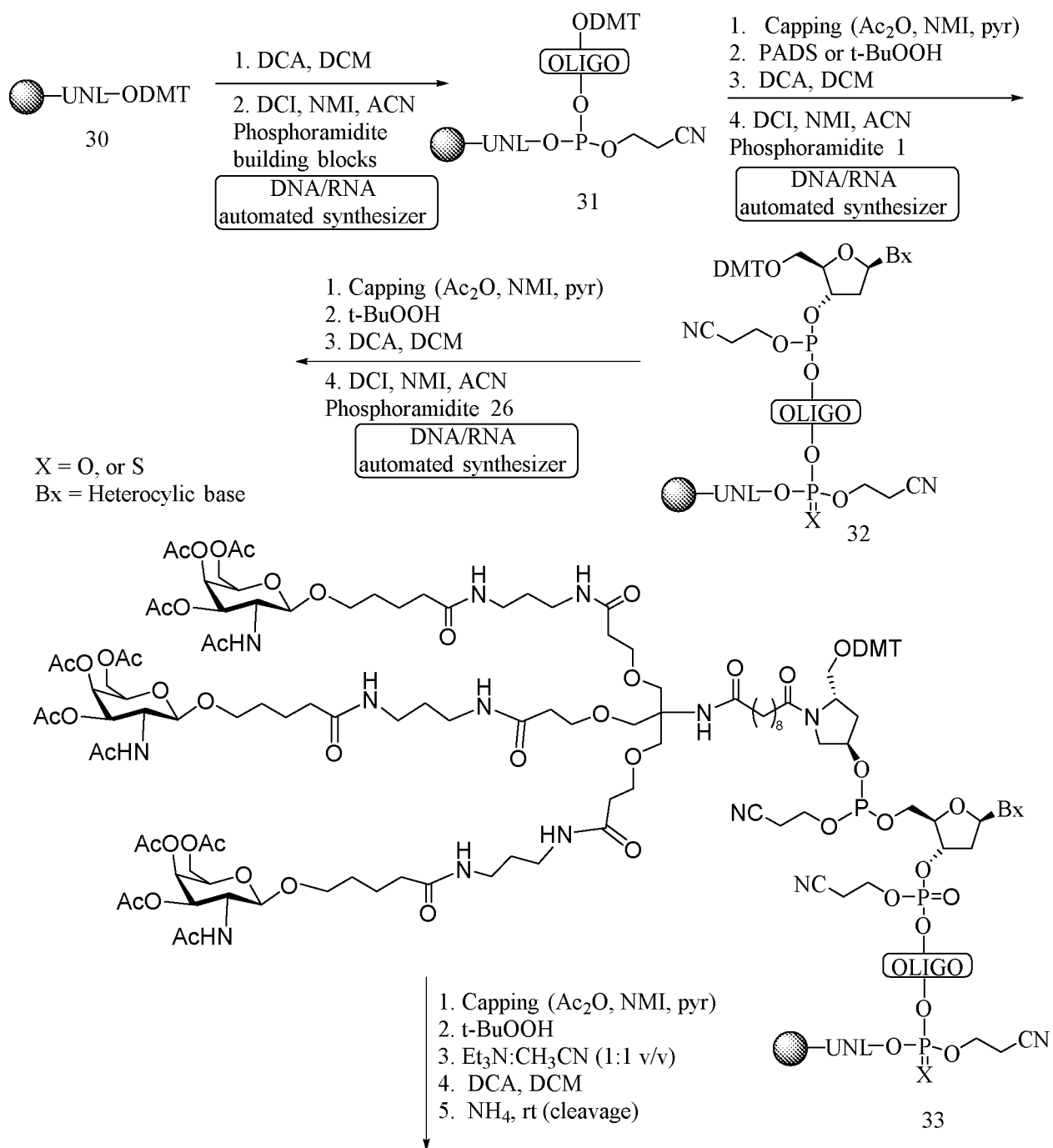


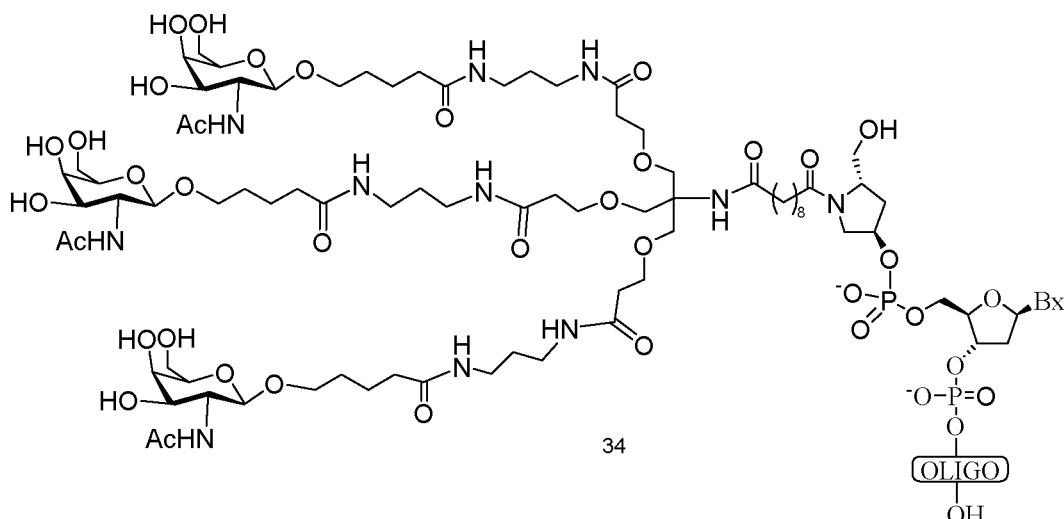
The GalNAc₃ cluster portion of the conjugate group GalNAc₃-1 (GalNAc₃-1_a) can be combined with
 5 any cleavable moiety to provide a variety of conjugate groups. Wherein GalNAc₃-1_a has the formula:



The solid support bound protected **GalNAc₃-1**, Compound 25, was prepared as per the procedures illustrated in Example 7. Oligomeric Compound 29 comprising **GalNAc₃-1** at the 3' terminus was prepared using standard procedures in automated DNA/RNA synthesis (see Dupouy *et al.*, *Angew. Chem. Int. Ed.*, 2006, 45, 3623-3627). Phosphoramidite building blocks, Compounds 1 and 1a were prepared as per the procedures illustrated in Example 1. The phosphoramidites illustrated are meant to be representative and not intended to be limiting as other phosphoramidite building blocks can be used to prepare oligomeric compounds having a predetermined sequence and composition. The order and quantity of phosphoramidites added to the solid support can be adjusted to prepare gapped oligomeric compounds as described herein. Such gapped oligomeric compounds can have predetermined composition and base sequence as dictated by any given target.

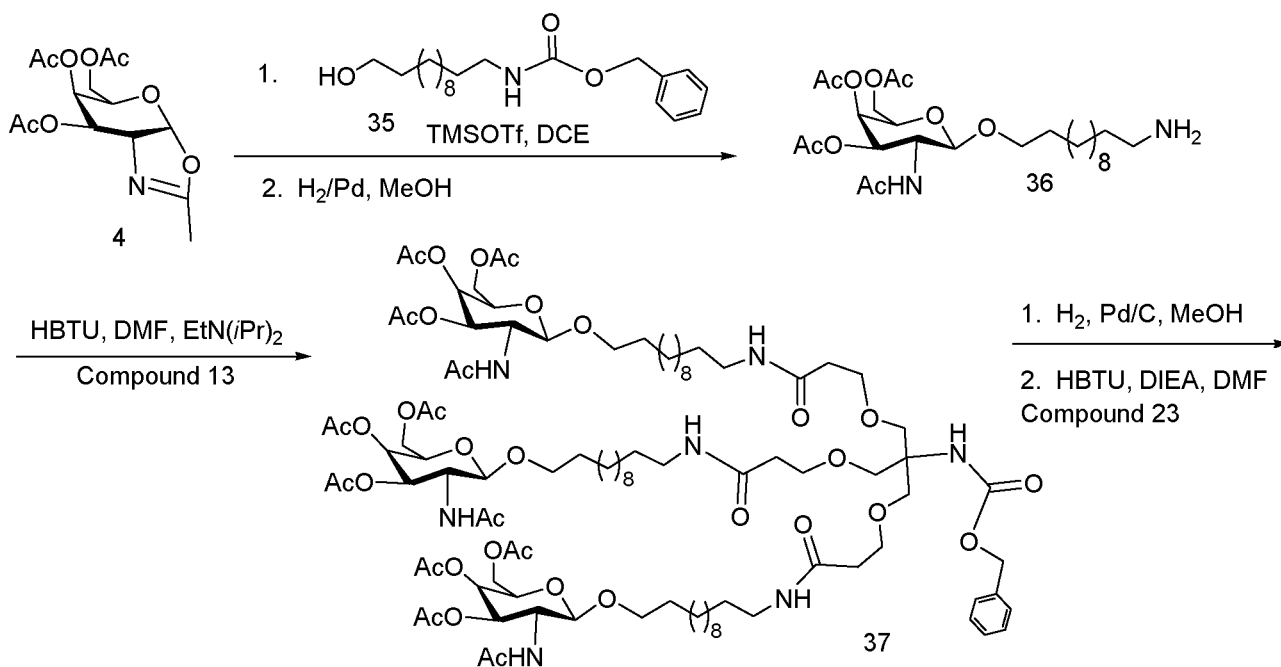
Example 10: General preparation conjugated ASOs comprising GalNAc₃-1 at the 5' terminus, Compound 34

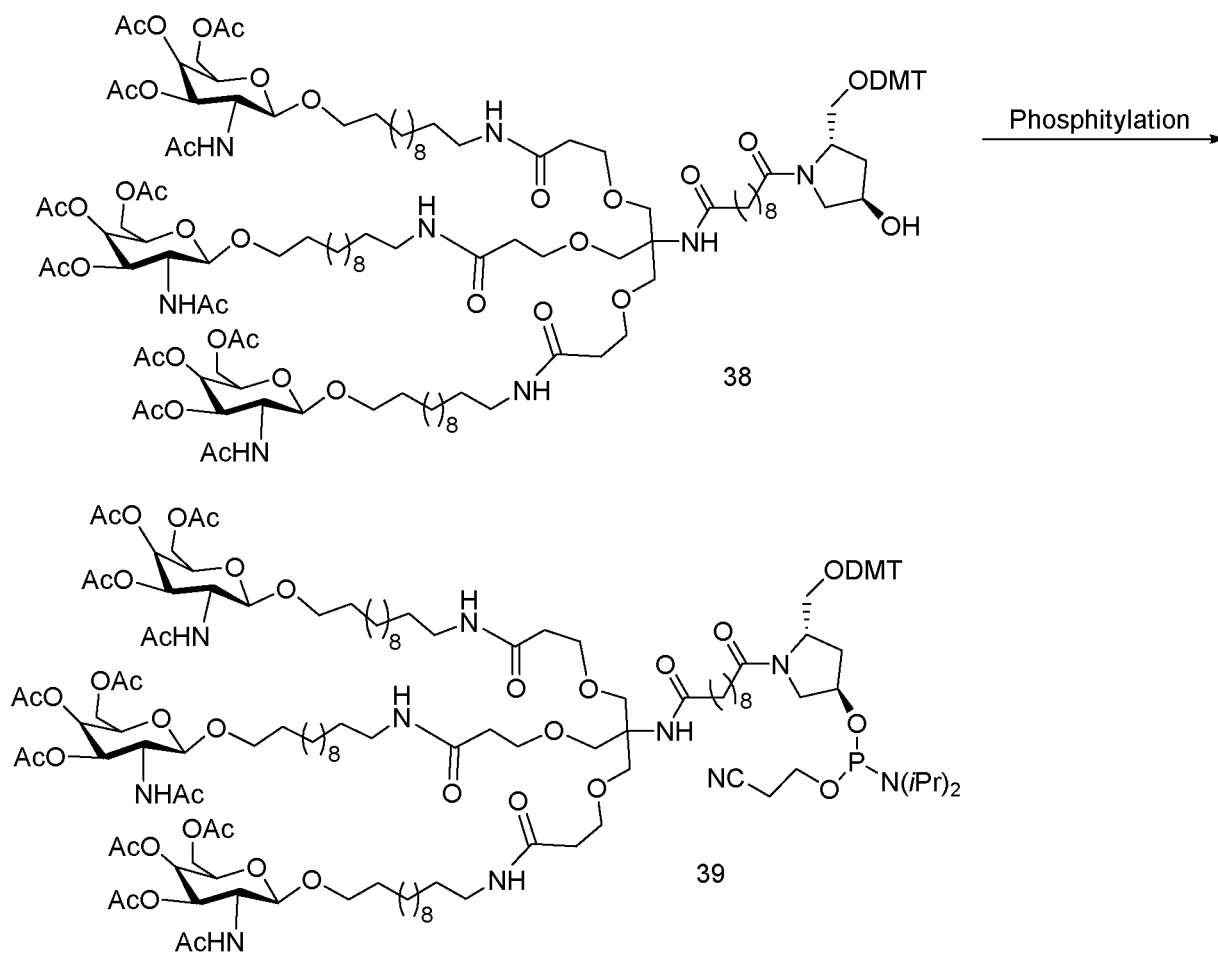




The UnylinkerTM 30 is commercially available. Oligomeric Compound 34 comprising a **GalNAc₃-1** cluster at the 5' terminus is prepared using standard procedures in automated DNA/RNA synthesis (see Dupouy *et al.*, *Angew. Chem. Int. Ed.*, 2006, 45, 3623-3627). Phosphoramidite building blocks, Compounds 1 and 1a were prepared as per the procedures illustrated in Example 1. The phosphoramidites illustrated are meant to be representative and not intended to be limiting as other phosphoramidite building blocks can be used to prepare an oligomeric compound having a predetermined sequence and composition. The order and quantity of phosphoramidites added to the solid support can be adjusted to prepare gapped oligomeric compounds as described herein. Such gapped oligomeric compounds can have predetermined composition and base sequence as dictated by any given target.

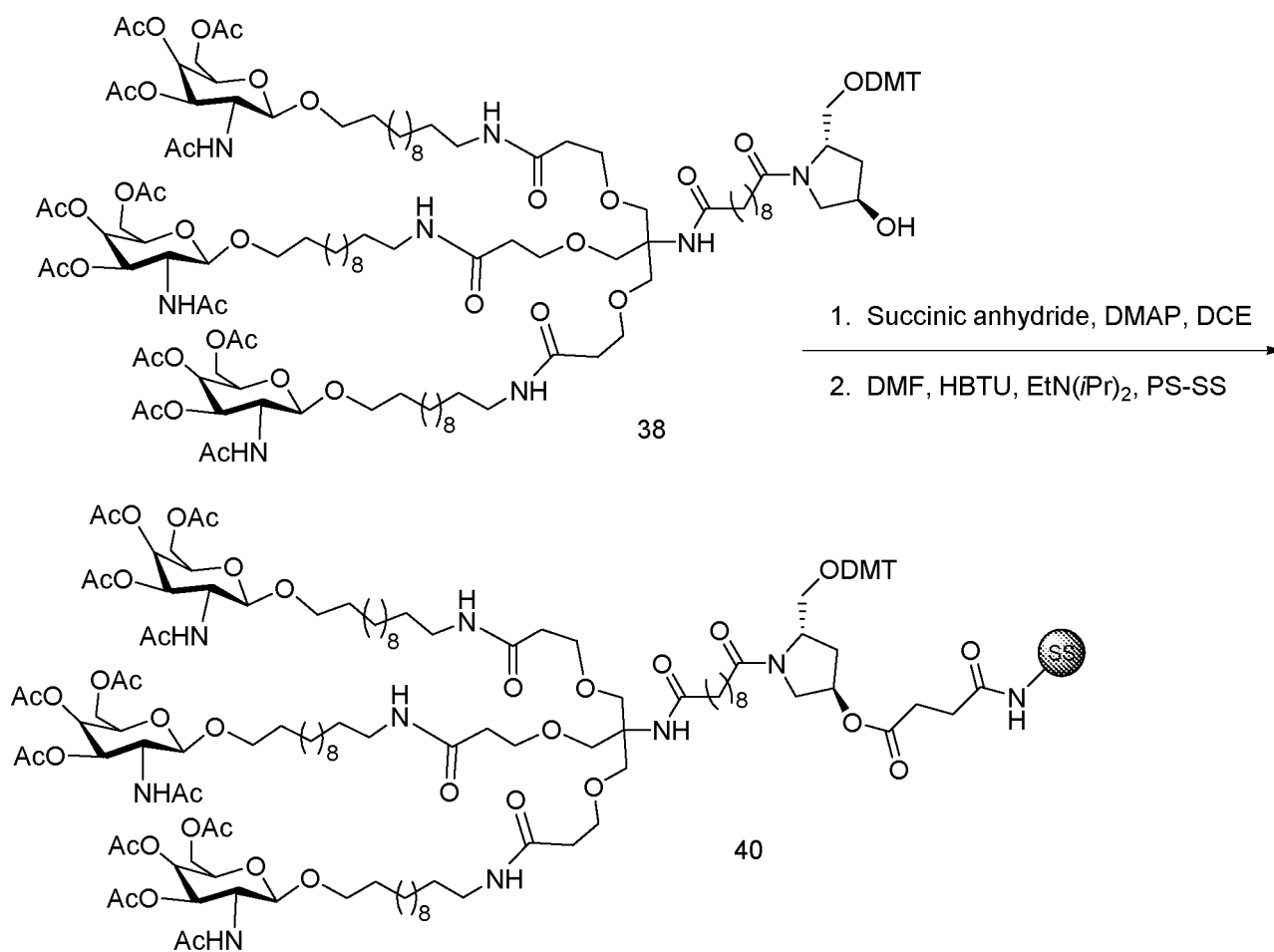
Example 11: Preparation of Compound 39





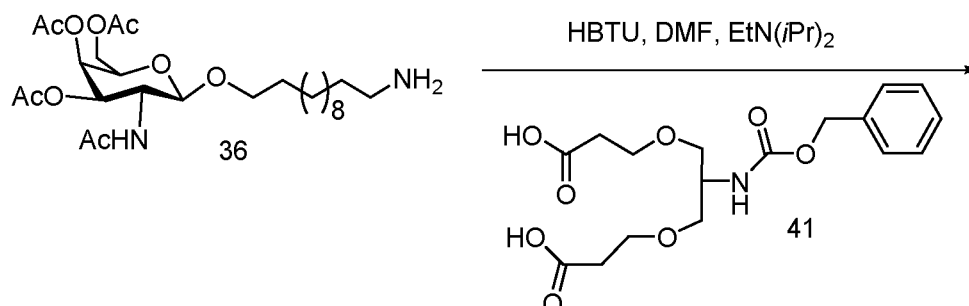
Compounds 4, 13 and 23 were prepared as per the procedures illustrated in Examples 2, 4, and 5. Compound 35 is prepared using similar procedures published in Rouchaud *et al.*, *Eur. J. Org. Chem.*, 2011, 12, 2346-2353.

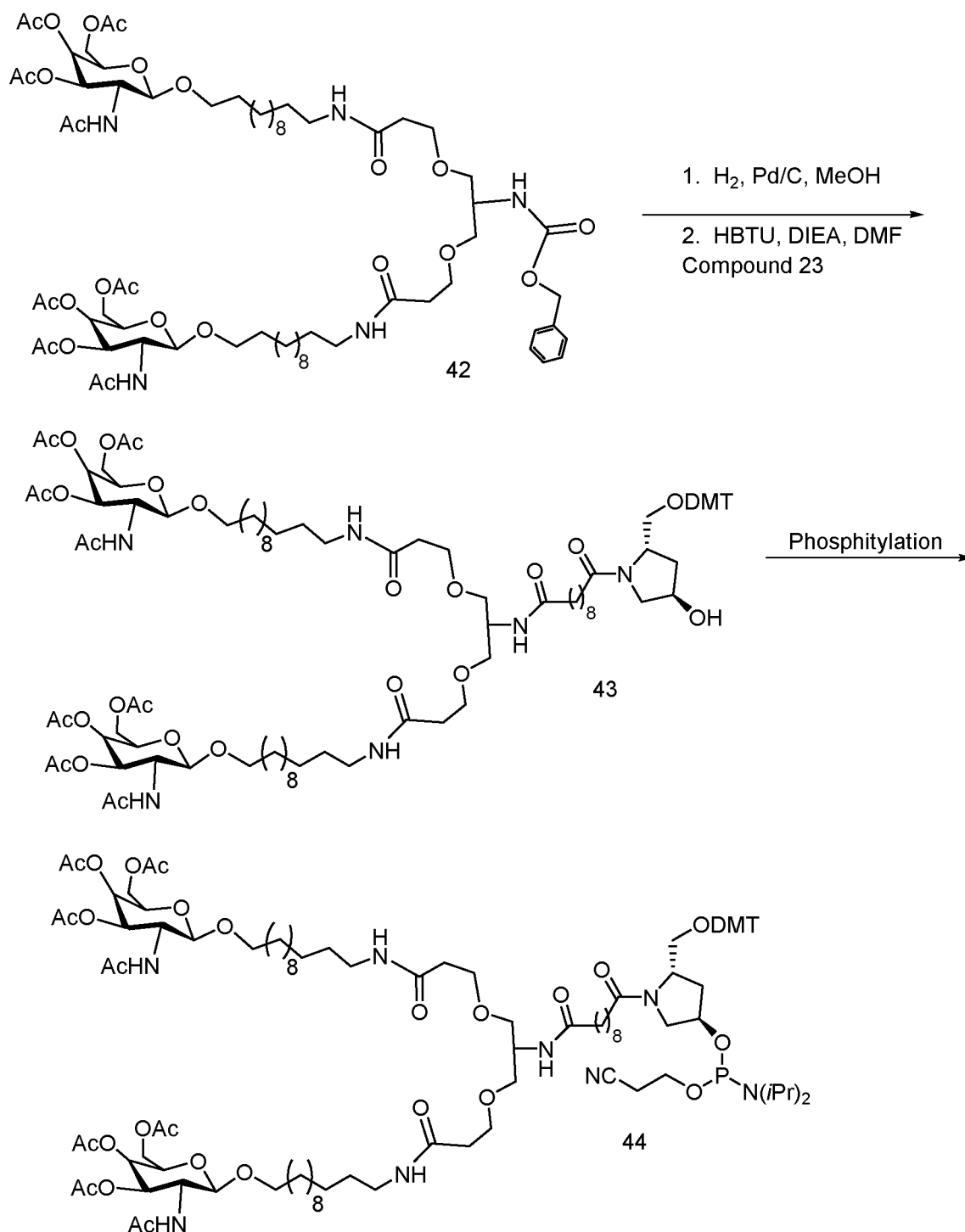
Example 12: Preparation of Compound 40



Compound 38 is prepared as per the procedures illustrated in Example 11.

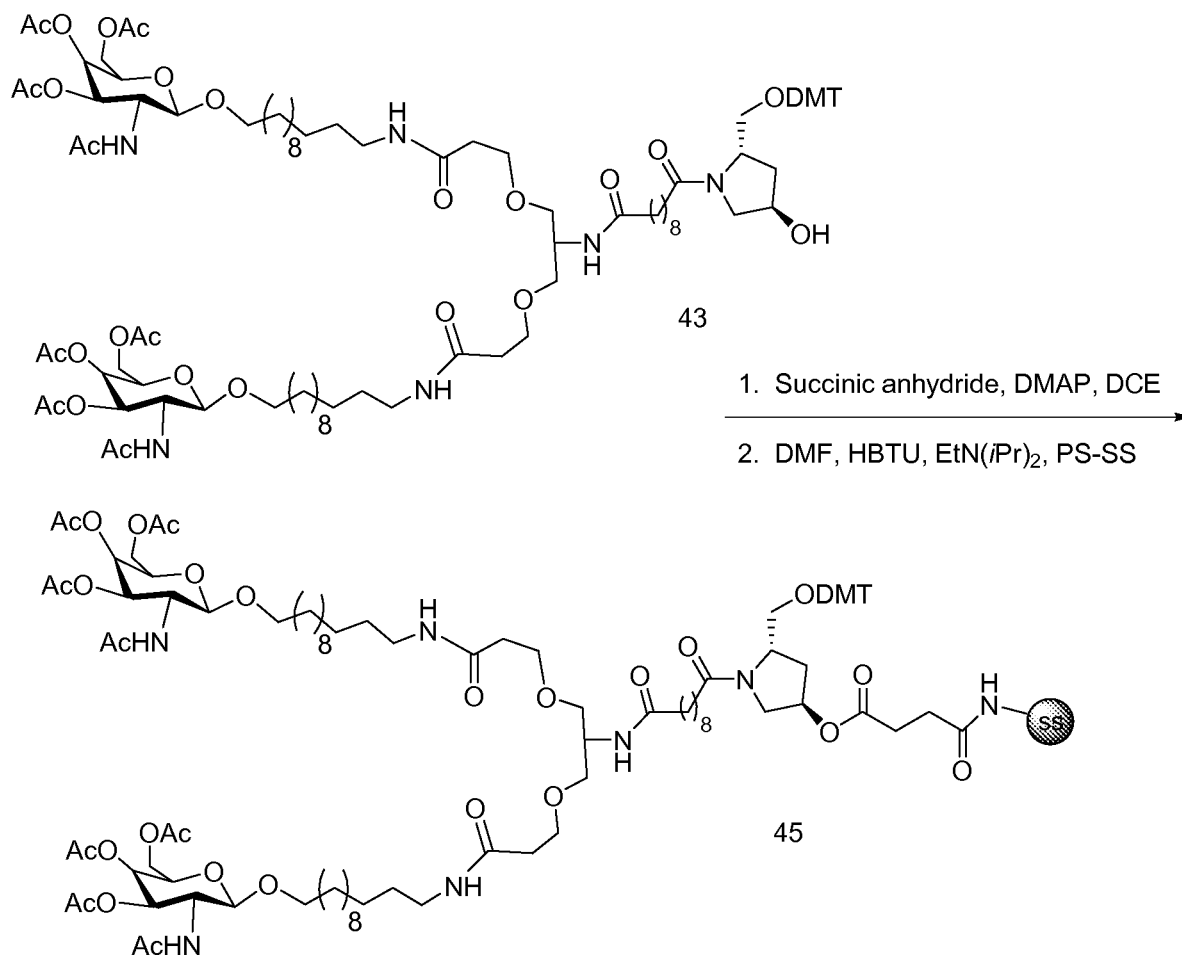
5 Example 13: Preparation of Compound 44





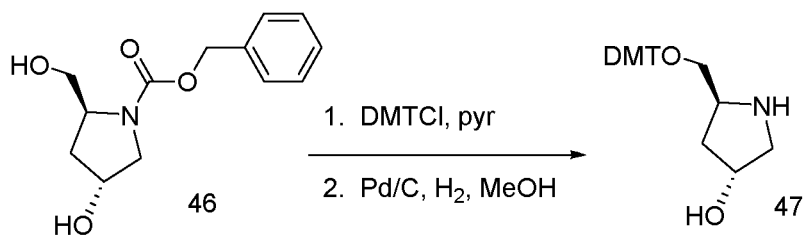
Compounds 23 and 36 are prepared as per the procedures illustrated in Examples 5 and 11. Compound 41 is prepared using similar procedures published in WO 2009082607.

Example 14: Preparation of Compound 45

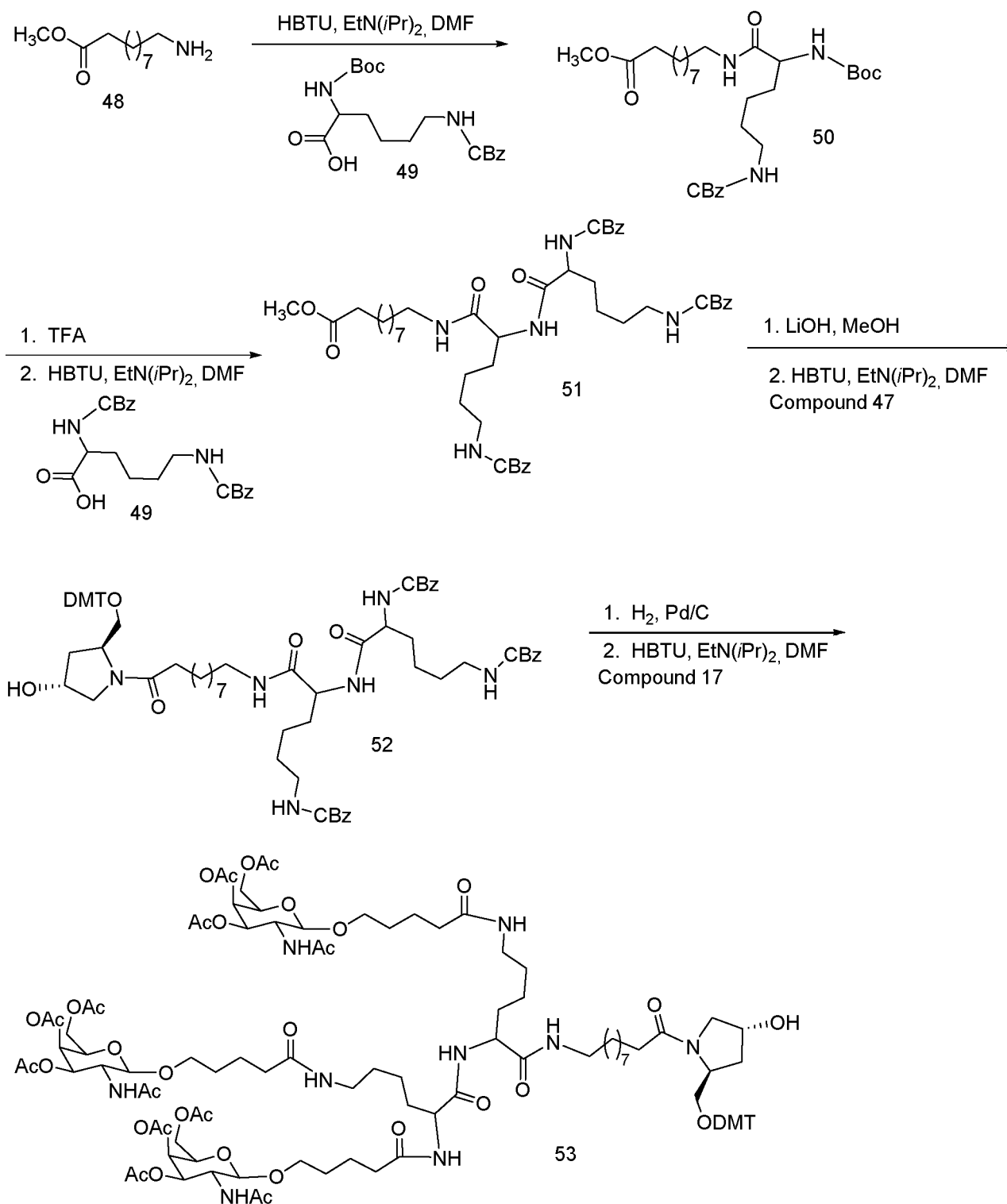


Compound 43 is prepared as per the procedures illustrated in Example 13.

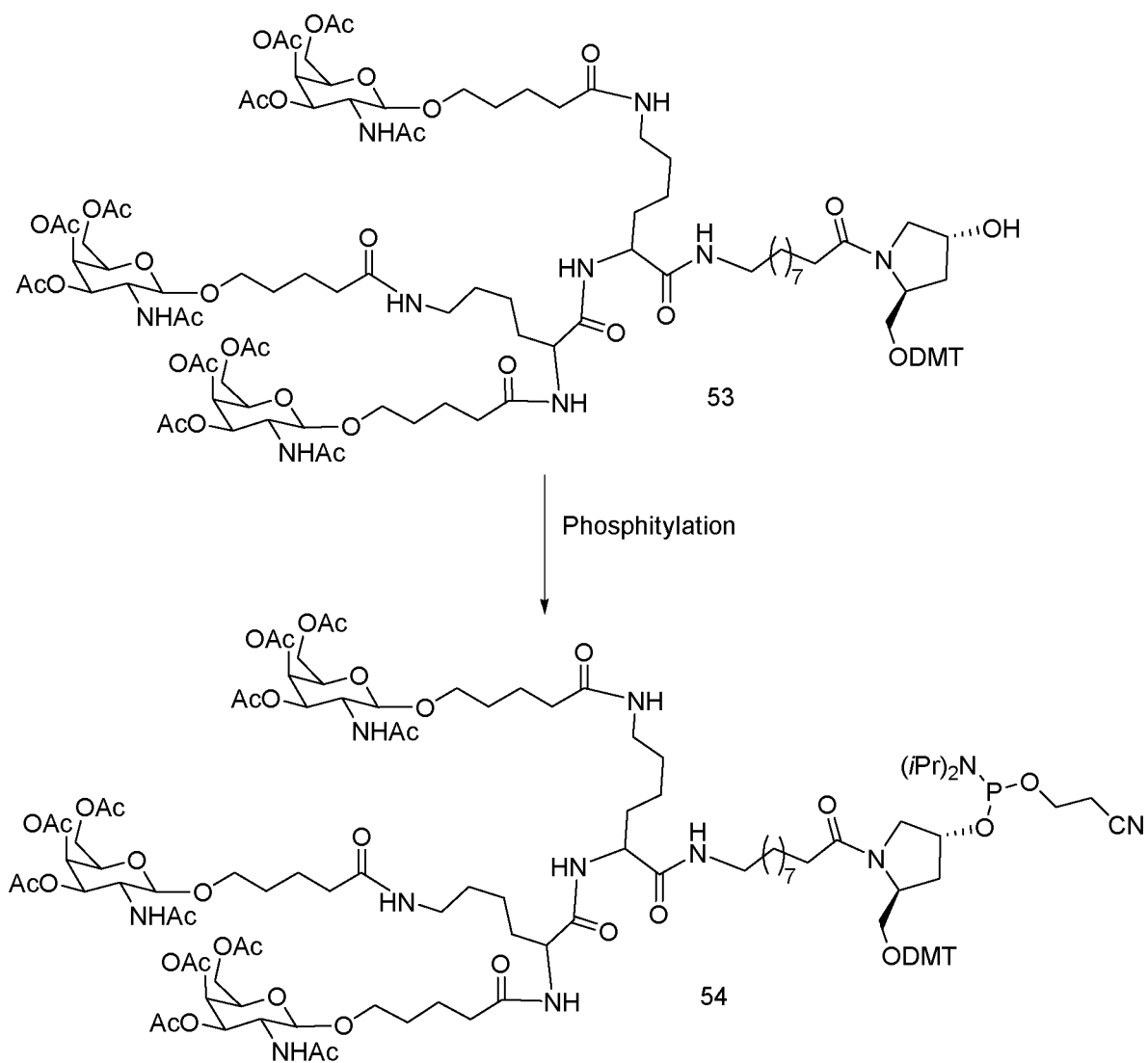
5 Example 15: Preparation of Compound 47



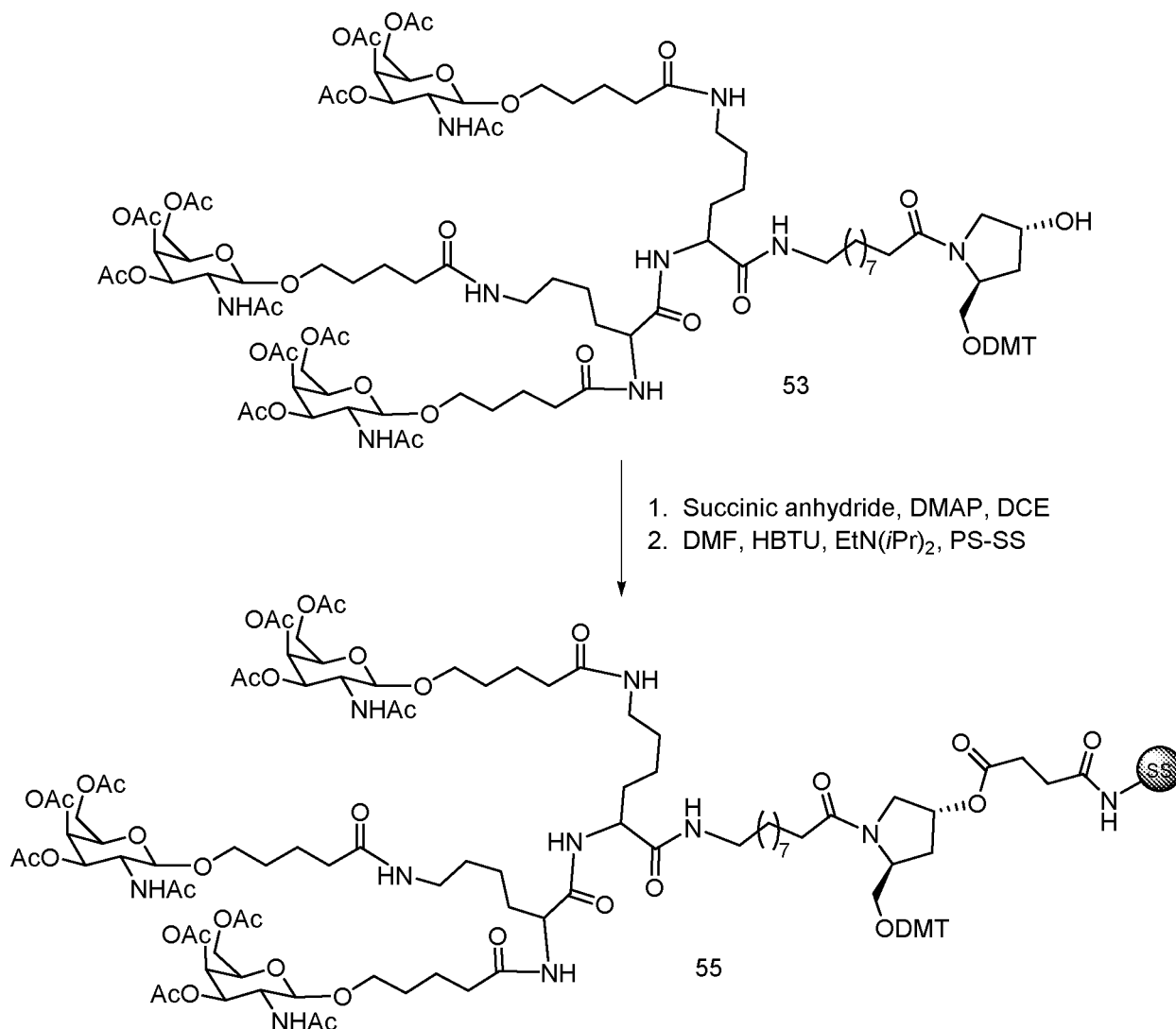
Compound 46 is commercially available.

Example 16: Preparation of Compound 53

Compounds 48 and 49 are commercially available. Compounds 17 and 47 are prepared as per the procedures illustrated in Examples 4 and 15.

Example 17: Preparation of Compound 54

Compound 53 is prepared as per the procedures illustrated in Example 16.

Example 18: Preparation of Compound 55

Compound 53 is prepared as per the procedures illustrated in Example 16.

Example 19: General method for the preparation of conjugated ASOs comprising GalNAc₃-1 at the 3' position via solid phase techniques (preparation of ISIS 647535, 647536 and 651900)

Unless otherwise stated, all reagents and solutions used for the synthesis of oligomeric compounds are purchased from commercial sources. Standard phosphoramidite building blocks and solid support are used for incorporation nucleoside residues which include for example T, A, G, and ^mC residues. A 0.1 M solution of phosphoramidite in anhydrous acetonitrile was used for β-D-2'-deoxyribonucleoside and 2'-MOE.

The ASO syntheses were performed on ABI 394 synthesizer (1-2 μmol scale) or on GE Healthcare Bioscience ÄKTA oligopilot synthesizer (40-200 μmol scale) by the phosphoramidite coupling method on an **GalNAc₃-1** loaded VIMAD solid support (110 μmol/g, Guzaev *et al.*, 2003) packed in the column. For the

coupling step, the phosphoramidites were delivered 4 fold excess over the loading on the solid support and phosphoramidite condensation was carried out for 10 min. All other steps followed standard protocols supplied by the manufacturer. A solution of 6% dichloroacetic acid in toluene was used for removing dimethoxytrityl (DMT) group from 5'-hydroxyl group of the nucleotide. 4,5-Dicyanoimidazole (0.7 M) in anhydrous CH₃CN was used as activator during coupling step. Phosphorothioate linkages were introduced by sulfurization with 0.1 M solution of xanthane hydride in 1:1 pyridine/CH₃CN for a contact time of 3 minutes. A solution of 20% *tert*-butylhydroperoxide in CH₃CN containing 6% water was used as an oxidizing agent to provide phosphodiester internucleoside linkages with a contact time of 12 minutes.

After the desired sequence was assembled, the cyanoethyl phosphate protecting groups were deprotected using a 1:1 (v/v) mixture of triethylamine and acetonitrile with a contact time of 45 minutes. The solid-support bound ASOs were suspended in aqueous ammonia (28-30 wt %) and heated at 55 °C for 6 h.

The unbound ASOs were then filtered and the ammonia was boiled off. The residue was purified by high pressure liquid chromatography on a strong anion exchange column (GE Healthcare Bioscience, Source 30Q, 30 μm, 2.54 x 8 cm, A = 100 mM ammonium acetate in 30% aqueous CH₃CN, B = 1.5 M NaBr in A, 0-40% of B in 60 min, flow 14 mL min⁻¹, λ = 260 nm). The residue was desalted by HPLC on a reverse phase column to yield the desired ASOs in an isolated yield of 15-30% based on the initial loading on the solid support. The ASOs were characterized by ion-pair-HPLC coupled MS analysis with Agilent 1100 MSD system.

Antisense oligonucleotides not comprising a conjugate were synthesized using standard oligonucleotide synthesis procedures well known in the art.

Using these methods, three separate antisense compounds targeting ApoC III were prepared. As summarized in Table 17, below, each of the three antisense compounds targeting ApoC III had the same nucleobase sequence; ISIS 304801 is a 5-10-5 MOE gapmer having all phosphorothioate linkages; ISIS 647535 is the same as ISIS 304801, except that it had a **GalNAc₃-1** conjugated at its 3'-end; and ISIS 647536 is the same as ISIS 647535 except that certain internucleoside linkages of that compound are phosphodiester linkages. As further summarized in Table 17, two separate antisense compounds targeting SRB-1 were synthesized. ISIS 440762 was a 2-10-2 cEt gapmer with all phosphorothioate internucleoside linkages; ISIS 651900 is the same as ISIS 440762, except that it included a **GalNAc₃-1** at its 3'-end.

Table 17

Modified ASO targeting ApoC III and SRB-1

ASO	Sequence (5' to 3')	Target	Calcd Mass	Observed Mass	SEQ ID No.
ISIS 304801	A _{cs} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{es} T _{es} T _{es} A _{cs} T _e	ApoC III	7165.4	7164.4	135
ISIS 647535	A _{cs} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{es} T _{es} T _{es} A _{cs} T _{eo} A_{do}'- GalNAc₃-1_a	ApoC III	9239.5	9237.8	136
ISIS 647536	A _{cs} G _{eo} ^m C _{eo} T _{eo} T _{eo} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{eo} T _{eo} T _{es} A _{cs} T _{eo} A_{do}'- GalNAc₃-1_a	ApoC III	9142.9	9140.8	136

ISIS 440762	$T_{ks}^m C_{ks} A_{ds} G_{ds} T_{ds}^m C_{ds} A_{ds} T_{ds} G_{ds} A_{ds}^m C_{ds} T_{ds} T_{ks}^m C_k$	SRB- 1	4647.0	4646.4	137
ISIS 651900	$T_{ks}^m C_{ks} A_{ds} G_{ds} T_{ds}^m C_{ds} A_{ds} T_{ds} G_{ds} A_{ds}^m C_{ds} T_{ds} T_{ks}^m C_{ko} A_{do} - \text{GalNAc}_3\text{-1}_a$	SRB- 1	6721.1	6719.4	138

Subscripts: “e” indicates 2’-MOE modified nucleoside; “d” indicates β -D-2’-deoxyribonucleoside; “k” indicates 6’-(*S*)-CH₃ bicyclic nucleoside (e.g. cEt); “s” indicates phosphorothioate internucleoside linkages (PS); “o” indicates phosphodiester internucleoside linkages (PO); and “o” indicates -O-P(=O)(OH)-. Superscript “m” indicates 5-methylcytosines. “GalNAc₃-1” indicates a conjugate group having the structure shown previously in Example 9. Note that GalNAc₃-1 comprises a cleavable adenosine which links the ASO to remainder of the conjugate, which is designated “GalNAc₃-1_a.” This nomenclature is used in the above table to show the full nucleobase sequence, including the adenosine, which is part of the conjugate. Thus, in the above table, the sequences could also be listed as ending with “GalNAc₃-1” with the “A_{do}” omitted. This convention of using the subscript “a” to indicate the portion of a conjugate group lacking a cleavable nucleoside or cleavable moiety is used throughout these Examples. This portion of a conjugate group lacking the cleavable moiety is referred to herein as a “cluster” or “conjugate cluster” or “GalNAc₃ cluster.” In certain instances it is convenient to describe a conjugate group by separately providing its cluster and its cleavable moiety.

Example 20: Dose-dependent antisense inhibition of human ApoC III in huApoC III transgenic mice

ISIS 304801 and ISIS 647535, each targeting human ApoC III and described above, were separately tested and evaluated in a dose-dependent study for their ability to inhibit human ApoC III in human ApoC III transgenic mice.

Treatment

Human ApoCIII transgenic mice were maintained on a 12-hour light/dark cycle and fed *ad libitum* Teklad lab chow. Animals were acclimated for at least 7 days in the research facility before initiation of the experiment. ASOs were prepared in PBS and sterilized by filtering through a 0.2 micron filter. ASOs were dissolved in 0.9% PBS for injection.

Human ApoC III transgenic mice were injected intraperitoneally once a week for two weeks with ISIS 304801 or 647535 at 0.08, 0.25, 0.75, 2.25 or 6.75 μ mol/kg or with PBS as a control. Each treatment group consisted of 4 animals. Forty-eight hours after the administration of the last dose, blood was drawn from each mouse and the mice were sacrificed and tissues were collected.

ApoC III mRNA Analysis

ApoC III mRNA levels in the mice’s livers were determined using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. ApoC III mRNA levels were determined relative to total RNA (using Ribogreen), prior to normalization to PBS-treated control. The results below are presented as the average percent of ApoC III

mRNA levels for each treatment group, normalized to PBS-treated control and are denoted as “% PBS”. The half maximal effective dosage (ED₅₀) of each ASO is also presented in Table 18, below.

As illustrated, both antisense compounds reduced ApoC III RNA relative to the PBS control. Further, the antisense compound conjugated to **GalNAc₃-1** (ISIS 647535) was substantially more potent than the antisense compound lacking the **GalNAc₃-1** conjugate (ISIS 304801).

Table 18

Effect of ASO treatment on ApoC III mRNA levels in human ApoC III transgenic mice

ASO	Dose (μmol/kg)	% PBS	ED ₅₀ (μmol/kg)	3' Conjugate	Internucleoside linkage/Length	SEQ ID No.
PBS	0	100	--	-	--	
ISIS 304801	0.08	95	0.77	None	PS/20	135
	0.75	42				
	2.25	32				
	6.75	19				
ISIS 647535	0.08	50	0.074	GalNAc₃-1	PS/20	136
	0.75	15				
	2.25	17				
	6.75	8				

ApoC III Protein Analysis (Turbidometric Assay)

Plasma ApoC III protein analysis was determined using procedures reported by Graham *et al*, *Circulation Research*, published online before print March 29, 2013.

Approximately 100 μl of plasma isolated from mice was analyzed without dilution using an Olympus Clinical Analyzer and a commercially available turbidometric ApoC III assay (Kamiya, Cat# KAI-006, Kamiya Biomedical, Seattle, WA). The assay protocol was performed as described by the vendor.

As shown in the Table 19 below, both antisense compounds reduced ApoC III protein relative to the PBS control. Further, the antisense compound conjugated to **GalNAc₃-1** (ISIS 647535) was substantially more potent than the antisense compound lacking the **GalNAc₃-1** conjugate (ISIS 304801).

Table 19

Effect of ASO treatment on ApoC III plasma protein levels in human ApoC III transgenic mice

ASO	Dose (μmol/kg)	% PBS	ED ₅₀ (μmol/kg)	3' Conjugate	Internucleoside Linkage/Length	SEQ ID No.
PBS	0	100	--	--	--	
ISIS 304801	0.08	86	0.73	None	PS/20	135
	0.75	51				

	2.25	23				
	6.75	13				
ISIS 647535	0.08	72	0.19	GalNAc₃-1	PS/20	136
	0.75	14				
	2.25	12				
	6.75	11				

Plasma triglycerides and cholesterol were extracted by the method of Bligh and Dyer (Bligh, E.G. and Dyer, W.J. *Can. J. Biochem. Physiol.* 37: 911-917, 1959)(Bligh, E and Dyer, W, *Can J Biochem Physiol*, 37, 911-917, 1959)(Bligh, E and Dyer, W, *Can J Biochem Physiol*, 37, 911-917, 1959) and measured by using a Beckmann Coulter clinical analyzer and commercially available reagents.

The triglyceride levels were measured relative to PBS injected mice and are denoted as “% PBS”. Results are presented in Table 20. As illustrated, both antisense compounds lowered triglyceride levels. Further, the antisense compound conjugated to **GalNAc₃-1** (ISIS 647535) was substantially more potent than the antisense compound lacking the **GalNAc₃-1** conjugate (ISIS 304801).

Table 20

Effect of ASO treatment on triglyceride levels in transgenic mice

ASO	Dose (μmol/kg)	% PBS	ED ₅₀ (μmol/kg)	3' Conjugate	Internucleoside Linkage/Length	SEQ ID No.
PBS	0	100	--	--	--	
ISIS 304801	0.08	87	0.63	None	PS/20	135
	0.75	46				
	2.25	21				
	6.75	12				
ISIS 647535	0.08	65	0.13	GalNAc₃-1	PS/20	136
	0.75	9				
	2.25	8				
	6.75	9				

Plasma samples were analyzed by HPLC to determine the amount of total cholesterol and of different fractions of cholesterol (HDL and LDL). Results are presented in Tables 21 and 22. As illustrated, both antisense compounds lowered total cholesterol levels; both lowered LDL; and both raised HDL. Further, the antisense compound conjugated to **GalNAc₃-1** (ISIS 647535) was substantially more potent than the antisense compound lacking the **GalNAc₃-1** conjugate (ISIS 304801). An increase in HDL and a decrease in LDL levels is a cardiovascular beneficial effect of antisense inhibition of ApoC III.

Table 21

Effect of ASO treatment on total cholesterol levels in transgenic mice

ASO	Dose (μmol/kg)	Total Cholesterol (mg/dL)	3' Conjugate	Internucleoside Linkage/Length	SEQ ID No.
PBS	0	257	--	--	
ISIS 304801	0.08	226	None	PS/20	135
	0.75	164			
	2.25	110			
	6.75	82			
ISIS 647535	0.08	230	GalNAc ₃ -1	PS/20	136
	0.75	82			
	2.25	86			
	6.75	99			

Table 22

Effect of ASO treatment on HDL and LDL cholesterol levels in transgenic mice

ASO	Dose (μmol/kg)	HDL (mg/dL)	LDL (mg/dL)	3' Conjugate	Internucleoside Linkage/Length	SEQ ID No.
PBS	0	17	28	--	--	
ISIS 304801	0.08	17	23	None	PS/20	135
	0.75	27	12			
	2.25	50	4			
	6.75	45	2			
ISIS 647535	0.08	21	21	GalNAc ₃ -1	PS/20	136
	0.75	44	2			
	2.25	50	2			
	6.75	58	2			

Pharmacokinetics Analysis (PK)

The PK of the ASOs was also evaluated. Liver and kidney samples were minced and extracted using standard protocols. Samples were analyzed on MSD1 utilizing IP-HPLC-MS. The tissue level (μg/g) of full-length ISIS 304801 and 647535 was measured and the results are provided in Table 23. As illustrated, liver concentrations of total full-length antisense compounds were similar for the two antisense compounds. Thus, even though the GalNAc₃-1 -conjugated antisense compound is more active in the liver (as demonstrated by the RNA and protein data above), it is not present at substantially higher concentration in the liver. Indeed, the calculated EC₅₀ (provided in Table 23) confirms that the observed increase in potency of the conjugated compound cannot be entirely attributed to increased accumulation. This result suggests that

the conjugate improved potency by a mechanism other than liver accumulation alone, possibly by improving the productive uptake of the antisense compound into cells.

The results also show that the concentration of **GalNAc₃-1** conjugated antisense compound in the kidney is lower than that of antisense compound lacking the GalNAc conjugate. This has several beneficial therapeutic implications. For therapeutic indications where activity in the kidney is not sought, exposure to kidney risks kidney toxicity without corresponding benefit. Moreover, high concentration in kidney typically results in loss of compound to the urine resulting in faster clearance. Accordingly, for non-kidney targets, kidney accumulation is undesired. These data suggest that **GalNAc₃-1** conjugation reduces kidney accumulation.

Table 23

PK analysis of ASO treatment in transgenic mice

ASO	Dose (μmol/kg)	Liver (μg/g)	Kidney (μg/g)	Liver EC ₅₀ (μg/g)	3' Conjugate	Internucleoside Linkage/Length	SEQ ID No.
ISIS 304801	0.1	5.2	2.1	53	None	PS/20	135
	0.8	62.8	119.6				
	2.3	142.3	191.5				
	6.8	202.3	337.7				
ISIS 647535	0.1	3.8	0.7	3.8	GalNAc₃-1	PS/20	136
	0.8	72.7	34.3				
	2.3	106.8	111.4				
	6.8	237.2	179.3				

Metabolites of ISIS 647535 were also identified and their masses were confirmed by high resolution mass spectrometry analysis. The cleavage sites and structures of the observed metabolites are shown below. The relative % of full length ASO was calculated using standard procedures and the results are presented in Table 23a. The major metabolite of ISIS 647535 was full-length ASO lacking the entire conjugate (i.e. ISIS 304801), which results from cleavage at cleavage site A, shown below. Further, additional metabolites resulting from other cleavage sites were also observed. These results suggest that introducing other cleavable bonds such as esters, peptides, disulfides, phosphoramidates or acyl-hydrazones between the **GalNAc₃-1** sugar and the ASO, which can be cleaved by enzymes inside the cell, or which may cleave in the reductive environment of the cytosol, or which are labile to the acidic pH inside endosomes and lysosomes, can also be useful.

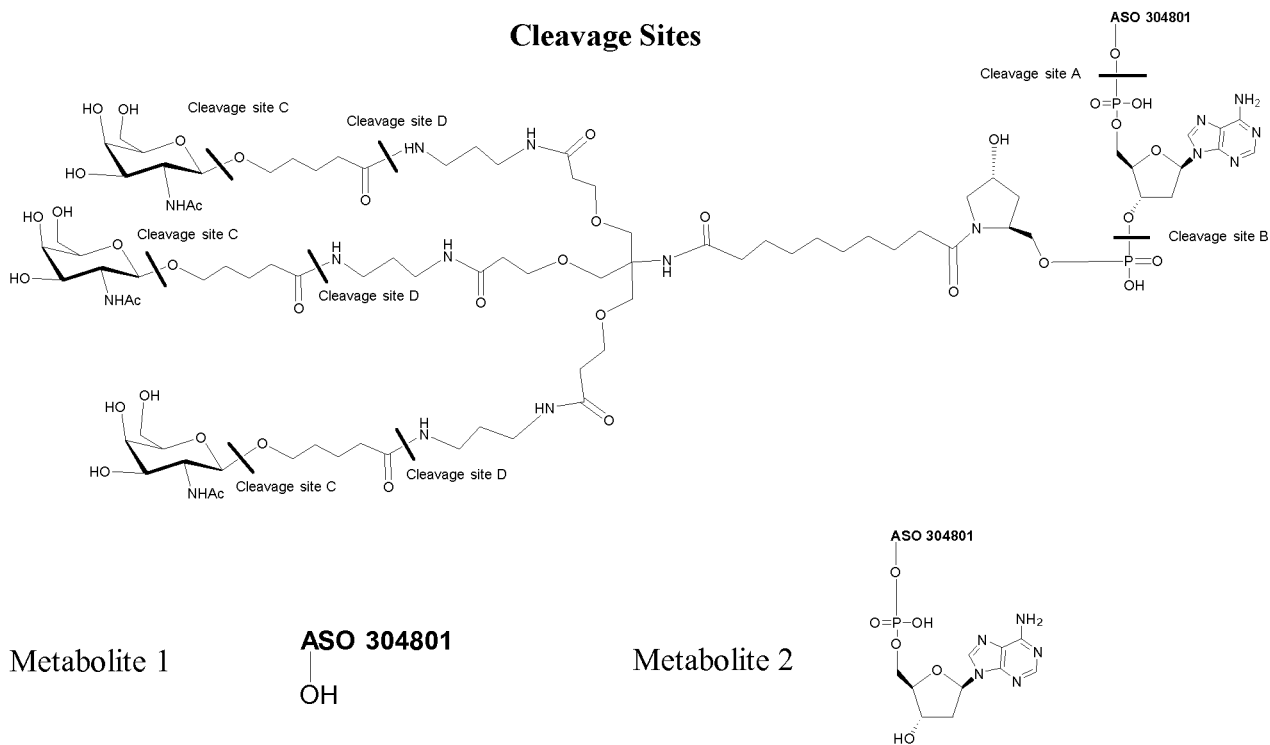
Table 23a

Observed full length metabolites of ISIS 647535

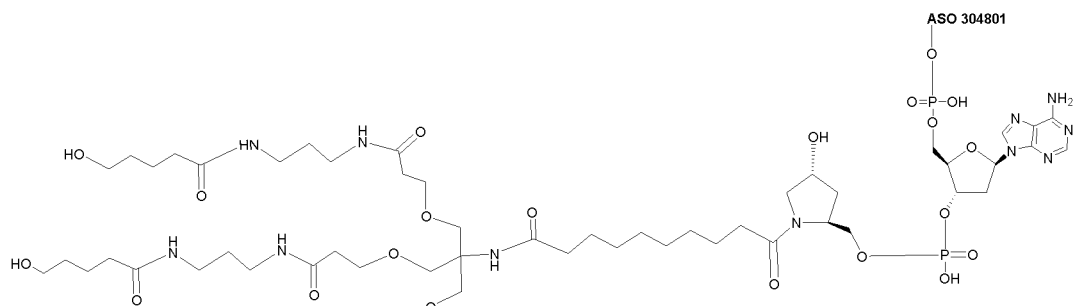
Metabolite	ASO	Cleavage site	Relative %
1	ISIS 304801	A	36.1

2	ISIS 304801 + dA	B	10.5
3	ISIS 647535 minus [3 GalNAc]	C	16.1
4	ISIS 647535 minus [3 GalNAc + 1 5-hydroxy-pentanoic acid tether]	D	17.6
5	ISIS 647535 minus [2 GalNAc + 2 5-hydroxy-pentanoic acid tether]	D	9.9
6	ISIS 647535 minus [3 GalNAc + 3 5-hydroxy-pentanoic acid tether]	D	9.8

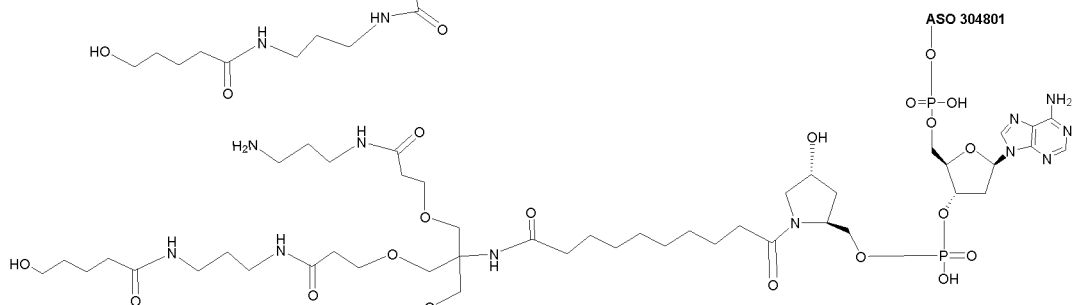
Cleavage Sites



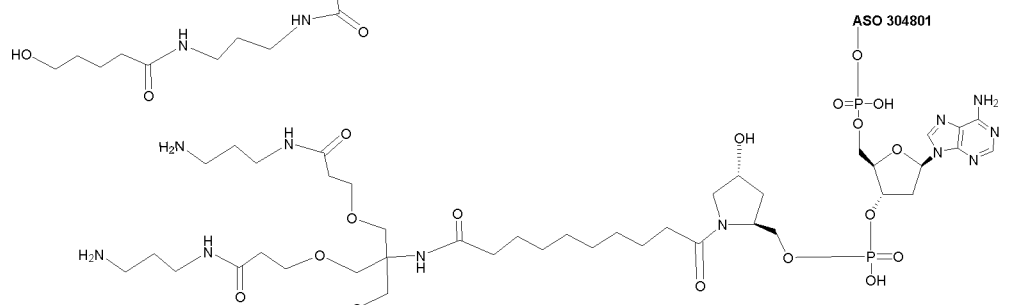
Metabolite 3



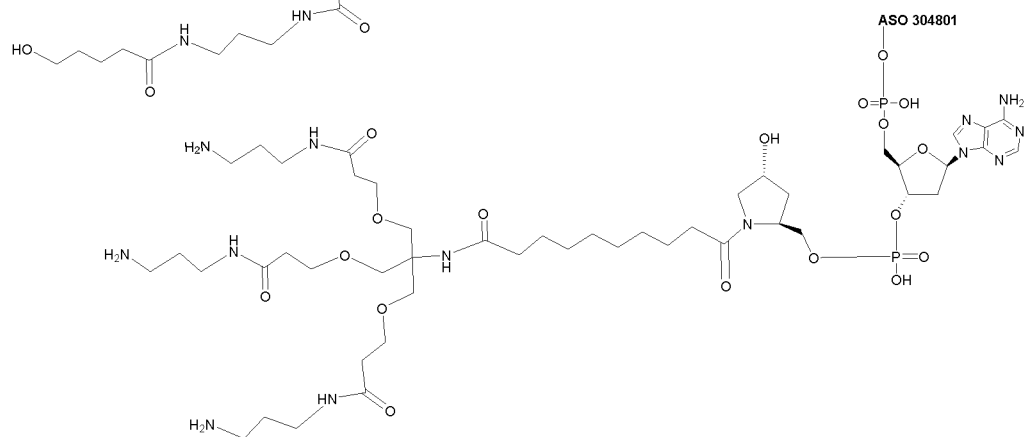
Metabolite 4



Metabolite 5



Metabolite 6



Example 21: Antisense inhibition of human ApoC III in human ApoC III transgenic mice in single administration study

ISIS 304801, 647535 and 647536 each targeting human ApoC III and described in Table 17, were further evaluated in a single administration study for their ability to inhibit human ApoC III in human ApoC

Treatment

Human ApoCIII transgenic mice were maintained on a 12-hour light/dark cycle and fed *ad libitum* Teklad lab chow. Animals were acclimated for at least 7 days in the research facility before initiation of the experiment. ASOs were prepared in PBS and sterilized by filtering through a 0.2 micron filter. ASOs were dissolved in 0.9% PBS for injection.

Human ApoC III transgenic mice were injected intraperitoneally once at the dosage shown below with ISIS 304801, 647535 or 647536 (described above) or with PBS treated control. The treatment group consisted of 3 animals and the control group consisted of 4 animals. Prior to the treatment as well as after the last dose, blood was drawn from each mouse and plasma samples were analyzed. The mice were sacrificed 72 hours following the last administration .

Samples were collected and analyzed to determine the ApoC III mRNA and protein levels in the liver; plasma triglycerides; and cholesterol, including HDL and LDL fractions were assessed as described above (Example 20). Data from those analyses are presented in Tables 24-28, below. Liver transaminase levels, alanine aminotransferase (ALT) and aspartate aminotransferase (AST), in serum were measured relative to saline injected mice using standard protocols. The ALT and AST levels showed that the antisense compounds were well tolerated at all administered doses.

These results show improvement in potency for antisense compounds comprising a **GalNAc₃-1** conjugate at the 3' terminus (ISIS 647535 and 647536) compared to the antisense compound lacking a **GalNAc₃-1** conjugate (ISIS 304801). Further, ISIS 647536, which comprises a **GalNAc₃-1** conjugate and some phosphodiester linkages was as potent as ISIS 647535, which comprises the same conjugate and all internucleoside linkages within the ASO are phosphorothioate.

Table 24

Effect of ASO treatment on ApoC III mRNA levels in human ApoC III transgenic mice

ASO	Dose (mg/kg)	% PBS	ED ₅₀ (mg/kg)	3' Conjugate	Internucleoside linkage/Length	SEQ ID No.
PBS	0	99	--	-	--	
ISIS 304801	1	104	13.2	None	PS/20	135
	3	92				
	10	71				
	30	40				
ISIS 647535	0.3	98	1.9	GalNAc₃-1	PS/20	136
	1	70				
	3	33				

	10	20				
ISIS 647536	0.3	103	1.7	GalNAc₃-1	PS/PO/20	136
	1	60				
	3	31				
	10	21				

Table 25

Effect of ASO treatment on ApoC III plasma protein levels in human ApoC III transgenic mice

ASO	Dose (mg/kg)	% PBS	ED ₅₀ (mg/kg)	3' Conjugate	Internucleoside Linkage/Length	SEQ ID No.
PBS	0	99	--	--	--	
ISIS 304801	1	104	23.2	None	PS/20	135
	3	92				
	10	71				
	30	40				
ISIS 647535	0.3	98	2.1	GalNAc₃-1	PS/20	136
	1	70				
	3	33				
	10	20				
ISIS 647536	0.3	103	1.8	GalNAc₃-1	PS/PO/20	136
	1	60				
	3	31				
	10	21				

5

Table 26

Effect of ASO treatment on triglyceride levels in transgenic mice

ASO	Dose (mg/kg)	% PBS	ED ₅₀ (mg/kg)	3' Conjugate	Internucleoside Linkage/Length	SEQ ID No.
PBS	0	98	--	--	--	
ISIS 304801	1	80	29.1	None	PS/20	135
	3	92				
	10	70				
	30	47				
ISIS 647535	0.3	100	2.2	GalNAc₃-1	PS/20	136
	1	70				
	3	34				
	10	23				
ISIS 647536	0.3	95	1.9	GalNAc₃-1	PS/PO/20	136
	1	66				
	3	31				
	10	23				

Table 27

Effect of ASO treatment on total cholesterol levels in transgenic mice

ASO	Dose (mg/kg)	% PBS	3' Conjugate	Internucleoside Linkage/Length	SEQ ID No.
-----	--------------	-------	--------------	--------------------------------	------------

PBS	0	96	--	--	
ISIS 304801	1	104	None	PS/20	135
	3	96			
	10	86			
	30	72			
ISIS 647535	0.3	93	GalNAc₃-1	PS/20	136
	1	85			
	3	61			
	10	53			
ISIS 647536	0.3	115	GalNAc₃-1	PS/PO/20	136
	1	79			
	3	51			
	10	54			

Table 28

Effect of ASO treatment on HDL and LDL cholesterol levels in transgenic mice

ASO	Dose (mg/kg)	HDL % PBS	LDL % PBS	3' Conjugate	Internucleoside Linkage/Length	SEQ ID No.
PBS	0	131	90	--	--	
ISIS 304801	1	130	72	None	PS/20	135
	3	186	79			
	10	226	63			
	30	240	46			
ISIS 647535	0.3	98	86	GalNAc₃-1	PS/20	136
	1	214	67			
	3	212	39			
	10	218	35			
ISIS 647536	0.3	143	89	GalNAc₃-1	PS/PO/20	136
	1	187	56			
	3	213	33			
	10	221	34			

5 These results confirm that the **GalNAc₃-1** conjugate improves potency of an antisense compound. The results also show equal potency of a **GalNAc₃-1** conjugated antisense compounds where the antisense oligonucleotides have mixed linkages (ISIS 647536 which has six phosphodiester linkages) and a full phosphorothioate version of the same antisense compound (ISIS 647535).

10 Phosphorothioate linkages provide several properties to antisense compounds. For example, they resist nuclease digestion and they bind proteins resulting in accumulation of compound in the liver, rather than in the kidney/urine. These are desirable properties, particularly when treating an indication in the liver. However, phosphorothioate linkages have also been associated with an inflammatory response. Accordingly, reducing the number of phosphorothioate linkages in a compound is expected to reduce the risk of

inflammation, but also lower concentration of the compound in liver, increase concentration in the kidney and urine, decrease stability in the presence of nucleases, and lower overall potency. The present results show that a **GalNAc₃-1** conjugated antisense compound where certain phosphorothioate linkages have been replaced with phosphodiester linkages is as potent against a target in the liver as a counterpart having full phosphorothioate linkages. Such compounds are expected to be less proinflammatory (See Example 24 describing an experiment showing reduction of PS results in reduced inflammatory effect).

Example 22: Effect of GalNAc₃-1 conjugated modified ASO targeting SRB-1 *in vivo*

ISIS 440762 and 651900, each targeting SRB-1 and described in Table 17, were evaluated in a dose-dependent study for their ability to inhibit SRB-1 in Balb/c mice.

Treatment

Six week old male Balb/c mice (Jackson Laboratory, Bar Harbor, ME) were injected subcutaneously once at the dosage shown below with ISIS 440762, 651900 or with PBS treated control. Each treatment group consisted of 4 animals. The mice were sacrificed 48 hours following the final administration to determine the SRB-1 mRNA levels in liver using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. SRB-1 mRNA levels were determined relative to total RNA (using Ribogreen), prior to normalization to PBS-treated control. The results below are presented as the average percent of SRB-1 mRNA levels for each treatment group, normalized to PBS-treated control and is denoted as “% PBS”.

As illustrated in Table 29, both antisense compounds lowered SRB-1 mRNA levels. Further, the antisense compound comprising the **GalNAc₃-1** conjugate (ISIS 651900) was substantially more potent than the antisense compound lacking the **GalNAc₃-1** conjugate (ISIS 440762). These results demonstrate that the potency benefit of **GalNAc₃-1** conjugates are observed using antisense oligonucleotides complementary to a different target and having different chemically modified nucleosides, in this instance modified nucleosides comprise constrained ethyl sugar moieties (a bicyclic sugar moiety).

Table 29

Effect of ASO treatment on SRB-1 mRNA levels in Balb/c mice

ASO	Dose (mg/kg)	Liver % PBS	ED ₅₀ (mg/kg)	3' Conjugate		Internucleoside linkage/Length	SEQ ID No.
PBS	0	100		-		--	
ISIS 440762	0.7	85	2.2	None		PS/14	137
	2	55					
	7	12					
	20	3					

ISIS 651900	0.07	98	0.3	GalNAc₃-1		PS/14	138
	0.2	63					
	0.7	20					
	2	6					
	7	5					

Example 23: Human Peripheral Blood Mononuclear Cells (hPBMC) Assay Protocol

The hPBMC assay was performed using BD Vacutainer CPT tube method. A sample of whole blood from volunteered donors with informed consent at US HealthWorks clinic (Faraday & El Camino Real, Carlsbad) was obtained and collected in 4-15 BD Vacutainer CPT 8 ml tubes (VWR Cat.# BD362753). The approximate starting total whole blood volume in the CPT tubes for each donor was recorded using the PBMC assay data sheet.

The blood sample was remixed immediately prior to centrifugation by gently inverting tubes 8-10 times. CPT tubes were centrifuged at rt (18-25 °C) in a horizontal (swing-out) rotor for 30 min. at 1500-1800 RCF with brake off (2700 RPM Beckman Allegra 6R). The cells were retrieved from the buffy coat interface (between Ficoll and polymer gel layers); transferred to a sterile 50 ml conical tube and pooled up to 5 CPT tubes/50 ml conical tube/donor. The cells were then washed twice with PBS (Ca⁺⁺, Mg⁺⁺ free; GIBCO). The tubes were topped up to 50 ml and mixed by inverting several times. The sample was then centrifuged at 330 x g for 15 minutes at rt (1215 RPM in Beckman Allegra 6R) and aspirated as much supernatant as possible without disturbing pellet. The cell pellet was dislodged by gently swirling tube and resuspended cells in RPMI+10% FBS+pen/strep (~1 ml / 10 ml starting whole blood volume). A 60 µl sample was pipette into a sample vial (Beckman Coulter) with 600 µl VersaLyse reagent (Beckman Coulter Cat# A09777) and was gently vortexed for 10-15 sec. The sample was allowed to incubate for 10 min. at rt and being mixed again before counting. The cell suspension was counted on Vicell XR cell viability analyzer (Beckman Coulter) using PBMC cell type (dilution factor of 1:11 was stored with other parameters). The live cell/ml and viability were recorded. The cell suspension was diluted to 1 x 10⁷ live PBMC/ml in RPMI+ 10% FBS+pen/strep.

The cells were plated at 5 x 10⁵ in 50 µl/well of 96-well tissue culture plate (Falcon Microtest). 50 µl/well of 2x concentration oligos/controls diluted in RPMI+10% FBS+pen/strep. was added according to experiment template (100 µl/well total). Plates were placed on the shaker and allowed to mix for approx. 1 min. After being incubated for 24 hrs at 37 °C; 5% CO₂, the plates were centrifuged at 400 x g for 10 minutes before removing the supernatant for MSD cytokine assay (i.e. human IL-6, IL-10, IL-8 and MCP-1).

Example 24: Evaluation of Proinflammatory Effects in hPBMC Assay for GalNAc₃-1 conjugated ASOs

The antisense oligonucleotides (ASOs) listed in Table 30 were evaluated for proinflammatory effect in hPBMC assay using the protocol described in Example 23. ISIS 353512 is an internal standard known to be a high responder for IL-6 release in the assay. The hPBMCs were isolated from fresh, volunteered donors

and were treated with ASOs at 0, 0.0128, 0.064, 0.32, 1.6, 8, 40 and 200 μ M concentrations. After a 24 hr treatment, the cytokine levels were measured.

The levels of IL-6 were used as the primary readout. The EC_{50} and E_{max} was calculated using standard procedures. Results are expressed as the average ratio of E_{max}/EC_{50} from two donors and is denoted as “ E_{max}/EC_{50} .” The lower ratio indicates a relative decrease in the proinflammatory response and the higher ratio indicates a relative increase in the proinflammatory response.

With regard to the test compounds, the least proinflammatory compound was the PS/PO linked ASO (ISIS 616468). The **GalNAc₃-1** conjugated ASO, ISIS 647535 was slightly less proinflammatory than its non-conjugated counterpart ISIS 304801. These results indicate that incorporation of some PO linkages reduces proinflammatory reaction and addition of a **GalNAc₃-1** conjugate does not make a compound more proinflammatory and may reduce proinflammatory response. Accordingly, one would expect that an antisense compound comprising both mixed PS/PO linkages and a **GalNAc₃-1** conjugate would produce lower proinflammatory responses relative to full PS linked antisense compound with or without a **GalNAc₃-1** conjugate. These results show that **GalNAc₃-1** conjugated antisense compounds, particularly those having reduced PS content are less proinflammatory.

Together, these results suggest that a **GalNAc₃-1** conjugated compound, particularly one with reduced PS content, can be administered at a higher dose than a counterpart full PS antisense compound lacking a **GalNAc₃-1** conjugate. Since half-life is not expected to be substantially different for these compounds, such higher administration would result in less frequent dosing. Indeed such administration could be even less frequent, because the **GalNAc₃-1** conjugated compounds are more potent (See Examples 20-22) and re-dosing is necessary once the concentration of a compound has dropped below a desired level, where such desired level is based on potency.

Table 30

Modified ASOs

ASO	Sequence (5' to 3')	Target	SEQ ID No.
ISIS 104838	G ^m _{es} C ^m _{es} T ^m _{es} G ^m _{es} A ^m _{es} T ^m _{ds} T ^m _{ds} A ^m _{ds} G ^m _{ds} A ^m _{ds} G ^m _{ds} A ^m _{ds} G ^m _{ds} A ^m _{ds} G ^m _{ds} G ^m _{es} T ^m _{es} C ^m _{es} C ^m _{es} C ^m _e	TNF α	139
ISIS 353512	T ^m _{es} C ^m _{es} C ^m _{es} C ^m _{ds} A ^m _{ds} T ^m _{ds} T ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} G ^m _{ds} G ^m _{ds} A ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} C ^m _{ds} T ^m _{es} G ^m _{es} G ^m _e	CRP	140
ISIS 304801	A ^m _{es} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{ds} T ^m _{ds} T ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} C ^m _{ds} A ^m _{ds} G ^m _{ds} C ^m _{ds} T ^m _{es} T ^m _{es} T ^m _{es} A ^m _{es} T ^m _e	ApoC III	135
ISIS 647535	A ^m _{es} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{ds} T ^m _{ds} T ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} C ^m _{ds} A ^m _{ds} G ^m _{ds} C ^m _{ds} T ^m _{es} T ^m _{es} T ^m _{es} A ^m _{es} T ^m _{eo} A ^m _{do} - GalNAc₃-1_a	ApoC III	136
ISIS 616468	A ^m _{es} G ^m _{eo} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{ds} T ^m _{ds} T ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} C ^m _{ds} A ^m _{ds} G ^m _{ds} C ^m _{ds} T ^m _{eo} T ^m _{eo} T ^m _{es} A ^m _{es} T ^m _e	ApoC III	135

Subscripts: “e” indicates 2'-MOE modified nucleoside; “d” indicates β -D-2'-deoxyribonucleoside; “k” indicates 6'-(S)-CH₃ bicyclic nucleoside (e.g. cEt); “s” indicates phosphorothioate

internucleoside linkages (PS); “o” indicates phosphodiester internucleoside linkages (PO); and “o” indicates -O-P(=O)(OH)-. Superscript “m” indicates 5-methylcytosines. “**A_{do}-GalNAc₃-1_a**” indicates a conjugate having the structure **GalNAc₃-1** shown in Example 9 attached to the 3'-end of the antisense oligonucleotide, as indicated.

5

Table 31

Proinflammatory Effect of ASOs targeting ApoC III in hPBMC assay

ASO	EC ₅₀ (μM)	E _{max} (μM)	E _{max} /EC ₅₀	3' Conjugate	Internucleoside Linkage/Length	SEQ ID No.
ISIS 353512 (high responder)	0.01	265.9	26,590	None	PS/20	140
ISIS 304801	0.07	106.55	1,522	None	PS/20	135
ISIS 647535	0.12	138	1,150	GalNAc₃-1	PS/20	136
ISIS 616468	0.32	71.52	224	None	PS/PO/20	135

Example 25: Effect of GalNAc₃-1 conjugated modified ASO targeting human ApoC III *in vitro*

ISIS 304801 and 647535 described above were tested *in vitro*. Primary hepatocyte cells from transgenic mice at a density of 25,000 cells per well were treated with 0.03, 0.08, 0.24, 0.74, 2.22, 6.67 and 20 μM concentrations of modified oligonucleotides. After a treatment period of approximately 16 hours, RNA was isolated from the cells and mRNA levels were measured by quantitative real-time PCR and the hApoC III mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN.

The IC₅₀ was calculated using the standard methods and the results are presented in Table 32. As illustrated, comparable potency was observed in cells treated with ISIS 647535 as compared to the control, ISIS 304801.

Table 32

Modified ASO targeting human ApoC III in primary hepatocytes

ASO	IC ₅₀ (μM)	3' Conjugate	Internucleoside linkage/Length	SEQ ID No.
ISIS 304801	0.44	None	PS/20	135
ISIS 647535	0.31	GalNAc₃-1	PS/20	136

In this experiment, the large potency benefits of **GalNAc₃-1** conjugation that are observed *in vivo* were not observed *in vitro*. Subsequent free uptake experiments in primary hepatocytes *in vitro* did show increased potency of oligonucleotides comprising various GalNAc conjugates relative to oligonucleotides that lacking the GalNAc conjugate.(see Examples 60, 82, and 92)

5 **Example 26: Effect of PO/PS linkages on ApoC III ASO Activity**

Human ApoC III transgenic mice were injected intraperitoneally once at 25 mg/kg of ISIS 304801, or ISIS 616468 (both described above) or with PBS treated control once per week for two weeks. The treatment group consisted of 3 animals and the control group consisted of 4 animals. Prior to the treatment as well as after the last dose, blood was drawn from each mouse and plasma samples were analyzed. The mice
10 were sacrificed 72 hours following the last administration.

Samples were collected and analyzed to determine the ApoC III protein levels in the liver as described above (Example 20). Data from those analyses are presented in Table 33, below.

These results show reduction in potency for antisense compounds with PO/PS (ISIS 616468) in the wings relative to full PS (ISIS 304801).

15

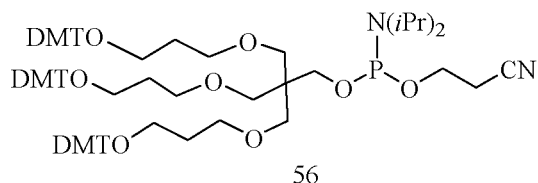
Table 33

Effect of ASO treatment on ApoC III protein levels in human ApoC III transgenic mice

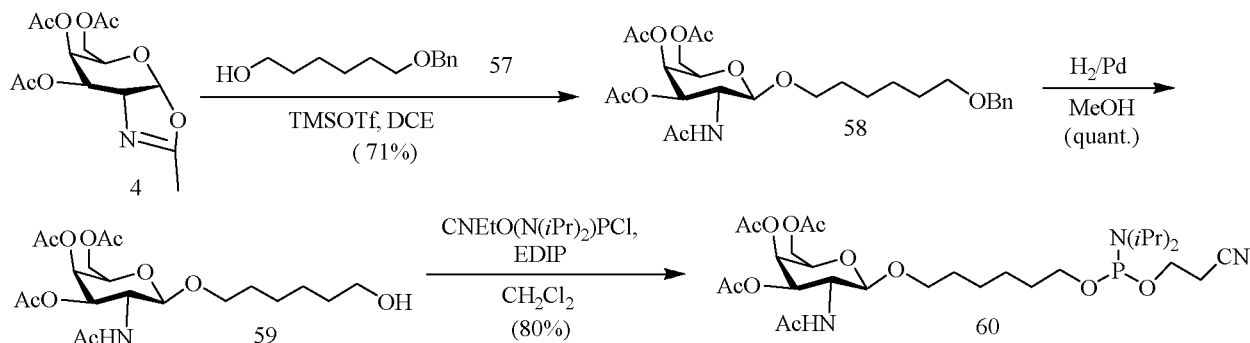
ASO	Dose (mg/kg)	% PBS	3' Conjugate	Internucleoside linkage/Length	SEQ ID No.
PBS	0	99	-	--	
ISIS 304801	25 mg/kg/wk for 2 wks	24	None	Full PS	135
ISIS 616468	25 mg/kg/wk for 2 wks	40	None	14 PS/6 PO	135

Example 27: Compound 56

20

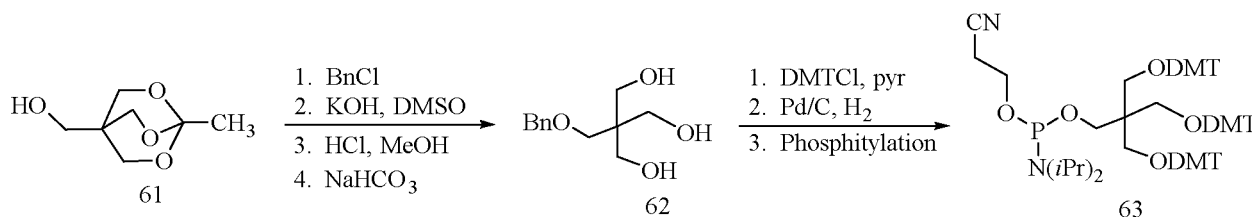


Compound 56 is commercially available from Glen Research or may be prepared according to published procedures reported by Shchepinov *et al.*, *Nucleic Acids Research*, 1997, 25(22), 4447-4454.

Example 28: Preparation of Compound 60

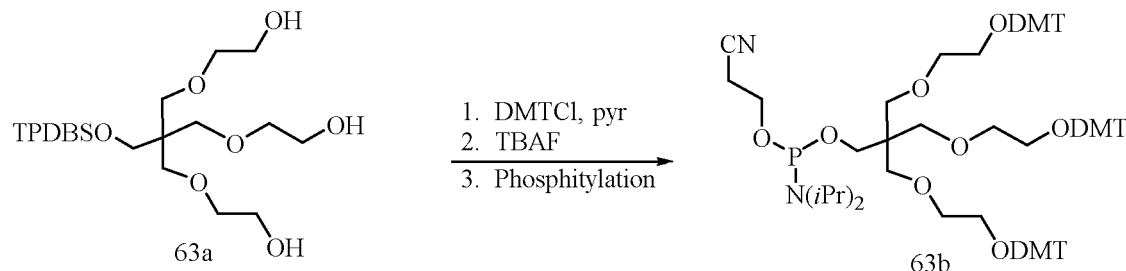
Compound 4 was prepared as per the procedures illustrated in Example 2. Compound 57 is commercially available. Compound 60 was confirmed by structural analysis.

Compound 57 is meant to be representative and not intended to be limiting as other monoprotected substituted or unsubstituted alkyl diols including but not limited to those presented in the specification herein can be used to prepare phosphoramidites having a predetermined composition.

Example 29: Preparation of Compound 63

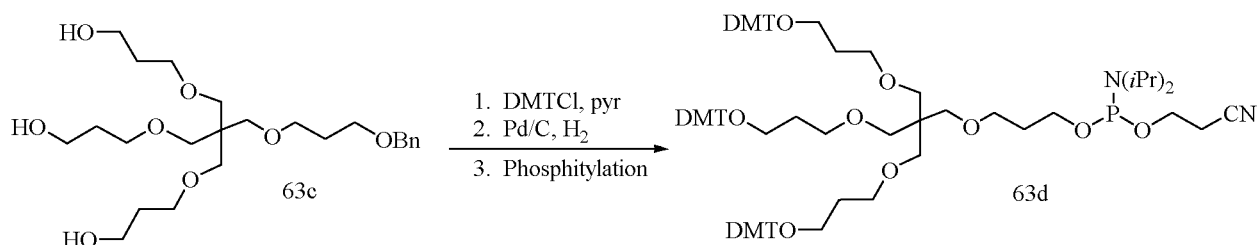
Compounds 61 and 62 are prepared using procedures similar to those reported by Tober *et al.*, *Eur. J. Org. Chem.*, 2013, 3, 566-577; and Jiang *et al.*, *Tetrahedron*, 2007, 63(19), 3982-3988.

Alternatively, Compound 63 is prepared using procedures similar to those reported in scientific and patent literature by Kim *et al.*, *Synlett*, 2003, 12, 1838-1840; and Kim *et al.*, published PCT International Application, WO 2004063208.

Example 30: Preparation of Compound 63b

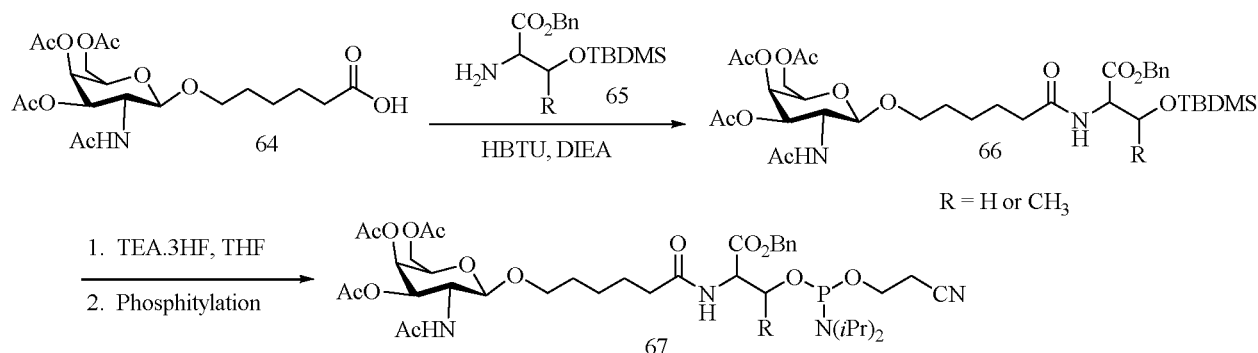
Compound 63a is prepared using procedures similar to those reported by Hanessian *et al.*, *Canadian Journal of Chemistry*, 1996, 74(9), 1731-1737.

Example 31: Preparation of Compound 63d



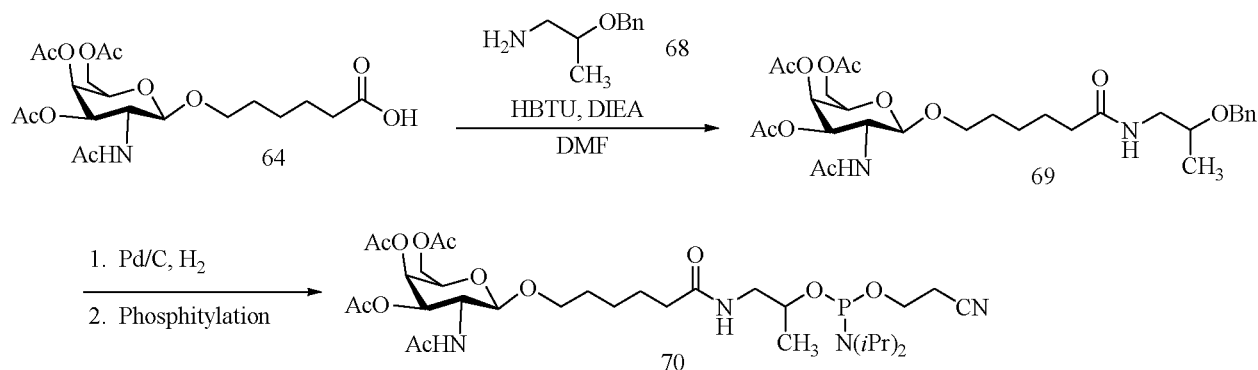
Compound 63c is prepared using procedures similar to those reported by Chen *et al.*, *Chinese Chemical Letters*, 1998, 9(5), 451-453.

5 Example 32: Preparation of Compound 67

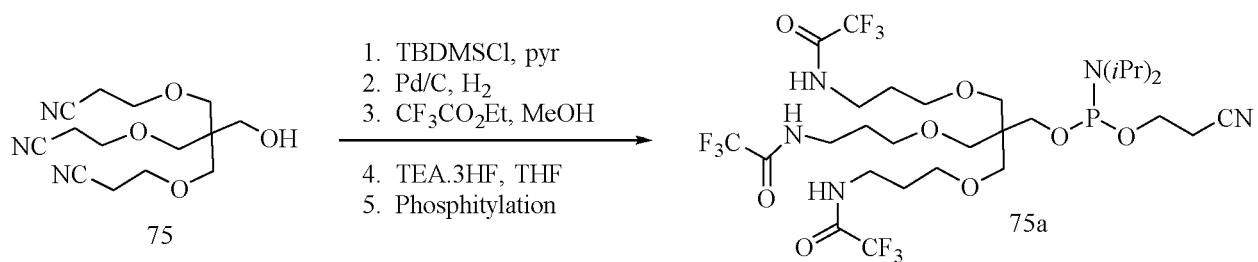


Compound 64 was prepared as per the procedures illustrated in Example 2. Compound 65 is prepared using procedures similar to those reported by Or *et al.*, published PCT International Application, WO 2009003009. The protecting groups used for Compound 65 are meant to be representative and not intended to be limiting as other protecting groups including but not limited to those presented in the specification herein can be used.

Example 33: Preparation of Compound 70

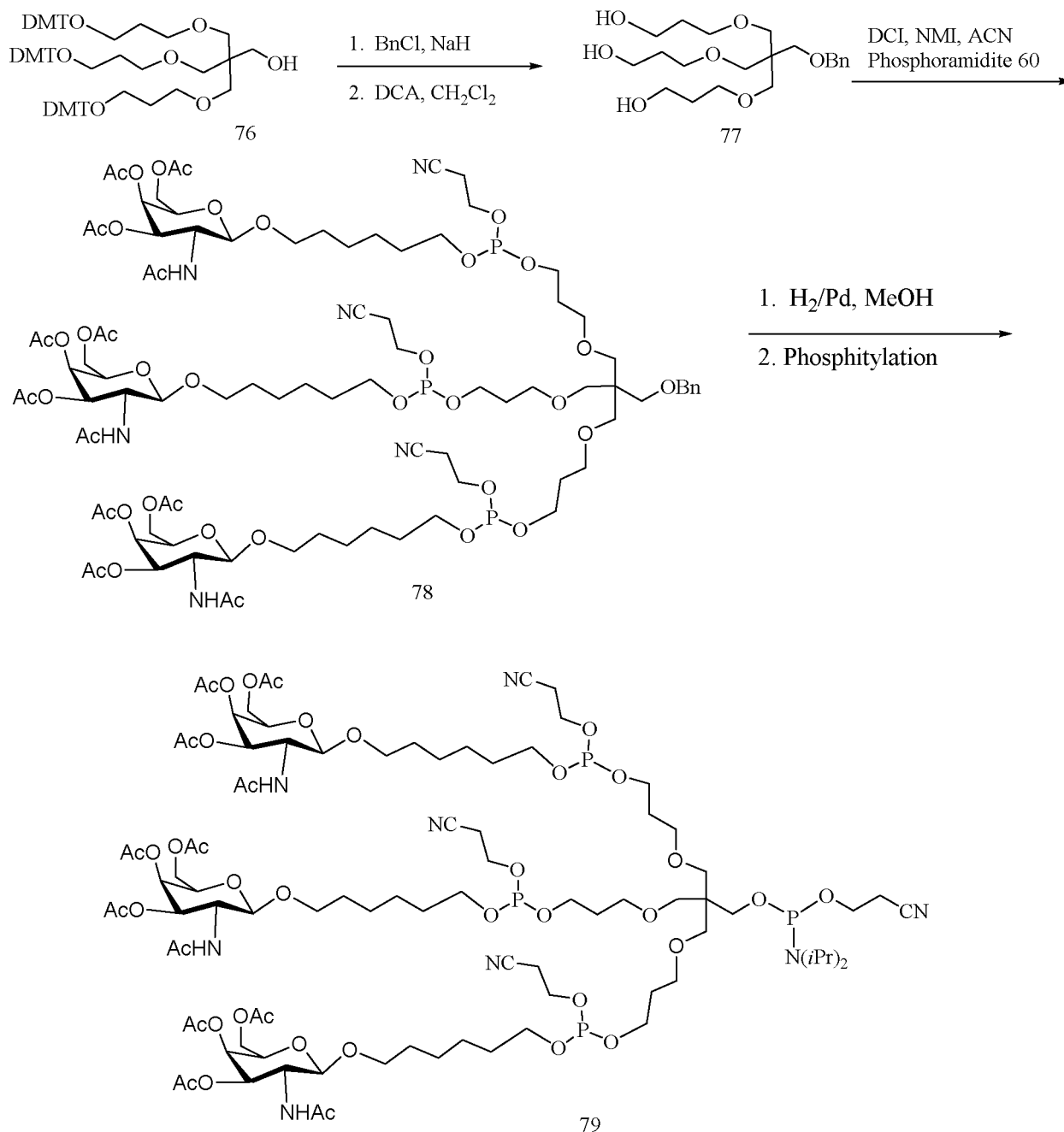


Compound 64 was prepared as per the procedures illustrated in Example 2. Compound 68 is commercially available. The protecting group used for Compound 68 is meant to be representative and not intended to be limiting as other protecting groups including but not limited to those presented in the specification herein can be used.

Example 34: Preparation of Compound 75a

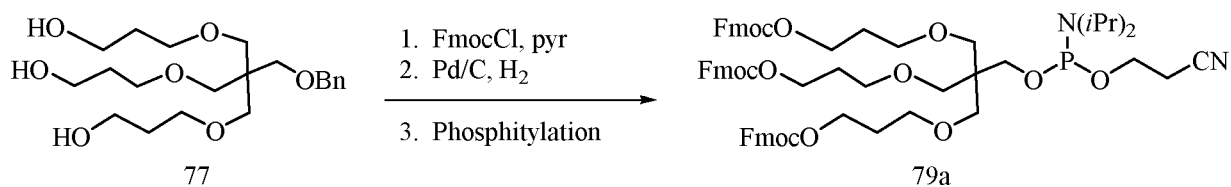
Compound 75 is prepared according to published procedures reported by Shchepinov *et al.*, *Nucleic*

5 *Acids Research*, 1997, 25(22), 4447-4454.

Example 35: Preparation of Compound 79

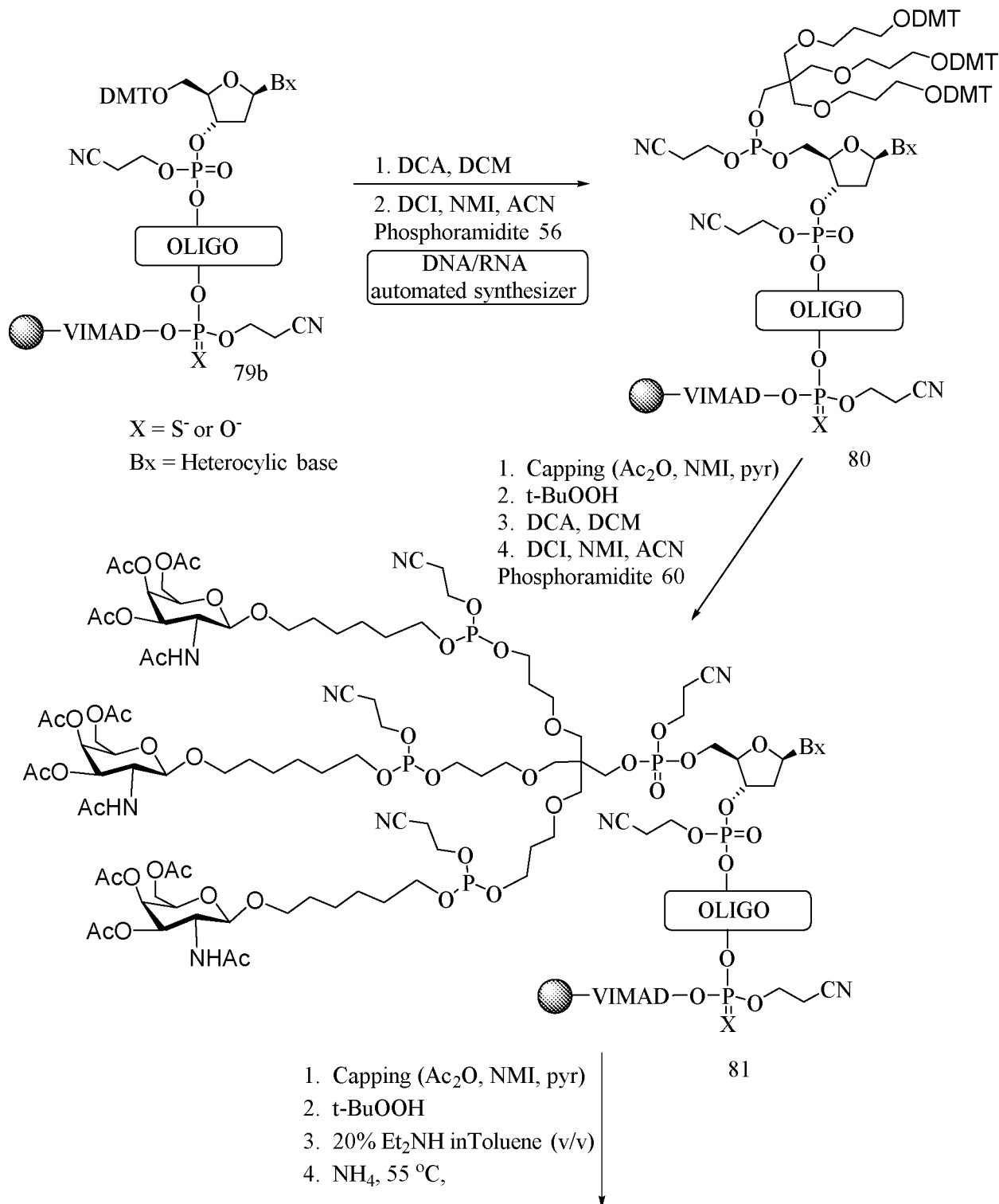
Compound 76 was prepared according to published procedures reported by Shchepinov *et al.*, *Nucleic Acids Research*, 1997, 25(22), 4447-4454.

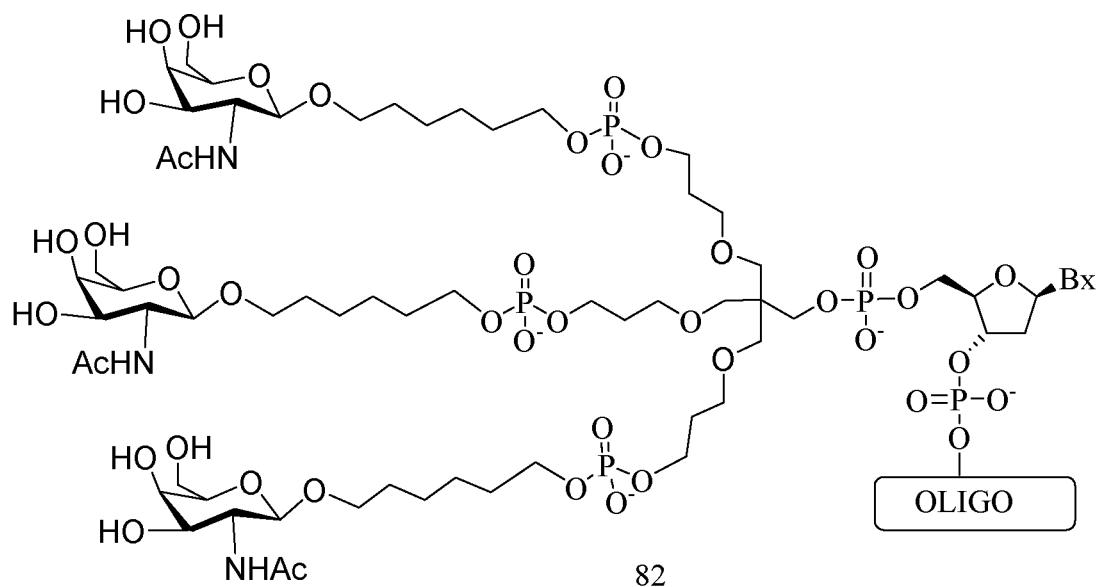
5

Example 36: Preparation of Compound 79a

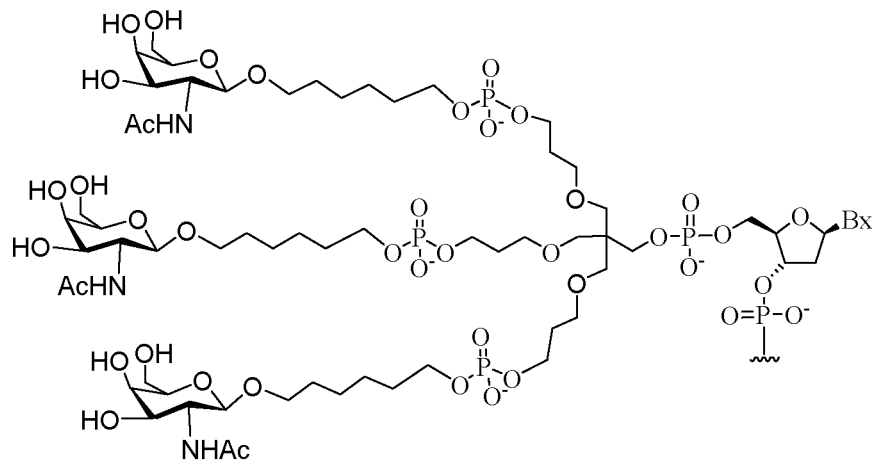
Compound 77 is prepared as per the procedures illustrated in Example 35.

Example 37: General method for the preparation of conjugated oligomeric compound 82 comprising a phosphodiester linked GalNAc₃-2 conjugate at 5' terminus *via* solid support (Method I)

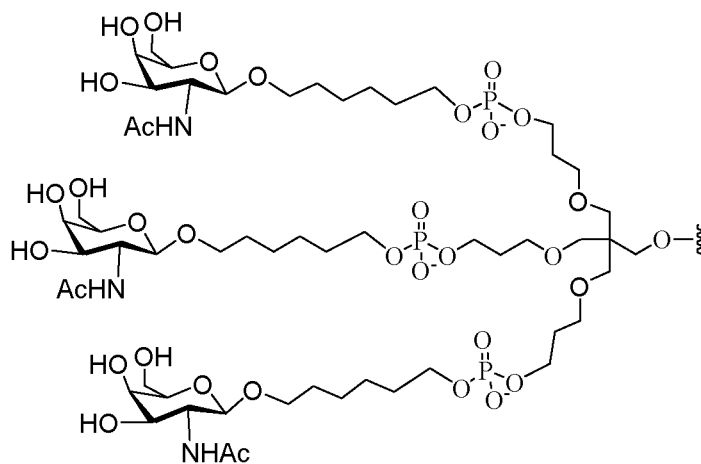




wherein GalNAc₃-2 has the structure:

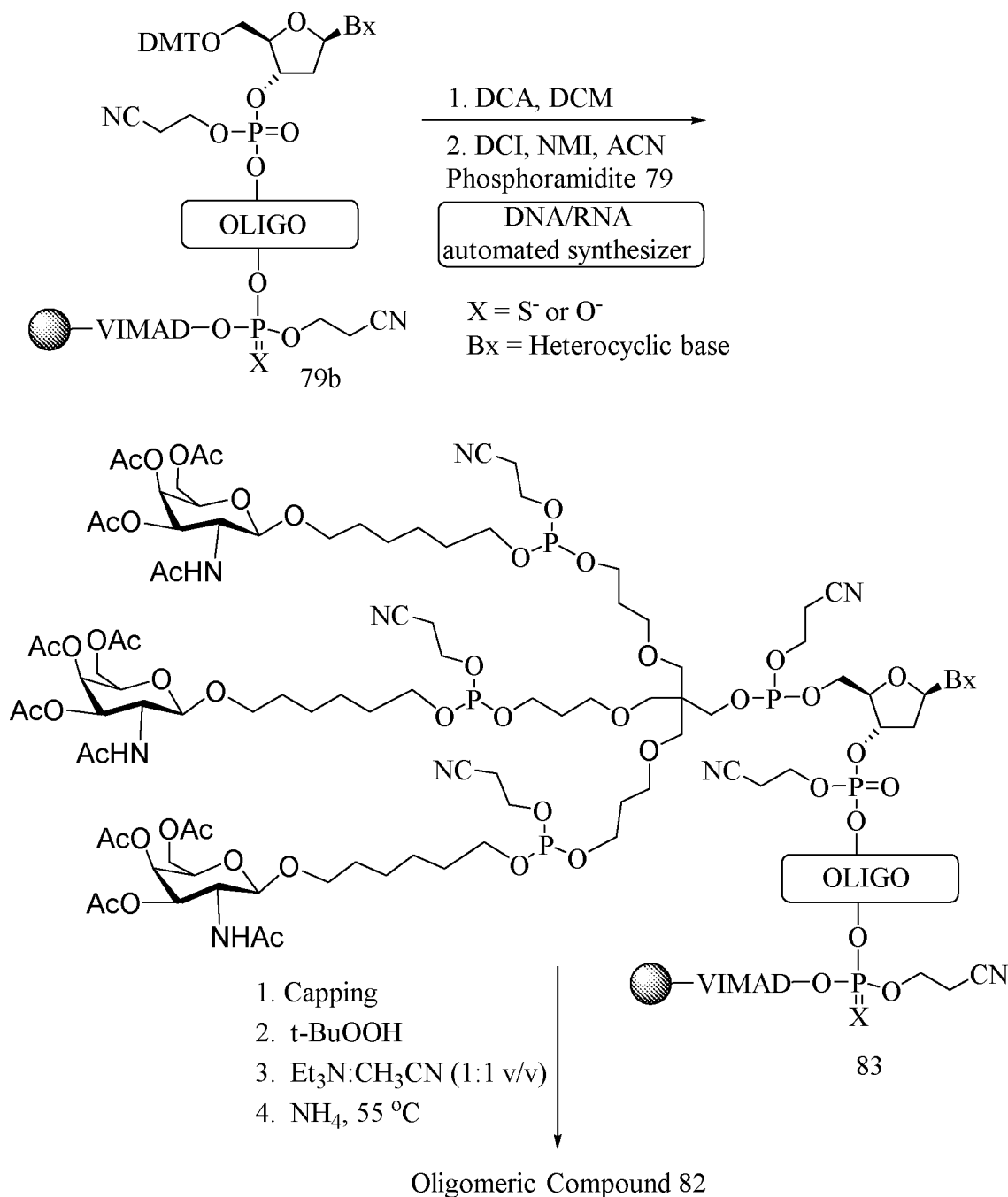


5 The GalNAc₃ cluster portion of the conjugate group GalNAc₃-2 (GalNAc₃-2_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. Wherein GalNAc₃-2_a has the formula:



The VIMAD-bound oligomeric compound 79b was prepared using standard procedures for automated DNA/RNA synthesis (see Dupouy *et al.*, *Angew. Chem. Int. Ed.*, 2006, 45, 3623-3627). The phosphoramidite Compounds 56 and 60 were prepared as per the procedures illustrated in Examples 27 and 28, respectively. The phosphoramidites illustrated are meant to be representative and not intended to be limiting as other phosphoramidite building blocks including but not limited those presented in the specification herein can be used to prepare an oligomeric compound having a phosphodiester linked conjugate group at the 5' terminus. The order and quantity of phosphoramidites added to the solid support can be adjusted to prepare the oligomeric compounds as described herein having any predetermined sequence and composition.

Example 38: Alternative method for the preparation of oligomeric compound 82 comprising a phosphodiester linked GalNAc₃-2 conjugate at 5' terminus (Method II)

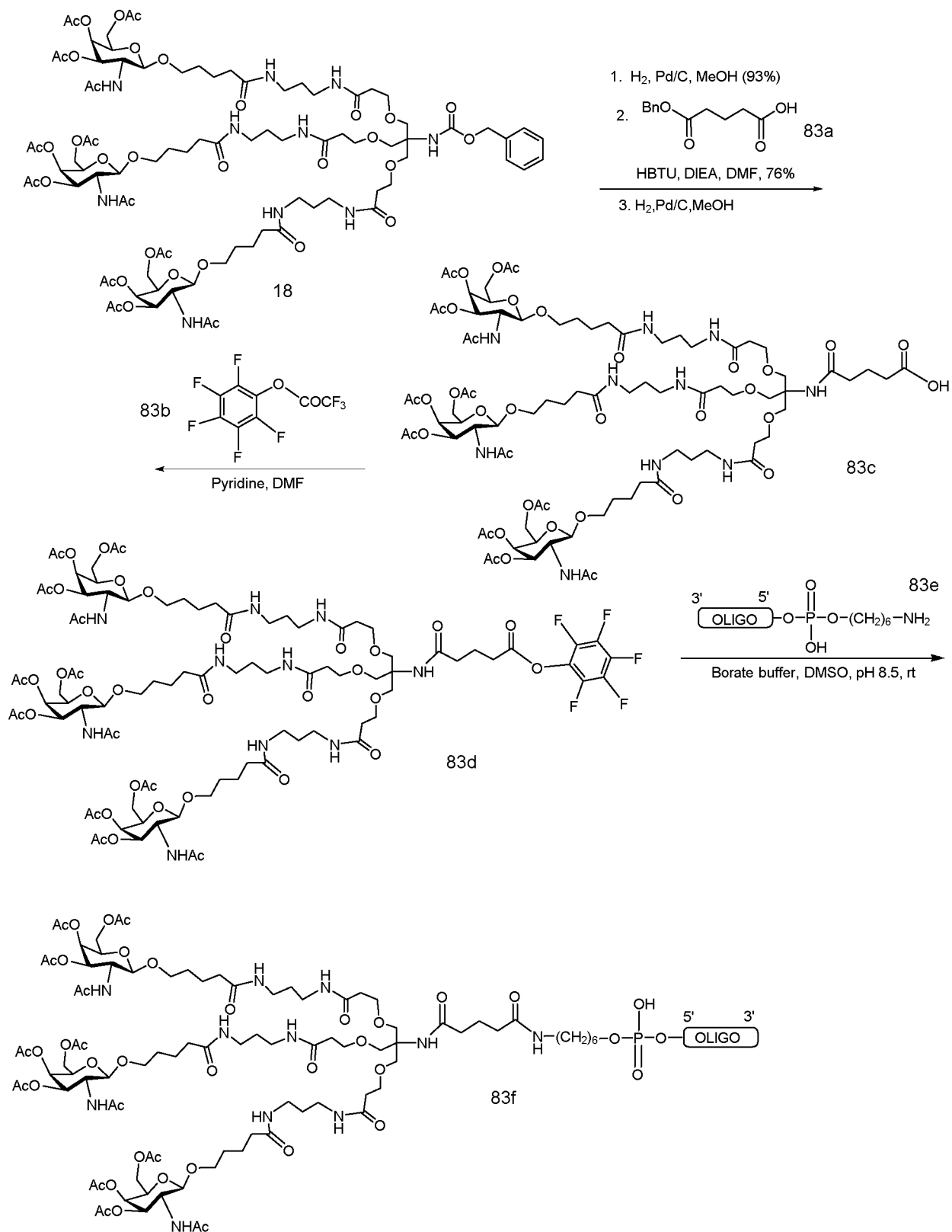


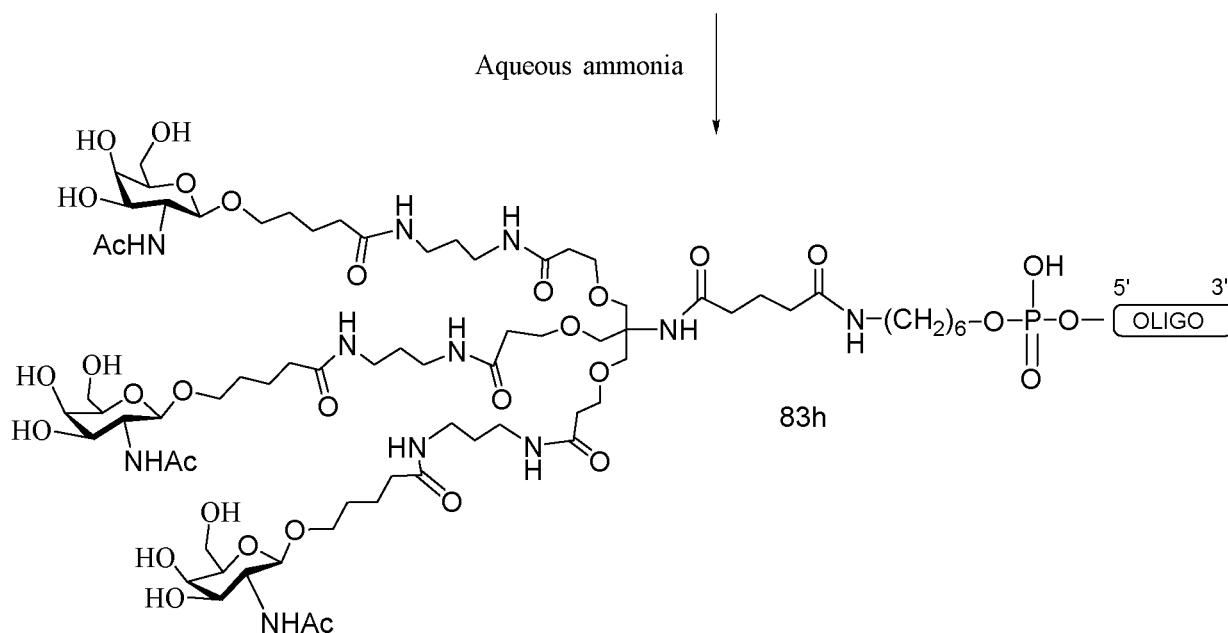
5 The VIMAD-bound oligomeric compound 79b was prepared using standard procedures for automated DNA/RNA synthesis (see Dupouy *et al.*, *Angew. Chem. Int. Ed.*, 2006, 45, 3623-3627). The GalNAc₃-2 cluster phosphoramidite, Compound 79 was prepared as per the procedures illustrated in Example 35. This alternative method allows a one-step installation of the phosphodiester linked GalNAc₃-2 conjugate to the oligomeric compound at the final step of the synthesis. The phosphoramidites illustrated are meant to

be representative and not intended to be limiting, as other phosphoramidite building blocks including but not limited to those presented in the specification herein can be used to prepare oligomeric compounds having a phosphodiester conjugate at the 5' terminus. The order and quantity of phosphoramidites added to the solid support can be adjusted to prepare the oligomeric compounds as described herein having any predetermined sequence and composition.

5

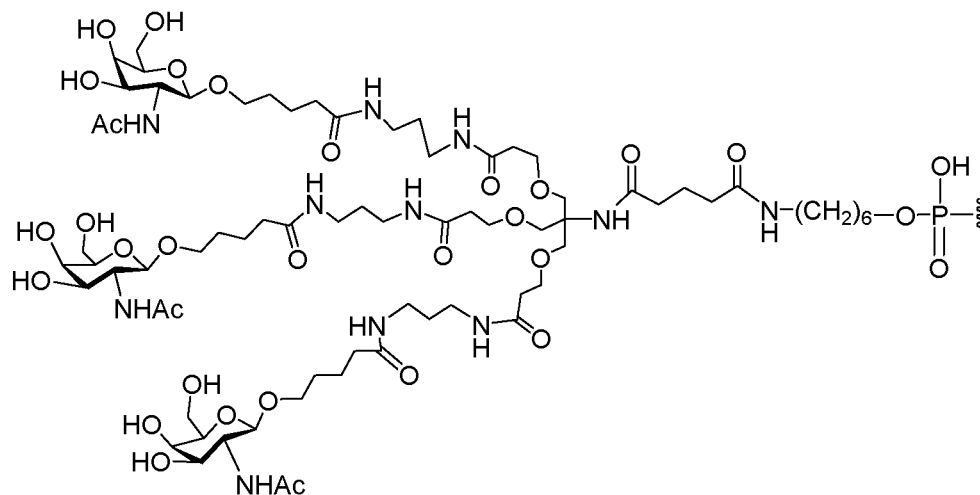
Example 39: General method for the preparation of oligomeric compound 83h comprising a GalNAc₃-3 Conjugate at the 5' Terminus (GalNAc₃-1 modified for 5' end attachment) *via* Solid Support





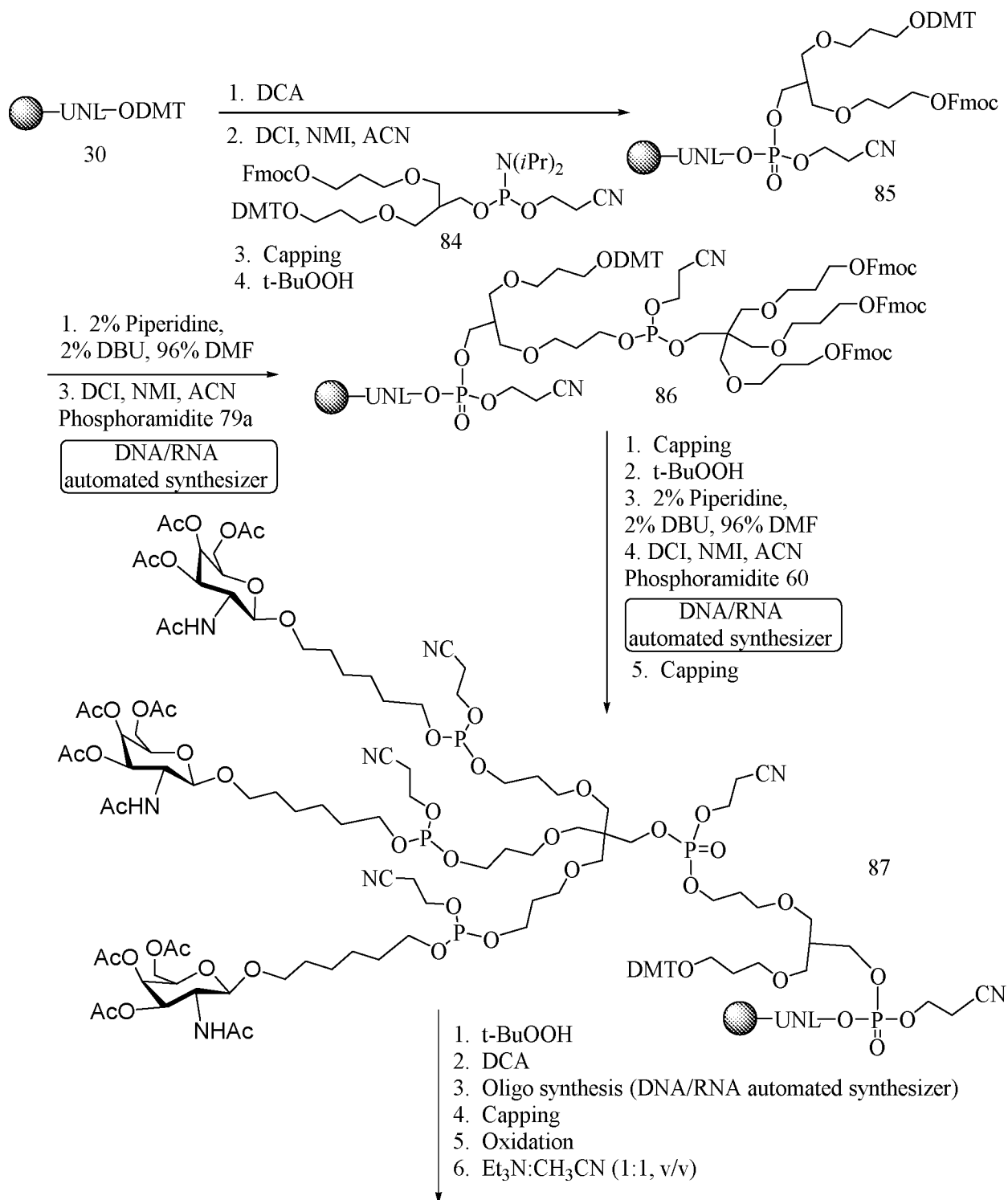
Compound 18 was prepared as per the procedures illustrated in Example 4. Compounds 83a and 83b are commercially available. Oligomeric Compound 83e comprising a phosphodiester linked hexylamine was prepared using standard oligonucleotide synthesis procedures. Treatment of the protected oligomeric compound with aqueous ammonia provided the 5'-GalNAc₃-3 conjugated oligomeric compound (83h).

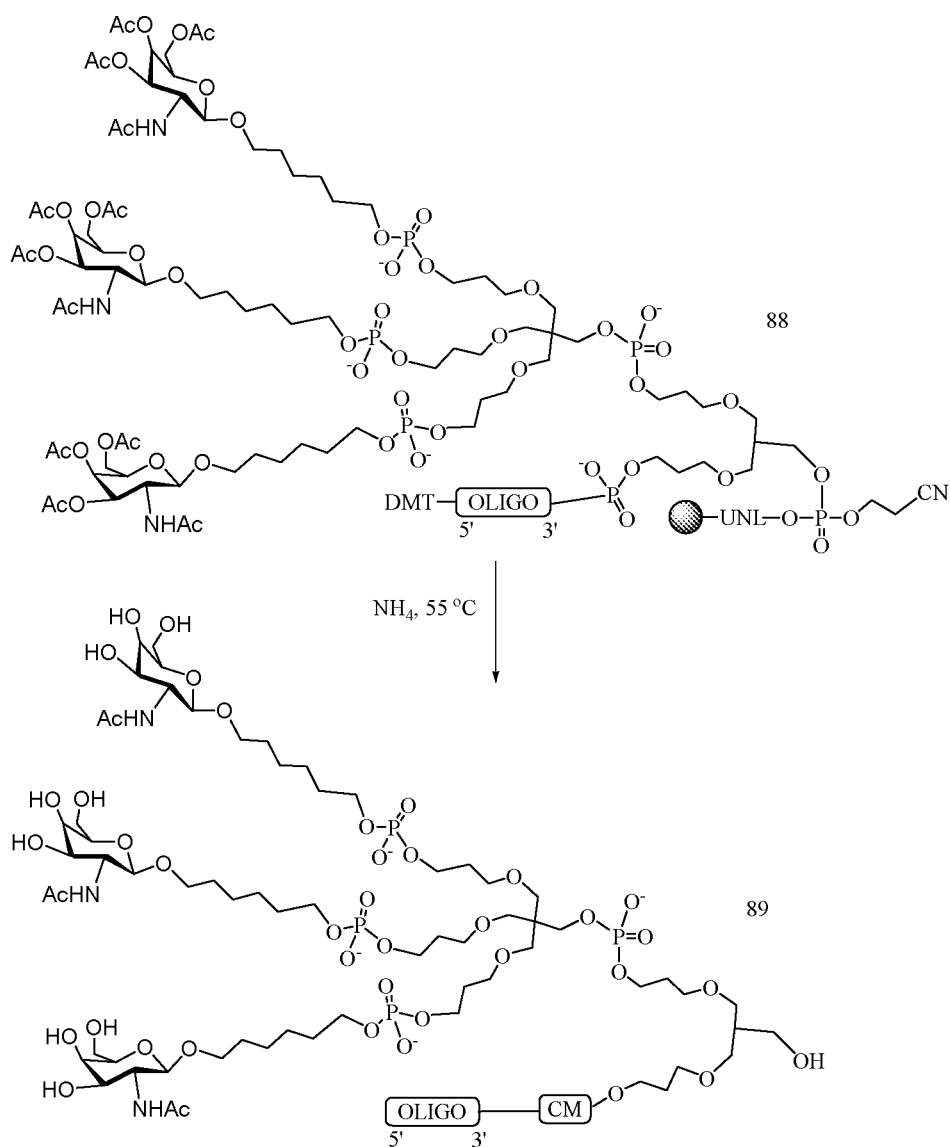
Wherein GalNAc₃-3 has the structure:



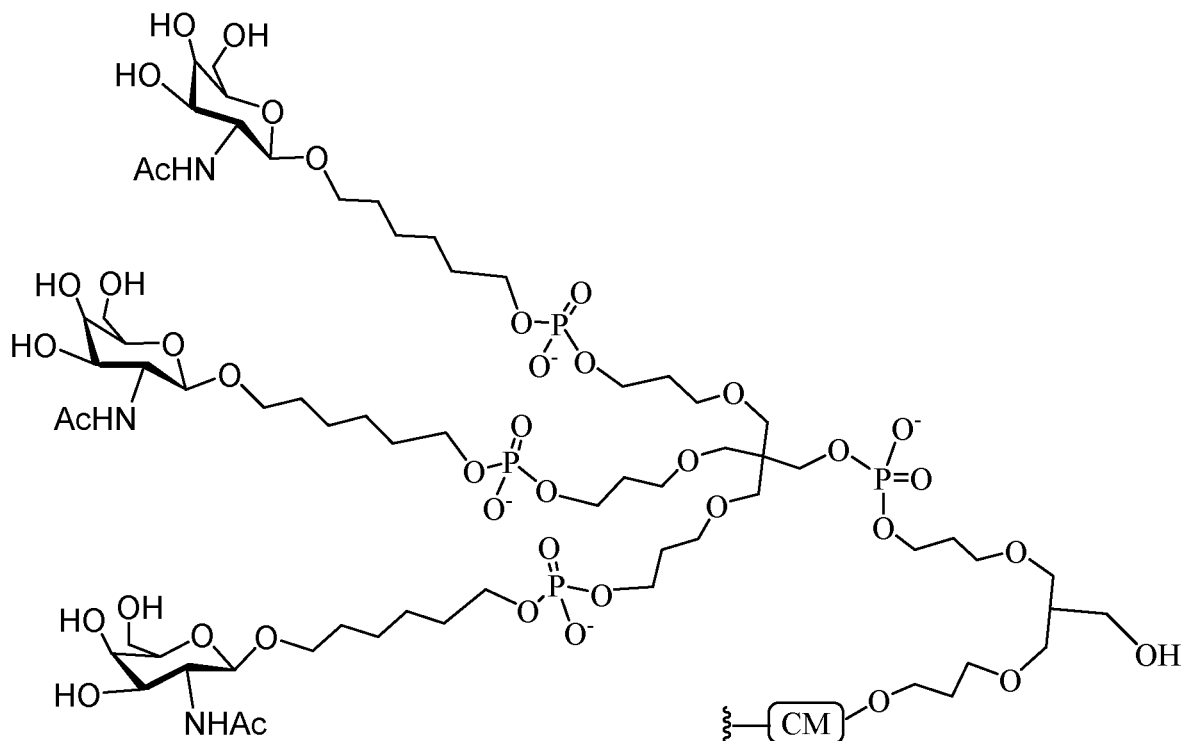
The GalNAc₃ cluster portion of the conjugate group GalNAc₃-3 (GalNAc₃-3_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. Wherein GalNAc₃-3_a has the formula:

Example 40: General method for the preparation of oligomeric compound 89 comprising a phosphodiester linked GalNAc₃-4 conjugate at the 3' terminus *via* solid support

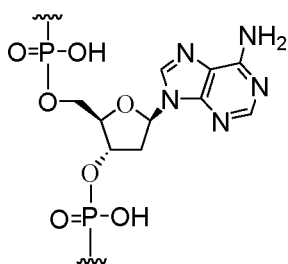




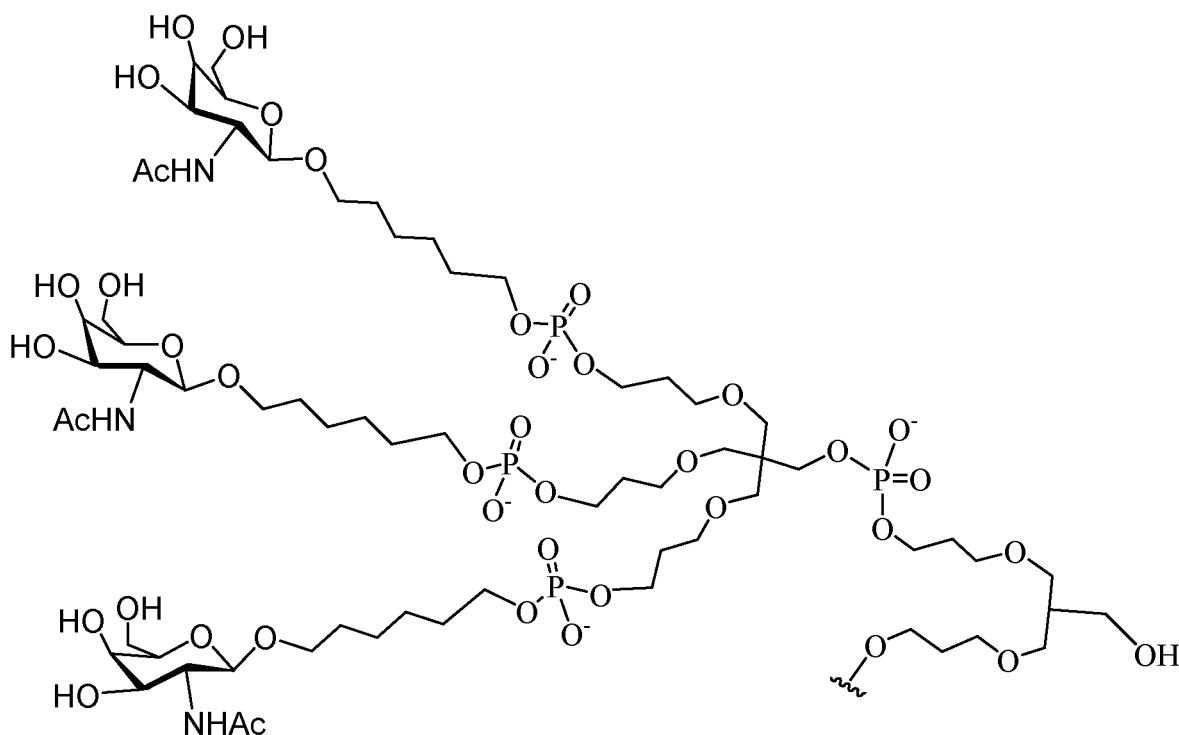
Wherein GalNAc₃-4 has the structure:



Wherein CM is a cleavable moiety. In certain embodiments, cleavable moiety is:



- 5 The GalNAc₃ cluster portion of the conjugate group GalNAc₃-4 (GalNAc₃-4_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. Wherein GalNAc₃-4_a has the formula:



The protected Unylinker functionalized solid support Compound 30 is commercially available.

5 Compound 84 is prepared using procedures similar to those reported in the literature (*see* Shchepinov *et al.*, *Nucleic Acids Research*, 1997, 25(22), 4447-4454; Shchepinov *et al.*, *Nucleic Acids Research*, 1999, 27, 3035-3041; and Horner *et al.*, *Nucleic Acids Research*, 1997, 25, 4842-4849).

10 The phosphoramidite building blocks, Compounds 60 and 79a are prepared as per the procedures illustrated in Examples 28 and 36. The phosphoramidites illustrated are meant to be representative and not intended to be limiting as other phosphoramidite building blocks can be used to prepare an oligomeric compound having a phosphodiester linked conjugate at the 3' terminus with a predetermined sequence and composition. The order and quantity of phosphoramidites added to the solid support can be adjusted to prepare the oligomeric compounds as described herein having any predetermined sequence and composition.

15 **Example 41: General method for the preparation of ASOs comprising a phosphodiester linked GalNAc₃-2 (see Example 37, Bx is adenine) conjugate at the 5' position *via* solid phase techniques (preparation of ISIS 661134)**

20 Unless otherwise stated, all reagents and solutions used for the synthesis of oligomeric compounds are purchased from commercial sources. Standard phosphoramidite building blocks and solid support are used for incorporation nucleoside residues which include for example T, A, G, and ^mC residues. Phosphoramidite compounds 56 and 60 were used to synthesize the phosphodiester linked GalNAc₃-2

conjugate at the 5' terminus. A 0.1 M solution of phosphoramidite in anhydrous acetonitrile was used for β -D-2'-deoxyribonucleoside and 2'-MOE.

The ASO syntheses were performed on ABI 394 synthesizer (1-2 μ mol scale) or on GE Healthcare Bioscience \ddot{A} KTA oligopilot synthesizer (40-200 μ mol scale) by the phosphoramidite coupling method on VIMAD solid support (110 μ mol/g, Guzaev *et al.*, 2003) packed in the column. For the coupling step, the phosphoramidites were delivered at a 4 fold excess over the initial loading of the solid support and phosphoramidite coupling was carried out for 10 min. All other steps followed standard protocols supplied by the manufacturer. A solution of 6% dichloroacetic acid in toluene was used for removing the dimethoxytrityl (DMT) groups from 5'-hydroxyl groups of the nucleotide. 4,5-Dicyanoimidazole (0.7 M) in anhydrous CH_3CN was used as activator during the coupling step. Phosphorothioate linkages were introduced by sulfurization with 0.1 M solution of xanthane hydride in 1:1 pyridine/ CH_3CN for a contact time of 3 minutes. A solution of 20% *tert*-butylhydroperoxide in CH_3CN containing 6% water was used as an oxidizing agent to provide phosphodiester internucleoside linkages with a contact time of 12 minutes.

After the desired sequence was assembled, the cyanoethyl phosphate protecting groups were deprotected using a 20% diethylamine in toluene (v/v) with a contact time of 45 minutes. The solid-support bound ASOs were suspended in aqueous ammonia (28-30 wt %) and heated at 55 $^{\circ}\text{C}$ for 6 h.

The unbound ASOs were then filtered and the ammonia was boiled off. The residue was purified by high pressure liquid chromatography on a strong anion exchange column (GE Healthcare Bioscience, Source 30Q, 30 μ m, 2.54 x 8 cm, A = 100 mM ammonium acetate in 30% aqueous CH_3CN , B = 1.5 M NaBr in A, 0-40% of B in 60 min, flow 14 mL min⁻¹, λ = 260 nm). The residue was desalted by HPLC on a reverse phase column to yield the desired ASOs in an isolated yield of 15-30% based on the initial loading on the solid support. The ASOs were characterized by ion-pair-HPLC coupled MS analysis with Agilent 1100 MSD system.

Table 34

ASO comprising a phosphodiester linked GalNAc₃-2 conjugate at the 5' position targeting SRB-1

ISIS No.	Sequence (5' to 3')	CalCd Mass	Observed Mass	SEQ ID No.
661134	GalNAc₃-2_a-o'A_{do}T_{ks}^mC_{ks}A_{ds}G_{ds}T_{ds}^mC_{ds}A_{ds}T_{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _k	6482.2	6481.6	141

Subscripts: "e" indicates 2'-MOE modified nucleoside; "d" indicates β -D-2'-deoxyribonucleoside; "k" indicates 6'-(*S*)-CH₃ bicyclic nucleoside (e.g. cEt); "s" indicates phosphorothioate internucleoside linkages (PS); "o" indicates phosphodiester internucleoside linkages (PO); and "o'" indicates -O-P(=O)(OH)-. Superscript "m" indicates 5-methylcytosines. The structure of GalNAc₃-2_a is shown in Example 37.

Example 42: General method for the preparation of ASOs comprising a GalNAc₃-3 conjugate at the 5' position *via* solid phase techniques (preparation of ISIS 661166)

The synthesis for ISIS 661166 was performed using similar procedures as illustrated in Examples 39 and 41.

ISIS 661166 is a 5-10-5 MOE gapmer, wherein the 5' position comprises a GalNAc₃-3 conjugate. The ASO was characterized by ion-pair-HPLC coupled MS analysis with Agilent 1100 MSD system.

Table 34a
ASO comprising a GalNAc₃-3 conjugate at the 5' position via a hexylamino phosphodiester linkage targeting Malat-1

ISIS No.	Sequence (5' to 3')	Conjugate	Calcd Mass	Observed Mass	SEQ ID No.
661166	5'-GalNAc ₃ -3 _{a-o'} ^m C _{es} G _{es} G _{es} T _{es} G _{es} ^m C _{ds} A _{ds} A _{ds} G _{ds} G _{ds} ^m C _{ds} T _{ds} T _{ds} A _{ds} G _{ds} G _{es} A _{es} A _{es} T _{es} T _e	5'-GalNAc ₃ -3	8992.16	8990.51	142

Subscripts: "e" indicates 2'-MOE modified nucleoside; "d" indicates β-D-2'-deoxyribonucleoside; "s" indicates phosphorothioate internucleoside linkages (PS); "o" indicates phosphodiester internucleoside linkages (PO); and "o'" indicates -O-P(=O)(OH)-. Superscript "m" indicates 5-methylcytosines. The structure of "5'-GalNAc₃-3a" is shown in Example 39.

Example 43: Dose-dependent study of phosphodiester linked GalNAc₃-2 (see examples 37 and 41, Bx is adenine) at the 5' terminus targeting SRB-1 *in vivo*

ISIS 661134 (see Example 41) comprising a phosphodiester linked GalNAc₃-2 conjugate at the 5' terminus was tested in a dose-dependent study for antisense inhibition of SRB-1 in mice. Unconjugated ISIS 440762 and 651900 (GalNAc₃-1 conjugate at 3' terminus, see Example 9) were included in the study for comparison and are described previously in Table 17.

Treatment

Six week old male Balb/c mice (Jackson Laboratory, Bar Harbor, ME) were injected subcutaneously once at the dosage shown below with ISIS 440762, 651900, 661134 or with PBS treated control. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration to determine the liver SRB-1 mRNA levels using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. SRB-1 mRNA levels were determined relative to total RNA (using Ribogreen), prior to normalization to PBS-treated control. The results below are presented as the average percent of SRB-1 mRNA levels for each treatment group, normalized to PBS-treated control and is denoted as "% PBS". The ED₅₀s were measured using similar methods as described previously and are presented below.

As illustrated in Table 35, treatment with antisense oligonucleotides lowered SRB-1 mRNA levels in a dose-dependent manner. Indeed, the antisense oligonucleotides comprising the phosphodiester linked GalNAc₃-2 conjugate at the 5' terminus (ISIS 661134) or the GalNAc₃-1 conjugate linked at the 3' terminus (ISIS 651900) showed substantial improvement in potency compared to the unconjugated antisense oligonucleotide (ISIS 440762). Further, ISIS 661134, which comprises the phosphodiester linked GalNAc₃-2 conjugate at the 5' terminus was equipotent compared to ISIS 651900, which comprises the GalNAc₃-1 conjugate at the 3' terminus.

Table 35

ASOs containing GalNAc₃-1 or GalNAc₃-2 targeting SRB-1

ISIS No.	Dosage (mg/kg)	SRB-1 mRNA levels (% PBS)	ED ₅₀ (mg/kg)	Conjugate	SEQ ID No.
PBS	0	100	--	--	
440762	0.2	116	2.58	No conjugate	137
	0.7	91			
	2	69			
	7	22			
	20	5			
651900	0.07	95	0.26	3' GalNAc ₃ -1	138
	0.2	77			
	0.7	28			
	2	11			
	7	8			
661134	0.07	107	0.25	5' GalNAc ₃ -2	141
	0.2	86			
	0.7	28			
	2	10			
	7	6			

Structures for 3' GalNAc₃-1 and 5' GalNAc₃-2 were described previously in Examples 9 and 37.

Pharmacokinetics Analysis (PK)

The PK of the ASOs from the high dose group (7 mg/kg) was examined and evaluated in the same manner as illustrated in Example 20. Liver sample was minced and extracted using standard protocols. The full length metabolites of 661134 (5' GalNAc₃-2) and ISIS 651900 (3' GalNAc₃-1) were identified and their masses were confirmed by high resolution mass spectrometry analysis. The results showed that the major metabolite detected for the ASO comprising a phosphodiester linked GalNAc₃-2 conjugate at the 5' terminus (ISIS 661134) was ISIS 440762 (data not shown). No additional metabolites, at a detectable level, were observed. Unlike its counterpart, additional metabolites similar to those reported previously in Table 23a were observed for the ASO having the GalNAc₃-1 conjugate at the 3' terminus (ISIS 651900). These results suggest that having the phosphodiester linked GalNAc₃-1 or GalNAc₃-2 conjugate may improve the PK profile of ASOs without compromising their potency.

Example 44: Effect of PO/PS linkages on antisense inhibition of ASOs comprising GalNAc₃-1 conjugate (see Example 9) at the 3' terminus targeting SRB-1

ISIS 655861 and 655862 comprising a GalNAc₃-1 conjugate at the 3' terminus each targeting SRB-1 were tested in a single administration study for their ability to inhibit SRB-1 in mice. The parent unconjugated compound, ISIS 353382 was included in the study for comparison.

The ASOs are 5-10-5 MOE gapmers, wherein the gap region comprises ten 2'-deoxyribonucleosides and each wing region comprises five 2'-MOE modified nucleosides. The ASOs were prepared using similar methods as illustrated previously in Example 19 and are described Table 36, below.

Table 36

Modified ASOs comprising GalNAc₃-1 conjugate at the 3' terminus targeting SRB-1

ISIS No.	Sequence (5' to 3')	Chemistry	SEQ ID No.
353382 (parent)	G ^{es} _{es} C ^{es} _{es} T ^{es} _{es} T ^{es} _{es} C ^{es} _{es} A ^{ds} _{ds} G ^{ds} _{ds} T ^{ds} _{ds} C ^{ds} _{ds} A ^{ds} _{ds} T ^{ds} _{ds} G ^{ds} _{ds} A ^{ds} _{ds} mC ^{ds} _{ds} T ^{ds} _{ds} T ^{es} _{es} mC ^{es} _{es} mC ^{es} _{es} T ^{es} _{es} T ^e _e	Full PS no conjugate	143
655861	G ^{es} _{es} C ^{es} _{es} T ^{es} _{es} T ^{es} _{es} C ^{es} _{es} A ^{ds} _{ds} G ^{ds} _{ds} T ^{ds} _{ds} C ^{ds} _{ds} A ^{ds} _{ds} T ^{ds} _{ds} G ^{ds} _{ds} A ^{ds} _{ds} mC ^{ds} _{ds} T ^{ds} _{ds} T ^{es} _{es} mC ^{es} _{es} mC ^{es} _{es} T ^{eo} _{eo} A_{do}-GalNAc₃-1_a	Full PS with GalNAc ₃ -1 conjugate	144
655862	G ^{es} _{es} C ^{eo} _{eo} T ^{eo} _{eo} T ^{eo} _{eo} mC ^{eo} _{eo} A ^{ds} _{ds} G ^{ds} _{ds} T ^{ds} _{ds} C ^{ds} _{ds} A ^{ds} _{ds} T ^{ds} _{ds} G ^{ds} _{ds} A ^{ds} _{ds} mC ^{ds} _{ds} T ^{ds} _{ds} T ^{eo} _{eo} mC ^{eo} _{eo} mC ^{es} _{es} T ^{eo} _{eo} A_{do}-GalNAc₃-1_a	Mixed PS/PO with GalNAc ₃ -1 conjugate	144

Subscripts: “e” indicates 2'-MOE modified nucleoside; “d” indicates β-D-2'-deoxyribonucleoside; “s” indicates phosphorothioate internucleoside linkages (PS); “o” indicates phosphodiester internucleoside linkages (PO); and “o” indicates -O-P(=O)(OH)-. Superscript “m” indicates 5-methylcytosines. The structure of “GalNAc₃-1” is shown in Example 9.

Treatment

Six week old male Balb/c mice (Jackson Laboratory, Bar Harbor, ME) were injected subcutaneously once at the dosage shown below with ISIS 353382, 655861, 655862 or with PBS treated control. Each treatment group consisted of 4 animals. Prior to the treatment as well as after the last dose, blood was drawn from each mouse and plasma samples were analyzed. The mice were sacrificed 72 hours following the final administration to determine the liver SRB-1 mRNA levels using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. SRB-1 mRNA levels were determined relative to total RNA (using Ribogreen), prior to normalization to PBS-treated control. The results below are presented as the average percent of SRB-1 mRNA levels for each treatment group, normalized to PBS-treated control and is denoted as “% PBS”. The ED₅₀s were measured using similar methods as described previously and are reported below.

As illustrated in Table 37, treatment with antisense oligonucleotides lowered SRB-1 mRNA levels in a dose-dependent manner compared to PBS treated control. Indeed, the antisense oligonucleotides

comprising the GalNAc₃-1 conjugate at the 3' terminus (ISIS 655861 and 655862) showed substantial improvement in potency comparing to the unconjugated antisense oligonucleotide (ISIS 353382). Further, ISIS 655862 with mixed PS/PO linkages showed an improvement in potency relative to full PS (ISIS 655861).

5

Table 37
Effect of PO/PS linkages on antisense inhibition of ASOs
comprising GalNAc₃-1 conjugate at 3' terminus targeting SRB-1

ISIS No.	Dosage (mg/kg)	SRB-1 mRNA levels (% PBS)	ED ₅₀ (mg/kg)	Chemistry	SEQ ID No.
PBS	0	100	--	--	
353382 (parent)	3	76.65	10.4	Full PS without conjugate	143
	10	52.40			
	30	24.95			
655861	0.5	81.22	2.2	Full PS with GalNAc ₃ -1 conjugate	144
	1.5	63.51			
	5	24.61			
	15	14.80			
655862	0.5	69.57	1.3	Mixed PS/PO with GalNAc ₃ -1 conjugate	144
	1.5	45.78			
	5	19.70			
	15	12.90			

10

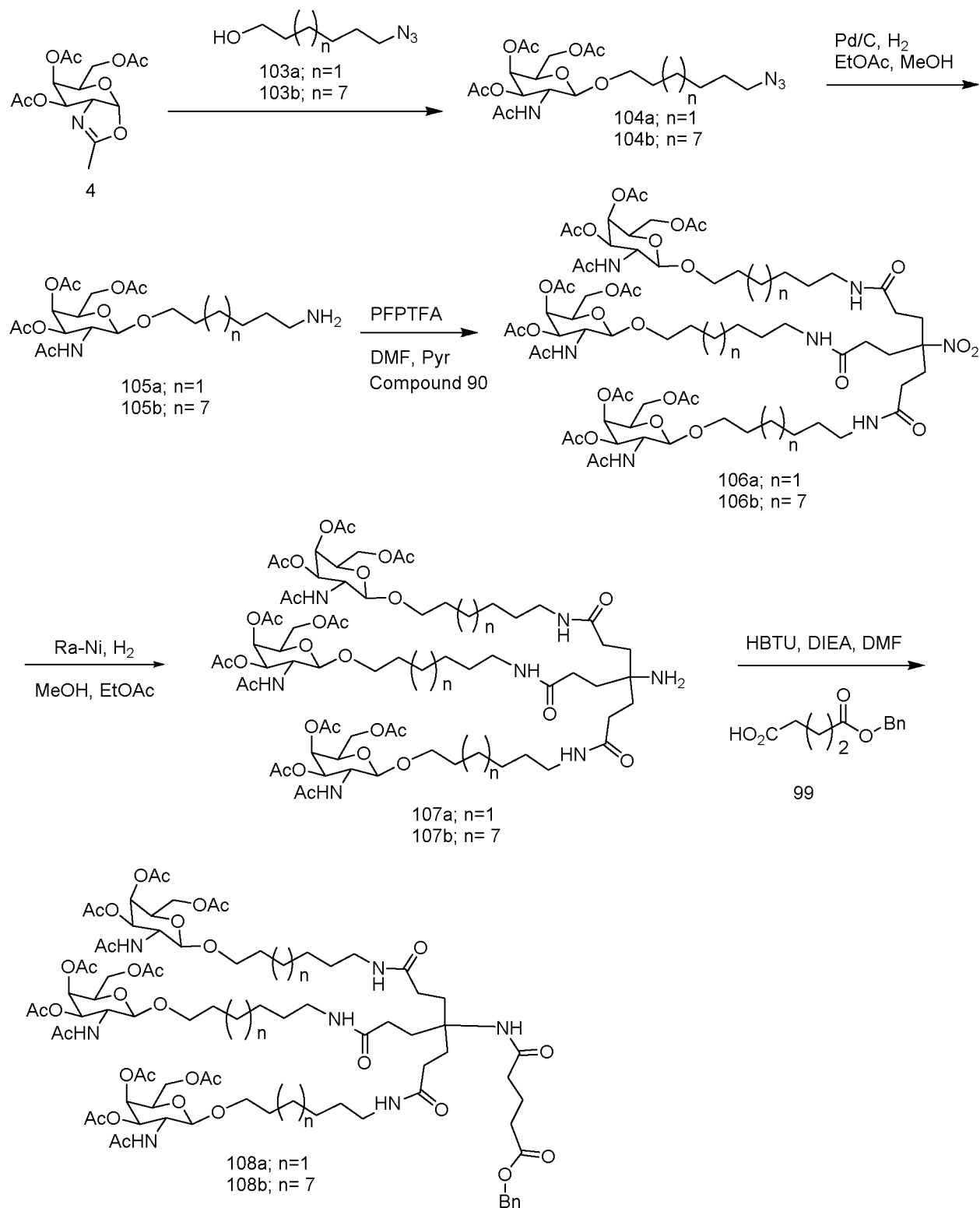
Liver transaminase levels, alanine aminotransferase (ALT) and aspartate aminotransferase (AST), in serum were measured relative to saline injected mice using standard protocols. Organ weights were also evaluated. The results demonstrated that no elevation in transaminase levels (Table 38) or organ weights (data not shown) were observed in mice treated with ASOs compared to PBS control. Further, the ASO with mixed PS/PO linkages (ISIS 655862) showed similar transaminase levels compared to full PS (ISIS 655861).

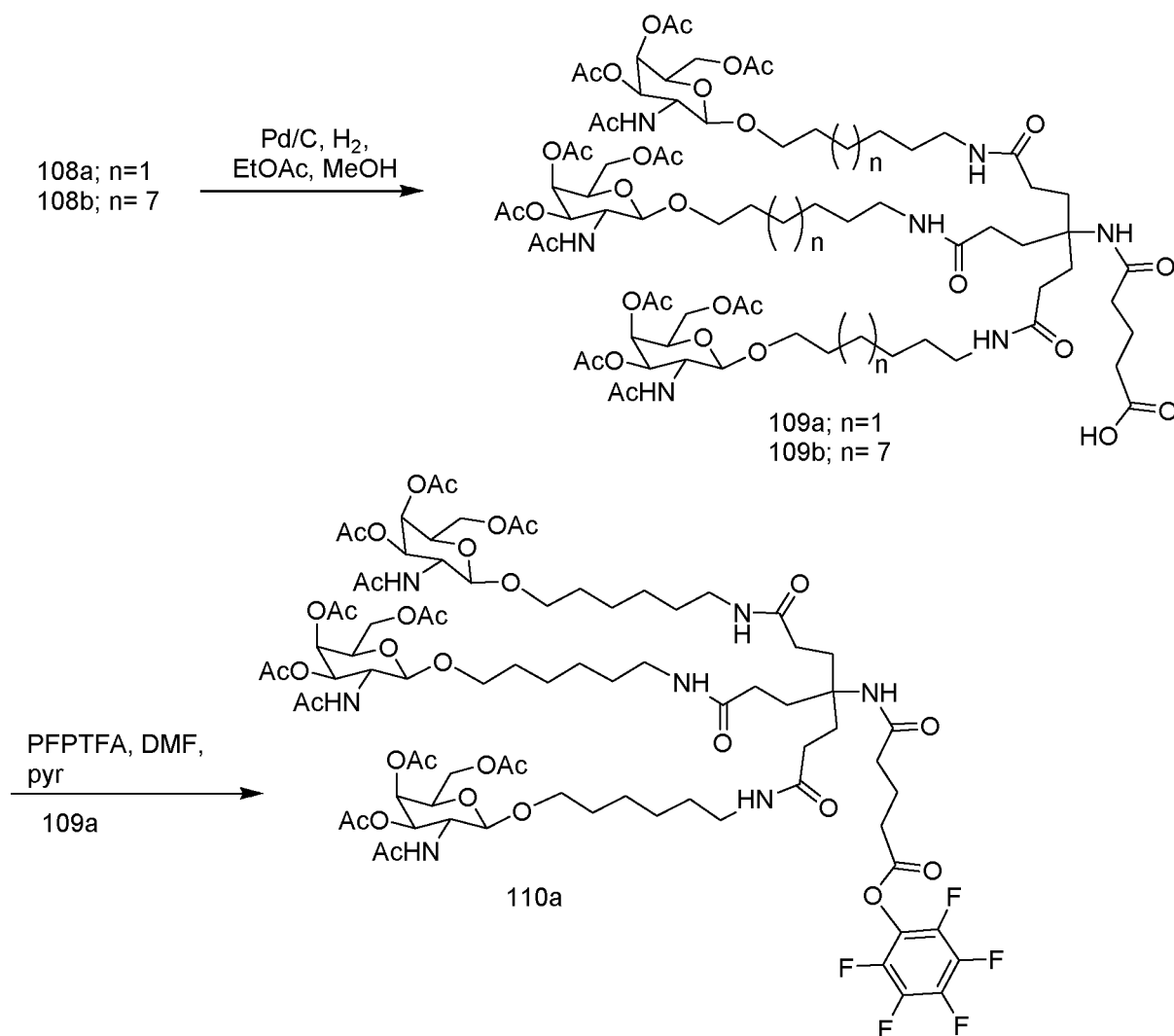
15

Table 38
Effect of PO/PS linkages on transaminase levels of ASOs
comprising GalNAc₃-1 conjugate at 3' terminus targeting SRB-1

ISIS No.	Dosage (mg/kg)	ALT (U/L)	AST (U/L)	Chemistry	SEQ ID No.
PBS	0	28.5	65	--	
353382 (parent)	3	50.25	89	Full PS without conjugate	143
	10	27.5	79.3		
	30	27.3	97		
655861	0.5	28	55.7	Full PS with GalNAc₃-1	144
	1.5	30	78		
	5	29	63.5		
	15	28.8	67.8		
655862	0.5	50	75.5	Mixed PS/PO with GalNAc₃-1	144
	1.5	21.7	58.5		
	5	29.3	69		

	15	22	61		
--	----	----	----	--	--

Example 45: Preparation of PFP Ester, Compound 110a



Compound 4 (9.5g, 28.8 mmols) was treated with compound 103a or 103b (38 mmols), individually, and TMSOTf (0.5 eq.) and molecular sieves in dichloromethane (200 mL), and stirred for 16 hours at room temperature. At that time, the organic layer was filtered thru celite, then washed with sodium bicarbonate, water and brine. The organic layer was then separated and dried over sodium sulfate, filtered and reduced under reduced pressure. The resultant oil was purified by silica gel chromatography (2%-->10% methanol/dichloromethane) to give compounds 104a and 104b in >80% yield. LCMS and proton NMR was consistent with the structure.

Compounds 104a and 104b were treated to the same conditions as for compounds 100a-d (Example 47), to give compounds 105a and 105b in >90% yield. LCMS and proton NMR was consistent with the structure.

Compounds 105a and 105b were treated, individually, with compound 90 under the same conditions as for compounds 901a-d, to give compounds 106a (80%) and 106b (20%). LCMS and proton NMR was consistent with the structure.

Compounds 106a and 106b were treated to the same conditions as for compounds 96a-d (Example 47), to give 107a (60%) and 107b (20%). LCMS and proton NMR was consistent with the structure.

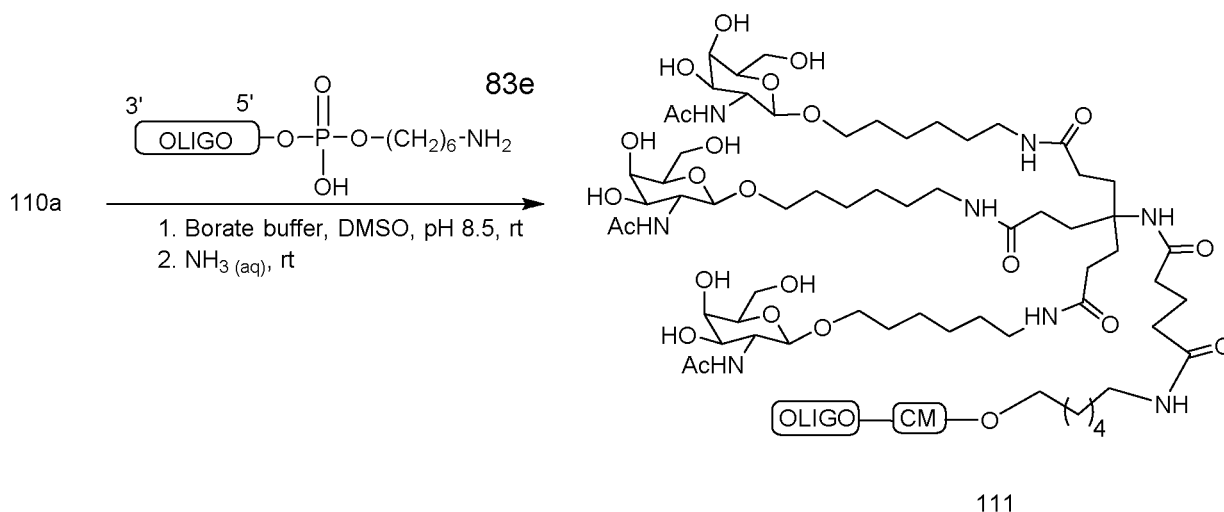
Compounds 107a and 107b were treated to the same conditions as for compounds 97a-d (Example 47), to give compounds 108a and 108b in 40-60% yield. LCMS and proton NMR was consistent with the structure.

Compounds 108a (60%) and 108b (40%) were treated to the same conditions as for compounds 100a-d (Example 47), to give compounds 109a and 109b in >80% yields. LCMS and proton NMR was consistent with the structure.

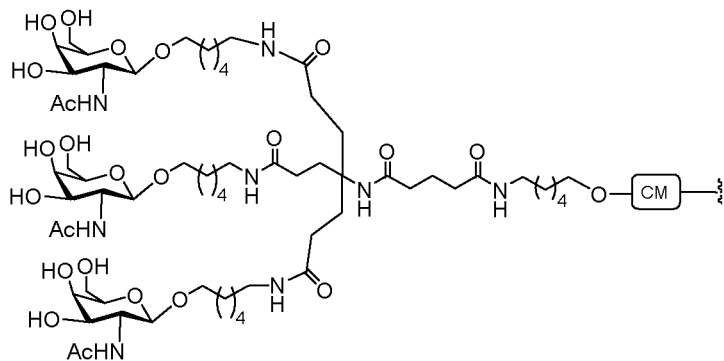
Compound 109a was treated to the same conditions as for compounds 101a-d (Example 47), to give Compound 110a in 30-60% yield. LCMS and proton NMR was consistent with the structure. Alternatively, Compound 110b can be prepared in a similar manner starting with Compound 109b.

Example 46: General Procedure for Conjugation with PFP Esters (Oligonucleotide 111); Preparation of ISIS 666881 (GalNAc₃-10)

A 5'-hexylamino modified oligonucleotide was synthesized and purified using standard solid-phase oligonucleotide procedures. The 5'-hexylamino modified oligonucleotide was dissolved in 0.1 M sodium tetraborate, pH 8.5 (200 μ L) and 3 equivalents of a selected PFP esterified GalNAc₃ cluster dissolved in DMSO (50 μ L) was added. If the PFP ester precipitated upon addition to the ASO solution DMSO was added until all PFP ester was in solution. The reaction was complete after about 16 h of mixing at room temperature. The resulting solution was diluted with water to 12 mL and then spun down at 3000 rpm in a spin filter with a mass cut off of 3000 Da. This process was repeated twice to remove small molecule impurities. The solution was then lyophilized to dryness and redissolved in concentrated aqueous ammonia and mixed at room temperature for 2.5 h followed by concentration *in vacuo* to remove most of the ammonia. The conjugated oligonucleotide was purified and desalted by RP-HPLC and lyophilized to provide the GalNAc₃ conjugated oligonucleotide.



