



(86) **Date de dépôt PCT/PCT Filing Date:** 2014/12/24
(87) **Date publication PCT/PCT Publication Date:** 2015/07/02
(85) **Entrée phase nationale/National Entry:** 2016/05/24
(86) **N° demande PCT/PCT Application No.:** US 2014/072303
(87) **N° publication PCT/PCT Publication No.:** 2015/100394
(30) **Priorité/Priority:** 2013/12/24 (US61/920,652)

(51) **Cl.Int./Int.Cl. C12N 15/113** (2010.01)
(71) **Demandeur/Applicant:**
IONIS PHARMACEUTICALS, INC., US
(72) **Inventeurs/Inventors:**
FREIER, SUSAN M., US;
GRAHAM, MARK J., US;
CROOKE, ROSANNE M., US
(74) **Agent:** NORTON ROSE FULBRIGHT CANADA
LLP/S.E.N.C.R.L., S.R.L.

(54) **Titre : MODULATION DE L'EXPRESSION DE LA PROTEINE ANGPTL3**
(54) **Title: MODULATION OF ANGIOPOIETIN-LIKE 3 EXPRESSION**

(57) **Abrégé/Abstract:**

Provided herein are methods, compounds, and compositions for reducing expression of an ANGPTL3 mRNA and protein in an animal. Also provided herein are methods, compounds, and compositions for reducing lipids and/or glucose in an animal. Such methods, compounds, and compositions are useful to treat, prevent, delay, or ameliorate any one or more of cardiovascular disease and/or metabolic disease, or a symptom thereof, in an individual in need thereof.



(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization
International Bureau

(43) International Publication Date
2 July 2015 (02.07.2015)





(10) International Publication Number
WO 2015/100394 A1

- (51) International Patent Classification:
C12N 15/113 (2010.01)

(21) International Application Number:
PCT/US2014/072303

(22) International Filing Date:
24 December 2014 (24.12.2014)

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data:
61/920,652 24 December 2013 (24.12.2013) US

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

(72) Inventors: **FREIER, Susan, M.**; 2855 Gazelle Ct, Carlsbad, CA 92010 (US). **GRAHAM, Mark, J.**; 2855 Gazelle Ct, Carlsbad, CA 92010 (US). **CROOKE, Rosanne, M.**; 2855 Gazelle Ct, Carlsbad, CA 92010 (US).

(74) Agents: **GRANT, Bruce D.** et al.; Grant IP, 2714 Loker Ave W, Suite 110, 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, 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, ST, 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: MODULATION OF ANGIOPOIETIN-LIKE 3 EXPRESSION

(57) Abstract: Provided herein are methods, compounds, and compositions for reducing expression of an ANGPTL3 mRNA and protein in an animal. Also provided herein are methods, compounds, and compositions for reducing lipids and/or glucose in an animal. Such methods, compounds, and compositions are useful to treat, prevent, delay, or ameliorate any one or more of cardiovascular disease and/or metabolic disease, or a symptom thereof, in an individual in need thereof.

WO 2015/100394 A1

MODULATION OF ANGIOPOIETIN-LIKE 3 EXPRESSION

Sequence Listing

The present application is being filed along with a Sequence Listing in electronic format. The
5 Sequence Listing is provided as a file entitled BIOL0179WO_ST25.txt, created on December 22, 2014 which
is .98 MB in size. The information in the electronic format of the sequence listing is incorporated herein by
reference in its entirety.

Field of the Invention

10 Provided herein are methods, compounds, and compositions for reducing expression of angiotensin-
like 3 (ANGPTL3) mRNA and protein in an animal. Also, provided herein are methods, compounds, and
compositions having an ANGPTL3 inhibitor for reducing ANGPTL3 related diseases or conditions in an
animal. Such methods, compounds, and compositions are useful, for example, to treat, prevent, delay or
ameliorate any one or more of cardiovascular disease or metabolic syndrome, or a symptom thereof, in an
15 animal.

Background

Diabetes and obesity (sometimes collectively referred to as “diabesity”) are interrelated in that
obesity is known to exacerbate the pathology of diabetes and greater than 60% of diabetics are obese. Most
20 human obesity is associated with insulin resistance and leptin resistance. In fact, it has been suggested that
obesity may have an even greater impact on insulin action than diabetes itself (Sindelka et al., *Physiol Res.*,
2002, 51, 85-91). Additionally, several compounds on the market for the treatment of diabetes are known to
induce weight gain, a very undesirable side effect to the treatment of this disease.

Cardiovascular disease is also interrelated to obesity and diabetes. Cardiovascular disease
25 encompasses a wide variety of etiologies and has an equally wide variety of causative agents and interrelated
players. Many causative agents contribute to symptoms such as elevated plasma levels of cholesterol,
including non-high density lipoprotein cholesterol (non-HDL-C), as well as other lipid-related disorders.
Such lipid-related disorders, generally referred to as dyslipidemia, include hyperlipidemia,
hypercholesterolemia and hypertriglyceridemia among other indications. Elevated non-HDL cholesterol is
30 associated with atherogenesis and its sequelae, including cardiovascular diseases such as arteriosclerosis,
coronary artery disease, myocardial infarction, ischemic stroke, and other forms of heart disease. These rank
as the most prevalent types of illnesses in industrialized countries. Indeed, an estimated 12 million people in
the United States suffer with coronary artery disease and about 36 million require treatment for elevated
cholesterol levels.

Epidemiological and experimental evidence has shown that high levels of circulating triglyceride (TG) can contribute to cardiovascular disease and a myriad of metabolic disorders (Valdivielso et al., 2009, *Atherosclerosis* Zhang et al., 2008, *Circ Res.* 1;102(2):250-6). TG derived from either exogenous or endogenous sources is incorporated and secreted in chylomicrons from the intestine or in very low density lipoproteins (VLDL) from the liver. Once in circulation, TG is hydrolyzed by lipoprotein lipase (LpL) and the resulting free fatty acids can then be taken up by local tissues and used as an energy source. Due to the profound effect LpL has on plasma TG and metabolism in general, discovering and developing compounds that affect LpL activity are of great interest.

Metabolic syndrome is a combination of medical disorders that increase one's risk for cardiovascular disease and diabetes. The symptoms, including high blood pressure, high triglycerides, decreased HDL and obesity, tend to appear together in some individuals. It affects a large number of people in a clustered fashion. In some studies, the prevalence in the USA is calculated as being up to 25% of the population. Metabolic syndrome is known under various other names, such as (metabolic) syndrome X, insulin resistance syndrome, Reaven's syndrome or CHAOS. With the high prevalence of cardiovascular disorders and metabolic disorders there remains a need for improved approaches to treat these conditions

The angiopoietins are a family of secreted growth factors. Together with their respective endothelium-specific receptors, the angiopoietins play important roles in angiogenesis. One family member, angiopoietin-like 3 (also known as angiopoietin-like protein 3, ANGPT5, ANGPTL3, or angiopoietin 5), is predominantly expressed in the liver, and is thought to play a role in regulating lipid metabolism (Kaplan et al., *J. Lipid Res.*, **2003**, 44, 136-143). Genome-wide association scans (GWAS) surveying the genome for common variants associated with plasma concentrations of HDL, LDL and triglyceride found an association between triglycerides and single-nucleotide polymorphisms (SNPs) near ANGPTL3 (Willer et al., *Nature Genetics*, 2008, 40(2):161-169). Individuals with homozygous ANGPTL3 loss-of-function mutations present with low levels of all atherogenic plasma lipids and lipoproteins, such as total cholesterol (TC) and TG, low density lipoprotein cholesterol (LDL-C), apolipoprotein B (apoB), non-HDL-C, as well as HDL-C (Romeo et al. 2009, *J Clin Invest*, 119(1):70-79; Musunuru et al. 2010 *N Engl J Med*, 363:2220-2227; Martin-Campos et al. 2012, *Clin Chim Acta*, 413:552-555; Minicocci et al. 2012, *J Clin Endocrinol Metab*, 97:e1266-1275; Noto et al. 2012, *Arterioscler Thromb Vasc Biol*, 32:805-809; Pisciotta et al. 2012, *Circulation Cardiovasc Genet*, 5:42-50). This clinical phenotype has been termed familial combined hypolipidemia (FHBL2). Despite reduced secretion of VLDL, subjects with FHBL2 do not have increased hepatic fat content. They also appear to have lower plasma glucose and insulin levels, and importantly, both diabetes and cardiovascular disease appear to be absent from these subjects. No adverse clinical phenotypes have been reported to date (Minicocci et al. 2013, *J of Lipid Research*, 54:3481-3490). Reduction of ANGPTL3 has been shown to lead to a decrease in TG, cholesterol and LDL levels in animal models (U.S. Serial Number 13/520,997; PCT Publication WO 2011/085271). Mice deficient in ANGPTL3 have very low plasma triglyceride (TG) and cholesterol levels, while overexpression produces the opposite effects (Koishi

et al. 2002; Koster 2005; Fujimoto 2006). Accordingly, the potential role of ANGPTL3 in lipid metabolism makes it an attractive target for therapeutic intervention.

To date, therapeutic strategies to treat cardiometabolic disease by directly targeting ANGPTL3 levels have been limited. ANGPTL3 polypeptide fragments (U.S. Serial Number 12/128,545), anti-ANGPTL3
 5 antibodies (U.S. Serial Number 12/001,012) and ANGPTL3 nucleic acid inhibitors including antisense oligonucleotides (U.S. Serial Number 13/520,997; PCT Publication WO 2011/085271; incorporated by reference herein, in their entirety) have previously been suggested or developed, but none of the compounds directly targeting ANGPTL3 have been approved for treating cardiometabolic disease. Accordingly, there is an unmet need for highly potent and tolerable compounds to inhibit ANGPTL3. The invention disclosed
 10 herein relates to the discovery of novel, highly potent inhibitors of ANGPTL3 expression and their use in treatment.

Summary of the Invention

Provided herein are compositions and methods for modulating expression of ANGPTL3 mRNA and protein. In certain embodiments, the composition is an ANGPTL3 specific inhibitor. In certain embodiments,
 15 the ANGPTL3 specific inhibitor decreases expression of ANGPTL3 mRNA and protein.

In certain embodiments, the composition is an ANGPTL3 specific inhibitor. In certain embodiments, the ANGPTL3 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 ANGPTL3 specific inhibitor is a modified oligonucleotide consisting of
 20 12 to 30 linked nucleosides and having 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: 77.

In certain embodiments, the ANGPTL3 specific inhibitor is a modified oligonucleotide consisting of 12 to 30 linked nucleosides and comprising a nucleobase sequence comprising a portion of at least 8
 25 contiguous nucleobases complementary to an equal length portion of nucleobases 1140-1159 of SEQ ID NO: 1, wherein the nucleobase sequence of the modified oligonucleotide is at least 80% complementary to SEQ ID NO: 1.

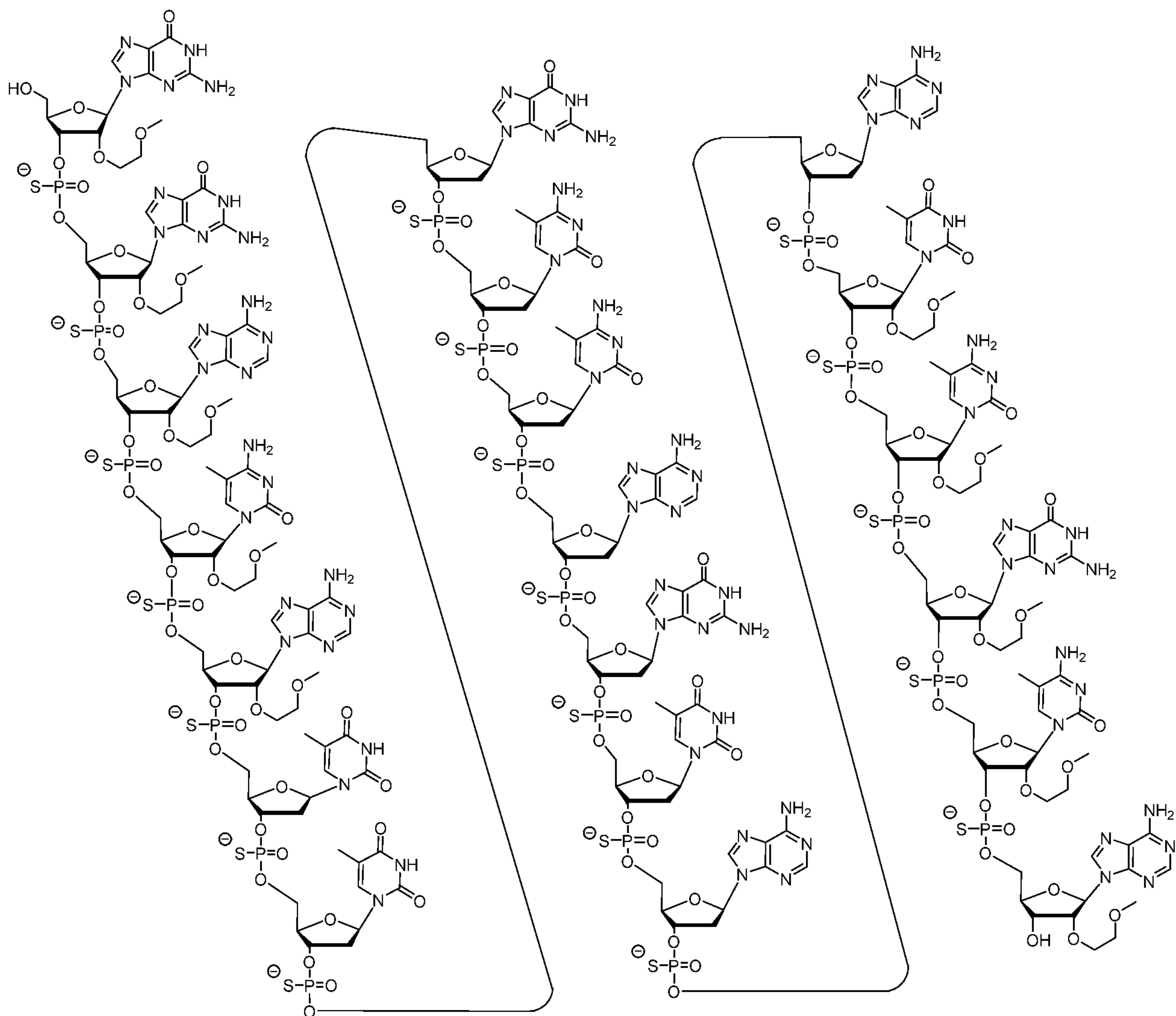
In certain embodiments, the ANGPTL3 specific inhibitor is a modified oligonucleotide consisting of 12 to 30 linked nucleosides and comprising a nucleobase sequence comprising a portion of at least 8
 30 contiguous nucleobases complementary to an equal length portion of nucleobases 9715-9734 of SEQ ID NO: 2, wherein the nucleobase sequence of the modified oligonucleotide is at least 80% complementary to SEQ ID NO: 2.

In certain embodiments, the ANGPTL3 specific inhibitor is a modified oligonucleotide consisting of 20 linked nucleosides and having a nucleobase sequence comprising at least 8 contiguous nucleobases of
 35 SEQ ID NO: 77, 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 each internucleoside linkage is a phosphorothioate linkage and wherein each
5 cytosine residue is a 5-methylcytosine.

In certain embodiments, the ANGPTL3 specific inhibitor is a modified oligonucleotide consisting of 20 linked nucleosides and having a nucleobase sequence consisting of at least 8 contiguous nucleobases of SEQ ID NO: 77, wherein the modified oligonucleotide consists of: (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 each internucleoside linkage is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

In certain embodiments, the ANGPTL3 specific inhibitor is a modified oligonucleotide represented
15 by the following structure and the designation ISIS 563580. In certain embodiments, the ANGPTL3 specific inhibitor comprises the modified oligonucleotide ISIS 563580 represented by the following structure.



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

In certain embodiments, the modulation of ANGPTL3 expression occurs in a cell or tissue. In certain
 5 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 ANGPTL3 mRNA level. In certain embodiments, the modulation is a reduction in ANGPTL3 protein level. In certain embodiments, both ANGPTL3 mRNA and protein levels are reduced. Such reduction may occur in a time-dependent or in a dose-dependent manner.

Certain embodiments provide compositions and methods for use in therapy. Certain embodiments
 10 provide compositions and methods for preventing, treating, delaying, slowing the progression and/or ameliorating ANGPTL3 related diseases, disorders, and conditions. In certain embodiments, such diseases, disorders, and conditions are cardiovascular and/or metabolic diseases, disorders, and conditions. In certain embodiments, the compositions and methods for therapy include administering an ANGPTL3 specific
 15 inhibitor to an individual in need thereof. In certain embodiments, the ANGPTL3 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.

Detailed Description of the Invention

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 invention, as claimed. 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 for the portions of the document discussed herein, as well as in their entirety.

15

Definitions

Unless specific definitions are provided, the nomenclature utilized 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 can be used for chemical synthesis, and chemical analysis. Where permitted, all patents, applications, published applications and other publications, GENBANK Accession Numbers and associated sequence information obtainable through databases such as National Center for Biotechnology Information (NCBI) and other data referred to throughout in the disclosure herein are incorporated by reference for the portions of the document discussed herein, as well as in their entirety.

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

“2’-O-methoxyethyl” (also 2’-MOE and 2’-O(CH₂)₂-OCH₃) refers to an O-methoxy-ethyl modification of the 2’ position of a furosyl ring. A 2’-O-methoxyethyl modified sugar is a modified sugar.

“2’-O-methoxyethyl nucleotide” means a nucleotide comprising a 2’-O-methoxyethyl modified sugar moiety.

“3’ target site” or “3’ stop site” refers to the nucleotide of a target nucleic acid which is complementary to the 3’-most nucleotide of a particular antisense compound.

“5’ target site” or “5 start site” refers to the nucleotide of a target nucleic acid which is complementary to the 5’-most nucleotide of a particular antisense compound.

“5-methylcytosine” means a cytosine modified with a methyl group attached to the 5’ position. A 5-methylcytosine is a modified nucleobase.

“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%

“Active pharmaceutical agent” means the substance or substances in a pharmaceutical composition that provide a therapeutic benefit when administered to an individual. For example, in certain embodiments an antisense oligonucleotide targeted to ANGPTL3 is an active pharmaceutical agent.

“Active target region” or “target region” means a region to which one or more active antisense compounds is targeted.

“Active antisense compounds” means antisense compounds that reduce target nucleic acid levels or protein levels.

“Adipogenesis” means the development of fat cells from preadipocytes. “Lipogenesis” means the production or formation of fat, either fatty degeneration or fatty infiltration.

“Adiposity” or “Obesity” refers to the state of being obese or an excessively high amount of body fat or adipose tissue in relation to lean body mass. The amount of body fat includes concern for both the distribution of fat throughout the body and the size and mass of the adipose tissue deposits. Body fat distribution can be estimated by skin-fold measures, waist-to-hip circumference ratios, or techniques such as ultrasound, computed tomography, or magnetic resonance imaging. According to the Center for Disease Control and Prevention, individuals with a body mass index (BMI) of 30 or more are considered obese. The term “Obesity” as used herein includes conditions where there is an increase in body fat beyond the physical requirement as a result of excess accumulation of adipose tissue in the body. The term “obesity” includes, but is not limited to, the following conditions: adult-onset obesity; alimentary obesity; endogenous or metabolic obesity; endocrine obesity; familial obesity; hyperinsular obesity; hyperplastic-hypertrophic obesity; hypogonadal obesity; hypothyroid obesity; lifelong obesity; morbid obesity and exogenous obesity.

“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.

“Administering” means providing an agent to an animal, and includes, but is not limited to, administering by a medical professional and self-administering.

“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 ANGPTL3. “Second agent” means a second therapeutic compound of the invention (e.g. a second antisense oligonucleotide targeting ANGPTL3) and/or a non-ANGPTL3 therapeutic compound.

“Amelioration” 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.

“ANGPTL3” means any nucleic acid or protein of ANGPTL3.

5 “ANGPTL3 expression” means the level of mRNA transcribed from the gene encoding ANGPTL3 or the level of protein translated from the mRNA. ANGPTL3 expression can be determined by art known methods such as a Northern or Western blot.

10 “ANGPTL3 nucleic acid” means any nucleic acid encoding ANGPTL3. For example, in certain embodiments, an ANGPTL3 nucleic acid includes a DNA sequence encoding ANGPTL3, a RNA sequence transcribed from DNA encoding ANGPTL3 (including genomic DNA comprising introns and exons), and a mRNA sequence encoding ANGPTL3. “ANGPTL3 mRNA” means a mRNA encoding an ANGPTL3 protein.

“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.

15 “Antisense activity” means any detectable or measurable activity attributable to the hybridization of an antisense compound to its target nucleic acid. In certain embodiments, antisense activity is a decrease in the amount or expression of a target nucleic acid or protein encoded by such target nucleic acid.

“Antisense compound” means an oligomeric compound that is capable of undergoing hybridization to a target nucleic acid through hydrogen bonding.

20 “Antisense inhibition” means reduction of target nucleic acid levels or target protein levels in the presence of an antisense compound complementary to a target nucleic acid compared to target nucleic acid levels or target protein levels in the absence of the antisense compound.

“Antisense oligonucleotide” means a single-stranded oligonucleotide having a nucleobase sequence that permits hybridization to a corresponding region or segment of a target nucleic acid.

25 “ApoB-containing lipoprotein” means any lipoprotein that has apolipoprotein B as its protein component, and is understood to include LDL, VLDL, IDL, and lipoprotein(a) and can be generally targeted by lipid lowering agent and therapies. “ApoB-100-containing LDL” means ApoB-100 isoform containing LDL.

30 “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.

“Bicyclic sugar” means a furosyl ring modified by the bridging of two non-geminal ring atoms. A bicyclic sugar is a modified sugar.

35 “Bicyclic nucleic acid” or “BNA” refers to a nucleoside or nucleotide wherein the furanose portion of the nucleoside or nucleotide includes a bridge connecting two carbon atoms on the furanose ring, thereby forming a bicyclic ring system.

“Cap structure” or “terminal cap moiety” means chemical modifications, which have been incorporated at either terminus of an antisense compound.

“Cardiovascular disease” or “cardiovascular disorder” refers to a group of conditions related to the heart, blood vessels, or the circulation. Examples of cardiovascular diseases or disorders include, but are not limited to, aneurysm, angina, arrhythmia, atherosclerosis, cerebrovascular disease (stroke), coronary heart disease, hypertension, dyslipidemia, hyperlipidemia, and hypercholesterolemia.

“Cardiometabolic disease” or “cardiometabolic disorder” are diseases or disorders concerning both the cardiovascular system and the metabolic system. Examples of cardiometabolic diseases or disorders include, but are not limited to, diabetes and dyslipidemias.

“Chemically distinct region” refers to a region of an antisense compound that is in some way chemically different than another region of the same antisense compound. For example, a region having 2'-O-methoxyethyl nucleotides is chemically distinct from a region having nucleotides without 2'-O-methoxyethyl modifications.

“Chimeric antisense compound” means an antisense compound that has at least two chemically distinct regions.

“Co-administration” means administration of two or more agents to an individual. The two or more agents can be in a single pharmaceutical composition, or can be in separate pharmaceutical compositions. Each of the two or more agents can be administered through the same or different routes of administration. Co-administration encompasses parallel or sequential administration.

“Cholesterol” is a sterol molecule found in the cell membranes of all animal tissues. Cholesterol must be transported in an animal's blood plasma by lipoproteins including very low density lipoprotein (VLDL), intermediate density lipoprotein (IDL), low density lipoprotein (LDL), and high density lipoprotein (HDL). “Plasma cholesterol” refers to the sum of all lipoproteins (VDL, IDL, LDL, HDL) esterified and/or non-estrified cholesterol present in the plasma or serum.

“Cholesterol absorption inhibitor” means an agent that inhibits the absorption of exogenous cholesterol obtained from diet.

“Complementarity” means the capacity for pairing between nucleobases of a first nucleic acid and a second nucleic acid. In certain embodiments, complementarity between the first and second nucleic acid may be between two DNA strands, between two RNA strands, or between a DNA and an RNA strand. In certain embodiments, some of the nucleobases on one strand are matched to a complementary hydrogen bonding base on the other strand. In certain embodiments, all of the nucleobases on one strand are matched to a complementary hydrogen bonding base on the other strand. In certain embodiments, a first nucleic acid is an antisense compound and a second nucleic acid is a target nucleic acid. In certain such embodiments, an antisense oligonucleotide is a first nucleic acid and a target nucleic acid is a second nucleic acid.

“Contiguous nucleobases” means nucleobases immediately adjacent to each other.

“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.

“Deoxyribonucleotide” means a nucleotide having a hydrogen at the 2' position of the sugar portion of the nucleotide. Deoxyribonucleotides may be modified with any of a variety of substituents.

5 “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
10 lethargy.

“Diabetic dyslipidemia” or “type 2 diabetes with dyslipidemia” means a condition characterized by Type 2 diabetes, reduced HDL-C, elevated triglycerides, and elevated small, dense LDL particles.

“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,
15 e.g. saline solution.

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

“Dosage unit” means a form in which a pharmaceutical agent is provided, e.g. pill, tablet, or other
20 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.

“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,
25 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 be expressed as mg/kg or g/kg.

30 “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.

35 “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 target nucleic acid is a second nucleic acid.

“Gapmer” means a chimeric antisense compound in which an internal region having a plurality of nucleosides that support RNase H cleavage is positioned between external regions having one or more nucleosides, wherein the nucleosides comprising the internal region are chemically distinct from the nucleoside or nucleosides comprising the external regions. The internal region can be referred to as a “gap segment” and the external regions can be referred to as “wing segments.”