Oligonucleotide 111 is conjugated with GalNAc₃-10. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-10 (GalNAc₃-10_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)- as shown in the oligonucleotide (ISIS 666881) synthesized with GalNAc₃-10 below. The structure of GalNAc₃-10 (GalNAc₃-10_a-CM-) is shown below:



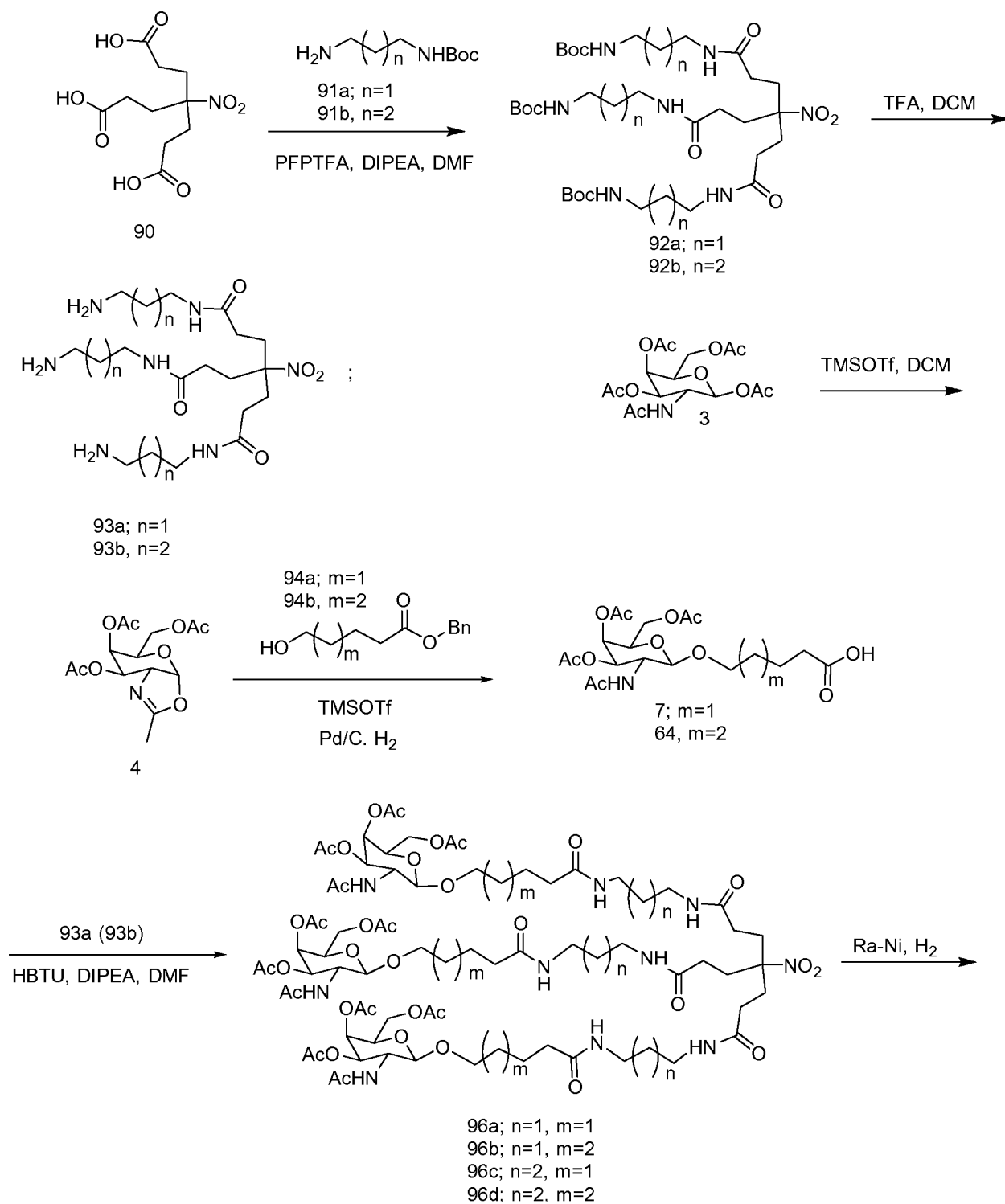
Following this general procedure ISIS 666881 was prepared. 5'-hexylamino modified oligonucleotide, ISIS 660254, was synthesized and purified using standard solid-phase oligonucleotide procedures. ISIS 660254 (40 mg, 5.2 μmol) was dissolved in 0.1 M sodium tetraborate, pH 8.5 (200 μL) and 3 equivalents PFP ester (Compound 110a) dissolved in DMSO (50 μL) was added. The PFP ester precipitated upon addition to the ASO solution requiring additional DMSO (600 μL) to fully dissolve the PFP ester. The reaction was complete after 16 h of mixing at room temperature. The solution was diluted with water to 12 mL total volume and spun down at 3000 rpm in a spin filter with a mass cut off of 3000 Da. This process was repeated twice to remove small molecule impurities. The solution was lyophilized to dryness and redissolved in concentrated aqueous ammonia with mixing at room temperature for 2.5 h followed by

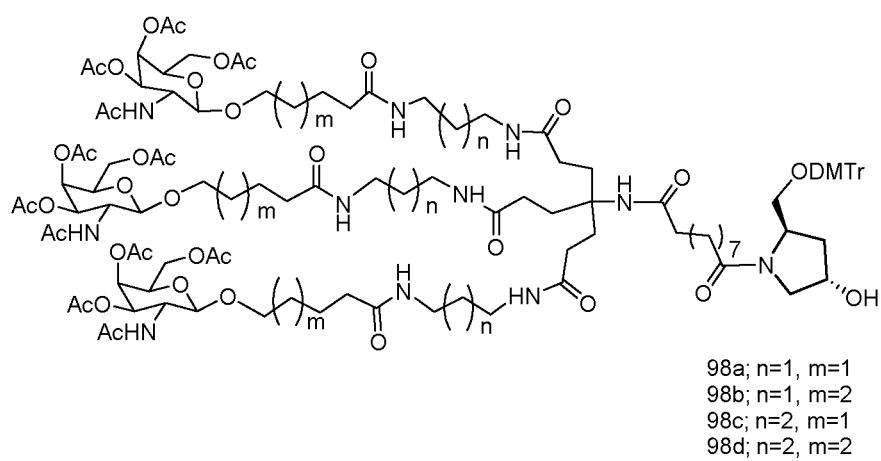
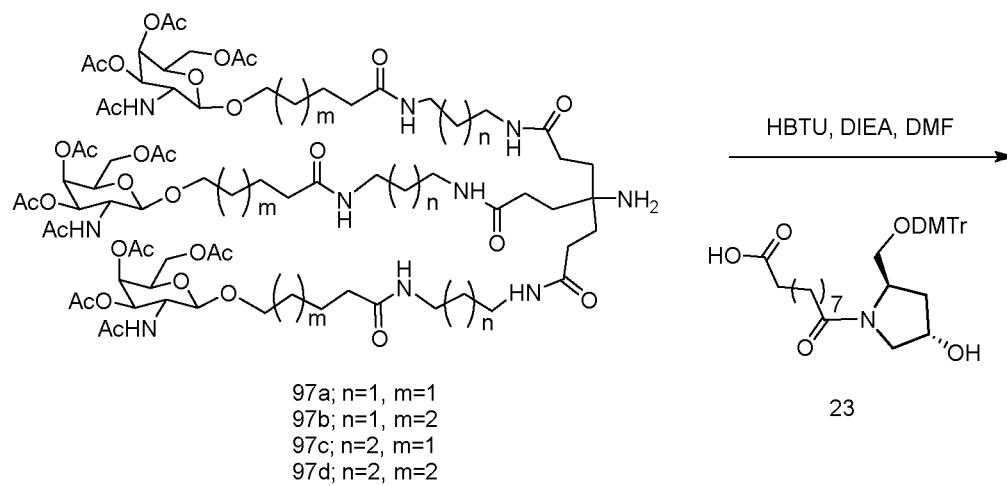
concentration *in vacuo* to remove most of the ammonia. The conjugated oligonucleotide was purified and desalted by RP-HPLC and lyophilized to give ISIS 666881 in 90% yield by weight (42 mg, 4.7 μ mol).

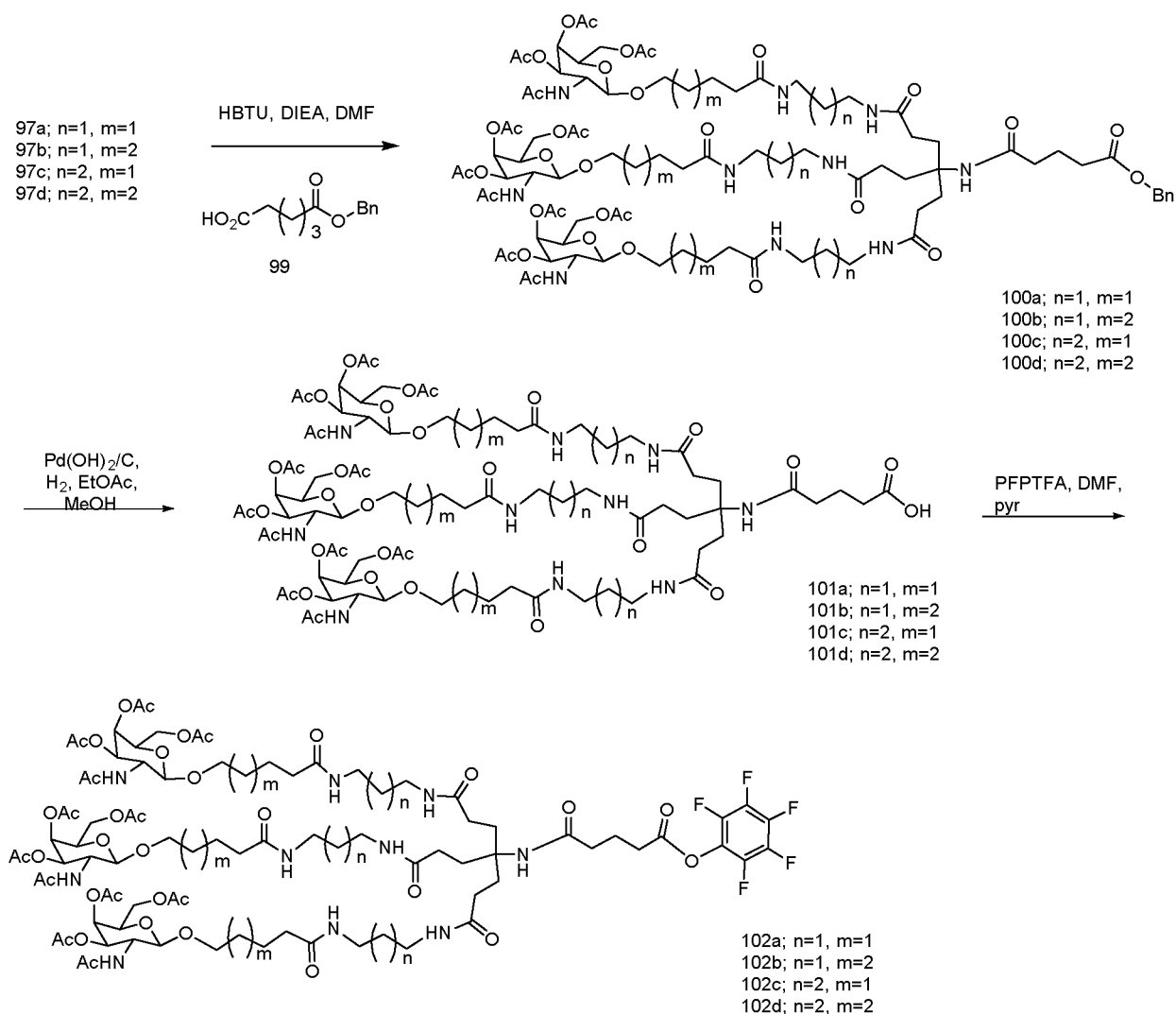
GalNAc₃-10 conjugated oligonucleotide

ASO	Sequence (5' to 3')	5' group	SEQ ID No.
ISIS 660254	NH ₂ (CH ₂) ₆ -oA _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	Hexylamine	145
ISIS 666881	GalNAc₃-10 _a -oA _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	GalNAc₃-10	145

- 5 Capital letters indicate the nucleobase for each nucleoside and ^mC indicates a 5-methyl cytosine. Subscripts: “e” indicates a 2'-MOE modified nucleoside; “d” indicates a β -D-2'-deoxyribonucleoside; “s” indicates a phosphorothioate internucleoside linkage (PS); “o” indicates a phosphodiester internucleoside linkage (PO); and “o” indicates -O-P(=O)(OH)-. Conjugate groups are in bold.

Example 47: Preparation of Oligonucleotide 102 Comprising GalNAc₃-8





The triacid 90 (4 g, 14.43 mmol) was dissolved in DMF (120 mL) and *N,N*-Diisopropylethylamine (12.35 mL, 72 mmoles). Pentafluorophenyl trifluoroacetate (8.9 mL, 52 mmoles) was added dropwise, under argon, and the reaction was allowed to stir at room temperature for 30 minutes. Boc-diamine 91a or 91b (68.87 mmol) was added, along with *N,N*-Diisopropylethylamine (12.35 mL, 72 mmoles), and the reaction was allowed to stir at room temperature for 16 hours. At that time, the DMF was reduced by >75% under reduced pressure, and then the mixture was dissolved in dichloromethane. The organic layer was washed with sodium bicarbonate, water and brine. The organic layer was then separated and dried over sodium sulfate, filtered and reduced to an oil under reduced pressure. The resultant oil was purified by silica gel chromatography (2%-->10% methanol/dichloromethane) to give compounds 92a and 92b in an approximate 80% yield. LCMS and proton NMR were consistent with the structure.

Compound 92a or 92b (6.7 mmoles) was treated with 20 mL of dichloromethane and 20 mL of trifluoroacetic acid at room temperature for 16 hours. The resultant solution was evaporated and then

dissolved in methanol and treated with DOWEX-OH resin for 30 minutes. The resultant solution was filtered and reduced to an oil under reduced pressure to give 85-90% yield of compounds 93a and 93b.

Compounds 7 or 64 (9.6 mmoles) were treated with HBTU (3.7g, 9.6 mmoles) and *N,N*-Diisopropylethylamine (5 mL) in DMF (20 mL) for 15 minutes. To this was added either compounds 93a or 93b (3 mmoles), and allowed to stir at room temperature for 16 hours. At that time, the DMF was reduced by >75% under reduced pressure, and then the mixture was dissolved in dichloromethane. The organic layer was washed with sodium bicarbonate, water and brine. The organic layer was then separated and dried over sodium sulfate, filtered and reduced to an oil under reduced pressure. The resultant oil was purified by silica gel chromatography (5%-->20% methanol/dichloromethane) to give compounds 96a-d in 20-40% yield. LCMS and proton NMR was consistent with the structure.

Compounds 96a-d (0.75 mmoles), individually, were hydrogenated over Raney Nickel for 3 hours in Ethanol (75 mL). At that time, the catalyst was removed by filtration thru celite, and the ethanol removed under reduced pressure to give compounds 97a-d in 80-90% yield. LCMS and proton NMR were consistent with the structure.

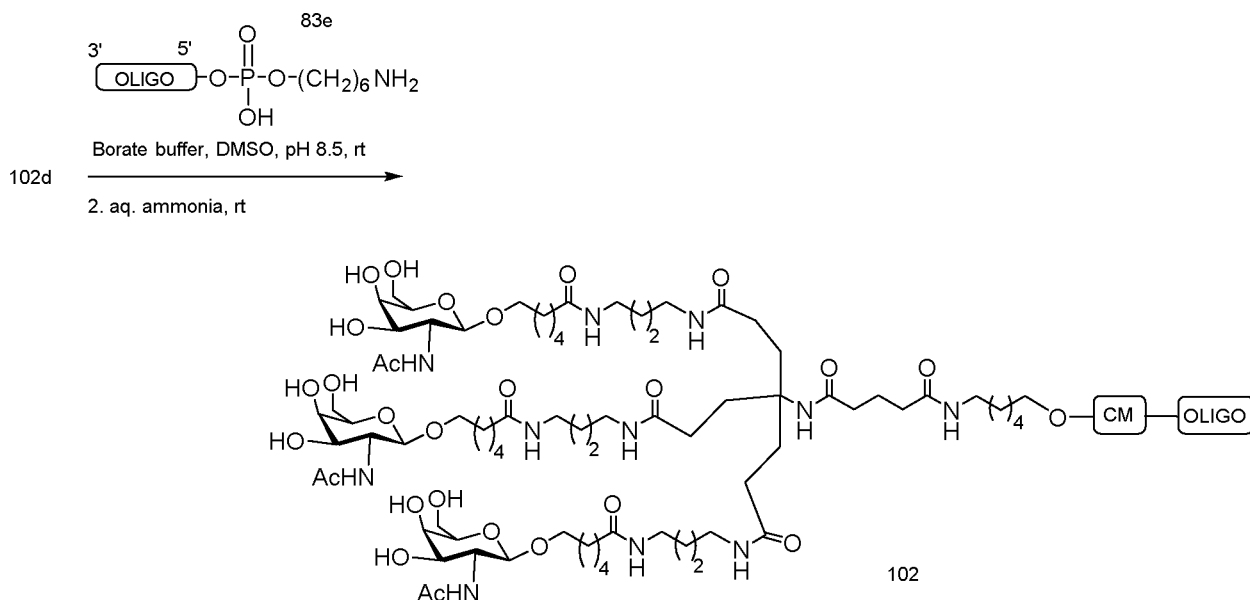
Compound 23 (0.32g, 0.53 mmoles) was treated with HBTU (0.2g, 0.53 mmoles) and *N,N*-Diisopropylethylamine (0.19 mL, 1.14 mmoles) in DMF (30mL) for 15 minutes. To this was added compounds 97a-d (0.38 mmoles), individually, and allowed to stir at room temperature for 16 hours. At that time, the DMF was reduced by >75% under reduced pressure, and then the mixture was dissolved in dichloromethane. The organic layer was washed with sodium bicarbonate, water and brine. The organic layer was then separated and dried over sodium sulfate, filtered and reduced to an oil under reduced pressure. The resultant oil was purified by silica gel chromatography (2%-->20% methanol/dichloromethane) to give compounds 98a-d in 30-40% yield. LCMS and proton NMR was consistent with the structure.

Compound 99 (0.17g, 0.76 mmoles) was treated with HBTU (0.29 g, 0.76 mmoles) and *N,N*-Diisopropylethylamine (0.35 mL, 2.0 mmoles) in DMF (50mL) for 15 minutes. To this was added compounds 97a-d (0.51 mmoles), individually, and allowed to stir at room temperature for 16 hours. At that time, the DMF was reduced by >75% under reduced pressure, and then the mixture was dissolved in dichloromethane. The organic layer was washed with sodium bicarbonate, water and brine. The organic layer was then separated and dried over sodium sulfate, filtered and reduced to an oil under reduced pressure. The resultant oil was purified by silica gel chromatography (5%-->20% methanol/ dichloromethane) to give compounds 100a-d in 40-60% yield. LCMS and proton NMR was consistent with the structure.

Compounds 100a-d (0.16 mmoles), individually, were hydrogenated over 10% Pd(OH)₂/C for 3 hours in methanol/ethyl acetate (1:1, 50 mL). At that time, the catalyst was removed by filtration thru celite, and the organics removed under reduced pressure to give compounds 101a-d in 80-90% yield. LCMS and proton NMR was consistent with the structure.

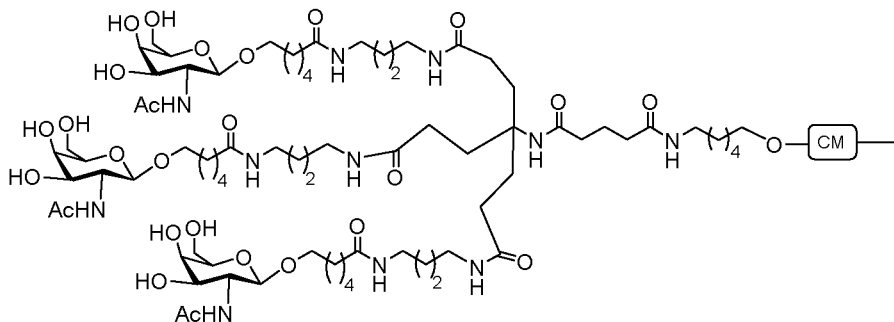
Compounds 101a-d (0.15 mmoles), individually, were dissolved in DMF (15 mL) and pyridine (0.016 mL, 0.2 mmoles). Pentafluorophenyl trifluoroacetate (0.034 mL, 0.2 mmoles) was added dropwise,

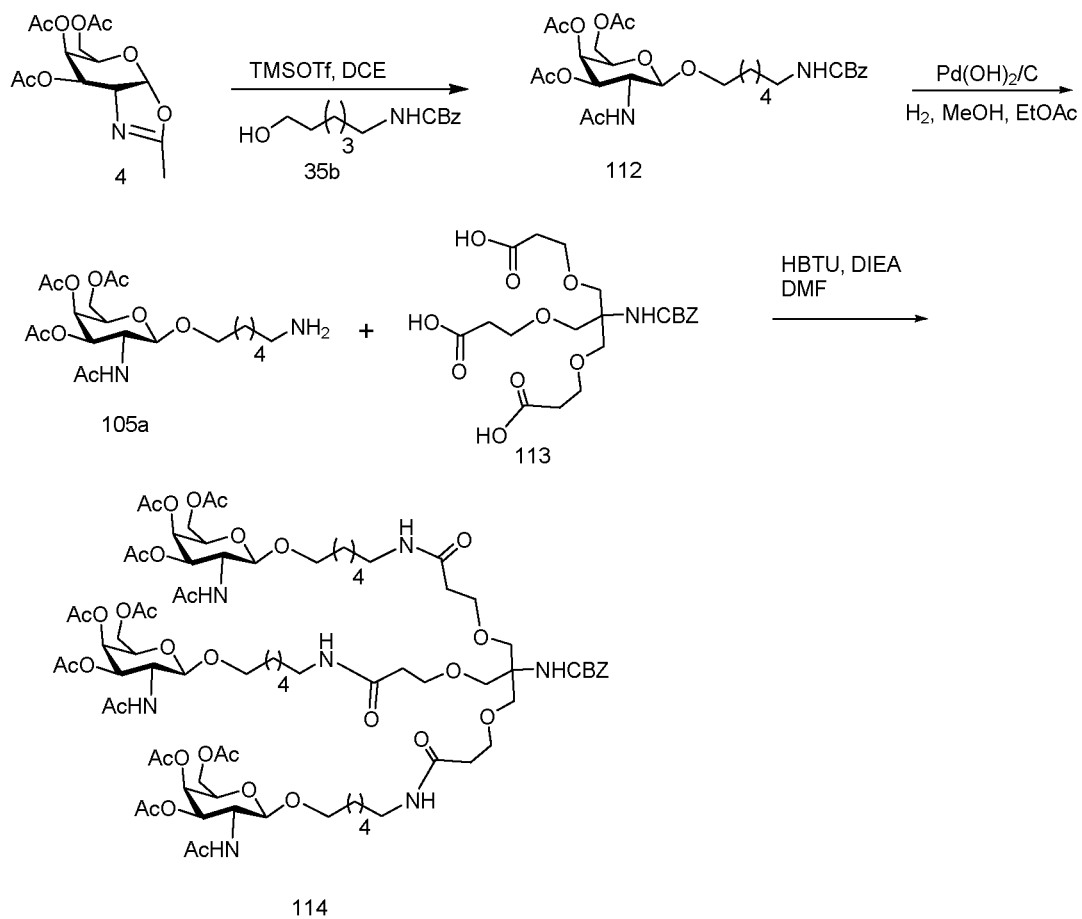
under argon, and the reaction was allowed to stir at room temperature for 30 minutes. At that time, the DMF was reduced by >75% under reduced pressure, and then the mixture was dissolved in dichloromethane. The organic layer was washed with sodium bicarbonate, water and brine. The organic layer was then separated and dried over sodium sulfate, filtered and reduced to an oil under reduced pressure. The resultant oil was purified by silica gel chromatography (2%-->5% methanol/dichloromethane) to give compounds 102a-d in an approximate 80% yield. LCMS and proton NMR were consistent with the structure.

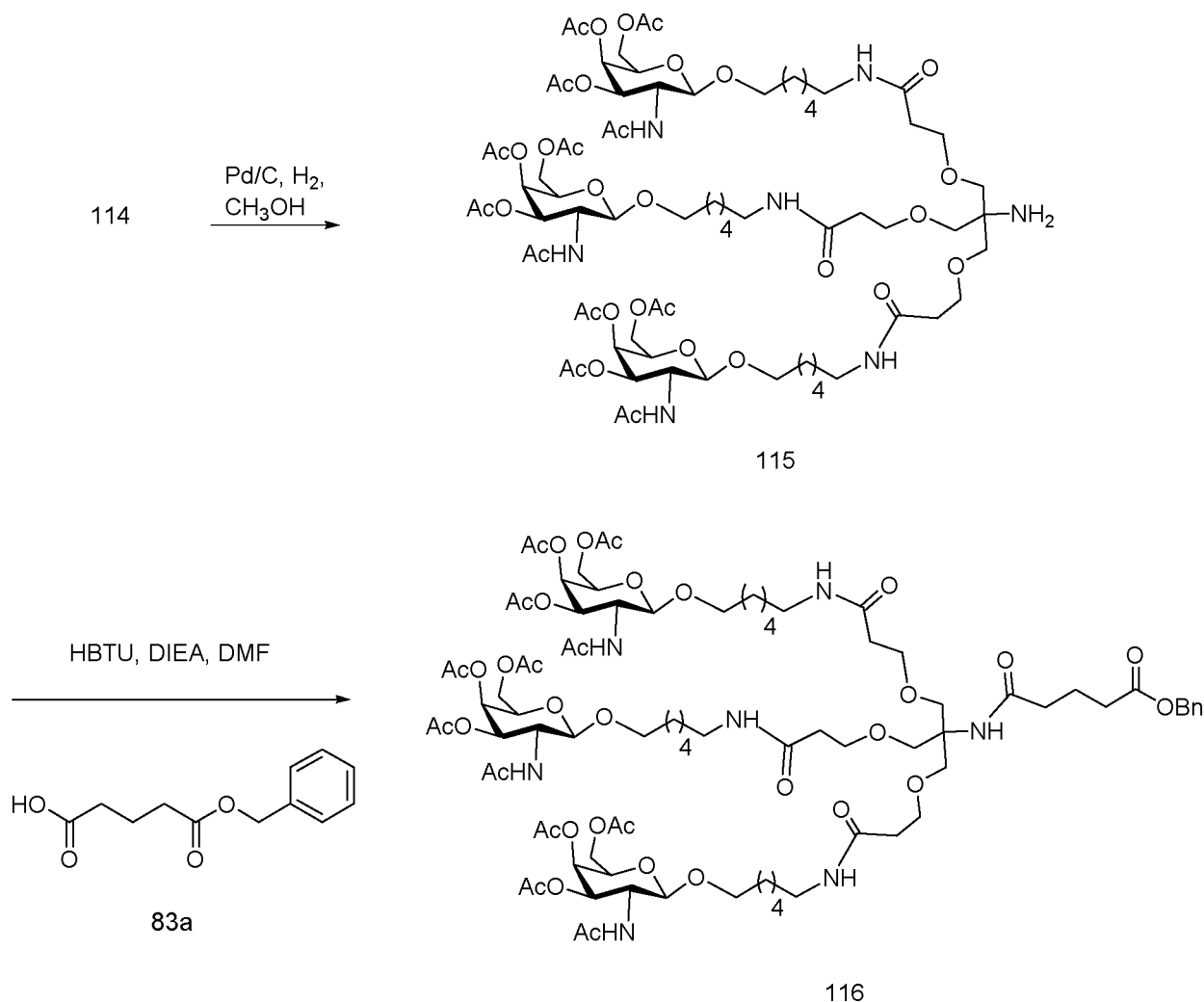


Oligomeric Compound 102, comprising a GalNAc₃-8 conjugate group, was prepared using the general procedures illustrated in Example 46. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-8 (GalNAc₃-8_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In a preferred embodiment, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-.

The structure of GalNAc₃-8 (GalNAc₃-8_a-CM-) is shown below:



Example 48: Preparation of Oligonucleotide 119 Comprising GalNAc₃-7



Compound 112 was synthesized following the procedure described in the literature (*J. Med. Chem.* 2004, 47, 5798-5808).

5 Compound 112 (5 g, 8.6 mmol) was dissolved in 1:1 methanol/ethyl acetate (22 mL/22 mL). Palladium hydroxide on carbon (0.5 g) was added. The reaction mixture was stirred at room temperature under hydrogen for 12 h. The reaction mixture was filtered through a pad of celite and washed the pad with 1:1 methanol/ethyl acetate. The filtrate and the washings were combined and concentrated to dryness to yield Compound 105a (quantitative). The structure was confirmed by LCMS.

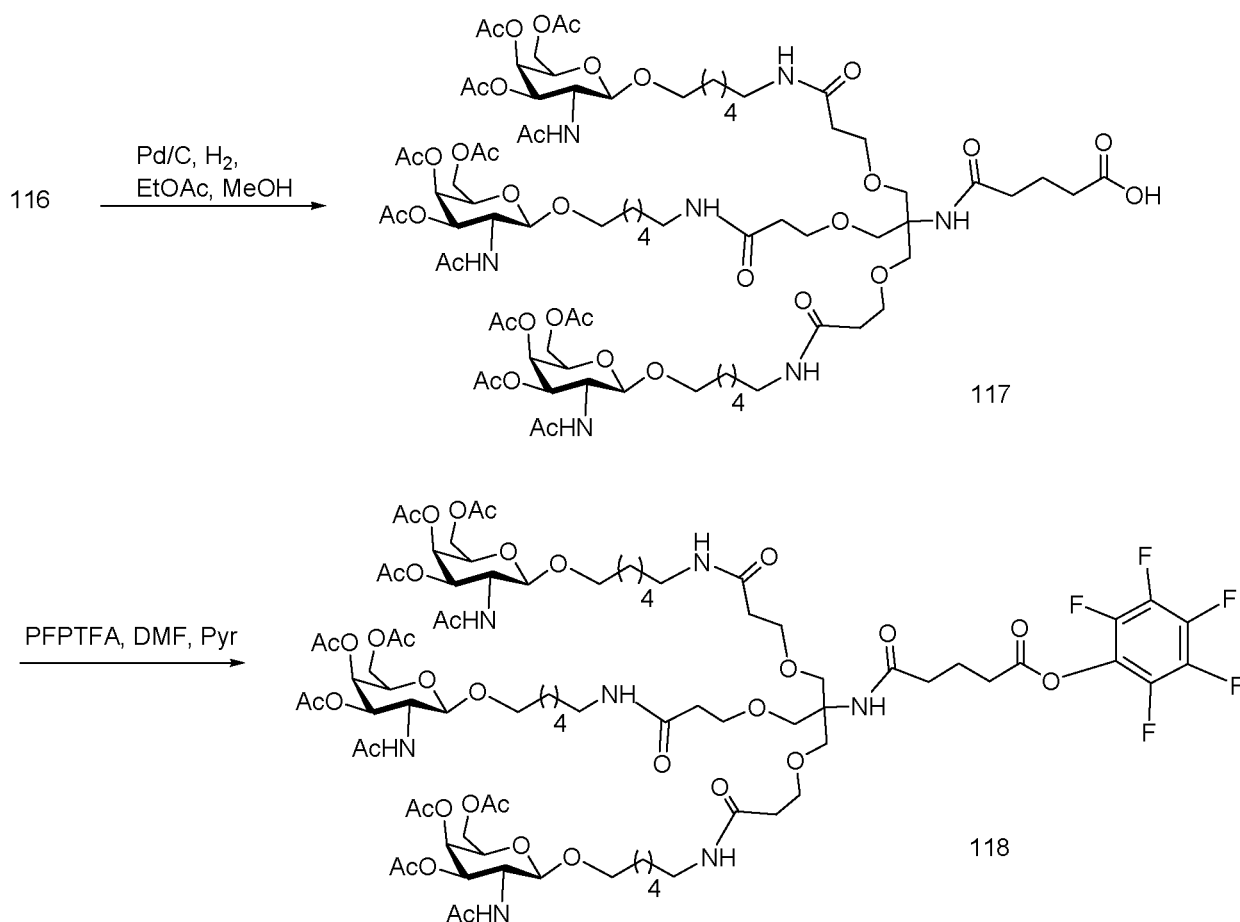
10 Compound 113 (1.25 g, 2.7 mmol), HBTU (3.2 g, 8.4 mmol) and DIEA (2.8 mL, 16.2 mmol) were dissolved in anhydrous DMF (17 mL) and the reaction mixture was stirred at room temperature for 5 min. To this a solution of Compound 105a (3.77 g, 8.4 mmol) in anhydrous DMF (20 mL) was added. The reaction was stirred at room temperature for 6 h. Solvent was removed under reduced pressure to get an oil. The residue was dissolved in CH_2Cl_2 (100 mL) and washed with aqueous saturated NaHCO_3 solution (100 mL) and brine (100 mL). The organic phase was separated, dried (Na_2SO_4), filtered and evaporated. The residue

15

was purified by silica gel column chromatography and eluted with 10 to 20 % MeOH in dichloromethane to yield Compound 114 (1.45 g, 30%). The structure was confirmed by LCMS and ^1H NMR analysis.

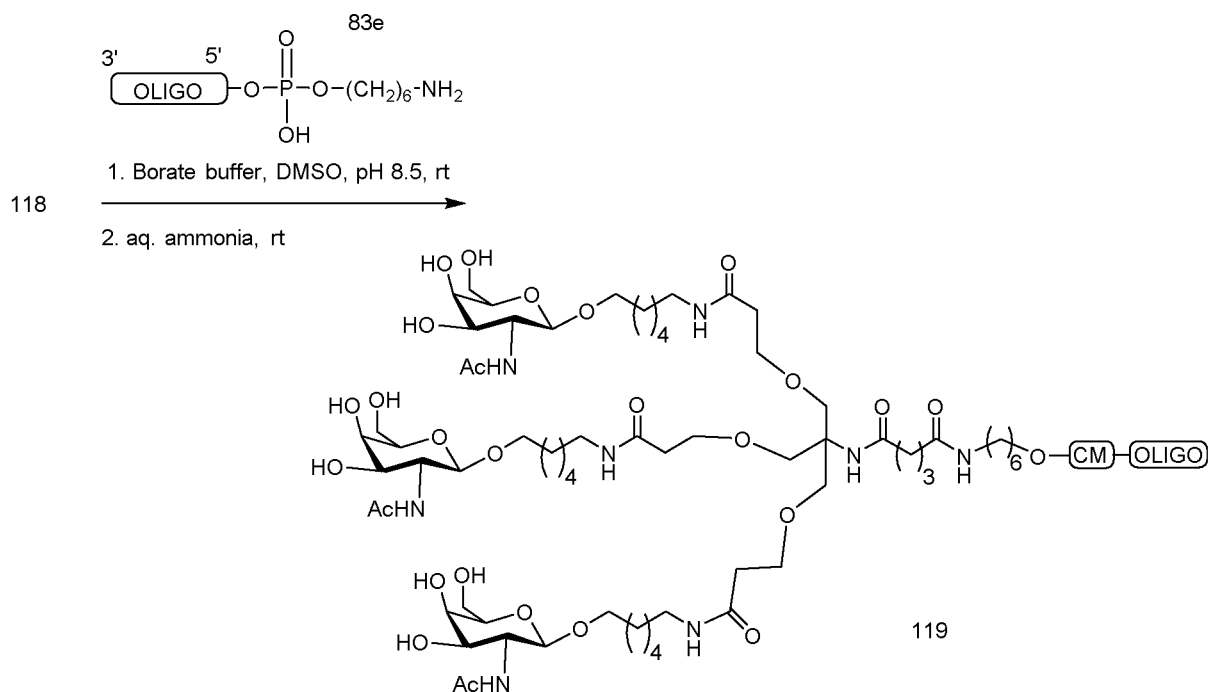
Compound 114 (1.43 g, 0.8 mmol) was dissolved in 1:1 methanol/ethyl acetate (4 mL/4 mL). Palladium on carbon (wet, 0.14 g) was added. The reaction mixture was flushed with hydrogen and stirred at room temperature under hydrogen for 12 h. The reaction mixture was filtered through a pad of celite. The celite pad was washed with methanol/ethyl acetate (1:1). The filtrate and the washings were combined together and evaporated under reduced pressure to yield Compound 115 (quantitative). The structure was confirmed by LCMS and ^1H NMR analysis.

Compound 83a (0.17 g, 0.75 mmol), HBTU (0.31 g, 0.83 mmol) and DIEA (0.26 mL, 1.5 mmol) were dissolved in anhydrous DMF (5 mL) and the reaction mixture was stirred at room temperature for 5 min. To this a solution of Compound 115 (1.22 g, 0.75 mmol) in anhydrous DMF was added and the reaction was stirred at room temperature for 6 h. The solvent was removed under reduced pressure and the residue was dissolved in CH_2Cl_2 . The organic layer was washed aqueous saturated NaHCO_3 solution and brine and dried over anhydrous Na_2SO_4 and filtered. The organic layer was concentrated to dryness and the residue obtained was purified by silica gel column chromatography and eluted with 3 to 15 % MeOH in dichloromethane to yield Compound 116 (0.84 g, 61%). The structure was confirmed by LC MS and ^1H NMR analysis.



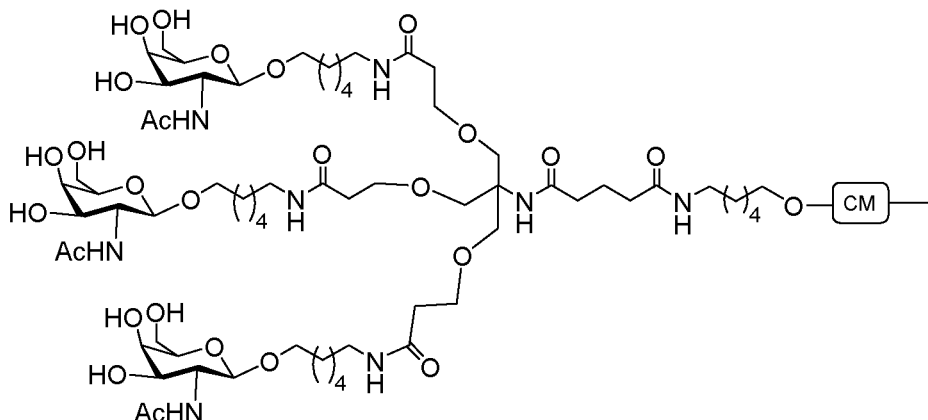
Compound 116 (0.74 g, 0.4 mmol) was dissolved in 1:1 methanol/ethyl acetate (5 mL/5 mL). Palladium on carbon (wet, 0.074 g) was added. The reaction mixture was flushed with hydrogen and stirred at room temperature under hydrogen for 12 h. The reaction mixture was filtered through a pad of celite. The celite pad was washed with methanol/ethyl acetate (1:1). The filtrate and the washings were combined together and evaporated under reduced pressure to yield compound 117 (0.73 g, 98%). The structure was confirmed by LCMS and ^1H NMR analysis.

Compound 117 (0.63 g, 0.36 mmol) was dissolved in anhydrous DMF (3 mL). To this solution *N,N*-Diisopropylethylamine (70 μL , 0.4 mmol) and pentafluorophenyl trifluoroacetate (72 μL , 0.42 mmol) were added. The reaction mixture was stirred at room temperature for 12 h and poured into a aqueous saturated NaHCO_3 solution. The mixture was extracted with dichloromethane, washed with brine and dried over anhydrous Na_2SO_4 . The dichloromethane solution was concentrated to dryness and purified with silica gel column chromatography and eluted with 5 to 10 % MeOH in dichloromethane to yield compound 118 (0.51 g, 79%). The structure was confirmed by LCMS and ^1H and ^{19}F NMR.

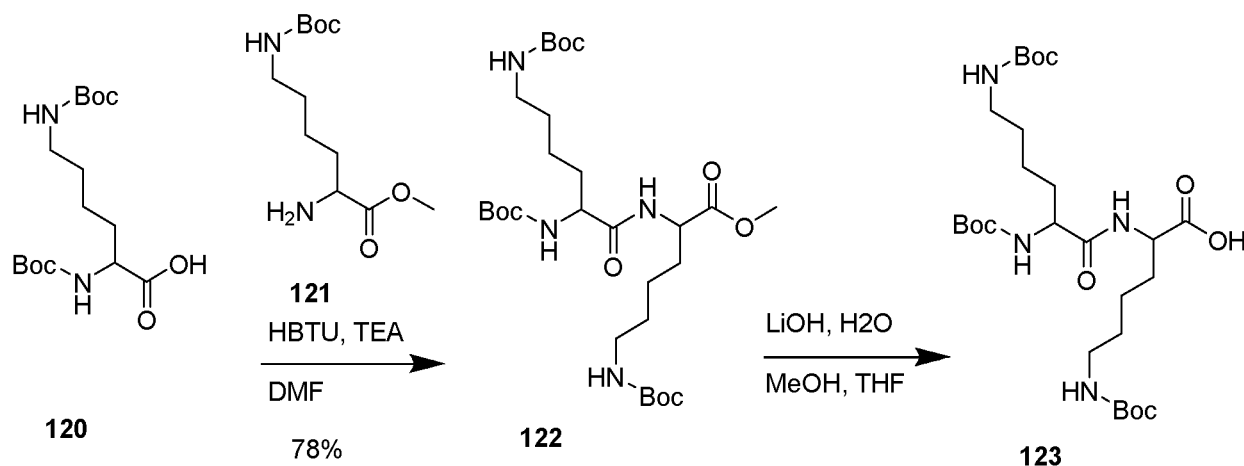


Oligomeric Compound 119, comprising a $\text{GalNAc}_3\text{-7}$ conjugate group, was prepared using the general procedures illustrated in Example 46. The GalNAc_3 cluster portion of the conjugate group $\text{GalNAc}_3\text{-7}$ ($\text{GalNAc}_3\text{-7}_a$) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is $-\text{P}(=\text{O})(\text{OH})-\text{A}_d-\text{P}(=\text{O})(\text{OH})-$.

The structure of $\text{GalNAc}_3\text{-7}$ ($\text{GalNAc}_3\text{-7}_a\text{-CM-}$) is shown below:



Example 49: Preparation of Oligonucleotide 132 Comprising GalNAc₃-5



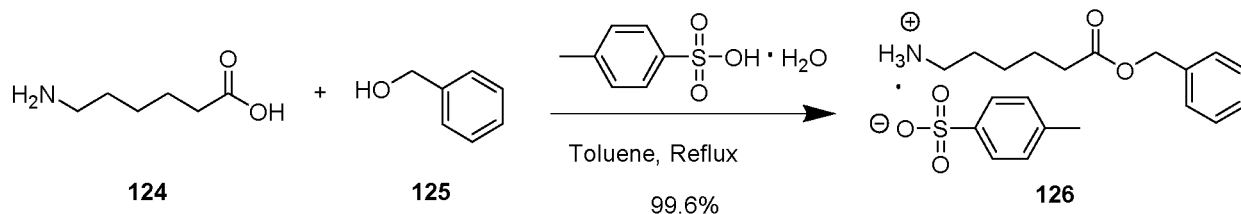
5

Compound 120 (14.01 g, 40 mmol) and HBTU (14.06 g, 37 mmol) were dissolved in anhydrous DMF (80 mL). Triethylamine (11.2 mL, 80.35 mmol) was added and stirred for 5 min. The reaction mixture was cooled in an ice bath and a solution of compound 121 (10 g, mmol) in anhydrous DMF (20 mL) was added. Additional triethylamine (4.5 mL, 32.28 mmol) was added and the reaction mixture was stirred for 18 h under an argon atmosphere. The reaction was monitored by TLC (ethyl acetate:hexane; 1:1; $R_f = 0.47$). The solvent was removed under reduced pressure. The residue was taken up in EtOAc (300 mL) and washed with 1M NaHSO₄ (3 x 150 mL), aqueous saturated NaHCO₃ solution (3 x 150 mL) and brine (2 x 100 mL). Organic layer was dried with Na₂SO₄. Drying agent was removed by filtration and organic layer was concentrated by rotary evaporation. Crude mixture was purified by silica gel column chromatography and eluted by using 35 – 50% EtOAc in hexane to yield a compound 122 (15.50 g, 78.13%). The structure was confirmed by LCMS and ¹H NMR analysis. Mass m/z 589.3 [M + H]⁺.

15

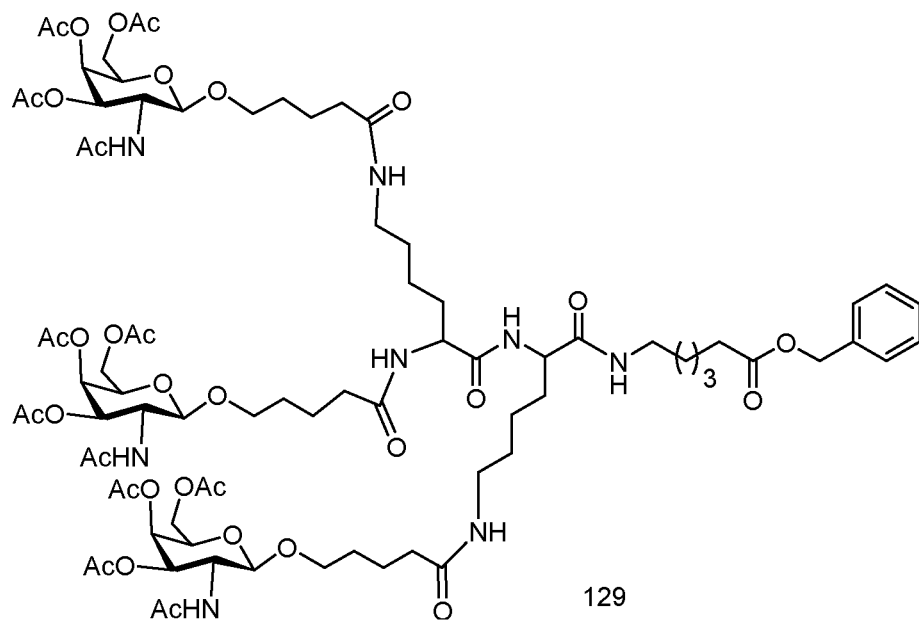
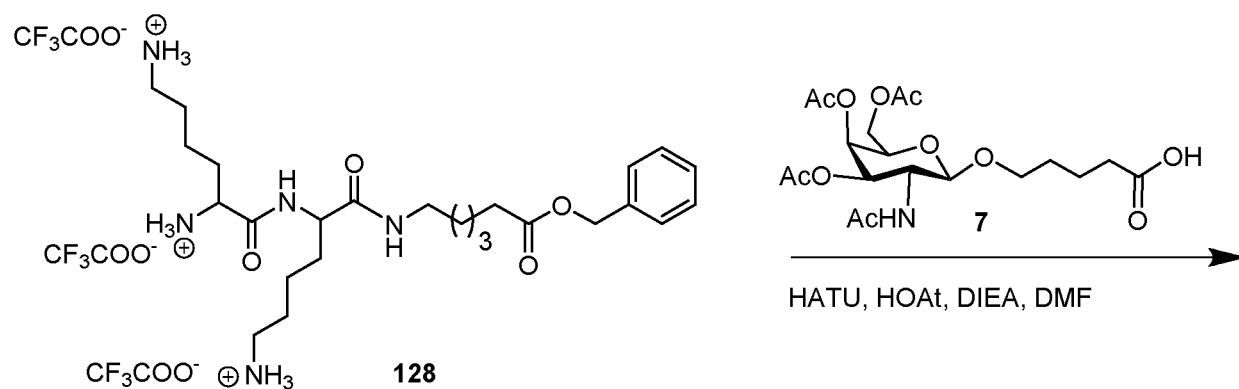
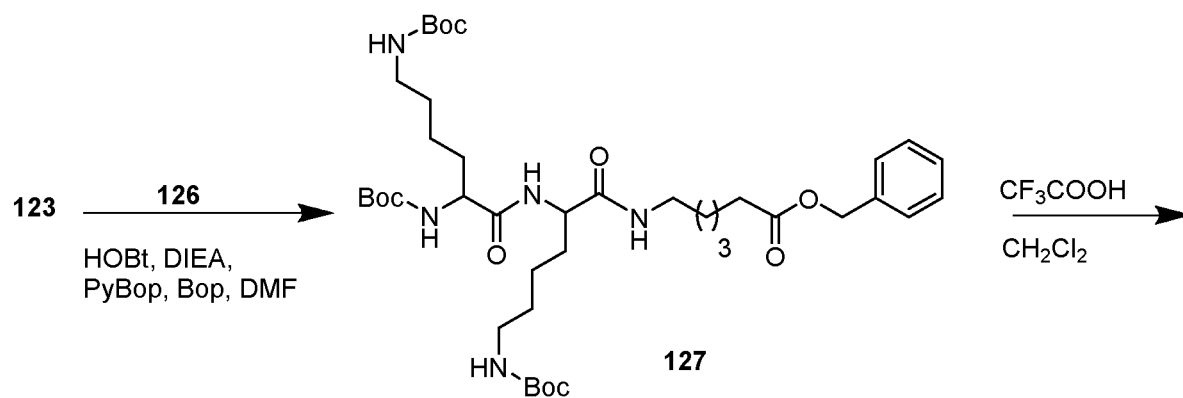
A solution of LiOH (92.15 mmol) in water (20 mL) and THF (10 mL) was added to a cooled solution of Compound 122 (7.75 g, 13.16 mmol) dissolved in methanol (15 mL). The reaction mixture was stirred at

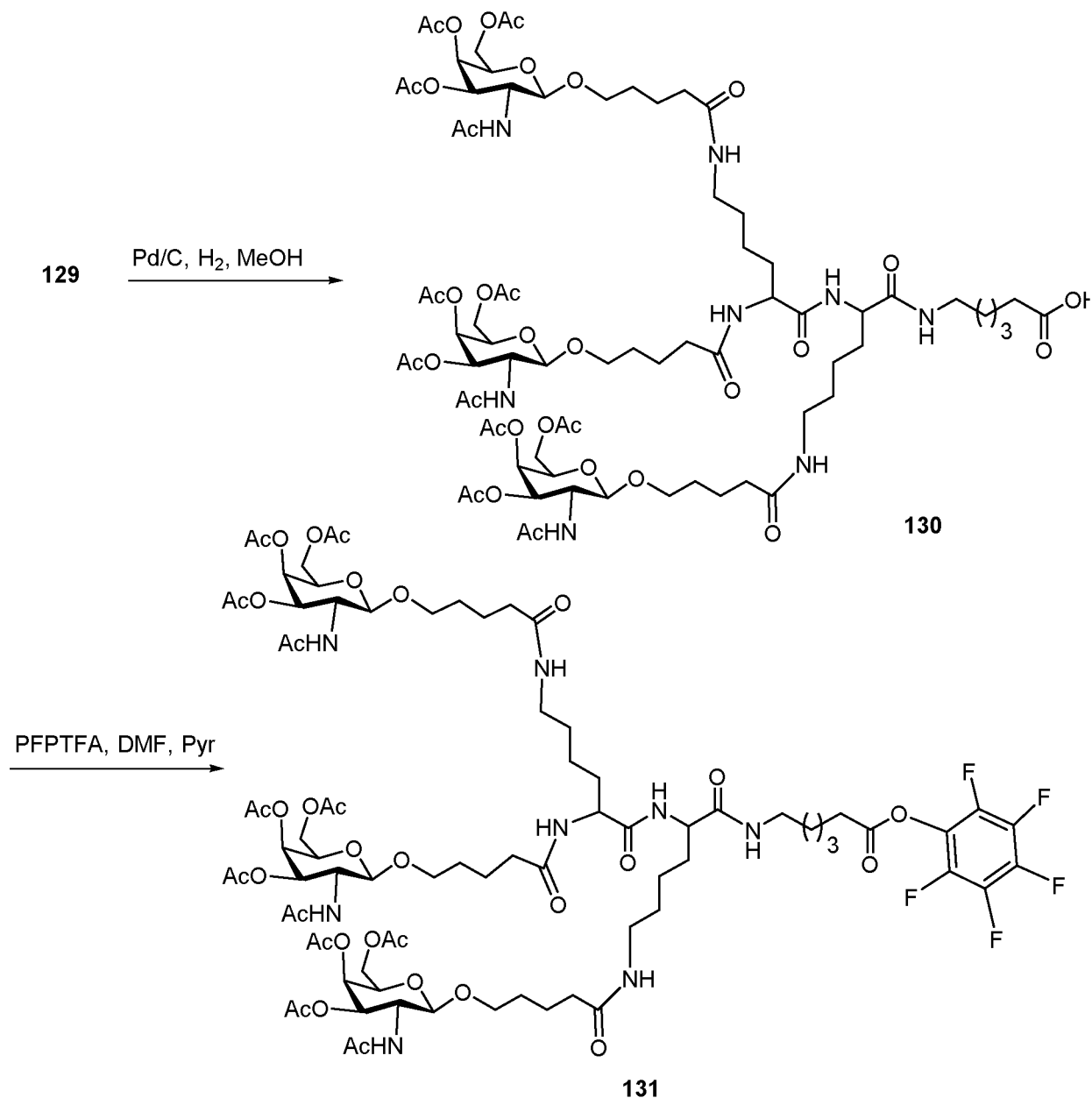
room temperature for 45 min. and monitored by TLC (EtOAc:hexane; 1:1). The reaction mixture was concentrated to half the volume under reduced pressure. The remaining solution was cooled in an ice bath and neutralized by adding concentrated HCl. The reaction mixture was diluted, extracted with EtOAc (120 mL) and washed with brine (100 mL). An emulsion formed and cleared upon standing overnight. The organic layer was separated, dried (Na_2SO_4), filtered and evaporated to yield Compound 123 (8.42 g). Residual salt is the likely cause of excess mass. LCMS is consistent with structure. Product was used without any further purification. M.W.cal:574.36; M.W.f.d:575.3 $[\text{M} + \text{H}]^+$.



Compound 126 was synthesized following the procedure described in the literature (*J. Am. Chem.*

Soc. 2011, 133, 958-963).





Compound 123 (7.419 g, 12.91 mmol), HOBt (3.49 g, 25.82 mmol) and compound 126 (6.33 g, 16.14 mmol) were dissolved in and DMF (40 mL) and the resulting reaction mixture was cooled in an ice bath. To this *N,N*-Diisopropylethylamine (4.42 mL, 25.82 mmol), PyBop (8.7 g, 16.7 mmol) followed by Bop coupling reagent (1.17 g, 2.66 mmol) were added under an argon atmosphere. The ice bath was removed and the solution was allowed to warm to room temperature. The reaction was completed after 1 h as determined by TLC (DCM:MeOH:AA; 89:10:1). The reaction mixture was concentrated under reduced pressure. The residue was dissolved in EtOAc (200 mL) and washed with 1 M NaHSO₄ (3x100 mL), aqueous saturated NaHCO₃ (3x100 mL) and brine (2x100 mL). The organic phase separated dried (Na₂SO₄), filtered and concentrated. The residue was purified by silica gel column chromatography with a gradient of 50% hexanes/EtOAc to 100% EtOAc to yield Compound 127 (9.4 g) as a white foam. LCMS and ¹H NMR

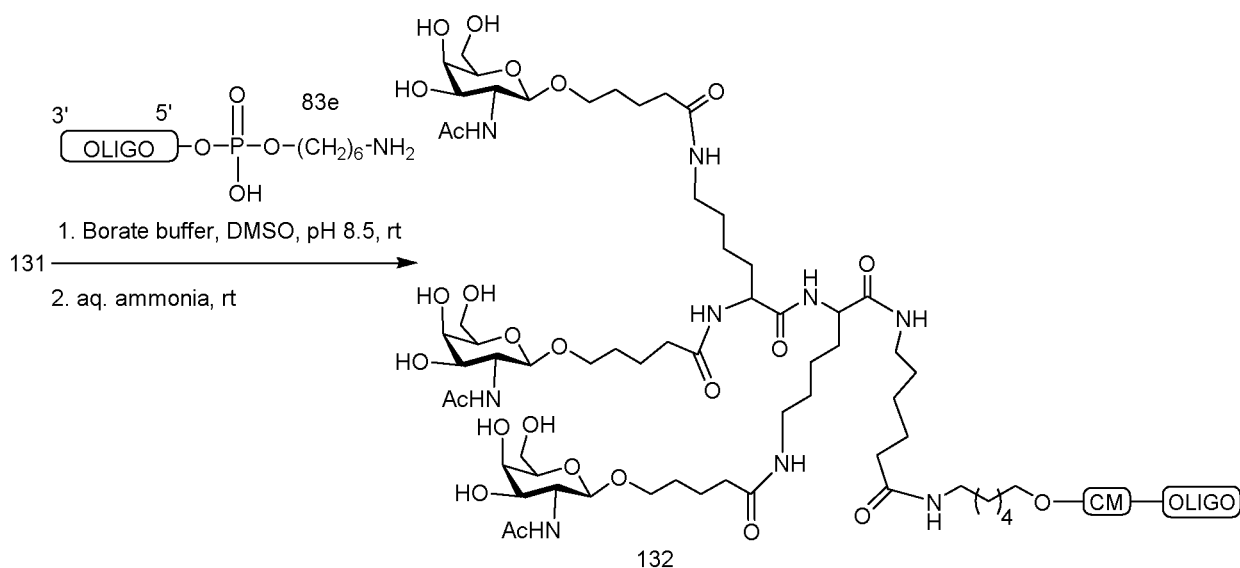
were consistent with structure. Mass m/z 778.4 $[M + H]^+$.

Trifluoroacetic acid (12 mL) was added to a solution of compound 127 (1.57 g, 2.02 mmol) in dichloromethane (12 mL) and stirred at room temperature for 1 h. The reaction mixture was co-evaporated with toluene (30 mL) under reduced pressure to dryness. The residue obtained was co-evaporated twice with acetonitrile (30 mL) and toluene (40 mL) to yield Compound 128 (1.67 g) as trifluoro acetate salt and used for next step without further purification. LCMS and 1H NMR were consistent with structure. Mass m/z 478.2 $[M + H]^+$.

Compound 7 (0.43 g, 0.963 mmol), HATU (0.35 g, 0.91 mmol), and HOAt (0.035 g, 0.26 mmol) were combined together and dried for 4 h over P_2O_5 under reduced pressure in a round bottom flask and then dissolved in anhydrous DMF (1 mL) and stirred for 5 min. To this a solution of compound 128 (0.20 g, 0.26 mmol) in anhydrous DMF (0.2 mL) and N,N -Diisopropylethylamine (0.2 mL) was added. The reaction mixture was stirred at room temperature under an argon atmosphere. The reaction was complete after 30 min as determined by LCMS and TLC (7% MeOH/DCM). The reaction mixture was concentrated under reduced pressure. The residue was dissolved in DCM (30 mL) and washed with 1 M $NaHSO_4$ (3x20 mL), aqueous saturated $NaHCO_3$ (3 x 20 mL) and brine (3x20 mL). The organic phase was separated, dried over Na_2SO_4 , filtered and concentrated. The residue was purified by silica gel column chromatography using 5-15% MeOH in dichloromethane to yield Compound 129 (96.6 mg). LC MS and 1H NMR are consistent with structure. Mass m/z 883.4 $[M + 2H]^+$.

Compound 129 (0.09 g, 0.051 mmol) was dissolved in methanol (5 mL) in 20 mL scintillation vial. To this was added a small amount of 10% Pd/C (0.015 mg) and the reaction vessel was flushed with H_2 gas. The reaction mixture was stirred at room temperature under H_2 atmosphere for 18 h. The reaction mixture was filtered through a pad of Celite and the Celite pad was washed with methanol. The filtrate washings were pooled together and concentrated under reduced pressure to yield Compound 130 (0.08 g). LCMS and 1H NMR were consistent with structure. The product was used without further purification. Mass m/z 838.3 $[M + 2H]^+$.

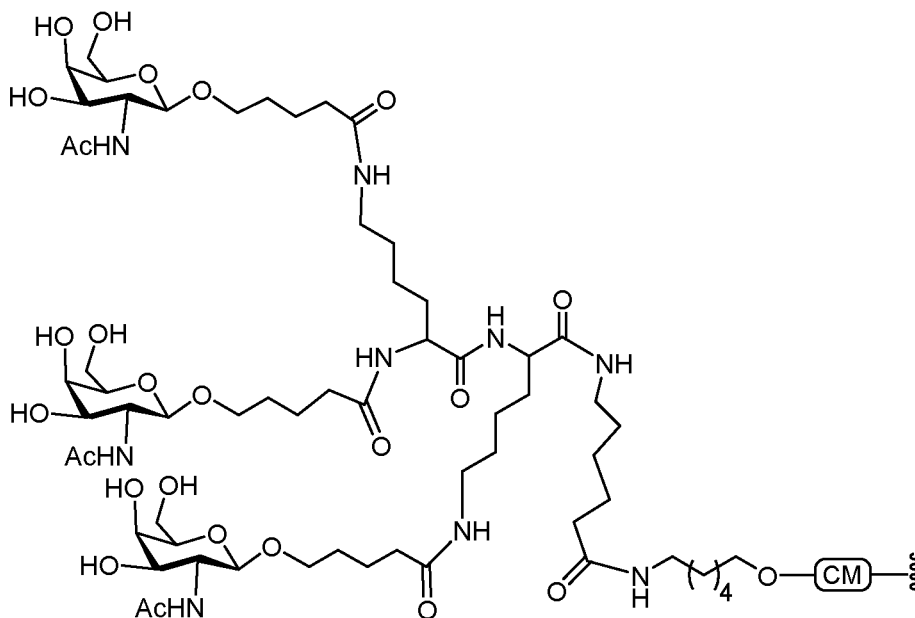
To a 10 mL pointed round bottom flask were added compound 130 (75.8 mg, 0.046 mmol), 0.37 M pyridine/DMF (200 μ L) and a stir bar. To this solution was added 0.7 M pentafluorophenyl trifluoroacetate/DMF (100 μ L) drop wise with stirring. The reaction was completed after 1 h as determined by LC MS. The solvent was removed under reduced pressure and the residue was dissolved in $CHCl_3$ (~ 10 mL). The organic layer was partitioned against $NaHSO_4$ (1 M, 10 mL), aqueous saturated $NaHCO_3$ (10 mL) and brine (10 mL) three times each. The organic phase separated and dried over Na_2SO_4 , filtered and concentrated to yield Compound 131 (77.7 mg). LCMS is consistent with structure. Used without further purification. Mass m/z 921.3 $[M + 2H]^+$.

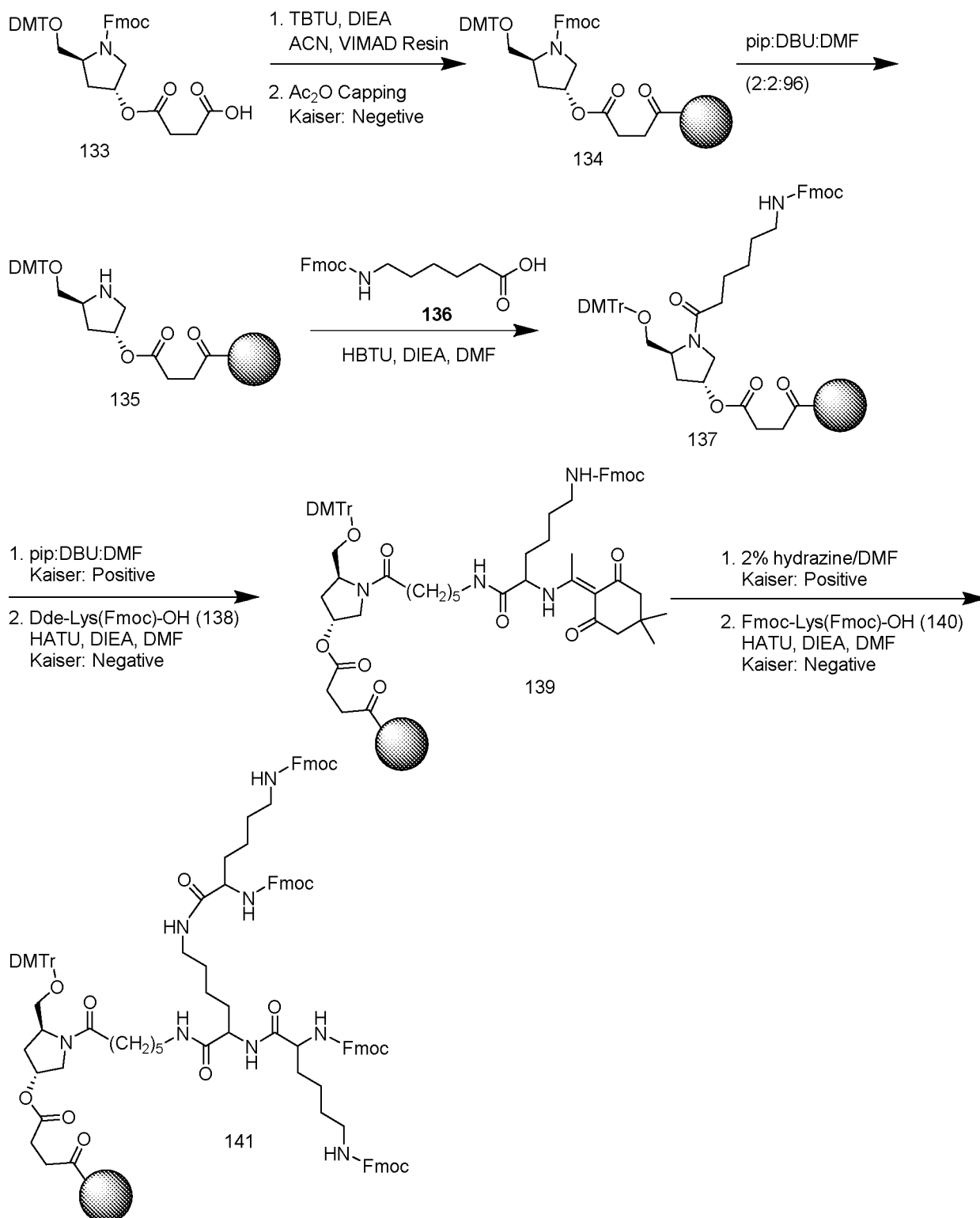


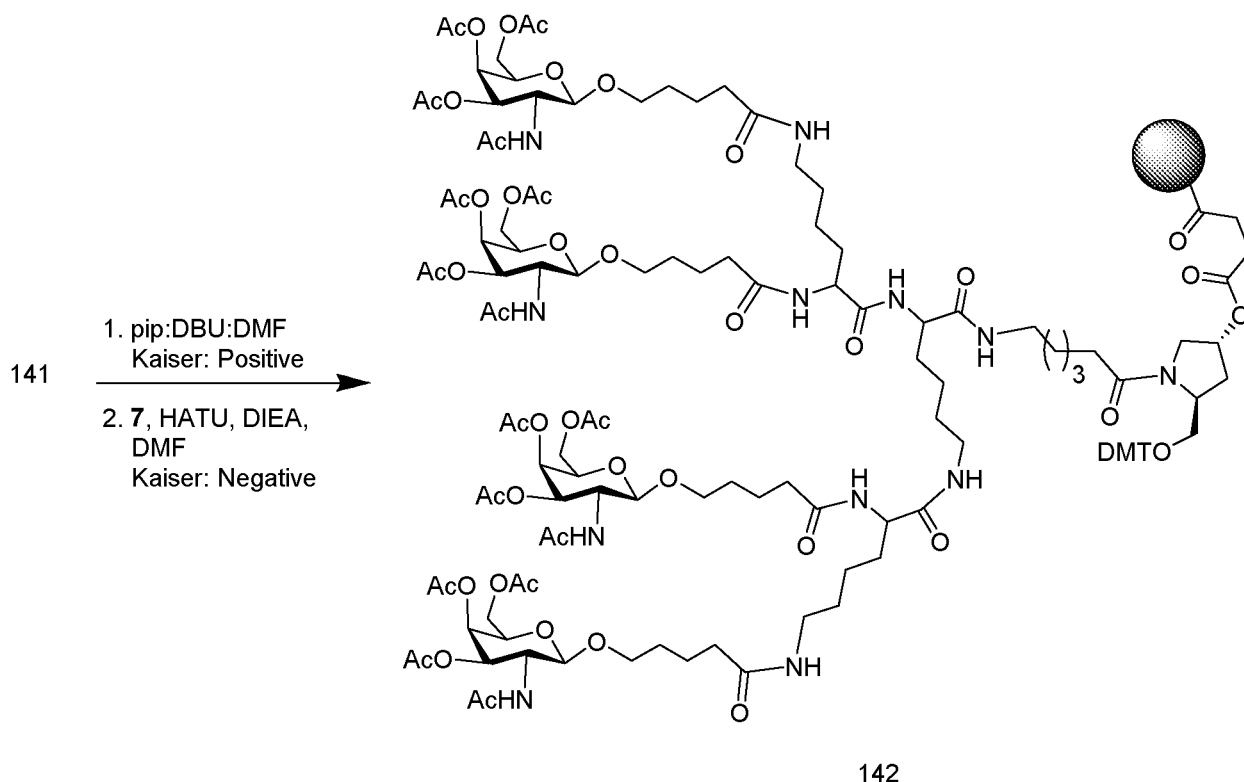
5 Oligomeric Compound 132, comprising a GalNAc₃-5 conjugate group, was prepared using the general procedures illustrated in Example 46. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-5 (GalNAc₃-5_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-.

The structure of GalNAc₃-5 (GalNAc₃-5_a-CM-) is shown below:

10



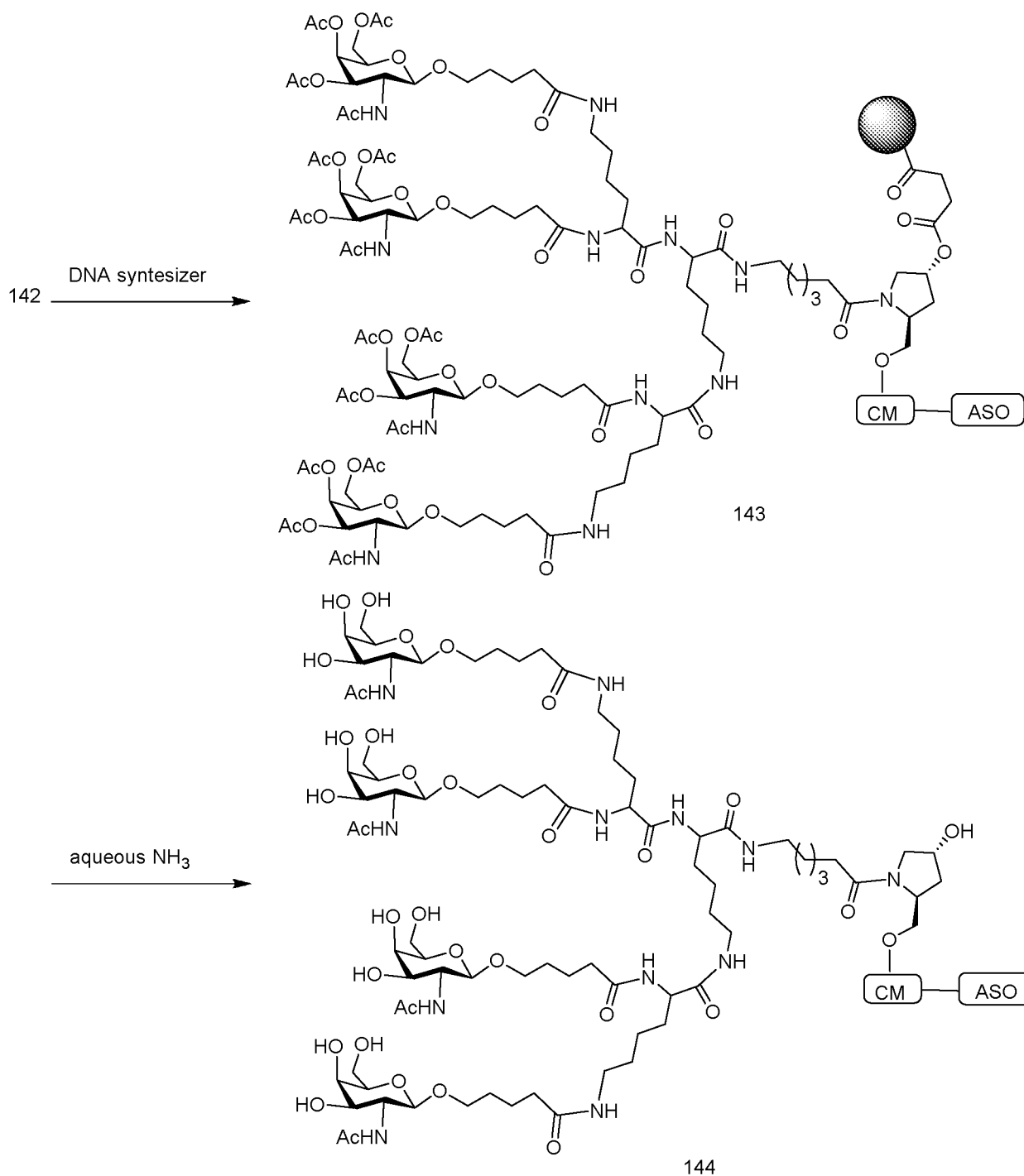
Example 50: Preparation of Oligonucleotide 144 Comprising GalNAc₄-11



Synthesis of Compound 134. To a Merrifield flask was added aminomethyl VIMAD resin (2.5 g, 450 $\mu\text{mol/g}$) that was washed with acetonitrile, dimethylformamide, dichloromethane and acetonitrile. The resin was swelled in acetonitrile (4 mL). Compound 133 was pre-activated in a 100 mL round bottom flask by adding 20 (1.0 mmol, 0.747 g), TBTU (1.0 mmol, 0.321 g), acetonitrile (5 mL) and DIEA (3.0 mmol, 0.5 mL). This solution was allowed to stir for 5 min and was then added to the Merrifield flask with shaking. The suspension was allowed to shake for 3 h. The reaction mixture was drained and the resin was washed with acetonitrile, DMF and DCM. New resin loading was quantitated by measuring the absorbance of the DMT cation at 500 nm (extinction coefficient = 76000) in DCM and determined to be 238 $\mu\text{mol/g}$. The resin was capped by suspending in an acetic anhydride solution for ten minutes three times.

The solid support bound compound 141 was synthesized using iterative Fmoc-based solid phase peptide synthesis methods. A small amount of solid support was withdrawn and suspended in aqueous ammonia (28-30 wt%) for 6 h. The cleaved compound was analyzed by LC-MS and the observed mass was consistent with structure. Mass m/z 1063.8 $[M + 2H]^+$.

The solid support bound compound 142 was synthesized using solid phase peptide synthesis methods.



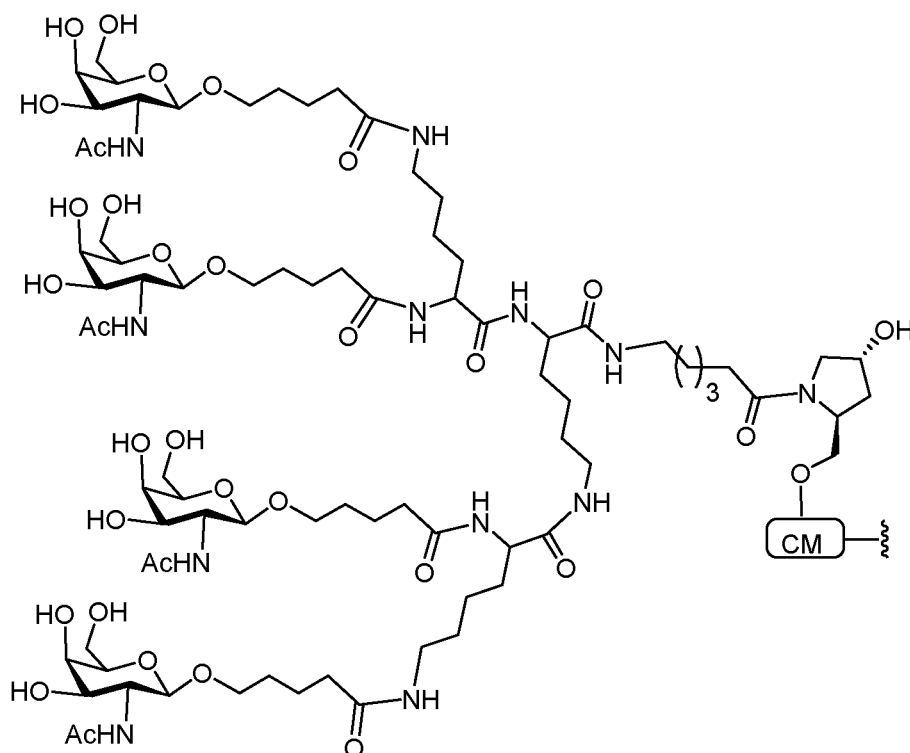
The solid support bound compound 143 was synthesized using standard solid phase synthesis on a DNA synthesizer.

5 The solid support bound compound 143 was suspended in aqueous ammonia (28-30 wt%) and heated at 55 °C for 16 h. The solution was cooled and the solid support was filtered. The filtrate was concentrated and the residue dissolved in water and purified by HPLC on a strong anion exchange column. The fractions containing full length compound 144 were pooled together and desalted. The resulting GalNAc₄-11

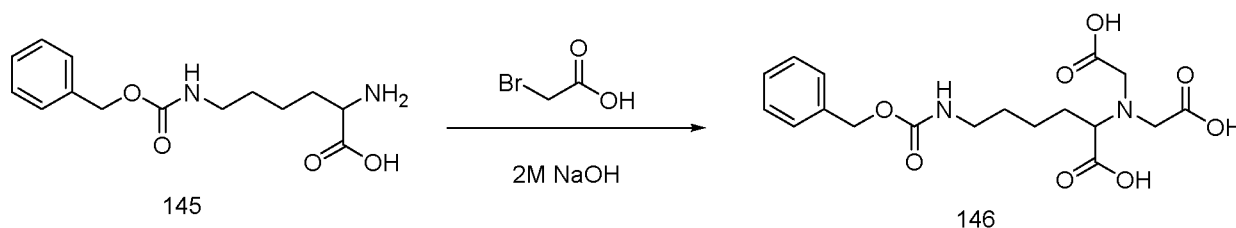
conjugated oligomeric compound was analyzed by LC-MS and the observed mass was consistent with structure.

The GalNAc₄ cluster portion of the conjugate group GalNAc₄-11 (GalNAc₄-11_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-.

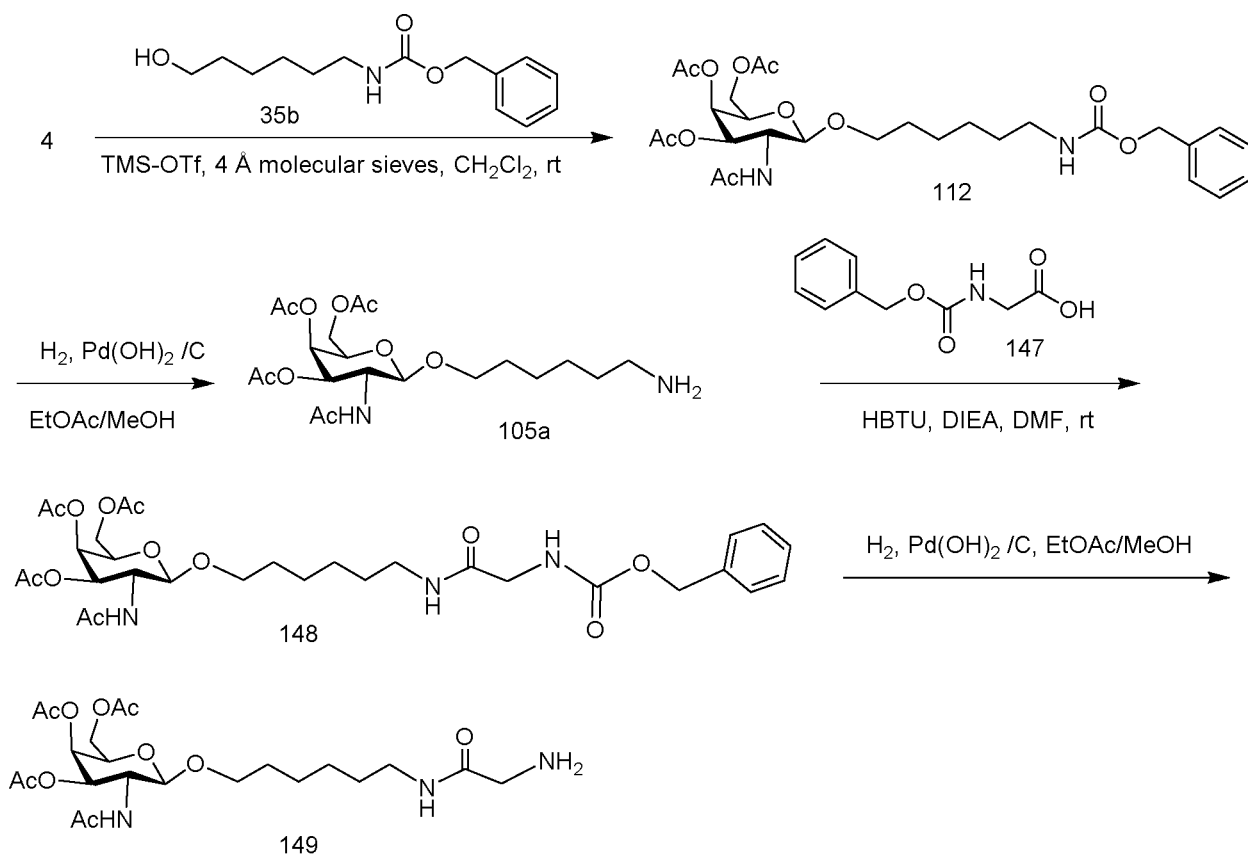
The structure of GalNAc₄-11 (GalNAc₄-11_a-CM) is shown below:



Example 51: Preparation of Oligonucleotide 155 Comprising GalNAc₃-6



Compound 146 was synthesized as described in the literature (*Analytical Biochemistry* 1995, 229, 54-60).



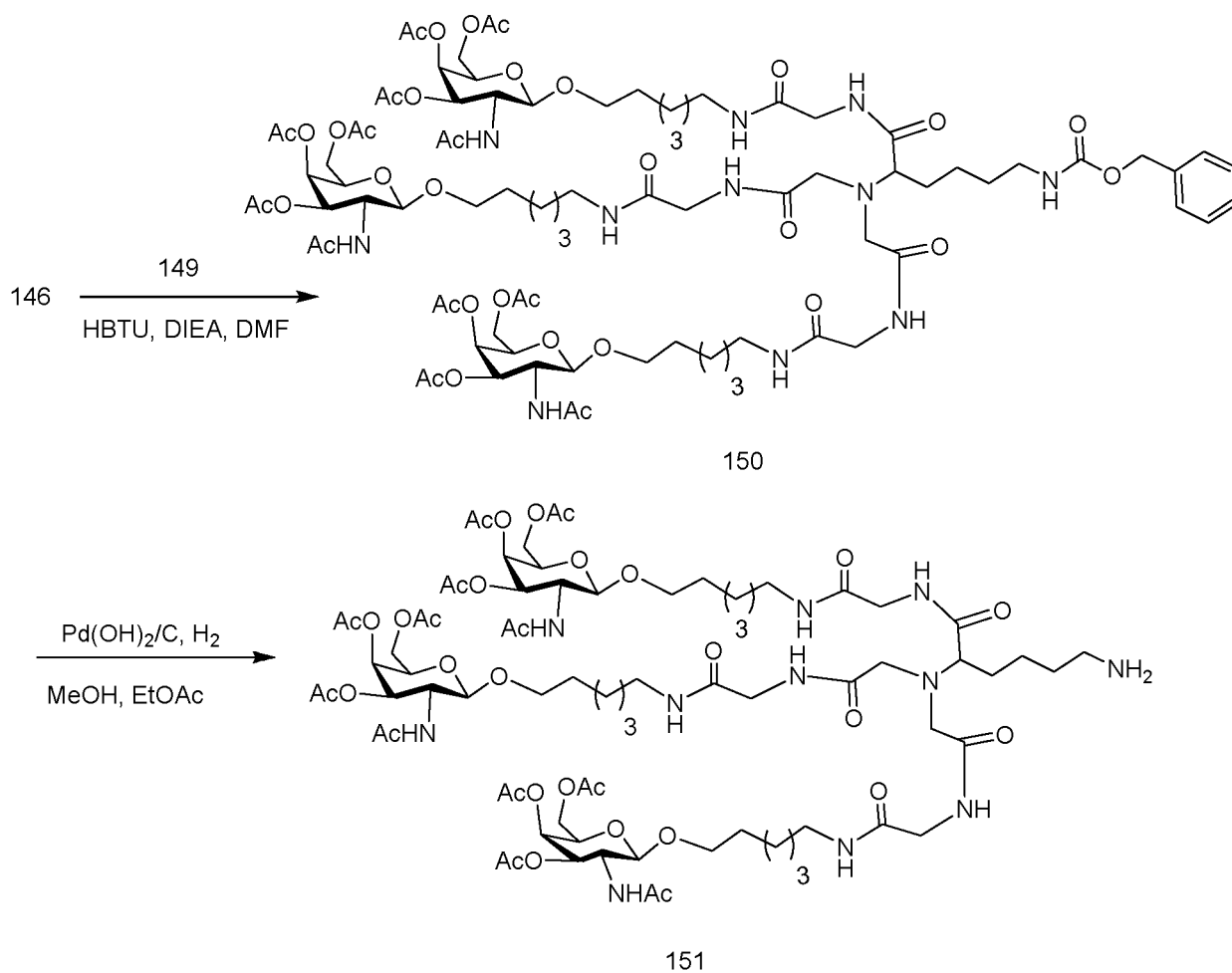
Compound 4 (15 g, 45.55 mmol) and compound 35b (14.3 grams, 57 mmol) were dissolved in CH₂Cl₂ (200 ml). Activated molecular sieves (4 Å, 2 g, powdered) were added, and the reaction was allowed to stir for 30 minutes under nitrogen atmosphere. TMS-OTf was added (4.1 ml, 22.77 mmol) and the reaction was allowed to stir at room temp overnight. Upon completion, the reaction was quenched by pouring into solution of saturated aqueous NaHCO₃ (500 ml) and crushed ice (~ 150 g). The organic layer was separated, washed with brine, dried over MgSO₄, filtered, and was concentrated to an orange oil under reduced pressure. The crude material was purified by silica gel column chromatography and eluted with 2-10 % MeOH in CH₂Cl₂ to yield Compound 112 (16.53 g, 63 %). LCMS and ¹H NMR were consistent with the expected compound.

Compound 112 (4.27 g, 7.35 mmol) was dissolved in 1:1 MeOH/EtOAc (40 ml). The reaction mixture was purged by bubbling a stream of argon through the solution for 15 minutes. Pearlman's catalyst (palladium hydroxide on carbon, 400 mg) was added, and hydrogen gas was bubbled through the solution for 30 minutes. Upon completion (TLC 10% MeOH in CH₂Cl₂, and LCMS), the catalyst was removed by filtration through a pad of celite. The filtrate was concentrated by rotary evaporation, and was dried briefly under high vacuum to yield Compound 105a (3.28 g). LCMS and ¹H NMR were consistent with desired product.

Compound 147 (2.31 g, 11 mmol) was dissolved in anhydrous DMF (100 mL). *N,N*-Diisopropylethylamine (DIEA, 3.9 mL, 22 mmol) was added, followed by HBTU (4 g, 10.5 mmol). The

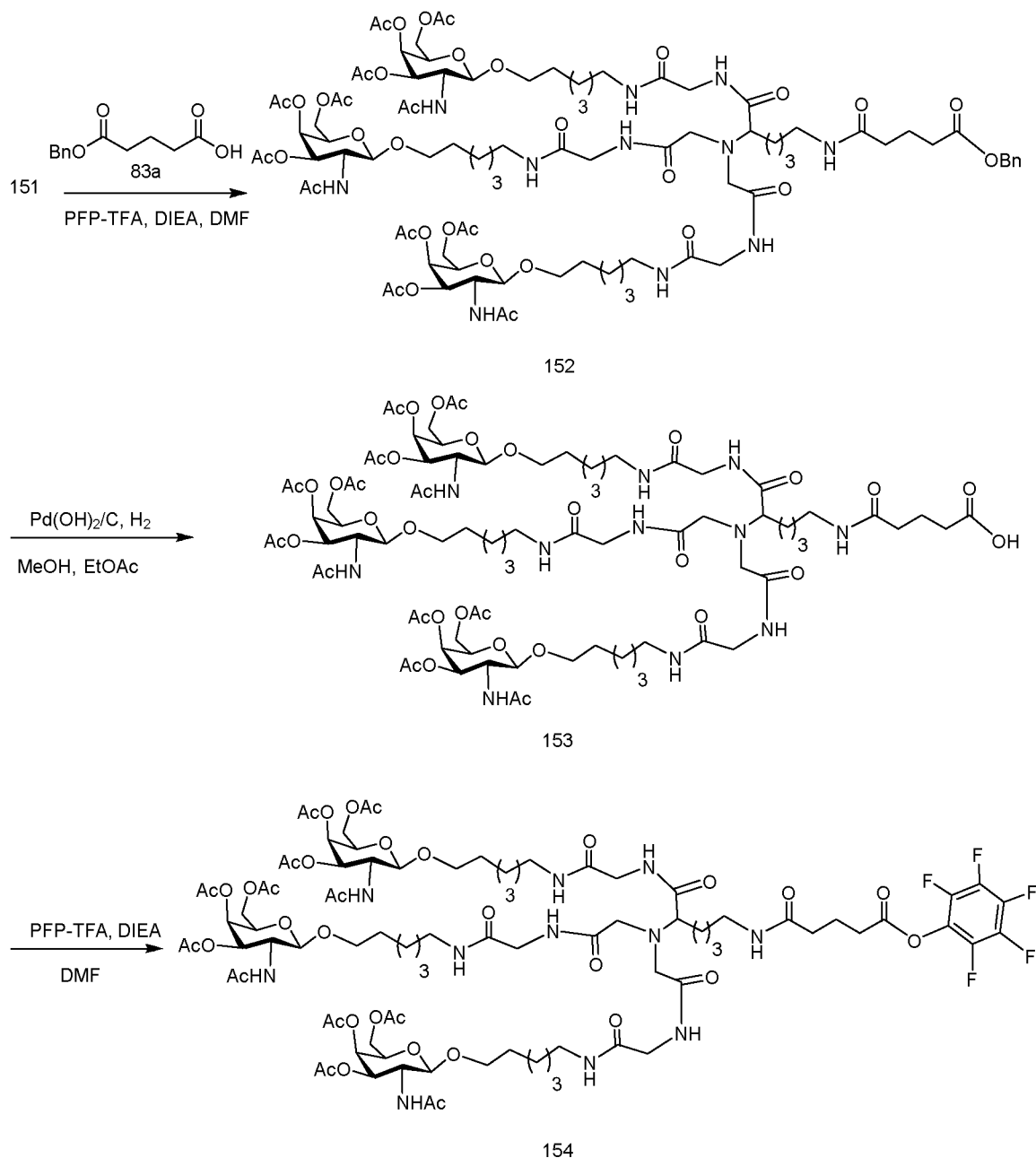
reaction mixture was allowed to stir for ~ 15 minutes under nitrogen. To this a solution of compound 105a (3.3 g, 7.4 mmol) in dry DMF was added and stirred for 2 h under nitrogen atmosphere. The reaction was diluted with EtOAc and washed with saturated aqueous NaHCO₃ and brine. The organics phase was separated, dried (MgSO₄), filtered, and concentrated to an orange syrup. The crude material was purified by column chromatography 2-5 % MeOH in CH₂Cl₂ to yield Compound 148 (3.44 g, 73 %). LCMS and ¹H NMR were consistent with the expected product.

Compound 148 (3.3 g, 5.2 mmol) was dissolved in 1:1 MeOH/EtOAc (75 ml). The reaction mixture was purged by bubbling a stream of argon through the solution for 15 minutes. Pearlman's catalyst (palladium hydroxide on carbon) was added (350 mg). Hydrogen gas was bubbled through the solution for 30 minutes. Upon completion (TLC 10% MeOH in DCM, and LCMS), the catalyst was removed by filtration through a pad of celite. The filtrate was concentrated by rotary evaporation, and was dried briefly under high vacuum to yield Compound 149 (2.6 g). LCMS was consistent with desired product. The residue was dissolved in dry DMF (10 ml) was used immediately in the next step.



Compound 146 (0.68 g, 1.73 mmol) was dissolved in dry DMF (20 mL). To this DIEA (450 μ L, 2.6 mmol, 1.5 eq.) and HBTU (1.96 g, 0.5.2 mmol) were added. The reaction mixture was allowed to stir for 15 minutes at room temperature under nitrogen. A solution of compound 149 (2.6 g) in anhydrous DMF (10 mL) was added. The pH of the reaction was adjusted to pH = 9-10 by addition of DIEA (if necessary). The reaction was allowed to stir at room temperature under nitrogen for 2 h. Upon completion the reaction was diluted with EtOAc (100 mL), and washed with aqueous saturated aqueous NaHCO_3 , followed by brine. The organic phase was separated, dried over MgSO_4 , filtered, and concentrated. The residue was purified by silica gel column chromatography and eluted with 2-10 % MeOH in CH_2Cl_2 to yield Compound 150 (0.62 g, 20 %). LCMS and ^1H NMR were consistent with the desired product.

Compound 150 (0.62 g) was dissolved in 1:1 MeOH/ EtOAc (5 L). The reaction mixture was purged by bubbling a stream of argon through the solution for 15 minutes. Pearlman's catalyst (palladium hydroxide on carbon) was added (60 mg). Hydrogen gas was bubbled through the solution for 30 minutes. Upon completion (TLC 10% MeOH in DCM, and LCMS), the catalyst was removed by filtration (syringe-tip Teflon filter, 0.45 μ m). The filtrate was concentrated by rotary evaporation, and was dried briefly under high vacuum to yield Compound 151 (0.57 g). The LCMS was consistent with the desired product. The product was dissolved in 4 mL dry DMF and was used immediately in the next step.

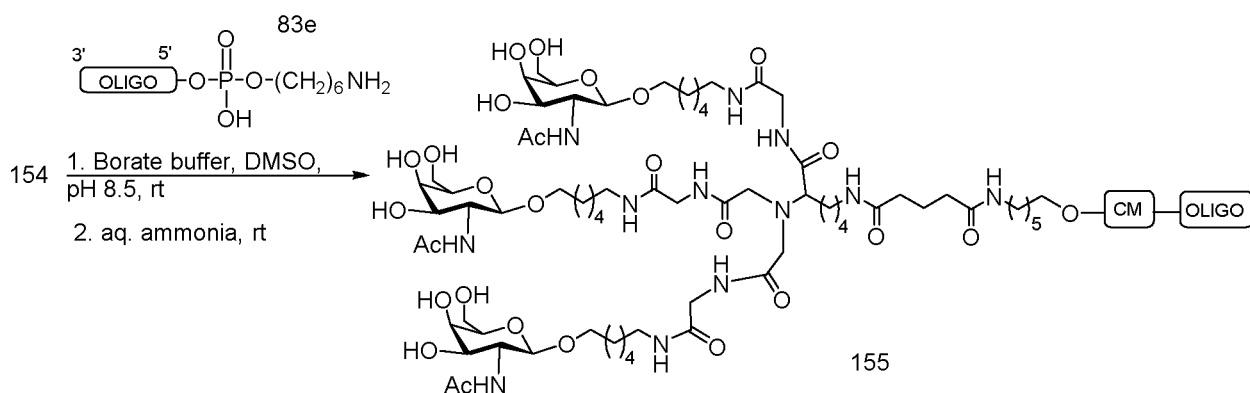


Compound 83a (0.11 g, 0.33 mmol) was dissolved in anhydrous DMF (5 mL) and *N,N*-Diisopropylethylamine (75 μL , 1 mmol) and PFP-TFA (90 μL , 0.76 mmol) were added. The reaction mixture turned magenta upon contact, and gradually turned orange over the next 30 minutes. Progress of reaction was monitored by TLC and LCMS. Upon completion (formation of the PFP ester), a solution of compound 151 (0.57 g, 0.33 mmol) in DMF was added. The pH of the reaction was adjusted to pH = 9-10 by addition of *N,N*-Diisopropylethylamine (if necessary). The reaction mixture was stirred under nitrogen for ~30 min. Upon completion, the majority of the solvent was removed under reduced pressure. The residue was diluted with CH_2Cl_2 and washed with aqueous saturated NaHCO_3 , followed by brine. The organic phase separated, dried over MgSO_4 , filtered, and concentrated to an orange syrup. The residue was purified by

silica gel column chromatography (2-10 % MeOH in CH₂Cl₂) to yield Compound 152 (0.35 g, 55 %). LCMS and ¹H NMR were consistent with the desired product.

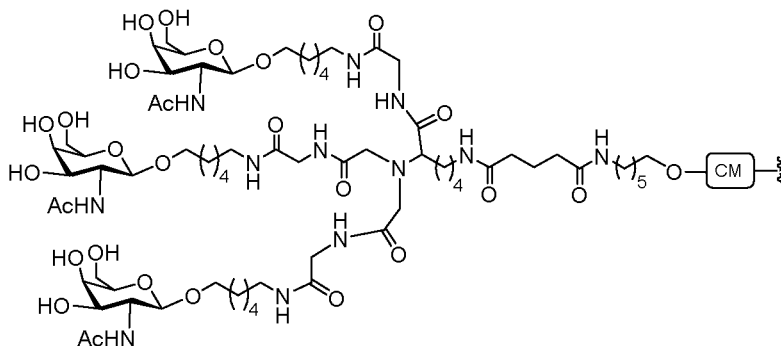
Compound 152 (0.35 g, 0.182 mmol) was dissolved in 1:1 MeOH/EtOAc (10 mL). The reaction mixture was purged by bubbling a stream of argon thru the solution for 15 minutes. Pearlman's catalyst (palladium hydroxide on carbon) was added (35 mg). Hydrogen gas was bubbled thru the solution for 30 minutes. Upon completion (TLC 10% MeOH in DCM, and LCMS), the catalyst was removed by filtration (syringe-tip Teflon filter, 0.45 μm). The filtrate was concentrated by rotary evaporation, and was dried briefly under high vacuum to yield Compound 153 (0.33 g, quantitative). The LCMS was consistent with desired product.

Compound 153 (0.33 g, 0.18 mmol) was dissolved in anhydrous DMF (5 mL) with stirring under nitrogen. To this *N,N*-Diisopropylethylamine (65 μL, 0.37 mmol) and PFP-TFA (35 μL, 0.28 mmol) were added. The reaction mixture was stirred under nitrogen for ~ 30 min. The reaction mixture turned magenta upon contact, and gradually turned orange. The pH of the reaction mixture was maintained at pH = 9-10 by adding more *N,N*-Diisopropylethylamine. The progress of the reaction was monitored by TLC and LCMS. Upon completion, the majority of the solvent was removed under reduced pressure. The residue was diluted with CH₂Cl₂ (50 mL), and washed with saturated aqueous NaHCO₃, followed by brine. The organic layer was dried over MgSO₄, filtered, and concentrated to an orange syrup. The residue was purified by column chromatography and eluted with 2-10 % MeOH in CH₂Cl₂ to yield Compound 154 (0.29 g, 79 %). LCMS and ¹H NMR were consistent with the desired product.

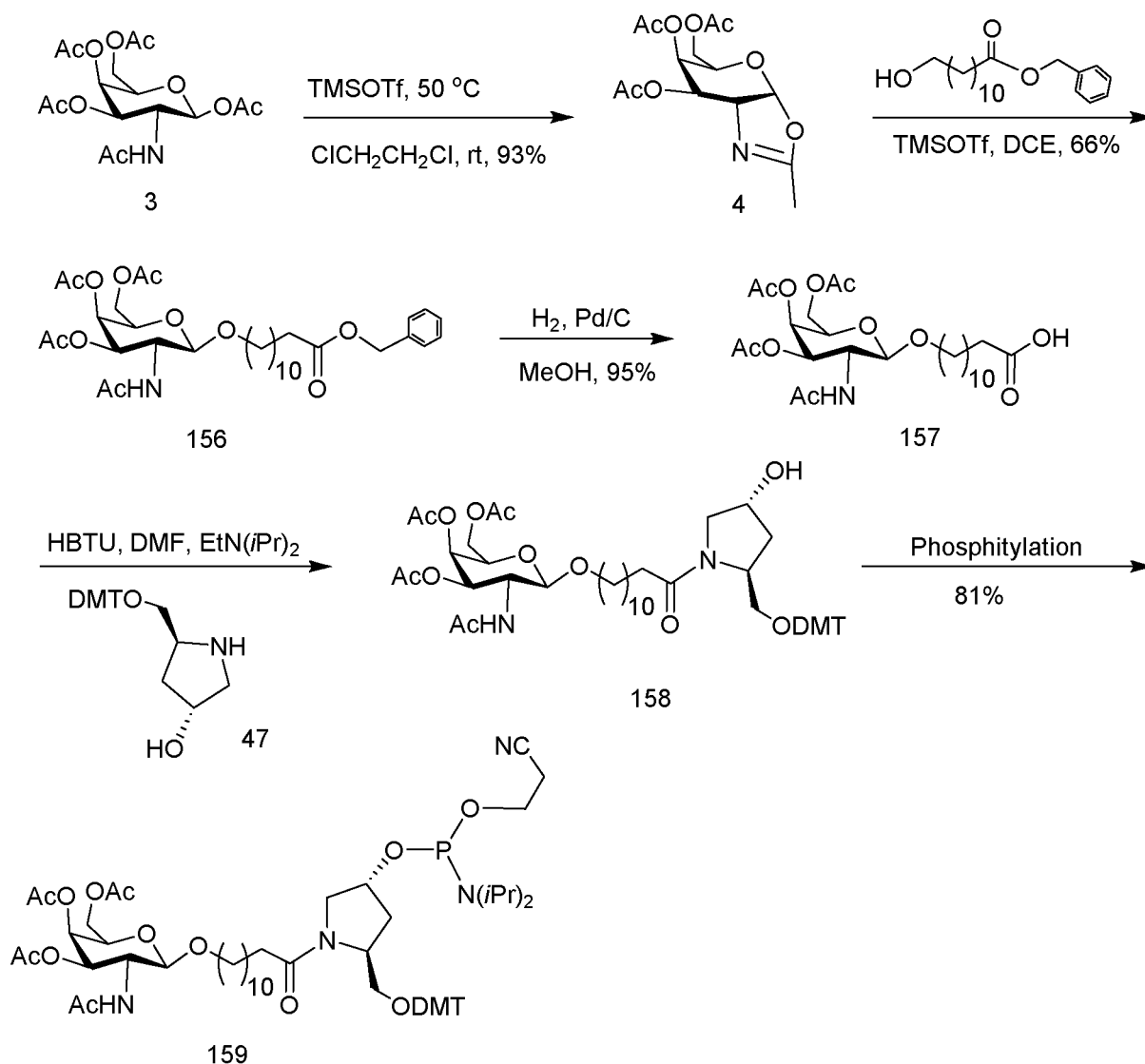


Oligomeric Compound 155, comprising a GalNAc₃-6 conjugate group, was prepared using the general procedures illustrated in Example 46. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-6 (GalNAc₃-6_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-.

The structure of GalNAc₃-6 (GalNAc₃-6_a-CM-) is shown below:



Example 52: Preparation of Oligonucleotide 160 Comprising GalNAc₃-9



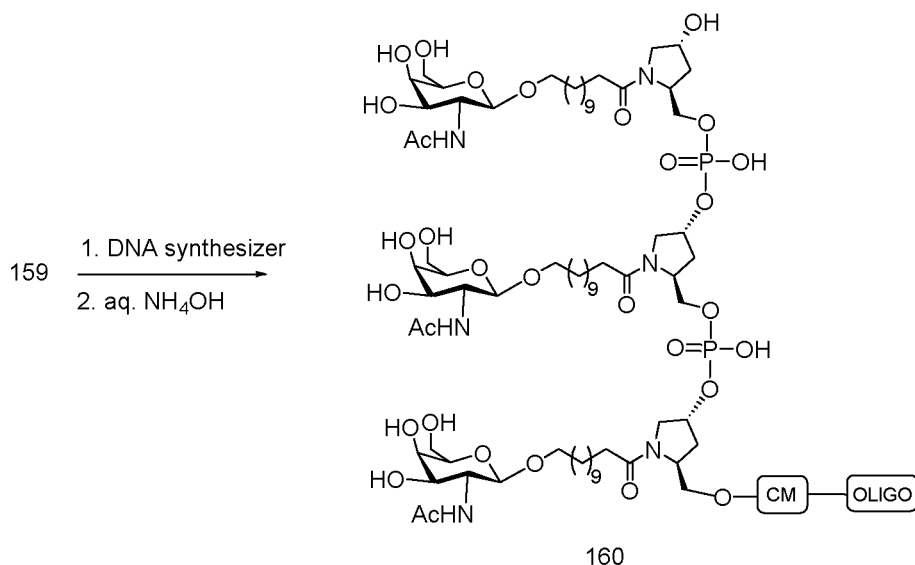
Compound 156 was synthesized following the procedure described in the literature (*J. Med. Chem.* 2004, 47, 5798-5808).

Compound 156, (18.60 g, 29.28 mmol) was dissolved in methanol (200 mL). Palladium on carbon (6.15 g, 10 wt%, loading (dry basis), matrix carbon powder, wet) was added. The reaction mixture was stirred at room temperature under hydrogen for 18 h. The reaction mixture was filtered through a pad of

celite and the celite pad was washed thoroughly with methanol. The combined filtrate was washed and concentrated to dryness. The residue was purified by silica gel column chromatography and eluted with 5-10 % methanol in dichloromethane to yield Compound 157 (14.26 g, 89%). Mass m/z 544.1 $[M-H]^+$.

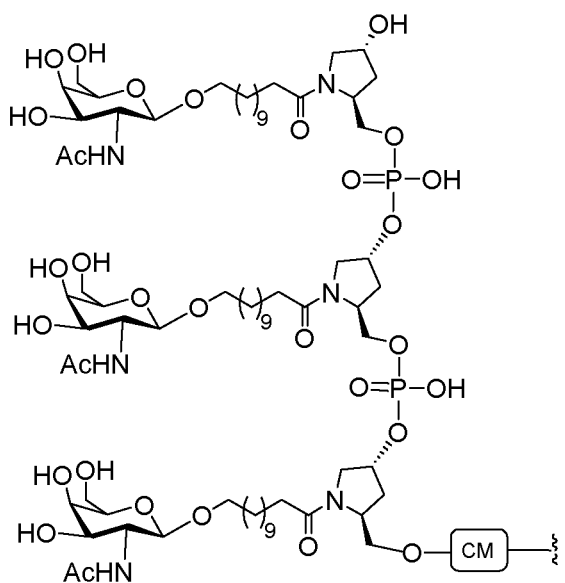
Compound 157 (5 g, 9.17 mmol) was dissolved in anhydrous DMF (30 mL). HBTU (3.65 g, 9.61 mmol) and *N,N*-Diisopropylethylamine (13.73 mL, 78.81 mmol) were added and the reaction mixture was stirred at room temperature for 5 minutes. To this a solution of compound 47 (2.96 g, 7.04 mmol) was added. The reaction was stirred at room temperature for 8 h. The reaction mixture was poured into a saturated NaHCO_3 aqueous solution. The mixture was extracted with ethyl acetate and the organic layer was washed with brine and dried (Na_2SO_4), filtered and evaporated. The residue obtained was purified by silica gel column chromatography and eluted with 50% ethyl acetate in hexane to yield compound 158 (8.25g, 73.3%). The structure was confirmed by MS and ^1H NMR analysis.

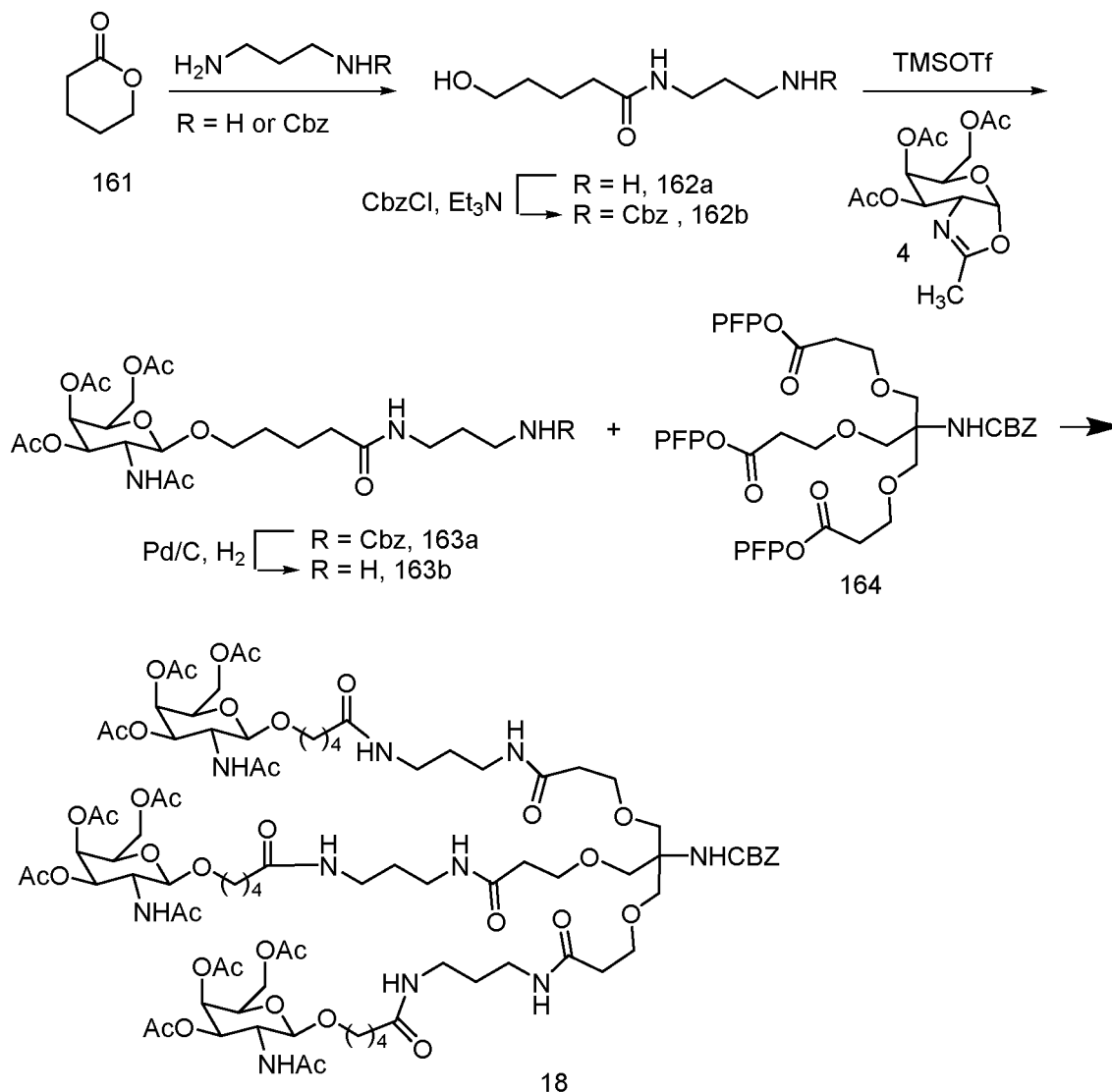
Compound 158 (7.2 g, 7.61 mmol) was dried over P_2O_5 under reduced pressure. The dried compound was dissolved in anhydrous DMF (50 mL). To this 1H-tetrazole (0.43 g, 6.09 mmol) and *N*-methylimidazole (0.3 mL, 3.81 mmol) and 2-cyanoethyl-*N,N,N',N'*-tetraisopropyl phosphorodiamidite (3.65 mL, 11.50 mmol) were added. The reaction mixture was stirred under an argon atmosphere for 4 h. The reaction mixture was diluted with ethyl acetate (200 mL). The reaction mixture was washed with saturated NaHCO_3 and brine. The organic phase was separated, dried (Na_2SO_4), filtered and evaporated. The residue was purified by silica gel column chromatography and eluted with 50-90 % ethyl acetate in hexane to yield Compound 159 (7.82 g, 80.5%). The structure was confirmed by LCMS and ^{31}P NMR analysis.



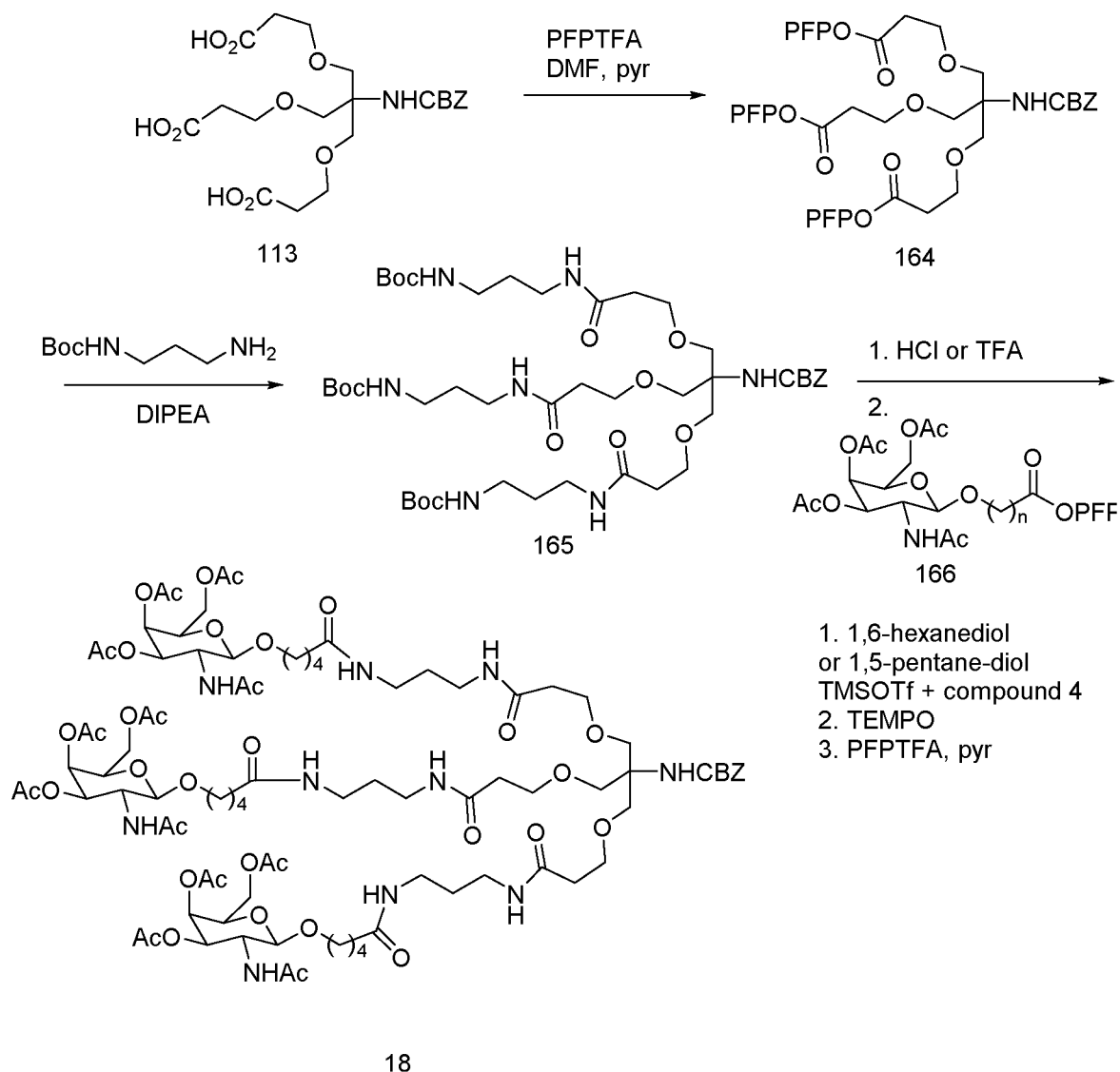
Oligomeric Compound 160, comprising a GalNAc_3 -9 conjugate group, was prepared using standard oligonucleotide synthesis procedures. Three units of compound 159 were coupled to the solid support, followed by nucleotide phosphoramidites. Treatment of the protected oligomeric compound with aqueous ammonia yielded compound 160. The GalNAc_3 cluster portion of the conjugate group GalNAc_3 -9 (GalNAc_3 -

9_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-. The structure of GalNAc₃-9 (GalNAc₃-9_a-CM) is shown below:



Example 53: Alternate procedure for preparation of Compound 18 (GalNAc₃-1a and GalNAc₃-3a)

Lactone 161 was reacted with diamino propane (3-5 eq) or Mono-Boc protected diamino propane (1
 5 eq) to provide alcohol 162a or 162b. When unprotected propanediamine was used for the above reaction, the
 excess diamine was removed by evaporation under high vacuum and the free amino group in 162a was
 protected using CbzCl to provide 162b as a white solid after purification by column chromatography.
 Alcohol 162b was further reacted with compound 4 in the presence of TMSOTf to provide 163a which was
 converted to 163b by removal of the Cbz group using catalytic hydrogenation. The pentafluorophenyl (PFP)
 10 ester 164 was prepared by reacting triacid 113 (see Example 48) with PFPTFA (3.5 eq) and pyridine (3.5 eq)
 in DMF (0.1 to 0.5 M). The triester 164 was directly reacted with the amine 163b (3-4 eq) and DIPEA (3-4
 eq) to provide Compound 18. The above method greatly facilitates purification of intermediates and
 minimizes the formation of byproducts which are formed using the procedure described in Example 4.

Example 54: Alternate procedure for preparation of Compound 18 (GalNAc₃-1a and GalNAc₃-3a)

The triPFP ester 164 was prepared from acid 113 using the procedure outlined in example 53 above and reacted with mono-Boc protected diamine to provide 165 in essentially quantitative yield. The Boc groups were removed with hydrochloric acid or trifluoroacetic acid to provide the triamine which was reacted with the PFP activated acid 166 in the presence of a suitable base such as DIPEA to provide Compound 18.

The PFP protected Gal-NAc acid 166 was prepared from the corresponding acid by treatment with PFPTFA (1-1.2 eq) and pyridine (1-1.2 eq) in DMF. The precursor acid in turn was prepared from the corresponding alcohol by oxidation using TEMPO (0.2 eq) and BAIB in acetonitrile and water. The precursor alcohol was prepared from sugar intermediate 4 by reaction with 1,6-hexanediol (or 1,5-pentanediol or other diol for other n values) (2-4 eq) and TMSOTf using conditions described previously in example 47.

Example 55: Dose-dependent study of oligonucleotides comprising either a 3' or 5'-conjugate group (comparison of GalNAc₃-1, 3, 8 and 9) targeting SRB-1 *in vivo*

The oligonucleotides listed below were tested in a dose-dependent study for antisense inhibition of SRB-1 in mice. Unconjugated ISIS 353382 was included as a standard. Each of the various GalNAc₃ conjugate groups was attached at either the 3' or 5' terminus of the respective oligonucleotide by a phosphodiester linked 2'-deoxyadenosine nucleoside (cleavable moiety).

Table 39
Modified ASO targeting SRB-1

ASO	Sequence (5' to 3')	Motif	Conjugate	SEQ ID No.
ISIS 353382 (parent)	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	none	143
ISIS 655861	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _{eo} A_{do}'-GalNAc₃-1_a	5/10/5	GalNAc₃-1	144
ISIS 664078	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _{eo} A_{do}'-GalNAc₃-9_a	5/10/5	GalNAc₃-9	144
ISIS 661161	GalNAc₃-3_a-o'-A_{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	GalNAc₃-3	145
ISIS 665001	GalNAc₃-8_a-o'-A_{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	GalNAc₃-8	145

Capital letters indicate the nucleobase for each nucleoside and ^mC indicates a 5-methyl cytosine. Subscripts: "e" indicates a 2'-MOE modified nucleoside; "d" indicates a β-D-2'-deoxyribonucleoside; "s" indicates a phosphorothioate internucleoside linkage (PS); "o" indicates a phosphodiester internucleoside linkage (PO); and "o'" indicates -O-P(=O)(OH)-. Conjugate groups are in bold.

The structure of GalNAc₃-1_a was shown previously in Example 9. The structure of GalNAc₃-9 was shown previously in Example 52. The structure of GalNAc₃-3 was shown previously in Example 39. The structure of GalNAc₃-8 was shown previously in Example 47.

Treatment

Six week old male Balb/c mice (Jackson Laboratory, Bar Harbor, ME) were injected subcutaneously once at the dosage shown below with ISIS 353382, 655861, 664078, 661161, 665001 or with saline. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration to determine the liver SRB-1 mRNA levels using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. The results below are presented as the average percent of SRB-1 mRNA levels for each treatment group, normalized to the saline control.

As illustrated in Table 40, treatment with antisense oligonucleotides lowered SRB-1 mRNA levels in a dose-dependent manner. Indeed, the antisense oligonucleotides comprising the phosphodiester linked GalNAc₃-1 and GalNAc₃-9 conjugates at the 3' terminus (ISIS 655861 and ISIS 664078) and the GalNAc₃-3 and GalNAc₃-8 conjugates linked at the 5' terminus (ISIS 661161 and ISIS 665001) showed substantial improvement in potency compared to the unconjugated antisense oligonucleotide (ISIS 353382). Furthermore, ISIS 664078, comprising a GalNAc₃-9 conjugate at the 3' terminus was essentially equipotent compared to ISIS 655861, which comprises a GalNAc₃-1 conjugate at the 3' terminus. The 5' conjugated antisense oligonucleotides, ISIS 661161 and ISIS 665001, comprising a GalNAc₃-3 or GalNAc₃-9, respectively, had increased potency compared to the 3' conjugated antisense oligonucleotides (ISIS 655861 and ISIS 664078).

Table 40
ASOs containing GalNAc₃-1, 3, 8 or 9 targeting SRB-1

ISIS No.	Dosage (mg/kg)	SRB-1 mRNA (% Saline)	Conjugate
Saline	n/a	100	
353382	3	88	none
	10	68	
	30	36	
655861	0.5	98	GalNAc ₃ -1 (3')
	1.5	76	
	5	31	
	15	20	
664078	0.5	88	GalNAc ₃ -9 (3')
	1.5	85	
	5	46	
	15	20	
661161	0.5	92	GalNAc ₃ -3 (5')
	1.5	59	
	5	19	
	15	11	
665001	0.5	100	GalNAc ₃ -8 (5')
	1.5	73	
	5	29	
	15	13	

Liver transaminase levels, alanine aminotransferase (ALT) and aspartate aminotransferase (AST), in serum were measured relative to saline injected mice using standard protocols. Total bilirubin and BUN were also evaluated. The change in body weights was evaluated with no significant change from the saline group. ALTs, ASTs, total bilirubin and BUN values are shown in the table below.

Table 41

ISIS No.	Dosage mg/kg	ALT	AST	Total Bilirubin	BUN	Conjugate
Saline		24	59	0.1	37.52	
353382	3	21	66	0.2	34.65	none
	10	22	54	0.2	34.2	
	30	22	49	0.2	33.72	
655861	0.5	25	62	0.2	30.65	GalNac ₃ -1 (3')
	1.5	23	48	0.2	30.97	
	5	28	49	0.1	32.92	
	15	40	97	0.1	31.62	
664078	0.5	40	74	0.1	35.3	GalNac ₃ -9 (3')
	1.5	47	104	0.1	32.75	
	5	20	43	0.1	30.62	
	15	38	92	0.1	26.2	
661161	0.5	101	162	0.1	34.17	GalNac ₃ -3 (5')
	1.5 g	42	100	0.1	33.37	
	5 g	23	99	0.1	34.97	
	15	53	83	0.1	34.8	
665001	0.5	28	54	0.1	31.32	GalNac ₃ -8 (5')
	1.5	42	75	0.1	32.32	
	5	24	42	0.1	31.85	
	15	32	67	0.1	31.	

Example 56: Dose-dependent study of oligonucleotides comprising either a 3' or 5'-conjugate group (comparison of GalNac₃-1, 2, 3, 5, 6, 7 and 10) targeting SRB-1 *in vivo*

The oligonucleotides listed below were tested in a dose-dependent study for antisense inhibition of SRB-1 in mice. Unconjugated ISIS 353382 was included as a standard. Each of the various GalNac₃ conjugate groups was attached at the 5' terminus of the respective oligonucleotide by a phosphodiester linked 2'-deoxyadenosine nucleoside (cleavable moiety) except for ISIS 655861 which had the GalNac₃ conjugate group attached at the 3' terminus.

Table 42

Modified ASO targeting SRB-1

ASO	Sequence (5' to 3')	Motif	Conjugate	SEQ ID No.
ISIS 353382 (parent)	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} mC _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	no conjugate	143
ISIS 655861	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} mC _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _{eo} A _{do} '-GalNac ₃ -1 _a	5/10/5	GalNac ₃ -1	144
ISIS 664507	GalNac ₃ -2 _a -o'-A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} mC _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	GalNac ₃ -2	145
ISIS 661161	GalNac ₃ -3 _a -o'-A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} mC _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	GalNac ₃ -3	145
ISIS 666224	GalNac ₃ -5 _a -o'-A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds}	5/10/5	GalNac ₃ -5	145

	$^mC_{ds}A_{ds}T_{ds}G_{ds}A_{ds}^mC_{ds}T_{ds}T_{es}^mC_{es}^mC_{es}T_{es}T_e$			
ISIS 666961	GalNAc₃-6_a-o'-A_{do} $G_{es}^mC_{es}T_{es}T_{es}^mC_{es}A_{ds}G_{ds}T_{ds}$ $^mC_{ds}A_{ds}T_{ds}G_{ds}A_{ds}^mC_{ds}T_{ds}T_{es}^mC_{es}^mC_{es}T_{es}T_e$	5/10/5	GalNAc₃-6	145
ISIS 666981	GalNAc₃-7_a-o'-A_{do} $G_{es}^mC_{es}T_{es}T_{es}^mC_{es}A_{ds}G_{ds}T_{ds}$ $^mC_{ds}A_{ds}T_{ds}G_{ds}A_{ds}^mC_{ds}T_{ds}T_{es}^mC_{es}^mC_{es}T_{es}T_e$	5/10/5	GalNAc₃-7	145
ISIS 666881	GalNAc₃-10_a-o'-A_{do} $G_{es}^mC_{es}T_{es}T_{es}^mC_{es}A_{ds}G_{ds}T_{ds}$ $^mC_{ds}A_{ds}T_{ds}G_{ds}A_{ds}^mC_{ds}T_{ds}T_{es}^mC_{es}^mC_{es}T_{es}T_e$	5/10/5	GalNAc₃-10	145

Capital letters indicate the nucleobase for each nucleoside and mC indicates a 5-methyl cytosine. Subscripts: “e” indicates a 2'-MOE modified nucleoside; “d” indicates a β -D-2'-deoxyribonucleoside; “s” indicates a phosphorothioate internucleoside linkage (PS); “o” indicates a phosphodiester internucleoside linkage (PO); and “o'” indicates -O-P(=O)(OH)-. Conjugate groups are in bold.

The structure of GalNAc₃-1_a was shown previously in Example 9. The structure of GalNAc₃-2_a was shown previously in Example 37. The structure of GalNAc₃-3_a was shown previously in Example 39. The structure of GalNAc₃-5_a was shown previously in Example 49. The structure of GalNAc₃-6_a was shown previously in Example 51. The structure of GalNAc₃-7_a was shown previously in Example 48. The structure of GalNAc₃-10_a was shown previously in Example 46.

Treatment

Six week old male Balb/c mice (Jackson Laboratory, Bar Harbor, ME) were injected subcutaneously once at the dosage shown below with ISIS 353382, 655861, 664507, 661161, 666224, 666961, 666981, 666881 or with saline. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration to determine the liver SRB-1 mRNA levels using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. The results below are presented as the average percent of SRB-1 mRNA levels for each treatment group, normalized to the saline control.

As illustrated in Table 43, treatment with antisense oligonucleotides lowered SRB-1 mRNA levels in a dose-dependent manner. Indeed, the conjugated antisense oligonucleotides showed substantial improvement in potency compared to the unconjugated antisense oligonucleotide (ISIS 353382). The 5' conjugated antisense oligonucleotides showed a slight increase in potency compared to the 3' conjugated antisense oligonucleotide.

Table 43

ISIS No.	Dosage (mg/kg)	SRB-1 mRNA (% Saline)	Conjugate
Saline	n/a	100.0	
353382	3	96.0	none
	10	73.1	
	30	36.1	
655861	0.5	99.4	GalNAc₃-1 (3')

	1.5	81.2	
	5	33.9	
	15	15.2	
664507	0.5	102.0	GalNac₃-2 (5')
	1.5	73.2	
	5	31.3	
	15	10.8	
661161	0.5	90.7	GalNac₃-3 (5')
	1.5	67.6	
	5	24.3	
	15	11.5	
666224	0.5	96.1	GalNac₃-5 (5')
	1.5	61.6	
	5	25.6	
	15	11.7	
666961	0.5	85.5	GalNac₃-6 (5')
	1.5	56.3	
	5	34.2	
	15	13.1	
666981	0.5	84.7	GalNac₃-7 (5')
	1.5	59.9	
	5	24.9	
	15	8.5	
666881	0.5	100.0	GalNac₃-10 (5')
	1.5	65.8	
	5	26.0	
	15	13.0	

Liver transaminase levels, alanine aminotransferase (ALT) and aspartate aminotransferase (AST), in serum were measured relative to saline injected mice using standard protocols. Total bilirubin and BUN were also evaluated. The change in body weights was evaluated with no significant change from the saline group.

5 ALTs, ASTs, total bilirubin and BUN values are shown in Table 44 below.

Table 44

ISIS No.	Dosage mg/kg	ALT	AST	Total Bilirubin	BUN	Conjugate
Saline		26	57	0.2	27	
353382	3	25	92	0.2	27	none
	10	23	40	0.2	25	
	30	29	54	0.1	28	
655861	0.5	25	71	0.2	34	GalNac₃-1 (3')
	1.5	28	60	0.2	26	
	5	26	63	0.2	28	
	15	25	61	0.2	28	
664507	0.5	25	62	0.2	25	GalNac₃-2 (5')
	1.5	24	49	0.2	26	
	5	21	50	0.2	26	
	15	59	84	0.1	22	

661161	0.5	20	42	0.2	29	GalNac₃-3 (5')
	1.5 g	37	74	0.2	25	
	5 g	28	61	0.2	29	
	15	21	41	0.2	25	
666224	0.5	34	48	0.2	21	GalNac₃-5 (5')
	1.5	23	46	0.2	26	
	5	24	47	0.2	23	
	15	32	49	0.1	26	
666961	0.5	17	63	0.2	26	GalNac₃-6 (5')
	1.5	23	68	0.2	26	
	5	25	66	0.2	26	
	15	29	107	0.2	28	
666981	0.5	24	48	0.2	26	GalNac₃-7 (5')
	1.5	30	55	0.2	24	
	5	46	74	0.1	24	
	15	29	58	0.1	26	
666881	0.5	20	65	0.2	27	GalNac₃-10 (5')
	1.5	23	59	0.2	24	
	5	45	70	0.2	26	
	15	21	57	0.2	24	

Example 57: Duration of action study of oligonucleotides comprising a 3'-conjugate group targeting ApoC III *in vivo*

Mice were injected once with the doses indicated below and monitored over the course of 42 days for ApoC-III and plasma triglycerides (Plasma TG) levels. The study was performed using 3 transgenic mice that express human APOC-III in each group.

Table 45
Modified ASO targeting ApoC III

ASO	Sequence (5' to 3')	Linkages	SEQ ID No.
ISIS 304801	A _{es} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{es} T _{es} T _{es} A _{es} T _e	PS	135
ISIS 647535	A _{es} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{es} T _{es} T _{es} A _{es} T _{eo} A_{do}-GalNac₃-1_a	PS	136
ISIS 647536	A _{es} G _{eo} ^m C _{eo} T _{eo} T _{eo} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{eo} T _{eo} T _{es} A _{es} T _{eo} A_{do}-GalNac₃-1_a	PO/PS	136

Capital letters indicate the nucleobase for each nucleoside and ^mC indicates a 5-methyl cytosine. Subscripts: “e” indicates a 2'-MOE modified nucleoside; “d” indicates a β-D-2'-deoxyribonucleoside; “s” indicates a phosphorothioate internucleoside linkage (PS); “o” indicates a phosphodiester internucleoside linkage (PO); and “o” indicates -O-P(=O)(OH)-. Conjugate groups are in bold.

The structure of GalNac₃-1_a was shown previously in Example 9.

Table 46

ApoC III mRNA (% Saline on Day 1) and Plasma TG Levels (% Saline on Day 1)

ASO	Dose	Target	Day 3	Day 7	Day 14	Day 35	Day 42
Saline	0 mg/kg	ApoC-III	98	100	100	95	116
ISIS 304801	30 mg/kg	ApoC-III	28	30	41	65	74
ISIS 647535	10 mg/kg	ApoC-III	16	19	25	74	94
ISIS 647536	10 mg/kg	ApoC-III	18	16	17	35	51
Saline	0 mg/kg	Plasma TG	121	130	123	105	109
ISIS 304801	30 mg/kg	Plasma TG	34	37	50	69	69
ISIS 647535	10 mg/kg	Plasma TG	18	14	24	18	71
ISIS 647536	10 mg/kg	Plasma TG	21	19	15	32	35

As can be seen in the table above the duration of action increased with addition of the 3'-conjugate group compared to the unconjugated oligonucleotide. There was a further increase in the duration of action for the conjugated mixed PO/PS oligonucleotide 647536 as compared to the conjugated full PS oligonucleotide 647535.

Example 58: Dose-dependent study of oligonucleotides comprising a 3'-conjugate group (comparison of GalNAc₃-1 and GalNAc₄-11) targeting SRB-1 *in vivo*

The oligonucleotides listed below were tested in a dose-dependent study for antisense inhibition of SRB-1 in mice. Unconjugated ISIS 440762 was included as an unconjugated standard. Each of the conjugate groups were attached at the 3' terminus of the respective oligonucleotide by a phosphodiester linked 2'-deoxyadenosine nucleoside cleavable moiety.

The structure of GalNAc₃-1_a was shown previously in Example 9. The structure of GalNAc₃-11_a was shown previously in Example 50.

Treatment

Six week old male Balb/c mice (Jackson Laboratory, Bar Harbor, ME) were injected subcutaneously once at the dosage shown below with ISIS 440762, 651900, 663748 or with saline. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration to determine the liver SRB-1 mRNA levels using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. The results below are presented as the average percent of SRB-1 mRNA levels for each treatment group, normalized to the saline control.

As illustrated in Table 47, treatment with antisense oligonucleotides lowered SRB-1 mRNA levels in a dose-dependent manner. The antisense oligonucleotides comprising the phosphodiester linked GalNAc₃-1 and GalNAc₄-11 conjugates at the 3' terminus (ISIS 651900 and ISIS 663748) showed substantial

improvement in potency compared to the unconjugated antisense oligonucleotide (ISIS 440762). The two conjugated oligonucleotides, GalNac₃-1 and GalNac₄-11, were equipotent.

Table 47
Modified ASO targeting SRB-1

ASO	Sequence (5' to 3')	Dose mg/kg	% Saline control	SEQ ID No.
Saline			100	
ISIS 440762	T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _k	0.6	73.45	137
		2	59.66	
		6	23.50	
ISIS 651900	T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _{ko} A_{do}'-GalNac₃-1_a	0.2	62.75	138
		0.6	29.14	
		2	8.61	
		6	5.62	
ISIS 663748	T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _{ko} A_{do}'-GalNac₄-11_a	0.2	63.99	138
		0.6	33.53	
		2	7.58	
		6	5.52	

5

Capital letters indicate the nucleobase for each nucleoside and ^mC indicates a 5-methyl cytosine. Subscripts: “e” indicates a 2'-MOE modified nucleoside; “k” indicates 6'-(S)-CH₃ bicyclic nucleoside; “d” indicates a β-D-2'-deoxyribonucleoside; “s” indicates a phosphorothioate internucleoside linkage (PS); “o” indicates a phosphodiester internucleoside linkage (PO); and “o'” indicates -O-P(=O)(OH)-. Conjugate groups are in bold.

10

Liver transaminase levels, alanine aminotransferase (ALT) and aspartate aminotransferase (AST), in serum were measured relative to saline injected mice using standard protocols. Total bilirubin and BUN were also evaluated. The change in body weights was evaluated with no significant change from the saline group. ALTs, ASTs, total bilirubin and BUN values are shown in Table 48 below.

15

Table 48

ISIS No.	Dosage mg/kg	ALT	AST	Total Bilirubin	BUN	Conjugate
Saline		30	76	0.2	40	
440762	0.60	32	70	0.1	35	none
	2	26	57	0.1	35	
	6	31	48	0.1	39	
651900	0.2	32	115	0.2	39	GalNac ₃ -1 (3')
	0.6	33	61	0.1	35	
	2	30	50	0.1	37	
	6	34	52	0.1	36	
663748	0.2	28	56	0.2	36	GalNac ₄ -11 (3')
	0.6	34	60	0.1	35	
	2	44	62	0.1	36	

	6	38	71	0.1	33	
--	---	----	----	-----	----	--

Example 59: Effects of GalNAc₃-1 conjugated ASOs targeting FXI *in vivo*

The oligonucleotides listed below were tested in a multiple dose study for antisense inhibition of FXI in mice. ISIS 404071 was included as an unconjugated standard. Each of the conjugate groups was attached at the 3' terminus of the respective oligonucleotide by a phosphodiester linked 2'-deoxyadenosine nucleoside cleavable moiety.

Table 49
Modified ASOs targeting FXI

ASO	Sequence (5' to 3')	Linkages	SEQ ID No.
ISIS 404071	T _{es} G _{es} G _{es} T _{es} A _{es} A _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ds} T _{ds} ^m C _{ds} A _{es} G _{es} A _{es} G _{es} G _e	PS	146
ISIS 656172	T _{es} G _{es} G _{es} T _{es} A _{es} A _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ds} T _{ds} ^m C _{ds} A _{es} G _{es} A _{es} G _{es} G _{eo} A_{do}'-GalNAc₃-1_a	PS	147
ISIS 656173	T _{es} G _{eo} G _{eo} T _{eo} A _{eo} A _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ds} T _{ds} ^m C _{ds} A _{eo} G _{eo} A _{es} G _{es} G _{eo} A_{do}'-GalNAc₃-1_a	PO/PS	147

Capital letters indicate the nucleobase for each nucleoside and ^mC indicates a 5-methyl cytosine. Subscripts: "e" indicates a 2'-MOE modified nucleoside; "d" indicates a β-D-2'-deoxyribonucleoside; "s" indicates a phosphorothioate internucleoside linkage (PS); "o" indicates a phosphodiester internucleoside linkage (PO); and "o'" indicates -O-P(=O)(OH)-. Conjugate groups are in bold.

The structure of GalNAc₃-1_a was shown previously in Example 9.

Treatment

Six week old male Balb/c mice (Jackson Laboratory, Bar Harbor, ME) were injected subcutaneously twice a week for 3 weeks at the dosage shown below with ISIS 404071, 656172, 656173 or with PBS treated control. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration to determine the liver FXI mRNA levels using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. Plasma FXI protein levels were also measured using ELISA. FXI mRNA levels were determined relative to total RNA (using RIBOGREEN®), prior to normalization to PBS-treated control. The results below are presented as the average percent of FXI mRNA levels for each treatment group. The data was normalized to PBS-treated control and is denoted as "% PBS". The ED₅₀s were measured using similar methods as described previously and are presented below.

Table 50
Factor XI mRNA (% Saline)

ASO	Dose mg/kg	% Control	Conjugate	Linkages
-----	------------	-----------	-----------	----------

Saline		100	none	
ISIS 404071	3	92	none	PS
	10	40		
	30	15		
ISIS 656172	0.7	74	GalNAc₃-1	PS
	2	33		
	6	9		
ISIS 656173	0.7	49	GalNAc₃-1	PO/PS
	2	22		
	6	1		

As illustrated in Table 50, treatment with antisense oligonucleotides lowered FXI mRNA levels in a dose-dependent manner. The oligonucleotides comprising a 3'-GalNAc₃-1 conjugate group showed substantial improvement in potency compared to the unconjugated antisense oligonucleotide (ISIS 404071).
 5 Between the two conjugated oligonucleotides an improvement in potency was further provided by substituting some of the PS linkages with PO (ISIS 656173).

As illustrated in Table 50a, treatment with antisense oligonucleotides lowered FXI protein levels in a dose-dependent manner. The oligonucleotides comprising a 3'-GalNAc₃-1 conjugate group showed substantial improvement in potency compared to the unconjugated antisense oligonucleotide (ISIS 404071).
 10 Between the two conjugated oligonucleotides an improvement in potency was further provided by substituting some of the PS linkages with PO (ISIS 656173).

Table 50a
Factor XI protein (% Saline)

ASO	Dose mg/kg	Protein Control)	(%	Conjugate	Linkages
Saline		100		none	
ISIS 404071	3	127		none	PS
	10	32			
	30	3			
ISIS 656172	0.7	70		GalNAc₃-1	PS
	2	23			
	6	1			
ISIS 656173	0.7	45		GalNAc₃-1	PO/PS
	2	6			
	6	0			

15 Liver transaminase levels, alanine aminotransferase (ALT) and aspartate aminotransferase (AST), in serum were measured relative to saline injected mice using standard protocols. Total bilirubin, total albumin, CRE and BUN were also evaluated. The change in body weights was evaluated with no significant change from the saline group. ALTs, ASTs, total bilirubin and BUN values are shown in the table below.

Table 51

ISIS No.	Dosage mg/kg	ALT	AST	Total Albumin	Total Bilirubin	CRE	BUN	Conjugate
Saline		71.8	84.0	3.1	0.2	0.2	22.9	
404071	3	152.8	176.0	3.1	0.3	0.2	23.0	none
	10	73.3	121.5	3.0	0.2	0.2	21.4	
	30	82.5	92.3	3.0	0.2	0.2	23.0	
656172	0.7	62.5	111.5	3.1	0.2	0.2	23.8	GalNac ₃ -1 (3')
	2	33.0	51.8	2.9	0.2	0.2	22.0	
	6	65.0	71.5	3.2	0.2	0.2	23.9	
656173	0.7	54.8	90.5	3.0	0.2	0.2	24.9	GalNac ₃ -1 (3')
	2	85.8	71.5	3.2	0.2	0.2	21.0	
	6	114.0	101.8	3.3	0.2	0.2	22.7	

Example 60: Effects of conjugated ASOs targeting SRB-1 *in vitro*

The oligonucleotides listed below were tested in a multiple dose study for antisense inhibition of SRB-1 in primary mouse hepatocytes. ISIS 353382 was included as an unconjugated standard. Each of the conjugate groups were attached at the 3' or 5' terminus of the respective oligonucleotide by a phosphodiester linked 2'-deoxyadenosine nucleoside cleavable moiety.

Table 52

Modified ASO targeting SRB-1

ASO	Sequence (5' to 3')	Motif	Conjugate	SEQ ID No.
ISIS 353382	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	none	143
ISIS 655861	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _{eo} A _{do} '-GalNac ₃ -1 _a	5/10/5	GalNac ₃ -1	144
ISIS 655862	G _{es} ^m C _{eo} T _{eo} T _{eo} ^m C _{eo} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{eo} ^m C _{eo} ^m C _{es} T _{es} T _{eo} A _{do} '-GalNac ₃ -1 _a	5/10/5	GalNac ₃ -1	144
ISIS 661161	GalNac ₃ -3 _{a-o} 'A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	GalNac ₃ -3	145
ISIS 665001	GalNac ₃ -8 _{a-o} 'A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	GalNac ₃ -8	145
ISIS 664078	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _{eo} A _{do} '-GalNac ₃ -9 _a	5/10/5	GalNac ₃ -9	144
ISIS 666961	GalNac ₃ -6 _{a-o} 'A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	GalNac ₃ -6	145
ISIS 664507	GalNac ₃ -2 _{a-o} 'A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	GalNac ₃ -2	145
ISIS 666881	GalNac ₃ -10 _{a-o} 'A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	GalNac ₃ -10	145
ISIS 666224	GalNac ₃ -5 _{a-o} 'A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	GalNac ₃ -5	145
ISIS 666981	GalNac ₃ -7 _{a-o} 'A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	5/10/5	GalNac ₃ -7	145

Capital letters indicate the nucleobase for each nucleoside and ^mC indicates a 5-methyl cytosine. Subscripts: “e” indicates a 2'-MOE modified nucleoside; “d” indicates a β-D-2'-deoxyribonucleoside; “s” indicates a phosphorothioate internucleoside linkage (PS); “o” indicates a phosphodiester internucleoside linkage (PO); and “o” indicates -O-P(=O)(OH)-. Conjugate groups are in bold.

5 The structure of GalNAc₃-1_a was shown previously in Example 9. The structure of GalNAc₃-3a was shown previously in Example 39. The structure of GalNAc₃-8a was shown previously in Example 47. The structure of GalNAc₃-9a was shown previously in Example 52. The structure of GalNAc₃-6a was shown previously in Example 51. The structure of GalNAc₃-2a was shown previously in Example 37. The structure of GalNAc₃-10a was shown previously in Example 46. The structure of GalNAc₃-5a was shown previously in Example 49. The structure of GalNAc₃-7a was shown previously in Example 48.

Treatment

15 The oligonucleotides listed above were tested *in vitro* in primary mouse hepatocyte cells plated at a density of 25,000 cells per well and treated with 0.03, 0.08, 0.24, 0.74, 2.22, 6.67 or 20 nM modified oligonucleotide. After a treatment period of approximately 16 hours, RNA was isolated from the cells and mRNA levels were measured by quantitative real-time PCR and the SRB-1 mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN®.

20 The IC₅₀ was calculated using standard methods and the results are presented in Table 53. The results show that, under free uptake conditions in which no reagents or electroporation techniques are used to artificially promote entry of the oligonucleotides into cells, the oligonucleotides comprising a GalNAc conjugate were significantly more potent in hepatocytes than the parent oligonucleotide (ISIS 353382) that does not comprise a GalNAc conjugate.

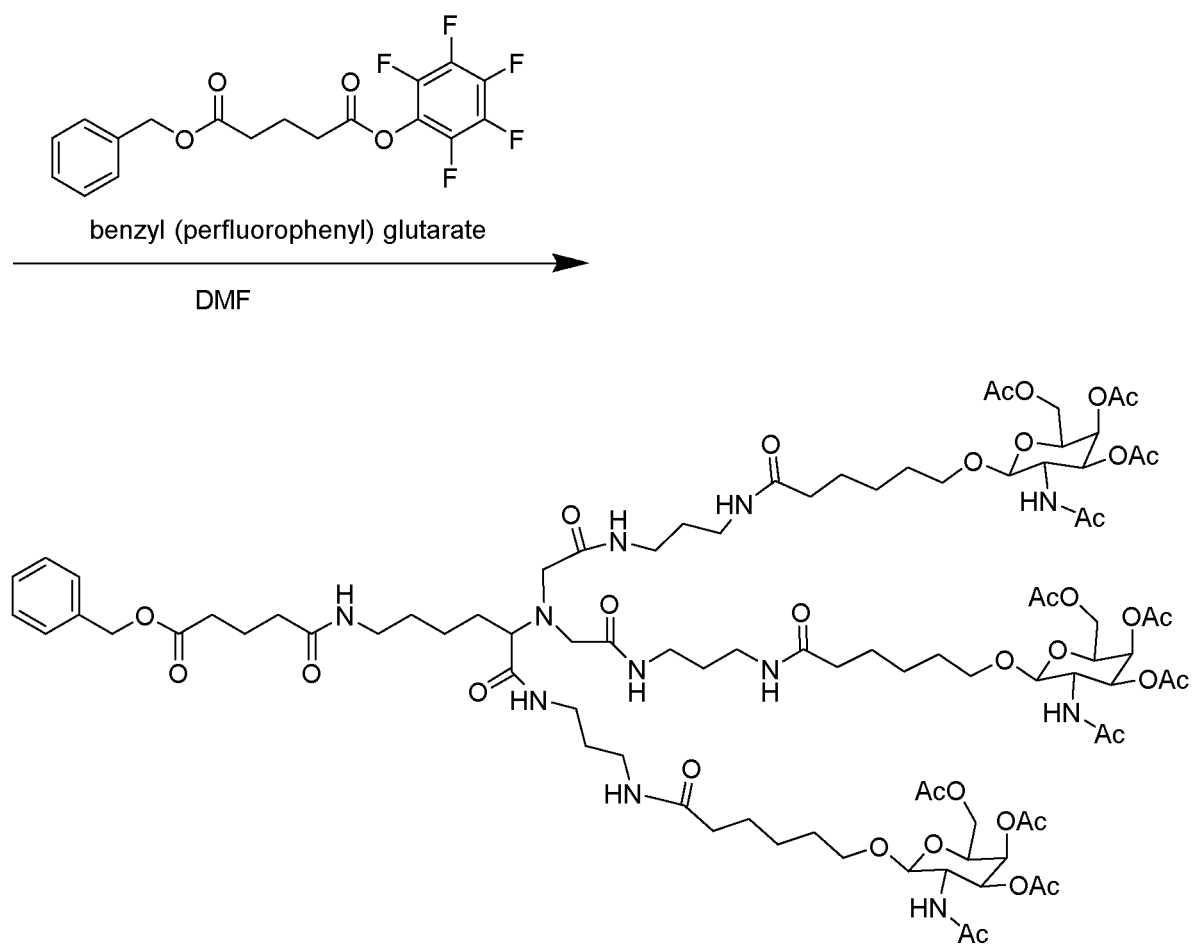
Table 53

ASO	IC ₅₀ (nM)	Internucleoside linkages	Conjugate	SEQ ID No.
ISIS 353382	190 ^a	PS	none	143
ISIS 655861	11 ^a	PS	GalNAc₃-1	144
ISIS 655862	3	PO/PS	GalNAc₃-1	144
ISIS 661161	15 ^a	PS	GalNAc₃-3	145
ISIS 665001	20	PS	GalNAc₃-8	145
ISIS 664078	55	PS	GalNAc₃-9	144
ISIS 666961	22 ^a	PS	GalNAc₃-6	145
ISIS 664507	30	PS	GalNAc₃-2	145
ISIS 666881	30	PS	GalNAc₃-10	145
ISIS 666224	30 ^a	PS	GalNAc₃-5	145
ISIS 666981	40	PS	GalNAc₃-7	145

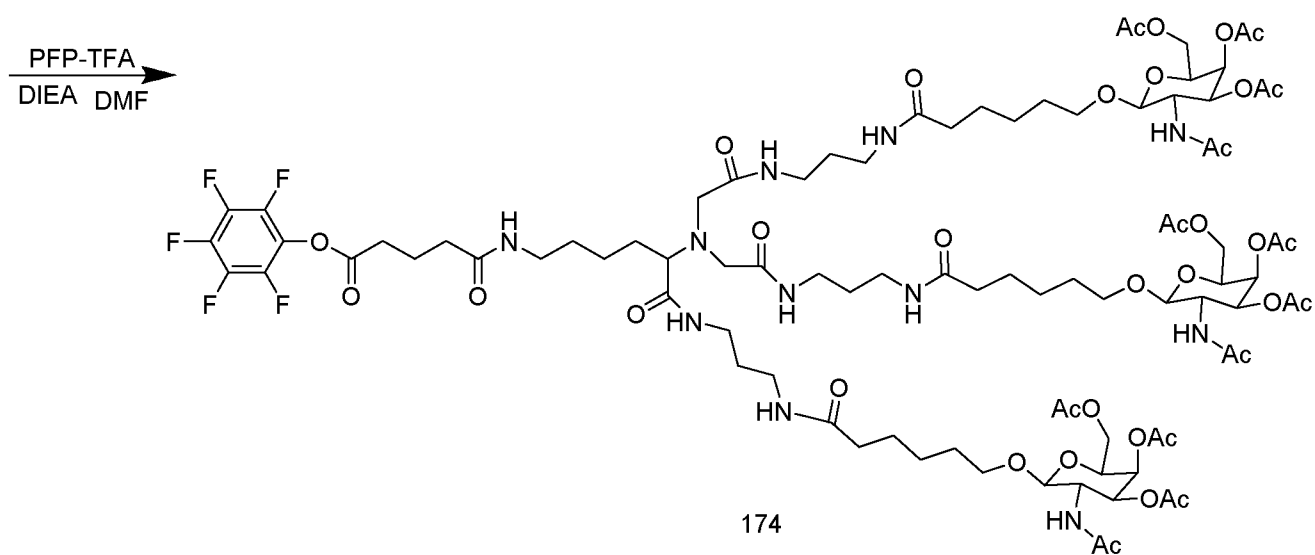
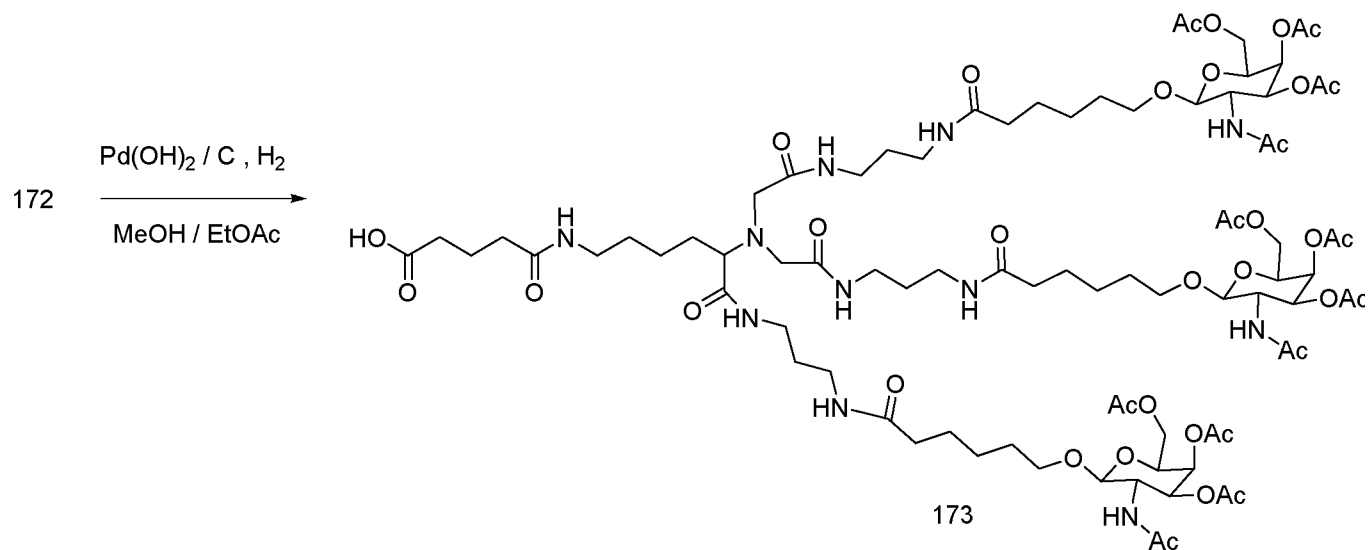
^aAverage of multiple runs.