“Gap-widened” means a chimeric antisense compound having a gap segment of 12 or more contiguous 2'-deoxyribonucleosides positioned between and immediately adjacent to 5' and 3' wing segments having from one to six nucleosides.

“Glucose” is a monosaccharide used by cells as a source of energy and metabolic intermediate. “Plasma glucose” refers to glucose present in the plasma.

“High density lipoprotein-C (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.

“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.

“Hybridization” means the annealing of complementary nucleic acid molecules. In certain embodiments, complementary nucleic acid molecules include an antisense compound and a target nucleic acid.

“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).

“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 and triglycerides.

“Hypertriglyceridemia” means a condition characterized by elevated triglyceride levels.

“Identifying” or “selecting a subject having a metabolic or cardiovascular disease” means identifying or selecting a subject having been diagnosed with a metabolic disease, a cardiovascular disease, or a metabolic syndrome; or, identifying or selecting a subject having any symptom of 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 may be accomplished by any method, including but not limited to, standard

clinical tests or assessments, such as measuring serum or circulating (plasma) cholesterol, 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.

“Identifying” or “selecting a diabetic subject” means identifying or selecting a subject having been
 5 identified as diabetic or identifying or selecting a subject having any symptom of diabetes (type 1 or type 2) such as, but not limited to, having a fasting glucose of at least 110 mg/dL, glycosuria, polyuria, polydipsia, increased insulin resistance, and/or decreased insulin sensitivity.

“Identifying” or “selecting an obese subject” means identifying or selecting a subject having been
 10 diagnosed as obese or identifying or selecting a subject with a BMI over 30 and/or a waist circumference of greater than 102 cm in men or greater than 88 cm in women.

“Identifying” or “selecting a subject having dyslipidemia” means identifying or selecting a subject diagnosed with a disorder of lipid and/or lipoprotein metabolism, including lipid and/or lipoprotein overproduction or deficiency. Dyslipidemias may be manifested by elevation of lipids such as cholesterol and triglycerides as well as lipoproteins such as low-density lipoprotein (LDL) cholesterol.

15 “Identifying” or “selecting” a subject having increased adiposity” means identifying or selecting a subject having an increased amount of body fat (or adiposity) that includes concern for one or both the distribution of fat throughout the body and the size and mass of the adipose tissue deposits. Body fat distribution can be estimated by skin-fold measures, waist-to-hip circumference ratios, or techniques such as ultrasound, computer tomography, or magnetic resonance imaging. According to the Center for Disease
 20 Control and Prevention, individuals with a body mass index (BMI) of 30 or more are considered obese.

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

“Immediately adjacent” means there are no intervening elements between the immediately adjacent
 25 elements.

“Individual” or “subject” or “animal” means a human or non-human animal selected for treatment or therapy.

“Inhibiting the expression or activity” refers to a reduction or blockade of the expression or activity and does not necessarily indicate a total elimination of expression or activity.

30 “Insulin resistance” is defined as the condition in which normal amounts of insulin are inadequate to produce a normal insulin response from cells, e.g., fat, muscle and/or 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
 35 resistance often leads to metabolic syndrome and type 2 diabetes.

“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.

“Internucleoside linkage” refers to the chemical bond between nucleosides.

5 “Intravenous administration” means administration into a vein.

“Linked nucleosides” means adjacent nucleosides which are bonded together.

“Lipid-lowering” means a reduction in one or more lipids in a subject. Lipid-lowering can occur with one or more doses over time.

10 “Lipid-lowering agent” means an agent, for example, an ANGPTL3-specific modulator, provided to a subject to achieve a lowering of lipids in the subject. For example, in certain embodiments, a lipid-lowering agent is provided to a subject to reduce one or more of apoB, apoC-III, total cholesterol, LDL-C, VLDL-C, IDL-C, non-HDL-C, triglycerides, small dense LDL particles, and Lp(a) in a subject.

15 “Lipid-lowering therapy” 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 apoB, apoC-III, total cholesterol, LDL-C, VLDL-C, IDL-C, non-HDL-C, triglycerides, small dense LDL particles, and Lp(a) in a subject.

20 “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 in that the HDL has a higher proportion of protein versus lipid, whereas the VLDL has a lower proportion of protein versus lipid.

“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.

25 “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 limitation, cigarette smoking, hypertension, low HDL-C, family history of coronary heart disease, age, and other factors disclosed herein.

30 “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. Metabolic disorders include, but are not limited to, hyperglycemia, prediabetes, diabetes (type I and type 2), obesity, insulin resistance, metabolic syndrome and dyslipidemia due to type 2 diabetes.

35 “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 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).

“Mismatch” or “non-complementary nucleobase” refers to the case when a nucleobase of a first nucleic acid is not capable of pairing with the corresponding nucleobase of a second or target nucleic acid.

5 “Mixed dyslipidemia” means a condition characterized by elevated cholesterol and elevated triglycerides.

“Modified internucleoside linkage” refers to a substitution or any change from a naturally occurring internucleoside bond (i.e. a phosphodiester internucleoside bond).

10 “Modified nucleobase” refers to any nucleobase other than adenine, cytosine, guanine, thymidine, or uracil. An “unmodified nucleobase” means the purine bases adenine (A) and guanine (G), and the pyrimidine bases thymine (T), cytosine (C), and uracil (U).

“Modified nucleoside” means a nucleoside having, independently, one or more modified sugar moiety or modified nucleobase.

15 “Modified nucleotide” means a nucleotide having, independently, one or more modified sugar moiety, modified internucleoside linkage, or modified nucleobase. A “modified nucleoside” means a nucleoside having, independently, one or more modified sugar moiety or modified nucleobase.

“Modified oligonucleotide” means an oligonucleotide comprising at least one modified nucleotide.

“Modified sugar” refers to a substitution or change from a natural sugar.

“Motif” means the pattern of chemically distinct regions in an antisense compound.

20 “MTP inhibitor” means an agent inhibits the enzyme microsomal triglyceride transfer protein.

“Naturally occurring internucleoside linkage” means a 3' to 5' phosphodiester linkage.

“Natural sugar moiety” means a sugar found in DNA (2'-H) or RNA (2'-OH).

25 “Non-alcoholic fatty liver disease” or “NAFLD” means a condition characterized by fatty inflammation of the liver that is not due to excessive alcohol use (for example, alcohol consumption of over 20 g/day). In certain embodiments, NAFLD is related to insulin resistance and metabolic syndrome. NAFLD encompasses a disease spectrum ranging from simple triglyceride accumulation in hepatocytes (hepatic steatosis) to hepatic steatosis with inflammation (steatohepatitis), fibrosis, and cirrhosis.

30 “Nonalcoholic steatohepatitis” (NASH) occurs from progression of NAFLD beyond deposition of triglycerides. A “second hit” capable of inducing necrosis, inflammation, and fibrosis is required for development of NASH. Candidates for the second-hit can be grouped into broad categories: factors causing an increase in oxidative stress and factors promoting expression of proinflammatory cytokines. It has been suggested that increased liver triglycerides lead to increased oxidative stress in hepatocytes of animals and humans, indicating a potential cause-and-effect relationship between hepatic triglyceride accumulation, oxidative stress, and the progression of hepatic steatosis to NASH (Browning and Horton, *J Clin Invest*,
35 **2004**, 114, 147-152). Hypertriglyceridemia and hyperfattyacidemia can cause triglyceride accumulation in peripheral tissues (Shimamura et al., *Biochem Biophys Res Commun*, **2004**, 322, 1080-1085).

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

5 “Nucleobase” means a heterocyclic moiety capable of pairing with a base of another nucleic acid.

“Nucleobase sequence” means the order of contiguous nucleobases independent of any sugar, linkage, or nucleobase modification.

“Nucleoside” means a nucleobase linked to a sugar.

10 “Nucleoside mimetic” includes those structures used to replace the sugar or the sugar and the base and not necessarily the linkage at one or more positions of an oligomeric compound such as for example nucleoside mimetics having morpholino, cyclohexenyl, cyclohexyl, tetrahydropyranyl, bicyclo or tricyclo sugar mimetics e.g. non furanose sugar units.

“Nucleotide” means a nucleoside having a phosphate group covalently linked to the sugar portion of the nucleoside.

15 “Nucleotide mimetic” includes those structures used to replace the nucleoside and the linkage at one or more positions of an oligomeric compound such as for example peptide nucleic acids or morpholinos (morpholinos linked by -N(H)-C(=O)-O- or other non-phosphodiester linkage).

20 “Oligomeric compound” or “oligomer” refers to a polymeric structure comprising two or more sub-structures and capable of hybridizing to a region of a nucleic acid molecule. In certain embodiments, oligomeric compounds are oligonucleosides. In certain embodiments, oligomeric compounds are oligonucleotides. In certain embodiments, oligomeric compounds are antisense compounds. In certain embodiments, oligomeric compounds are antisense oligonucleotides. In certain embodiments, oligomeric compounds are chimeric oligonucleotides.

25 “Oligonucleotide” means a polymer of linked nucleosides each of which can be modified or unmodified, independent one from another.

30 “Parenteral administration” means administration by a manner other than through the digestive tract. Parenteral administration includes topical administration, subcutaneous administration, intravenous administration, intramuscular administration, intraarterial administration, intraperitoneal administration, or intracranial administration, e.g. intrathecal or intracerebroventricular administration. Administration can be continuous, or chronic, or short or intermittent.

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

35 “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 ANGPTL3 is pharmaceutical agent.

“Pharmaceutical composition” means a mixture of substances suitable for administering to an individual. For example, a pharmaceutical composition can comprise one or more active agents and a sterile aqueous solution.

“Pharmaceutically acceptable carrier” means a medium or diluent that does not interfere with the structure or function of the oligonucleotide. Certain, of such carriers enable pharmaceutical compositions to be formulated as, for example, tablets, pills, dragees, capsules, liquids, gels, syrups, slurries, suspension and lozenges for the oral ingestion by a subject. Certain of such carriers enable pharmaceutical compositions to be formulated for injection or infusion. For example, a pharmaceutically acceptable carrier can be a sterile aqueous solution.

“Pharmaceutically acceptable salts” means physiologically and pharmaceutically acceptable salts of antisense compounds, i.e., salts that retain the desired biological activity of the parent oligonucleotide and do not impart undesired toxicological effects thereto.

“Phosphorothioate linkage” means a linkage between nucleosides where the phosphodiester bond is modified by replacing one of the non-bridging oxygen atoms with a sulfur atom. A phosphorothioate linkage is a modified internucleoside linkage.

“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.

“Prevent” 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.

“Prodrug” means a therapeutic agent that is prepared in an inactive form that is converted to an active form within the body or cells thereof by the action of endogenous enzymes or other chemicals or conditions.

“Side effects” means physiological responses attributable to a treatment other than the desired effects. In certain embodiments, side effects include injection site reactions, liver function test abnormalities, renal function abnormalities, liver toxicity, renal toxicity, central nervous system abnormalities, myopathies, and malaise. For example, increased aminotransferase levels in serum can indicate liver toxicity or liver function abnormality. For example, increased bilirubin can indicate liver toxicity or liver function abnormality.

“Single-stranded oligonucleotide” means an oligonucleotide which is not hybridized to a complementary strand.

“Specifically hybridizable” refers to an antisense compound having a sufficient degree of complementarity with a target nucleic acid to induce a desired effect, while exhibiting minimal or no effects on non-target nucleic acids under conditions in which specific binding is desired, i.e. under physiological conditions in the case of *in vivo* assays and therapeutic treatments.

“Statin” means an agent that inhibits the activity of HMG-CoA reductase.

“Subcutaneous administration” means administration just below the skin.

“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.

“Target nucleic acid,” “target RNA,” and “target RNA transcript” all refer to a nucleic acid capable of being targeted by antisense compounds.

5 “Target region” is defined as a portion of the target nucleic acid having at least one identifiable structure, function, or characteristic.

“Target segment” means the sequence of nucleotides of a target nucleic acid to which one or more antisense compound is targeted. “5’ target site” or “5’ start site” refers to the 5’-most nucleotide of a target segment. “3’ target site” or “3’ stop site” refers to the 3’-most nucleotide of a target segment.

10 “Therapeutically effective amount” means an amount of an agent that provides a therapeutic benefit to an individual.

“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 include recommendations for dietary intake of total daily calories, total fat, saturated fat, polyunsaturated fat, 15 monounsaturated fat, carbohydrate, protein, cholesterol, insoluble fiber, as well as recommendations for physical activity.

“Triglyceride” means a lipid or neutral fat consisting of glycerol combined with three fatty acid molecules.

20 “Type 2 diabetes” (also known as “type 2 diabetes mellitus” or “diabetes mellitus, type 2”, and formerly called “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.

“Treat” refers to administering a pharmaceutical composition to effect an alteration or improvement of a disease, disorder, or condition.

25 “Unmodified nucleotide” means a nucleotide composed of naturally occurring nucleobases, sugar moieties, and internucleoside linkages. In certain embodiments, an unmodified nucleotide is a RNA nucleotide (i.e. β -D-ribonucleosides) or a DNA nucleotide (i.e. β -D-deoxyribonucleoside).

Certain Embodiments

In certain embodiments disclosed herein, ANGPTL3 has the sequence as set forth in GenBank 30 Accession No. NM_014495.2 (incorporated herein as SEQ ID NO: 1). In certain embodiments, ANGPTL3 has the sequence as set forth in GenBank Accession No. NT_032977.9 nucleotides 33032001 to 33046000 (incorporated herein as SEQ ID NO: 2).

Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide consisting of 12 to 30 nucleosides having a nucleobase sequence comprising at least 8 35 contiguous nucleobases complementary to an equal length portion of SEQ ID NOs: 1-2.

Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide 12 to 30 linked nucleosides in length targeted to ANGPTL3. The ANGPTL target can have a sequence selected from any one of SEQ ID NOs: 1-2.

5 Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and comprising a nucleobase sequence comprising a portion of at least 8 contiguous nucleobases complementary to an equal length portion of nucleobases 1140 to 1159 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 modified oligonucleotide is 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
10 contiguous nucleobases complementary to an equal length portion of nucleobases 1140 to 1159 of SEQ ID NO: 1.

Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and comprising a nucleobase sequence complementary to nucleobases 1140 to 1159 of SEQ ID NO: 1, wherein the nucleobase sequence of the
15 modified oligonucleotide is at least 80% complementary to SEQ ID NO: 1.

Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and comprising a nucleobase sequence comprising a portion of at least 8 contiguous nucleobases complementary to an equal length portion of nucleobases 1907 to 1926 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 modified oligonucleotide is 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
20 contiguous nucleobases complementary to an equal length portion of nucleobases 1907 to 1926 of SEQ ID NO: 1.

Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and comprising a nucleobase sequence complementary to nucleobases 1907 to 1926 of SEQ ID NO: 1, wherein the nucleobase sequence of the
25 modified oligonucleotide is at least 80% complementary to SEQ ID NO: 1.

Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and comprising a nucleobase sequence comprising a portion of at least 8 contiguous nucleobases complementary to an equal length portion of nucleobases 147 to 162 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 modified oligonucleotide is at least 8, least 9, least 10, least 11, at least 12, least 13, at least 14, at least 15, or 16 contiguous nucleobases complementary to an equal length portion of nucleobases 147 to 162 of SEQ ID NO: 1.
30

35 Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and comprising a nucleobase sequence

complementary to nucleobases 147 to 162 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 modified oligonucleotide consists of 12 to 30, 15 to 30, 18 to 24, 19 to 22, 13 to 25, 14 to 25, 15 to 25 or 16 to 24 linked nucleosides. In certain embodiments, the modified
 5 oligonucleotide consists of 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29 or 30 linked nucleosides or a range defined by any two of these values. In certain embodiments, the modified oligonucleotide is 16 linked nucleosides in length. In certain embodiments, the modified oligonucleotide is 20 linked nucleosides in length.

In certain embodiments, the modified oligonucleotide comprises a nucleobase sequence comprising a
 10 portion of 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 complementary to an equal length portion of SEQ ID NO: 1 or 2.

Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and having a nucleobase sequence comprising at
 15 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 a nucleobase sequence selected from any one of SEQ ID NOs: 15-27, 30-73, 75-85, 87-232, 238, 240-243, 245-247, 249-262, 264-397, 399-469, 471-541, 543-600, 604-760, 762-819, 821-966, 968-971, 973-975, 977-990, 992-1110, 1112-1186, 1188-1216, 1218-1226, 1228-1279, 1281-1293, 1295-1304, 1306-1943, 1945-1951, 1953-1977, 1979-1981, 1983-2044, 2046-2097, 2099-
 20 2181, 2183-2232, 2234-2238, 2240-2258, 2260-2265, 2267-2971, 2973-2976, 2978-4162, 4164-4329, 4331-4389, 4391-4394, 4396-4877.

Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and having 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,
 25 least 19, or 20 contiguous nucleobases of the nucleobase sequences of SEQ ID NOs: 77.

Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and having 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: 20.

Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and having a nucleobase sequence comprising at
 30 least 8, least 9, least 10, least 11, at least 12, least 13, at least 14, at least 15, or 16 contiguous nucleobases of the nucleobase sequence of SEQ ID NO: 110.

In certain embodiments, the nucleobase sequence of the modified oligonucleotide is at least 70%,
 35 75%, 80%, 85%, 90%, 95% or 100% complementary to any one of SEQ ID NO: 1-2 as measured over the entirety of the modified oligonucleotide.

In certain embodiments, the compound disclosed herein is a single-stranded oligonucleotide. In certain embodiments, the compound disclosed herein is a single-stranded modified oligonucleotide.

In certain embodiments, at least one internucleoside linkage of said modified oligonucleotide is a modified internucleoside linkage. In certain embodiments, each internucleoside linkage is a phosphorothioate internucleoside linkage.

In certain embodiments, at least one nucleoside of the modified oligonucleotide comprises a modified sugar. In certain embodiments, at least one modified sugar is a bicyclic sugar. In certain embodiments, 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.

In certain embodiments, at least one nucleoside of said modified oligonucleotide comprises a modified nucleobase. In certain embodiments, the modified nucleobase is a 5-methylcytosine.

Certain embodiments disclosed herein provide compounds or compositions comprising a modified oligonucleotide with: a) a gap segment consisting of linked deoxynucleosides; b) a 5' wing segment consisting of linked nucleosides; and c) a 3' wing segment consisting of linked nucleosides. The gap segment is positioned between the 5' wing segment and the 3' wing segment and each nucleoside of each wing segment comprises a modified sugar.

In certain embodiments, the modified oligonucleotide consists of 12 to 30 linked nucleosides and comprises: 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.

In certain embodiments, the compounds or compositions disclosed herein comprise a modified oligonucleotide consisting of 20 linked nucleosides having a nucleobase sequence comprising at least 8 contiguous nucleobases complementary to an equal length portion of SEQ ID NO: 1-2, wherein the modified oligonucleotide comprises: a gap segment consisting of ten linked deoxynucleosides; a 5' wing segment consisting of five linked nucleosides; and 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 is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

In certain embodiments, the modified oligonucleotide consists of 20 linked nucleosides and 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 is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

In certain embodiments, the compounds or compositions disclosed herein comprise a modified oligonucleotide consisting of 20 linked nucleosides having a nucleobase sequence comprising at least 8 contiguous nucleobases of a nucleobase sequence selected of SEQ ID NO: 77, wherein the modified oligonucleotide comprises: a gap segment consisting of ten linked deoxynucleosides; a 5' wing segment
 5 consisting of five linked nucleosides; and 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 is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

In certain embodiments, the modified oligonucleotide consists of 20 linked nucleosides with the
 10 nucleobase sequence of SEQ ID NO: 77 and 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 is a phosphorothioate linkage and wherein each cytosine residue is a 5-
 15 methylcytosine.

In certain embodiments, the modified oligonucleotide is ISIS 563580. In certain embodiments, ISIS 563580 is characterized as a 5-10-5 MOE gapmer, having a sequence of (from 5' to 3')
 GGACATTGCCAGTAATCGCA (incorporated herein as SEQ ID NO: 77), wherein each internucleoside linkage is a phosphorothioate linkage, each cytosine is a 5'-methylcytosine, each of nucleosides 1-5 and 16-
 20 20 are 2'-O-methoxyethyl modified nucleosides, and each of nucleosides 6-15 are 2'-deoxynucleosides.

In certain embodiments, ISIS 563580 is described by the following chemical notation: Ges Ges Aes mCes Aes Tds Tds Gds mCds mCds Ads Gds Tds Ads Ads Tes mCes Ges mCes Ae; wherein,

A = an adenine,

mC = a 5'-methylcytosine

25 G = a guanine,

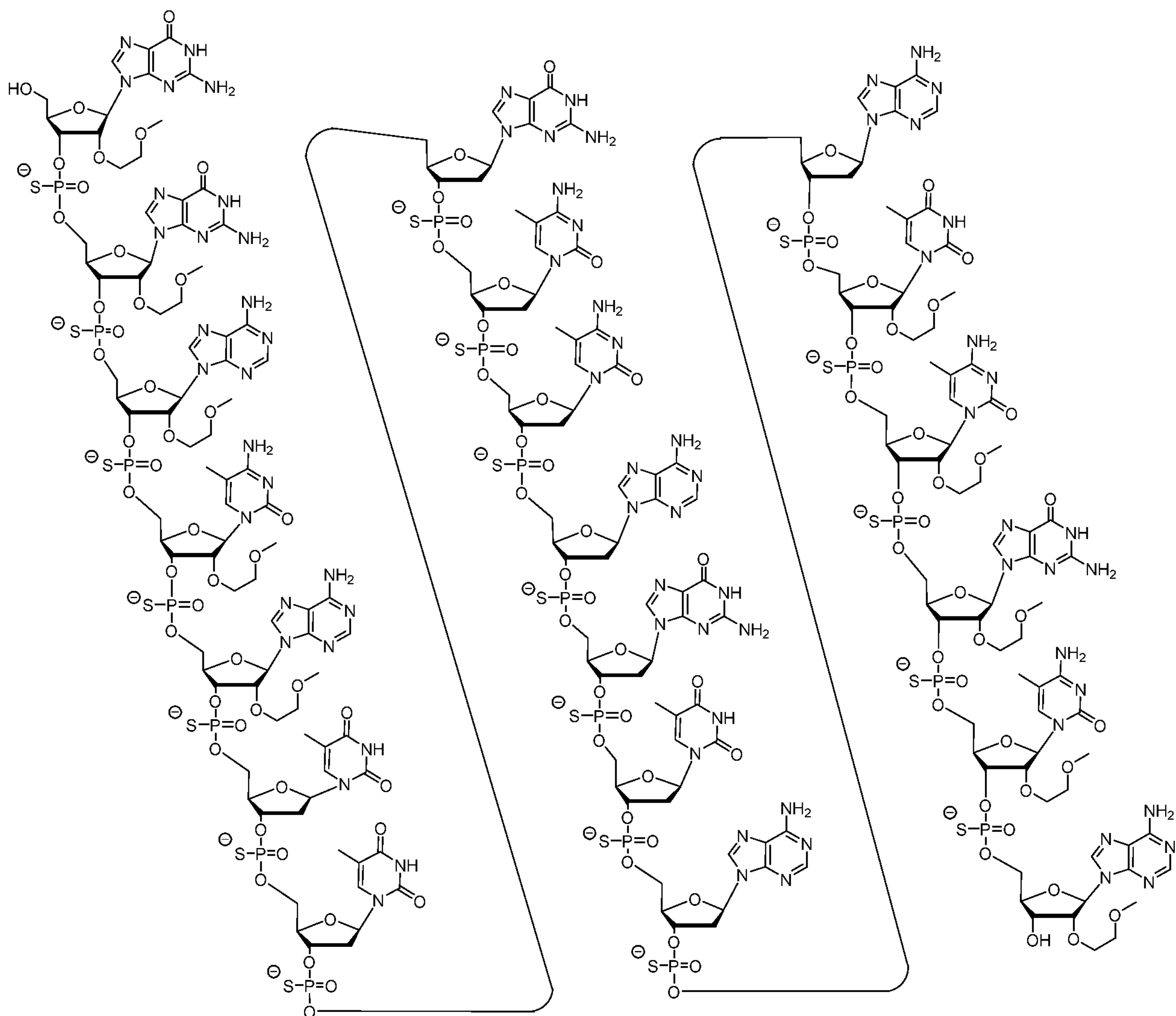
T = a thymine,

e = a 2'-O-methoxyethyl modified nucleoside,

d = a 2'-deoxynucleoside, and

s = a phosphorothioate internucleoside linkage.

30 In certain embodiments, ISIS 563580 is described by the following chemical structure:



In certain embodiments, the modified oligonucleotide comprises ISIS 563580 represented by the preceding chemical structure.

In certain embodiments, the compounds or compositions disclosed herein comprise a modified oligonucleotide consisting of 20 linked nucleosides having a nucleobase sequence comprising at least 8 contiguous nucleobases of a nucleobase sequence selected of SEQ ID NO: 20, wherein the modified oligonucleotide comprises: a gap segment consisting of ten linked deoxynucleosides; a 5' wing segment consisting of five linked nucleosides; and 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 is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

In certain embodiments, the modified oligonucleotide consists of 20 linked nucleosides with the nucleobase sequence of SEQ ID NO: 20 and 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 is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

In certain embodiments, the compounds or compositions disclosed herein comprise a modified
 5 oligonucleotide consisting of 16 linked nucleosides having a nucleobase sequence comprising at least 8
 contiguous nucleobases of a nucleobase sequence of SEQ ID NO: 110, wherein the modified oligonucleotide
 comprises: a gap segment consisting of ten linked deoxynucleosides; a 5' wing segment consisting of three
 linked nucleosides; and a 3' wing segment consisting of three linked nucleosides; wherein the gap segment is
 positioned between the 5' wing segment and the 3' wing segment; wherein each wing segment comprises at
 10 least one 2'-O-methoxyethyl sugar and at least one cEt sugar; wherein each internucleoside linkage is a
 phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

In certain embodiments, the modified oligonucleotide consists of 16 linked nucleosides with the
 nucleobase sequence of SEQ ID NO: 110 and comprises: a gap segment consisting of ten linked
 deoxynucleosides; a 5' wing segment consisting of three linked nucleosides; a 3' wing segment consisting of
 15 three linked nucleosides; wherein the gap segment is positioned between the 5' wing segment and the 3' wing
 segment; wherein each wing segment comprises at least one 2'-O-methoxyethyl sugar and at least one cEt
 sugar; wherein each internucleoside linkage is a phosphorothioate linkage and wherein each cytosine residue
 is a 5-methylcytosine.