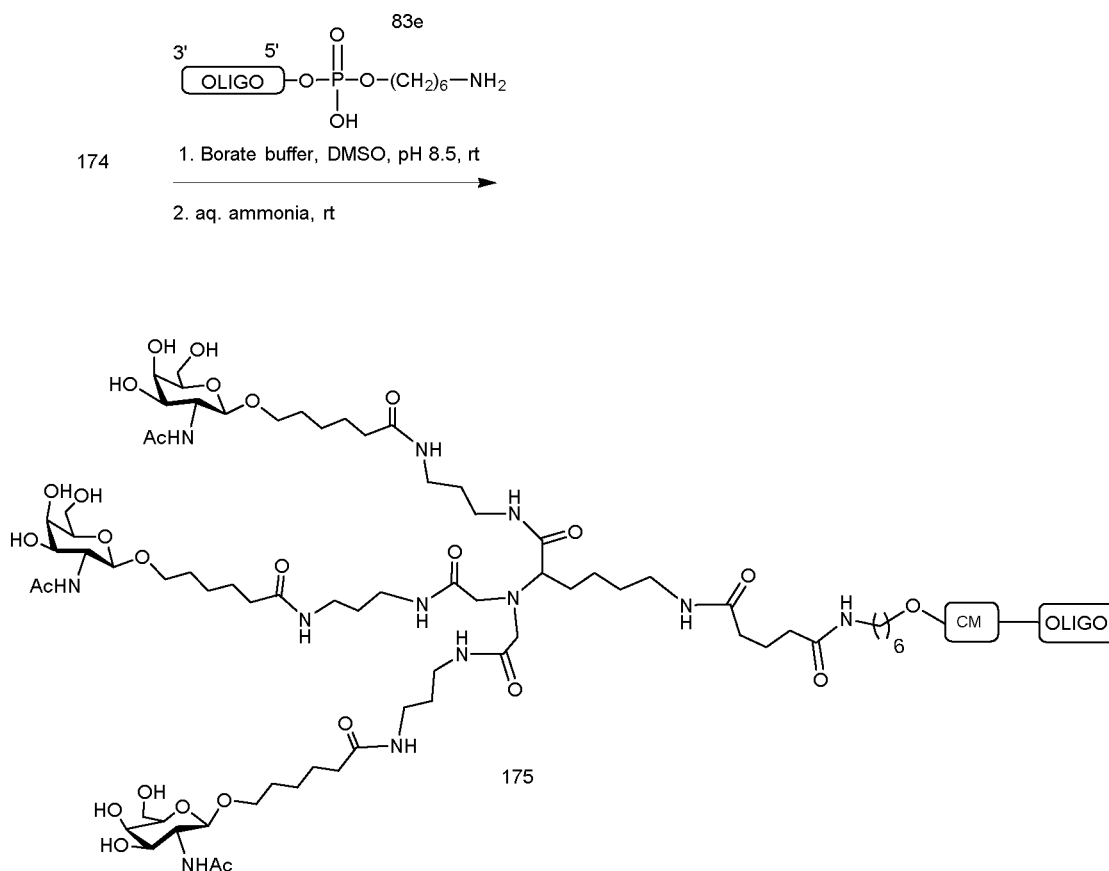
25 **Example 61: Preparation of oligomeric compound 175 comprising GalNAc₃-12**



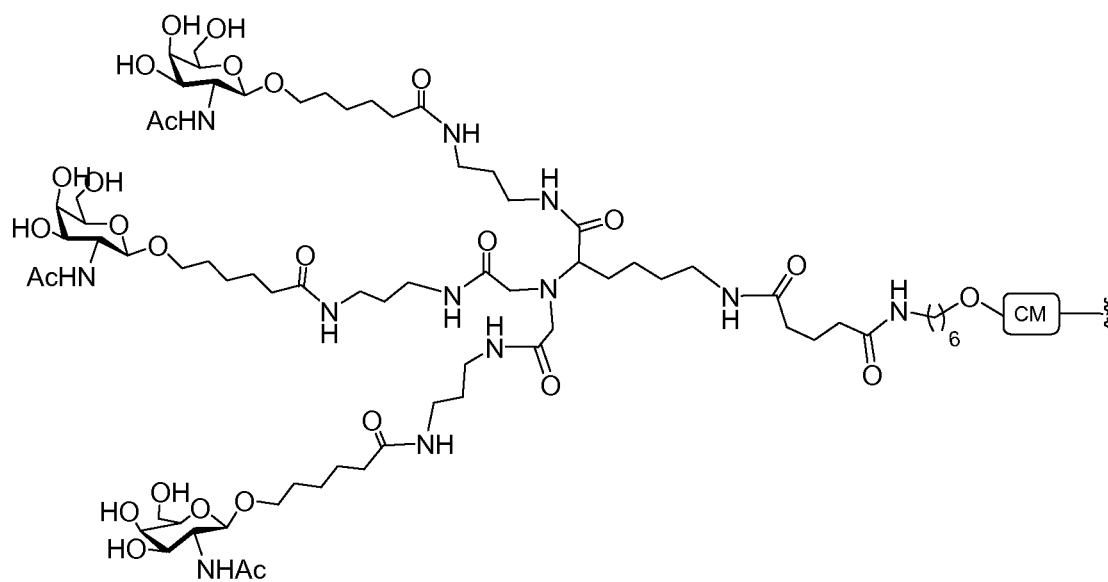


172

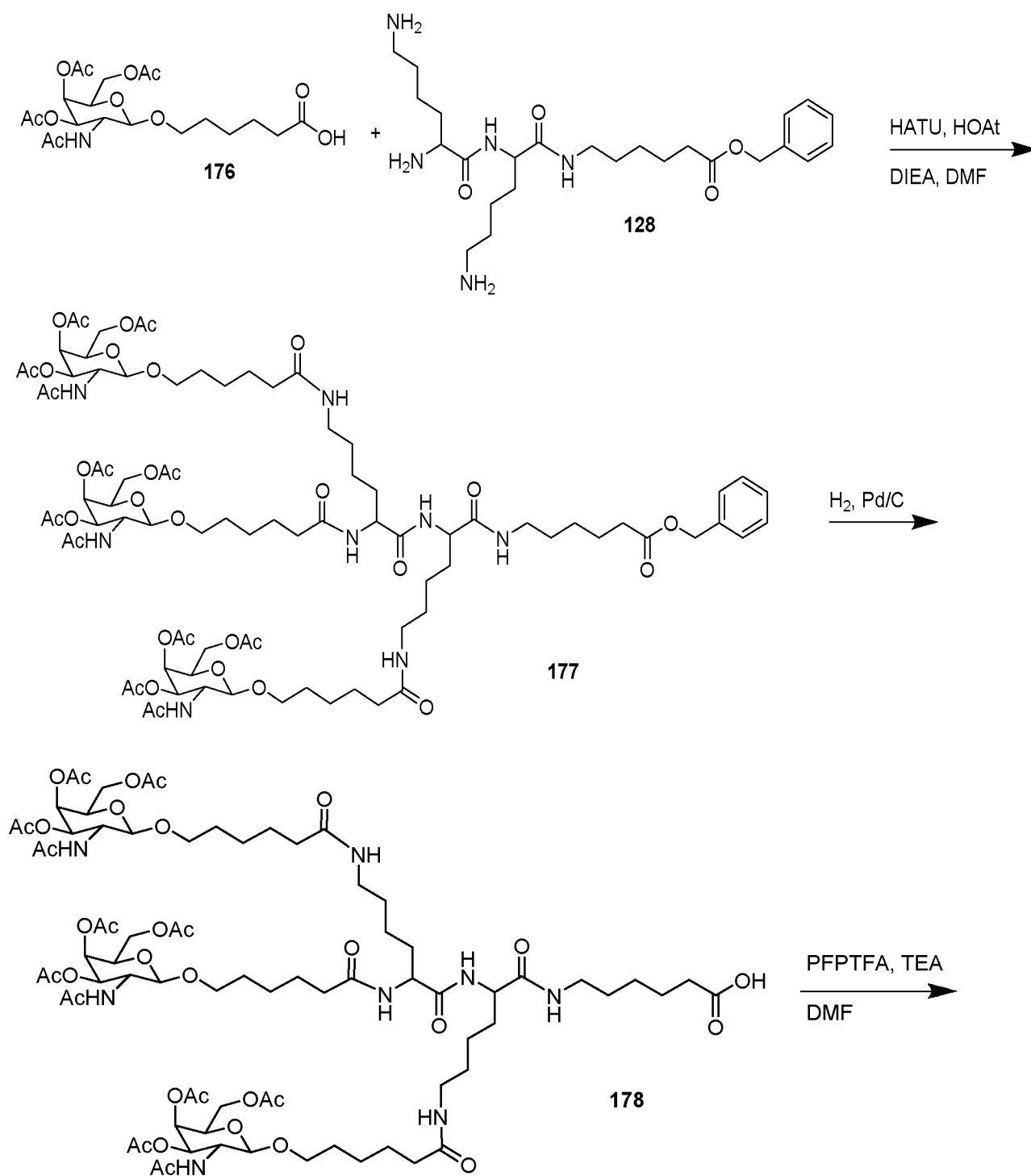


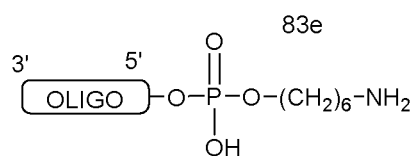
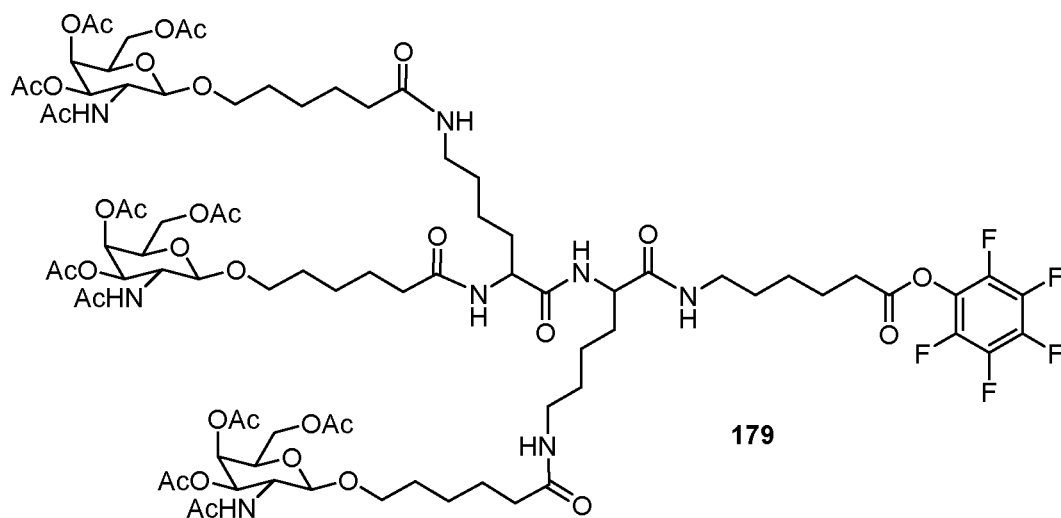


Compound 169 is commercially available. Compound 172 was prepared by addition of benzyl (perfluorophenyl) glutarate to compound 171. The benzyl (perfluorophenyl) glutarate was prepared by adding PFP-TFA and DIEA to 5-(benzyloxy)-5-oxopentanoic acid in DMF. Oligomeric compound 175, comprising a GalNAc₃-12 conjugate group, was prepared from compound 174 using the general procedures illustrated in Example 46. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-12 (GalNAc₃-12_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In a certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-. The structure of GalNAc₃-12 (GalNAc₃-12_a-CM-) is shown below:



Example 62: Preparation of oligomeric compound 180 comprising GalNAc₃-13

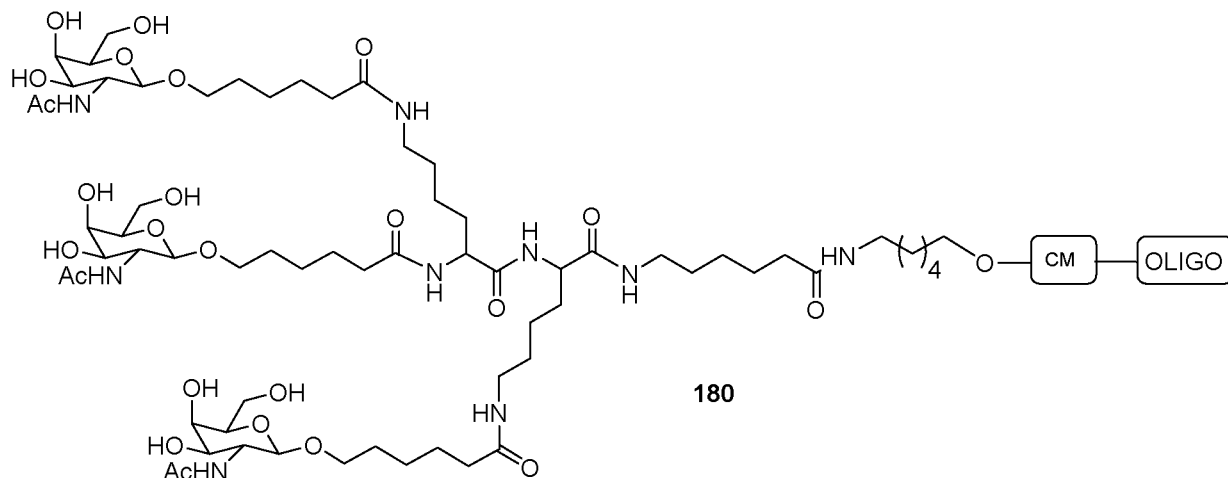




1. Borate buffer, DMSO, pH 8.5, rt



2. aq. ammonia, rt

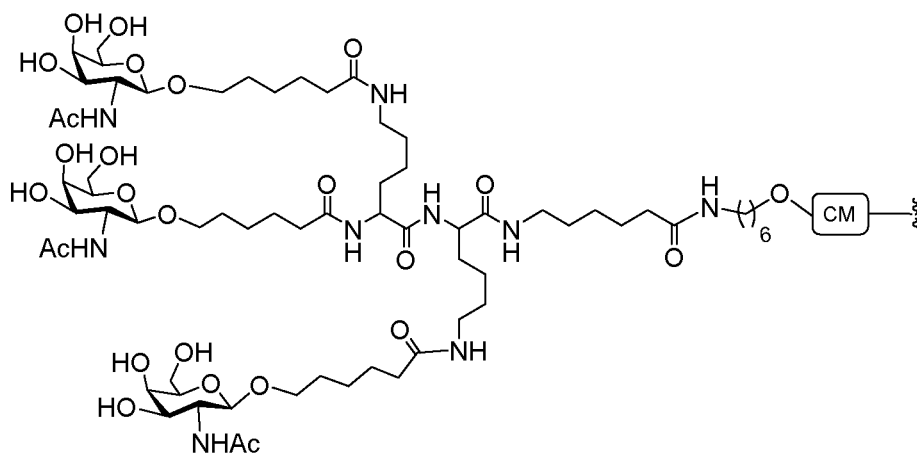


5

Compound 176 was prepared using the general procedure shown in Example 2. Oligomeric compound 180, comprising a GalNAc₃-13 conjugate group, was prepared from compound 177 using the general procedures illustrated in Example 49. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-13 (GalNAc₃-13_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In a

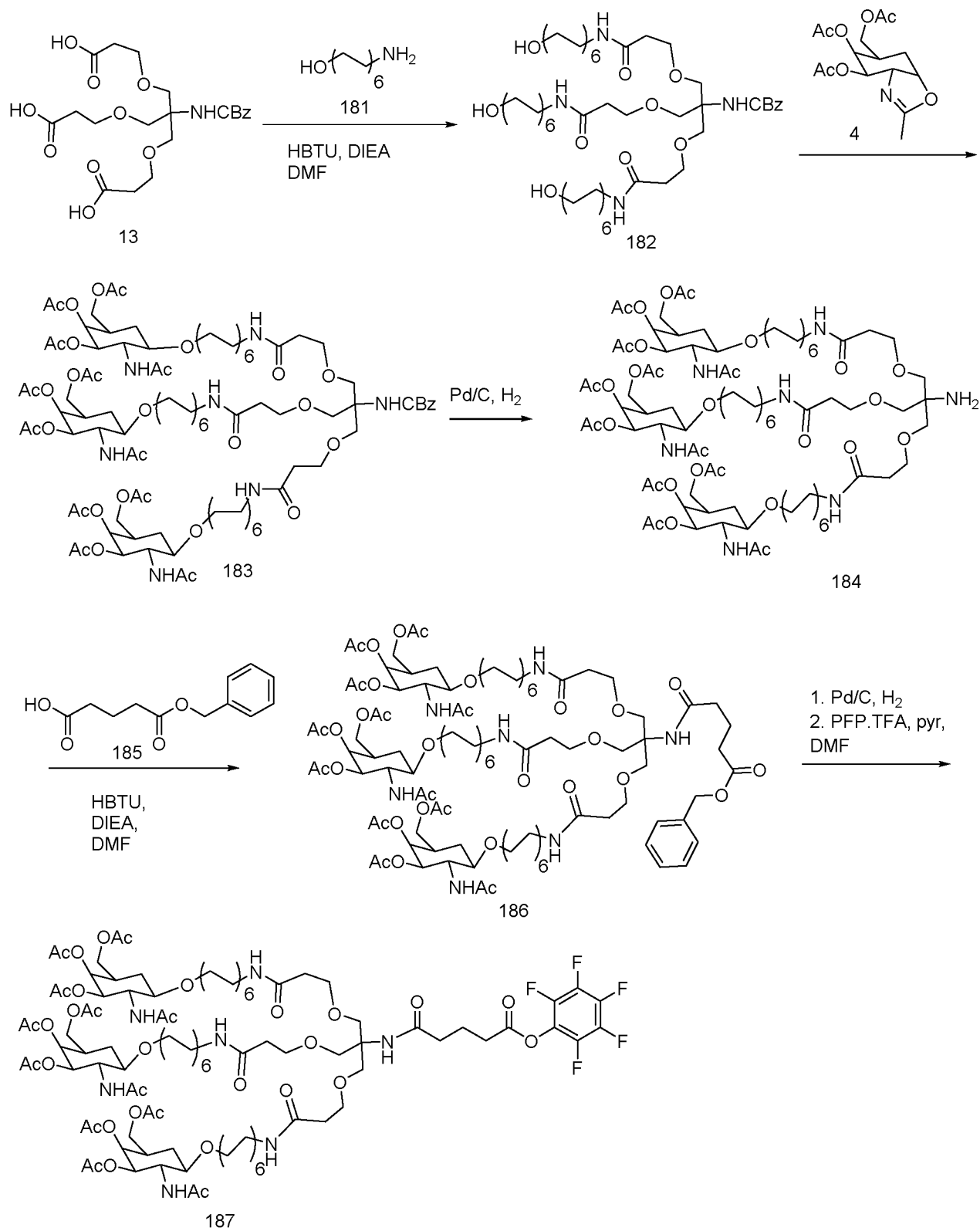
10

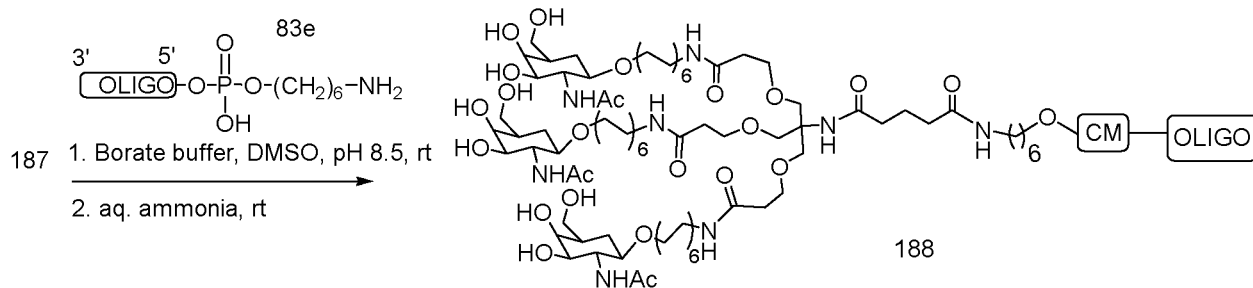
certain embodiments, the cleavable moiety is $\text{-P(=O)(OH)-A}_d\text{-P(=O)(OH)-}$. The structure of GalNAc₃-13 (GalNAc₃-13_a-CM-) is shown below:



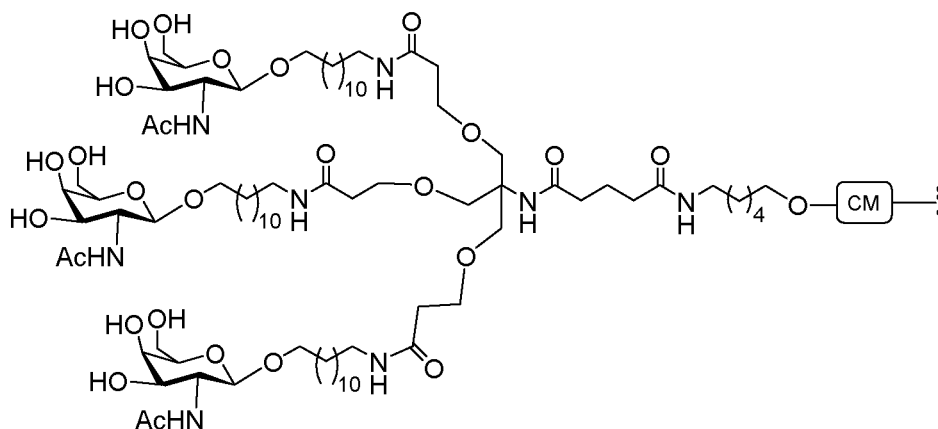
Example 63: Preparation of oligomeric compound 188 comprising GalNAc₃-14

5

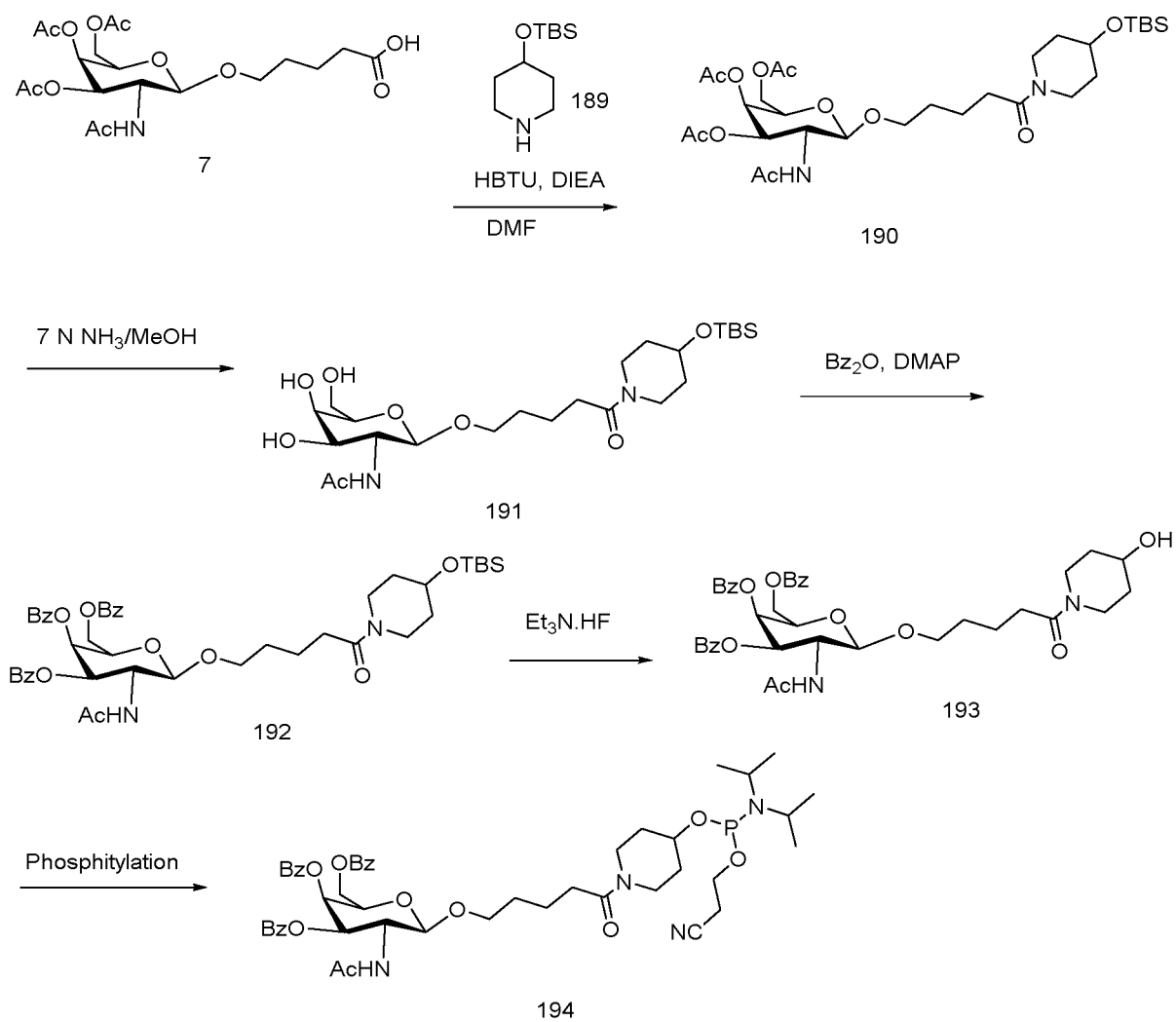


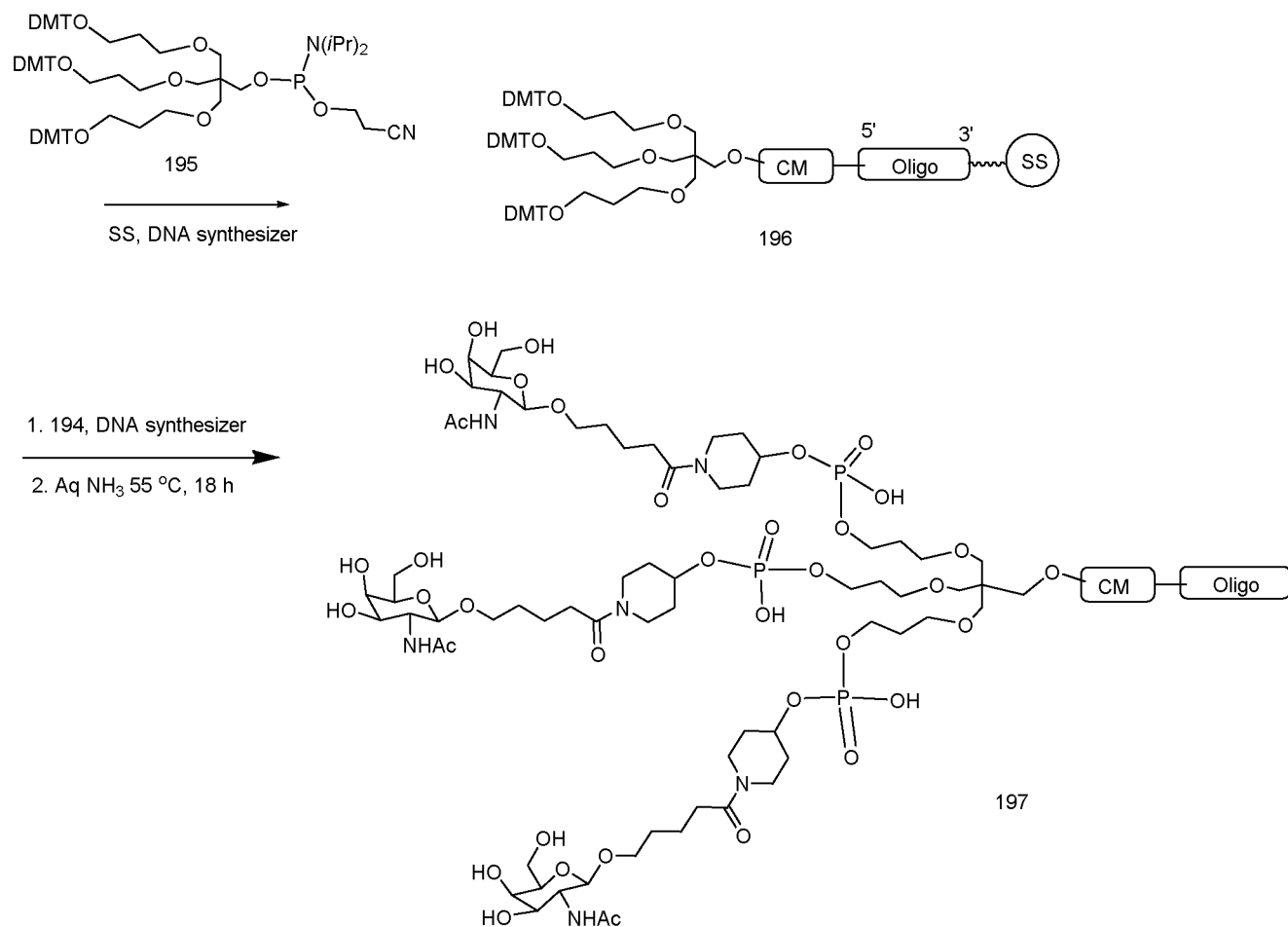


Compounds 181 and 185 are commercially available. Oligomeric compound 188, comprising a GalNAc₃-14 conjugate group, was prepared from compound 187 using the general procedures illustrated in Example 46. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-14 (GalNAc₃-14_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-. The structure of GalNAc₃-14 (GalNAc₃-14_a-CM-) is shown below:

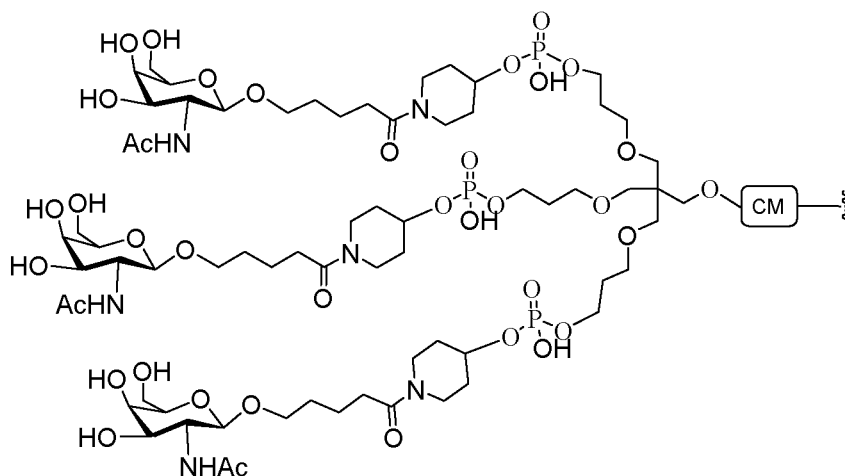


Example 64: Preparation of oligomeric compound 197 comprising GalNAc₃-15





Compound 189 is commercially available. Compound 195 was prepared using the general procedure shown in Example 31. Oligomeric compound 197, comprising a GalNAc₃-15 conjugate group, was prepared from compounds 194 and 195 using standard oligonucleotide synthesis procedures. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-15 (GalNAc₃-15_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-. The structure of GalNAc₃-15 (GalNAc₃-15_a-CM-) is shown below:



Example 65: Dose-dependent study of oligonucleotides comprising a 5'-conjugate group (comparison of GalNAc₃-3, 12, 13, 14, and 15) targeting SRB-1 *in vivo*

5 The oligonucleotides listed below were tested in a dose-dependent study for antisense inhibition of SRB-1 in mice. Unconjugated ISIS 353382 was included as a standard. Each of the GalNAc₃ conjugate groups was attached at the 5' terminus of the respective oligonucleotide by a phosphodiester linked 2'-deoxyadenosine nucleoside (cleavable moiety).

Table 54

Modified ASOs targeting SRB-1

ISIS No.	Sequences (5' to 3')	Conjugate	SEQ ID No.
353382	G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	none	143
661161	GalNAc₃-3_{a-o} ·A ^m _{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	GalNAc ₃ -3	145
671144	GalNAc₃-12_{a-o} ·A ^m _{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	GalNAc ₃ -12	145
670061	GalNAc₃-13_{a-o} ·A ^m _{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	GalNAc ₃ -13	145
671261	GalNAc₃-14_{a-o} ·A ^m _{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	GalNAc ₃ -14	145
671262	GalNAc₃-15_{a-o} ·A ^m _{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	GalNAc ₃ -15	145

Capital letters indicate the nucleobase for each nucleoside and ^mC indicates a 5-methyl cytosine. Subscripts: “e” indicates a 2'-MOE modified nucleoside; “d” indicates a β-D-2'-deoxyribonucleoside; “s” indicates a phosphorothioate internucleoside linkage (PS); “o” indicates a phosphodiester internucleoside linkage (PO); and “o” indicates -O-P(=O)(OH)-. Conjugate groups are in bold.

The structure of GalNAc₃-3_a was shown previously in Example 39. The structure of GalNAc₃-12_a was shown previously in Example 61. The structure of GalNAc₃-13_a was shown previously in Example 62. The structure of GalNAc₃-14_a was shown previously in Example 63. The structure of GalNAc₃-15_a was shown previously in Example 64.

5

Treatment

Six to eight week old C57bl6 mice (Jackson Laboratory, Bar Harbor, ME) were injected subcutaneously once or twice at the dosage shown below with ISIS 353382, 661161, 671144, 670061, 671261, 671262, or with saline. Mice that were dosed twice received the second dose three days after the first dose. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration to determine the liver SRB-1 mRNA levels using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. The results below are presented as the average percent of SRB-1 mRNA levels for each treatment group, normalized to the saline control.

As illustrated in Table 55, treatment with antisense oligonucleotides lowered SRB-1 mRNA levels in a dose-dependent manner. No significant differences in target knockdown were observed between animals that received a single dose and animals that received two doses (see ISIS 353382 dosages 30 and 2 x 15 mg/kg; and ISIS 661161 dosages 5 and 2 x 2.5 mg/kg). The antisense oligonucleotides comprising the phosphodiester linked GalNAc₃-3, 12, 13, 14, and 15 conjugates showed substantial improvement in potency compared to the unconjugated antisense oligonucleotide (ISIS 335382).

Table 55
SRB-1 mRNA (% Saline)

ISIS No.	Dosage (mg/kg)	SRB-1 mRNA (% Saline)	ED ₅₀ (mg/kg)	Conjugate
Saline	n/a	100.0	n/a	n/a
353382	3	85.0	22.4	none
	10	69.2		
	30	34.2		
	2 x 15	36.0		
661161	0.5	87.4	2.2	GalNAc ₃ -3
	1.5	59.0		
	5	25.6		
	2 x 2.5	27.5		
	15	17.4		
671144	0.5	101.2	3.4	GalNAc ₃ -12
	1.5	76.1		
	5	32.0		
	15	17.6		
670061	0.5	94.8	2.1	GalNAc ₃ -13
	1.5	57.8		
	5	20.7		

	15	13.3		
671261	0.5	110.7	4.1	GalNAc ₃ -14
	1.5	81.9		
	5	39.8		
	15	14.1		
671262	0.5	109.4	9.8	GalNAc ₃ -15
	1.5	99.5		
	5	69.2		
	15	36.1		

Liver transaminase levels, alanine aminotransferase (ALT) and aspartate aminotransferase (AST), in serum were measured relative to saline injected mice using standard protocols. Total bilirubin and BUN were also evaluated. The changes in body weights were evaluated with no significant differences from the saline group (data not shown). ALTs, ASTs, total bilirubin and BUN values are shown in Table 56 below.

Table 56

ISIS No.	Dosage (mg/kg)	ALT (U/L)	AST (U/L)	Total Bilirubin (mg/dL)	BUN (mg/dL)	Conjugate
Saline	n/a	28	60	0.1	39	n/a
353382	3	30	77	0.2	36	none
	10	25	78	0.2	36	
	30	28	62	0.2	35	
	2 x 15	22	59	0.2	33	
661161	0.5	39	72	0.2	34	GalNAc ₃ -3
	1.5	26	50	0.2	33	
	5	41	80	0.2	32	
	2 x 2.5	24	72	0.2	28	
	15	32	69	0.2	36	
671144	0.5	25	39	0.2	34	GalNAc ₃ -12
	1.5	26	55	0.2	28	
	5	48	82	0.2	34	
	15	23	46	0.2	32	
670061	0.5	27	53	0.2	33	GalNAc ₃ -13
	1.5	24	45	0.2	35	
	5	23	58	0.1	34	
	15	24	72	0.1	31	
671261	0.5	69	99	0.1	33	GalNAc ₃ -14
	1.5	34	62	0.1	33	
	5	43	73	0.1	32	
	15	32	53	0.2	30	
671262	0.5	24	51	0.2	29	GalNAc ₃ -15
	1.5	32	62	0.1	31	
	5	30	76	0.2	32	
	15	31	64	0.1	32	

Example 66: Effect of various cleavable moieties on antisense inhibition *in vivo* by oligonucleotides targeting SRB-1 comprising a 5'-GalNAc₃ cluster

The oligonucleotides listed below were tested in a dose-dependent study for antisense inhibition of SRB-1 in mice. Each of the GalNAc₃ conjugate groups was attached at the 5' terminus of the respective oligonucleotide by a phosphodiester linked nucleoside (cleavable moiety (CM)).

Table 57

Modified ASOs targeting SRB-1

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
661161	GalNAc₃-3_a-o · A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A _{ds} G _{ds} T ^m _{ds} C ^m _{ds} A _{ds} T _{ds} G _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T _e	GalNAc ₃ -3a	A _d	145
670699	GalNAc₃-3_a-o · T_{do} G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{eo} A _{ds} G _{ds} T ^m _{ds} C ^m _{ds} A _{ds} T _{ds} G _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{eo} C ^m _{eo} C ^m _{es} T ^m _{es} T _e	GalNAc ₃ -3a	T _d	148
670700	GalNAc₃-3_a-o · A_{eo} G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{eo} A _{ds} G _{ds} T ^m _{ds} C ^m _{ds} A _{ds} T _{ds} G _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{eo} C ^m _{eo} C ^m _{es} T ^m _{es} T _e	GalNAc ₃ -3a	A _e	145
670701	GalNAc₃-3_a-o · T_{eo} G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{eo} A _{ds} G _{ds} T ^m _{ds} C ^m _{ds} A _{ds} T _{ds} G _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{eo} C ^m _{eo} C ^m _{es} T ^m _{es} T _e	GalNAc ₃ -3a	T _e	148
671165	GalNAc₃-13_a-o · A_{do} G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{eo} A _{ds} G _{ds} T ^m _{ds} C ^m _{ds} A _{ds} T _{ds} G _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{eo} C ^m _{eo} C ^m _{es} T ^m _{es} T _e	GalNAc ₃ -13a	A _d	145

Capital letters indicate the nucleobase for each nucleoside and ^mC indicates a 5-methyl cytosine. Subscripts: “e” indicates a 2'-MOE modified nucleoside; “d” indicates a β-D-2'-deoxyribonucleoside; “s” indicates a phosphorothioate internucleoside linkage (PS); “o” indicates a phosphodiester internucleoside linkage (PO); and “o” indicates -O-P(=O)(OH)-. Conjugate groups are in bold.

The structure of GalNAc₃-3_a was shown previously in Example 39. The structure of GalNAc₃-13a was shown previously in Example 62.

Treatment

Six to eight week old C57bl6 mice (Jackson Laboratory, Bar Harbor, ME) were injected subcutaneously once at the dosage shown below with ISIS 661161, 670699, 670700, 670701, 671165, or with saline. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration to determine the liver SRB-1 mRNA levels using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. The results below are presented as the average percent of SRB-1 mRNA levels for each treatment group, normalized to the saline control.

As illustrated in Table 58, treatment with antisense oligonucleotides lowered SRB-1 mRNA levels in a dose-dependent manner. The antisense oligonucleotides comprising various cleavable moieties all showed similar potencies.

Table 58

SRB-1 mRNA (% Saline)

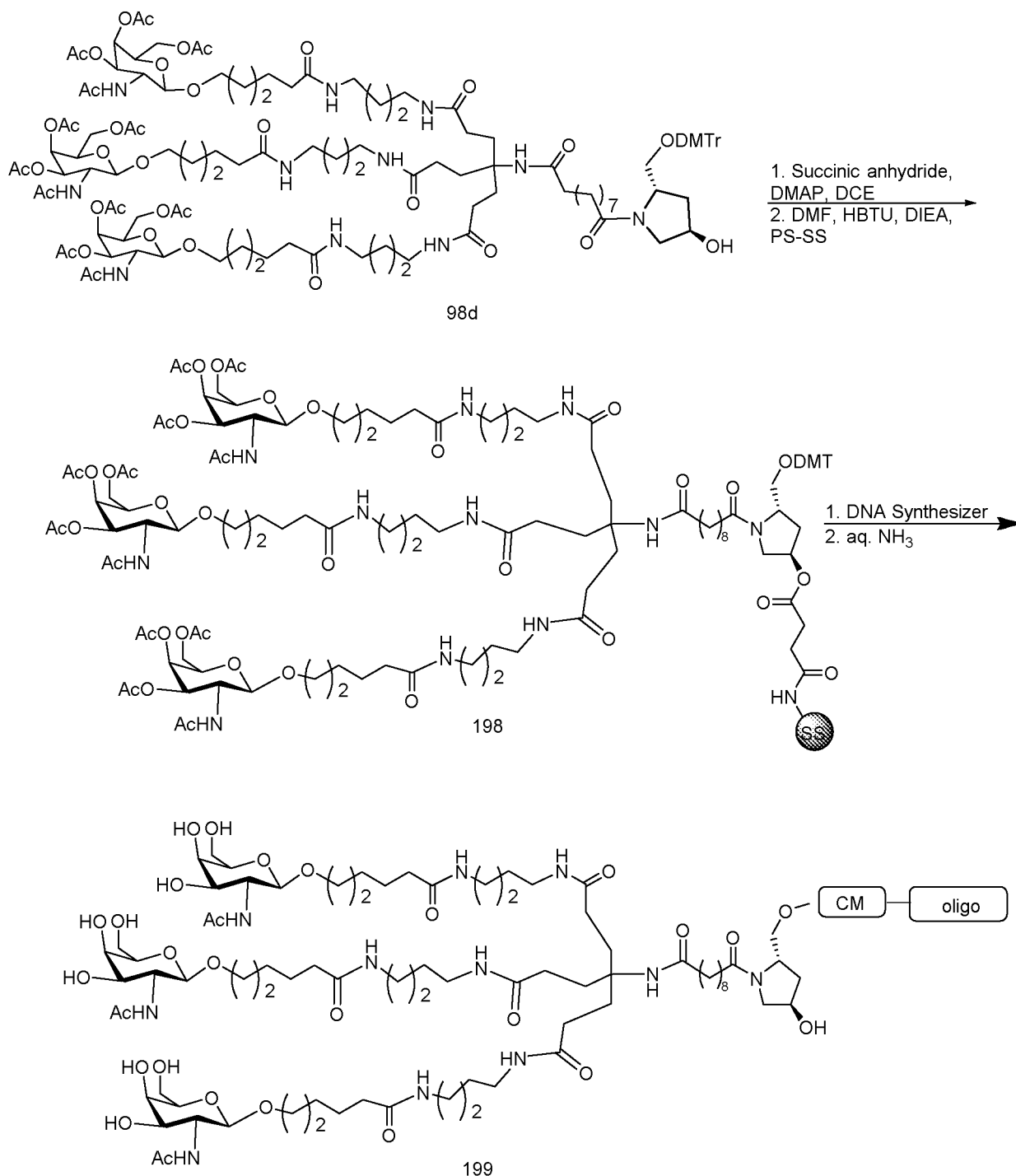
ISIS No.	Dosage (mg/kg)	SRB-1 mRNA (% Saline)	GalNAc ₃ Cluster	CM
Saline	n/a	100.0	n/a	n/a
661161	0.5	87.8	GalNAc ₃ -3a	A _d
	1.5	61.3		
	5	33.8		
	15	14.0		
670699	0.5	89.4	GalNAc ₃ -3a	T _d
	1.5	59.4		
	5	31.3		
	15	17.1		
670700	0.5	79.0	GalNAc ₃ -3a	A _e
	1.5	63.3		
	5	32.8		
	15	17.9		
670701	0.5	79.1	GalNAc ₃ -3a	T _e
	1.5	59.2		
	5	35.8		
	15	17.7		
671165	0.5	76.4	GalNAc ₃ -13a	A _d
	1.5	43.2		
	5	22.6		
	15	10.0		

Liver transaminase levels, alanine aminotransferase (ALT) and aspartate aminotransferase (AST), in serum were measured relative to saline injected mice using standard protocols. Total bilirubin and BUN were also evaluated. The changes in body weights were evaluated with no significant differences from the saline group (data not shown). ALTs, ASTs, total bilirubin and BUN values are shown in Table 56 below.

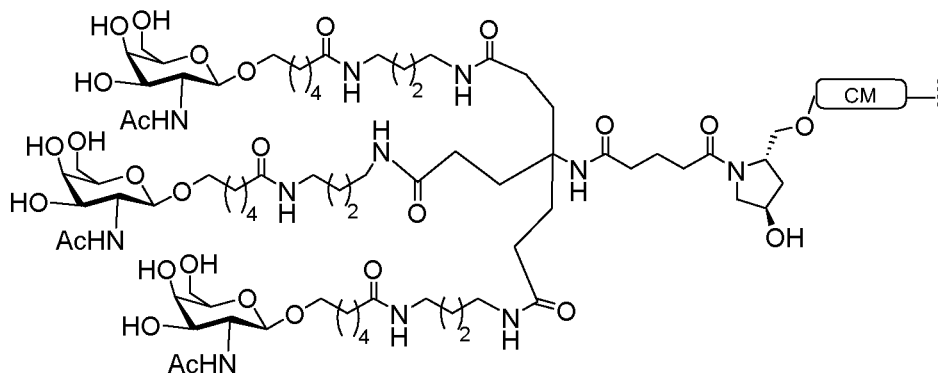
Table 59

ISIS No.	Dosage (mg/kg)	ALT (U/L)	AST (U/L)	Total Bilirubin (mg/dL)	BUN (mg/dL)	GalNAc ₃ Cluster	CM
Saline	n/a	24	64	0.2	31	n/a	n/a
661161	0.5	25	64	0.2	31	GalNAc ₃ -3a	A _d
	1.5	24	50	0.2	32		
	5	26	55	0.2	28		
	15	27	52	0.2	31		
670699	0.5	42	83	0.2	31	GalNAc ₃ -3a	T _d
	1.5	33	58	0.2	32		
	5	26	70	0.2	29		
	15	25	67	0.2	29		
670700	0.5	40	74	0.2	27	GalNAc ₃ -3a	A _e
	1.5	23	62	0.2	27		
	5	24	49	0.2	29		
	15	25	87	0.1	25		
670701	0.5	30	77	0.2	27	GalNAc ₃ -3a	T _e
	1.5	22	55	0.2	30		

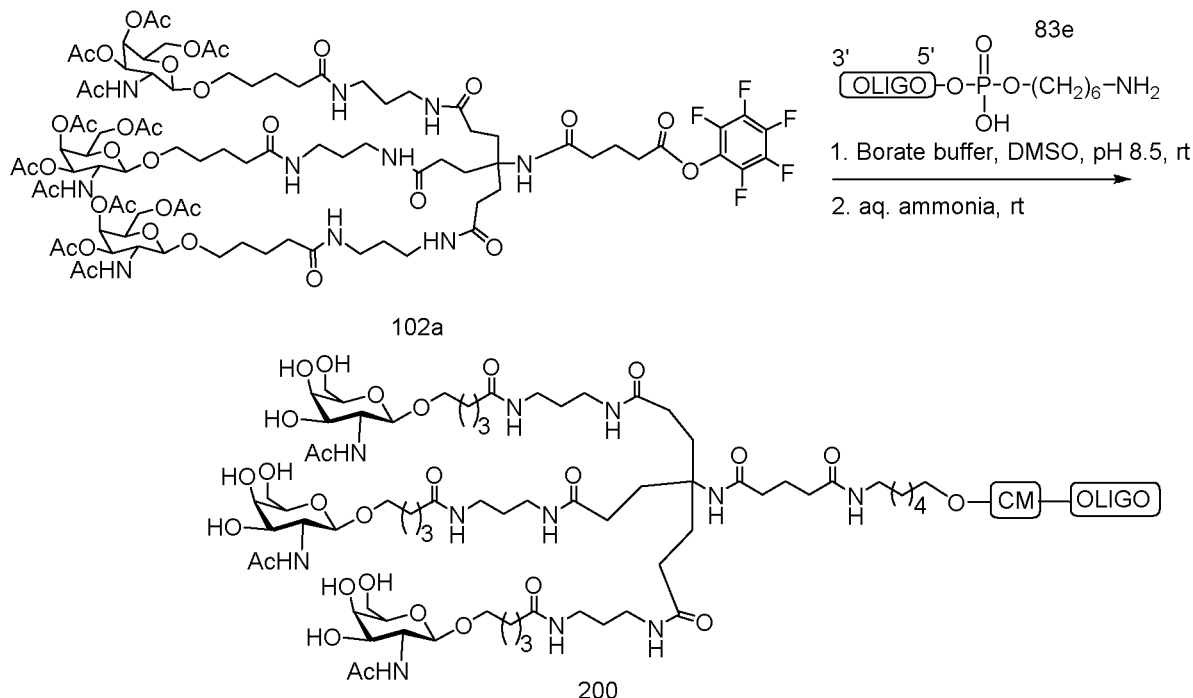
	5	81	101	0.2	25		
	15	31	82	0.2	24		
671165	0.5	44	84	0.2	26	GalNAc ₃ -13a	A _d
	1.5	47	71	0.1	24		
	5	33	91	0.2	26		
	15	33	56	0.2	29		

Example 67: Preparation of oligomeric compound 199 comprising GalNAc₃-16

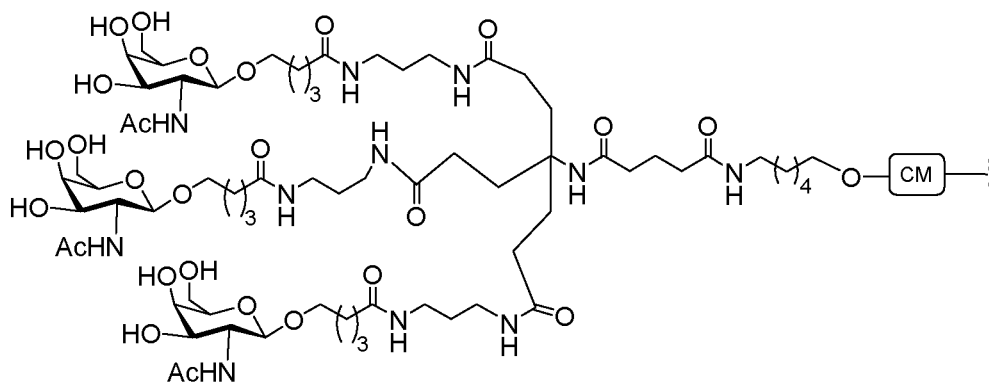
Oligomeric compound 199, comprising a GalNAc₃-16 conjugate group, is prepared using the general procedures illustrated in Examples 7 and 9. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-16 (GalNAc₃-16_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-. The structure of GalNAc₃-16 (GalNAc₃-16_a-CM-) is shown below:



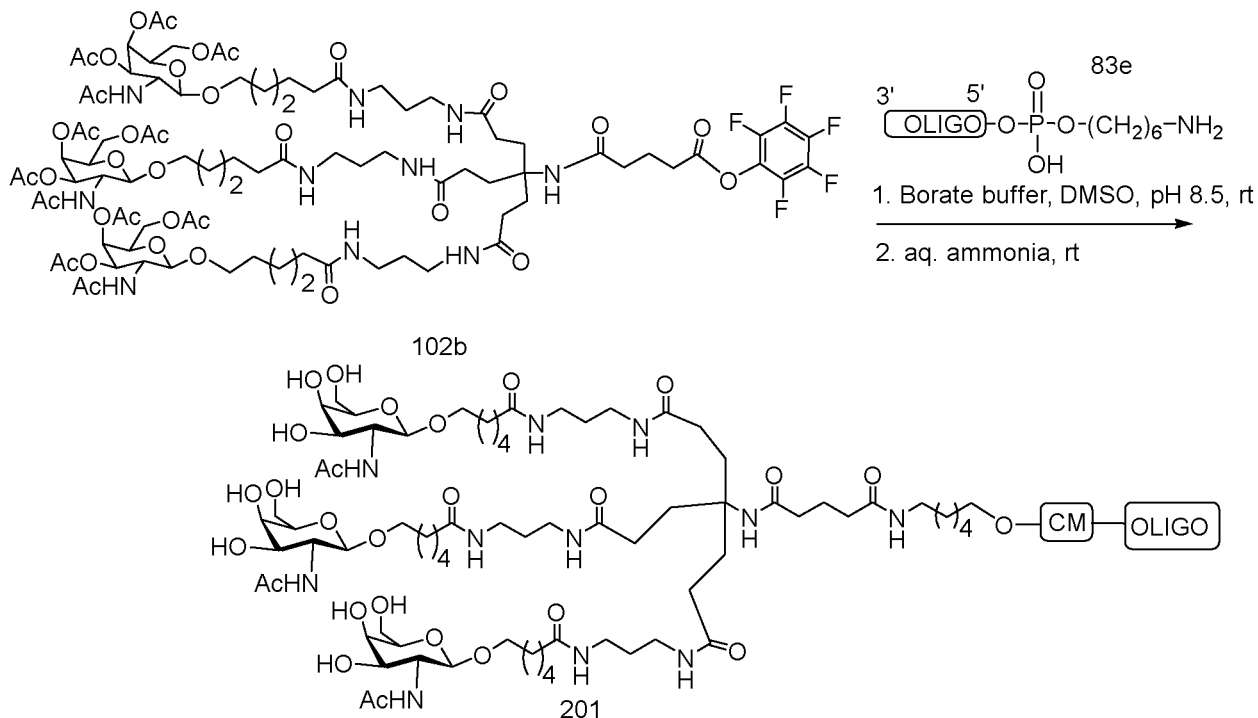
Example 68: Preparation of oligomeric compound 200 comprising GalNAc₃-17



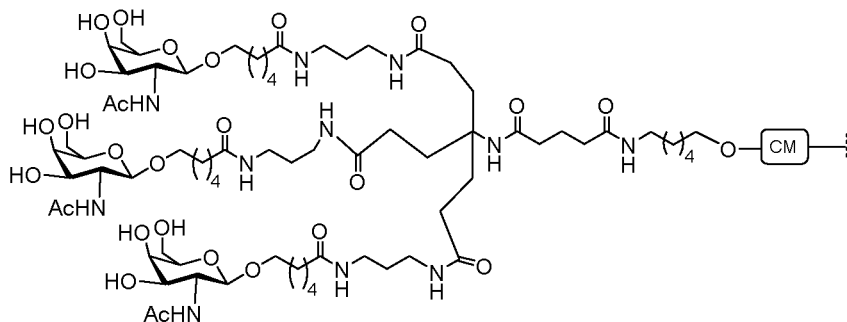
Oligomeric compound 200, comprising a GalNAc₃-17 conjugate group, was prepared using the general procedures illustrated in Example 46. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-17 (GalNAc₃-17_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A₄-P(=O)(OH)-. The structure of GalNAc₃-17 (GalNAc₃-17_a-CM-) is shown below:

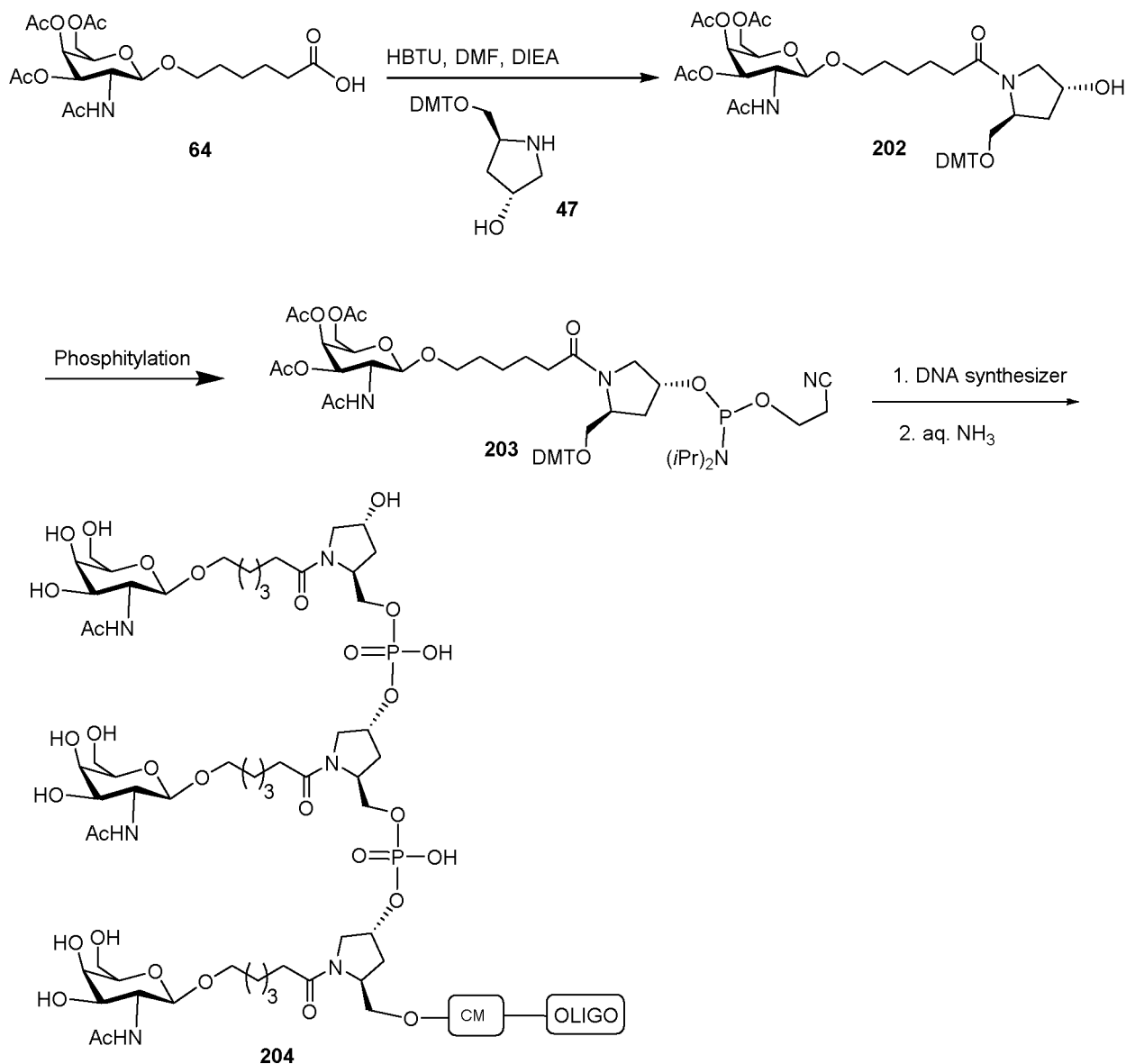


Example 69: Preparation of oligomeric compound 201 comprising GalNAc₃-18

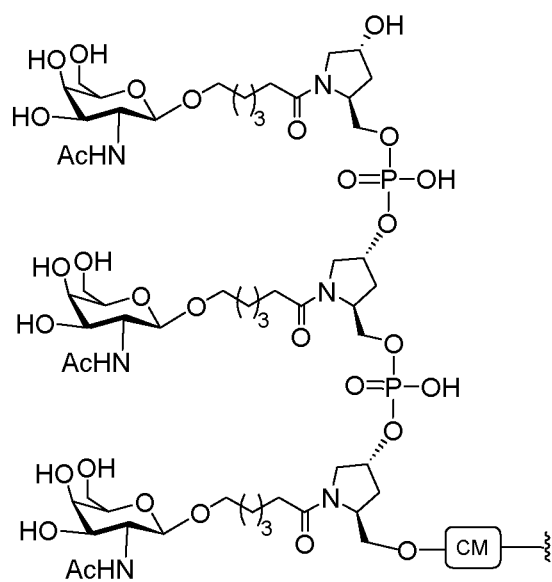


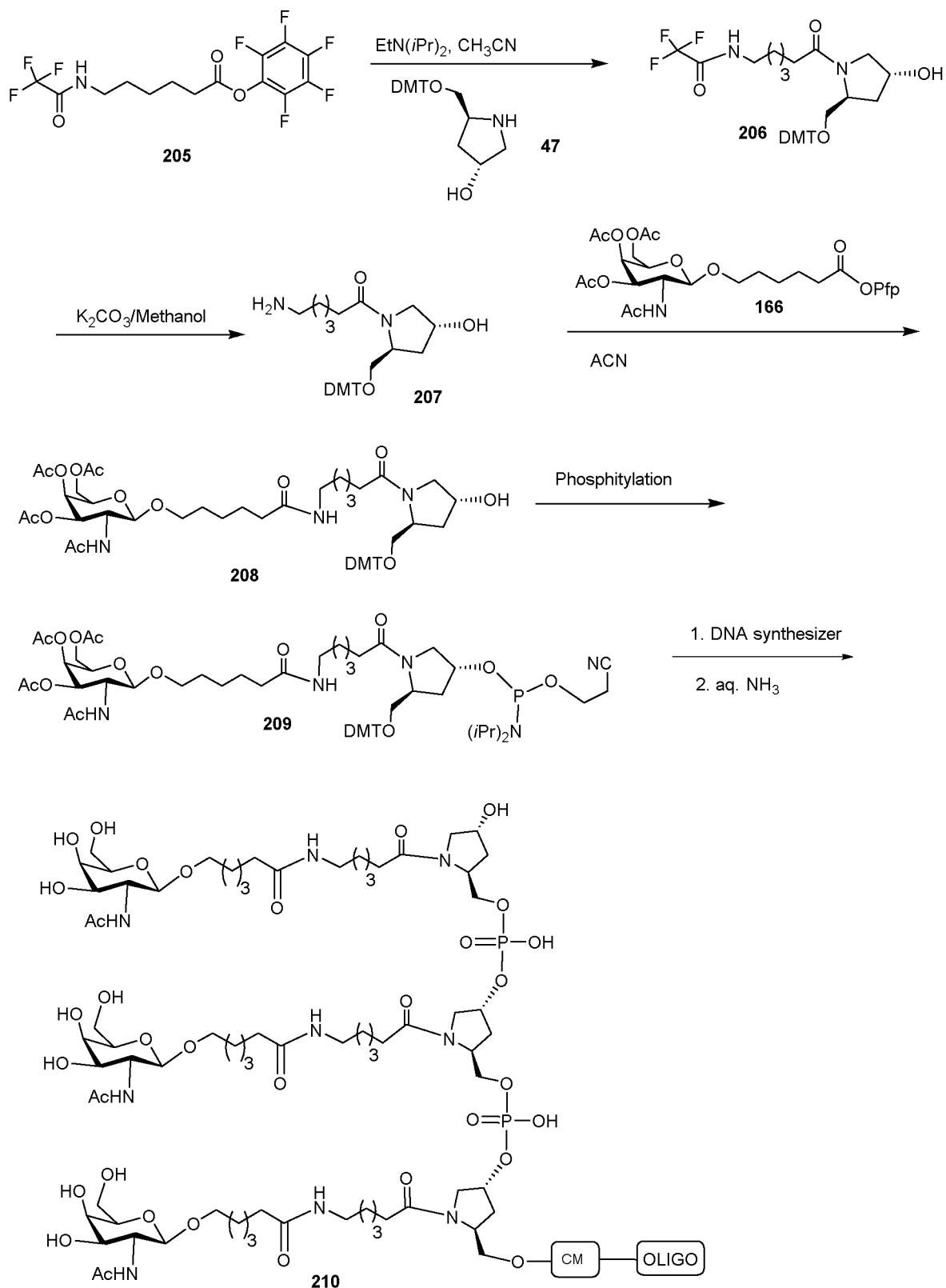
Oligomeric compound 201, comprising a GalNAc₃-18 conjugate group, was prepared using the general procedures illustrated in Example 46. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-18 (GalNAc₃-18_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-. The structure of GalNAc₃-18 (GalNAc₃-18_a-CM-) is shown below:



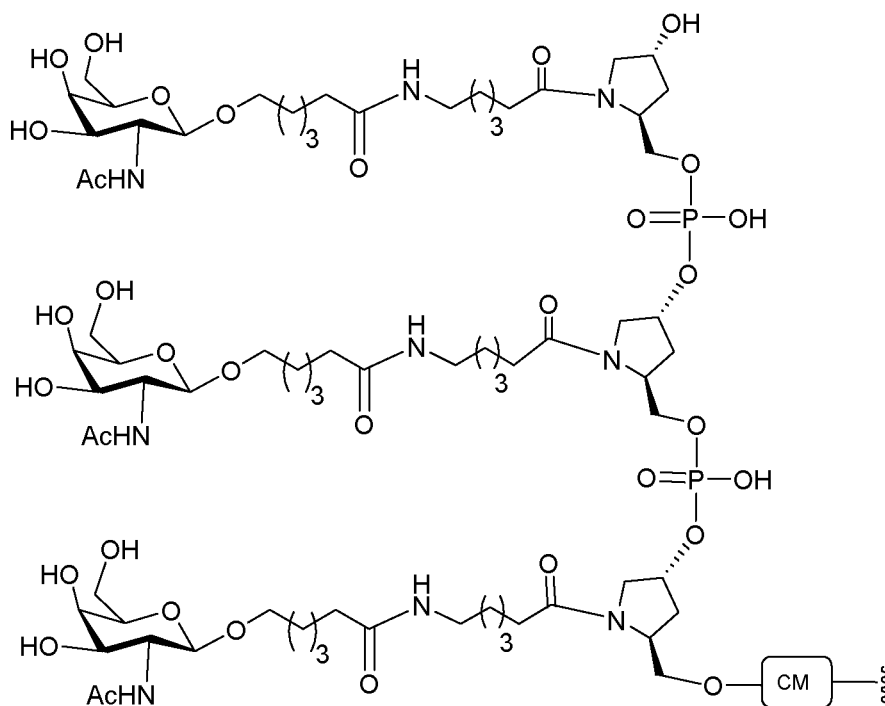
Example 70: Preparation of oligomeric compound 204 comprising GalNAc₃-19

Oligomeric compound 204, comprising a GalNAc₃-19 conjugate group, was prepared from compound 64 using the general procedures illustrated in Example 52. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-19 (GalNAc₃-19_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A₄-P(=O)(OH)-. The structure of GalNAc₃-19 (GalNAc₃-19_a-CM-) is shown below:

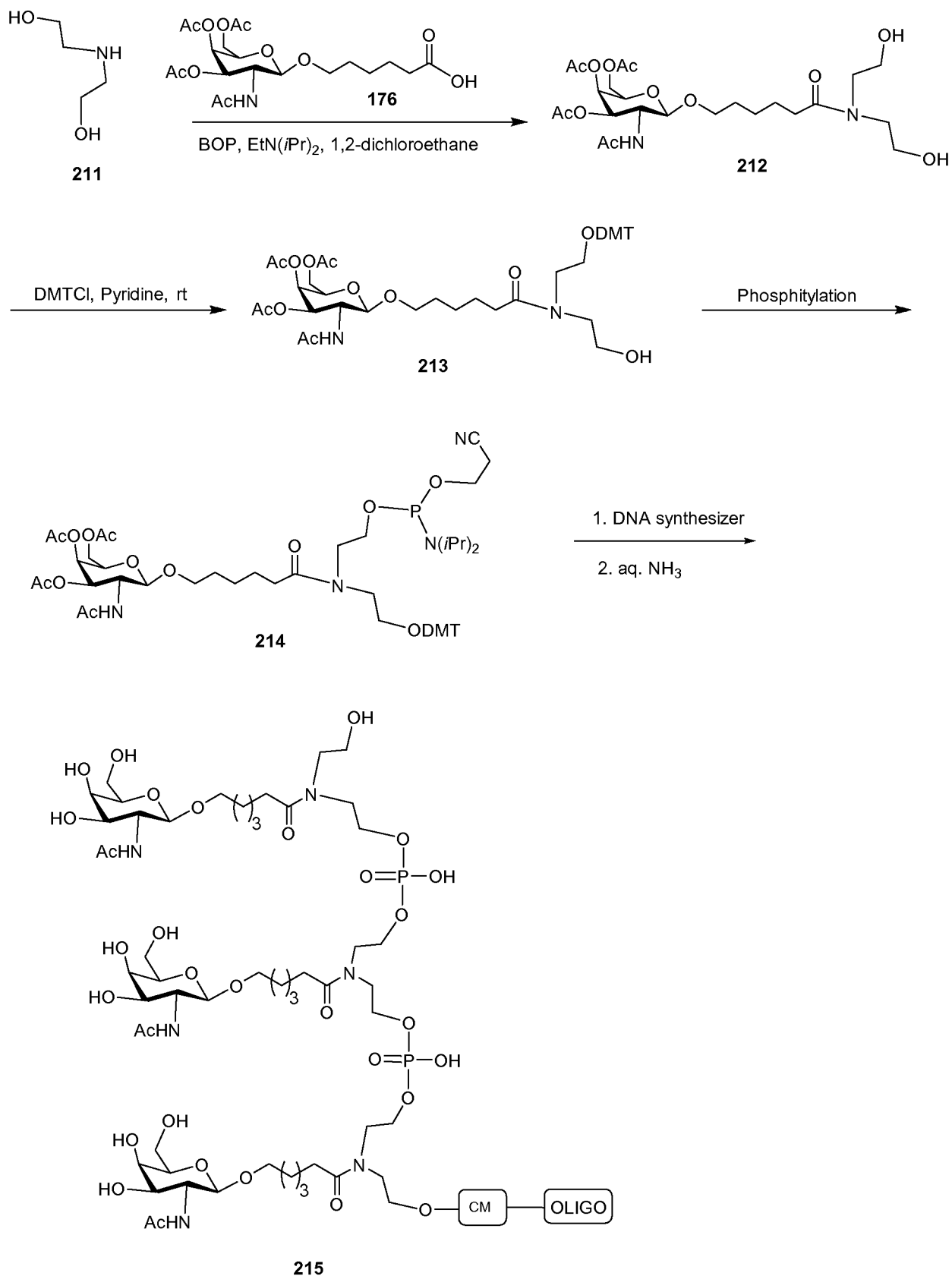


Example 71: Preparation of oligomeric compound 210 comprising GalNAc₃-20

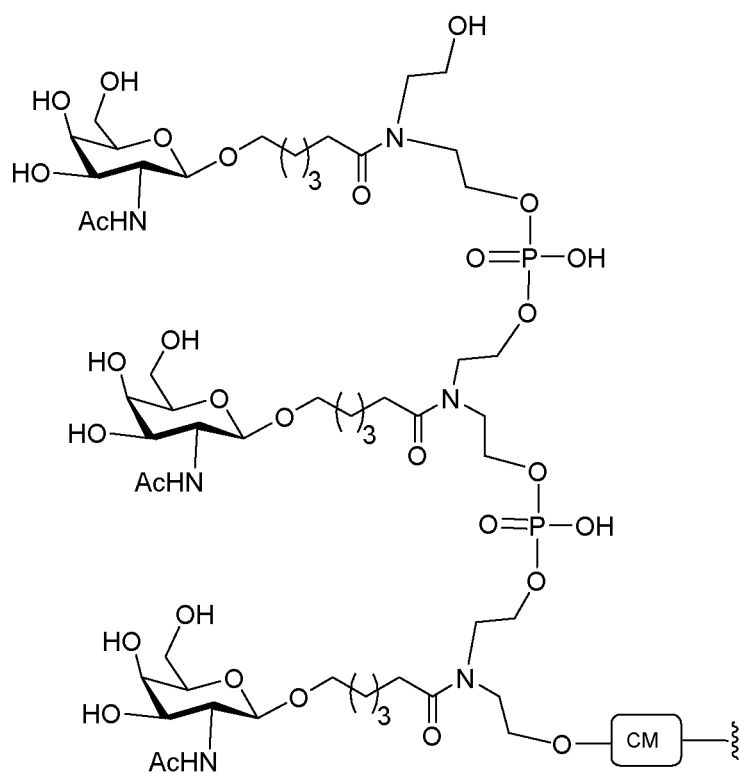
- Compound 205 was prepared by adding PFP-TFA and DIEA to 6-(2,2,2-trifluoroacetamido)hexanoic acid in acetonitrile, which was prepared by adding triflic anhydride to 6-aminohexanoic acid. The reaction mixture was heated to 80 °C, then lowered to rt. Oligomeric compound 210, comprising a GalNAc₃-20 conjugate group, was prepared from compound 208 using the general procedures illustrated in Example 52. The
- 5 GalNAc₃ cluster portion of the conjugate group GalNAc₃-20 (GalNAc₃-20_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-. The structure of GalNAc₃-20 (GalNAc₃-20_a-CM-) is shown below:

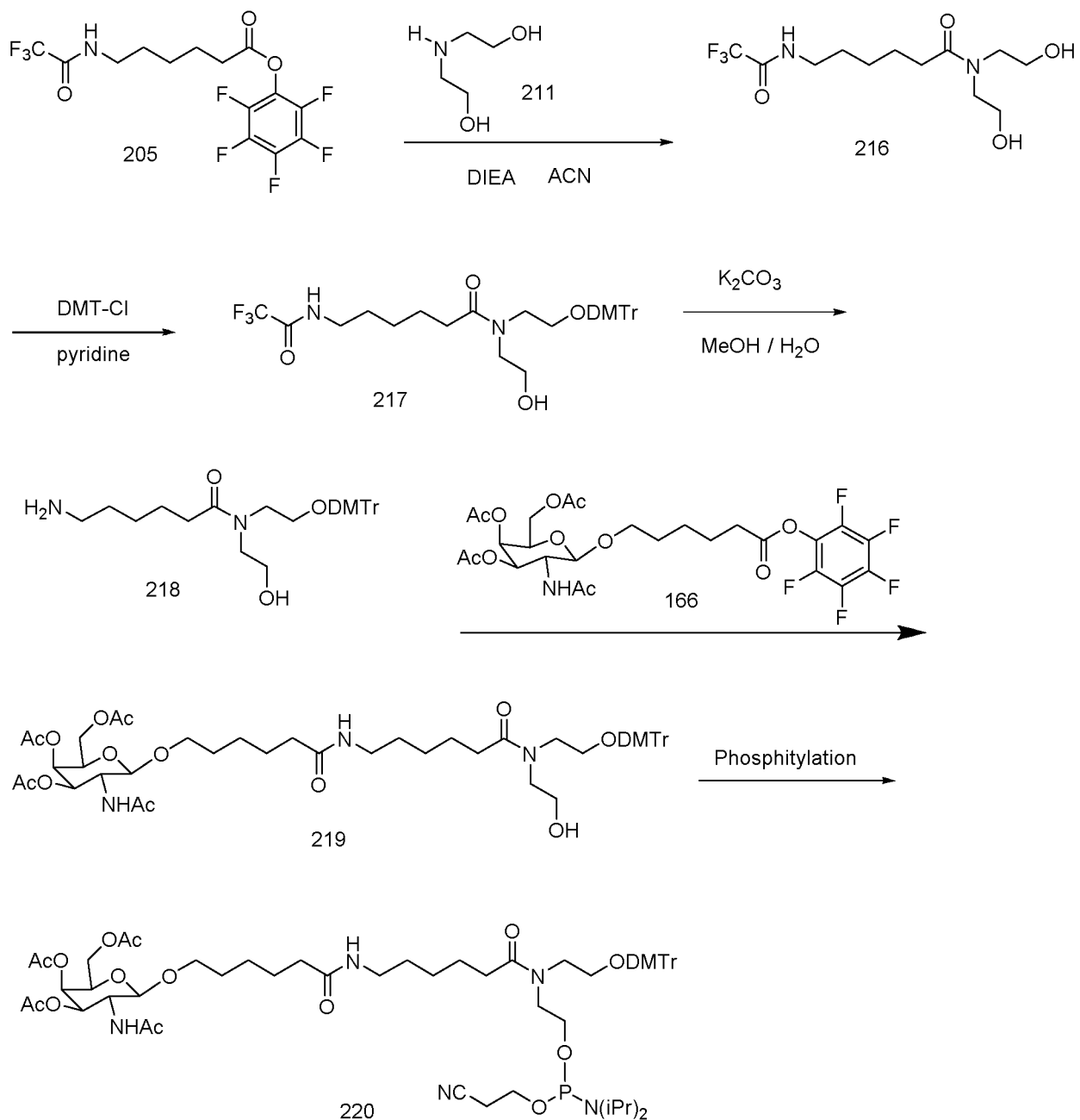


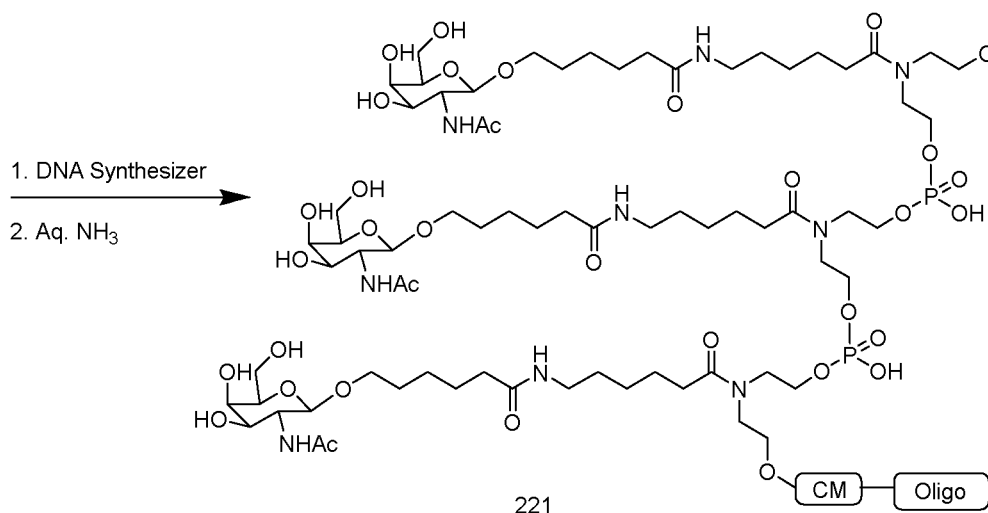
Example 72: Preparation of oligomeric compound 215 comprising GalNAc₃-21



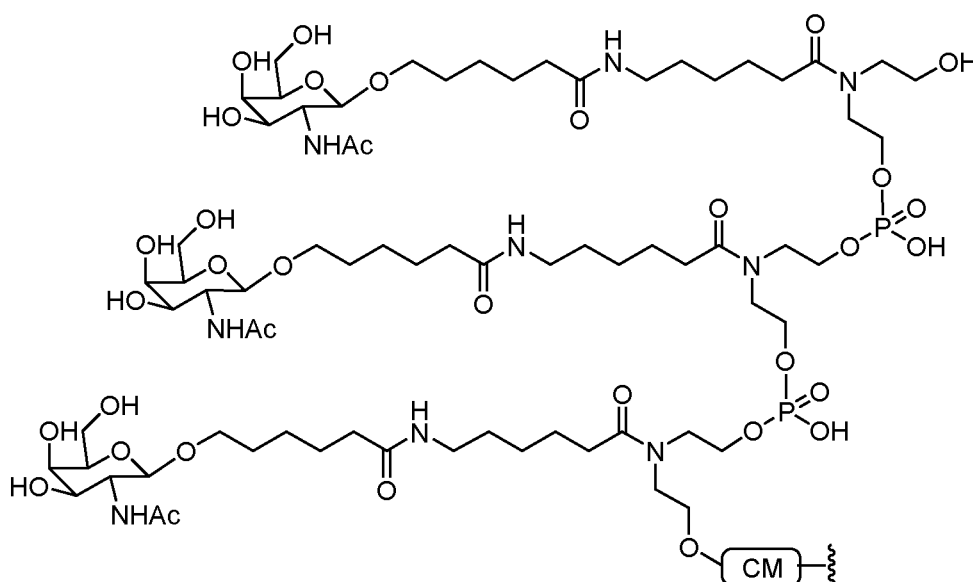
Compound 211 is commercially available. Oligomeric compound 215, comprising a GalNAc₃-21 conjugate group, was prepared from compound 213 using the general procedures illustrated in Example 52. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-21 (GalNAc₃-21_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-. The structure of GalNAc₃-21 (GalNAc₃-21_a-CM-) is shown below:



Example 73: Preparation of oligomeric compound 221 comprising GalNAc₃-22



Compound 220 was prepared from compound 219 using diisopropylammonium tetrazolide. Oligomeric compound 221, comprising a GalNAc₃-21 conjugate group, is prepared from compound 220 using the general procedure illustrated in Example 52. The GalNAc₃ cluster portion of the conjugate group GalNAc₃-22 (GalNAc₃-22_a) can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the cleavable moiety is -P(=O)(OH)-A_d-P(=O)(OH)-. The structure of GalNAc₃-22 (GalNAc₃-22_a-CM-) is shown below:



Example 74: Effect of various cleavable moieties on antisense inhibition *in vivo* by oligonucleotides targeting SRB-1 comprising a 5'-GalNAc₃ conjugate

The oligonucleotides listed below were tested in a dose-dependent study for antisense inhibition of SRB-1 in mice. Each of the GalNAc₃ conjugate groups was attached at the 5' terminus of the respective oligonucleotide.

Table 60
Modified ASOs targeting SRB-1

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
353382	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} C _{es} ^m C _{es} T _{es} T _e	n/a	n/a	143
661161	GalNAc₃-3_a-o ·A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	GalNAc ₃ -3a	A _d	145
666904	GalNAc₃-3_a-o ·G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	GalNAc ₃ -3a	PO	143
675441	GalNAc₃-17_a-o ·A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	GalNAc ₃ -17a	A _d	145
675442	GalNAc₃-18_a-o ·A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	GalNAc ₃ -18a	A _d	145

In all tables, capital letters indicate the nucleobase for each nucleoside and ^mC indicates a 5-methyl cytosine. Subscripts: “e” indicates a 2'-MOE modified nucleoside; “d” indicates a β-D-2'-deoxyribonucleoside; “s” indicates a phosphorothioate internucleoside linkage (PS); “o” indicates a phosphodiester internucleoside linkage (PO); and “o” indicates -O-P(=O)(OH)-. Conjugate groups are in bold.

The structure of GalNAc₃-3_a was shown previously in Example 39. The structure of GalNAc₃-17a was shown previously in Example 68, and the structure of GalNAc₃-18a was shown in Example 69.

10 Treatment

Six to eight week old C57BL/6 mice (Jackson Laboratory, Bar Harbor, ME) were injected subcutaneously once at the dosage shown below with an oligonucleotide listed in Table 60 or with saline. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration to determine the SRB-1 mRNA levels using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. The results below are presented as the average percent of SRB-1 mRNA levels for each treatment group, normalized to the saline control.

As illustrated in Table 61, treatment with antisense oligonucleotides lowered SRB-1 mRNA levels in a dose-dependent manner. The antisense oligonucleotides comprising a GalNAc conjugate showed similar potencies and were significantly more potent than the parent oligonucleotide lacking a GalNAc conjugate.

Table 61
SRB-1 mRNA (% Saline)

ISIS No.	Dosage (mg/kg)	SRB-1 mRNA (% Saline)	GalNAc ₃ Cluster	CM
Saline	n/a	100.0	n/a	n/a

353382	3	79.38	n/a	n/a
	10	68.67		
	30	40.70		
661161	0.5	79.18	GalNAc ₃ -3a	A _d
	1.5	75.96		
	5	30.53		
	15	12.52		
666904	0.5	91.30	GalNAc ₃ -3a	PO
	1.5	57.88		
	5	21.22		
	15	16.49		
675441	0.5	76.71	GalNAc ₃ -17a	A _d
	1.5	63.63		
	5	29.57		
	15	13.49		
675442	0.5	95.03	GalNAc ₃ -18a	A _d
	1.5	60.06		
	5	31.04		
	15	19.40		

Liver transaminase levels, alanine aminotransferase (ALT) and aspartate aminotransferase (AST), in serum were measured relative to saline injected mice using standard protocols. Total bilirubin and BUN were also evaluated. The change in body weights was evaluated with no significant change from the saline group (data not shown). ALTs, ASTs, total bilirubin and BUN values are shown in Table 62 below.

Table 62

ISIS No.	Dosage (mg/kg)	ALT (U/L)	AST (U/L)	Total Bilirubin (mg/dL)	BUN (mg/dL)	GalNAc ₃ Cluster	CM
Saline	n/a	26	59	0.16	42	n/a	n/a
353382	3	23	58	0.18	39	n/a	n/a
	10	28	58	0.16	43		
	30	20	48	0.12	34		
661161	0.5	30	47	0.13	35	GalNAc ₃ -3a	A _d
	1.5	23	53	0.14	37		
	5	26	48	0.15	39		
	15	32	57	0.15	42		
666904	0.5	24	73	0.13	36	GalNAc ₃ -3a	PO
	1.5	21	48	0.12	32		
	5	19	49	0.14	33		
	15	20	52	0.15	26		
675441	0.5	42	148	0.21	36	GalNAc ₃ -17a	A _d
	1.5	60	95	0.16	34		
	5	27	75	0.14	37		
	15	24	61	0.14	36		
675442	0.5	26	65	0.15	37	GalNAc ₃ -18a	A _d
	1.5	25	64	0.15	43		
	5	27	69	0.15	37		
	15	30	84	0.14	37		

Example 75: Pharmacokinetic analysis of oligonucleotides comprising a 5'-conjugate group

The PK of the ASOs in Tables 54, 57 and 60 above was evaluated using liver samples that were obtained following the treatment procedures described in Examples 65, 66, and 74. The liver samples were minced and extracted using standard protocols and analyzed by IP-HPLC-MS alongside an internal standard.

5 The combined tissue level ($\mu\text{g/g}$) of all metabolites was measured by integrating the appropriate UV peaks, and the tissue level of the full-length ASO missing the conjugate ("parent," which is Isis No. 353382 in this case) was measured using the appropriate extracted ion chromatograms (EIC).

Table 63**PK Analysis in Liver**

ISIS No.	Dosage (mg/kg)	Total Tissue Level by UV ($\mu\text{g/g}$)	Parent ASO Tissue Level by EIC ($\mu\text{g/g}$)	GalNAc ₃ Cluster	CM
353382	3	8.9	8.6	n/a	n/a
	10	22.4	21.0		
	30	54.2	44.2		
661161	5	32.4	20.7	GalNAc ₃ -3a	A _d
	15	63.2	44.1		
671144	5	20.5	19.2	GalNAc ₃ -12a	A _d
	15	48.6	41.5		
670061	5	31.6	28.0	GalNAc ₃ -13a	A _d
	15	67.6	55.5		
671261	5	19.8	16.8	GalNAc ₃ -14a	A _d
	15	64.7	49.1		
671262	5	18.5	7.4	GalNAc ₃ -15a	A _d
	15	52.3	24.2		
670699	5	16.4	10.4	GalNAc ₃ -3a	T _d
	15	31.5	22.5		
670700	5	19.3	10.9	GalNAc ₃ -3a	A _e
	15	38.1	20.0		
670701	5	21.8	8.8	GalNAc ₃ -3a	T _e
	15	35.2	16.1		
671165	5	27.1	26.5	GalNAc ₃ -13a	A _d
	15	48.3	44.3		
666904	5	30.8	24.0	GalNAc ₃ -3a	PO
	15	52.6	37.6		
675441	5	25.4	19.0	GalNAc ₃ -17a	A _d
	15	54.2	42.1		
675442	5	22.2	20.7	GalNAc ₃ -18a	A _d
	15	39.6	29.0		

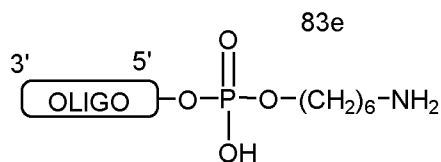
The results in Table 63 above show that there were greater liver tissue levels of the oligonucleotides comprising a GalNAc₃ conjugate group than of the parent oligonucleotide that does not comprise a GalNAc₃ conjugate group (ISIS 353382) 72 hours following oligonucleotide administration, particularly when taking

15 into consideration the differences in dosing between the oligonucleotides with and without a GalNAc₃ conjugate group. Furthermore, by 72 hours, 40-98% of each oligonucleotide comprising a GalNAc₃ conjugate

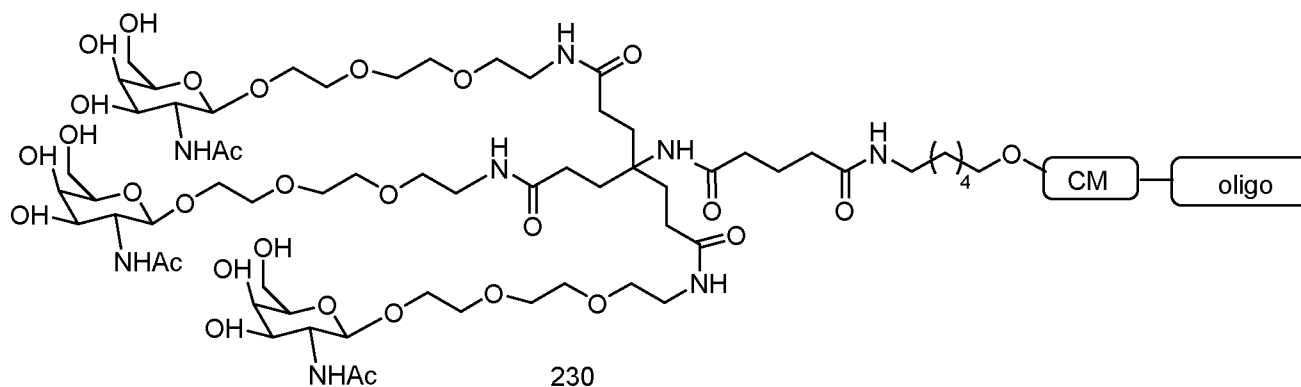
group was metabolized to the parent compound, indicating that the GalNAc₃ conjugate groups were cleaved from the oligonucleotides.

Example 76: Preparation of oligomeric compound 230 comprising GalNAc₃-23





1. Borate buffer, DMSO, pH 8.5, rt
 2. aq. ammonia, rt

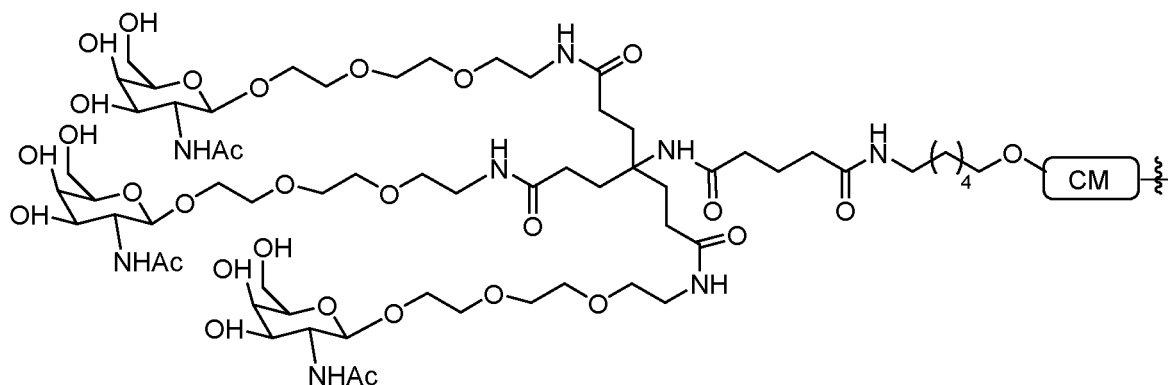


Compound 222 is commercially available. 44.48 ml (0.33 mol) of compound 222 was treated with
 5 tosyl chloride (25.39 g, 0.13 mol) in pyridine (500mL) for 16 hours. The reaction was then evaporated to an
 oil, dissolved in EtOAc and washed with water, sat. NaHCO₃, brine, and dried over Na₂SO₄. The ethyl
 acetate was concentrated to dryness and purified by column chromatography, eluted with EtOAc/hexanes
 (1:1) followed by 10% methanol in CH₂Cl₂ to give compound 223 as a colorless oil. LCMS and NMR were
 consistent with the structure. 10 g (32.86 mmol) of 1-Tosyltriethylene glycol (compound 223) was treated
 10 with sodium azide (10.68 g, 164.28 mmol) in DMSO (100mL) at room temperature for 17 hours. The
 reaction mixture was then poured onto water, and extracted with EtOAc. The organic layer was washed with
 water three times and dried over Na₂SO₄. The organic layer was concentrated to dryness to give 5.3g of
 compound 224 (92%). LCMS and NMR were consistent with the structure. 1-Azidotriethylene glycol
 (compound 224, 5.53 g, 23.69 mmol) and compound 4 (6 g, 18.22 mmol) were treated with 4A molecular
 15 sieves (5g), and TMSOTf (1.65 ml, 9.11 mmol) in dichloromethane (100mL) under an inert atmosphere.
 After 14 hours, the reaction was filtered to remove the sieves, and the organic layer was washed with sat.
 NaHCO₃, water, brine, and dried over Na₂SO₄. The organic layer was concentrated to dryness and purified
 by column chromatography, eluted with a gradient of 2 to 4% methanol in dichloromethane to give
 compound 225. LCMS and NMR were consistent with the structure. Compound 225 (11.9 g, 23.59 mmol)
 20 was hydrogenated in EtOAc/Methanol (4:1, 250mL) over Pearlman's catalyst. After 8 hours, the catalyst was

removed by filtration and the solvents removed to dryness to give compound 226. LCMS and NMR were consistent with the structure.

In order to generate compound 227, a solution of nitromethanetrispropionic acid (4.17 g, 15.04 mmol) and Hunig's base (10.3 ml, 60.17 mmol) in DMF (100mL) were treated dropwise with
5 pentafluorotrifluoro acetate (9.05 ml, 52.65 mmol). After 30 minutes, the reaction was poured onto ice water and extracted with EtOAc. The organic layer was washed with water, brine, and dried over Na₂SO₄. The organic layer was concentrated to dryness and then recrystallized from heptane to give compound 227 as a white solid. LCMS and NMR were consistent with the structure. Compound 227 (1.5 g, 1.93 mmol) and compound 226 (3.7 g, 7.74 mmol) were stirred at room temperature in acetonitrile (15 mL) for 2 hours. The
10 reaction was then evaporated to dryness and purified by column chromatography, eluting with a gradient of 2 to 10% methanol in dichloromethane to give compound 228. LCMS and NMR were consistent with the structure. Compound 228 (1.7 g, 1.02 mmol) was treated with Raney Nickel (about 2g wet) in ethanol (100mL) in an atmosphere of hydrogen. After 12 hours, the catalyst was removed by filtration and the organic layer was evaporated to a solid that was used directly in the next step. LCMS and NMR were
15 consistent with the structure. This solid (0.87 g, 0.53 mmol) was treated with benzylglutaric acid (0.18 g, 0.8 mmol), HBTU (0.3 g, 0.8 mmol) and DIEA (273.7 μ l, 1.6 mmol) in DMF (5mL). After 16 hours, the DMF was removed under reduced pressure at 65°C to an oil, and the oil was dissolved in dichloromethane. The organic layer was washed with sat. NaHCO₃, brine, and dried over Na₂SO₄. After evaporation of the organic layer, the compound was purified by column chromatography and eluted with a gradient of 2 to 20%
20 methanol in dichloromethane to give the coupled product. LCMS and NMR were consistent with the structure. The benzyl ester was deprotected with Pearlman's catalyst under a hydrogen atmosphere for 1 hour. The catalyst was then removed by filtration and the solvents removed to dryness to give the acid. LCMS and NMR were consistent with the structure. The acid (486 mg, 0.27 mmol) was dissolved in dry DMF (3 mL). Pyridine (53.61 μ l, 0.66 mmol) was added and the reaction was purged with argon.
25 Pentafluorotrifluoro acetate (46.39 μ l, 0.4 mmol) was slowly added to the reaction mixture. The color of the reaction changed from pale yellow to burgundy, and gave off a light smoke which was blown away with a stream of argon. The reaction was allowed to stir at room temperature for one hour (completion of reaction was confirmed by LCMS). The solvent was removed under reduced pressure (rotovap) at 70 °C. The residue was diluted with DCM and washed with 1N NaHSO₄, brine, saturated sodium bicarbonate and brine
30 again. The organics were dried over Na₂SO₄, filtered, and were concentrated to dryness to give 225 mg of compound 229 as a brittle yellow foam. LCMS and NMR were consistent with the structure.

Oligomeric compound 230, comprising a GalNAc₃-23 conjugate group, was prepared from compound 229 using the general procedure illustrated in Example 46. The GalNAc₃ cluster portion of the GalNAc₃-23 conjugate group (GalNAc₃-23_a) can be combined with any cleavable moiety to provide a variety
35 of conjugate groups. The structure of GalNAc₃-23 (GalNAc₃-23_a-CM) is shown below:



Example 77: Antisense inhibition *in vivo* by oligonucleotides targeting SRB-1 comprising a GalNAc₃ conjugate

The oligonucleotides listed below were tested in a dose-dependent study for antisense inhibition of SRB-1 in mice.

Table 64
Modified ASOs targeting SRB-1

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
661161	GalNAc₃-3_a-o ·A _{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} G ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} T ^m _{es} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{ds} C ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{ds}	GalNAc ₃ -3 _a	A _d	145
666904	GalNAc₃-3_a-o ·G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} G ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} T ^m _{es} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{ds} C ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{ds}	GalNAc ₃ -3 _a	PO	143
673502	GalNAc₃-10_a-o ·A _{do} G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{eo} A ^m _{es} G ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} T ^m _{es} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{ds} C ^m _{eo} C ^m _{es} T ^m _{es} T ^m _{es}	GalNAc ₃ -10 _a	A _d	145
677844	GalNAc₃-9_a-o ·A _{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} G ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} T ^m _{es} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{ds} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es}	GalNAc ₃ -9 _a	A _d	145
677843	GalNAc₃-23_a-o ·A _{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} G ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} T ^m _{es} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{ds} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es}	GalNAc ₃ -23 _a	A _d	145
655861	G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} G ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} T ^m _{es} G ^m _{es} A ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} A _{do} ·-GalNAc ₃ -1 _a	GalNAc ₃ -1 _a	A _d	144
677841	G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} G ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} T ^m _{es} G ^m _{es} A ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} A _{do} ·-GalNAc ₃ -19 _a	GalNAc ₃ -19 _a	A _d	144
677842	G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} G ^m _{es} T ^m _{es} C ^m _{es} A ^m _{es} T ^m _{es} G ^m _{es} A ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} A _{do} ·-GalNAc ₃ -20 _a	GalNAc ₃ -20 _a	A _d	144

The structure of GalNAc₃-1_a was shown previously in Example 9, GalNAc₃-3_a was shown in Example 39, GalNAc₃-9_a was shown in Example 52, GalNAc₃-10_a was shown in Example 46, GalNAc₃-19_a was shown in Example 70, GalNAc₃-20_a was shown in Example 71, and GalNAc₃-23_a was shown in Example 76.

Treatment

Six to eight week old C57BL/6 mice (Jackson Laboratory, Bar Harbor, ME) were each injected subcutaneously once at a dosage shown below with an oligonucleotide listed in Table 64 or with saline. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration to determine the SRB-1 mRNA levels using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. The results below are presented as the average percent of SRB-1 mRNA levels for each treatment group, normalized to the saline control.

As illustrated in Table 65, treatment with antisense oligonucleotides lowered SRB-1 mRNA levels in a dose-dependent manner.

Table 65**SRB-1 mRNA (% Saline)**

ISIS No.	Dosage (mg/kg)	SRB-1 mRNA (% Saline)	GalNAc ₃ Cluster	CM
Saline	n/a	100.0	n/a	n/a
661161	0.5	89.18	GalNAc ₃ -3a	A _d
	1.5	77.02		
	5	29.10		
	15	12.64		
666904	0.5	93.11	GalNAc ₃ -3a	PO
	1.5	55.85		
	5	21.29		
	15	13.43		
673502	0.5	77.75	GalNAc ₃ -10a	A _d
	1.5	41.05		
	5	19.27		
	15	14.41		
677844	0.5	87.65	GalNAc ₃ -9a	A _d
	1.5	93.04		
	5	40.77		
	15	16.95		
677843	0.5	102.28	GalNAc ₃ -23a	A _d
	1.5	70.51		
	5	30.68		
	15	13.26		
655861	0.5	79.72	GalNAc ₃ -1a	A _d
	1.5	55.48		
	5	26.99		
	15	17.58		
677841	0.5	67.43	GalNAc ₃ -19a	A _d
	1.5	45.13		
	5	27.02		
	15	12.41		
677842	0.5	64.13	GalNAc ₃ -20a	A _d
	1.5	53.56		
	5	20.47		
	15	10.23		

Liver transaminase levels, alanine aminotransferase (ALT) and aspartate aminotransferase (AST), in serum were also measured using standard protocols. Total bilirubin and BUN were also evaluated. Changes in body weights were evaluated, with no significant change from the saline group (data not shown). ALTs, ASTs, total bilirubin and BUN values are shown in Table 66 below.

Table 66

ISIS No.	Dosage (mg/kg)	ALT (U/L)	AST (U/L)	Total Bilirubin (mg/dL)	BUN (mg/dL)	GalNAc ₃ Cluster	CM
Saline	n/a	21	45	0.13	34	n/a	n/a
661161	0.5	28	51	0.14	39	GalNAc ₃ -3a	A _d
	1.5	23	42	0.13	39		
	5	22	59	0.13	37		
	15	21	56	0.15	35		
666904	0.5	24	56	0.14	37	GalNAc ₃ -3a	PO
	1.5	26	68	0.15	35		
	5	23	77	0.14	34		
	15	24	60	0.13	35		
673502	0.5	24	59	0.16	34	GalNAc ₃ -10a	A _d
	1.5	20	46	0.17	32		
	5	24	45	0.12	31		
	15	24	47	0.13	34		
677844	0.5	25	61	0.14	37	GalNAc ₃ -9a	A _d
	1.5	23	64	0.17	33		
	5	25	58	0.13	35		
	15	22	65	0.14	34		
677843	0.5	53	53	0.13	35	GalNAc ₃ -23a	A _d
	1.5	25	54	0.13	34		
	5	21	60	0.15	34		
	15	22	43	0.12	38		
655861	0.5	21	48	0.15	33	GalNAc ₃ -1a	A _d
	1.5	28	54	0.12	35		
	5	22	60	0.13	36		
	15	21	55	0.17	30		
677841	0.5	32	54	0.13	34	GalNAc ₃ -19a	A _d
	1.5	24	56	0.14	34		
	5	23	92	0.18	31		
	15	24	58	0.15	31		
677842	0.5	23	61	0.15	35	GalNAc ₃ -20a	A _d
	1.5	24	57	0.14	34		
	5	41	62	0.15	35		
	15	24	37	0.14	32		

Example 78: Antisense inhibition *in vivo* by oligonucleotides targeting Angiotensinogen comprising a GalNAc₃ conjugate

The oligonucleotides listed below were tested in a dose-dependent study for antisense inhibition of Angiotensinogen (AGT) in normotensive Sprague Dawley rats.

Table 67
Modified ASOs targeting AGT

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
552668	^m C _{es} A _{es} ^m C _{es} T _{es} G _{es} A _{ds} T _{ds} T _{ds} T _{ds} T _{ds} G _{ds} ^m C _{ds} ^m C _{ds} ^m C _{ds} A _{es} G _{es} G _{es} A _{es} T _e	n/a	n/a	149
669509	^m C _{es} A _{es} ^m C _{es} T _{es} G _{es} A _{ds} T _{ds} T _{ds} T _{ds} T _{ds} G _{ds} ^m C _{ds} ^m C _{ds} ^m C _{ds} A _{es} G _{es} G _{es} A _{es} T _{eo} A _{do} ⁺ -GalNAc ₃ -1 _a	GalNAc ₃ -1 _a	A _d	150

5 The structure of GalNAc₃-1_a was shown previously in Example 9.

Treatment

10 Six week old, male Sprague Dawley rats were each injected subcutaneously once per week at a dosage shown below, for a total of three doses, with an oligonucleotide listed in Table 67 or with PBS. Each treatment group consisted of 4 animals. The rats were sacrificed 72 hours following the final dose. AGT liver mRNA levels were measured using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. AGT plasma protein levels were measured using the Total Angiotensinogen ELISA (Catalog # JP27412, IBL International, Toronto, ON) with plasma diluted 1:20,000. The results below are presented as the average percent of AGT mRNA levels in liver or AGT protein levels in plasma for each treatment group, normalized to the PBS control.

As illustrated in Table 68, treatment with antisense oligonucleotides lowered AGT liver mRNA and plasma protein levels in a dose-dependent manner, and the oligonucleotide comprising a GalNAc conjugate was significantly more potent than the parent oligonucleotide lacking a GalNAc conjugate.

Table 68
AGT liver mRNA and plasma protein levels

ISIS No.	Dosage (mg/kg)	AGT liver mRNA (% PBS)	AGT plasma protein (% PBS)	GalNAc ₃ Cluster	CM
PBS	n/a	100	100	n/a	n/a
552668	3	95	122	n/a	n/a
	10	85	97		
	30	46	79		
	90	8	11		
669509	0.3	95	70	GalNAc ₃ -1 _a	A _d
	1	95	129		
	3	62	97		
	10	9	23		

Liver transaminase levels, alanine aminotransferase (ALT) and aspartate aminotransferase (AST), in plasma and body weights were also measured at time of sacrifice using standard protocols. The results are shown in Table 69 below.

Table 69

Liver transaminase levels and rat body weights

ISIS No.	Dosage (mg/kg)	ALT (U/L)	AST (U/L)	Body Weight (% of baseline)	GalNAc ₃ Cluster	CM
PBS	n/a	51	81	186	n/a	n/a
552668	3	54	93	183	n/a	n/a
	10	51	93	194		
	30	59	99	182		
	90	56	78	170		
669509	0.3	53	90	190	GalNAc ₃ -1a	A _d
	1	51	93	192		
	3	48	85	189		
	10	56	95	189		

Example 79: Duration of action *in vivo* of oligonucleotides targeting APOC-III comprising a GalNAc₃ conjugate

The oligonucleotides listed in Table 70 below were tested in a single dose study for duration of action in mice.

Table 70

Modified ASOs targeting APOC-III

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
304801	A _{es} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{es} T _{es} T _{es} A _{es} T _e	n/a	n/a	135
647535	A _{es} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{es} T _{es} T _{es} A _{es} T _{eo} A _{do} -GalNAc ₃ -1a	GalNAc ₃ -1a	A _d	136
663083	GalNAc ₃ -3a-o'-A _{do} A _{es} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{es} T _{es} T _{es} A _{es} T _e	GalNAc ₃ -3a	A _d	151
674449	GalNAc ₃ -7a-o'-A _{do} A _{es} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{es} T _{es} T _{es} A _{es} T _e	GalNAc ₃ -7a	A _d	151
674450	GalNAc ₃ -10a-o'-A _{do} A _{es} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{es} T _{es} T _{es} A _{es} T _e	GalNAc ₃ -10a	A _d	151
674451	GalNAc ₃ -13a-o'-A _{do} A _{es} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{es} T _{es} T _{es} A _{es} T _e	GalNAc ₃ -13a	A _d	151

The structure of GalNAc₃-1_a was shown previously in Example 9, GalNAc₃-3_a was shown in Example 39, GalNAc₃-7_a was shown in Example 48, GalNAc₃-10_a was shown in Example 46, and GalNAc₃-13_a was shown in Example 62.

Treatment

Six to eight week old transgenic mice that express human APOC-III were each injected subcutaneously once with an oligonucleotide listed in Table 70 or with PBS. Each treatment group consisted of 3 animals. Blood was drawn before dosing to determine baseline and at 72 hours, 1 week, 2 weeks, 3 weeks, 4 weeks, 5 weeks, and 6 weeks following the dose. Plasma triglyceride and APOC-III protein levels were measured as described in Example 20. The results below are presented as the average percent of plasma triglyceride and APOC-III levels for each treatment group, normalized to baseline levels, showing that the oligonucleotides comprising a GalNAc conjugate group exhibited a longer duration of action than the parent oligonucleotide without a conjugate group (ISIS 304801) even though the dosage of the parent was three times the dosage of the oligonucleotides comprising a GalNAc conjugate group.

Table 71
Plasma triglyceride and APOC-III protein levels in transgenic mice

ISIS No.	Dosage (mg/kg)	Time point (days post-dose)	Triglycerides (% baseline)	APOC-III protein (% baseline)	GalNAc ₃ Cluster	CM
PBS	n/a	3	97	102	n/a	n/a
		7	101	98		
		14	108	98		
		21	107	107		
		28	94	91		
		35	88	90		
		42	91	105		
304801	30	3	40	34	n/a	n/a
		7	41	37		
		14	50	57		
		21	50	50		
		28	57	73		
		35	68	70		
		42	75	93		
647535	10	3	36	37	GalNAc ₃ -1a	A _d
		7	39	47		
		14	40	45		
		21	41	41		
		28	42	62		
		35	69	69		
		42	85	102		
663083	10	3	24	18	GalNAc ₃ -3a	A _d
		7	28	23		
		14	25	27		
		21	28	28		
		28	37	44		
		35	55	57		
		42	60	78		
674449	10	3	29	26	GalNAc ₃ -7a	A _d
		7	32	31		

		14	38	41		
		21	44	44		
		28	53	63		
		35	69	77		
		42	78	99		
674450	10	3	33	30	GalNAc ₃ -10a	A _d
		7	35	34		
		14	31	34		
		21	44	44		
		28	56	61		
		35	68	70		
		42	83	95		
674451	10	3	35	33	GalNAc ₃ -13a	A _d
		7	24	32		
		14	40	34		
		21	48	48		
		28	54	67		
		35	65	75		
		42	74	97		

Example 80: Antisense inhibition *in vivo* by oligonucleotides targeting Alpha-1 Antitrypsin (A1AT) comprising a GalNAc₃ Conjugate

The oligonucleotides listed in Table 72 below were tested in a study for dose-dependent inhibition of A1AT in mice.

Table 72
Modified ASOs targeting A1AT

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
476366	A _{es} ^m C _{es} ^m C _{es} ^m C _{es} A _{es} A _{ds} T _{ds} T _{ds} ^m C _{ds} A _{ds} G _{ds} A _{ds} A _{ds} G _{ds} G _{ds} A _{es} A _{es} G _{es} G _{es} A _e	n/a	n/a	152
656326	A _{es} ^m C _{es} ^m C _{es} ^m C _{es} A _{es} A _{ds} T _{ds} T _{ds} ^m C _{ds} A _{ds} G _{ds} A _{ds} A _{ds} G _{ds} G _{ds} A _{es} A _{es} G _{es} G _{es} A _{eo} A_{do}-GalNAc₃-1_a	GalNAc ₃ -1a	A _d	153
678381	GalNAc₃-3_a-o A_{do} A _{es} ^m C _{es} ^m C _{es} ^m C _{es} A _{es} A _{ds} T _{ds} T _{ds} ^m C _{ds} A _{ds} G _{ds} A _{ds} A _{ds} G _{ds} G _{ds} A _{es} A _{es} G _{es} G _{es} A _e	GalNAc ₃ -3a	A _d	154
678382	GalNAc₃-7_a-o A_{do} A _{es} ^m C _{es} ^m C _{es} ^m C _{es} A _{es} A _{ds} T _{ds} T _{ds} ^m C _{ds} A _{ds} G _{ds} A _{ds} A _{ds} G _{ds} G _{ds} A _{es} A _{es} G _{es} G _{es} A _e	GalNAc ₃ -7a	A _d	154
678383	GalNAc₃-10_a-o A_{do} A _{es} ^m C _{es} ^m C _{es} ^m C _{es} A _{es} A _{ds} T _{ds} T _{ds} ^m C _{ds} A _{ds} G _{ds} A _{ds} A _{ds} G _{ds} G _{ds} A _{es} A _{es} G _{es} G _{es} A _e	GalNAc ₃ -10a	A _d	154
678384	GalNAc₃-13_a-o A_{do} A _{es} ^m C _{es} ^m C _{es} ^m C _{es} A _{es} A _{ds} T _{ds} T _{ds} ^m C _{ds} A _{ds} G _{ds} A _{ds} A _{ds} G _{ds} G _{ds} A _{es} A _{es} G _{es} G _{es} A _e	GalNAc ₃ -13a	A _d	154

The structure of GalNAc₃-1_a was shown previously in Example 9, GalNAc₃-3_a was shown in Example 39, GalNAc₃-7_a was shown in Example 48, GalNAc₃-10_a was shown in Example 46, and GalNAc₃-13_a was shown in Example 62.

Treatment

Six week old, male C57BL/6 mice (Jackson Laboratory, Bar Harbor, ME) were each injected subcutaneously once per week at a dosage shown below, for a total of three doses, with an oligonucleotide listed in Table 72 or with PBS. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration. A1AT liver mRNA levels were determined using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. A1AT plasma protein levels were determined using the Mouse Alpha 1-Antitrypsin ELISA (catalog # 41-A1AMS-E01, Alpco, Salem, NH). The results below are presented as the average percent of A1AT liver mRNA and plasma protein levels for each treatment group, normalized to the PBS control.

As illustrated in Table 73, treatment with antisense oligonucleotides lowered A1AT liver mRNA and A1AT plasma protein levels in a dose-dependent manner. The oligonucleotides comprising a GalNAc conjugate were significantly more potent than the parent (ISIS 476366).

Table 73**A1AT liver mRNA and plasma protein levels**

ISIS No.	Dosage (mg/kg)	A1AT liver mRNA (% PBS)	A1AT plasma protein (% PBS)	GalNAc ₃ Cluster	CM
PBS	n/a	100	100	n/a	n/a
476366	5	86	78	n/a	n/a
	15	73	61		
	45	30	38		
656326	0.6	99	90	GalNAc ₃ -1a	A _d
	2	61	70		
	6	15	30		
	18	6	10		
678381	0.6	105	90	GalNAc ₃ -3a	A _d
	2	53	60		
	6	16	20		
	18	7	13		
678382	0.6	90	79	GalNAc ₃ -7a	A _d
	2	49	57		
	6	21	27		
	18	8	11		
678383	0.6	94	84	GalNAc ₃ -10a	A _d
	2	44	53		
	6	13	24		
	18	6	10		
678384	0.6	106	91	GalNAc ₃ -13a	A _d
	2	65	59		
	6	26	31		
	18	11	15		

Liver transaminase and BUN levels in plasma were measured at time of sacrifice using standard protocols. Body weights and organ weights were also measured. The results are shown in Table 74 below.

Body weight is shown as % relative to baseline. Organ weights are shown as % of body weight relative to the PBS control group.

Table 74

ISIS No.	Dosage (mg/kg)	ALT (U/L)	AST (U/L)	BUN (mg/dL)	Body weight (% baseline)	Liver weight (Rel % BW)	Kidney weight (Rel % BW)	Spleen weight (Rel % BW)
PBS	n/a	25	51	37	119	100	100	100
476366	5	34	68	35	116	91	98	106
	15	37	74	30	122	92	101	128
	45	30	47	31	118	99	108	123
656326	0.6	29	57	40	123	100	103	119
	2	36	75	39	114	98	111	106
	6	32	67	39	125	99	97	122
	18	46	77	36	116	102	109	101
678381	0.6	26	57	32	117	93	109	110
	2	26	52	33	121	96	106	125
	6	40	78	32	124	92	106	126
	18	31	54	28	118	94	103	120
678382	0.6	26	42	35	114	100	103	103
	2	25	50	31	117	91	104	117
	6	30	79	29	117	89	102	107
	18	65	112	31	120	89	104	113
678383	0.6	30	67	38	121	91	100	123
	2	33	53	33	118	98	102	121
	6	32	63	32	117	97	105	105
	18	36	68	31	118	99	103	108
678384	0.6	36	63	31	118	98	103	98
	2	32	61	32	119	93	102	114
	6	34	69	34	122	100	100	96
	18	28	54	30	117	98	101	104

Example 81: Duration of action *in vivo* of oligonucleotides targeting A1AT comprising a GalNAc₃ cluster

The oligonucleotides listed in Table 72 were tested in a single dose study for duration of action in mice.

Treatment

Six week old, male C57BL/6 mice were each injected subcutaneously once with an oligonucleotide listed in Table 72 or with PBS. Each treatment group consisted of 4 animals. Blood was drawn the day before dosing to determine baseline and at 5, 12, 19, and 25 days following the dose. Plasma A1AT protein levels were measured via ELISA (see Example 80). The results below are presented as the average percent of plasma A1AT protein levels for each treatment group, normalized to baseline levels. The results show that the oligonucleotides comprising a GalNAc conjugate were more potent and had longer duration of action than the parent lacking a GalNAc conjugate (ISIS 476366). Furthermore, the oligonucleotides comprising a 5'-

GalNAc conjugate (ISIS 678381, 678382, 678383, and 678384) were generally even more potent with even longer duration of action than the oligonucleotide comprising a 3'-GalNAc conjugate (ISIS 656326).

Table 75**Plasma A1AT protein levels in mice**

ISIS No.	Dosage (mg/kg)	Time point (days post-dose)	A1AT (% baseline)	GalNAc ₃ Cluster	CM
PBS	n/a	5	93	n/a	n/a
		12	93		
		19	90		
		25	97		
476366	100	5	38	n/a	n/a
		12	46		
		19	62		
		25	77		
656326	18	5	33	GalNAc ₃ -1a	A _d
		12	36		
		19	51		
		25	72		
678381	18	5	21	GalNAc ₃ -3a	A _d
		12	21		
		19	35		
		25	48		
678382	18	5	21	GalNAc ₃ -7a	A _d
		12	21		
		19	39		
		25	60		
678383	18	5	24	GalNAc ₃ -10a	A _d
		12	21		
		19	45		
		25	73		
678384	18	5	29	GalNAc ₃ -13a	A _d
		12	34		
		19	57		
		25	76		

Example 82: Antisense inhibition *in vitro* by oligonucleotides targeting SRB-1 comprising a GalNAc₃ conjugate

Primary mouse liver hepatocytes were seeded in 96 well plates at 15,000 cells/well 2 hours prior to treatment. The oligonucleotides listed in Table 76 were added at 2, 10, 50, or 250 nM in Williams E medium and cells were incubated overnight at 37 °C in 5% CO₂. Cells were lysed 16 hours following oligonucleotide addition, and total RNA was purified using RNease 3000 BioRobot (Qiagen). SRB-1 mRNA levels were determined using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. IC₅₀ values were determined using Prism 4 software

(GraphPad). The results show that oligonucleotides comprising a variety of different GalNAc conjugate groups and a variety of different cleavable moieties are significantly more potent in an *in vitro* free uptake experiment than the parent oligonucleotides lacking a GalNAc conjugate group (ISIS 353382 and 666841).

Table 76

Inhibition of SRB-1 expression *in vitro*

ISIS No.	Sequence (5' to 3')	Linkages	GalNAc cluster	CM	IC ₅₀ (nM)	SEQ ID No.
353382	G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	n/a	n/a	250	143
655861	G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _{eo} GalNAc₃-1_a	PS	GalNAc ₃ -1 _a	A _d	40	144
661161	GalNAc₃-3_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -3 _a	A _d	40	145
661162	GalNAc₃-3_a-o'-A_{do} G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{eo} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{eo} C ^m _{eo} C ^m _{es} T ^m _{es} T ^m _e	PO/PS	GalNAc ₃ -3 _a	A _d	8	145
664078	G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _{eo} GalNAc₃-9_a	PS	GalNAc ₃ -9 _a	A _d	20	144
665001	GalNAc₃-8_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -8 _a	A _d	70	145
666224	GalNAc₃-5_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -5 _a	A _d	80	145
666841	G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{eo} C ^m _{eo} C ^m _{es} T ^m _{es} T ^m _e	PO/PS	n/a	n/a	>250	143
666881	GalNAc₃-10_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -10 _a	A _d	30	145
666904	GalNAc₃-3_a-o'-G_{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -3 _a	PO	9	143
666924	GalNAc₃-3_a-o'-T_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -3 _a	T _d	15	148
666961	GalNAc₃-6_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -6 _a	A _d	150	145
666981	GalNAc₃-7_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -7 _a	A _d	20	145
670061	GalNAc₃-13_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -13 _a	A _d	30	145
670699	GalNAc₃-3_a-o'-T_{do} G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{eo} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{eo} C ^m _{eo} C ^m _{es} T ^m _{es} T ^m _e	PO/PS	GalNAc ₃ -3 _a	T _d	15	148
670700	GalNAc₃-3_a-o'-A_{eo} G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{eo} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{eo} C ^m _{eo} C ^m _{es} T ^m _{es} T ^m _e	PO/PS	GalNAc ₃ -3 _a	A _e	30	145
670701	GalNAc₃-3_a-o'-T_{eo} G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{eo} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{eo} C ^m _{eo} C ^m _{es} T ^m _{es} T ^m _e	PO/PS	GalNAc ₃ -3 _a	T _e	25	148
671144	GalNAc₃-12_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -12 _a	A _d	40	145

671165	GalNAc₃-13_a-o'-A_{do} G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{eo} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{eo} C ^m _{eo} C ^m _{es} T ^m _{es} T ^m _e	PO/PS	GalNAc ₃ -13 _a	A _d	8	145
671261	GalNAc₃-14_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -14 _a	A _d	>250	145
671262	GalNAc₃-15_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -15 _a	A _d	>250	145
673501	GalNAc₃-7_a-o'-A_{do} G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{eo} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{eo} C ^m _{eo} C ^m _{es} T ^m _{es} T ^m _e	PO/PS	GalNAc ₃ -7 _a	A _d	30	145
673502	GalNAc₃-10_a-o'-A_{do} G ^m _{es} C ^m _{eo} T ^m _{eo} T ^m _{eo} C ^m _{eo} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{eo} C ^m _{eo} C ^m _{es} T ^m _{es} T ^m _e	PO/PS	GalNAc ₃ -10 _a	A _d	8	145
675441	GalNAc₃-17_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -17 _a	A _d	30	145
675442	GalNAc₃-18_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -18 _a	A _d	20	145
677841	GalNAc₃-19_a G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _{eo} A ^m _{do}	PS	GalNAc ₃ -19 _a	A _d	40	144
677842	GalNAc₃-20_a G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _{eo} A ^m _{do}	PS	GalNAc ₃ -20 _a	A _d	30	144
677843	GalNAc₃-23_a-o'-A_{do} G ^m _{es} C ^m _{es} T ^m _{es} T ^m _{es} C ^m _{es} A ^m _{ds} G ^m _{ds} T ^m _{ds} C ^m _{ds} A ^m _{ds} T ^m _{ds} G ^m _{ds} A ^m _{ds} C ^m _{ds} T ^m _{ds} T ^m _{es} C ^m _{es} C ^m _{es} T ^m _{es} T ^m _e	PS	GalNAc ₃ -23 _a	A _d	40	145

The structure of GalNAc₃-1_a was shown previously in Example 9, GalNAc₃-3_a was shown in Example 39, GalNAc₃-5_a was shown in Example 49, GalNAc₃-6_a was shown in Example 51, GalNAc₃-7_a was shown in Example 48, GalNAc₃-8_a was shown in Example 47, GalNAc₃-9_a was shown in Example 52, GalNAc₃-10_a was shown in Example 46, GalNAc₃-12_a was shown in Example 61, GalNAc₃-13_a was shown in Example 62, GalNAc₃-14_a was shown in Example 63, GalNAc₃-15_a was shown in Example 64, GalNAc₃-17_a was shown in Example 68, GalNAc₃-18_a was shown in Example 69, GalNAc₃-19_a was shown in Example 70, GalNAc₃-20_a was shown in Example 71, and GalNAc₃-23_a was shown in Example 76.

Example 83: Antisense inhibition *in vivo* by oligonucleotides targeting Factor XI comprising a GalNAc₃ cluster

The oligonucleotides listed in Table 77 below were tested in a study for dose-dependent inhibition of Factor XI in mice.

Table 77
Modified oligonucleotides targeting Factor XI

ISIS No.	Sequence (5' to 3')	GalNAc cluster	CM	SEQ ID No.
404071	T _{es} G _{es} G _{es} T _{es} A _{es} A _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ds} T _{ds} ^m C _{ds} A _{es} G _{es} A _{es} G _{es} G _e	n/a	n/a	146
656173	T _{es} G _{eo} G _{eo} T _{eo} A _{eo} A _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ds} T _{ds} ^m C _{ds} A _{eo} G _{eo} A _{es} G _{es} G _{eo} A _{do} '-GalNAc ₃ -1 _a	GalNAc ₃ -1 _a	A _d	147
663086	GalNAc₃-3_a-o'-A_{do} T _{es} G _{eo} G _{eo} T _{eo} A _{eo} A _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} ^m C _{ds} T _{ds}	GalNAc ₃ -3 _a	A _d	155

	$T_{ds}T_{ds}^mC_{ds}A_{eo}G_{eo}A_{es}G_{es}G_e$			
678347	GalNAc₃-7_a -o'-A _{do} T _{es} G _{eo} G _{eo} T _{eo} A _{eo} A _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} ^m C _{ds} T _{ds} $T_{ds}T_{ds}^mC_{ds}A_{eo}G_{eo}A_{es}G_{es}G_e$	GalNAc ₃ -7 _a	A _d	155
678348	GalNAc₃-10_a -o'-A _{do} T _{es} G _{eo} G _{eo} T _{eo} A _{eo} A _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} ^m C _{ds} $T_{ds}T_{ds}^mC_{ds}A_{eo}G_{eo}A_{es}G_{es}G_e$	GalNAc ₃ -10 _a	A _d	155
678349	GalNAc₃-13_a -o'-A _{do} T _{es} G _{eo} G _{eo} T _{eo} A _{eo} A _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} ^m C _{ds} $T_{ds}T_{ds}^mC_{ds}A_{eo}G_{eo}A_{es}G_{es}G_e$	GalNAc ₃ -13 _a	A _d	155

The structure of GalNAc₃-1_a was shown previously in Example 9, GalNAc₃-3_a was shown in Example 39, GalNAc₃-7_a was shown in Example 48, GalNAc₃-10_a was shown in Example 46, and GalNAc₃-13_a was shown in Example 62.

5 Treatment

Six to eight week old mice were each injected subcutaneously once per week at a dosage shown below, for a total of three doses, with an oligonucleotide listed below or with PBS. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final dose. Factor XI liver mRNA levels were measured using real-time PCR and normalized to cyclophilin according to standard protocols. Liver transaminases, BUN, and bilirubin were also measured. The results below are presented as the average percent for each treatment group, normalized to the PBS control.

As illustrated in Table 78, treatment with antisense oligonucleotides lowered Factor XI liver mRNA in a dose-dependent manner. The results show that the oligonucleotides comprising a GalNAc conjugate were more potent than the parent lacking a GalNAc conjugate (ISIS 404071). Furthermore, the oligonucleotides comprising a 5'-GalNAc conjugate (ISIS 663086, 678347, 678348, and 678349) were even more potent than the oligonucleotide comprising a 3'-GalNAc conjugate (ISIS 656173).

Table 78

Factor XI liver mRNA, liver transaminase, BUN, and bilirubin levels

ISIS No.	Dosage (mg/kg)	Factor XI mRNA (% PBS)	ALT (U/L)	AST (U/L)	BUN (mg/dL)	Bilirubin (mg/dL)	GalNAc ₃ Cluster	SEQ ID No.
PBS	n/a	100	63	70	21	0.18	n/a	n/a
404071	3	65	41	58	21	0.15	n/a	146
	10	33	49	53	23	0.15		
	30	17	43	57	22	0.14		
656173	0.7	43	90	89	21	0.16	GalNAc ₃ -1a	147
	2	9	36	58	26	0.17		
	6	3	50	63	25	0.15		
663086	0.7	33	91	169	25	0.16	GalNAc ₃ -3a	155
	2	7	38	55	21	0.16		
	6	1	34	40	23	0.14		
678347	0.7	35	28	49	20	0.14	GalNAc ₃ -7a	155
	2	10	180	149	21	0.18		
	6	1	44	76	19	0.15		
678348	0.7	39	43	54	21	0.16	GalNAc ₃ -10a	155
	2	5	38	55	22	0.17		

	6	2	25	38	20	0.14		
678349	0.7	34	39	46	20	0.16	GalNAc ₃ -13a	155
	2	8	43	63	21	0.14		
	6	2	28	41	20	0.14		

Example 84: Duration of action *in vivo* of oligonucleotides targeting Factor XI comprising a GalNAc₃ Conjugate

The oligonucleotides listed in Table 77 were tested in a single dose study for duration of action in mice.

Treatment

Six to eight week old mice were each injected subcutaneously once with an oligonucleotide listed in Table 77 or with PBS. Each treatment group consisted of 4 animals. Blood was drawn by tail bleeds the day before dosing to determine baseline and at 3, 10, and 17 days following the dose. Plasma Factor XI protein levels were measured by ELISA using Factor XI capture and biotinylated detection antibodies from R & D Systems, Minneapolis, MN (catalog # AF2460 and # BAF2460, respectively) and the OptEIA Reagent Set B (Catalog # 550534, BD Biosciences, San Jose, CA). The results below are presented as the average percent of plasma Factor XI protein levels for each treatment group, normalized to baseline levels. The results show that the oligonucleotides comprising a GalNAc conjugate were more potent with longer duration of action than the parent lacking a GalNAc conjugate (ISIS 404071). Furthermore, the oligonucleotides comprising a 5'-GalNAc conjugate (ISIS 663086, 678347, 678348, and 678349) were even more potent with an even longer duration of action than the oligonucleotide comprising a 3'-GalNAc conjugate (ISIS 656173).

Table 79

Plasma Factor XI protein levels in mice

ISIS No.	Dosage (mg/kg)	Time point (days post-dose)	Factor XI (% baseline)	GalNAc ₃ Cluster	CM	SEQ ID No.
PBS	n/a	3	123	n/a	n/a	n/a
		10	56			
		17	100			
404071	30	3	11	n/a	n/a	146
		10	47			
		17	52			
656173	6	3	1	GalNAc ₃ -1a	A _d	147
		10	3			
		17	21			
663086	6	3	1	GalNAc ₃ -3a	A _d	155
		10	2			
		17	9			
678347	6	3	1	GalNAc ₃ -7a	A _d	155
		10	1			
		17	8			

678348	6	3	1	GalNAc ₃ -10a	A _d	155
		10	1			
		17	6			
678349	6	3	1	GalNAc ₃ -13a	A _d	155
		10	1			
		17	5			

Example 85: Antisense inhibition *in vivo* by oligonucleotides targeting SRB-1 comprising a GalNAc₃ Conjugate

Oligonucleotides listed in Table 76 were tested in a dose-dependent study for antisense inhibition of SRB-1 in mice.

Treatment

Six to eight week old C57BL/6 mice were each injected subcutaneously once per week at a dosage shown below, for a total of three doses, with an oligonucleotide listed in Table 76 or with saline. Each treatment group consisted of 4 animals. The mice were sacrificed 48 hours following the final administration to determine the SRB-1 mRNA levels using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. The results below are presented as the average percent of liver SRB-1 mRNA levels for each treatment group, normalized to the saline control.

As illustrated in Tables 80 and 81, treatment with antisense oligonucleotides lowered SRB-1 mRNA levels in a dose-dependent manner.

Table 80
SRB-1 mRNA in liver

ISIS No.	Dosage (mg/kg)	SRB-1 mRNA (% Saline)	GalNAc ₃ Cluster	CM
Saline	n/a	100	n/a	n/a
655861	0.1	94	GalNAc ₃ -1a	A _d
	0.3	119		
	1	68		
	3	32		
661161	0.1	120	GalNAc ₃ -3a	A _d
	0.3	107		
	1	68		
	3	26		
666881	0.1	107	GalNAc ₃ -10a	A _d
	0.3	107		
	1	69		
	3	27		
666981	0.1	120	GalNAc ₃ -7a	A _d
	0.3	103		
	1	54		
	3	21		
670061	0.1	118	GalNAc ₃ -13a	A _d
	0.3	89		

677842	1	52	GalNAc ₃ -20a	A _d
	3	18		
	0.1	119		
	0.3	96		
	1	65		
	3	23		

Table 81**SRB-1 mRNA in liver**

ISIS No.	Dosage (mg/kg)	SRB-1 mRNA (% Saline)	GalNAc ₃ Cluster	CM
661161	0.1	107	GalNAc ₃ -3a	A _d
	0.3	95		
	1	53		
	3	18		
677841	0.1	110	GalNAc ₃ -19a	A _d
	0.3	88		
	1	52		
	3	25		

5 Liver transaminase levels, total bilirubin, BUN, and body weights were also measured using standard protocols. Average values for each treatment group are shown in Table 82 below.

Table 82

ISIS No.	Dosage (mg/kg)	ALT (U/L)	AST (U/L)	Bilirubin (mg/dL)	BUN (mg/dL)	Body Weight (% baseline)	GalNAc ₃ Cluster	CM
Saline	n/a	19	39	0.17	26	118	n/a	n/a
655861	0.1	25	47	0.17	27	114	GalNAc ₃ -1a	A _d
	0.3	29	56	0.15	27	118		
	1	20	32	0.14	24	112		
	3	27	54	0.14	24	115		
661161	0.1	35	83	0.13	24	113	GalNAc ₃ -3a	A _d
	0.3	42	61	0.15	23	117		
	1	34	60	0.18	22	116		
	3	29	52	0.13	25	117		
666881	0.1	30	51	0.15	23	118	GalNAc ₃ -10a	A _d
	0.3	49	82	0.16	25	119		
	1	23	45	0.14	24	117		
	3	20	38	0.15	21	112		
666981	0.1	21	41	0.14	22	113	GalNAc ₃ -7a	A _d
	0.3	29	49	0.16	24	112		
	1	19	34	0.15	22	111		
	3	77	78	0.18	25	115		
670061	0.1	20	63	0.18	24	111	GalNAc ₃ -13a	A _d
	0.3	20	57	0.15	21	115		
	1	20	35	0.14	20	115		
	3	27	42	0.12	20	116		
677842	0.1	20	38	0.17	24	114	GalNAc ₃ -20a	A _d
	0.3	31	46	0.17	21	117		
	1	22	34	0.15	21	119		

	3	41	57	0.14	23	118		
--	---	----	----	------	----	-----	--	--

Example 86: Antisense inhibition *in vivo* by oligonucleotides targeting TTR comprising a GalNAc₃ cluster

Oligonucleotides listed in Table 83 below were tested in a dose-dependent study for antisense inhibition of human transthyretin (TTR) in transgenic mice that express the human TTR gene.

Treatment

Eight week old TTR transgenic mice were each injected subcutaneously once per week for three weeks, for a total of three doses, with an oligonucleotide and dosage listed in the tables below or with PBS. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration. Tail bleeds were performed at various time points throughout the experiment, and plasma TTR protein, ALT, and AST levels were measured and reported in Tables 85-87. After the animals were sacrificed, plasma ALT, AST, and human TTR levels were measured, as were body weights, organ weights, and liver human TTR mRNA levels. TTR protein levels were measured using a clinical analyzer (AU480, Beckman Coulter, CA). Real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) were used according to standard protocols to determine liver human TTR mRNA levels. The results presented in Tables 84-87 are the average values for each treatment group. The mRNA levels are the average values relative to the average for the PBS group. Plasma protein levels are the average values relative to the average value for the PBS group at baseline. Body weights are the average percent weight change from baseline until sacrifice for each individual treatment group. Organ weights shown are normalized to the animal's body weight, and the average normalized organ weight for each treatment group is then presented relative to the average normalized organ weight for the PBS group.

In Tables 84-87, "BL" indicates baseline, measurements that were taken just prior to the first dose. As illustrated in Tables 84 and 85, treatment with antisense oligonucleotides lowered TTR expression levels in a dose-dependent manner. The oligonucleotides comprising a GalNAc conjugate were more potent than the parent lacking a GalNAc conjugate (ISIS 420915). Furthermore, the oligonucleotides comprising a GalNAc conjugate and mixed PS/PO internucleoside linkages were even more potent than the oligonucleotide comprising a GalNAc conjugate and full PS linkages.

Table 83
Oligonucleotides targeting human TTR

Isis No.	Sequence 5' to 3'	Linkages	GalNAc cluster	CM	SEQ ID No.
420915	T _{es} ^m C _{es} T _{es} T _{es} G _{es} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{es} T _{es} ^m C _{es} ^m C _{es} ^m C _e	PS	n/a	n/a	156
660261	T _{es} ^m C _{es} T _{es} T _{es} G _{es} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{es} T _{es} ^m C _{es} ^m C _{es} ^m C _{eo} A _{do} '-GalNAc ₃ -1 _a	PS	GalNAc ₃ -1a	A _d	157
682883	GalNAc ₃ -3 _{a-o} '-T _{es} ^m C _{eo} T _{eo} T _{eo} G _{eo} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds}	PS/PO	GalNAc ₃ -3a	PO	156

	$T_{ds}G_{ds}A_{ds}A_{ds}A_{eo}T_{eo}^mC_{es}^mC_{es}^mC_e$				
682884	GalNAc₃-7_{a-o} , $T_{es}^mC_{eo}T_{eo}T_{eo}G_{eo}G_{ds}T_{ds}T_{ds}A_{ds}^mC_{ds}A_{ds}$ $T_{ds}G_{ds}A_{ds}A_{ds}A_{eo}T_{eo}^mC_{es}^mC_{es}^mC_e$	PS/PO	GalNAc ₃ -7a	PO	156
682885	GalNAc₃-10_{a-o} , $T_{es}^mC_{eo}T_{eo}T_{eo}G_{eo}G_{ds}T_{ds}T_{ds}A_{ds}^mC_{ds}$ $A_{ds}T_{ds}G_{ds}A_{ds}A_{ds}A_{eo}T_{eo}^mC_{es}^mC_{es}^mC_e$	PS/PO	GalNAc ₃ -10a	PO	156
682886	GalNAc₃-13_{a-o} , $T_{es}^mC_{eo}T_{eo}T_{eo}G_{eo}G_{ds}T_{ds}T_{ds}A_{ds}^mC_{ds}$ $A_{ds}T_{ds}G_{ds}A_{ds}A_{ds}A_{eo}T_{eo}^mC_{es}^mC_{es}^mC_e$	PS/PO	GalNAc ₃ -13a	PO	156
684057	$T_{es}^mC_{eo}T_{eo}T_{eo}G_{eo}G_{ds}T_{ds}T_{ds}A_{ds}^mC_{ds}A_{ds}T_{ds}G_{ds}A_{ds}A_{ds}$ $A_{eo}T_{eo}^mC_{es}^mC_{es}^mC_{eo}A_{do}$ - GalNAc₃-19_a	PS/PO	GalNAc ₃ -19a	A _d	157

The legend for Table 85 can be found in Example 74. The structure of GalNAc₃-1 was shown in Example 9.

The structure of GalNAc₃-3_a was shown in Example 39. The structure of GalNAc₃-7_a was shown in Example 48. The structure of GalNAc₃-10_a was shown in Example 46. The structure of GalNAc₃-13_a was shown in Example 62. The structure of GalNAc₃-19_a was shown in Example 70.

5

Table 84

Antisense inhibition of human TTR *in vivo*

Isis No.	Dosage (mg/kg)	TTR mRNA (% PBS)	Plasma TTR protein (% PBS)	GalNAc cluster	CM	SEQ ID No.
PBS	n/a	100	100	n/a	n/a	
420915	6	99	95	n/a	n/a	156
	20	48	65			
	60	18	28			
660261	0.6	113	87	GalNAc ₃ -1a	A _d	157
	2	40	56			
	6	20	27			
	20	9	11			

Table 85

Antisense inhibition of human TTR *in vivo*

Isis No.	Dosage (mg/kg)	TTR mRNA (% PBS)	Plasma TTR protein (% PBS at BL)				GalNAc cluster	CM	SEQ ID No.
			BL	Day 3	Day 10	Day 17 (After sac)			
PBS	n/a	100	100	96	90	114	n/a	n/a	
420915	6	74	106	86	76	83	n/a	n/a	156
	20	43	102	66	61	58			
	60	24	92	43	29	32			
682883	0.6	60	88	73	63	68	GalNAc ₃ -3a	PO	156
	2	18	75	38	23	23			
	6	10	80	35	11	9			
682884	0.6	56	88	78	63	67	GalNAc ₃ -7a	PO	156
	2	19	76	44	25	23			
	6	15	82	35	21	24			
682885	0.6	60	92	77	68	76	GalNAc ₃ -10a	PO	156
	2	22	93	58	32	32			
	6	17	85	37	25	20			
682886	0.6	57	91	70	64	69	GalNAc ₃ -	PO	156

	2	21	89	50	31	30	13a		
	6	18	102	41	24	27			
684057	0.6	53	80	69	56	62	GalNAc ₃ - 19a	A _d	157
	2	21	92	55	34	30			
	6	11	82	50	18	13			

Table 86

Transaminase levels, body weight changes, and relative organ weights

Isis No.	Dose (mg/kg)	ALT (U/L)				AST (U/L)				Body (% BL)	Liver (% PBS)	Spleen (% PBS)	Kidney (% PBS)	SEQ ID No.
		BL	Day 3	Day 10	Day 17	BL	Day 3	Day 10	Day 17					
PBS	n/a	33	34	33	24	58	62	67	52	105	100	100	100	n/a
420915	6	34	33	27	21	64	59	73	47	115	99	89	91	156
	20	34	30	28	19	64	54	56	42	111	97	83	89	
	60	34	35	31	24	61	58	71	58	113	102	98	95	
660261	0.6	33	38	28	26	70	71	63	59	111	96	99	92	157
	2	29	32	31	34	61	60	68	61	118	100	92	90	
	6	29	29	28	34	58	59	70	90	114	99	97	95	
	20	33	32	28	33	64	54	68	95	114	101	106	92	

5

Table 87

Transaminase levels, body weight changes, and relative organ weights

Isis No.	Dose (mg/kg)	ALT (U/L)				AST (U/L)				Body (% BL)	Liver (% PBS)	Spleen (% PBS)	Kidney (% PBS)	SEQ ID No.
		BL	Day 3	Day 10	Day 17	BL	Day 3	Day 10	Day 17					
PBS	n/a	32	34	37	41	62	78	76	77	104	100	100	100	n/a
420915	6	32	30	34	34	61	71	72	66	102	103	102	105	156
	20	41	34	37	33	80	76	63	54	106	107	135	101	
	60	36	30	32	34	58	81	57	60	106	105	104	99	
682883	0.6	32	35	38	40	53	81	74	76	104	101	112	95	156
	2	38	39	42	43	71	84	70	77	107	98	116	99	
	6	35	35	41	38	62	79	103	65	105	103	143	97	
682884	0.6	33	32	35	34	70	74	75	67	101	100	130	99	156
	2	31	32	38	38	63	77	66	55	104	103	122	100	
	6	38	32	36	34	65	85	80	62	99	105	129	95	
682885	0.6	39	26	37	35	63	63	77	59	100	109	109	112	156
	2	30	26	38	40	54	56	71	72	102	98	111	102	
	6	27	27	34	35	46	52	56	64	102	98	113	96	
682886	0.6	30	40	34	36	58	87	54	61	104	99	120	101	156
	2	27	26	34	36	51	55	55	69	103	91	105	92	
	6	40	28	34	37	107	54	61	69	109	100	102	99	
684057	0.6	35	26	33	39	56	51	51	69	104	99	110	102	157
	2	33	32	31	40	54	57	56	87	103	100	112	97	
	6	39	33	35	40	67	52	55	92	98	104	121	108	

Example 87: Duration of action *in vivo* by single doses of oligonucleotides targeting TTR comprising a GalNAc₃ cluster

ISIS numbers 420915 and 660261 (see Table 83) were tested in a single dose study for duration of action in mice. ISIS numbers 420915, 682883, and 682885 (see Table 83) were also tested in a single dose study for duration of action in mice.

5

Treatment

Eight week old, male transgenic mice that express human TTR were each injected subcutaneously once with 100 mg/kg ISIS No. 420915 or 13.5 mg/kg ISIS No. 660261. Each treatment group consisted of 4 animals. Tail bleeds were performed before dosing to determine baseline and at days 3, 7, 10, 17, 24, and 39 following the dose. Plasma TTR protein levels were measured as described in Example 86. The results below are presented as the average percent of plasma TTR levels for each treatment group, normalized to baseline levels.

10

Table 88
Plasma TTR protein levels

ISIS No.	Dosage (mg/kg)	Time point (days post-dose)	TTR (% baseline)	GalNAc ₃ Cluster	CM	SEQ ID No.
420915	100	3	30	n/a	n/a	156
		7	23			
		10	35			
		17	53			
		24	75			
		39	100			
660261	13.5	3	27	GalNAc ₃ -1a	A _d	157
		7	21			
		10	22			
		17	36			
		24	48			
		39	69			

15

Treatment

Female transgenic mice that express human TTR were each injected subcutaneously once with 100 mg/kg ISIS No. 420915, 10.0 mg/kg ISIS No. 682883, or 10.0 mg/kg 682885. Each treatment group consisted of 4 animals. Tail bleeds were performed before dosing to determine baseline and at days 3, 7, 10, 17, 24, and 39 following the dose. Plasma TTR protein levels were measured as described in Example 86. The results below are presented as the average percent of plasma TTR levels for each treatment group, normalized to baseline levels.

20

Table 89
Plasma TTR protein levels

ISIS No.	Dosage (mg/kg)	Time point (days post-dose)	TTR (% baseline)	GalNAc ₃ Cluster	CM	SEQ ID No.
420915	100	3	48	n/a	n/a	156
		7	48			
		10	48			
		17	66			
		31	80			
682883	10.0	3	45	GalNAc ₃ -3a	PO	156
		7	37			
		10	38			
		17	42			
		31	65			
682885	10.0	3	40	GalNAc ₃ -10a	PO	156
		7	33			
		10	34			
		17	40			
		31	64			

The results in Tables 88 and 89 show that the oligonucleotides comprising a GalNAc conjugate are more potent with a longer duration of action than the parent oligonucleotide lacking a conjugate (ISIS 420915).

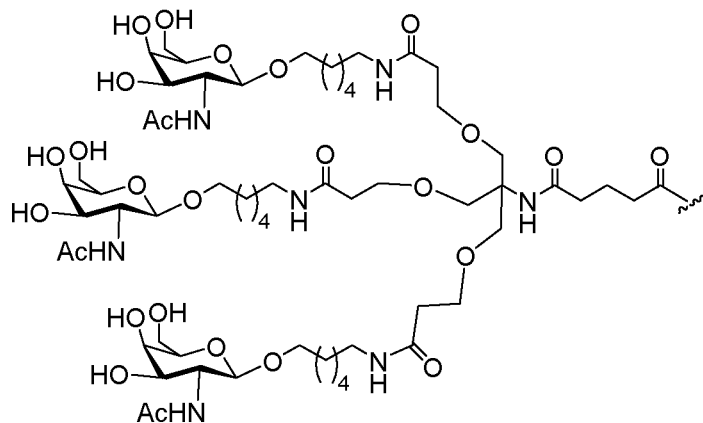
Example 88: Splicing modulation *in vivo* by oligonucleotides targeting SMN comprising a GalNAc₃ conjugate

The oligonucleotides listed in Table 90 were tested for splicing modulation of human survival of motor neuron (SMN) in mice.

Table 90
Modified ASOs targeting SMN

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
387954	A _{es} T _{es} T _{es} ^m C _{es} A _{es} ^m C _{es} T _{es} T _{es} T _{es} ^m C _{es} A _{es} T _{es} A _{es} A _{es} T _{es} G _{es} ^m C _{es} T _{es} G _{es} G _e	n/a	n/a	158
699819	GalNAc₃-7_a-o ·A _{es} T _{es} T _{es} ^m C _{es} A _{es} ^m C _{es} T _{es} T _{es} T _{es} ^m C _{es} A _{es} T _{es} A _{es} A _{es} T _{es} G _{es} ^m C _{es} T _{es} G _{es} G _e	GalNAc ₃ -7a	PO	158
699821	GalNAc₃-7_a-o ·A _{es} T _{eo} T _{eo} ^m C _{eo} A _{eo} ^m C _{eo} T _{eo} T _{eo} T _{eo} ^m C _{eo} A _{eo} T _{eo} A _{eo} A _{eo} T _{eo} G _{eo} ^m C _{eo} T _{es} G _{es} G _e	GalNAc ₃ -7a	PO	158
700000	A _{es} T _{es} T _{es} ^m C _{es} A _{es} ^m C _{es} T _{es} T _{es} T _{es} ^m C _{es} A _{es} T _{es} A _{es} A _{es} T _{es} G _{es} ^m C _{es} T _{es} G _{es} G _{eo} A _{do} ·-GalNAc ₃ -1 _a	GalNAc ₃ -1a	A _d	157
703421	X-ATT ^m CA ^m CTTT ^m CATAATG ^m CTGG	n/a	n/a	158
703422	GalNAc₃-7_b -X-ATT ^m CA ^m CTTT ^m CATAATG ^m CTGG	GalNAc ₃ -7b	n/a	158

The structure of GalNAc₃-7_a was shown previously in Example 48. “X” indicates a 5' primary amine generated by Gene Tools (Philomath, OR), and GalNAc₃-7_b indicates the structure of GalNAc₃-7_a lacking the –NH-C₆-O portion of the linker as shown below:



ISIS numbers 703421 and 703422 are morpholino oligonucleotides, wherein each nucleotide of the two oligonucleotides is a morpholino nucleotide.

5 Treatment

Six week old transgenic mice that express human SMN were injected subcutaneously once with an oligonucleotide listed in Table 91 or with saline. Each treatment group consisted of 2 males and 2 females. The mice were sacrificed 3 days following the dose to determine the liver human SMN mRNA levels both with and without exon 7 using real-time PCR according to standard protocols. Total RNA was measured using Ribogreen reagent. The SMN mRNA levels were normalized to total mRNA, and further normalized to the averages for the saline treatment group. The resulting average ratios of SMN mRNA including exon 7 to SMN mRNA missing exon 7 are shown in Table 91. The results show that fully modified oligonucleotides that modulate splicing and comprise a GalNAc conjugate are significantly more potent in altering splicing in the liver than the parent oligonucleotides lacking a GalNAc conjugate. Furthermore, this trend is maintained for multiple modification chemistries, including 2'-MOE and morpholino modified oligonucleotides.

Table 91
Effect of oligonucleotides targeting human SMN *in vivo*

ISIS No.	Dose (mg/kg)	+Exon 7 / -Exon 7	GalNAc ₃ Cluster	CM	SEQ ID No.
Saline	n/a	1.00	n/a	n/a	n/a
387954	32	1.65	n/a	n/a	158
387954	288	5.00	n/a	n/a	158
699819	32	7.84	GalNAc ₃ -7a	PO	158
699821	32	7.22	GalNAc ₃ -7a	PO	158
700000	32	6.91	GalNAc ₃ -1a	A _d	159
703421	32	1.27	n/a	n/a	158
703422	32	4.12	GalNAc ₃ -7b	n/a	158

Example 89: Antisense inhibition *in vivo* by oligonucleotides targeting Apolipoprotein A (Apo(a)) comprising a GalNAc₃ conjugate

The oligonucleotides listed in Table 92 below were tested in a study for dose-dependent inhibition of Apo(a) in transgenic mice.

5

Table 92
Modified ASOs targeting Apo(a)

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
494372	T _{es} G _{es} ^m C _{es} T _{es} ^m C _{es} ^m C _{ds} G _{ds} T _{ds} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{es} G _{es} T _{es} ^m C _e	n/a	n/a	58
681257	GalNAc₃-7_a-o' T _{es} G _{eo} ^m C _{eo} T _{eo} ^m C _{eo} ^m C _{ds} G _{ds} T _{ds} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{eo} G _{eo} T _{es} ^m C _e	GalNAc ₃ -7a	PO	58

The structure of GalNAc₃-7_a was shown in Example 48.

Treatment

10 Eight week old, female C57BL/6 mice (Jackson Laboratory, Bar Harbor, ME) were each injected subcutaneously once per week at a dosage shown below, for a total of six doses, with an oligonucleotide listed in Table 92 or with PBS. Each treatment group consisted of 3-4 animals. Tail bleeds were performed the day before the first dose and weekly following each dose to determine plasma Apo(a) protein levels. The mice were sacrificed two days following the final administration. Apo(a) liver mRNA levels were determined using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) according to standard protocols. Apo(a) plasma protein levels were determined using ELISA, and liver transaminase levels were determined. The mRNA and plasma protein results in Table 93 are presented as the treatment group average percent relative to the PBS treated group. Plasma protein levels were further normalized to the baseline (BL) value for the PBS group. Average absolute transaminase levels and body weights (% relative to baseline averages) are reported in Table 94.

15 As illustrated in Table 93, treatment with the oligonucleotides lowered Apo(a) liver mRNA and plasma protein levels in a dose-dependent manner. Furthermore, the oligonucleotide comprising the GalNAc conjugate was significantly more potent with a longer duration of action than the parent oligonucleotide lacking a GalNAc conjugate. As illustrated in Table 94, transaminase levels and body weights were unaffected by the oligonucleotides, indicating that the oligonucleotides were well tolerated.

25

Table 93
Apo(a) liver mRNA and plasma protein levels

ISIS No.	Dosage (mg/kg)	Apo(a) mRNA (% PBS)	Apo(a) plasma protein (% PBS)						
			BL	Week 1	Week 2	Week 3	Week 4	Week 5	Week 6
PBS	n/a	100	100	120	119	113	88	121	97
494372	3	80	84	89	91	98	87	87	79
	10	30	87	72	76	71	57	59	46

	30	5	92	54	28	10	7	9	7
681257	0.3	75	79	76	89	98	71	94	78
	1	19	79	88	66	60	54	32	24
	3	2	82	52	17	7	4	6	5
	10	2	79	17	6	3	2	4	5

Table 94

ISIS No.	Dosage (mg/kg)	ALT (U/L)	AST (U/L)	Body weight (% baseline)
PBS	n/a	37	54	103
494372	3	28	68	106
	10	22	55	102
	30	19	48	103
681257	0.3	30	80	104
	1	26	47	105
	3	29	62	102
	10	21	52	107

Example 90: Antisense inhibition *in vivo* by oligonucleotides targeting TTR comprising a GalNAc₃ cluster

Oligonucleotides listed in Table 95 below were tested in a dose-dependent study for antisense inhibition of human transthyretin (TTR) in transgenic mice that express the human TTR gene.

Treatment

TTR transgenic mice were each injected subcutaneously once per week for three weeks, for a total of three doses, with an oligonucleotide and dosage listed in Table 96 or with PBS. Each treatment group consisted of 4 animals. Prior to the first dose, a tail bleed was performed to determine plasma TTR protein levels at baseline (BL). The mice were sacrificed 72 hours following the final administration. TTR protein levels were measured using a clinical analyzer (AU480, Beckman Coulter, CA). Real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc. Eugene, OR) were used according to standard protocols to determine liver human TTR mRNA levels. The results presented in Table 96 are the average values for each treatment group. The mRNA levels are the average values relative to the average for the PBS group. Plasma protein levels are the average values relative to the average value for the PBS group at baseline. "BL" indicates baseline, measurements that were taken just prior to the first dose. As illustrated in Table 96, treatment with antisense oligonucleotides lowered TTR expression levels in a dose-dependent manner. The oligonucleotides comprising a GalNAc conjugate were more potent than the parent lacking a GalNAc conjugate (ISIS 420915), and oligonucleotides comprising a phosphodiester or deoxyadenosine cleavable moiety showed significant improvements in potency compared to the parent lacking a conjugate (see ISIS numbers 682883 and 666943 vs 420915 and see Examples 86 and 87).

Table 95
Oligonucleotides targeting human TTR

Isis No.	Sequence 5' to 3'	Linkages	GalNAc cluster	CM	SEQ ID No.
420915	T _{es} ^m C _{es} T _{es} T _{es} G _{es} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{es} T _{es} ^m C _{es} ^m C _{es} ^m C _e	PS	n/a	n/a	156
682883	GalNAc₃-3_{a-o} ·T _{es} ^m C _{eo} T _{eo} T _{eo} G _{eo} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{eo} T _{eo} ^m C _{es} ^m C _{es} ^m C _e	PS/PO	GalNAc ₃ -3a	PO	156
666943	GalNAc₃-3_{a-o}·A_{do} ·T _{es} ^m C _{eo} T _{eo} T _{eo} G _{eo} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{eo} T _{eo} ^m C _{es} ^m C _{es} ^m C _e	PS/PO	GalNAc ₃ -3a	A _d	160
682887	GalNAc₃-7_{a-o}·A_{do} ·T _{es} ^m C _{eo} T _{eo} T _{eo} G _{eo} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{eo} T _{eo} ^m C _{es} ^m C _{es} ^m C _e	PS/PO	GalNAc ₃ -7a	A _d	160
682888	GalNAc₃-10_{a-o}·A_{do} ·T _{es} ^m C _{eo} T _{eo} T _{eo} G _{eo} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{eo} T _{eo} ^m C _{es} ^m C _{es} ^m C _e	PS/PO	GalNAc ₃ -10a	A _d	160
682889	GalNAc₃-13_{a-o}·A_{do} ·T _{es} ^m C _{eo} T _{eo} T _{eo} G _{eo} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{eo} T _{eo} ^m C _{es} ^m C _{es} ^m C _e	PS/PO	GalNAc ₃ -13a	A _d	160

The legend for Table 95 can be found in Example 74. The structure of GalNAc₃-3_a was shown in Example 39. The structure of GalNAc₃-7_a was shown in Example 48. The structure of GalNAc₃-10_a was shown in Example 46. The structure of GalNAc₃-13_a was shown in Example 62.

Table 96
Antisense inhibition of human TTR *in vivo*

Isis No.	Dosage (mg/kg)	TTR mRNA (% PBS)	TTR protein (% BL)	GalNAc cluster	CM
PBS	n/a	100	124	n/a	n/a
420915	6	69	114	n/a	n/a
	20	71	86		
	60	21	36		
682883	0.6	61	73	GalNAc ₃ -3a	PO
	2	23	36		
	6	18	23		
666943	0.6	74	93	GalNAc ₃ -3a	A _d
	2	33	57		
	6	17	22		
682887	0.6	60	97	GalNAc ₃ -7a	A _d
	2	36	49		
	6	12	19		
682888	0.6	65	92	GalNAc ₃ -10a	A _d
	2	32	46		
	6	17	22		
682889	0.6	72	74	GalNAc ₃ -13a	A _d
	2	38	45		
	6	16	18		

Example 91: Antisense inhibition *in vivo* by oligonucleotides targeting Factor VII comprising a GalNAc₃ conjugate in non-human primates

Oligonucleotides listed in Table 97 below were tested in a non-terminal, dose escalation study for antisense inhibition of Factor VII in monkeys.

Treatment

Non-naïve monkeys were each injected subcutaneously on days 0, 15, and 29 with escalating doses of an oligonucleotide listed in Table 97 or with PBS. Each treatment group consisted of 4 males and 1 female. Prior to the first dose and at various time points thereafter, blood draws were performed to determine plasma Factor VII protein levels. Factor VII protein levels were measured by ELISA. The results presented in Table 98 are the average values for each treatment group relative to the average value for the PBS group at baseline (BL), the measurements taken just prior to the first dose. As illustrated in Table 98, treatment with antisense oligonucleotides lowered Factor VII expression levels in a dose-dependent manner, and the oligonucleotide comprising the GalNAc conjugate was significantly more potent in monkeys compared to the oligonucleotide lacking a GalNAc conjugate.

Table 97
Oligonucleotides targeting Factor VII

Isis No.	Sequence 5' to 3'	Linkages	GalNAc cluster	CM	SEQ ID No.
407935	A _{es} T _{es} G _{es} ^m C _{es} A _{es} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} A _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} T _{es} G _{es} A _e	PS	n/a	n/a	161
686892	GalNAc₃-10_{a-o} A _{es} T _{es} G _{es} ^m C _{es} A _{es} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} A _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} T _{es} G _{es} A _e	PS	GalNAc ₃ -10a	PO	161

The legend for Table 97 can be found in Example 74. The structure of GalNAc₃-10_a was shown in Example 46.

Table 98
Factor VII plasma protein levels

ISIS No.	Day	Dose (mg/kg)	Factor VII (% BL)
407935	0	n/a	100
	15	10	87
	22	n/a	92
	29	30	77
	36	n/a	46
	43	n/a	43
686892	0	3	100
	15	10	56
	22	n/a	29
	29	30	19
	36	n/a	15
	43	n/a	11

Example 92: Antisense inhibition in primary hepatocytes by antisense oligonucleotides targeting ApoC-III comprising a GalNAc₃ conjugate

Primary mouse hepatocytes were seeded in 96-well plates at 15,000 cells per well, and the oligonucleotides listed in Table 99, targeting mouse ApoC-III, were added at 0.46, 1.37, 4.12, or 12.35, 37.04, 111.11, or 333.33 nM or 1.00 μ M. After incubation with the oligonucleotides for 24 hours, the cells were lysed and total RNA was purified using RNeasy (Qiagen). ApoC-III mRNA levels were determined using real-time PCR and RIBOGREEN® RNA quantification reagent (Molecular Probes, Inc.) according to standard protocols. IC₅₀ values were determined using Prism 4 software (GraphPad). The results show that regardless of whether the cleavable moiety was a phosphodiester or a phosphodiester-linked deoxyadenosine, the oligonucleotides comprising a GalNAc conjugate were significantly more potent than the parent oligonucleotide lacking a conjugate.

Table 99

Inhibition of mouse APOC-III expression in mouse primary hepatocytes

ISIS No.	Sequence (5' to 3')	CM	IC ₅₀ (nM)	SEQ ID No.
440670	^m C _{es} A _{es} G _{es} ^m C _{es} T _{es} T _{ds} T _{ds} A _{ds} T _{ds} T _{ds} A _{ds} G _{ds} G _{ds} G _{ds} A _{ds} ^m C _{es} A _{es} G _{es} ^m C _{es} A _e	n/a	13.20	162
661180	^m C _{es} A _{es} G _{es} ^m C _{es} T _{es} T _{ds} T _{ds} A _{ds} T _{ds} T _{ds} A _{ds} G _{ds} G _{ds} G _{ds} A _{ds} ^m C _{es} A _{es} G _{es} ^m C _{es} A _{eo} A _{do} '-GalNAc ₃ -1 _a	A _d	1.40	163
680771	GalNAc ₃ -3 _{a-o} ' ^m C _{es} A _{es} G _{es} ^m C _{es} T _{es} T _{ds} T _{ds} A _{ds} T _{ds} T _{ds} A _{ds} G _{ds} G _{ds} G _{ds} A _{ds} ^m C _{es} A _{es} G _{es} ^m C _{es} A _e	PO	0.70	162
680772	GalNAc ₃ -7 _{a-o} ' ^m C _{es} A _{es} G _{es} ^m C _{es} T _{es} T _{ds} T _{ds} A _{ds} T _{ds} T _{ds} A _{ds} G _{ds} G _{ds} G _{ds} A _{ds} ^m C _{es} A _{es} G _{es} ^m C _{es} A _e	PO	1.70	162
680773	GalNAc ₃ -10 _{a-o} ' ^m C _{es} A _{es} G _{es} ^m C _{es} T _{es} T _{ds} T _{ds} A _{ds} T _{ds} T _{ds} A _{ds} G _{ds} G _{ds} G _{ds} A _{ds} ^m C _{es} A _{es} G _{es} ^m C _{es} A _e	PO	2.00	162
680774	GalNAc ₃ -13 _{a-o} ' ^m C _{es} A _{es} G _{es} ^m C _{es} T _{es} T _{ds} T _{ds} A _{ds} T _{ds} T _{ds} A _{ds} G _{ds} G _{ds} G _{ds} A _{ds} ^m C _{es} A _{es} G _{es} ^m C _{es} A _e	PO	1.50	162
681272	GalNAc ₃ -3 _{a-o} ' ^m C _{es} A _{eo} G _{eo} ^m C _{eo} T _{eo} T _{ds} T _{ds} A _{ds} T _{ds} T _{ds} A _{ds} G _{ds} G _{ds} G _{ds} A _{ds} ^m C _{eo} A _{eo} G _{es} ^m C _{es} A _e	PO	< 0.46	162
681273	GalNAc ₃ -3 _{a-o} 'A _{do} ^m C _{es} A _{es} G _{es} ^m C _{es} T _{es} T _{ds} T _{ds} A _{ds} T _{ds} T _{ds} A _{ds} G _{ds} G _{ds} G _{ds} A _{ds} ^m C _{es} A _{es} G _{es} ^m C _{es} A _e	A _d	1.10	164
683733	^m C _{es} A _{es} G _{es} ^m C _{es} T _{es} T _{ds} T _{ds} A _{ds} T _{ds} T _{ds} A _{ds} G _{ds} G _{ds} G _{ds} A _{ds} ^m C _{es} A _{es} G _{es} ^m C _{es} A _{eo} A _{do} '-GalNAc ₃ -19 _a	A _d	2.50	163

The structure of GalNAc₃-1_a was shown previously in Example 9, GalNAc₃-3_a was shown in Example 39, GalNAc₃-7_a was shown in Example 48, GalNAc₃-10_a was shown in Example 46, GalNAc₃-13_a was shown in Example 62, and GalNAc₃-19_a was shown in Example 70.

Example 93: Antisense inhibition *in vivo* by oligonucleotides targeting SRB-1 comprising mixed wings and a 5'-GalNAc₃ conjugate

The oligonucleotides listed in Table 100 were tested in a dose-dependent study for antisense inhibition of SRB-1 in mice.

Table 100
Modified ASOs targeting SRB-1

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
449093	T _{ks} T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _{ks} ^m C _k	n/a	n/a	165
699806	GalNAc₃-3_a-o , T _{ks} T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _{ks} ^m C _k	GalNAc ₃ -3 _a	PO	165
699807	GalNAc₃-7_a-o , T _{ks} T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _{ks} ^m C _k	GalNAc ₃ -7 _a	PO	165
699809	GalNAc₃-7_a-o , T _{ks} T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _e	GalNAc ₃ -7 _a	PO	165
699811	GalNAc₃-7_a-o , T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _{ks} ^m C _k	GalNAc ₃ -7 _a	PO	165
699813	GalNAc₃-7_a-o , T _{ks} T _{ds} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _{ds} ^m C _k	GalNAc ₃ -7 _a	PO	165
699815	GalNAc₃-7_a-o , T _{es} T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _{ks} ^m C _e	GalNAc ₃ -7 _a	PO	165

The structure of GalNAc₃-3_a was shown previously in Example 39, and the structure of GalNAc₃-7_a was shown previously in Example 48. Subscripts: “e” indicates 2'-MOE modified nucleoside; “d” indicates β-D-2'-deoxyribonucleoside; “k” indicates 6'-(S)-CH₃ bicyclic nucleoside (cEt); “s” indicates phosphorothioate internucleoside linkages (PS); “o” indicates phosphodiester internucleoside linkages (PO). Superscript “m” indicates 5-methylcytosines.

Treatment

Six to eight week old C57BL/6 mice (Jackson Laboratory, Bar Harbor, ME) were injected subcutaneously once at the dosage shown below with an oligonucleotide listed in Table 100 or with saline. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration. Liver SRB-1 mRNA levels were measured using real-time PCR. SRB-1 mRNA levels were normalized to cyclophilin mRNA levels according to standard protocols. The results are presented as the average percent of SRB-1 mRNA levels for each treatment group relative to the saline control group. As illustrated in Table 101, treatment with antisense oligonucleotides lowered SRB-1 mRNA levels in a dose-dependent manner, and the gapmer oligonucleotides comprising a GalNAc conjugate and having wings that were either full cEt or mixed sugar modifications were significantly more potent than the parent oligonucleotide lacking a conjugate and comprising full cEt modified wings.

Body weights, liver transaminases, total bilirubin, and BUN were also measured, and the average values for each treatment group are shown in Table 101. Body weight is shown as the average percent body weight relative to the baseline body weight (% BL) measured just prior to the oligonucleotide dose.

Table 101

SRB-1 mRNA, ALT, AST, BUN, and total bilirubin levels and body weights

ISIS No.	Dosage (mg/kg)	SRB-1 mRNA (% PBS)	ALT (U/L)	AST (U/L)	Bil	BUN	Body weight (% BL)
PBS	n/a	100	31	84	0.15	28	102
449093	1	111	18	48	0.17	31	104
	3	94	20	43	0.15	26	103
	10	36	19	50	0.12	29	104
699806	0.1	114	23	58	0.13	26	107
	0.3	59	21	45	0.12	27	108
	1	25	30	61	0.12	30	104
699807	0.1	121	19	41	0.14	25	100
	0.3	73	23	56	0.13	26	105
	1	24	22	69	0.14	25	102
699809	0.1	125	23	57	0.14	26	104
	0.3	70	20	49	0.10	25	105
	1	33	34	62	0.17	25	107
699811	0.1	123	48	77	0.14	24	106
	0.3	94	20	45	0.13	25	101
	1	66	57	104	0.14	24	107
699813	0.1	95	20	58	0.13	28	104
	0.3	98	22	61	0.17	28	105
	1	49	19	47	0.11	27	106
699815	0.1	93	30	79	0.17	25	105
	0.3	64	30	61	0.12	26	105
	1	24	18	41	0.14	25	106

Example 94: Antisense inhibition *in vivo* by oligonucleotides targeting SRB-1 comprising 2'-sugar modifications and a 5'-GalNAc₃ conjugate

The oligonucleotides listed in Table 102 were tested in a dose-dependent study for antisense inhibition of SRB-1 in mice.

Table 102

Modified ASOs targeting SRB-1

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
353382	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	n/a	n/a	143
700989	G _{ms} C _{ms} U _{ms} U _{ms} C _{ms} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} U _{ms} C _{ms} C _{ms} U _{ms} U _m	n/a	n/a	166
666904	GalNAc₃-3a-o ·G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	GalNAc ₃ -3a	PO	143
700991	GalNAc₃-7a-o ·G _{ms} C _{ms} U _{ms} U _{ms} C _{ms} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} U _{ms} C _{ms} C _{ms} U _{ms} U _m	GalNAc ₃ -7a	PO	166

Subscript "m" indicates a 2'-O-methyl modified nucleoside. See Example 74 for complete table legend. The structure of GalNAc₃-3_a was shown previously in Example 39, and the structure of GalNAc₃-7_a was shown previously in Example 48.

Treatment

The study was completed using the protocol described in Example 93. Results are shown in Table 103 below and show that both the 2'-MOE and 2'-OMe modified oligonucleotides comprising a GalNAc conjugate were significantly more potent than the respective parent oligonucleotides lacking a conjugate. The results of the body weights, liver transaminases, total bilirubin, and BUN measurements indicated that the compounds were all well tolerated.