Certain embodiments provide methods of using the compounds and compositions described herein
 20 for inhibiting ANGPTL3 expression. In certain embodiments, the compounds or compositions inhibit
 ANGPTL3 by at least 5%, at least 10%, at least 20%, at least 30%, at least 35%, at least 40%, at least 45%, at
 least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at
 least 90% or at least 95%. In a preferred embodiment, the antisense compound comprising a modified
 oligonucleotide decreases ANGPTL3 by at least 50%. In a preferred embodiment, the antisense compound
 25 comprising a modified oligonucleotide decreases ANGPTL3 by at least 55%. In a preferred embodiment the
 antisense compound comprising a modified oligonucleotide decreases ANGPTL3 by at least 60%. In a
 preferred embodiment, the antisense compound comprising a modified oligonucleotide decreases ANGPTL3
 by at least 65%. In a preferred embodiment, the antisense compound comprising a modified oligonucleotide
 decreases ANGPTL3 by at least 70%. In a preferred embodiment, the antisense compound comprising a
 30 modified oligonucleotide decreases ANGPTL3 by at least 75%. In a preferred embodiment, the antisense
 compound comprising a modified oligonucleotide decreases ANGPTL3 by at least 80%. In a preferred
 embodiment, the antisense compound comprising a modified oligonucleotide decreases ANGPTL3 by at least
 85%. In a preferred embodiment, the antisense compound comprising a modified oligonucleotide decreases
 ANGPTL3 by at least 90%. In a preferred embodiment, the antisense compound comprising a modified
 35 oligonucleotide decreases ANGPTL3 by at least 95%.

Certain embodiments provide methods of using the compounds and compositions described herein for reducing one or more of triglycerides, LDL-cholesterol, non-HDL cholesterol, VLDL-cholesterol, total cholesterol, ApoB and ApoC-III. In certain embodiments, the compounds or compositions reduce one or more of triglycerides, LDL-cholesterol, non-HDL cholesterol, VLDL-cholesterol, total cholesterol, ApoB and ApoC-III by at least 5%, at least 10%, at least 20%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90% or at least 95%.

In certain embodiments, the compounds or compositions disclosed herein have an IC_{50} of less than 20 μ M, less than 10 μ M, less than 8 μ M, less than 5 μ M, less than 2 μ M, less than 1 μ M, or less than 0.8 μ M, when tested human cells, for example, in the Hep3B cell line as described in Examples 2-3 and 7-10.

In certain embodiments, the compounds or compositions disclosed 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 13.

In certain embodiments, the compounds or compositions disclosed herein are highly tolerable, as demonstrated by the *in vivo* tolerability measurements described in the examples. In certain embodiments, the antisense compounds as described herein are highly tolerable, as demonstrated by having an increase in ALT and/or AST value of no more than 4 fold, 3 fold, 2 fold or 1.5 fold over saline treated animals.

In certain embodiments, the compounds or compositions disclosed herein comprise a salt of the modified oligonucleotide.

In certain embodiments, the compounds or compositions disclosed herein further comprise a pharmaceutically acceptable carrier or diluent.

In certain embodiments, the animal is a human.

Certain embodiments provide methods of using the compounds and compositions described herein in therapy. In certain embodiments, the therapy is used in treating, preventing, or slowing progression of a disease related to elevated ANGPTL3. In certain embodiments, the disease is a cardiovascular and/or metabolic disease, disorder or condition. In certain embodiments, the metabolic and/or cardiovascular disease includes, but is not limited to, obesity, diabetes, atherosclerosis, dyslipidemia, lipodystrophy, coronary heart disease, non-alcoholic fatty liver disease (NAFLD), nonalcoholic steatohepatitis (NASH) hyperfattyacidemia or metabolic syndrome, or a combination thereof. The dyslipidemia can be hyperlipidemia. The hyperlipidemia can be combined hyperlipidemia, familial combined hyperlipidemia (FCHL), hypercholesterolemia, hypertriglyceridemia, or both hypercholesterolemia and hypertriglyceridemia. The hypercholesterolemia can be familial homozygous hypercholesterolemia (HoFH), familial heterozygous hypercholesterolemia (HeFH). The hypertriglyceridemia can be familial chylomicronemia syndrome (FCS) or hyperlipoproteinemia Type IV. The NAFLD can be hepatic steatosis or steatohepatitis. The diabetes can be type 2 diabetes or type 2 diabetes with dyslipidemia.

In certain embodiments, the compounds or compositions disclosed herein are designated as a first agent and the methods or uses disclosed herein further comprise administering a second agent. In certain embodiments, the first agent and the second agent are co-administered. In certain embodiments the first agent and the second agent are co-administered sequentially or concomitantly.

5 In certain embodiments, the second agent is a glucose-lowering agent. The glucose lowering agent can include, but is not limited to, a therapeutic lifestyle change, PPAR agonist, a dipeptidyl peptidase (IV) inhibitor, a GLP-1 analog, insulin or an insulin analog, an insulin secretagogue, a SGLT2 inhibitor, a human amylin analog, a biguanide, an alpha-glucosidase inhibitor, or a combination thereof. The glucose-lowering agent can include, but is not limited to metformin, sulfonylurea, rosiglitazone, meglitinide, thiazolidinedione, 10 alpha-glucosidase inhibitor or a combination thereof. The sulfonylurea can be acetohexamide, chlorpropamide, tolbutamide, tolazamide, glimepiride, a glipizide, a glyburide, or a gliclazide. The meglitinide can be nateglinide or repaglinide. The thiazolidinedione can be pioglitazone or rosiglitazone. The alpha-glucosidase can be acarbose or miglitol.

In certain embodiments, the second agent is a lipid-lowering therapy. In certain embodiments the 15 lipid lowering therapy can include, but is not limited to, a therapeutic lifestyle change, HMG-CoA reductase inhibitor, cholesterol absorption inhibitor, MTP inhibitor (e.g., a small molecule, polypeptide, antibody or antisense compound targeted to MTP), ApoB inhibitor (e.g., a small molecule, polypeptide, antibody or antisense compound targeted to ApoB), ApoC3 inhibitor (e.g., a small molecule, polypeptide, antibody or antisense compound targeted to ApoC3), PCSK9 inhibitor (e.g., a small molecule, polypeptide, antibody or 20 antisense compound targeted to PCSK9), CETP inhibitor (e.g., a small molecule, polypeptide, antibody or antisense compound targeted to CETP), fibrate, beneficial oil (e.g., krill or fish oils (e.g., Vascepa^R), flaxseed oil, or other oils rich in omega-3 fatty acids such as α -linolenic acid (ALA), docosahexaenoic acid (DHA) or eicosapentaenoic acid (EPA)), or any combination thereof. The HMG-CoA reductase inhibitor can be atorvastatin, rosuvastatin, fluvastatin, lovastatin, pravastatin, or simvastatin. The cholesterol absorption 25 inhibitor can be ezetimibe. The fibrate can be fenofibrate, bezafibrate, ciprofibrate, clofibrate, gemfibrozil and the like.

In certain embodiments, administration comprises parenteral administration.

In certain embodiments, administering a compound disclosed herein results in a reduction of lipid levels, including triglyceride levels, cholesterol levels, insulin resistance, glucose levels or a combination 30 thereof. One or more of the levels can be independently reduced by at least 5%, at least 10%, at least 20%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90% or at least 95%. Administering the compound can result in improved insulin sensitivity or hepatic insulin sensitivity. Administering the compound disclosed herein can result in a reduction in atherosclerotic plaques, obesity, glucose, lipids, glucose 35 resistance, cholesterol, or improvement in insulin sensitivity or any combination thereof.

Certain embodiments provide the use of a compound as described herein in the manufacture of a medicament for treating, ameliorating, delaying or preventing one or more of a metabolic disease or a cardiovascular disease.

5 Certain embodiments provide a kit for treating, preventing, or ameliorating one or more of a metabolic disease or a cardiovascular disease as described herein wherein the kit comprises: a) a compound as described herein; and optionally b) an additional agent or therapy as described herein. The kit can further include instructions or a label for using the kit to treat, prevent, or ameliorate one or more of a metabolic disease or a cardiovascular disease.

Antisense Compounds

10 Oligomeric compounds include, but are not limited to, oligonucleotides, oligonucleosides, oligonucleotide analogs, oligonucleotide mimetics, antisense compounds, antisense oligonucleotides, and siRNAs. An oligomeric compound can be “antisense” to a target nucleic acid, meaning that is capable of undergoing hybridization to a target nucleic acid through hydrogen bonding.

15 In certain embodiments, an antisense compound has a nucleobase sequence that, when written in the 5' to 3' direction, comprises the reverse complement of the target segment of a target nucleic acid to which it is targeted. In certain such embodiments, an antisense oligonucleotide has a nucleobase sequence that, when written in the 5' to 3' direction, comprises the reverse complement of the target segment of a target nucleic acid to which it is targeted.

In certain embodiments, an antisense compound targeted to ANGPTL3 nucleic acid is 10 to 30
20 nucleotides in length. In other words, antisense compounds are from 10 to 30 linked nucleobases. In other embodiments, the antisense compound comprises a modified oligonucleotide consisting of 8 to 80, 10 to 80, 12 to 50, 12 to 30, 15 to 30, 18 to 24, 19 to 22, or 20 linked nucleobases. In certain such embodiments, the antisense compound comprises a modified oligonucleotide consisting of 8, 9, 10, 11, 12, 13, 14, 15, 16, 17,
25 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, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, or 80 linked nucleobases in length, or a range defined by any two of the above values.

In certain embodiments, the antisense compound comprises a shortened or truncated modified oligonucleotide. The shortened or truncated modified oligonucleotide can have a single nucleoside deleted from the 5' end (5' truncation), or alternatively from the 3' end (3' truncation). A shortened or truncated
30 oligonucleotide can have two or more nucleosides deleted from the 5' end, or alternatively can have two or more nucleosides deleted from the 3' end. Alternatively, the deleted nucleosides can be dispersed throughout the modified oligonucleotide, for example, in an antisense compound having one or more nucleoside deleted from the 5' end and one or more nucleoside deleted from the 3' end.

When a single additional nucleoside is present in a lengthened oligonucleotide, the additional
35 nucleoside can be located at the 5', 3' end or central portion of the oligonucleotide. When two or more

additional nucleosides are present, the added nucleosides can be adjacent to each other, for example, in an oligonucleotide having two nucleosides added to the 5' end (5' addition), or alternatively to the 3' end (3' addition) or the central portion, of the oligonucleotide. Alternatively, the added nucleoside can be dispersed throughout the antisense compound, for example, in an oligonucleotide having one or more nucleoside added to the 5' end, one or more nucleoside added to the 3' end, and/or one or more nucleoside added to the central portion.

It is possible to increase or decrease the length of an antisense compound, such as an antisense oligonucleotide, and/or introduce mismatch bases without eliminating activity. For example, in Woolf et al. (Proc. Natl. Acad. Sci. USA 89:7305-7309, 1992), a series of antisense oligonucleotides 13-25 nucleobases in length were tested for their ability to induce cleavage of a target RNA in an oocyte injection model. Antisense oligonucleotides 25 nucleobases in length with 8 or 11 mismatch bases near the ends of the antisense oligonucleotides were able to direct specific cleavage of the target mRNA, albeit to a lesser extent than the antisense oligonucleotides that contained no mismatches. Similarly, target specific cleavage was achieved using 13 nucleobase antisense oligonucleotides, including those with 1 or 3 mismatches.

Gautschi et al (J. Natl. Cancer Inst. 93:463-471, March 2001) demonstrated the ability of an oligonucleotide having 100% complementarity to the bcl-2 mRNA and having 3 mismatches to the bcl-xL mRNA to reduce the expression of both bcl-2 and bcl-xL in vitro and in vivo. Furthermore, this oligonucleotide demonstrated potent anti-tumor activity in vivo.

Maher and Dolnick (Nuc. Acid. Res. 16:3341-3358, 1988) tested a series of tandem 14 nucleobase antisense oligonucleotides, and a 28 and 42 nucleobase antisense oligonucleotides comprised of the sequence of two or three of the tandem antisense oligonucleotides, respectively, for their ability to arrest translation of human DHFR in a rabbit reticulocyte assay. Each of the three 14 nucleobase antisense oligonucleotides alone was able to inhibit translation, albeit at a more modest level than the 28 or 42 nucleobase antisense oligonucleotides.

Certain Antisense Compound Motifs and Mechanisms

In certain embodiments, antisense compounds have chemically modified subunits arranged in patterns, or motifs, to confer to the antisense compounds properties such as enhanced inhibitory activity, increased binding affinity for a target nucleic acid, or resistance to degradation by *in vivo* nucleases.

Chimeric antisense compounds typically contain at least one region modified so as to confer increased resistance to nuclease degradation, increased cellular uptake, increased binding affinity for the target nucleic acid, and/or increased inhibitory activity. A second region of a chimeric antisense compound may confer another desired property e.g., serve as a substrate for the cellular endonuclease RNase H, which cleaves the RNA strand of an RNA:DNA duplex.

Antisense activity may result from any mechanism involving the hybridization of the antisense compound (e.g., oligonucleotide) with a target nucleic acid, wherein the hybridization ultimately results in a biological effect. In certain embodiments, the amount and/or activity of the target nucleic acid is modulated. In certain embodiments, the amount and/or activity of the target nucleic acid is reduced. In certain
5 embodiments, hybridization of the antisense compound to the target nucleic acid ultimately results in target nucleic acid degradation. In certain embodiments, hybridization of the antisense compound to the target nucleic acid does not result in target nucleic acid degradation. In certain such embodiments, the presence of the antisense compound hybridized with the target nucleic acid (occupancy) results in a modulation of antisense activity. In certain embodiments, antisense compounds having a particular chemical motif or
10 pattern of chemical modifications are particularly suited to exploit one or more mechanisms. In certain embodiments, antisense compounds function through more than one mechanism and/or through mechanisms that have not been elucidated. Accordingly, the antisense compounds described herein are not limited by particular mechanism.

Antisense mechanisms include, without limitation, RNase H mediated antisense; RNAi mechanisms,
15 which utilize the RISC pathway and include, without limitation, siRNA, ssRNA and microRNA mechanisms; and occupancy based mechanisms. Certain antisense compounds may act through more than one such mechanism and/or through additional mechanisms.

RNase H-Mediated Antisense

20 In certain embodiments, antisense activity results at least in part from degradation of target RNA by RNase H. RNase H is a cellular endonuclease that 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. Accordingly, antisense compounds comprising at least a portion of DNA or DNA-like nucleosides may activate RNase H, resulting in cleavage of the target nucleic acid. In certain embodiments,
25 antisense compounds that utilize RNase H comprise one or more modified nucleosides. In certain embodiments, such antisense compounds comprise at least one block of 1-8 modified nucleosides. In certain such embodiments, the modified nucleosides do not support RNase H activity. In certain embodiments, such antisense compounds are gapmers, as described herein. In certain such embodiments, the gap of the gapmer comprises DNA nucleosides. In certain such embodiments, the gap of the gapmer comprises DNA-like
30 nucleosides. In certain such embodiments, the gap of the gapmer comprises DNA nucleosides and DNA-like nucleosides.

Certain antisense compounds having a gapmer motif are considered chimeric antisense compounds. In a gapmer an internal region having a plurality of nucleotides that supports RNaseH cleavage is positioned between external regions having a plurality of nucleotides that are chemically distinct from the nucleosides of
35 the internal region. In the case of an antisense oligonucleotide having a gapmer motif, the gap segment generally serves as the substrate for endonuclease cleavage, while the wing segments comprise modified

nucleosides. In certain embodiments, the regions of a gapmer are differentiated by the types of sugar moieties comprising each distinct region. The types of sugar moieties that are used to differentiate the regions of a gapmer may in some embodiments include β -D-ribonucleosides, β -D-deoxyribonucleosides, 2'-modified nucleosides (such 2'-modified nucleosides may include 2'-MOE and 2'-O-CH₃, among others), and
 5 bicyclic sugar modified nucleosides (such bicyclic sugar modified nucleosides may include those having a constrained ethyl). In certain embodiments, nucleosides in the wings may include several modified sugar moieties, including, for example 2'-MOE and bicyclic sugar moieties such as constrained ethyl or LNA. In certain embodiments, wings may include several modified and unmodified sugar moieties. In certain embodiments, wings may include various combinations of 2'-MOE nucleosides, bicyclic sugar moieties such
 10 as constrained ethyl nucleosides or LNA nucleosides, and 2'-deoxynucleosides.

Each distinct region may comprise uniform sugar moieties, variant, or alternating sugar moieties. The wing-gap-wing motif is frequently described as "X-Y-Z", where "X" represents the length of the 5'-wing, "Y" represents the length of the gap, and "Z" represents the length of the 3'-wing. "X" and "Z" may comprise uniform, variant, or alternating sugar moieties. In certain embodiments, "X" and "Y" may include
 15 one or more 2'-deoxynucleosides. "Y" may comprise 2'-deoxynucleosides. As used herein, a gapmer described as "X-Y-Z" has a configuration such that the gap is positioned immediately adjacent to each of the 5'-wing and the 3' wing. Thus, no intervening nucleotides exist between the 5'-wing and gap, or the gap and the 3'-wing. Any of the antisense compounds described herein can have a gapmer motif. In certain embodiments, "X" and "Z" are the same; in other embodiments they are different. In certain embodiments,
 20 "Y" is between 8 and 15 nucleosides. X, Y, or Z can be any of 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 25, 30 or more nucleosides.

In certain embodiments, the antisense compound targeted to an ANGPTL3 nucleic acid has a gapmer motif in which the gap consists of 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, or 16 linked nucleosides.

In certain embodiments, the antisense oligonucleotide has a sugar motif described by Formula A as
 25 follows: (J)_m-(B)_n-(J)_p-(B)_r-(A)_t-(D)_g-(A)_v-(B)_w-(J)_x-(B)_y-(J)_z

wherein:

each A is independently a 2'-substituted nucleoside;

each B is independently a bicyclic nucleoside;

each J is independently either a 2'-substituted nucleoside or a 2'-deoxynucleoside;

30 each D is a 2'-deoxynucleoside;

m is 0-4; n is 0-2; p is 0-2; r is 0-2; t is 0-2; v is 0-2; w is 0-4; x is 0-2; y is 0-2; z is 0-4; g is 6-14;

provided that:

at least one of m, n, and r is other than 0;

at least one of w and y is other than 0;

35 the sum of m, n, p, r, and t is from 2 to 5; and

the sum of v, w, x, y, and z is from 2 to 5.

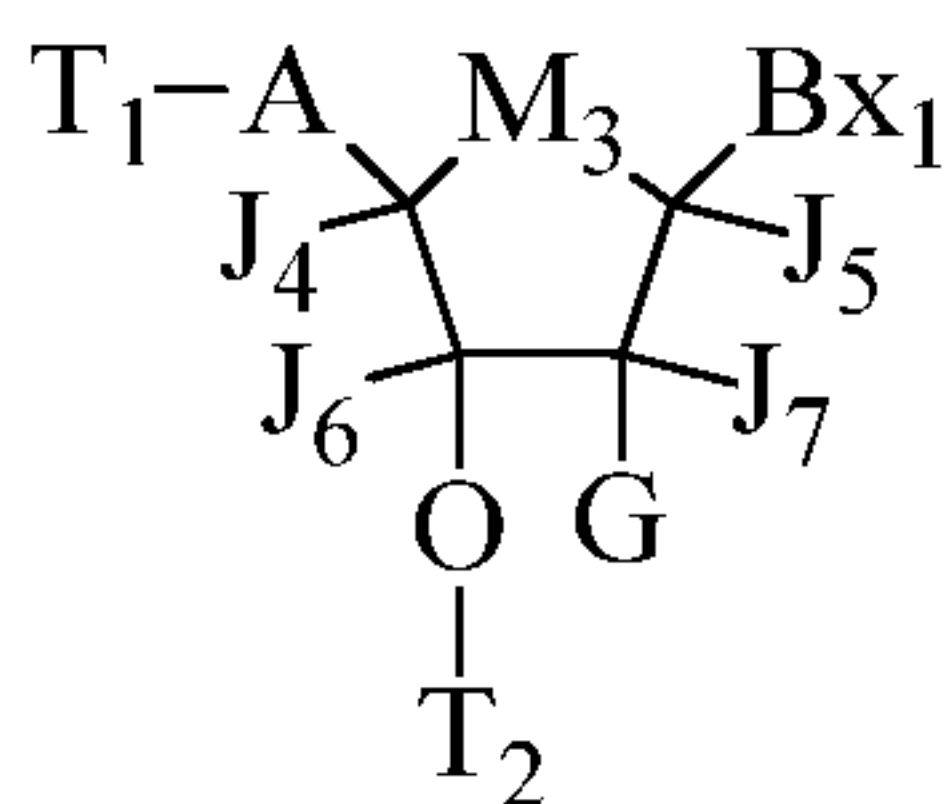
RNAi Compounds

In certain embodiments, antisense compounds are interfering RNA compounds (RNAi), which include double-stranded RNA compounds (also referred to as short-interfering RNA or siRNA) and single-stranded RNAi compounds (or ssRNA). Such compounds work at least in part through the RISC pathway to degrade and/or sequester a target nucleic acid (thus, include microRNA/microRNA-mimic compounds). In certain embodiments, antisense compounds comprise modifications that make them particularly suited for such mechanisms.

i. ssRNA compounds

In certain embodiments, antisense compounds including those particularly suited for use as single-stranded RNAi compounds (ssRNA) comprise a modified 5'-terminal end. In certain such embodiments, the 5'-terminal end comprises a modified phosphate moiety. In certain embodiments, such modified phosphate is stabilized (e.g., resistant to degradation/cleavage compared to unmodified 5'-phosphate). In certain embodiments, such 5'-terminal nucleosides stabilize the 5'-phosphorous moiety. Certain modified 5'-terminal nucleosides may be found in the art, for example in WO/2011/139702.

In certain embodiments, the 5'-nucleoside of an ssRNA compound has Formula IIc:



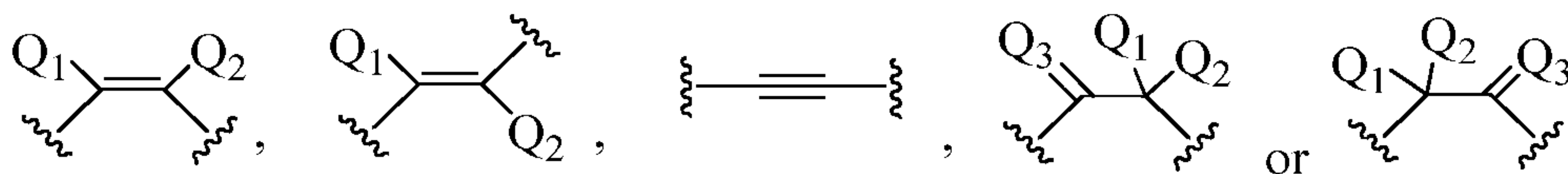
IIc

wherein:

T₁ is an optionally protected phosphorus moiety;

T₂ is an internucleoside linking group linking the compound of Formula IIc to the oligomeric compound;

A has one of the formulas:



Q₁ and Q₂ are each, independently, H, halogen, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₁-C₆ alkoxy, substituted C₁-C₆ alkoxy, C₂-C₆ alkenyl, substituted C₂-C₆ alkenyl, C₂-C₆ alkynyl, substituted C₂-C₆ alkynyl or N(R₃)(R₄);

Q₃ is O, S, N(R₅) or C(R₆)(R₇);

each R₃, R₄, R₅, R₆ and R₇ is, independently, H, C₁-C₆ alkyl, substituted C₁-C₆ alkyl or C₁-C₆ alkoxy;

M₃ is O, S, NR₁₄, C(R₁₅)(R₁₆), C(R₁₅)(R₁₆)C(R₁₇)(R₁₈), C(R₁₅)=C(R₁₇), OC(R₁₅)(R₁₆) or OC(R₁₅)(Bx₂);

R₁₄ is H, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₁-C₆ alkoxy, substituted C₁-C₆ alkoxy, C₂-C₆ alkenyl, substituted C₂-C₆ alkenyl, C₂-C₆ alkynyl or substituted C₂-C₆ alkynyl;

5 R₁₅, R₁₆, R₁₇ and R₁₈ are each, independently, H, halogen, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₁-C₆ alkoxy, substituted C₁-C₆ alkoxy, C₂-C₆ alkenyl, substituted C₂-C₆ alkenyl, C₂-C₆ alkynyl or substituted C₂-C₆ alkynyl;

Bx₁ is a heterocyclic base moiety;

10 or if Bx₂ is present then Bx₂ is a heterocyclic base moiety and Bx₁ is H, halogen, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₁-C₆ alkoxy, substituted C₁-C₆ alkoxy, C₂-C₆ alkenyl, substituted C₂-C₆ alkenyl, C₂-C₆ alkynyl or substituted C₂-C₆ alkynyl;

J₄, J₅, J₆ and J₇ are each, independently, H, halogen, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₁-C₆ alkoxy, substituted C₁-C₆ alkoxy, C₂-C₆ alkenyl, substituted C₂-C₆ alkenyl, C₂-C₆ alkynyl or substituted C₂-C₆ alkynyl;