Table 103
SRB-1 mRNA

ISIS No.	Dosage (mg/kg)	SRB-1 mRNA (% PBS)
PBS	n/a	100
353382	5	116
	15	58
	45	27
700989	5	120
	15	92
	45	46
666904	1	98
	3	45
	10	17
700991	1	118
	3	63
	10	14

Example 95: Antisense inhibition *in vivo* by oligonucleotides targeting SRB-1 comprising bicyclic nucleosides and a 5'-GalNAc₃ conjugate

The oligonucleotides listed in Table 104 were tested in a dose-dependent study for antisense inhibition of SRB-1 in mice.

Table 104
Modified ASOs targeting SRB-1

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No
440762	T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _k	n/a	n/a	137
666905	GalNAc₃-3_a-o' T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _k	GalNAc ₃ -3 _a	PO	137
699782	GalNAc₃-7_a-o' T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _k	GalNAc ₃ -7 _a	PO	137
699783	GalNAc₃-3_a-o' T _{ls} ^m C _{ls} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ls} ^m C _l	GalNAc ₃ -3 _a	PO	137
653621	T _{ls} ^m C _{ls} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ls} ^m C _{lo} A_{do}'-GalNAc₃-1_a	GalNAc ₃ -1 _a	A _d	138
439879	T _{gs} ^m C _{gs} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _d G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{gs} ^m C _g	n/a	n/a	137
699789	GalNAc₃-3_a-o' T _{gs} ^m C _{gs} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _d G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{gs} ^m C _g	GalNAc ₃ -3 _a	PO	137

Subscript "g" indicates a fluoro-HNA nucleoside, subscript "l" indicates a locked nucleoside comprising a 2'-O-CH₂-4' bridge. See the Example 74 table legend for other abbreviations. The structure of GalNAc₃-1_a was

shown previously in Example 9, the structure of GalNAc₃-3_a was shown previously in Example 39, and the structure of GalNAc₃-7a was shown previously in Example 48.

Treatment

5 The study was completed using the protocol described in Example 93. Results are shown in Table 105 below and show that oligonucleotides comprising a GalNAc conjugate and various bicyclic nucleoside modifications were significantly more potent than the parent oligonucleotide lacking a conjugate and comprising bicyclic nucleoside modifications. Furthermore, the oligonucleotide comprising a GalNAc conjugate and fluoro-HNA modifications was significantly more potent than the parent lacking a conjugate and comprising fluoro-HNA modifications. The results of the body weights, liver transaminases, total bilirubin, and BUN measurements indicated that the compounds were all well tolerated.

Table 105
SRB-1 mRNA, ALT, AST, BUN, and total bilirubin levels and body weights

ISIS No.	Dosage (mg/kg)	SRB-1 mRNA (% PBS)
PBS	n/a	100
440762	1	104
	3	65
	10	35
666905	0.1	105
	0.3	56
	1	18
699782	0.1	93
	0.3	63
	1	15
699783	0.1	105
	0.3	53
	1	12
653621	0.1	109
	0.3	82
	1	27
439879	1	96
	3	77
	10	37
699789	0.1	82
	0.3	69
	1	26

15 **Example 96: Plasma protein binding of antisense oligonucleotides comprising a GalNAc₃ conjugate group**

Oligonucleotides listed in Table 70 targeting ApoC-III and oligonucleotides in Table 106 targeting Apo(a) were tested in an ultra-filtration assay in order to assess plasma protein binding.

Table 106

Modified oligonucleotides targeting Apo(a)

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No
494372	$T_{es}G_{es}^mC_{es}T_{es}^mC_{es}^mC_{ds}G_{ds}T_{ds}T_{ds}G_{ds}G_{ds}T_{ds}G_{ds}^mC_{ds}T_{ds}T_{es}G_{es}T_{es}$ $T_{es}^mC_e$	n/a	n/a	58
693401	$T_{es}G_{eo}^mC_{eo}T_{eo}^mC_{eo}^mC_{ds}G_{ds}T_{ds}T_{ds}G_{ds}G_{ds}T_{ds}G_{ds}^mC_{ds}T_{ds}T_{eo}G_{eo}T_{es}$ $T_{es}^mC_e$	n/a	n/a	58
681251	GalNAc₃-7_a-o' $T_{es}G_{es}^mC_{es}T_{es}^mC_{es}^mC_{ds}G_{ds}T_{ds}T_{ds}G_{ds}G_{ds}T_{ds}G_{ds}^mC_{ds}$ $T_{ds}T_{es}G_{es}T_{es}T_{es}^mC_e$	GalNAc ₃ -7 _a	PO	58
681257	GalNAc₃-7_a-o' $T_{es}G_{eo}^mC_{eo}T_{eo}^mC_{eo}^mC_{ds}G_{ds}T_{ds}T_{ds}G_{ds}G_{ds}T_{ds}G_{ds}^mC_{ds}$ $T_{ds}T_{eo}G_{eo}T_{es}T_{es}^mC_e$	GalNAc ₃ -7 _a	PO	58

See the Example 74 for table legend. The structure of GalNAc₃-7a was shown previously in Example 48.

Ultrafree-MC ultrafiltration units (30,000 NMWL, low-binding regenerated cellulose membrane, Millipore, Bedford, MA) were pre-conditioned with 300 μ L of 0.5% Tween 80 and centrifuged at 2000 g for 10 minutes, then with 300 μ L of a 300 μ g/mL solution of a control oligonucleotide in H₂O and centrifuged at 2000 g for 16 minutes. In order to assess non-specific binding to the filters of each test oligonucleotide from Tables 70 and 106 to be used in the studies, 300 μ L of a 250 ng/mL solution of oligonucleotide in H₂O at pH 7.4 was placed in the pre-conditioned filters and centrifuged at 2000 g for 16 minutes. The unfiltered and filtered samples were analyzed by an ELISA assay to determine the oligonucleotide concentrations. Three replicates were used to obtain an average concentration for each sample. The average concentration of the filtered sample relative to the unfiltered sample is used to determine the percent of oligonucleotide that is recovered through the filter in the absence of plasma (% recovery).

Frozen whole plasma samples collected in K3-EDTA from normal, drug-free human volunteers, cynomolgus monkeys, and CD-1 mice, were purchased from Bioreclamation LLC (Westbury, NY). The test oligonucleotides were added to 1.2 mL aliquots of plasma at two concentrations (5 and 150 μ g/mL). An aliquot (300 μ L) of each spiked plasma sample was placed in a pre-conditioned filter unit and incubated at 37°C for 30 minutes, immediately followed by centrifugation at 2000 g for 16 minutes. Aliquots of filtered and unfiltered spiked plasma samples were analyzed by an ELISA to determine the oligonucleotide concentration in each sample. Three replicates per concentration were used to determine the average percentage of bound and unbound oligonucleotide in each sample. The average concentration of the filtered sample relative to the concentration of the unfiltered sample is used to determine the percent of oligonucleotide in the plasma that is not bound to plasma proteins (% unbound). The final unbound oligonucleotide values are corrected for non-specific binding by dividing the % unbound by the % recovery for each oligonucleotide. The final % bound oligonucleotide values are determined by subtracting the final % unbound values from 100. The results are shown in Table 107 for the two concentrations of oligonucleotide tested (5 and 150 μ g/mL) in each species of plasma. The results show that GalNAc conjugate groups do not have a significant impact on plasma protein binding. Furthermore, oligonucleotides with full PS

internucleoside linkages and mixed PO/PS linkages both bind plasma proteins, and those with full PS linkages bind plasma proteins to a somewhat greater extent than those with mixed PO/PS linkages.

Table 107
Percent of modified oligonucleotide bound to plasma proteins

ISIS No.	Human plasma		Monkey plasma		Mouse plasma	
	5 µg/mL	150 µg/mL	5 µg/mL	150 µg/mL	5 µg/mL	150 µg/mL
304801	99.2	98.0	99.8	99.5	98.1	97.2
663083	97.8	90.9	99.3	99.3	96.5	93.0
674450	96.2	97.0	98.6	94.4	94.6	89.3
494372	94.1	89.3	98.9	97.5	97.2	93.6
693401	93.6	89.9	96.7	92.0	94.6	90.2
681251	95.4	93.9	99.1	98.2	97.8	96.1
681257	93.4	90.5	97.6	93.7	95.6	92.7

5

Example 97: Modified oligonucleotides targeting TTR comprising a GalNAc₃ conjugate group

The oligonucleotides shown in Table 108 comprising a GalNAc conjugate were designed to target TTR.

Table 108

Modified oligonucleotides targeting TTR

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No
666941	GalNAc₃-3_{a-o}'A_{do} T _{es} ^m C _{es} T _{es} T _{es} G _{es} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{es} T _{es} ^m C _{es} ^m C _{es} ^m C _e	GalNAc ₃ -3	A _d	160
666942	T _{es} ^m C _{eo} T _{eo} T _{eo} G _{eo} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{eo} T _{eo} ^m C _{es} ^m C _{es} ^m C _{eo} A_{do}'-GalNAc₃-3_a	GalNAc ₃ -1	A _d	157
682876	GalNAc₃-3_{a-o}'T_{es} ^m C _{es} T _{es} T _{es} G _{es} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{es} T _{es} ^m C _{es} ^m C _{es} ^m C _e	GalNAc ₃ -3	PO	156
682877	GalNAc₃-7_{a-o}'T_{es} ^m C _{es} T _{es} T _{es} G _{es} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{es} T _{es} ^m C _{es} ^m C _{es} ^m C _e	GalNAc ₃ -7	PO	156
682878	GalNAc₃-10_{a-o}'T_{es} ^m C _{es} T _{es} T _{es} G _{es} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{es} T _{es} ^m C _{es} ^m C _{es} ^m C _e	GalNAc ₃ -10	PO	156
682879	GalNAc₃-13_{a-o}'T_{es} ^m C _{es} T _{es} T _{es} G _{es} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{es} T _{es} ^m C _{es} ^m C _{es} ^m C _e	GalNAc ₃ -13	PO	156
682880	GalNAc₃-7_{a-o}'A_{do} T _{es} ^m C _{es} T _{es} T _{es} G _{es} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{es} T _{es} ^m C _{es} ^m C _{es} ^m C _e	GalNAc ₃ -7	A _d	160
682881	GalNAc₃-10_{a-o}'A_{do} T _{es} ^m C _{es} T _{es} T _{es} G _{es} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{es} T _{es} ^m C _{es} ^m C _{es} ^m C _e	GalNAc ₃ -10	A _d	160
682882	GalNAc₃-13_{a-o}'A_{do} T _{es} ^m C _{es} T _{es} T _{es} G _{es} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{es} T _{es} ^m C _{es} ^m C _{es} ^m C _e	GalNAc ₃ -13	A _d	160
684056	T _{es} ^m C _{es} T _{es} T _{es} G _{es} G _{ds} T _{ds} T _{ds} A _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} A _{ds} A _{es} T _{es} ^m C _{es} ^m C _{es} ^m C _{eo} A_{do}'-GalNAc₃-19_a	GalNAc ₃ -19	A _d	157

The legend for Table 108 can be found in Example 74. The structure of GalNAc₃-1 was shown in Example 9. The structure of GalNAc₃-3_a was shown in Example 39. The structure of GalNAc₃-7_a was shown in Example 48. The structure of GalNAc₃-10_a was shown in Example 46. The structure of GalNAc₃-13_a was shown in Example 62. The structure of GalNAc₃-19_a was shown in Example 70.

Example 98: Evaluation of pro-inflammatory effects of oligonucleotides comprising a GalNAc conjugate in hPMBC assay

The oligonucleotides listed in Table 109 and were tested for pro-inflammatory effects in an hPMBC assay as described in Examples 23 and 24. (See Tables 30, 83, 95, and 108 for descriptions of the oligonucleotides.) ISIS 353512 is a high responder used as a positive control, and the other oligonucleotides are described in Tables 83, 95, and 108. The results shown in Table 109 were obtained using blood from one volunteer donor. The results show that the oligonucleotides comprising mixed PO/PS internucleoside linkages produced significantly lower pro-inflammatory responses compared to the same oligonucleotides having full PS linkages. Furthermore, the GalNAc conjugate group did not have a significant effect in this assay.

Table 109

ISIS No.	E _{max} /EC ₅₀	GalNAc ₃ cluster	Linkages	CM
353512	3630	n/a	PS	n/a
420915	802	n/a	PS	n/a
682881	1311	GalNAc ₃ -10	PS	A _d
682888	0.26	GalNAc ₃ -10	PO/PS	A _d
684057	1.03	GalNAc ₃ -19	PO/PS	A _d

Example 99: Binding affinities of oligonucleotides comprising a GalNAc conjugate for the asialoglycoprotein receptor

The binding affinities of the oligonucleotides listed in Table 110 (see Table 76 for descriptions of the oligonucleotides) for the asialoglycoprotein receptor were tested in a competitive receptor binding assay. The competitor ligand, α 1-acid glycoprotein (AGP), was incubated in 50 mM sodium acetate buffer (pH 5) with 1 U neuraminidase-agarose for 16 hours at 37°C, and > 90% desialylation was confirmed by either sialic acid assay or size exclusion chromatography (SEC). Iodine monochloride was used to iodinate the AGP according to the procedure by Atsma et al. (see J Lipid Res. 1991 Jan; 32(1):173-81.) In this method, desialylated α 1-acid glycoprotein (de-AGP) was added to 10 mM iodine chloride, Na¹²⁵I, and 1 M glycine in 0.25 M NaOH. After incubation for 10 minutes at room temperature, ¹²⁵I-labeled de-AGP was separated from free ¹²⁵I by concentrating the mixture twice utilizing a 3 KDMWCO spin column. The protein was tested for labeling efficiency and purity on a HPLC system equipped with an Agilent SEC-3 column (7.8x300mm) and a β -RAM counter. Competition experiments utilizing ¹²⁵I-labeled de-AGP and various GalNAc-cluster containing ASOs were performed as follows. Human HepG2 cells (10⁶ cells/ml) were plated on 6-well plates in 2 ml of appropriate growth media. MEM media supplemented with 10% fetal bovine serum (FBS), 2 mM L-Glutamine and 10mM HEPES was used. Cells were incubated 16-20 hours @ 37°C with 5% and 10% CO₂ respectively. Cells were washed with media without FBS prior to the experiment. Cells were incubated for 30

min @37°C with 1ml competition mix containing appropriate growth media with 2% FBS, 10^{-8} M ^{125}I - labeled de-AGP and GalNAc-cluster containing ASOs at concentrations ranging from 10^{-11} to 10^{-5} M. Non-specific binding was determined in the presence of 10^{-2} M GalNAc sugar. Cells were washed twice with media without FBS to remove unbound ^{125}I -labeled de-AGP and competitor GalNAc ASO. Cells were lysed using Qiagen's RLT buffer containing 1% β -mercaptoethanol. Lysates were transferred to round bottom assay tubes after a brief 10 min freeze/thaw cycle and assayed on a γ -counter. Non-specific binding was subtracted before dividing ^{125}I protein counts by the value of the lowest GalNAc-ASO concentration counts. The inhibition curves were fitted according to a single site competition binding equation using a nonlinear regression algorithm to calculate the binding affinities (K_D 's).

The results in Table 110 were obtained from experiments performed on five different days. Results for oligonucleotides marked with superscript "a" are the average of experiments run on two different days. The results show that the oligonucleotides comprising a GalNAc conjugate group on the 5'-end bound the asialoglycoprotein receptor on human HepG2 cells with 1.5 to 16-fold greater affinity than the oligonucleotides comprising a GalNAc conjugate group on the 3'-end.

Table 110
Asialoglycoprotein receptor binding assay results

ISIS No.	GalNAc conjugate	Oligonucleotide end to which GalNAc conjugate is attached	K_D (nM)
661161 ^a	GalNAc ₃ -3	5'	3.7
666881 ^a	GalNAc ₃ -10	5'	7.6
666981	GalNAc ₃ -7	5'	6.0
670061	GalNAc ₃ -13	5'	7.4
655861 ^a	GalNAc ₃ -1	3'	11.6
677841 ^a	GalNAc ₃ -19	3'	60.8

Example 100: Antisense inhibition *in vivo* by oligonucleotides comprising a GalNAc conjugate group targeting Apo(a) *in vivo*

The oligonucleotides listed in Table 111a below were tested in a single dose study for duration of action in mice.

Table 111a
Modified ASOs targeting APO(a)

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
681251	GalNAc ₃ -7 _a -o'-T _{es} G _{es} ^m C _{es} T _{es} ^m C _{es} ^m C _{ds} G _{ds} T _{ds} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{es} G _{es} T _{es} T _{es} ^m C _e	GalNAc ₃ -7a	PO	58
681257	GalNAc ₃ -7 _a -o'-T _{es} G _{eo} ^m C _{eo} T _{eo} ^m C _{eo} ^m C _{ds} G _{ds} T _{ds} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{eo} G _{eo} T _{es} T _{es} ^m C _e	GalNAc ₃ -7a	PO	58

The structure of GalNAc₃-7_a was shown in Example 48.

Treatment

Female transgenic mice that express human Apo(a) were each injected subcutaneously once per week, for a total of 6 doses, with an oligonucleotide and dosage listed in Table 111b or with PBS. Each treatment group consisted of 3 animals. Blood was drawn the day before dosing to determine baseline levels of Apo(a) protein in plasma and at 72 hours, 1 week, and 2 weeks following the first dose. Additional blood draws will occur at 3 weeks, 4 weeks, 5 weeks, and 6 weeks following the first dose. Plasma Apo(a) protein levels were measured using an ELISA. The results in Table 111b are presented as the average percent of plasma Apo(a) protein levels for each treatment group, normalized to baseline levels (% BL). The results show that the oligonucleotides comprising a GalNAc conjugate group exhibited potent reduction in Apo(a) expression. This potent effect was observed for the oligonucleotide that comprises full PS internucleoside linkages and the oligonucleotide that comprises mixed PO and PS linkages.

Table 111b
Apo(a) plasma protein levels

ISIS No.	Dosage (mg/kg)	Apo(a) at 72 hours (% BL)	Apo(a) at 1 week (% BL)	Apo(a) at 3 weeks (% BL)
PBS	n/a	116	104	107
681251	0.3	97	108	93
	1.0	85	77	57
	3.0	54	49	11
	10.0	23	15	4
681257	0.3	114	138	104
	1.0	91	98	54
	3.0	69	40	6
	10.0	30	21	4

Example 101: Antisense inhibition by oligonucleotides comprising a GalNAc cluster linked via a stable moiety

The oligonucleotides listed in Table 112 were tested for inhibition of mouse APOC-III expression *in vivo*. C57Bl/6 mice were each injected subcutaneously once with an oligonucleotide listed in Table 112 or with PBS. Each treatment group consisted of 4 animals. Each mouse treated with ISIS 440670 received a dose of 2, 6, 20, or 60 mg/kg. Each mouse treated with ISIS 680772 or 696847 received 0.6, 2, 6, or 20 mg/kg. The GalNAc conjugate group of ISIS 696847 is linked via a stable moiety, a phosphorothioate linkage instead of a readily cleavable phosphodiester containing linkage. The animals were sacrificed 72 hours after the dose. Liver APOC-III mRNA levels were measured using real-time PCR. APOC-III mRNA levels were normalized to cyclophilin mRNA levels according to standard protocols. The results are presented in Table 112 as the average percent of APOC-III mRNA levels for each treatment group relative to the saline control group. The results show that the oligonucleotides comprising a GalNAc conjugate group were significantly more potent than the oligonucleotide lacking a conjugate group. Furthermore, the

oligonucleotide comprising a GalNAc conjugate group linked to the oligonucleotide via a cleavable moiety (ISIS 680772) was even more potent than the oligonucleotide comprising a GalNAc conjugate group linked to the oligonucleotide via a stable moiety (ISIS 696847).

Table 112

Modified oligonucleotides targeting mouse APOC-III

ISIS No.	Sequences (5' to 3')	CM	Dosage (mg/kg)	APOC-III mRNA (% PBS)	SEQ ID No.
440670	${}^m\text{C}_{\text{es}}\text{A}_{\text{es}}\text{G}_{\text{es}}{}^m\text{C}_{\text{es}}\text{T}_{\text{es}}\text{T}_{\text{ds}}\text{T}_{\text{ds}}\text{A}_{\text{ds}}\text{T}_{\text{ds}}\text{T}_{\text{ds}}\text{A}_{\text{ds}}$ $\text{G}_{\text{ds}}\text{G}_{\text{ds}}\text{G}_{\text{ds}}\text{A}_{\text{ds}}{}^m\text{C}_{\text{es}}\text{A}_{\text{es}}\text{G}_{\text{es}}{}^m\text{C}_{\text{es}}\text{A}_{\text{e}}$	n/a	2	92	162
			6	86	
			20	59	
			60	37	
680772	GalNAc₃-7_{a-o} , ${}^m\text{C}_{\text{es}}\text{A}_{\text{es}}\text{G}_{\text{es}}{}^m\text{C}_{\text{es}}\text{T}_{\text{es}}\text{T}_{\text{ds}}\text{T}_{\text{ds}}\text{A}_{\text{ds}}$ $\text{T}_{\text{ds}}\text{T}_{\text{ds}}\text{A}_{\text{ds}}\text{G}_{\text{ds}}\text{G}_{\text{ds}}\text{G}_{\text{ds}}\text{A}_{\text{ds}}{}^m\text{C}_{\text{es}}\text{A}_{\text{es}}\text{G}_{\text{es}}{}^m\text{C}_{\text{es}}\text{A}_{\text{e}}$	PO	0.6	79	162
			2	58	
			6	31	
			20	13	
696847	GalNAc₃-7_{a-s} , ${}^m\text{C}_{\text{es}}\text{A}_{\text{es}}\text{G}_{\text{es}}{}^m\text{C}_{\text{es}}\text{T}_{\text{es}}\text{T}_{\text{ds}}\text{T}_{\text{ds}}\text{A}_{\text{ds}}\text{T}_{\text{ds}}$ $\text{T}_{\text{ds}}\text{A}_{\text{ds}}\text{G}_{\text{ds}}\text{G}_{\text{ds}}\text{G}_{\text{ds}}\text{A}_{\text{ds}}{}^m\text{C}_{\text{es}}\text{A}_{\text{es}}\text{G}_{\text{es}}{}^m\text{C}_{\text{es}}\text{A}_{\text{e}}$	n/a (PS)	0.6	83	162
			2	73	
			6	40	
			20	28	

The structure of GalNAc₃-7_a was shown in Example 48.

Example 102: Distribution in liver of antisense oligonucleotides comprising a GalNAc conjugate

The liver distribution of ISIS 353382 (see Table 36) that does not comprise a GalNAc conjugate and ISIS 655861 (see Table 36) that does comprise a GalNAc conjugate was evaluated. Male balb/c mice were subcutaneously injected once with ISIS 353382 or 655861 at a dosage listed in Table 113. Each treatment group consisted of 3 animals except for the 18 mg/kg group for ISIS 655861, which consisted of 2 animals. The animals were sacrificed 48 hours following the dose to determine the liver distribution of the oligonucleotides. In order to measure the number of antisense oligonucleotide molecules per cell, a Ruthenium (II) tris-bipyridine tag (MSD TAG, Meso Scale Discovery) was conjugated to an oligonucleotide probe used to detect the antisense oligonucleotides. The results presented in Table 113 are the average concentrations of oligonucleotide for each treatment group in units of millions of oligonucleotide molecules per cell. The results show that at equivalent doses, the oligonucleotide comprising a GalNAc conjugate was present at higher concentrations in the total liver and in hepatocytes than the oligonucleotide that does not comprise a GalNAc conjugate. Furthermore, the oligonucleotide comprising a GalNAc conjugate was present at lower concentrations in non-parenchymal liver cells than the oligonucleotide that does not comprise a GalNAc conjugate. And while the concentrations of ISIS 655861 in hepatocytes and non-parenchymal liver cells were similar per cell, the liver is approximately 80% hepatocytes by volume. Thus, the majority of the ISIS 655861 oligonucleotide that was present in the liver was found in hepatocytes, whereas the majority of the ISIS 353382 oligonucleotide that was present in the liver was found in non-parenchymal liver cells.

Table 113

ISIS No.	Dosage (mg/kg)	Concentration in whole liver (molecules*10 ⁶ per cell)	Concentration in hepatocytes (molecules*10 ⁶ per cell)	Concentration in non-parenchymal liver cells (molecules*10 ⁶ per cell)
353382	3	9.7	1.2	37.2
	10	17.3	4.5	34.0
	20	23.6	6.6	65.6
	30	29.1	11.7	80.0
	60	73.4	14.8	98.0
	90	89.6	18.5	119.9
655861	0.5	2.6	2.9	3.2
	1	6.2	7.0	8.8
	3	19.1	25.1	28.5
	6	44.1	48.7	55.0
	18	76.6	82.3	77.1

Example 103: Duration of action *in vivo* of oligonucleotides targeting APOC-III comprising a GalNAc₃ conjugate

The oligonucleotides listed in Table 114 below were tested in a single dose study for duration of action in mice.

Table 114

Modified ASOs targeting APOC-III

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
304801	A _{es} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{es} T _{es} T _{es} A _{es} T _e	n/a	n/a	135
663084	GalNAc₃-3_a-o ·A _{d0} A _{es} G _{eo} ^m C _{eo} T _{eo} T _{eo} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{eo} T _{eo} T _{es} A _{es} T _e	GalNAc ₃ -3 _a	A _d	151
679241	A _{es} G _{eo} ^m C _{eo} T _{eo} T _{eo} ^m C _{ds} T _{ds} T _{ds} G _{ds} T _{ds} ^m C _{ds} ^m C _{ds} A _{ds} G _{ds} ^m C _{ds} T _{eo} T _{eo} T _{es} A _{es} T _{eo} A_{d0}-GalNAc₃-19_a	GalNAc ₃ -19 _a	A _d	136

The structure of GalNAc₃-3_a was shown in Example 39, and GalNAc₃-19_a was shown in Example 70.

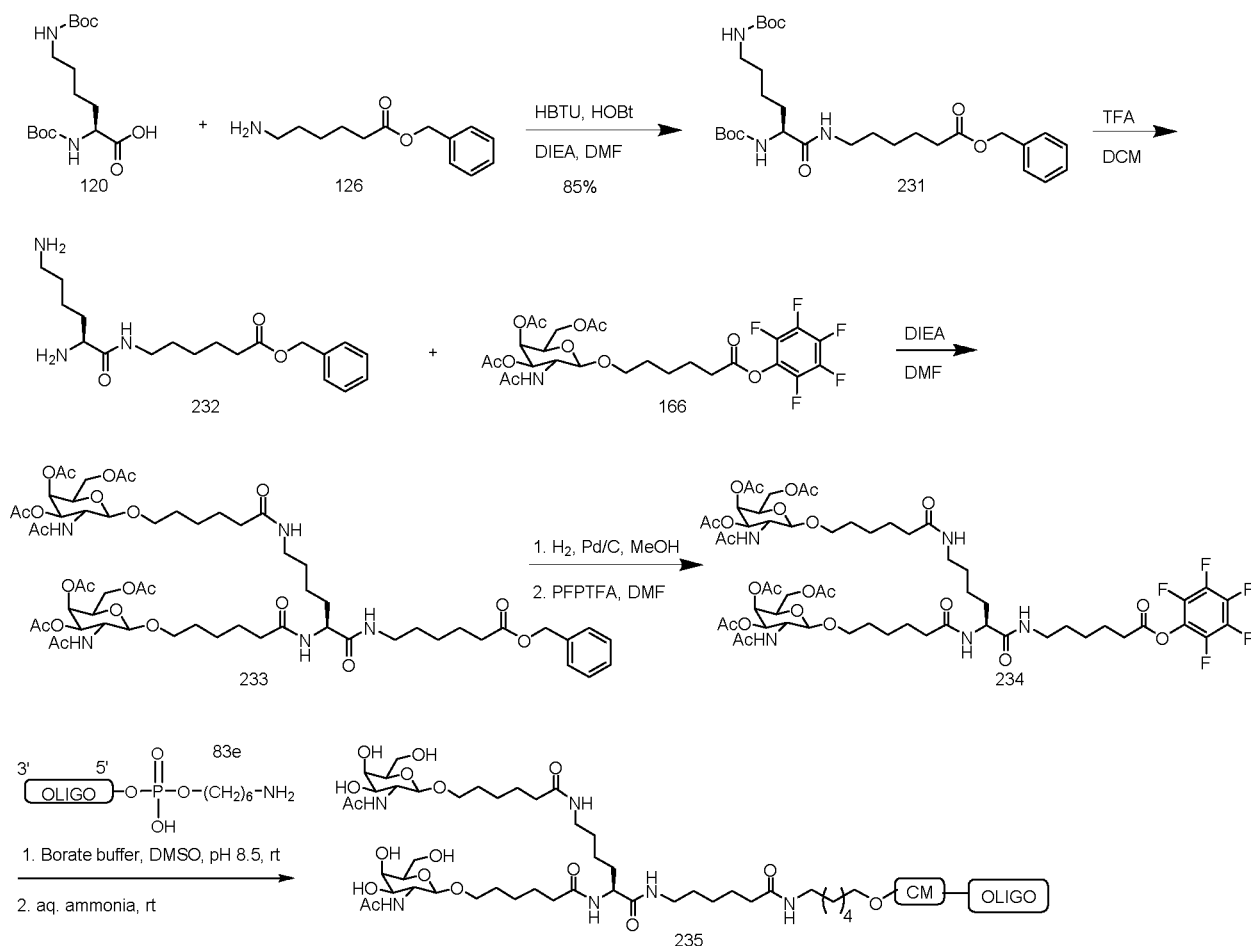
Treatment

Female transgenic mice that express human APOC-III were each injected subcutaneously once with an oligonucleotide listed in Table 114 or with PBS. Each treatment group consisted of 3 animals. Blood was drawn before dosing to determine baseline and at 3, 7, 14, 21, 28, 35, and 42 days following the dose. Plasma triglyceride and APOC-III protein levels were measured as described in Example 20. The results in Table 115 are presented as the average percent of plasma triglyceride and APOC-III levels for each treatment group, normalized to baseline levels. A comparison of the results in Table 71 of example 79 with the results in Table 115 below show that oligonucleotides comprising a mixture of phosphodiester and phosphorothioate

internucleoside linkages exhibited increased duration of action than equivalent oligonucleotides comprising only phosphorothioate internucleoside linkages.

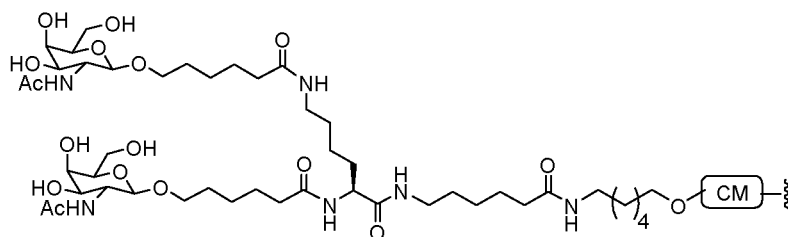
Table 115
Plasma triglyceride and APOC-III protein levels in transgenic mice

ISIS No.	Dosage (mg/kg)	Time point (days post-dose)	Triglycerides (% baseline)	APOC-III protein (% baseline)	GalNAc ₃ Cluster	CM
PBS	n/a	3	96	101	n/a	n/a
		7	88	98		
		14	91	103		
		21	69	92		
		28	83	81		
		35	65	86		
		42	72	88		
304801	30	3	42	46	n/a	n/a
		7	42	51		
		14	59	69		
		21	67	81		
		28	79	76		
		35	72	95		
		42	82	92		
663084	10	3	35	28	GalNAc ₃ -3a	A _d
		7	23	24		
		14	23	26		
		21	23	29		
		28	30	22		
		35	32	36		
		42	37	47		
679241	10	3	38	30	GalNAc ₃ -19a	A _d
		7	31	28		
		14	30	22		
		21	36	34		
		28	48	34		
		35	50	45		
		42	72	64		

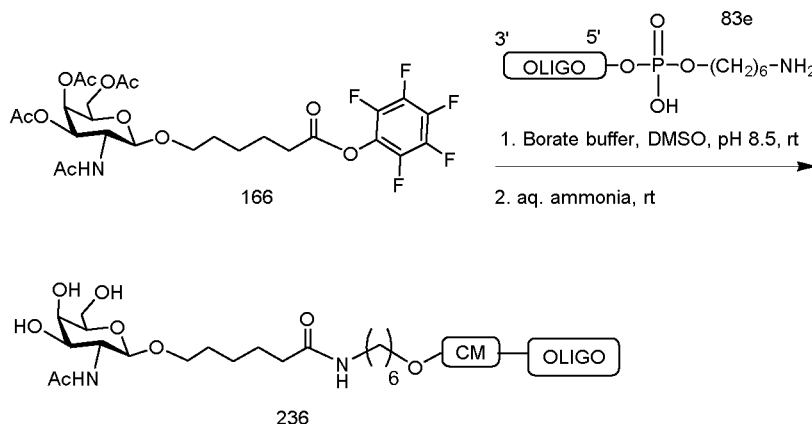
Example 104: Synthesis of oligonucleotides comprising a 5'-GalNAc₂ conjugate

Compound 120 is commercially available, and the synthesis of compound 126 is described in Example 49. Compound 120 (1 g, 2.89 mmol), HBTU (0.39 g, 2.89 mmol), and HOBT (1.64 g, 4.33 mmol) were dissolved in DMF (10 mL) and *N,N*-diisopropylethylamine (1.75 mL, 10.1 mmol) were added. After about 5 min, aminohexanoic acid benzyl ester (1.36 g, 3.46 mmol) was added to the reaction. After 3h, the reaction mixture was poured into 100 mL of 1 M NaHSO₄ and extracted with 2 x 50 mL ethyl acetate. Organic layers were combined and washed with 3 x 40 mL sat NaHCO₃ and 2 x brine, dried with Na₂SO₄, filtered and concentrated. The product was purified by silica gel column chromatography (DCM:EA:Hex, 1:1:1) to yield compound 231. LCMS and NMR were consistent with the structure. Compound 231 (1.34 g, 2.438 mmol) was dissolved in dichloromethane (10 mL) and trifluoroacetic acid (10 mL) was added. After stirring at room temperature for 2h, the reaction mixture was concentrated under reduced pressure and co-evaporated with toluene (3 x 10 mL). The residue was dried under reduced pressure to yield compound 232 as the trifluoroacetate salt. The synthesis of compound 166 is described in Example 54. Compound 166 (3.39 g, 5.40 mmol) was dissolved in DMF (3 mL). A solution of compound 232 (1.3 g, 2.25 mmol) was dissolved in DMF (3 mL) and *N,N*-diisopropylethylamine (1.55 mL) was added. The reaction was stirred at room temperature for 30 minutes, then poured into water (80 mL) and the aqueous layer was extracted with EtOAc (2x100 mL). The organic phase was separated and washed with sat. aqueous NaHCO₃ (3 x 80 mL), 1

M NaHSO₄ (3 x 80 mL) and brine (2 x 80 mL), then dried (Na₂SO₄), filtered, and concentrated. The residue was purified by silica gel column chromatography to yield compound 233. LCMS and NMR were consistent with the structure. Compound 233 (0.59 g, 0.48 mmol) was dissolved in methanol (2.2 mL) and ethyl acetate (2.2 mL). Palladium on carbon (10 wt% Pd/C, wet, 0.07 g) was added, and the reaction mixture was stirred under hydrogen atmosphere for 3 h. The reaction mixture was filtered through a pad of Celite and concentrated to yield the carboxylic acid. The carboxylic acid (1.32 g, 1.15 mmol, cluster free acid) was dissolved in DMF (3.2 mL). To this *N,N*-diisopropylethylamine (0.3 mL, 1.73 mmol) and PFPTFA (0.30 mL, 1.73 mmol) were added. After 30 min stirring at room temperature the reaction mixture was poured into water (40 mL) and extracted with EtOAc (2 x 50 mL). A standard work-up was completed as described above to yield compound 234. LCMS and NMR were consistent with the structure. Oligonucleotide 235 was prepared using the general procedure described in Example 46. The GalNAc₂ cluster portion (GalNAc₂-24_a) of the conjugate group GalNAc₂-24 can be combined with any cleavable moiety present on the oligonucleotide to provide a variety of conjugate groups. The structure of GalNAc₂-24 (GalNAc₂-24_a-CM) is shown below:

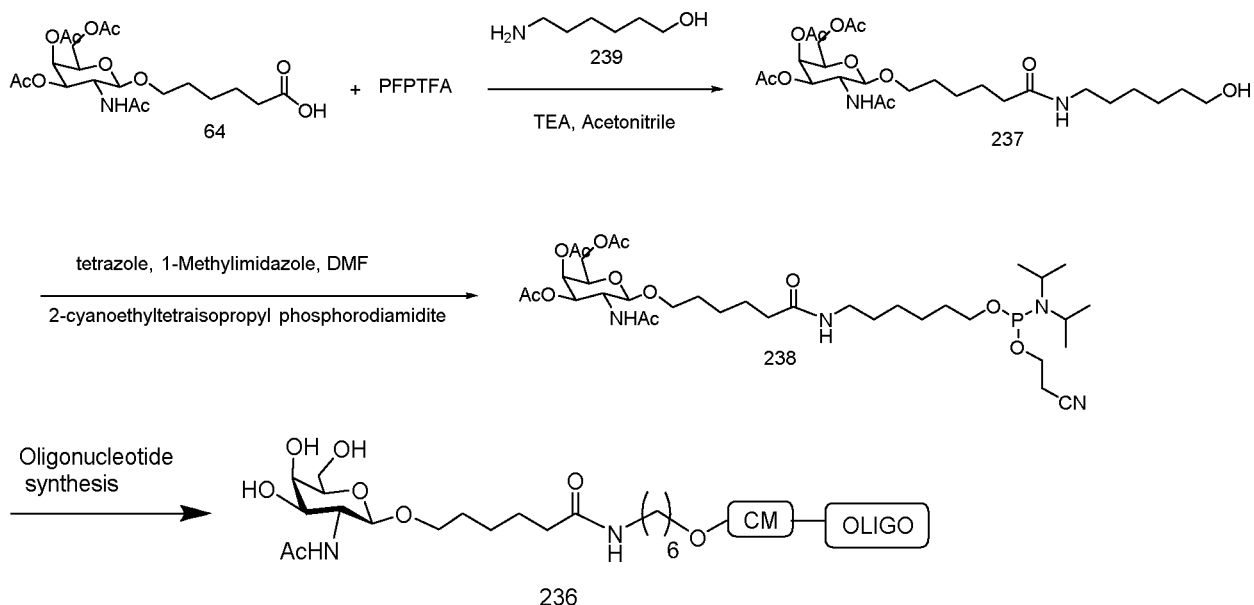


Example 105: Synthesis of oligonucleotides comprising a GalNAc₁-25 conjugate

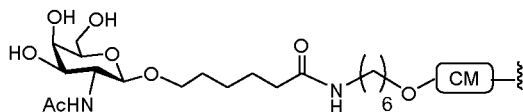


The synthesis of compound 166 is described in Example 54. Oligonucleotide 236 was prepared using the general procedure described in Example 46.

Alternatively, oligonucleotide 236 was synthesized using the scheme shown below, and compound 238 was used to form the oligonucleotide 236 using procedures described in Example 10.



The GalNAc₁ cluster portion (GalNAc₁-25_a) of the conjugate group GalNAc₁-25 can be combined with any cleavable moiety present on the oligonucleotide to provide a variety of conjugate groups. The structure of GalNAc₁-25 (GalNAc₁-25_a-CM) is shown below:



Example 106: Antisense inhibition *in vivo* by oligonucleotides targeting SRB-1 comprising a 5'-GalNAc₂ or a 5'-GalNAc₃ conjugate

Oligonucleotides listed in Tables 116 and 117 were tested in dose-dependent studies for antisense inhibition of SRB-1 in mice.

Treatment

Six to week old, male C57BL/6 mice (Jackson Laboratory, Bar Harbor, ME) were injected subcutaneously once with 2, 7, or 20 mg/kg of ISIS No. 440762; or with 0.2, 0.6, 2, 6, or 20 mg/kg of ISIS No. 686221, 686222, or 708561; or with saline. Each treatment group consisted of 4 animals. The mice were sacrificed 72 hours following the final administration. Liver SRB-1 mRNA levels were measured using real-time PCR. SRB-1 mRNA levels were normalized to cyclophilin mRNA levels according to standard protocols. The antisense oligonucleotides lowered SRB-1 mRNA levels in a dose-dependent manner, and the ED₅₀ results are presented in Tables 116 and 117. Although previous studies showed that trivalent GalNAc-conjugated oligonucleotides were significantly more potent than divalent GalNAc-conjugated oligonucleotides, which were in turn significantly more potent than monovalent GalNAc conjugated oligonucleotides (*see, e.g., Khorev et al., Bioorg. & Med. Chem.*, Vol. 16, 5216-5231 (2008)), treatment with

antisense oligonucleotides comprising monovalent, divalent, and trivalent GalNAc clusters lowered SRB-1 mRNA levels with similar potencies as shown in Tables 116 and 117.

Table 116
Modified oligonucleotides targeting SRB-1

ISIS No.	Sequences (5' to 3')	GalNAc Cluster	ED ₅₀ (mg/kg)	SEQ ID No
440762	T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _k	n/a	4.7	137
686221	GalNAc₂-24_a -o'-A _{do} T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _k	GalNAc ₂ -24 _a	0.39	141
686222	GalNAc₃-13_a -o'-A _{do} T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _k	GalNAc ₃ -13 _a	0.41	141

5 See Example 93 for table legend. The structure of GalNAc₃-13a was shown in Example 62, and the structure of GalNAc₂-24a was shown in Example 104.

Table 117
Modified oligonucleotides targeting SRB-1

ISIS No.	Sequences (5' to 3')	GalNAc Cluster	ED ₅₀ (mg/kg)	SEQ ID No
440762	T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _k	n/a	5	137
708561	GalNAc₁-25_a -o'-T _{ks} ^m C _{ks} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{ks} ^m C _k	GalNAc ₁ -25 _a	0.4	137

See Example 93 for table legend. The structure of GalNAc₁-25a was shown in Example 105.

10 The concentrations of the oligonucleotides in Tables 116 and 117 in liver were also assessed, using procedures described in Example 75. The results shown in Tables 117a and 117b below are the average total antisense oligonucleotide tissues levels for each treatment group, as measured by UV in units of μg oligonucleotide per gram of liver tissue. The results show that the oligonucleotides comprising a GalNAc conjugate group accumulated in the liver at significantly higher levels than the same dose of the oligonucleotide lacking a GalNAc conjugate group. Furthermore, the antisense oligonucleotides comprising one, two, or three GalNAc ligands in their respective conjugate groups all accumulated in the liver at similar levels. This result is surprising in view of the Khorev et al. literature reference cited above and is consistent with the activity data shown in Tables 116 and 117 above.

Table 117a

Liver concentrations of oligonucleotides comprising a GalNAc₂ or GalNAc₃ conjugate group

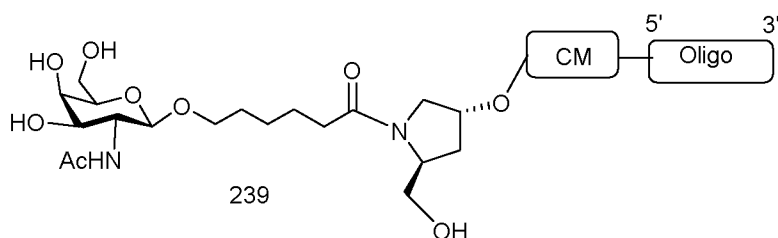
ISIS No.	Dosage (mg/kg)	[Antisense oligonucleotide] (μg/g)	GalNAc cluster	CM
440762	2	2.1	n/a	n/a
	7	13.1		
	20	31.1		
686221	0.2	0.9	GalNAc ₂ -24 _a	A _d
	0.6	2.7		
	2	12.0		
	6	26.5		

686222	0.2	0.5	GalNAc ₃ -13 _a	A _d
	0.6	1.6		
	2	11.6		
	6	19.8		

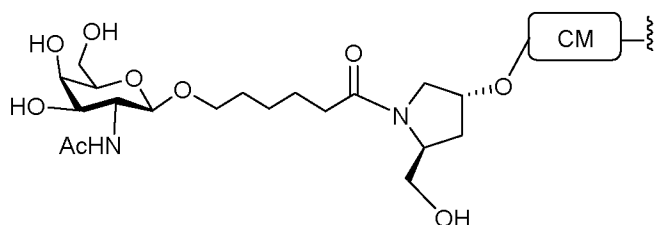
Table 117b

Liver concentrations of oligonucleotides comprising a GalNAc₁ conjugate group

ISIS No.	Dosage (mg/kg)	[Antisense oligonucleotide] (μg/g)	GalNAc cluster	CM
440762	2	2.3	n/a	n/a
	7	8.9		
	20	23.7		
708561	0.2	0.4	GalNAc ₁ -25 _a	PO
	0.6	1.1		
	2	5.9		
	6	23.7		
	20	53.9		

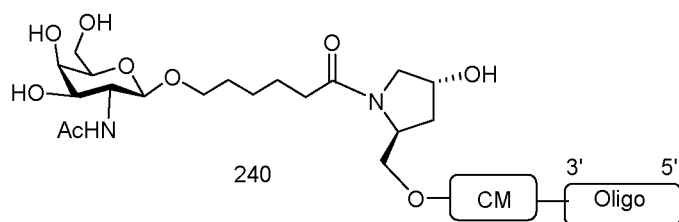
5 Example 107: Synthesis of oligonucleotides comprising a GalNAc₁-26 or GalNAc₁-27 conjugate

Oligonucleotide 239 is synthesized via coupling of compound 47 (see Example 15) to acid 64 (see Example 32) using HBTU and DIEA in DMF. The resulting amide containing compound is phosphitylated, then added to the 5'-end of an oligonucleotide using procedures described in Example 10. The GalNAc₁ cluster portion (GalNAc₁-26_a) of the conjugate group GalNAc₁-26 can be combined with any cleavable moiety present on the oligonucleotide to provide a variety of conjugate groups. The structure of GalNAc₁-26 (GalNAc₁-26_a-CM) is shown below:

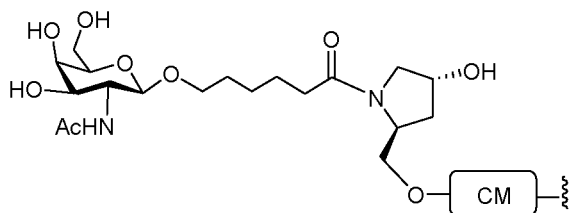


In order to add the GalNAc₁ conjugate group to the 3'-end of an oligonucleotide, the amide formed from the reaction of compounds 47 and 64 is added to a solid support using procedures described in Example

7. The oligonucleotide synthesis is then completed using procedures described in Example 9 in order to form oligonucleotide 240.



The GalNAc₁ cluster portion (GalNAc_{1-27a}) of the conjugate group GalNAc₁₋₂₇ can be combined with any
 5 cleavable moiety present on the oligonucleotide to provide a variety of conjugate groups. The structure of GalNAc₁₋₂₇ (GalNAc_{1-27a}-CM) is shown below:



Example 108: Antisense inhibition *in vivo* by oligonucleotides comprising a GalNAc conjugate group targeting Apo(a) *in vivo*

The oligonucleotides listed in Table 118 below were tested in a single dose study in mice.

Table 118
Modified ASOs targeting APO(a)

ISIS No.	Sequences (5' to 3')	GalNAc ₃ Cluster	CM	SEQ ID No.
494372	T _{es} G _{es} ^m C _{es} T _{es} ^m C _{es} ^m C _{ds} G _{ds} T _{ds} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{es} G _{es} T _{es} T _{es} ^m C _e	n/a	n/a	58
681251	GalNAc₃-7a-o' T _{es} G _{es} ^m C _{es} T _{es} ^m C _{es} ^m C _{ds} G _{ds} T _{ds} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{es} G _{es} T _{es} T _{es} ^m C _e	GalNAc ₃ -7a	PO	58
681255	GalNAc₃-3a-o' T _{es} G _{eo} ^m C _{eo} T _{eo} ^m C _{eo} ^m C _{ds} G _{ds} T _{ds} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{eo} G _{eo} T _{es} T _{es} ^m C _e	GalNAc ₃ -3a	PO	58
681256	GalNAc₃-10a-o' T _{es} G _{eo} ^m C _{eo} T _{eo} ^m C _{eo} ^m C _{ds} G _{ds} T _{ds} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{eo} G _{eo} T _{es} T _{es} ^m C _e	GalNAc ₃ -10a	PO	58
681257	GalNAc₃-7a-o' T _{es} G _{eo} ^m C _{eo} T _{eo} ^m C _{eo} ^m C _{ds} G _{ds} T _{ds} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{eo} G _{eo} T _{es} T _{es} ^m C _e	GalNAc ₃ -7a	PO	58
681258	GalNAc₃-13a-o' T _{es} G _{eo} ^m C _{eo} T _{eo} ^m C _{eo} ^m C _{ds} G _{ds} T _{ds} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{eo} G _{eo} T _{es} T _{es} ^m C _e	GalNAc ₃ -13a	PO	58
681260	T _{es} G _{eo} ^m C _{eo} T _{eo} ^m C _{eo} ^m C _{ds} G _{ds} T _{ds} T _{ds} G _{ds} G _{ds} T _{ds} G _{ds} ^m C _{ds} T _{ds} T _{eo} G _{eo} T _{es} T _{es} ^m C _{eo} A₄₀'-GalNAc₃-19	GalNAc ₃ -19a	A _d	167

The structure of GalNAc₃-7a was shown in Example 48.

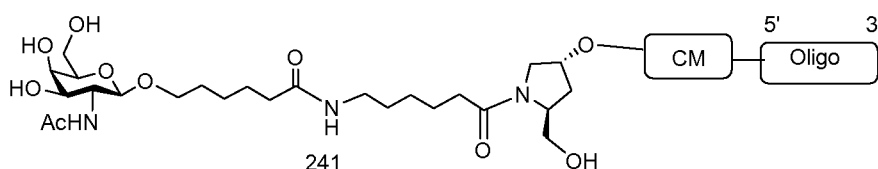
Treatment

Male transgenic mice that express human Apo(a) were each injected subcutaneously once with an oligonucleotide and dosage listed in Table 119 or with PBS. Each treatment group consisted of 4 animals. Blood was drawn the day before dosing to determine baseline levels of Apo(a) protein in plasma and at 1 week following the first dose. Additional blood draws will occur weekly for approximately 8 weeks. Plasma Apo(a) protein levels were measured using an ELISA. The results in Table 119 are presented as the average percent of plasma Apo(a) protein levels for each treatment group, normalized to baseline levels (% BL). The results show that the antisense oligonucleotides reduced Apo(a) protein expression. Furthermore, the oligonucleotides comprising a GalNAc conjugate group exhibited even more potent reduction in Apo(a) expression than the oligonucleotide that does not comprise a conjugate group.

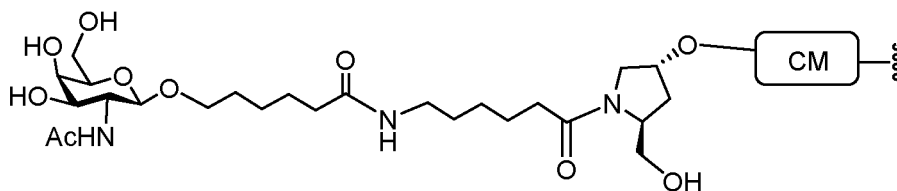
Table 119
Apo(a) plasma protein levels

ISIS No.	Dosage (mg/kg)	Apo(a) at 1 week (% BL)
PBS	n/a	143
494372	50	58
681251	10	15
681255	10	14
681256	10	17
681257	10	24
681258	10	22
681260	10	26

Example 109: Synthesis of oligonucleotides comprising a GalNAc₁-28 or GalNAc₁-29 conjugate

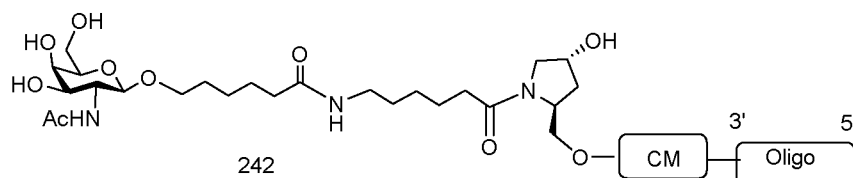


Oligonucleotide 241 is synthesized using procedures similar to those described in Example 71 to form the phosphoramidite intermediate, followed by procedures described in Example 10 to synthesize the oligonucleotide. The GalNAc₁ cluster portion (GalNAc₁-28_a) of the conjugate group GalNAc₁-28 can be combined with any cleavable moiety present on the oligonucleotide to provide a variety of conjugate groups. The structure of GalNAc₁-28 (GalNAc₁-28_a-CM) is shown below:

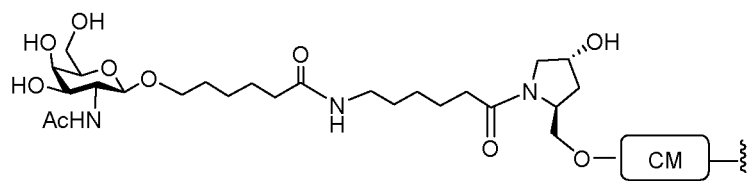


In order to add the GalNAc₁ conjugate group to the 3'-end of an oligonucleotide, procedures similar to those described in Example 71 are used to form the hydroxyl intermediate, which is then added to the solid support using procedures described in Example 7. The oligonucleotide synthesis is then completed using

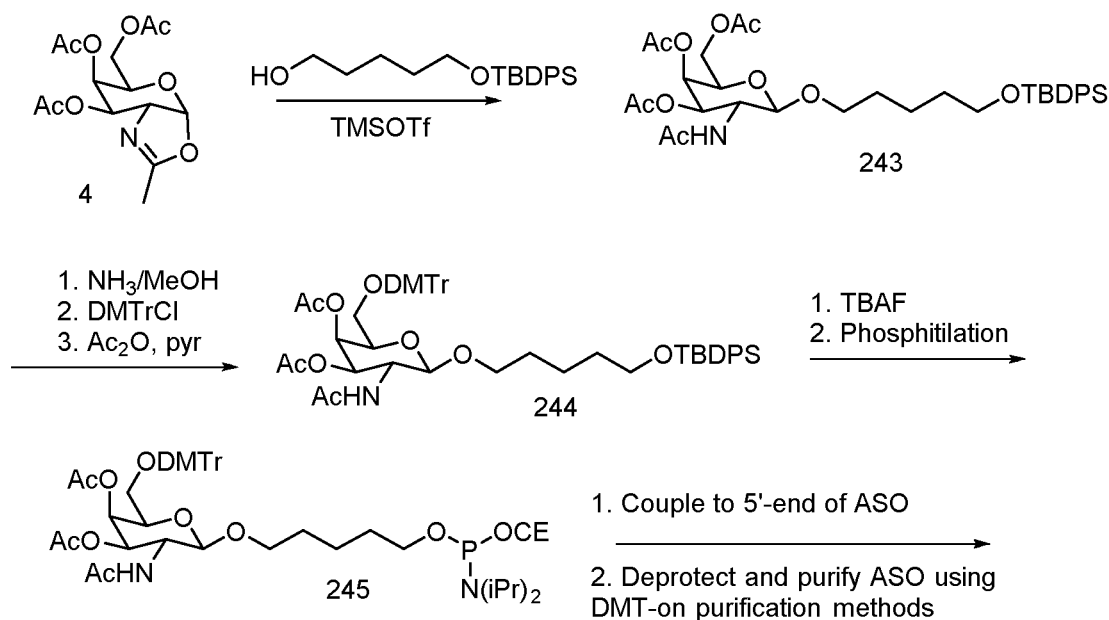
procedures described in Example 9 in order to form oligonucleotide 242.

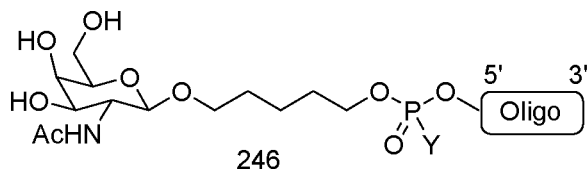


The GalNAc₁ cluster portion (GalNAc₁-29_a) of the conjugate group GalNAc₁-29 can be combined with any cleavable moiety present on the oligonucleotide to provide a variety of conjugate groups. The structure of GalNAc₁-29 (GalNAc₁-29_a-CM) is shown below:

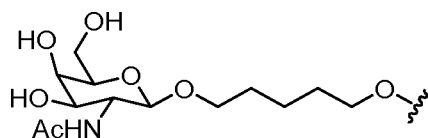


Example 110: Synthesis of oligonucleotides comprising a GalNAc₁-30 conjugate

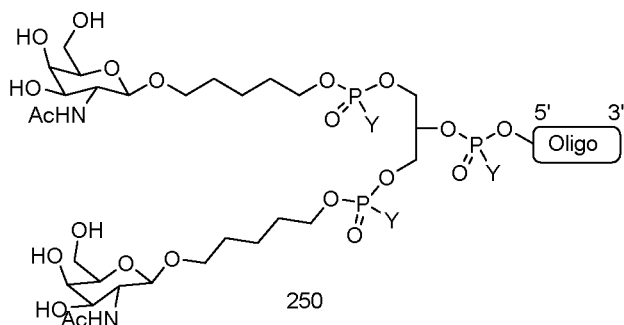
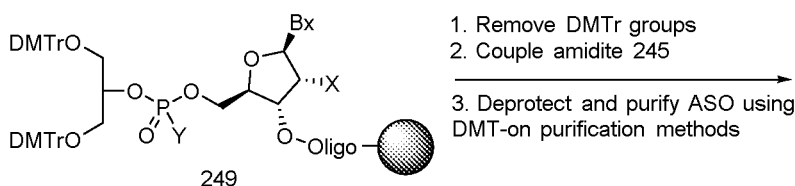
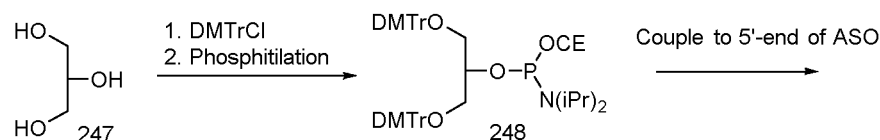




Oligonucleotide 246 comprising a GalNAc₁-30 conjugate group, wherein Y is selected from O, S, a substituted or unsubstituted C₁-C₁₀ alkyl, amino, substituted amino, azido, alkenyl or alkynyl, is synthesized as shown above. The GalNAc₁ cluster portion (GalNAc₁-30_a) of the conjugate group GalNAc₁-30 can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, Y is part of the cleavable moiety. In certain embodiments, Y is part of a stable moiety, and the cleavable moiety is present on the oligonucleotide. The structure of GalNAc₁-30_a is shown below:

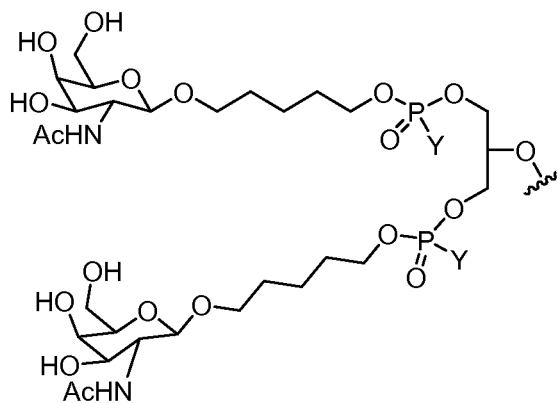


Example 111: Synthesis of oligonucleotides comprising a GalNAc₂-31 or GalNAc₂-32 conjugate

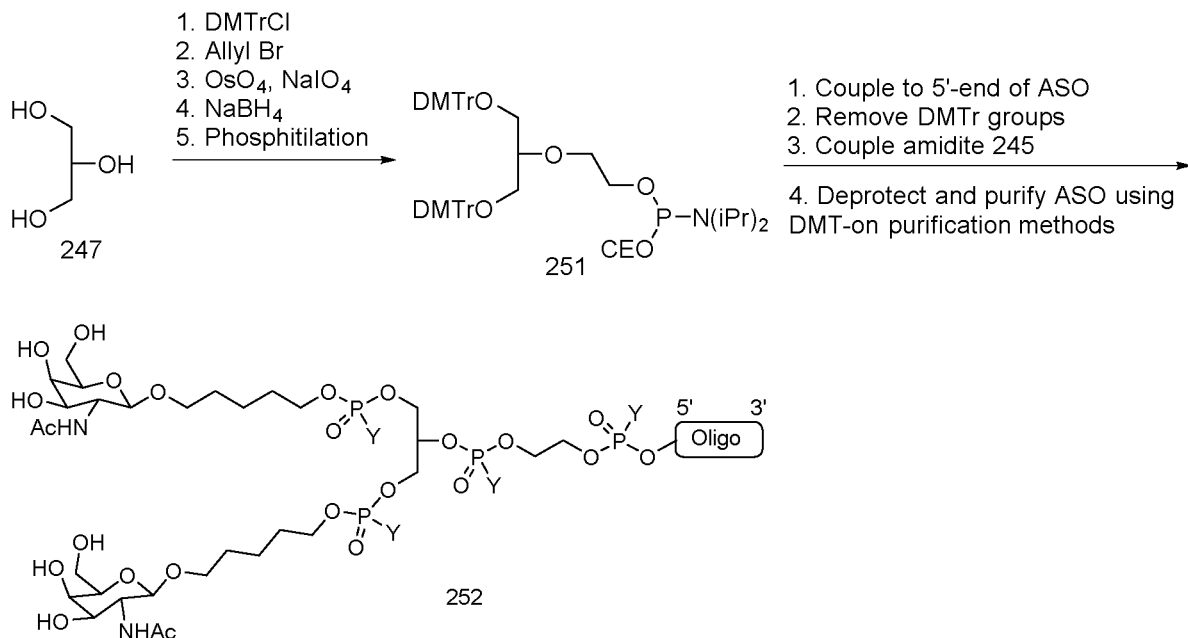


Oligonucleotide 250 comprising a GalNAc₂-31 conjugate group, wherein Y is selected from O, S, a substituted or unsubstituted C₁-C₁₀ alkyl, amino, substituted amino, azido, alkenyl or alkynyl, is synthesized as shown above. The GalNAc₂ cluster portion (GalNAc₂-31_a) of the conjugate group GalNAc₂-31 can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the Y-containing group directly adjacent to the 5'-end of the oligonucleotide is part of the cleavable moiety. In certain embodiments, the Y-containing group directly adjacent to the 5'-end of the oligonucleotide is part of a

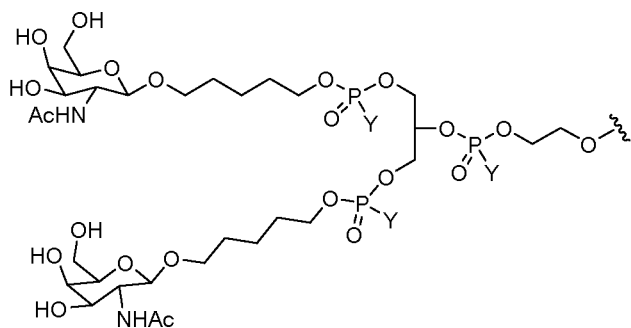
stable moiety, and the cleavable moiety is present on the oligonucleotide. The structure of GalNAc₂-31_a is shown below:



The synthesis of an oligonucleotide comprising a GalNAc₂-32 conjugate is shown below.



Oligonucleotide 252 comprising a GalNAc₂-32 conjugate group, wherein Y is selected from O, S, a substituted or unsubstituted C₁-C₁₀ alkyl, amino, substituted amino, azido, alkenyl or alkynyl, is synthesized as shown above. The GalNAc₂ cluster portion (GalNAc₂-32_a) of the conjugate group GalNAc₂-32 can be combined with any cleavable moiety to provide a variety of conjugate groups. In certain embodiments, the Y-containing group directly adjacent to the 5'-end of the oligonucleotide is part of the cleavable moiety. In certain embodiments, the Y-containing group directly adjacent to the 5'-end of the oligonucleotide is part of a stable moiety, and the cleavable moiety is present on the oligonucleotide. The structure of GalNAc₂-32_a is shown below:



Example 112: Modified oligonucleotides comprising a GalNAc₁ conjugate

The oligonucleotides in Table 120 targeting SRB-1 were synthesized with a GalNAc₁ conjugate group in order to further test the potency of oligonucleotides comprising conjugate groups that contain one GalNAc ligand.

Table 120

ISIS No.	Sequence (5' to 3')	GalNAc cluster	CM	SEQ ID NO.
711461	GalNAc₁-25_{a-o} .A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	GalNAc ₁ -25 _a	A _d	145
711462	GalNAc₁-25_{a-o} .G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	GalNAc ₁ -25 _a	PO	143
711463	GalNAc₁-25_{a-o} .G _{es} ^m C _{eo} T _{eo} T _{eo} ^m C _{eo} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{eo} ^m C _{eo} ^m C _{es} T _{es} T _e	GalNAc ₁ -25 _a	PO	143
711465	GalNAc₁-26_{a-o} .A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	GalNAc ₁ -26 _a	A _d	145
711466	GalNAc₁-26_{a-o} .G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	GalNAc ₁ -26 _a	PO	143
711467	GalNAc₁-26_{a-o} .G _{es} ^m C _{eo} T _{eo} T _{eo} ^m C _{eo} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{eo} ^m C _{eo} ^m C _{es} T _{es} T _e	GalNAc ₁ -26 _a	PO	143
711468	GalNAc₁-28_{a-o} .A _{do} G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	GalNAc ₁ -28 _a	A _d	145
711469	GalNAc₁-28_{a-o} .G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _e	GalNAc ₁ -28 _a	PO	143
711470	GalNAc₁-28_{a-o} .G _{es} ^m C _{eo} T _{eo} T _{eo} ^m C _{eo} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{eo} ^m C _{eo} ^m C _{es} T _{es} T _e	GalNAc ₁ -28 _a	PO	143
713844	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _{eo} . GalNAc₁-27_a	GalNAc ₁ -27 _a	PO	143
713845	G _{es} ^m C _{eo} T _{eo} T _{eo} ^m C _{eo} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{eo} ^m C _{eo} ^m C _{es} T _{es} T _{eo} . GalNAc₁-27_a	GalNAc ₁ -27 _a	PO	143
713846	G _{es} ^m C _{eo} T _{eo} T _{eo} ^m C _{eo} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{eo} ^m C _{eo} ^m C _{es} T _{es} T _{eo} . A_{do}.GalNAc₁-27_a	GalNAc ₁ -27 _a	A _d	144
713847	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _{eo} . GalNAc₁-29_a	GalNAc ₁ -29 _a	PO	143
713848	G _{es} ^m C _{eo} T _{eo} T _{eo} ^m C _{eo} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{eo} ^m C _{eo} ^m C _{es} T _{es} T _{eo} . GalNAc₁-29_a	GalNAc ₁ -29 _a	PO	143
713849	G _{es} ^m C _{es} T _{es} T _{es} ^m C _{es} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{es} ^m C _{es} ^m C _{es} T _{es} T _{eo} . A_{do}.GalNAc₁-29_a	GalNAc ₁ -29 _a	A _d	144
713850	G _{es} ^m C _{eo} T _{eo} T _{eo} ^m C _{eo} A _{ds} G _{ds} T _{ds} ^m C _{ds} A _{ds} T _{ds} G _{ds} A _{ds} ^m C _{ds} T _{ds} T _{eo} ^m C _{eo} ^m C _{es} T _{es} T _{eo} . A_{do}.GalNAc₁-29_a	GalNAc ₁ -29 _a	A _d	144

Example 113: Dose-dependent antisense inhibition of human apolipoprotein (a) (apo(a)) in human primary hepatocytes

- 5 Selected gapmer antisense oligonucleotides from a previous publication (WO2005/000201, the content of which is incorporated by reference in its entirety herein) were tested in a single dose assay in human primary hepatocytes. Cells were obtained from Tissue Transformation Technologies (BD Biosciences, Franklin Lakes, NJ) and treated with 150 nM of antisense oligonucleotide. After a treatment period of

approximately 16 hours, RNA was isolated from the cells and apo(a) mRNA levels were measured by quantitative real-time PCR. Human apo(a) primer probe set hAPO(a)3' (forward sequence ACAGCAATCAAACGAAGACACTG, designated herein as SEQ ID NO: 5; reverse sequence AGCTTATACACAAAAATACCAAAAATGC, designated herein as SEQ ID NO: 6; probe sequence TCCCAGCTACCAGCTATGCCAAACCTT, designated herein as SEQ ID NO: 7) was used to measure mRNA levels. Additionally, mRNA levels were also measured using human apo(a) primer probe set hAPO(a)12kB (forward sequence CCACAGTGGCCCCGGT, designated herein as SEQ ID NO: 8; reverse sequence ACAGGGCTTTTCTCAGGTGGT, designated herein as SEQ ID NO: 9; probe sequence CCAAGCACAGAGGCTCCTTCTGAACAAG, designated herein as SEQ ID NO: 10). Apo(a) mRNA levels were normalized to GAPDH mRNA expression. Results are presented in the table below as percent inhibition of apo(a), relative to untreated control cells.

Table 121

Antisense inhibition of human apo(a) in human primary hepatocytes

ISIS No	% inhibition (hAPO(a)3' PPset)	% inhibition (hAPO(a)12kB PPset)
144367	68	77
144368	42	59
144369	43	69
144370	80	75
144371	42	57
144372	87	54
144373	63	49
144374	45	80
144375	33	11
144376	62	82
144377	42	72
144378	0	72
144379	73	46
144380	75	78
144381	63	64
144382	0	58
144383	63	79
144384	38	0

144385	40	94
144386	47	61
144387	38	60
144388	0	57
144389	52	39
144390	12	0
144391	73	57
144392	43	50
144393	83	82
144394	40	76
144395	80	84
144396	53	72
144397	23	64
144398	7	33
144399	43	44
144400	70	75
144401	87	72

Several antisense oligonucleotides were selected for further testing in a dose response assay.

The selected antisense oligonucleotides were tested in human primary hepatocytes with 25 nM, 50 nM, 150 nM, or 300 nM concentrations of antisense oligonucleotide, as specified in the table below. After a treatment period of approximately 16 hours, RNA was isolated from the cells and apo(a) mRNA levels were measured by quantitative real-time PCR. Human apo(a) primer probe set hAPO(a)3' was used to measure mRNA levels. Apo(a) mRNA levels were normalized to GAPDH mRNA expression. Results are presented as percent inhibition of apo(a), relative to untreated control cells.

Table 122

Dose-dependent antisense inhibition of human apo(a) in human primary hepatocytes, as measured with hAPO(a)3'

ISIS No	25 nM	50 nM	150 nM	300 nM
144367	52	78	76	74
144370	64	74	68	66
144385	0	15	43	5
144393	0	9	39	25

144395	17	9	8	32
--------	----	---	---	----

ISIS 144367 demonstrated better efficacy and dose-dependency than the other antisense oligonucleotides. Hence, ISIS 144367 was considered the benchmark antisense oligonucleotide to compare the potency of newly designed antisense oligonucleotides disclosed herein.

5 **Example 114: Antisense inhibition of human apo(a) in transgenic mouse primary hepatocytes**

Antisense oligonucleotides were newly designed targeting an apo(a) nucleic acid and were tested for their effects on apo(a) mRNA *in vitro*. The antisense oligonucleotides were tested for potency in a series of parallel experiments that had similar culture conditions. Primary hepatocytes from human apo(a) transgenic mice (Frazer, K.A. et al., Nat. Genet. 1995. 9: 424-431) were used in this study. Hepatocytes at a density of 10 35,000 cells per well were transfected using electroporation with 1,000 nM antisense oligonucleotide. After a treatment period of approximately 24 hours, RNA was isolated from the cells and apo(a) mRNA levels were measured by quantitative real-time PCR. Human primer probe set hAPO(a)12kB was used to measure mRNA levels. Apo(a) mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN®. The results for each experiment are presented in separate tables shown below. ISIS 144367 15 from was used as a benchmark for the new antisense oligonucleotides and also included in the studies. Results are presented as percent inhibition of apo(a), relative to untreated control cells. A total of 1,511 gapmers were tested under these culture conditions. Only those antisense oligonucleotides that were selected for further study are presented in the table below with each table representing a separate experiment.

The newly designed chimeric antisense oligonucleotides were designed as 5-10-5 MOE gapmers. 20 The gapmers are 20 nucleosides in length, wherein the central gap segment comprises of ten 2'-deoxynucleosides and is flanked by wing segments on the 5' direction and the 3' direction comprising five nucleosides each. Each nucleoside in the 5' wing segment and each nucleoside in the 3' wing segment has a 2'-MOE modification. The internucleoside linkages throughout each gapmer are phosphorothioate (P=S) linkages. All cytosine residues throughout each gapmer are 5-methylcytosines.

25 The apo(a) target sequence contains multiple Kringle repeat sequences, therefore, an antisense oligonucleotide may target one or more regions of apo(a) depending whether on the oligonucleotide targets a Kringle sequence or not. "Start site" indicates the 5'-most nucleoside to which the gapmer is targeted in the human sequence. "Stop site" indicates the 3'-most nucleoside to which the gapmer is targeted human sequence. An apo(a) antisense oligonucleotide may have more than one "Start site" or "Stop site" depending 30 on whether or not it targets a Kringle repeat.

Most gapmers listed in the tables are targeted with 100% complementarity to one or more regions of either the human apo(a) mRNA, designated herein as SEQ ID NO: 1 (GENBANK Accession No.

NM_005577.2) or the human apo(a) genomic sequence, designated herein as SEQ ID NO: 2 (GENBANK Accession No. NT_007422.12 truncated from nucleotides 3230000 to 3380000), or both. 'n/a' indicates that the antisense oligonucleotide does not target that particular sequence with 100% complementarity.

5

Table 123

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	90	21210	21229	11
494157	238	257	CCTGTGACAGTGGTGGAGTA	95	21199	21218	12
	580	599			26690	26709	
	922	941			32237	32256	
	1606	1625			43330	43349	
	1948	1967			48874	48893	
	2290	2309			54420	54439	
	3316	3335			72037	72056	
494158	239	258	TCCTGTGACAGTGGTGGAGT	95	21200	21219	13
	581	600			26691	26710	
	923	942			32238	32257	
	1607	1626			43331	43350	
	1949	1968			48875	48894	
	2291	2310			54421	54440	
	3317	3336			72038	72057	
494159	241	260	CTTCCTGTGACAGTGGTGGGA	97	21202	21221	14
	583	602			26693	26712	
	925	944			32240	32259	
	1609	1628			43333	43352	
	1951	1970			48877	48896	
	2293	2312			54423	54442	
	3319	3338			72040	72059	
	4663	4682			94404	94423	
	5005	5024			115515	115534	
494160	242	261	CCTTCCTGTGACAGTGGTGG	97	21203	21222	15
	4664	4683			94405	94424	
	5006	5025			115516	115535	
494161	243	262	TCCTTCCTGTGACAGTGGTG	96	21204	21223	16
	4665	4684			94406	94425	
	5007	5026			115517	115536	
494162	244	263	GTCCTTCCTGTGACAGTGGT	95	21205	21224	17
	3664	3683			77585	77604	
	4666	4685			94407	94426	
	5008	5027			115518	115537	
494163	245	264	GGTCCTTCCTGTGACAGTGG	96	21206	21225	18

	4667	4686			94408	94427	
494164	246	265	AGGTCCTTCCTGTGACAGTG	93	21207	21226	19
	4668	4687			94409	94428	
494165	247	266	CAGGTCCTTCCTGTGACAGT	91	21208	21227	20
	4669	4688			94410	94429	
494166	248	267	GCAGGTCCTTCCTGTGACAG	89	21209	21228	21
494167	250	269	TGGCAGGTCCTTCCTGTGAC	92	21211	21230	22
494168	251	270	TTGGCAGGTCCTTCCTGTGA	89	21212	21231	23
494169	252	271	CTTGGCAGGTCCTTCCTGTG	92	21213	21232	24
494170	253	272	GCTTGGCAGGTCCTTCCTGT	88	21214	21233	25

Table 124

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	91	21210	21229	11
				84			
494283	584	603	TCTTCCTGTGACAGTGGTGG	93	26694	26713	26
	926	945			32241	32260	
	1610	1629			43334	43353	
	1952	1971			48878	48897	
	2294	2313			54424	54443	
	3320	3339			72041	72060	
494284	585	604	TTCTTCCTGTGACAGTGGTG	95	26695	26714	27
	927	946			32242	32261	
	1611	1630			43335	43354	
	1953	1972			48879	48898	
	2295	2314			54425	54444	
	3321	3340			72042	72061	
494285	586	605	GTTCTTCCTGTGACAGTGGT	95	26696	26715	28
	928	947			32243	32262	
	1612	1631			43336	43355	
	1954	1973			48880	48899	
	2296	2315			54426	54445	
	3322	3341			72043	72062	
494286	587	606	GGTTCTTCCTGTGACAGTGG	95	26697	26716	29
	929	948			32244	32263	
	1613	1632			43337	43356	
	1955	1974			48881	48900	
	2297	2316			54427	54446	
494287	588	607	AGGTTCTTCCTGTGACAGTG	95	26698	26717	30
	930	949			32245	32264	
	1614	1633			43338	43357	
	1956	1975			48882	48901	
	2298	2317			54428	54447	
494288	589	608	CAGGTTCTTCCTGTGACAGT	91	26699	26718	31
	931	950			32246	32265	
	1615	1634			43339	43358	

	1957	1976			48883	48902	
	2299	2318			54429	54448	
	2983	3002			66500	66519	
494290	592	611	TGGCAGGTTCTTCCTGTGAC	90	26702	26721	32
	934	953			32249	32268	
	1618	1637			43342	43361	
	1960	1979			48886	48905	
	2302	2321			54432	54451	
	2986	3005			66503	66522	
494291	593	612	TTGGCAGGTTCTTCCTGTGA	89	26703	26722	33
	935	954			32250	32269	
	1619	1638			43343	43362	
	1961	1980			48887	48906	
	2303	2322			54433	54452	
	2987	3006			66504	66523	
494292	594	613	CTTGGCAGGTTCTTCCTGTG	94	26704	26723	35
	936	955			32251	32270	
	1620	1639			43344	43363	
	1962	1981			48888	48907	
	2304	2323			54434	54453	
	2988	3007			66505	66524	
494294	596	615	AGCTTGGCAGGTTCTTCCTG	90	26706	26725	36
	938	957			32253	32272	
	1622	1641			43346	43365	
	1964	1983			48890	48909	
	2306	2325			54436	54455	
	2990	3009			66507	66526	
494299	626	645	ACTATGCGAGTGTGGTGTCA	91	26736	26755	37
	968	987			32283	32302	
	1310	1329			37830	37849	
	1652	1671			43376	43395	
	1994	2013			48920	48939	
	2336	2355			54466	54485	
	2678	2697			60021	60040	
	3020	3039			66537	66556	
494300	627	646	GACTATGCGAGTGTGGTGTGTC	93	26737	26756	38
	969	988			32284	32303	
	1311	1330			37831	37850	
	1653	1672			43377	43396	
	1995	2014			48921	48940	
	2337	2356			54467	54486	
	2679	2698			60022	60041	
	3021	3040			66538	66557	
494301	628	647	CGACTATGCGAGTGTGGTGT	93	26738	26757	39

	970	989			32285	32304	
	1312	1331			37832	37851	
	1654	1673			43378	43397	
	1996	2015			48922	48941	
	2338	2357			54468	54487	
	2680	2699			60023	60042	
	3022	3041			66539	66558	
494302	629	648	CCGACTATGCGAGTGTGGTG	94	26739	26758	40
	971	990			32286	32305	
	1313	1332			37833	37852	
	1655	1674			43379	43398	
	1997	2016			48923	48942	
	2339	2358			54469	54488	
	2681	2700			60024	60043	
	3023	3042			66540	66559	
494303	630	649	TCCGACTATGCGAGTGTGGT	93	26740	26759	41
	972	991			32287	32306	
	1314	1333			37834	37853	
	1656	1675			43380	43399	
	1998	2017			48924	48943	
	2340	2359			54470	54489	
	2682	2701			60025	60044	
	3024	3043			66541	66560	
494304	631	650	GTCCGACTATGCGAGTGTGG	94	26741	26760	42
	973	992			32288	32307	
	1315	1334			37835	37854	
	1657	1676			43381	43400	
	1999	2018			48925	48944	
	2341	2360			54471	54490	
	2683	2702			60026	60045	
	3025	3044			66542	66561	
494305	632	651	GGTCCGACTATGCGAGTGTG	93	26742	26761	43
	974	993			32289	32308	
	1316	1335			37836	37855	
	1658	1677			43382	43401	
	2000	2019			48926	48945	
	2342	2361			54472	54491	
	2684	2703			60027	60046	
	3026	3045			66543	66562	
494306	633	652	GGGTCCGACTATGCGAGTGT	92	26743	26762	44
	975	994			32290	32309	
	1317	1336			37837	37856	
	1659	1678			43383	43402	

	2001	2020			48927	48946	
	2343	2362			54473	54492	
	2685	2704			60028	60047	
	3027	3046			66544	66563	
494307	1190	1209	CTGCTCAGTCGGTGCTTGTT	91	n/a	n/a	45
	2558	2577					
494310	1193	1212	CCTCTGCTCAGTCGGTGCTT	90	n/a	n/a	46
	2561	2580					
494311	1194	1213	GCCTCTGCTCAGTCGGTGCT	88	37714	37733	47
	2562	2581			59905	59924	
494334	1267	1286	CTTCCAGTGACAGTGGTGGA	90	37787	37806	48
	2635	2654			59978	59997	
494336	1269	1288	TTCTTCCAGTGACAGTGGTG	90	37789	37808	49
	2637	2656			59980	59999	
494337	1270	1289	GTTCTTCCAGTGACAGTGGT	95	37790	37809	50
	2638	2657			59981	60000	
494338	1271	1290	GGTTCTTCCAGTGACAGTGG	91	37791	37810	133
	2639	2658			59982	60001	
494521	6393	6412	GACCTTAAAAGCTTATACAC	82	140049	140068	51
494525	6397	6416	GTCAGACCTTAAAAGCTTAT	84	140053	140072	52
494530	6402	6421	TGTCAGTCAGACCTTAAAAG	82	140058	140077	53
494535	6407	6426	GAATTTGTCAGTCAGACCTT	85	140063	140082	54
494536	6408	6427	AGAATTTGTCAGTCAGACCT	83	140064	140083	55
494544	6417	6436	CCTTAATACAGAATTTGTCA	82	140073	140092	56

Table 125

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	84	21210	21229	11
494371	3900	3919	GCTCCGTTGGTGCTTGTTCA	93	n/a	n/a	57
494372	3901	3920	TGCTCCGTTGGTGCTTGTTT	93	n/a	n/a	58
494373	3902	3921	TTGCTCCGTTGGTGCTTGTT	83	n/a	n/a	59
494374	3903	3922	TTTGCTCCGTTGGTGCTTGT	89	n/a	n/a	60
494375	3904	3923	CTTTGCTCCGTTGGTGCTTG	85	n/a	n/a	61
494386	3977	3996	TCCTGTAACAGTGGTGGAGA	86	81985	82004	62
494387	3978	3997	TTCCTGTAACAGTGGTGGAG	82	81986	82005	63
494388	3979	3998	CTTCCTGTAACAGTGGTGGGA	86	81987	82006	64
494389	3980	3999	CCTTCCTGTAACAGTGGTGG	92	81988	82007	65
494390	3981	4000	TCCTTCCTGTAACAGTGGTG	92	81989	82008	66
494391	3982	4001	GTCCTTCCTGTAACAGTGGT	84	81990	82009	67
494392	3983	4002	TGTCCTTCCTGTAACAGTGG	81	81991	82010	68

Table 126

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	86	21210	21229	11
498369	3203	3222	TGGAGCCAGAATAACATTCTG	91	70667	70686	69
498379	3213	3232	CCTCTAGGCTTGGAGCCAGA	85	70677	70696	70
498408	3323	3342	AGTTCTTCCTGTGACAGTGG	86	72044	72063	71
498433	3367	3386	GTCCGACTATGCTGGTGTGG	87	72088	72107	72
498434	3368	3387	GGTCCGACTATGCTGGTGTG	86	72089	72108	73
498435	3369	3388	GGGTCCGACTATGCTGGTGT	83	72090	72109	74

Table 127

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	90	21210	21229	11
498229	2871	2890	CCTCTAGGCTTGAATCGGG	90	65117	65136	75
498238	2883	2902	GTTTCAGAAGGAGCCTCTAGG	93	65129	65148	76
498239	2884	2903	TGTTTCAGAAGGAGCCTCTAG	94	65130	65149	77
498240	2887	2906	GCTTGTTTCAGAAGGAGCCTC	98	n/a	n/a	78
	4573	4592					
498241	2888	2907	TGCTTGTTTCAGAAGGAGCCT	94	n/a	n/a	79
	4574	4593					
498242	2889	2908	GTGCTTGTTTCAGAAGGAGCC	96	n/a	n/a	80
	4575	4594					
498243	2890	2909	GGTGCTTGTTTCAGAAGGAGC	97	n/a	n/a	81
	4576	4595					
498244	2891	2910	TGGTGCTTGTTTCAGAAGGAG	92	n/a	n/a	82
	4577	4596					
498251	2898	2917	GCTCAGTTGGTGCTTGTTCA	90	n/a	n/a	83
498252	2899	2918	TGCTCAGTTGGTGCTTGTTTC	90	n/a	n/a	84

Table 128

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	91	21210	21229	11
498517	3548	3567	GCTTGATCTGGGACCACCG	89	76233	76252	85

5

Table 129

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	94	21210	21229	11

498833	4900	4919	GCCTCCATGCTTGGAAGTGG	94	114205	114224	86
498859	4926	4945	GCTCAGTTGGTGCTGCTTCA	92	n/a	n/a	87
498868	4978	4997	CCTCGATAACTCTGGCCATT	94	115488	115507	88
498875	5003	5022	TCCTGTGACAGTGGTGGAGA	94	115513	115532	89

Table 130

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	92	21210	21229	11
499020	6257	6276	GTAGGTTGATGCTTCACTCT	91	139913	139932	90
499041	6318	6337	CGTTTGATTGCTGTCTATTA	90	139974	139993	91

Table 131

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	91	21210	21229	11
498523	3554	3573	CTCTGTGCTTGGATCTGGGA	94	76239	76258	92
498524	3555	3574	CCTCTGTGCTTGGATCTGGG	96	76240	76259	93
498525	3556	3575	GCCTCTGTGCTTGGATCTGG	94	76241	76260	94
498529	3560	3579	AGAAGCCTCTGTGCTTGGAT	89	76245	76264	95
498535	3566	3585	TTCAGAAGAAGCCTCTGTGC	89	76251	76270	96
498550	3582	3601	GCTCCGTTGGTGCTTCTTCA	90	n/a	n/a	97
498553	3585	3604	TTTGCTCCGTTGGTGCTTCT	87	n/a	n/a	98
498555	3587	3606	GCTTTGCTCCGTTGGTGCTT	90	n/a	n/a	99
	3905	3924					
498556	3588	3607	GGCTTTGCTCCGTTGGTGCT	89	77509	77528	100
	3906	3925			81914	81933	
498557	3589	3608	GGGCTTTGCTCCGTTGGTGCT	89	77510	77529	101
	3907	3926			81915	81934	
498579	3662	3681	CCTTCCTGTGACAGTGGTAG	87	77583	77602	102
498580	3663	3682	TCCTTCCTGTGACAGTGGTA	92	77584	77603	103
498581	3665	3684	TGTCTTCCTGTGACAGTGG	94	77586	77605	104
	5009	5028			115519	115538	

5

Table 132

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	100	21210	21229	11
494230	477	496	CCTCTAGGCTTGAACCGGG	95	25380	25399	105
	819	838			30927	30946	
	1161	1180			36471	36490	

	1503	1522			42020	42039	
	1845	1864			47564	47583	
	2187	2206			53110	53129	
	2529	2548			58662	58681	
494243	494	513	TGCTTGTTTCGGAAGGAGCCT	93	n/a	n/a	106
	836	855					
	1178	1197					
	1520	1539					
	1862	1881					
	2204	2223					
	2546	2565					
494244	495	514	GTGCTTGTTTCGGAAGGAGCC	95	n/a	n/a	107
	837	856					
	1179	1198					
	1521	1540					
	1863	1882					
	2205	2224					
	2547	2566					

Table 133

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	96	21210	21229	11
494466	4208	4227	GCTTGGAAGTGGGACCACCG	95	85138	85157	108
494470	4212	4231	CTGTGCTTGGAAGTGGGACC	94	85142	85161	109
494472	4214	4233	CTCTGTGCTTGGAAGTGGGA	92	85144	85163	110

5 Example 115: Dose-dependent antisense inhibition of apo(a) in transgenic mouse primary hepatocytes

Gapmers from the studies described above exhibiting significant *in vitro* inhibition of apo(a) mRNA were selected and tested at various doses in transgenic mouse primary hepatocytes in a series of parallel studies with similar culture conditions. Cells were plated at a density of 35,000 per well and transfected using electroporation with 0.0625 μ M, 0.125 μ M, 0.25 μ M, 0.500 μ M, or 1.000 μ M concentrations of antisense oligonucleotide. After a treatment period of approximately 16 hours, RNA was isolated from the cells and apo(a) mRNA levels were measured by quantitative real-time PCR. Apo(a) primer probe set hAPO(a)12kB was used to measure mRNA levels. Apo(a) mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN®. Results are presented as percent inhibition of apo(a), relative to untreated control cells.

The results of each of the studies are depicted in the tables presented below with each table representing a separate experiment. The half maximal inhibitory concentration (IC₅₀) of each oligonucleotide is also presented in the tables. Apo(a) mRNA levels were significantly reduced in a dose-dependent manner in antisense oligonucleotide treated cells. The potency of the newly designed oligos was compared with the benchmark oligonucleotide ISIS 144367.

Table 134

ISIS No	0.0625 μM	0.125 μM	0.250 μM	0.500 μM	1.000 μM	IC ₅₀ (μM)
144367	11	27	46	62	80	0.31
494157	11	47	53	76	87	0.23
494158	19	57	75	84	88	0.13
494159	41	65	77	84	92	0.07
494160	44	69	76	85	91	0.06
494161	40	64	74	85	91	0.08
494162	36	63	76	87	88	0.09
494163	20	59	75	85	92	0.13
494164	3	45	62	74	90	0.21
494165	25	39	57	71	75	0.19
494166	17	30	47	59	76	0.31
494167	30	43	55	72	80	0.18
494168	25	36	44	59	75	0.28
494169	19	39	51	61	81	0.25

Table 135

ISIS No	0.0625 μM	0.125 μM	0.250 μM	0.500 μM	1.000 μM	IC ₅₀ (μM)
144367	23	40	58	76	88	0.19
494170	38	34	60	76	84	0.13
494230	55	71	89	95	97	0.03
494243	47	73	87	92	97	0.05
494244	58	73	86	92	96	0.03
494283	54	70	84	93	94	0.05
494284	45	62	83	92	95	0.07
494285	56	70	84	92	95	0.04
494286	51	70	87	93	95	0.05
494287	32	60	67	87	91	0.11
494288	26	41	61	79	88	0.17
494290	30	43	64	81	87	0.15
494291	29	40	56	75	85	0.18

Table 136

ISIS No	0.0625 μM	0.125 μM	0.250 μM	0.500 μM	1.000 μM	IC ₅₀ (μM)
144367	10	38	62	68	84	0.23
494292	17	36	74	85	90	0.17
494294	10	34	53	80	91	0.22
494299	32	29	56	77	88	0.16
494300	34	46	76	86	90	0.12
494301	44	56	72	86	89	0.09
494302	42	59	78	88	89	0.08
494303	37	58	70	86	89	0.10
494304	46	71	78	89	90	0.05
494305	39	58	62	85	87	0.10
494306	31	52	65	79	88	0.13
494307	23	23	39	65	78	0.34
494310	14	29	62	70	88	0.25

Table 137

ISIS No	0.0625 μM	0.125 μM	0.250 μM	0.500 μM	1.000 μM	IC ₅₀ (μM)
144367	0	29	45	73	92	0.27
494311	28	53	65	85	95	0.13
494334	20	44	66	86	96	0.16
494336	15	38	54	84	97	0.20
494337	28	50	77	90	98	0.12
494338	21	40	68	91	98	0.15
494371	19	0	71	89	97	0.15
494372	33	44	77	91	97	0.12
494373	15	36	65	83	95	0.19
494374	3	17	51	83	90	0.24
494375	1	34	56	80	93	0.23
494386	13	26	46	73	91	0.25
494387	17	27	45	67	88	0.28

Table 138

ISIS No	0.0625 μM	0.125 μM	0.250 μM	0.500 μM	1.000 μM	IC ₅₀ (μM)
144367	35	42	62	70	91	0.15
494537	19	34	54	79	90	0.21
494544	10	38	73	86	94	0.17

498229	36	58	80	92	97	0.10
498238	41	57	75	91	97	0.09
498239	56	71	79	90	94	0.03
498240	91	94	98	99	100	<0.06
498241	75	84	91	96	98	<0.06
498242	11	27	42	47	63	0.49
498243	91	93	96	98	99	<0.06
498244	4	0	0	13	43	>1.00
498251	30	30	42	73	89	0.26
498252	37	33	58	80	92	0.20
498369	22	22	10	22	34	>1.00

Table 139

ISIS No	0.0625 μM	0.125 μM	0.250 μM	0.500 μM	1.000 μM	IC ₅₀ (μM)
144367	15	32	54	75	90	0.22
498379	29	48	71	80	95	0.13
498408	38	57	77	88	96	0.09
498433	29	36	70	88	96	0.15
498434	49	43	50	78	90	0.19
498435	27	39	57	78	93	0.18
498517	64	72	82	93	98	<0.06
498721	77	84	88	96	97	<0.06
498833	73	78	91	95	99	<0.06
498859	7	24	37	62	75	0.36
498868	7	14	39	63	81	0.36
498875	16	21	33	55	81	0.39
499020	7	24	23	55	78	0.36
499041	6	16	33	64	83	0.35

Table 140

ISIS No	0.0625 μM	0.125 μM	0.250 μM	0.500 μM	1.000 μM	IC ₅₀ (μM)
144367	14	47	64	79	91	0.14
498523	36	50	80	87	95	0.11
498524	43	79	87	93	97	0.01
498525	32	49	75	86	96	0.12
498529	21	49	57	78	90	0.17
498535	20	34	55	76	86	0.21
498550	12	50	69	84	96	0.11

498553	8	43	55	77	91	0.21
498555	13	35	68	86	94	0.19
498556	27	37	71	85	91	0.15
498557	18	42	75	89	95	0.16
498579	16	38	67	89	95	0.16
498580	36	57	81	91	96	0.10
498581	34	64	75	93	97	0.05

Table 141

ISIS No	0.0625 μM	0.125 μM	0.250 μM	0.500 μM	1.000 μM	IC ₅₀ (μM)
144367	0	9	26	49	77	0.47
494388	0	0	21	33	55	0.89
494389	0	15	22	50	79	0.46
494390	5	20	37	68	81	0.33
494391	7	20	32	54	68	0.46
494392	18	24	40	57	76	0.35
494466	33	45	58	69	82	0.16
494470	45	58	68	79	87	0.08
494472	37	50	60	69	83	0.13
494521	0	0	0	15	54	0.17
494525	0	0	2	28	65	0.85
494530	0	6	27	51	80	0.46
494535	0	7	24	53	74	0.49
494536	0	2	15	42	67	0.63

Table 142

ISIS No	0.0625 μM	0.125 μM	0.250 μM	0.500 μM	1.000 μM	IC ₅₀ (μM)
144367	0	4	16	26	77	0.65
498379	12	18	27	32	63	0.81
498408	0	11	46	50	77	0.41
498433	22	30	46	60	83	0.27
498434	39	29	25	47	78	0.40
498435	21	28	26	43	73	0.50
498517	44	48	63	70	84	0.11
498721	54	54	66	75	89	<0.06
498833	44	51	58	67	83	0.11
498859	0	29	14	35	66	0.69
498868	0	12	9	26	60	1.07
498875	0	30	31	53	78	0.40

499020	0	27	19	45	74	0.51
499041	0	12	10	37	65	0.77

As presented in the tables above, ISIS 494157 (SEQ ID NO: 12), ISIS 494158 (SEQ ID NO:13), ISIS 494159 (SEQ ID NO:14), ISIS 494160 (SEQ ID NO: 15), ISIS 494161 (SEQ ID NO:16), ISIS 494162 (SEQ ID NO: 17), ISIS 494163 (SEQ ID NO: 18), ISIS 494164 (SEQ ID NO: 19), ISIS 494165 (SEQ ID NO: 20), ISIS 494167 (SEQ ID NO: 22), ISIS 494168 (SEQ ID NO: 23), ISIS 494169 (SEQ ID NO: 24), ISIS 494170 (SEQ ID NO: 25), ISIS 494230 (SEQ ID NO: 105), ISIS 494243 (SEQ ID NO: 106), ISIS 494244 (SEQ ID NO: 107), ISIS 494283 (SEQ ID NO: 26), ISIS 494284 (SEQ ID NO: 27), ISIS 494285 (SEQ ID NO: 28), ISIS 494286 (SEQ ID NO: 29), ISIS 494287 (SEQ ID NO: 30), ISIS 494288 (SEQ ID NO: 31), ISIS 494290 (SEQ ID NO: 32), ISIS 494291 (SEQ ID NO: 33), ISIS 494292 (SEQ ID NO: 35), ISIS 494294 (SEQ ID NO: 36), ISIS 494299 (SEQ ID NO: 37), ISIS 494300 (SEQ ID NO: 38), ISIS 494301 (SEQ ID NO: 39), ISIS 494302 (SEQ ID NO: 40), ISIS 494303 (SEQ ID NO: 41), ISIS 494304 (SEQ ID NO: 42), ISIS 494305 (SEQ ID NO:43), ISIS 494306 (SEQ ID NO: 44), ISIS 494311 (SEQ ID NO: 47), ISIS 494334 (SEQ ID NO: 48), ISIS 494336 (SEQ ID NO: 49), ISIS 494337 (SEQ ID NO: 50), ISIS 494338 (SEQ ID NO: 133), ISIS 494371 (SEQ ID NO: 57), ISIS 494372 (SEQ ID NO: 58), ISIS 494373 (SEQ ID NO: 59), ISIS 494374 (SEQ ID NO: 60), ISIS 494375 (SEQ ID NO: 61), ISIS 494386 (SEQ ID NO: 62), ISIS 494389 (SEQ ID NO: 65), ISIS 494390 (SEQ ID NO: 66), ISIS 494392 (SEQ ID NO: 68), ISIS 494466 (SEQ ID NO: 108), ISIS 494470 (SEQ ID NO: 109), ISIS 494472 (SEQ ID NO: 110), ISIS 494521 (SEQ ID NO: 51), ISIS 494530 (SEQ ID NO: 53), ISIS 498229 (SEQ ID NO: 75), ISIS 498238 (SEQ ID NO: 76), ISIS 498239 (SEQ ID NO: 77), ISIS 498240 (SEQ ID NO: 78), ISIS 498241 (SEQ ID NO: 79), ISIS 498243 (SEQ ID NO: 81), ISIS 498379 (SEQ ID NO: 70), ISIS 498408 (SEQ ID NO: 71), ISIS 498433 (SEQ ID NO: 72), ISIS 498434 (SEQ ID NO: 73), ISIS 498435 (SEQ ID NO: 74), ISIS 498517 (SEQ ID NO: 85), ISIS 498523 (SEQ ID NO: 92), ISIS 498524 (SEQ ID NO: 93), ISIS 498525 (SEQ ID NO: 94), ISIS 498550 (SEQ ID NO: 97), ISIS 498580 (SEQ ID NO: 103), ISIS 498581 (SEQ ID NO: 104), ISIS 498721 (ATGCCTCGATAACTCCGTCC; SEQ ID NO: 134), ISIS 498833 (SEQ ID NO: 86), ISIS 498875 (SEQ ID NO: 89), and ISIS 499020 (SEQ ID NO: 90) were more potent than ISIS 144367 (SEQ ID NO: 11).

Example 116: Dose-dependent antisense inhibition of apo(a) in transgenic mouse primary hepatocytes

Potent gapmers from the studies described above were further selected and tested at various doses in transgenic mouse primary hepatocytes in a series of studies with similar culture conditions. Cells were plated at a density of 35,000 per well and transfected using electroporation with 0.049 μ M, 0.148 μ M, 0.444 μ M, 1.333 μ M, or 4.000 μ M concentrations of antisense oligonucleotide, as specified in tables below. After a treatment period of approximately 16 hours, RNA was isolated from the cells and apo(a) mRNA levels were measured by quantitative real-time PCR. Apo(a) primer probe set hAPO(a)12kB was used to measured

mRNA levels. Apo(a) mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN®. Results are presented as percent inhibition of apo(a), relative to untreated control cells.

The results of each of the studies are depicted in the tables presented below with each table representing a separate experiment. The half maximal inhibitory concentration (IC₅₀) of each oligonucleotide is also presented in the tables. Apo(a) mRNA levels were significantly reduced in a dose-dependent manner in antisense oligonucleotide treated cells. The potency of the newly designed oligos was compared with the benchmark oligonucleotide, ISIS 144367. As presented in the tables below, ISIS 494157 (SEQ ID NO: 12), ISIS 494158 (SEQ ID NO:13), ISIS 494159 (SEQ ID NO:14), ISIS 494160 (SEQ ID NO: 15), ISIS 494161 (SEQ ID NO:16), ISIS 494162 (SEQ ID NO: 17), ISIS 494163 (SEQ ID NO: 18), ISIS 494164 (SEQ ID NO: 19), ISIS 494230 (SEQ ID NO: 105), ISIS 494243 (SEQ ID NO: 106), ISIS 494244 (SEQ ID NO: 107), ISIS 494283 (SEQ ID NO: 26), ISIS 494284 (SEQ ID NO: 27), ISIS 494285 (SEQ ID NO: 28), ISIS 494286 (SEQ ID NO: 29), ISIS 494287 (SEQ ID NO: 30), ISIS 494290 (SEQ ID NO: 32), ISIS 494292 (SEQ ID NO: 35), ISIS 494300 (SEQ ID NO: 38), ISIS 494301 (SEQ ID NO: 39), ISIS 494302 (SEQ ID NO: 40), ISIS 494303 (SEQ ID NO: 41), ISIS 494304 (SEQ ID NO: 42), ISIS 494305 (SEQ ID NO: 43), ISIS 494306 (SEQ ID NO: 44), ISIS 494310 (SEQ ID NO: 46), ISIS 494311 (SEQ ID NO: 47), ISIS 494337 (SEQ ID NO: 50), ISIS 494371 (SEQ ID NO: 57), ISIS 494372 (SEQ ID NO: 58), ISIS 494375 (SEQ ID NO: 61), ISIS 494388 (SEQ ID NO: 64), ISIS 494389 (SEQ ID NO: 65), ISIS 494390 (SEQ ID NO: 66), ISIS 494392 (SEQ ID NO: 68), ISIS 494466 (SEQ ID NO: 108), ISIS 494470 (SEQ ID NO: 109), ISIS 494472 (SEQ ID NO: 110), ISIS 498238 (SEQ ID NO: 76), ISIS 498239 (SEQ ID NO: 77), ISIS 498433 (SEQ ID NO: 72), ISIS 498434 (SEQ ID NO: 73), ISIS 498435 (SEQ ID NO: 74), ISIS 498523 (SEQ ID NO: 92), ISIS 498524 (SEQ ID NO: 93), ISIS 498525 (SEQ ID NO: 94), ISIS 498580 (SEQ ID NO: 103), and ISIS 498581 (SEQ ID NO: 104) were more potent than ISIS 144367 (SEQ ID NO: 11).

Table 143

ISIS No	0.049 μM	0.148 μM	0.444 μM	1.333 μM	4.000 μM	IC ₅₀ (μM)
144367	0	26	67	89	92	0.32
494157	23	50	83	96	96	0.15
494158	26	62	85	96	96	0.11
494159	42	65	87	95	94	0.07
494160	51	70	88	94	94	<0.05
494161	36	67	87	95	96	0.08
494162	40	69	89	94	95	0.07
494163	41	57	87	95	94	0.08
494164	15	43	75	93	96	0.20
494230	39	77	94	99	99	0.05
494243	39	76	92	98	99	0.06
494244	58	79	91	97	99	0.02

494283	18	45	80	93	91	0.18
494284	9	53	80	95	94	0.18

Table 144

ISIS No	0.049 μM	0.148 μM	0.444 μM	1.333 μM	4.000 μM	IC ₅₀ (μM)
144367	21	40	79	94	93	0.18
494285	53	68	90	97	97	<0.05
494286	46	69	89	96	97	0.05
494287	31	38	79	94	95	0.15
494290	22	53	74	93	94	0.16
494292	37	51	81	93	95	0.11
494294	22	40	72	91	94	0.19
494299	15	43	75	93	95	0.20
494300	25	38	79	95	95	0.17
494301	23	48	82	92	95	0.15
494302	26	59	86	93	94	0.12
494303	10	58	84	92	91	0.16
494304	25	62	83	93	93	0.12

Table 145

ISIS No	0.049 μM	0.148 μM	0.444 μM	1.333 μM	4.000 μM	IC ₅₀ (μM)
144367	23	40	70	90	94	0.19
494305	20	48	82	93	95	0.16
494306	26	53	78	91	92	0.14
494310	36	50	79	88	92	0.12
494311	38	50	74	93	95	0.12
494334	20	42	73	90	94	0.19
494336	5	39	74	92	95	0.23
494337	23	51	87	96	96	0.14
494338	12	42	82	93	95	0.19
494371	28	49	82	94	94	0.14
494372	28	54	81	93	88	0.13
494373	21	28	67	86	92	0.25
494375	26	40	77	85	92	0.18

Table 146

ISIS No	0.049 μM	0.148 μM	0.444 μM	1.333 μM	4.000 μM	IC ₅₀ (μM)
144367	5	33	65	78	81	0.32
494388	30	32	60	82	86	0.25
494389	30	45	69	84	84	0.17
494390	32	47	67	83	87	0.16
494392	23	38	54	79	82	0.31
494466	48	67	86	91	95	0.04
494470	74	87	92	96	98	<0.05
494472	69	84	92	96	97	<0.05
494544	5	18	49	74	79	0.48
498238	25	51	76	92	96	0.15
498239	25	62	83	93	97	0.12
498379	5	21	53	71	81	0.55
498408	1	38	63	79	80	0.32
498433	23	43	70	77	79	0.21

Table 147

ISIS No	0.049 μM	0.148 μM	0.444 μM	1.333 μM	4.000 μM	IC ₅₀ (μM)
144367	0	40	76	90	93	0.26
498434	32	44	64	78	84	0.20
498435	24	42	64	77	79	0.23
498517	28	23	53	81	85	0.45
498523	50	64	81	90	93	<0.05
498524	53	70	84	93	96	<0.05
498525	38	55	80	92	96	0.09
498550	12	18	62	81	83	0.33
498557	13	33	67	79	83	0.33
498579	6	42	69	80	85	0.31
498580	6	46	76	82	83	0.23
498581	5	40	78	81	84	0.25
498721	40	31	58	78	83	0.35
498833	21	20	58	80	90	0.44

Example 117: Antisense inhibition of human apo(a) in transgenic mouse primary hepatocytes

5 Additional antisense oligonucleotides were newly designed targeting an apo(a) nucleic acid and were tested for their effects on apo(a) mRNA *in vitro*. The antisense oligonucleotides were tested in a series of experiments that had similar culture conditions. Primary hepatocytes from human apo(a) transgenic mice were used in this study. Hepatocytes at a density of 35,000 cells per well were transfected using

electroporation with 1,000 nM antisense oligonucleotide. After a treatment period of approximately 24 hours, RNA was isolated from the cells and apo(a) mRNA levels were measured by quantitative real-time PCR. Human primer probe set hAPO(a)12kB was used to measure mRNA levels. Apo(a) mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN®. The results for each experiment are presented in separate tables shown below. ISIS 144367 was also included in the studies for comparison. Results are presented as percent inhibition of apo(a), relative to untreated control cells. A total of 231 antisense oligonucleotides were tested under these culture conditions. Only those antisense oligonucleotides that were selected for further studies are presented below.

The newly designed chimeric antisense oligonucleotides were designed as 3-10-4 MOE gapmers.

The gapmers are 17 nucleosides in length, wherein the central gap segment comprises of ten 2'-deoxynucleosides and is flanked by wing segments on the 5' direction and the 3' direction comprising three nucleosides and four nucleosides respectively. Each nucleoside in the 5' wing segment and each nucleoside in the 3' wing segment has a 2'-MOE modification. The internucleoside linkages throughout each gapmer are phosphorothioate (P=S) linkages. All cytosine residues throughout each gapmer are 5-methylcytosines.

The apo(a) target sequence contains multiple Kringle repeat sequences, therefore, an antisense oligonucleotide may target one or more regions of apo(a) depending whether on the oligonucleotide targets a Kringle sequence or not. "Start site" indicates the 5'-most nucleoside to which the gapmer is targeted in the human sequence. "Stop site" indicates the 3'-most nucleoside to which the gapmer is targeted human sequence. An apo(a) antisense oligonucleotide may have more than one "Start site" or "Stop site" depending on whether or not it targets a Kringle repeat.

Most gapmers listed in the tables are targeted with 100% complementarity to multiple regions of either the human apo(a) mRNA, designated herein as SEQ ID NO: 1 (GENBANK Accession No. NM_005577.2) or the human apo(a) genomic sequence, designated herein as SEQ ID NO: 2 (GENBANK Accession No. NT_007422.12 truncated from nucleotides 3230000 to 3380000), or both. 'n/a' indicates that the antisense oligonucleotide does not target that particular sequence with 100% complementarity.

Table 148

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	64	21210	21229	11
510542	241	257	CCTGTGACAGTGGTGGA	79	21202	21218	111
	583	599	CCTGTGACAGTGGTGGA		26693	26709	
	925	941	CCTGTGACAGTGGTGGA		32240	32256	
	1609	1625	CCTGTGACAGTGGTGGA		43333	43349	
	1951	1967	CCTGTGACAGTGGTGGA		48877	48893	
	2293	2309	CCTGTGACAGTGGTGGA		54423	54439	

	3319	3335	CCTGTGACAGTGGTGGGA		72040	72056	
	4663	4679	CCTGTGACAGTGGTGGGA		94404	94420	
	5005	5021	CCTGTGACAGTGGTGGGA		115515	115531	
510543	242	258	TCCTGTGACAGTGGTGG	75	21203	21219	112
	584	600	TCCTGTGACAGTGGTGG		26694	26710	
	926	942	TCCTGTGACAGTGGTGG		32241	32257	
	1610	1626	TCCTGTGACAGTGGTGG		43334	43350	
	1952	1968	TCCTGTGACAGTGGTGG		48878	48894	
	2294	2310	TCCTGTGACAGTGGTGG		54424	54440	
	3320	3336	TCCTGTGACAGTGGTGG		72041	72057	
	4664	4680	TCCTGTGACAGTGGTGG		94405	94421	
	5006	5022	TCCTGTGACAGTGGTGG		115516	115532	
510544	243	259	TTCCTGTGACAGTGGTG	73	21204	21220	113
	585	601	TTCCTGTGACAGTGGTG		26695	26711	
	927	943	TTCCTGTGACAGTGGTG		32242	32258	
	1611	1627	TTCCTGTGACAGTGGTG		43335	43351	
	1953	1969	TTCCTGTGACAGTGGTG		48879	48895	
	2295	2311	TTCCTGTGACAGTGGTG		54425	54441	
	3321	3337	TTCCTGTGACAGTGGTG		72042	72058	
	4665	4681	TTCCTGTGACAGTGGTG		94406	94422	
	5007	5023	TTCCTGTGACAGTGGTG		115517	115533	
510545	244	260	CTTCCTGTGACAGTGGT	65	21205	21221	114
	586	602	CTTCCTGTGACAGTGGT		26696	26712	
	928	944	CTTCCTGTGACAGTGGT		32243	32259	
	1612	1628	CTTCCTGTGACAGTGGT		43336	43352	
	1954	1970	CTTCCTGTGACAGTGGT		48880	48896	
	2296	2312	CTTCCTGTGACAGTGGT		54426	54442	
	3322	3338	CTTCCTGTGACAGTGGT		72043	72059	
	3664	3680	CTTCCTGTGACAGTGGT		77585	77601	
	4666	4682	CTTCCTGTGACAGTGGT		94407	94423	
	5008	5024	CTTCCTGTGACAGTGGT		115518	115534	
510546	245	261	CCTTCCTGTGACAGTGG	74	21206	21222	115
	3665	3681	CCTTCCTGTGACAGTGG		77586	77602	
	4667	4683	CCTTCCTGTGACAGTGG		94408	94424	
	5009	5025	CCTTCCTGTGACAGTGG		115519	115535	
510547	246	262	TCCTTCCTGTGACAGTG	77	21207	21223	116
	3666	3682	TCCTTCCTGTGACAGTG		77587	77603	
	4668	4684	TCCTTCCTGTGACAGTG		94409	94425	
	5010	5026	TCCTTCCTGTGACAGTG		115520	115536	
510548	247	263	GTCCTTCCTGTGACAGT	73	21208	21224	117
	3667	3683	GTCCTTCCTGTGACAGT		77588	77604	
	4669	4685	GTCCTTCCTGTGACAGT		94410	94426	
	5011	5027	GTCCTTCCTGTGACAGT		115521	115537	

510549	248	264	GGTCCTTCCTGTGACAG	67	21209	21225	118
	4670	4686	GGTCCTTCCTGTGACAG		94411	94427	
510595	632	648	CCGACTATGCGAGTGTG	76	26742	26758	119
	974	990	CCGACTATGCGAGTGTG		32289	32305	
	1316	1332	CCGACTATGCGAGTGTG		37836	37852	
	1658	1674	CCGACTATGCGAGTGTG		43382	43398	
	2000	2016	CCGACTATGCGAGTGTG		48926	48942	
	2342	2358	CCGACTATGCGAGTGTG		54472	54488	
	2684	2700	CCGACTATGCGAGTGTG		60027	60043	
	3026	3042	CCGACTATGCGAGTGTG		66543	66559	
510597	634	650	GTCCGACTATGCGAGTG	70	26744	26760	120
	976	992	GTCCGACTATGCGAGTG		32291	32307	
	1318	1334	GTCCGACTATGCGAGTG		37838	37854	
	1660	1676	GTCCGACTATGCGAGTG		43384	43400	
	2002	2018	GTCCGACTATGCGAGTG		48928	48944	
	2344	2360	GTCCGACTATGCGAGTG		54474	54490	
	2686	2702	GTCCGACTATGCGAGTG		60029	60045	
	3028	3044	GTCCGACTATGCGAGTG		66545	66561	
510598	635	651	GGTCCGACTATGCGAGT	70	26745	26761	121
	977	993	GGTCCGACTATGCGAGT		32292	32308	
	1319	1335	GGTCCGACTATGCGAGT		37839	37855	
	1661	1677	GGTCCGACTATGCGAGT		43385	43401	
	2003	2019	GGTCCGACTATGCGAGT		48929	48945	
	2345	2361	GGTCCGACTATGCGAGT		54475	54491	
	2687	2703	GGTCCGACTATGCGAGT		60030	60046	
	3029	3045	GGTCCGACTATGCGAGT		66546	66562	

Table 149

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	83	21210	21229	11
510783	6400	6416	GTCAGACCTTAAAAGCT	75	140056	140072	122
512944	3561	3577	AAGCCTCTGTGCTTGGGA	81	76246	76262	123
512947	3560	3576	AGCCTCTGTGCTTGGAT	85	76245	76261	124
512958	3559	3575	GCCTCTGTGCTTGGATC	82	76244	76260	125
512959	3585	3601	GCTCCGTTGGTGCTTCT	77	n/a	n/a	126

Table 150

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
144367	249	268	GGCAGGTCCTTCCTGTGACA	76	21210	21229	11
510701	4217	4233	CTCTGTGCTTGGAACTG	78	85147	85163	127

510702	219	235	TGCCTCGATAACTCTGT	79	21180	21196	128
	561	577			26671	26687	
	903	919			32218	32234	
	1245	1261			37765	37781	
	1587	1603			43311	43327	
	1929	1945			48855	48871	
	2271	2287			54401	54417	
	2613	2629			59956	59972	
	4299	4315			86472	86488	
510704	563	579	TGTGCCTCGATAACTCT	80	26673	26689	129
	905	921			32220	32236	
	1247	1263			37767	37783	
	1589	1605			43313	43329	
	1931	1947			48857	48873	
	2273	2289			54403	54419	
	2615	2631			59958	59974	
	4301	4317			86474	86490	
	4985	5001			115495	115511	
510757	4929	4945	GCTCAGTTGGTGCTGCT	74	n/a	n/a	130

Example 118: Dose-dependent antisense inhibition of apo(a) in transgenic mouse primary hepatocytes

Potent gapmers from the studies described above were further selected and tested at various doses in transgenic mouse primary hepatocytes in a series of studies with similar culture conditions. Cells were plated at a density of 35,000 per well and transfected using electroporation with 0.156 μ M, 0.313 μ M, 0.625 μ M, 1.250 μ M, 2.500 μ M, or 5.000 μ M concentrations of antisense oligonucleotide, as specified in the tables below. After a treatment period of approximately 16 hours, RNA was isolated from the cells and apo(a) mRNA levels were measured by quantitative real-time PCR. Apo(a) primer probe set hAPO(a)12kB was used to measured mRNA levels. Apo(a) mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN®. Results are presented as percent inhibition of apo(a), relative to untreated control cells.

The results of each of the studies are depicted in the tables presented below with each study represented in a separate table. The half maximal inhibitory concentration (IC_{50}) of each oligonucleotide is also presented in the tables.

Table 151

ISIS No	0.156 μ M	0.312 μ M	0.625 μ M	1.250 μ M	2.500 μ M	5.000 μ M	IC_{50} (μ M)
144367	28	55	70	83	90	92	0.31
510542	33	58	75	87	89	90	0.27

510543	33	45	68	78	89	89	0.34
510544	33	50	65	78	88	90	0.33
510545	33	58	76	87	91	90	0.26
510546	39	62	76	87	89	91	0.22
510547	36	66	82	84	86	91	0.22
510548	50	70	82	91	88	90	0.13
510549	32	59	73	85	86	90	0.27
510595	26	57	78	88	90	90	0.29
510597	30	53	76	85	89	89	0.30

Table 152

ISIS No	0.156 μM	0.312 μM	0.625 μM	1.250 μM	2.500 μM	5.000 μM	IC ₅₀ (μM)
144367	36	52	78	87	93	94	0.26
510598	48	58	81	88	93	92	0.18
510701	45	59	78	87	95	95	0.18
510702	49	63	75	90	94	95	0.15
510704	55	67	80	93	94	95	<0.16
510757	34	48	68	79	90	93	0.33
510783	21	32	51	58	78	84	0.69
512944	57	72	81	91	96	97	<0.16
512947	64	74	86	92	96	97	<0.16
512958	48	69	83	91	96	97	0.13
512959	39	59	76	84	93	93	0.22

Table 153

ISIS No	0.156 μM	0.312 μM	0.625 μM	1.250 μM	2.500 μM	5.000 μM	IC ₅₀ (μM)
144367	41	58	75	81	88	87	0.22
510542	38	54	69	74	85	83	0.27
510545	21	43	73	77	80	78	0.39
510546	37	58	73	81	83	81	0.24
510547	38	58	72	79	84	86	0.24
510548	40	63	77	79	81	84	0.21
510549	37	47	67	77	81	83	0.31
510595	34	66	73	81	80	75	0.23
510597	39	59	74	83	76	77	0.23

Table 154

ISIS No	0.156 μM	0.312 μM	0.625 μM	1.250 μM	2.500 μM	5.000 μM	IC ₅₀ (μM)
144367	33	60	72	83	81	81	0.26
510598	47	62	75	75	76	76	0.18
510701	41	67	80	87	92	91	0.19
510702	51	64	77	80	80	83	0.13
510704	54	61	77	84	89	80	0.12
512944	71	74	81	88	92	94	0.02
512947	65	77	86	90	93	95	0.03
512958	63	73	84	92	93	96	0.06
512959	39	62	80	82	86	82	0.22

Apo(a) mRNA levels were significantly reduced in a dose-dependent manner in antisense oligonucleotide-treated cells. The potency of the newly designed oligonucleotides was compared with the benchmark oligonucleotide, ISIS 144367. As presented in the tables above, ISIS 510542 (SEQ ID NO: 111), ISIS 510545 (SEQ ID NO: 114), ISIS 510546 (SEQ ID NO: 115), ISIS 510547 (SEQ ID NO: 116), ISIS 510548 (SEQ ID NO: 117), ISIS 510549 (SEQ ID NO: 118), ISIS 510595 (SEQ ID NO: 119), ISIS 510597 (SEQ ID NO: 120), ISIS 510598 (SEQ ID NO: 121), ISIS 510701 (SEQ ID NO: 127), ISIS 510702 (SEQ ID NO: 128), ISIS 510704 (SEQ ID NO: 129), ISIS 512944 (SEQ ID NO: 123), ISIS 512947 (SEQ ID NO: 124), ISIS 512958 (SEQ ID NO: 125), and ISIS 512959 (SEQ ID NO: 126) were more potent than ISIS 144367 (SEQ ID NO: 11).

Example 119: Effect of *in vivo* antisense inhibition of human apo(a) in human apo(a) transgenic mice

Transgenic mice with the human apo(a) gene (Frazer, K.A. et al., Nat. Genet. 1995. 9: 424-431) were utilized in the studies described below. ISIS antisense oligonucleotides that demonstrated statistically significant inhibition of apo(a) mRNA *in vitro* as described above were evaluated further in this model.

Study 1

Female human apo(a) transgenic mice were maintained on a 12-hour light/dark cycle and fed *ad libitum* normal lab chow. The mice were divided into treatment groups consisting of 4 mice each. The groups received intraperitoneal injections of ISIS 494159, ISIS 494160, ISIS 494161, ISIS 494162, ISIS 494163, ISIS 494230, ISIS 494243, ISIS 494244, ISIS 494283, ISIS 494284, ISIS 494285, ISIS 494286, ISIS 494301, ISIS 494302, ISIS 494304, ISIS 494466, ISIS 494470, ISIS 494472, ISIS 498239, ISIS 498408, ISIS 498517, ISIS 494158, ISIS 494311, ISIS 494337, ISIS 494372, ISIS 498238, ISIS 498523, ISIS 498525, ISIS 510548, ISIS 512944, ISIS 512947, or ISIS 512958 at a dose of 25 mg/kg twice a week for 2 weeks. One group of mice received intraperitoneal injections of PBS twice a week for 2 weeks. The PBS group served as the

control group. Two days following the final dose, the mice were euthanized, organs harvested and analyses done.

Inhibition of human apo(a) mRNA

Total RNA was extracted from the livers of some of the treatment groups, and human apo(a) mRNA was quantitated by RT-PCR. The results are presented in the table below, expressed as percent inhibition of apo(a) mRNA compared to the PBS control.

Table 155

Percent inhibition of human apo(a) mRNA in transgenic mice

ISIS No	% inhibition
144367	98
494159	100
494160	95
494161	98
494162	100
494163	100
494230	96
494243	99
494244	99
494283	100
494284	100
494285	100
494286	98
494301	99
494302	96
494304	94
494466	97
494470	93
494472	98
498239	72
498408	100
498517	98

The data demonstrates significant inhibition of apo(a) mRNA by several ISIS oligonucleotides. ISIS 494159 (SEQ ID NO: 14), ISIS 494162 (SEQ ID NO: 17), ISIS 494163 (SEQ ID NO: 18), ISIS 494243 (SEQ ID NO: 106), ISIS 494244 (SEQ ID NO: 107), ISIS 494283 (SEQ ID NO: 26), ISIS 494284 (SEQ ID NO: 27), ISIS 494285 (SEQ ID NO: 28), ISIS 494301 (SEQ ID NO: 39), and ISIS 498408 (SEQ ID NO: 71) were more potent than the benchmark ISIS 144367 (SEQ ID NO: 11).

Inhibition of human apo(a) protein

Plasma human apo(a) protein was measured from all treatment groups using an Apo(a) ELISA kit (Mercodia 10-1106-01, Uppsala, Sweden). The results are presented in the table below, expressed as percent inhibition of apo(a) mRNA compared to the PBS control.

5

Table 156

Percent inhibition of human apo(a) protein in transgenic mice

ISIS No	% inhibition
144367	86
494159	86
494160	0
494161	82
494162	84
494163	82
494230	60
494243	84
494244	87
494283	98
494284	98
494285	89
494286	89
494301	93
494302	88
494304	83
494466	76
494470	73
494472	72
498239	54
498408	84
498517	56
494158	71
494311	83
494337	80
494372	78
498238	58
498523	47
498525	58
510548	74
512944	18
512947	65
512958	72

The data demonstrates significant inhibition of apo(a) mRNA by several ISIS oligonucleotides. ISIS 494159 (SEQ ID NO: 14), ISIS 494244 (SEQ ID NO: 82), ISIS 494283 (SEQ ID NO: 26), ISIS 494284 (SEQ ID NO: 27), ISIS 494285 (SEQ ID NO: 28), ISIS 494286 (SEQ ID NO: 29), ISIS 494301 (SEQ ID NO: 39), and ISIS 494302 (SEQ ID NO: 40) were as potent as or more potent than the benchmark ISIS 144367 (SEQ ID NO: 11)..

Study 2

ISIS 494159, ISIS 494161, ISIS 494162, ISIS 494163, and ISIS 494243 were further evaluated in this transgenic model. ISIS 144367 was included for comparison.

Treatment

Female human apo(a) transgenic mice were divided into treatment groups consisting of 4 mice each. The groups received intraperitoneal injections of ISIS 144367, ISIS 494159, ISIS 494161, ISIS 494162, ISIS 494163, or ISIS 494243 at doses of 1.5 mg/kg, 5 mg/kg, 15 mg/kg, or 50 mg/kg twice a week for 2 weeks. One group of mice received intraperitoneal injections of PBS twice a week for 2 weeks. The PBS group served as the control group. Two days following the final dose, the mice were euthanized, organs harvested and analyses done.

Inhibition of human apo(a) mRNA

Total RNA was extracted from the livers of the treatment groups, and human apo(a) mRNA was quantitated by RT-PCR. The results are presented in the table below, expressed as percent inhibition of apo(a) mRNA compared to the PBS control.

Table 157

Dose-dependent inhibition of human apo(a) mRNA in transgenic mice

ISIS No	Dose (mg/kg/wk)	% inhibition	ED ₅₀
144367	100	71	31
	30	42	
	10	0	
	3	5	
494159	100	91	5
	30	67	
	10	48	
	3	39	
494161	100	82	6

	30	49	
	10	61	
	3	30	
494162	100	90	5
	30	67	
	10	58	
	3	25	
494163	100	83	5
	30	66	
	10	58	
	3	21	
494243	100	80	32
	30	26	
	10	0	
	3	6	

The data demonstrates significant inhibition of apo(a) mRNA by several ISIS oligonucleotides. ISIS 494159 (SEQ ID NO: 14), ISIS 494161 (SEQ ID NO: 16), 494162 (SEQ ID NO:17), and ISIS 94163 (SEQ ID NO: 18) were more efficacious than the benchmark ISIS 144367 (SEQ ID NO: 11). *Reduction of human apo(a) protein levels*

Blood was collected from the treatment groups, and human apo(a) protein levels were quantitated by an Apo(a) ELISA kit (Mercodia 10-1106-01, Uppsala, Sweden). The results are presented in the table below, expressed as percent reduction of apo(a) protein levels compared to the PBS control.

Table 158

Dose-dependent inhibition of human apo(a) protein in transgenic mice

ISIS No	Dose (mg/kg/wk)	% inhibition	ED ₅₀
144367	100	73	71
	30	0	
	10	6	
	3	69	
494159	100	88	2
	30	88	
	10	85	
	3	36	

494161	100	90	2
	30	85	
	10	73	
	3	44	
494162	100	89	3
	30	78	
	10	76	
	3	24	
494163	100	90	3
	30	86	
	10	60	
	3	37	
494243	100	61	174
	30	0	
	10	0	
	3	0	

The data demonstrates significant reduction of apo(a) plasma protein levels by several ISIS oligonucleotides. ISIS 494159 (SEQ ID NO: 14), ISIS 494161 (SEQ ID NO: 16), ISIS 494162 (SEQ ID NO: 17), and ISIS 494163 (SEQ ID NO: 18) were more efficacious than the benchmark ISIS 144367 (SEQ ID NO: 11).

5 Study 3

ISIS 494244, ISIS 494283, and ISIS 494284 were further evaluated in this model. ISIS 144367 was included for comparison.

Treatment

10 Female human apo(a) transgenic mice were divided into treatment groups consisting of 4 mice each. The groups received intraperitoneal injections of ISIS 144367, ISIS 494244, ISIS 494283, or ISIS 494284 at doses of 0.75 mg/kg, 2.5 mg/kg, 7.5 mg/kg, or 25 mg/kg twice a week for 2 weeks. One group of mice received intraperitoneal injections of PBS twice a week for 2 weeks. The PBS group served as the control group. Two days following the final dose, the mice were euthanized, organs harvested and analyses done.

15 *Inhibition of human apo(a) mRNA*

Total RNA was extracted from the livers of the treatment groups, and human apo(a) mRNA was quantitated by RT-PCR. The results are presented in the table below, expressed as percent inhibition of apo(a) mRNA compared to the PBS control.

Table 159

Dose-dependent inhibition of human apo(a) mRNA in transgenic mice

ISIS No	Dose (mg/kg/wk)	% inhibition	ED ₅₀
144367	50	75	22
	15	60	
	5	0	
	1.5	0	
494244	50	73	18
	15	41	
	5	34	
	1.5	0	
494283	50	74	16
	15	52	
	5	24	
	1.5	0	
494284	50	73	16
	15	58	
	5	17	
	1.5	2	

The data demonstrates significant inhibition of apo(a) mRNA by several ISIS oligonucleotides. ISIS 494244 (SEQ ID NO: 107), ISIS 494283 (SEQ ID NO: 26), and ISIS 494284 (SEQ ID NO: 27) were more efficacious than the benchmark, ISIS 144367 (SEQ ID NO: 11).

Reduction of human apo(a) protein levels

Blood was collected from the treatment groups, and human apo(a) protein levels were quantitated by an Apo(a) ELISA kit (Mercodia 10-1106-01, Uppsala, Sweden). The results are presented in the table below, expressed as percent reduction of apo(a) protein levels compared to the PBS control.

Table 160

Dose-dependent inhibition of human apo(a) plasma protein in transgenic mice

ISIS No	Dose (mg/kg/wk)	% inhibition	ED ₅₀
144367	50	64	16
	15	14	
	5	0	
	1.5	0	
494244	50	67	2
	15	60	
	5	58	
	1.5	0	
494283	50	64	4

	15	65	
	5	64	
	1.5	69	
494284	50	66	4
	15	63	
	5	51	
	1.5	54	

The data demonstrates significant reduction of apo(a) plasma protein levels by several ISIS oligonucleotides. ISIS 494244 (SEQ ID NO: 107), ISIS 494283 (SEQ ID NO: 26), and ISIS 494284 (SEQ ID NO: 27) were more efficacious than the benchmark, ISIS 144367 (SEQ ID NO: 11).

5 Study 4

ISIS 494285, ISIS 494286, ISIS 494301, ISIS 494302, and ISIS 494311 were further evaluated in this model.

Treatment

Male human apo(a) transgenic mice were divided into treatment groups consisting of 4 mice each. Each such group received intraperitoneal injections of ISIS 494285, ISIS 494286, ISIS 494301, ISIS 494302, or ISIS 494311 at doses of 5 mg/kg, 15 mg/kg, or 50 mg/kg once a week for 2 weeks. One group of 3 mice received intraperitoneal injections of PBS once a week for 2 weeks. The PBS group served as the control group. Two days following the final dose, the mice were euthanized, organs harvested and analyses done.

Inhibition of human apo(a) mRNA

Total RNA was extracted from the livers of the treatment groups, and human apo(a) mRNA was quantitated by RT-PCR. The results are presented in the table below, expressed as percent inhibition of apo(a) mRNA compared to the PBS control. The data demonstrates significant inhibition of apo(a) mRNA by ISIS 494285 (SEQ ID NO: 28), ISIS 494286 (SEQ ID NO: 29), ISIS 494301 (SEQ ID NO: 39), ISIS 494302 (SEQ ID NO: 40) and ISIS 494311 (SEQ ID NO: 47).

Table 161

Dose-dependent inhibition of human Apo(a) mRNA in transgenic mice

ISIS No	Dose (mg/kg/wk)	% inhibition	ED ₅₀
494285	50	98	1
	15	97	
	5	79	
494286	50	97	1
	15	91	

	5	80	
494301	50	98	3
	15	96	
	5	59	
494302	50	98	2
	15	88	
	5	72	
494311	50	99	1
	15	96	
	5	87	

Reduction of human apo(a) protein levels

Blood was collected from the treatment groups, and human apo(a) protein levels were quantitated by an Apo(a) ELISA kit (Mercodia 10-1106-01, Uppsala, Sweden). The results are presented in the table below, expressed as percent reduction of apo(a) protein levels compared to the PBS control. The data demonstrates significant reduction of apo(a) plasma protein levels by ISIS 494285, ISIS 494286, ISIS 494301, ISIS 494302 and ISIS 494311.

Table 162

Dose-dependent inhibition of human apo(a) protein in transgenic mice

ISIS No	Dose (mg/kg/wk)	% inhibition	ED ₅₀
494285	50	88	2
	15	88	
	5	72	
494286	50	90	2
	15	85	
	5	75	
494301	50	89	5
	15	86	
	5	38	
494302	50	90	3
	15	82	
	5	61	
494311	50	90	3
	15	82	
	5	69	

10 Study 5

ISIS 494372, ISIS 498524, ISIS 498581, ISIS 498721, and ISIS 498833 were further evaluated in this model.

Treatment

Female human apo(a) transgenic mice were divided into treatment groups consisting of 4 mice each. The groups received intraperitoneal injections of ISIS 494372, ISIS 498524, ISIS 498581, ISIS 498721, or ISIS 498833 at doses of 5 mg/kg, 15 mg/kg, or 50 mg/kg once a week for 2 weeks. One group of 3 mice received intraperitoneal injections of PBS once a week for 2 weeks. The PBS group served as the control group. Two days following the final dose, the mice were euthanized, organs harvested and analyses done.

Inhibition of human apo(a) mRNA

Total RNA was extracted from the livers of the treatment groups, and human apo(a) mRNA was quantitated by RT-PCR. The results are presented in the table below, expressed as percent inhibition of apo(a) mRNA compared to the PBS control. The data demonstrates significant inhibition of apo(a) mRNA by ISIS 494372 (SEQ ID NO: 28), ISIS 498524 (SEQ ID NO: 93), ISIS 498581 (SEQ ID NO: 104), and ISIS 498721 (ATGCCTCGATAACTCCGTCC; SEQ ID NO: 134).

Table 163

Dose-dependent inhibition of human Apo(a) mRNA in transgenic mice

ISIS No	Dose (mg/kg/wk)	% inhibition	ED ₅₀
494372	50	88	18
	15	49	
	5	0	
498524	50	83	8
	15	74	
	5	34	
498581	50	98	7
	15	58	
	5	48	
498721	50	97	14
	15	68	
	5	0	
498833	50	61	155
	15	0	
	5	17	

Reduction of human apo(a) protein levels

Blood was collected from the treatment groups, and human apo(a) protein levels were quantitated by an Apo(a) ELISA kit (Mercodia 10-1106-01, Uppsala, Sweden). The results are presented in the table below, expressed as percent reduction of apo(a) protein levels compared to the PBS control. The data demonstrates significant reduction of apo(a) plasma protein levels by ISIS 494372 (SEQ ID NO: 28), ISIS 498581 (SEQ ID NO: 104), and ISIS 498721 (ATGCCTCGATAACTCCGTCC; SEQ ID NO: 134).

Table 164

Dose-dependent inhibition of human apo(a) protein in transgenic mice

ISIS No	Dose (mg/kg/wk)	% inhibition	ED ₅₀
494372	50	68	32
	15	25	
	5	12	
498524	50	38	118
	15	0	
	5	0	
498581	50	79	9
	15	52	
	5	49	
498721	50	81	10
	15	63	
	5	29	
498833	50	15	738
	15	0	
	5	67	

10 **Example 120: Tolerability of antisense oligonucleotides targeting human apo(a) in rodent models**

Gapmer antisense oligonucleotides targeting human apo(a) were selected from the studies described above for tolerability studies in CD1 mice and in Sprague Dawley rats. Rodents do not express endogenous apo(a), hence these studies tested the tolerability of each human antisense oligonucleotide in an animal rather than any phenotypic changes that may be caused by inhibiting apo(a) in the animal.

15 Tolerability in CD1 mice: Study 1

CD1® mice (Charles River, MA) are a multipurpose mice model, frequently utilized for safety and efficacy testing. The mice were treated with ISIS antisense oligonucleotides selected from studies described above and evaluated for changes in the levels of various plasma chemistry markers.

Treatment

Groups of male CD1 mice were injected subcutaneously twice a week for 6 weeks with 50 mg/kg of ISIS 494159, ISIS 494161, ISIS 494162, ISIS 494244, ISIS 494283, ISIS 494284, ISIS 494285, ISIS 494286, ISIS 494301, ISIS 494302, ISIS 494311, ISIS 494337, ISIS 494372, and ISIS 510548. One group of six-week old male CD1 mice was injected subcutaneously twice a week for 6 weeks with PBS. Mice were euthanized 48 hours after the last dose, and organs and plasma were harvested for further analysis.

Plasma chemistry markers

To evaluate the effect of ISIS oligonucleotides on liver and kidney function, plasma levels of transaminases, bilirubin, albumin, creatinine, and BUN were measured using an automated clinical chemistry analyzer (Hitachi Olympus AU400e, Melville, NY). The results are presented in the table below. ISIS oligonucleotides that caused changes in the levels of any of the liver or kidney function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 165

Plasma chemistry markers of CD1 mice

	ALT (IU/L)	AST (IU/L)	Albumin (g/dL)	BUN (mg/dL)	Creatinine (mg/dL)	Bilirubin (mg/dL)
PBS	38	71	2.9	25.2	0.16	0.15
ISIS 494159	615	525	2.7	23.9	0.11	0.20
ISIS 494161	961	670	2.6	23.7	0.15	0.14
ISIS 494162	1373	1213	2.7	23.7	0.14	0.18
ISIS 494283	237	242	2.5	26.2	0.14	0.13
ISIS 494284	192	307	2.3	27.1	0.14	0.10
ISIS 494285	582	436	2.3	25.4	0.16	0.11
ISIS 494286	191	227	2.5	21.1	0.12	0.15
ISIS 494301	119	130	2.7	26.4	0.15	0.12
ISIS 494302	74	96	2.8	24.8	0.14	0.15
ISIS 494311	817	799	2.7	28.7	0.12	0.17
ISIS 494337	722	397	2.5	20.0	0.13	0.11
ISIS 494372	73	164	2.6	28.5	0.16	0.11
ISIS 510548	2819	2245	3.1	26.0	0.15	0.15

Organ weights

Liver, spleen and kidney weights were measured at the end of the study, and are presented in the table below. ISIS oligonucleotides that caused any changes in organ weights outside the expected range for antisense oligonucleotides were excluded from further studies.

Table 166

Organ weights of CD1 mice (g)

	Kidney	Liver	Spleen
PBS	0.68	2.0	0.13
ISIS 494159	0.68	3.0	0.21
ISIS 494161	0.62	3.5	0.20
ISIS 494162	0.60	3.3	0.20
ISIS 494283	0.65	2.8	0.24
ISIS 494284	0.69	2.7	0.29
ISIS 494285	0.59	3.2	0.21
ISIS 494286	0.64	2.8	0.25
ISIS 494301	0.72	3.0	0.43
ISIS 494302	0.63	2.3	0.23
ISIS 494311	0.61	3.2	0.19
ISIS 494337	0.56	2.3	0.17
ISIS 494372	0.60	2.5	0.27
ISIS 510548	0.55	3.7	0.20

5 Tolerability in Sprague Dawley rats

Sprague-Dawley rats are a multipurpose model used for safety and efficacy evaluations. The rats were treated with ISIS antisense oligonucleotides selected from studies described above and evaluated for changes in the levels of various plasma chemistry markers.

Treatment

- 10 Groups of male Sprague Dawley rats were injected subcutaneously twice a week for 8 weeks with 30 mg/kg of ISIS 494159, ISIS 494161, ISIS 494162, ISIS 494244, ISIS 494283, ISIS 494284, ISIS 494285, ISIS 494286, ISIS 494301, ISIS 494302, ISIS 494311, ISIS 494337, ISIS 494372, and ISIS 510548. One group of six male Sprague Dawley rats was injected subcutaneously twice a week for 8 weeks with PBS. Rats were euthanized 48 hours after the last dose, and organs and plasma were harvested for further analysis.

15 *Plasma chemistry markers*

To evaluate the effect of ISIS oligonucleotides on liver and kidney function, plasma levels of transaminases, bilirubin, albumin, creatinine, and BUN were measured using an automated clinical chemistry analyzer (Hitachi Olympus AU400e, Melville, NY). The results are presented in the table below. ISIS

oligonucleotides that caused changes in the levels of any of the liver or kidney function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 167

Plasma chemistry markers of Sprague Dawley rats

	ALT (IU/L)	AST (IU/L)	Bilirubin (mg/dL)	Albumin (mg/dL)	BUN (mg/dL)	Creatinine (mg/dL)
PBS	30	82	0.09	3.2	19	0.28
ISIS 494159	182	208	0.14	3.4	22	0.35
ISIS 494161	36	86	0.13	3.4	23	0.35
ISIS 494162	102	158	0.17	2.6	28	0.32
ISIS 494283	53	156	0.13	2.9	24	0.32
ISIS 494284	34	113	0.08	2.0	28	0.32
ISIS 494285	110	294	0.10	1.4	110	0.52
ISIS 494286	40	83	0.07	1.6	48	0.44
ISIS 494301	38	132	0.08	3.0	18	0.33
ISIS 494302	47	105	0.09	3.2	19	0.34
ISIS 494311	93	185	0.51	2.7	23	0.30
ISIS 494372	54	119	0.12	3.0	19	0.33
ISIS 510548	116	181	0.11	1.7	65	0.66

5

Kidney function

To evaluate the effect of ISIS oligonucleotides on kidney function, urine levels of total protein and creatinine were measured using an automated clinical chemistry analyzer (Hitachi Olympus AU400e, Melville, NY). Results are presented in the table below, expressed in mg/dL.

10

Table 168

Kidney function markers (mg/dL) in Sprague-Dawley rats

	Creatinine	Total protein
PBS	103	118
ISIS 494159	70	279
ISIS 494161	105	315

ISIS 494162	58	925
ISIS 494283	114	1091
ISIS 494284	97	2519
ISIS 494285	38	2170
ISIS 494286	51	625
ISIS 494301	62	280
ISIS 494302	101	428
ISIS 494311	48	1160
ISIS 494372	46	154
ISIS 510548	55	2119

Organ weights

Liver, spleen and kidney weights were measured at the end of the study, and are presented in the table below. ISIS oligonucleotides that caused any changes in organ weights outside the expected range for antisense oligonucleotides were excluded from further studies.

Table 169

Organ weights of Sprague Dawley rats (g)

	Kidney	liver	Spleen
PBS	3.5	13.1	0.9
ISIS 494159	3.1	11.7	1.6
ISIS 494161	2.8	12.5	2
ISIS 494162	3.1	14.2	1.6
ISIS 494283	3.3	12.9	2.3
ISIS 494284	4.1	15.8	2.7
ISIS 494285	3.8	13.4	0.8
ISIS 494286	4.2	16.7	2.5
ISIS 494301	3.2	12.1	2.3
ISIS 494302	3.4	13.3	2.4
ISIS 494311	3.5	17.4	3.2
ISIS 494372	3.6	12.9	3.2
ISIS 510548	6.4	21.2	1.5

The finding from the rodent tolerability studies showed that in general, taking into consideration all the tolerability markers screened, ISIS 494372 was the best tolerated antisense compound in both the CD1 mouse model and the Sprague Dawley rat model.

5

Example 121: Pharmacokinetics of antisense oligonucleotide in CD1 mice

CD1 mice were treated with ISIS oligonucleotides and the oligonucleotide concentrations in the liver and kidney were evaluated.

Treatment

10 Groups of four CD1 mice each were injected subcutaneously twice per week for 6 weeks with 50 mg/kg of ISIS 494283, ISIS 494284, ISIS 494286, ISIS 494301, ISIS 494302, or ISIS 494372. The mice were sacrificed 2 days following the final dose. Livers were harvested for analysis.

Measurement of oligonucleotide concentration

15 The concentration of the total oligonucleotide concentration was measured. The method used is a modification of previously published methods (Leeds et al., 1996; Geary et al., 1999) which consist of a phenol-chloroform (liquid-liquid) extraction followed by a solid phase extraction. An internal standard (ISIS 355868, a 27-mer 2'-O-methoxyethyl modified phosphorothioate oligonucleotide, GCGTTTGCTCTTCTTCTTTCGTTTTT, designated herein as SEQ ID NO: 131) was added prior to extraction. Tissue sample concentrations were calculated using calibration curves, with a lower limit of
20 quantitation (LLOQ) of approximately 1.14 µg/g. Half-lives were then calculated using WinNonlin software (PHARSIGHT).

The results are presented in the table below, expressed as µg/g liver or kidney tissue. The data indicates that ISIS 494372 was at an acceptable concentration in the liver and kidneys.

Table 170

25 Oligonucleotide concentration (µg/g tissue) of ISIS oligonucleotides in CD1 mice

ISIS No	Liver	Kidney
494283	581	549
494284	511	678
494286	368	445
494301	812	347
494302	617	263
494372	875	516

Example 122: Pharmacokinetics of antisense oligonucleotide in Sprague Dawley rats

Male Sprague Dawley rats were treated with ISIS oligonucleotides and the oligonucleotide concentrations in the liver and kidney were evaluated.

5 *Treatment*

Groups of four rats each were injected subcutaneously twice per week for 3 weeks with 10 mg/kg of ISIS 494283, ISIS 494284, ISIS 494286, ISIS 494301, ISIS 494302, or ISIS 494372. The rats were sacrificed 2 days following the final dose. Livers were harvested for analysis.

Measurement of oligonucleotide concentration

10 The concentration of the total oligonucleotide concentration was measured. The method used is a modification of previously published methods (Leeds et al., 1996; Geary et al., 1999) which consist of a phenol-chloroform (liquid-liquid) extraction followed by a solid phase extraction. An internal standard (ISIS 355868, a 27-mer 2'-O-methoxyethyl modified phosphorothioate oligonucleotide, GCGTTTGCTCTTCTTCTTTCGTTTTTT, designated herein as SEQ ID NO: 131) was added prior to
15 extraction. Tissue sample concentrations were calculated using calibration curves, with a lower limit of quantitation (LLOQ) of approximately 1.14 µg/g. Half-lives were then calculated using WinNonlin software (PHARSIGHT).

The results are presented in the table below, expressed as µg/g liver or kidney tissue. The data indicates that ISIS 494372 was at an acceptable concentration in the liver and kidneys.

20

Table 171

Oligonucleotide concentration (µg/g tissue) of ISIS oligonucleotides in Sprague Dawley rats

ISIS No	Liver	Kidney
494283	220	434
494284	178	573
494286	234	448
494301	279	540
494302	205	387
494372	288	663

Example 123: Effect of ISIS antisense oligonucleotides targeting human apo(a) in cynomolgus monkeys

25

Cynomolgus monkeys were treated with ISIS antisense oligonucleotides selected from studies described above. At the time this study was undertaken, the cynomolgus monkey genomic sequence was not

available in the National Center for Biotechnology Information (NCBI) database; therefore, cross-reactivity with the cynomolgus monkey gene sequence could not be confirmed. Instead, the sequences of the ISIS antisense oligonucleotides used in the cynomolgus monkeys was compared to a rhesus monkey sequence for homology. It is expected that ISIS oligonucleotides with homology to the rhesus monkey sequence are fully cross-reactive with the cynomolgus monkey sequence as well.

The human antisense oligonucleotides tested are also cross-reactive with the rhesus mRNA sequence (XM_001098061.1; designated herein as SEQ ID NO: 132). The greater the complementarity between the human oligonucleotide and the rhesus monkey sequence, the more likely the human oligonucleotide can cross-react with the rhesus monkey sequence. The start and stop sites of each oligonucleotide to SEQ ID NO: 132 is presented in the table below. Each antisense oligonucleotide targets more than one region in SEQ ID NO:132 and has multiple start sites. "Start site" indicates the 5'-most nucleotide to which the gapmer is targeted in the rhesus monkey sequence. 'Mismatches' indicates the number of nucleotides mismatched between the human oligonucleotide sequence and the rhesus sequence.

Antisense oligonucleotide tolerability, as well as their pharmacokinetic profile in the liver and kidney, was evaluated.

Table 172

Antisense oligonucleotides complementary to SEQ ID NO: 132

ISIS No	Start Site	Mismatches
494283	278	2
	620	2
	923	2
	1265	2
	1607	1
	1949	1
	2267	1
	2609	1
	2951	1
	3293	1
494284	279	1
	621	1
	924	1
	1266	1
	1608	1
	1950	1
	2268	1
	2610	1
	2952	1
	3294	1

494286	281	1
	623	1
	926	1
	1268	1
	1610	2
	1952	2
	2270	2
	2612	2
	2954	2
	3296	2
494301	322	2
	664	2
	967	2
	1309	1
	1651	2
494302	323	2
	968	2
	1310	1
	1652	2
494372	1186	2
	1870	1
	2188	1

Treatment

Prior to the study, the monkeys were kept in quarantine for at least a 30-day period, during which the animals were observed daily for general health. The monkeys were 2-4 years old and weighed between 2 and 4 kg. Seven groups of four randomly assigned male cynomolgus monkeys each were injected subcutaneously with ISIS oligonucleotide or PBS using a stainless steel dosing needle and syringe of appropriate size into the one of four sites on the back of the monkeys. The injections were given in clock-wise rotation; one site per dosing. The monkeys were dosed four times a week for the first week (days 1, 3, 5, and 7) as loading doses, and subsequently once a week for weeks 2-12, with 40 mg/kg of ISIS 494283, ISIS 494284, ISIS 494286, ISIS 494301, ISIS 494302, or ISIS 494372. A control group of 8 cynomolgus monkeys was injected with PBS subcutaneously thrice four times a week for the first week (days 1, 3, 5, and 7), and subsequently once a week for weeks 2-12.

During the study period, the monkeys were observed at least once daily for signs of illness or distress. Any animal experiencing more than momentary or slight pain or distress due to the treatment, injury or illness was treated by the veterinary staff with approved analgesics or agents to relieve the pain after consultation with the Study Director. Any animal in poor health or in a possible moribund condition was identified for further monitoring and possible euthanasia. For instance, one animal in the treatment group of

ISIS 494302 was found moribund on day 56 and was euthanized. Scheduled euthanasia of the animals was conducted on days 86 and 87 by exsanguination under deep anesthesia. The protocols described in the Example were approved by the Institutional Animal Care and Use Committee (IACUC).

Target Reduction

5 *RNA analysis*

On day 86, RNA was extracted from liver tissue for real-time PCR analysis of apo(a) using human primer probe set ABI Hs00916691_m1 (Applied Biosystems, Carlsbad CA). Results are presented as percent inhibition of apo(a) mRNA, relative to PBS control. As shown in the table below, treatment with ISIS antisense oligonucleotides resulted in significant reduction of apo(a) mRNA in comparison to the PBS control.

The mRNA levels of plasminogen, another kringle-containing protein, were also measured. Treatment with ISIS 494372 did not alter the mRNA levels of plasminogen.

Table 173

Percent Inhibition of apo(a) mRNA in the cynomolgus monkey liver relative to the PBS control

ISIS No	% inhibition
494283	91
494284	99
494286	96
494301	88
494302	89
494372	93

15 *Protein analysis*

On different days, one mL of blood was collected from the cephalic, saphenous, or femoral vein of all study monkeys. The blood samples were put into tubes containing K2-EDTA for plasma separation. The tubes were centrifuged at 3,000 rpm for 10 min at room temperature to obtain plasma. Apo(a) protein levels were analyzed by an Apo(a) ELISA kit (Mercodia 10-1106-01, Uppsala, Sweden). Results are presented as percentage change of levels from the baseline. As shown in the table below, treatment with several ISIS antisense oligonucleotides resulted in significant reduction of apo(a) protein levels in comparison to the PBS control. Specifically, treatment with ISIS 494372 reduced cynomolgus plasma protein levels of apo(a).

The protein levels of apoB were also measured in the study groups. Antisense inhibition of apo(a) had no effect on apoB levels.

Table 174

Apo(a) plasma protein levels (% inhibition over baseline values) in the cynomolgus monkey

	Day 16	Day 30	Day 44	Day 56	Day 72	Day 86
PBS	0	0	10	0	0	0
ISIS 494283	78	79	81	66	66	70
ISIS 494284	92	95	95	93	93	94
ISIS 494286	92	95	96	94	94	94
ISIS 494301	41	45	52	20	17	29
ISIS 494302	17	0	2	0	0	20
ISIS 494372	67	80	83	79	78	81

Tolerability studies

Body and organ weight measurements

- 5 To evaluate the effect of ISIS oligonucleotides on the overall health of the animals, body and organ weights were measured at day 86. Body weights were measured and are presented in the table below. Organ weights were measured and the data is presented in the table below. The results indicate that treatment with ISIS 494372 was well tolerated in terms of the body and organ weights of the monkeys.

Table 175

10

Body weights (g) in the cynomolgus monkey

	Day 14	Day 35	Day 49	Day 56	Day 70	Day 84
PBS	2637	2691	2748	2733	2739	2779
ISIS 494283	2591	2670	2698	2656	2704	2701
ISIS 494284	2559	2661	2676	2675	2662	2646
ISIS 494286	2693	2770	2838	2800	2796	2816
ISIS 494301	2587	2604	2627	2591	2596	2604
ISIS 494302	2759	2760	2839	2825	3113	3122
ISIS 494372	2719	2877	2985	2997	3037	3036

Table 176

Organ weights (% body weight) in the cynomolgus monkey

	Spleen	Kidneys	Liver	Heart	Lungs
PBS	0.14	0.38	2.2	0.33	0.51
ISIS 494283	0.24	0.95	2.8	0.33	0.49
ISIS 494284	0.19	0.60	2.6	0.36	0.55
ISIS 494286	0.22	0.63	2.7	0.38	0.55
ISIS 494301	0.38	0.81	3.0	0.36	0.61
ISIS 494302	0.17	0.95	2.5	0.39	0.57
ISIS 494372	0.18	1.16	2.6	0.36	0.56

15 *Liver function*

To evaluate the effect of ISIS oligonucleotides on hepatic function, monkeys were fasted overnight prior to blood collection. Approximately 1.5 mL of blood was collected from each animal and put into tubes without anticoagulant for serum separation. The tubes were kept at room temperature for a minimum of 90 min and then centrifuged at 3,000 rpm for 10 min at room temperature to obtain serum. Levels of various liver function markers were measured using a Toshiba 200FR NEO chemistry analyzer (Toshiba Co., Japan). Plasma levels of ALT and AST were measured and the results are presented in the table below, expressed in IU/L. Bilirubin, a liver function marker, was similarly measured and is presented in the table below, expressed in mg/dL. The results indicate that treatment with ISIS 494372 was well tolerated in terms of the liver function in monkeys.

Table 177

Liver function markers in cynomolgus monkey plasma

	ALT (IU/L)	AST (IU/L)	Bilirubin (mg/dL)
PBS	33	43	0.20
ISIS 494283	75	73	0.12
ISIS 494284	115	79	0.17
ISIS 494286	67	73	0.13
ISIS 494301	129	90	0.15
ISIS 494302	141	75	0.15
ISIS 494372	46	75	0.17

C-reactive protein level analysis

To evaluate any inflammatory effect of ISIS oligonucleotides in cynomolgus monkeys, blood samples were taken for analysis. The monkeys were fasted overnight prior to blood collection. Approximately 1.5 mL of blood was collected from each animal and put into tubes without anticoagulant for serum separation. The tubes were kept at room temperature for a minimum of 90 min and then centrifuged at 3,000 rpm for 10 min at room temperature to obtain serum. C-reactive protein (CRP), which is synthesized in the liver and which serves as a marker of inflammation, was measured using a Toshiba 200FR NEO chemistry analyzer (Toshiba Co., Japan). The results indicate that treatment with ISIS 494372 did not cause any inflammation in monkeys.

Table 178

C-reactive protein levels (mg/L) in cynomolgus monkey plasma

	CRP
--	-----

PBS	1.4
ISIS 494283	14.7
ISIS 494284	7.7
ISIS 494286	4.4
ISIS 494301	3.5
ISIS 494302	2.4
ISIS 494372	10.2

Complement C3 analysis

To evaluate any effect of ISIS oligonucleotides on the complement pathway in cynomolgus monkeys, blood samples were taken for analysis on day 84 (pre-dose) and day 85 (24 hours post-dose). Approximately 0.5 mL of blood was collected from each animal and put into tubes without anticoagulant for serum separation. The tubes were kept at room temperature for a minimum of 90 min and then centrifuged at 3,000 rpm for 10 min at room temperature to obtain serum. C3 was measured using a Toshiba 200FR NEO chemistry analyzer (Toshiba Co., Japan). The results indicate that treatment with ISIS 494372 did not cause any effect on the complement pathway in monkeys.

Table 179

Complement C3 levels (mg/dL) in cynomolgus monkey plasma

	Pre-dose	Post-dose
PBS	140	139
ISIS 494283	127	101
ISIS 494284	105	75
ISIS 494286	84	38
ISIS 494301	118	76
ISIS 494302	98	58
ISIS 494372	123	109

Hematology

To evaluate any effect of ISIS oligonucleotides in cynomolgus monkeys on hematologic parameters, blood samples of approximately 0.5 mL of blood was collected on day 87 from each of the available study animals in tubes containing K₂-EDTA. Samples were analyzed for red blood cell (RBC) count, white blood cells (WBC) count, as well as for platelet count, using an ADVIA120 hematology analyzer (Bayer, USA). The data is presented in the table below.

The data indicate that treatment with ISIS 494372 was well tolerated in terms of the hematologic parameters of the monkeys.

Table 180

Blood cell counts in cynomolgus monkeys

	WBC (x 10 ³ /μL)	RBC (x 10 ⁶ /μL)	Platelet (x 10 ³ /μL)
PBS	15	6.3	329
ISIS 494283	16	5.3	456
ISIS 494284	13	6.3	330
ISIS 494286	14	5.5	304
ISIS 494301	15	6.0	392
ISIS 494302	12	6.3	305
ISIS 494372	11	6.1	447

5 Example 124: Characterization of the pharmacological activity of ISIS 494372 in cynomolgus monkeys

The pharmacological activity of ISIS 494372 was characterized by measuring liver apo(a) mRNA and plasma apo(a) levels in monkeys administered the compound over 13 weeks and allowed to recover for another 13 weeks.

10 Treatment

Five groups of 14 randomly assigned male and female cynomolgus monkeys each were injected subcutaneously with ISIS oligonucleotide or PBS using a stainless steel dosing needle and syringe of appropriate size into the one of four sites on the back (scapular region) of the monkeys. The monkeys were dosed four times a week for the first week (days 1, 3, 5, and 7) as loading doses, and subsequently once a week for weeks 2-13 as maintenance doses, as shown in the table below. The loading dose during the first week is expressed as mg/kg/dose, while the maintenance doses on weeks 2-13 are expressed as mg/kg/week.

Table 181

Dosing groups in cynomolgus monkeys

Group	Test Article	Dose	Number of animals for necropsy		
			Interim	Terminal	Recovery
1	PBS	-	4	6	4
2	ISIS 494372	4	-	6	-
3		8	-	6	-
4		12	4	6	4
5		40	4	6	4

Liver samples from animals were taken at the interim, terminal and recovery phases of the study for the analyses of apo(a) mRNA. In addition, plasma samples were collected on different days to measure apo(a) protein levels. This non-clinical study was conducted in accordance with the United States Food and Drug Administration (FDA) Good Laboratory Practice (GLP) Regulations, 21 CFR Part 58.

5 *RNA analysis*

Liver samples were collected from monkeys on days 30, 93, and 182, and frozen. Briefly, a piece (0.2 g) of frozen liver was homogenized in 2 mL of RLT solution (Qiagen). The resulting lysate was applied to Qiagen RNeasy mini columns. After purification and quantification, the tissues were subjected to RT-PCR analysis. The Perkin-Elmer ABI Prism 7700 Sequence Detection System, which uses real-time fluorescent RT-PCR detection, was used to quantify apo(a) mRNA. The assay is based on a target-specific probe labeled with fluorescent reporter and quencher dyes at opposite ends. The probe was hydrolyzed through the 5'-exonuclease activity of Taq DNA polymerase, leading to an increasing fluorescence emission of the reporter dye that can be detected during the reaction. A probe set (ABI Rhesus LPA probe set ID Rh02789275_m1, Applied Biosystems, Carlsbad CA) targeting position 1512 of the rhesus monkey apo(a) mRNA transcript
 10
 15
 GENBANK Accession No XM_001098061.2 (SEQ ID NO: 132) sequence was used to measure cynomolgus monkey liver apo(a) mRNA expression levels. Apo(a) expression was normalized using RIBOGREEN®. Results are presented as percent inhibition of apo(a) mRNA, relative to PBS control.

As shown in the table below, treatment with ISIS 494372 resulted in a dose-dependent reduction of apo(a) mRNA in comparison to the PBS control. At day 30, hepatic apo(a) mRNA expression was reduced in
 20
 a dose-dependent manner by 74% and 99% in the 12 mg/kg/week and 40 mg/kg/week dosing cohorts, respectively. These reductions are statistically significant by one-way ANOVA (Dunnett's multiple comparison test, $P < 0.05$).

Apo(a) mRNA levels were also measured during the recovery phase. Liver expression levels at day 88 after the last dose were still reduced 49% and 69% in the 12 mg/kg/week and 40 mg/kg/week dosing
 25
 cohorts, respectively.

Table 182

Percent inhibition levels of liver apo(a) mRNA in the dosing phase in cynomolgus monkeys treated with ISIS 494372

Day	Dose (mg/kg/wk)	% inhibition
30	12	73
	40	99

93	4	44
	8	43
	12	53
	40	93

Protein analysis

Approximately 20 μ l of plasma was analyzed using a commercially available apo(a) ELISA kit (Mercodia 10-1106-01, Uppsala, Sweden). The assay protocol was performed as described by the manufacturer. The results are presented in the tables below as percentage change from Day 1 pre-dose apo(a) plasma protein concentrations. Statistically significant differences from Day 1 baseline plasma apo(a) using the Dunnett's multicomparison test are marked with an asterisk.