15 or J₄ forms a bridge with one of J₅ or J₇ wherein said bridge comprises from 1 to 3 linked biradical groups selected from O, S, NR₁₉, C(R₂₀)(R₂₁), C(R₂₀)=C(R₂₁), C[=C(R₂₀)(R₂₁)] and C(=O) and the other two of J₅, J₆ and J₇ are each, independently, H, halogen, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₁-C₆ alkoxy, substituted C₁-C₆ alkoxy, C₂-C₆ alkenyl, substituted C₂-C₆ alkenyl, C₂-C₆ alkynyl or substituted C₂-C₆ alkynyl;

20 each R₁₉, R₂₀ and R₂₁ is, independently, H, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₁-C₆ alkoxy, substituted C₁-C₆ alkoxy, C₂-C₆ alkenyl, substituted C₂-C₆ alkenyl, C₂-C₆ alkynyl or substituted C₂-C₆ alkynyl;

G is H, OH, halogen or O-[C(R₈)(R₉)]_n-(C=O)_m-X₁]-Z;

each R₈ and R₉ is, independently, H, halogen, C₁-C₆ alkyl or substituted C₁-C₆ alkyl;

25 X₁ is O, S or N(E₁);

Z is H, halogen, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₂-C₆ alkenyl, substituted C₂-C₆ alkenyl, C₂-C₆ alkynyl, substituted C₂-C₆ alkynyl or N(E₂)(E₃);

E₁, E₂ and E₃ are each, independently, H, C₁-C₆ alkyl or substituted C₁-C₆ alkyl;

n is from 1 to about 6;

30 m is 0 or 1;

j is 0 or 1;

each substituted group comprises one or more optionally protected substituent groups independently selected from halogen, OJ₁, N(J₁)(J₂), =NJ₁, SJ₁, N₃, CN, OC(=X₂)J₁, OC(=X₂)N(J₁)(J₂) and C(=X₂)N(J₁)(J₂);

X₂ is O, S or NJ₃;

35 each J₁, J₂ and J₃ is, independently, H or C₁-C₆ alkyl;

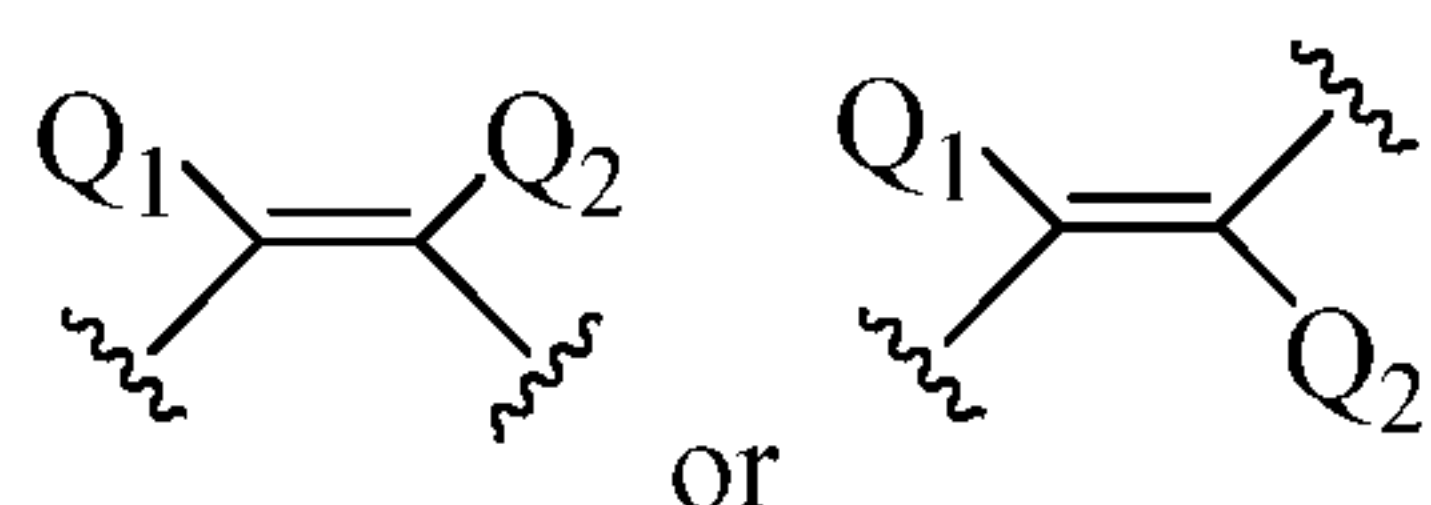
when j is 1 then Z is other than halogen or N(E₂)(E₃); and

wherein said oligomeric compound comprises from 8 to 40 monomeric subunits and is hybridizable to at least a portion of a target nucleic acid.

In certain embodiments, M_3 is O, CH=CH, OCH₂ or OC(H)(Bx₂). In certain embodiments, M_3 is O.

In certain embodiments, J_4 , J_5 , J_6 and J_7 are each H. In certain embodiments, J_4 forms a bridge with one of J_5 or J_7 .

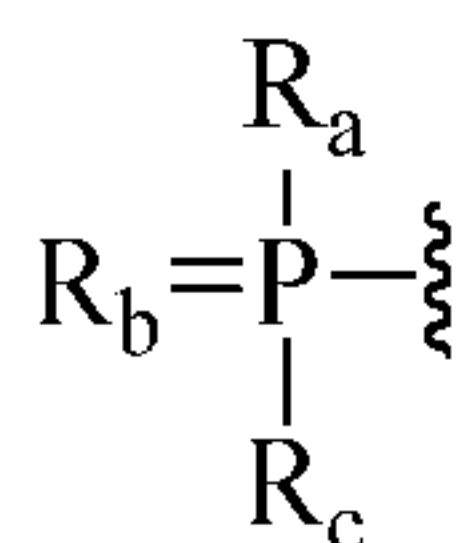
In certain embodiments, A has one of the formulas:



wherein:

Q_1 and Q_2 are each, independently, H, halogen, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₁-C₆ alkoxy or substituted C₁-C₆ alkoxy. In certain embodiments, Q_1 and Q_2 are each H. In certain embodiments, Q_1 and Q_2 are each, independently, H or halogen. In certain embodiments, Q_1 and Q_2 is H and the other of Q_1 and Q_2 is F, CH₃ or OCH₃.

In certain embodiments, T_1 has the formula:



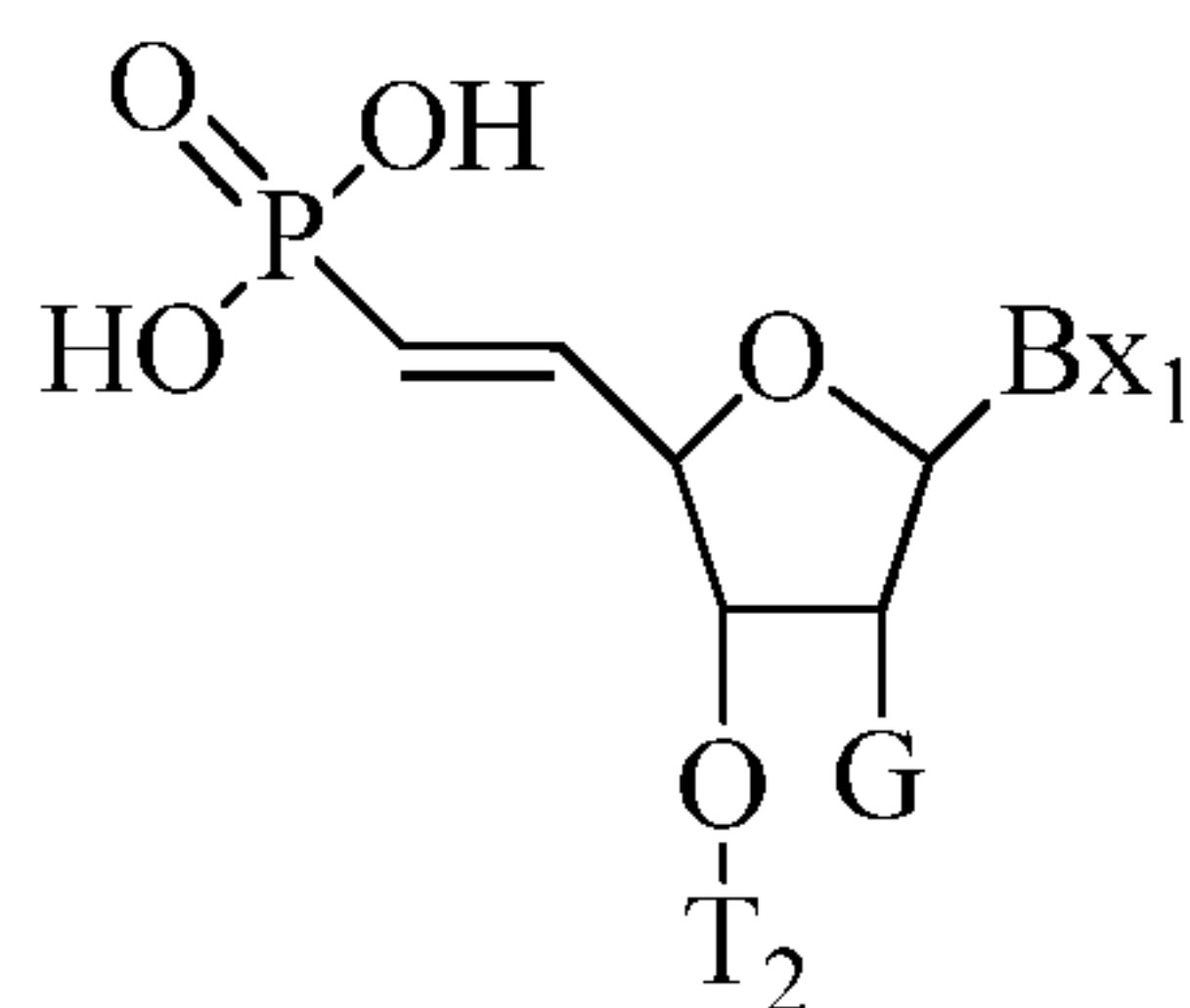
wherein:

R_a and R_c are each, independently, protected hydroxyl, protected thiol, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₁-C₆ alkoxy, substituted C₁-C₆ alkoxy, protected amino or substituted amino; and

R_b is O or S. In certain embodiments, R_b is O and R_a and R_c are each, independently, OCH₃, OCH₂CH₃ or CH(CH₃)₂.

In certain embodiments, G is halogen, OCH₃, OCH₂F, OCHF₂, OCF₃, OCH₂CH₃, O(CH₂)₂F, OCH₂CHF₂, OCH₂CF₃, OCH₂-CH=CH₂, O(CH₂)₂-OCH₃, O(CH₂)₂-SCH₃, O(CH₂)₂-OCF₃, O(CH₂)₃-N(R₁₀)(R₁₁), O(CH₂)₂-ON(R₁₀)(R₁₁), O(CH₂)₂-O(CH₂)₂-N(R₁₀)(R₁₁), OCH₂C(=O)-N(R₁₀)(R₁₁), OCH₂C(=O)-N(R₁₂)-(CH₂)₂-N(R₁₀)(R₁₁) or O(CH₂)₂-N(R₁₂)-C(=NR₁₃)[N(R₁₀)(R₁₁)] wherein R₁₀, R₁₁, R₁₂ and R₁₃ are each, independently, H or C₁-C₆ alkyl. In certain embodiments, G is halogen, OCH₃, OCF₃, OCH₂CH₃, OCH₂CF₃, OCH₂-CH=CH₂, O(CH₂)₂-OCH₃, O(CH₂)₂-O(CH₂)₂-N(CH₃)₂, OCH₂C(=O)-N(H)CH₃, OCH₂C(=O)-N(H)-(CH₂)₂-N(CH₃)₂ or OCH₂-N(H)-C(=NH)NH₂. In certain embodiments, G is F, OCH₃ or O(CH₂)₂-OCH₃. In certain embodiments, G is O(CH₂)₂-OCH₃.

In certain embodiments, the 5'-terminal nucleoside has Formula IIe:



Ile

In certain embodiments, antisense compounds, including those particularly suitable for ssRNA 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.

In certain embodiments, the oligonucleotides comprise or consist of a region having uniform sugar modifications. In certain such embodiments, each nucleoside of the region comprises the same RNA-like sugar modification. In certain embodiments, each nucleoside of the region is a 2'-F nucleoside. In certain embodiments, each nucleoside of the region is a 2'-OMe nucleoside. In certain embodiments, each nucleoside of the region is a 2'-MOE nucleoside. In certain embodiments, each nucleoside of the region is a cEt nucleoside. In certain embodiments, each nucleoside of the region is an LNA nucleoside. In certain embodiments, the uniform region constitutes all or essentially all of the oligonucleotide. In certain embodiments, the region constitutes the entire oligonucleotide except for 1-4 terminal nucleosides.