Maximal reduction in plasma apo(a) protein was observed in all dosing cohorts by Day 93. In the recovery phase, apo(a) plasma protein levels in the 40 mg/kg/week dosing cohort were at 22% and 93% of the baseline after 4 and 13 weeks (Days 121 and 182) of recovery, respectively. The rate of recovery in the 12 mg/kg/week cohort was similar to that seen in the 40 mg/kg/week cohort.

Table 183

Apo(a) plasma protein levels as a percent of Day 1 levels in the dosing phase in cynomolgus monkeys treated with ISIS 494372

Day	Dose (mg/kg/wk)	%
30	4	93
	8	70
	12	49
	40	15*
93	4	73
	8	56
	12	32*
	40	11*

Table 184

Apo(a) plasma protein levels as a percent of Day 1 levels in the recovery phase in cynomolgus monkeys treated with ISIS 494372

Day	Dose (mg/kg/wk)	%
121	12	38*
	40	22*
182	12	84
	40	93

5 **Example 125: Measurement of viscosity of ISIS antisense oligonucleotides targeting human Apo(a)**

The viscosity of select antisense oligonucleotides from the studies described above was measured with the aim of screening out antisense oligonucleotides which have a viscosity more than 40 centipoise (cP). Oligonucleotides having a viscosity greater than 40 cP would have less than optimal viscosity.

ISIS oligonucleotides (32-35 mg) were weighed into a glass vial, 120 μ L of water was added and the antisense oligonucleotide was dissolved into solution by heating the vial at 50°C. Part (75 μ L) of the pre-heated sample was pipetted to a micro-viscometer (Cambridge). The temperature of the micro-viscometer was set to 25°C and the viscosity of the sample was measured. Another part (20 μ L) of the pre-heated sample was pipetted into 10 mL of water for UV reading at 260 nm at 85°C (Cary UV instrument). The results are presented in the table below and indicate that most of the antisense oligonucleotides solutions are optimal in their viscosity under the criterion stated above. Those that were not optimal are marked as 'viscous'. Specifically, ISIS 494372 was optimal in its viscosity under the criterion stated above.

Table 185

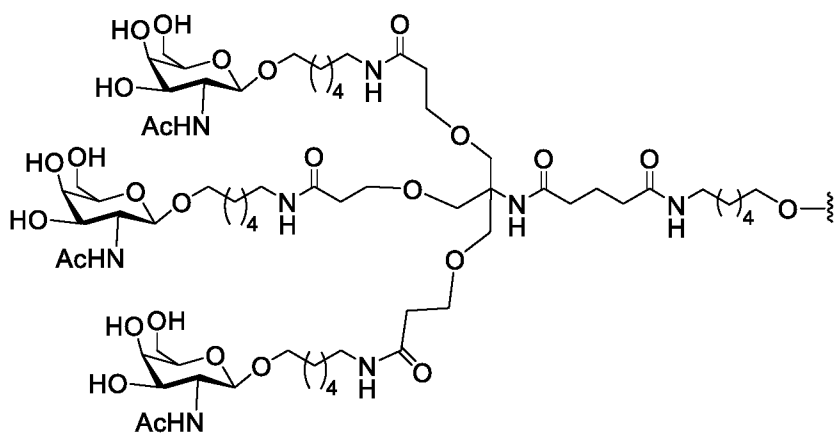
Viscosity and concentration of ISIS antisense oligonucleotides targeting human Apo(a)

ISIS No	Motif	Viscosity (cP)	Concentration (mg/mL)
494158	5-10-5 MOE	9.0	350
494159	5-10-5 MOE	11.7	325
494161	5-10-5 MOE	12.0	350
494162	5-10-5 MOE	25.8	350
494163	5-10-5 MOE	Viscous	275
494243	5-10-5 MOE	28.4	325
494244	5-10-5 MOE	19.2	300

494283	3-10-4 MOE	13.4	300
494284	5-10-5 MOE	13.4	350
494285	5-10-5 MOE	23.1	350
494286	5-10-5 MOE	16.5	275
494301	5-10-5 MOE	17.1	325
494302	5-10-5 MOE	24.3	350
494304	5-10-5 MOE	49.3	275
494311	5-10-5 MOE	10.8	325
494337	5-10-5 MOE	29.5	325
494372	5-10-5 MOE	12.5	350
494466	5-10-5 MOE	Viscous	275
494470	5-10-5 MOE	16.7	350
494472	5-10-5 MOE	23.6	350
498408	5-10-5 MOE	31.5	300
510548	5-10-5 MOE	9.0	350
512947	3-10-4 MOE	6.8	350
512958	5-10-5 MOE	26.0	350

CLAIMS:

1. A compound comprising a modified oligonucleotide and a conjugate group, wherein the modified oligonucleotide consists of 20 contiguous nucleobases complementary to an equal length portion of nucleobases 3901 to 3920 of SEQ ID NO: 1, wherein the nucleobase sequence of the modified oligonucleotide is at least 80% complementary to SEQ ID NO: 1; and wherein the conjugate group comprises:



2. The compound of claim 1, wherein:
- (i) the nucleobase sequence of the modified oligonucleotide is at least 85%, at least 90%, at least 95%, or 100% complementary to SEQ ID NO: 1,
 - (ii) the modified oligonucleotide comprises at least one modified sugar, optionally wherein: (a) at least one modified sugar is a bicyclic sugar, or (b) at least one modified sugar comprises a 2'-O-methoxyethyl, a constrained ethyl, a 3'-fluoro-HNA or a 4'-((CH₂)_n-O-2') bridge, wherein n is 1 or 2,
 - (iii) at least one nucleoside comprises a modified nucleobase, optionally wherein the modified nucleobase is a 5-methylcytosine, and/or
 - (iv) each internucleoside linkage of the modified oligonucleotide is selected from a phosphodiester internucleoside linkage and a phosphorothioate internucleoside linkage, optionally wherein the

modified oligonucleotide comprises: (a) at least 5 phosphodiester internucleoside linkages or (b) at least 2 phosphorothioate internucleoside linkages.

3. The compound of claim 1 or claim 2, wherein the modified oligonucleotide is single-stranded.

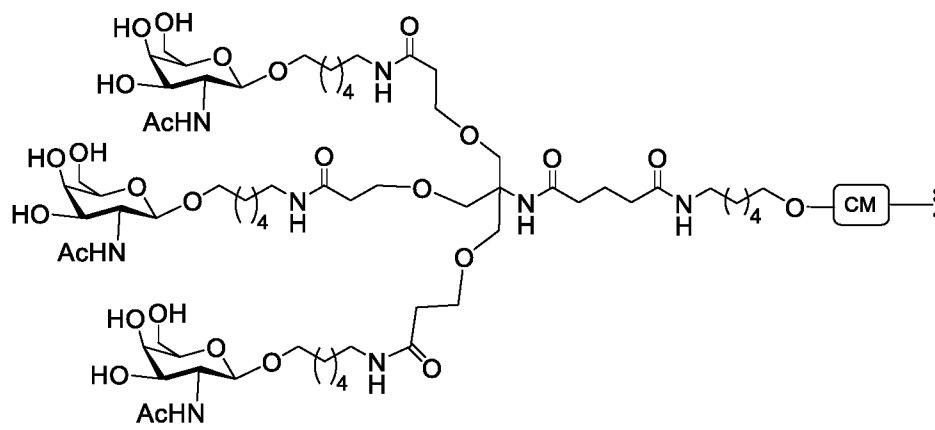
5

4. The compound of any one of claims 1-3, wherein the conjugate group is linked to the modified oligonucleotide at the 5' end of the modified oligonucleotide.

5. The compound of any one of claims 1-3, wherein the conjugate group is linked to the modified

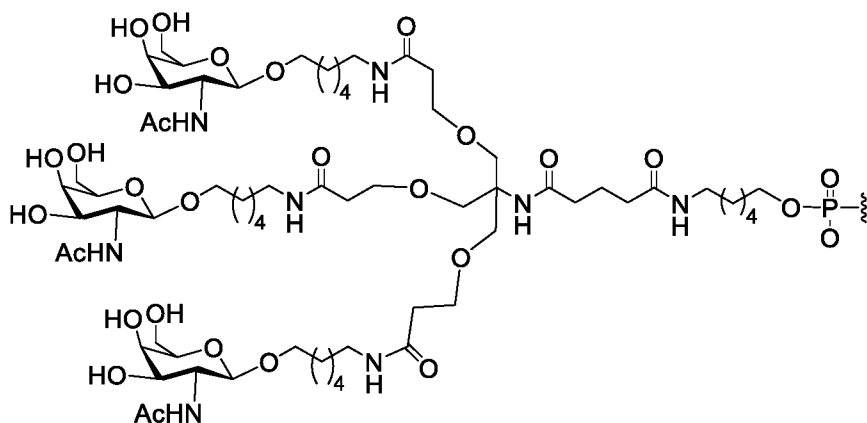
10 oligonucleotide at the 3' end of the modified oligonucleotide.

6. The compound of any one of claims 1-5, wherein the conjugate group comprises:



15 wherein the cleavable moiety (CM) is a bond or group that is capable of being split under physiological conditions.

7. The compound of any one of claims 1-6, wherein the conjugate group comprises:



8. The compound of claim 1, wherein the modified oligonucleotide comprises:

- 5 a gap segment consisting of linked deoxynucleosides;
- a 5' wing segment consisting of linked nucleosides;
- a 3' wing segment consisting of linked nucleosides;
- wherein the gap segment is positioned between the 5' wing segment and the 3' wing segment
- and wherein each nucleoside of each wing segment comprises a modified sugar.

10

9. The compound of claim 1, wherein the modified oligonucleotide comprises:

- a gap segment consisting of ten linked deoxynucleosides;
- a 5' wing segment consisting of five linked nucleosides;
- a 3' wing segment consisting of five linked nucleosides;
- 15 wherein the gap segment is positioned between the 5' wing segment and the 3' wing segment, wherein each nucleoside of each wing segment comprises a 2'-O-methoxyethyl sugar,
- and wherein each cytosine residue is a 5-methylcytosine.

10. The compound of claim 8 or claim 9, wherein each internucleoside linkage in the gap segment of
 20 the modified oligonucleotide is a phosphorothioate linkage.

11. The compound of claim 10, wherein the modified oligonucleotide further comprises at least one phosphorothioate internucleoside linkage in each wing segment.

12. The compound of claim 1, wherein the modified oligonucleotide consists of 20 linked nucleosides with the nucleobase sequence of SEQ ID NO: 58, and wherein the modified oligonucleotide comprises:

a gap segment consisting of ten linked deoxynucleosides;

a 5' wing segment consisting of five linked nucleosides;

a 3' wing segment consisting of five linked nucleosides;

wherein the gap segment is positioned between the 5' wing segment and the 3' wing segment, wherein each nucleoside of each wing segment comprises a 2'-O-methoxyethyl sugar, wherein at least one internucleoside linkage is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

13. The compound of claim 1, wherein the modified oligonucleotide comprises at least 8, at least 9, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or 20 contiguous nucleobases of the nucleobase sequence of SEQ ID NO: 58, and wherein the modified oligonucleotide comprises:

a gap segment consisting of ten linked deoxynucleosides;

a 5' wing segment consisting of five linked nucleosides;

a 3' wing segment consisting of five linked nucleosides;

wherein the gap segment is positioned between the 5' wing segment and the 3' wing segment, wherein each nucleoside of each wing segment comprises a 2'-O-methoxyethyl sugar, wherein each internucleoside linkage in the gap segment is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

14. The compound of claim 13, wherein:

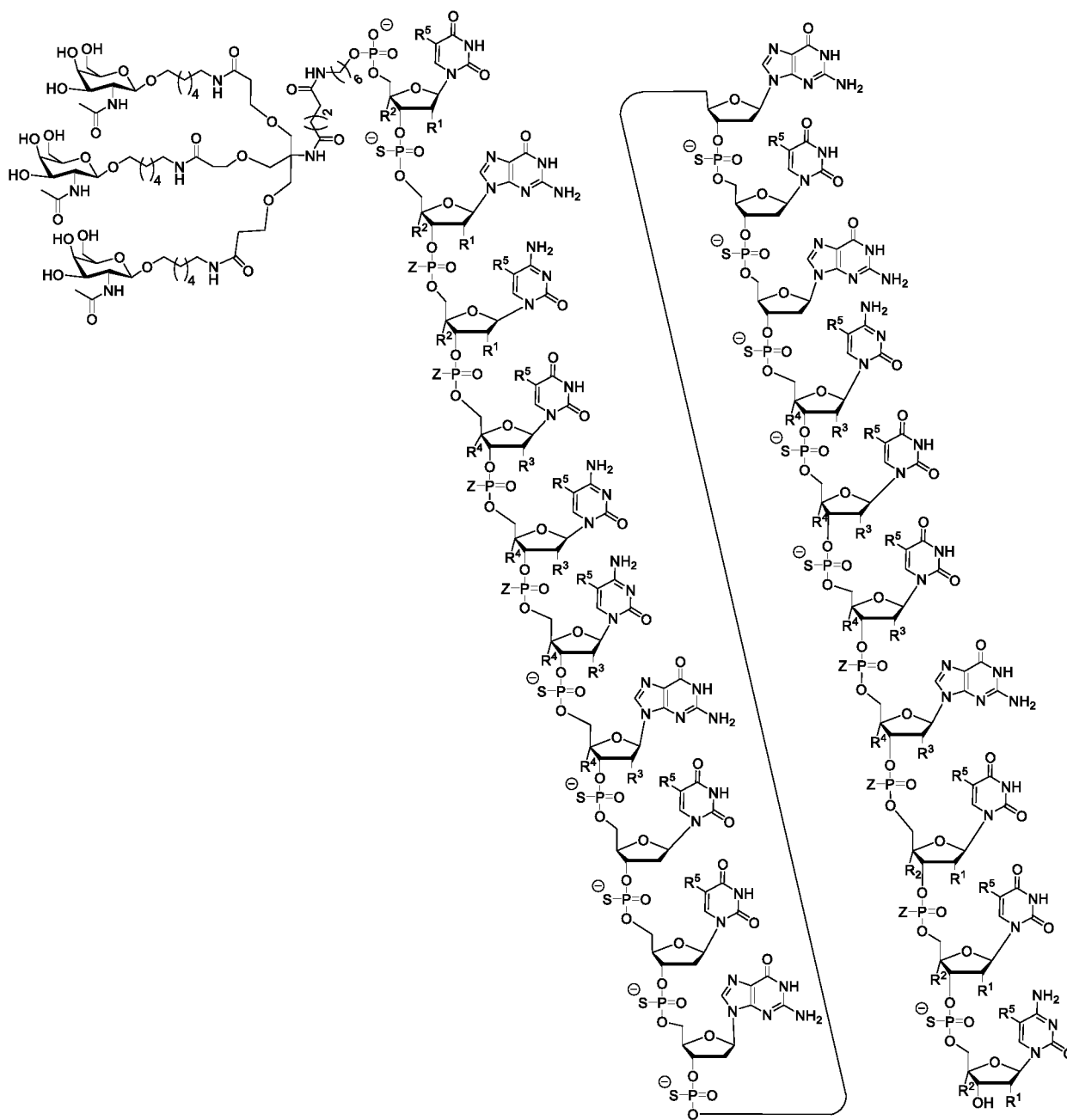
(i) the modified oligonucleotide further comprises at least one phosphorothioate internucleoside linkage in each wing segment,

(ii) the internucleoside linkages are phosphorothioate linkages between nucleosides 1-2, nucleosides 6-16 and nucleosides 18-20 of the modified oligonucleotide, wherein nucleosides 1-20 are positioned 5' to 3', or

(iii) the 2nd, 3rd, 4th, and 5th internucleoside linkage from the 5'-end is a phosphodiester internucleoside linkage, wherein the 3rd and 4th internucleoside linkage from the 3'-end is a phosphodiester internucleoside linkage, and wherein each remaining internucleoside linkage is a phosphorothioate internucleoside linkage.

15. The compound of claim 1, having the formula:

10

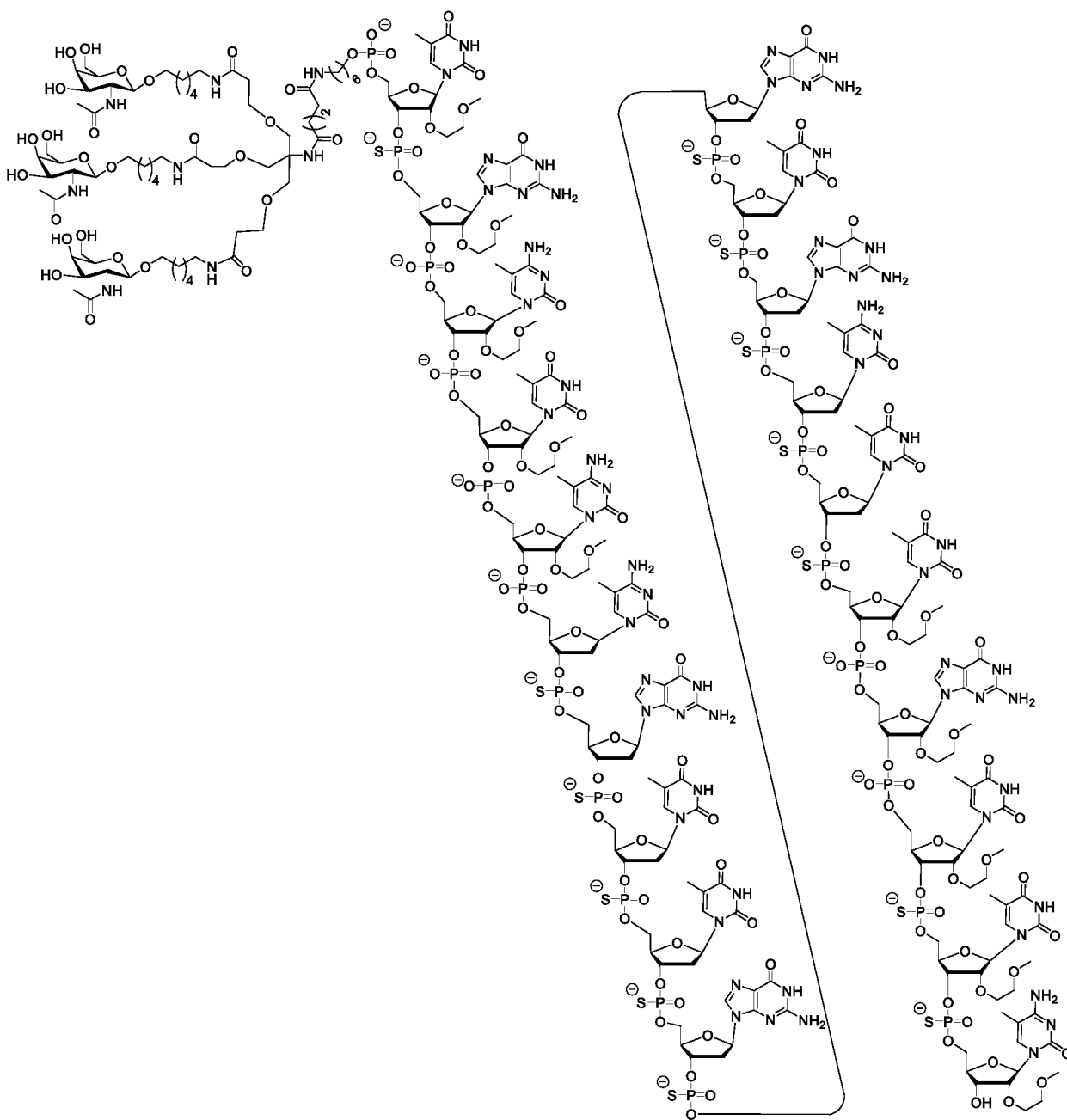


wherein either R^1 is $-\text{OCH}_2\text{CH}_2\text{OCH}_3$ (MOE) and R^2 is H; or R^1 and R^2 together form a bridge, wherein R^1 is $-\text{O}-$ and R^2 is $-\text{CH}_2-$, $-\text{CH}(\text{CH}_3)-$, or $-\text{CH}_2\text{CH}_2-$, and R^1 and R^2 are directly connected such that the resulting bridge is selected from: $-\text{O}-\text{CH}_2-$, $-\text{O}-\text{CH}(\text{CH}_3)-$, and $-\text{O}-\text{CH}_2\text{CH}_2-$;

and for each pair of R^3 and R^4 on the same ring, independently for each ring: either R^3 is selected from H and $-OCH_2CH_2OCH_3$ and R^4 is H; or R^3 and R^4 together form a bridge, wherein R^3 is $-O-$, and R^4 is $-CH_2-$, $-CH(CH_3)-$, or $-CH_2CH_2-$ and R^3 and R^4 are directly connected such that the resulting bridge is selected from: $-O-CH_2-$, $-O-CH(CH_3)-$, and $-O-CH_2CH_2-$;

- 5 and R^5 is selected from H and $-CH_3$;
and Z is selected from S^- and O^- .

16. The compound of claim 1, having the formula:



17. The compound of any preceding claim, wherein the compound is in a salt form, optionally
 5 wherein the compound is in the form of a sodium salt and/or potassium salt.

18. A pharmaceutical composition comprising a compound of any one of claims 1-17, and a pharmaceutically acceptable diluent or carrier.

5 19. A method comprising administering to an animal a compound according to any one of claims 1-17 or a composition according to claim 18, wherein administering the compound or composition treats, prevents or slows progression of a disease related to elevated apo(a) and/or elevated Lp(a).

20. Use of the compound according to any one of claims 1-17, or use of the composition according
10 to claim 18, in the manufacture of a medicament for treating, preventing, or slowing progression of a disease related to elevated apo(a) and/or elevated Lp(a).

21. The method of claim 19 or the use of claim 20, wherein the disease is an inflammatory, cardiovascular or metabolic disease, disorder or condition.

15

22. The method of claim 19 or the use of claim 20, wherein the disease is aortic stenosis or angina.

BI OL0250W0SEQ_ST25. txt
SEQUENCE LISTING

<110> Isis Pharmaceutical s, Inc.

<120> COMPOSITIONS AND METHODS FOR MODULATING APOLIPOPROTEIN (a) EXPRESSION

<130> BI OL0250W0

<150> 61/818,442

<151> 2013-05-01

<150> 61/823,826

<151> 2013-05-15

<150> 61/843,887

<151> 2013-07-08

<150> 61/871,673

<151> 2013-08-29

<150> 61/880,790

<151> 2013-09-20

<150> 61/976,991

<151> 2014-04-08

<150> 61/986,867

<151> 2014-04-30

<160> 167

<170> PatentIn version 3.5

<210> 1

<211> 6489

<212> DNA

<213> Homo sapiens

<400> 1

aggtagcttt ggggctggct ttctcaagga agcccagctc cctgtgattg agaatgaagt 60

gtgcaatcgc tatgactggg attgggacac actttctggg cactgctggc cagtcccaaa 120

atggaacata aggaagtggg tcttctactt cttttatttc tgaaatcagc agcacctgag 180

caaagccatg tgggtccagga ttgctacat ggtgatggac agagttatcg aggcacgtac 240

tccaccactg tcacaggaag gacctgccaa gcttgggtcat ctatgacacc acatcaacat 300

aataggacca cagaaaacta cccaaatgct ggcttgatca tgaactactg caggaatcca 360

gatgctgtgg cagctcctta ttgttatacg agggatcccg gtgtcaggtg ggagtactgc 420

aacctgacgc aatgctcaga cgcagaaggg actgccgtcg cgcctccgac tgttaccccg 480

gttccaagcc tagaggctcc ttccgaacaa gcaccgactg agcaaaggcc tggggtgcag 540

gagtgtacc atggtaatgg acagagttat cgaggcacat actccaccac tgtcacagga 600

agaacctgcc aagcttgggtc atctatgaca ccacactcgc atagtctggac ccagaatac 660

tacccaaatg ctggcttgat catgaactac tgcaggaatc cagatgtgtg ggcagctcct 720

tattgttata cgagggatcc cgggtgtcagg tgggagtact gcaacctgac gcaatgtctca 780

gacgcagaag ggactgccgt cgcgcctccg actgttaccg cggttccaag cctagaggct 840

BI OL0250W0SEQ_ST25. txt

ccttccgaac	aagcaccgac	tgagcaaagg	cctgggggtgc	aggagtgcta	ccatggtaat	900
ggacagagtt	atcgaggcac	atactccacc	actgtcacag	gaagaacctg	ccaagcttgg	960
tcatctatga	caccacactc	gcatagtcgg	accccagaat	actacccaaa	tgctggcttg	1020
atcatgaact	actgcaggaa	tccagatgct	gtggcagctc	cttattgtta	tacgagggat	1080
cccgggtgtca	gggtgggagta	ctgcaacctg	acgcaatgct	cagacgcaga	agggactgcc	1140
gtcgcgcctc	cgactgttac	cccggttcca	agcctagagg	ctccttccga	acaagcaccg	1200
actgagcaga	ggcctgggggt	gcaggagtgc	taccacggta	atggacagag	ttatcgaggc	1260
acatactcca	ccactgtcac	tggaagaacc	tgccaagctt	ggtcacttat	gacaccacac	1320
tcgcatagtc	ggaccccaga	atactacca	aatgctggct	tgatcatgaa	ctactgcagg	1380
aatccagatg	ctgtggcagc	tccttattgt	tatacgaggg	atcccgggtgt	cagggtgggag	1440
tactgcaacc	tgacgcaatg	ctcagacgca	gaagggactg	ccgtcgcgcc	tccgactgtt	1500
accccggttc	caagcctaga	ggctccttcc	gaacaagcac	cgactgagca	aaggcctggg	1560
gtgcaggagt	gctaccatgg	taatggacag	agttatcgag	gcacatactc	caccactgtc	1620
acaggaagaa	cctgccaagc	ttggtcactt	atgacaccac	actcgcatag	tcggacccca	1680
gaatactacc	caaatgctgg	cttgatcatg	aactactgca	ggaatccaga	tgctgtggca	1740
gctccttatt	gttatacgag	ggatcccgggt	gtcagggtggg	agtactgcaa	cctgacgcaa	1800
tgctcagacg	cagaagggac	tgccgtcgcg	cctccgactg	ttaccccggt	tccaagccta	1860
gaggctcctt	ccgaacaagc	accgactgag	caaaggcctg	gggtgcagga	gtgctaccat	1920
ggtaatggac	agagttatcg	aggcacatac	tccaccactg	tcacaggaag	aacctgccaa	1980
gcttgggtcat	ctatgacacc	acactcgcac	agtcggaccc	cagaatacta	cccaaagtct	2040
ggcttgatca	tgaactactg	caggaatcca	gatgctgtgg	cagctcctta	ttgtttatacg	2100
agggatccccg	gtgtcagggtg	ggagtactgc	aacctgacgc	aatgctcaga	cgcagaaggg	2160
actgccgtcg	cgctccgac	tgttacccccg	gttccaagcc	tagaggctcc	ttccgaacaa	2220
gcaccgactg	agcaaaggcc	tgggggtgcag	gagtgtctacc	atggtaatgg	acagagttat	2280
cgaggcacat	actccaccac	tgtcacagga	agaacctgcc	aagcttggtc	atctatgaca	2340
ccacactcgc	atagtcggac	cccagaatac	tacccaaatg	ctggcttgat	catgaactac	2400
tgaggaatc	cagatgctgt	ggcagctcct	tattgtttata	cgagggatcc	cgggtgtcagg	2460
tgggagtact	gcaacctgac	gcaatgctca	gacgcagaag	ggactgccgt	cgcgccctccg	2520
actgttacc	cggttccaag	cctagagggt	ccttccgaac	aagcaccgac	tgagcagagg	2580
cctgggggtgc	aggagtgcta	ccacggtaat	ggacagagtt	atcgaggcac	atactccacc	2640
actgtcactg	gaagaacctg	ccaagcttgg	tcatctatga	caccacactc	gcatagtcgg	2700
accccagaat	actacccaaa	tgctggcttg	atcatgaact	actgcaggaa	tccagatcct	2760
gtggcagccc	cttattgtta	tacgagggat	cccagtgtca	gggtgggagta	ctgcaacctg	2820
acacaatgct	cagacgcaga	agggactgcc	gtcgcgcctc	caactattac	cccgattcca	2880

BI OL0250W0SEQ_ST25. txt

agcctagagg	ctccttctga	acaagcacca	actgagcaaa	ggcctggggt	gcaggagtgc	2940
taccacggaa	atggacagag	ttatcaaggc	acatacttca	ttactgtcac	aggaagaacc	3000
tgccaagctt	ggatcatctat	gacaccacac	tcgcatagtc	ggaccccagc	atactaccca	3060
aatgctggct	tgatcaagaa	ctactgccga	aatccagatc	ctgtggcagc	cccttggtgt	3120
tatacaacag	atcccagtgt	caggtgggag	tactgcaacc	tgacacgatg	ctcagatgca	3180
gaatggactg	ccttcgtccc	tccgaatgtt	attctggctc	caagcctaga	ggcttttttt	3240
gaacaagcac	tgactgagga	aacccccggg	gtacaggact	gctactacca	ttatggacag	3300
agttaccgag	gcacatactc	caccactgtc	acaggaagaa	cttgccaagc	ttggtcatct	3360
atgacaccac	accagcatag	tcggacccca	gaaaactacc	caaagtctgg	cctgaccagg	3420
aactactgca	ggaatccaga	tgctgagatt	cgcccttggg	gttacaccat	ggatcccagt	3480
gtcaggtggg	agtactgcaa	cctgacacaa	tgcttgggtg	cagaatcaag	tgtccttgca	3540
actctcacgg	tggtcccaga	tccaagcaca	gaggcttctt	ctgaagaagc	accaacggag	3600
caaagccccg	gggtccagga	ttgctaccat	ggtgatggac	agagttatcg	aggctcattc	3660
tctaccactg	tcacaggaag	gacatgtcag	tcttggctct	ctatgacacc	acactggcat	3720
cagaggacaa	cagaatatta	tccaaatggg	ggcctgacca	ggaactactg	caggaatcca	3780
gatgctgaga	ttagtccttg	gtgttatacc	atggatccca	atgtcagatg	ggagtactgc	3840
aacctgacac	aatgtccagt	gacagaatca	agtgtccttg	cgacgtccac	ggctgtttct	3900
gaacaagcac	caacggagca	aagccccaca	gtccaggact	gctaccatgg	tgatggacag	3960
agttatcgag	gctcattctc	caccactgtt	acaggaagga	catgtcagtc	ttggtcctct	4020
atgacaccac	actggcatca	gagaaccaca	gaatactacc	caaattggtg	cctgaccagg	4080
aactactgca	ggaatccaga	tgctgagatt	cgcccttggg	gttataccat	ggatcccagt	4140
gtcagatggg	agtactgcaa	cctgacgcaa	tgtccagtga	tggaatcaac	tctcctcaca	4200
actcccacgg	tggtcccagt	tccaagcaca	gagcttcctt	ctgaagaagc	accaactgaa	4260
aacagcactg	gggtccagga	ctgctaccga	ggtgatggac	agagttatcg	aggcacactc	4320
tccaccacta	tcacaggaag	aacatgtcag	tcttggctgt	ctatgacacc	acattggcat	4380
cggaggatcc	cattatacta	tccaaatgct	ggcctgacca	ggaactactg	caggaatcca	4440
gatgctgaga	ttcgcccttg	gtgttacacc	atggatccca	gtgtcaggtg	ggagtactgc	4500
aacctgacac	gatgtccagt	gacagaatcg	agtgtcctca	caactcccac	agtggccccg	4560
gttccaagca	cagaggctcc	ttctgaacaa	gcaccacctg	agaaaagccc	tgtggtccag	4620
gattgctacc	atggtgatgg	acggagtatt	cgaggcatat	cctccaccac	tgtcacagga	4680
aggacctgtc	aatcttgggt	atctatgata	ccacactggc	atcagaggac	cccagaaaac	4740
tacccaaatg	ctggcctgac	cgagaactac	tgcaggaatc	cagattcttg	gaaacaaccc	4800
tggtgttaca	caaccgatcc	gtgtgtgagg	tggtgagtact	gcaatctgac	acaatgtctca	4860
gaaacagaat	caggtgtcct	agagactccc	actgttgttc	cagttccaag	catggagggt	4920

BI OL0250W0SEQ_ST25. txt

cattctgaag cagcaccaac tgagcaaacc cctgtggtcc ggcagtgcta ccatggtaat	4980
ggccagagtt atcgaggcac attctccacc actgtcacag gaaggacatg tcaatcttgg	5040
tcatccatga caccacaccg gcatcagagg accccagaaa actacccaaa tgatggcctg	5100
acaatgaact actgcaggaa tccagatgcc gatacaggcc cttggtgttt taccatggac	5160
cccagcatca ggtgggagta ctgcaacctg acgcgatgct cagacacaga agggactgtg	5220
gtcgtcctc cgactgtcat ccaggttcca agcctagggc ctcttctga acaagactgt	5280
atgtttggga atgggaaagg ataccggggc aagaaggcaa ccactgttac tgggacgcca	5340
tgccaggaat gggctgcccc ggagcccat agacacagca cgttcattcc agggacaaat	5400
aaatgggcag gtctggaaaa aaattactgc cgtaaccctg atggtgacat caatggtccc	5460
tgggtgtaca caatgaatcc aagaaaactt ttgactact gtgatatccc tctctgtgca	5520
tcctcttcat ttgattgtgg gaagcctcaa gtggagccga agaaatgtcc tggaagcatt	5580
gtaggggggt gtgtggcccc cccacattcc tggccctggc aagtcagtct cagaacaagg	5640
tttgaaagc acttctgtgg aggcacctta atatccccag agtgggtgct gactgtgct	5700
cactgcttga agaagtcctc aaggccttca tcctacaagg tcctcctggg tgcacaccaa	5760
gaagtgaacc tcgaatctca tgttcaggaa atagaagtgt ctaggctgtt cttggagccc	5820
acacaagcag atattgcctt gctaaagcta agcaggcctg ccgtcatcac tgacaaagta	5880
atgccagctt gtctgccatc cccagactac atggtcaccg ccaggactga atgttacatc	5940
actggctggg gagaaaccca aggtaccttt gggactggcc ttctcaagga agcccagctc	6000
cttggtattg agaatgaagt gtgcaatcac tataagtata tttgtgctga gcatttggcc	6060
agaggcactg acagttgccg ggggtgacagt ggagggcctc tggtttgctt cgagaaggac	6120
aaatacattt tacaaggagt cacttcttgg ggtcttggct gtgcacgccc caataagcct	6180
ggtgtctatg ctcgtgtttc aaggtttggt acttggattg agggaatgat gagaaataat	6240
taattggacg ggagacagag tgaagcatca acctacttag aagctgaaac gtgggtaagg	6300
atttagcatg ctggaaataa tagacagcaa tcaaacgaag aactgttcc cagctaccag	6360
ctatgccaaa ccttggcatt tttggtattt ttgtgtataa gcttttaagg tctgactgac	6420
aaattctgta ttaagggtgc atagctatga catttgtaa aaataaactc tgcacttatt	6480
ttgatttga	6489

<210> 2
 <211> 150001
 <212> DNA
 <213> Homo sapiens

<400> 2	
atctttcagc ctctatatta ttttattgtg atttttaatt tccttgaatt ggattttgcc	60
attgtgctaa atcttgatga tcttcatttg tatccgtagt ctgaattata tttctgtcat	120
ttgagttagc tcagtcttgt taagaaccct tgttggaaaa ctggtgcagt tgtttggagg	180

BI OL0250W0SEQ_ST25. txt

acatatgacc ttctggccat ttgatttatt ggagttctta cgttggttct ttctcatgtc	240
tctgtgtggg tgtttcttta actgcagtgt agattgagta cagccaatag acttcttctt	300
tggaggtttt cacagggcca aggccttgta cagggtcttt atttgtagct gacttcttgt	360
ctttggtttc atagtggggc atgttagcaa aatagttttg ctgttgaagt tttggggtgt	420
gatccatttt ttattttaat gatttgttat ttcctttata cctaaaacaa gcagaaaacc	480
agtaaaggtc tttgagtctc tgaattcata actccagcat tcatattgct tcctcaggta	540
agtggggttt tcacccagcc ctttaagggtg ttagattatt ttttatgtga aattagccag	600
attgtatttc taaacatgat gtaaaacaat aatgacaaaa gttataataa actagccttc	660
ttaccaaadc cacatgtcta atgtgtgtgg gaggtgtta ggcaggggac ctgcagctaa	720
gggagaggca gacaggcccc atggcccaa atctaggata gtatttggtg ttggttgatg	780
ggtgagagaa agagaggga catctgtgca ggatgtggtg tcagcacctg gactacatct	840
tagggattcc ttcttcatth ttcagtatgc cctgacaata attatatcta tcagacttac	900
ccccttgacc actggaacac taagactgtt ttgggatctc tgcctgactt tctcagaggt	960
gctggtgagg acattatgag tctggaacct agaaaagcgt tctgactctg ctgactttct	1020
cagaggtgct ggtgaggaca ttatgagtct ggagccctag aaaagcggtc tgactctgcc	1080
actagccaga cagacctgga ctaggcacgt taactctttg tatgacttga ctccaacccc	1140
tcatttgtaa aaccagcatt ttcaagtggg tttttccaca tcagcctttt gcataagctg	1200
tcatttgaag aaaggthttt gtttgtttgt tttttgttta acaaaaaggt taaaaaccac	1260
tggcttagat aattgcaaag tttgctttcc tttttctgtg ctttttctac tatttttaaa	1320
atgtcatcct ctttggtttc ttgatcccc tttctgcact cctgagtctg ggaacactga	1380
ggccaactaa aaggaaactt ggcaaaagag gaacaccttt ggggtgtgcca ggctgctccc	1440
agtgttttgc acttataaaa atttaaatgc tgcaaacctc taagacttag atattattgt	1500
tcctatttta caagttagga acctgaggct cagagaaggt gcaggatggc acaggagagac	1560
ctgaattgga accctgggtc ccacttactg gctgtcggga cttagaaaag tcatgaactc	1620
tcattgattg ttttcttata tgaaatggg gctgcagggt tgtcggggga gaaacaataa	1680
gaatgtgcat caagtgtcga gcacgtgcta cgcactccat catggcagct cctactaata	1740
tacagaatag agttgtatct aacatgactc tttcttgcaa gtgacagaaa atccaactta	1800
agatggatta agcaaaaaag gggatttctt gttgagctga aaagtcttta ggctcacatg	1860
atggccccag ggcccaggcc ctgtccagcc atgcagtagg catcatcctt gggcaciaag	1920
gtgagattct tgtggtggca gatgctgtgg cagctcttgc tttgccagga aagactgagg	1980
aaggccactg tccccattaa gtgaacaata gttggccagg tctgagaggt tgaacttggg	2040
tcacaggcct gtccctgaac ccatcactga ttggctccaa cctgcatcag ctattacatg	2100
ctagagggtg aggcaggacc ccactcatac ccagaagggc aaagggtgga tccctcaaca	2160
ggattatggg atgtagggtg atagactgct gggcagccag aaagcaaaca gatcctctcc	2220

BI OL0250W0SEQ_ST25. txt

aatacctcaa ctgatgaaag caccaagcta aaatcataag gatctgggtg tgaattctgg	2280
ctctaccatc ttccatgtga cattgggcag ttatttaatc tcttttagcc ttggctttct	2340
tacctgtact aacatataag gtgattgtga tgagcatcat catcgtcaac atcatcatca	2400
ccatccacat tgccaccacc actcccatta tcatcttcat caacatcatc accaccgcca	2460
ccatcaccat tatcattacc accaccgcta tcaactattat catcacctc aacatcatca	2520
ccatcatcac tatcatcacc accaccatca tcgttactac cactaccacc accatcatca	2580
ccacagccac caccaccatc accatcatta ctactcagca ccaccatcat cattccacca	2640
ccatcaccat cattccacca tcaccattat cattaccacc accactgtca ctattatcat	2700
cacctcaac atcatcacca ccaccatcat cattactacc accaccacca ccatcaccat	2760
catcatcatt ctaccacat caccattatc atcaccatca ccatcaccac cgctatcatc	2820
atgataatca ttatcattac caccaccatt agcattatca ttaccaccac catcactatc	2880
actatcacca tcaccagac cactaccacc atcaccaaca ccatcattac taccaccac	2940
caccatcatc atcattccac caccatcaca attattacca ccaccacat caccaccacc	3000
accaccatca ctatcatcat cagtagacat catataacca gtttgtagct ggcccagagc	3060
ctacttgctg tttcttctgc ccacaacca tccacacatt tctaaccacc atccccact	3120
aggcttctgc ctgcctgggt ctcacctgca ggtccactga gaaaatgatt ctcagaacac	3180
taactagacc atgagggtgcc aaaaacata actcaggcct gttcatcaat tttctacatg	3240
tcaataatga catcagggtca attggcggtc tcagcctctg agagggaggt caaagttttc	3300
ctgctctccc cttcatgttt ccagggtgtc cctgacttgg atcaaatgca gagtttgag	3360
gtgttgaggc caaggggatt ttccagggtca gtcgtcatcc acaatcaatg gactgatcct	3420
gccgctggac ttaccctgct gccctctccc caaggcccca tcaggaggagg cttcaatcct	3480
cttgtcacct gtggcctacc tgccctcaga gatgacatct ctatgtcggc cactggatgg	3540
cagcacctac tcgcagacca catcaacttt cctggcaact tcggttaggt tttcaccatt	3600
atcaggatgt ttgccttgct caaatagcag attctagaga acggtgctcc ctcacacaac	3660
tatgtagtcc aggtgatgca ccctctgccc gatgcttggg agtcagaaac ttccatcatg	3720
cagctctgcc cagattgagc tgagctggcc tctggagtga ggtgctggga caaacatctt	3780
ccatgctgct catgtcaact ccagatgcag tcaggtttct gaaccaaagt caatgatcta	3840
agtgcagtca aaggctctgg ggaagaaag agagagtgcc tcatctcttg cctgtgccat	3900
gctcgcaaag caaggatttt tgcaaaattc taatgaaagc tgggcttgca aaattagaaa	3960
actggattat ttgtgagaac actgaaacat ccctgggtgt gtccatctgg aaaaacagca	4020
tttcctctgg caattttgca accgttctat ttgaatttgg caaagaaaat aaagcagttt	4080
ttcacaaaag aataaacaca accaggagaa tcttcactct ccaaattgt caaagaagta	4140
taaattagaa aatgaatcag gacaatttca acctgttaga ttagctaata tttaaaaatt	4200
gaacactcat acaagtgtgg tgaagtgatt gttttctagt gacattttac actgtcataa	4260

BI OL0250W0SEQ_ST25. txt

ccttctagaa aataaattgg cagtgttatt gggagacaga aatatgtcta tataatztat	4320
gggaacttag gctcagaaaa tattaaggaa taagaatgaa ctttatgaac aaagatgtgg	4380
agggttggaa gcaagagggg ggccaacgcg cacggggagg aagcatttgg gcagtgactc	4440
cgcagaccca ggctcagggt gaactagaca acctccttac acctcagttt ccttaactgt	4500
agagcaggag tgatggaact gcctgtttca taggactgtt gtgaggatga agtgagatac	4560
accacattat aagcttgtgc ctggaaagga taatgcttag taaatgatga ctattctttt	4620
ttattgcaat aaaatgtaca cagcgtaaga gttactatit taaccattit tgcagggtag	4680
caccaagtgg catttagtac attcacagtg gtgtgcaacc atcatcatat ttccagaata	4740
ttttcctcat ccccaaagga aacctcatgc tcattaatca gtagctctcc tttaaaatat	4800
tagttatgaa gatcatagca ctatacaaaa ctcatatgt aatgttgagt gaaaaaatca	4860
gggtgtgaaa ttttgtgata tgatgtaatt agtgaaagaa gcatacaaaa agtctgaaaa	4920
tataaaaaca atagcaattg cttttctcag actctacatt taaacattat tctttatggg	4980
tttaaaagca aagaaaaagg taaagaaaca acaaccaacc gcaaagcacc atgacaaagc	5040
tcagattgtt aaatccagggt ttttggaaca tagactctta tatgacgttt acactctcca	5100
gggttcagag agtctggcag cattgggagc tgccttgtgt tctacagcct cacggacaga	5160
caggagggtcc atcaccactg ctctgttctt ctggagtttc cttgtgaaca tgttgtggac	5220
gtagttacca tttctttcat ctttttaaac acaggtagct ttggggctgg ctttctcaag	5280
gaagcccagc tccctgtgat tgagaatgaa gtgtgcaatc gctatgagtt tctgaatgga	5340
agagtcaaat cactgagct ctgtgctggg catttggtg gaggcattga cagttgcaag	5400
gtaagaaaag atcaagagac caaagttagt cttgtgtctt cctgtctcag tctcagtcct	5460
ttagacttga gtcccaaagt agcgaattca agtaggattt aatcaatgga agaccccagt	5520
ctaagtgttg ctcaaaaact ccctagatct gtcccaaagt tatattcaga tcatccaagg	5580
ggacttcttg gggcttgagt tccagatcag cagcaagga gccataagtg ccataactac	5640
ctcagaccac tcacctctct ggggtgtccc ggtggccagg gactaaagtg gtgatttttc	5700
tggtagggaa ggaggtagag ggtacaggac agagactaac tgcacacaat atctgagact	5760
ggagctcaga tattgtctgat gatcagagtt ggcgtgtctc cccaattgat ttacaactgg	5820
ggcttgata ctgttttaaa cgggaggagc ctcctaacca tcttgacaca accactgacg	5880
tgactacact agagatagac tctttccact taattctacc actcttgctt tacttcatga	5940
gaacgaaaat gtaagattgc accatgaatt catttgcgga aagattgata ctatgctttt	6000
attttatttt attttatttt attttatttt attttatttt attgagactc tcaccccggg	6060
tgaagtgcac tgacgtgatt ttggctcact gcaacttcca cctcctgggt tcaagtgaat	6120
actccagcct ccctagtagc tgggattaca ggtgccacc accacgcctg gctaattttt	6180
gtatttttag tagagatggg gtttcaccac attggcctgg ctggtctcaa actcctgacc	6240
ttgtgatcca cctgtcttgg cctcccaaag tgctgggatt acagagttga gccaccgcac	6300

BI OL0250W0SEQ_ST25. txt

tcgaccctat gttttatatt taaaaatatt tttttatatt ttttaagccac aactactaga	6360
ataggaagga ttgatatttt attaatattt tttggatatt attatttttt tttctttcct	6420
gagacattct tgctctgtca cccaggctgg agtgcagtgg cacattcttg gctcactgca	6480
acctccatct cctgtgttca agcaattcta gtgcctcagc ctacttagta gctgggatga	6540
ctggcatgtg cctccacacc cagctaattt ttgtattttt tgtagagaca gggttttggc	6600
atgttgccca ggcttgtctc aaactcctgg cctcaggtga tccatctgcc gtggcctccc	6660
aaaatgctgg gattataggc atgagccacc acccctcctt ggaaggattg atatcttata	6720
acataattta taattacaga aaacatgtga gttcactagg aataaataaa ttttgaagat	6780
aataaaagat tttcacttat gttgtcattt cggcacagtt tggatatagga tgtggagatg	6840
ttaacattta tacctagctt gctcgtaaac taagacctga aagggttggtg tctatcagct	6900
gcaccctgg gtagcgacac aacctcggga aggcctcagc cccctcctcg tacagcactg	6960
cctgttgga agcttgaggg aggctatgga tgtgcagcac ttggcagagg gtctggatcat	7020
ggaagttacc agcaaataatg agctactttt atgattttat tttatccaaa agaaagagaa	7080
tgaaagaaga ggggaggaaa caagactaat caggaaagat gaagggtctag gggtgaggga	7140
aggagtaagg agacataaag gcaatgtgga gcagctgagg ggggaaatgg ctttcaccac	7200
ttcccagcat ctattgacat tgcactctca aatattttat aagactctat attcaaggta	7260
atgtttgaac cctgctgagc cagtggcatg ggtctctgag agaatcatta acttaatttg	7320
actatctggg ttgtgggtgc gtttactctc atgtaagtca acaatgtcct gggattggga	7380
cacactttct gggcactgct ggccagtccc aaaatggaac ataaggaagt ggttcttcta	7440
cttcttttat ttctgaaatc aggttaagaca tagttttttt aaattataag aattattttt	7500
tctcccacaa tgtagtaaaa atacatatgc catggcttta tgtgcaattc atttaatttt	7560
tgattcatga aattcccagt tcaaaatctt gtatatgatt gaaaaattct taaaaaata	7620
agtttaattt ccccgatgaag actgtcacgg tgctggaatg aatgggcaga aaaaataatg	7680
gttgattttt ctaatctaaa agagtgtgcc tacatgatgg ccagtctggc tgaaaaataa	7740
atagccattg tagctaacta tgcaaaggat ggctaagctc ttcgcttggt tctcagtttc	7800
attaatttat atcatctctg ttcagggtgcc atgctcccct cactagcaag ttgaaacaat	7860
gaaataactc tttgaatatg tttggttcct tgacctgttc atggagtggg actcagcatt	7920
tctctctttg ttatggcctg agtaaggctt tccatcggtg tacatttgct tcttatccct	7980
ggagaaatta tacacatcca tttgccagat gatatacgca tataatgatt caacaaatac	8040
tcagggtatt tgttgagtgg gttaggtccc cacattttta tacatacata cacacataca	8100
caccgtgtgt gattgtgaat gtaagtgtgt gtcctttaca aatactagct tatttagctc	8160
atgggtatagg tagggtagca tagtcatccc cattttataa acaaagaaat ctagacttag	8220
gaaaatcatg ttatttgtct cgtgaccaa ttcccaaatc aaggaaataa agaaacctgg	8280
atttaagcca gattccaag aaaaaatcta gggctcttct cactttttca tctttgttcc	8340

BI OL0250W0SEQ_ST25. txt

aacatttgaa	aaaataaatc	taaacacatt	ccaatgtaac	tgaagagcag	gttaattggt	8400
tgccacttgc	agaatccaat	taagaagaga	gaagtctggt	ataaagaaag	tgatttgctt	8460
ccaaagctag	cttaggggaa	gaaatgcagc	agtcctgccg	tactacttca	ctttaggagc	8520
agaaagtggc	acttttaaaa	ggcaacagag	gaggcgagca	aggattcagg	ggtccatgct	8580
agcttgggca	ccttatccac	caggtagtgt	agcagttgcc	tgctggtgcc	tttgtgagca	8640
gggtgttgtc	ccttgaggca	aatctctgga	gggtgagagt	tttgtagtgg	gcatgctttg	8700
gtttataaat	cacctgtgaa	ctcaggagt	ccatcttgaa	gcacatacat	agttagatga	8760
acttgccctg	cagggagagt	ctgatgaaag	ggaggtagat	gcttgcaatt	taatctataa	8820
attaccagat	aaaattttac	aagttgactt	taaagtcaaa	cacatttgaa	tttagtgga	8880
gccattcaag	aaaatatcaa	agaaaataca	gagcaggaga	agattaagca	aagagttttt	8940
tggggaaatt	ggtgtctatg	tctgtgtgtg	tagggagtgc	aggggatatg	aatattctat	9000
ttcagcccat	ggaaactagg	atgtagatca	ctgtgaactt	attcagcagg	ctacacccaa	9060
aggctagaac	aaacttctct	gccacaggat	taacatatgt	tttaatcgac	ctggggggca	9120
cattctctga	taagctcttt	tggaaagcca	ggctttctgt	ggacgtgtta	tctttccaat	9180
gtgtgctgga	atgcccgggg	agaggaaaaa	gtttctttta	cagccatgct	cagtgagaag	9240
cggagaaaca	tcttctattc	acaaattgct	aagtctttta	cacatgcaaa	tatgcataca	9300
cattcacaca	ccacagttag	gaagaaattc	tcacaccatt	aataaaaatac	atttacttca	9360
gtagcaatat	acatctacat	tttgcctata	atataaaagt	atttttccta	ttaaaagatt	9420
tgtttaatgt	ttcttcacca	acaaataaac	cctattaaat	ccccattgcc	atatgagccc	9480
tggaggtgaa	tcagagaaac	aaaaggattg	tggaaaaatc	atcagggttaa	aaaaagaaaa	9540
attgattctg	ttttgggata	tttcctagca	acatgagctg	gggaggggat	ctcagcagtg	9600
atgctctatg	aagcataata	aaatgacaca	gttacaggta	acttagttaa	agggggaaat	9660
aaatggaagt	ttcctctttt	tgaatatcaa	ttgtagcctg	ctctgctaca	tttcaaaaac	9720
actcttcaaa	atgtttaact	gaactcactg	taggaagcac	cttattaatt	tattgtgtgt	9780
tttgaagtca	cactgtgagc	tatagaattt	acccaagcac	aactcttcct	ggaaaagaga	9840
gttcaaatga	gaaacagtgc	gggtggaaga	catggatatg	ggcctaaaat	atctatttct	9900
caatgatatt	ttgatataat	tatcaagtgc	tttttagtgg	attaggttca	gaatgcatca	9960
gccaatgcct	gttcaataat	ccagttttcc	agcatagagc	atattaaatt	gaggaaggac	10020
aaagtcacag	aggtggggag	caggtggact	gtggccaagg	actttgcatg	aaacagttag	10080
cgtgcatcct	cctccttgcc	ctgccctcat	ggtctgtgta	ctctcaggag	gtcaggacag	10140
gcctttctga	gaatgagaat	ctgttcatct	gcctttctac	tgatacttg	tcatcggcac	10200
acaaacacat	gttctctgca	gtgtgtcatc	tttcagaacc	tcccctgacc	ctgtattccc	10260
tagaagtctc	gctgctttca	gagccaggct	tctctcctgc	tgccaccccc	actgctcttc	10320
tagtcactct	ttaaccact	ccatctgcat	gtggccccc	ccacaccct	caaagtggtc	10380

BI OL0250W0SEQ_ST25. txt

aaggttgtcc	tgttgcttaa	ttccatggaa	gcttggctat	cttcatttta	ttagcctctt	10440
ttggcctctc	accctgtgaa	aatcactaca	ttttgtgcca	gagatggagc	tggcatctcc	10500
aggcttggaa	gagggctgct	gaagctcagc	caggtgtcct	aaggagcctc	aggacagggg	10560
atgctcagta	gccttgcaat	gggaacacag	ctgagcccca	cttggccacc	ctttgccaca	10620
accaggcaga	aagcagcttt	tgaacagatt	tgttgccctca	gatttgatct	caaagaaaaa	10680
tcgtgggcag	tatttggtccc	aggttctgct	tttttacaat	ttcctctgaa	atctggatgc	10740
ctatcaacac	cttggaaaaa	ctgaattctc	cccaactaat	agtgggtgtgt	cactgtagta	10800
agcctagtac	aaaaatggcc	ttctttgtgg	aggagcttca	tatcctccat	tttttttttg	10860
cttaattttt	gccaagatg	agaacataat	ttagttcact	ttttatttat	tccaacatc	10920
atccatgcac	caacattttt	gtaactaaag	gagggaccat	tcagaagatg	cttatcaact	10980
gtcaaagtga	cagtgttaca	accaatgcac	atattgtaag	aatcaaaca	atggcctcca	11040
aggttcattt	ctacacaggg	attagcagat	caacatcaat	cttggcaaca	cagttgccac	11100
tgatggtgtc	ttattttttt	tatcatgaca	tggcaatcaa	gagcaaacad	gattttattct	11160
tatttaagat	tttatggtta	gactaggcag	atagctagat	atgagcagga	ggtggaagcc	11220
cctgagagaa	tggaggtctg	gagaatctga	aaccccgagag	attacccaag	tcctgcatgc	11280
tagacatgag	tggaggaggg	ggaataccta	ggtagaaaag	aatgcccctt	aagatgcccc	11340
gcagtcgctc	actgtgcagt	taacttttca	gaatgctgct	agatacatgc	tgatagggag	11400
ggaagagggc	aaaggagaaa	ttcctaagag	atacacggtt	gcagttagta	tacatctgag	11460
tgctatacaa	ccttcttttg	gtggtggcaa	gaagcaatgc	agccattacg	tagaattcat	11520
atcaaacacc	tgtatcacag	gtgttaaaga	aacaagaaac	attgtacttc	ttgtattctt	11580
aataatgatt	tgcaatattg	tctttagtat	cactgcaaac	ctctataaat	atgattttta	11640
aaaagtattt	ctttaggttg	gaattacttc	tacgcattga	cttatcttcc	tgggtttcat	11700
tagccgtacc	cgttgtactt	tcttccttac	cactgtttat	ctcaaactct	tgagattaaa	11760
gtatgggctc	aggaggggagc	gaggagcttc	aggactctca	cggacctcca	gcacagtgtg	11820
gctgccttat	ggaaaagtgg	ccacactgtt	ttctgactg	gtccctgccc	ctactattcc	11880
tcactgggca	gagcacagcc	accctggccc	tgctgaaca	ttttagtcag	tgttggctct	11940
gtgcttctct	ggggaggaaa	tccaagagac	aaccacagc	ccctctgcca	tttcagctgc	12000
agcagtacca	ccgttaatgc	ccttgggctt	gagaaagaag	ggacctggcc	acttccctga	12060
cacctccagc	acacagcagg	gaaagaattc	cagtttctct	ttcttgtgag	ctttcacctg	12120
ctactcttca	ccaggcaagg	ctcctggctt	gggcccacag	tgaggccacc	tcgaactcag	12180
ttgaacattt	ccactggctg	cactctgtgt	ttttgtgggg	tgaagctccc	agaggtgact	12240
gaaagtcctt	ctgccactaa	cactgcagtc	atactgccct	tgctgtactt	ggactagggg	12300
aggaaaaaag	atcctgagtg	ctttactcac	accccagtgt	gccccagcca	ccctatggaa	12360
aagaggccag	tgtgtcatcc	ctgcaagcac	cctgaggccc	ctgcccctgc	tgccccaag	12420

BI OL0250W0SEQ_ST25. txt

ctgtagagcc	agaatataaa	gctggcagaa	aaatgtaaaa	aggctagact	ggcttagcct	12480
cccagcctac	atctttctcc	tgtgctggat	ccttcctgct	cttgaacatc	ggactccaag	12540
ttcttcagct	gtgggacttg	gactgtcttc	cttgctcctc	agattgcagg	tggcctatta	12600
tgggaccttg	taatcttgtg	agttaatacc	acttaataag	ctcccccttg	tgtgagtata	12660
tctatatcta	tagatagata	taggtatact	cactatatat	acacatatat	acatatactc	12720
tctctctctc	tctctcatat	atatatatat	ataatctcct	attagttctg	tccctctaga	12780
gaaccccgac	taatacagat	tttcatacca	gaagtggttc	ttgaggaaca	gaatattaag	12840
gatggaattc	tttcattggg	tttgggactt	ctgggtgttg	ctgattaata	tgattagacc	12900
aaaaaatgct	aaggactcta	cttctaatag	tatggagaac	actgatagta	cttggcctga	12960
attgtttaga	gagttatgca	aaataaatgc	atttgacact	actgattcat	cacttatgag	13020
aggcaaggag	tttagtgact	ctatacataa	tacctttgac	tatatgtgga	gaaccaagga	13080
acataatgaa	gttggttgat	tgctcctaag	ttctctggag	aaagagatga	aagaaaatga	13140
tgatctcagg	ggatctgtct	cccaccttca	gaagcagata	ctgagccaca	aatctgctaa	13200
gattgccctg	aatgagagtt	ttaactcctg	tagagaaaga	gttgaaattg	tgaaaaaaca	13260
gagacaagct	gttatcatgc	gagtagctga	tctgcaacaa	gaggtgcatg	cacagccttg	13320
ccagggtgtt	actgttaaa	tgagggcatt	gactggaaaa	aaatgggacc	ctggaacttg	13380
gagtggggat	gtgtggggaga	accctgatga	agctgaggac	actgagtttg	tgaactctga	13440
tgaaactttt	ttgccagaag	aaacagtttc	cccatcccca	gtagtggtaa	catcccctcc	13500
ctgaccctg	ctgccattag	cctttccacc	tttgtctgag	gatgtaaacc	ctgcactgct	13560
tgaggcaaca	gtgatggcct	tccctgaggc	agctgccagg	caagataatg	ttgattctcc	13620
tcaagaggca	cccctaata	ccctgaatgc	ttctagacct	ataactaggc	taaattcctt	13680
gcgggcccc	gaggtgaggt	tcagagtgtg	acccatgagg	aggtgcatta	tactctaaaa	13740
gaactgctta	agctttctaa	tttatatttg	cagaaatctg	gagaacaggc	atgggaatgg	13800
atattaaggg	taagggataa	tgggtggaag	gacatagagt	tggatcaagc	tgaatttatt	13860
ggtttgccc	tactaagtag	ggattctgca	tttaatgttg	cagctcgggg	acttagaaaa	13920
ggttctgata	gggccgggag	cagtggctca	cgctgtaat	cccagcacct	tgggaggcgg	13980
gggcgggcag	atcacgagat	caggagattg	agacaattct	ggctaaaatg	gtgaaacccc	14040
atctctgcta	aaaatacaaa	aattagctgg	gcatggtgat	gcgtaactgt	aatctcatct	14100
acttgggagg	ctgaggcaag	agaactgctt	gaacctgtga	ggcagagatt	gcagtgaacc	14160
aagatcgccc	cactgcattc	cagcctggta	acagagcaag	actccatttc	caaaaaaaaa	14220
aaaaaaaaag	ttataatagt	ttatttgctt	ggttagctga	aatatggatt	aaaagatggt	14280
ccaatgttag	tgagctggaa	atgccttggt	ttaatgtaga	ggaagtgatc	caaaggctta	14340
gggagattag	gatggtggag	tggattagtc	actttagacc	tactcatccc	agctgggagg	14400
gtccagaaga	tacacccttg	gccgaagctt	tgtgaaatag	atttgtgaga	gcagcacctg	14460

BI OL0250W0SEQ_ST25. txt

tatTTTTgaa	gagcccgtaa	ttgctcttct	ctgtatgtca	gatctaacag	taggaaccac	14520
agtcactcaa	ctacaaaatt	taaatacaat	gggaataatt	ggatcctgag	gtggcagggg	14580
ccaagtgttg	gcactgaacc	atcaaaggca	aggtgggcat	aactaccata	atagacagca	14640
gaggcaaagc	agccatcaga	atagtctgac	tcatgtagag	ctctggcatt	ggctaattaa	14700
tcatgggtgtt	cctagaagtg	aaattgatgg	gaaacctact	gtattcctac	ttgatttata	14760
taaacaaaaa	actgccaggt	agaatggact	aaagactaat	ctgaattata	aaaacagaga	14820
atcatggggc	ctcaatcaat	ttccagactc	gaacctgtta	cagttccaga	acccactgaa	14880
tgaagggggag	gctggatccc	cttgaggaag	gacaccacta	ggctactgac	aacttatgct	14940
gttactcttt	ctccatcct	tccctaagga	gacctctggc	cttttaccag	ggtaactgtg	15000
tgtactggag	aaagggaggt	aatgagacat	ttcagaaagt	actggacact	ggctctgagc	15060
tgacgttgat	tccagggtag	ccaaaacgtt	attgtggttc	cccagttaaa	gtaggggctt	15120
atggagggtta	ggtaattaat	ggagttttag	ctcatttctg	acttacagtg	gttccagtgg	15180
gtccctggac	ttatcctctg	gtcattttcc	cagtgccaaa	atgcataatt	tgtatagaca	15240
tacttattag	ctggcagaaa	tgccacattg	gctccctgac	tggtaggatg	agggttatta	15300
tggtgggaaa	ggccaaacag	aagccattag	agctgtctct	acctagaaaa	ataaaaaaat	15360
caaaaacaat	atcccatccc	tggagggact	gaagtgatta	gtgtcaccat	caaggacttg	15420
aaagacgcag	gggtgggtgat	tcccaccaca	tccctgttca	actctcccat	ttgacctgtg	15480
cagaggacag	atggatcttg	gaaaatgatg	gtggattatt	ttaagcttaa	ccaagtgggtg	15540
actccaattg	cagctgctct	accagttgtg	gttttgttgc	ttgagcaaat	taacacatct	15600
cctgggtgcct	ggtatgcagc	cattggcttg	gcaagtggct	ttttctccat	tcctgtccat	15660
aagaccacc	agaagcaatt	tgccttcagc	tgacaaggcc	agcattatac	ctttaccacc	15720
ctacctcagg	ggtgtatcaa	ctctccagct	ttgtgtcata	atcttatttg	gagagacctt	15780
gctcgctttt	cacttccacg	agatataaca	ctgggtccatt	acattcatga	cattatgatg	15840
attggataca	gtgagcaaga	agtagcaaac	acactgaact	tattgggtgag	acatttgtat	15900
gccagaggat	gggaaataaa	tccagctaaa	atttagggac	tttctacctc	ggtaaaaattt	15960
ctaggggttcc	agtggcatga	gacctatgga	gatattcctt	ctaaggtgaa	gcataacttg	16020
ctgcgttttg	cccctcttac	aaccaagaaa	gaggcacaat	gcctgggtggg	cctattttgga	16080
ttttggaggc	aacacattcc	tcgtttgggt	gtgttactct	ggccatttta	tcgagtgacc	16140
tgaaaggctg	ccagatttaa	gtgcagtcta	gaacaaaaga	aggctctgaa	acagggtccag	16200
gctgctgtga	aagctgctct	gccatttggg	ccacatgacc	ccgcagatcc	aatgggtgctt	16260
gagggtgtcag	tggcagatag	ggatgctgtt	tggagccttt	ggcaggcccc	cataggtgaa	16320
tcacagtgga	gacctctagg	attttggagc	aaggccctgc	cacttctgca	gataactact	16380
ctccttttga	gagacagcta	ttggtctgtt	attgggcttt	ggtggtaact	gaacgtttga	16440
ctgtgggtca	taaagtcacc	atgctacctg	aacctgccta	tcatgaactg	gttgctttct	16500

BI 0L0250W0SEQ_ST25. txt

gacccatcta gccatgaagt gggtcagcac agcggcattt catcatcaaa ttgaagtgg	16560
gtgtatgtga tcgggcttga gcaggctctg aaggcacaag taagttacat aaggaagtgg	16620
ctcaaatgcc catgttctcc actcatgcca ccctgccttc cctccccag cctgcaccaa	16680
tggcctcatg gggagttccc tatgatcagt tgacagagga agggaagact aaggactgg	16740
tcatagatgg ttctgcacga tatgcaggca ccacccgaaa gtggacagct gcagcactat	16800
atccactttc taaatgcatg tgtacacttg tgctaagaaa atatctttat tttatttcct	16860
ttatttttcc tttatcatgt gaccttagat ttatggactt cacatcagca ttttaagcatt	16920
taagtgttgt tcatatcagc atttaaatat tgttaacctt atgtaataac ttttggtttg	16980
gggattgggtg cgtttctggt tgtatgagga tagttgtatt atattaggca taattatgac	17040
cttattattg tctttatttg aagattatgt atgatttcag gatgtgtgta tgggttcaag	17100
ttgacaagga gttggacttg tgatggttaa tactgtcaac ttgattggat tgaaagatgc	17160
aaagtattaa tctcggttat gtctgtgagg gtgtggcaaa aggagattaa catttgagtc	17220
agtgggctgg gaaggcagac ccacccttaa tctgggtaca caccatctaa tcaagttcca	17280
gtgtggccag attgtaaagc agggagaaaa atgtgaaaag actagactga attagcttcc	17340
cagcctacat ctttctcctg tgccaaatgc ttctgtctct tgaacatcgg actccaagtt	17400
cttcagcggt gggagttgga ctggctttct tgctcctcag cttgcagagg gcctgttgtg	17460
gaaccttgtg atccgctgag ttaatactac ttaataagat cccctttata tacatataat	17520
atattatatt atatataata tatataatat atattatata taatatatat aatatattat	17580
atattatata taatatatat tatatattat atataatata tattatatat aatatatatt	17640
atatattata tattatatat aatatatatt atatataata tatataaaat atatatatat	17700
cctattagtt ctgtccctct agagaaccct gactaataca atttatgtca ttaatctcat	17760
ttattgattt gtatacattg aaccaacctt atatcccagg aataaaacct acttgattgt	17820
gggtggattag ctttttgatg tactcttgga ttcaattgct ggtattttat tgagaatttt	17880
tgcatctgtg ttcataagg atattggctt gaagttttct ttttttgttg ttccatatca	17940
gaatgatgac gacctcatag aatgagttag tctgtcctct tttatctttt ggaattgttt	18000
caggaggctt gatatcagct cttctttata tgactggtat actttggcta ggaatctctc	18060
tgggtccagg gtttttctgg tgtaggtttt taattactga ttcaacttca gaactcatta	18120
ctcattattg agttctaaaa ctcactttca tgtactcttc aaaagactgt cttcttctgt	18180
tgttgagcgg ggtgttctct caaggctggt taggtgaagg tggttgctgg tgttcttctg	18240
tatccttact gcttgtcttt ctcttttttt attgactact gaggattaat ggtgatgtgt	18300
ccaactttaa ctctagatta gtctatttct cttttagatt gtaactctgt tttatatatt	18360
ttgaagctct gttgttaggc atgtgtattt ggattgttag gtcttcttga tgatgacctt	18420
tatcattatg taatgtttct tcttatctct ggaagtattc gttgttctga agtctatttg	18480
tgctgatatg aatacagcct tcacagctct attttacta gtatttgtat atctttttct	18540

BI OL0250W0SEQ_ST25. txt

cagcttttaa attgagatgt tcagaccatt tgcattaaag tagttgttaa taggattaaa	18600
tttaaactta ccattaagtt gggtatttct ctttgtccca tttaaacttt gttccttttt	18660
tcataatttt ctgccttcat ttatattgag tttatctcca cgacttactt attaaattaa	18720
ttttaaatgg ttttagtatt ttccacaatg tttataatat atactttgat tttttcacat	18780
tccaccttca aatgacagaa ttatactgga tatatagaaa tcttacatca ttgcacttct	18840
ccttcctccc tctcaaaatg ttgtgctatt gctctttgta atagaggctt acttctatta	18900
tgttatagct ctcataatac attgacacta tttttaccct gaataatcag ttgtttttta	18960
aagtgattat gactacaaat attttgaata atttctttat tttaccattt ctggtgctcc	19020
ttatctttta cagtagatcc caatttccat ctggagtcac attctttctg tgaaaaacaa	19080
cctttagcat ttcttatagc acgggactgc tgttgctggt gtctttcagc ttttctttgt	19140
ctgaagaagt ctttattttg ccttcagttt taaaagtga ttttgctgag tatagatact	19200
gggttgagag ttcatctct tgtatcattt taacaatgat gttccattat attccgtttt	19260
gaatagtttc tgactagaaa tctgatcttt gtttctttgt attcaatagt tcctttttct	19320
ctgactgcct ttaagatatt ctcatctttg ttttcaaca gtttgactat aatttgttta	19380
ttattaactt ttgtatttta ttctgcttga ggtttctga gctccttgga ttgacagatt	19440
gttgattttt attgtttttg taaaattcat agccattatc tattctactg ttttgttttt	19500
tttttactt ctctctctct gtattcttct ttttgactg taagtattca aatgttagat	19560
cattcatatt gcttcataaa ccttatatgc ttcttctgct tttttttttt tgtcaggaac	19620
tctttttttg tatctgtggt gggttgata agttctagta gactatgttc aagtttatgg	19680
attattttgt tagttgtgtc taattgactc ctgagtgcac tcagagaatt ctcatctct	19740
gatattataa atctcttctt agcattttca tgttactctt ttctatagtt tccatctctt	19800
tgctgaaatt ctccccctat ccatggatat tgtccacctt taccacaaga ttctttaaca	19860
tattaacata ggtatcatc aaacccaaac tgatagtttc cagatggtgt cttttctgag	19920
tctgtctgtc ttgattgctt tattatttaa cagtgactta tcttccctct tcagcttttg	19980
gtgtgtcttg taattgttta atcaaacact gggatcata aatggaggaa cagtagagat	20040
tgagtaaat attatttatg ctttgaaatg ggcacccatc ttctgttgaa aatatgtttt	20100
gtggtcaatt gagtcaacct agtaactggt tgaactgaat ttggcatttg tgcttggtgc	20160
ttttatctta aatgcaccac aggtttaaat tcctccagtg atgggttgct gctatctttt	20220
gcttagagtg gggcctgggg tgtggaagaa ttttctcagt gttcctatct attattagat	20280
tttagcagtc actgcatgcc tgcactacag aggggatatc ttcatacaca taatctaacc	20340
ccattgaaac tgctgtttct tcttaatgaa tgctcaatct ttggtggaaa taaacaaatg	20400
ctgtatctcc tggagccact tcagtcttag tcaggttctg cagggctttg aagggaatgc	20460
attctcagta ttcttggtcc ttatttggat ggaacttgaa cctgtggtgg gtttggagag	20520
aaagagtagc agacgtctgc tatgttgcaa tgcaggatgc tgggcacaag aaaatttcca	20580

BI OL0250W0SEQ_ST25. txt

gtctctcctc	caaggaaata	agatttgatc	atctacctat	ccctgagaag	tgaagggcct	20640
tgcctgcggt	gctagatgca	aaaccatttt	tctcccccca	ttgccagaa	acttaaggct	20700
ttggcctttc	tgagcagtgg	tctaggggaat	tgtgcaaggt	tttcatattt	gaccctgaca	20760
gcccatacacc	acctacagct	tgcagtgccca	aatgtatctc	cctctgatct	ctcctgtcct	20820
gtggtcctca	tgaacattaa	gaagagattt	ctaaaaaaga	gcttgcacat	gagcatagtt	20880
tctggtgaga	agaattctga	tatgttaact	tcctctaaac	ttttaataaa	aatattttcta	20940
agaattaaat	aaagtcttag	aatgatatga	atctattcct	ttggtttttt	gcacgtctgt	21000
ctgcctgcta	atcaagagaa	gagaatggtc	gtaattctca	gagacttttt	cctgtttgtg	21060
tcataaatga	cttcacattt	ttttctgttc	taagaactat	tcagcttgat	ttcttctgtt	21120
ttaatttttag	cagcacctga	gcaaagccat	gtgggtccagg	attgctacca	tggatgatga	21180
cagagttatc	gaggcacgta	ctccaccact	gtcacaggaa	ggacctgccca	agcttgggtca	21240
tctatgacac	cacatcaaca	taataggacc	acagaaaact	acccaaatgc	gtatgtcatt	21300
aatcttacag	taagcaaaac	aagggtccaag	taaaatttgt	cttagaaaag	gtgtgcgtca	21360
agctaacttc	ttatgattaa	atttttctca	cacatagaat	gcatggcaaaa	atgtctgaga	21420
aacattactt	tgagcaaaga	gtatgataga	agagaaatgt	taagctggct	ctctttcctg	21480
agagtttgat	aaaatcagga	gaatatctgg	cggtggtgag	gccacaataa	tggaaaatca	21540
gaatgttttag	acagagtcag	cttcaacaac	actcactaaa	ggatcaatgtg	atctttaccc	21600
cttgaaattc	tataattcta	atctccaatt	cctgaagtga	aggttgtgtt	ggccttttct	21660
gtcttggctc	acaagtaaat	gatatgtgca	tatctatgga	aaggcgaatc	tatctttttc	21720
tatatctatg	tctattccaa	cgggtagaaa	caccctgggt	cctgagcacc	agtgggtctga	21780
aggaatacgg	gttgccagga	agagagaagc	aaaggcagga	aggcagatga	aagtaagaaa	21840
tgagacagat	gctaaacaat	aaaaagtgcg	ggaagataga	cagaagctgg	ggtctgacca	21900
caccatggcc	agtctttcac	acataagtga	ctaccaaaaga	caagaaaaaa	tgattttccgc	21960
ttgttggaca	atagatggta	gaggaccaag	ggaattgcga	gagagagaac	aatgagatca	22020
actcaacaga	tgacttggtt	ttcttcctgg	agacccttcc	tgactgaag	ggcaggagat	22080
ggagcccaaa	aaaaactgta	gccatcttgc	tgaacagagg	aggacattg	gagtttggga	22140
ttattcaggt	ggctaggatt	ttctaggcct	gctaacaatg	agaacagatt	tgtggaggaa	22200
aggagtctta	gaaatatgca	tagaaatctc	ctcgagtcac	tggctaaaca	tgaagctgca	22260
tgtacacaga	aaatagatcc	acaagaaagt	agggcaaaga	acatctacgg	aagagcagca	22320
actacaatgg	aacagtgagc	tcaataaaca	tgacagagct	caaatagcac	taagggatat	22380
tggagtittg	accacacaga	ggagagagac	ttcactgaac	atcttgggca	ttcagtagag	22440
accaggaaa	agccatactt	taggagtaga	attagtatat	tcttagaata	aaggcagctc	22500
cacacaaaca	atagcaaaac	tgaaaaggaa	gtctccaagc	atcagaatga	tgtccaagtc	22560
aatgaactgc	ctctgagagg	aaaactcaac	catctttaga	ggtaaacatc	aaagtcaagt	22620

BI OL0250W0SEQ_ST25. txt

ggctcagcta	tgcagtatcc	acagtgtgag	gcctaaatat	aaaacttgac	tacacataga	22680
aaccttttag	tgtgaccac	aagcaggagg	aaaatcagcc	aatacaaaca	gacccagaag	22740
agacagaaat	gattagaatg	gcataaaaaat	ttgacatatc	actatataat	aattgagttc	22800
taggatttaa	gaaaacatga	atatagaatg	caacagacac	cttatccaga	gacagtaaga	22860
gtataaagag	ccaaatcgaa	gaactactaa	gagatatgtc	ttaaatgaaa	aaattactag	22920
atggcctccc	catctagtta	gacatttcag	aagaaaatac	caaataaaaa	ataattgcat	22980
agaacctaca	gaaccagata	cacacataca	aaacacacgc	atgcatacac	acacactcaa	23040
acatgtataa	gcttacaaac	acacacacac	atccacaaat	gctgaaaaat	gaaatcaacc	23100
gagccacaca	gacataaagg	aaaacataaa	aagatttcct	acatgtggga	agcaagtcac	23160
agaaaggggg	aaggagattg	gaacagaaat	atatactgaa	agcaaggatg	gctgaaaatt	23220
ttccaaatat	aaagaagatt	aaaaaatcac	ggactcaaga	agctcaatgg	atcagaaaaa	23280
taatttctaa	aatgacaatt	ataggatgcc	actgggtaca	tagcagttca	actgtcagag	23340
ggcaaagaca	taatacacag	aaaaatctcg	taaggaacgg	gaaaaacaaa	aagctgtgtc	23400
ttgctagagg	aacagtgata	caagtgacta	atgtgttccc	atcagaaaca	ctgcaacctg	23460
gacacaaaag	aataacatta	aagtaataaa	cgtaagaaag	aagagctcaa	ctgagaaggc	23520
tacatccagc	aataaaatgc	cttgaagttc	atccatgttg	gaggaatgca	cattgtgcac	23580
tcccctaaac	aaagaaaccg	gaaactgtaa	gactttggaa	tcagcaggct	tatgtaacaa	23640
aagaggtgac	cctaaggaat	taaggagaag	aagaatagaa	caagaaggga	actttctgca	23700
gcctatataa	tgaagaacct	agcaattggc	aaatgtagat	gaaaatgcta	catgttttct	23760
tgatcaaacg	tttatatctt	tttaaattgag	agttgacgag	ttgaagcaaa	atgataccaa	23820
tatatttaac	tttaccatat	gtagaagtaa	aaatttgaac	atgtagcata	aatcatgtag	23880
ggattaattg	gaagtgtacc	actgtaagtt	tcttacctca	tgcacgatag	tatgtaatac	23940
taataaaagg	ttaatgtgtg	ggttcaaagg	gatattgcaa	atcctagagc	aatcacaaag	24000
tttttaactc	tgaggtttgt	tgtataataa	caatatTTTta	tgtattcaaa	agaggggaagc	24060
caaggaagaa	aaaaaagtct	ttaaagagct	ctggctctta	gtacatccag	ttgctcattg	24120
aatgagcttc	ctggaatgga	gggtctggga	ctgagactag	gccacatgtg	tagagccact	24180
agagacacaa	tgttggatcc	ccatggccca	taatacatTTt	cccatttttct	caggcagcca	24240
caggatcatga	atgtgaggat	actgagaggt	tggagcaacg	ttcttgggag	gcataaggaa	24300
gagcgaatgc	ttcaagatcc	ccgcagccca	aactcctcag	ctgctttgcc	tcctaattca	24360
ttgttttttg	ctcctccata	gctgtccgac	ctcttcagat	ctcttagtct	tcctgccatc	24420
ttcctttatg	ccatgggacc	cactgttctt	tcaactcatc	ccccagttct	ggagtggctg	24480
tggacagcag	aggatagact	gagagcagga	gagaaggtcc	tgcccaggaa	cccatttctag	24540
agatactgca	ttctgcctgg	gagcaagttt	tccagggcag	ctttgagaag	tcttgcagaa	24600
acaaacctac	ttgaccgaca	tgatatggga	atgacagaca	gtaatactat	ttgcacaatg	24660

BI OL0250W0SEQ_ST25. txt

cttttccatg ggaaaggtag agccttttca ctaggttttg agtacatgga gtgtgagagt	24720
tgacctggaa aggttatcct ccttgatgcc atgttttctc tgaagaacta catgttcgtt	24780
gcaactccca cattagaata tgaagtccta ccgagagaga tacggagact agacagatac	24840
agatgcattt gcatgtgaat acacaatccc acaatacaga cgtcaaaacc cataccagtt	24900
attccagaga gatggattgg gcagaaggca gaaggagaat actctgatcg tttttcggcc	24960
acgtgtgtgt gttatctcag tgtttctaag aagcgtttgc tacttttagat tttttattta	25020
aaaaaaatag taataatcta ttaagtatga gagatgtgca gagaggatta gtgatcgaga	25080
gccatttttg ctggtggcaa tcatatggta cttttaatgg gaatattaga aaggcaccgg	25140
taatgacctt gttgcagcac aaaggagaga gtgtgggggtg cccctgcatg ttgtcccacc	25200
tcttgtgacg tgtatcgttt tgggaatttcc agtggccttga tcatgaacta ctgcaggaat	25260
ccagatgctg tggcagctcc ttattgttat acgagggatc ccggtgtcag gtgggagtac	25320
tgcaacctga cgcaatgctc agacgcagaa gggactgccg tcgcgcctcc gactgttacc	25380
ccggttccaa gcctagaggc tccttccgaa caaggttaagg agtctgtggc cagacatcta	25440
cacgcttcga tgctgggatg aaaagccatg gaaattccca ctgatgcagc cgccttcaat	25500
ggtaaacgga tgctcgagt tggcctgagt tctaccatgt aggaggaagc ctccgtgcac	25560
tctctggggg agccagcgga gtgatttctg gtgcaacgtg gttgggcttt gtcttttagga	25620
tgggcacaaa ccctccaggg ggatcgactt caaaattcac cttgtttgtaa aacgggctac	25680
ctcagtgtcc cagccaaaat ttttattgta acatgctgtc aggtgtgtca ctctttccaa	25740
gccagtaagc ttttccgggg atttcttcaa gtagccagca ttcagagcaa tcttcagcat	25800
tgcagattct gagaaatgtg gctctggagc ctgtcaccct cgagaaacct aagagggctg	25860
cattgattcc atgtggccct gggctctatg agcagtacat gagctcccag tgctctaagg	25920
ctcttcagcc ctaggctttg aaggagtgta tttctcagta ttcttaaacc tctttctgat	25980
gacacttgta cctgtgaggg gtctagagag aaagagtagt agactcctac ttactacaa	26040
ttcaggatgc agggcatgag aggattccct ctctcctcca agggaagaag cttttggcgt	26100
gcacacatcc ctgagaagca aagtgtcttt gtcttcagtc agatacatag gaccgttttc	26160
tgcccatg ccggaagcc aaaggccttg gctttcatga tcaacggtct agggaaacat	26220
gcaaaatttc catgtctgtc ccaaactctg cccccgacag ccaattacca cctgcagccc	26280
gcattgccaa atgcggtgcc gtttgcagta agattcagta gagtttccta gaaaggtgct	26340
acctcgtgag ctactttcc aatgaggaat ctgatctgtt gtgtttctct aaggtgtcag	26400
gtgaaatatt tccaagaact tactacagtt ctagaatggg aggaatctgt tgctttggtg	26460
tttgtttgtt ggtcggtttt ctacatcca tctgcctatg gataaggaaa agagaacggt	26520
cgtaattctc atagactcct ttctggttgt gtcacaaatg gcttcacatg tttctctatg	26580
ctcagagata ctacagcttga tttcccgtgt tttcatttca gcaccgactg agcaaaggcc	26640
tgggggtgcag gagtgtacc atggtaatgg acagagttat cgaggcacat actccaccac	26700

BI OL0250W0SEQ_ST25. txt

tgtcacagga	agaacctgcc	aagcttggtc	atctatgaca	ccacactcgc	atagtcggac	26760
cccagaatac	tacccaaatg	cgtatgtctt	tgttctttac	cataagagaa	gaaagggcca	26820
agtgaagttt	ctgttacaag	agatgtgtct	caagctgagt	tctccgaact	caacttgtga	26880
cagatgcaga	tggcgtagca	aatgtctca	ggatgattgc	cttgagacta	agggctctgag	26940
agaagggaaa	tgtaagctc	cctctccttc	ctcctagttc	tattgagcag	aagggaaatc	27000
tggaggtgag	gagatcacat	tatgaagaaa	gtcagaatga	caaaggacca	gacacttaga	27060
ttacccttcc	acaacaccaa	ctaaacgtca	atggagactt	tccagttgga	attccgttat	27120
tctggcttcc	acttcctgaa	gggaagggtg	cgtttgcctt	ttctctctgg	gttcaagagg	27180
aaagaatagg	tgcttattta	tggacaggtg	aattgatctg	tttctatatc	tacgtatatt	27240
ccgattgtca	gaaaaacact	cgttcctaag	taccagtggc	ctgaagggat	acaggttccc	27300
agcaagagaa	gatccaagga	aggaaggcag	atgagagtca	gcacagagag	ggatgctgaa	27360
aagtaaaagg	gatgggtgga	tggagagaag	cccgggtctg	accaccaat	ggccaatatt	27420
ttggccacaa	gcgactacca	gagacatgga	aaaatggttt	ctacatgtgg	gacaacagat	27480
ggtagaggac	ctagagaatt	gagagagggg	caatgatggg	ctccactccg	cagatgcctt	27540
ggctttcttc	ctggataccc	ttcctgcact	gaatagcaag	gagatggagc	ccaagcagac	27600
tgtagccatc	ttgctgaatg	gaggagaggg	attggagttt	gggatgactg	tggtagctga	27660
aatttttcta	ggtctgctag	aaataagaac	tggtttgtgt	ggaggaaaag	agctctacaa	27720
atacgcatag	aagtctcctc	cagtcgttgg	cctgacatga	cgctgcctgt	gcacaggaaa	27780
tggttccacg	agaaagtgtg	gcaaagaaca	tttactgaga	aacagcaagt	acaagagcac	27840
aggaagctca	ataaagaaga	gagagatcac	atagcactct	gggatactgg	agttcttccc	27900
agctagacca	gagagtcctc	acggagcaca	ttgccaattc	agtggagacc	ccagaacagc	27960
cgtaatttaa	aggtacactt	agtatattac	tagaataaag	tcagctgcag	acaaccctt	28020
gcacagctgg	aaagcaagtg	tccaagcatc	aaatcggttt	ccaatcaatg	aagtgcctgt	28080
gagaggaaat	ctcaactctc	tttagaagta	aacaacaaag	tcgattgcct	cagctatgcg	28140
gtatccgcag	agtgagtcct	aaatttaaaa	tctgactaca	tgtagaaaag	cgtttcgtgt	28200
gacccatgac	caggaaataa	atcgggtaat	acaacaggc	tcaggaatga	gagaaatgat	28260
tagaattgcg	tgaaaatttg	acatatcagt	atgataactg	atttcaaata	tttaaaaaaa	28320
caacatgcaa	gaaagcagat	atcatatcaa	gagaaattaa	cagtacagaa	tagccaaatt	28380
aaattaaaga	ggtagtataa	aaaaagtatg	tcttaattga	aaaaaattac	tgtatggccg	28440
gctgatcaat	ttagacgttt	cagaggaaaa	cattacccaa	cacacaattc	tagagaacct	28500
acagaatgag	ctacacacac	acacacacac	acacacacac	acactgaaaa	cacaccata	28560
ctcacacaca	cgcagaaact	cacaagttct	aacacacaca	gacacgcgca	cccctgaaga	28620
aacagtgaat	tataaaatta	agcgagcctc	acagacatgt	aggaaaatat	gaaaagattt	28680
cctgcatgtg	ggaagcaagt	cacagtaaag	agcaagggag	tttataatag	aaacaaatac	28740

BI OL0250W0SEQ_ST25. txt

cagaatcaag gatggctgat aacttttcaa ttacgaagaa cattaataaaaa aatcacagaa	28800
tcgtgaaact caagggatca tatagggaat ttcggaaaaa aaaccaacc tgtatgatgt	28860
acttttgtac atcacagttc gaaggtaaca aggcaaatat gtaataagaa gaaacctgtc	28920
acgagaaact ggaggaaaa gagctgtgtc ttcctacaag tacactgata caaatgcca	28980
atgtgttcac ctacagaaaca ctggaagcca gataccaggg aatattgtta aaatgataat	29040
caggaacaaa aagagatcaa ccgggaatgc tgaatccagc aataaaatgc cttgaaggtc	29100
atccatgtcg gataaatgca tattgtgcac tgcccaaag aaagaaaccg gaaactgtaa	29160
gaattgaaa tcagcaggct tatgtaacaa gagagggtgac ccgaaggaat taggtagaag	29220
aagaattgaa caagaaagga actttctgca gccacgtaa tgaagaatcc agcaattggc	29280
aaatgtagat agatgtaaat gcaaaatatt ttcttgatca aatttctata tctttgtaaa	29340
tgagagtga ctactgaaa caaatgata gcaagatatt taacttcagc atatgtagag	29400
gtaagaattt gaaatgtag cataaatcac gaagggatta attcgaagtg taccgttgta	29460
agtttcttta cctcatgcac gatggtgtgt catattaata aaagggtact gtgcgggttc	29520
gaagggatat tgcaaactct agagcaatca caaagggttg aactctgagg tttttggtat	29580
aataagaata gtccatgcat tcaaaagagg gaagccaagg aagaactaga agtctttcaa	29640
gagctcaggc tcttatacat ccagttgctc attgaaccag cttcctggaa tggagggctc	29700
ggggttgaga ctaggccaca agtctagagt ctctagagag acagtgttgg aaccccatgg	29760
cccataatac atttcccatt ttctcaggca gccagaggtc atgaatgtga ggatactggg	29820
aggttggagc aacgttcttg ggaggcataa ggaagagcga atgcttcaag atccccgcag	29880
cccaaactac tcgcctgctt tgccccctaa tgcatttttc tctgctgctc cgtagctgtc	29940
cgacctcttc agatctctta gtccaccctg ccgtcttctt ttatgcatg ggtccactg	30000
ttctttcaac tcatccccct ttccctcagt cccggagtag ctgcggccag cagagggtag	30060
actgagagca ggagagaagg acctgcctag gaacccttc tagagatact gcatcctgcc	30120
tgggagcaag tttccaggg cagctttgag aagtcttgga gaaacaaacc tactaaacct	30180
gacagacagt aatactatct gcacaatgct tttctgtggg aaaggtagag ctttttact	30240
acgtattgag tacatagagt gtgagggttg acctggaacg gctatcctcc tggatgacgt	30300
gtgttttctg aagaactaca tgttcgttgc aactcccaca ttagaatatg aagtcctacc	30360
gagagagata cggagactag acagatacag atgcatttgc atgtgaatac acaatcccac	30420
aatacagacg tcaaaaccca taccagttat tccagagaga tggattgggc agaaggcaga	30480
aggagaatac tctgatcgtt tttcggccac gtgtgtgtgt tatctcagtg tttctaagaa	30540
gcgtttgcta ctttagattt tttatttaaa aaaatagtaa taatctatta agtatgagag	30600
atgtgcagag aggattagtg atcgagagcc atttttgctg gtggcaatca tatggtactt	30660
ttaatgggaa tattagaaag gcaccggtaa tgacctgtt gcagcacaaa ggagagagtg	30720
tggggtgccc ctgcatgttg tcccacctct tgtgacgtgt atcgttttgg aatttccagt	30780

BI OL0250W0SEQ_ST25. txt

ggcttgatca tgaactactg caggaatcca gatgctgtgg cagctcctta ttgtttatacg	30840
agggatcccc gtgtcagggtg ggagtactgc aacctgacgc aatgctcaga cgcagaaggg	30900
actgccgtcg cgcctccgac tgttaccccc gttccaagcc tagaggctcc ttccgaacaa	30960
ggtaaggagt ctgtggccag acatctacac gcttcgatgc tgggatgaaa agccatggaa	31020
attcccactg atgcagccgc cttcaatggg aaacggatgc tcgagtgttg cctgagttct	31080
accatgtagg aggaagcctc cgtgcactct ctgggggagc cagcggagtg atttctgggtg	31140
caacgtgggtt gggctttgtc tttaggatgg gcacaaaccc tccaggggga tcgacttcaa	31200
aattcacctt gttgtaaaac gggctacctc agtgtcccag ccaaaatttt tattgtaaca	31260
tgctgtcagg tgtgtcactc tttccaagcc agtaagcttt tccggggatt tcttcaagta	31320
gccagcattc agagcaatct tcagcattgc agattctgag aaatgtggct ctggagcctg	31380
tcaccctcga gaaacctaag agggctgcat tgattccatg tggccctggg tctatggagc	31440
agtacatgag ctcccagtgc tctaaggctc ttcagcccta ggctttgaag ggagtgtatt	31500
ctcagtattc ttaaacctct ttctgatgac acttgtacct gtgaggggtc tagagagaaa	31560
gagtagtaga ctctactttt actacaattc aggatgcagg gcatgagagg attccctctc	31620
tcctccaagg gaagaagctt ttggcgtgca cacatccctg agaagcaaag tgtctttgtc	31680
ttcagtcaga tacataggac cgttttctgc cccatggccc ggaagccaaa ggccttggct	31740
ttcatgatca acggtctagg gaaacatgca aaatttccat gtctgtccca aactctgccc	31800
ccgacagcca attaccacct gcagcccgc ttgccaaatg cggtgccgtt tgcatagaaga	31860
ttcagtagag tttcctagaa aggtgctacc tcgtgagctc actttccaat gaggaatctg	31920
atctgtttgtg tttctctaag gtgtcagggtg aaatatattc aagaacttac tacagttcta	31980
gaatgggagg aatctgttgc tttgggtgtt gtttgttggg cggttttctc acatccatct	32040
gcctatggat aaggaaaaga gaacggtcgt aattctcata gactcctttc tggttgtgtc	32100
acaaatggct tcacatgttt ctctatgctc agagatactc agcttgattt cccgtgtttt	32160
catttcagca ccgactgagc aaaggcctgg ggtgcaggag tgctaccatg gtaatggaca	32220
gagttatcga ggcacatact ccaccactgt cacaggaaga acctgccaag cttggtcatc	32280
tatgacacca cactcgcata gtcggacccc agaatactac ccaaatacgt atgtctttgt	32340
tctttacat aagagaagaa agggccaagt gaagtttctg ttacaagaga tgtgtctcaa	32400
gctgagttct ccgaactcaa cttgtgacag atgcagatgg cgtagcaaaa tgtctcagga	32460
tgattgcctt ggagctaagg gtctgagaga agggaaatgt taagctccct ctcttctctc	32520
ctagttctat tgagcagaag ggaaatctgg aggtgaggag atcacattat gaagaaagtc	32580
agaatgacaa aggaccagac acttagatta cccttcaca acaccaacta aacgtcaatg	32640
gagactttcc agttggaatt ccgttattct ggcttccact tcctgaaggg aaggttgctg	32700
ttgccttttc tctctgggtt caagaggaaa gaataggtgc ttatttatgg acaggtgaat	32760
tgatctgttt ctatatctac gtatatccg attgtcagaa aaacactcgt tcctaagtac	32820

BI OL0250W0SEQ_ST25. txt

cagtggcctg	aagggataca	ggttcccagc	aagagaagat	ccaaggaagg	aaggcagatg	32880
agagtcagca	cagagagggg	tgctgaaaag	taaaagggat	gggtggatgg	agagaagccc	32940
gggtctgacc	acccaatggc	caatatTTTT	gccacaagcg	actaccagag	acatggaaaa	33000
atggtttcta	catgtgggac	aacagatggt	agaggaccta	gagaattgag	agaggggcaa	33060
tgatgggctc	cactccgcag	atgccttggc	tttcttcctg	gatacccttc	ctgcactgaa	33120
tagcaaggag	atggagccca	agcagactgt	agccatcttg	ctgaatggag	gagagggatt	33180
ggagtttggg	atgactgtgg	tagctgaaat	ttttctaggt	ctgctagaaa	taagaactgg	33240
tttgtggagg	aaaagagctc	tacaaatacg	catagaagtc	tcctccagtc	gttggcctga	33300
catgacgctg	cctgtgcaca	ggaaatgggt	ccacgagaaa	gtgtggcaaa	gaacattttac	33360
tgagaaacag	caagtacaag	agcacaggaa	gctcaataaa	gaagagagag	atcacatagc	33420
actctgggat	actggagttc	ttcccagcta	gaccagagag	tcctcacgga	gcacattgcc	33480
aattcagtgg	agaccccaga	acagccgtaa	tttaaaggta	cacttagtat	attactagaa	33540
taaagtcagc	tgagacaac	cccttgaca	gctggaaagc	aagtgtccaa	gcatcaaadc	33600
ggtttccaat	caatgaagtg	cctgtgagag	gaaatctcaa	ctctcttttag	aagtaaacaa	33660
caaagtcgat	tgccctagct	atgcggtatc	cgcagagtga	gtcctaaatt	taaaatctga	33720
ctacatgtag	aaaagcgttt	cgtgtgaccc	atgaccagga	aataaatcgg	gtaatacaaa	33780
caggctcagg	aatgagagaa	atgattagaa	ttgcgtgaaa	atttgacata	tcagtatgat	33840
aactgatttc	aaatatTTAA	aaaaacaaca	tgcaagaaag	cagatatcat	atcaagagaa	33900
attaacagta	cagaatagcc	aaattaaatt	aaagagctag	tataaaaaaa	gtatgtctta	33960
attgaaaaaa	attactgtat	ggccggctga	tcaattttaga	cgtttcagag	gaaaacatta	34020
cccaacacac	aattctagag	aacctacaga	atgagctaca	cacacacaca	cacacacaca	34080
cacaaactga	aaacacaccc	atactcacac	acacgcagaa	actcacaagt	tctaacacac	34140
acagacacgc	gcacccctga	agaaacagtg	aaatataaaa	ttaagcgagc	ctcacagaca	34200
tgtaggaaaa	tatgaaaaga	tttcctgcat	gtgggaagca	agtcacagta	aagagcaagg	34260
gagtttggaa	tagaaacaaa	taccggaatc	aaggatggct	gataactttt	caattacgaa	34320
gaacattaaa	aaaaatcaca	gaatcgtgaa	actcaagggg	tcacataggg	aatttcggaa	34380
aaaaaaccca	acctgtatga	tgtacttttg	tacatcacag	ttcgaaggta	acaaggcaaa	34440
gatataataa	gaagaaacct	gtcacgagaa	actggaggaa	aaagagctgt	gtcttcctac	34500
aagtacactg	atacaaattg	ccaatgtgtt	cacctcagaa	acactggaag	ccagatacca	34560
gggaatattg	ttaaaatgat	aatcaggaac	aaaaagagat	caaccgggaa	tgctgaatcc	34620
agcaataaaa	tgcttgaag	atcatccatg	tcggataaat	gcatattgtg	cactgccccca	34680
aagaaagaaa	ccggaaactg	taagaattgg	aaatcagcag	gcttatgtaa	caagagaggt	34740
gacccgaagg	aattaggtag	aagaagaatt	gaacaagaaa	ggaactttct	gcagcccacg	34800
taatgaagaa	tccagcaatt	ggcaaatgta	gatagatgta	aatgcaaaat	attttcttga	34860

BI OL0250W0SEQ_ST25. txt

tcaaatttct	atatctttgt	aatgagagat	tgactacttg	aaacaaaatg	atagcaagat	34920
atttaacttc	agcatatgta	gaggtaagaa	tttgaaatgg	tagcataaat	cacgaaggga	34980
ttaattcgaa	gtgtaccgtt	gtaagtttct	ttacctcatg	cacgatgggtg	tgtcatatta	35040
ataaaagggg	actgtgcggg	ttcgaaggga	tattgcaaat	cctagagcaa	tcacaaaggt	35100
ttgaactctg	aggtttttgg	tataataaga	atagtccatg	cattcaaaag	agggaagcca	35160
aggaagaact	agaagtcttt	caagagctca	ggctcttata	catccagttg	ctcattgaac	35220
cagcttcctg	gaatggaggg	tctgggggtt	agactaggcc	acaagtctag	agtctctaga	35280
gagacagtgt	tggaacccca	tggcccataa	tacatttccc	atcttctcag	gcagccagag	35340
gtcatgaatg	tgaggatact	gggaggttgg	agcaacgttc	ttgggaggca	taaggaagag	35400
cgaatgcttc	aagatccccg	cagcccaaac	tactcgcttg	ctttgcccc	taatgcattt	35460
ttctctgctg	ctccgtagct	gtccgacctc	ttcagatctc	ttagtccacc	ctgccgtctt	35520
cctttatgcc	atgggtccca	ctgttctttc	aactcatccc	cctttccctc	agtcccggag	35580
tagctgcggc	cagcagaggg	tagactgaga	gcaggagaga	aggacctgcc	taggaacccc	35640
ttctagagat	actgcatcct	gcctgggagc	aagttttcca	gggcagcttt	gagaagtctt	35700
ggagaaacaa	acctactaaa	cctgacagac	agtaatacta	tttgacaaat	gcttttctgt	35760
gggaaaggta	gagccttttc	actacgtatt	gagtacatag	agtgtgaggg	ttgacctgga	35820
acggctatcc	tcctggatga	cgtgtgtttt	ctgaagaact	acatgttcgt	tgcaactccc	35880
acattagaat	atgaagtcct	accgagagag	atacggagac	tagacagata	cagatgcatt	35940
tgcatgtgaa	tacacaatcc	cacaatacag	acgtcaaaac	ccataccagt	tattccagag	36000
agatggattg	ggtaggaggg	agaaggagaa	tactctgatc	gtttttcggc	cacgtgtgtg	36060
tgttatctca	gtgtttctaa	gaagcgtttg	ctactttaga	ttttttatct	aaaaaaaata	36120
gtaataatct	attaagtatg	agagatgtgc	agagaggatt	agtgatcgag	agccattttt	36180
gctggtggca	atcatatggg	acttttaatg	ggaatattag	aaaggcaccg	gtaatgacct	36240
tgttgcagca	caaaggagag	agtgtggggg	gccctgcat	gttgtccac	ctcttgtgac	36300
gtgtatcggt	ttggaatttc	cagtggcttg	atcatgaact	actgcaggaa	tccagatgct	36360
gtggcagctc	cttattgtta	tacgagggat	cccgggtgca	gggtgggagta	ctgcaacctg	36420
acgcaatgct	cagacgcaga	agggactgcc	gtcgcgcctc	cgactgttac	cccggttcca	36480
agcctagagg	ctccttccga	acaaggtaag	gagtctgtgg	ccagacatct	acacgcttcg	36540
atgctgggat	gaaaagccat	ggaaattccc	actgatgcag	ccgccttcaa	tggtaaacgg	36600
atgctcgagt	gttgccggag	ttctgccatg	ttgggggaag	cctccgtgta	ctctctgggg	36660
gagccagcgg	agtgatttct	ggtgcaactt	gggtgggctt	tgtctttaga	atgggcacaa	36720
accttccagg	gtgatgggct	tcacaactca	cctccttcta	aaatgggcta	tctcagtgtc	36780
ttagccaaaa	tttttattgt	aacgtgctgt	caggtgtgtg	attctttctg	tcgcagtaag	36840
cttttctggg	gatttcttca	agtagccagc	agtcagtgtg	atcttcagca	ttgcagattt	36900

BI OL0250W0SEQ_ST25. txt

caaaaaatgt	ggctctggag	cctgtcatcc	tcgagaaacc	taacagggct	gcattaattc	36960
catatggtcc	tgggtctatg	gagcagtata	tgagctccca	atgctctaag	gctcttcagt	37020
cctaggcctt	gaaggagtg	atttctcagt	gttcttaaac	ctctttctga	tggcacttgt	37080
acctgtgagg	ggtctagaga	gaaaggttag	tagacttctc	ctttactgca	attcaggatg	37140
cagggcatga	gaagattccc	tccctcctcc	aagggagaa	ggttttggcg	tgcacacatc	37200
cttgagaagc	aaagtgtctt	tgcccttcagt	cagatatata	ggatcgtttt	ctgccccatg	37260
gcctggaagc	cagaggcctt	ggctttcatg	atcaacgatc	tagggaaaca	tgcaaaattt	37320
ccatgtcttt	cccctcctct	gccctcgaca	gcccaattacc	acctgcatcc	tgcatgtcca	37380
aatgcagtgc	cctttgtatg	aacattcagt	agagtttcat	agaaagggtgc	tacttcgtga	37440
gcgcactttg	cagtgagaag	gagtctgttc	tgttctgttt	ttctaaggat	ttcagggtgaa	37500
atatttccta	gaacttacta	cagttctaga	ttggtaggaa	tctgtagggt	tgctgtatgt	37560
tttttggttg	gttttctccc	atccatctgc	ctacaggtaa	gggaaagata	acgttcgtaa	37620
ttctcataga	ctcctttctg	gttgtgtcat	aaatggcttc	acatatttcg	ttattctcag	37680
agatactcag	tttatttctt	gtgttttcat	ttcagcaccg	actgagcaga	ggcctgggggt	37740
gcaggagtgc	taccacggta	atggacagag	ttatcgaggc	acatactcca	ccactgtcac	37800
tggaagaacc	tgccaagctt	ggtcatctat	gacaccacac	tcgcatagtc	ggaccccaga	37860
atactacca	aatgcgtatg	tctttgttct	ttaccataag	agaagaaagg	gccaaagtga	37920
gtttctgtta	caagagatgt	gtctcaagct	gagttctccg	aactcaactt	gtgacagatg	37980
cagatggcgt	agcaaaatgt	ctcaggatga	ttgccttgga	gctaaggggtc	tgagagaagg	38040
gaaatgttaa	gctccctctc	cttcctccta	gttctattga	gcagaaggga	aatctggagg	38100
tgaggagatc	acattatgaa	gaaagtcaga	atgacaaagg	accagacact	tagattaccc	38160
ttccacaaca	ccaactaaac	gtcaatggag	actttccagt	tggaattccg	ttattctggc	38220
ttccacttcc	tgaagggaag	gttgcgtttg	ccttttctct	ctgggttcaa	gaggaaagaa	38280
taggtgctta	tttatggaca	ggtgaattga	tctgtttcta	tatctacgta	tattccgatt	38340
gtcagaaaaa	cactcgttcc	taagtaccag	tggcctgaag	ggatacaggt	tcccagcaag	38400
agaagatcca	aggaaggaag	gcagatgaga	gccagcacag	agagggatgc	tgaaaagtaa	38460
aagggatggg	tggatggaga	gaagcccggg	tctgaccacc	caatggccaa	tattttggcc	38520
acaagcgact	accagagaca	tggaaaaatg	gtttctacat	gtgggacaac	agatggtaga	38580
ggacctagag	aattgagaga	ggggcaatga	tgggctccac	tccgcagatg	ccttggcttt	38640
cttcctggat	acccttcctg	cactgaatag	caaggagatg	gagcccaagc	agactgtagc	38700
catcttgctg	aatggaggag	agggattgga	gtttgggatg	actgtggtag	ctgaaatfff	38760
tctaggctctg	ctagaaataa	gaactggttt	gtggaggaaa	agagctctac	aaatacgcat	38820
agaagtctcc	tccagtcgtt	ggcctgacat	gacgctgcct	gtgcacagga	aatggttcca	38880
cgagaaagtg	tggcaaagaa	catttactga	gaaacagcaa	gtacaagagc	acaggaagct	38940

BI OL0250W0SEQ_ST25. txt

caataaagaa gagagagatc acatagcact ctgggatact ggagttcttc ccagctagac	39000
cagagagtcc tcacggagca cattgccaat tcagtggaga cccagaaca gccgtaattt	39060
aaaggtacac ttagtatatt actagaataa agtcagctgc agacaacccc ttgcacagct	39120
ggaaagcaag tgtccaagca tcaaatcggg ttccaatcaa tgaagtgcct gtgggaggaa	39180
atctcaactc tctttagaag taaacaacaa agtcgattgc ctgagctatg cggtatccgc	39240
agagtgagtc ctaaatttaa aatctgacta catgtagaaa agcgtttcgt gtgacccatg	39300
accaggaaat aaatcgggta atacaaacag gctcaggaat gagagaaatg attagaattg	39360
cgtgaaaatt tgacatatca gtatgataac tgatttcaaa tatttaaaaa aacaacatgc	39420
aagaaagcag atatcatatc aagagaaatt aacagtacag aatagccaaa ttaaattaaa	39480
gagctagtat aaaaaagta tgtcttaatt gaaaaaatt actgtatggc cggctgatca	39540
aattagacgt ttgagaggaa aacattaccc aacacacaat tctagagAAC ctacagaatg	39600
agctacacac acacacacac acacacacac acacactgaa aacacacca tactcacaca	39660
cacgcagaaa ctcaacaagt ctaacacaca cagacacgag caccctgaa gaaacagtga	39720
aatataaaat taagcgagcc tcacagacat gtaggaaaat atgaaaagat ttcttgcagt	39780
tggaagcaa gtcacagtaa agagcaaggg agtttggaat agaaacaaat accggaatca	39840
aggatggctg ataacttttc aattacgaag aacattaaaa aaaatcacag aatcgtgaaa	39900
ctcaagggat catatagggg atttcggaaa aaaaacccaa cctgtatgat gtacttttgt	39960
acatcacagt tcgaaggtaa caaggcaaag atataataag aagaaacctg tcacgagaaa	40020
ctggaggaaa aagagctgtg tcttcctaca agtacactga tacaattgc caatgtgttc	40080
acctcagaaa cactggaagc cagataccag ggaatattgt taaaatgata atcaggaaca	40140
aaaagagatc aaccgggaat gctgaatcca gcaataaaat gccttgaaga tcatccatgt	40200
cggataaatg catattgtgc actgccccaa agaaagaaac cggaaactgt cagaattgga	40260
aatcagcagg ctatgtaac aagagagggtg acccgaagga attaggtaga agaagaattg	40320
aacaagaaag gaactttctg cagcccacgt aatgaagaat ccagcaattg gcaaatgtag	40380
atagatgtaa atgcaaaata ttttcttgat caaatttcta tatctttgta aatgagagtt	40440
gactacttga aacaaaatga tagcaagata tttaacttca gcatatgtag aggtagaagt	40500
ttgaaatggg agcataaatc acgaagggat taattcgaag tgtaccgttg taagtttctt	40560
tacctcatgc acgatgggtg gtcatattaa taaaagggtg ctgtgcgggt tcgaagggat	40620
attgcaaatc ctagagcaat cacaagggtt tgaactctga ggtttttggg ataataagaa	40680
tagtccatgc attcaaaaga gggaagccaa ggaagaacta gaagtctttc aagagctcag	40740
gctcttatac atccagttgc tcattgaacc agcttcctgg aatggagggt ctggggttga	40800
gactaggcca caagtctaga gtctctagag agacagtgtt ggaaccccat ggcccataat	40860
acatttccca ttttctcagg cagccagagg tcatgaatgt gaggatactg ggaggttgga	40920
gcaacgttct tgggaggcat aaggaagagc gaatgcttca agatccccgc agcccaaact	40980

BI OL0250W0SEQ_ST25. txt

actcgctgc	tttgcacct	aatgcatttt	tctctgctgc	tccgtagctg	tccgacctct	41040
tcagatctct	tagtccaccc	tgccgtcttc	ctttatgcca	tgggtcccat	tgttctttca	41100
actcatcccc	ctttccctca	gtcccggagt	agctgcggcc	agcagagggg	agactgagag	41160
caggagagaa	ggacctgcct	aggaaccct	tctagagata	ctgcatcctg	cctgggagca	41220
agttttccag	ggcagctttg	agaagtcttg	gagaaacaaa	cctactaaac	ctgacagaca	41280
gtaatactat	ttgcacaatg	cttttctgtg	ggaaaggtag	agccttttca	ctacgtattg	41340
agtacataga	gtgtgagggg	tgacctggaa	cggctatcct	cctggatgac	gtgcgttttc	41400
tgaagaacta	catgttcgtt	gcaactccca	cattagaata	tgaagtccta	ccgagagaga	41460
tacggagact	agacagatac	agatgcattt	gcatgtgaat	acacaatccc	acaatacaga	41520
cgtcaaaacc	cataccagtt	attccagaga	gatggattgg	gcagaaggca	gaaggagaat	41580
actctgatcg	tttttcggcc	acgtgtgtgt	gttatctcag	tgtttctaag	aagcgtttgc	41640
tactttagat	tttttattta	aaaaaaatag	taataatcta	ttaagtatga	gagatgtgca	41700
gagaggatta	gtgatcgaga	gccatttttg	ctggtggcaa	tcatatggta	cttttaattgg	41760
gaatattaga	aaggcaccgg	taatgacctt	gttgcagcac	aaaggagaga	gtgtgggggtg	41820
cccctgcatg	ttgtcccacc	tcttgtgacg	tgtatcgttt	tgggaatttcc	agtggcttga	41880
tcatgaacta	ctgcaggaat	ccagatgctg	tggcagctcc	ttattgttat	acgagggatc	41940
ccggtgtcag	gtgggagtag	tgcaacctga	cgcaatgctc	agacgcagaa	gggactgccg	42000
tcgcgcctcc	gactgttacc	ccggttccaa	gcctagaggc	tccttccgaa	caaggttaagg	42060
agtctgtggc	cagacatcta	cacgcttcga	tgctgggatg	aaaagccatg	gaaattccca	42120
ctgatgcagc	cgccttcaat	ggtaaaccga	tgctcgagtg	ttgcctgagt	tctaccatgt	42180
aggaggaagc	ctccgtgcac	tctctggggg	agccagcgga	gtgatttctg	gtgcaacgtg	42240
gttgggcttt	gtcttttagga	tgggcacaaa	ccctccaggg	ggatcgactt	caaaattcac	42300
cttgttgtaa	aacgggctac	ctcagtgtcc	cagccaaaat	ttttattgta	acatgctgtc	42360
agggtgtgtca	ctctttccaa	gccagtaagc	ttttccgggg	atttcttcaa	gtagccagca	42420
ttcagagcaa	tcttcagcat	tgcaatttct	gagaaatgtg	gctctggagc	ctgtcacctt	42480
cgagaaacct	aagagggctg	cattgattcc	atgtggccct	gggtctatgg	agcagtacat	42540
gagctcccag	tgctctaagg	ctcttcagcc	ctaggctttg	aaggagtgta	tttctcagta	42600
ttcttaaacc	tctttctgat	gacacttgta	cctgtgaggg	gtctagagag	aaagagtagt	42660
agactcctac	tttactacaa	ttcaggatgc	agggcatgag	aggattccct	ctctcctcca	42720
aggaagaag	cttttggcgt	gcacacatcc	ctgagaagca	aagtgtcttt	gtcttcagtc	42780
agatacatag	gaccgttttc	tgcccatg	cccgaagcc	aaaggccttg	gctttcatga	42840
tcaacggtct	agggaacat	gcaaaatttc	catgtctgtc	ccaaactctg	ccccgcagag	42900
ccaattacca	cctgcagccc	gcattgccaa	atgcggtgcc	gtttgcatga	agattcagta	42960
gagtttccta	gaaaggtgct	acctcgtgag	ctcactttcc	aatgaggaat	ctgatctggt	43020

BI OL0250W0SEQ_ST25. txt

gtgttttctct aaggtgtcag gtgaaatatt tccaagaact tactacagtt ctagaatggg	43080
aggaatctgt tgctttggtg tttgtttgtt ggtcggtttt ctcacatcca tctgcctatg	43140
gataaggaaa agagaacggt cgtaattctc atagactcct ttctggttgt gtcacaaatg	43200
gcttcacatg tttctctatg ctcagagata ctcagcttga tttcccgtgt tttcatttca	43260
gcaccgactg agcaaaggcc tggggtgcag gagtgtacc atggtaatgg acagagttat	43320
cgaggcacat actccaccac tgtcacagga agaacctgcc aagcttggtc atctatgaca	43380
ccacactcgc atagtcggac cccagaatac tacccaaagtg cgtatgtctt tgttctttac	43440
cataagagaa gaaagggcca agtgaagttt ctgttacaag agatgtgtct caagctgagt	43500
tctccgaact caacttgtga cagatgcaga tggcgtagca aaatgtctca ggatgattgc	43560
cttgagacta agggctctgag agaagggaaa tgtaagctc cctctccttc ctcctagttc	43620
tattgagcag aagggaaatc tggagggtgag gagatcacat tatgaagaaa gtcagaatga	43680
caaaggacca gacacttaga ttacccttcc acaacaccaa ctaaactgca atggagactt	43740
tccagttgga attccgttat tctggcttcc acttcctgaa ggaaggttg cgtttgcctt	43800
ttctctctgg gttcaagagg aaagaatagg tgcttattta tggacagggtg aattgatctg	43860
tttctatatac tacgtatatt ccgattgtca gaaaaacact cgttcctaag taccagtggc	43920
ctgaagggat acaggttccc agcaagagaa gatccaagga aggaaggcag atgagagtca	43980
gcacagagag ggatgctgaa aagtaaaagg gatgggtgga tggagagaag cccgggtctg	44040
accaccaat ggccaatatt ttggccacaa gcgactacca gagacatgga aaaatggttt	44100
ctacatgtgg gacaacagat ggtagaggac ctagagaatt gagagagggg caatgatggg	44160
ctccactccg cagatgcctt ggctttcttc ctggataccc ttctgcact gaatagcaag	44220
gagatggagc ccaagcagac thtagccatc ttgctgaatg gaggagaggg attggagttt	44280
gggatgactg tggtagctga aatttttcta ggtctgctag aaataagaac tggtttgtgg	44340
aggaaaagag ctctacaaat acgcatagaa gtctcctcca gtcgttggcc tgacatgacg	44400
ctgcctgtgc acaggaaatg gttccacgag aaagtgtggc aaagaacatt tactgagaaa	44460
cagcaagtac aagagcacag gaagctcaat aaagaagaga gagatcacat agcactctgg	44520
gatactggag ttcttcccag ctagaccaga gagtctcac ggagcacatt gccaatcag	44580
tggagacccc agaacagccg taatttaaag gtacacttag tatattacta gaataaagtc	44640
agctgcagac aacccttgc acagctggaa agcaagtgtc caagcatcaa atcggtttcc	44700
aatcaatgaa gtgcctgtga gaggaatct caactctctt tagaagtaaa caacaaagtc	44760
gattgcctca gctatgcggt atccgcagag tgagtcctaa atttaaaatc tgactacatg	44820
tagaaaagcg tttcgtgtga cccatgacca ggaaataaat cgggtaatac aaacaggctc	44880
aggaatgaga gaaatgatta gaattgcgtg aaaatttgaa atatcagtat gataactgat	44940
ttcaaatatt taaaaaaaca acatgcaaga aagcagatat catatcaaga gaaattaaca	45000
gtacagaata gccaaattaa attaaagagc tagtataaaa aaagtatgtc ttaattgaaa	45060

BI OL0250W0SEQ_ST25. txt

aaaattactg	tatggccggc	tgatcaat	tt agacgttt	ca gagggaaa	aca ttaccaaca	45120	
cacaattcta	gagaacctac	agaatgagct	acacacacac	acacacacac	acacacaaac	45180	
tgaaaacaca	cccatactca	cacacacgca	gaaactcaca	agttctaaca	cacacagaca	45240	
cgcgcacccc	tgaagaaaca	gtgaaatata	aaattaagcg	agcctcacag	acatgtagga	45300	
aaatatgaaa	agatttcctg	catgtgggaa	gcaagtcaca	gtaaagagca	agggagtttg	45360	
gaatagaaac	aaataccaga	atcaaggatg	gctgataact	tttcaattac	gaagaacatt	45420	
aaaaaaaaatc	acagaatcgt	gaaactcaag	ggatcacata	gggaatttcg	gaaaaaaaaac	45480	
ccaacctgta	tgatgtactt	ttgtacatca	cagttcgaag	gtaacaaggc	aaagatatata	45540	
taagaagaaa	cctgtcacga	gaaactggag	gaaaaagagc	tgtgtcttcc	tacaagtaca	45600	
ctgatacaaa	ttgccaatgt	gttcacctca	gaaacactgg	aagccagata	ccagggaata	45660	
ttgttaaaat	gataatcagg	aacaaaaaga	gatcaaccgg	gaatgctgaa	tccagcaata	45720	
aaatgccttg	aagatcatcc	atgtcggata	aatgcatatt	gtgcaactgcc	caaagaaaag	45780	
aaaccggaaa	ctgtaagaat	tggaaatcag	caggcttatg	taacaagaga	ggtgacccga	45840	
aggaattagg	tagaagaaga	attgaacaag	aaaggaactt	tctgcagccc	acgtaatgaa	45900	
gaatccagca	attggcaaat	gtagatagat	gtaaatgcaa	aatatcttct	tgatcaaatt	45960	
tctatatctt	tgtaaagag	agttgactac	ttgaaacaaa	atgatagcaa	gatatttaac	46020	
ttcagcatat	gtagaggtaa	gaatttgaaa	tggtagcata	aatcacgaag	ggattaattc	46080	
gaagtgtacc	gttgtaagtt	tctttacctc	atgcacgatg	gtgtgtcata	ttaataaaag	46140	
ggtactgtgc	gggttcgaag	ggatattgca	aatcctagag	caatcacaaa	ggtttgaact	46200	
ctgagggtttt	tggtataata	agaatagtcc	atgcattcaa	aagagggaag	ccaaggaaga	46260	
actagaagtc	tttcaagagc	tcaggctctt	atacatccag	ttgctcattg	aaccagcttc	46320	
ctggaatgga	gggtctgggg	ttgagactag	gccacaagtc	tagagtctct	agagagacag	46380	
tgttggaacc	ccatggccca	taatacat	ttt cccatt	tttctt	caggcagcca	gagggtcatga	46440
atgtgaggat	actgggaggt	tggagcaacg	ttcttgggag	gcataaggaa	gagcgaatgc	46500	
ttcaagatcc	ccgcagccca	aactactcgc	ctgctttgcc	ccctaata	gca ttttct	ctg	46560
ctgctccgta	gctgtccgac	ctcttcagat	ctcttagtcc	accctgccgt	cttcctttat	46620	
gccatgggtc	ccactgtttc	ttcaactcat	ccccctttcc	ctcagtcccc	gagtagctgc	46680	
ggccagcaga	gggtagactg	agagcaggag	agaaggacct	gcctaggaac	cccttctaga	46740	
gatactgcat	cctgcctggg	agcaagtttt	ccagggcagc	tttgagaagt	cttgagagaaa	46800	
caaacctact	aaacctgaca	gacagtaata	ctatttgcac	aatgcttttc	tgtgggaaag	46860	
gtagagcctt	ttcactacgt	attgagtaca	tagagtgtga	gggttgacct	ggaacggcta	46920	
tcctcctgga	tgacgtgtgt	tttctgaaga	actacatgtt	cgttgcaact	cccacattag	46980	
aatatgaagt	cctaccgaga	gagatacgga	gactagacag	atacagatgc	atttgcattg	47040	
gaatacacaa	tcccacaata	cagacgtcaa	aaccataacc	agttattcca	gagagatgga	47100	

BI OL0250W0SEQ_ST25. txt

ttgggcagaa ggcagaagga gaatactctg atcgtttttc ggccacgtgt gtgtgttatac	47160
tcagtgtttc taagaagcgt ttgctacttt agatttttta tttaaaaaaa atagtaataa	47220
tctattaagt atgagagatg tgcagagagg attagtgatc gagagccatt tttgctggtg	47280
gcaatcatat ggtactttta atgggaatat tagaaaggca ccggtaatga ccttgttgca	47340
gcacaaagga gagagtgtgg ggtgcccctg catgttgtcc cacctcttgt gacgtgtatc	47400
gttttggaat ttccagtggc ttgatcatga actactgcag gaatccagat gctgtggcag	47460
ctccttattg ttatacgagg gatcccgggtg tcagggtggga gtactgcaac ctgacgcaat	47520
gctcagacgc agaagggact gccgtcgcgc ctccgactgt taccgccggtt ccaagcctag	47580
aggctccttc cgaacaaggt aaggagtctg tggccagaca tctacacgct tcgatgctgg	47640
gatgaaaagc catggaaatt cccactgatg cagccgcctt caatggtaaa cggatgctcg	47700
agtgttgctc gagttctacc atgtaggagg aagcctccgt gcactctctg ggggagccag	47760
cggagtgatt tctggtgcaa cgtggttggg ctttgtcttt aggatgggca caaacctcc	47820
agggggatcg acttcaaaat tcaccttggt gtaaaacggg ctacctcagt gtcccagcca	47880
aaatttttat tgtaacatgc tgtcagggtg gtcactcttt ccaagccagt aagcttttcc	47940
ggggatttct tcaagtagcc agcattcaga gcaatcttca gcattgcaga ttctgagaaa	48000
tgtggctctg gagcctgtca ccctcgagaa acctaagagg gctgcattga ttccatgtgg	48060
ccctgggtct atggagcagt acatgagctc ccagtgtctt aaggctcttc agccctaggc	48120
tttgaaggga gtgatttctc agtattctta aacctctttc tgatgacact tgtacctgtg	48180
aggggtctag agagaaagag tagtagactc ctactttact acaattcagg atgcagggca	48240
tgagaggatt ccctctctcc tccaaggga gaagcttttg gcgtgcacac atccctgaga	48300
agcaaagtgt ctttgtcttc agtcagatac ataggaccgt tttctgcccc atggcccga	48360
agccaaaggc cttggctttc atgatcaacg gtctagggaa acatgcaaaa tttccatgtc	48420
tgtcccaaac tcttcccccg acagccaatt accacctgca gcccgcatg ccaaatgcgg	48480
tgccgtttgc atgaagattc agtagagttt cctagaaagg tgctacctcg tgagctcact	48540
ttccaatgag gaatctgatc tgttgtgttt ctctaagggtg tcaggtgaaa tatttccaag	48600
aacttactac agttctagaa tgggaggaat ctgttgcttt ggtgtttgtt tgttggtcgg	48660
ttttctcaca tccatctgcc tatggataag gaaaagagaa cggtcgtaat tctcatagac	48720
tcctttctgg ttgtgtcaca aatggcttca catgtttctc tatgctcaga gatactcagc	48780
ttgatttccc gtgttttcat ttcagcaccg actgagcaaa ggcctgggggt gcaggagtgc	48840
taccatggta atggacagag ttatcgaggc acatactcca ccactgtcac aggaagaacc	48900
tgccaagctt ggtcatctat gacaccacac tcgcatagtc ggaccccaga atactacca	48960
aatgcgtatg tctttgttct ttaccataag agaagaaagg gccaagtga gtttctgtta	49020
caagagatgt gtctcaagct gagttctccg aactcaactt gtgacagatg cagatggcgt	49080
agcaaaatgt ctcaggatga ttgccttggg gctaagggtc tgagagaagg gaaatgttaa	49140

BI OL0250W0SEQ_ST25. txt

gctccctctc	cttcctccta	gttctattga	gcagaagggg	aatctggagg	tgagaagatc	49200
acattatgaa	gaaagtcaga	atgacaaagg	accagacact	tagattaccc	ttccacaaca	49260
ccaactaaac	gtcaatggag	actttccagt	tggaattccg	ttattctggc	ttccacttcc	49320
tgaaggggaag	gttgcgtttg	ccttttctct	ctgggttcaa	gaggaaagaa	taggtgctta	49380
tttatggaca	ggtgaattga	tctgtttcta	tatctacgta	tattccgatt	gtcagaaaaa	49440
cactcgttcc	taagtaccag	tggcctgaag	ggatacaggt	tcccagcaag	agaagatcca	49500
aggaaggaag	gcagatgaga	gtcagcacag	agagggatgc	tgaaaagtaa	aagggatggg	49560
tggaatggaga	gaagcccggg	tctgaccacc	caatggccaa	tattttggcc	acaagcgact	49620
accagagaca	tggaaaaatg	gtttctacat	gtgggacaac	agatggtaga	ggacctagag	49680
aattgagaga	ggggcaatga	tgggctccac	tccgcagatg	ccttggcttt	cttcctggat	49740
acccttcctg	cactgaatag	caaggagatg	gagcccaagc	agactgtagc	catcttgctg	49800
aatggaggag	agggattgga	gtttgggatg	actgtggtag	ctgaaatfff	tctaggtctg	49860
ctagaaataa	gaactggttt	gtgtggagga	aaagagctct	acaaatacgc	atagaagtct	49920
cctccagtcg	ttggcctgac	atgacgctgc	ctgtgcacag	gaaatggttc	cacgagaaag	49980
tgtggcaaag	aacatttact	gagaaacagc	aagtacaaga	gcacaggaag	ctcaataaag	50040
aagagagaga	tcacatagca	ctctgggata	ctggagtctt	tcccagctag	accagagagt	50100
cctcacggag	cacattgcca	attcagtgga	gaccccgaaa	cagccgtaat	ttaaaggtag	50160
acttagtata	ttactagaat	aaagtcagct	gcagacaacc	ccttgcacag	ctggaaagca	50220
agtgtccaag	catcaaatcg	gtttccaatc	aatgaagtgc	ctgtgagagg	aaatctcaac	50280
tctctttaga	agtaaacaac	aaagtcgatt	gcctcagcta	tgcggtatcc	gcagagtgag	50340
tcctaaatff	aaaatctgac	tacatgtaga	aaagcgtttc	gtgtgaccca	tgaccaggaa	50400
ataaatcggg	taatacaaac	aggctcagga	atgagagaaa	tgattagaat	tgcgtagaaa	50460
tttgacatat	cagtatgata	actgatttca	aatattfata	aaaacaacat	gcaagaaagc	50520
agatatcata	tcaagagaaa	ttaacagtac	agaatagcca	aattaaatta	aagaggtagt	50580
ataaaaaaag	tatgtcttaa	ttgaaaaaaa	ttactgtatg	gccggctgat	caatttagac	50640
gtttcagagg	aaaacattac	ccaacacaca	attctagaga	acctacagaa	tgagctacac	50700
acacacacac	acacacacac	acaaactgaa	aacacaccca	tactcacaca	cacgcagaaa	50760
ctcacaagtt	ctaacacaca	cagacacgcg	cacccttgaa	gaaacagtga	aatataaaat	50820
taagcgagcc	tcacagacat	gtaggaaaat	atgaaaagat	ttcctgcatg	tgggaagcaa	50880
gtcacagtaa	agagcaaggg	agtttggaat	agaaacaaat	accggaatca	aggatggctg	50940
ataacttttc	aattacgaag	aacattaaaa	aaaatcacag	aatcgtgaaa	ctcaagggat	51000
cacatagggg	atttcggaaa	aaaaacccaa	cctgtatgat	gtacttttgt	acatcacagt	51060
tcgaaggtaa	caaggcaaag	atataataag	aagaaacctg	tcacagagaaa	ctggaggaaa	51120
aagagctgtg	tcttcctaca	agtacactga	tacaaattgc	caatgtgttc	acctcagaaa	51180

BI OL0250W0SEQ_ST25. txt

cactggaagc	cagataccag	ggaatattgt	taaaatgata	atcaggaaca	aaaagagatc	51240
aaccgggaat	gctgaatcca	gcaataaaat	gccttgaagg	tcatccatgt	cggataaatg	51300
catattgtgc	actgccccaa	agaaagaaac	cggaaactgt	aagaattgga	aatcagcagg	51360
cttatgtaac	aagagagggtg	acccgaagga	attaggtaga	agaagaattg	aacaagaaag	51420
gaactttctg	cagcccacgt	aatgaagaat	ccagcaattg	gcaaattgtag	atagatgtaa	51480
atgcaaaata	ttttcttgat	caaatttcta	tatctttgta	aatgagagtt	gactacttga	51540
aacaaaatga	tagcaagata	tttaacttca	gcatatgtag	aggtaagaat	ttgaaatggt	51600
agcataaatc	acgaagggat	taattcgaag	tgtaccgttg	taagtttctt	tacctcatgc	51660
acgatggtgt	gtcatattaa	taaaagggta	ctgtgcgggt	tcgaagggat	attgcaaatc	51720
ctagagcaat	cacaaagggt	tgaactctga	ggtttttgggt	ataataagaa	tagtccatgc	51780
attcaaaaga	gggaagccaa	ggaagaacta	gaagtctttc	aagagctcag	gctcttatac	51840
atccagttgc	tattgaacc	agcttcctgg	aatggagggt	ctggggttga	gactaggcca	51900
caagtctaga	gtctctagag	agacagtgtt	ggaaccccat	ggcccataat	acatttccca	51960
ttttctcagg	cagccagagg	tcatgaatgt	gaggatactg	ggaggttgga	gcaacgttct	52020
tgggaggcat	aaggaagagc	gaatgcttca	agatccccgc	agcccaaact	actcgcctgc	52080
tttgccccct	aatgcatttt	tctctgctgc	tccgtagctg	tccgacctct	tcagatctct	52140
tagtccaccc	tgccgtcttc	ctttatgcca	tgggtccac	tgttctttca	actcatcccc	52200
ctttccctca	gtcccggagt	agctgcggcc	agcagagggt	agactgagag	caggagagaa	52260
ggacctgcct	aggaaccctt	tctagagata	ctgcatactg	cctgggagca	agttttccag	52320
ggcagctttg	agaagtcttg	gagaaacaaa	cctactaaac	ctgacagaca	gtaatactat	52380
ttgcacaatg	cttttctgtg	ggaaaggtag	agccttttca	ctacgtattg	agtacataga	52440
gtgtgagggt	tgacctggaa	cggctatcct	cctggatgac	gtgcgttttc	tgaagaacta	52500
catgttcgtt	gcaactccca	cattagaata	tgaagtccta	ccgagagaga	tacggagact	52560
agacagatac	agatgcattt	gcatgtgaat	acacaatccc	acaatacaga	cgtcaaaacc	52620
cataccagtt	attccagaga	gatggattgg	gcagaaggca	gaaggagaat	actctgatcg	52680
tttttcggcc	acgtgtgtgt	gttatctcag	tgtttctaag	aagcgtttgc	tacttttagat	52740
tttttattta	aaaaaaatag	taataatcta	ttaagtatga	gagatgtgca	gagaggatta	52800
gtgatcgaga	gccatttttg	ctggtggcaa	tcatatggta	cttttaatgg	gaatattaga	52860
aaggcaccgg	taatgacctt	gttgacgcac	aaaggagaga	gtgtgggggtg	cccctgcatg	52920
ttgtcccacc	tcttgtgacg	tgtatcgttt	tggaaatttc	agtggcttga	tcatgaacta	52980
ctgcaggaat	ccagatgctg	tggcagctcc	ttattgttat	acgagggatc	ccggtgtcag	53040
gtgggagtac	tgcaacctga	cgcaatgctc	agacgcagaa	gggactgccg	tcgcgcctcc	53100
gactgttacc	ccggttccaa	gcctagaggc	tccttccgaa	caaggtaagg	agtctgtggc	53160
cagacatcta	cacgcttcga	tgctgggatg	aaaagccatg	gaaattccca	ctgatgcagc	53220

BI OL0250W0SEQ_ST25. txt

cgcccttcaat	ggtaaacgga	tgctcgagtg	ttgcctgagt	tctacatgt	aggaggaagc	53280
ctccgtgcac	tctctggggg	agccagcgga	gtgatttctg	gtgcaacgtg	gttgggcttt	53340
gtcttttagga	tgggcacaaa	ccctccaggg	ggatcgactt	caaaattcac	cttggtgtaa	53400
aacgggctac	ctcagtgtcc	cagccaaaat	ttttattgta	acatgctgtc	aggtgtgtca	53460
ctctttccaa	gccagtaagc	ttttccgggg	atttcttcaa	gtagccagca	ttcagagcaa	53520
tcttcagcat	tgagattct	gagaaatgtg	gctctggagc	ctgtcacct	cgagaaacct	53580
aagagggctg	cattgattcc	atgtggccct	gggtctatgg	agcagtacat	gagctcccag	53640
tgctctaagg	ctcttcagcc	ctaggctttg	aaggagtgta	tttctcagta	ttcttaaacc	53700
tctttctgat	gacacttgta	cctgtgaggg	gtctagagag	aaagagtagt	agactcctac	53760
tttactacaa	ttcaggatgc	agggcatgag	aggattccct	ctctcctcca	aggaagaag	53820
cttttgcggt	gcacacatcc	ctgagaagca	aagtgtcttt	gtcttcagtc	agatacatag	53880
gaccgttttc	tgcccatgg	cccgaagcc	aaaggccttg	gctttcatga	tcaacggtct	53940
agggaaacat	gcaaaatttc	catgtctgtc	ccaaactctg	ccccgacag	ccaattacca	54000
cctgcagccc	gcattgccaa	atgcggtgcc	gtttgcatga	agattcagta	gagtttccta	54060
gaaaggtgct	acctcgtgag	ctcactttcc	aatgaggaat	ctgatctggt	gtgtttctct	54120
aaggtgtcag	gtgaaatatt	tccaagaact	tactacagtt	ctagaatggg	aggaatctgt	54180
tgctttgggtg	tttgtttggt	ggtcggtttt	ctcacatcca	tctgcctatg	gataaggaaa	54240
agagaacggt	cgtaattctc	atagactcct	ttctggttgt	gtcacaaatg	gcttcacatg	54300
tttctctatg	ctcagagata	ctcagcttga	tttcccggtg	tttcatttca	gcaccgactg	54360
agcaaaggcc	tggggtgcag	gagtgtctacc	atggtaatgg	acagagttaa	cgaggcacat	54420
actccaccac	tgtcacagga	agaacctgcc	aagcttggtc	atctatgaca	ccacactcgc	54480
atagtcggac	cccagaatac	tacccaaatg	cgtatgtctt	tgttctttac	cataagagaa	54540
gaaagggcca	agtgaagttt	ctgttacaag	agatgtgtct	caagctgagt	tctccgaact	54600
caacttgtga	cagatgcaga	tggcgtagca	aaatgtctca	ggatgattgc	cttggagcta	54660
agggctctgag	agaaggga	tgtaagctc	cctctccttc	ctcctagttc	tattgagcag	54720
aagggaatc	tggaggtgag	gagatcacat	tatgaagaaa	gtcagaatga	caaaggacca	54780
gacacttaga	ttacccttcc	acaacaccaa	ctaaacgtca	atggagactt	tccagttgga	54840
attccgttat	tctggcttcc	acttctgaa	gggaagggtg	cgtttgccct	ttctctctgg	54900
gttcaagagg	aaagaatagg	tgcttattta	tggacaggtg	aattgatctg	tttctatatc	54960
tacgtatatt	ccgattgtca	gaaaaacact	cgttcctaag	taccagtggc	ctgaagggat	55020
acaggttccc	agcaagagaa	gatccaagga	aggaaggcag	atgagagtca	gcacagagag	55080
ggatgctgaa	aagtaaaagg	gatgggtgga	tggagagaag	cccgggtctg	accaccaat	55140
ggccaatatt	ttggccacaa	gcgactacca	gagacatgga	aaaatggttt	ctacatgtgg	55200
gacaacagat	ggtagaggac	ctagagaatt	gagagagggg	caatgatggg	ctccactccg	55260

BI OL0250W0SEQ_ST25. txt

cagatgcctt	ggctttcttc	ctggataccc	ttcctgcact	gaatagcaag	gagatggagc	55320
ccaagcagac	tgtagccatc	ttgctgaatg	gaggagaggg	attggagttt	gggatgactg	55380
tggtagctga	aatttttcta	ggtctgctag	aaataagaac	tggttttgtgt	ggaggaaaag	55440
agctctacaa	atacgcatag	aagtctcctc	cagtcgttgg	cctgacatga	cgctgcctgt	55500
gcacaggaaa	tggttccacg	agaaagtgtg	gcaaagaaca	tttactgaga	aacagcaagt	55560
acaagagcac	aggaagctca	ataaagaaga	gagagatcac	atagcactct	gggatactgg	55620
agttcttccc	agctagacca	gagagtcctc	acggagcaca	ttgccaattc	agtggagacc	55680
ccagaacagc	cgtaatttaa	aggtacactt	agaatattac	tagaataaag	tcagctgcag	55740
acaaccctt	gcacagctgg	aaagcaagtg	tccaagcatc	aatcggttt	ccaatcaatg	55800
aagtcctgt	gagaggaaat	ctcaactctc	tttagaagta	aacaacaaag	tcgattgcct	55860
cagctatgcg	gtatccgcag	agtgagtcct	aaatttaaaa	tctgactaca	tgtagaaaag	55920
cgtttcgtgt	gacccatgac	caggaaataa	atcgggtaat	acaaacaggc	tcaggaatga	55980
gagaaatgat	tagaattgcg	tgaaaatttg	acatatcagt	atgataactg	atttcaaata	56040
tttaaaaaaa	caacatgcaa	gaaagcagat	atcatatcaa	gagaaattaa	cagtacagaa	56100
tagccaaatt	aaattaaaga	gctagtataa	aaaaagtatg	tcttaattga	aaaaaattac	56160
tgtatggccg	gctgatcaaa	ttagacgttt	cagaggaaaa	cattacccaa	cacacaattt	56220
tagagaacct	acagaatgag	ctacacacac	acacacacac	acacacacac	acacaaactg	56280
aaaacacacc	catactcaca	cacacgcaga	aactcacaag	ttctaacaca	cacagacacg	56340
cgcaccctg	aagaaacagt	gaaatataaa	attaagcgag	cctcacagac	atgtaggaaa	56400
atatgaaaag	atttcctgca	tgtgggaagc	aagtcacagt	aaagagcaag	ggagttttata	56460
atagaaacaa	ataccagaat	caaggatggc	tgataacttt	tcaattacga	agaacattaa	56520
aaaaaatcac	agaatcgtga	aactcaaggg	atcatatagg	gaatttcgga	aaaaaaaccc	56580
aacctgtatg	atgtactttt	gtacatcaca	gttcgaaggt	aacaaggcaa	agatgtaata	56640
agaagaaacc	tgtcacgaga	aactggagga	aaaagagctg	tgtcttccta	caagtacact	56700
gatacaaatt	gccaatgtgt	tcacctcaga	aacactggaa	gccagatacc	agggaatatt	56760
gttaaaatga	taatcaggaa	caaaaagaga	tcaaccggga	atgctgaatc	cagcaataaa	56820
atgccttgaa	ggcatccat	gtcggataaa	tgcatattgt	gcactgcccc	aaagaaagaa	56880
accggaaact	gtaagaattg	gaaatcagca	ggcttatgta	acaagagagg	tgacctgaag	56940
gaattaggta	gaagaagaat	tgaacaagaa	aggaactttc	tgacagccac	gtaatgaaga	57000
atccagcaat	tggaatgt	agatagatgt	aaatgcaaaa	tattttcttg	atcaaatttc	57060
tatatctttg	taaatgagag	ttgactactt	gaaacaaaat	gatagcaaga	tatttaactt	57120
cagcatatgt	agaggtaaga	atttgaaatg	gtagcataaa	tcacgaaggg	attaattcga	57180
agtgtaccgt	tgtaatgttc	tttacctcat	gcacgatggg	gtgtcatatt	aataaaaggg	57240
tactgtgcgg	gttcgaaggg	atattgcaaa	tcctagagca	atcacaaggg	tttgaactct	57300

BI OL0250W0SEQ_ST25. txt

gagggtttttg	gtataataag	aatagtccat	gcattcaaaa	gaggggaagcc	aaggaagaac	57360
tagaagtctt	tcaagagctc	aggctcttat	acatccagtt	gctcattgaa	ccagcttcct	57420
ggaatggagg	gtctgggggt	gagactaggc	cacaagtcta	gagtctctag	agagacagtg	57480
ttggaacccc	atggcccata	atacatttcc	cattttctca	ggcagccaga	ggatcatgaat	57540
gtgaggatac	tgggagggtg	gagcaacggt	cttgggaggc	ataaggaaga	gcgaatgctt	57600
caagatcccc	gcagcccaaa	ctactcgcct	gctttgcccc	ctaatagcatt	tttctctgct	57660
gctccgtagc	tgtccgacct	cttcagatct	cttagtccac	cctgccgtct	tcctttatgc	57720
catgggtccc	actgtttctt	caactcatcc	ccctttccct	cagtcccgga	gtagctgcgg	57780
ccagcagagg	gtagactgag	agcaggagag	aaggacctgc	ctaggaaccc	cttctagaga	57840
tactgcatcc	tgcctgggag	caagttttcc	agggcagctt	tgagaagtct	tggagaaaca	57900
aacctactaa	acctgacaga	cagtaatact	atttgacaaa	tgcttttctg	tgggaaagggt	57960
agagcctttt	cactacgtat	tgagtacata	gagtgtgagg	gttgacctgg	aacggctatc	58020
ctcctggatg	acgtgcgttt	tctgaagaac	tacatgttcg	ttgcaactcc	cacattagaa	58080
tatgaagtcc	taccgagaga	gatacggaga	ctagacagat	acagatgcat	ttgcatgtga	58140
atacacaatc	ccacaatata	gacgtcaaaa	cccataccag	ttattccaga	gagatggatt	58200
gggcagaagg	cagaaggaga	atactctgat	cgtttttcgg	ccacgtgtgt	gtgttatctc	58260
agtgtttcta	agaagcggtt	gctacttttag	attttttatt	taaaaaaaaat	agtaataatc	58320
tattaagtat	gagagatgtg	cagagacgat	tagtgatcga	gagccatttt	tgctggtggc	58380
aatcatatgg	tacttttaat	gggaatatta	gaaaggcacc	ggtaatgacc	ttgttgccgc	58440
acaaaggaga	gagtgtgggg	tgcccctgca	tgttgtccca	cctcttgtga	cgtgtatcgt	58500
tttggaaatt	ccagtggcct	gatcatgaac	tactgcagga	atccagatgc	tgtggcagct	58560
ccttattgtt	atacagagga	tcccgggtgc	aggtgggagt	actgcaacct	gacgcaatgc	58620
tcagacgcag	aagggaactgc	cgtcgcgcct	ccgactgtta	ccccgggttc	aagcctagag	58680
gctccttccg	aacaaggtaa	ggagtctgtg	gccagacatc	tacacgcttc	gatgctggga	58740
tgaaaagcca	tggaaattcc	cactgatgca	gccgccttca	atggtaaacg	gatgctcgag	58800
tgttgcctga	gttctacat	gtaggaggaa	gcctccgtgc	actctctggg	ggagccagcg	58860
gagtgatatt	tggtgcaacg	tggttgggct	ttgtcttttag	gatgggcaca	aaccctccag	58920
ggggatcgac	ttcaaaattc	accttgttgt	aaaacgggct	acctcagtgt	cccagccaaa	58980
atttttattg	taacatgctg	tcaggtgtgt	cactctttcc	aagccagtaa	gcttttccgg	59040
ggatttcttc	aagtagccag	cattcagagc	aatcttcagc	attgcagatt	ctgagaaatg	59100
tggctctgga	gcctgtcatc	ctcgagaaac	ctaacagggc	tgcatthaatt	ccatatggtc	59160
ctgggtctat	ggagcagtat	atgagctccc	aatgctctaa	ggctcttcag	tcctaggcct	59220
tgaagggagt	gatttctcag	tggtcttaaa	cctctttctg	atggcacttg	tacctgtgag	59280
gggtctagag	agaaagggtta	gtagacttct	cctttactgc	aattcaggat	gcagggcatg	59340

BI OL0250W0SEQ_ST25. txt

agaagattcc	ctccctcctc	caaggaaga	aggttttggc	gtgcacacat	ccttgagaag	59400
caaagtgtct	ttgccttcag	tcagatatat	aggatcgttt	tctgccccat	ggcctggaag	59460
ccagaggcct	tggctttcat	gatcaacgat	ctagggaaac	atgcaaaatt	tccatgtctt	59520
tccctcctc	tgccctcgac	agccaattac	cacctgcac	ctgcattgcc	aaatgcagtg	59580
ccctttgtat	gaacattcag	tagagtttca	tagaaagggt	ctacttcgtg	agcgcacttt	59640
gcagtgagaa	ggagtctggt	ctgttctggt	tttctaagga	tttcagggtga	aatatttcct	59700
agaacttact	acagttctag	attggttagga	atctgtaggt	ttgctgtatg	ttttttgggt	59760
ggttttctcc	catccatctg	cctacaggta	agggaagat	aacgttcata	attctcatag	59820
actcctttct	ggttgtgtca	taaatggctt	cacatatttc	gttattctca	gagatactca	59880
gtttatttct	tgtgttttca	tttcagcacc	gactgagcag	aggcctgggg	tgcaggagtg	59940
ctaccacggt	aatggacaga	gttatcgagg	cacatactcc	accactgtca	ctggaagaac	60000
ctgccaagct	tggatcatcta	tgacaccaca	ctcgcatagt	cggaccccag	aatactaccc	60060
aaatgcgtat	gtctttgttc	tttaccataa	gagaataaag	ggccaactga	agtttctgtg	60120
acaagagaca	tgcttcaagc	tgagtctctc	gaactcaact	tgtgtcagat	tcagatgggtg	60180
tagcaaaatg	tctcaggatg	atttccttgg	agctaagggt	ctgagagaag	agaaatgtta	60240
agctgcctca	ccttcctcct	agttttgtgg	agcagaaggg	aaatgaggag	gcgaggagat	60300
caccttatga	agaaagtcag	aatgacgaac	caccaaacac	ttagattacc	cttgcccaac	60360
accactaag	cgtaaatgaa	gactttccag	ttggaattcc	gttattctga	cttccaattc	60420
ctgaagggaa	gatttgtgtt	gccttttctg	tctgggctca	tgaggaaagt	ttatgtgctt	60480
acttatggac	agggtgaattg	atctgtttct	atttctacct	gtattccaat	agggagaaaa	60540
tctcttggtc	ctaagtacca	gtggcctgaa	aggatagagg	ttcccagcaa	gagaagatcc	60600
aaggaaggaa	ggcagatgag	agtcagcaca	gagagggatg	ctgaaaagta	aaagggatgg	60660
gtagatggat	agaagccctg	gtctgaccac	cccatggcca	atcatttggc	cataatcaac	60720
aaccaaagac	atggaaaaat	ggtttctaca	tgtgggacaa	cagatggtag	aggacctaga	60780
gaattgagag	agggccaatg	atgagctcaa	ctccatagat	gccttggctt	tcttcctgga	60840
tacccttcct	gcactgaata	gcaaggagat	ggagctcaag	cagcctgtag	ccatctagct	60900
gagcagagga	gagggattgg	agtttgggat	gactctggta	ttttctaggt	ccgctacaaa	60960
taagaactgg	tttgtggagg	aaaggagctc	tacaaatacg	catagaagtc	tcctccagta	61020
gttggcctca	catgacactg	catgtgcaca	gaaaatgggt	ctacagaaag	tgtggcaaag	61080
aacatttact	gagaaacagc	aactacaaga	gaacagcaag	ctcaattaag	aagatagaga	61140
tcacatagca	ctctgtgtta	ttggagtctt	taccagctag	atgagagagt	gctcacggaa	61200
cacattgcca	attcagtgga	gacccagaa	cagccataat	ttcaaagtac	aattagtata	61260
ttactagaat	aaaggcagct	gcagacaacc	ccttgcacag	ctgaaaagca	agtgtccaag	61320
catcaaatgg	gtttccaatc	aatgaagtgc	ctgtgagagg	aaatctcaac	tctcttcaga	61380

BI OL0250W0SEQ_ST25. txt

agtaaacaac	aaagtcaatt	gcctcagcta	tgcggtatcc	ccagagtgag	tcctaaatta	61440
aaaatttgac	tacgtgtaga	aaagaatttc	gtgtgatcca	tgaccagaaa	ataaatcagg	61500
caatacaaac	aggctcagaa	atgacatcga	taattagaat	tgcatgaaaa	tttgacatat	61560
cagtatgata	actgatttca	gatattttaa	aaaagtgcaa	caaagcaggt	atcatatcaa	61620
gacaaattaa	tagtatagaa	tagccaaatc	aaattaaaga	actattatac	aaaaagtatg	61680
tcttaaatga	agaaattact	gtatgtccgc	ctgaaaaatt	tagatgtttc	agaagaaaaa	61740
attaaccaa	aacaattctg	cagaacctac	agaatgagcc	acacacacac	acattcaaaa	61800
cacacccata	cacacacaca	tgcaaaaact	cacaagttct	aacacacaca	caaacacaca	61860
cacacatgca	catccctaaa	gaaataggga	aatataaaat	taaccgaccc	tcagagacat	61920
gcaggaaaat	ataagaagat	ttcctgcatg	tgggaagcaa	gtcacagtaa	agagcaaggg	61980
agtttgagat	agatacaaat	accggaatca	cggatggctg	ataacttttc	aattatgaag	62040
aacgttagaa	aaatcacaga	ttcatgaaac	taaagggatc	aaataggaaa	tttcgagaaa	62100
aaaaactaca	tgatgcactt	ctctacatca	cagttcaaag	gtaacaaggc	aaggatataa	62160
gaagaagaaa	catctcacga	gaaactggag	aaaaaagagc	tgtgtcttcc	tagagtacag	62220
tgatacaaat	tgctaatgcg	ttcacctcag	aaacactgga	agccagatac	cagggaatat	62280
tattaaaatg	ataatgagga	acaagaagag	atcaaccgag	aatgctgaat	ccagcaataa	62340
aatgccttga	agatcatcca	tgttggataa	atgcatattg	tgactgccc	aaaacaaaga	62400
aactggaaag	tgtaagactt	tggaatcagc	aggcttatgt	agcaacagag	gtgacccgaa	62460
agaattaggt	ataagaagaa	tagaagaatt	gcatgaaaat	ttgacatatg	actaagataa	62520
ctatttcaaa	tatttaaaaa	aagatgaata	tgtaataaaa	cagataaaaat	atcaaaaagaa	62580
agtaacagta	ttgactagcc	aatcaaatt	aaagacttag	tgtaaaaagc	tatgtcttaa	62640
aagaaaaaat	tactggatgg	ctgcctgatc	aatttagaca	tttctgaata	ggaaactaac	62700
caaaaatcaa	ttctacagaa	ccaactacac	acatatatac	acatacaaca	cacccataca	62760
caccacgca	aaaactcaca	agttcacaca	cacacacaca	cacacacaac	cctcaagaaa	62820
tagtgaaata	gaaaaccaac	cgaacctcac	agacatgttg	caaaatagga	aaagatttcc	62880
tgcatatggg	aagcaagtca	cagaaaagag	aacgggagat	tggaaacaga	aacaaatacc	62940
ggaatcaagg	atggccgaaa	acttttcatt	gatcaagaat	attaacaaaa	tcgcaaaaac	63000
acgaaattca	atgcatcaaa	taggcgtttc	gaaaaaaaga	aaaaatctgg	tatgatgcac	63060
ttttgtactt	cacattttca	cggtaagaag	acaaagatat	aataacaaga	aacttcttat	63120
gagaaactgg	ggaaaaacaa	gctgtttctt	gctagaagaa	cagtgataca	aattgctaata	63180
gcattctcgt	caaaaacact	ggaagccaga	taccgggaat	gttattaatg	tggtaaacag	63240
gaacaagaag	agatcaacca	agaatgctaa	atccagcaat	aaaatgcctt	gaagatcatc	63300
catgctgcat	aatgtatgt	tgtgcactgc	cccaaacaaa	gaaaccggaa	actgtaagaa	63360
tttggaatca	gcaggctgat	gtaacaagag	aggtgaccca	aaggaattag	gtagaagaag	63420

BI OL0250W0SEQ_ST25. txt

aatagtacaa	gaaggggaact	ttctgcagcc	catgtaatga	agaacccagc	aattggcaaa	63480
tgtagatgta	aatgcaaaat	atcttcttga	ccaaatttct	atatatTTTT	aaatgagcgt	63540
tgactactgg	aaacaaaatg	atagcaatat	atttaatttt	agcatatgta	gaggtaagaa	63600
tttgaacaag	tagcgtaaat	catgtaggga	ataattagaa	gtgtaccatt	gtaagtttct	63660
tacctcatgc	acaatgggat	gtaatatata	taaaatgtta	ctgtgtgggt	tcaaggagat	63720
attgcaaadc	ctagagcaat	cacaaagttt	tgaactctga	ggtatatatt	ataataagaa	63780
tattccatgt	attcaaaaaga	gagaagccaa	ggaagaaaga	aatttgtcac	gagtttgggc	63840
tcttagtaca	tcctgtagct	cattgaacca	gcttcctgga	atggaggggtc	tgggattgac	63900
actaggccac	atgtatagag	tctctagaga	gacagtgttt	catcccatg	gcccgtaata	63960
catttcccat	tttctcaggc	agccacaggt	catgaatgtg	aggatagaga	gaggttggag	64020
caacgttctt	gggaggcata	aggaagagca	aatgcttcaa	gatccccgca	gccc aaactc	64080
ctacctgtt	tgccccctaa	tgcaagtgtc	ctccgtagct	gtccgacctc	ttcagatctc	64140
ttagtctacc	ctgccatctt	cctttatgcc	atgggtccca	ctgttctttc	aactcatccc	64200
cctttccctc	agtgcagagt	agctgcggcc	agcagagggg	agactgagag	caggagagaa	64260
ggtcctgccc	aggaacccat	tctagagatg	ctgcattctg	cctgggagca	agttttccag	64320
ggcagctttg	agaagtcttg	cagaaacaaa	cctatttgac	ccacatgata	tgggaatgac	64380
agaaagtaat	acaatttgca	cagtgtcttt	ccatgggaaa	agtagagcct	tttcgcgagg	64440
ttttgagtac	atagagagtg	aaggttgacc	tggaaagggt	atcctcctgg	atcccatggt	64500
ttttctgaag	aactacctgt	tagttgcaac	ttgcacatta	gaatatgaag	tcctaccgag	64560
agagatacgg	agaactagat	aaatacagat	acttttgtat	gtgaataaac	gattccacaa	64620
tacacacatc	aaaatccata	ccagttattc	cagagagatg	gattgggcag	aaggcagaag	64680
gagaatactc	tgatcgtttt	ttgccacagt	gtatgtatta	tctcagtgtt	tctaagaagc	64740
gtttgctact	ttagatTTTT	ttttataata	ataatctttt	aagtatgaga	aatgtgcaga	64800
caggattagt	gattgagagc	catttgtgct	tgtggcaatc	atatggtact	tttatgggaa	64860
tattagaaaag	gcactggtaa	tgaccttggt	gcagcacaaa	ggagaggggtg	tggggtgccc	64920
ctgcatattg	tcccacctct	tgtgacgtgt	atcgtttttg	aatttccagt	ggcttgatca	64980
tgaactactg	caggaatcca	gacctgtgtg	cagccccctta	ttgttatacg	agggatccca	65040
gtgtcagggtg	ggagtactgc	aacctgacac	aatgctcaga	cgcagaaggg	actgccgtcg	65100
cgcctccaac	tattacccccg	attccaagcc	tagaggctcc	ttctgaacaa	ggtaaggagc	65160
ctgtggccag	aaacctacac	gtttcgatgc	tgggatgaaa	agccatggaa	attcccactg	65220
atgcagcagc	ctccaatggg	aaacggatgc	tcgagtgttg	actgagttct	gtcatgtagg	65280
aggaagcctc	cgtgcactct	ctgggggagc	cagcggattg	atttctggta	caacgttggg	65340
tgggctgtgt	ctttagaatt	ggcacaacc	ctccaggggtg	atcgacttca	caactcacct	65400
cgttgaaaaa	tgggctatct	cagtgtctta	gccaaaattt	ttattgtaac	atgctgtcag	65460

BI OL0250W0SEQ_ST25. txt

atgtgtgact	ctttccaagc	cagtaagctt	ttcctgggac	ttcttcaatt	agccagcatt	65520
cagtgcaatc	ttcagcattg	cagattcaga	gaaatgtggc	tctggagcct	gtcacccttg	65580
agaaacaggg	ctaacagggg	tgcatthaatt	ccaaatcacc	ctgggttctat	ggagcagtac	65640
atgaactccc	aatgatctat	gtttcaggac	ttcctcagtc	ataggtgggc	tctgcagccc	65700
taggttttta	agtgagtac	tgccccgtgt	tctggtggca	gttgtacctg	tgagcgggtct	65760
ggatagaaaag	agtcggagac	ttctgtatta	ttgcaactca	ggatgtgggt	catgagagga	65820
tttcatctct	cctgcagggg	agtaagctgt	tcgcctccac	ccatccctga	taactgaagt	65880
gtctttgtct	gcagtcctag	acgaaggact	gttgtctctc	ccatggccca	gaagctgaag	65940
accttgcctt	ttgttatgaa	acgttcattg	ttttcatgtc	tgtccgtttc	tctgccccta	66000
acaccaatc	accatgtatg	gcctgtaccc	ccaaatgcat	cgtgctttgc	tgtttgctgc	66060
cccatagtcc	tcatgaacat	tcagtagaaa	ttcccataaa	tgtgcttgca	cgtgagcaca	66120
gtttccattg	agaagccctc	tcatttgtcc	tttttttcta	agcttttatg	tgaaatatatt	66180
ctaagaactt	actacagttc	taaagtgtta	ggaatttggt	tctttggtgt	ttttgtttgt	66240
tggttggttg	ttgcttttct	caagtccatc	tgccctacaa	taaagaaaca	agaatgttac	66300
ttgtcatatt	ctcctgaggt	cataattctc	agagactttt	ttctggtttg	tgccataagt	66360
ggcttcacat	gtttgtctct	tcttggaac	actcagtttg	atttcttttc	ttttcatttc	66420
agcaccaact	gagcaaaggc	ctgggggtgca	ggagtgtctac	cacggaaatg	gacagagtta	66480
tcaaggcaca	tacttcatta	ctgtcacagg	aagaacctgc	caagcttggt	catctatgac	66540
accacactcg	catagtcgga	ccccagcata	ctacccaaat	gcgtatgtct	attttcttta	66600
ccataagtga	aggaaggggtc	agtggaaatt	tctgttagta	gagtcatgct	tcaagctgag	66660
tgttcaggac	tcaagttgtc	tcagatgaac	agtgcatagc	aaaatgtctc	aggaacattg	66720
tctttgagca	aagagtctaa	gagaagacaa	atgttaattct	ggctctcctt	cctcctagtt	66780
taatggagca	gaaaggtatc	tggaggcaag	gatatcacat	taagaaacaa	gtcaagatga	66840
caaatgatga	aactcttaga	gtacccttcc	acaacaccca	ctaaggttca	atgcagcctt	66900
ttctccttgg	aattctatta	aactaaactc	caattcctga	agtgaagggt	ctgttggggg	66960
tttctgtttt	ggcttacaag	gaaagtatat	atgtatatct	atggagaggc	aaatctatct	67020
ctttctatat	ctacgtctat	tccaatatgt	agaaacacag	tcggttctga	ccaccagtgg	67080
tctgaaggga	tactggttgt	tagagaataa	aaatggcagg	aaggcagatg	agagtcagca	67140
aagagagaga	tcctgtaaag	taaaaggggtg	gatagatgga	cagaagccca	ggtctgacca	67200
gccccatggc	aggctttagg	ccataagtga	caccaaagac	atggaaaaat	ggtttctaca	67260
tgttggacaa	cagacagtag	tggacaaaaa	gaatagtgac	agggggaaca	atgagatcaa	67320
ctccatagat	accttggctt	tcttcctgga	ggcccttctt	gcactgaaga	gcaaggtgat	67380
ggagcccaga	tggactgtag	ccatcttcct	gaatgcagga	gagagattgg	aatttgggac	67440
tactgtggta	gctaggattt	tataggcctg	ctgagaatga	gaatggattt	gtggatgaaa	67500

BI OL0250W0SEQ_ST25. txt

ggagctccag	gggcacgcat	agtagtctcc	tcgaatcttt	ggctaaacat	gacgttgcatt	67560
gtgcccagaa	aaagggtcca	caagaaagta	gagaaaagaa	tatatcctga	ggaatagcaa	67620
ctgcgattga	acagtgcgct	caataaagag	gacagagccc	tcatagcatt	ctgggatact	67680
ggagtcttga	ccagctggag	gagagacctc	actgaacctc	ttgggaatac	agtagagact	67740
ccagaaaagt	catactttag	gagtagaatt	agtaaatttc	tagaaaaaaa	ggcagctcta	67800
gacaaaccct	ggcaaaactg	aaaagcaagt	ctccaagcat	taaaatcatt	tccaagtcaa	67860
ttaactgcct	gggagaggaa	aaccctcttt	agaggtaaac	aacaaagtca	agtggctcag	67920
ctatgtggtg	ttcacagtgt	gagttctaaa	tttaaaactt	gactacacat	agagaagctt	67980
ttagtatgaa	ccatgaccag	gtgaaaaatc	agtcaataca	aatagacctt	gaaatgacag	68040
aaatgattag	aatggcaaaa	aatttgacat	atcaatatgt	caactgagtt	ttaggtttta	68100
agaaaacatg	aatacggagt	gaagcagata	ccatatcaag	agacagtaac	agtatagaag	68160
agccaaatta	aattaaagaa	ctagtataag	aaggatatgt	ttaaatgaaa	aaattactgg	68220
atgtattccc	aatggagtga	gatgtttcag	aagtaaaaac	taactgaaaa	acaattttat	68280
accacctaca	gaaccagcta	cacatacaca	aatgacacac	acataacac	acatactcac	68340
acatgcacag	gcttagaaac	atgcacgcac	acacacacac	acacacacac	acacctccac	68400
aaatactaaa	aatgaaatc	caatgatcct	cacagacagg	cgggaaaata	taaaaagatt	68460
tcctgcatgt	gggtaggaag	tcacagaagg	agaggaagga	gagattgcta	caggaacaaa	68520
tactggaagc	aaggatagct	aaaaactttt	caaataagaa	gaatattaaa	aaccacagat	68580
tcaagaagct	gaatgaatca	gacagggaat	ttccaaaaaa	aaaaaaaaaa	aaactgtatg	68640
attcactttt	gtacatcacc	gttcaacagt	cagaaggcaa	agatataata	acaagaaaca	68700
tctcatgaga	aactggagga	aaaagagctg	tgtcttgcta	gaagaacagt	gatacaaatt	68760
gctaattgcat	tctcatcaga	aacactggaa	cccagttaac	aggggatatc	attaaaatga	68820
taaactagaa	aaaaaagaga	tcaaatgaga	atgctacatc	cagcaataaa	atgccttgaa	68880
gatcatccat	gttgataaaa	tgcatattgt	gcactgcccc	aaataaataa	acaaaaaact	68940
aataatttgg	aatcagcagg	cttgtgtaac	aagagatggt	gcccaaagaa	aattagctag	69000
aagaagaata	gttcaagagg	agaactttct	gcagcccacg	taatgaagaa	cccagcaaat	69060
ggcaaatgta	gatgtaaatg	caaaatattt	tcttgatcaa	atttctatat	ctttttaaat	69120
gagagttgac	tacttgaagc	aaaatgatag	caatatattt	aacttttagca	tatgtagagg	69180
taaaaatttg	aacatataga	ctaaatcatg	tggggaataa	ttggaagtgt	accattgtaa	69240
gtttcttacc	ttatccacga	tggtatgtaa	tattaatgaa	aggttgaatt	tgtgggtcca	69300
aagggatatt	gtaaatccta	aagcaatcat	aaaattttga	attctgaggg	atattatata	69360
ataagaattt	tccatgtatc	caaaagaggg	aagccaagga	agaaaaagaa	gtctttcaag	69420
tactcaagct	ctgagcacat	ccagttgctc	attgaaccag	cttcctggaa	tggagggctc	69480
gggcttgaga	ctaggtcaca	tgtgtagagt	ctctagagag	acagtgttgg	atcccatgg	69540

BI OL0250W0SEQ_ST25. txt

cccataatac	atttcccgtt	ttcccaggca	gccacaggtc	acgaatggga	ggattctgag	69600
aggttggagc	aatgttctta	ggaggcataa	ggaggagtga	atgctctgag	atttccccag	69660
cctgagggtcc	tccatagctg	cccgaacctct	tcagacctca	tagtctgccc	agctgtctcc	69720
ctttatgcca	tgagtgccac	tgttctttca	actcatcccc	cattccctca	gtcccggaat	69780
tgctgtggcc	agcagaggat	ggactgagag	caggagagga	agtcctgacc	aggaacccat	69840
cctagagata	ctgcatcctg	cctgaaagct	aggtttccag	ggcagctttg	agaagtcttg	69900
cagaaagaaa	cccacttgac	ccacctgata	cggtatcgac	agacaggaat	actttttgtg	69960
caatggtttt	acatgctgaa	catagagcct	tttggctaca	ttttgagtac	attgaatgag	70020
actgctggcc	tgggaaggat	atcatgctgg	atgccatttt	tttctctgga	gaactatgtg	70080
ttagttccaa	ctcgcacatt	actatatgaa	gtcctacaca	gagagatacg	gagagctaga	70140
cagatagaga	tacttttgta	tgtgcataac	caattccaca	atacacacgt	caaaatccat	70200
accagttatt	ccagagagat	ggattgggca	gaaggcagaa	ggaggatatt	ctgatccctt	70260
tttggccaca	tgtatgtata	atctcagtgt	ttctaggaag	tgtgtgctgc	attagatttt	70320
ttttctttaa	aaaaagtgat	aatatattaa	gtatgagaaa	tgtgcagaga	ggattagaga	70380
ttgagagcca	tttgtcattg	tggcaattgt	atggtatctc	ttttgggaat	atttcaaagg	70440
caccagtaat	gaccttgttg	tagcaaaata	tacagtgttc	ctgcatatgt	acccattttt	70500
tgtgatgtgt	attcttttgg	aatttccagt	ggcttgatca	agaactactg	ccgaaatcca	70560
gatcctgtgg	cagccccttg	gtgttataca	acagatccca	gtgtcagggtg	ggagtactgc	70620
aacctgacac	gatgctcaga	tgcagaatgg	actgccttcg	tccctccgaa	tgttattctg	70680
gctccaagcc	tagaggcttt	ttttgaacaa	ggtaagaagt	tgtgccagac	atttacctgc	70740
ttggatgctg	ggatgaaaag	ccatggatac	cccactgac	gcacaaccct	tcagtgtctac	70800
actggttctc	gtgtgttggg	tctgggtctg	ccatgtggga	ggaagcctta	gcgcactctc	70860
tgggggagcc	agagggtgtga	tttttgggtgc	aacctgtgcg	agctgtgtct	ttaggatggg	70920
cggaaaccat	tctgggtgct	cgacttcacc	actcccctca	ttgtaaaagg	ggctatctca	70980
ttgtcctaga	caaaattctt	attgtaatat	gctgtcagat	gtgtgtgtct	ttccaagcca	71040
gtaaactttt	ccagggattt	cttcaagtag	acagcattca	gtgcaatctt	cagcattgca	71100
gattccgaga	aatgtggctc	tagatcctgt	tatccttgag	aaacctaact	gggttgcatt	71160
aattccatat	ctccctgggt	ctgtggagta	gtacatgagc	tcccgaagct	ctatctctca	71220
ggcttttttc	agtccgaggc	aggttgtgca	gttcttagct	ttgaaggagg	tgattttttc	71280
gtgtgctttt	gcctctttct	gatggaactt	gtacctgcgg	ggggctctgga	gaaaaagagt	71340
agtagacttt	tgctttattg	caatgcatta	tgctgggcac	gagaggattc	cctatcttat	71400
tgtaggatgat	aagcttttgg	cctccactca	tccctgagaa	gtgaagtgtt	gttgccctaca	71460
gttttagctg	caggactgtt	gtctgccccca	tcaccaggag	tttaatgctt	tcttttttga	71520
gcaatcatct	agggacacat	gcaaggtttt	tatatgtcct	tgccctctcc	ccaaaaaacc	71580

BI OL0250W0SEQ_ST25. txt

attttaatgc ttggagactt gcttttcagc ttgccaaat gcatcaccct ttctttctatg	71640
ctgttccatg tcgtcatgaa cactctgtag agattcctag aaatgagctt ccatgttagt	71700
ggagtttccg atgagaagca atctgatatt tcttttccac taagttttac atgaaatatt	71760
tctaagaact tactacagtt ctagaatggg aggcattctt tactttcgtg ttgttttgtg	71820
tgttttctca tgtccatttg cctattaata aagaatagag aatggttgta aatctcagtg	71880
actctttttt ggtttatgtc ataaatggct tcctgtattt ttctgttcta ggaaataata	71940
agcttgatgt cttctgtttt aatttcagca ctgactgagg aaacccccgg ggtacaggac	72000
tgctactacc attatggaca gagttaccga ggcacatact ccaccactgt cacaggaaga	72060
acttgccaag cttgggtcatc tatgacacca caccagcata gtcggacccc agaaaactac	72120
ccaaatgcgt acgtctttgt tctttaccat aagcgaagga agggccaatg gaagtttctg	72180
ttagaagagt catgcttcaa ggtgactgct caggactcaa cttgggtcag atgcagagga	72240
acatttcctg tgagcaaaag ttcttagaga agactttgtt tttttgagac agagtcttgc	72300
ttgtttgcc aggctggagt gcagtggcat gatctcggct cactgcaagc tccgcctccc	72360
gggttcacac cattctcctg cttcagcctc tctagcagct gggactacag gcacccacca	72420
ccacaccgg ctaatTTTTT gtatTTTTtag tagagacagg gtttactgt tctagccagg	72480
atggctttgg tctcctgacc tcgtgatccg cctgcctcag cctcccaaag tgctgggatt	72540
acaggcgtga gccaccgtgc ctggctgaga agacattttt taagctggct ctcttctctc	72600
ctagttttat ggaagcagaa ggatatatgg agttgagaag atcttattaa taaaacagcc	72660
gggatgacaa atgaccaaag agttagagta tccttctaca acatcggctg agggttaata	72720
caaccttttc accttggat tctatcattc taagctctag tccctgaagt gaatgttgtg	72780
ttggcctttt gcatcttggg tcacagggaa ttgatacttg cacatctatg gagaggcaaa	72840
tctttttcta tctacttctt tttcaatggg taaaaacaca cttgggtcctg agcaccagtg	72900
gtctgaagag atacggtctg cccagaggag aagaacaaag gcaggaaagc agatgagagt	72960
cagcaaaggg gcgatgctga aaagtaaaag gggcgggtag atggacagaa gccatgatct	73020
ggccattcta tggccagtct ttcggccata agtgactacc aaagacacgg caaaacggtt	73080
tccacatgtt gaacaacaga tgctagagga ccaagagtat tgcaagaggg agaaaatgag	73140
atcaacccat caatgccttg gctttcttca aggagaccct tcctgcactg aagagcaagg	73200
agatggagcc caagctgact gtagccatgt tgctgaacag aggagagtga ttggactttg	73260
ggattactca ggtagttagg attttctagc catgctaaga gtaagaatgg acttgtggag	73320
gataggagct ccaggcatag aagtctctc aagtgttagt ctaaacataa agcagcactt	73380
gcatagaaga ttttcacaa gaaaatatgg caaaaaaaca ccatatattg aggaacaaca	73440
actacaaggg aacagtgagc ttaataaagg tgacagagct cacatagtgc tctggaatat	73500
tggagttttg accagctaga gagaagagac ctcatgaaa atcttgggca ttcagtagag	73560
acctcagaaa agtcagactt tatgagtaga ctttgtatat tcctagaata aaggcagctc	73620

BI OL0250W0SEQ_ST25. txt

cagaaaaaac	ctagcaaagc	tgaaaagcaa	atctccaagc	attaaaatgg	tgctcctagtc	73680
aattaactgc	cttctagaag	aaaactcaac	actctttaca	ggatgaacaac	aaagttaagt	73740
tgctgagcta	tgcaatatcc	acagtgtgag	tcctaaattt	ataacttttac	tacacataaa	73800
aaagcattta	gtgtgaacca	taaccaggaa	aataatcagt	caataaaaaat	agaaccagga	73860
atgatagaaa	tgattttaa	ggcatgagaa	tttgacatat	tagtatcata	actgcattgc	73920
tggtatttaag	aaaacataaa	catggaacgt	aacagatatc	atatcaaggg	aaagtaaaag	73980
gataaaagag	tcaaatcaaa	ttaaaggact	attaaaaggt	atatctttaa	tgaaaaattc	74040
actggatggt	ctcccaatca	ggttagtgt	ttccagggaa	aaaattaact	gaaaaataat	74100
tcaatagaat	ctacagaaat	agctgcacat	atatacacac	aatggcacac	gtgcacacac	74160
ccacacccac	acaggtgtga	atcctagagc	cacacgagca	ttgaaacata	gagaagtaaa	74220
aattgttcat	tgaggaatat	gtagcaatgc	tcaatgtgtt	ttacccta	aagagctttt	74280
gtgatgtatg	attgaaaaac	tgacacaact	gaagagagaa	atagataagc	ccacactctg	74340
agttagagat	ttccttgatt	ctctcactat	ggttataaat	ctttcccaa	cacaacaggc	74400
tagaacaat	atgcagaaaa	ttagacatag	tatctttgtt	ctcaataaaa	acgtcgacct	74460
atttaacatt	ataccgaact	accgagtaca	cattaaagt	tgcatggagc	attcactgag	74520
gtgtactcta	cacatgacct	tccagcaagt	ctccatagat	ttaaaagaat	taaagtcata	74580
cagagtgtgt	cactttattc	tcccagaata	aagttagata	tgaataatga	gaagtttgcc	74640
agcttctcaa	atatttgga	gtcatacggt	gcatttcaa	atactctttg	ggacaaagaa	74700
aacatcacta	aggaatttag	aaaagttttg	aactgagtaa	gaatataaca	caatttatcc	74760
aaacttagga	gatgcagtga	atgtcttttag	gcttttacat	aatttttagat	gctcttaggg	74820
aaaaacagaa	gcatgtaata	atcaagattt	caaactgcaa	ttctcaaagt	gtagtctaga	74880
gaaacctgag	gacctttgag	taccttcaga	gacagtccat	gaggttaaag	gactttgcta	74940
cgtgaaaagt	aagatgctat	tggccctttt	tactttcatt	ttccaacaag	agaagagggg	75000
agttttccag	cagttacata	atatgtaatg	gcatcatgtc	tctgatggct	aagaaaatgg	75060
gcaattgttg	actttgtgtg	ttaaaaaat	tctcagtgtt	ggtttcttat	actataaata	75120
ttcatcttgt	gttttgaaaa	agaaaagctc	tttggaatcc	cctatgaaca	aagactttga	75180
cagttgttga	tctaagacca	cagcttaa	atctacacaa	gaaaaaaaaa	aaaagcaaat	75240
aagagccaag	gaaagcagat	ggaaggaagt	agtccaaacc	agtgcatttc	agtgaacaag	75300
aaaagagacc	aacaaggagg	taaactcttg	aaacagaaag	ttgattcttt	gaaaagatcc	75360
atatgattga	acacagtctg	gctaaacaaa	tgacagacca	atgaggggtgc	acaaccatca	75420
ccatctggag	taacagagga	gaggtgccat	tactatagca	tcttccagtt	ctgaaagctg	75480
aaaagaagat	tttgagaaca	attgtatgtg	aataaattca	ggaatgttaa	tcatgtgggc	75540
caattcctga	ggaagacaac	aatcagcaa	accagatgct	gaatagttag	tgtagtcctg	75600
tagagagaca	tacagagagg	ctgacagaga	aatatttgta	tgtgcataaa	acaatctaca	75660

BI OL0250W0SEQ_ST25. txt

agacacactt	caaaatcaat	ctcagttaat	ctggaggaac	atatttcaca	gaaggtggaa	75720
ggaggggtatt	ctgatcctct	tgtacattgt	acaacattgt	acaatgtaca	gagtataatt	75780
gtacaagtac	aattgaagtt	gtacaagtac	aagtgcact	tgcacaatgt	acagagtaaa	75840
cattgatgtt	tactctcaat	tttcttatgg	agcacagatg	actttggatg	tgttacaata	75900
tgaatgataa	tttgtctttg	agatgttcgc	agttgttttag	aagttgagga	ccatttgtgc	75960
atattatggg	accttttagtg	aaaatatattc	aaagtctctt	tttacacttt	gttacagcaa	76020
aatgtagagg	gcgctaagtg	cccttgaatc	ttctcccatc	tctggtgacc	tgtgttgttt	76080
tgaaatttgc	agtggcctga	ccaggaacta	ctgcaggaat	ccagatgctg	agattcgccc	76140
ttggtgttac	accatggatc	ccagtgtcag	gtgggagtag	tgcaacctga	cacaatgcct	76200
ggtgacagaa	tcaagtgtcc	ttgcaactct	cacggtggtc	ccagatccaa	gcacagaggc	76260
ttcttctgaa	gaaggtagga	agtctatggc	cagacaacca	caccctagga	cgttgggatg	76320
aaaagagttg	caaaatctta	gtgatataga	agccttccat	gctcacacaa	ttccaagtag	76380
aatgtggact	cagggtcagc	caactgggaag	gaacactcag	cgcttctct	gggagaacca	76440
gagctgtgat	gtttggtacc	ctgtgaaagg	gtggtatcta	taggaagggg	gcagaccctc	76500
tagggcactg	gacttaccac	tcccctgggt	attcaaagga	tcattttagt	gtcttagcca	76560
gaagaatatt	ctaacatttt	gccaaatttg	tgaagattta	ccaagctcat	gataagcctt	76620
tcattggtatt	tcttcaagta	gtcagtgttc	attgcatctt	tggctttgcg	gtttcggagg	76680
aatgcggttt	ttgagtctgt	catccttgag	aaacctaata	tgacttttct	tagttccata	76740
tacttctggg	tccaggtagc	agtacatagc	caacaaatgc	tccatcgttc	tggcctatct	76800
ccatcttaag	ccagtcctgc	acaactaggc	tttgatggga	gggatctctc	agtgttcttg	76860
cccctccttc	tcatggaaca	tatatctgtg	ttggtctctg	agaagaagag	tagtggatat	76920
ctactttgtt	gcaatgcaga	atcctggggc	aaagatacca	gccatccctc	caagggaata	76980
aaattttggc	cagtagccct	ctctgagaga	caatttgtct	ttgcctacga	gtcctagatg	77040
caggaccgct	tcctgcccc	tcttcaagaa	gctgaaggct	ttggctttgg	aggatcagca	77100
gtctagggaa	atgtgtgacg	gtttcatgtc	tgtccccact	gacagtcaat	caccacctac	77160
aacctgcaca	gcctgatgca	tagcagtcta	gtttcctgcc	ttattctcag	gaacacccag	77220
aagatgtcta	tattaaagag	catgcacatg	agtgcaattt	tgactgatag	gcactctgat	77280
ctttcctttg	gtgcctgtgt	tttaaaggaa	atctttctaa	gaactcgtt	aagttctaga	77340
atgctatgaa	tctttgggtt	ttattattgg	tatgtccatc	tgcttgctag	tacagaacag	77400
agcatggtag	tctttctcag	agacaatgat	cctgtttcag	tcacagattt	cttctgatgc	77460
ttctgtgttc	tagaaattac	tcagcttgat	ttctcctctt	tgaatttcag	caccaacgga	77520
gcaaagcccc	ggggtccagg	attgctacca	tggtgatgga	cagagttatc	gaggctcatt	77580
ctctaccact	gtcacaggaa	ggacatgtca	gtcttgggtc	tctatgacac	cacactggca	77640
tcagaggaca	acagaatatt	atccaaatgg	gtacaacctt	gagttttctt	caaagacaga	77700

BI OL0250W0SEQ_ST25. txt

cagcagcccc	cttacatttc	tcttggaagg	gccatgcttc	caactaactt	cttatgacaa	77760
atttatctca	gatctggaat	gttgggtaga	atgtctcagg	cttctttctt	caggcacagt	77820
gtctgaaagg	agagaaatgt	caggccagct	ctcttttctc	atagttgaca	gaagcaggag	77880
gatatttgaa	ggtggtgagt	tctcatgaat	agaaagctca	ggacacatgg	ccacgtgctt	77940
agaaatagca	ccattccaca	atgccacta	aagaccaatg	caatagttca	accagggatt	78000
tctgtcattc	taatctccaa	gtcctgaagt	gaagggttgta	ttagccatgt	tcatcttggg	78060
caacaaataa	aggatatcta	tgttgacatc	cagatcttcc	aatcactttc	tcctctaacc	78120
tgtacctggg	ttctgagaac	aaggatatctg	aagagctatg	tgttgccagc	acatgagggg	78180
caaaagtagg	aaggcagctg	agagtcagga	agtataaaga	ttctgaagag	ttacacatgc	78240
aggaagatgg	acagaaaccc	agttcagacc	acgtcagcgt	ttctgccatg	aaggactatc	78300
aaatacatag	gaaaagtgtt	ttcatagggt	ggacaacaga	catgacaggc	ctgagaaaaat	78360
tcagaaaggg	aatcaaagga	gatcaacctt	atcatgtccc	tggcatcctt	ccttgagacc	78420
cttgaagggc	aagcagatgg	agcccagctg	accacagcag	tcttgcttaa	ctgaggagag	78480
agactggagt	ttgtgatgcc	tcaggcatct	gacgtattct	aggctggcta	agaatgagag	78540
gggattttgtg	gaggaaagga	gctccaagaa	tacacaccga	agtctttctca	aggctttggc	78600
taaatacaaa	gctgcgtatg	cacaaggaga	gttttcacaa	agaaagaaca	ataaagaaaa	78660
gctactgggg	aaagaacaac	tgcaagggaa	cagtgaagctc	aatggagatg	ctagagctca	78720
catagcactg	ggggatattt	gagttctgac	cactcagagg	agagacacct	cactgaacat	78780
cttgggcatt	cagtagaggt	caaagaaagc	cataatttgg	gagtaggatc	ttcggattcc	78840
tagaaataag	gtgactccag	aaacactcca	gcaacccttc	ttccaagcca	gtctaaaagg	78900
atccaaatga	tttccaagta	aattaactgc	cttccagaaa	aaagtaaact	caaccctcct	78960
tagaggtaag	gaacgaatac	aagtttctca	gttatatgac	atccccagag	tgcaacttgc	79020
atttaaaaaat	ttactagaca	caaaagaagt	tttactgtg	atccataact	gggagaaaaa	79080
tcactcaaca	caaataggcc	cagaaataat	agaaattatg	gcattggcaa	gaacatttaa	79140
aatgcacctc	tgagaactgt	gtttcaggaa	aatgtcagca	aaagctgacc	atgagagaaa	79200
tgaatgcata	atatcagaaa	agaaaagaat	tgaagagcca	aatggaaatt	taaaaactga	79260
gaaaagttaa	atctgtaatg	aggaattcac	tggatggcct	tataaccagt	ttagatatta	79320
tggtaggaaa	aggatgaacg	gaaaatgatt	caattaaagc	tagacaaacc	acaagacaga	79380
cagacagaca	caaatacaca	tacacacaat	gactgaacca	attaatcaac	agagcctcaa	79440
ggacatctag	gaaaacatcc	acacatttaa	tatatgtgtt	aggcaagtca	cagaaagaga	79500
ggaaaaagat	aatgtgacag	aagttatact	tgaagccatg	acggctgaca	aatttccaaa	79560
catacagaaa	atgagaaatt	catagtcatg	aagctcaatg	actcaggtat	agatttttaa	79620
agagcaaaac	tctgatttac	tggggtacat	catagttaaa	ttgtctgatt	tcaaagctaa	79680
gaagaaaaaa	aggggggttcc	tatgaacaaa	cattttgaca	gttgatctaa	gaccacagct	79740

BI OL0250W0SEQ_ST25. txt

taaatatcta	ggcaaggaaa	agcaaataag	acacaaggaa	aggggatgga	tggaaatagt	79800
ccaaaccaat	gacattcagt	gaacaagaaa	atagaccaac	aaaggagtaa	atccatgaaa	79860
cagaaagttg	gttctttgaa	aagattcatg	tgattgacca	cagtctggct	gaacagatga	79920
cagaccaagg	agggagtaca	accatcacca	tttgaagtaa	caggggagag	gagccattgc	79980
tataccatac	tccagggtctg	aaagctgaca	agaagatatc	aagaaaaact	gtatgtgaat	80040
aaattcatga	atgtagatca	tgtggatcaa	ttccttaggt	aaacaacaaa	tcagcaaacc	80100
agatactgaa	tagattgggt	actcctatag	aaagacatac	agatagccag	acagagaaac	80160
atttgtacgt	gcataaaaca	atctacaaga	ctcacttcaa	aatctctcag	ttaatccaaa	80220
gtaacatatt	tggcagaagg	tggaaggagg	gtattctgat	cctttcttgt	acacattgat	80280
gttttctctc	ggttttctta	tggagtatag	acgagtttgg	atgtgtttaca	ataagaatga	80340
taatctgtct	ttgaaatgtt	cacagttgtt	tagaagttga	ggacgatttg	tgattgttac	80400
aggaccttta	gtgagaatat	ttcaaagtca	ctttttacca	ctttgtttaca	acaaaatgta	80460
gaggatgtct	ggtgcccttg	tatcttctcc	catctctggg	gaactgtatt	gttttgtaat	80520
ttgcagtggc	ctgaccagga	actactgcag	gaatccagat	gctgagatta	gtccttgggtg	80580
ttataccatg	gatcccaatg	tcagatggga	gtactgcaac	ctgacacaat	gtccagtgcac	80640
agaatcaagt	gtccttgcca	cgtccacggc	tgtttctgaa	caaggtaaga	agtctctggc	80700
cagacaacca	cacccttgga	cgttgggata	aaaagagttg	caaaatctta	gtgatacaga	80760
agccttccat	gctgcacggg	aatctgaatg	tggactcagg	gtcagccaat	gggaaggaag	80820
cctcagcgcc	ttctctgggg	gaaccagggc	tgagatTTTT	ggcaccccgt	gacaggggtg	80880
tgtcttttagg	aagcgtgcag	accttctagg	gcactggatt	taccactccc	ctgggttattc	80940
aatagattat	ttcagtgtcc	tagtgaaaat	ggatattcta	acatcctgcc	aaatttgtga	81000
tgatttacca	agctcatcat	gagcctttcc	tggtatttct	tcaagtagac	agtactcatt	81060
gcaaacttca	gctttacagt	ttcagaggaa	tgtgggtttt	gagtctgtca	tccttgagaa	81120
acctgatatg	actttactta	gttccatata	ctcctgggtc	taggtaacag	tacatagcca	81180
gcaaatgctc	tatctccctg	tctaccttaa	tcttaggcag	gtgctgcaca	cctaggcttt	81240
gatggaaggg	atttcttagt	gttcttgccc	ctccttctca	tggaacacgt	atctgtgttg	81300
ctgtttgtga	agaagagtag	tggatgtcta	ctttgttgca	atgcaggatc	ctgggcccaa	81360
gatttcccg	cgtccctcca	aggaataaaa	atTTTggcca	gtacccctct	ctgagagaca	81420
atgtgtcttt	gcctggaagt	cctagatgga	ggaccacttc	ctgccccatc	ttccagaaac	81480
ttaaggcttt	ggctttggag	gatcagtgtc	ctggagaaat	gtgtgacggg	ttcatgtctg	81540
ccccactga	caaccaccac	ctacagcctg	caccgcctga	tgcatggcac	tctggtctcc	81600
tgcttgttc	tcaggaacac	ccaaaagaga	tctttgcca	agaacaggca	catgagtgca	81660
atTTtgactg	ataggcactc	tgatctgtcc	tttggtgccc	aggttttaaa	gaaaatcttt	81720
ctaaaaactc	attgaagttc	cagaatgcta	tgaatctttg	agctttgtta	ttggcatgtc	81780

BI OL0250W0SEQ_ST25. txt

catctgccta ctaatgtaga acagagcatg gtcgtcattt tcagagatga tgtcctgttt	81840
ctatcatgga ttttttttct catgcttctg tgttctggaa attactcagt ttgttttctc	81900
ctctttgaat ttcagcacca acggagcaaa gccccacagt ccaggactgc taccatggtg	81960
atggacagag ttatcgaggc tcatttctcca ccactgttac aggaaggaca tgtcagtcctt	82020
ggtcctctat gacaccacac tggcatcaga gaaccacaga atactacca aatgggtatg	82080
tctttgagtt ttctcccaag agaaacagcc acccacttaa atttctcctg gaagagccat	82140
gcttccagct aacttcttat gaccaatttt ctctcagacc cagaatgttg gacagaatgt	82200
ctcaggcttc ttgctttggg cacaggggtct gagaggagag aaatgtcagg ccagctctct	82260
tttctcatag ttgatagaag taggaggata cttggagggtg gtgagggtctc atgaatagaa	82320
agctcagaag aacatatgac catgtgctta gaaatagcac cattccacaa tgcccactaa	82380
agaccagtga aatagttcaa ccagggaatt ctgtcattct aatctccaag ccctggagtg	82440
aaggtttgtt ttgccatgtt tgtcttgggt aacaagtga ggatatctat attgacttcg	82500
agatcttccg atcactttct cctctaacct gtataaacac attgggttct gagaacaagg	82560
tgtctgaaaa gctatgtgtt gccagcccat gaggggcaaa aggaggaagg cagctgagag	82620
tcaggaagta tagagatgct gaagagttac acattcagga agatggacag aaacccatgt	82680
ctggctatgc cagcctttct gccatgaagg actatcaaat acatgagaaa acagttttca	82740
caggttggac aacagatatg gtaggcttga gagaactgag aaagggaatc aaaggagatc	82800
aacttcatca ttaacctgtc ttccttcctg gacacagtgt tggattgaag gacaagcaga	82860
tggagcccag ctgaccacag cagtcttgct taactgagga gagagactgg agtctgcat	82920
gcctcaggca gctgatgtgt tctaggctgg ctaagaatga gaagggattt gtggaagaaa	82980
ggagctccag gaatacacac agaagtctcc tcaaggcttt ggctaaatac aaagctgcgt	83040
atgcacaggg agagttttca taaagaaaga acaacaaaga aaagctactt gggaaagaac	83100
aactgcaggg gaacagtaag ctcaatggag atgccagagc tcacatagca ctgggggata	83160
tttgaattct gaccactcag aggagaaaca cctcactaca ttttgggcat tcagtagaga	83220
ccaaagaaa ctgtattttg ggattgggat catcttattc ctagaatcaa ggtgactcca	83280
gaaaaactcc aacaaccctt cttccaagcc agtctaaaag gatccaaatg atctccaagt	83340
aaattaactg cattccacaa gaaaaaaaaa actcaacccc ccttagaggc aagggacaaa	83400
tacaagttgc tcagttatat ggcatccta ttgcgttact tctattttaa aatttaatag	83460
agacacaaga agctttcact gtgatacata actgggagaa aaaatcactc aacacaaaca	83520
ggcccagaaa ttatagaatt gatgacattg gtgagaacat ttaaaatgca cctctgagaa	83580
ctgtgtttca ggaaaatgtc agcaaaagct gaccatgaga gaaacaaaag cagaatagca	83640
agagaaaaga aaagaaccgg agagccaaat gaaaattaaa gaactgagaa aagggtacatc	83700
tctaataag aactcactgg atggccttat catcacttta gacattacgg taggaaaggt	83760
gacctagaaa ataattcaat aggagctaca caaatcacag gacagacaga cagaccaaca	83820

BI OL0250W0SEQ_ST25. txt

gacagaaaca	cacacacaca	cacacacaca	cacacacaca	cacacacaca	cacacaaaga	83880
ctgaacctat	taatcaacag	agcctcaagg	gcatctagga	aaaatccaca	cattttaatat	83940
atgtgttagg	caagtcacag	aaggagaaga	aaaagatatc	atgacagaca	ttatacttga	84000
agcgatgatg	gctcgcaaca	cgccaaatat	acagaaaaca	agaaactcat	agtcaagaag	84060
ctaaatgact	caggtataga	attttaaaga	gcaaaaactct	atgattttact	gggatatatc	84120
atagttaagt	tgcctcaatt	caaagctaaa	aagaaaaaaa	gggggttcct	atgaacaaca	84180
gctttgacag	ctgttgatct	aagaccacag	cttaaataatc	taggcaagga	aaagcaaata	84240
aggcacaagg	aaagaggatg	gaaggaaata	gtccaaacca	atgacattca	gtggaaaaga	84300
aaatagacca	acaaaggagt	aatccatga	aacagaaagt	taggtttcttt	gaaaagtcta	84360
tatgattggc	caaagtctgg	ctaaacagat	gacagaccaa	ggaggggagca	tatccatcac	84420
catcatgagt	aacaggagag	agatgccatt	gctatagcat	cctccagggtg	tgaaagctga	84480
gaagtagata	ttgagatcaa	ctgtatgtaa	ataaatcat	gaatgtagat	catgtggatg	84540
gattgcttag	gtaaataaca	aatcagcaaa	tcaaactg	aatagatcat	gcagttttat	84600
agagacttac	agacagcctg	acagataaac	atttgtatgt	acgtgaaaca	atctccaaga	84660
cacacttcaa	aatccctctc	ggttaatcca	aaggaatgta	tttggcagaa	ggtagaagga	84720
gggtattctg	atcctttctg	gtacacattg	atgttttctc	tcagttttct	tataaagcat	84780
agattacttt	gaatgtgtta	caataagaat	cataagctgt	ctttgaaatg	ttgacagttg	84840
tttagaagtt	gaggaccatt	tgtgagtgtt	atgggacttt	agtgagaata	tttcaaattt	84900
gcttgtttac	actttgttac	aagaaaacat	agagggtgcc	aggtggtgct	gtatcttctc	84960
caatctctgg	tgacctgtat	tgttttggaa	tttgcagtgg	cctgaccagg	aactactgca	85020
ggaatccaga	tgctgagatt	cgcccttgg	gttataccat	ggatcccagt	gtcagatggg	85080
agtactgcaa	cctgacgcaa	tgtccagtga	tggaatcaac	tctcctcaca	actcccacgg	85140
tggtdccagt	tccaagcaca	gagcttctt	ctgaagaagg	taagaagcct	gcagtcagac	85200
aaccataccc	tcggacattg	ggataaaaag	atttgcaaaa	tctttgtgat	gcagaaaact	85260
tccatgctgc	acaggaagtc	gaaggatgaag	tcatggacag	ccaatgggaa	ggaagcttca	85320
gtgccttctc	tggggggacc	agagctggga	tgttgagtgc	cttgtgaggg	atggtgtctt	85380
taaaaggggc	acagaccctc	taggacactg	gatttatcac	ttccctgtta	tcaaacgaat	85440
catattagt	tcctagccaa	gatggatatt	ctaacatcct	gccaaacttg	tgaagatata	85500
ccaagctcct	aagcctgtcc	agccctttct	tcaagtaggc	agtgtttatt	gcagtcttca	85560
gctttacat	tttgaaggaa	tgccattttt	gaggctgttg	ttcttgagaa	acctaactg	85620
tcttcattag	atccgtattg	tcctgagact	ttgaagcagt	acatagccac	caaattgttt	85680
atctccccag	cctaccttca	tcttgggcat	gccttcacaa	cctaggattt	gaggggaagg	85740
atttctcagt	gttctcatcc	ctgcttctca	tggaaacattt	atctccgttg	ttttttgaga	85800
agaagagtag	tggatgtcag	ctttcttgta	atgagggatc	ctgggcccaa	gattccctgt	85860

BI OL0250W0SEQ_ST25. txt

ctccccctcct	aggctataaa	atthttggcct	gtactccttc	tccctgagag	gcaatgtgtc	85920
tttacctaca	agtcctagat	gcaagatcct	tttctgcccc	acaccccaga	atctgaaggc	85980
ttttgctttg	gaggagcagt	ggtctagtgt	gcaagggttt	catgtatacc	ccccactaac	86040
agccaatcac	cacctatagc	ctgaacagct	tgatgcatgg	caccctggtc	tcctgccttg	86100
ttctcatgaa	caccagaag	aggtgtaagc	aaaagaccat	tcacatgagt	gtaattttga	86160
agtataggca	ctctgatctg	ttttttgttt	gtttctttgt	ttgtttgttt	tccagggttg	86220
aattaaaata	tttatgacta	cttattaaat	ttctagaatc	ctataagtct	atthgtatth	86280
ttattctaca	tttcaatthg	catgctaata	tagaagagtg	taaattgtta	atcctcagat	86340
tattccactt	tgtgtgtcat	aatthttttc	acatthccct	tttctaggca	atactgagct	86400
tgatthttct	ttttaatthc	agcaccaact	gaaaacagca	ctgggggtcca	ggactgctac	86460
cgagggtgatg	gacagagtta	tcgaggcaca	ctctccacca	ctatcacagg	aagaacatgt	86520
cagtcttgggt	cgtctatgac	accacattgg	catcgaggga	tcccattata	ctatccaaat	86580
gcgtatgtct	atcatgttag	ccataaaaagg	aacaatagtc	aactaaaatt	tctcttagct	86640
ggcccatgct	acaagctcac	ttcctaggtc	caaatthttc	atagactcag	agthttgtagc	86700
aaaatgtctc	aggaaactta	ctthttgagca	aaagggtctga	atgaagagaa	gtthtttaggat	86760
tgctatctth	cataacaatt	tgatggaagc	agcaggatat	atggagggtgg	tgaagtctca	86820
ttaatgtaaa	gctaaggaga	tcaaattgacc	aatgtctgag	acaaagtatc	attccacaat	86880
gccactaaa	ggtccatgca	gtctthtcaac	catgcaatth	tatcattcta	tcctccattc	86940
cctgaagtga	aatthgtgtt	tgccatthtt	gacacgaatc	agaagtaaca	aattcaggct	87000
gggtgcagtg	gctcaggcct	gtgatcccaa	cactthtggga	ggacaagacg	ggcagatcac	87060
cagagggtcag	gagttcaaga	ccagcctggc	taacatggca	aaaccccatc	tctacgaaaa	87120
attaaaaaat	tagccggtca	tggtggtggg	tacctgtaat	tccaactact	tgggagggtg	87180
aggcaggaga	aacacttgag	cctgggattc	agagthttgt	gtgagccgag	aacatgccac	87240
tgactccag	cctgggtgac	agagcaagac	tcaatctcaa	aaaaaaaaaa	aaagaagaag	87300
aagaagaaaa	gaagaagagg	aagaagaaga	agaggaagaa	gaagaagaag	aagaagagga	87360
agaggaagag	gaggaggagg	aggaggagga	agaagaagaa	gaagaagaag	aagaagaaga	87420
agaagaagaa	gaagaagaag	aagaagaaga	agaagaaaat	agaaatgagt	gcatatatth	87480
atatatgagt	actagcctgt	atgaacacac	tgggttctaa	gcaccagtht	tctgaaggga	87540
tatgggttgt	caggcagagt	aaaagcagga	atgcagatga	gagtcaggaa	gtaaacagat	87600
gtggtgatta	aatgggcag	gtacatggac	aaaaaatgc	atgtctgaca	aaaactggcc	87660
tcttgccata	agtgagtatg	aataatatgg	aaaaactgtt	tgacatgtt	gaacagcaga	87720
cagtacaacc	tgagatagth	tagaaaggga	aacaaataag	atcaacccca	taattaccct	87780
tcctagactt	aagggc aaag	agthtttaacc	aaagcattcc	acagcagtht	tgctaaactg	87840
gggagagaga	ctggagthtt	gtthtactaat	aaaaccgaga	thttctaggt	taggtaataa	87900

BI OL0250W0SEQ_ST25. txt

tgagaaagta	tttgtggaga	aaaggagctc	caggaataca	cacagaagtc	tcttcaagtc	87960
tctggctgaa	cagaaagctg	tgtatgcaca	gaaagagttt	ccagagagaa	aggagaacaa	88020
agaacagcta	ctggggaaag	aacaactgct	ggggaacagt	gagctcaatg	aagatgccag	88080
agctcacata	gcactgggag	gtatttgagc	tctgaccagc	ctgaggagag	acacttcatt	88140
gaacatcttg	ggcattcagc	aaagacccca	aaaaaccata	cttcaggagt	agaattaatg	88200
cattcctaga	ataaagtcta	ctccagaaac	accctagaaa	agcttagaaa	ccaagtctaa	88260
aaagatccaa	atgatctcca	agtaaattaa	ttgcctgtca	gaagaaaaca	acctcttcag	88320
aggtaaacaa	caaaattaaa	ttgctcaatt	atatagtatg	cacaatgtgt	ggcatacatt	88380
taaaaatttg	ctaaacatac	aaaaagcatt	tagtgtgacc	cataaccagg	agaaaaatca	88440
gtcaatacaa	atagacccaa	aatgataaaa	aataacagaa	ttggcaagga	gattttaaata	88500
gtatgtatca	taattgtgtt	caaggattta	aagaaagcgt	ggacaagaaa	taaataaatg	88560
gataatatca	acagaaagaa	aaattgtaaa	aggaccaaata	ggagagtcaa	gaactgaaaa	88620
aaaagacatc	tctttaatga	gaaaatcact	acatggcctt	ataatcatat	tagatagtac	88680
agatgataaa	gctaactaga	aatatttagg	gtggtgcaaa	ccatagcacg	cttatacaaa	88740
gcctgagaag	ataaacagag	cctcaaggac	atctatgaaa	atatcaaaat	atttaatat	88800
tgtttaaagc	aagtcacaga	ggaagggaaa	gagatattgg	aacagaaaaa	atacttgaag	88860
cagtgatggc	tgatgacttt	ctaaatatgg	aaaaaatgat	aaactcacat	agtcaagaag	88920
ctcaatggat	cagatatagg	attttaaaaa	gtaaagctgt	atgatttatt	tggacacatc	88980
ataattaaat	tgtccataat	caaagataga	aagtaaaatc	ttatttgaag	ccaaggggaa	89040
aaaacatacc	tttacaatga	gtaacagtga	cacaaatgac	tgatgccttc	tcatacagaaa	89100
caacacaaat	cagaaacaat	agaataacac	ctttagagtg	gtaagaagaa	aaaaagatca	89160
aatcagaaac	aacaaaataa	cacgtttaga	gtggttaagga	ggaaaacaag	atcaaatcag	89220
aaacaatgga	ataacacctt	tagagtgtaa	gaaagaaaaa	aagatcaaata	caggaacaac	89280
agaataacgc	cttcagagtg	gtaagaagga	aaacaagata	aatcagaaaa	caatgaaata	89340
acacctttag	agtagtaaga	agaagaaaag	atcaggtcag	aaaaaatgga	ataatatgct	89400
aagaagaaaa	aaaaagatca	agtcagaaac	aatggaataa	cacctttaga	gtgaaaagaa	89460
ggaaaaaaaac	ccagcaagct	taaacgctat	gcacagcaaa	caattccact	gaaaatgaat	89520
gttacgtaag	tacatatctt	gtcctcctaa	aaacaaagaa	caaataaaaag	aatgtttcat	89580
cagcaggatt	atgtaataaa	agatgtgaaa	gaatgctatg	taagtagaag	aaaaataata	89640
ccatatggga	attggcatca	aaaccacaaa	atactatcaa	aacaaaaaaa	ctttattgat	89700
aaatttaaca	caatatgcaa	aagaactata	ccatgtatac	tacataacat	tggtgagaag	89760
aaaattagaa	gatctaaata	aagacacatc	atgcttatag	attaaaaaat	ccaatgtcac	89820
ttttcacaaa	actgatcttt	agtttcaacc	cacacccaag	cagaattcct	gcagtctttt	89880
cttgaaaacc	taacagaatg	tatatgctag	aatcaccaag	acaatcttta	aaaagaataa	89940

BI OL0250W0SEQ_ST25. txt

aaaacttgga	ataaaatcac	aagtttgtgg	gatagatgca	tatggtaata	tggaatttct	90000
cataaagaca	cagtaatcaa	gacatgtggt	attggctggg	acgcttggct	gtaatcctaa	90060
cactttggga	ggccaagatg	agaggattgc	ctgagatgag	gagttgcaga	caagcctggg	90120
caacatagca	agaccctcat	ctctacaaat	atttaaaaaa	attagccagg	tttggtgcca	90180
tgtgcctgta	gtcccagcta	ttcaggaagc	tgagggtggga	ggatcactgg	agcccatgag	90240
gtggaggctg	aaatgagcca	tgattgtgct	actgaacttt	agcctgggag	acagattaaa	90300
accttccctc	tctctctcaa	acaaacaaac	aaaaaataca	tagtattggg	caaaacatat	90360
gcaaacaaaa	acagaaaagg	gtcagcataa	atttacatat	atggtcaatt	tattttcaat	90420
acaggtagca	aagcaattta	atgaggaaat	ttttttccaa	aattgggtctg	aaacaactgg	90480
atagccatag	aaaaaaacta	taacaaatgt	gacgcttgaa	tcctactgta	tgactcaaat	90540
taaattaatt	tgagatagct	cttagacctc	aatgtaacag	ctaattctga	ggctgaaata	90600
taagactgct	atgaaaaagt	atagtatctt	ataaccttgg	agaaggaaaa	atTTTTtgag	90660
ggaagaacca	gaaaacacta	actgtaaaag	aaaacaaatg	ataatgtgga	cattcattga	90720
ataaaaaactt	atgctcacca	aatatgactg	ttaagaaaat	aaataagtaa	gtaacacact	90780
ggaagaaaaa	cactctcatc	catatatctg	acaaatggcc	tgtatccaga	gtatagaaac	90840
atttctccca	ctcactaatc	agaggacaaa	caacctaatc	aaaatgggca	acaggcttga	90900
atagtcatTT	cttaggagaa	gatgcacaca	gagccaacaa	tcacctgaaa	aagtgcacaa	90960
catcttagcc	atcaaaaatc	aagagttata	accctcataa	gatgacactg	aacatccagt	91020
gtacatggat	atcattaaga	agacacaata	ataagtgggtg	tcaccgattt	ggagctagaa	91080
tgtgccactc	tctcatatgc	tggtggaagt	tcaaaatcat	acaacaaatt	aaaaaatcag	91140
tctgatgctt	tcttataaag	ttcgataaat	atgcatctat	cctacaaacc	tgtaattcta	91200
ttcttgaata	tttaccctcc	aaaatgaaaa	cataagtcca	caaaaatcta	tataaatatt	91260
catagcagct	ttatgtttta	taaactcaaa	ataaaaaacta	tttcaatgtt	ttcatcaaaa	91320
gaaaatgaaa	actattttaa	tggtttcatc	aaaagaaaat	gaaaaaagaa	tttccagtat	91380
atttatacaa	aggaataacta	ttcatcaaca	aggaacaagt	tactgatagt	ctcagaagca	91440
tgaacaaacc	tcaaaaatat	attaaggaaa	gaagccagac	gtcaaagtgt	atagtctgta	91500
tgagtccatt	catgtgagtt	tatagaaaac	acaatttatg	gtgaaagaaa	ccaatagcat	91560
ttgacactgg	ccgtgggaag	agggtagcag	agattgattg	agcagccaca	caagggagtt	91620
tctgggggtg	tgaaaatgtt	ctgcattgtg	agggcagtgt	gggctacaca	agtatatgta	91680
tttatcaaat	ctcatccagc	tacatttaag	atctgtgcat	ctcactctat	gtgaaaatat	91740
actcaactga	aaaacagagc	aggtatctgt	ttcagggtgct	acatcacttg	atacgtccag	91800
ttgtgttaaa	aaccactgcc	taacatcctc	aaatggggga	tctgggcttg	agactaggctc	91860
acatgtgtag	agtctctaca	gagaccgtgt	tggattccca	tgctccataa	tacgttccaa	91920
gttttctcag	acagccacag	gtcatgaatg	tgaggattct	gagaggttgg	agcaacgttc	91980

BI OL0250W0SEQ_ST25. txt

ttgggaggca	taatggggaa	ggcattctcc	aagattcctc	cagcctgggg	tcttcacctg	92040
ctgtgcctct	tactgcattg	ttttctgact	catccatagc	cacttgaccc	cttcagatcc	92100
catagtctac	ctagccgtct	ccctttatgc	cttgggtccc	gctgttcttt	caactcatca	92160
cccattcctt	cagtcccaga	gtggctgcag	ccagcagagg	atggactgag	agcaggagag	92220
gaggctcgtc	ccatgaaccc	atcctagaga	agcagcatcc	tgccctgggag	ctagttttcc	92280
aggggaagctt	ttataagtcc	tgtagacca	aaccctctg	ctctaccaga	tacagtatct	92340
atagtaatac	tattttcatg	attatcttat	attgcaaagt	tagagcattt	atgctacact	92400
atgagtaaat	agagtaaggg	ggctggcatg	ggaattatat	aatcttggat	gccacttctt	92460
ccttggggaa	atgtatttga	gttccaactt	acatattact	atatagtctt	atagagagag	92520
agacaaagag	ctagacagac	agagatatct	ttgtatgtgc	attaaaaaat	ctaagataca	92580
tatttcaaaa	tctgtgtcat	ttattctgga	ggaaagtatt	tggcagaagg	tgaaaggaag	92640
atattctgat	cctttcttgt	acagacatgt	attatctcag	ttttcataga	gagcatatac	92700
tacttttgat	gttttaaaac	aaaaattata	atctgtgatg	tgtccacagt	tgtttaaaag	92760
ttgaagctga	agaccatttg	tgcttgtggc	aatattattg	tgggtataatg	ggaatatttc	92820
aaaggcactt	gttaacactt	tgttacagca	aaatgtagag	ggcgctaagt	gcccttgaat	92880
attctcccat	ctctggtgac	ctgtgttgtt	ttgaaatttg	cagtggcctg	accaggaact	92940
actgcaggaa	tccagatgct	gagattcgcc	cttgggtgta	caccatggat	cccagtgtca	93000
ggtgggagta	ctgcaacctg	acacgatgtc	cagtgcagag	atcgagtgtc	ctcacaactc	93060
ccacagtggc	cccggttcca	agcacagagg	ctccttctga	acaaggtaag	aaatttgtgg	93120
ttagacatct	atatactggg	atgaaaaacc	atggaaaatc	ttactgatgc	agaagccttc	93180
agtggtagac	tggaggggtg	gttgagggtc	tgcaatgtgg	aggaaagcct	cagcgccctc	93240
tctgggggat	ccagaactgt	gatttttggc	acgctgtgag	gaggcagtgt	ctttaggaag	93300
ggcacggtgt	ctttaggaag	ggcacagacc	cgccagggca	ctggacttac	cactcccctg	93360
gttattaaat	gggtcatttc	agtgtcctag	ccaaaatgga	tattctaaca	gcctgccaaa	93420
tatgtgaaga	tttccaagcc	aataagcctt	tccagtgatt	taaagtagac	ttttttcatt	93480
gcaatctaca	gtttgcagtt	tcttaagaac	atggcctttg	agtatgatat	cctagagaaa	93540
cctaaggaga	ctgcattatt	tttctattgt	cctggggctg	catagcagga	ggtaaccaac	93600
gaatgctgtc	tctccctggc	ctatctcagt	ctttcacagg	ctctgttcac	ctcagctttg	93660
aagttagaaa	tttctaggtg	ttcttgccct	ttcttctcat	gaaacctgca	ttggcagtga	93720
gtctacagaa	gaagaggaag	agaattctgc	tttgttacia	ttcaggactc	tgggcactag	93780
aagattccct	atctctctc	caagggaata	agttgtttgt	ctctaaccct	ccttgagaaa	93840
caatgagtct	ttgcctgcac	tcctaaatgt	aggatgattt	cctgcccata	ttttcaaaag	93900
attaagcctt	ttgccttggg	atgagcaatg	gtctagggaa	atgcgcaagg	gtcttgtgtc	93960
ggcccctgac	tgaccaccag	tcacctccta	cagcctgcac	caagggaatg	attgcattct	94020

BI OL0250W0SEQ_ST25. txt

ggtcttctgc cctgtggttc tcatgaaaac cagcagagat tcatatgatg gagctgcaca	94080
tgaatgtaat ttccaatgtc cagcattctc ctctgttctt tatctttaga tttaaaaata	94140
atgtttctat gaacttatta aaattctaga atactatgaa tctactgggt cttttcacat	94200
ccttttgcta ctagtagaaa aaagaatagt aataattttc agaggctact gtccagtatg	94260
tgacataaat tgtctcccat gtttctctgc tcatgcaatt actgagtatg atttatttta	94320
ttttaatttc agcaccacct gagaaaagcc ctgtgggtcca ggattgctac catgggtgatg	94380
gacggagtta tcgaggcata tcctccacca ctgtcacagg aaggacctgt caatcttggt	94440
catctatgat accacactgg catcagagga cccagaaaa ctacccaaat gcgtatgtat	94500
ttgattaaaa ccataagagg agcaacagcc aactcaaata ttggttagaa gacccatgct	94560
ttaagctcac ttcttaggga caaatttctc ttagactcac attttgcaa aatgtctcag	94620
gacctttgct tttgagcaaa gagtctaaga gaagagaaat tttaggcctg ctatttttcc	94680
taatagtttt atggaaggag tagaatatac ggaagtggcg aagtcataatt aatgtaaagc	94740
tcagaagata aatgaccaa gcttaaacac agcaccattc cacaatgccc actaaaaatc	94800
aatgtcatct ttactcgtg caattctgtc attctaaatt tcaattcccg aaggtttggt	94860
tgccattttt gtcatgggta ataagtaaaa aaaaaaaaaat taagatgtgt atatatatat	94920
atatatatat atatatacac acacacacac acacacaaac atctgaatat ttatatatat	94980
gtctgaatat ttatatactt gtgtataaaa cttatattta aatttttgca taaattttata	95040
tatttttaat atttcattaa aaattatatt gtttcactat gtatgtctga gtatttttat	95100
atattttaat ataacatttt aaatatttat atataaatat tcaggatatgt aactgaatat	95160
tcatttacac acacaaatat atgtgtgcat gtgtgtatat atatatatat ccatatatat	95220
atatatatat atatatacat atatatatat atatatatat gtatatatat atatatatat	95280
atatatacac acacacacac acacacatac atacaggat aaacacactg ggcctgaagc	95340
accagtgggtc tgaaaggaca tgtgttgcca ggacttgaag agcaaaaagca ggaaggcgga	95400
tgagagtcag gaggtacaca aacgctgaaa agtaaaatgg acaagtacat ggacaaaaag	95460
caggataag cataacagcc ttttggaagt aaatgactat aaaatatatg aaaatactgt	95520
tttcacaagt tgcacaacag atagtagtgt attgagataa tttagaacag aaaacaaatg	95580
tgatcaaccc cataagtgtg ctgtatttca tcatggattg aaggaaaaag agatggagcc	95640
caagaagacc acagcagtct tgatgaactg agagacacca gagtttgga ttacaaaggc	95700
agctgggatt ttctacactt ggtaataatg agaaagaatt tgtggagata aagagctaca	95760
gtcatgtacc tagaagtcac ctcaagttaa tataaatctg catatgcaca gggagtgatt	95820
ccacaatgaa agtaggacaa agaacagcta ctggggaaag aataactaca aggaacaat	95880
gagttcaatg gagatggcag agctcacaaa gcactggggg atatttgagt tcttaccagc	95940
tagaaaagag acctcattgc aaatcttggg cattcagtag agacccaga aaagccactc	96000
tttggaaca gagttgatgt attttaagag caaaatctac tccacaaaaa tcctagcaaa	96060

BI OL0250W0SEQ_ST25. txt

attgaaaagc	aagtcagaaa	gaccaaatac	ctctcaacat	aaattagttg	cccatcagaa	96120
gaaagcttaa	cctcttcata	ggtaaacaat	aaaatcaaat	tgctcagtta	tctggcatcc	96180
acaatatgtg	acataaattt	aaaaatttac	tagacataca	agaagcattt	agtgtgatcc	96240
ataaccagga	gaaaaatcat	tcaatacaaa	tagacccaga	aatgacagaa	atgatagaat	96300
tagcaaaaac	atttaaaata	tacatatgat	catttgatct	tgtgatcaga	tatcacaaga	96360
gaagaaagag	atacttgaac	agaaaaaatg	cctgaagcaa	tgatggctga	aaactttcca	96420
aatatgaaga	aaaaaaagct	cacagattca	agaaaactaa	tcaatcagaa	atatgatttt	96480
gaaaagtaaa	aatgtatgat	ttactttggc	aaatcttctt	ggttaaattg	tctaaaatca	96540
aagaaagcta	ggaaaatttt	ataagccaga	ggaaaaaaga	ttgtttatat	aaaggaacag	96600
ttacacaaat	gactgatgcc	ttctcatcag	aaacaatgaa	agtcagaaac	aataaagtaa	96660
catcttttaa	gtaatagaag	aaaaacccaa	gaggtgaggg	atcgtggcag	acaggaggca	96720
ggactagatt	gcagctctgg	acagagcagc	atgcagaggc	tcatattgtg	aattttagcc	96780
ccatattgac	tgcaagaaca	gaccagcaat	cctgagagga	cccacagacc	gtgtgaagga	96840
agcagactgc	tcctgcagga	taagggagac	accccaaata	ctgtgagttc	cccaactgca	96900
gaagtggaaa	agggaggcct	tactccctca	aacacacccc	acaactggag	aagctgaaag	96960
tctgtttgca	ggagaagttc	ccaactttac	ctgggcctca	gtaaatttag	agagctgagc	97020
caagcaaaat	ataggggtag	aggaagcagc	agagaagacc	tcagagcttg	ctggatcccc	97080
aagcagctca	ttcctgcctg	gcaccacaga	gatccatcag	aagtgtggcc	aaaggaacag	97140
agggtaaaac	tccacatgga	ggactgctct	acctgaactt	tctaacaatt	tgaacagggg	97200
gagaagcctc	ctggccagaa	cttggggggag	ggcatgaatc	tggtttgcag	acttcacagg	97260
tgggggaagg	actaaagccc	ttttctttca	cagctgggag	gtggaaagcc	tcaggcaagt	97320
tttcaagcct	gactttcccc	ccacctggaa	acagacttgg	agctgttgcg	gggttggggg	97380
catggtggga	gtaagaccag	cccttcagtt	tgcatgggtg	ctgggtgagg	cctgtgactg	97440
acagcttccc	tccacttccc	cgacaactca	gatgactcag	cagaggcagc	cataatcctc	97500
ctaggtacac	aactccagtg	acctgggaac	ttcaccccca	caccatacag	aagcttcagt	97560
aagacgtgcc	caaggaaagt	ctgagctcag	acacgcctag	tcccaccccc	aactgatggt	97620
ccttccctac	ccaccctggt	agcagaagac	aaagagcata	taatctttgg	agttctaggg	97680
cccaccacc	tctagtccct	ctccacacta	gtatagctga	tgaggaggc	caaccagcac	97740
aaaaatagag	cattaaacca	ccaaagctag	gaacccttat	ggagtccatt	gcaccctcct	97800
ccacctccac	cagaacaggc	actggtatcc	acagctgaga	gacccataga	tggttcacat	97860
cacaggactc	tgtacagaca	gtccccagta	ccagcccaga	gctgggtaga	cttgctaggt	97920
ggcaagaccc	agaagacagg	caataatcac	tgagtttcag	ctcacaggaa	gccacatcca	97980
taggaaaaga	gggagagtac	tacatcaagg	gaacacccca	tgggataaaa	acatctgaac	98040
aacagccttc	agccctacct	tccctctgac	acagtctacc	caaatgagaa	ggaaccagaa	98100

BI OL0250W0SEQ_ST25. txt

aaccaaccct	ggtaatatga	caaaacaagg	ctcatcacac	tcccagttca	ccagcaatgg	98160
atccaaacca	agaagaaatc	cctgatttac	ctgaaagaga	attcaggagg	ttagttatta	98220
agctaatacag	ggagggacca	gagaaaggca	aagcccaatg	caaggaaatc	caaaaaaaaa	98280
aaggtataag	aagtaaaagg	tgaaatattc	aacaaaatag	atagcttaat	aaaaaaacaa	98340
taaaaaattc	agtagacttt	ggacacacct	ttggaaatgt	gacatgctct	ggaaagtctc	98400
agcaatagaa	ctgaacaagt	agaaaaaata	aattcagagc	tcaaagacaa	ggacttcaaa	98460
ttaacccaat	ccaacaaaga	caaagaataa	aggataagaa	aatatgaaca	aagccttcaa	98520
gatgtctggg	attatgttaa	atgaccaaat	ataagaataa	tcgtggctcc	tgaggaaaaa	98580
gacaatacta	aaagcttggg	aaacatattt	gggggaataa	ctggggaaaa	cttacctggc	98640
cttgctggac	acctagacat	gcaaatacaa	gaaacacaaa	gaacatgtaa	atacaagcag	98700
cacaaagaac	acctgggaaa	ttcatcacaa	aaagatctta	gcctaggcac	attctcatca	98760
ggttatgcaa	agttaagacg	aaggcaagaa	tcttaagagc	tgtgagacag	aagcaccagg	98820
taatgtataa	aggaaaccct	atcagattaa	cagccagttt	ttcagcagga	actgtacaag	98880
ctataaagga	ttggagccct	atcatagcct	cctcaaacaa	aacaattatc	agtcaagaat	98940
tttgtatcca	gcgaaagtaa	gcatcatata	tgaaggaaag	atacagtcgt	ttttggacaa	99000
acaaatgcta	agagaattca	ccattaccaa	gtcaccacta	gaagaactgc	taaaaggagc	99060
tctaaatctt	gaaacaaatc	ctagaaacac	atgaaaacag	aatctcttta	aagcataaat	99120
cacacaggac	ctataaaaca	aaagtacaag	ttaaaaaaca	aaaacaaaaa	acaaaaccaa	99180
agtacggagg	caataaagaa	tatgatgaat	gcagtggcac	ctcacatttc	aatgctaaaa	99240
ttgaatctaa	atggcctaaa	tgctccactt	aaaggataca	aaaagagttg	gtggctggca	99300
agatggctga	ataggaacag	ctccagtctg	ccgctccccg	tgagatcaac	acatagggtg	99360
ggtcatttct	gcatttccaa	ccaaggtacc	cggctcatct	cattgggact	ggttagacag	99420
tgggtgcagc	ccacagaggg	tgacctgaag	caggggtggg	tgtcacctca	cctgggaagt	99480
ggaaggggtc	agggaaactcc	ctcccctagc	caaaggaagc	cgtgaggggac	tgtgccgtga	99540
agaccagtgc	attctggcac	aaatactatg	cttttcccac	ggtctttgca	acctgaagac	99600
caggagattc	ccttgggtgc	ctacaccacc	agggccctgg	atttcaagcc	caaaactggg	99660
ctggcatttg	ggcagacact	aagctagctg	caggagtttt	ttttcatacc	ccagtgggtc	99720
ctggaatgcc	agcaagacag	aaccattcac	ccccgtgaag	aaagggctga	agccagggag	99780
ctaagtggtc	tttctcagtg	gatcccaccc	ccatggagcc	cagcaagcta	agctccactg	99840
gcttgaaatt	cttgctgcc	gcacagcagt	ctgaagttga	cctgggacgc	tcaagcttgg	99900
tgggaggagg	ggtatccaca	aatactgggg	cttgagtagg	aggttttccc	ctcacagtgt	99960
aagcaaaacc	gctaggaagt	ttgaactggg	caggggtgcac	tgcagcttgg	caaagccatt	100020
gtagcaagag	tgcctctcta	gattcctcct	ctctgggcag	ggcatctctg	aaagaaaggc	100080
agcagcccca	gtcagaagct	tatagataaa	actcccatct	ccctgggaca	gagcaactgg	100140

BI OL0250W0SEQ_ST25. txt

aggaaggggt	ggctgtgagt	gcagctccag	cagacttagt	ttcctgcctg	ccagctctga	100200
aaagagcacc	agatcccca	acacagcact	agagctctga	taaggacag	actgcctcct	100260
caagtgggtc	ctggtttcag	aagataataa	gaaactcctc	tgagctaaag	gagcatgttc	100320
taacacaatg	caaggaagct	aagaaccttg	aaaaaggtca	gaggaattgc	taactacagt	100380
aagcagttta	gagaagaaca	taaatgacct	tagggagctg	aaaaacacag	cacgagaact	100440
tcattgacaca	tacacaagta	tcaatagcaa	aatcgatcaa	gtggaagaaa	ggatatcaga	100500
gattgaaaat	caacttaatg	aagtaaagcg	tgaaaacaag	attaaggaat	aaagaatgaa	100560
aaggaatgaa	caaatcctcc	aagtatggga	ctatgtgaaa	agattgaacc	tacgtttgat	100620
tggtgtacct	gaaagtgatg	ggagaatgga	accaagttgg	aaaacactct	tcaggatatt	100680
atccaggaga	acttcccaa	cctagcaaga	caggccaaca	ttcaaattaa	ggaaatacag	100740
agaataccac	attcaaattc	aggaaataca	gagaacacca	caaagatact	cctcaagaag	100800
agcaacctga	agacacataa	tcgtcagatt	caccaaggtt	gaaatgaagg	aaaaaaatgt	100860
tgagggcagc	cagagagaaa	gtttgggtta	cccacaaagg	gaaccccatc	agactaacag	100920
tggatcttcc	tgacagaaact	ctacaagcca	gaagagagtg	ggaggccaat	attcaacatt	100980
cttttttact	attattatac	tttaagttct	agggtacatg	tgacacaggt	gcaggtttgt	101040
tacatatgta	tacatgtgcc	atgttggtgt	gctgcacca	ttaactcttc	atttacatta	101100
ggtatatctc	ctaatactat	ccctccccac	tcccccatc	ccatgacagg	ccccggtgtg	101160
tgatgttccc	cactctgtgt	ccatgtactc	tcattgttca	attcccacct	atgagtgaga	101220
acattcgggtg	tttgatttcc	tgtccttgtg	atagtttgct	gagaatgatg	gtttccagct	101280
tcattccacat	ccctacaaag	gacatgaagt	catccttctt	tatggctgca	tagtattcca	101340
tggtgtatat	gtgccacatt	ttcttaatcc	agtctacat	tgatggacgt	ttgtgttggt	101400
tccaagtctt	tgctattgtg	aatagtgccg	caataaacat	atgtgtgcat	gtgtctttat	101460
agcagcatga	tttataatcc	tttagatata	tatccagtaa	ttgtatggct	gtgtcaaatg	101520
gtatttctag	ttctaaatcc	ttgaggaatc	accgcactgt	cttccacaat	ggttgaacta	101580
gtttacagtc	ccaccaccag	tgtaaaaatg	ttcctatttc	tccacatcct	ctctagcatc	101640
tggtgtttcc	tgacttttta	atgatcacca	ttctaactgg	tatgagatgg	tatctcattg	101700
tggttttgat	ttgcatttct	ctgatggcca	gtgatgggtg	gcactttttc	atgtgtctct	101760
tgactgcata	aaagttttct	tttgagaatt	gtctgttaat	atcctttgcc	aactttttga	101820
tggtgttggt	tgattttttt	tcttgtaaatt	ttgtttatgt	tctttgtaga	ttctggatat	101880
tagccctttg	tcagatgggt	agattgtaaa	aattttctcc	cattctgtag	cttgccgtgt	101940
cattctgagg	gtagtttctt	ttgctgtgca	gaagctcttt	agtttaatta	gatcccatg	102000
gtcaattttg	gcttttggtg	ctattgcttt	tggtgattta	gtcatgaagt	ccttgcccat	102060
gcctatgtcc	tgaatgggtat	tgcttaggtt	ttcttctagg	gtttatatgg	ttttaggtct	102120
aacatttaag	tctttaatcc	atcttgaatt	aatttttata	taaggtgtaa	ggaagggatc	102180

BI OL0250W0SEQ_ST25. txt

cagtttcagc	tttctacata	tggttaggca	gttttcccag	cacatgtat	taaataggga	102240
aacctttccc	tatttcttgt	ttttgtcagg	tttgtcatag	atcagatggt	tgtagatgtg	102300
tggtattatt	tctgagggct	ctgttctgtt	ccattggtct	atatctctgt	tttggtacca	102360
gtacatgct	gttttgggta	ctgtagcctt	gtaatgtagt	ttgaagtcag	gcagagtgat	102420
gcctccagct	ttgctttttt	ggcttaggat	tgtcttggca	atgcatgctc	ttttttgttc	102480
catatgaact	ttaaagtagt	tttttccaat	tctgtgaaga	aagtcattgg	tagcttgatg	102540
gggatggcat	tgaatctata	aattacctta	ggcagtatgg	ccattttcac	aatattgatt	102600
cttcctatcc	atgagcatgg	aatgttcttc	catttgtttg	tgtcctcttt	tatttcatta	102660
agcagtgggt	tgtagttctc	cttgaagagg	tccttcccat	cccttgtaag	ttggattcct	102720
aggtatttta	ttctctttga	agcaattgtg	aatgggagtt	catccatgtc	cctacaaagg	102780
acatgaagtc	atgtatggga	atgcttgtga	tttttgcaca	ttgattttgt	atcttgagac	102840
tttgctgaag	ttgcttatca	gcttaaggag	attttgggtct	gagaagatgg	ggttttctaa	102900
atatacaatc	atgtcatctg	caaacaggga	caatttaact	tcctcttttc	ctaactgaat	102960
accttttatt	tccttctcct	gcctaattgc	cctggccaga	acttccaaca	ctatgttgaa	103020
taggagtggg	gagagagggc	atccctgtct	tgtgccagtt	ttcaaaggga	atgcttccag	103080
tttttgccca	ttcagtatga	tattggctat	gggtttgtca	taaatagctc	ttattatttt	103140
gagatatgtc	ccatcaatac	atagtttatt	gagagttcag	catggagagc	tgttgaattt	103200
tgtcaaaggc	cttttctgca	tctattgaga	taatcatgtg	gtttttgtct	ttggttctgt	103260
ttatatgatg	gattacattt	attgatttgc	atatgttgaa	ccagccttgc	atcccaggga	103320
taaagccaac	ttgatcatgg	tggataagct	ttttgatgtg	ctgctggatt	cggtttgcca	103380
gtattttatt	gaggattttt	gcatcaatgt	tcatcatgga	tgttgggtcta	aaattctcat	103440
ttttgttgtg	tctctgccag	gatttgggtat	caggatgatg	ctggcctcat	aaaatgagtt	103500
agggaggatt	ccctcttttt	ctatgattgg	aatagtttca	gaagaattgg	taccagctcc	103560
tctttgtatc	tgtggtagaa	ttcggctatg	aatctctcct	ggactttttt	tggttggtag	103620
gctcttaatt	attgcctcaa	tttcagagcc	tgttattggg	ctattcaagg	attcaatttc	103680
tttctgggtt	agtcttggta	gggtgtatgt	gtccagggaat	ttttccattt	cttctagatt	103740
ttctagttta	tttgcacaga	gggtgtttata	atattctctg	atggtagttt	gtatttctgt	103800
gggattggta	gtgatatccc	ctttatcatt	ttttattgca	tctatttgat	tcttctctct	103860
tttcttcttt	attagtcttg	ctagtgggtct	atcaattttg	ttgatctttt	caaaaaacca	103920
gctcctggat	tcattgatgt	tttgaagggt	tttttgtgtc	tctatctcct	tcagttctgc	103980
tctgggtcta	gttatttctt	gccttctgct	agctttttta	tgtgtttgct	cttgcttctc	104040
tagttctttt	aatgggtgatg	ttaggggtgtc	aatttttagat	ctttcctgct	ttctcttggtg	104100
ggcatttagt	gctgtaaatc	tccccctaca	cactgcttta	aatgtgtccc	agagattctg	104160
gtatgttgtg	tctttgttgt	cattgggtttc	aaagaatatc	tttatttctg	ccttcatttc	104220

BI OL0250W0SEQ_ST25. txt

gttacatacc	cagtagtcac	tcaggtgcag	gttggttcagt	ttccatatag	ttgagcagtt	104280
tttaatgagt	ttcttaatcc	tgagtcctag	tttgattgca	ctgtgggtctg	agagacagtt	104340
tgttataaatt	tctgtttcttt	tacatttgct	gaggaatgcc	tcactttccaa	ctatctggtc	104400
aatttcagaa	taagtgcgat	gtgggtgctga	gaagaatgta	tattctgttg	atttgggggtg	104460
gagagttctg	tagatgtcta	ttaggtctgc	ttgggtgcaga	gctgagttca	attcctggat	104520
atccatgtta	acttttctgtc	tcattgatct	gtctaattgtt	gacagtgggg	tgttaaagtc	104580
tcccattatt	attgtgtggg	agtctaagtc	tctttgtagg	tctctaagga	cttgctttat	104640
gaatctaggt	gctcctgtat	tgggtgcata	tatatattagg	atagttagct	cttcttggtta	104700
aattgggtccc	tttaccatta	tgtaatggcc	ttctttgtct	cttttgatct	ttgttagttt	104760
aaagtctgtt	ttatcagaga	ctaggattgc	aaccctgtct	ttttttgttg	ttttccattt	104820
gcttggtaga	tcttcctcca	tccctttatt	ttgagcctat	gtgtgtctct	gcacgtgaga	104880
tgtgtcttca	gaatacagca	cactgatgga	tcttgactct	ttatccaatt	ttccagtcctg	104940
tgtcttttaa	ttggagcatt	tagcccatth	acatttaagg	ttaatatttt	tatgtgtgaa	105000
tttgatcctg	tcatcatgat	gttcgctgg	tattttgctc	attagttgat	gcagtttctt	105060
cctagcatcg	atggttttta	caatttggca	tgtttgtgca	gtggctgata	ccgattgttt	105120
ctttccatgt	ttagtgttc	cttcaggagc	tcttgtaagg	caggcctgg	ggtgacaaaa	105180
tctctcagca	tttgcttgct	tgtaaaggat	tttatttctc	cttcacttat	gaagcttagt	105240
ttggctggat	atgatattct	cagttgaaaa	ttcttttctt	taagaatgtt	gaatattggc	105300
tgccactctc	ttctggcttg	tagagtttct	gctgagagat	ctgctgttag	tctgatgggc	105360
ttccctttgt	gggtaacccg	acctttctgg	tgaatctgac	aattatgtgt	cttggagtta	105420
ctcttctcga	ggagtatttt	tgtggcattc	tctgtatttc	ctgaatttga	atgttggcct	105480
gcctttgtag	gttggggaag	ttctcctgga	taatatcctg	aagagtgttt	tccaacttgg	105540
ttccattctc	ctcgtcactt	tcaggtacac	caagcagatg	tagatttgg	cttttcacat	105600
agtcccatat	ttattggagg	ctttgttcat	ttctttttac	tccttttttt	ctctaaactt	105660
ctcttctcgc	ttcatttcat	tcatttgatc	tttaatcact	gatacccttt	cttccacttg	105720
attgaatcaa	ctactgaaac	ttgttcatgt	gtcacgtagt	tctcgtgcca	tggttttcag	105780
ctccattaga	tcatttaagg	tcttctctat	gctgtttatt	ttagtctgcc	attcatctaa	105840
actttttcaa	ggtttttagc	ttctttgcaa	tgggttcgaa	catccttctt	tagctcggag	105900
aaatttggtta	ttacagatcg	tctgaagcct	tcttctctca	actcatcaaa	gtcatttctt	105960
gtccagcttt	gttctgttgc	tcgtgaggag	ctgcgttcct	tcggaggaga	agaggcaccc	106020
tgatttttag	aattttcagc	tgttctgctc	tggtttctcc	ccatctttgt	ggtttatcta	106080
cctttgggtc	ttgatgatgg	tgatgtacag	atggggtttt	ggtgtggatg	tcttttctgt	106140
ttgttagttt	tccttctaac	agtcaggacc	ctcagctgca	ggtctgttgg	agtttgcctg	106200
aggtccactc	cagtccctgt	ttgcctgggt	attaccagtg	gaggctgcag	aacagcaaat	106260

BI OL0250W0SEQ_ST25. txt

attacagaac	agcaaagtgt	gctgcctgat	tcttcctctg	gaagcttcat	ctcagagggg	106320
caccagctg	tatgaggtgt	cagttggccc	ctactgggag	gtgtcccca	gttaggctac	106380
tcgggggtca	cggaccact	tgaggaggca	gtctgtccat	tctcagatct	caaactctct	106440
gctgggagaa	ccactactct	cttcaaagct	gtcagacagg	gatgtttaag	tctgcagaag	106500
tttctgctgc	cttttgttca	gctatgccct	gccccagag	gtggagtcta	cagaggcagg	106560
caggctctct	tgagctgtgg	tgggtccac	ccagtttgag	cttcctggtc	gctttgttta	106620
cctactcaag	tctcagcaat	ggcagacgcc	cctccccag	ctttgctgcc	gccttgacgt	106680
tcggtctcag	actactgtgc	tagcagttca	atctcagact	gctgtactag	cagtgcagaa	106740
ggctctgtgg	gcatgggacc	ctctgagcca	tgtgcaggat	ataatctcct	ggtgtgccgt	106800
ttgctaagac	cattggaaaa	gtgcaatatt	agggtgggag	tgtcccgatt	ttccgggtac	106860
atctgtcatg	gcttccttg	gctaggaaag	ggaattccct	gaccccttac	acttccggg	106920
tgaggcaata	tccgccttg	cttcggctca	ctctccgtgg	gctgcacca	ctgtctgaca	106980
agccccggtg	agatgaacc	agtacctcag	ctggaaatgc	agaaaccacc	catcttctgc	107040
tttgctcatg	ctgggaactg	tggactggag	ctgttcttat	tcggccatct	tgaaacctcc	107100
cctctctcac	gatcacaagg	tcccacaata	ggccgtctgc	aggctgagga	gcaagaaaag	107160
ccagtctgaa	ttccaaaact	gaagaaattg	gagtctgatg	ttcaagggca	ggaaacatcc	107220
agtgccaaag	aaagatgtag	aatattcaac	attcttaaag	aaaataattt	tcaacctaga	107280
atttcatatc	cagccaaact	aagctttata	acaaggaga	agtaaaatcc	tttacaacaa	107340
agcaaatgct	gaggaatttt	gtcaacacca	ggcctgcctt	acaagaggtc	ctgaagaaaa	107400
cactaaatat	ggaaaggaaa	aaccagtaac	agctactgca	aaaacatacc	aaattgtaaa	107460
caccatcaac	actataaaga	aactgcatca	actaatgggc	aaaatagcca	gctagcatca	107520
taatgacagg	atcaaattca	cacataacaa	tattaacctt	aatgtaaat	gggctaaatg	107580
ccccaattaa	aagacacaga	ctgggaaatt	gaataaagag	tcaagacca	ttggtttgct	107640
gtgttcagaa	gacccatctc	agggtgaaaa	gacatacatg	ggctcaaaat	aaagaaatga	107700
aggaatat	accaagcaaa	tggaaagaaa	aaaaaagcag	cggttgcaat	cttagtcttt	107760
gatgaaacag	actttaaac	atcaaagatc	aaaagagaca	aaggagggca	ttacctaatg	107820
gtaaaagtat	caatgcaaca	agaagatctg	actgtcctac	ttatatatgc	accaataaca	107880
ggagcaccca	gattaataaa	gcaagttctt	agagacctac	aaagagactt	agacttcac	107940
acaaaaatag	tgggagactt	taacaccca	cagccaatat	tagatcgacg	tgacagaaaa	108000
ttaacaagga	tattcaggac	gtgaattcag	ctctggacca	agctgacctt	atagacatct	108060
acagaactcg	acaccacaaa	tcaacagaat	atacattctt	ctcagacca	cattgcactt	108120
attctaaaat	tgaccacata	attggaagta	aaacatttct	cagcaaatgc	cgtagaatgg	108180
aatcataac	aaacagtctc	tcagaccaa	gtgcaatcaa	actagaactc	aggattaata	108240
aactcactca	aaaccacaca	actatatgga	aactgaacaa	cctgctcctg	aattactact	108300

BI OL0250W0SEQ_ST25. txt

gggtaaataa	caaaattaag	gcagaagtag	ataagttctt	agaaaccaa	gagaacaaag	108360
acacaatgtg	ccagaatctc	tggtacacag	ctaaagccat	gttttagaggg	aaatttatag	108420
cactaaatgc	ccacaggaga	aagcgggaaa	gatctaaaat	caacacccta	acatcacaat	108480
tcaaagaacc	agagaagcaa	gagcaaacaa	atacaaaagc	tagcagaaga	caagaaataa	108540
ctaagatcag	agcagaactg	aaggggataa	agacacgaaa	acccttttaa	aaattaataa	108600
atccaagagc	tggttttttg	aaaagattaa	caaaatacat	agaagcctag	ccagactaat	108660
aaagaagaaa	atagagaaga	atcaaataga	cacaataaag	aataataaag	gggatatcac	108720
caatgatgcc	acagaaatac	aaactacat	cagagaatac	tttaaaccac	tctatgcaaa	108780
taaaatagaa	aatctaaaag	aatggataa	attcctggac	acatacaccc	tccaagact	108840
aaaccaggaa	gaagtcaaat	ccctgaatag	accaataaca	agttctgaaa	tcgaggcagt	108900
aattaatagc	ttaccaacca	aaaaaagccc	agaccagagg	gattaacagt	caaactctaa	108960
cagaggtaca	aagaagagct	agtactattc	cttctgaaac	tattccacac	aatagaaaaa	109020
gagggactcc	tgccctaactc	attttatgag	gccagcatca	ttctgatacc	aaaacctggc	109080
agagacacaa	caagaaaaga	aaatttcagg	ccaacatccc	tgatgaacat	caatgtgaaa	109140
atcctcaata	aaatactggc	aaactgaatc	cagcagcaca	tcaaaaagct	tatccaccat	109200
gatcaagttg	gcttcatccc	tgggatgcaa	ggctgggttca	acataattcaa	atcaataaac	109260
ataatccatc	acataaacag	aaccaatgac	aaaaaccgta	tgattatcgc	aatagacgca	109320
gaaaaggcct	ttgataaaat	tcaataccca	atcatgctaa	aaactcttaa	taaactaggt	109380
attgatggag	catgtctcaa	aataataaga	gctacttatg	acaaatgcat	agccaatatc	109440
atactgaatg	agcagaagct	ggaagcattc	cctttgaaaa	ccagcacaag	acaaggatgc	109500
cctctctcac	cactcctatt	caacatagta	ttggaaattc	tgtccagggc	aatcaggcaa	109560
gagaaagaaa	taaaggtatt	caagtgggaa	gagagggagt	caaattatit	ctctttgcag	109620
atgacatgat	tgtatattta	gaaaactcta	tcatctcagc	ccaaaatctc	cttaagctga	109680
taagcaactt	cagcaaagtc	tcaggataca	aatcaatgt	gcaaaaatca	caagcattcc	109740
tatacaccaa	taagagacac	agagccaaat	cctgagtga	ttccatttca	caattgctac	109800
aaagagaata	aatatacct	aggaatccaa	cttacaaggg	atgtgaagga	cctcttcaag	109860
gagaactaca	aaccactgct	caaggaaata	agataggaca	caaacaaatg	gaaaaacatt	109920
ccatgctaata	ggattggaag	aatcaatatt	gtgaaaattg	ccatactgcc	caaagtgatt	109980
tatagattca	atgttatccc	catcaagcta	ccattgatit	cttcacataa	ttagaaaaaa	110040
ctactttcaa	tttcatatgg	aatagaaaaa	gggcctgtat	atccaagaca	acctaagcaa	110100
aaagaacaaa	gctggaggca	tcatgctatc	tgacttcaaa	atatactaca	aggctacagt	110160
aacaaaaaca	gcatggtatg	gtactggtac	caaaacagat	atatagacca	atagaacaga	110220
acagaggcct	cagaaataac	accacacatc	tacaactatt	ggatctttga	caaactggac	110280
aaaaataagc	aatggggaaa	ggattcccta	tttaataaat	ggtgttggga	aaactggcta	110340

BI OL0250W0SEQ_ST25. txt

gccatatgca	gaaaactgaa	actggatccc	ttccttacac	cttatacaca	aattaactca	110400
agatagatta	aagaattaaa	tgtaagacct	aaaaccataa	aaaccctaga	agacactttg	110460
ggaggccgag	gtggatggat	cacgaggtca	ggagatcgag	accatcttgg	ctaacacagt	110520
gaaagcccat	ctctactaaa	aatacaaaaa	attagctggg	tgtggtcgtg	ggcacctgta	110580
gtcccagcta	cttgggaggc	tgaggcagga	gaatggcatg	agctgaggag	gttgagcttg	110640
cagcaagcca	agattgtgcc	actgcactcc	agcctgggca	acagagtggg	actccatcaa	110700
aaaaacaaaa	acaaaaacaa	aaaatcaaac	cctagaagaa	aacataggca	ataccattca	110760
ggacataggc	atgggagaag	acttcatgac	taaaacagca	aaaccaatgg	caacaaaagc	110820
caaaatttac	aatcagatc	taattaaaat	aaagagcttc	tgcacagcaa	aaaactctca	110880
tcagagtga	aaagcaacct	atggagaaaa	attctgtggt	ctagccatct	gacaaaagggc	110940
taatgtttag	aatgtacaag	caacttaaac	aaatgtacaa	gaaaaaaaaa	acaaccccat	111000
caaaaagtgg	gcaaaggata	tgaacagaca	cttctgacag	gaagaccttt	atgtggctga	111060
caaacatgaa	aaaagctcat	catcactgtt	aattagagaa	atgcaaatcg	aaaccacaat	111120
gagataccat	ctcatgcccg	ttagaatggc	gatcattaaa	aagtcaggaa	acaacagatg	111180
ctgaagagga	tgtgtggaga	aagaggaaca	catttacact	gttgggtggga	gtgtaaatta	111240
gttcaacat	tgtggaagac	agtgcggtga	ttcctcaagg	atctagaacc	agaagtacca	111300
tttgacccag	caatcccatt	actgggtata	tacccaaagg	attataaatc	attctacaat	111360
aaagacacat	gcacacgtat	gtttattgta	gcactattca	caatagcaaa	gacttggaac	111420
caactgaaat	gcccatacat	gatagactgg	ataaagaaaa	tgtggcacat	atacactgtg	111480
gaatactatg	cagccataaa	acaggatgag	ttcatgtctt	ttgcaggggac	atggatgaag	111540
ctggaaacca	tcattctcag	caaactaaca	caagaacaga	aaaccaaaca	ccatatgttc	111600
tcactcataa	gtgtgagttg	aacaatgaga	acacatggac	acaggaaggg	gaacatcaca	111660
cacagggggc	tgttggggag	ttgaggctag	gggagggatt	ggattaggag	aaatacctaa	111720
tgtagatgat	gggttgctgg	gtgcagcaaa	ccaccatgac	acgtgtatac	ctatgtaaca	111780
aaccacacac	ttctacacat	gtatctcaga	acttaaagta	taataataat	aagatacaga	111840
actgcagaat	gaataagaac	tcaccaacca	tctgtgcctt	tcaggagact	catttaagac	111900
ataaggactc	acataaacct	aaagtaaagt	ggtggaaata	ataataagtg	gtgtcactga	111960
tgtggaggta	gattataaaa	ctcttatcat	atgctggtgg	aagatcaaaa	tgataaaacg	112020
aattaaaaaa	tcagtcagat	ggtttcttaa	aaagttccat	caatatgcct	ctatcttaca	112080
aacctgcaat	tctattcctg	aatctttatc	ccaaggaaat	gaaaaagtaa	gtccacaaag	112140
agttctatat	gaatatttat	aggagcttta	tttattataa	ttcaaactgt	aaaaataatt	112200
tcaatgttca	tcaataacaa	aatgaaaaaa	taatttgcaa	cctactggta	cacttgaata	112260
ctattcagca	ctgagtatct	taaatagcat	ggatggagct	caaaaatata	ctcaggaaag	112320
aagccatgta	tattctgtat	gagttcattt	acatgagatc	atttacattt	cctccaaaag	112380

BI OL0250W0SEQ_ST25. txt

aggaaaaact	aatttctgtt	gaaagaaacc	aatgtatttg	cctctggcag	tggttaagggg	112440
gtagcacaga	ttaattgggt	agggactcaa	gagagtttct	ggggtcacag	aaatgttccg	112500
tgtggtgatg	ggagtttggg	ctccacaggt	ataggtgttg	atccaaaatc	atcaaaaaaa	112560
caacattgca	gatctgtgca	tctcactctg	tgggaaagta	tatctcaact	gtaaaaaggg	112620
cagaaattgc	ttttaaacgc	tcagcctttt	agcacatcca	gttgcttgga	gaaccagctt	112680
actcaaatgg	gggtctaggc	tggagactag	gtcacaggca	tagagtctct	aaactttccc	112740
atggcacata	atacgtttca	ggtttttctca	gagagctgca	ggtttagtaat	ctgaggattc	112800
tgacaagttg	ggtcaacggt	cctaggaggc	atgaatggga	gtgcattctc	taagatccct	112860
ccaccccagg	gtccttgctt	tctgtgcctc	ttactccatt	gttttctgac	tcctctgtag	112920
ccactcgacc	tcttcagatc	ccattgtcta	cccagccatc	gccctttatg	acttgggtcc	112980
cactgttctt	tcattctatc	ctccattccc	tcagtttcgg	agtggctgcc	gctagcacag	113040
gatggactga	gagcaggaga	ggtggctctg	cccaggaacc	catcctagag	aaatggcatc	113100
ctgtctggga	gctagttttt	tagggcaggt	tttataagtc	ttgtaaagcc	agacacactt	113160
gatctacctg	gtatgttatt	tacagtaata	ctattttcat	aattgctttt	cactctaaaa	113220
gtagagcctt	ttagctacac	tgtgagtaaa	taaaggggct	ggcctgggaa	tggtatcatg	113280
ttggatgttg	tttcttccct	gaagtaatat	atatcagtta	caattttacat	gttactgcag	113340
agtcctagag	agagacacag	agaatgagac	agataccaat	acattttttat	gtgcattaaa	113400
aaaatctaag	gccaggcgca	gtggctcaca	cctgtaatcc	cagcactttg	ggaggccgag	113460
gtgggtggat	cacgaggcca	ggagattgag	accatcctgg	ctaacacggt	gaaaccctgt	113520
cttactataa	aatacaaaaa	attagccagg	cgtggtggcg	ggcgcttgta	gtcccagcta	113580
ctcaggagac	tgaggcagga	gaatggcttg	aaccaggag	gcagaccttg	cagttagccg	113640
agattgcgcc	actgcactcc	agtctgggcg	acagagcgag	actccgtcac	aaaaaaaaaa	113700
aaaaatctaa	aatgcactct	tcaaaatcta	tgtcattttat	tctggaggaa	tgcagttggc	113760
agaaggagga	agatattccg	aatttttctt	gtatacattt	atgtatgatc	tcagtttttt	113820
tatggatcat	agaccaattt	tgatatttta	aaataaaaaat	tataatctat	cttggaaatt	113880
tacatggttc	tttagaactt	gaggaccgtt	tttgcttttc	ggaatattat	tgtacctaaa	113940
atgggaatat	tacaacgtca	ctttttaaca	ctttgttata	acaaagttaa	gacagcgctg	114000
ggtgcccctg	aattttttcc	cgcctcttgt	gacctgtgtt	gttttggaat	ttgcagtggc	114060
ctgaccgaga	actactgcag	gaatccagat	tctgggaaac	aaccctgggtg	ttacacaacc	114120
gatccgtgtg	tgaggtggga	gtactgcaat	ctgacacaat	gctcagaaac	agaatcaggt	114180
gtcctagaga	ctcccactgt	tgttccagtt	ccaagcatgg	aggctcattc	tgaagcaggt	114240
aagaagtctg	tggccagata	tctacacatt	tgaacattgg	gatgaaaaga	gatggaaaat	114300
ctgactgatg	cagaagcctt	ccatgctaca	cagaaacttg	agggtatggc	aggtggaaag	114360
aagcctcagc	actctctctg	gtggagcaat	ttttggcgca	acgtgcgtgg	gcggtgactt	114420

BI OL0250W0SEQ_ST25. txt

caggaatggt	gcaaaccac	ctgggcactt	gacttaccac	tcactttggt	atgaaagggg	114480
ttatctcgg	gttcagaca	aaattccaat	tctaacaat	ggccaaattt	gtgccaaatt	114540
tcacactagt	gagtgtttcc	aggcatttat	taaaatggac	agtgttcatt	gcaatcttca	114600
gcattgcagt	tgctgaggt	tgtggccgct	gagtttgtca	tcctggggaa	acctaataatg	114660
atgatattta	ttccatctaa	tcctggggct	atttggcagt	aaataccaca	gaatacacta	114720
tttctctggc	ttatttcagt	cttaggtagg	ctctgcacac	ctatgcttgg	aaggcaggaa	114780
tttcttggtg	ttcttgtgcc	ttcttctcat	ggaacgtgca	tctttggtgt	gtgttgagag	114840
gaagggtagt	agacttctgc	ttgtttgcaa	tgcaggatgc	tggaacaaga	ggattccctg	114900
tctctactgt	aagggaataa	gattttagcc	tccatccttc	tctaagaagc	aatgtgtctt	114960
tgctccaag	tactagatgc	aggaccatga	actgccccgt	ccaccagaag	cttaaggctt	115020
tggcttttca	ggagcaatca	tctaggggaa	tgtgcagggt	tttcatgtct	gtcccctact	115080
gacagccaat	caccatacag	cctgcataac	ctaataccatc	atcgtctggt	ttcctgcctc	115140
attgttttca	tgaacaacca	gtagagagcc	atacgaaaga	gcttgacacat	gagtctttgt	115200
tccaattgta	agagcactga	taggtccttt	tcccaccagg	ttttgaatat	aaaatttcta	115260
agaacttatt	aaaatattag	aatgttatta	atctattggt	tttgcttcag	catgtccttc	115320
tgcttgtgag	tatactaaag	agaacagtca	taattctgaa	actactgtcc	tgtttgtgtc	115380
ataaattgct	tcacatgttt	ctgcatacta	gtagttactc	agcttgattt	tgtctatttt	115440
cagcaccaac	tgagcaaacc	cctgtggtcc	ggcagtgcta	ccatggtaat	ggccagagtt	115500
atcgaggcac	attctccacc	actgtcacag	gaaggacatg	tcaatcttgg	tcattccatga	115560
caccacaccg	gcatcagagg	acccagaaa	actacccaaa	tgagtatgtc	tttgatgtta	115620
cttgtaagag	gagcaacagc	caacttaagt	tcctcctaga	agagccttgc	ttcaagctaa	115680
cttgtagga	caaatttccc	ttagaccag	aagggtgtgtc	aaaatgtcca	gacaactttg	115740
cttttgatca	aagagtctga	gagaataggt	attttaggct	tgctatcttt	tctaatagtc	115800
tgatggaagc	agaaggctac	atggagctga	tgaggctttt	ttaatataaa	gctcaagaga	115860
tcaaatgatc	aaatacttag	agtgccattc	tacaaggctc	ataaaagatc	aatgcactct	115920
ttcacccatg	caattctatc	attctaacct	cccttctctg	aaatgaaggc	tttttgccat	115980
ttttgtcatg	ggtcacaagt	aaataattca	catgtatatg	agtatatata	taaccagggtg	116040
tgtttattca	gactagtatg	tatatatata	catatatatg	ttcatataag	ttagtattca	116100
tatatatggt	catatatata	tgttcataca	gactagtatt	catatatata	tacatatata	116160
tatacacaca	catatatata	tatatatata	tgttctaggg	aaacatgcaa	ggtttttatg	116220
tctgtccctg	actgatgacc	aaatacccta	tagcctgcac	agctgcaagc	tgtatagcca	116280
tacaatttgc	aggacacaca	cacatacaca	cacacacaca	cacacacaca	cactaacata	116340
taatataata	taatataata	taatataata	taatataata	taatataatt	aatatatata	116400
aacctgtgtg	aacacactgg	gttctaagct	ccagttttct	gaagggatat	gggttgccag	116460

BI OL0250W0SEQ_ST25. txt

gagaggaaga	gcaaaagcaa	gaatgtagat	gagaattagg	aagtaaacag	atatggagat	116520
taaaatgggc	aggtacatgg	acaaaaaacc	aggtctgaca	aaaactggct	ttctgccata	116580
aatgactata	aaagatatta	aaaaacactt	tccacatggt	ggacaagaga	cagtacagga	116640
ctgagataat	ttagaaaagg	aatgaatga	gcgcaactcc	gtaactatta	tgactttctt	116700
cctggagaac	cttcctggac	tgaagggcaa	ggaattggag	ccaaagccaa	ccacagcagt	116760
cttgctgaac	tgaggaaaga	gactggagtt	tggtatagct	aagaaaatgt	gtattttcta	116820
tgctaggtaa	taatgagaaa	gaatttgtgg	tgaaaaggag	ctgaaggaat	atgcatggaa	116880
gtctaataata	aactgcatat	gcacagggag	aaattctaca	aagtgggaca	gagaaccact	116940
actggggaaa	ggacaaattc	agggaaacag	tgagctcaat	ggtgacgcca	gagctcacgt	117000
agcactgggg	gataccgggg	ttctgatcag	cccaggagaga	gacacctcat	tgaacatctc	117060
gggcattcag	tagagacccc	agaaaagtca	tacttttagga	gtaggattta	tgcccttcta	117120
gaataaagac	taccccagaa	acaccctagt	aaagcttaaa	aaccaagtct	aaaaggaccc	117180
aaatgatctc	caagtaaatt	aactgcctga	cagaagaaaa	ctcaaccatc	actggaggta	117240
aataacatga	ttacagtgc	ctgtaatggt	gcattcacaa	ggagtgcac	catttaaaaa	117300
tttatgaggc	aggaaaaagc	aattagtgtg	atccataact	aggagaaaaa	ccagtcaata	117360
caaatagacc	aagaaatagt	agaaacgatg	gaattgacaa	agaaattaaa	actgtatata	117420
tgataattgt	gttcaaagat	ttaaagaaaa	catgaacatg	agggaacaaa	atgcagaata	117480
taaaaaaaaag	caaatgctga	aaacaaccaa	atggaaatta	aagaactaca	aaaaagtata	117540
accttaataa	aatactcact	ggatggcctt	aatattagtt	tatacattac	agaagaaaaa	117600
gtgaaccaga	agataactca	atgaaagcca	tacaatctgt	aagacacaca	cacacgcaca	117660
cgcgcgcgcg	cgcacacaca	cacacacaca	gagagagaga	gagagagaaa	gagagagaga	117720
gaaaggctga	aaaaaataaa	tagaacctta	aggatatcag	tgaaaatagc	aaaagattta	117780
atatatgggt	aaagcaagtc	acagaaggac	gggaaggaga	tattgggaca	gaaaaaaata	117840
ctcaaagcaa	tgatggctga	agactttaca	cgtatgaaga	aatgataaaa	ctcacagtca	117900
agaagctcaa	tgaatcagaa	atagtatttt	taaaagcaaa	actctatgat	ttacttgggt	117960
acattataga	taaatcgtcc	aacatcaaag	ataacaagga	taatcttata	agccagagga	118020
aaacaatatc	atttacatag	agggacagta	atgaaagtga	ccgatgcctt	ctccttggaa	118080
acaatggcat	aacatcttta	aagtataaaa	gagaaataaa	aacagatcaa	cctaggacga	118140
catgtccagc	caaaacaaac	aaataaacia	aaaaaccctt	taaaataaac	gtgatgtaaa	118200
tacgtattct	gccacctcca	gaggaaacia	gcaaaaaaac	aaaagaatgt	ttccaaggca	118260
ggcttctgta	ttaaaagatt	ttaaggaaag	ttattcaggt	agaagaaaaa	taataccaga	118320
tggaactttt	aatccatact	aagtaatgaa	gagccctgga	aatggcaaat	ggcaatgtca	118380
atataaaata	ctcttattta	tctaattttt	aatgtatttt	aaaggacaat	ttgtgatatt	118440
aattaaaata	ataggaatat	attgtttgtt	caacgtatgt	agtagtaaaa	ttcataaaaa	118500

BI OL0250W0SEQ_ST25. txt

cagtagcaca	aataatgcag	atgataactg	gaagtatact	gttaatgagt	tttttgcatt	118560
atccatgaag	ttatataata	ttaatagatg	gttgaatgtg	atagtttaag	gtgggatatt	118620
ataaatccta	ggacaaccaa	aaaaatttaa	actgagagga	atggatagta	agaggaatag	118680
tccttttatg	caaaagaagg	aagaaaaaga	ggaataaaga	atataaaaga	tatggtgtaa	118740
acagaaaata	catagcatta	ttgtagacac	aaactgaact	accttatgag	tatattaaat	118800
ataaaaggat	taagcattac	aaataaaagg	cagagattgt	aaattgaata	aaaaccacag	118860
ctaagtgtgt	tctttttaga	ataaatactc	tttaagtgtg	aagatctact	ttaaacacca	118920
aaatatgaaa	aaggatatat	accatgaaaa	cctgaatcat	aaataagctg	gagtgggtgat	118980
taatggatgc	aggcactcct	aaagactaat	aagtgaatgt	ggtcaaattg	aagaaacaaa	119040
agtatatacg	tgctcaatgt	gcaaaaactt	tttctgtata	catgctatga	tcctttggaa	119100
aattaaagtt	ttaaagcaat	atcactgaca	atagtatcaa	aaccaaaaaa	tatttagtga	119160
taaatttcac	acactatgct	caaggactat	acaccttgca	ctagaaaaca	atgttgagga	119220
aagaattaaa	agatctaaat	atacaccatg	cttatagatt	aaaagactcc	atatcagttc	119280
tcgtgaaatt	gatctttgga	tgaaaccac	accaagcac	tattgcaaca	gtcctttttt	119340
ggaaaaaaaa	attggaggac	ttatatacct	taatataaag	acttataaaa	gtacaggaat	119400
caagacatgt	ggtattggcc	tggccccttg	gctcatgcct	gttaccctaa	cattttggga	119460
ggctgagtct	ggaggatggc	ttgagcccag	atgttcaaga	ccagccttag	caacagagtg	119520
agaccctctc	tctacaaaaa	ataaacaatt	agatcgatgt	gatgacttgc	acatgtagtt	119580
tcagctactc	ggaatgctga	ggtgagagga	ttgcttgact	caggaggtct	agccatgagt	119640
gagcattgat	catgcctctg	cattccagcc	tggatgatgg	aatgagacac	tgtctcaaaa	119700
aaaaaaaaaa	aaaaggatat	gtgttattgg	ccaaaaaagt	atgcaaacct	aaaaagggat	119760
ggcccaccac	cagaccaca	tacatatatg	gtaaatggat	tttccgtata	gatggcaaag	119820
caattcaatg	gagacaaaaa	tgtttttaca	aatcattctg	aaccatttgg	atatccatga	119880
tacaaaacaa	aagcagaact	tgacttttgc	ttttcatctc	aaattatfff	gatatctctt	119940
ccacctaagt	gtcagagcta	aaactgaacc	tgaaatatga	aagttccatg	aaaaaatata	120000
aaatcttcac	aaccttggag	aaggcaaact	tttttgaggc	aggagtctgt	aaacactcac	120060
tataaaaata	aacaaattat	aatgtgggct	ttcatgaaaa	ctcatgctta	ccaaaagtca	120120
ttgttaagaa	aataaatagg	caagtaacac	atgagaagaa	aaatgctctc	tgtccatata	120180
tctgacaaat	ggcttgtgtc	cagaatatag	gaacatttct	cccactcact	aaacagagga	120240
caaacaacta	atgggcaaca	gattgaatag	gcatttcttg	gggatagata	gatgtacaca	120300
tagccaataa	gcacctgaaa	aatgtccag	tatctcagcc	atgaaaaata	aagagttata	120360
atcatcatga	gatgtcacca	aacacccaat	ggacatggat	attattaaga	agacaccaca	120420
gtaactgatg	tactgatgt	agagcaagga	tgtgaaactc	tctcatatgc	tggtgaaagt	120480
gcaaaatgat	acaaccactt	ttgaaatcag	tctgatagtt	tctccaaaag	ttcaataaat	120540

BI OL0250W0SEQ_ST25. txt

gcacttttac	cctacaaacc	tgcaatcctg	tttgtgaata	tttacccccac	agaaatggaa	120600
acataagtcc	acgaagacat	ctccaagaat	attcatagca	gctttatfff	ttataacccc	120660
aaactgtaga	caatttcaat	gtcaatcaat	aagaaaatga	ataaataatt	tgtgaactag	120720
tcatacaatg	gcatactgtt	cagcaataaa	agggagcatg	tttttgatac	tctcaaatag	120780
tatggaagat	gctcaaaaat	attacattaa	agaaagatgc	cagataacaa	aatgaacat	120840
tatgtatgag	tctattgatg	taaggttcca	gaaaggtaaa	actaatttct	ggtgaaagaa	120900
accaatatca	tttgcctctg	gccatgggaa	gagagtagca	gagattgatt	gagcagtaaa	120960
acgaagtttt	tttctggggg	gatgtaaatg	tcctgtattg	tgattgaagt	gtgagttaca	121020
caagtgtaca	tgttcatcag	aagtcacaa	actacatcta	agatctgtgc	atttgactat	121080
acatgaaaat	atacctcagt	tgaaaataga	tcaataacct	ccctcatata	ctatacttgc	121140
taacacagcc	agctgcttgg	agaaccagct	tgctggaatg	gagaatctgg	gcttgagact	121200
gggtcacatg	tatagagtct	ctacagagac	aatgttgcac	tcccacggta	cataatacat	121260
ttcaaggttt	ctcagacagc	cacatgtcat	gaatgtgagg	attctgagag	gttggagcaa	121320
cattcctggg	aggaacgaag	gggagcacat	tctccaagat	ccccaccac	cggggtcctc	121380
accggctgtg	cttttttttt	tttttttctt	gacagagtct	cgctctgtcg	ccaggcagga	121440
gtgtaatggc	ccaatctcgg	ctgattgcag	cctccaactc	cagggttcaa	gagattctcc	121500
tgctcagct	tcatgagtag	ctgggactac	agatgtgcgc	cactgcgccc	agctaatttt	121560
tgtatfifta	gtagagacgg	ggttttgcca	tgttggccaa	gatggctctg	ctctgttgac	121620
ctcgtgatcc	acccgccttg	gcttcccaa	gtgctgggat	tacaggcgtg	agccaaagca	121680
cccagcctgt	gcctctcact	tactcaattg	tttttctgaa	ccctccatag	ctgggtggacc	121740
ttttcagatc	ccatagtcta	gccagccctc	tcactttatg	ccttgggtcc	cactgttctt	121800
tcatctcatc	ccccttctgt	cagtcccgca	gtggctgtgg	ccagtagagg	atggactgag	121860
agtaggagag	gaggttctgc	ccaggaaccc	atcctagaga	aacagcatcc	tgcttgggac	121920
ctagtcttcc	aggtcagctt	ttataagtct	tttagactca	aactcacttg	accacactga	121980
agtggatttg	acaataatgc	tattttcatg	gttgtttttc	actgtaaattg	cagagccttt	122040
tagctacacg	actagtacag	agagtaaggg	aggctggcct	gggaatgata	tcatcttgga	122100
tggcatttcc	tccttggaga	aatatatgtt	agttccaact	cacatgttac	tatacagtcc	122160
tgtagaaaga	gatacagaga	gtagacagg	tatagacgca	tttgtatatg	cataacaatc	122220
tataagacac	acatcaaaa	ccgtataacc	gttcctctag	gggtatgtgc	ttggcagaag	122280
gtagaaggag	ggtattctgg	ttcctttctt	ttgcacattt	atgtatgatc	tcagttttta	122340
tatggagcat	tgatagggtt	tggctatgtc	cccacccaaa	atctcatctt	gacttghtaat	122400
ctctataatc	ctgataatcc	ccatgtgtca	agggcaggac	caggtggagg	taactggatc	122460
atgggggcag	tttctcccag	gctgttctca	tgacagttag	agagtctcct	gagatctgat	122520
ggttttgtaa	gtgtctggca	tttcccctac	ttgcacttac	tctgtcctgc	cgcctgtgaa	122580

BI OL0250W0SEQ_ST25. txt

gaaggtgcct	gtttctccct	tgccttctgc	catgactgta	aatttccaga	ggcctcccca	122640
gcaatgtgga	actgtgagtc	aattaaaact	cttttctttg	taacttacct	agtctgtctc	122700
gggtatttcc	tcatagcaat	gtgagaacgg	gctaatacaa	gcatatacta	cttttgatat	122760
tttaaaataa	aaattatcat	ctatctttga	aaggcatgca	caaattggga	gttgaggaa	122820
atttgtgttg	tggcaattgt	atgatacctt	taatgggaat	atttcaaaga	cacttggtta	122880
gactttgtta	gaacaaaatg	tagagggtgc	tggatgtccc	tgaatattct	tccgcctcct	122940
gtaacttgta	ttgctttgga	atttccagt	gcctgacaat	gaactactgc	aggaatccag	123000
atgccgatac	aggcccttgg	tgttttacca	tggacccag	catcagggtg	gagtactgca	123060
acctgacg	atgctcagac	acagaaggga	ctgtggctgc	tcctccgact	gtcatccagg	123120
ttccaagcct	agggcctcct	tctgaacaag	gtaagaagtc	tgtgtcttac	cttgtctagc	123180
acatacctct	ctatgtgctt	ggacaacggg	atgaaaagac	atgaaaaacc	acactgatgc	123240
agaagccttt	agtgtacac	gggagctcga	gtgttggttg	aggttctgcc	atgaccaagg	123300
aagtctcagt	gccgtccctg	ggaaagccag	agctgtgatt	tttggcaca	cttgtgggag	123360
tagtgacttt	aggactggcg	caaaacctcc	agggtgctca	acttaaccac	tcaccttatt	123420
ctaaaatggg	ttatttcagt	gtcccagtca	aattcctatt	ctaactgct	gtcaactgtg	123480
tgattatttc	caagccaata	agcatttcca	gtaatttctt	aaaatagtgt	tcattgcagt	123540
cttcagcggt	gtggctcctg	agggatgtgg	cccctgattc	tgtcgtccta	gagaagcctg	123600
acatgactgc	attgattctg	tatcgtcctg	ggtctatgtg	gctgcctggc	tgtctgtaat	123660
catctgtttt	atttttattt	ttttctacag	actgtatgtt	tgggaatggg	aaaggatacc	123720
ggggcaagaa	ggcaaccact	gttactggga	cgccatgcc	ggaatgggct	gccaggagc	123780
cccatagaca	cagcacgttc	attccaggga	caaataaatg	ggcaggctctg	gaaaaaatg	123840
taagccactt	tgatttggac	tctttttccc	tttctgaca	aatcttttca	aacagaagag	123900
gggcagagga	aaatactgga	aagacttcag	gaggctaagc	gtaattagcc	ttagcatgga	123960
aagtgaagc	agcacaggcc	agcaaagccc	cacgcgtgtg	ggggttctca	ggcctcttct	124020
cttttgacat	ttctttactg	tttccattgt	tgggtgctgt	ttctcgtttc	tagtgcttgt	124080
cctctaagcc	agggttcccc	actccagtac	tggactgggt	actggtactg	gaactggtaa	124140
ttatctgtgg	cctgttagga	actgggctgc	acagcaggag	gtgagcttcg	ggggagcaaa	124200
caaagcttca	tctgtatttt	ctgctgcttc	ccatcactct	catagctgcc	tgagctctgc	124260
cagctgtcag	atcagaggca	gcattagatt	atcatagcac	aaaccctatt	gtgaactgca	124320
catgtgagga	atctagattg	catgctcctt	atgagaatct	aatgcctgat	gatctgtcat	124380
gcttccatca	ccccagatg	ggaccaccta	cttgcaggaa	aattagctca	gggctcccac	124440
tgattttacc	ttatgggtgag	atgcacattt	atttcattat	atattacaat	gtaataataa	124500
ttgaaataaa	gtgcacgata	aatggaagg	acttgagtca	tcctttaacc	atcgccccct	124560
caccccagg	gcacagaaaa	attgcctttt	atgaaactgg	tctctgggtgc	caaaaaagtt	124620

BI OL0250W0SEQ_ST25. txt

ggggaaccac	actgctctgg	gttctagtag	tcagagatgc	cctctatgag	gcttaagtca	124680
gatttttcta	gaaaagattt	ggatgggcca	tcaggtcacc	atgagacttc	ccttagcctc	124740
atgcattctc	tgtgatgggt	tactttgggg	cctatgaata	gggaagactg	agatatagga	124800
aaaaccaaag	tgtctgtgtt	ccccactct	cacacccatg	taacataaca	cttctcacac	124860
cagatatggg	gggatttctc	ctcacacccc	aagcgagtct	ccagcagata	ccagctgggt	124920
gtcctacaat	gtaactcggg	cctgacactc	tatctggaga	cagtgtcaga	tcccacaagt	124980
taaggctcag	tcctacaaga	ctgccccact	gcagatgcca	atcccaagtt	gcaggctgtg	125040
acctgtactt	ctgcccagct	ggataaagat	ctgtttttct	atatgaccct	ccatgggttt	125100
gattactttg	ctagagtggc	tcacagaact	cagggaaaca	cgttactttt	atttaccat	125160
ttattataaa	agatattaaa	aaggatcctg	gtgaacagcc	aggtggaaga	gatgcacagg	125220
gcaaggcacg	tgggaagggg	ctcagagcct	ctatgccctc	tccagtgcac	cagtccccag	125280
taccctaagt	gttcagcaac	ccagaagctc	tccaagtga	gtcttggttg	gtttttatgg	125340
aggcttcatt	acagaggcac	agttgattac	atcattggcc	atcgggtgatc	ggctcacctt	125400
cggccccctc	tccctccctg	gaggttggag	ggtggggctg	aacagttcca	accctcaagt	125460
cacatgggtg	gttcccttgg	caaccagccc	ctggggctat	ccaggaaccc	accaagagtt	125520
gcttcattgc	agctcccttc	accaggaaaa	ctccaaggga	tttaggagct	ctgtgttaag	125580
aactgggggg	cagagaccca	atatacattt	cttattctat	cacaatatca	caggaagcta	125640
aggatgatac	tgcctttgtg	tgtcttggct	gtggatggtg	cataatgcat	ggaagtaagc	125700
atttctgaat	caacagcaaa	caggctttat	caggtagaag	accctcagc	gccccaggga	125760
caaagctcat	caatgatgtc	ccactgtcct	ctgaggctct	agctctaaga	cctccagtgg	125820
gtcaagctcc	tggagaagtg	gcacattctc	caaagaccct	tcagggtcac	cacaccctgg	125880
ttaaggggtg	ggcctcataa	ctccttttga	ctatgactga	tggcttacag	catagaaaga	125940
aataactttg	tcaaaaaata	taataatgat	agaaaggaag	aaggaacgct	cccttttgtc	126000
ttctaagaat	agatgtgaaa	tgtgtgtgcc	ttagaatatc	ttctccctct	cctgctccac	126060
gtgagctgga	gcttacatgc	ctgcttgttt	tcagtactgc	cgtaaccctg	atggtgacat	126120
caatggtccc	tgggtgctaca	caatgaatcc	aagaaaactt	tttgactact	gtgatatccc	126180
tctctgtggt	aagttgcctt	ctgttttggg	aaggaaactg	cttccttaat	atggatttgg	126240
aaaaaaaaaa	gcaaaaaaaaa	cagaaaatgg	cttttagact	gagtgttct	ggggaggaga	126300
tggctgccct	ctccaccaga	gcctgctttt	catcatggcc	accttgaacc	tgccctacta	126360
ttggccccat	ttgttaggaa	aacacccgcc	cctcccacca	cacacacata	aataaaataa	126420
atgtcaaatt	cccaaagggc	aaacttagag	gtgatcta	cagcccggga	tagtcccacc	126480
gaacccttct	ttgtctagcg	tgggatgcat	gaaaaacaaa	tttagagtca	ttatgatgaa	126540
aaactgtcct	cttctgcagc	tgagaagaaa	aaaaaaatac	gagcagcagg	aaacagctaa	126600
gcatgtaatg	cacattgtaa	acctcagatg	gccatcctag	gaaatcaatg	aagggtagtg	126660

BI OL0250W0SEQ_ST25. txt

cagctcttta	gccccagatg	gcctttctcg	taagattact	actcatgagt	cccattagcg	126720
acattgctta	gagactgctt	gtaggttcc	ttcctcattg	ctctgagact	cttattggga	126780
gtatgaggct	tggatcaggg	gaaggggaat	tgacattaga	tcttaaataga	ttggggtaac	126840
aaatccatgg	gggaaaaaaa	gccacttgta	cttgttccct	attttcttcc	tgctgaccaa	126900
tcaacttgtc	tgtccgagtt	acagaacacc	accctggact	tttcttttgt	gtaatttggt	126960
tgcttggtg	tggttctgcc	atgtgaagg	accttgagct	gggggaagaa	ggttggcctc	127020
caagtccact	gaagaccagc	atcctgagat	tgcttgggga	ggtggtacag	ggcagtgatg	127080
aagatcatgg	gagccacact	gcccatcgtc	acatttgggc	cactcctggg	gagagcaaga	127140
gggaagaagg	agaggttagg	gtgataggaa	agattctact	tggccaatat	tattataatg	127200
tggcattgtg	gtctctggat	ttagtgtgag	ttgatagctg	acttttttct	cgagtgggtg	127260
cttttgttct	attttgtcgg	tgctattgca	gaagcatctt	ggtggttcct	ctacctcaaa	127320
gtctcttgat	ggggtcagtt	ccagttctcc	gcttctggcc	ccatctagta	cacgccactg	127380
cctctcactg	cctgggctct	ctatccttga	caggctgcct	tgaatttaag	cccagtctga	127440
cttacctgcc	tcaaacaccc	acagtagtgc	ctgggactca	tgcacctttg	actcccatgg	127500
aaggaagtgt	cagtagcttc	ccaggtgcaa	ttctgctgtc	ctcaccaca	ttgaggatgt	127560
atgagaatca	ggttcttaga	gattggagaa	agaaggaaga	atgggaacaa	gatttcttcc	127620
aatggactgt	gaggttcccc	accttacttt	gatgtaagac	aagtgagggt	aacccaagc	127680
ctggtgagga	gggttcccat	cagacacttg	gaaatcctga	ggactgtttc	ctgcagaagg	127740
atgtggttgg	tgggatattc	aggtttgact	catgattgag	aaagttagag	cctctggttg	127800
gagaaagagt	ttaataacta	tttcatttcc	accaacacat	tcagtacgaa	taataaataa	127860
gtaaaaataa	atagaaacat	tcagttttat	tttgaatagt	aggagtaggg	tataatttct	127920
gtagttactc	ttttagtaca	atgatgcatg	tttactgtat	gtaaggcata	ctagcagaaa	127980
ttgagctcag	cactagaaaa	gatgattgca	ttccatgcc	tgcttctttt	ttacaaaaga	128040
cttctataga	tagatttctc	aaacaaccca	cagcaaataga	aaagtatttt	ggaaaactca	128100
ggttccagat	tactggagt	gtagaatctc	tggttggttg	gggaggaatt	tcctcttgca	128160
gttggttatta	ataattatat	gaataattat	taactatatt	aataatttata	gttttgaaga	128220
ccttgaaggg	ctggagacaa	cagagaagca	tttttgaaca	ccctctgtag	cccctgcact	128280
gtttagaggca	ttgatgggtg	gtaccaaaga	tgggacactt	tccctacctc	cagagacctt	128340
gtgggcttgc	tgacagagaga	aggcaggag	gaggaaaaga	agaatagagg	cacatgtgtg	128400
taaattaccc	ccacagcagt	cagttagtca	tgaggaggctc	cccagaagaa	ctgtcctgaa	128460
gctggctgag	agaaggcaac	atttcaacat	aggacagtta	tccttgctac	ataaaatcac	128520
atacacacat	gcacatatgt	ccacacacag	agactcacat	gcaaaagaat	cctttgtgcc	128580
tttcagtaaa	ctttacatgg	tttagaaaga	acttatattt	ccttgaaagg	agagtgtcct	128640
ttgttgttta	ctaccacttt	ttaaacttag	aaagaaaaat	ctaaagagtg	tttatgattt	128700

BI OL0250W0SEQ_ST25. txt

taccatttaa	tttcaccttt	gagatgtgaa	aaactagtagc	ttggaattcg	tcctgaatta	128760
aacgacacaa	ttgctaactt	ggactcaa	gcgacttctt	ttccacactt	gtgccacagc	128820
atcctcttca	tttgattgtg	ggaagcctca	agtggagccg	aagaaatgtc	ctggaagcat	128880
tgtagggggg	tgtgtggccc	accacattc	ctggccctgg	caagtcagtc	tcagaacaag	128940
gtaagaacag	gcccagaaac	catctatact	gtccttccat	gtaagcccca	caaaaccctt	129000
ctacatttac	acagaaccca	cacagctgat	gcacaaatc	ctgcctctct	gttttctgaa	129060
ggaggaaaaa	atatagaaaa	attaaaaaaa	gttatattat	tataggttct	ctacttggaa	129120
aatagccaaa	atacaaatct	ttttcttgat	ctgggcagtt	ccatcaaaat	ctgtaggcac	129180
agtgatttgc	accaagttcc	aatacttttg	gaaaatattg	aagatgtctt	gagggtttct	129240
atggatatcc	attgtctcac	tgtcagatga	aaagaaaggg	aagtttttag	aaatgtgaca	129300
ctttgcagtg	aggaggagca	agagcaaact	tacctacagt	ctatcacagg	cacagatttt	129360
tttttacact	tttgtgaatc	attgaattca	atgccgaggc	tattcatcta	ttcacaacaa	129420
catgaacaaa	ttatgggttg	tgatcccat	aaatgaagag	taatcagtc	gaaccacag	129480
aacctggaca	ttttgggtat	cgtttcagtg	gaacatgcaa	ttcgtaagtt	cagtttgctt	129540
gggtgtctct	taggaagaac	acataggaca	cagacccatc	tgccctgcatg	ttttgcttcc	129600
tcatctcctt	tctacaccag	ggcacctgtg	ctcaattgct	gttctcctct	aaagagactt	129660
ccttctgtaa	gtttgtgaaa	tgccatcgac	aaacctgac	gcacgcatt	tcactctgct	129720
gttgagtga	ttttcttta	ctttatcggt	tgtaacttct	tgctctacag	agctttcacc	129780
ttccacatat	ttcagattca	ttctttccta	aactgtgtgg	tggtctatgt	cctcactgac	129840
tatcaacata	ctgccatcat	gcacttccta	tctctattcc	tcttcgttgc	aatctggctc	129900
caagtggctc	acaccattat	tctgatctat	caactgccta	cacagtccta	gaaagtaagt	129960
gagtcaagaa	acatcccca	aaagtaaact	tttcaggtaa	gatcagaaga	ccctcatgag	130020
tcactgctgc	tcaggatcgt	atctggctcc	ttgaagagtg	accttgcata	gatcttgtca	130080
taaaaaatga	aagagacctt	gggaaggtct	tgggctggct	acttttgtca	gagtccaggg	130140
ctgtgggggtg	aaagccacag	ctatagagct	tcattctgga	gtcacttagc	tttgctctcc	130200
tggggacagg	ctgtgcctat	tcttgcccta	ggcatcaaaa	aaagtggcac	agatggggcc	130260
ttctgaaaaa	tctcactact	ggagcacagc	tcgaagtttc	tactatcctg	acgttggggc	130320
gtagtccttt	gctttgggaa	tatgaacatg	atcaaaactg	agtgaacttg	tcttcctggc	130380
tttctgtaca	atgaagtaga	acaaaccatc	caatttgacc	aaagccttgg	catgttttct	130440
ttctaggttt	ggaaagcact	tctgtggagg	caccttaata	tccccagagt	gggtgctgac	130500
tgctgctcac	tgcttgaaga	agtacgttta	agggaact	gacatggggt	cttatcttca	130560
agactttttt	cctccctctc	ttcctccatc	ccttctttct	tcccaccctc	cccttccttc	130620
ctccccacct	ctcttccttt	tctggaagga	acactaggaa	ccagggaaatg	catgcagaat	130680
cctgaggcag	aatttcagg	gcaattggat	gagagaggag	ggaagtgttt	ctagagggaa	130740

BI OL0250W0SEQ_ST25. txt

tctgcagagg	gaagacccag	tgcaagtgat	tttttgacc	tgtataaacc	gcaggacaga	130800
gctgttact	accagaggca	tcaatctgta	ttgcattgct	ctagagcaat	atctgaggct	130860
gaataattta	taaagaaaag	agtttaattg	gcacatgttt	ctgcaggctt	tacaggaagc	130920
aggatgctgt	catctcctct	gcttctgtgt	gggcctaagg	aagattacaa	tcatggtgga	130980
gggcaaagt	ggagcaggca	tgtcacatgg	ccagagcagg	agcaagagac	agagagagat	131040
ggggtggggg	tgctgcacaa	taccaaata	ccagactttg	caagaactaa	gagtgcagagc	131100
tcactgatca	ccatgaagat	gtggcccaag	ccattcaaga	gggatgcacc	tctatgatcc	131160
aaaccccttt	cacaggccat	agctccatca	ctggggacta	cagttgaaca	cgagatttag	131220
gtggggacaa	atatacaaac	tatatcacag	tctctgatga	aacagattga	gaacagacct	131280
taactgtcag	tttccagcaa	attgtgaatt	ttgtttcttg	ccactcataa	gtcactgatt	131340
ctgggtggcc	gaggtgtca	gaggacagc	gccaaattca	tggcacagag	gatacctgaa	131400
ggggtgggac	catatctttc	tcttgacatc	ctcatctttt	ctaggtcctc	aaggccttca	131460
tcctacaagg	tcctcctggg	tgcacaccaa	gaagtgaacc	tcgaatctca	tgttcaggaa	131520
atagaagtgt	ctaggctgtt	cttgagccc	acacaagcag	atattgcctt	gctaaagcta	131580
agcaggctact	cgctcacctg	tggctctcac	cccacgctgg	tgaagatatt	tgctttatgt	131640
ctgggtttta	tgggcatgg	ccactgcatg	gcagtgggga	ggaactgtct	atcacatgaa	131700
aggctcaagg	gctttgggga	cagcatcaat	cttcaacccc	agccctgcca	catgttagtt	131760
gtgctcttta	aaaaggcaga	aggattcggt	tcctcacgtg	gaaaaagaga	taccctgtta	131820
cccgtaaaac	ttacttaatg	ttcaccagtt	catccacatt	catgatcagg	gaaaggttgt	131880
tattccaggc	taactattct	cctttcataa	taatatgctg	gagagaaatca	aatgagattg	131940
catttcaaag	cgcttgaaaa	accaccatat	cgagccatgc	ttagtgtggg	cgcttcta	132000
cactgctatt	caggaggctg	acgaggaaga	attgcttgag	cccaggactt	caaggctgta	132060
ggcagctatg	attgtgccac	tgcactccag	gctgggtgac	agatcaagac	cctgtctcaa	132120
caaaagaaaa	gaaaacaaaa	caaatgaaca	gaaatattcc	acaatgtcaa	aaaaaaaaaa	132180
aaccacaca	acatacaatt	tacaaatgca	aataataata	ttattgttgt	cttctttgat	132240
tttctctttc	ctgggtgaaat	ttgtttttat	taagcctgac	aaagtgatac	ctttgcttac	132300
atcacttaaa	gttagtctat	ttggacctag	gtgacagtac	aatcagctaa	gaaacagtat	132360
ttgtaggaga	ggcagggttg	ggacagggtga	caaggcatgt	ggggtgctcg	ctgtgctggt	132420
ggctctggaa	ggcagggtgt	caatgcagac	agggatgagc	atggcctggg	tgggaaggca	132480
tggggcaggc	aggagcctga	gctgctctcc	tgggcctggg	cacaagccca	tggcagcttc	132540
tctgggtctg	tgaactgagg	ggtgatgtcc	tggaaatcctc	tgacactcta	ggaaggagag	132600
aagggccttt	ctggctcagc	ctttataaac	agtagctgat	ctccctcttg	ctccccaggg	132660
tcctccccac	catcccagca	aatgtgcaaa	tacaagatct	ctgctcctca	tggtcctcag	132720
agagctgggg	tgttctgatg	gcttgaacaa	gtcacttagg	aaatgtgggg	ttttggaggc	132780

BI OL0250W0SEQ_ST25. txt

attctctgat	aggctgatac	gttttgagtt	tagagttccc	accgcacatc	cccacacccc	132840
tagagtctag	ggcatttagt	gctccatgag	ggaacctgta	gagtgaggac	atctgcatca	132900
caggctgggc	cttctagtgt	ccagaagcag	aaagtgtgtc	tgcttcaaag	ttggtgctaa	132960
tgatgatttt	tggtcagaat	acggcatttc	tcatttccat	tcctttatcc	ccttgaactt	133020
actaaagtag	aatcagggtct	aaaaaccaga	gttctaattct	ttaagagtcc	ctgggattct	133080
aaggtatatg	aatgtccttg	gaaaacaata	ccatttagtt	catgcaaggt	gcttattttcc	133140
catcctcttt	catttgatgt	ctagcatttt	actgcattct	taccaccacg	gtttagtaac	133200
attcacgagg	aggaagtgga	ggatccagat	ggagcaactt	gctctgggca	cacaaggcat	133260
ttgcaatttt	ataccctctt	gatgatgtct	cagccagaca	ttctgcccag	tcatcaatgc	133320
cctcttcaat	taatatgaaa	ggacacactt	ggcatgagat	tccaatcgtg	cacagaatat	133380
acatgagaag	tgtgcctttg	tcatccctac	tttcaaaggc	taaggccacc	ctcagtttct	133440
tgcatgcaac	tgatgccttt	caaatgaaac	cttacatctg	tgtagtccat	aggcaaccac	133500
aggcaaatgt	gagggtgaaa	cgctgtgttc	tacattgttc	tgtgtcagtg	aagcaaggca	133560
gtgccagctc	agagggctct	ggggcttcaa	ggcagggatg	cctggttgta	ggtactgcca	133620
cttcagctg	ggcagtgaia	cataactgct	aatactttcc	ttacaggcct	gccgtcatca	133680
ctgacaaaagt	aatgccagct	tgtctgccat	ccccagacta	catggtcacc	gccaggactg	133740
aatgttacat	cactggctgg	ggagaaaccc	aaggtgagat	caattccatt	gccacgtaa	133800
caaattgttt	ttgaccttca	gtgcatgtta	caaaatgagc	atitttgaga	tagttgtaca	133860
aattcctacc	catgaatgtg	gtctaccac	tcctgacttt	gcctggacac	ctgtctatgt	133920
ctccataatc	agtcttcaag	ggacttgggc	aaggggagcg	gtgccatttc	cttgagtctc	133980
tctctttttt	gttttcagaa	tcttttaatt	ttttttgtaa	tgattgtatg	tttcccttac	134040
aacaaaaaca	aacaccagta	gaggtctttg	agtctcttaa	tcataatttc	agcattcata	134100
ttgcttcccc	aggtaagtgg	ggttttgacc	cagccctcaa	gttaagggtg	ttagattatt	134160
tttcatgtga	aattagacag	actgcgtttc	taaacatggt	gcaaaacagt	aacgacaaaa	134220
gttgtaatta	aactattctt	cttcccaa	atccacatgt	ctaattgtgtg	tgtgaggggtg	134280
ttaggcaggg	gacctgaagc	tgggggagag	gcagacagtt	cccatggccc	caagtctagg	134340
atggcatttg	gtattgggtg	atgggtgaga	gcaagagagg	gaatattttt	gtgcatgatg	134400
tggatatcagc	acctgtacta	cattttatgg	attccttctt	ctctttgcgg	tatgccctga	134460
caataattat	atccgtcagc	cttaccctt	tggcagtagg	aaaactgaaa	ctgtcttaaa	134520
gtctcagctc	tactttctca	gaggtgcagg	caagggcact	gggagtctgg	ggccctggaa	134580
aactgttctg	actctgccac	ttgccagata	gacctgaact	agacacgtta	cctctttgta	134640
ccacttggct	ctaattccctt	atctgtaaaa	ccagcatttt	caaattggtgc	tttgacatc	134700
agccttttgc	ataagctttg	atttgataaa	atgttttttg	tgtttttaaa	aagattaaaa	134760
accacaggtt	tagataattt	caaagtaggc	ttcccttttt	ctgtcatttt	cctattattt	134820

BI OL0250W0SEQ_ST25. txt

ttaaaacctc	acctccttga	ctccttggtc	cctttttctg	cactgctgag	tctgggagca	134880
ctgaggccag	gtaaaaggaa	acttggcaaa	tgagggggcac	ctatgggtgt	gggaggctgc	134940
tcctgggtgtt	tgcatatttt	aaaattttaa	tgctacaaac	cactgtgagt	taggtattat	135000
tgttcctatt	ttaccattga	ggaagctggg	gctcagagaa	ggtggagggt	ggtacagaca	135060
aacctgaatt	ggaaccctgg	ctcctgccta	tggtgtgtca	ggacttagaa	aagtcgtgag	135120
ctctcgctga	ttgtttcctc	agctgatgtg	ggctgcaggg	ctgttatggg	ggaaataata	135180
agaaagtgca	tcaagtgtct	agcacatcct	aagcactcca	tcatggcagc	tcctactact	135240
aataaagaat	agaattatat	ctaocatgat	tctttcttgc	aagtgcagag	aaatccaact	135300
caaattggat	taagcaaaac	aagggaatt	cttagtgagc	tgcaaagttt	tcaggctcac	135360
atgatggccc	caaatcccag	gtcctcccaa	tcatggagta	ggcactatit	gggggcacaa	135420
aggtgacatt	cccatggctg	cagatgctgt	ggtgctgtgg	ctgtaccggg	aaagaataag	135480
aaaggccact	ctcccaatta	tgtgaacaat	agtctgccca	ctctgagaag	tcaaacttgg	135540
gtcacagtcc	tgcccctgaa	cccatcactg	actggctctg	acctgcacca	attgttccat	135600
gttggagggtg	aaggcaagac	cccactaata	cccataaggg	gcaaaagtta	gatagatcct	135660
tcaagaggat	tatgggaggt	agggcaaaaa	gctgctgggc	agccagaaaag	caaacagagc	135720
ctctatgata	cctcaactga	tgaaagcatg	aagctaaaat	cataaggatc	tgggtgtgag	135780
ttctggctct	cccatcttcc	atgtgacatt	gggcagttat	ttaatctctt	ttagcctccg	135840
ctttctcatc	ttacatatga	gataattgtg	aggattaaga	ttacacataa	tcatcatcat	135900
caccgtccac	cactaccacc	atcatcccca	tcaacatcat	cgccaccact	atcatcattc	135960
ttactggcac	taccatcacc	atcaccacca	ttccaccacc	atcaccaata	tcatcactgt	136020
caacatcatt	accaccatca	ccatcaccac	caccatcatc	attactacca	ctaccactac	136080
taccaccatc	accatcacca	ccattccacc	accatcacca	atatcatcac	tctcaacatc	136140
atcaccatca	ccatcaccac	caccatcatc	atcattacta	ccactaccac	tactaccacc	136200
atcaccatca	ccactgtccc	actactatca	gcatgacatc	accatcacca	ccaccatcat	136260
cattaccacc	gctactacca	acatcaccat	caccacaatt	ctactgccat	caccattaac	136320
attaccacca	ccatcatcac	tatcaccatc	accaccatca	tcaccactgc	cattatcact	136380
gccaccatca	tcactatcct	ctatatttcc	tcatctgtat	tatcattact	accaccatca	136440
ctatcaccac	catcgtcacc	atcataatca	ccatcaacac	catctccaat	accaccatca	136500
ctgtaaccat	catcaccacc	accatgatca	ctatcaccat	catcacaatg	atcactgtaa	136560
ccatcattac	taccaccac	catcaccact	actccaccac	catcaccatt	atcattacca	136620
tcaccattat	caccaccatc	atcatcacca	gcaccaccat	catcaccagc	accaccatca	136680
ccatcaccat	cattaacacc	atcactatca	ccattggttt	aatcatcacc	accatcatca	136740
taaataaaca	tcacataacc	aggggtgtagc	tgggtgttga	ccccagagcc	cactcactgt	136800
ttcctctctc	ccaccccat	ccacacattt	ctaaccacca	tcctgcactg	ggctcccagt	136860

BI OL0250W0SEQ_ST25. txt

ctcctctggt	ctcaccacaca	tgtccactga	gaaaaggatt	ttcagaacac	caactagacc	136920
aggaggagcc	acatacataa	ctcaggcctg	cttatcaact	ttctacatgt	taataatgac	136980
atcagatcaa	tgggtgttct	cagcttctca	gaaggaggtc	aaaattctcc	ccctctcccc	137040
ttcatgtgtc	cagaccttcc	cggatttggg	tgtaccaagt	gcagagtggg	gttgaggcca	137100
aggggctcat	ccatgtaagt	ctcatctgca	atcactgggc	tgatcccgtg	gccctgtctc	137160
cagggcgcca	tcagagaggg	cttcaatcct	caggttacct	gtggcccacc	ctgccctcag	137220
aggtgccatc	tctacattgg	ccacgagatg	gcagcacata	ctcatagact	gcattaattt	137280
cccagcaact	cctggtgggt	tttccctctt	atcaggatgt	ttgccttgct	cagagagcaa	137340
atctgagagc	agtacacct	aacttaactt	tcagcaaaat	attttgagaa	gggtgcccct	137400
ttacacatct	gtgcagtcca	ggtgatgcat	cccatgccca	atgctcggtg	gtcaggagga	137460
gcttcctcca	tgcagctctg	cggaagagac	tcttccacgc	tgctcatgta	aactccagat	137520
tcggtgtcag	ttttctgaca	ccgaagacaa	tgatctaagt	gcagtcaagg	gctttgggga	137580
aagcaggaga	gagtgcctca	gttctagcct	gtgccatgct	tgcaaagttt	tgcaaaattc	137640
taatgagagc	tgggcttgca	acattggaaa	cttggattat	ttgtgagagc	actgagaaat	137700
ccctgggcat	gtccatctgg	aaaaacagca	tttccctctg	cacttttagca	gaggttctgt	137760
ttcaatttgg	cgaaggaaat	taagcagttt	ttcacaaaag	aagaactaca	acgaggagaa	137820
ttgtccctag	tatttcttct	ccctaattgt	caaggaagtg	taaattagaa	aatgaatcag	137880
gacaatttcc	acctactatg	ttagctaata	ttttaaaaat	tgaatatcac	aagggtgagg	137940
caaagtaatt	gttttccagt	gacattttcc	actgtcacac	ccttttagag	aataatttgg	138000
caatgttact	gtgagataga	aatatgtcta	tataattatg	ggaactgaga	cttcagaaag	138060
taataaggaa	taagaatgaa	atttatgaac	aaacatgtgg	aaggttggaa	gcaagagtgg	138120
ggccaacacg	catggggagg	aagcatttgg	gcagcgactc	cgagaccca	gactcaagct	138180
gagctataca	acctccttac	gcctcagttt	cctcaactga	agaacaggaa	tgacaagtgc	138240
ctgtttcata	ggaccgttgt	gaggattaag	tgagatatac	cacattatga	gcttgtgcct	138300
ggaaagggtg	attcttagta	aatgatgact	attctttttt	attgcaataa	aatttataca	138360
acatagagtt	actattttaa	ccatttttgc	aggtaccact	gagtggcatt	cagtacattc	138420
acaatggtgt	gcaaccgtca	ccatatttcc	aggacatttt	tctcatcccc	aaaggaaacc	138480
tcatgcccac	taagcagtca	ctcctcatta	aaatattagt	tatgaagact	gtagcatttt	138540
tttaaaaact	catgatataa	cattgattga	aaaaatcagt	ataggaaatt	gtgcattatg	138600
atgtaatagt	aaaagaagca	tataaaaatc	tgaaaaaagt	atataaaaag	aatagcaatt	138660
gtatttctca	gactctcttt	acattgtaaa	aatcattttg	atagcttcaa	aagaaaagca	138720
aaaagtacac	aaacaacaac	caaccccaaa	gcagcatgac	aaagcccaga	ttgttgaatc	138780
caggtcttgg	gaacataaaa	tcttatatga	catttgcact	ttaatgggtc	agagagtcca	138840
gtggcattgg	gagctgcctt	gtgttctgca	gcctcacgga	cagacaggag	gtccagctcc	138900

BI OL0250W0SEQ_ST25. txt

actgctctgt	tcttctggaa	tttcctcgtg	aacaagcttt	ggcctcagta	accatttctt	138960
tcatcttttt	aaacacaggt	acctttggga	ctggccttct	caaggaagcc	cagctccttg	139020
ttattgagaa	tgaagtgtgc	aatcactata	agtatatattg	tgctgagcat	ttggccagag	139080
gcactgacag	ttgccaggta	agaaaagatc	aatagatcaa	agtcttgtgc	tctcccgtct	139140
cagtctcagt	cccttagacg	tcagtcccaa	agtggcaaata	tcaggaaggt	tttgtcagtg	139200
gaagaccca	gtctaagtgt	tgctcagaaa	ctccccagat	ctgtccctga	atgcatattc	139260
agatcatcta	aggagacgtc	ttggggcctt	agttccagat	ccatagcaag	ggagccgtaa	139320
gtgccataac	tacctcaggc	cactcacctt	cctgggtgtgt	gctggtcacc	agtgactgaa	139380
gtgggtggctt	ttccagtaga	gaggaaggta	gagggtacag	gaccgagaca	aattacacac	139440
acttaacaat	gatgtccagg	ctagcccagt	ctaaaggaaa	caccaagtta	ggaagcaatg	139500
catgcaggat	tcacaaggga	ttatTTTTTT	tcccaggaaa	aaactaagtg	atgtggTTTT	139560
gttgaataga	ctttgctaag	tacttaagca	ctgcagatgc	ttgagtaata	tgctcataag	139620
ttcctttctg	atttgaatta	ctgggaaaat	gtacatatgg	ataagagaag	gatggcatcc	139680
catattaaaa	ggttggcagc	ttaaagctca	catgaatttt	cccctacctc	tgtttagggg	139740
gacagtggag	ggcctctggt	ttgcttcgag	aaggacaaat	acattttaca	aggagtcact	139800
tcttggggtc	ttggctgtgc	acgccccaat	aagcctgggt	tctatgctcg	tgtttcaagg	139860
tttgttactt	ggattgaggg	aatgatgaga	aataattaat	tggacgggag	acagagtgaa	139920
gcatcaacct	acttagaagc	tgaacgtgg	gtaaggattt	agcatgctgg	aaataataga	139980
cagcaatcaa	acgaagacac	tgttcccagc	taccagctat	gccaaacctt	ggcatttttg	140040
gtatTTTTgt	gtataagctt	ttaaggctcg	actgacaaat	tctgtattaa	ggtgtcatag	140100
ctatgacatt	tgtaaaaaat	aaactctgca	cttattttga	tttgaattaa	ttttggTTTT	140160
ggtcttcaaa	attttcatgc	tcttttcata	ccatctatTT	ttatTTTTat	tttttagact	140220
ttacgtcctg	gggtacatgt	gcagaatgtg	caggtttgtt	acatagatgt	acacgtgcca	140280
tggtagtttg	ctgcacccat	caacctgtca	tctaattcgg	tatttctttt	agttctatcc	140340
ctcccctagc	cctccacccc	ttgacaggcc	cagggtgtgtg	atgttgccct	ccctgtgtcc	140400
atgtgttctc	attgttcaac	tcacacttat	gagtgagaac	atgccgtgtt	tgTTTTtctg	140460
ttcttgtgtt	agtttgctga	gaatgatagt	ttccagcttc	atccatgtcc	ctgcaaagga	140520
catgaactca	tcctTTTTta	tggctgcata	gaattccatg	gtgtatatgt	gccacatttt	140580
atccaatcta	acattgatgg	gcaattgggt	tggttccaac	tctttgctat	tgtgaatagt	140640
gccacaataa	acatacgtgt	gcatgtgttt	tcatagcaga	atgatttata	atcctctggg	140700
tatatacca	gtaatgggat	tgcaagggtca	aatgggtgtt	ctggtgctag	atctttgagg	140760
aatcaccaca	ctgtcttcca	caatgggtga	actaatttat	gctcccacca	acaatatcaa	140820
ggcattccta	tttctccaca	tcctctccag	catctgttgt	ttcctgactt	tttaatgata	140880
gccattctaa	ctggcatgag	atggtatctc	attgtgggtt	tgatttgcat	ttctctaata	140940

BI OL0250W0SEQ_ST25. txt

atcagtgatg	atgagctttt	ctcatatggt	tggtggctgc	ataaatgcct	tttttgaga	141000
agcatctgtt	catatccttt	gccactttt	tgatggtgtt	gttttttct	ggtaaatttg	141060
tttaagttct	ttgtagattc	tggaattag	ccttttgta	gatggataga	tggaataat	141120
tttatcctat	tatgtaggtt	gcctgttcac	tccgatgata	gtttcttttg	ctgtgcagaa	141180
gctctttggt	ttaattagat	ctcatttgtc	tattttggct	tttgttacca	ttgcttttag	141240
tgtttttagt	atgaagtctt	ctcccatgct	atgtcctgaa	tggtattgcc	taagttttct	141300
tccagggttt	ttatggtttt	aggttttgca	tttaagtctt	taatccatct	tgagttaatt	141360
tttgataaag	taatgccctt	ctttgtctct	tttgatcttt	gttggcttaa	agtatatattt	141420
atcagagact	agaattgcaa	tccctgcttt	tttttttctt	tttgctttcc	ttttgcttgg	141480
taaatattct	tccatccctt	tattttgagc	ctatgtatgt	ctgcacatga	gataggtttc	141540
ctgaatacag	cacaccaatg	ggtcttgact	ctttattcaa	tttgccagtc	tgtgtctttt	141600
aattgggggc	atttagtcca	tttacattta	aggttaatat	tgttatgtgt	gaatttgatc	141660
ctgtcattat	gatgctagcg	ggttattttg	cccattagtt	gatgcagttt	cttcatagtg	141720
tggaatggcct	ttacaatttg	gtagtttttg	cagtggctgg	taccaattgt	tcctttccat	141780
gttagtgct	tcgttcagga	gctcttgatg	ggcaggcctt	gtggtgacaa	aatctttcag	141840
catttgcttg	tctgtaaagg	attttatttc	tcctttgctt	atgaagctta	gtttcgctgg	141900
gtatgaaatt	ctgggttgaa	aattattttc	ttttagaatg	ttgaatattg	gccccactc	141960
tcttcgggct	tggtgggttt	ctgcagagag	atccactggt	agtctgattg	gcttcccttt	142020
ccgggtaacc	caacctttct	ctctggctgc	ccttagaaat	ttttccttca	tttcaacctt	142080
ggtgaatctg	acgattatgt	cttgagggtg	ctcttctcga	ggagtatctt	tgtggtgttc	142140
tctgtatttc	ctgaatttga	atgttggtct	gtcttgctag	gttggggaag	ctctccttga	142200
taatatcctg	aagagtgttt	tccaacttgg	ttctattctc	cccatcactt	tcaggtagat	142260
caatcaaagt	tagatttggg	cttttcacat	agtcccatat	ttcttggagc	ctttgtttat	142320
tccttttcat	tctttatcct	ctattcttgt	cttcttgctt	tatttcatta	agttgatctt	142380
caatctctga	tatcctttct	tttgcttgat	cgatttggct	attgatactt	gtatatgctt	142440
cacaaagttc	ttatgctgtg	tttttcagtc	agatcaggct	atttatgttc	ttctctaaac	142500
tggttattct	acttagcaat	tcagtgaacc	ttttttcaag	gttcttagct	tctttgcatt	142560
gggttagaac	atgctgcttt	agctcggagg	attttgttat	tatacacctt	atataatagc	142620
ctgatataac	tataagattt	ttttgtaagc	accatcgtaa	ccacaaagca	aaaacctaaa	142680
gtagatatac	aaaagataaa	aaggaatcaa	agcataccac	tagagaaaat	cacttaatac	142740
caaataaaga	tacgaagagt	ggaataaagg	aacgaagggt	ctacaaaaca	accagaaagc	142800
aattaacaaa	atggtgatag	cagatcttac	ctataaataa	ttatcttgaa	tggaatgga	142860
ttaaattttc	caataaaaag	acatacagt	gccaaataga	ttaaaaaata	agatccaact	142920
atatgatgcc	tataacacac	tcacttcacc	tgtaaggact	caaacagact	gaaagtaaag	142980

BI OL0250W0SEQ_ST25. txt

ggatggaaaa	aatattctat	gcaaattggaa	acaagaagat	agaggggtag	ttatacagat	143040
tgagtatcac	taatccaaac	atctgaaatc	tgaaatactc	caaaattaaa	aatgttttaag	143100
tgccaacatg	atgttcaaag	gaaatgttct	tcggagcatt	ttggattttt	gtgttttaggg	143160
atgcaaaaac	agtaaata	taatttgtat	tagtccattc	tcacactgct	ataaagaata	143220
ctacaaagag	actgagtaat	tataaaggaa	agatgtttta	ttaactcaga	gttccacagg	143280
cttaacagga	agcatggcta	aggaggccac	aggaaactta	taatcatggc	ggaagatgaa	143340
ggagaagcag	gcaccttctt	cacaaggtgg	caggacggag	tgtgagtgtg	tgaaggagga	143400
actgtcaaac	acttataaaa	ccatcagatc	ttgtgggaac	tcactcactc	tcacaagaac	143460
agcatagggg	aaaccgcccc	catgatccaa	ccccctccca	ctgggctcct	cccttgacac	143520
atggggatca	tgagggttac	aattcacgat	gagatttggg	tgggacacag	ccaaccata	143580
tcataatgca	aacattgcaa	aaacaattca	aaattcaaaa	catttctggt	ttcaggcatt	143640
ttggataagg	gaaactcaac	tcaacatgag	gtaaagcaga	ctttaagtca	aaaactgtaa	143700
aaagagacga	agaagaatgt	aataataagg	agatcagttc	attacaaata	tatagcaatt	143760
ataaatatat	attaatatat	atacccaaaa	ttgtagtacc	tacatatagt	aactaaaaca	143820
aacattaata	gatctcacag	gagagctaca	ctgtaatata	atcatagtag	cacacttgaa	143880
tagctccact	ttcactaatg	gacagatcat	ccagacagag	aatcaatatg	gaaacacgag	143940
acttaaacta	cacttttagcc	aagtagacct	aacagaaata	tatagaacat	tccatccaac	144000
agcagtagaa	tacacattat	tctcaagtgc	acaggggaata	ttctccagaa	tagatcatat	144060
gtaggtcac	aaaactagtc	aaaaaatgta	agaagattga	aatcatatca	ggtttttttt	144120
ttagatcata	atcgtatgaa	actagaaatc	aataatgggg	gaatattgga	aaatccacaa	144180
atagatagaa	attaatcaat	atgctcctga	acaatcaatg	agtcgaagaa	gatattaaaa	144240
gaggaaattt	taaaaaatca	agacatgagt	tcatgtcctt	tgcagggaca	tgaatgaagc	144300
tggaaaccat	cattctcagc	aaactatcat	aaggacagaa	atccaaacac	cgcatgttct	144360
cactcatagg	taggaattga	acaatgagaa	cacttggcca	cagggcgggg	aacatcacac	144420
accagggctt	gtcagggggt	gggaagctgg	tgaagggata	gcattaggag	aaatatctaa	144480
tgtaaataac	gagttgatgg	gtgcagcaaa	ccaacacggc	acatgtatac	ctatgtaaca	144540
aacctgcacg	ttgtgcacat	gtaccccaga	acttaaagta	taataataaa	aaaagaaata	144600
tttgtttttg	atttatatgc	caatcagaca	aaatgtgaaa	agccctactg	aaattaagta	144660
tcacatgaa	agataaattc	tggataattt	tttcaagttt	taacaatgta	gctttaattg	144720
gagaaagcta	tcatttggaa	tgagttaatc	tatcctatac	taaaataagt	cacttgcttt	144780
aaaacataat	aaatatgatt	ttgaattgaa	aacaaaaaca	actcaagaca	aaggaaaatg	144840
gacacactaa	cataccaata	atttatagta	tgcagcaaaa	gtggttttta	gaggggaagct	144900
tttaccaata	aacacttcca	ttaaaaaaga	agatctcaaa	taagcaacct	aagattacac	144960
ctcaacaaac	tagacaaaga	actaactaac	ccaaaagtta	gtagaaggaa	agaaataata	145020

BI OL0250W0SEQ_ST25. txt

aagatcacat	cagaaatagt	aaagactaaa	aaactgatac	caaaaagaaa	taaaactact	145080
agttggtttt	caataaaata	acaaaattga	ccaactttta	gctagattaa	gaaaaacaga	145140
gaatactcaa	ataaaaaccag	aaagaggaga	cattacaata	gatactacag	aagtacaaac	145200
gatcataaga	gactactatg	aataattaca	tgccaacaaa	ttggataact	tagaagaaat	145260
ggatgaattc	ctagagcaaa	aaacctacaa	agactgactc	agaaagaaat	agaaaatctg	145320
aacagaccaa	taatgtgtac	atgattgtat	cagtaataac	aagtctccca	tcaatgaaaa	145380
ggccaggacc	taatggcttc	actgctgaag	cataccaaac	attacaaaga	ctaatatcaa	145440
ccctcctcaa	actcttctta	aaaactaaaa	agaaggaatg	ctttcacatt	cattttatga	145500
ggatagcatt	acactgatac	taaacacaga	aaaataatac	gctaataaaa	gaacattaca	145560
ggcaatatcc	ctgataaaca	tatgtgcaaa	aatccgcaac	aaaatactag	aaaactgaat	145620
ccagtagcac	tttaaaaaga	tcattcacca	tgatcaagtg	cgatttgttt	cacgaatgca	145680
agaatagttc	aacttacaca	aataaataaa	tgaaaggatg	gatgataaaa	atgtgtatct	145740
atatatatat	gttttataca	cacacacaca	cacacacaca	cacacacaca	cagaggaata	145800
ttattcagcc	ttaatgaaga	agaaaatcct	gcctttgcat	caacctggag	gacattataa	145860
taagtgaat	aagccagaca	cagaaaggca	aatactgtgt	gatctcgctt	acatatggaa	145920
tctaagaaa	tcaaattcct	agaaatagag	agtagcttag	tgattgccag	agccgtggaa	145980
gggggaaatg	gagagatgtt	gatcaaagga	tacaactgta	tagctttgca	agataaatag	146040
gttctggaga	tctaattgtc	agaatggtga	ctagagttaa	taatactgta	ttgcatactt	146100
gaaatttgct	aaaagagttg	atcttaagtg	tcctcaccat	atacacaaaa	gtattatgtg	146160
aggtggtgaa	tattttaatt	agcttatgat	aataatttca	cagtgtacat	ctatatatta	146220
gcattacatt	gtacatctta	aatatatata	atttttattt	gtgaagtgtg	cctcaataaa	146280
actggaaaaa	ataattgaaa	agtaatgaaa	aaaattaaaa	gctattatgt	gtcaaatgac	146340
attatcaaga	aagtgaaaag	caacctactg	atgaagcaaa	cctattgaca	aaggcctggt	146400
gtccagaata	tattaagatc	tctaggctgg	gagcagtggc	tcacacctgt	aatcccagca	146460
cttggggagg	ccaaggtggg	aggatcactt	gagcctggga	gttcgacact	gcagtgagct	146520
atgattgggc	cactgccctc	caggctgcgt	gacagagtga	gactgccatc	tcttaaccca	146580
cttcttattt	agaaaaagaa	aatatgtagc	ttgctgcctg	catagtattc	ttggggcaaa	146640
tgggaaatga	gttaaaaaaa	aaaaaaagaa	ctcttacaac	tacaacaata	aaagaaaaac	146700
aagaacgtga	atagacattt	tttccaaaaa	agatatataa	ataggcaata	agtacatgaa	146760
atgatggtca	acatcattag	tcattaagaa	aatgccataa	aatcacaaat	gaaataagac	146820
ttcatatcca	ttaaaatgtc	tataatttaa	aaaatggaaa	ataacaagca	tttgtgagga	146880
tgtggagaaa	ttagaatcct	gtatatgtct	ggtgggaatg	tacagggaaa	atggtttggc	146940
cactgtggaa	aacaatttga	cagttcctta	aaatgctaaa	catagaatta	ccatgtgatc	147000
taacaatttt	actcttaggt	gtatatatac	aagaattgaa	aacaagtgcc	caaacagata	147060

BI OL0250W0SEQ_ST25. txt

ccttgc	catga	gaatgtt	cat	agcagc	actg	ttacaac	agc	cacaccc	aaa	tgtcaat	caa	147120	
tagatg	aggg	gataa	acaaa	ttgtg	gttta	tacagc	taca	aaaagga	aatg	aagtact	ggt	147180	
atccg	ctaca	tggct	gaaac	ttgaa	agcaa	gggct	gggat	ggggtc	atgg	aaagtac	cag	147240	
cttatt	gggt	actgc	attgt	gcttt	gggg	catgaa	aatg	ttttg	gaact	ggatg	gaggt	147300	
ggtggt	tgcc	aatgt	gaaca	tactaa	atac	aacgc	attgt	tcacta	tataag	actgct	actt	147360	
ttctt	atgag	aatttc	actt	caattaa	aaaa	atacct	tcca	tgtatc	ccttt	ctaagg	atga	147420	
tactaga	aata	tttgct	tttg	caaaat	gagg	aagta	acttt	ttttaaa	aaag	gaagat	gtgg	147480	
gatcc	atgaa	acggg	atcaa	atatc	agaga	ggaaag	gggg	tcttct	ggat	gacagt	ccat	147540	
ggagat	ccca	caactg	caca	gcagg	ccgg	tgtgc	accca	ggccac	acca	gagcag	agcc	147600	
ggtggt	tccc	gaggag	ctct	ctgga	agaaa	aacgc	tagat	ggcct	gattg	gtttg	ggggc	147660	
atattg	aaaa	ggtat	atac	tgaga	attttg	gagtg	gaatt	aggaa	acaga	cataaa	agct	147720	
tacagaa	ag	aaaata	atga	attct	aggga	gaaata	ataa	aggata	actac	aggcct	cagt	147780	
tacataa	aca	ctgaat	at	actta	accaa	aattac	aata	taattac	ata	attat	tttag	147840	
gtacat	atgg	caaaag	gatg	tgtgg	gtgta	tgtag	tatgt	acgg	tgtgtg	aagt	gtatgt	147900	
gtgtg	gatg	tggac	ggtat	gtgtat	gctg	tgtat	gcaa	taaa	atcaca	atgaa	ataag	147960	
acttcat	atc	cattaa	aatg	tctata	at	aatgt	ctat	aatttt	aaaa	atgga	aaaaca	148020	
cttctcat	at	ggcagg	agca	ggagca	aggg	tgggg	gaggt	accac	acaca	cttaa	acaac	148080	
cagatct	cct	gagaac	tcac	tatcag	gaga	acagc	acctg	gagaag	gtgc	taaacc	attc	148140	
atgag	ttact	gccct	atgag	ccaat	cacct	cccat	cagac	ccgcct	tcca	cacta	aggat	148200	
tacaatt	tga	cttgaa	at	gggcat	gaac	acagat	cgaa	accata	tcaa	taggta	atga	148260	
ctaaa	actga	aaaaa	gaagt	accac	agtca	gaaag	ttatt	tagag	agctg	aaggta	aatg	148320	
ccaat	aggat	cagtt	gaaag	aattg	gaggt	ggccg	ggtgc	ggtgg	ctcag	gcctg	taatc	148380	
ccagc	acttt	gggagg	cgga	ggtgg	gtgga	tcgcc	ctgag	gtcagg	agtt	tgagac	cagc	148440	
ctggc	caaca	tgg	tgaacc	cagtct	ctac	taaaa	ataca	aaaatt	tagcc	aggcct	ggtg	148500	
gtggac	gccg	tagtcc	cagc	tactca	agag	gctgag	gcag	gagaat	cgct	tgaacc	aggg	148560	
aggtga	aggt	tgcagt	gaac	cgagat	ctgt	ccactg	cact	ccagc	ctggg	tgacag	agca	148620	
aaact	ccatc	tcaaaa	ataa	atgaa	ataaa	gaattg	gaag	tgtttg	cctc	tggaga	gaag	148680	
gaaac	gcagt	aattct	gtaa	aaacaga	act	ttttact	tttt	tttct	ttttt	ttttt	ttttt	148740	
tgagac	agag	tctc	ttctg	tcacc	caggc	tggagt	gcag	tgg	tcagtc	ttgg	ctcacc	148800	
gcaac	ctctg	cctct	tgggt	tcaag	caatt	cccgt	gcctc	agcct	cccaa	gtagc	tagga	148860	
ttacag	atat	gggct	gctat	atccag	ctaa	tttttt	tttt	ttttat	tag	agatga	agtt	148920	
tcacc	atg	tt	ggcca	atctg	gtctca	agct	cctgg	actca	tgatc	ctcct	gcctc	ggcct	148980
tccaa	attgc	tatgat	taca	ggtgt	gagcc	accat	gcctg	gacaga	actt	tttgact	cctt	149040	
taaact	atgt	gcatat	ataa	agctga	ttta	aaaaaa	acca	agtaaa	ataa	ttttaaa	atg	149100	

BI OL0250W0SEQ_ST25. txt

ttccaaaaca	gattggatgg	gtacacactt	catcatgagt	ggttgagggg	gactgggtta	149160
gagatgagga	aattccaggg	actggggaaa	agttaaaatg	acaaactgtt	cacaattgtt	149220
aactgcaggt	tgtgggaaa	ttggttaagt	gctacagtgt	ttgttcctc	tgtaggtttg	149280
catatatatta	acatttctta	aattagcata	ataatgaact	gtgtaatcag	ctgtagagtt	149340
gaggggtgtgg	agctggcaca	ggacagctga	gctactgggt	taaaataaat	gacatttaaa	149400
aaaatggcta	ttttagaagt	taacagatat	aagacaccct	gatcaaggga	tgataagaaa	149460
ggactccagg	gctctgtctc	agctgtcttg	gcaacacctg	gaagacatgg	gcctctgcaa	149520
ggtctcatac	tttcaggagg	tgttgatgaa	ggatatggac	agatctgaag	ctctgggcac	149580
tgcattgtct	gagaagagaa	gctccggaaa	cgcgaggagt	gagtgcagat	gcagaagggc	149640
tgtcatccag	cagaggggta	ggtgacaact	ggcctagcga	gtgaccctta	tcatggctac	149700
atttgttgat	cactttcttt	gtatgaggca	ctgctgtgat	tgatttaa	ttccacttac	149760
ctaaatccaa	cgttgtgcac	ttgtgaattt	ctactcttac	aaaaaacaca	acggcaacaa	149820
cctcaaacca	gtaatctagt	caaaaaagca	attccaagg	catgacattc	agattcatca	149880
gcactcacag	agactacagt	gattgctgat	aacgccaact	taatacctgg	ccaacagcat	149940
ggatcctgac	ctccactttt	cttgtgtgtt	tacagaacca	caaaaagggtg	cagtgttttc	150000
a						150001

<210> 3
 <211> 138001
 <212> DNA
 <213> Homo sapiens

<400> 3	
ctctcccaaa	ttgtcaaaga agtataaatt agaaaatgaa tcaggacaat ttcaacctgt 60
tagattagct	aatatttaaa aattgaacac tcatacaagt gtggtgaagt gattgttttc 120
tagtgacatt	ttacactgtc ataaccttct agaaaataaa ttggcagtgt tattgggaga 180
cagaaatatg	tctatataat ttatgggaac ttaggctcag aaaatattaa ggaataagaa 240
tgaactttat	gaacaaagat gtggaggggt ggaagcaaga ggggggccaa cgcgcacggg 300
gaggaagcat	ttgggcagtg actccgcaga cccagggtca ggttgaacta gacaacctcc 360
ttacacctca	gtttccttaa ctgtagagca ggagtgatgg aactgcctgt ttcataggac 420
tgttgtgagg	atgaagtgag atacaccaca ttataagctt gtgcctggaa aggataatgc 480
ttagtaaatg	atgactattc ttttttattg caataaaatg tacacagcgt aagagttact 540
attttaacca	tttttgcagg gtaccaccaa gtggcattta gtacattcac agtgggtgtgc 600
aaccatcatc	atatttccag aatattttcc tcatcccaa aggaaacctc atgctcatta 660
atcagtagct	ctccttttaa atattagtta tgaagatcat agcactatac aaaactcatt 720
atgtaatgtt	gagtgaaaaa atcagggtgt gaaattttgt gatatgatgt aattagttaa 780
agaagcatac	aaaaagtctg aaaatataaa aacaatagca attgcatttc tcagactcta 840
catttaaaca	ttattcttta tggtttttaa agcaaagaaa aaggtaaaga aacaacaacc 900

BI OL0250W0SEQ_ST25. txt

aaccgcaaag	cacatgaca	aagctcagat	tgtaaatacc	aggttttttg	aacatagact	960
cttatatgac	gtttacactc	tccaggggttc	agagagtctg	gcagcattgg	gagctgcctt	1020
gtgttctaca	gcctcacgga	cagacaggag	gtccatcacc	actgctctgt	tcttctggag	1080
tttccttggtg	aacatgttgt	ggacgtagtt	accatttctt	tcattctttt	aaacacaggt	1140
acctttgggg	ctggctttct	caaggaagcc	cagctccctg	tgattgagaa	tgaagtgtgc	1200
aatcgctatg	agtttctgaa	tggaagagtc	aaatccactg	agctctgtgc	tgggcatttg	1260
gctggaggca	ttgacagttg	caaggtaaga	aaagatcaag	agaccaaagt	tagtcttggtg	1320
ctctcctgtc	tcagtctcag	tcccttagac	ttgagtccca	aagtagcgaa	ttcaagtagg	1380
atttaataca	tggaagaccc	cagtctaagt	gttgctcaga	aactccctag	atctgtccca	1440
aatgtatatt	cagatcatcc	aaggggactt	cttggggctt	gagttccaga	tcagcagcaa	1500
gggagccata	agtgccataa	ctacctcaga	ccactcacc	tcctgggggtg	tcccggtggc	1560
cagggactaa	agtgggtgatt	tttctggtag	ggaaggaggt	agaggggtaca	ggacagagac	1620
taactgcaca	caatatctga	gactggagct	cagatattgc	tgatgatcag	agttggcgtg	1680
tctccccaat	tgattttaca	ctgggggttg	gatactgttt	taaacgggag	gagcctccta	1740
accatcttga	cacaaccact	gacgtgacta	cactagagat	agactctttc	cacttaattc	1800
taccactctt	gctttacttc	atgagaacga	aaatgtaaga	ttgcaccatg	aattcatttg	1860
cggaaagatt	gatactatgc	ttttatttta	ttttatttta	ttttatttta	ttttatttta	1920
ttttattgag	actctcacc	cggttgaagt	gcactgacgt	gattttggct	cactgcaact	1980
tccacctcct	gggttcaagt	gaatactcca	gcctccctag	tagctgggat	tacaggtgcc	2040
caccaccacg	cctgggcta	ttttgtattt	ttagtagaga	tgggggtttca	ccacattggc	2100
ctggctggtc	tcaaactcct	gaccttgtga	tccacctgtc	ttggcctccc	aaagtgtctg	2160
gattacagag	ttgagccacc	gcactcgacc	ctatgtttta	tttttaaaaa	tattttattta	2220
tttatttaag	ccacaactac	tagaatagga	aggattgata	ttttattaat	tttatttggt	2280
atttattatt	tttttttctt	tcctgagaca	ttcttgctct	gtcaccagag	ctggagtgc	2340
gtggcacatt	cttggctcac	tgcaacctcc	atctcctgtg	ttcaagcaat	tctagtgcct	2400
cagcctactt	agtagctggg	atgactggca	tgtgcctcca	caccagcta	atttttgtat	2460
ttttttaga	gacaggggtt	tggcatgttg	cccaggcttg	tctcaaactc	ctggcctcag	2520
gtgatccatc	tgccgtggcc	tccaaaatg	ctgggattat	aggcatgagc	caccaccccc	2580
tcctggaagg	attgatattc	tataacataa	tttataatta	cagaaaacat	gtgagttcac	2640
taggaataaa	taaattttga	agataataaa	agattttcac	ttatgttgct	atttcggcac	2700
agtttggtat	aggatgtgga	gatgttaaca	tttataccta	gcttgctcgt	aaactaagac	2760
ctgaaagggt	tgtgtctatc	agctgcaccc	ctgggtagcg	acacaacctc	gggaaggcct	2820
cagccccctc	ctcgtacagc	actgcctgtt	ggaaagcttg	agggaggcta	tggatgtgca	2880
gcacttggca	gaggggtctg	tcattggaagt	taccagcaaa	tatgagctac	ttttatgatt	2940

BI OL0250W0SEQ_ST25. txt

ttattttatc	caaaagaaag	agaatgaaag	aagaggggag	gaaacaagac	taatcaggaa	3000
agatgaaggt	ctaggggtga	gggaaggagt	aaggagacat	aaaggcaatg	tggagcagct	3060
gaggggggaa	atggctttca	ccacttccca	gcattctattg	acattgcact	ctcaaataatt	3120
ttataagact	ctatatcaa	ggtaatgttt	gaaccctgct	gagccagtgg	catgggtctc	3180
tgagagaatc	attaacttaa	tttgactatc	tggtttgtgg	gtgcgttttac	tctcatgtaa	3240
gtcaacaatg	tcctgggatt	gggacacact	ttctgggcac	tgctggccag	tcccaaatg	3300
gaacataagg	aagtggttct	tctacttctt	ttatttctga	aatcaggtaa	gacatagttt	3360
ttttaaat	taagaattat	tttttctccc	acaatgtagt	aaaaatacat	atgccatggc	3420
tttatgtgca	attcatttaa	tttttgattc	atgaaattcc	cagttcaaaa	tcttgtatat	3480
gattgaaaa	ttcttaaaaa	aataagttaa	atttccccgt	gaagactgtc	acgggtgctgg	3540
aatgaatggg	cagaaaaaat	aatggttgat	ttttctaatac	taaaagagt	tgccatagct	3600
atggccagtc	tggctgaaaa	ataaatagcc	attgtagcta	actatgcaaa	ggatggctaa	3660
gctcttcgct	tggttctcag	tttcattaat	ttatatcatc	tctgttcagg	tgccatgctc	3720
ccctcactag	caagttgaaa	caatgaaata	actctttgaa	tatgtttggg	tccttgacct	3780
gttcatggag	tgggactcag	catttctctc	tttgttatgg	cctgagtaag	gctttccatc	3840
ggtatacatt	tgcttcttat	ccctggagaa	attatacaca	tccatttgcc	agatgatata	3900
cgcataata	gattcaacaa	atactcaggg	tatttgttga	gtggggttagg	tccccacatt	3960
tttatacata	catacacaca	tacacaccgt	gtgtgattgt	gaatgtaagt	gtgtgtcctt	4020
tacaaatact	agcttattta	gctcatggta	taggtagggg	agcatagtca	tccccatttt	4080
ataaacaag	aaatctagac	ttaggaaaat	catgttattt	gtctcgtgac	caaattccca	4140
aatcaaggaa	ataaagaaac	ctggatttaa	gccagatttc	caagaaaaaa	tctagggctc	4200
ttctcacttt	ttcatctttg	ttccaacatt	tgaaaaaata	aatctaaaca	cattccaatg	4260
taactgaaga	gcagggtta	tgtttgccac	ttgcagaatc	caattaagaa	gagagaagtc	4320
tggataaaag	aaagtgattt	gcttccaaag	ctagcttagg	ggaagaaatg	cagcagtcct	4380
gccgtactac	ttacttttag	gagcagaaa	tggcactttt	aaaaggcaac	agaggaggcg	4440
agcaaggatt	caggggtcca	tgctagcttg	ggcaccttat	ccaccaggta	gttgagcagt	4500
tgctgtctgg	tgcttttgtg	agcaggggtg	tgtcccttga	ggcaaatctc	tggaggggtga	4560
gagttttgta	gtgggcatgc	tttggtttat	aaatcacctg	tgaactcagg	agttccatct	4620
tgaagcacat	acatagttag	atgaacttgc	cctgcaggga	gagtcctgatg	aaaggagggt	4680
agatgcttgc	aatttaattct	ataaattacc	agataaaatt	ttacaagttg	actttaaagt	4740
caaacacatt	tgaatttagt	ggaagccatt	caagaaaata	tcaaagaaaa	tacagagcag	4800
gagaagatta	agcaaagagt	tttttgggga	aattgggtgtc	tatgtctgtg	tgtgtagggga	4860
gtgcagggga	tatgaatatt	ctatttcagc	ccatggaaac	taggatgtag	atcactgtga	4920
acttattcag	caggctacac	ccaaaggcta	gaacaaactt	ctctgccaca	ggattaacat	4980

BI OL0250W0SEQ_ST25. txt

atgttttaaat	cgacctgggg	ggcacattct	ctgataagct	cttttgga	gccaggcttt	5040
ctgtggacgt	gttatctttc	caatgtgtgc	tggaaatgcc	ggggagagga	aaaagtttct	5100
tttacagcca	tgctcagtga	gaagcggaga	aacatcttct	attcacaaat	tgctaagtct	5160
tttacacatg	caaatatgca	tacacattca	cacaccacag	tgaggaagaa	attctcacac	5220
cattaataaa	atacatttac	ttcagtagca	atatacatct	acattttgcc	tataatataa	5280
aagtatTTTT	cctattaaaa	gatttgTTTA	atgtttcttc	accaacaaat	aaaccctatt	5340
aaatcccat	tgccatatga	gccctggagg	tgaatcagag	aaacaaaagg	attgtggaaa	5400
aatcatcagg	ttaaaaaaag	aaaaattgat	tctgttttgg	gatatttcct	agcaacatga	5460
gctggggagg	ggatctcagc	agtgatgctc	tatgaagcat	aataaaatga	cacagttaca	5520
ggtaacttag	ttaaaggggg	aaataaatgg	aagtttcctc	tttttgaata	tcaattgtag	5580
cctgctctgc	tacatttcaa	aaacactctt	caaaatgttt	aactgaactc	actgtaggaa	5640
gcaccttatt	aatttattgt	gtgttttgaa	gtcacactgt	gagctataga	atttacccaa	5700
gcacaactct	tcctggaaaa	gagagttcaa	atgagaaaca	gtgcgggggtg	aagacatgga	5760
tatgggccta	aaatatctat	ttctcaatga	tattttgata	tatctatcaa	gtgcttttta	5820
gtggattagg	ttcagaatgc	atcagccaat	gcctgttcaa	taatccagtt	ttccagcata	5880
gagcatatta	aattgaggaa	ggacaaagtc	acagagggtg	ggagcagggtg	gactgtggcc	5940
aaggactttg	catgaaacag	tgagcgtgca	tcctcctcct	tgccctgccc	tcatggctctg	6000
tgtactctca	ggaggctcagg	acaggccttt	ctgagaatga	gaatctgttc	atctgccttt	6060
ctactggata	cttgtcatcg	gcatacaaac	acatgttctc	tgcagtgtgt	catctttcag	6120
aacctcccct	gaccctgtat	tccttagaag	tctcgctgct	ttcagagcca	ggcttctctc	6180
ctgctgccac	ccccactgct	cttctagtca	ctctttaacc	cactccatct	gcattgtggcc	6240
cccaccacac	ccctcaaagt	ggtcaagggt	gtcctgttgc	ttaattccat	ggaagcttgg	6300
ctatcttcat	tttattagcc	tcttttggcc	tctcaccttg	tgaaaatcac	tacattttgt	6360
gccagagatg	gagctggcat	ctccaggctt	ggaagagggc	tgctgaagct	cagccagggtg	6420
tcctaaggag	cctcaggaca	ggggatgctc	agtagccttg	caatgggaac	acagctgagc	6480
cccacttggc	caccctttgc	cacaaccagg	cagaaagcag	cttttgaaca	gatttgttgc	6540
ctcagatttg	atctcaaaga	aaaatcgtgg	gcagtattgg	tcccaggttc	tgctttttta	6600
caatttcctc	tgaaatctgg	atgcctatca	acaccttggg	aaaactgaat	tctccccaac	6660
taatagtgg	gtgtcactgt	agtaagccta	gtacaaaaat	ggccttcttt	gtggaggagc	6720
ttcatatcct	ccattttttt	tttgcttaat	ttttgcccaa	gatgagaaca	taatttagtt	6780
cactttttat	ttattcccaa	catcatccat	gcaccaacat	ttttgtaact	aaaggaggga	6840
ccattcagaa	gatgcttata	aactgtcaaa	gtgacagtgt	tacaaccaat	gcacatatgt	6900
taagaaatca	aacaatggcc	tccaagggtc	atttctacac	agggattagc	agatcaacat	6960
caatcttggc	aacacagttg	ccactgatgg	tgtcttattt	tttttatcat	gacatggcaa	7020

BI OL0250W0SEQ_ST25. txt

tcaagagcaa	acatgattta	ttcttattta	agattttatg	gtagactag	gcagatagct	7080
agatatgagc	aggaggtgga	agcccctgag	agaatggagg	tctggagaat	ctgaaacccc	7140
agagattacc	caagtcctgc	atgctagaca	tgagtggagg	agggggaata	cctaggtaga	7200
aaagaatgcc	ccttaagatg	cccagcagtc	gctcactgtg	cagttaactt	ttcagaatgc	7260
tgctagatac	atgctgatag	ggaggggaaga	gggcaaagga	gaaattccta	agagatacac	7320
ggttgacgtt	agtatacatc	tgagtgctat	acaaccttct	ttgggtggtg	gcaagaagca	7380
atgcagccat	tacgtagaat	tcatatcaaa	cacctgtatc	acaggtgtta	aagaaacaag	7440
aaacattgta	cttcttgtat	tcttaataat	gatttgcaat	attgtcttta	gtatcactgc	7500
aaacctctat	aaatatgatt	tttaaaaagt	atttcttttag	gttggaatta	cttctacgca	7560
ttgacttatc	ttcctggggt	tcattagccg	taccctgtgt	actttcttcc	ttaccactgt	7620
ttatctcaaa	ctcttgagat	taaagtatgg	gctcaggagg	gagcgaggag	cttcaggact	7680
ctcacggacc	tccagcacag	tgtagctgcc	ttatggaaaa	gtggccacac	tgttttctgc	7740
actggtccct	gcccctacta	ttcctcactg	ggcagagcac	agccaccctg	gccctgcctg	7800
aacatttttag	tcagtgtttg	ctctgtgctt	ctctggggag	gaaatccaag	agacaaccca	7860
cagcccctct	gccatttcag	ctgcagcagt	accaccgtta	atgcccttgg	gcttgagaaa	7920
gaagggacct	ggccacttcc	ctgacacctc	cagcacacag	cagggaaaga	attccagttt	7980
ctctttcttg	tgagctttca	cctgtacttc	ttcaccaggc	aaggctcctg	gcttggggccc	8040
acagtgcagg	cacctcgaac	tcagttgaac	atttccactg	gctgcactct	gtgtttttgt	8100
ggggtgaagc	tcccagaggt	gactgaaagt	ccttctgccca	ctaacactgc	agtcatactg	8160
cccttgctgt	acttggaacta	gggaaggaaa	aaagatcctg	agtgcctttac	tcacacccca	8220
gtgtgccccca	gccaccctat	ggaaaagagg	ccagtgtgtc	atccctgcaa	gcaccctgag	8280
gcccctgccc	ctgctgcccc	caagctgtag	agccagaata	taaagctggc	agaaaaatgt	8340
aaaaaggcta	gactggctta	gcctcccagc	ctacatcttt	ctcctgtgct	ggatccttcc	8400
tgctcttgaa	catcggactc	caagttcttc	agctgtggga	cttggactgt	cttccttgct	8460
cctcagattg	caggtggcct	attatgggac	cttgtaatct	tgtgagttaa	taccacttaa	8520
taagctcccc	tttgtgtgag	tatatctata	tctatagata	gatataggta	tactcactat	8580
atatacacat	atatacatat	actctctctc	tctctctctc	atatatatat	atatataatc	8640
tcctattagt	tctgtccctc	tagagaaccc	cgactaatac	agattttcat	accagaagtg	8700
gttcttgagg	aacagaatat	taaggatgga	attctttcat	tggttttggg	acttctggtg	8760
ttggctgatt	aatatgatta	gaccaaaaaa	tgctaaggac	tctacttcta	atagtatgga	8820
gaacactgat	agtacttggc	ctgaattgtt	tagagagtta	tgcaaaaataa	atgcatttga	8880
cactactgat	tcatcactta	tgagaggcaa	ggagtttagt	gactctatac	ataatacctt	8940
tgactatatg	tggagaacca	aggaacataa	tgaagttggt	tgattgctcc	taagttctct	9000
ggagaaaagag	atgaaagaaa	atgatgatct	caggggatct	gtctcccacc	ttcagaagca	9060

BI OL0250W0SEQ_ST25. txt

gatactgagc	cacaaatctg	ctaagattgc	cctgaatgag	agttttaact	cctgtagaga	9120
aagagttgaa	attgtgaaaa	aacagagaca	agctgttatc	atgcgagtag	ctgatctgca	9180
acaagaggtg	catgcacagc	cttgccaggt	gtttactgtt	aaagtgaggg	cattgactgg	9240
aaaaaaatgg	gaccctggaa	cttggagtgg	ggatgtgtgg	gagaaccctg	atgaagctga	9300
ggacactgag	tttgtgaact	ctgatgaaac	ttttttgcc	gaagaaacag	tttccccatc	9360
cccagtagtg	gtaacatccc	ctccctgacc	cgtgctgcc	ttagcctttc	cacctttgtc	9420
tgaggatgta	aaccctgcac	tgcttgaggc	aacagtgatg	gccttccctg	aggcagctgc	9480
caggcaagat	aatgttgatt	ctcctcaaga	ggcaccctta	atgcccctga	atgcttctag	9540
acctataact	aggctaaatt	ccttgcgggc	cccagaggtg	aggttcagag	tgtgacccat	9600
gaggaggtgc	attatactct	aaaagaactg	cttaagcttt	ctaatttata	ttggcagaaa	9660
tctggagaac	aggcatggga	atggatatta	agggttaagg	ataatggtgg	aagggaacata	9720
gagttggatc	aagctgaatt	tattggtttg	gccctactaa	gtagggattc	tgcatttaat	9780
gttgacgctc	ggggacttag	aaaaggttct	gatagggccg	ggagcagtg	ctcacgcctg	9840
taatcccagc	accttgggag	gcgggggagg	gcagatcacg	agatcaggag	attgagacaa	9900
ttctggctaa	aatggtgaaa	ccccatctct	gctaaaaata	caaaaattag	ctgggcatgg	9960
tgatgcgtaa	ctgtaatctc	atctacttgg	gaggctgagg	caagagaact	gcttgaacct	10020
gtgaggcaga	gattgcagtg	agccaagatc	gccccactgc	attccagcct	ggtaacagag	10080
caagactcca	tttccaaaaa	aaaaaaaaaa	aaagttataa	tagttttattt	gcttgggttag	10140
ctgaaatatg	gattaaaaga	tggccaatg	ttagtgagct	ggaaatgcct	tggtttaatg	10200
tagaggaagt	gatccaaagg	cttagggaga	ttaggatggt	ggagtggatt	agtcacttta	10260
gacctactca	tcccagctgg	gaggggtccag	aagatacacc	cttggccgaa	gctttgtgaa	10320
atagatttgt	gagagcagca	cctgtatttt	tgaagagccc	gtaattgctc	ttctctgtat	10380
gtcagatcta	acagtaggaa	ccacagtcac	tcaactacaa	aatttaaata	caatgggaat	10440
aattggatcc	tgaggtggca	ggggccaagt	gttggcactg	aaccatcaaa	ggcaaggtgg	10500
gcataactac	cataatagac	agcagaggca	aagcagccat	cagaatagtc	tgactcatgt	10560
agagctctgg	cattggctaa	ttaatcatgg	tgttcctaga	agtgaatttg	atgggaaacc	10620
tactgtattc	ctacttgatt	tatataaaca	aaaaactgcc	aggtagaatg	gactaaagac	10680
taatctgaat	tataaaaaca	gagaatcatg	ggccctcaat	caatttccag	actcgaacct	10740
gttacagttc	cagaaccac	tgaatgaagg	ggaggctgga	tccccttgag	gaaggacacc	10800
actaggctac	tgacaactta	tgctgttact	ctttctccca	tccttcccta	aggagacctc	10860
tggcctttta	ccagggtaac	tgtgtgtact	ggagaaaggg	aagtaatgag	acatttcaga	10920
aagtactgga	cactggctct	gagctgacgt	tgattccagg	gtacccaaaa	cgttattgtg	10980
gttccccagt	taaagtaggg	gcttatggag	gttaggtaat	taatggagtt	ttagctcatt	11040
tctgacttac	agtggttcca	gtgggtccct	ggacttatcc	tctggtcatt	ttcccagctg	11100

BI OL0250W0SEQ_ST25. txt

caaaatgcat	aatttgtata	gacatactta	ttagctggca	gaaatgccac	attggctccc	11160
tgactggtag	gatgagggct	attatggttg	gaaaggccaa	acagaagcca	ttagagctgt	11220
ctctacctag	aaaaataaaa	aaatcaaaaa	caatatccca	tccctggagg	gactgaagtg	11280
attagtgtca	ccatcaagga	cttgaaagac	gcaggggtgg	tgattcccac	cacatccctg	11340
ttcaactctc	ccatttgacc	tgtgcagagg	acagatggat	cttggaataa	gatggtggat	11400
tattttaagc	ttaaccaagt	ggtgactcca	attgcagctg	ctctaccagt	tgtggttttg	11460
ttgcttgagc	aaattaacac	atctcctggt	gcctggatat	cagccattgg	cttggaagtg	11520
ggctttttct	ccattcctgt	ccataagacc	caccagaagc	aatttgcctt	cagctgacaa	11580
ggccagcatt	atacctttac	caccctacct	caggggtgta	tcaactctcc	agctttgtgt	11640
cataatctta	tttgagaga	ccttgctcgc	ttttcacttc	cacgagatat	aacactggtc	11700
cattacattc	atgacattat	gatgattgga	tacagtgagc	aagaagtagc	aaacacactg	11760
aacttattgg	tgagacattt	gtatgccaga	ggatgggaaa	taaatccagc	taaaatttag	11820
ggactttcta	cctcggtaaa	atttctaggg	ttccagtggc	atgagacctt	tggagatatt	11880
ccttctaagg	tgaagcataa	cttgctgcgt	ttggcccctc	ttacaaccaa	gaaagaggca	11940
caatgcctgg	tgggcctatt	tggatttttg	aggcaacaca	ttcctcgttt	gggtgtgtta	12000
ctctggccca	tttatcgagt	gacctgaaag	gctgccagat	ttaagtgcag	tctagaacaa	12060
aagaaggctc	tgaacaggt	ccaggctgct	gtgaaagctg	ctctgccatt	tgggccacat	12120
gaccccgag	atccaatggt	gcttgagggt	tcagtggcag	atagggatgc	tgtttgagc	12180
ctttggcagg	cccccatagg	tgaatcacag	tggagacctc	taggattttg	gagcaaggcc	12240
ctgccacttc	tgagataac	tactctcctt	ttgagagaca	gctattgggt	tgttattggg	12300
ctttggtggt	aactgaacgt	ttgactgtgg	gtcataaagt	caccatgcta	cctgaacctg	12360
cctatcatga	actggttgct	ttctgacca	tctagccatg	aagtgggtca	gcacagcggc	12420
atttcatcat	caaattgaag	tgggtgtgtat	gtgatcgggc	ttgagcaggt	cctgaaggca	12480
caagtaagtt	acataaggaa	gtggctcaaa	tgcccatggt	ctccactcat	gccaccctgc	12540
cttccctccc	ccagcctgca	ccaatggcct	catggggagt	tccctatgat	cagttgacag	12600
aggaagggaa	gactaaggac	tggttcatag	atggttctgc	acgatatgca	ggcaccaccc	12660
gaaagtggac	agctgcagca	ctatatccac	tttctaaatg	catgtgtaca	cttgtgctaa	12720
gaaaatatct	ttattttatt	tcctttattt	ttcctttatc	atgtgacctt	agattttatg	12780
acttcacatc	agcatttaag	catttaagt	ttgttcatat	cagcatttaa	atattgttaa	12840
ccttatgtaa	taacttttgg	tttggggatt	ggtgcgtttc	tggttgtatg	aggatagttg	12900
tattatatta	ggcataatta	tgaccttatt	attgtcttta	tttgaagatt	atgtatgatt	12960
tcaggatgtg	tgtatgggtt	caagttgaca	aggagtggga	cttgtgatgg	ttaatactgt	13020
caacttgatt	ggattgaaag	atgcaaagta	ttaatctcgg	ttatgtctgt	gaggggtgtg	13080
caaaaggaga	ttaacatttg	agtcagtggg	ctgggaaggc	agaccacccc	ttaatctggg	13140

BI OL0250W0SEQ_ST25. txt

tacacaccat	ctaatacaagt	tccagtgtgg	ccagattgta	aagcagggag	aaaaatgtga	13200
aaagactaga	ctgaattagc	ttcccagcct	acatctttct	cctgtgccaa	atgcttcctg	13260
ctcttgaaca	tcggactcca	agttcttcag	cgttgggagt	tggactggct	ttcttgcctc	13320
tcagcttgca	gagggcctgt	tgtggaacct	tgtgatccgc	tgagttaata	ctacttaata	13380
agatccccct	tatatacata	taatataatta	tattatataat	aatatatata	atatataatta	13440
tatataatat	atataatata	ttatatatta	tatataatat	atattatata	ttatatataa	13500
tatatattat	atataatata	tattatataat	tatatattat	atataatata	tattatataat	13560
aatatatata	aaatatataat	atatactatt	agttctgtcc	ctctagagaa	ccctgactaa	13620
tacaatttat	gtcattaatc	tcattttattg	atttgtatac	attgaaccaa	ccttatatcc	13680
caggaataaa	acctacttga	ttgtgggtga	ttagcttttt	gatgtactct	tggattcaat	13740
tgctgggtatt	ttattgagaa	tttttgcata	tgtgttcata	aaggatattg	gcttgaagtt	13800
ttcttttttt	gttgttccat	atcagaatga	tgacgacctc	atagaatgag	ttagtctgtc	13860
ctcttttatc	ttttggaatt	gtttcaggag	gcttgatata	agctcttctt	tatatgactg	13920
gtatactttg	gctaggaatc	tctctgggtcc	aggggttttt	ctgggtgtagg	tttttaatta	13980
ctgattcaac	ttcagaactc	attactcatt	attgagttct	aaaactcact	ttcatgtact	14040
cttcaaaaaga	ctgtcttctt	ctgttggtga	gcgggggtgtt	ctctcaaggt	cgtttaggtg	14100
aagggtggtg	ctgggtgttct	tctgtatcct	tactgcttgt	ctttctcttt	ttttattgac	14160
tactgaggat	taatgggtgat	gtgtccaact	ttaactctag	attagtctat	ttctctttta	14220
gattgtaact	ctgttttata	tattttgaag	ctctgttggt	aggcatgtgt	atttggattg	14280
ttaggctctc	ttgatgatga	cctttatcat	tatgtaatgt	ttcttcttat	ctctggaagt	14340
attcgttgtt	ctgaagtcta	tttgtgctga	tatgaataca	gccttcacag	ctctattttc	14400
actagtattt	gtatatcttt	ttctcagctt	ttaaattgag	atgttcagac	catttgcatt	14460
aaagtagttg	ttaataggat	taaatttaaa	tctaccatta	agttgggttat	ttctctttgt	14520
cccatttaaa	ctttgttccct	tttttcataat	ttttctgcct	tcatttatat	tgagtttatc	14580
tccacgactt	acttatataa	ttaattttta	atgggttttag	tattttccac	aatgtttata	14640
atatatactt	tgattttttc	acattccacc	ttcaaattgac	agaattatac	tggatatata	14700
gaaatcttac	atcattgcac	ttctccttcc	tccctctcaa	aatgttgtgc	tattgctctt	14760
tgtaatagag	gcttacttct	attatgttat	agctctcata	atacattgac	actattttta	14820
ccctgaataa	tcagttgttt	tttaaagtga	ttatgactac	aaatattttg	aataatttct	14880
ttattttacc	atttctgggtg	ctccttatct	tttacagtag	atcccaattt	ccatctggag	14940
tcacattctt	tctgtgaaaa	acaaccttta	gcatttctta	tagcacggga	ctgctgttgc	15000
tgttgtcttt	cagcttttct	ttgtctgaag	aagtccttat	tttgccttca	gtttttaaaa	15060
gtgattttgc	tgagtataga	tactgggttg	agagtttcat	tccttgtatc	attttaacaa	15120
tgatgttcca	ttatatcccg	ttttgaatag	tttctgacta	gaaatctgat	ctttgtttct	15180

BI OL0250W0SEQ_ST25. txt

ttgtattcaa tagttccttt ttctctgact gcctttaaga tattctcatc tttgtttttc	15240
aacagtttga ctataatttg tttattatta actttttgta tttattctgc ttgaggtttc	15300
ctgagctcct tggatttgca gattgttgat ttttattggt tttgtaaaat tcatagccat	15360
tatctattct actgttttgt ttttttttc acttctctct ctctgtattc ttctttttgg	15420
actgtaagta ttcaaagtgt agatcattca tattgcttca taaaccttat atgcttcttc	15480
tgcttttttt ttttgtcag gaactctttt tttgtatctg tgttggtttg gataagttct	15540
agtagactat gttcaagttt atggattatt ttgttagttg tgtctaattg actcctcagt	15600
gcattcagag aattcttcat ctctgatatt ataaatctct tcctagcatt ttcatgttac	15660
tcttttctat agtttccatc tctttgctga aattctcccc ctatccatgg atattgtcca	15720
cctttaccac aagattcttt aacatattaa cataggtatc atacaaaccc aaactgatag	15780
ttccagatg gtgtcttttc tgagtctgtc tgtcttgatt gctttattat ttaacagtga	15840
cttatcttcc ctcttcagct tttggtgtgt cttgtaattg tttaatcaaa cactgggtat	15900
cataaatgga ggaacagtag agattgcagt aaatattatt tatgctttga aatgggcacc	15960
catcttctgt tgaaaatatg ttttgtggtc aattgagtca acctagtaac tggttgaact	16020
gaatttggca tttgtgcttg ttgcttttat cttaaagca ccacaggttt aaattcctcc	16080
agtgatgggt tgctgctatc ttttgcttag agtggggcct ggggtgtgga agaattttct	16140
cagtgttcct atctattatt agatttttagc agtcactgca tgcctgcact acagagggga	16200
tatcttcata cacataatct aaccccatg aaactgctgt ttcttcttaa tgaatgctca	16260
atctttggtg gaaataaaca aatgctgtat ctcttgagc cacttcagtc ttagtcaggt	16320
tctgcagggc tttgaaggga atgcattctc agtattcttg tgccttattt ggatggaact	16380
tgaacctgtg gtgggttttg agagaaagag tagcagacgt ctgctatggt gcaatgcagg	16440
atgctgggca caagaaaatt tccagtctct cctccaagga aataagattt gatcatctac	16500
ctatccctga gaagtgaagg gctttgcctg cgggtgctaga tgcaaaacca ttttctccc	16560
cccattgccc agaaacttaa ggctttggct tttctgagca gtggcttagg gaattgtgca	16620
aggttttcat atttgaccct gacagcccat caccacctac agcttgcagt gccaaatgta	16680
tctccctctg atctctctg tcctgtggtc ctcatgaaca ttaagaagag atttctaaaa	16740
aagagcttgc acatgagcat agtttctggt gagaagaatt ctgatatggt aacttctct	16800
aaacttttaa ataaaatatt tctaagaatt aaataaagtt ctagaatgat atgaatctat	16860
tcctttggtt tttgacgt ctgtctgcct gctaatacag agaagagaat ggtcgtaatt	16920
ctcagagact ttttctgtt tgtgtcataa atgacttcac attttttct gttctaagaa	16980
ctattcagct tgatttcttc tgttttaatt ttagcagcac ctgagcaaag ccatgtggtc	17040
caggattgct accatggtga tggacagagt tatcgaggca cgtactccac cactgtcaca	17100
ggaaggacct gccaaagtgt gtcatttatg acaccacatc aacataatag gaccacagaa	17160
aactacccaa atgcgtatgt cattaatctt acagtaagca aaacaaggtc caagtaaaat	17220

BI OL0250W0SEQ_ST25. txt

ttgtcttaga	aaaggtgtgc	gtcaagctaa	cttcttatga	ttaaattttt	ctcacacata	17280
gaatgcatgg	caaatgtct	gagaaacatt	acttttagca	aagagtatga	tagaagagaa	17340
atgttaagct	ggctctcttt	cctgagagtt	tgataaaatc	aggagaatat	ctggcggtgg	17400
tgaggccaca	ataatggaaa	atcagaatgt	ttagacagag	tcagcttcaa	caacactcac	17460
taaaggtcaa	tgtgatcttt	accccttgaa	attctataat	tctaattctc	aattcctgaa	17520
gtgaagggtg	tgttggcctt	ttctgtcttg	gctcacaagt	aatgatatg	tgcatatcta	17580
tggaaaggcg	aatctatctt	tttctatata	tatgtctatt	ccaacgggta	gaaacaccct	17640
gggtcctgag	caccagtggg	ctgaaggaat	acgggttgcc	aggaagagag	aagcaaaggc	17700
aggaaggcag	atgaaagtaa	gaaatgagac	agatgctaaa	caataaaaag	tgcggaaga	17760
tagacagaag	ctggggctg	accacacat	ggccagtctt	tcacacataa	gtgactacca	17820
aagacaagaa	aaaatgattt	ccgcttggtg	gacaatagat	ggtagaggac	caaggaatt	17880
gcgagagaga	gaacaatgag	atcaactcaa	cagatgcact	ggttttcttc	ctggagaccc	17940
ttcctgcact	gaagggcagg	agatggagcc	caaaaaaac	tgtagccatc	ttgctgaaca	18000
gaggagggac	attggagttt	gggattattc	aggtggctag	gattttctag	gcctgctaac	18060
aatgagaaca	gatttgtgga	ggaaaggagt	tctagaaata	tgcatagaaa	tctcctcgag	18120
tcattggcta	aacatgaagc	tgcatgtaca	cagaaaatag	atccacaaga	aagtagggca	18180
aagaacatct	acggaagagc	agcaactaca	atggaacagt	gagctcaata	aacatgacag	18240
agctcaaata	gcactaaggg	atattggagt	ttggaccaca	cagaggagag	agacttcact	18300
gaacatcttg	ggcattcagt	agagaccag	gaaaagccat	actttaggag	tagaattagt	18360
atattcttag	aataaaggca	gctccacaca	aacaatagca	aaactgaaaa	ggaagtctcc	18420
aagcatcaga	atgatgtcca	agtcaatgaa	ctgcctctga	gaggaaaact	caaccatctt	18480
tagaggtaaa	catcaaagtc	aagtggctca	gctatgcagt	atccacagtg	tgaggcctaa	18540
atataaaact	tgactacaca	tagaaacctt	ttagtgtgac	ccacaagcag	gaggaaaatc	18600
agccaatata	aacagaccca	gaagagacag	aaatgattag	aatggcataa	aaatttgaca	18660
tatcactata	taataattga	gttctaggat	ttaagaaaac	atgaatatag	aatgcaacag	18720
acaccttatc	cagagacagt	aagagtataa	agagccaaat	cgaagaacta	ctaagagata	18780
tgtcttaaata	gaaaaaatta	ctagatggcc	tcccatctta	gtagacatt	tcagaagaaa	18840
ataccaaatg	aaaaataatt	gcatagaacc	tacagaacca	gatacacaca	tacaaaacac	18900
acgcatgcat	acacacacac	tcaaactatg	ataagcttac	aaacacacac	acacatccac	18960
aaatgctgaa	aaatgaaatc	aaccgagcca	cacagacata	aaggaaaaca	taaaaagatt	19020
tcctacatgt	gggaagcaag	tcacagaaag	ggggaaggag	attggaacag	aaatatatac	19080
tgaaagcaag	gatggctgaa	aattttccaa	atataaagaa	gattaaaaaa	tcacggactc	19140
aagaagctca	atggatcaga	aaaataattt	ctaaaatgac	aattatagga	tgccactggg	19200
tacatagcag	ttcaactgtc	agagggcaaa	gacataatac	acagaaaaat	ctcgtaagga	19260

BI OL0250W0SEQ_ST25. txt

acgggaaaaa	caaaaagctg	tgtcttgcta	gaggaacagt	gatacaagtg	actaatgtgt	19320
tcccatcaga	aacactgcaa	cctggacaca	aaagaataac	attaaagtaa	taaacgtaag	19380
aaagaagagc	tcaactgaga	aggctacatc	cagcaataaa	atgccttgaa	gttcatccat	19440
gttggaggaa	tgacatttgt	gcactcccct	aaacaaagaa	accggaaact	gtaagacttt	19500
ggaatcagca	ggcttatgta	acaaaagagg	tgaccctaag	gaattaagga	gaagaagaat	19560
agaacaagaa	gggaactttc	tgcagcctat	ataatgaaga	acctagcaat	tggtcaaatgt	19620
agatgaaaat	gctacatgtt	ttcttgatca	aacgtttata	tctttttaaa	tgagagttga	19680
cgagttgaag	caaaatgata	ccaatatatt	taactttacc	atatgtagaa	gtaaaaattt	19740
gaacatgtag	cataaatcat	gtagggatta	attggaagtg	taccactgta	agtttcttac	19800
ctcatgcacg	atagtatgta	atactaataa	aaggttaatg	tgtgggttca	aagggatatt	19860
gcaaatccta	gagcaatcac	aaagttttta	actctgaggt	ttgttgata	ataacaatat	19920
tttatgtatt	caaagagggg	aagccaagga	agaaaaaaaa	gtctttaaag	agctctggct	19980
cttagtacat	ccagttgctc	attgaatgag	cttcctggaa	tggaggggtct	gggactgaga	20040
ctaggccaca	tgtgtagagc	cactagagac	acaatgttgg	atccccatgg	cccataatac	20100
atttcccatt	ttctcaggca	gccacaggtc	atgaatgtga	ggatactgag	aggttggagc	20160
aacgttcttg	ggaggcataa	ggaagagcga	atgcttcaag	atccccgcag	cccaaactcc	20220
tcagctgctt	tgccctctaa	ttcattgttt	tttgctcctc	catagctgtc	cgacctcttc	20280
agatctctta	gtcttcctgc	catcttcctt	tatgccatgg	gacccactgt	tctttcaact	20340
catccccag	ttctggagtg	gctgtggaca	gcagaggata	gactgagagc	aggagagaag	20400
gtcctgcca	ggaaccatt	ctagagatac	tgcatctgc	ctgggagcaa	gttttccagg	20460
gcagctttga	gaagtcttgc	agaaacaaac	ctacttgacc	gacatgatat	gggaatgaca	20520
gacagtaata	ctatttgcac	aatgcttttc	catgggaaag	gtagagcctt	ttcactaggt	20580
tttgagtaca	tggagtgtga	gagttgacct	ggaaaggtta	tcctccttga	tgccatgttt	20640
tctctgaaga	actacatgtt	cgttgcaact	cccacattag	aatatgaagt	cctaccgaga	20700
gagatacgga	gactagacag	atacagatgc	atttgcatgt	gaatacacaa	tcccacaata	20760
cagacgtcaa	aaccataacc	agttattcca	gagagatgga	ttgggcagaa	ggcagaagga	20820
gaatactctg	atcgtttttc	ggccacgtgt	gtgtgttatc	tcagtgtttc	taagaagcgt	20880
ttgctacttt	agatttttta	tttaaaaaaa	atagtaataa	tctattaagt	atgagagatg	20940
tgcagagagg	attagtgatc	gagagccatt	tttgctgggtg	gcaatcatat	ggtactttta	21000
atgggaatat	tagaaaggca	ccggtaatga	ccttgttgca	gcacaaagga	gagagtgtgg	21060
ggtgcccctg	catgttgtcc	cacctcttgt	gacgtgtatc	gttttggaat	ttccagtggc	21120
ttgatcatga	actactgcag	gaatccagat	gctgtggcag	ctccttattg	ttatacgagg	21180
gatcccgggtg	tcaggtggga	gtactgcaac	ctgacgcaat	gctcagacgc	agaagggact	21240
gccgtcgcgc	ctccgactgt	taccccggtt	ccaagcctag	aggctccttc	cgaacaaggt	21300

BI OL0250W0SEQ_ST25. txt

aaggagtctg	tggccagaca	tctacacgct	tcgatgctgg	gatgaaaagc	catggaaatt	21360
cccactgatg	cagccgcctt	caatggtaaa	cggatgctcg	agtgttgcct	gagttctacc	21420
atgtaggagg	aagcctccgt	gcactctctg	ggggagccag	cggagtgatt	tctggtgcaa	21480
cgtggttggg	ctttgtcttt	aggatgggca	caaaccctcc	agggggatcg	acttcaaaat	21540
tcaccttggt	gtaaaacggg	ctacctcagt	gtcccagcca	aaatTTTTat	tgtaacatgc	21600
tgtcaggtgt	gtcactcttt	ccaagccagt	aagcttttcc	ggggatttct	tcaagtagcc	21660
agcattcaga	gcaatcttca	gcattgcaga	ttctgagaaa	tgtggctctg	gagcctgtca	21720
ccctcgagaa	acctaagagg	gctgcattga	ttccatgtgg	ccctgggtct	atggagcagt	21780
acatgagctc	ccagtgtctt	aaggctcttc	agccctaggc	tttgaaggga	gtgatttctc	21840
agtattctta	aacctctttc	tgatgacact	tgtacctgtg	aggggtctag	agagaaagag	21900
tagtagactc	ctactttact	acaattcagg	atgcagggca	tgagaggatt	ccctctctcc	21960
tccaagggaa	gaagcttttg	gcgtgcacac	atccctgaga	agcaaagtgt	ctttgtcttc	22020
agtcagatac	ataggaccgt	tttctgcccc	atggcccgga	agccaaaggc	cttggctttc	22080
atgatcaacg	gtctagggaa	acatgcaaaa	tttccatgtc	tgtcccaaac	tctgcccccg	22140
acagccaatt	accacctgca	gcccgcattg	ccaaatgcgg	tgccgtttgc	atgaagattc	22200
agtagagttt	cctagaaagg	tgctacctcg	tgagctcact	ttccaatgag	gaatctgatc	22260
tgttggtgtt	ctctaagggt	tcaggtgaaa	tatttccaag	aacttactac	agttctagaa	22320
tgggaggaat	ctgttgcttt	ggtgtttgtt	tgttggctcg	ttttctcaca	tccatctgcc	22380
tatggataag	gaaaagagaa	cggtcgtaat	tctcatagac	tcctttctgg	ttgtgtcaca	22440
aatggcttca	catgtttctc	tatgctcaga	gatactcagc	ttgatttccc	gtgttttcat	22500
ttcagcaccg	actgagcaaa	ggcctggggg	gcaggagtgc	taccatggta	atggacagag	22560
ttatcgaggc	acatactcca	ccactgtcac	aggaagaacc	tgccaagctt	ggtcatctat	22620
gacaccacac	tcgcatagtc	ggaccccaga	atactacca	aatgcgtatg	tctttgttct	22680
ttaccataag	agaagaaagg	gccaagtga	gtttctgtta	caagagatgt	gtctcaagct	22740
gagttctccg	aactcaactt	gtgacagatg	cagatggcgt	agcaaaatgt	ctcaggatga	22800
ttgccttggg	gctaagggtc	tgagagaagg	gaaatgttaa	gctccctctc	cttccctcta	22860
gttctattga	gcagaaggga	aatctggagg	tgaggagatc	acattatgaa	gaaagtcaga	22920
atgacaaagg	accagacact	tagattaccc	ttccacaaca	ccaactaaac	gtcaatggag	22980
actttccagt	tggaattccg	ttattctggc	ttccacttcc	tgaagggaag	gttgcgtttg	23040
ccttttctct	ctgggttcaa	gaggaaagaa	taggtgctta	tttatggaca	ggtgaattga	23100
tctgtttcta	tatctacgta	tattccgatt	gtcagaaaaa	cactcgttcc	taagtaccag	23160
tggcctgaag	ggatacaggt	tcccagcaag	agaagatcca	aggaagggaag	gcagatgaga	23220
gtcagcacag	agagggatgc	tgaaaagtaa	aagggatggg	tggatggaga	gaagcccggg	23280
tctgaccacc	caatggccaa	tattttggcc	acaagcgact	accagagaca	tggaaaaatg	23340

BI OL0250W0SEQ_ST25. txt

gtttctacat	gtgggacaac	agatggtaga	ggacctagag	aattgagaga	ggggcaatga	23400
tgggctccac	tccgcagatg	ccttggcttt	cttcttgat	acccttctg	cactgaatag	23460
caaggagatg	gagcccaagc	agactgtagc	catcttgctg	aatggaggag	agggattgga	23520
gtttgggatg	actgtggtag	ctgaaatfff	tctaggtctg	ctagaaataa	gaactggfff	23580
gtgtggagga	aaagagctct	acaaatacgc	atagaagtct	cctccagtcg	ttggcctgac	23640
atgacgctgc	ctgtgcacag	gaaatggttc	cacgagaaag	tgtggcaaag	aacattttact	23700
gagaaacagc	aagtacaaga	gcacaggaag	ctcaataaag	aagagagaga	tcacatagca	23760
ctctgggata	ctggagtfff	tcccagctag	accagagagt	cctcacggag	cacattgcca	23820
attcagtgga	gaccccgaa	cagccgtaat	ttaaaggtag	acttagtata	ttactagaat	23880
aaagtcagct	gcagacaacc	ccttgacacag	ctggaaagca	agtgtccaag	catcaaatcg	23940
gtttccaatc	aatgaagtgc	ctgtgagagg	aaatctcaac	tctctttaga	agtaaacaac	24000
aaagtcgatt	gcctcagcta	tgcggtatcc	gcagagttag	tcctaaatff	aaaatctgac	24060
tacatgtaga	aaagcgtttc	gtgtgacca	tgaccaggaa	ataaatcggg	taatacaaac	24120
aggctcagga	atgagagaaa	tgattagaat	tgcgtgaaaa	tttgacatat	cagtatgata	24180
actgatttca	aatattftaa	aaaacaacat	gcaagaaagc	agatatcata	tcaagagaaa	24240
ttaacagtac	agaatagcca	aattaaatta	aagaggtagt	ataaaaaaag	tatgtcttaa	24300
ttgaaaaaaa	ttactgtatg	gccggctgat	caatfttagac	gtttcagagg	aaaacattac	24360
ccaacacaca	attctagaga	acctacagaa	tgagctacac	acacacacac	acacacacac	24420
acacacactg	aaaacacacc	catactcaca	cacacgcaga	aactcacaag	ttctaacaca	24480
cacagacacg	cgcaccctg	aagaaacagt	gaaatataaa	attaagcgag	cctcacagac	24540
atgtaggaaa	atatgaaaag	atttcttgca	tgtgggaagc	aagtcacagt	aaagagcaag	24600
ggagttttata	atagaaacaa	ataccagaat	caaggatggc	tgataactff	tcaattacga	24660
agaacattaa	aaaaaatcac	agaatcgtga	aactcaaggg	atcatatagg	gaatttcgga	24720
aaaaaaaccc	aacctgtatg	atgtactfff	gtacatcaca	gttcgaagg	aacaaggcaa	24780
agatgtaata	agaagaaacc	tgtcacgaga	aactggagga	aaaagagctg	tgtcttccta	24840
caagtacact	gatacaaatt	gccaatgtgt	tcacctcaga	aacactggaa	gccagatacc	24900
agggaatatt	gttaaaatga	taatcaggaa	caaaaagaga	tcaaccggga	atgctgaatc	24960
cagcaataaa	atgccttgaa	ggcatccat	gtcggataaa	tgcatattgt	gactgcccc	25020
aaagaaagaa	accggaaact	gtaagaattg	gaaatcagca	ggcttatgta	acaagagagg	25080
tgacccgaag	gaattaggta	gaagaagaat	tgaacaagaa	aggaaactff	tgcagcccac	25140
gtaatgaaga	atccagcaat	tggcaaattg	agatagatgt	aaatgcaaaa	tattttcttg	25200
atcaaatttc	tatatctftg	taaatgagag	ttgactactt	gaaacaaaat	gatagcaaga	25260
tatttaactt	cagcatatgt	agaggtaaga	atttgaaatg	gtagcataaa	tcacgaaggg	25320
attaattcga	agtgtaccgt	tgtaaatftc	tttacctcat	gcacgatgg	gtgtcatatt	25380

BI OL0250W0SEQ_ST25. txt

aataaaaggg	tactgtgcgg	gttcgaaggg	atattgcaaa	tcctagagca	atcacaaagg	25440
tttgaactct	gaggtttttg	gtataataag	aatagtccat	gcattcaaaa	gaggggaagcc	25500
aaggaagaac	tagaagtctt	tcaagagctc	aggctcttat	acatccagtt	gctcattgaa	25560
ccagcttcct	ggaatggagg	gtctgggggt	gagactaggc	cacaagtcta	gagtctctag	25620
agagacagtg	ttggaacccc	atggcccata	atacatttcc	catttttctca	ggcagccaga	25680
ggcatgaat	gtgaggatac	tgggaggttg	gagcaacgtt	cttgggaggc	ataaggaaga	25740
gcgaatgctt	caagatcccc	gcagcccaaa	ctactcgcct	gctttgcccc	ctaatgcatt	25800
tttctctgct	gctccgtagc	tgtccgacct	cttcagatct	cttagtccac	cctgccgtct	25860
tcctttatgc	catgggtccc	actgttcttt	caactcatcc	ccctttccct	cagtcccgga	25920
gtagctgcgg	ccagcagagg	gtagactgag	agcaggagag	aaggacctgc	ctaggaaccc	25980
cttctagaga	tactgcatcc	tgcctgggag	caagttttcc	agggcagctt	tgagaagtct	26040
tggagaaaca	aacctactaa	acctgacaga	cagtaatact	atttgcacaa	tgcttttctg	26100
tgggaaaggt	agagcctttt	cactacgtat	tgagtacata	gagtgtgagg	gttgacctgg	26160
aacggctatc	ctcctggatg	acgtgtgttt	tctgaagaac	tacatgttcg	ttgcaactcc	26220
cacattagaa	tatgaagtcc	taccgagaga	gatacggaga	ctagacagat	acagatgcat	26280
ttgcatgtga	atacacaatc	ccacaataca	gacgtcaaaa	cccataccag	ttattccaga	26340
gagatggatt	gggcagaagg	cagaaggaga	atactctgat	cgtttttctg	ccacgtgtgt	26400
gtgttatctc	agtgtttcta	agaagcggtt	gctactttag	attttttatt	taaaaaaata	26460
gtaataatct	attaagtatg	agagatgtgc	agagaggatt	agtgatcgag	agccattttt	26520
gctgggtggca	atcatatggt	acttttaatg	ggaatattag	aaaggcaccg	gtaatgacct	26580
tgttgcagca	caaaggagag	agtgtggggg	gccctgcat	gttgtccac	ctcttgtgac	26640
gtgtatcggt	ttggaatttc	cagtggcttg	atcatgaact	actgcaggaa	tccagatgct	26700
gtggcagctc	cttattgtta	tacgagggat	cccgggtgtca	ggtgggagta	ctgcaacctg	26760
acgcaatgct	cagacgcaga	agggactgcc	gtcgcgcctc	cgactgttac	cccggttcca	26820
agcctagagg	ctccttccga	acaaggtaag	gagtctgtgg	ccagacatct	acacgcttcg	26880
atgctgggat	gaaaagccat	ggaaattccc	actgatgcag	ccgccttcaa	tggtaaacgg	26940
atgctcgagt	gttgccctgag	ttctaccatg	taggaggaag	cctccgtgca	ctctctgggg	27000
gagccagcgg	agtgatttct	ggtgcaacgt	ggttgggctt	tgtcttttag	atgggcacaa	27060
accctccagg	gggatcgact	tcaaaattca	ccttgttgta	aaacgggcta	cctcagtgtc	27120
ccagccaaaa	tttttattgt	aacatgctgt	caggtgtgtc	actctttcca	agccagtaag	27180
cttttccggg	gatttcttca	agtagccagc	attcagagca	atcttcagca	ttgcagattc	27240
tgagaaatgt	ggctctggag	cctgtcaccc	tcgagaaacc	taagagggct	gcattgattc	27300
catgtggccc	tgggtctatg	gagcagtaca	tgagctccca	gtgctctaag	gctcttcagc	27360
cctaggcctt	gaaggggagt	atttctcagt	attcttaaac	ctctttctga	tgacacttgt	27420

BI OL0250W0SEQ_ST25. txt

acctgtgagg	ggtctagaga	gaaagagtag	tagactccta	ctttactaca	attcaggatg	27480
cagggcatga	gaggattccc	tctctcctcc	aaggggaagaa	gctttttggcg	tgcacacatc	27540
cctgagaagc	aaagtgtctt	tgtcttcagt	cagatacata	ggaccgtttt	ctgccccatg	27600
gcccggaagc	caaaggcctt	ggctttcatg	atcaacggtc	tagggaaaca	tgcaaaattt	27660
ccatgtctgt	cccaaactct	gcccccgaca	gccaaattacc	acctgcagcc	cgcattgccca	27720
aatgcggtgc	cgtttgcatg	aagattcagt	agagtttcct	agaaagggtgc	tacctcgtga	27780
gctcactttc	caatgaggaa	tctgatctgt	tgtgtttctc	taagggtgtca	ggtgaaatat	27840
ttccaagaac	ttactacagt	tctagaatgg	gaggaatctg	ttgctttgggt	gtttgtttgt	27900
tggtcggttt	tctcacatcc	atctgcctat	ggataaggaa	aagagaacgg	tcgtaattct	27960
catagactcc	tttctggttg	tgtcacaaat	ggcttcacat	gtttctctat	gctcagagat	28020
actcagcttg	atttcccgtg	ttttcatttc	agcaccgact	gagcaaaggc	ctgggggtgca	28080
ggagtgcctac	catggtaatg	gacagagtta	tcgaggcaca	tactccacca	ctgtcacagg	28140
aagaacctgc	caagcttggg	catctatgac	accacactcg	catagtcgga	ccccagaata	28200
ctacccaaat	gcgtatgtct	ttgttcttta	ccataagaga	agaaagggcc	aagtgaagtt	28260
tctgttacia	gagatgtgtc	tcaagctgag	ttctccgaac	tcaacttggtg	acagatgcag	28320
atggcgtagc	aaaatgtctc	aggatgattg	ccttgagact	aaggggtctga	gagaagggaa	28380
atgttaagct	ccctctcctt	cctcctagtt	ctattgagca	gaagggaaat	ctggagggtga	28440
ggagatcaca	ttatgaagaa	agtcagaatg	acaaaggacc	agacacttag	attacccttc	28500
cacaacacca	actaaacgtc	aatggagact	ttccagttgg	aattccgtta	ttctggcttc	28560
cacttcctga	aggggaaggtt	gcgtttgcct	tttctctctg	ggttcaagag	gaaagaatag	28620
gtgcttattt	atggacaggt	gaattgatct	gtttctatat	ctacgtatat	tccgattgtc	28680
agaaaaacac	tcgttcctaa	gtaccagtgg	cctgaaggga	tacaggttcc	cagcaagaga	28740
agatccaagg	aaggaaggca	gatgagagtc	agcacagaga	gggatgctga	aaagtaaaag	28800
ggatgggtgg	atggagagaa	gcccgggtct	gaccacccaa	tggccaatat	tttggccaca	28860
agcgactacc	agagacatgg	aaaaatgggt	tctacatgtg	ggacaacaga	tggtagagga	28920
cctagagaat	tgagagaggg	gcaatgatgg	gctccactcc	gcagatgcct	tggctttctt	28980
cctggatacc	cttcctgcac	tgaatagcaa	ggagatggag	ccaagcaga	ctgtagccat	29040
cttgctgaat	ggaggagagg	gattggagtt	tgggatgact	gtggtagctg	aaatttttct	29100
aggctctgta	gaaataagaa	ctggtttgtg	gaggaaaaga	gctctacaaa	tacgcataga	29160
agtctcctcc	agtcgttggc	ctgacatgac	gctgcctgtg	cacaggaaat	ggttccacga	29220
gaaagtgtgg	caaagaacat	ttactgagaa	acagcaagta	caagagcaca	ggaagctcaa	29280
taaagaagag	agagatcaca	tagcactctg	ggatactgga	gttcttccca	gctagaccag	29340
agagtcctca	cggagcacat	tgccaattca	gtggagaccc	cagaacagcc	gtaatttaaa	29400
ggtacactta	gtatattact	agaataaagt	cagctgcaga	caacccttg	cacagctgga	29460

BI OL0250W0SEQ_ST25. txt

aagcaagtgt	ccaagcatca	aatcggtttc	caatcaatga	agtcctgtg	agaggaaatc	29520
tcaactctct	ttagaagtaa	acaacaaagt	cgattgcctc	agctatgcgg	tatccgcaga	29580
gtgagtccta	aatttaaaat	ctgactacat	gtagaaaagc	gtttcgtgtg	acccatgacc	29640
aggaaataaa	tcgggtaata	caaacaggct	caggaatgag	agaaatgatt	agaattgcgt	29700
gaaaatttga	catatcagta	tgataactga	tttcaaata	ttaaaaaac	aacatgcaag	29760
aaagcagata	tcatatcaag	agaaattaac	agtacagaat	agccaaatta	aattaaagag	29820
ctagtataaa	aaaagtatgt	cttaattgaa	aaaaattact	gtatggccgg	ctgatcaatt	29880
tagacgtttc	agaggaaaac	attacccaac	acacaattct	agagaacctc	cagaatgagc	29940
tacacacaca	cacacacaca	cacacacaaa	ctgaaaacac	accatactc	acacacacgc	30000
agaaactcac	aagtictaac	acacacagac	acgcgcaccc	ctgaagaaac	agtgaatat	30060
aaaattaagc	gagcctcaca	gacatgtagg	aaaatatgaa	aagatttcct	gcatgtggga	30120
agcaagtcac	agtaaagagc	aaggaggttt	ggaatagaaa	caaataccgg	aatcaaggat	30180
ggctgataac	ttttcaatta	cgaagaacat	taaaaaaat	cacagaatcg	tgaaactcaa	30240
gggatcacat	aggggaatttc	ggaaaaaaa	cccaacctgt	atgatgtact	tttgtacatc	30300
acagttcgaa	ggtaacaagg	caaagatata	ataagaagaa	acctgtcacg	agaaactgga	30360
ggaaaaagag	ctgtgtcttc	ctacaagtac	actgatacaa	attgccaatg	tgttcacctc	30420
agaaacactg	gaagccagat	accaggggaat	attgttataaa	tgataatcag	gaacaaaaag	30480
agatcaaccg	ggaatgctga	atccagcaat	aaaatgcctt	gaagatcatc	catgtcggat	30540
aatgcatat	tgtgactgc	cccaaagaaa	gaaaccggaa	actgtaagaa	ttggaaatca	30600
gcaggcttat	gtaacaagag	aggtgacctg	aaggaattag	gtagaagaag	aattgaacaa	30660
gaaaggaact	ttctgcagcc	cacgtaatga	agaatccagc	aattggcaaa	tgtagataga	30720
tgtaaatagca	aaatatatttc	ttgatcaaat	ttctatatct	ttgtaaata	gagttgacta	30780
cttgaaacaa	aatgatagca	agatatatta	cttcagcata	tgtagaggta	agaatttgaa	30840
atggtagcat	aatcacgaa	gggattaatt	cgaagtgtac	cgttgtaagt	ttctttacct	30900
catgcacgat	ggtgtgtcat	attaataaaa	gggtactgtg	cgggttcgaa	gggatattgc	30960
aaatcctaga	gcaatcacia	aggtttgaac	tctgaggttt	ttggtataat	agaatagtc	31020
catgcattca	aaagagggaa	gccaaggaag	aactagaagt	ctttcaagag	ctcaggctct	31080
tatacatcca	gttgctcatt	gaaccagctt	cctggaatgg	agggctctggg	gttgagacta	31140
ggccacaagt	ctagagtctc	tagagagaca	gtgttggaac	cccatggccc	ataatacatt	31200
tccatttttc	tcaggcagcc	agaggtcatg	aatgtgagga	tactgggagg	ttggagcaac	31260
gttcttggga	ggcataagga	agagcgaatg	cttcaagatc	cccgcagccc	aaactactcg	31320
cctgctttgc	cccctaata	atctttctct	gctgctccgt	agctgtccga	cctcttcaga	31380
tctcttagtc	caccctgccg	tcttccttta	tgccatgggt	cccactgttc	tttcaactca	31440
tccccctttc	cctcagtccc	ggagtagctg	cggccagcag	agggtagact	gagagcagga	31500

BI OL0250W0SEQ_ST25. txt

gagaaggacc	tgccntagaa	ccccttctag	agatactgca	tcctgcctgg	gagcaagttt	31560
tccagggcag	ctttgagaag	tcttgagaa	acaaacctac	taaacctgac	agacagtaat	31620
actatttgca	caatgctttt	ctgtgggaaa	ggtagagcct	tttactacg	tattgagtac	31680
atagagtgtg	agggttgacc	tggaacggct	atcctcctgg	atgacgtgtg	ttttctgaag	31740
aactacatgt	tcgttgcaac	tcccacatta	gaatatgaag	tcctaccgag	agagatacgg	31800
agactagaca	gatacagatg	catttgcatg	tgaatacaca	atcccacaat	acagacgtca	31860
aaaccatac	cagttattcc	agagagatgg	attgggtagg	aggcagaagg	agaatactct	31920
gatcgttttt	cggccacgtg	tgtgtgttat	ctcagtgttt	ctaagaagcg	tttgctactt	31980
tagatttttt	atttaaaaaa	aatagtaata	atctattaag	tatgagagat	gtgcagagag	32040
gattagtgat	cgagagccat	ttttgctgg	ggcaatcata	tgggtactttt	aatgggaata	32100
ttagaaaggc	accggtaatg	accttggtgc	agcacaaagg	agagagtgtg	gggtgccctt	32160
gcatgttgtc	ccacctcttg	tgacgtgtat	cgttttggaa	tttccagtgg	cttgatcatg	32220
aactactgca	ggaatccaga	tgctgtggca	gctccttatt	gttatacgag	ggatcccgg	32280
gtcagggtggg	agtactgcaa	cctgacgcaa	tgctcagacg	cagaagggac	tgccgtcgcg	32340
cctccgactg	ttaccccgg	tccaagccta	gaggctcctt	ccgaacaagg	taaggagtct	32400
gtggccagac	atctacacgc	ttcgatgctg	ggatgaaaag	ccatggaaat	tcccactgat	32460
gcagccgcct	tcaatggtaa	acggatgctc	gagtgttgcc	ggagtctctgc	catgttgggg	32520
gaagcctccg	tgtactctct	gggggagcca	gcggagtgat	ttctggtgca	acttgggtgg	32580
gctttgtctt	tagaatgggc	acaaaccttc	cagggtgatg	ggcttcacaa	ctcacctcct	32640
tctaaaatgg	gctatctcag	tgtcttagcc	aaaattttta	ttgtaacgtg	ctgtcagggtg	32700
tgtgattctt	tctgtcgcag	taagcttttc	tggggatttc	ttcaagtagc	cagcagtcag	32760
tgcaatcttc	agcattgcag	atttcaaaaa	atgtggctct	ggagcctgtc	atcctcgaga	32820
aacctaacag	ggctgcatta	attccatatg	gtcctgggtc	tatggagcag	tatatgagct	32880
cccaatgctc	taaggctctt	cagtcctagg	ctttgaagg	agtgatttct	cagtgttctt	32940
aaacctcttt	ctgatggcac	ttgtacctgt	gaggggtcta	gagagaaagg	ttagtagact	33000
tctcctttac	tgcaattcag	gatgcagggc	atgagaagat	tccctccctc	ctccaaggga	33060
agaaggtttt	ggcgtgcaca	catccttgag	aagcaaagtg	tctttgcctt	cagtcagata	33120
tataggatcg	ttttctgccc	catggcctgg	aagccagagg	ccttggcttt	catgatcaac	33180
gatctaggga	aacatgcaaa	atttccatgt	ctttcccctc	ctctgccctc	gacagccaat	33240
taccacctgc	atcctgcatt	gccaaatgca	gtgccctttg	tatgaacatt	cagtagagtt	33300
tcatagaaag	gtgctacttc	gtgagcgcac	tttgacgtga	gaaggagtct	gttctgttct	33360
gtttttctaa	ggatttcagg	tgaaatattt	cctagaactt	actacagttc	tagattggta	33420
ggaatctgta	ggtttgctgt	atgttttttg	gttggttttc	tcccatccat	ctgcctacag	33480
gtaagggaaa	gataacgttc	gtaattctca	tagactcctt	tctggttgtg	tcataaatgg	33540

BI OL0250W0SEQ_ST25. txt

cttcacatat	ttcgttattc	tcagagatac	tcagttttatt	tcttgtgttt	tcatttcagc	33600
accgactgag	cagaggcctg	gggtgcagga	gtgctaccac	ggtaatggac	agagttatcg	33660
aggcacatac	tccaccactg	tcactggaag	aacctgccaa	gcttgggtcat	ctatgacacc	33720
acactcgcat	agtcggaccc	cagaatacta	cccaaatgcg	tatgtctttg	ttctttacca	33780
taagagaaga	aagggccaag	tgaagtttct	gttacaagag	atgtgtctca	agctgagttc	33840
tccgaactca	acttgtgaca	gatgcagatg	gcgtagcaaa	atgtctcagg	atgattgcct	33900
tggagctaag	ggtctgagag	aagggaaatg	ttaagctccc	tctccttcct	cctagttcta	33960
ttgagcagaa	gggaaatctg	gaggtgagga	gatcacatta	tgaagaaagt	cagaatgaca	34020
aaggaccaga	cacttagatt	acccttccac	aacaccaact	aaacgtcaat	ggagactttc	34080
cagttggaat	tccgttattc	tggcttccac	ttcctgaagg	gaaggttgcg	tttgcctttt	34140
ctctctgggt	tcaagaggaa	agaataggtg	cttattttatg	gacaggtgaa	ttgatctggt	34200
tctatatcta	cgtatattcc	gattgtcaga	aaaacactcg	ttcctaagta	ccagtggcct	34260
gaagggatac	aggttcccag	caagagaaga	tccaaggaag	gaaggcagat	gagagccagc	34320
acagagaggg	atgctgaaaa	gtaaaagggg	tgggtggatg	gagagaagcc	cgggtctgac	34380
cacccaatgg	ccaatatttt	ggccacaagc	gactaccaga	gacatggaaa	aatggtttct	34440
acatgtggga	caacagatgg	tagaggacct	agagaattga	gagaggggca	atgatgggct	34500
ccactccgca	gatgccttgg	ctttcttcct	ggataccctt	cctgcactga	atagcaagga	34560
gatggagccc	aagcagactg	tagccatctt	gctgaatgga	ggagagggat	tggagtttgg	34620
gatgactgtg	gtagctgaaa	tttttctagg	tctgctagaa	ataagaactg	gtttgtggag	34680
gaaaagagct	ctacaaatac	gcatagaagt	ctcctccagt	cgttggcctg	acatgacgct	34740
gcctgtgcac	aggaaatggt	tccacgagaa	agtgtggcaa	agaacattta	ctgagaaaca	34800
gcaagtacaa	gagcacagga	agctcaataa	agaagagaga	gatcacatag	cactctggga	34860
tactggagtt	cttcccagct	agaccagaga	gtcctcacgg	agcacattgc	caattcagtg	34920
gagaccccag	aacagccgta	atttaaaggt	acacttagta	tattactaga	ataaagtcag	34980
ctgcagacaa	ccccttgcac	agctggaaag	caagtgtcca	agcatcaaat	cggttttcaa	35040
tcaatgaagt	gcctgtggga	ggaaatctca	actctcttta	gaagtaaaca	acaaagtcga	35100
ttgcctcagc	tatgcggtat	ccgcagagtg	agtcctaaat	ttaaaatctg	actacatgta	35160
gaaaagcggt	tcgtgtgacc	catgaccagg	aaataaatcg	ggtaatacaa	acaggctcag	35220
gaatgagaga	aatgattaga	attgcgtgaa	aatttgacat	atcagtatga	taactgattt	35280
caaatattta	aaaaaacaac	atgcaagaaa	gcagatatca	tatcaagaga	aattaacagt	35340
acagaatagc	caaattaaat	taaagagcta	gtataaaaaa	agtatgtctt	aattgaaaaa	35400
aattactgta	tggccggctg	atcaaattag	acgtttcaga	ggaaaacatt	acccaacaca	35460
caattctaga	gaacctacag	aatgagctac	acacacacac	acacacacac	acacacacac	35520
tgaaaacaca	cccatactca	cacacacgca	gaaactcaca	agtttctaaca	cacacagaca	35580

BI OL0250W0SEQ_ST25. txt

cgcgaccccc	tgaagaaaca	gtgaaatata	aaattaagcg	agcctcacag	acatgtagga	35640
aaatatgaaa	agatttcctg	catgtgggaa	gcaagtcaca	gtaaagagca	agggagtttg	35700
gaatagaaac	aaataccgga	atcaaggatg	gctgataact	tttcaattac	gaagaacatt	35760
aaaaaaaaatc	acagaatcgt	gaaactcaag	ggatcatata	gggaatttcg	gaaaaaaaaac	35820
ccaacctgta	tgatgtactt	ttgtacatca	cagttcgaag	gtaacaaggc	aaagatataa	35880
taagaagaaa	cctgtcacga	gaaactggag	gaaaaagagc	tgtgtcttcc	tacaagtaca	35940
ctgatacaaa	ttgccaatgt	gttcacctca	gaaacactgg	aagccagata	ccagggaata	36000
ttgttaaaat	gataatcagg	aacaaaaaga	gatcaaccgg	gaatgctgaa	tccagcaata	36060
aaatgccttg	aagatcatcc	atgtcggata	aatgcatatt	gtgcactgcc	ccaaagaaag	36120
aaaccggaag	ctgtcagaat	tggaaatcag	caggcttatg	taacaagaga	ggtgaccgga	36180
aggaattagg	tagaagaaga	attgaacaag	aaaggaactt	tctgcagccc	acgtaatgaa	36240
gaatccagca	attggcaaat	gtagatagat	gtaaatgcaa	aatatcttct	tgatcaaatt	36300
tctatatctt	tgtaaagag	agttgactac	ttgaaacaaa	atgatagcaa	gatatttaac	36360
ttcagcatat	gtagaggtaa	gaatttgaaa	tggtagcata	aatcacgaag	ggattaattc	36420
gaagtgtacc	gttgtaagtt	tctttacctc	atgcacgatg	gtgtgtcata	ttaataaaag	36480
ggtactgtgc	gggttcgaag	ggatattgca	aatcctagag	caatcacaaa	ggtttgaact	36540
ctgaggtttt	tggtataata	agaatagtcc	atgcattcaa	aagaggggaag	ccaaggaaga	36600
actagaagtc	tttcaagagc	tcaggctctt	atacatccag	ttgctcattg	aaccagcttc	36660
ctggaatgga	gggtctgggg	ttgagactag	gccacaagtc	tagagtctct	agagagacag	36720
tgttggaacc	ccatggccca	taatacattt	cccattttct	caggcagcca	gaggtcatga	36780
atgtgaggat	actgggaggt	tggagcaacg	ttcttgggag	gcataaggaa	gagcgaatgc	36840
ttcaagatcc	ccgcagccca	aactactcgc	ctgctttgcc	ccctaattgca	tttttctctg	36900
ctgctccgta	gctgtccgac	ctcttcagat	ctcttagtcc	accctgccgt	cttcctttat	36960
gccatgggtc	ccattgttct	ttcaactcat	ccccctttcc	ctcagtcccg	gagtagctgc	37020
ggccagcaga	gggtagactg	agagcaggag	agaaggacct	gcctaggaac	cccttctaga	37080
gatactgcat	cctgcctggg	agcaagtttt	ccagggcagc	tttgagaagt	cttgagagaaa	37140
caaacctact	aaacctgaca	gacagtaata	ctatttgcac	aatgcttttc	tgtgggaaag	37200
gtagagcctt	ttcactacgt	attgagtaca	tagagtgtga	gggttgacct	ggaacggcta	37260
tcctcctgga	tgacgtgcgt	tttctgaaga	actacatgtt	cgttgcaact	cccacattag	37320
aatatgaagt	cctaccgaga	gagatacgga	gactagacag	atacagatgc	atttgcattg	37380
gaatacacaa	tcccacaata	cagacgtcaa	aaccataacc	agttattcca	gagagatgga	37440
ttgggcagaa	ggcagaagga	gaatactctg	atcgtttttc	ggccacgtgt	gtgtgttata	37500
tcagtgtttc	taagaagcgt	ttgctacttt	agatttttta	tttaaaaaaa	atagtaataa	37560
tctattaagt	atgagagatg	tgacagagag	attagtgatc	gagagccatt	tttgcctggg	37620

BI OL0250W0SEQ_ST25. txt

gcaatcatat	ggtactttta	atgggaatat	tagaaaggca	ccggtaatga	ccttgttgca	37680
gcacaaagga	gagagtgtgg	ggtgcccctg	catgttgtcc	cacctcttgt	gacgtgtatc	37740
gttttggaat	ttccagtggc	ttgatcatga	actactgcag	gaatccagat	gctgtggcag	37800
ctccttattg	ttatacgagg	gatcccgggtg	tcaggtggga	gtactgcaac	ctgacgcaat	37860
gctcagacgc	agaagggact	gccgtcgcgc	ctccgactgt	taccccgggtt	ccaagcctag	37920
aggctccttc	cgaacaaggt	aaggagtctg	tggccagaca	tctacacgct	tcgatgctgg	37980
gatgaaaagc	catggaaatt	cccactgatg	cagccgcctt	caatggtaaa	cggatgctcg	38040
agtgttgccct	gagttctacc	atgtaggagg	aagcctccgt	gcactctctg	ggggagccag	38100
cggagtgatt	tctggtgcaa	cgtggttggg	ctttgtcttt	aggatgggca	caaaccctcc	38160
agggggatcg	acttcaaaat	tcaccttggt	gtaaaacggg	ctacctcagt	gtcccagcca	38220
aaatttttat	tgtaacatgc	tgtcaggtgt	gtcactcttt	ccaagccagt	aagcttttcc	38280
ggggatttct	tcaagtagcc	agcattcaga	gcaatcttca	gcattgcaga	ttctgagaaa	38340
tgtggctctg	gagcctgtca	ccctcgagaa	acctaagagg	gctgcattga	ttccatgtgg	38400
ccctgggtct	atggagcagt	acatgagctc	ccagtgtctt	aaggctcttc	agccctaggc	38460
tttgaaggga	gtgatttctc	agtattctta	aacctctttc	tgatgacact	tgtacctgtg	38520
aggggtctag	agagaaagag	tagtagactc	ctactttact	acaattcagg	atgcagggca	38580
tgagaggatt	ccctctctcc	tccaagggaa	gaagcttttg	gcgtgcacac	atccctgaga	38640
agcaaagtgt	ctttgtcttc	agtcagatac	ataggaccgt	tttctgcccc	atggcccgga	38700
agccaaaggc	cttggtcttc	atgatcaacg	gtctagggaa	acatgcaaaa	tttccatgtc	38760
tgtcccaaac	tctgcccccg	acagccaatt	accacctgca	gcccgcattg	ccaaatgcgg	38820
tgccgtttgc	atgaagattc	agtagagttt	cctagaaagg	tgctacctcg	tgagctcact	38880
ttccaatgag	gaatctgatc	tgttgtgttt	ctctaagggtg	tcaggtgaaa	tattttcaag	38940
aacttactac	agttctagaa	tgggaggaat	ctgttgcttt	ggtgttttgtt	tgttggtcgg	39000
ttttctcaca	tccatctgcc	tatggataag	gaaaagagaa	cggtcgtaat	tctcatagac	39060
tcctttctgg	ttgtgtcaca	aatggcttca	catgtttctc	tatgctcaga	gatactcagc	39120
ttgatttccc	gtgttttcat	ttcagcaccg	actgagcaaa	ggcctgggggt	gcaggagtgc	39180
taccatggta	atggacagag	ttatcgaggc	acatactcca	ccactgtcac	aggaagaacc	39240
tgccaagctt	ggtcatctat	gacaccacac	tcgcatagtc	ggaccccaga	atactaccca	39300
aatgcgtatg	tctttgttct	ttaccataag	agaagaaagg	gccaagtga	gtttctgtta	39360
caagagatgt	gtctcaagct	gagttctccg	aactcaactt	gtgacagatg	cagatggcgt	39420
agcaaaatgt	ctcaggatga	ttgccttggga	gctaaggggtc	tgagagaagg	gaaatgttaa	39480
gctccctctc	cttctccta	gttctattga	gcagaaggga	aatctggagg	tgaggagatc	39540
acattatgaa	gaaagtcaga	atgacaaagg	accagacact	tagattaccc	ttccacaaca	39600
ccaactaaac	gtcaatggag	actttccagt	tgggaattccg	ttattctggc	ttccacttcc	39660

BI OL0250W0SEQ_ST25. txt

tgaaggaag	gttgcgtttg	ccttttctct	ctgggttcaa	gaggaaagaa	taggtgctta	39720
tttatggaca	ggtgaattga	tctgtttcta	tatctacgta	tattccgatt	gtcagaaaaa	39780
cactcgttcc	taagtaccag	tggcctgaag	ggatacaggt	tcccagcaag	agaagatcca	39840
aggaaggaag	gcagatgaga	gtcagcacag	agagggatgc	tgaaaagtaa	aagggatggg	39900
tggatggaga	gaagcccggg	tctgaccacc	caatggccaa	tattttggcc	acaagcgact	39960
accagagaca	tggaaaaatg	gtttctacat	gtgggacaac	agatggtaga	ggacctagag	40020
aattgagaga	ggggcaatga	tgggctccac	tccgcagatg	ccttggcttt	cttcctggat	40080
acccttcctg	cactgaatag	caaggagatg	gagcccaagc	agactgtagc	catcttgctg	40140
aatggaggag	agggattgga	gtttgggatg	actgtggtag	ctgaaatfff	tctaggtctg	40200
ctagaaataa	gaactggttt	gtggaggaaa	agagctctac	aaatacgcac	agaagtctcc	40260
tccagtcgtt	ggcctgacac	gacgctgcct	gtgcacagga	aatggttcca	cgagaaagtg	40320
tggcaaagaa	catttactga	gaaacagcaa	gtacaagagc	acaggaagct	caataaagaa	40380
gagagagatc	acatagcact	ctgggatact	ggagttcttc	ccagctagac	cagagagtcc	40440
tcacggagca	cattgccaat	tcagtggaga	ccccagaaca	gccgtaatff	aaaggtacac	40500
ttagtatatt	actagaataa	agtcagctgc	agacaacccc	ttgcacagct	ggaaagcaag	40560
tgtccaagca	tcaaactcgt	ttccaatcaa	tgaagtgcct	gtgagaggaa	atctcaactc	40620
tctttagaag	taaacaacaa	agtcgattgc	ctcagctatg	cggatatccg	agagttagtc	40680
ctaaatttaa	aatctgacta	catgtagaaa	agcgtttcgt	gtgacccatg	accaggaaat	40740
aaatcgggta	atacaaacag	gctcaggaat	gagagaaatg	attagaattg	cgtgaaaatt	40800
tgaaatatca	gtatgataac	tgatttcaaa	tatttaaaaa	aacaacatgc	aagaaagcag	40860
atatcatatc	aagagaaatt	aacagtacag	aatagccaaa	ttaaattaaa	gagctagtat	40920
aaaaaaagta	tgtcttaatt	gaaaaaaatt	actgtatggc	cggctgatca	atttagacgt	40980
ttcagaggaa	aacattaccc	aacacacaat	tctagagaac	ctacagaatg	agctacacac	41040
acacacacac	acacacacac	aaactgaaaa	cacaccata	ctcacacaca	cgcagaaact	41100
cacaagttct	aacacacaca	gacacgcgca	cccctgaaga	aacagtgaaa	tataaaatta	41160
agcgagcctc	acagacatgt	aggaaaatat	gaaaagatff	cctgcatgtg	ggaagcaagt	41220
cacagtaaag	agcaaggag	tttggaatag	aaacaaatac	cagaatcaag	gatggctgat	41280
aacttttcaa	ttacgaagaa	cattaaaaaa	aatcacagaa	tcgtgaaact	caagggatca	41340
cataggggaat	ttcggaaaaa	aaaccaacc	tgtatgatgt	acttttgtac	atcacagttc	41400
gaaggtaaca	aggcaaagat	ataataagaa	gaaacctgtc	acgagaaact	ggaggaaaaa	41460
gagctgtgtc	ttcctacaag	tacactgata	caaattgcca	atgtgttcac	ctcagaaaca	41520
ctggaagcca	gataccaggg	aatatgttta	aaatgataat	caggaacaaa	aagagatcaa	41580
ccgggaatgc	tgaatccagc	aataaaatgc	cttgaagatc	atccatgtcg	gataaatgca	41640
tattgtgcac	tgccccaag	aaagaaaccg	gaaactgtaa	gaattggaaa	tcagcaggct	41700

BI OL0250W0SEQ_ST25. txt

tatgtaacaa	gagaggtgac	ccgaaggaat	taggtagaag	aagaattgaa	caagaaagga	41760
actttctgca	gcccacgtaa	tgaagaatcc	agcaattggc	aaatgtagat	agatgtaaat	41820
gcaaaatatt	ttcttgatca	aatttctata	tctttgtaaa	tgagagttga	ctacttgaaa	41880
caaaatgata	gcaagatatt	taacttcagc	atatgtagag	gtaagaattt	gaaatggtag	41940
cataaatcac	gaagggatta	attcgaagt	taccgttgta	agtttcttta	cctcatgcac	42000
gatggtgtgt	catattaata	aaagggtagt	gtgcgggttc	gaagggatat	tgcaaactct	42060
agagcaatca	caaaggtttg	aactctgagg	tttttggtat	aataagaata	gtccatgcat	42120
tcaaaagagg	gaagccaagg	aagaactaga	agtctttcaa	gagctcaggc	tcttatacat	42180
ccagttgctc	attgaaccag	cttcctggaa	tggagggctt	ggggttgaga	ctaggccaca	42240
agtctagagt	ctctagagag	acagtgttgg	aaccccatgg	cccataatac	atttcccatt	42300
ttctcaggca	gccagaggtc	atgaatgtga	ggatactggg	aggttggagc	aacgtttctg	42360
ggaggcataa	ggaagagcga	atgcttcaag	atccccgcag	cccaaactac	tcgcctgctt	42420
tgccccctaa	tgcatTTTTc	tctgctgctc	cgtagctgtc	cgacctcttc	agatctctta	42480
gtccaccctg	ccgtcttctt	ttatgccatg	gggtccactg	ttctttcaac	tcattccccct	42540
ttccctcagt	cccggagtag	ctgcggccag	cagagggtag	actgagagca	ggagagaagg	42600
acctgcctag	gaacccttct	tagagatact	gcattcctgcc	tgggagcaag	ttttccaggg	42660
cagctttgag	aagtcttggg	gaaacaaacc	tactaaacct	gacagacagt	aatactatTT	42720
gcacaatgct	tttctgtggg	aaaggtagag	ccttttctact	acgtattgag	tacatagagt	42780
gtgaggggtg	acctggaacg	gctatcctcc	tggatgacgt	gtgttttctg	aagaactaca	42840
tgttcgttgc	aactcccaca	ttagaatatg	aagtcctacc	gagagagata	cggagactag	42900
acagatacag	atgcatttgc	atgtgaatac	acaatcccac	aatacagacg	tcaaaaccca	42960
taccagttat	tccagagaga	tggattgggc	agaaggcaga	aggagaatac	tctgatcggt	43020
tttcggccac	gtgtgtgtgt	tatctcagt	tttctaagaa	gcgtttgcta	cttttagatTT	43080
tttatTTaaa	aaaaatagta	ataatctatt	aagtatgaga	gatgtgcaga	gaggattagt	43140
gatcgagagc	catttttgct	ggtggcaatc	atatggtact	tttaatggga	atattagaaa	43200
ggcaccggta	atgaccttgt	tgcagcacia	aggagagagt	gtgggggtgcc	cctgcatggt	43260
gtcccacctc	ttgtgacgtg	tatcgTTTTg	gaatttccag	tggcttgatc	atgaactact	43320
gcaggaatcc	agatgctgtg	gcagctcctt	attgtttatac	gagggatccc	ggtgtcagggt	43380
gggagtactg	caacctgacg	caatgctcag	acgcagaagg	gactgccgtc	gcgcctccga	43440
ctgttacccc	ggttccaagc	ctagaggctc	cttccgaaca	aggtaaggag	tctgtggcca	43500
gacatctaca	cgcttcgatg	ctgggatgaa	aagccatgga	aattcccact	gatgcagccg	43560
ccttcaatgg	taaacggatg	ctcgagtgtt	gcctgagttc	taccatgtag	gaggaagcct	43620
ccgtgcactc	tctgggggag	ccagcggagt	gatttctggt	gcaacgtggt	tgggctttgt	43680
ctttaggatg	ggcacaaacc	ctccaggggg	atcgacttca	aaattcacct	tgttgtaaaa	43740

BI OL0250W0SEQ_ST25. txt

cggtgtacct	cagtgtccca	gccaaaattt	ttattgtaac	atgctgtcag	gtgtgtcact	43800
ctttccaagc	cagtaagctt	ttccggggat	ttcttcaagt	agccagcatt	cagagcaatc	43860
ttcagcattg	cagattctga	gaaatgtggc	tctggagcct	gtcacccctc	agaaacctaa	43920
gagggctgca	ttgattccat	gtggccctgg	gtctatggag	cagtacatga	gctcccagtg	43980
ctctaaggct	cttcagccct	aggctttgaa	gggagtgatt	tctcagtatt	cttaaaccctc	44040
tttctgatga	cacttgtagc	tgtgaggggt	ctagagagaa	agagtagtag	actcctactt	44100
tactacaatt	caggatgcag	ggcatgagag	gattccctct	ctcctccaag	ggaagaagct	44160
tttggcgtgc	acacatccct	gagaagcaaa	gtgtctttgt	cttcagtcag	atacatagga	44220
ccgttttctg	ccccatggcc	cggaagccaa	aggccttggc	tttcatgata	aacgggtctag	44280
ggaacatgc	aaaatttcca	tgtctgtccc	aaactcttcc	cccgacagcc	aattaccacc	44340
tgcagcccg	attgccaaat	gcggtgccgt	ttgcatgaag	attcagtaga	gtttcctaga	44400
aaggtgctac	ctcgtgagct	cactttccaa	tgaggaatct	gatctgttgt	gtttctctaa	44460
gggtgcaggt	gaaatatttc	caagaactta	ctacagttct	agaatgggag	gaatctgttg	44520
ctttgggtgt	tggtttgttg	tcggttttct	cacatccatc	tgcttatgga	taaggaaaag	44580
agaacggctc	taattctcat	agactccttt	ctggttgtgt	cacaaatggc	ttcacatggt	44640
tctctatgct	cagagatact	cagcttgatt	tcccgtgttt	tcatttcagc	accgactgag	44700
caaaggcctg	gggtgcagga	gtgctacat	ggtaatggac	agagttatcg	aggcacatac	44760
tccaccactg	tcacaggaag	aacctgccaa	gcttgggtcat	ctatgacacc	acactcgcat	44820
agtcggaccc	cagaatacta	cccaaagtcg	tatgtctttg	ttctttacca	taagagaaga	44880
aagggccaag	tgaagtttct	gttacaagag	atgtgtctca	agctgagttc	tccgaactca	44940
acttgtgaca	gatgcagatg	gcgtagcaaa	atgtctcagg	atgattgcct	tggagctaag	45000
ggctctgagag	aagggaaatg	ttaagctccc	tctccttcct	cctagtttcta	ttgagcagaa	45060
gggaaatctg	gaggtgagaa	gatcacatta	tgaagaaagt	cagaatgaca	aaggaccaga	45120
cacttagatt	acccttccac	aacaccaact	aaacgtcaat	ggagactttc	cagttggaat	45180
tccgttattc	tggcttccac	ttcctgaagg	gaaggttgag	tttgcccttt	ctctctgggt	45240
tcaagaggaa	agaataggtg	cttattttat	gacaggtgaa	ttgatctgtt	tctatatcta	45300
cgtatattcc	gattgtcaga	aaaacactcg	ttcctaagta	ccagtggcct	gaagggatac	45360
aggttcccag	caagagaaga	tccaaggaag	gaaggcagat	gagagtcagc	acagagaggg	45420
atgctgaaaa	gtaaaagga	tgggtggatg	gagagaagcc	cgggtctgac	cacccaatgg	45480
ccaatatttt	ggccacaagc	gactaccaga	gacatggaaa	aatggtttct	acatgtggga	45540
caacagatgg	tagaggacct	agagaattga	gagaggggca	atgatgggct	ccactccgca	45600
gatgccttgg	ctttcttcct	ggataccctt	cctgcactga	atagcaagga	gatggagccc	45660
aagcagactg	tagccatctt	gctgaatgga	ggagagggat	tggagtttgg	gatgactgtg	45720
gtagctgaaa	tttttctagg	tctgctagaa	ataagaactg	gtttgtgtgg	aggaaaagag	45780

BI OL0250W0SEQ_ST25. txt

ctctacaaat	acgcatagaa	gtctcctcca	gtcgttggcc	tgacatgacg	ctgcctgtgc	45840
acaggaaatg	gttccacgag	aaagtgtggc	aaagaacatt	tactgagaaa	cagcaagtac	45900
aagagcacag	gaagctcaat	aaagaagaga	gagatcacat	agcactctgg	gatactggag	45960
ttcttcccag	ctagaccaga	gagtcctcac	ggagcacatt	gccaattcag	tggagacccc	46020
agaacagccg	taatttaaag	gtacacttag	tatattacta	gaataaagtc	agctgcagac	46080
aaccccttgc	acagctggaa	agcaagtgtc	caagcatcaa	atcggtttcc	aatcaatgaa	46140
gtgcctgtga	gaggaaatct	caactctctt	tagaagtaaa	caacaaagtc	gattgcctca	46200
gctatgcggt	atccgcagag	tgagtcctaa	atttaaaatc	tgactacatg	tagaaaagcg	46260
tttcgtgtga	cccatgacca	ggaaataaat	cgggtaatac	aaacaggctc	aggaatgaga	46320
gaaatgatta	gaattgcgtg	aaaatttgac	atatcagtat	gataactgat	ttcaaataat	46380
taaaaaaaca	acatgcaaga	aagcagatat	catatcaaga	gaaattaaca	gtacagaata	46440
gccaaattaa	attaaagagg	tagtataaaa	aaagtatgtc	ttaattgaaa	aaaattactg	46500
tatggccggc	tgatcaattt	agacgtttca	gaggaaaaca	ttaccaaca	cacaattcta	46560
gagaacctac	agaatgagct	acacacacac	acacacacac	acacacaaac	tgaaaacaca	46620
cccatactca	cacacacgca	gaaactcaca	agttctaaca	cacacagaca	cgcgcacccc	46680
tgaagaaaca	gtgaaatata	aaattaagcg	agcctcacag	acatgtagga	aaatatgaaa	46740
agatttcctg	catgtgggaa	gcaagtcaca	gtaaagagca	agggagtttg	gaatagaaac	46800
aaataccgga	atcaaggatg	gctgataact	tttcaattac	gaagaacatt	aaaaaaaaatc	46860
acagaatcgt	gaaactcaag	ggatcacata	gggaatttcg	gaaaaaaaaac	ccaacctgta	46920
tgatgtactt	ttgtacatca	cagttcgaag	gtaacaaggc	aaagatataa	taagaagaaa	46980
cctgtcacga	gaaactggag	gaaaaagagc	tgtgtcttcc	tacaagtaca	ctgatacaaa	47040
ttgccaatgt	gttcacctca	gaaacactgg	aagccagata	ccagggaata	ttgttaaaat	47100
gataatcagg	aacaaaaaga	gatcaaccgg	gaatgctgaa	tccagcaata	aaatgccttg	47160
aaggatcatcc	atgtcggata	aatgcatatt	gtgactgcc	ccaaagaaag	aaaccggaaa	47220
ctgtaagaat	tggaaatcag	caggcttatg	taacaagaga	ggtgaccgga	aggaattagg	47280
tagaagaaga	attgaacaag	aaaggaactt	tctgcagccc	acgtaatgaa	gaatccagca	47340
attggcaaata	gtagatagat	gtaaatacaa	aatattttct	tgatcaaatt	tctatatctt	47400
tgtaaataag	agttgactac	ttgaaacaaa	atgatagcaa	gatattttaac	ttcagcatat	47460
gtagaggtaa	gaatttgaaa	tggtagcata	aatcacgaag	ggattaattc	gaagtgtacc	47520
gttgtaagtt	tctttacctc	atgcacgatg	gtgtgtcata	ttaataaaaag	ggtactgtgc	47580
gggttcgaag	ggatattgca	aatcctagag	caatcacaaa	ggtttgaact	ctgaggtttt	47640
tggtataata	agaatagtcc	atgcattcaa	aagagggaag	ccaaggaaga	actagaagtc	47700
tttcaagagc	tcaggctctt	atacatccag	ttgctcattg	aaccagcttc	ctggaatgga	47760
gggtctgggg	ttgagactag	gccacaagtc	tagagtctct	agagagacag	tgttggaacc	47820

BI OL0250W0SEQ_ST25. txt

ccatggccca	taatacattt	cccattttct	caggcagcca	gaggtcatga	atgtgaggat	47880
actgggaggt	tggagcaacg	ttcttgggag	gcataaggaa	gagcgaatgc	ttcaagatcc	47940
ccgcagccca	aactactcgc	ctgctttgcc	ccctaattgca	tttttctctg	ctgctccgta	48000
gctgtccgac	ctcttcagat	ctcttagtcc	accctgccgt	cttcctttat	gccatgggtc	48060
ccactgttct	ttcaactcat	ccccctttcc	ctcagtcccc	gagtagctgc	ggccagcaga	48120
gggtagactg	agagcaggag	agaaggacct	gcctaggaac	cccttctaga	gatactgcat	48180
cctgcctggg	agcaagtttt	ccagggcagc	tttgagaagt	cttgagaaaa	caaacctact	48240
aaacctgaca	gacagtaata	ctatttgcac	aatgcttttc	tgtgggaaag	gtagagcctt	48300
ttcactacgt	attgagtaca	tagagtgtga	gggttgacct	ggaacggcta	tcctcctgga	48360
tgacgtgcgt	tttctgaaga	actacatgtt	cgttgcaact	cccacattag	aatatgaagt	48420
cctaccgaga	gagatacgga	gactagacag	atacagatgc	atttgcatgt	gaatacacaa	48480
tcccacaata	cagacgtcaa	aaccataacc	agttattcca	gagagatgga	ttgggcagaa	48540
ggcagaagga	gaatactctg	atcgtttttc	ggccacgtgt	gtgtgttatc	tcagtgtttc	48600
taagaagcgt	ttgctacttt	agatttttta	tttaaaaaaa	atagtaataa	tctattaagt	48660
atgagagatg	tgcagagagg	attagtgatc	gagagccatt	tttgctgggtg	gcaatcatat	48720
ggtactttta	atgggaatat	tagaaaggca	ccggtaatga	ccttgttgca	gcacaaagga	48780
gagagtgtgg	ggtgcccttg	catgttgtcc	cacctcttgt	gacgtgtatc	gttttggaat	48840
ttccagtggc	ttgatcatga	actactgcag	gaatccagat	gctgtggcag	ctccttattg	48900
ttatacgagg	gatcccgggtg	tcaggtggga	gtactgcaac	ctgacgcaat	gctcagacgc	48960
agaagggact	gccgtcgcgc	ctccgactgt	taccccggtt	ccaagcctag	aggctccttc	49020
cgaacaaggt	aaggagtctg	tggccagaca	tctacacgct	tcgatgctgg	gatgaaaagc	49080
catggaaatt	cccactgatg	cagccgcctt	caatggtaaa	cggatgctcg	agtgttgcc	49140
gagttctacc	atgtaggagg	aagcctccgt	gcactctctg	ggggagccag	cggagtgatt	49200
tctggtgcaa	cgtggttggg	ctttgtcttt	aggatgggca	caaaccctcc	agggggatcg	49260
acttcaaaat	tcaccttggt	gtaaaacggg	ctacctcagt	gtcccagcca	aaatttttat	49320
tgtaacatgc	tgtcagggtg	gtcactcttt	ccaagccagt	aagcttttcc	ggggatttct	49380
tcaagtagcc	agcattcaga	gcaatcttca	gcattgcaga	ttctgagaaa	tgtggctctg	49440
gagcctgtca	ccctcgagaa	acctaagagg	gctgcattga	ttccatgtgg	ccctgggtct	49500
atggagcagt	acatgagctc	ccagtgtctt	aaggctcttc	agccctaggc	tttgaaggga	49560
gtgatttctc	agtattctta	aacctctttc	tgatgacact	tgtacctgtg	aggggtctag	49620
agagaaagag	tagtagactc	ctactttact	acaattcagg	atgcagggca	tgagaggatt	49680
ccctctctcc	tccaagggaa	gaagcttttg	gcgtgcacac	atccctgaga	agcaaagtgt	49740
ctttgtcttc	agtcagatac	ataggaccgt	tttctgcccc	atggcccggg	agccaaaggc	49800
cttggctttc	atgatcaacg	gtctagggaa	acatgcaaaa	tttccatgtc	tgtcccaaac	49860

BI OL0250W0SEQ_ST25. txt

tctgcccccg	acagccaatt	accacctgca	gccccgattg	ccaaatgcgg	tgccgtttgc	49920
atgaagattc	agtagagttt	cctagaaagg	tgctacctcg	tgagctcact	ttccaatgag	49980
gaatctgata	tggtgtgttt	ctctaagggt	tcagggtgaaa	tatttccaag	aacttactac	50040
agttctagaa	tgaggaggaat	ctgttgcttt	ggtgtttgtt	tggttggtcgg	ttttctcaca	50100
tccatctgcc	tatggataag	gaaaagagaa	cggctgtaat	tctcatagac	tcctttctgg	50160
ttgtgtcaca	aatggcttca	catgtttctc	tatgctcaga	gatactcagc	ttgatttccc	50220
gtgttttcat	ttcagcaccg	actgagcaaa	ggcctgggggt	gcaggagtg	taccatggta	50280
atggacagag	ttatcgaggc	acatactcca	ccactgtcac	aggaagaacc	tgccaagctt	50340
ggtcatctat	gacaccacac	tcgcatagtc	ggaccccaga	atactacca	aatgcgtatg	50400
tctttgttct	ttaccataag	agaagaaagg	gccaagtga	gtttctgtta	caagagatgt	50460
gtctcaagct	gagttctccg	aactcaactt	gtgacagatg	cagatggcgt	agcaaaatgt	50520
ctcaggatga	ttgccttggg	gctaaggggtc	tgagagaagg	gaaatgttaa	gctccctctc	50580
cttcctccta	gttctattga	gcagaaggga	aatctggagg	tgaggagatc	acattatgaa	50640
gaaagtcaga	atgacaaagg	accagacact	tagattaccc	ttccacaaca	ccaactaaac	50700
gtcaatggag	actttccagt	tgggaattccg	ttattctggc	ttccacttcc	tgaagggag	50760
gttgcgtttg	ccttttctct	ctgggttcaa	gaggaaagaa	taggtgctta	tttatggaca	50820
ggatgaattga	tctgttttcta	tatctacgta	tattccgatt	gtcagaaaaa	cactcgttcc	50880
taagtaccag	tgccctgaag	ggatacaggt	tcccagcaag	agaagatcca	aggaaggaag	50940
gcagatgaga	gtcagcacag	agagggatgc	tgaaaagtaa	aagggatggg	tggatggaga	51000
gaagcccggg	tctgaccacc	caatggccaa	tattttggcc	acaagcgact	accagagaca	51060
tggaaaaatg	gtttctacat	gtgggacaac	agatggtaga	ggacctagag	aattgagaga	51120
ggggcaatga	tggtgtccac	tccgcagatg	ccttggtttt	cttcctggat	acccttcctg	51180
cactgaatag	caaggagatg	gagcccaagc	agactgtagc	catcttgctg	aatggaggag	51240
agggattgga	gtttgggatg	actgtggtag	ctgaaatttt	tctaggtctg	ctagaaataa	51300
gaactggttt	gtgtggagga	aaagagctct	acaatacgc	atagaagtct	cctccagtcg	51360
ttggcctgac	atgacgctgc	ctgtgcacag	gaaatggttc	cacgagaaa	tgtggcaaag	51420
aacatttact	gagaaacagc	aagtacaaga	gcacaggaag	ctcaataaag	aagagagaga	51480
tcacatagca	ctctgggata	ctggagttct	tcccagctag	accagagagt	cctcacggag	51540
cacattgcca	attcagtgga	gaccccagaa	cagccgtaat	ttaaaggtac	acttagaata	51600
ttactagaat	aaagtcagct	gcagacaacc	ccttgacacag	ctggaaagca	agtgtccaag	51660
catcaaatcg	gtttccaatc	aatgaagtgc	ctgtgagagg	aaatctcaac	tctctttaga	51720
agtaaacaac	aaagtcgatt	gcctcagcta	tgcggtatcc	gcagagttag	tcctaaattt	51780
aaaatctgac	tacatgtaga	aaagcgtttc	gtgtgaccca	tgaccaggaa	ataaatcggg	51840
taatacaaac	aggctcagga	atgagagaaa	tgattagaat	tgcttgaaaa	tttgacatat	51900

BI OL0250W0SEQ_ST25. txt

cagtatgata	actgatttca	aatattttaa	aaaacaacat	gcaagaaagc	agatatcata	51960
tcaagagaaa	ttaacagtac	agaatagcca	aattaaatta	aagagctagt	ataaaaaaag	52020
tatgtcttaa	ttgaaaaaaa	ttactgtatg	gccggctgat	caaattagac	gtttcagagg	52080
aaaacattac	ccaacacaca	attttagaga	acctacagaa	tgagctacac	acacacacac	52140
acacacacac	acacacacaa	actgaaaaca	caccatact	cacacacacg	cagaaactca	52200
caagttctaa	cacacacaga	cacgcgcacc	cctgaagaaa	cagtgaata	taaaattaag	52260
cgagcctcac	agacatgtag	gaaaatatga	aaagatttcc	tgcatgtggg	aagcaagtca	52320
cagtaaagag	caagggagtt	tataatagaa	acaatatcca	gaatcaagga	tggctgataa	52380
cttttcaatt	acgaagaaca	ttaaaaaaaaa	tcacagaatc	gtgaaactca	agggatcata	52440
tagggaattt	cggaaaaaaa	acccaacctg	tatgatgtac	ttttgtacat	cacagttcga	52500
aggtacaag	gcaaagatgt	aataagaaga	aacctgtcac	gagaaactgg	aggaaaaaga	52560
gctgtgtctt	cctacaagta	caactgataca	aattgccaat	gtgttcacct	cagaaacact	52620
ggaagccaga	taccagggaa	tattgttaaa	atgataatca	ggaacaaaaa	gagatcaacc	52680
gggaatgctg	aatccagcaa	taaaatgcct	tgaaggctcat	ccatgtcggg	taaatgcata	52740
ttgtgcactg	cccaaagaa	agaaaccgga	aactgtaaga	attggaaatc	agcaggctta	52800
tgtaacaaga	gaggtgaccc	gaaggaatta	ggtagaagaa	gaattgaaca	agaaaggaac	52860
tttctgcagc	ccacgtaatg	aagaatccag	caattggcaa	atgtagatag	atgtaaatgc	52920
aaaatatattt	cttgatcaaa	tttctatatc	tttgtaaattg	agagttgact	acttgaaaca	52980
aaatgatagc	aagatatatta	acttcagcat	atgtagaggt	aagaatttga	aatggtagca	53040
taaatcacga	agggtattaat	tcgaagtgtg	ccgttgtaag	tttctttacc	tcatgcacga	53100
tgggtgtgtca	tattaataaa	agggtactgt	gcgggttcga	agggatattg	caaatcctag	53160
agcaatcaca	aaggtttgaa	ctctgagggt	tttggtataa	taagaatagt	ccatgcattc	53220
aaaagagggg	agccaaggaa	gaactagaag	tctttcaaga	gctcaggctc	ttatacatcc	53280
agttgctcat	tgaaccagct	tcctggaatg	gaggggtctg	ggttgagact	aggccacaag	53340
tctagagtct	ctagagagac	agtgttgga	ccccatggcc	cataatacat	ttcccatttt	53400
ctcaggcagc	cagaggctcat	gaatgtgagg	atactgggag	gttgagcaa	cgttcttggg	53460
aggcataagg	aagagcgaat	gcttcaagat	ccccgcagcc	caaactactc	gcctgctttg	53520
ccccctaattg	catttttctc	tgctgtccg	tagctgtccg	acctcttcag	atctcttagt	53580
ccaccctgcc	gtcttccttt	atgccatggg	tcccactggt	ctttcaactc	atcccccttt	53640
ccctcagtcc	cggagtagct	gcggccagca	gagggtagac	tgagagcagg	agagaaggac	53700
ctgcctagga	accctttcta	gagatactgc	atcctgcctg	ggagcaagtt	ttccagggca	53760
gctttgagaa	gtcttgagaa	aacaaacctg	ctaaacctga	cagacagtaa	tactattttgc	53820
acaatgcttt	tctgtgggaa	aggtagagcc	ttttcactac	gtattgagta	catagagtgt	53880
gaggggttgac	ctggaacggc	tatcctcctg	gatgacgtgc	gttttctgaa	gaactacatg	53940

BI OL0250W0SEQ_ST25. txt

ttcgttgcaa	ctccacatt	agaatatgaa	gtcctaccga	gagagatacg	gagactagac	54000
agatacagat	gcatttgc	gtgaatacac	aatcccacaa	tacagacgtc	aaaacccata	54060
ccagttattc	cagagagatg	gattgggcag	aaggcagaag	gagaatactc	tgatcgtttt	54120
tcggccacgt	gtgtgtgtta	tctcagtgtt	tctaagaagc	gtttgctact	ttagattttt	54180
tatttaaaaa	aaatagtaat	aatctattaa	gtatgagaga	tgtgcagaga	cgattagtga	54240
tcgagagcca	tttttgctgg	tggcaatcat	atggtacttt	taatgggaat	attagaaagg	54300
caccggtaat	gaccttggtg	cagcaciaag	gagagagtgt	ggggtgcccc	tgcatgttgt	54360
cccacctctt	gtgacgtgta	tcgttttgga	atttccagt	gcttgatcat	gaactactgc	54420
aggaatccag	atgctgtggc	agctccttat	tgttatacga	gggatccccg	tgtcaggtgg	54480
gagtactgca	acctgacgca	atgctcagac	gcagaaggga	ctgccgtcgc	gcctccgact	54540
gttaccgccg	ttccaagcct	agaggctcct	tccgaacaag	gtaaggagtc	tgtggccaga	54600
catctacacg	cttcgatgct	gggatgaaaa	gccatggaaa	ttcccactga	tgcagccgcc	54660
ttcaatggta	aacggatgct	cgagtgttgc	ctgagttcta	ccatgtagga	ggaagcctcc	54720
gtgcactctc	tgggggagcc	agcggagtga	tttctggtgc	aacgtggttg	ggctttgtct	54780
ttaggatggg	cacaaacctt	ccagggggat	cgacttcaaa	attcaccttg	ttgtaaaacg	54840
ggctacctca	gtgtcccagc	caaaattttt	attgtaacat	gctgtcaggt	gtgtcactct	54900
ttccaagcca	gtaagctttt	ccggggattt	cttcaagtag	ccagcattca	gagcaatctt	54960
cagcattgca	gattctgaga	aatgtggctc	tggagcctgt	catcctcgag	aaacctaaca	55020
gggctgcatt	aattccatat	ggtcctgggt	ctatggagca	gtatatgagc	tcccaatgct	55080
ctaaggctct	tcagtcctag	gctttgaagg	gagtgatttc	tcagtgttct	taaacctctt	55140
tctgatggca	cttgtacctg	tgaggggtct	agagagaaag	gttagtagac	ttctccttta	55200
ctgcaattca	ggatgcaggg	catgagaaga	ttccctccct	cctccaaggg	aagaaggttt	55260
tggcgtgcac	acatccttga	gaagcaaagt	gtctttgcct	tcagtcagat	atataggatc	55320
gttttctgcc	ccatggcctg	gaagccagag	gccttggctt	tcatgatcaa	cgatctaggg	55380
aaacatgcaa	aatttccatg	tctttcccct	cctctgccct	cgacagccaa	ttaccacctg	55440
catcctgcat	tgccaaatgc	agtgcccttt	gtatgaacat	tcagtagagt	ttcatagaaa	55500
ggtgctactt	cgtgagcgca	ctttgcagt	agaaggagtc	tgttctgttc	tgtttttcta	55560
aggatttcag	gtgaaatatt	tcctagaact	tactacagtt	ctagattggg	aggaatctgt	55620
aggtttgctg	tatgtttttt	ggttggtttt	ctcccatcca	tctgcctaca	ggtaaggga	55680
agataacgtt	cataattctc	atagactcct	ttctggttgt	gtcataaatg	gcttcacata	55740
tttcgttatt	ctcagagata	ctcagtttat	ttcttgtgtt	ttcatttcag	caccgactga	55800
gcagaggcct	ggggtgcagg	agtgtacca	cggtaatgga	cagagttatc	gaggcacata	55860
ctccaccact	gtcactggaa	gaacctgcca	agcttgggtca	tctatgacac	cacactcgca	55920
tagtcggacc	ccagaatact	acccaaagtc	gtatgtcttt	gttctttacc	ataagagaat	55980

BI OL0250W0SEQ_ST25. txt

aaagggccaa	ctgaagtttc	tgtgacaaga	gacatgcttc	aagctgagtt	ctccgaactc	56040
aacttggtgc	agattcagat	ggtgtagcaa	aatgtctcag	gatgatttcc	ttggagctaa	56100
gggtctgaga	gaagagaaat	gttaagctgc	ctcaccttcc	tcctagtttt	gtggagcaga	56160
agggaaatga	ggaggcgagg	agatcacctt	atgaagaaag	tcagaatgac	gaaccaccaa	56220
acacttagat	tacccttgcc	caacacccac	taagcgtcaa	tgaagacttt	ccagttggaa	56280
ttccgttatt	ctgacttcca	attcctgaag	ggaagattgt	gtttgccttt	tctgtctggg	56340
ctcatgagga	aagtttatgt	gcttacttat	ggacaggtga	attgatctgt	ttctatttct	56400
acctgtattc	caatagggag	aaaatctctt	ggctctaagt	accagtggcc	tgaaaggata	56460
gaggttccca	gcaagagaag	atccaaggaa	ggaaggcaga	tgagagtcag	cacagagagg	56520
gatgctgaaa	agtaaaaggg	atgggtagat	ggatagaagc	cctgggtctga	ccaccccatg	56580
gccaatcatt	tggccataat	caacaaccaa	agacatggaa	aaatggtttc	tacatgtggg	56640
acaacagatg	gtagaggacc	tagagaattg	agagagggcc	aatgatgagc	tcaactccat	56700
agatgccttg	gctttcttcc	tggataccct	tcctgcactg	aatagcaagg	agatggagct	56760
caagcagcct	gtagccatct	agctgagcag	aggagagggg	ttggagtttg	ggatgactct	56820
ggtattttct	agggtccgcta	caaataagaa	ctggtttgtg	gaggaaagga	gctctacaaa	56880
tacgcataga	agtctcctcc	agtagttggc	ctcacatgac	actgcatgtg	cacagaaaat	56940
ggttctacag	aaagtgtggc	aaagaacatt	tactgagaaa	cagcaactac	aagagaacag	57000
caagctcaat	taagaagata	gagatcacat	agcactctgt	gttattggag	ttcttaccag	57060
ctagatgaga	gagtgtcac	ggaacacatt	gccaatcag	tggagacccc	agaacagcca	57120
taatttcaaa	gtacaattag	tatattacta	gaataaaggc	agctgcagac	aacccttgc	57180
acagctgaaa	agcaagtgtc	caagcatcaa	atgggtttcc	aatcaatgaa	gtgcctgtga	57240
gaggaaatct	caactctctt	cagaagtaaa	caacaaagtc	aattgcctca	gctatgcggt	57300
atccccagag	tgagtcctaa	attaaaaatt	tgactacgtg	tagaaaagaa	tttcgtgtga	57360
tccatgacca	gaaaataaat	caggcaatac	aaacaggctc	agaaatgaca	tcgataatta	57420
gaattgcatg	aaaatttgac	atatcagtat	gataactgat	ttcagatatt	taaaaaaagt	57480
gcaacaaagc	aggtatcata	tcaagacaaa	ttaatagtat	agaatagcca	aatcaaatta	57540
aagaactatt	atacaaaaag	tatgtcttaa	atgaagaaat	tactgtatgt	ccgcctgaaa	57600
aatttagatg	tttcagaaga	aaaaattaac	caaaaacaat	tctgcagaac	ctacagaatg	57660
agccacacac	acacacattc	aaaacacacc	catacacaca	cacatgcaaa	aactcacaag	57720
ttctaacaca	cacacaaaca	cacacacaca	tgcacatccc	taaagaaata	gggaaatata	57780
aaattaaccg	accctcagag	acatgcagga	aaatataaga	agatttcctg	catgtgggaa	57840
gcaagtcaca	gtaaagagca	agggagtttg	gagtagatac	aaataccgga	atcacggatg	57900
gctgataact	tttcaattat	gaagaacggt	agaaaaatca	cagattcatg	aaactaaagg	57960
gatcaaatag	gaaatttcga	gaaaaaaaac	tacatgatgc	acttctctac	atcacagttc	58020

BI OL0250W0SEQ_ST25. txt

aaaggtaaca	aggcaaggat	ataagaagaa	gaaacatctc	acgagaaact	ggagaaaaaa	58080
gagctgtgtc	ttcctagagt	acagtgatac	aaattgctaa	tgcgttcacc	tcagaaacac	58140
tggaagccag	ataccaggga	atattattaa	aatgataatg	aggaacaaga	agagatcaac	58200
cgagaatgct	gaatccagca	ataaaatgcc	ttgaagatca	tccatgttgg	ataaatgcat	58260
attgtgcact	gcccaaaaca	aagaaactgg	aaagtgtaa	actttggaat	cagcaggcct	58320
atgtagcaac	agaggtgacc	cgaaagaatt	aggtataaga	agaatagaag	aattgcatga	58380
aaatttgaca	tatgactaag	ataactatct	caaatactta	aaaaaagatg	aatatgtaat	58440
aaaacagata	aaatatcaaa	agaaagtaac	agtattgact	agccaaatca	aattaaagac	58500
ttagtgtaaa	aagctatgtc	ttaaaagaaa	aaattactgg	atggctgcct	gatcaattta	58560
gacatttctg	aataggaaac	taaccaaaaa	tcaattctac	agaaccaact	acacacatat	58620
atacacatac	aacacaccca	tacacaccca	cgcaaaaact	cacaagttca	cacacacaca	58680
cacacacaca	caaccctcaa	gaaatagtga	aatagaaaac	caaccgaacc	tcacagacat	58740
gttgcaaaat	aggaaaagat	ttcctgcata	tggaagcaa	gtcacagaaa	agagaacggg	58800
agattggaaa	cagaaacaaa	taccggaatc	aaggatggcc	gaaaactttt	cattgatcaa	58860
gaatattaac	aaaatcgcaa	aaacacgaaa	ttcaatgcat	caaataggcg	tttcgaaaaa	58920
aagaaaaaat	ctggtatgat	gcacttttgt	acttcacatt	ttcacggtaa	gaagacaaag	58980
atataataac	aagaaacttc	ttatgagaaa	ctggggaaaa	acaagctgtt	tcttgctaga	59040
agaacagtga	tacaaattgc	taatgcattc	tcgtcaaaaa	caactggaagc	cagataccgg	59100
gaatgttatt	aatgtggtaa	acaggaacaa	gaagagatca	accaagaatg	ctaaatccag	59160
caataaaatg	ccttgaagat	catccatgct	gcataaatgt	atgttgtgca	ctgccccaaa	59220
caaagaaacc	ggaaactgta	agaatttgga	atcagcaggc	tgatgtaaca	agagaggtga	59280
cccaaaggaa	ttaggtagaa	gaagaatagt	acaagaaggg	aactttctgc	agcccatgta	59340
atgaagaacc	cagcaattgg	caaattgtaga	tgtaaattgca	aaatattttc	ttgaccaa	59400
ttctatatat	ttttaaatga	gcgttgacta	ctggaaacaa	aatgatagca	atatatttaa	59460
ttttagcata	tgtagaggta	agaatttgaa	caagtagcgt	aatcatgta	gggaataatt	59520
agaagtgtac	cattgtaagt	ttcttacctc	atgcacaatg	gtatgtaata	ttaataaaat	59580
gttactgtgt	gggttcaagg	agatattgca	aatcctagag	caatcacaaa	gttttgaact	59640
ctgaggtata	ttgtataata	agaatattcc	atgtattcaa	aagagagaag	ccaaggaaga	59700
aagaaatttg	tcacgagttt	gggctcttag	tacatcctgt	agctcattga	accagcttcc	59760
tggaatggag	ggtctgggat	tgacactagg	ccacatgtat	agagtctcta	gagagacagt	59820
gtttcatccc	catggcccg	aatacatctc	ccattttctc	aggcagccac	aggatcatgaa	59880
tgtgaggata	gagagagggt	ggagcaacgt	tcttgggagg	cataaggaag	agcaaatgct	59940
tcaagatccc	cgcagcccaa	actcctacct	gctttgcccc	ctaatagcagt	gttcctccgt	60000
agctgtccga	cctcttcaga	tctcttagtc	taccctgcca	tcttccttta	tgccatgggt	60060

BI OL0250W0SEQ_ST25. txt

cccactgttc	tttcaactca	tccccctttc	cctcagtgca	gagtagctgc	ggccagcaga	60120
gggtagactg	agagcaggag	agaaggtcct	gcccaggaac	ccatttctaga	gatgctgcat	60180
tctgcctggg	agcaagtttt	ccagggcagc	tttgagaagt	cttgcagaaa	caaacctatt	60240
tgaccacat	gatatgggaa	tgacagaaa	taatacaatt	tgacacagtc	ttttccatgg	60300
gaaaagtaga	gccttttcgc	gaggttttga	gtacatagag	agtgaagggt	gacctggaaa	60360
ggttatcctc	ctggatccca	tgttttttct	gaagaactac	ctgttagttg	caacttgcac	60420
attagaatat	gaagtcctac	cgagagagat	acggagaact	agataaatac	agatactttt	60480
gtatgtgaat	aaacgattcc	acaatacaca	catcaaaatc	cataccagtt	attccagaga	60540
gatggattgg	gcagaaggca	gaaggagaat	actctgatcg	ttttttgccc	acgtgtatgt	60600
attatctcag	tgtttctaag	aagcgtttgc	tacttttagat	ttttttttat	aataataatc	60660
ttttaagtat	gagaaatgtg	cagacaggat	tagtgattga	gagccatttg	tgcttgtggc	60720
aatcatatgg	tacttttatg	ggaatattag	aaaggcactg	gtaatgacct	tgttgcagca	60780
caaaggagag	ggtgtggggg	gcccctgcat	attgtcccac	ctcttgtgac	gtgtatcggt	60840
ttggaatttc	cagtggcttg	atcatgaact	actgcaggaa	tccagatcct	gtggcagccc	60900
cttattgtta	tacgagggat	cccagtggtc	ggtgggagta	ctgcaacctg	acacaatgct	60960
cagacgcaga	agggactgcc	gtcgcgcctc	caactattac	cccgattcca	agcctagagg	61020
ctccttctga	acaaggtaag	gagcctgtgg	ccagaaacct	acacgtttcg	atgctgggat	61080
gaaaagccat	ggaaattccc	actgatgcag	cagcctccaa	tggtaaacgg	atgctcgagt	61140
gttgactgag	ttctgtcatg	taggaggaag	cctccgtgca	ctctctgggg	gagccagcgg	61200
attgatttct	ggtacaacgt	tgggtgggct	gtgtcttttag	aattggcaca	aaccctccag	61260
ggtgatcgac	ttcacaactc	acctcgttga	aaaatgggct	atctcagtgt	cttagccaaa	61320
atttttattg	taacatgctg	tcagatgtgt	gactctttcc	aagccagtaa	gcttttcctg	61380
ggacttcttc	aattagccag	cattcagtcg	aatcttcagc	attgcagatt	cagagaaatg	61440
tggctctgga	gcctgtcacc	cttgagaaac	agggctaaca	gggttgcat	aattccaaat	61500
caccctgggt	ctatggagca	gtacatgaac	tccaatgat	ctatgtttca	ggacttcctc	61560
agtcataagg	gggctctgca	gccctagggt	tttaagttag	tgactgcccc	gtgttctggt	61620
ggcagttgta	cctgtgagcg	gtctggatag	aaagagtcgg	agacttctgt	attattgcaa	61680
ctcaggatgt	gggtcatgag	aggatttcat	ctctcctgca	ggggagtaag	ctgttcgcct	61740
ccacccatcc	ctgataactg	aagtgtcttt	gtctgcagtc	ctagacgaag	gactgttgct	61800
tctcccatgg	cccagaagct	gaagaccttg	ccttttggtta	tgaaacgttc	attgttttca	61860
tgtctgtccg	tttctctgcc	cctaacaccc	aatcaccatg	tatggcctgt	acccccaaat	61920
gcatcgtgct	ttgctgtttg	ctgccccata	gtcctcatga	acattcagta	gaaattccca	61980
taaatgtgct	tgacgtgag	cacagtttcc	attgagaagc	cctctcattt	gtcctttttt	62040
tctaagcttt	tatgtgaaat	atttctaaga	acttactaca	gttctaaagt	gttaggaatt	62100

BI OL0250W0SEQ_ST25. txt

tgtttctttg	gtgtttttgt	ttgttggttg	gttggtgctt	ttctcaagtc	catctgccta	62160
caaataaaga	aacaagaatg	ttacttgtca	tattctcctg	aggtcataat	tctcagagac	62220
ttttttctgg	tttggtccat	aagtggcttc	acatgtttgt	ctcttcttgg	aaacactcag	62280
tttgatttct	tttcttttca	tttcagcacc	aactgagcaa	aggcctgggg	tgcaggagtg	62340
ctaccacgga	aatggacaga	gttatcaagg	cacatacttc	attactgtca	caggaagaac	62400
ctgccaagct	tggatcatcta	tgacaccaca	ctcgcatagt	cggaccccag	catactaccc	62460
aaatgcgtat	gtctattttc	tttaccataa	gtgaaggaag	ggtcagtgga	aatttctgtt	62520
agtagagtca	tgcttcaagc	tgagtgttca	ggactcaagt	tgtctcagat	gaacagtgca	62580
tagcaaaatg	tctcaggaac	attgtctttg	agcaaagagt	ctaagagaag	acaaatgtta	62640
atctggctct	ccttcctcct	agtttaatgg	agcagaaagg	tatctggagg	caaggatatc	62700
acattaagaa	acaagtcaag	atgacaaatg	atgaaactct	tagagtaccc	ttccacaaca	62760
cccactaagg	ttcaatgcag	ccttttctcc	ttggaattct	attaaactaa	actccaattc	62820
ctgaagtga	ggttctgttg	gggttttctg	ttttggctta	caaggaaagt	atatatgtat	62880
atctatggag	aggcaaactc	atctctttct	atatctacgt	ctattccaat	atgtagaaac	62940
acagtcggtt	ctgaccacca	gtggtctgaa	gggatactgg	ttgttagaga	ataaaaatgg	63000
caggaaggca	gatgagagtc	agcaaagaga	gagatcctgt	aaagtaaaag	ggtggataga	63060
tggacagaag	cccaggtctg	accagcccat	ggccaggctt	taggccataa	gtgacaccaa	63120
agacatggaa	aatgggtttc	tacatgttgg	acaacagaca	gtagtggacc	aaaagaatag	63180
tgacaggggg	aacaatgaga	tcaactccat	agataccttg	gctttcttcc	tggaggccct	63240
tcttgactg	aagagcaagg	tgatggagcc	cagatggact	gtagccatct	tcctgaatgc	63300
aggagagaga	ttggaatttg	ggactactgt	ggtagctagg	attttatagg	cctgctgaga	63360
atgagaatgg	atttgtggat	gaaaggagct	ccagggggcac	gcatagtagt	ctcctcgaat	63420
ctttggctaa	acatgacgtt	gcatgtgccc	agaaaaaggt	tccacaagaa	agtagagaaa	63480
agaatatatc	ctgaggaata	gcaactgcga	ttgaacagtg	agctcaataa	agaggacaga	63540
gccctcatag	cattctggga	tactggagtt	ctgaccagct	ggaggagaga	cctcactgaa	63600
cctcttggga	atacagtaga	gactccagaa	aagtcatact	ttaggagtag	aattagtaaa	63660
tttctagaaa	aaaaggcagc	tctagacaaa	ccctggcaaa	actgaaaagc	aagtctccaa	63720
gcattaaaat	catttccaag	tcaattaact	gcctggggaga	ggaaaaccct	ctttagaggt	63780
aaacaacaaa	gtcaagtggc	tcagctatgt	ggtgttcaca	gtgtgagttc	taaattttaa	63840
acttgactac	acatagagaa	gcttttagta	tgaacatga	ccaggtgaaa	aatcagtcaa	63900
tacaaataga	cctagaaatg	acagaaatga	ttagaatggc	aaaaaatttg	acatatcaat	63960
atgtcaactg	agttttaggt	tttaagaaaa	catgaatacg	gaatgaagca	gataccatat	64020
caagagacag	taacagtata	gaagagccaa	attaaattaa	agaactagta	taagaaggta	64080
tgtcttaa	gaaaaatta	ctggatgtat	tccaatgga	gtgagatgtt	tcagaagtaa	64140

BI OL0250W0SEQ_ST25. txt

aaactaactg	aaaaacaatt	ttataccacc	tacagaacca	gctacacata	cacaaatgac	64200
acacacatat	acacacatac	tcacacatgc	acaggcttag	aaacatgcac	gcacacacac	64260
acacacacac	acacacacct	ccacaaatac	taaaaaatga	aatccactga	tcctcacaga	64320
caggcgggaa	aataataaaa	gatttcctgc	atgtgggtag	gaagtcacag	aaggagagga	64380
aggagagatt	gctacaggaa	caaatactgg	aagcaaggat	agctaaaaac	ttttcaaata	64440
agaagaatat	taaaaaccac	agattcaaga	agctgaatga	atcagacagg	gaatttccaa	64500
aaaaaaaaaa	aaaaaaactg	tatgattcac	ttttgtacat	caccgttcaa	cagtcagaag	64560
gcaaagatat	aataacaaga	aacatctcat	gagaaactgg	aggaaaaaga	gctgtgtctt	64620
gctagaagaa	cagtataca	aattgcta	gcattctcat	cagaaacact	ggaacccagt	64680
taacagggga	tatcattaaa	atgataaact	agaaaaaaaa	gagatcaa	gagaatgcta	64740
catccagcaa	taaaatgcct	tgaagatcat	ccatgttggga	taaatgcata	ttgtgactg	64800
ccccaataa	ataaaccaa	aactaataat	ttggaatcag	caggcttgtg	taacaagaga	64860
tgttgcccaa	agaaaattag	ctagaagaag	aatagttcaa	gaggagaact	ttctgcagcc	64920
cacgtaatga	agaaccagc	aatggcaaa	tgtagatgta	aatgcaaaat	atcttcttga	64980
tcaaatttct	atatcttttt	aatgagagt	tgactacttg	aagcaaaatg	atagcaatat	65040
atttaacttt	agcatatgta	gaggtaaaaa	tttgaacata	tagactaaat	catgtgggga	65100
ataattggaa	gtgtaccatt	gtaagtttct	taccttatcc	acgatgggtat	gtaatattaa	65160
tgaaagggtg	aatttgtggg	tccaaaggga	tattgtaaat	cctaaagcaa	tcataaaatt	65220
ttgaattctg	agggatatta	tataataaga	atcttccatg	tatccaaaag	agggaagcca	65280
aggaagaaaa	agaagtcttt	caagtactca	agctctgagc	acatccagtt	gctcattgaa	65340
ccagcttcct	ggaatggagg	gtctgggctt	gagactaggt	cacatgtgta	gagtctctag	65400
agagacagtg	ttggatcccc	atggcccata	atacatttcc	cgttttccca	ggcagccaca	65460
ggtcacgaat	gggaggattc	tgagagggtg	gagcaatggt	cttaggaggc	ataaggagga	65520
gtgaatgctc	tgagatttcc	ccagcctgag	gtcctccata	gctgcccagc	ctcttcagac	65580
ctcatagtct	gccagctgt	ctccctttat	gccatgagtg	ccactgttct	ttcaactcat	65640
ccccattcc	ctcagtccc	gaattgctgt	ggccagcaga	ggatggactg	agagcaggag	65700
aggaagtcct	gaccaggaac	ccatcctaga	gatactgcat	cctgcctgaa	agctaggttt	65760
ccagggcagc	tttgagaagt	cttgcaaaa	gaaaccact	tgaccacact	gatacggtat	65820
cgacagacag	gaatactttt	tgtgcaatgg	ttttacatgc	tgaacataga	gccttttggc	65880
tacattttga	gtacattgaa	tgagactgct	ggcctgggaa	ggatatcatg	ctggatgcca	65940
ttttttctc	tggagaacta	tgtgttagtt	ccaactcgca	cattactata	tgaagtccta	66000
cacagagaga	tacggagagc	tagacagata	gagatacttt	tgtatgtgca	taaccaattc	66060
cacaatacac	acgtcaaaat	ccataccagt	tattccagag	agatggattg	ggcagaaggc	66120
agaaggagga	tattctgatc	cctttttggc	cacatgtatg	tataatctca	gtgtttctag	66180

BI OL0250W0SEQ_ST25. txt

gaagtgtgtg	ctgcattaga	tttttttct	ttaaaaaaag	tgataatata	ttaagtatga	66240
gaaatgtgca	gagaggatta	gagattgaga	gccatttgtc	atttgtggcaa	ttgtatggta	66300
tctcttttgg	gaatatttca	aaggcaccag	taatgacctt	gtttagtagcaa	aatatacagt	66360
gttcctgcat	atgtacccat	tttttgtgat	gtgtattctt	ttggaatttc	cagtggcctg	66420
atcaagaact	actgccgaaa	tccagatcct	gtggcagccc	cttgggtgtta	tacaacagat	66480
cccagtgca	gggtgggagta	ctgcaacctg	acacgatgct	cagatgcaga	atggactgcc	66540
ttcgtccctc	cgaatgttat	tctggctcca	agcctagagg	ctttttttga	acaaggtaag	66600
aagttgtgcc	agacatttac	ctgcttggat	gctgggatga	aaagccatgg	atacccccac	66660
tgacgcacaa	cccttcagtg	ctacactggg	tctcgtgtgt	tggttctggg	tctgccatgt	66720
gggaggaagc	cttagcgcac	tctctggggg	agccagaggt	gtgatttttg	gtgcaacctg	66780
tgcgagctgt	gtcttttagga	tgggcggaaa	ccattctggg	tgctcgactt	caccactccc	66840
ctcattgtaa	aaggggctat	ctcattgtcc	tagacaaaat	tcttattgta	atatgctgtc	66900
agatgtgtgt	gtctttccaa	gccagtaaac	ttttccaggg	atttcttcaa	gtagacagca	66960
ttcagtgcaa	tcttcagcat	tgcagattcc	gagaaatgtg	gctctagatc	ctgttatcct	67020
tgagaaacct	aactgggttg	cattaattcc	atatctccct	gggtctgtgg	agtagtacat	67080
gagctcccga	agctctatct	ctcaggtctt	tttcagtccg	aggcaggttg	tgcagttctt	67140
agctttgaag	ggagtgattt	tttcgtgtgc	ttttgcctct	ttctgatgga	acttgtacct	67200
gcgggggggtc	tggagaaaaa	gagtagtaga	cttttgcttt	attgcaatgc	attatgctgg	67260
gcacgagagg	attccctatc	ttattgtagg	tgataagctt	ttggcctcca	ctcatccctg	67320
agaagtgaag	tgttgttgcc	tacagtttta	gctgcaggac	tgttgtctgc	cccatcacca	67380
ggagtttaat	gctttctttt	ttgagcaatc	atctagggac	acatgcaagg	tttttatatg	67440
tccttgccctc	ctccccaaaa	aaccatttta	atgcttggag	acttgctttt	cagctttgcc	67500
aaatgcatca	ccctttcttc	tatgctgttc	catgtcgtca	tgaacactct	gtagagattc	67560
ctagaaatga	gcttccatgt	tagtggagtt	tccgatgaga	agcaatctga	tatttctttt	67620
ccactaagtt	ttacatgaaa	tatttctaag	aacttactac	agttctagaa	tggtaggcat	67680
ctcttacttt	cgtgttttgt	tgtgtgtttt	ctcatgtcca	tttgcctatt	aataaagaat	67740
agagaatggg	tgtaaatctc	agtgactctt	ttttggttta	tgtcataaat	ggcttcctgt	67800
atttttctgt	tctaggaaat	aataagcttg	atgtcttctg	ttttaatttc	agcactgact	67860
gaggaaaccc	ccgggggtaca	ggactgctac	taccattatg	gacagagtta	ccgaggcaca	67920
tactccacca	ctgtcacagg	aagaacttgc	caagcttggg	catctatgac	accacaccag	67980
catagtcgga	ccccagaaaa	ctacccaaat	gcgtacgtct	ttgttcttta	ccataagcga	68040
aggaagggcc	aatggaagtt	tctgttagaa	gagtcatgct	tcaaggtgac	tgctcaggac	68100
tcaacttggc	tcagatgcag	aggaacattt	cctgtgagca	aaagtcttta	gagaagactt	68160
tgtttttttg	agacagagtc	ttgcttttgt	gccaggctg	gagtgacgtg	gcatgatctc	68220

BI OL0250W0SEQ_ST25. txt

ggctcactgc	aagctccgcc	tcccgggttc	acaccattct	cctgcttcag	cctctctagc	68280
agctgggact	acaggcaccc	accaccacac	ccggctaatt	ttttgtatit	ttagtagaga	68340
cagggtttca	ctgttctagc	caggatggtc	ttgggtctct	gacctcgtga	tccgcctgcc	68400
tcagcctccc	aaagtgcctg	gattacaggc	gtgagccacc	gtgcctggct	gagaagacat	68460
ttttaagct	ggctctcctt	cctcctagtt	ttatggaagc	agaaggatat	atggagttga	68520
gaagatctta	ttaataaaac	agccgggatg	acaaatgacc	aaagagttag	agtatccttc	68580
tacaacatcg	gctgaggggt	aatacaacct	tttcaccttg	gaattctatc	attctaagct	68640
ctagtccctg	aagtgaatgt	tgtgttggcc	ttttgcatct	tgggtcacag	ggaattgata	68700
cttgcacatc	tatggagagg	caaatctttt	tctatctact	tctttttcaa	tgggtacaaa	68760
cacacttggg	cctgagcacc	agtggctctga	agagatacgg	tctgcccaga	ggagaagaac	68820
aaaggcagga	aagcagatga	gagtcagcaa	aggggcatg	ctgaaaagta	aaaggggctg	68880
gtagatggac	agaagccatg	atctggccat	tctatggcca	gtctttcggc	cataagtgc	68940
taccaaagac	acggcaaaac	ggtttcaca	tgttgaacaa	cagatgctag	aggaccaaga	69000
gtattgcaag	aggagaaaa	tgagatcaac	ccatcaatgc	cttggctttc	ttcaaggaga	69060
cccttcctgc	actgaagagc	aaggagatgg	agcccaagct	gactgtagcc	atgttgctga	69120
acagaggaga	gtgattggac	tttgggatta	ctcaggtagt	taggattttc	tagccatgct	69180
aagagtaaga	atggacttgt	ggaggatagg	agctccaggc	atagaagtct	cctcaagtgt	69240
tagtctaaac	ataaagcagc	acttgcatag	aagattttcc	acaagaaaat	atggcaaaaa	69300
aacaccatat	attgaggaac	aacaactaca	agggaacagt	gagcttaata	aaggtgacag	69360
agctcacata	gtgctctgga	atattggagt	tttgaccagc	tagagagaag	agacctcatt	69420
gaaaatcttg	ggcattcagt	agagacctca	gaaaagtcag	actttatgag	tagactttgt	69480
atattcctag	aataaaggca	gctccagaaa	aaacctagca	aagctgaaaa	gcaaactctc	69540
aagcattaaa	atgggtgtct	agtcaattaa	ctgccttcta	gaagaaaact	caacactctt	69600
tacaggtgaa	caacaaagtt	aagttgctga	gctatgcaat	atccacagtg	tgagtcctaa	69660
atttataact	ttactacaca	taaaaaagca	tttagtgtga	accataacca	ggaaaataat	69720
cagtcaataa	aaatagaacc	aggaatgata	gaaatgattt	aaatggcatg	agaatttgac	69780
atattagtat	cataactgca	ttgctggatt	taagaaaaca	taaacatgga	acgtaacaga	69840
tatcatatca	agggaagta	aaaggataaa	agagtcaaat	caaattaaag	gactattaaa	69900
aggatatct	taaatgaaaa	attcactgga	tgggtctcca	atcagggttag	ttgtttccag	69960
ggaaaaaatt	aactgaaaaa	taattcaata	gaatctacag	aaatagctgc	acatatatac	70020
acacaatggc	acacgtgcac	acaccacac	ccacacaggt	gtgaatccta	gagccacacg	70080
agcattgaaa	catagagaag	taaaaattgt	tcattgagga	atatgtagca	atgctcaatg	70140
tgttttacc	taataagagc	ttttgtgatg	tatgattgaa	aaactgacac	aactgaagag	70200
agaaatagat	aagccacac	tctgagttag	agatttcctt	gattctctca	ctatggttat	70260

BI OL0250W0SEQ_ST25. txt

aaatctttcc	caaacacaac	aggctagaac	aaatatgcag	aaaattagac	atagtatctt	70320
tgttctcaat	aaaaacgtcg	acctatttta	cattataccg	aactaccgag	tacacattaa	70380
agtgtgcatg	gagcattcac	tgaggtgtac	tctacacatg	accttccagc	aagtctccat	70440
agatttaaaa	gaattaaagt	catacagagt	gtgtcacttt	attctcccag	aataaagtga	70500
gatatgaata	atgagaagtt	tgccagcttc	tcaaataattt	gggagtcata	cgggtgcattt	70560
caaaatactc	tttgggacaa	agaaaacatc	actaaggaat	ttagaaaagt	tttgaactga	70620
gtaagaatat	aacacaattt	atccaaactt	aggagatgca	gtgaatgtct	ttaggcctttt	70680
acataatttt	agatgctctt	agggaaaaac	agaagcatgt	aataatcaag	atttcaaact	70740
gcaatttctca	aagtgtagtc	tagagaaacc	tgaggacctt	tgagtacctt	cagagacagt	70800
ccatgagggt	aaaggacttt	gctacgtgaa	aagtaagatg	ctattggccc	tttttacttt	70860
cattttccaa	caagagaaga	ggggagtttt	ccagcagtta	cataatatgt	aatggcatca	70920
tgtctctgat	ggctaagaaa	atgggcaatt	gttgactttg	tgtgttaaaa	aaattctcag	70980
tgttggtttc	ttatactata	aatattcatc	ttgtgttttg	aaaaagaaaa	gctctttgga	71040
atcccctatg	aacaaagact	ttgacagttg	ttgatctaag	accacagctt	aaatatctac	71100
acaagaaaaa	aaaaaaaaagc	aaataagagc	caaggaaagc	agatggaagg	aagtagtcca	71160
aaccagtgc	attcagtga	caagaaaaga	gaccaacaag	ggagttaaact	cttgaaacag	71220
aaagttgatt	ctttgaaaag	atccatatga	ttgaacacag	tctggctaaa	caaatgcacag	71280
accaatgagg	gtgcacaacc	atcaccatct	ggagtaacag	aggagaggtg	ccattactat	71340
agcatcttcc	agttctgaaa	gctgaaaaga	agattttgag	aacaattgta	tgtgaataaa	71400
ttcaggaatg	ttaatcatgt	gggccaattc	ctgaggaaga	caacaaatca	gcaaaccaga	71460
tgctgaatag	ttagtgtagt	cctgtagaga	gacatacaga	gaggctgaca	gagaaatatt	71520
tgatatgtgca	taaaacaatc	tacaagacac	acttcaaaat	caatctcagt	taatctggag	71580
gaacatattt	cacagaaggt	ggaaggaggg	tattctgatc	ctcttgtaca	ttgtacaaca	71640
ttgtacaatg	tacagagtat	aattgtacaa	gtacaattga	agttgtacaa	gtacaagtgc	71700
aacttgcaca	atgtacagag	taaacattga	tgtttactct	caattttctt	atggagcaca	71760
gatgactttg	gatgtgttac	aatatgaatg	ataatttgtc	tttgagatgt	tcgcagttgt	71820
ttagaagttg	aggaccattt	gtgcatatta	tgggaccttt	agtgaaaata	tttcaaagtc	71880
tctttttaca	ctttgttaca	gcaaaatgta	gagggcgcta	agtgcccttg	aatcttctcc	71940
catctctggt	gacctgtgtt	gttttgaaat	ttgcagtggc	ctgaccagga	actactgcag	72000
gaatccagat	gctgagattc	gcccttggtg	ttacaccatg	gatcccagtg	tcaggtggga	72060
gtactgcaac	ctgacacaat	gcctgggtgac	agaatcaagt	gtccttgcaa	ctctcacggt	72120
ggtcccagat	ccaagcacag	aggcttcttc	tgaagaaggt	aggaagtcta	tggccagaca	72180
accacaccct	aggacgttgg	gatgaaaaga	gttgcaaaat	cttagtgata	tagaagcctt	72240
ccatgctcac	acaattccaa	gtagaatgtg	gactcagggg	cagccactgg	gaaggaacac	72300

BI OL0250W0SEQ_ST25. txt

tcagcgcctt	ctctgggaga	accagagctg	tgatgtttgg	taccctgtga	aaggggtgga	72360
tctataggaa	gggtgcagac	cctctagggc	actggactta	ccactcccct	ggttattcaa	72420
aggatcattt	tagtgtctta	gccagaagaa	tattctaaca	ttttgccaaa	tttgtgaaga	72480
tttaccaagc	tcatgataag	cctttcatgg	tatttcttca	agtagtcagt	gttcattgca	72540
tctttggcct	tgcggtttcg	gaggaatgcg	gtttttgagt	ctgtcatcct	tgagaaacct	72600
aatatgactt	ttcttagttc	catatacttc	tgggtccagg	tagcagtaca	tagccaacaa	72660
atgctccatc	gttctggcct	atctccatct	taagccagtc	ctgcacaact	aggctttgat	72720
gggaggggatc	tctcagtgtt	cttgcccctc	cttctcatgg	aacatatatc	tgtgttggtc	72780
tctgagaaga	agagtagtgg	atatctactt	tgttgcaatg	cagaatcctg	ggccaaagat	72840
accagccatc	cctccaaggg	aataaaattt	tggccagtag	ccctctctga	gagacaattt	72900
gtctttgcct	acgagtccta	gatgcaggac	cgcttctgc	cccatcttca	agaagctgaa	72960
ggctttggct	ttggaggatc	agcagtctag	ggaaatgtgt	gacggtttca	tgtctgtccc	73020
cactgacagt	caatcaccac	ctacaacctg	cacagcctga	tgcatagcag	tctagtttcc	73080
tgccattattc	tcaggaacac	ccagaagatg	tctatattaa	agagcatgca	catgagtgca	73140
attttgactg	ataggcactc	tgatctttcc	tttgggtgcct	gtgtttttaa	ggaaatcttt	73200
ctaagaactc	gttaaagttc	tagaatgcta	tgaatctttg	ggttttatta	ttggtatgtc	73260
catctgcctg	ctagtacaga	acagagcatg	gtagtctttc	tcagagacaa	tgatcctgtt	73320
tcagtcacag	atttcttctg	atgcttctgt	gttctagaaa	ttactcagct	tgatttctcc	73380
tctttgaatt	tcagcaccaa	cggagcaaag	ccccggggtc	caggattgct	accatggtga	73440
tggaacagagt	tatcgaggct	cattctctac	cactgtcaca	ggaaggacat	gtcagtcttg	73500
gtcctctatg	acaccacact	ggcatcagag	gacaacagaa	tattatccaa	atgggtacaa	73560
ccttgagttt	tcttcaaaga	cagacagcag	cccccttaca	tttctcttgg	aagggccatg	73620
cttccaacta	acttcttatg	acaaatttat	ctcagatctg	gaatgtttgg	tagaatgtct	73680
caggcttctt	tcttcaggca	cagtgtctga	aaggagagaa	atgtcaggcc	agctctcttt	73740
tctcatagtt	gacagaagca	ggaggatatt	tgaaggtggt	gagtttctcat	gaatagaaag	73800
ctcaggacac	atggccacgt	gcttagaaat	agcaccattc	cacaatgccc	actaaagacc	73860
aatgcaatag	ttcaaccagg	gatttctgtc	attctaattct	ccaagtcctg	aagtgaaggt	73920
tgtattagcc	atgttcatct	tgggcaacaa	ataaaggata	tctatgttga	catccagatc	73980
ttccaatcac	tttctcctct	aacctgtacc	tgggttctga	gaacaaggta	tctgaagagc	74040
tatgtgttgc	cagcacatga	ggggcaaaaag	taggaaggca	gctgagagtc	aggaagtata	74100
aagattctga	agagttacac	atgcaggaag	atggacagaa	accagttca	gaccacgtca	74160
gcgtttctgc	catgaaggac	tatcaaatac	ataggaaaag	tgttttcata	ggttggacaa	74220
cagacatgac	aggcctgaga	aaattcagaa	agggaatcaa	aggagatcaa	ccttatcatg	74280
tccctggcat	ccttccttga	gacccttgaa	gggcaagcag	atggagccca	gctgaccaca	74340

BI OL0250W0SEQ_ST25. txt

gcagtcttgc	ttaactgagg	agagagactg	gagtttgtga	tgccctcaggc	atctgacgta	74400
ttctaggctg	gctaagaatg	agaggggatt	tgtggaggaa	aggagctcca	agaatacaca	74460
ccgaagtctt	ctcaaggctt	tggctaaata	caaagctgcg	tatgcacaag	gagagttttc	74520
acaaagaaag	aacaataaag	aaaagctact	ggggaaagaa	caactgcaag	ggaacagtga	74580
gctcaatgga	gatgctagag	ctcacatagc	actgggggat	atttgagttc	tgaccactca	74640
gaggagagac	acctcactga	acatcttggg	cattcagtag	aggtcaaaga	aagccataat	74700
ttgggagtag	gatcttcgga	ttcctagaaa	taagggtgact	ccagaaacac	tccagcaacc	74760
cttcttccaa	gccagtctaa	aaggatccaa	atgatttcca	agtaaattaa	ctgccttcca	74820
gaaaaaagta	aactcaaccc	tccttagagg	taaggaacga	atacaagttt	ctcagttata	74880
tgacatcccc	agagtgaac	ttgcatttaa	aaatttacta	gacacaaaag	aagttttcac	74940
tgtgatccat	aactgggaga	aaaatcactc	aacacaaata	ggcccagaaa	taatagaaat	75000
tatggcattg	gcaagaacat	ttaaaatgca	cctctgagaa	ctgtgtttca	ggaaaatgtc	75060
agcaaaagct	gaccatgaga	gaaatgaatg	cataatatca	gaaaagaaaa	gaattgaaga	75120
gccaaatgga	aatttaaaaa	ctgagaaaag	ttatatctgt	aatgaggaat	tcactggatg	75180
gccttataac	cagtttagat	attatggtag	gaaaagggtga	acgagaaaat	gattcaatta	75240
aagctagaca	aaccacaaga	cagacagaca	gacacaaata	cacatacaca	caatgactga	75300
accaattaat	caacagagcc	tcaaggacat	ctagggaaaac	atccacacat	ttaatatatg	75360
tgtaggcaa	gtcacagaaa	gagaggaaaa	agataatgtg	acagaagtta	tacttgaagc	75420
catgacggct	gacaaatttc	caaacataca	gaaaatgaga	aattcatagt	catgaagctc	75480
aatgactcag	gtatagattt	ttaaagagca	aaactctgat	ttactggggt	acatcatagt	75540
taaattgtct	gatttcaaag	ctaagaagaa	aaaaaggggg	ttcctatgaa	caaacatttt	75600
gacagttgat	ctaagaccac	agcttaaata	tctaggcaag	gaaaagcaaa	taagacacaa	75660
ggaaagggga	tgatggaaa	tagtccaaac	caatgacatt	cagtgaacaa	gaaaatagac	75720
caacaaagga	gtaaatccat	gaaacagaaa	gttggttctt	tgaaaagatt	catgtgattg	75780
accacagtct	ggctgaacag	atgacagacc	aaggaggggag	tacaaccatc	accatttgaa	75840
gtaacagggg	agaggagcca	ttgctatacc	atactccagg	tctgaaagct	gacaagaaga	75900
tatcaagaaa	aactgtatgt	gaataaattc	atgaatgtag	atcatgtgga	tcaattcctt	75960
aggtaaacia	caaatcagca	aaccagatac	tgaatagatt	gggtactcct	atagaaagac	76020
atacagatag	ccagacagag	aaacatttgt	acgtgcataa	aacaatctac	aagactcact	76080
tcaaaatctc	tcagttaatc	caaagtaaca	tatttggcag	aagggtggaag	gagggatttc	76140
tgatcctttc	ttgtacacat	tgatgttttc	tctcggtttt	cttatggagt	atagacgagt	76200
ttggatgtgt	tacaataaga	atgataatct	gtctttgaaa	tgttcacagt	tgtttagaag	76260
ttgaggacga	tttgtgattg	ttacaggacc	tttagtgaga	atatttcaaa	gtcacttttt	76320
accactttgt	tacaacaaaa	tgtagaggat	gtctgggtgcc	cttgtatctt	ctcccatctc	76380

BI OL0250W0SEQ_ST25. txt

tggtgaactg	tattgttttg	taatttgcag	tggcctgacc	aggaactact	gcaggaatcc	76440
agatgctgag	attagtcctt	ggtgttatac	catggatccc	aatgtcagat	gggagtactg	76500
caacctgaca	caatgtccag	tgacagaatc	aagtgtcctt	gcgacgtcca	cggctgtttc	76560
tgaacaaggt	aagaagtctc	tggccagaca	accacaccct	tggacgttgg	gataaaaaga	76620
gttgcaaaat	cttagtgata	cagaagcctt	ccatgctgca	cgggaatctg	aatgtggact	76680
cagggtcagc	caatgggaag	gaagcctcag	cgccttctct	gggggaacca	gggctgagat	76740
ttttggcacc	ccgtgacagg	gtggtgtctt	taggaagcgt	gcagaccttc	tagggcactg	76800
gatttaccac	tcccctgggt	attcaataga	ttatttcagt	gtcctagtga	aaatggatat	76860
tctaacatcc	tgccaaatth	gtgatgattt	accaagctca	tcatgagcct	ttcctggtat	76920
ttcttcaagt	agacagtact	cattgcaaac	ttcagcttta	cagtttcaga	ggaatgtggt	76980
ttttgagtct	gtcatccttg	agaaacctga	tatgacttta	cttagttcca	tatcctcctg	77040
ggtctaggta	acagtacata	gccagcaaht	gctctatctc	cctgtctacc	ttaatcttag	77100
gcaggtgctg	cacacctagg	ctttgatgga	agggatttct	tagtgttctt	gcccctcctt	77160
ctcatggaac	acgtatctgt	gttgctgttt	gtgaagaaga	gtagtggatg	tctactttgt	77220
tgcaatgcag	gatcctgggc	ccaagatttc	ccgccgtccc	tccaagggaa	taaaattttg	77280
gccagtaccc	ctctctgaga	gacaatgtgt	ctttgcctgg	aagtcctaga	tggaggacca	77340
cttcctgccc	catcttcag	aaacttaagg	ctttggcttt	ggaggatcag	tgctctggag	77400
aaatgtgtga	cggtttcatg	tctgccccca	ctgacaacca	ccacctacag	cctgcaccgc	77460
ctgatgcatg	gcactctggt	ctcctgcctt	gttctcagga	acacccaaaa	gagatctttg	77520
ccaaagaaca	ggcacatgag	tgcaattttg	actgataggc	actctgatct	gtcctttggt	77580
gcccagggtt	taaagaaaat	ctttctaaaa	actcattgaa	gttccagaat	gctatgaatc	77640
tttgagcttt	gttattggca	tgtccatctg	cctactaatg	tagaacagag	catggtcgtc	77700
attttcagag	atgatgtcct	gtttctatca	tggatttttt	ttctcatgct	tctgtgttct	77760
ggaaattact	cagtttgttt	tctcctcttt	gaatttcagc	accaacggag	caaagcccca	77820
cagtccagga	ctgctacat	ggtgatggac	agagttaatc	aggctcattc	tccaccactg	77880
ttacaggaag	gacatgtcag	tcttggctct	ctatgacacc	acactggcat	cagagaacca	77940
cagaatacta	cccaaattgg	tatgtctttg	agttttctcc	caagagaaac	agccaccac	78000
ttaaattttct	cctggaagag	ccatgcttcc	agctaacttc	ttatgacca	atttctctca	78060
gaccagaat	gttggaacaga	atgtctcagg	cttcttgctt	tgggcacagg	gtctgagagg	78120
agagaaatgt	caggccagct	ctcttttctc	atagttgata	gaagtaggag	gatacttgga	78180
ggtggtgagg	tctcatgaat	agaaagctca	gaagaacata	tgaccatgtg	cttagaaata	78240
gcaccattcc	acaatgcccc	ctaaagacca	gtgaaatagt	tcaaccaggg	aattctgtca	78300
ttctaattct	caagccctgg	agtgaagggt	gtgtttgcca	tgtttgtctt	gggtaacaag	78360
tgaaggatat	ctatatgtac	ttcgagatct	tccgatcact	ttctcctcta	acctgtataa	78420

BI OL0250W0SEQ_ST25. txt

acacattggg	ttctgagaac	aaggtgtctg	aaaagctatg	tgttgccagc	ccatgagggg	78480
caaaaggagg	aaggcagctg	agagtcagga	agtatagaga	tgctgaagag	ttacacattc	78540
aggaagatgg	acagaaaccc	atgtctggct	atgccagcct	ttctgccatg	aaggactatc	78600
aaatacatga	gaaaacagtt	ttcacagggt	ggacaacaga	tatggtaggc	ttgagagAAC	78660
tgagaaaggg	aatcaaagga	gatcaacttc	atcattaacc	tgtcttcctt	cctggacaca	78720
gtgttggatt	gaaggacaag	cagatggagc	ccagctgacc	acagcagtct	tgcttaactg	78780
aggagagaga	ctggagtctg	cgatgcctca	ggcagctgat	gtgttctagg	ctggctaaga	78840
atgagaaggg	atttgtggaa	gaaaggagct	ccaggaatac	acacagaagt	ctcctcaagg	78900
ctttggctaa	atacaaagct	gcgtatgcac	agggagagtt	ttcataaaga	aagaacaaca	78960
aagaaaagct	acttgggaaa	gaacaactgc	aggggaacag	taagctcaat	ggagatgcca	79020
gagctcacat	agcactgggg	gatattttgaa	ttctgaccac	tcagaggaga	aacacctcac	79080
tacatttttg	gcattcagta	gagaccaaag	aaagctgtat	tttgggattg	ggatcatctt	79140
attcctagaa	tcaaggtgac	tccagaaaaa	ctccaacaac	ccttcttcca	agccagtcta	79200
aaaggatcca	aatgatctcc	aagtaaatta	actgcattcc	acaagaaaaa	aaaaactcaa	79260
cccccttag	aggcaagggg	caaatacaag	ttgctcagtt	atatggcatt	cctattgcgt	79320
tacttctatt	taaaaattta	atagagacac	aagaagcttt	cactgtgata	cataactggg	79380
agaaaaaatc	actcaacaca	aacaggccca	gaaattatag	aattgatgac	attggtgaga	79440
acatttaaaa	tgcacctctg	agaactgtgt	ttcaggaaaa	tgtcagcaaa	agctgaccat	79500
gagagaaaca	aaagcagaat	agcaagagaa	aagaaaagaa	ccggagagcc	aaatgaaaat	79560
taaagaactg	agaaaaggta	catctctaata	gaagaactca	ctggatggcc	ttatcatcac	79620
tttagacatt	acggtaggaa	aggtgacctt	gaaaataatt	caataggagc	tacacaaatc	79680
acaggacaga	cagacagacc	aacagacaga	aacacacaca	cacacacaca	cacacacaca	79740
cacacacaca	cacacacaca	aagactgaac	ctattaatca	acagagcctc	aagggcattct	79800
aggaaaaatc	cacacattta	atatatgtgt	taggcaagtc	acagaaggag	aagaaaaaga	79860
tatcatgaca	gacattatac	ttgaagcgat	gatggctcgc	aacacgccaa	atatacagaa	79920
aacaagaaac	tcatagtcaa	gaagctaaat	gactcaggta	tagaatttta	aagagcaaaa	79980
ctctatgatt	tactgggata	tatcatagtt	aagttgcctc	aattcaaagc	taaaaagaaa	80040
aaaagggggg	tcctatgaac	aacagctttg	acagctgttg	atctaagacc	acagcttaaa	80100
tatctaggca	aggaaaagca	aataaggcac	aaggaaagag	gatggaagga	aatagtccaa	80160
accaatgaca	ttcagtggaa	aagaaaatag	accaacaaag	gagtaaattcc	atgaaacaga	80220
aagttagggt	ctttgaaaag	tctatatgat	tgGCCaaagt	ctggctaaac	agatgacaga	80280
ccaaggaggg	agcatatcca	tcacatcat	gagtaacagg	agagagatgc	catttgctata	80340
gcatcctcca	ggtgtgaaag	ctgagaagta	gatattgaga	tcaactgtat	gtaaataaat	80400
tcatgaatgt	agatcatgtg	gatggattgc	ttaggtaaata	aacaaatcag	caaatcaaac	80460

BI OL0250W0SEQ_ST25. txt

actgaataga	tcatgcagtt	ttatagagac	ttacagacag	cctgacagat	aaacatttgt	80520
atgtacgtga	aacaatctcc	aagacacact	tcaaaatccc	tctcggttaa	tccaaaggaa	80580
tgtatttggc	agaaggtaga	aggagggtat	tctgatcctt	tctggtacac	attgatgttt	80640
tctctcagtt	ttcttataaa	gcatagatta	ctttgaatgt	gttacaataa	gaatcataag	80700
ctgtctttga	aatgttgaca	gttgtttaga	agttgaggac	catttgtag	tgttatggga	80760
ctttagttag	aatatttcaa	atttgcttgt	ttacactttg	ttacaagaaa	acatagaggg	80820
tgccagggtg	tgctgtatct	tctccaatct	ctggtgacct	gtattgtttt	ggaatttgca	80880
gtggcctgac	caggaactac	tgcaggaatc	cagatgctga	gattcgccct	tggtgttata	80940
ccatggatcc	cagtgtcaga	tgggagtact	gcaacctgac	gcaatgtcca	gtgatggaat	81000
caactctcct	cacaactccc	acggtggtcc	cagttccaag	cacagagctt	ccttctgaag	81060
aaggaagaa	gcctgcagtc	agacaacat	accctcggac	attgggataa	aaagatttgc	81120
aaaatctttg	tgatgcagaa	aacttccatg	ctgcacagga	agtcgaaggt	gaagtcatgg	81180
acagccaatg	ggaaggaagc	ttcagtgctt	tctctggggg	gaccagagct	gggatgttga	81240
gtgccttgtg	agggatggtg	tctttaaaag	gggcacagac	cctctaggac	actggattta	81300
tcacttccct	gttatcaaac	gaatcatatt	agtgtcctag	ccaagatgga	tattctaaca	81360
tcctgcaaaa	cttggaaga	tataccaagc	tcctaagcct	gtccagccct	ttcttcaagt	81420
aggcagtgtt	tattgcagtc	ttcagcttta	ccattttgaa	ggaatgccat	ttttgaggct	81480
gttgttcttg	agaaacctaa	catgtcttca	ttagatccgt	attgtcctga	gactttgaag	81540
cagtacatag	ccaccaaatt	gtttatctcc	ccagcctacc	ttcatcttgg	gcatgccttc	81600
cacacctagg	atttgaggga	agggatttct	cagtgttctc	atccctgctt	ctcatggaac	81660
atttatctcc	gttgtttttt	gagaagaaga	gtagtggatg	tcagctttct	tgtaatgagg	81720
gatcctgggc	ccaagattcc	ctgtctcccc	tcctaggcta	taaaattttg	gcctgtactc	81780
cttctccctg	agaggcaatg	tgtctttacc	tacaagtcct	agatgcaaga	tccttttctg	81840
ccccacaccc	cagaatctga	aggcttttgc	tttgaggag	cagtggctta	gtgtgcaagg	81900
gtttcatgta	tacccccac	taacagccaa	tcaccaccta	tagcctgaac	agcttgatgc	81960
atggcaccct	ggtctcctgc	cttgttctca	tgaacacca	gaagagggtg	aagcaaaaga	82020
ccattcacat	gagtgttaatt	ttgaagtata	ggcactctga	tctgtttttt	gtttgtttct	82080
ttgtttgttt	gttttccagg	gttgaattaa	aatatttatg	actacttatt	aaatttctag	82140
aatcctataa	gtctatttgt	atttttattc	tacatttcaa	tttgcattgct	aatatagaag	82200
agtgtaaaatt	gttaatcctc	agattattcc	actttgtgtg	tcataatttt	tttcacattt	82260
cccttttcta	ggcaatactg	agcttgattt	tctcttttaa	tttcagcacc	aactgaaaac	82320
agcactgggg	tccaggactg	ctaccgaggt	gatggacaga	gttatcgagg	cacactctcc	82380
accactatca	caggaagaac	atgtcagctt	tggtcgtcta	tgacaccaca	ttggcatcgg	82440
aggatcccat	tatactatcc	aatgcgtat	gtctatcatg	ttagccataa	aaggaacaat	82500

BI OL0250W0SEQ_ST25. txt

agtcaactaa aatttctctt agctggccca tgctacaagc tcacttccta ggtccaaatt	82560
tctcatagac tcagagtttg tagcaaaatg tctcaggaaa cttacttttg agcaaaaggt	82620
ctgaatgaag agaagtttta ggattgctat ctttcataac aatttgatgg aagcagcagg	82680
atatatggag gtggtgaagt ctcattaatg taaagctaag gagatcaaat gaccaaatgc	82740
tgagacaaag tatcattcca caatgcccac taaagggtcca tgcagtcttt caaccatgca	82800
attctatcat tctatcctcc attccctgaa gtgaaatttg tgtttgccat ttttgacacg	82860
aatcagaagt aacaaattca ggctgggtgc agtggctcag gcctgtgatc ccaacacttt	82920
gggaggacaa gacgggcaga tcaccagagg tcaggagttc aagaccagcc tggctaacat	82980
ggcaaaaccc catctctacg aaaaattaaa aaattagccg gtcattggtgg tgggtacctg	83040
taattccaac tacttgggag gctgaggcag gagaaacact tgagcctggg attcagagtt	83100
tgctgtgagc cgagaacatg ccactgcact ccagcctggg tgacagagca agactcaatc	83160
tcaaaaaaaaa aaaaaaagaa gaagaagaag aaaagaagaa gaggaagaag aagaagagga	83220
agaagaagaa gaagaagaag aggaagagga agaggaggag gaggaggagg aggaagaaga	83280
agaagaagaa gaagaagaag aagaagaaga agaagaagaa gaagaagaag aagaagaaga	83340
aaatagaaat gagtgcataat atttatatat gagtactagc ctgtatgaac aacttgggtt	83400
ctaagcacca gttttctgaa gggatatggg ttgtcaggca gagtaaaagc aggaatgcag	83460
atgagagtca ggaagtaaac agatgtggtg attaaaatgg gcagggtacat ggacaaaaaa	83520
atgcatgtct gacaaaaact ggcctcttgc cataagttag tatgaataat atggaaaaac	83580
tgtttgcaca tgttgaacag cagacagtac aacctgagat agtttagaaa gggaaacaaa	83640
taagatcaac ccataatta cccttcctag acttaagggc aaagagtttt aaccaaaagca	83700
ttccacagca gtcttgctaa actggggaga gagactggag ttttgtttac taataaaacc	83760
gagattttct aggttaggta ataattgagaa agtattttgt gagaaaagga gctccaggaa	83820
tacacacaga agtctcttca agtctctggc tgaacagaaa gctgtgtatg cacagaaaga	83880
gtttccagag agaaaggaga acaaagaaca gctactgggg aaagaacaac tgctggggaa	83940
cagttagctc aatgaagatg ccagagctca catagcactg ggagggtattt gagctctgac	84000
cagcctgagg agagacactt cattgaacat cttgggcatt cagcaaagac ccaaaaaaac	84060
catacttcag gagtagaatt aatgcattcc tagaataaag tctactccag aaacacccta	84120
gaaaagctta gaaaccaagt ctaaaaagat ccaaatgata tccaagtaaa ttaattgcct	84180
gtcagaagaa aacaacctct tcagaggtaa acaacaaaat taaattgctc aatttatatag	84240
tatgcacaat gtgtggcata catttaaaaa tttgctaaac atacaaaaag catttagtgt	84300
gaccataac caggagaaaa atcagtcaat acaaatagac ccaaaaatga taaaaataac	84360
agaattggca aggagattta aaatgtatgt atcataattg tgttcaagga tttaaagaaa	84420
gcgtggacaa gaaataaata aatggataat atcaacagaa agaaaaattg taaaaggacc	84480
aaatggagag tcaagaactg aaaaaaaga catctcttta atgagaaaat cactacatgg	84540

BI OL0250W0SEQ_ST25. txt

ccttataatc	atattagata	gtacagatga	taaagctaac	tagaaaatat	taggggtggtg	84600
caaaccatag	cacgcttata	caaagcctga	gaagataaac	agagcctcaa	ggacatctat	84660
gaaaatatca	aaatatttaa	tatttgttta	aagcaagtca	cagaggaagg	gaaagagata	84720
ttggaacaga	aaaaatactt	gaagcagtga	tggctgatga	ctttctaaat	atggaaaaaa	84780
tgataaactc	acatagtcaa	gaagctcaat	ggatcagata	taggatttta	aaaagtaaag	84840
ctgtatgatt	tatttggaac	catcataatt	aaattgtcca	taatcaaaga	tagaaagtaa	84900
aatcttattt	gaagcccaag	ggaaaaaaca	tacctttaca	tagagtaaca	gtgacacaaa	84960
tgactgatgc	cttctcatca	gaaacaacac	aaatcagaaa	caatagaata	acacctttag	85020
agtggtaaga	agaaaaaaag	atcaaatacag	aaacaacaaa	ataacacggt	tagagtggta	85080
aggaggaaaa	caagatcaaa	tcagaaacaa	tggataaaca	ccttttagagt	gtaagaaaga	85140
aaaaaagatc	aaatcaggaa	caacagaata	acgccttcag	agtggtaaga	aggaaaacaa	85200
gataaaatca	gaaacaatga	aataacacct	ttagagtagt	aagaagaaga	aaagatcagg	85260
tcagaaaaaa	tggataaata	tgctaagaag	aaaaaaaag	atcaagtcag	aaacaatgga	85320
ataacacctt	tagagtgaag	agaaggaaaa	aaaccagca	agcttaaacg	ctatgcacag	85380
caaacaattc	cactgaaaat	gaatgttacg	taagtacata	ttctgtcctc	ctaaaaacaa	85440
agaacaaata	aaagaatgtt	tcatcagcag	gattatgtaa	taaaagatgt	gaaagaatgc	85500
tatgtaagta	gaagaaaaat	aataccatat	gggaattggc	atcaaaacca	caaaatacta	85560
tcaaaacaaa	aaaactttat	tgataaat	aacacaatat	gcaaaagaac	tataccatgt	85620
atactacata	acattggtga	gaagaaaatt	agaagatcta	aataaagaca	catcatgctt	85680
atagattaaa	aaatccaatg	tcacttttca	caaaactgat	ctttagtttc	aaccacacacc	85740
caagcagaat	tcctgcagtc	ttttcttgaa	aacctaacag	aatgtatatg	ctagaatcac	85800
caagacaatc	tttaaaaaga	ataaaaaact	tggataaaaa	tcacaagttt	gtgggataga	85860
tgcatatggt	aatatggaaa	ttctcataaa	gacacagtaa	tcaagacatg	tggtattggc	85920
tgggacgctt	ggctgtaatc	ctaacacttt	gggaggccaa	gatgagagga	ttgcctgaga	85980
tgaggagttg	cagacaagcc	tgggcaacat	agcaagaccc	tcattctctac	aaatatttaa	86040
aaaaattagc	caggtttggt	gccatgtgcc	tgtagtccca	gctattcagg	aagctgaggt	86100
gggaggatca	ctggagccca	tgaggtggag	gctgaaatga	gccatgattg	tgctactgaa	86160
ctttagcctg	ggagacagat	taaaaccttc	cctctctctc	tcaaacaaac	aaacaaaaaa	86220
tacatagtat	tgggcaaaac	atatgcaaac	aaaaacagaa	aagggtcagc	ataaatttac	86280
atatatggtc	aatttatttt	caatacaggt	agcaaagcaa	tttaatgagg	aaattttttt	86340
ccaaaattgg	tctgaaacaa	ctggatagcc	atagaaaaaa	actataacaa	atgtgacgct	86400
tgaatcctac	tgatgactc	aaattaaatt	aatttgagat	agctcttaga	cctcaatgta	86460
acagctaatt	ctgaggctga	aatataagac	tgctatgaaa	aagtatagta	tcttataacc	86520
ttggagaagg	aaaaattttt	tgagggaaga	accagaaaac	actaactgta	aaagaaaaca	86580

BI OL0250W0SEQ_ST25. txt

aatgataatg	tggacattca	ttgaataaaa	acttatgctc	accaaatatg	actgttaaga	86640
aaataaataa	gtaagtaaca	cactggaaga	aaaacactct	catccatata	tctgacaaat	86700
ggcctgtatc	cagagtatag	aaacatttct	cccactcact	aatcagagga	caaacaacct	86760
aatcaaaaatg	ggcaacaggc	ttgaatagtc	atttcttagg	agaagatgca	cacagagcca	86820
acaatcacct	gaaaaagtgc	acaacatctt	agccatcaaa	aatcaagagt	tataaccctc	86880
ataagatgac	actgaacatc	cagtgtacat	ggatatcatt	aagaagacac	aataataagt	86940
gggtgcaccg	atttggagct	agaatgtgcc	actctctcat	atgctgggtg	aagttcaaaa	87000
tcatacaaca	aattaaaaaa	tcagtctgat	gctttcttat	aaagttcgat	aaatatgcat	87060
ctatcctaca	aacctgtaat	tctattcttg	aatattttacc	ccccaaaatg	aaaacataag	87120
tccacaaaaa	tctatataaa	tattcatagc	agctttatgt	tttataaact	caaaataaaa	87180
actattttcaa	tgttttcatc	aaaagaaaat	gaaaactatt	taaatggttt	catcaaaaga	87240
aatgaaaaa	agaattttcca	gtatattttat	acaaggaat	actatttcac	aacaaggaac	87300
aagttactga	tagtctcaga	agcatgaaca	aacctcaaaa	atatattaag	gaaagaagcc	87360
agacgtcaaa	gtgtatagtc	tgtatgagtc	cattcatgtg	agtttataga	aaacacaatt	87420
tatggtgaaa	gaaaccaata	gcatttgaca	ctggccgtgg	gaagagggta	gcagagattg	87480
attgagcagc	cacacaaggg	agtttctggg	gtggtgaaaa	tgttctgcat	tgtgagggca	87540
gtgtgggcta	cacaagtata	tgtattttatc	aaatctcatc	cagctacatt	taagatctgt	87600
gcattctcact	ctatgtgaaa	atatactcaa	ctgaaaaaca	gagcaggtat	ctgtttcagg	87660
tgctacatca	cttgatacgt	ccagtttgt	taaaaaccac	tgccatacat	cctcaaatgg	87720
gggatctggg	cttgagacta	ggtcacatgt	gtagagtctc	tacagagacc	gtgttggtt	87780
cccatgctcc	ataatacgtt	ccaagttttc	tcagacagcc	acaggtcatg	aatgtgagga	87840
ttctgagagg	ttggagcaac	gttcttgggg	ggcataatgg	ggaaggcatt	ctccaagatt	87900
cctccagcct	ggggctctta	cctgctgtgc	ctcttactgc	attgttttct	gactcatcca	87960
tagccacttg	acccttcag	atcccatagt	ctacctagcc	gtctcccttt	atgccttggg	88020
tcccgtgtt	ctttcaactc	atcacccatt	ccttcagtcc	cagagtggct	gcagccagca	88080
gaggatggac	tgagagcagg	agaggaggtc	gtgcccata	acccatccta	gagaagcagc	88140
atcctgcctg	ggagctagtt	ttccagggaa	gcttttataa	gtcctgtaga	cccaaaccba	88200
cttgctctac	cagatacagt	atttatagta	atactatttt	catgattatt	ttatattgca	88260
aatgtagagc	atttatgcta	cactatgagt	aaatagagta	agggggctgg	catgggaatt	88320
atataatctt	ggatgccact	tcttccttgg	ggaaatgtat	ttgagttcca	acttacatat	88380
tactatatag	tcttatagag	agagagacaa	agagctagac	agacagagat	atctttgtat	88440
gtgcattaaa	aaatctaaga	tacatatattc	aaaatctgtg	tcattttattc	tggaggaaag	88500
tatttggcag	aagggtgaaag	gaagatatc	tgatcctttc	ttgtacagac	atgtattatc	88560
tcagttttca	tagagagcat	atactacttt	tgatgtttta	aaacaaaaat	tataatctgt	88620

BI OL0250W0SEQ_ST25. txt

gatgtgtcca	cagttgttta	aaagttgaag	ctgaagacca	tttgtgcttg	tggcaatatt	88680
atttgtggtat	aatgggaata	tttcaaaggc	acttggttaac	actttgtttac	agcaaaatgt	88740
agagggcgct	aagtgccctt	gaatattctc	ccatctctgg	tgacctgtgt	tgttttgaaa	88800
tttgcagtgg	cctgaccagg	aactactgca	ggaatccaga	tgctgagatt	cgcccttggt	88860
gttacaccat	ggatcccagt	gtcaggtggg	agtactgcaa	cctgacacga	tgtccagtga	88920
cagaatcgag	tgtcctcaca	actcccacag	tggccccggt	tccaagcaca	gaggctcctt	88980
ctgaacaagg	taagaaatth	gtggtttagac	atctatatac	tgggatgaaa	aaccatggaa	89040
aatcttactg	atgcagaagc	cttcagtggg	acactggagg	gttggttgag	ggtctgcaat	89100
gtggaggaaa	gcctcagcgc	cctctctggg	ggatccagaa	ctgtgatttt	tggcacgctg	89160
tgaggaggca	gtgtcttttag	gaagggcacg	gtgtcttttag	gaagggcaca	gacccgccag	89220
ggcactggac	ttaccactcc	cctggttatt	aaatgggtca	tttcagtgtc	ctagccaaaa	89280
tggatattct	aacagcctgc	caaatatgtg	aagattttcca	agccaataag	cctttccagt	89340
gatttaaagt	agactttttt	cattgcaatc	tacagtttgc	agtttcttaa	gaacatggcc	89400
tttgagtatg	atatacctaga	gaaacctaag	gagactgcat	tatttttcta	ttgtcctggg	89460
gctgcatagc	aggaggtaac	caacgaatgc	tgtctctccc	tggcctatct	cagtctttca	89520
caggctctgt	tcacctcagc	tttgaagtta	gaaatttcta	ggtgttcttg	cctcttcttc	89580
tcatgaaacc	tgatttgga	gtgagtctac	agaagaagag	gaagagaatt	ctgctttggt	89640
acaattcagg	actctgggca	ctagaagatt	ccctatctct	cctccaaggg	aataagttgt	89700
ttgtctctaa	ccctccttga	gaaacaatga	gtctttgcct	gcactcctaa	atgtaggatg	89760
atttcctgcc	caaattttca	aaagattaag	ccttttgcct	tggatatgagc	aatgggtctag	89820
ggaaatgcgc	aaggggtcttg	tgtcggcccc	tgactgacca	ccagtcacct	cctacagcct	89880
gcaccaagga	atgcattgca	ttctgggtctt	ctgccctgtg	gttctcatga	aaaccagcag	89940
agattcatat	gatggagctg	cacatgaatg	taattttcaa	tgtccagcat	tctcctctgt	90000
tctttatctt	tagatttaaa	aataatgttt	ctatgaactt	attaaaattc	tagaatacta	90060
tgaatctact	gggtcttttc	acatcctttt	gctactagta	gaaaaaagaa	tagtaataat	90120
tttcagaggc	tactgtccag	tatgtgacat	aaattgtctc	ccatgtttct	ctgctcatgc	90180
aattactgag	tatgatttat	tttattttta	tttcagcacc	acctgagaaa	agccctgtgg	90240
tccaggattg	ctaccatggg	gatggacgga	gttatcgagg	catatcctcc	accactgtca	90300
caggaaggac	ctgtcaatct	tggatcatcta	tgataccaca	ctggcatcag	aggacccag	90360
aaaactaccc	aatgcgtat	gtatttgatt	aaaaccataa	gaggagcaac	agccaactca	90420
aatattgggt	agaagaccca	tgctttaagc	tcacttccta	gggacaaatt	tctcttagac	90480
tcacattttg	gcaaaatgtc	tcaggacctt	tgcttttgag	caaagagtct	aagagaagag	90540
aaattttagg	cctgctatth	ttcctaatag	ttttatggaa	ggagtagaat	atacggaagt	90600
ggcgaagtca	tattaatgta	aagctcagaa	gataaatgac	caaagcttaa	acacagcacc	90660

BI OL0250W0SEQ_ST25. txt

attccacaat	gcccaactaaa	aatcaatgtc	atctttcact	cgtgcaattc	tgtcattcta	90720
aatttcaatt	cccgaagggt	tgtttgccat	ttttgtcatg	ggtaataagt	aaaaaaaaa	90780
aaattaagat	gtgtatatat	atatatatat	atatatatat	acacacacac	acacacacac	90840
aaacatctga	atatttatat	atatgtctga	atatttatat	acttgtgtat	aaaacttata	90900
tttaaatttt	tgcataaatt	tatatatfff	taatatfffca	ttaaaaatta	tattgtttca	90960
ctatgtatgt	ctgagtatff	ttatatatff	taatataaca	ttttaaatat	ttatatataa	91020
atattcaggt	atgtaactga	atattcattt	acacacacaa	atatatgtgt	gcatgtgtgt	91080
atatatatat	atacccatat	atatatatat	atatatatat	acatatatat	atatatatat	91140
atatgtatat	atatatatat	atatatatat	acacacacac	acacacacac	atacatacag	91200
gtataaacac	actgggcctg	aagcaccagt	ggctctgaaag	gacatgtgtt	gccaggactt	91260
gaagagcaaa	agcaggaagg	cggatgagag	tcaggaggta	cacaaacgct	gaaaagtaaa	91320
atggacaagt	acatggacaa	aaagcaggta	taagcataac	agccttttgg	aagtaaataga	91380
ctataaaata	tatgaaaata	ctgttttcac	aagttgcaca	acagatagta	gtgtattgag	91440
ataatttaga	acagaaaaca	aatgtgatca	accccataag	tgtgctgtat	ttcatcatgg	91500
attgaaggaa	aaagagatgg	agcccaagaa	gaccacagca	gtcttgatga	actgagagac	91560
accagagttt	gggattacaa	aggcagctgg	gattttctac	acttggtaat	aatgagaaag	91620
aatttggtga	gataaagagc	tacagtcattg	tacctagaag	tcacctcagt	gtaataataa	91680
tctgcatatg	cacagggagt	gattccacaa	tgaaagtagg	acaaagaaca	gctactgggg	91740
aaagaataac	tacaagggaa	caatgagttc	aatggagatg	gcagagctca	caaagcactg	91800
ggggatattt	gagttcttac	cagctagaaa	agagacctca	ttgcaaattct	tgggcattca	91860
gtagagaccc	cagaaaagcc	actctttgga	aacagagttg	atgtatttta	agagcaaaat	91920
ctactccaca	aaaatcctag	caaaattgaa	aagcaagtca	gaaagaccaa	aatcctctca	91980
acataaatta	gttgcccatc	agaagaaagc	ttaacctctt	cataggtaaa	caataaaaatc	92040
aaattgctca	gttatctggc	atccacaata	tgtgacataa	atttaaaaat	ttactagaca	92100
tacaagaagc	atttagtgtg	atccataacc	aggagaaaaa	tcattcaata	caaataagacc	92160
cagaaatgac	agaaatgata	gaattagcaa	aaacatttaa	aatatacata	tgatcatttg	92220
atcttgatgat	cagatatcac	aagagaagaa	agagatactt	gaacagaaaa	aatgcctgaa	92280
gcaatgatgg	ctgaaaactt	tccaaatatg	aagaaaaaaa	agctcacaga	ttcaagaaaa	92340
ctaatcaatc	agaaatatga	ttttgaaaag	taaaaatgta	tgattttactt	tggcaaatct	92400
tcttggttaa	attgtctaaa	atcaaagaaa	gctaggaaaa	ttttataagc	cagaggaaaa	92460
aagattgttt	atataaagga	acagttacac	aaatgactga	tgctttctca	tcagaaacaa	92520
tgaaagtcag	aaacaataaa	gtaacatctt	taaagtaata	gaagaaaaac	ccaagaggtg	92580
agggatcgtg	gcagacagga	ggcaggacta	gattgcagct	ctggacagag	cagcatgcag	92640
aggctcatat	tgtgaatttt	agcccatat	tgactgcaag	aacagaccag	caatcctgag	92700

BI 0L0250W0SEQ_ST25. txt

aggacccaca	gaccgtgtga	aggaagcaga	ctgctcctgc	aggataaggg	agacacccca	92760
aatactgtga	gttccccaac	tgcagaagtg	gaaaagggag	gccttactcc	ctcaaacaca	92820
ccccacaact	ggagaagctg	aaagtctgtt	tgcaggagaa	gttcccaact	ttacctgggc	92880
ctcagtaaat	ttagagagct	gagccaagca	aaatataggg	gtagaggaag	cagcagagaa	92940
gacctcagag	cttgctggat	ccccaagcag	ctcattcctg	cctggcacca	cagagatcca	93000
tcagaagtgt	ggccaaagga	acagagggta	aaactccaca	tggaggactg	ctctacctga	93060
actttctaac	aatttgaaca	gggggagaag	cctcctggcc	agaacttggg	ggagggcatg	93120
aatctggttt	gcagacttca	caggtggggg	aaggactaaa	gcccttttct	ttcacagctg	93180
ggaggtggaa	agcctcaggc	aagttttcaa	gcctgacttt	ccccccacct	ggaaacagac	93240
ttggagctgt	tgcgggggtg	ggggcatggt	gggagtaaga	ccagcccttc	agtttgcatg	93300
ggtgctgggt	gaggcctgtg	actgacagct	tccctccact	tccccgacaa	ctcagatgac	93360
tcagcagagg	cagccataat	cctcctaggt	acacaactcc	agtgacctgg	gaacttcacc	93420
cccacaccat	acagaagctt	cagtaagacg	tgcccaagga	aagtctgagc	tcagacacgc	93480
ctagtcccac	ccccaaactga	tggtccttcc	ctaccacccc	tggtagcaga	agacaaagag	93540
catataatct	ttggagtctt	agggcccacc	cacctctagt	ccctctccac	actagtatag	93600
ctgatgcagg	aggccaacca	gcacaaaaat	agagcattaa	accaccaaag	ctaggaaccc	93660
ctatggagtc	cattgcaccc	tcctccacct	ccaccagaac	aggcactggt	atccacagct	93720
gagagaccca	tagatggttc	acatcacagg	actctgtaca	gacagtcccc	agtaccagcc	93780
cagagctggg	tagacttgct	aggtggcaag	accagaaga	caggcaataa	tactgcagt	93840
tcagtcaca	ggaagccaca	tccataggaa	aagagggaga	gtactacatc	aagggaacac	93900
cccatgggat	aaaaacatct	gaacaacagc	cttcagccct	accttccctc	tgacacagtc	93960
tacccaaatg	agaaggaacc	agaaaaccaa	ccctggtaat	atgacaaaac	aaggctcatc	94020
acactcccag	ttcaccagca	atggatccaa	accaagaaga	aatccctgat	ttacctgaaa	94080
gagaattcag	gaggttagtt	attaagctaa	tcagggaggg	accagagaaa	ggcaaagccc	94140
aatgcaagga	aatccaaaaa	aaaaaaggta	taagaagtaa	aaggtgaaat	attcaacaaa	94200
atagatagct	taataaaaaa	acaataaaaa	attcagtaga	ctttggacac	acctttggaa	94260
atgtgacatg	ctctggaaag	tctcagcaat	agaactgaac	aagtagaaaa	aataaattca	94320
gagctcaaag	acaaggactt	caaattaacc	caatccaaca	aagacaaaga	ataaaggata	94380
agaaaatatg	aacaaagcct	tcaagatgtc	tgggattatg	ttaaatgacc	aaatataaga	94440
ataatcgtgg	ctcctgagga	aaaagacaat	actaaaagct	tggaaaacat	atttggggga	94500
ataactgggg	aaaacttacc	tggccttgct	ggacacctag	acatgcaaat	acaagaaaca	94560
caaagaacat	gtaaatataa	gcagacaaaa	gaacacctgg	gaaattcatc	acaaaaagat	94620
cttagcctag	gcacattctc	atcaggttat	gcaaagttaa	gacgaaggca	agaatcttaa	94680
gagctgtgag	acagaagcac	caggtaatgt	ataaaggaaa	ccctatcaga	ttaacagcca	94740

BI OL0250W0SEQ_ST25. txt

gtttttcagc	aggaactgta	caagctataa	aggattggag	ccctatcata	gcctcctcaa	94800
acaaaacaat	tatcagtcaa	gaattttgta	tccagcgaaa	gtaagcatca	tatatgaagg	94860
aaagatacag	tcgtttttgg	acaaacaaat	gctaagagaa	ttcaccatta	ccaagtcacc	94920
actagaagaa	ctgctaaaag	gagctctaaa	tcttgaaaca	aatcctagaa	acacatgaaa	94980
acagaatctc	tttaaagcat	aatcacaca	ggacctataa	aacaaaagta	caagttaaaa	95040
aacaaaaaca	aaaaacaaaa	ccaaagtacg	gaggcaataa	agaatatgat	gaatgcagtg	95100
gcacctcaca	tttcaatgct	aaaattgaat	ctaaatggcc	taaatgctcc	acttaaagga	95160
tacaaaaaga	gttggtggct	ggcaagatgg	ctgaatagga	acagctccag	tctgccgctc	95220
cccgtgagat	caacacatag	ggtgggtcat	ttctgcattt	ccaaccaagg	tacccggctc	95280
atctcattgg	gactggttag	acagtgggtg	cagcccacag	agggtgacct	gaagcagggg	95340
ggggtgtcac	ctcacctggg	aagtggaagg	ggtcagggaa	ctccctcccc	tagccaaagg	95400
aagccgtgag	ggactgtgcc	gtgaagacca	gtgcattctg	gcacaaatac	tatgcttttc	95460
ccacgggtctt	tgcaacctga	agaccaggag	attcccttgg	gtgcctacac	caccagggcc	95520
ctggatttca	agcccaaaac	tgggctggca	tttgggcaga	cactaagcta	gctgcaggag	95580
ttttttttca	taccccagtg	gtccctggaa	tgccagcaag	acagaaccat	tcacccccgt	95640
gaagaaaggg	ctgaagccag	ggagctaagt	ggtctttctc	agtggatccc	acccccatgg	95700
agcccagcaa	gctaagctcc	actggcttga	aattcttgtc	gccagcacag	cagtctgaag	95760
ttgacctggg	acgtcaagc	ttggtgggag	gaggggtatc	cacaaatact	ggggcttgag	95820
taggagggtt	tcccctcaca	gtgtaagcaa	aaccgctagg	aagtttgaac	tgggcagggg	95880
gcactgcagc	ttggcaaagc	cattgtagca	agagtgcctc	tctagattcc	tcctctctgg	95940
gcagggcatc	tctgaaagaa	aggcagcagc	cccagtcaga	agcttataga	taaaactccc	96000
atctccctgg	gacagagcaa	ctggaggaag	gggtggctgt	gagtgcagct	ccagcagact	96060
tagtttcctg	cctgccagct	ctgaaaagag	caccagatcc	cccaacacag	cactagagct	96120
ctgataaggg	acagactgcc	tcctcaagtg	ggtcctgggt	tcagaagata	ataagaaact	96180
cctctgagct	aaaggagcat	gttctaacac	aatgcaagga	agctaagaac	cttgaaaaag	96240
gtcagaggaa	ttgctaacta	cagtaagcag	tttagagaag	aacataaatg	accttaggga	96300
gctgaaaaac	acagcacgag	aacttcatga	cacatacaca	agtatcaata	gcaaaatcga	96360
tcaagtggaa	gaaaggatat	cagagattga	aaatcaactt	aatgaagtaa	agcgtgaaaa	96420
caagattaag	gaataaagaa	tgaaaaggaa	tgaacaaatc	ctccaagtat	gggactatgt	96480
gaaaagattg	aacctacgtt	tgattgggtg	acctgaaagt	gatgggagaa	tggaaccaag	96540
ttggaaaaca	ctcttcagga	tattatccag	gagaacttcc	ccaacctagc	aagacaggcc	96600
aacattcaaa	ttaaggaaat	acagagaata	ccacattcaa	attcaggaaa	tacagagaac	96660
accacaaaga	tactcctcaa	gaagagcaac	ctgaagacac	ataatcgtca	gattcaccaa	96720
ggttgaaatg	aaggaaaaaa	atgttgaggg	cagccagaga	gaaagtttgg	gttaccacaa	96780

BI OL0250W0SEQ_ST25. txt

aaggggaaccc	catcagacta	acagtggatc	ttcctgcaga	aactctacaa	gccagaagag	96840
agtggggaggc	caatattcaa	cattcttttt	tactattatt	atactttaag	ttctagggta	96900
catgtgcaca	aggtgcaggt	ttgttacata	tgtatacatg	tgccatgttg	gtgtgctgca	96960
cccattaact	cttcatttac	attaggtata	tctcctaata	ctatccctcc	ccactccccc	97020
catcccatga	caggccccgg	tgtgtgatgt	tccccactct	gtgtccatgt	actctcattg	97080
ttcaattccc	acctatgagt	gagaacattc	gggtgtttgga	tttctgtcct	tgtgatagtt	97140
tgctgagaat	gatggtttcc	agcttcatcc	acatccctac	aaaggacatg	aagtcacctt	97200
tctttatggc	tgcatagtat	tccatgggtg	atatgtgcca	cattttctta	atccagttcta	97260
ccattgatgg	acgttttgtg	tggttccaag	tctttgctat	tgtgaatagt	gccgcaataa	97320
acatatgtgt	gcatgtgtct	ttatagcagc	atgatttata	atcctttaga	tatatatcca	97380
gtaattgtat	ggctgtgtca	aatgggtatt	ctagttctaa	atccttgagg	aatcaccgca	97440
ctgtcttcca	caatggttga	actagtttac	agtcccacca	ccagtgtaaa	aatgttccta	97500
tttctccaca	tcctctctag	catctgttgt	ttcctgactt	tttaatgatc	accatttctaa	97560
ctgggtatgag	atgggtatctc	attgtggttt	tgatttgcac	ttctctgatg	gccagtgatg	97620
gtgagcactt	tttcatgtgt	ctcttgactg	cataaaagtt	ttcttttgag	aattgtctgt	97680
taatatcctt	tgccaacttt	ttgatggggg	tgtttgattt	tttttcttgt	aaatttgttt	97740
atgttctttg	tagattctgg	atattagccc	tttgtcagat	gggtagattg	taaaaatttt	97800
ctcccattct	gtagcttgcc	tgttcattct	gagggtagtt	tcttttgctg	tgccagaagct	97860
ctttagttta	attagatccc	attgggtcaat	tttggctttt	gttgctattg	cttttggtga	97920
tttagtcatg	aagtccttgc	ccatgcctat	gtcctgaatg	gtattgctta	ggttttcttc	97980
tagggtttat	atggtttttag	gtctaacatt	taagtcttta	atccatcttg	aattaatttt	98040
tatataaggt	gtaaggaagg	gatccagttt	cagctttcta	catatggcta	ggcagttttc	98100
ccagcaccat	gtattaaata	gggaaacctt	tccctatttc	ttgtttttgt	caggtttgctc	98160
atagatcaga	tggttgtaga	tgtgtggtat	tatttctgag	ggctctgttc	tgttccattg	98220
gtctatatct	ctgttttggt	accagtacca	tgctgttttg	gttactgtag	ccttgtaatg	98280
tagtttgaag	tcaggcagag	tgatgcctcc	agctttgctt	ttttggctta	ggattgtctt	98340
ggcaatgcat	gctctttttt	gttccatatg	aacttttaaag	tagttttttc	caattctgtg	98400
aagaaagtca	ttggtagctt	gatggggatg	gcattgaatc	tataaattac	cttaggcagt	98460
atggccattt	tcacaatatt	gattcttcct	atccatgagc	atggaatggt	cttccatttg	98520
tttgtgtcct	cttttatttc	attaagcagt	ggtttgtagt	tctccttgaa	gaggtccttc	98580
ccatcccctg	taagttggat	tcctaggtat	tttattctct	ttgaagcaat	tgtgaatggg	98640
agttcatcca	tgtccctaca	aaggacatga	agtcattgtat	gggaatgctt	gtgatttttg	98700
cacattgatt	ttgtatcttg	agactttgct	gaagttgctt	atcagcttaa	ggagattttg	98760
gtctgagaag	atgggggtttt	ctaaatatac	aatcatgtca	tctgcaaaca	gggacaattt	98820

BI OL0250W0SEQ_ST25. txt

aacttcctct	tttcctaact	gaataccctt	tatttccttc	tcctgcctaa	ttgccctggc	98880
cagaacttcc	aacactatgt	tgaataggag	tggtagagaga	gggcatccct	gtctttgtgcc	98940
agttttcaaa	gggaatgctt	ccagtttttg	cccattcagt	atgatattgg	ctatggggtt	99000
gtcataaata	gctcttatta	ttttgagata	tgtcccatca	atacatagtt	tattgagagt	99060
tcagcatgga	gagctgttga	attttgtcaa	aggccttttc	tgcattctatt	gagataatca	99120
tgtggttttt	gtctttgggt	ctgtttatat	gatggattac	atttattgat	ttgcatatgt	99180
tgaaccagcc	ttgcatccca	gggataaagc	caacttgatc	atgggtggata	agctttttga	99240
tgtgctgctg	gattcgggtt	gccagtat	tattgaggat	ttttgcatca	atgttcatca	99300
tggatgttgg	tctaaaattc	tcatttttgt	tgtgtctctg	ccaggatttg	gtatcaggat	99360
gatgctggcc	tcataaaatg	agttagggag	gattccctct	ttttctatga	ttggaatagt	99420
ttcagaagaa	ttgggtaccg	ctcctctttg	tatctgtggg	agaattcggc	tatgaatctc	99480
tcctggactt	tttttggttg	gtaggctctt	aattattgcc	tcaatttcag	agcctgttat	99540
tgggtctattc	aaggattcaa	tttctttctg	gtttagtctt	ggtaggggtg	atgtgtccag	99600
gaatttttcc	atttcttcta	gattttctag	tttatttgca	cagaggtgtt	tataatat	99660
tctgatggta	gtttgtat	ctgtgggatt	ggtagtgata	tcccctttat	cattttttat	99720
tgcattctatt	tgattcttct	ctcttttctt	ctttattagt	cttgctagtg	gtctatcaat	99780
tttgttgatc	ttttcaaaaa	accagctcct	ggattcattg	atgttttgaa	ggtttttttg	99840
tgtctctatc	tccttcagtt	ctgctctggg	cttagttatt	tcttgccttc	tgctagcttt	99900
ttaatgtgtt	tgctcttgct	tctctagttc	ttttaatggg	gatgttaggg	tgtcaatttt	99960
agatctttcc	tgctttctct	tgtgggcatt	tagtgctgta	aatctcccc	tacacactgc	100020
tttaaatgtg	tcccagagat	tctgggatgt	tgtgtctttg	ttgtcattgg	tttcaaagaa	100080
tatctttatt	tctgccttca	tttcggtaca	taccagtag	tcactcagg	gcaggttgtt	100140
cagtttccat	atagttgagc	agttttta	gagtttctta	atcctgagtc	ctagtttgat	100200
tgcactgtgg	tctgagagac	agtttggtat	aatttctgtt	cttttacatt	tgctgaggaa	100260
tgcctcactt	ccaactatct	ggatcaattt	agaataagtg	cgatgtgggtg	ctgagaagaa	100320
tgtatatctt	gttgatttgg	ggtaggagat	tctgtagatg	tctattaggt	ctgcttgggtg	100380
cagagctgag	ttcaattcct	ggatatccat	gttaactttc	tgtctcattg	atctgtctaa	100440
tgttgacagt	ggggtgttaa	agtctcccat	tattattgtg	tgggagtcta	agtctctttg	100500
taggtctcta	aggacttgct	ttatgaatct	aggtgctcct	gtattgggtg	catatatatt	100560
taggatagtt	agctcttctt	gttaaatgg	tccctttacc	attatgta	ggccttcttt	100620
gtctcttttg	atctttgtta	gtttaaagtc	tgttttatca	gagactagga	ttgcaacccc	100680
tgcttttttt	gttgttttcc	atttgcttgg	tagatcttcc	tccatccctt	tattttgagc	100740
ctatgtgtgt	ctctgcacgt	gagatgtgtc	ttcagaatac	agcactga	tggatcttga	100800
ctcttttatcc	aattttccag	tctgtgtctt	ttaattggag	catttagccc	atttacattt	100860

BI OL0250W0SEQ_ST25. txt

aaggtaata	ttttatgtg	tgaatttgat	cctgtcatca	tgatgttcgc	tggttat	100920
gctcattagt	tgatgcagtt	tcttcctagc	atcgatgggt	tttacaattt	ggcatgtttg	100980
tgcatgggct	gataccgatt	gtttctttcc	atgttttagtg	cttccttcag	gagctcttgt	101040
aaggcaggcc	tggtggtgac	aaaatctctc	agcatttgct	tgtctgtaaa	ggat	101100
tctccttcac	ttatgaagct	tagtttggct	ggatatgata	ttctcagttg	aaaattcttt	101160
tctttaagaa	tgttgaatat	tggctgccac	tctcttctgg	cttgttagagt	ttctgctgag	101220
agatctgctg	ttagtctgat	gggcttcct	ttgtgggtaa	cccgaccttt	ctggtgaatc	101280
tgacaattat	gtgtcttggg	gttactcttc	tcgaggagta	tttttgtggc	attctctgta	101340
tttcctgaat	ttgaatgttg	gcctgccttt	gtaggttggg	gaagtctctcc	tgataaatat	101400
cctgaagagt	gtttccaac	ttggttccat	tctcctcgtc	actttcaggt	acaccaagca	101460
gatgtagatt	tggtcttttc	acatagtccc	atatttattg	gaggctttgt	tcatttcttt	101520
ttactccttt	ttttctctaa	acttctcttc	tcgcttcatt	tcattcattt	gatctttaat	101580
cactgatacc	ctttcttcca	cttgattgaa	tcaactactg	aaacttgttc	atgtgtcacg	101640
tagttctcgt	gccatgggtt	tcagctccat	tagatcattt	aaggcttctt	ctatgctggt	101700
tatttttagtc	tgccattcat	ctaaactttt	tcaaggtttt	tagcttcttt	gcaatggggt	101760
cgaacatcct	tctttagctc	ggagaaattt	gttattacag	atcgtctgaa	gccttcttct	101820
ctcaactcat	caaagtcatt	ctctgtccag	ctttgttctg	ttgctcgtga	ggagctgcgt	101880
tccttcggag	gagaagaggc	accctgattt	ttagaatttt	cagctgttct	gctctgggtt	101940
ctcccatct	ttgtgggtta	tctacctttg	gttcttgatg	atggtgatgt	acagatgggg	102000
ttttgggtg	gatgtctttt	ctgtttgtta	gttttccttc	taacagtcag	gaccctcagc	102060
tgcatgtctg	ttggagtttg	ctggagggtcc	actccagtc	ctgtttgcct	gggtattacc	102120
agtggaggct	gcagaacagc	aatattaca	gaacagcaaa	tggtgctgcc	tgattcttcc	102180
tctggaagct	tcatctcaga	ggggcaccca	gctgtatgag	gtgtcagttg	gcccctactg	102240
ggagggtgcc	cccagttagg	ctactcgggg	gtcacggacc	catttgagga	ggcagctctgt	102300
ccattctcag	atctcaaact	ctctgctggg	agaaccacta	ctctcttcaa	agctgtcaga	102360
cagggatgtt	taagtctgca	gaagtttctg	ctgccttttg	ttcagctatg	ccctgcccc	102420
agagggtggag	tctacagagg	caggcaggtc	tccttgagct	gtggtgggct	ccaccagtt	102480
tgagcttcct	ggtcgctttg	tttacctact	caagtctcag	caatggcaga	cgcccctccc	102540
ccagctttgc	tgccgccttg	cagttcggtc	tcagactact	gtgctagcag	ttcaatctca	102600
gactgctgta	ctagcagtga	gcaaggctct	gtgggcatgg	gaccctctga	gcatgtgca	102660
ggatataatc	tcctgggtgtg	ccgtttgcta	agaccatttg	aaaagtgcaa	tattaggggtg	102720
ggagtgtccc	gattttccgg	gtacatctgt	catggcttcc	cttggctagg	aaaggaatt	102780
ccctgacccc	ttacacttcc	cgggtgaggc	aatatccgc	cttgcttcgg	ctcactctcc	102840
gtgggctgca	cccactgtct	gacaagcccc	ggtgagatga	accagttacc	tcagctggaa	102900

BI OL0250W0SEQ_ST25. txt

atgcagaaac	cacccatctt	ctgctttgct	catgctggga	actgtggact	ggagctgttc	102960
ctattcggcc	atcttgaaac	ctcccctctc	tcacgatcac	aaggtcccac	aataggccgt	103020
ctgcaggctg	aggagcaaga	aaagccagtc	tgaattccaa	aactgaagaa	attggagtct	103080
gatgttcaag	ggcaggaaac	atccagtgcc	aaagaaagat	gtagaatatt	caacattctt	103140
aaagaaaata	attttcaacc	tagaatttca	tatccagcca	aactaagctt	tataacaaag	103200
gagaagtaaa	atcctttaca	aacaagcaaa	tgctgaggaa	ttttgtcaac	accaggcctg	103260
ccttacaaga	ggtcctgaag	aaaacactaa	atatggaaag	gaaaaaccag	taacagctac	103320
tgcaaaaaca	taccaaattg	taaacaccat	caacactata	aagaaactgc	atcaactaat	103380
gggcaaaaata	gccagctagc	atcataatga	caggatcaaa	ttcacacata	acaatattaa	103440
ccttaaattgt	aaatgggcta	aatgccccaa	ttaaaagaca	cagactggga	aattgaataa	103500
agagtcaaga	cccattgggt	tgctgtgttc	agaagaccca	tctcaggggtg	aaaagacata	103560
catgggctca	aaataaagaa	atgaaggaat	atttaccaag	caaattgaaa	gaaaaaaaaa	103620
gcagcgggtg	caatcttagt	ctttgatgaa	acagacttta	aaccatcaaa	gatcaaaaga	103680
gacaaaggag	ggcattacct	aatggtaaaa	gtatcaatgc	aacaagaaga	tctgactgtc	103740
ctacttatat	atgcacccaa	tacaggagca	cccagattaa	taaagcaagt	tcttagagac	103800
ctacaaagag	acttagactt	ccacacaaaa	atagtgggag	actttaacac	cccacagcca	103860
atattagatc	gacgtgacag	aaaattaaca	aggatattca	ggacgtgaat	tcagctctgg	103920
accaagctga	cctaatagac	atctacagaa	ctcgacacca	caaatacaac	gaatatacat	103980
tcttctcagc	accacattgc	acttattcta	aaattgacca	cataattgga	agtaaaacac	104040
ttctcagcaa	atgccgtaga	atggaaatca	taacaaacag	tctctcagac	caaagtgcaa	104100
tcaaactaga	actcaggatt	aataaactca	ctcaaaacca	cacaactata	tggaaactga	104160
acaacctgct	cctgaattac	tactgggtaa	ataacaaaat	taaggcagaa	gtagataagt	104220
tcttagaaac	caaagagaac	aaagacacaa	tgtgccagaa	tctctggtac	acagctaaag	104280
ccatgtttag	agggaatttt	atagcactaa	atgcccacag	gagaaagcgg	gaaagatcta	104340
aatcaacac	cctaacatca	caattcaaag	aaccagagaa	gcaagagcaa	acaaatacaa	104400
aagctagcag	aagacaagaa	ataactaaga	tcagagcaga	actgaagggg	ataaagacac	104460
gaaaaccctt	taaaaaatta	ataaatccaa	gagctggttt	tttgaaaaga	ttaacaaaat	104520
acatagaagc	ctagccagac	taataaagaa	gaaaatagag	aagaatcaaa	tagacacaat	104580
aaagaataat	aaaggggata	tcaccaatga	tgccacagaa	atacaaacta	ccatcagaga	104640
atactttaaa	cacctctatg	caaataaaaat	agaaaatcta	aaagaaatgg	ataaattcct	104700
ggacacatac	accctcccaa	gactaaacca	ggaagaagtc	aatccctga	atagaccaat	104760
aacaagttct	gaaatcgagg	cagtaattaa	tagcttacca	accaaaaaaa	gccagacca	104820
gagggattaa	cagtcaaadc	ctaacagagg	tacaaagaag	agctagtact	attccttctg	104880
aaactattcc	acacaataga	aaaagaggga	ctcctgccta	actcatttta	tgaggccagc	104940

BI OL0250W0SEQ_ST25. txt

atcattctga	taccaaacc	tggcagagac	acaacaagaa	aagaaaattt	caggccaaca	105000
tccctgatga	acatcaatgt	gaaaatcctc	aataaaatac	tggcaaactg	aatccagcag	105060
cacatcaaaa	agcttatcca	ccatgatcaa	gttggcttca	tccctgggat	gcaaggctgg	105120
ttcaacatat	tcaaatcaat	aaacataatc	catcacataa	acagaaccaa	tgacaaaaac	105180
cgtatgatta	tcgcaataga	cgagaaaaag	gcctttgata	aaattcaata	cccaatcatg	105240
ctaaaaactc	ttaataaact	aggtattgat	ggagcatgtc	tcaaaataat	aagagctact	105300
tatgacaaat	gcatagccaa	tatcactctg	aatgagcaga	agctggaagc	attccctttg	105360
aaaaccagca	caagacaagg	atgccctctc	tcaccactcc	tattcaacat	agtattggaa	105420
attctgtcca	gggcaatcag	gcaagagaaa	gaaataaagg	tattcaagtg	ggaagagagg	105480
gagtcaaatt	atttctcttt	gcagatgaca	tgattgtata	tttagaaaac	tctatcatct	105540
cagcccaaaa	tctccttaag	ctgataagca	acttcagcaa	agtctcagga	tacaaaatca	105600
atgtgcaaaa	atcacaagca	ttcctataca	ccaataagag	acacagagcc	aaatcctgag	105660
tgaattccca	ttcacaattg	ctacaaagag	aataaaatat	acctaggaat	ccaacttaca	105720
agggatgtga	aggacctctt	caaggagaac	tacaaaccac	tgctcaagga	aataagatag	105780
gacacaaaca	aatggaaaaa	cattccatgc	taatggattg	gaagaatcaa	tattgtgaaa	105840
attgccatac	tgcccaaagt	gatttataga	ttcaatgtta	tcccatcaa	gctaccattg	105900
atttcttcac	ataattagaa	aaaactactt	tcaatttcat	atggaataga	aaaagggcct	105960
gtatatccaa	gacaacctaa	gcaaaaagaa	caaagctgga	ggcatcatgc	tatctgactt	106020
caaaatatac	tacaaggcta	cagtaacaaa	aacagcatgg	tatggtactg	gtaccaaacc	106080
agatatatag	accaatagaa	cagaacagag	gcctcagaaa	taacaccaca	catctacaac	106140
tattggatct	ttgacaaact	ggacaaaaat	aagcaatggg	gaaaggattc	cctattttaat	106200
aaatggtgtt	gggaaaactg	gctagccata	tgagaaaaac	tgaaactgga	tcccttcctt	106260
acaccttata	cacaaattaa	ctcaagatag	attaaagaat	taaatgtaag	acctaaaacc	106320
ataaaaaccc	tagaagacac	tttgggaggc	cgaggtggat	ggatcacgag	gtcaggagat	106380
cgagaccatc	ttggctaaca	cagtgaagac	ccatctctac	taaaaataca	aaaaattagc	106440
tggtgtgtgt	cgtgggcacc	tgtagtccca	gctacttggg	aggctgaggc	aggagaatgg	106500
catgagctga	ggaggttgag	cttgacagca	gccaagattg	tgccactgca	ctccagcctg	106560
ggcaacagag	tgagactcca	tcaaaaaaac	aaaaacaaaa	acaaaaaatc	aaaccctaga	106620
agaaaacata	ggcaatacca	ttcaggacat	aggcatggga	gaagacttca	tgactaaaac	106680
agcaaaacca	atggcaacaa	aagccaaaat	ttacaaatca	gatctaatta	aaataaagag	106740
cttctgcaca	gcaaaaaact	ctcatcagag	tgaaaaagca	acctatggag	aaaaattctg	106800
tggtctagcc	atctgacaaa	gggctaattg	ttagaatgta	caagcaactt	aaacaaatgt	106860
acaagaaaaa	aaaaacaacc	ccatcaaaaa	gtgggcaaag	gatatgaaca	gacacttctg	106920
acaggaagac	ctttatgtgg	ctgacaaaca	tgaaaaaagc	tcatcatcac	tgtttaattag	106980

BI OL0250W0SEQ_ST25. txt

agaaatgcaa	atcgaaacca	caatgagata	ccatctcatg	cccgttagaa	tggcgatcat	107040
taaaaagtca	ggaaacaaca	gatgctgaag	aggatgtgtg	gagaaagagg	aacacattta	107100
cactgttgggt	gggagtgtaa	attagttcaa	ccattgtgga	agacagtgcg	gtgattcctc	107160
aaggatctag	aaccagaagt	accatttgac	ccagcaatcc	cattactggg	tatataccca	107220
aaggattata	aatcattcta	caataaagac	acatgcacac	gtatgtttat	tgtagcacta	107280
ttcacaatag	caaagacttg	gaaccaactg	aaatgcccat	caatgataga	ctggataaag	107340
aaaatgtggc	acataacac	tgtggaatac	tatgcagcca	taaaacagga	tgagttcatg	107400
tcttttgag	ggacatggat	gaagctggaa	accatcattc	tcagcaaact	aacacaagaa	107460
cagaaaacca	aacaccatat	gttctcactc	ataagtgtga	gttgaacaat	gagaacacat	107520
ggacacagga	agggaacat	cacacacagg	ggcctgttgg	ggagttgagg	ctaggggagg	107580
gattggatta	ggagaaatac	ctaattgtaga	tgatgggttg	ctgggtgcag	caaaccacca	107640
tgacacgtgt	atacctatgt	aacaaacca	cacattctac	acatgtatct	cagaacttaa	107700
agtataataa	taataagata	cagaactgca	gaatgaataa	gaactcacca	accatctgct	107760
gccttcagga	gactcattta	agacataagg	actcacataa	acttaaagta	aatgggtgga	107820
aataataata	agtgggtgtca	ctgatgtgga	ggtagattat	aaaactctta	tcatatgctg	107880
gtggaagatc	aaaatgataa	aacgaattaa	aaaatcagtc	agatggtttc	ttaaaaagtt	107940
ccatcaatat	gcctctatct	tacaaacctg	caattctatt	cctgaatctt	tatccaagg	108000
aatgaaaaa	gtaagtccac	aaagagttct	atatgaatat	ttataggagc	tttattttatt	108060
ataattcaaa	ctgtaaaaat	aatttcaatg	ttcatcaata	acaaaatgaa	aaaataattt	108120
gcaacctact	ggtacacttg	aatactattc	agcactgagt	atcttaaata	gcatggatgg	108180
agctcaaaaa	tatactcagg	aaagaagcca	tgtatatctt	gtatgagttc	atttacatga	108240
gatcatttac	atttcctcca	aaagaggaaa	aactaatttc	tgttgaaaga	aaccaatgta	108300
tttgccctctg	gcagtggtaa	gggggtagca	cagattaatt	gggtagggac	tcaagagagt	108360
ttctggggtc	acagaaatgt	tccgtgtggt	gatgggagtt	tgggctccac	aggtataggt	108420
gttgatccaa	aatcatcaaa	aaaacaacat	tgcagatctg	tgcactctac	tctgtgggaa	108480
agtatatctc	aactgtaaaa	agggcagaaa	ttgcttttaa	acgctcagcc	ttttagcaca	108540
tccagttgct	tggagaacca	gcttactcaa	atgggggtct	aggctggaga	ctaggtcaca	108600
ggcatagagt	ctctaaactt	tcccatggca	cataatacgt	ttcaggtttt	ctcagagagc	108660
tgcaggttag	taatctgagg	attctgacaa	gttgggtcaa	cgttcctagg	aggcatgaat	108720
gggagtgcac	tctctaagat	ccctccaccc	cagggtcctt	gctttctgtg	cctcttactc	108780
cattgttttc	tgactcctct	gtagccactc	gacctcttca	gatcccattg	tctaccacgc	108840
catcgccctt	tatgacttgg	gtcccactgt	tctttcatct	catcctccat	tccctcagtt	108900
tcggagtggc	tgccgctagc	agaggatgga	ctgagagcag	gagaggtggt	cctgcccagg	108960
aacccatcct	agagaaatgg	catcctgtct	gggagctagt	tttttagggc	aggttttata	109020

BI OL0250W0SEQ_ST25. txt

agtcttgtaa	agccagacac	acttgatcta	cctgggtatgt	tatttacagt	aatactat	109080
tcataattgc	ttttcactct	aaaagtagag	ccttttagct	acactgtgag	taaataaagg	109140
ggctggcctg	ggaatggtat	catgttggat	gttgtttctt	ccctgaagta	atatatatca	109200
gttacaat	acatgttact	gcagagtcct	agagagagac	acagagaatg	agacagatac	109260
caatacat	ttatgtgcat	taaaaaaatc	taaggccagg	cgcagtggct	cacacctgta	109320
atcccagcac	tttgggaggc	cgaggtgggt	ggatcacgag	gtcaggagat	tgagaccatc	109380
ctggctaaca	cggtgaaacc	ctgtctctac	taaaaataca	aaaaattagc	caggcgtggt	109440
ggcgggcgcc	tgtagtccca	gctactcagg	agactgaggc	aggagaatgg	cttgaaccca	109500
ggaggcagac	cttgacgtga	gccgagattg	cgccactgca	ctccagtctg	ggcgacagag	109560
cgagactccg	tcacaaaaaa	aaaaaaaaat	ctaaaatgca	ctcttcaaaa	tctatgtcat	109620
ttattctgga	ggaatgcagt	tggcagaagg	aggaagatat	tccgaat	tcttgatac	109680
atttatgtat	gatctcagtt	tttttatgga	tcatagacca	attttgatat	tttaaaataa	109740
aaattataat	ctatcttggg	aatttacatg	gttctttaga	acttgaggac	cgtttttgct	109800
tttcggaata	ttattgtacc	taaaatggga	atattacaac	gtcacttttt	aacactttgt	109860
tataacaaag	tttagacagc	gctgggtgcc	cctgaat	ttccgcctc	ttgtgacctg	109920
tgttgttttg	gaatttgcat	tggcctgacc	gagaactact	gcaggaatcc	agattctggg	109980
aaacaaccct	ggtgttacac	aaccgatccg	tgtgtgaggt	gggagtactg	caatctgaca	110040
caatgctcag	aaacagaatc	aggtgtccta	gagactccca	ctgttggttc	agttccaagc	110100
atggagggtc	attctgaagc	aggtagaag	tctgtggcca	gatattctaca	catttgaaca	110160
ttgggatgaa	aagagatgga	aaatctgact	gatgcagaag	ccttccatgc	tacacagaaa	110220
cttgagggtg	tggcaggtgg	aaagaagcct	cagcactctc	tctggtggag	caat	110280
cgcaacgtgc	gtgggcggtg	acttcaggaa	tggtgcaaac	ccacctgggc	acttgactta	110340
ccactcactt	tgttatgaaa	ggggttatct	cgggtgttcca	gacaaaattc	caattctaac	110400
atcaggccaa	atttgtgcca	aatttcacac	tagtgagtgt	ttccaggcat	ttattaaaat	110460
ggacagtgtt	cattgcaatc	ttcagcattg	cagttgctga	ggatgtggc	cgctgagttt	110520
gtcatcctgg	ggaaacctaa	tatgatgata	tttattccat	ctaatacctg	ggctatttgg	110580
cagtaaatac	cacagaatac	actatttctc	tggcttattt	cagtcttagg	taggctctgc	110640
acacctatgc	ttggaaggca	ggaatttctt	ggtgttcttg	tgccttcttc	tcatggaacg	110700
tgcatctttg	gtgtgtgttg	agaggaagg	tagtagactt	ctgctttgtt	gcaatgcagg	110760
atgctggaac	aagaggattc	cctgtctcta	ctgtaaggga	ataagatttt	agcctccatc	110820
cttctctaag	aagcaatgtg	tctttgcctc	caagtactag	atgcaggacc	atgaactgcc	110880
ccgtccacca	gaagcttaag	gctttggctt	ttcaggagca	atcatctagg	gaactgtgca	110940
gggttttcat	gtctgtcccc	tactgacagc	caatcaccat	acagcctgca	taacctaatc	111000
catcatcgtc	tggtttcctg	cctcattgtt	ttcatgaaca	accagtagag	agccatacga	111060

BI OL0250W0SEQ_ST25. txt

aagagcttgc	acatgagtct	ttgttccaat	tgtaagagca	ctgataggtc	cttttcccac	111120
caggttttga	atataaaatt	tctaagaact	tattaaaata	ttagaatgtt	attaatctat	111180
tgtttttgct	tcagcatgtc	cttctgcttg	tgagtatact	aaagagaaca	gtcataattc	111240
tgaaactact	gtcctgtttg	tgtcataaat	tgcttcacat	gtttctgcat	actagtagtt	111300
actcagcttg	atthttgtcta	ttttcagcac	caactgagca	aaccctgtg	gtccggcagt	111360
gctacatgg	taatggccag	agttatcgag	gcacattctc	caccactgtc	acaggaagga	111420
catgtcaatc	ttggatcatcc	atgacaccac	accggcatca	gaggaccca	gaaaactacc	111480
caaatgagta	tgtctttgat	gttacttgta	agaggagcaa	cagccaactt	aagttcctcc	111540
tagaagagcc	ttgcttcaag	ctaacttggt	aggacaaatt	tcccttagac	ccagaagggtg	111600
tgtcaaaatg	tccagacaac	tttgcttttg	atcaaagagt	ctgagagaat	aggtatttta	111660
ggcttgctat	cttttcta	agtctgatgg	aagcagaagg	ctacatggag	ctgatgaggt	111720
ctttttaata	taaagctcaa	gagatcaa	gatcaaatac	ttagagtgcc	attctacaag	111780
gctcataaaa	gatcaatgca	ctctttcacc	catgcaattc	tatcattcta	acctcccttc	111840
tctgaaatga	aggctttttg	ccatttttgt	catgggtcac	aagtaaataa	ttcacatgta	111900
tatgagtata	tatataacca	gggtgtgttt	ttcagactag	tatgtatata	tatacatata	111960
tatgttcata	taagttagta	ttcatatata	tgttcatata	tatatgttca	tacagactag	112020
tattcatata	tatatacata	tatatataca	cacacatata	tatatatata	tatatgttct	112080
agggaaacat	gcaaggtttt	tatgtctgtc	cctgactgat	gaccaaatac	cctatagcct	112140
gcacagctgc	aagctgtata	gccatacaat	ttgcaggaca	cacacacata	cacacacaca	112200
cacacacaca	cacacactaa	catataatat	aatataatat	aatataatat	aatataatat	112260
aatataatat	aattaatata	tataaacctg	tgtgaacaca	ctgggttcta	agctccagtt	112320
ttctgaaggg	atatgggttg	ccaggagagg	aagagcaaaa	gcaagaatgt	agatgagaat	112380
taggaagtaa	acagatatgg	agattaaaat	gggcaggtac	atggacaaaa	aaccaggtct	112440
gacaaaaact	ggctttctgc	cataaatgac	tataaaagat	attaaaaaac	actttccaca	112500
tgttggacaa	gagacagtac	aggactgaga	taatttagaa	aaggaaatga	atgagcgcaa	112560
ctccgtaact	attatgactt	tcttcttgga	gaaccttctt	ggactgaagg	gcaaggaatt	112620
ggagccaaag	ccaaccacag	cagtcttgct	gaactgagga	aagagactgg	agtttgggat	112680
agctaagaaa	atgtgtatth	tctatgctag	gtaataatga	gaaagaatth	gtggtgaaaa	112740
ggagctgaag	gaatatgcat	ggaagtctaa	tataaactgc	atatgcacag	ggagaaattc	112800
tacaaagtgg	gacagagaac	cactactggg	gaaaggacaa	attcagggaa	acagtgagct	112860
caatggtgac	gccagagctc	acgtagcact	gggggatacc	ggggttctga	tcagcccagag	112920
gagagacacc	tcattgaaca	tctcgggcat	tcagtagaga	ccccagaaaa	gtcatactth	112980
aggagtagga	tttatgcctt	cttagaataa	agactacccc	agaaacaccc	tagtaaagct	113040
taaaaaccaa	gtctaaaagg	acccaaataa	tctccaagta	aattaactgc	ctgacagaag	113100

BI OL0250W0SEQ_ST25. txt

aaaactcaac	catcactgga	ggtaaataac	atgattacag	tgctctgtaa	tgttgcattc	113160
acaaggagtg	acatcattta	aaaattttatg	aggcaggaaa	aagcaattag	tgtgatccat	113220
aactaggaga	aaaaccagtc	aatacaaata	gaccaagaaa	tagtagaaac	gatggaattg	113280
acaaagaaat	taaaactgta	tatatgataa	ttgtgttcaa	agattttaag	aaaacatgaa	113340
catgagggaa	acaaatgcag	aatataaaaa	aaagcaaattg	cgtaaaacaa	ccaaatggaa	113400
attaaagaac	tacaaaaaag	tataacctta	ataaaatact	cactggatgg	ccttaatat	113460
agtttataca	ttacagaaga	aaaagtgaac	cagaagataa	ctcaatgaaa	gccatacaat	113520
ctgtaagaca	cacacacacg	cacacgcgcg	cgcgcgacac	cacacacaca	cacagagaga	113580
gagagagaga	gaaagagaga	gagagaaagg	ctgaaaaaaaa	taaatagaac	cctaaggata	113640
tcagtgaaaa	tagcaaaaga	tttaatatat	gggtaaagca	agtcacagaa	ggacgggaag	113700
gagatatgtg	gacagaaaaa	aatactcaaa	gcaatgatgg	ctgaagactt	tacacgtatg	113760
aagaaaatga	taaactcaca	gtcaagaagc	tcaatgaatc	agaaatagta	tttttaaaag	113820
caaaactcta	tgatttactt	gggtacatta	tagataaatc	gtccaacatc	aaagataaca	113880
aggataatct	tataagccag	aggaaaacaa	tatcattttac	atagaggggac	agtaatgaaa	113940
gtgaccgatg	ccttctcctt	ggaaacaatg	gcataacatc	tttaaagtga	taaagagaaa	114000
taaaaacaga	tcaacctagg	acgacatgtc	cagccaaaac	aaacaaataa	acaaaaaac	114060
cctttaaaat	aaacgtgatg	taaatacgta	ttctgccacc	tccagaggaa	acaagcaaaa	114120
aaacaaaaga	atgtttccaa	ggcaggcttc	tgtattaaaa	gatttttaagg	aaagttattc	114180
aggtagaaga	aaaataatac	cagatgggaa	ctttaatcca	tactaagtaa	tgaagagccc	114240
tggaaatggc	aaatggcaat	gtcaatataa	aatactctta	tttatcta	ttttaaatgt	114300
atttaaagga	caatttgtga	tattaatata	aataatagga	atatattgtt	gtttcaacgt	114360
atgtagtagt	aaaattcata	aaaacagtag	cacaaataat	gcagatgata	actggaagta	114420
tactgttaat	gagttttttg	cattatccat	gaagttatat	aatattaata	gatggttgaa	114480
tgtgatagtt	taaggtggga	tattataaat	cctaggacaa	ccaaaaaat	ttaaactgag	114540
aggaatggat	agtaagagga	atagtccttt	tatgcaaaag	aaggaagaaa	aagaggaata	114600
aagaatataa	aagatatggg	gtaaacagaa	aatacatagc	attattgtag	acacaaactg	114660
aactacctta	tgagtatatt	aaatataaaa	ggattaagca	ttacaaataa	aaggcagaga	114720
ttgtaaattg	aataaaaacc	acagctaagt	gtgttctttt	tagaataaat	actctttaag	114780
tgtaaagatc	tactttaaac	acaaaatat	gaaaaaggat	atataccatg	aaaacctgaa	114840
tcataaataa	gctggagtgg	tgattaatgg	atgcaggcac	tcctaaagac	taataagtga	114900
atgtggtcaa	attgaagaaa	caaaagtata	tacgtgctca	atgtgcaaaa	actttttctg	114960
tatacatgct	atgatccttt	ggaaaattaa	agtttttaag	caatatcact	gacaatagta	115020
tcaaaaccaa	aaaatattta	gtgataaatt	tcacacacta	tgctcaagga	ctatacacct	115080
tgactagaa	aacaatgttg	aggaaagaat	taaaagatct	aaatatacac	catgcttata	115140

BI OL0250W0SEQ_ST25. txt

gattaaaaga	ctccatatca	gttctcgtga	aattgatctt	tggatgaaac	ccacacccaa	115200
gcactattgc	aacagtcctt	ttttggaaaa	aaaaattgga	ggacttatat	accttaatat	115260
aaagacttat	aaaagtacag	gaatcaagac	atgtggtatt	ggcctggccc	cttgggtcat	115320
gcctgttacc	ccaacatttt	gggaggctga	gtctggagga	tggcttgagc	ccagatgttc	115380
aagaccagcc	ttagcaacag	agtgagaccc	tctctctaca	aaaaataaac	aattagatcg	115440
atgtgatgac	ttgcacatgt	agtttcagct	actcggaatg	ctgaggtgag	aggattgctt	115500
gactcaggag	gtctagccat	gagtgagcat	tgatcatgcc	tctgcattcc	agcctggatg	115560
atggaatgag	acactgtctc	aaaaaaaaaa	aaaaaaaaag	atatgtgtta	ttggccaaaa	115620
aagtatgcaa	acataaaaag	ggatggccca	ccaccagacc	cacatacata	tatggtaaata	115680
ggattttccg	tatagatggc	aaagcaattc	aatggagaca	aaaatgtttt	acaaaatcat	115740
tctgaaccat	ttggatatcc	atgatacaaa	acaaaagcag	aacttgactt	ttgcttttca	115800
tctcaaatta	ttttgatatc	tcttccacct	aagtgtcaga	gctaaaactg	aacctgaaat	115860
atgaaagttc	catgaaaaaa	tataaatctt	tcacaacctt	ggagaaggca	aacttttttg	115920
aggcaggagt	ctgtaaacac	tcactataaa	ataaaaacaa	ttataatgtg	ggcttttcatg	115980
aaaactcatg	cttaccaaaa	gtcattgtta	agaaaataaa	taggcaagta	acacatgaga	116040
agaaaaatgc	tctctgtcca	tatatctgac	aaatggcttg	tgtccagaat	ataggaacat	116100
ttctcccact	cactaaacag	aggacaaaca	actaatgggc	aacagattga	ataggcattt	116160
cttggggata	gatagatgta	cacatagcca	ataagcacct	gaaaaaatgt	ccagtatctc	116220
agccatgaaa	aataaagagt	tataatcatc	atgagatgtc	accaaacc	caatggacat	116280
ggatattatt	aagaagacac	cacagtaact	gatgtcactg	atgtagagca	aggatgtgaa	116340
actctctcat	atgctggtga	aagtgcacaa	tgatacaacc	acttttgaaa	tcagtctgat	116400
agtttctcca	aaagttcaat	aatgcactt	ttaccctaca	aacctgcaat	cctgtttgtg	116460
aatatttacc	ccacagaaat	ggaaacataa	gtccacgaag	acatctccaa	gaatattcat	116520
agcagcttta	ttttttataa	ccccaaactg	tagacaattt	caatgtcaat	caataagaaa	116580
atgaataaat	aatttgtgaa	ctagtcatac	aatggcatac	tgttcagcaa	taaaaggag	116640
catgtttttg	atactctcaa	atagtatgga	agatgtctca	aaatattaca	ttaaagaaag	116700
atgccagata	acaaaaatga	acattatgta	tgagtctatt	gatgtaaggt	tccagaaagg	116760
taaaactaat	ttctggtgaa	agaaaccaat	atcatttgcc	tctggccatg	ggaagagagt	116820
agcagagatt	gattgagcag	taaaacgaag	tttttttctg	gggtgatgta	aatgtcctgt	116880
attgtgattg	aagtgtgagt	tacacaagtg	tacatgttca	tcagaagtca	tcaaactaca	116940
tctaagatct	gtgcatttga	ctatacatga	aaatatacct	cagttgaaaa	tagatcaata	117000
acctccctca	tatactatac	ttgctaacac	agccagctgc	ttggagaacc	agcttgctgg	117060
aatggagaat	ctgggcttga	gactgggtca	catgtataga	gtctctacag	agacaatgtt	117120
gcattcccac	ggtacataat	acatttcaag	gtttctcaga	cagccacatg	tcatgaatgt	117180

BI OL0250W0SEQ_ST25. txt

gaggattctg	agaggttggg	gcaacattcc	tgggaggaac	gaaggggagc	acattctcca	117240
agatccccc	ccaccgggg	cctcaccggc	tgtgcttttt	tttttttttt	tcttgacaga	117300
gtctcgctct	gtcgccaggc	aggagtgtaa	tggcccaatc	tcggctgatt	gcagcctcca	117360
actccagggt	tcaagagatt	ctcctgcctc	agcttcatga	gtagctggga	ctacagatgt	117420
gcgccactgc	gccagctaa	tttttgtatt	tttagtagag	acgggggttt	gccatgttgg	117480
ccaagatggt	ctcgctctgt	tgacctcgtg	atccacccgc	cttggcttcc	caaagtgctg	117540
ggattacagg	cgtgagccaa	agcaccagc	ctgtgcctct	cacttactca	attgtttttc	117600
tgaaccctcc	atagctgggt	gaccttttca	gatcccatag	tctagccagc	cctctcactt	117660
tatgccttgg	gtcccactgt	tccttcatct	catccccctt	ctgtcagtcc	cgcagtggct	117720
gtggccagta	gaggatggac	tgagagtagg	agaggaggtt	ctgccagga	acccatccta	117780
gagaaacagc	atcctgcctg	ggacctagtc	ttccagggtca	gctttttata	gtctttttaga	117840
ctcaaactca	cttgaccac	ctgaagtgg	attgacaata	atgctatttt	catggttgtt	117900
tttactgtga	aatgcagagc	cttttagcta	cacgactagt	acagagagta	agggaggctg	117960
gcctgggaat	gatatcatct	tggatggcat	ttcctccttg	gagaaatata	tgttagtctc	118020
aactcacatg	ttactataca	gtcctgtaga	aagagataca	gagagttaga	cagggtataga	118080
cgcatttgta	tatgcataac	aatctataag	acacacatca	aaatccgtat	accggttcct	118140
ctaggggtat	gtgcttggca	gaaggtagaa	ggagggtatt	ctggttcctt	tcttttgcac	118200
atttatgtat	gatctcagtt	tttatatgga	gcattgatag	ggtttggcta	tgtccccacc	118260
caaaatctca	tcttgacttg	taatctctat	aatcctgata	atcccatgt	gtcaagggca	118320
ggaccagggtg	gaggttaactg	gatcatgggg	gcagtttctc	ccaggctgtt	ctcatgacag	118380
tgagagagtc	tcctgagatc	tgatggtttt	gtaagtgtct	ggcatttccc	ctacttgcac	118440
ttactctgtc	ctgccgcctg	tgaagaagg	gcctgtttct	cccttgcctt	ctgccatgac	118500
tgtaaatttc	cagaggcctc	cccagcaatg	tggaactgtg	agtcaattaa	aactcttttc	118560
tttgtaactt	accagtcctg	tctcgggtat	ttcctcatag	caatgtgaga	acgggcta	118620
acaagcatat	actacttttg	atattttaaa	ataaaaatta	tcattctatct	ttgaaaggca	118680
tgacaaaatg	ggaagttag	gaacatttgt	gttgtggcaa	ttgtatgata	cctttaatgg	118740
gaatatttca	aagacacttg	ttaagacttt	gttagaacia	aatgtagagg	gtgctggatg	118800
tccctgaata	ttcttccgcc	tcctgtaact	tgtattgctt	tggaaatttcc	agtggcctga	118860
caatgaacta	ctgcaggat	ccagatgccg	atacaggccc	ttggtgtttt	accatggacc	118920
ccagcatcag	gtgggagtac	tgcaacctga	cgcatgctc	agacacagaa	gggactgtgg	118980
tcgctcctcc	gactgtcatc	caggttccaa	gcctagggcc	tccttctgaa	caaggtaaga	119040
agtctgtgtc	ttaccttgtc	tagcacatac	ctctctatgt	gcttggacaa	cgggatgaaa	119100
agacatgaaa	aaccacactg	atgcagaagc	ctttagtgtc	acacgggagc	tcgagtgttg	119160
gttgagggtt	tgccatgacc	aaggaagtct	cagtgccgtc	cctgggaaag	ccagagctgt	119220

BI OL0250W0SEQ_ST25. txt

gatttttggc	acaacttgtg	ggagtagtga	ctttaggact	ggcgcaaaac	ctccaggggtg	119280
ctcaacttaa	ccactcacct	tattctaaaa	tgggttattt	cagtgtccca	gtcaaattcc	119340
tattctaaac	tgctgtcaac	tgtgtgatta	tttccaagcc	aataagcatt	tccagtaatt	119400
tcttaaaata	gtgttcattg	cagtcttcag	cgttgtggct	cctgagggat	gtggcccctg	119460
attctgtcgt	cctagagaag	cctgacatga	ctgcattgat	tctgtatcgt	cctgggtcta	119520
tgtggctgcc	tggctgtctg	taatcatctg	ttttatTTTT	atTTTTtct	acagactgta	119580
tgtttgggaa	tgggaaagga	taccggggca	agaaggcaac	cactgttact	gggacgccat	119640
gccaggaatg	ggctgcccag	gagccccata	gacacagcac	gttcattcca	gggacaaaata	119700
aatgggcagg	tctggaaaaa	aatgtaagcc	actttgattt	ggactctttt	tccttttgct	119760
gacaaatctt	ttcaaacaga	agaggggcag	aggaaaatac	tggaaagact	tcaggaggct	119820
aagcgtaatt	agccttagca	tggaaagtgc	aagcagcaca	ggccagcaaa	gccccacgcg	119880
tgtgggggtt	ctcaggcctc	ttctcttttg	acatttcttt	actgtttcca	ttgttgggtg	119940
ctgtttctcg	tttctagtgc	ttgtcctcta	agccaggggt	ccccactcca	gtactggtac	120000
tgggtactgg	actggaactg	gtaattatct	gtggcctggt	aggaactggg	ctgcacagca	120060
ggaggtgagc	ttcggggggag	caaacaaagc	ttcatctgta	ttttctgctg	cttcccatca	120120
ctctcatagc	tgcttgagct	ctgccagctg	tcagatcaga	ggcagcatta	gattatcata	120180
gcacaaaccc	tattgtgaac	tgacatgtg	aggaatctag	attgcatgct	ccttatgaga	120240
atctaattgcc	tgatgatctg	tcatgcttcc	atcaccccca	gatgggacca	cctacttgca	120300
ggaaaattag	ctcagggtc	ccactgattt	taccttatgg	tgagatgcac	atttatttca	120360
ttatatatta	caatgtaata	ataattgaaa	taaagtgcac	gataaatgga	aggtacttga	120420
gtcatccttt	aaccatcgcc	ccctcacccc	aggtgcacag	aaaaattgcc	ttttatgaaa	120480
ctgggtctctg	gtgccaaaaa	agttggggaa	ccacactgct	ctgggttcta	gtagtcaag	120540
atgccctcta	tgaggcttaa	gtcagatttt	tctagaaaag	atttgatgg	gccatcaggt	120600
caccatgaga	cttcccttag	cctcatgcat	tctctgtgat	ggtttacttt	ggggcctatg	120660
aataggggaag	actgagatat	aggaaaaacc	aaagtgtctg	tgttccccca	ctctcacacc	120720
catgtaacat	aacacttctc	acaccagata	tggggggatt	tctcctcaca	ccccagcgga	120780
gtctccagca	gataccagct	gggtgtccta	caatgtaact	cggtcctgac	actctatctg	120840
gagacagtgt	cagatccac	aagttaaggc	tcagtcctac	aagactgccc	cactgcagat	120900
gccaatccca	agttgcaggc	tgtgacctgt	acttctgccc	agctggataa	agatctgttt	120960
ttctatatga	ccctccatgg	gtttgattac	tttgctagag	tggctcacag	aactcaggga	121020
aacacgttac	ttttattttac	ccatttatta	taaaagatat	taaaaaggat	cctggtgaac	121080
agccaggtgg	aagagatgca	cagggaagg	cacgtgggaa	ggggctcaga	gcctctatgc	121140
cctctccagt	gcaccagtcc	ccagtaccct	aagtgttcag	caaccagaa	gctctccaag	121200
tgcagtcttg	ttgggttttt	atggaggctt	cattacagag	gcacagttga	ttacatcatt	121260

BI OL0250W0SEQ_ST25. txt

ggccatcggt gatcggtca ccttcggccc ctcttcctc cctggagggt ggaggggtggg	121320
gctgaacagt tccaaccctc aagtcacatg gttgggttccc ttggcaacca gcccttgggg	121380
ctatccagga acccaccaag agttgcttca ttgcagctcc cttcacccag gaaactccaa	121440
gggatttagg agctctgtgt taagaactgg ggggcagaga cccaatatac atttcttatt	121500
ctatcacaat atcacaggaa gctaaggatg atactgcctt tgtgtgtctt ggctgtggat	121560
ggtgcataat gcatggaagt aagcatttct gaatcaacag caaacaggct ttatcaggta	121620
gaagaccct cagcgcccca gggacaaagc tcatcaatga tgtcccactg tcctctgagg	121680
ctctagctct aagacctcca gtgggtcaag ctcttgaga agtggcacat tctccaaaga	121740
cccttcaggg tcaccacacc ctgggtaagg gtgtggcctc ataactcctt ttgactatga	121800
ctgatggctt acagcataga aagaaataac tttgtcaaaa aatataataa tgatagaaag	121860
gaagaaggaa cgctcccttt tgtcttctaa gaatagatgt gaaatgtgtg tgccttagaa	121920
tatcttctcc ctctcctgct ccacgtgagc tggagcttac atgcctgctt gttttcagta	121980
ctgccgtaac cctgatgggtg acatcaatgg tccctgggtg tacacaatga atccaagaaa	122040
actttttgac tactgtgata tccctctctg tggttaagttg ctttctgttt tggttaaggaa	122100
actgcttcct taatatggat ttggaaaaaa aaaagcaaaa aaaacagaaa atggcttttg	122160
agctgagtgc ttctggggag gagatggctg ccctctccac cagagcctgc ttttcatcat	122220
ggccaccttg aacctgccct actattggcc ccatttggtta ggaaaacacc cgcccctccc	122280
accacacaca cataaataaa ataatgtca aattcccaaa gggcaaactt agaggtgac	122340
taatcagccc gggatagtcc caccgaacct ttctttgtct agcgtgggat gcatgaaaaa	122400
caaatttaga gtcattatga tgaaaaactg tcctcttctg cagctgagaa gaaaaaaaaa	122460
atacgagcag caggaaacag ctaagcatgt aatgcacatt gtaaacctca gatggccatc	122520
ctaggaaatc aatgaagggt agtgcagctc tttagcccca gatggccttt ctcgtaagat	122580
tactactcat gagtcccatt agcgacattg cttagagact gcttgttagg ttcttctc	122640
attgctctga gactcttatt gggagtatga ggcttggatc aggggaaggg gaattgacat	122700
tagatcttaa atgattgggg taacaaatcc atgggggaaa aaaagccact tgtacttggt	122760
ccctattttc ttctgctga ccaatcaact tgtctgtccg agttacagaa caccaccctg	122820
gacttttctt ttgtgtaatt tgggtgcttg tggttgggtc tgccatgtga agggacctg	122880
agctggggga agaagggttg cctccaagtc cactgaagac cagcatcctg agattgcctg	122940
gggagggtgg acagggcagt gatgaagatc atgggagcca cactgcccac cgtcacattt	123000
gggccactcc tggggagagc aagaggaag aaggagaggt tagggtgata ggaaagattc	123060
tacttggcca atattattat aatgtggcat tgtggtctct ggatttagtg tgagttgata	123120
gctgactttt ttctcgagtg ggtgcttttg ttctattttg tcggtgctat tgcagaagca	123180
tcttgggtgg tcctctacct caaagtctct tgatggggtc agttccagtt ctccgcttct	123240
ggcccatct agtacacgc actgcctctc actgcctggg ctctctatcc ttgacaggct	123300

BI 0L0250W0SEQ_ST25. txt

gccttgaatt taagcccagt ctgacttacc tgcctcaaac acccacagta gtgcctggga	123360
ctcatgcacc tttgactccc atggaaggga agtgcagtag cttcccaggt gcaattctgc	123420
tgtcctcacc cacattgagg atgtatgaga atcaggttct tagagattgg agaaagaagg	123480
aagaatggga acaagatttc ttccaatgga ctgtgaggtt ccccacctta ctttgatgta	123540
agacaagtga ggtaacccc aagcctgggtg aggaggggtc ccatcagaca cttggaaatc	123600
ctgaggactg tttcctgcag aaggatgtgg ttggtgggat attcaggttt gactcatgat	123660
tgagaaagt agagcctctg gttggagaaa gagtttaata actatttcat ttccaccaac	123720
acattcagta cgaataataa ataagtaaaa ataaatagaa acattcagtt ttattttgaa	123780
tagtaggagt aggggtataat ttctgtagtt actcttttag tacaatgatg catgtttact	123840
gtatgtaagg catactagca gaaattgagc tcagcactag aaaagatgat tgcattccat	123900
gccatgcttc ttttttacia aagacttcta tagatagatt ctcaaaacaa cccacagcaa	123960
atgaaaagt atttgaaaa ctcaggttcc agattcactg gagtgtagaa tctctggttg	124020
gttggggagg aatttcctct tgcagttgtt attaataatt atatgaataa ttattaacta	124080
tattaatatt tatagttttg aagacctga agggctggag acaacagaga agcatttttg	124140
aacaccctct gtagcccctg cactgttgta ggcattgatg ggtggtacca aagatgggac	124200
actttcccta cctccagaga ccttgtgggc ttgctgcaga gagaaggcag ggaggaggaa	124260
aagaagaata gaggcacatg tgtgtaaatt acccccacag cagtcagtta gtcatgggag	124320
gctccccaga agaactgtcc tgaagctggc tgagagaagg caacatttca acataggaca	124380
gttatccttg ctacataaaa tcacatacac acatgcacat atgtccacac acagagactc	124440
acatgcaaaa gaatcctttg tgcctttcag taaactttac atggttttaga aagaacttat	124500
atttccttga aaggagagtg tcctttgttg ttactacca ctttttaaac ttagaaagaa	124560
aaatctaaag agtgtttatg attttacat ttaatttcac ctttgagatg tgaaaaacta	124620
gtgcttggaa ttcgtcctga attaaacgac acaattgcta acttggaact aaatgcgact	124680
tcttttccca ccttgtgcca cagcatcctc ttcatttgat tgtgggaagc ctcaagtga	124740
gccgaagaaa tgccttgga gcatgtagg ggggtgtgtg gcccacccac attcctggcc	124800
ctggcaagtc agtctcagaa caaggtaaga acaggcccag aaaccatcta tactgtcctt	124860
ccatgtaagc cccacaaaac ccttctacat ttacacagaa cccacacagc tgatgcatca	124920
atacctgcct ctctgttttc tgaaggagga aaaaatatag aaaaattaaa aaaagttata	124980
ttattatagg ttctctactt ggaaaatagc caaaatacaa atctttttct tgatctgggc	125040
agttccatca aaatctgtag gcacagtgat ttgcaccaag ttccaatact tttggaaaat	125100
attgaagatg ctctgagggg ttctatggat atccattgtc tctactgtcag atgaaaagaa	125160
agggaagttt ttagaaatgt gacactttgc agtgaggagg gacaagagca aacttaccta	125220
cagtctatca caggcacaga ttttttttta cacttttgtg aatcattgaa ttcaatgccg	125280
aggctattca tctattcaca aacacatgaa caaattatgg gttgtgatcc ccataaatga	125340

BI OL0250W0SEQ_ST25. txt

agagtaatca	gtccgaaccc	acagaacctg	gacatttttg	gtatcgtttc	agtggaacat	125400
gcaattcgta	agttcagttt	gcttgggtgt	ctcttaggaa	gaacacatag	gacacagacc	125460
catctgcctg	catgttttgc	ttcctcatct	cctttctaca	ccagggcacc	tgtgctcaat	125520
tgctgttctc	ctctaaagag	acttccttct	gtaagtttgt	gaaatgccat	cgacaaacct	125580
gatcgcatcg	catttcactc	tgctgttgag	ttgatttttc	tttactttat	cgtttgtaac	125640
ttcttgctct	acagagcttt	caccttcac	atatttcaga	ttcattcttt	cctaaactgt	125700
gtgggtggtct	atgtcctcac	tgactatcaa	catactgcca	tcatgcactt	cctatctcta	125760
ttcctcttcg	ttgcaatctg	gctccaagt	gctcacacca	ttattctgat	ctatcaactg	125820
cctacacagt	cctagaaagt	aagtgagtca	agaaacatcc	cccaaaagta	aacttttcag	125880
gtaagatcag	aagaccctca	tgagtcactg	ctgctcagga	tcgtatctgg	ctccttgaag	125940
agtgcacttg	catagatctt	gtcataaaaa	atgaaagaga	ccttgggaag	gtcttgggct	126000
ggtcactttt	gtcagagtcc	agggctgtgg	ggtgaaagcc	acagctatag	agcttcattc	126060
tggagtcact	tagctttgct	ctcctgggga	caggctgtgc	ctattcttgc	ctcaggcatc	126120
aaaaaaagt	gcacagatgg	gcccttctga	aaaatctcac	tactggagca	cagctcgaag	126180
tttctactat	cctgacgttg	ggcggtagtc	ctttgctttg	ggaatatgaa	catgatcaaa	126240
actgagtga	cttgtcttcc	tggctttctg	tacaatgaag	tagaacaac	catccaattt	126300
gaccaaaagcc	ttggcatgtt	ttctttctag	gtttggaaag	cacttctgtg	gaggcacctt	126360
aatatcccca	gagtgggtgc	tgactgctgc	tcactgcttg	aagaagtacg	tttaagggaa	126420
aactgacatg	gggtcttata	ttcaagactt	ttttcctccc	tctcttctc	catcccttct	126480
ttcttccac	cctccccttc	cttctctccc	acctctcttc	cttttctgga	aggaacacta	126540
ggaaccaggg	aatgcatgca	gaatcctgag	gcagaatttc	cagggcaatt	ggatgagaga	126600
ggaggggaagt	gtttctagag	ggaatctgca	gaggggaagac	ccagtgaag	tgattttttg	126660
gacctgtata	aaccgcagga	cagagctgtt	cactaccaga	ggcatcaatc	tgtattgcat	126720
tgctctagag	caatatctga	ggctgaataa	tttataaaga	aaagagttta	attggcacat	126780
gtttctgcag	gctttacagg	aagcaggatg	ctgtcatctc	ctctgcttct	gtgtgggcct	126840
aaggaagatt	acaatcatgg	tggagggcaa	agtgggagca	ggcatgtcac	atggccagag	126900
caggagcaag	agacagagag	agatggggtg	ggggtgctgc	acaataccaa	atgaccagac	126960
tttgcaagaa	ctaagagtga	gagctcactg	atcaccatga	agatgtggcc	caagccattc	127020
aagagggatg	cacctctatg	atccaaaccc	ctttcacagg	ccatagctcc	atcactgggg	127080
actacagttg	aacacgagat	ttaggtgggg	acaaatatac	aaactatata	acagtctctg	127140
atgaaacaga	ttgagaacag	accttaactg	tcagtttcca	gcaaattgtg	aattttgttt	127200
cttgccactc	ataagtcact	gattctgggt	ggccgaggggt	gtcagaggga	cagcgccaag	127260
ttcatggcac	agaggatacc	tgaaggggct	ggacatatt	tttctcttga	catcctcatc	127320
ttttctaggt	cctcaaggcc	ttcatcctac	aaggctcatcc	tgggtgcaca	ccaagaagtg	127380

BI OL0250W0SEQ_ST25. txt

aacctgaat	ctcatgttca	ggaaatagaa	gtgtctaggc	tgttcttggga	gcccacacaa	127440
gcagatattg	ccttgctaaa	gctaagcagg	tactcgctca	cctgtggtct	tcaccccacg	127500
ctggtgaaga	tatttgcttt	atgtctgggt	tttatgggcc	atggccactg	catggcagtg	127560
gggaggaact	gtctatcaca	tgaaaggctc	aagggccttg	gggacagcat	caatcttcaa	127620
ccccagccct	gccacatgtt	agttgtgctc	tttaaaaagg	cagaaggatt	cgtttctca	127680
cgtggaaaaa	gagataccct	gttaccgta	aaacttactt	aatgttcacc	agttcatcca	127740
cattcatgat	cagggaaagg	ttgttattcc	aggctaacta	ttctcctttc	ataataatat	127800
gctggagaga	atcaaagtag	attgcatttc	aaagcgcttg	aaaaaccacc	atatcgagcc	127860
atgcttagtg	tgggcgcctc	taatcactgc	tattcaggag	gctgacgagg	aagaattgct	127920
tgagcccagg	acttcaaggc	tgtaggcagc	tatgattgtg	ccactgcact	ccaggctggg	127980
tgacagatca	agaccctgtc	tcaacaaaag	aaaagaaaac	aaaacaaatg	aacagaaata	128040
ttccacaatg	tcaaaaaaaaa	aaaaaaccca	cacaacatac	aatttacaaa	tgcaaataat	128100
aatattattg	ttgtcttctt	tgattttctc	tttctgggtg	aaattttggt	ttattaagcc	128160
tgacaaaagt	atacctttgc	ttacatcact	taaagttagt	ctatttggac	ctaggtgaca	128220
gtacaatcag	ctaagaaaca	gtattttag	gagaggcagg	tttgggacag	gtgacaaggc	128280
atgtggggtg	ctcgctgtgc	tgggtggctct	ggaaggcagg	gtgtcaatgc	agacagggat	128340
gagcatggcc	tggttgggaa	ggcatggggc	aggcaggagc	ctgagctgct	ctcctgggcc	128400
tggtcacaag	cccatggcag	cttctctggg	tctgtgaact	gaggggtgat	gtcctggaat	128460
cctctgacac	tctaggaagg	agagaagggc	cttctgggt	cagcctttat	aaacagtagc	128520
tgatctccct	cttgcctccc	agggtcctcc	ccaccatccc	agcaaagtgt	caaatacaag	128580
atctctgctc	ctcatggtcc	tcagagagct	ggggtgttct	gatggcttga	acaagtcact	128640
taggaaatgt	ggggttttgg	aggcattctc	tgataggctg	atacgttttg	agtttagagt	128700
tcccaccgca	catccccaca	cccctagagt	ctagggcatt	tagtgctcca	tgagggaacc	128760
tgtagagtga	ggacatctgc	atcacaggct	gggccttcta	gtgtccagaa	gcagaaagtg	128820
tgtctgcttc	aaagttgggt	ctaagtagta	tttttgggtca	gaatacggca	tttctcattt	128880
ccattccttt	atcccccttga	acttactaaa	gtagaatcag	gtctaaaaac	cagagttcta	128940
atctttaaga	gtccctggga	ttctaaggta	tatgaatgtc	cttggaaaac	aataccattt	129000
agttcatgca	agggtgcttat	ttcccatcct	ctttcatttg	atgtctagca	ttttactgca	129060
ttcttaccac	cacgggtttag	taacattcac	gaggaggaag	tggaggatcc	agatggagca	129120
acttgctctg	ggcacacaag	gcatttgcaa	ttttataccc	tcttgatgat	gtctcagcca	129180
gacattctgc	ccagtcatca	atgccctctt	caattaatat	gaaaggacac	acttggcatg	129240
agattccaat	cgtgcacaga	atatacatga	gaagtgtgcc	tttgtcatcc	ctactttcaa	129300
aggctaaggc	caccctcagt	ttcttgcatg	caactgatgc	ctttcaaagt	aaaccttaca	129360
tctgtgtagt	ccataggcaa	ccacaggcaa	atgtgagggt	gaaacgctgt	gttctacatt	129420

BI OL0250W0SEQ_ST25. txt

gttctgtgtc	agtgaagcaa	ggcagtgcc	gctcagaggg	ctctggggct	tcaaggcagg	129480
gatgcctggg	tgtaggtact	gccacttcca	gctgggcagt	gaaacataac	tgctaatact	129540
ttccttacag	gcctgccgtc	atcactgaca	aagtaatgcc	agcttgtctg	ccatccccag	129600
actacatggg	caccgccagg	actgaatgtt	acatcactgg	ctggggagaa	acccaaggtg	129660
agatcaattc	cattgcccac	gtaacaaatt	gtttttgacc	ttcagtgc	gttacaaaat	129720
gagcattttg	gagatagttg	tacaaattcc	tacctatgaa	tgtggtctac	ccactcctga	129780
ctttgcctgg	acacctgtct	atgtctccat	aatcagtcct	caagggactt	gggcaagggg	129840
agcggtgcca	tttcttgag	tctctctctt	ttttgttttc	agaatctttt	aatttttttt	129900
gtaatgattg	tatgtttccc	ttacaacaaa	aacaaacacc	agtagaggtc	tttgagtctc	129960
ttaatcataa	ttcagcatt	catattgctt	ccccaggtaa	gtggggtttt	gaccagccc	130020
tcaagttaag	gggtttagat	tatttttcat	gtgaaattag	acagactgcg	tttctaaaca	130080
tgggtgcaaaa	cagtaacgac	aaaagttgta	attaaactat	tcttcttccc	aaatacccac	130140
atgtctaata	tgtgtgtgag	gggtttagtc	aggggacctg	aagctggggg	agaggcagac	130200
agttcccatg	gccccaaagc	taggatggca	tttggtattg	gttgatgggt	gagagcaaga	130260
gaggggaatat	ttttgtgcat	gatgtggtat	cagcacctgt	actacatttt	atggattcct	130320
tcttctcttt	gcggtatgcc	ctgacaataa	ttatatccgt	cagccttacc	cccttggcag	130380
taggaaaact	gaaactgtct	taaagtctca	gctctacttt	ctcagagggtg	caggcaaggg	130440
cactgggagt	ctggggccct	ggaaaactgt	tctgactctg	ccacttgcca	gatagacctg	130500
aactagacac	gttacctctt	tgtaccactt	ggctctaata	ccttatctgt	aaaaccagca	130560
ttttcaaatg	gtgctttgca	catcagcctt	ttgcataagc	tttgatttga	taaaatgttt	130620
tttgtgtttt	taaaaagatt	aaaaaccaca	ggtttagata	atttcaaagt	aggcttcctt	130680
ttttctgtca	ttttcctatt	atttttaaaa	cctcacctcc	ttgactcctt	gttccctttt	130740
tctgcactgc	tgagtctggg	agcactgagg	ccaggtaaaa	ggaaacttgg	caaatgaggg	130800
gcacctatgg	gtgtgggagg	ctgctcctgg	tgtttgcata	ttttaaaatt	taaatgctac	130860
aaaccactgt	gagttaggta	ttattgttcc	tattttacca	ttgaggaagc	tggggctcag	130920
agaagggtga	gggtggtaca	gacaaacctg	aattggaacc	ctggctcctg	cctatgggct	130980
gtcaggactt	agaaaagtcg	tgagctctcg	ctgattgttt	cctcagctga	tgtgggctgc	131040
agggctgtta	tgggggaaat	aataagaaag	tgcatacaag	gctgagcaca	tcctaagcac	131100
tccatcatgg	cagctcctac	tactaataaa	gaatagaatt	atatctaaca	tgattctttc	131160
ttgcaagtga	cagaaaatcc	aactcaaatt	ggattaagca	aaacaaggga	aattcttagt	131220
gagctgcaaa	gttttcaggc	tcacatgatg	gccccaaatc	ccaggctcctc	ccaatcatgg	131280
agtaggcact	atttgggggc	acaaagggtga	cattcccatg	gctgcagatg	ctgtggtgct	131340
gtggctgtac	cgggaaagaa	taagaaaggc	cactctccca	attatgtgaa	caatagtctg	131400
cccactctga	gaagtcaaac	ttgggtcaca	gtcctgcccc	tgaacccatc	actgactggc	131460

BI OL0250W0SEQ_ST25. txt

tctgacctgc	accaattggt	ccatgttggga	ggtgaaggca	agacccact	aatacccata	131520
aggggcaaaa	gtagataga	tccttcaaga	ggattatggg	aggtagggca	aaaagctgct	131580
gggcagccag	aaagcaaaca	gagcctctat	gataacctca	ctgatgaaag	catgaagcta	131640
aaatcataag	gatctgggtg	tgagttctgg	ctctcccatc	ttccatgtga	cattgggcag	131700
ttatttaatc	tcttttagcc	tccgctttct	catcttacat	atgagataat	tgtgaggatt	131760
aagattacac	ataatcatca	tcatcaccgt	ccaccactac	caccatcatc	cccatcaaca	131820
tcatcgccac	cactatcatc	attcttactg	gcactacat	caccatcacc	accattccac	131880
caccatcacc	aatatcatca	ctgtcaacat	cattaccacc	atcaccatca	ccaccacat	131940
catcattact	accactacca	ctactaccac	catcaccatc	accaccattc	caccaccatc	132000
accaatatca	tcacttcaa	catcatcacc	atcaccatca	ccaccacat	catcatcatt	132060
actaccacta	ccactactac	caccatcacc	atcaccactg	tcccactact	atcagcatga	132120
catcaccatc	accaccacca	tcatcattac	caccgctact	accaacatca	ccatcaccac	132180
aattctactg	ccatcaccat	taacattacc	accaccatca	tcactatcac	catcaccacc	132240
atcatcacca	ctgccattat	cactgccacc	atcatcacta	tcctctatat	ttcctcatct	132300
gtattatcat	tactaccacc	atcactatca	ccaccatcgt	caccatcata	atcaccatca	132360
acaccatctc	caataccacc	atcactgtaa	ccatcatcac	caccaccatg	atcactatca	132420
ccatcatcac	aatgatcact	gtaaccatca	ttactacca	ccaccatcac	cactactcca	132480
ccaccatcac	cattatcatt	accatcacca	ttatcaccac	catcatcatc	accagcacca	132540
ccatcatcac	cagcaccacc	atcaccatca	ccatcattaa	caccatcact	atcaccattg	132600
gtttaatcat	caccaccatc	atcataaata	aacatcacat	aaccaggggtg	tagctgggtg	132660
ttgaccccag	agcccactca	ctgtttcctc	tctcccaccc	ccatccacac	atttctaacc	132720
accatcctgc	actgggctcc	cagtctcctc	tggtctcacc	cacatgtcca	ctgagaaaag	132780
gattttcaga	acaccaacta	gaccaggagg	agccacatac	ataactcagg	cctgcttatac	132840
aactttctac	atgttaataa	tgacatcaga	tcaatgggtg	ttctcagctt	ctcagaagga	132900
ggtcaaaaatt	ctccccctct	ccccttcatg	tgtccagacc	ttcccggatt	tggatgtacc	132960
aagtgcagag	tggtgttgag	gccaaggggc	tcatccatgt	aagtctcatc	tgcaatcact	133020
gggctgatcc	cgtggccctg	tctccagggc	gccatcagag	agggcttcaa	tcctcagggtt	133080
acctgtggcc	caccctgccc	tcagaggtgc	catctctaca	ttggccacga	gatggcagca	133140
catactcata	gactgcatta	atttcccagc	aactcctggt	gggttttccc	tcttatcagg	133200
atgtttgcct	tgctcagaga	gcaaactctga	gagcagtgac	acctaactta	actttcagca	133260
aaatatTTTTg	agaaggggtgc	ccctttacac	atctgtgcag	tccaggtgat	gcatcccatg	133320
cccaatgctc	ggtagtcagg	aggagcttcc	tccatgcagc	tctgcggaag	agactcttcc	133380
acgctgctca	tgtaaactcc	agattcgggtg	tcagttttct	gacaccgaag	acaatgatct	133440
aagtgcagtc	aagggctttg	gggaaagcag	gagagagtgc	ctcagttcta	gcctgtgcc	133500

BI OL0250W0SEQ_ST25. txt

tgcttgcaaa	gttttgcaaa	attctaata	gagctgggct	tgcaacattg	gaaacttgga	133560
ttatttgtga	gagcactgag	aatccctgg	gcatgtccat	ctggaaaaaac	agcatttctt	133620
ctggcacttt	agcagagggt	ctgtttcaat	ttggcgaagg	aaattaagca	gtttttcaca	133680
aaagaagaac	tacaacgagg	agaattgtcc	ctagtatttc	ttctccctaa	ttgtcaagga	133740
agtgtaaatt	agaaaatgaa	tcaggacaat	ttccacctac	tatgttagct	aatattttta	133800
aaattgaata	tcacaagggg	gaggcaaagt	aattgttttc	cagtgcatt	ttccactgtc	133860
acaccctttt	agagaataat	ttggcaatgt	tactgtgaga	tagaaatatg	tctatataat	133920
tatgggaact	gagacttcag	aaagtaataa	ggaataagaa	tgaaatttat	gaacaaacat	133980
gtggaagggt	ggaagcaaga	gtggggccaa	cacgcatggg	gaggaagcat	ttgggcagcg	134040
actccgcaga	cccagactca	agctgagcta	tacaacctcc	ttacgcctca	gtttcctcaa	134100
ctgaagaaca	ggaatgacaa	gtgcctgttt	cataggaccg	ttgtgaggat	taagtgaagt	134160
ataccacatt	atgagcttgt	gcctggaaag	gttgattctt	agtaaataat	gactattctt	134220
ttttattgca	ataaaattta	tacaacatag	agttactatt	ttaaccattt	ttgcaggtag	134280
cactgagtgg	cattcagtac	attcacaatg	gtgtgcaacc	gtcaccatat	ttccaggaca	134340
tttttctcat	cccaaagga	aacctcatgc	ccattaagca	gtcactcctc	attaaaatat	134400
tagttatgaa	gactgtagca	tttttttaaa	aactcatgat	ataacattga	ttgaaaaaat	134460
cagtatagga	aattgtgcat	tatgatgtaa	tagtaaaaga	agcatataaa	aatctgaaaa	134520
aagtatataa	aaagaatagc	aattgtattt	ctcagactct	ctttacattg	taaaaatcat	134580
tttgatagct	tcaaaagaaa	agcaaaaagt	acacaaacaa	caaccaaccc	caaagcagca	134640
tgacaaagcc	cagattgttg	aatccaggtc	ttgggaacat	aaaatcttat	atgacatttg	134700
cactttaatg	ggtcagagag	tccagtggca	ttgggagctg	ccttgtgttc	tgcagcctca	134760
cggacagaca	ggagggtccg	ctccactgct	ctgttcttct	ggaatttcct	cgtgaacaag	134820
ctttggcctc	agtaaccatt	tctttcatct	ttttaaacac	aggtaccttt	gggactggcc	134880
ttctcaagga	agcccagctc	cttggtattg	agaatgaagt	gtgcaatcac	tataagtata	134940
tttggtgctg	gcatttggcc	agaggcactg	acagttgcc	ggtaagaaaa	gatcaataga	135000
tcaaagtctt	gtgctctccc	gtctcagtct	cagtccctta	gacgtcagtc	caaagtggc	135060
aaattcagga	aggttttgtc	agtggaagac	cccagtctaa	gtgttgctca	gaaactcccc	135120
agatctgtcc	ctgaatgcat	attcagatca	tctaaggaga	cgtcttgggg	cttgagttcc	135180
agatccatag	caaggagacc	gtaagtgcc	taactacctc	aggccactca	ccttcctggt	135240
gtgtgctggt	caccagtgc	tgaagtggg	gcttttccag	tagagaggaa	ggtagagggt	135300
acaggaccga	gacaaattac	acacacttaa	caatgatgtc	caggctagcc	cagtctaaag	135360
gaaacaccaa	gttaggaagc	aatgcatgca	ggattcacia	gggattattt	tttttccag	135420
gaaaaaacta	agtgatgtgg	ttttgttgaa	tagactttgc	taagtactta	agcactgcag	135480
atgcttgagt	aatatgctca	taagttcctt	tctgatttga	attactggga	aaatgtacat	135540

BI OL0250W0SEQ_ST25. txt

atggataaga	gaaggatggc	atcccatatt	aaaaggttgg	cagcttaaag	ctcacatgaa	135600
ttttccccta	cctctgttta	gggtgacagt	ggagggcctc	tggtttgctt	cgagaaggac	135660
aaatacattt	tacaaggagt	cacttcttgg	ggtcttggct	gtgcacgccc	caataagcct	135720
ggtgtctatg	ctcgtgtttc	aagggttggt	acttggattg	agggaatgat	gagaaataat	135780
taattggacg	ggagacagag	tgaagcatca	acctacttag	aagctgaaac	gtgggtaagg	135840
atttagcatg	ctggaaataa	tagacagcaa	tcaaacgaag	acactgttcc	cagctaccag	135900
ctatgccaaa	ccttggcatt	tttgggtattt	ttgtgtataa	gcttttaagg	tctgactgac	135960
aaattctgta	ttaaggtgtc	atagctatga	catttggttaa	aaataaactc	tgcacttatt	136020
ttgatttgaa	ttaatttttg	ttttgggtctt	caaaattttc	atgctctttt	catcccatct	136080
atttttattt	ttatttttta	gactttacgt	cctggggtag	atgtgcagaa	tgtgcagggt	136140
tgttacatag	atgtacacgt	gccatggtag	tttgctgcac	ccatcaacct	gtcatctaata	136200
tcggtatttc	ttttagttct	atccctcccc	tagccctcca	ccccttgaca	ggcccagggtg	136260
tgtgatgttg	ccctccctgt	gtccatgtgt	tctcattgtt	caactcacac	ttatgagtga	136320
gaacatgccg	tgtttggttt	tctgttcttg	tgtttagttg	ctgagaatga	tagtttccag	136380
cttcatccat	gtccctgcaa	aggacatgaa	ctcatccttt	tttatggctg	catagaattc	136440
catggtgtat	atgtgccaca	ttttatccaa	tctaacattg	atgggcaatt	gggttggttc	136500
caactctttg	ctattgtgaa	tagtgccaca	ataaacatac	gtgtgcatgt	gttttcatag	136560
cagaatgatt	tataatcctc	tgggtatata	cccagtaatg	ggattgcagg	gtcaaatggt	136620
gtttctggtg	ctagatcttt	gaggaatcac	cacactgtct	tccacaatgg	ttgaactaat	136680
ttatgctccc	accaacaata	tcaaggcatt	cctattttctc	cacatcctct	ccagcatctg	136740
ttgtttcctg	actttttaat	gatcgccatt	ctaactggca	tgagatggta	tctcattgtg	136800
gttttgattt	gcatttctct	aatgatcagt	gatgatgagc	ttttctcata	tgtttggttg	136860
ctgcataaat	gccttttttg	gagaagcatc	tgttcatatc	ccttgcccac	tttttgatgg	136920
tgttgttttt	ttctggtaaa	tttgtttaag	ttctttgtag	attctggata	ttagcctttt	136980
gtcagatgga	tagatggcaa	aaattttatc	ctattatgta	ggttgcctgt	tcactccgat	137040
gatagtttct	tttgctgtgc	agaagctctt	tggtttaatt	agatctcatt	tgtctatttt	137100
ggcttttggt	accattgctt	ttagtgtttt	agtcatgaag	tcttctccca	tgctatgtcc	137160
tgaatgggat	tgccctaagt	ttcttccagg	gtttttatgg	ttttagggtt	tgcatttaag	137220
tctttaatcc	atcttgagtt	aatttttgta	taagtaatgc	ccttctttgt	ctcttttgat	137280
ccttggtggc	ttaaagtata	ttttatcaga	gactagaatt	gcaatccctg	cttttttttt	137340
tctttttgct	ttccttttgc	ttggtaaaata	ttcttccatc	cctttatttt	gagcctatgt	137400
atgtctgcac	atgagatagg	tttcctgaat	acagcacacc	aatgggtctt	gactctttat	137460
tcaatttgcc	agtctgtgtc	ttttaattgg	gggcatttag	tccatttaca	tttaaggtta	137520
atattgttat	gtgtgaattt	gatcctgtca	ttatgatgct	agcgggttat	tttgcccatt	137580

BI OL0250W0SEQ_ST25. txt

agttgatgca gtttcttcat agtgtggatg gcctttacaa tttggtagtt tttgcagtgg	137640
ctggtaccaa ttgttccttt ccatgttttag tgcttcgttc aggagctctt gtgaggcagg	137700
ccttgtggtg acaaaatctt tcagcatttg cttgtctgta aaggatttta tttctccttt	137760
gcttatgaag cttagtctcg ctgggtatga aattctgggt tgaaaattat tttcttttag	137820
aatgttgaat attggccccc actctcttcg ggcttgttgg gtttctgcag agagatccac	137880
tgtagtctg attggcttcc ctttccgggt aaccaacct ttctctctgg ctgcccttag	137940
aaatttttcc ttcatttcaa ccttggtgaa tctgacgatt atgtcttgag gtggctcttc	138000
t	138001

<210> 4
 <211> 13938
 <212> DNA
 <213> Homo sapiens

<400> 4	
ctgggattgg gacacacttt ctggacactg ctggccagtc caaaaatgga acataaggaa	60
gtggttcttc tacttctttt atttctgaaa tcagcagcac ctgagcaaag ccatgtggtc	120
caggattgct accatggtga tggacagagt tatcgaggca cgtactccac cactgtcaca	180
ggaaggacct gccaaacttg gtcattctatg acaccacatc aacataatag gaccacagaa	240
aactacccaa atgctggctt gatcatgaac tactgcagga atccagatgc tgtggcagct	300
ccttattgtt atacgagga tcccgggtgc aggtgggagt actgcaacct gacgcaatgc	360
tcagacgcag aagggactgc cgtcgcgcct ccgactgtta ccccggttcc aagcctagag	420
gctccttccg aacaagcacc gactgagcaa aggcctgggg tgcaggagtg ctaccatggt	480
aatggacaga gttatcgagg cacatactcc accactgtca caggaagaac ctgccaaact	540
tggatcatcta tgacaccaca ctgcgcatagt cggaccccag aatactaccc aaatgctggc	600
ttgatcatga actactgcag gaatccagat gctgtggcag ctcccttattg ttatacgagg	660
gatcccgggtg tcagggtggga gtactgcaac ctgacgcaat gctcagacgc agaagggact	720
gccgtcgcgc ctccgactgt taccgggtt ccaagcctag aggtccttc cgaacaagca	780
ccgactgagc aaaggcctgg ggtgcaggag tgctaccatg gtaatggaca gagttatcga	840
ggcacatact ccaccactgt cacaggaaga acctgccaaag cttgggtcatc tatgacacca	900
cactcgcata gtcggacccc agaatactac ccaaagtctg gcttgatcat gaactactgc	960
aggaatccag atgctgtggc agtccttat tgttatacga gggatcccgg tgtcagggtg	1020
gagtactgca acctgacgca atgctcagac gcagaaggga ctgccgtcgc gcctccgact	1080
gttaccgccg ttccaagcct agaggctcct tccgaacaag caccgactga gcaaaggcct	1140
ggggtgcagg agtgctacca tggtaatgga cagagttatc gaggcacata ctccaccact	1200
gtcacaggaa gaacctgcca agcttgggtca tctatgacac cacactcgca tagtcggacc	1260
ccagaatact acccaaagtc tggcttgatc atgaactact gcaggaatcc agatgctgtg	1320

BI OL0250W0SEQ_ST25. txt

gcagctcctt attgttatac gagggatccc ggtgtcaggt gggagtactg caacctgacg	1380
caatgctcag acgcagaagg gactgccgtc gcgcctccga ctgttacccc ggttccaagc	1440
ctagaggctc cttccgaaca agcaccgact gagcaaaggc ctgggggtgca ggagtgtctac	1500
catggtaatg gacagagtta tcgaggcaca tactccacca ctgtcacagg aagaacctgc	1560
caagcttggg catctatgac accacactcg catagtcgga cccagaata ctacccaaat	1620
gctggcttga tcatgaacta ctgcaggaat ccagatgtctg tggcagctcc ttattgttat	1680
acgagggatc ccggtgtcag gtgggagtac tgcaacctga cgcaatgctc agacgcagaa	1740
gggactgccg tcgcgcctcc gactgttacc ccggttccaa gcctagaggc tccttccgaa	1800
caagcaccga ctgagcaaag gcctgggggtg caggagtgtt accatggtaa tggacagagt	1860
tatcgaggca catactccac cactgtcaca ggaagaacct gccaagcttg gtcattatg	1920
acaccacact cgcatagtcg gacccagaa tactacccaa atgctggctt gatcatgaac	1980
tactgcagga atccagatgc tgtggcagct cttattgtt atacgaggga tcccggtgtc	2040
aggtgggagt actgcaacct gacgcaatgc tcagacgcag aagggactgc cgtcgcgcct	2100
ccgactgtta ccccggttcc aagcctagag gtccttccg aacaagcacc gactgagcaa	2160
aggcctgggg tgcaggagtg ctaccatggg aatggacaga gttatcgagg cacatactcc	2220
accactgtca caggaagaac ctgccaagct tggatcatcta tgacaccaca ctgcatagt	2280
cggaccccag aatactaccc aaatgctggc ttgatcatga actactgcag gaatccagat	2340
gctgtggcag ctccttattg ttatacgagg gatcccggtg tcagggtggga gtactgcaac	2400
ctgacgcaat gctcagacgc agaagggact gccgtcgcgc ctccgactgt taccggtt	2460
ccaagcctag aggtccttc cgaacaagca ccgactgagc aaaggcctgg ggtgcaggag	2520
tgctaccatg gtaatggaca gagttatcga ggcacatact ccaccactgt cacaggaaga	2580
acctgccaag cttggtcatc tatgacacca cactcgcata gtcggacccc agaatactac	2640
ccaaatgctg gcttgatcat gaactactgc aggaatccag atgctgtggc agctccttat	2700
tggtatacga gggatcccgg tgtcagggtg gactactgca acctgacgca atgctcagac	2760
gcagaaggga ctgccgtcgc gcctccgact gttaccccgg ttccaagcct agaggctcct	2820
tccgaacaag caccgactga gcaaaggcct ggggtgcagg agtgctacca tggtaatgga	2880
cagagttatc gaggcacata ctccaccact gtcacaggaa gaacctgcca agcttgggtca	2940
tctatgacac cacactcgca tagtcggacc ccagaatact acccaaatgc tggcttgatc	3000
atgaactact gcaggaatcc agatgctgtg gcagctcctt attgttatac gagggatccc	3060
ggtgtcaggt gggagtactg caacctgacg caatgctcag acgcagaagg gactgccgtc	3120
gcgcctccga ctgttacccc ggttccaagc ctagaggctc cttccgaaca agcaccgact	3180
gagcaaaggc ctgggggtgca ggagtgtctac catggtaatg gacagagtta tcgaggcaca	3240
tactccacca ctgtcacagg aagaacctgc caagcttggg catctatgac accacactcg	3300
catagtctgga cccagaata ctacccaaat gctggcttga tcatgaacta ctgcaggaat	3360

BI OL0250W0SEQ_ST25. txt

ccagatgctg	tggcagctcc	ttattgttat	acgagggatc	ccggtgtcag	gtgggagtac	3420
tgcaacctga	cgcaatgctc	agacgcagaa	gggactgccg	tcgcgctcc	gactgttacc	3480
ccggttccaa	gcctagaggc	tccttccgaa	caagcaccga	ctgagcaaag	gcctgggggtg	3540
caggagtgtc	accatggtaa	tggacagagt	tatcgaggca	catactccac	cactgtcaca	3600
ggaagaacct	gccaagcttg	gtcatctatg	acaccacact	cgcatagtcg	gaccccgaa	3660
tactaccaa	atgctggctt	gatcatgaac	tactgcagga	atccagatgc	tgtggcagct	3720
ccttattgtt	atacgaggga	tcccgggtgc	aggtgggagt	actgcaacct	gacgcaatgc	3780
tcagacgcag	aagggactgc	cgtcgcgct	ccgactgtta	ccccggttcc	aagcctagag	3840
gctccttccg	aacaagcacc	gactgagcaa	aggcctgggg	tgcaggagtg	ctaccatggt	3900
aatggacaga	gttatcgagg	cacatactcc	accactgtca	caggaagaac	ctgccaagct	3960
tggcatctta	tgacaccaca	ctcgcatagt	cggacccag	aatactacc	aaatgctggc	4020
ttgatcatga	actactgcag	gaatccagat	gctgtggcag	ctccttattg	ttatacgagg	4080
gatcccgggtg	tcaggtggga	gtactgcaac	ctgacgcaat	gctcagacgc	agaagggact	4140
gccgtcgcgc	ctccgactgt	taccccggtt	ccaagcctag	aggctccttc	cgaacaagca	4200
ccgactgagc	aaaggcctgg	ggtgcaggag	tgctaccatg	gtaatggaca	gagttatcga	4260
ggcacatact	ccaccactgt	cacaggaaga	acctgccaag	cttgggtcatc	tatgacacca	4320
cactcgcata	gtcggacccc	agaatactac	caaagtctg	gcttgatcat	gaactactgc	4380
aggaatccag	atgctgtggc	agctccttat	tgttatacga	gggatcccgg	tgtcagggtg	4440
gagtactgca	acctgacgca	atgctcagac	gcagaaggga	ctgccgtcgc	gcctccgact	4500
gttaccgccg	ttccaagcct	agaggctcct	tccgaacaag	caccgactga	gcaaaggcct	4560
ggggtgcagg	agtgtacca	tggtaatgga	cagagttatc	gaggcacata	ctccaccact	4620
gtcacaggaa	gaacctgcca	agcttgggtca	tctatgacac	cacactcgca	tagtcggacc	4680
ccagaatact	acccaaatgc	tggcttgatc	atgaactact	gcaggaatcc	agatgctgtg	4740
gcagtcctt	attgttatac	gagggatccc	ggtgtcagg	gggagtactg	caacctgacg	4800
caatgctcag	acgcagaagg	gactgccgtc	gcgcctccga	ctgttacc	ggttccaagc	4860
ctagaggctc	cttccgaaca	agcaccgact	gagcaaaggc	ctgggggtgca	ggagtgtctac	4920
catggtaatg	gacagagtta	tcgaggcaca	tactccacca	ctgtcacagg	aagaacctgc	4980
caagcttgg	catctatgac	accacactcg	catagtcgga	ccccagaata	ctacccaaat	5040
gctggcttga	tcatgaacta	ctgcaggaat	ccagatgctg	tggcagctcc	ttattgttat	5100
acgagggatc	ccggtgtcag	gtgggagtac	tgcaacctga	cgcaatgctc	agacgcagaa	5160
gggactgccg	tcgcgctcc	gactgttacc	ccggttccaa	gcctagaggc	tccttccgaa	5220
caagcaccga	ctgagcaaag	gcctgggggtg	caggagtgtc	accatggtaa	tggacagagt	5280
tatcgaggca	catactccac	cactgtcaca	ggaagaacct	gccaagcttg	gtcatctatg	5340
acaccacact	cgcatagtcg	gaccccgaa	tactaccaa	atgctggctt	gatcatgaac	5400

BI OL0250W0SEQ_ST25. txt

tactgcagga atccagatgc tgtggcagct ctttattgtt atacgagga tcccgggtgtc	5460
aggtgggagt actgcaacct gacgcaatgc tcagacgcag aagggactgc cgtcgcgcct	5520
ccgactgtta ccccggttcc aagcctagag gtccttccg aacaagcacc gactgagcaa	5580
aggcctgggg tgcaggagtg ctacatggt aatggacaga gttatcgagg cacatactcc	5640
accactgtca caggaagaac ctgccaagct tggatcatcta tgacaccaca ctcgcatagt	5700
cggaccccag aatactaccc aaatgctggc ttgatcatga actactgcag gaatccagat	5760
gctgtggcag ctccttattg ttatacgagg gatccccgtg tcagggtggga gtactgcaac	5820
ctgacgcaat gtcagacgc agaagggact gccgtcgcgc ctccgactgt taccggtt	5880
ccaagcctag aggtccttc cgaacaagca ccgactgagc aaaggcctgg ggtgcaggag	5940
tgctacatg gtaatggaca gagttatcga ggcacatact ccaccactgt cacaggaaga	6000
acctgccaag cttggtcatc tatgacacca cactcgcata gtcggacccc agaatactac	6060
ccaaatgctg gcttgatcat gaactactgc aggaatccag atgctgtggc agctccttat	6120
tgttatacga gggatcccg tgtcagggtg gactactgca acctgacgca atgctcagac	6180
gcagaagggga ctgccgtcgc gcctccgact gttaccccg ttccaagcct agaggctcct	6240
tccgaacaag caccgactga gcaaaggcct ggggtgcagg agtgctacca tggtaatgga	6300
cagagttatc gaggcacata ctccaccact gtcacaggaa gaacctgcca agcttggtca	6360
tctatgacac cacactcgca tagtcggacc ccagaatact acccaaatgc tggcttgatc	6420
atgaactact gcaggaatcc agatgctgtg gcagctcctt attgttatac gagggatccc	6480
gggtgcaggt gggagtactg caacctgacg caatgctcag acgcagaagg gactgccgtc	6540
gcgcctccga ctgttacccc ggttccaagc ctagaggctc cttccgaaca agcaccgact	6600
gagcaaaggc ctgggggtgca ggagtgtac catggtaatg gacagagtta tcgaggcaca	6660
tactccacca ctgtcacagg aagaacctgc caagcttggc catctatgac accacactcg	6720
catagtccga cccagaata ctacccaaat gctggcttga tcatgaacta ctgcaggaat	6780
ccagatgctg tggcagctcc ttattgttat acgagggatc ccggtgtcag gtgggagtac	6840
tgcaacctga cgcaatgctc agacgcagaa gggactgccg tcgcgcctcc gactgttacc	6900
ccggttccaa gcctagaggc tccttccgaa caagcaccga ctgagcaaag gcctgggggtg	6960
caggagtgtc accatggtaa tggacagagt tatcgaggca catactccac cactgtcaca	7020
ggaagaacct gccaaagcttg gtcatttatg acaccacact cgcatagtgc gacccagaa	7080
tactacccaa atgctggctt gatcatgaac tactgcagga atccagatgc tgtggcagct	7140
ccttattgtt atacgagga tcccgggtgtc aggtgggagt actgcaacct gacgcaatgc	7200
tcagacgcag aagggactgc cgtcgcgcct ccgactgtta ccccggttcc aagcctagag	7260
gctccttccg aacaagcacc gactgagcaa aggcctgggg tgcaggagtg ctacatggt	7320
aatggacaga gttatcgagg cacatactcc accactgtca caggaagaac ctgccaagct	7380
tggatcatcta tgacaccaca ctcgcatagt cggaccccag aatactaccc aaatgctggc	7440

BI OL0250W0SEQ_ST25. txt

ttgatcatga	actactgcag	gaatccagat	gctgtggcag	ctccttattg	ttatacgagg	7500
gatcccgggtg	tcaggtggga	gtactgcaac	ctgacgcaat	gctcagacgc	agaagggact	7560
gccgtcgcgc	ctccgactgt	taccccgggtt	ccaagcctag	aggctccttc	cgaacaagca	7620
ccgactgagc	aaaggcctgg	ggtgcaggag	tgctaccatg	gtaatggaca	gagttatcga	7680
ggcacatact	ccaccactgt	cacaggaaga	acctgccaaag	cttggtcatc	tatgacacca	7740
cactcgcata	gtcggacccc	agaatactac	ccaaatgctg	gcttgatcat	gaactactgc	7800
aggaatccag	atgctgtggc	agctccttat	tgttatacga	gggatcccgg	tgtcaggtgg	7860
gagtactgca	acctgacgca	atgctcagac	gcagaagggga	ctgccgtcgc	gcctccgact	7920
gttaccccgg	ttccaagcct	agaggctcct	tccgaacaag	caccgactga	gcagaggcct	7980
ggggtgcagg	agtgtacca	cggtaatgga	cagagttatc	gaggcacata	ctccaccact	8040
gtcactggaa	gaacctgcca	agcttgggtca	tctatgacac	cacactcgca	tagtcggacc	8100
ccagaatact	acccaaatgc	tggcttgatc	atgaactact	gcaggaatcc	agatgctgtg	8160
gcagctcctt	attgttatac	gagggatccc	ggtgtcaggt	gggagtactg	caacctgacg	8220
caatgctcag	acgcagaagg	gactgccgtc	gcgcctccga	ctgttacccc	ggttccaagc	8280
ctagaggctc	cttccgaaca	agcaccgact	gagcaaaggc	ctggggtgca	ggagtgtac	8340
catggtaatg	gacagagtta	tcgaggcaca	tactccacca	ctgtcacagg	aagaacctgc	8400
caagcttgggt	catctatgac	accacactcg	catagtcgga	ccccagaata	ctacccaaat	8460
gctggcttga	tcatgaacta	ctgcaggaat	ccagatgctg	tggcagctcc	ttattgttat	8520
acgagggatc	ccggtgtcag	gtgggagtag	tgcaacctga	cgcaatgctc	agacgcagaa	8580
gggactgccg	tcgcgcctcc	gactgttacc	ccggttccaa	gcctagaggc	tccttccgaa	8640
caagcaccga	ctgagcaaag	gcctgggggtg	caggagtgtc	accatggtaa	tggacagagt	8700
tatcgaggca	catactccac	cactgtcaca	ggaagaacct	gccaagcttg	gtcatctatg	8760
acaccacact	cgcatagtcg	gaccccagaa	tactacccaa	atgctggctt	gatcatgaac	8820
tactgcagga	atccagatgc	tgtggcagct	ccttattgtt	atagcagggga	tcccgggtgtc	8880
aggtgggagt	actgcaacct	gacgcaatgc	tcagacgcag	aagggactgc	cgtcgcgcct	8940
ccgactgtta	ccccggttcc	aagcctagag	gctccttccg	aacaagcacc	gactgagcag	9000
aggcctgggg	tgcaggagtg	ctaccacgggt	aatggacaga	gttatcgagg	cacatactcc	9060
accactgtca	ctggaagaac	ctgccaagct	tggtcatcta	tgacaccaca	ctcgcatagt	9120
cggaccccag	aatactaccc	aatgctggc	ttgatcatga	actactgcag	gaatccagat	9180
gctgtggcag	ctccttattg	ttatacgagg	gatcccgggtg	tcaggtggga	gtactgcaac	9240
ctgacgcaat	gctcagacgc	agaagggact	gccgtcgcgc	ctccgactgt	taccccgggtt	9300
ccaagcctag	aggctccttc	cgaacaagca	ccgactgagc	agaggcctgg	ggtgcaggag	9360
tgctaccacg	gtaatggaca	gagttatcga	ggcacatact	ccaccactgt	cactggaaga	9420
acctgccaaag	cttggtcatc	tatgacacca	cactcgcata	gtcggacccc	agaatactac	9480

BI OL0250W0SEQ_ST25. txt

ccaaatgctg	gcttgatcat	gaactactgc	aggaatccag	atgctgtggc	agctccttat	9540
tgttatacga	gggatcccg	tgtcaggtgg	gagtactgca	acctgacgca	atgctcagac	9600
gcagaaggga	ctgccgtcgc	gcctccgact	gttaccgccg	ttccaagcct	agaggctcct	9660
tccgaacaag	caccgactga	gcagaggcct	ggggtgcagg	agtgtctacca	cggtaatgga	9720
cagagttatc	gaggcacata	ctccaccact	gtcactggaa	gaacctgcca	agcttggtca	9780
tctatgacac	cacactcgca	tagtcggacc	ccagaatact	acccaaatgc	tggcttgatc	9840
atgaactact	gcaggaatcc	agatgctgtg	gcagctcctt	attgtttatac	gagggatccc	9900
ggtgtcaggt	gggagtactg	caacctgacg	caatgctcag	acgcagaagg	gactgccgtc	9960
gcgcctccga	ctgttaccgc	ggttccaagc	ctagaggctc	cttccgaaca	agcaccgact	10020
gagcagaggc	ctgggggtgca	ggagtgtctac	cacggtaatg	gacagagtta	tcgaggcaca	10080
tactccacca	ctgtcactgg	aagaacctgc	caagcttggt	catctatgac	accacactcg	10140
catagtcgga	ccccagaata	ctacccaaat	gctggcttga	tcatgaacta	ctgcaggaat	10200
ccagatcctg	tggcagcccc	ttattgttat	acgagggatc	ccagtgtcag	gtgggagtac	10260
tgcaacctga	cacaatgctc	agacgcagaa	gggactgccg	tcgcgcctcc	aactattacc	10320
ccgattccaa	gcctagaggc	tccttctgaa	caagcaccaa	ctgagcaaag	gcctgggggtg	10380
caggagtgtc	accacggaaa	tggacagagt	tatcaaggca	catacttcat	tactgtcaca	10440
ggaagaacct	gccaaacttg	gtcatctatg	acaccacact	cgcatagtcg	gaccccgaca	10500
tactacccaa	atgctggctt	gatcaagaac	tactgccgaa	atccagatcc	tgtggcagcc	10560
ccttgggtgt	atacaacaga	tcccagtgtc	aggtgggagt	actgcaacct	gacacgatgc	10620
tcagatgcag	aatggactgc	cttcgtccct	ccgaatgtta	ttctggctcc	aagcctagag	10680
gctttttttg	aacaagcact	gactgaggaa	acccccgggg	tacaggactg	ctactaccat	10740
tatggacaga	gttaccgagg	cacatactcc	accactgtca	caggaagaac	ttgccaagct	10800
tggatcatcta	tgacaccaca	ccagcatagt	cggacccag	aaaactaccc	aaatgctggc	10860
ctgaccagga	actactgcag	gaatccagat	gctgagattc	gcccttggtg	ttacaccatg	10920
gatcccagtg	tcaggtggga	gtactgcaac	ctgacacaat	gcctggtgac	agaatcaagt	10980
gtccttgcaa	ctctcacggt	ggtcccagat	ccaagcacag	aggcttcttc	tgaagaagca	11040
ccaacggagc	aaagccccgg	ggtccaggat	tgctaccatg	gtgatggaca	gagttatcga	11100
ggctcattct	ctaccactgt	cacaggaagg	acatgtcagt	cttggctctc	tatgacacca	11160
cactggcatc	agaggacaac	agaatattat	ccaaatggtg	gcctgaccag	gaactactgc	11220
aggaatccag	atgctgagat	tagtccttgg	tgttatacca	tggatcccaa	tgtcagatgg	11280
gagtactgca	acctgacaca	atgtccagtg	acagaatcaa	gtgtccttgc	gacgtccacg	11340
gctgtttctg	aacaagcacc	aacggagcaa	agccccacag	tccaggactg	ctaccatggt	11400
gatggacaga	gttatcgagg	ctcattctcc	accactgtta	caggaaggac	atgtcagtct	11460
tggtcctcta	tgacaccaca	ctggcatcag	agaaccacag	aatactaccc	aaatggtggc	11520

BI OL0250W0SEQ_ST25. txt

ctgaccagga	actactgcag	gaatccagat	gctgagattc	gcccttgggtg	ttataccatg	11580
gatcccagtg	tcagatggga	gtactgcaac	ctgacgcaat	gtccagtgat	ggaatcaact	11640
ctcctcacia	ctcccacggt	ggtcccagtt	ccaagcacag	agcttccttc	tgaagaagca	11700
ccaactgaaa	acagcactgg	ggtccaggac	tgctaccgag	gtgatggaca	gagttatcga	11760
ggcacactct	ccaccactat	cacaggaaga	acatgtcagt	cttggtcgtc	tatgacacca	11820
cattggcatc	ggaggatccc	attatactat	ccaaatgctg	gcctgaccag	gaactactgc	11880
aggaatccag	atgctgagat	tcgcccttgg	tgttacacca	tggatcccag	tgtcaggtgg	11940
gagtactgca	acctgacacg	atgtccagtg	acagaatcga	gtgtcctcac	aactcccaca	12000
gtggccccgg	ttccaagcac	agaggctcct	tctgaacaag	caccacctga	gaaaagccct	12060
gtggtccagg	attgctacca	tggatgatga	cggagttatc	gaggcatatc	ctccaccact	12120
gtcacaggaa	ggacctgtca	atcttgggtc	tctatgatac	cacactggca	tcagaggacc	12180
ccagaaaact	acccaaatgc	tggcctgacc	gagaactact	gcaggaatcc	agattctggg	12240
aaacaaccct	ggtgttacac	aaccgatccg	tgtgtgaggt	gggagtactg	caatctgaca	12300
caatgctcag	aaacagaatc	aggtgtccta	gagactccca	ctgttgttcc	agttccaagc	12360
atggaggctc	attctgaagc	agcaccaact	gagcaaacc	ctgtgggtccg	gcagtgtctac	12420
catggtaatg	gccagagtta	tcgaggcaca	ttctccacca	ctgtcacagg	aaggacatgt	12480
caatcttggg	catccatgac	accacaccgg	catcagagga	ccccagaaaa	ctacccaaat	12540
gatggcctga	caatgaacta	ctgcaggaat	ccagatgccg	atacaggccc	ttggtgtttt	12600
accatggacc	ccagcatcag	gtgggagtag	tgcaacctga	cgcgatgctc	agacacagaa	12660
gggactgtgg	tcgtctctcc	gactgtcatc	cagggttccaa	gcctagggcc	tccttctgaa	12720
caagactgta	tgtttgggaa	tgggaaagga	taccggggca	agaaggcaac	cactgttact	12780
gggacgcat	gccaggaatg	ggctgcccag	gagccccata	gacacagcac	gttcattcca	12840
gggacaaata	aatgggcagg	tctggaaaaa	aattactgcc	gtaaccctga	tggtagacatc	12900
aatggtccct	ggtgctacac	aatgaatcca	agaaaacttt	ttgactactg	tgatatccct	12960
ctctgtgcat	cctcttcatt	tgattgtggg	aagcctcaag	tggagccgaa	gaaatgtcct	13020
ggaagcattg	taggggggtg	tgtggcccac	ccacattcct	ggccctggca	agtcagtctc	13080
agaacaaggt	ttggaaagca	cttctgtgga	ggcaccttaa	tatccccaga	gtgggtgctg	13140
actgctgctc	actgcttgaa	gaagtcctca	aggccttcat	cctacaaggt	catcctgggt	13200
gcacaccaag	aagtgaacct	cgaatctcat	gttcaggaaa	tagaagtgtc	taggctgttc	13260
ttggagccca	cacaagcaga	tattgccttg	ctaaagctaa	gcaggcctgc	cgtcatcact	13320
gacaaagtaa	tgccagcttg	tctgccatcc	ccagactaca	tggtcaccgc	caggactgaa	13380
tgttacatca	ctggctgggg	agaaacccaa	ggtacctttg	ggactggcct	tctcaaggaa	13440
gccagctcc	ttgttattga	gaatgaagtg	tgcaatcact	ataagtatat	ttgtgctgag	13500
catttggcca	gaggcactga	cagttgccag	ggtgacagtg	gagggcctct	ggtttgcttc	13560

BI OL0250W0SEQ_ST25. txt

gagaaggaca aatacatttt acaaggagtc acttcttggg gtcttggctg tgcacgcccc 13620
aataagcctg gtgtctatgc tcgtgtttca aggtttgtta cttggattga gggaatgatg 13680
agaaataatt aattggacgg gagacagagt gaagcatcaa cctacttaga agctgaaacg 13740
tgggtaagga tttagcatgc tggaaataat agacagcaat caaacgaaga cactgttccc 13800
agctaccagc tatgccaaac cttggcattt ttggtatttt tgtgtataag cttttaaggt 13860
ctgactgaca aattctgtat taagggtgtca tagctatgac atttggtaaa aataaactct 13920
gcacttattt tgatttga 13938

<210> 5
<211> 23
<212> DNA
<213> Arti f i c i a l sequence

<220>
<223> Primer

<400> 5
acagcaatca aacgaagaca ctg 23

<210> 6
<211> 28
<212> DNA
<213> Arti f i c i a l sequence

<220>
<223> Primer

<400> 6
agcttatata caaaaatacc aaaaatgc 28

<210> 7
<211> 27
<212> DNA
<213> Arti f i c i a l sequence

<220>
<223> Probe

<400> 7
tcccagctac cagctatgcc aaacctt 27

<210> 8
<211> 16
<212> DNA
<213> Arti f i c i a l sequence

<220>
<223> Primer

<400> 8
ccacagtggc cccgggt 16

<210> 9
<211> 21
<212> DNA
<213> Arti f i c i a l sequence

<220>
 <223> Primer
 <400> 9
 acagggcttt tctcaggtgg t 21

<210> 10
 <211> 28
 <212> DNA
 <213> Arti fi ci al sequence
 <220>
 <223> Probe
 <400> 10
 ccaagcacag aggctccttc tgaacaag 28

<210> 11
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence
 <220>
 <223> Syntheti c ol i gonucl eoti de
 <400> 11
 ggcaggtcct tcctgtgaca 20

<210> 12
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence
 <220>
 <223> Syntheti c ol i gonucl eoti de
 <400> 12
 cctgtgacag tgggtggagta 20

<210> 13
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence
 <220>
 <223> Syntheti c ol i gonucl eoti de
 <400> 13
 tcctgtgaca gtgggtggagt 20

<210> 14
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence
 <220>
 <223> Syntheti c ol i gonucl eoti de
 <400> 14
 cttcctgtga cagtgggtgga 20

<210> 15

<211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 15
 ccttcctgtg acagtgggtg 20

 <210> 16
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 16
 tccttcctgt gacagtgggtg 20

 <210> 17
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 17
 gtccttcctg tgacagtgg 20

 <210> 18
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 18
 ggtccttcct gtgacagtgg 20

 <210> 19
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 19
 aggtccttcc tgtgacagtg 20

 <210> 20
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 20

caggtccttc ctgtgacagt 20

<210> 21
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 21
 gcaggtcctt cctgtgacag 20

<210> 22
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 22
 tggcaggtcc ttcctgtgac 20

<210> 23
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 23
 ttggcaggtc cttcctgtga 20

<210> 24
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 24
 cttggcaggt ccttcctgtg 20

<210> 25
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 25
 gcttggcagg tccttcctgt 20

<210> 26
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Synthetic oligonucleotide

 <400> 26
 tcttcctgtg acagtgggtg 20

 <210> 27
 <211> 20
 <212> DNA
 <213> Artificial sequence

 <220>
 <223> Synthetic oligonucleotide

 <400> 27
 ttcttcctgt gacagtgggtg 20

 <210> 28
 <211> 20
 <212> DNA
 <213> Artificial sequence

 <220>
 <223> Synthetic oligonucleotide

 <400> 28
 gttcttcctg tgacagtgggt 20

 <210> 29
 <211> 20
 <212> DNA
 <213> Artificial sequence

 <220>
 <223> Synthetic oligonucleotide

 <400> 29
 ggttcttcct gtgacagtgg 20

 <210> 30
 <211> 20
 <212> DNA
 <213> Artificial sequence

 <220>
 <223> Synthetic oligonucleotide

 <400> 30
 aggttcttcc tgtgacagtg 20

 <210> 31
 <211> 20
 <212> DNA
 <213> Artificial sequence

 <220>
 <223> Synthetic oligonucleotide

 <400> 31
 caggttcttc ctgtgacagt 20

 <210> 32

<211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 32
 tggcaggttc ttcctgtgac 20

<210> 33
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 33
 ttggcaggtt cttcctgtga 20

<210> 34
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 34
 cttggcagggt tcttcctgtg 20

<210> 35
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 35
 agcttggcag gttcttcctg 20

<210> 36
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 36
 actatgcgag tgtggtgtca 20

<210> 37
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 37

gactatgcga gtgtggtgtc 20

<210> 38
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 38
 cgactatgcg agtgtggtgt 20

<210> 39
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 39
 ccgactatgc gagtgtggtg 20

<210> 40
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 40
 tccgactatg cgagtgtggt 20

<210> 41
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 41
 gtccgactat gcgagtgtgg 20

<210> 42
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 42
 ggtccgacta tgcgagtgtg 20

<210> 43
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 43
 ggggccgact atgcgagtgt 20

<210> 44
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 44
 ctgctcagtc ggtgcttggt 20

<210> 45
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 45
 cctctgctca gtcggtgctt 20

<210> 46
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 46
 gcctctgctc agtcggtgct 20

<210> 47
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 47
 cttccagtga cagtgggtgga 20

<210> 48
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 48
 ttcttccagt gacagtgggtg 20

<210> 49

<211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 49
 gttcttccag tgacagtgg 20

<210> 50
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 50
 ggttcttcca gtgacagtgg 20

<210> 51
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 51
 gaccttaaaa gcttatacac 20

<210> 52
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 52
 gtcagacctt aaaagcttat 20

<210> 53
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 53
 tgtcagtcag accttaaaag 20

<210> 54
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 54

gaatttgtca gtcagacctt 20

<210> 55
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 55
 agaatttgtc agtcagacct 20

<210> 56
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 56
 ccttaataca gaatttgtca 20

<210> 57
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 57
 gctccgttgg tgcttggtca 20

<210> 58
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 58
 tgctccgttg gtgcttggtc 20

<210> 59
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 59
 ttgctccgtt ggtgcttggt 20

<210> 60
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 60
 ttgctccgt tgggtgcttg 20

<210> 61
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 61
 ctttgctccg ttggtgcttg 20

<210> 62
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 62
 tcctgtaaca gtggtggaga 20

<210> 63
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 63
 ttcctgtaac agtgggtggag 20

<210> 64
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 64
 cttcctgtaa cagtgggtgga 20

<210> 65
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 65
 ccttcctgta acagtgggtg 20

<210> 66

<211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 66
 tccttcctgt aacagtgggt 20

<210> 67
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 67
 gtccttcctg taacagtgggt 20

<210> 68
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 68
 tgtccttcct gtaacagtgg 20

<210> 69
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 69
 tggagccaga ataacattcg 20

<210> 70
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 70
 cctctaggct tggagccaga 20

<210> 71
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 71

agttcttcct gtgacagtgg 20

<210> 72
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 72
 gtccgactat gctggtgtgg 20

<210> 73
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 73
 ggtccgacta tgctggtgtg 20

<210> 74
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 74
 ggggccgact atgctggtgt 20

<210> 75
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 75
 cctctaggct tggaatcggg 20

<210> 76
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 76
 gttcagaagg agcctctagg 20

<210> 77
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 77
 tgttcagaag gaggcctctag 20

<210> 78
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 78
 gcttggtcag aaggagcctc 20

<210> 79
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 79
 tgcttggtca gaaggagcct 20

<210> 80
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 80
 gtgcttggtc agaaggagcc 20

<210> 81
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 81
 ggtgcttggt cagaaggagc 20

<210> 82
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 82
 tggtgcttgt tcagaaggag 20

<210> 83

<211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 83
 gctcagttgg tgcttggtca 20

<210> 84
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 84
 tgctcagttg gtgcttggtc 20

<210> 85
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 85
 gcttggatct gggaccaccg 20

<210> 86
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 86
 gcctccatgc ttggaactgg 20

<210> 87
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 87
 gctcagttgg tgctgcttca 20

<210> 88
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 88

cctcgataac tctggccatt 20

<210> 89
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 89
 tcctgtgaca gtggtggaga 20

<210> 90
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 90
 gtaggttgat gcttcactct 20

<210> 91
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 91
 cgtttgattg ctgtctatta 20

<210> 92
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 92
 ctctgtgctt ggatctggga 20

<210> 93
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 93
 cctctgtgct tggatctggg 20

<210> 94
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 94
 gcctctgtgc ttgatctgg 20

<210> 95
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 95
 agaagcctct gtgcttgat 20

<210> 96
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 96
 ttcagaagaa gcctctgtgc 20

<210> 97
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 97
 gctccgttg tgcttctca 20

<210> 98
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 98
 ttgctccgt tgggtgttct 20

<210> 99
 <211> 20
 <212> DNA
 <213> Artificial sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 99
 gctttgctcc gttggtgctt 20

<210> 100

<211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 100
 ggctttgctc cgttggtgct 20

<210> 101
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 101
 gggctttgct ccgttggtgc 20

<210> 102
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 102
 ccttcctgtg acagtggtag 20

<210> 103
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 103
 tccttcctgt gacagtggta 20

<210> 104
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 104
 tgccttcct gtgacagtgg 20

<210> 105
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 105

cctctaggct tggaaccggg 20

<210> 106
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 106
 tgcttggttcg gaaggagcct 20

<210> 107
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 107
 gtgcttggttc ggaaggagcc 20

<210> 108
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 108
 gcttggaact gggaccaccg 20

<210> 109
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 109
 ctgtgcttgg aactgggacc 20

<210> 110
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 110
 ctctgtgctt ggaactggga 20

<210> 111
 <211> 17
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 111
 cctgtgacag tgggtgga 17

<210> 112
 <211> 17
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 112
 tcctgtgaca gtggtgg 17

<210> 113
 <211> 17
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 113
 ttcctgtgac agtgggtg 17

<210> 114
 <211> 17
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 114
 cttcctgtga cagtgggt 17

<210> 115
 <211> 17
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 115
 ccttcctgtg acagtgg 17

<210> 116
 <211> 17
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 116
 tccttcctgt gacagtg 17

<210> 117

<211> 17
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 117
 gtccttcctg tgacagt 17

<210> 118
 <211> 17
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 118
 ggtccttcct gtgacag 17

<210> 119
 <211> 17
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 119
 ccgactatgc gagtgtg 17

<210> 120
 <211> 17
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 120
 gtccgactat gcgagt 17

<210> 121
 <211> 17
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 121
 ggtccgacta tgcgagt 17

<210> 122
 <211> 17
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 122

gtcagacctt aaaagct 17

<210> 123
 <211> 17
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 123
 aagcctctgt gcttgga 17

<210> 124
 <211> 17
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 124
 agcctctgtg cttggat 17

<210> 125
 <211> 17
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 125
 gcctctgtgc ttggatc 17

<210> 126
 <211> 17
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 126
 gctccgttgg tgcttct 17

<210> 127
 <211> 17
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Syntheti c ol i gonucl eoti de

<400> 127
 ctctgtgctt ggaactg 17

<210> 128
 <211> 17
 <212> DNA
 <213> Arti fi ci al sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 128
 tgcctcgata actctgt 17

<210> 129
 <211> 17
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 129
 tgtgcctcga taactct 17

<210> 130
 <211> 17
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 130
 gctcagttgg tgctgct 17

<210> 131
 <211> 27
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 131
 gcgtttgctc ttcttcttgc gtttttt 27

<210> 132
 <211> 3987
 <212> DNA
 <213> Macaca mulatta
 <400> 132
 atgtatcgtt ttggaatttc cagtggcttg atcaggaact actgcaggaa tccagatcct 60
 gtggcagccc ctattgtta tacgatggat cccaatgtca ggtgggagta ctgcaacctg 120
 acacaatgct cagatgcaga agggactgcc gtcgcacctc cgaatgtcac cctggttcca 180
 agcctagagg ctcttccga acaatcaccg actgagcaaa ggcctgggggt gcaggagtgc 240
 taccacggtta atggacagag ttatcgaggc acatacttca cactgtgac aggaagaacc 300
 tgccaagctt ggtcatctat gacaccgcac tctcatagtc ggaccccgga aaactaccca 360
 aatggtggct tgatcaggaa ctactgcagg aatccagatc ctgtggcagc cccttattgt 420
 tataccatgg atcccaatgt caggtgggag tactgcaacc taacacaatg ctgagacgca 480
 gaagggattg ccgtcacacc tctgactgtt accccggttc caagcctaga ggctccttcc 540
 aagcaagcac caactgagca aaggcctgggt gtccaggagt gttaccatgg taatggacag 600

BI OL0250W0SEQ_ST25. txt

agttatcgag	gcacatactt	caccactgtg	acaggaagaa	cctgccaagc	ttggatcatct	660
atgacaccac	attctcatag	tcgtacccca	gaaaactacc	caaattggcag	tccgacctct	720
tcagatctct	tagtctaccc	tgccgtcttc	cttgatgcca	tgggtccac	tggtctttca	780
actcatccgc	tttcctcag	tcccggagt	gctgcgacca	gcagaggata	tattgagagc	840
aagagagaag	caccgactga	gcaaaggcct	ggggtgcagg	agtgtacca	cggtaatgga	900
cagagttatc	gaggcacata	cttcaccact	gtgacaggaa	gaacctgcca	agcttgggtca	960
tctatgacac	cgactctca	tagtcggacc	ccggaaaact	acccaaatgg	tggcttgatc	1020
aggaactact	gcaggaatcc	agatcctgtg	gcagcccctt	attgtttatac	catggatccc	1080
agtgtcaggt	gggagtactg	caacctgaca	caatgtctcag	acgcagaagg	gactgccgtc	1140
gcacctccga	atgtacccc	ggttccaagc	ctagaggctc	cttctgagca	agcaccaact	1200
gagcaaaggc	ttgggggtga	ggagtgtac	cacagtaatg	gacagagtta	tcgaggcaca	1260
tacttcacca	ctgtgacagg	aagaacctgc	caagcttgggt	catctatgac	accacactct	1320
catagtcgga	ccccagaaaa	ctacccaaat	gctggcttgg	tcaagaacta	ctgccgaaat	1380
ccagatcctg	tggcagcccc	ttgggtgttat	acaacggatc	ccagtgtcag	gtgggagtac	1440
tgcaacctga	cacgatgctc	agatgcagaa	gggactgctg	tcgtgcctcc	aaatattatt	1500
ccggttccaa	gcctagaggc	ttttcttgaa	caagaaccga	ctgaggaaac	ccccggggta	1560
caggagtgtc	actaccatta	tggacagagt	tatagaggca	catactccac	cactgttaca	1620
ggaagaactt	gccaaagtgt	gtcatctatg	acaccacacc	agcatagtcg	gacccccaaa	1680
aactatccaa	atgttggcct	gaccaggaac	tactgcagga	atccagatgc	tgagattcgc	1740
ccttgggtgtt	ataccatgga	tcccagtgtc	aggtgggagt	actgcaacct	gacacaatgt	1800
ctggtgacag	aatcaagtgt	ccttgaaact	ctcacagtgg	tcccagatcc	aagcacacag	1860
gcttcttctg	aagaagcacc	aacggagcaa	agtcccagagg	tccaggactg	ctaccatggt	1920
gatggacaga	gttatcgagg	ctcattctcc	accactgtca	caggaaggac	atgtcagtct	1980
tggctctcta	tgacaccaca	ctggcatcag	aggacaacag	aatattatcc	agatgggtggc	2040
ctgaccagga	actactgcag	gaatccagat	gctgagattc	gcccttgggtg	ttataccatg	2100
gatcccagt	tcaggtggga	gtactgcaac	ctgacacaat	gtccagtgc	agaatcaagt	2160
gtcctcgcaa	cgtccatggc	tgtttctgaa	caagcaccaa	tggagcaaag	ccccgggggtc	2220
caggactgtc	accatggtga	tggacagagt	tatcgagggtt	cattctccac	cactgtcaca	2280
ggaaggacat	gtcagtcttg	gtcctctatg	acaccacact	ggcatcagag	gaccatagaa	2340
tactacccaa	atggtggcct	gaccaagaac	tactgcagga	atccagatgc	tgagattcgc	2400
ccttgggtgtt	ataccatgga	tcccagagtc	agatgggagt	actgcaacct	gacacaatgt	2460
gtggtgatgg	aatcaagtgt	ccttgcaact	cccatggtgg	tcccagttcc	aagcagagag	2520
gttccttctg	aagaagcacc	aactgaaaac	agccctgggg	tccaggactg	ctaccaaggt	2580
gatggacaga	gttatcgagg	cacattctcc	accactatca	caggaagaac	atgtcagtct	2640

BI OL0250W0SEQ_ST25. txt

tggttgctta	tgacaccaca	tcggcatcgg	aggatcccat	tacgctatcc	aaatgctggc	2700
ctgaccagga	actattgcag	aatccagat	gctgagattc	gcccttggtg	ttacaccatg	2760
gatcccagtg	tcaggtggga	gtactgcaac	ctgacacaat	gtccagtgac	agaatcaagt	2820
gtcctcacia	ctcccacggt	ggtcccgggt	ccaagcacag	aggctccttc	tgaacaagca	2880
ccacctgaga	aaagccctgt	ggtccaggat	tgctaccatg	gtgatggaca	gagttatcga	2940
ggcacatcct	ccaccactgt	cacaggaagg	aactgtcagt	cttggtcatc	tatgatacca	3000
cactggcatc	agaggacccc	agaaaactac	ccaaatgctg	gcctgaccag	gaactactgc	3060
aggaatccag	attctgggaa	acaaccctgg	tgttacacga	ctgatccatg	tgtgagggtg	3120
gagtactgca	acctgacaca	atgctcagaa	acagaatcag	gtgtcctaga	gactcccact	3180
gttggtccgg	ttccaagcat	ggaagctcat	tctgaagcag	caccaactga	gcaaaccctt	3240
gtgggtccagc	agtgttacca	tggtaatgga	cagagttatc	gaggcacatt	ctccaccact	3300
gtcacaggaa	ggacatgtca	atcttggtca	tccatgacac	cacaccagca	taagaggacc	3360
ccggaaaacc	acccaaatga	tggtttgaca	atgaactact	gcaggaatcc	agatgctgac	3420
acaggccctt	ggtgttttac	catggacccc	agcgtcaggc	gggagtactg	caacctgacg	3480
cgatgctcag	acacagaagg	gactgtggtc	acacctccga	ctgttatccc	ggttccaagc	3540
ctagaggctc	cttctgaaca	agtgtttgga	attcatcctg	aattaaacga	cacaattgct	3600
aacttggaact	caaaggtgaa	ttcttttcca	ccttgtgcca	cagcatcctc	ttcatttgat	3660
tgtgggaagc	ctcaagtgga	gccaaagaaa	tgtcctggaa	gcattgtagg	tgggtgtgtg	3720
gccacccac	attcctggcc	ctggcaagtc	agtcttagaa	caaggtttgg	aaagcacttc	3780
tgtggaggca	ccttaatatc	cccagagtgg	gtgctgactg	ctgcttgctg	cttgagagacg	3840
ttctcaaggc	cttcttcta	caaggtcatc	ctgggtgcac	accaagaagt	gaatctcgaa	3900
tctcatgttc	aagaaataga	agtgtctagg	ttgttcttgg	agcccatagg	agcagatatt	3960
gccttgctaa	agctaagcag	gtactaa				3987

<210> 133
 <211> 20
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 133
 ggttcttcca gtgacagtgg

20

<210> 134
 <211> 20
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide

<400> 134
atgcctcgat aactccgtcc 20

<210> 135
<211> 20
<212> DNA
<213> Arti fi ci al sequence

<220>
<223> Syntheti c ol i gonucl eoti de

<400> 135
agcttcttgt ccagctttat 20

<210> 136
<211> 21
<212> DNA
<213> Arti fi ci al sequence

<220>
<223> Syntheti c ol i gonucl eoti de

<400> 136
agcttcttgt ccagctttat a 21

<210> 137
<211> 14
<212> DNA
<213> Arti fi ci al sequence

<220>
<223> Syntheti c ol i gonucl eoti de

<400> 137
tcagtcatga cttc 14

<210> 138
<211> 15
<212> DNA
<213> Arti fi ci al sequence

<220>
<223> Syntheti c ol i gonucl eoti de

<400> 138
tcagtcatga cttca 15

<210> 139
<211> 20
<212> DNA
<213> Arti fi ci al sequence

<220>
<223> Syntheti c ol i gonucl eoti de

<400> 139
gctgattaga gagaggtccc 20

<210> 140
<211> 20
<212> DNA
<213> Arti fi ci al sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 140
 tccatttca ggagacctgg 20

<210> 141
 <211> 15
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 141
 atcagtcag acttc 15

<210> 142
 <211> 20
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 142
 cggtgcaagg ctaggaatt 20

<210> 143
 <211> 20
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 143
 gcttcagtc tgacttcctt 20

<210> 144
 <211> 21
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 144
 gcttcagtc tgacttcctt a 21

<210> 145
 <211> 21
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 145
 agcttcagtc atgacttcct t 21

<210> 146
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 146
 tggtaatcca ctttcagagg 20

<210> 147
 <211> 21
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 147
 tggtaatcca ctttcagagg a 21

<210> 148
 <211> 21
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 148
 tgcttcagtc atgacttcct t 21

<210> 149
 <211> 20
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 149
 cactgatttt tgcccaggat 20

<210> 150
 <211> 21
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 150
 cactgatttt tgcccaggat a 21

<210> 151
 <211> 21
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

<400> 151
aagcttcttg tccagcttta t 21

<210> 152
<211> 20
<212> DNA
<213> Arti fi ci al sequence

<220>
<223> Syntheti c ol i gonucl eoti de

<400> 152
acccaattca gaaggaagga 20

<210> 153
<211> 21
<212> DNA
<213> Arti fi ci al sequence

<220>
<223> Syntheti c ol i gonucl eoti de

<400> 153
acccaattca gaaggaagga a 21

<210> 154
<211> 21
<212> DNA
<213> Arti fi ci al sequence

<220>
<223> Syntheti c ol i gonucl eoti de

<400> 154
aaccaattc agaaggaagg a 21

<210> 155
<211> 21
<212> DNA
<213> Arti fi ci al sequence

<220>
<223> Syntheti c ol i gonucl eoti de

<400> 155
atggtaatcc actttcagag g 21

<210> 156
<211> 20
<212> DNA
<213> Arti fi ci al sequence

<220>
<223> Syntheti c ol i gonucl eoti de

<400> 156
tcttggttac atgaaatccc 20

<210> 157
<211> 21
<212> DNA
<213> Arti fi ci al sequence

<220>
 <223> Synthetic oligonucleotide
 <400> 157
 tcttggttac atgaaatccc a 21

<210> 158
 <211> 20
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 158
 attcactttc ataatgctgg 20

<210> 159
 <211> 21
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 159
 attcactttc ataatgctgg a 21

<210> 160
 <211> 21
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 160
 atcttgggta catgaaatcc c 21

<210> 161
 <211> 20
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 161
 atgcatgggtg atgcttctga 20

<210> 162
 <211> 20
 <212> DNA
 <213> Artificial sequence
 <220>
 <223> Synthetic oligonucleotide
 <400> 162
 cagctttatt aggacagca 20

<210> 163
 <211> 21
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 163
 cagctttatt agggacagca a 21

<210> 164
 <211> 21
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 164
 acagctttat tagggacagc a 21

<210> 165
 <211> 16
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 165
 ttcagtcag acttcc 16

<210> 166
 <211> 18
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

<220>
 <221> mi sc_feature
 <222> (1)..(4)
 <223> bases at these posi ti ons are RNA

<220>
 <221> mi sc_feature
 <222> (15)..(18)
 <223> bases at these posi ti ons are RNA

<400> 166
 gcuucagtca tgactucc 18

<210> 167
 <211> 21
 <212> DNA
 <213> Arti fi ci al sequence

 <220>
 <223> Syntheti c ol i gonucl eoti de

 <400> 167

tgctccgttg gtgcttggtc a

BI 0L0250W0SEQ_ST25. txt

21