In certain embodiments, oligonucleotides comprise one or more regions of alternating sugar modifications, wherein the nucleosides alternate between nucleotides having a sugar modification of a first type and nucleotides having a sugar modification of a second type. In certain embodiments, nucleosides of both types are RNA-like nucleosides. In certain embodiments the alternating nucleosides are selected from: 2'-OMe, 2'-F, 2'-MOE, LNA, and cEt. In certain embodiments, the alternating modifications are 2'-F and 2'-OMe. Such regions may be contiguous or may be interrupted by differently modified nucleosides or conjugated nucleosides.

In certain embodiments, the alternating region of alternating modifications each consist of a single nucleoside (i.e., the pattern is $(AB)_x A_y$ wherein A is a nucleoside having a sugar modification of a first type and B is a nucleoside having a sugar modification of a second type; x is 1-20 and y is 0 or 1). In certain embodiments, one or more alternating regions in an alternating motif includes more than a single nucleoside of a type. For example, oligonucleotides may include one or more regions of any of the following nucleoside motifs:

AABBAA;

ABBABB;

AABAAB;

ABBABAABB;

ABABAA;

AABABAB;

ABABAA;

5 ABBAABBABABAA;

BABBAABBABABAA; or

ABABBAABBABABAA;

wherein A is a nucleoside of a first type and B is a nucleoside of a second type. In certain embodiments, A and B are each selected from 2'-F, 2'-OMe, BNA, and MOE.

10 In certain embodiments, oligonucleotides having such an alternating motif also comprise a modified 5' terminal nucleoside, such as those of formula IIc or IIe.

In certain embodiments, oligonucleotides comprise a region having a 2-2-3 motif. Such regions comprises the following motif:

$-(A)_2-(B)_x-(A)_2-(C)_y-(A)_3-$

15 wherein: A is a first type of modified nucleoside;

B and C, are nucleosides that are differently modified than A, however, B and C may have the same or different modifications as one another;

x and y are from 1 to 15.

In certain embodiments, A is a 2'-OMe modified nucleoside. In certain embodiments, B and C are both 2'-F modified nucleosides. In certain embodiments, A is a 2'-OMe modified nucleoside and B and C are both 2'-F modified nucleosides.

In certain embodiments, oligonucleosides have the following sugar motif:

$5'-(Q)-(AB)_xA_y-(D)_z$

wherein:

25 Q is a nucleoside comprising a stabilized phosphate moiety. In certain embodiments, Q is a nucleoside having Formula IIc or IIe;

A is a first type of modified nucleoside;

B is a second type of modified nucleoside;

D is a modified nucleoside comprising a modification different from the nucleoside adjacent to it.

30 Thus, if y is 0, then D must be differently modified than B and if y is 1, then D must be differently modified than A. In certain embodiments, D differs from both A and B.

X is 5-15;

Y is 0 or 1;

Z is 0-4.

35 In certain embodiments, oligonucleosides have the following sugar motif:

5'- (Q)- (A)_x-(D)_z

wherein:

Q is a nucleoside comprising a stabilized phosphate moiety. In certain embodiments, Q is a nucleoside having Formula IIc or IIe;

5 A is a first type of modified nucleoside;

D is a modified nucleoside comprising a modification different from A.

X is 11-30;

Z is 0-4.

10 In certain embodiments A, B, C, and D in the above motifs are selected from: 2'-OMe, 2'-F, 2'-MOE, LNA, and cEt. In certain embodiments, D represents terminal nucleosides. In certain embodiments, such terminal nucleosides are not designed to hybridize to the target nucleic acid (though one or more might hybridize by chance). In certain embodiments, the nucleobase of each D nucleoside is adenine, regardless of the identity of the nucleobase at the corresponding position of the target nucleic acid. In certain embodiments the nucleobase of each D nucleoside is thymine.

15 In certain embodiments, antisense compounds, including those particularly suited for use as ssRNA 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 comprise a region of uniformly modified internucleoside linkages. In certain such embodiments, the
20 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 phosphorothioate and at least one internucleoside linkage is phosphorothioate.

25 In certain embodiments, the oligonucleotide comprises at least 6 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises at least 8 phosphorothioate internucleoside linkages. In certain embodiments, the oligonucleotide comprises at least 10 phosphorothioate internucleoside linkages. 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
30 one block of at least 8 consecutive 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
35 within 3 nucleosides of the 3' end of the oligonucleotide.

Oligonucleotides having any of the various sugar motifs described herein, may have any linkage

motif. For example, the oligonucleotides, including but not limited to those described above, may have a linkage motif selected from non-limiting the table below:

5' most linkage	Central region	3'-region
PS	Alternating PO/PS	6 PS
PS	Alternating PO/PS	7 PS
PS	Alternating PO/PS	8 PS

5 *ii. siRNA compounds*

In certain embodiments, antisense compounds are double-stranded RNAi compounds (siRNA). In such embodiments, one or both strands may comprise any modification motif described above for ssRNA. In certain embodiments, ssRNA compounds may be unmodified RNA. In certain embodiments, siRNA compounds may comprise unmodified RNA nucleosides, but modified internucleoside linkages.

10 Several embodiments relate to double-stranded compositions wherein each strand comprises a motif defined by the location of one or more modified or unmodified nucleosides. In certain embodiments, compositions are provided comprising a first and a second oligomeric compound that are fully or at least partially hybridized to form a duplex region and further comprising a region that is complementary to and hybridizes to a nucleic acid target. It is suitable that such a composition comprise a first oligomeric
15 compound that is an antisense strand having full or partial complementarity to a nucleic acid target and a second oligomeric compound that is a sense strand having one or more regions of complementarity to and forming at least one duplex region with the first oligomeric compound.

The compositions of several embodiments modulate gene expression by hybridizing to a nucleic acid target resulting in loss of its normal function. In some embodiments, the target nucleic acid is ANGPTL3. In
20 certain embodiment, the degradation of the targeted ANGPTL3 is facilitated by an activated RISC complex that is formed with compositions disclosed herein.

Several embodiments are directed to double-stranded compositions wherein one of the strands is useful in, for example, influencing the preferential loading of the opposite strand into the RISC (or cleavage) complex. The compositions are useful for targeting selected nucleic acid molecules and modulating the
25 expression of one or more genes. In some embodiments, the compositions of the present invention hybridize to a portion of a target RNA resulting in loss of normal function of the target RNA.

Certain embodiments are drawn to double-stranded compositions wherein both the strands comprises a hemimer motif, a fully modified motif, a positionally modified motif or an alternating motif. Each strand of the compositions of the present invention can be modified to fulfil a particular role in for example the siRNA
30 pathway. Using a different motif in each strand or the same motif with different chemical modifications in each strand permits targeting the antisense strand for the RISC complex while inhibiting the incorporation of

the sense strand. Within this model, each strand can be independently modified such that it is enhanced for its particular role. The antisense strand can be modified at the 5'-end to enhance its role in one region of the RISC while the 3'-end can be modified differentially to enhance its role in a different region of the RISC.

The double-stranded oligonucleotide molecules can be a double-stranded polynucleotide molecule comprising self-complementary sense and antisense regions, wherein the antisense region comprises nucleotide sequence that is complementary to nucleotide sequence in a target nucleic acid molecule or a portion thereof and the sense region having nucleotide sequence corresponding to the target nucleic acid sequence or a portion thereof. The double-stranded oligonucleotide molecules can be assembled from two separate oligonucleotides, where one strand is the sense strand and the other is the antisense strand, wherein the antisense and sense strands are self-complementary (i.e. each strand comprises nucleotide sequence that is complementary to nucleotide sequence in the other strand; such as where the antisense strand and sense strand form a duplex or double-stranded structure, for example wherein the double-stranded region is about 15 to about 30, e.g., about 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29 or 30 base pairs; the antisense strand comprises nucleotide sequence that is complementary to nucleotide sequence in a target nucleic acid molecule or a portion thereof and the sense strand comprises nucleotide sequence corresponding to the target nucleic acid sequence or a portion thereof (e.g., about 15 to about 25 or more nucleotides of the double-stranded oligonucleotide molecule are complementary to the target nucleic acid or a portion thereof). Alternatively, the double-stranded oligonucleotide is assembled from a single oligonucleotide, where the self-complementary sense and antisense regions of the siRNA are linked by means of a nucleic acid based or non-nucleic acid-based linker(s).

The double-stranded oligonucleotide can be a polynucleotide with a duplex, asymmetric duplex, hairpin or asymmetric hairpin secondary structure, having self-complementary sense and antisense regions, wherein the antisense region comprises nucleotide sequence that is complementary to nucleotide sequence in a separate target nucleic acid molecule or a portion thereof and the sense region having nucleotide sequence corresponding to the target nucleic acid sequence or a portion thereof. The double-stranded oligonucleotide can be a circular single-stranded polynucleotide having two or more loop structures and a stem comprising self-complementary sense and antisense regions, wherein the antisense region comprises nucleotide sequence that is complementary to nucleotide sequence in a target nucleic acid molecule or a portion thereof and the sense region having nucleotide sequence corresponding to the target nucleic acid sequence or a portion thereof, and wherein the circular polynucleotide can be processed either in vivo or in vitro to generate an active siRNA molecule capable of mediating RNAi.

In certain embodiments, the double-stranded oligonucleotide comprises separate sense and antisense sequences or regions, wherein the sense and antisense regions are covalently linked by nucleotide or non-nucleotide linkers molecules as is known in the art, or are alternately non-covalently linked by ionic interactions, hydrogen bonding, van der waals interactions, hydrophobic interactions, and/or stacking interactions. In certain embodiments, the double-stranded oligonucleotide comprises nucleotide sequence that

is complementary to nucleotide sequence of a target gene. In another embodiment, the double-stranded oligonucleotide interacts with nucleotide sequence of a target gene in a manner that causes inhibition of expression of the target gene.

As used herein, double-stranded oligonucleotides need not be limited to those molecules containing only RNA, but further encompasses chemically modified nucleotides and non-nucleotides. In certain embodiments, the short interfering nucleic acid molecules lack 2'-hydroxy (2'-OH) containing nucleotides. In certain embodiments short interfering nucleic acids optionally do not include any ribonucleotides (e.g., nucleotides having a 2'-OH group). Such double-stranded oligonucleotides that do not require the presence of ribonucleotides within the molecule to support RNAi can however have an attached linker or linkers or other attached or associated groups, moieties, or chains containing one or more nucleotides with 2'-OH groups. Optionally, double-stranded oligonucleotides can comprise ribonucleotides at about 5, 10, 20, 30, 40, or 50% of the nucleotide positions. As used herein, the term siRNA is meant to be equivalent to other terms used to describe nucleic acid molecules that are capable of mediating sequence specific RNAi, for example short interfering RNA (siRNA), double-stranded RNA (dsRNA), micro-RNA (miRNA), short hairpin RNA (shRNA), short interfering oligonucleotide, short interfering nucleic acid, short interfering modified oligonucleotide, chemically modified siRNA, post-transcriptional gene silencing RNA (ptgsRNA), and others. In addition, as used herein, the term RNAi is meant to be equivalent to other terms used to describe sequence specific RNA interference, such as post transcriptional gene silencing, translational inhibition, or epigenetics. For example, double-stranded oligonucleotides can be used to epigenetically silence genes at both the post-transcriptional level and the pre-transcriptional level. In a non-limiting example, epigenetic regulation of gene expression by siRNA molecules of the invention can result from siRNA mediated modification of chromatin structure or methylation pattern to alter gene expression (see, for example, Verdel et al., 2004, Science, 303, 672-676; Pal-Bhadra et al., 2004, Science, 303, 669-672; Allshire, 2002, Science, 297, 1818-1819; Volpe et al., 2002, Science, 297, 1833-1837; Jenuwein, 2002, Science, 297, 2215-2218; and Hall et al., 2002, Science, 297, 2232-2237).

It is contemplated that compounds and compositions of several embodiments provided herein can target ANGPTL3 by a dsRNA-mediated gene silencing or RNAi mechanism, including, e.g., "hairpin" or stem-loop double-stranded RNA effector molecules in which a single RNA strand with self-complementary sequences is capable of assuming a double-stranded conformation, or duplex dsRNA effector molecules comprising two separate strands of RNA. In various embodiments, the dsRNA consists entirely of ribonucleotides or consists of a mixture of ribonucleotides and deoxynucleotides, such as the RNA/DNA hybrids disclosed, for example, by WO 00/63364, filed Apr. 19, 2000, or U.S. Ser. No. 60/130,377, filed Apr. 21, 1999. The dsRNA or dsRNA effector molecule may be a single molecule with a region of self-complementarity such that nucleotides in one segment of the molecule base pair with nucleotides in another segment of the molecule. In various embodiments, a dsRNA that consists of a single molecule consists entirely of ribonucleotides or includes a region of ribonucleotides that is complementary to a region of

deoxyribonucleotides. Alternatively, the dsRNA may include two different strands that have a region of complementarity to each other.

In various embodiments, both strands consist entirely of ribonucleotides, one strand consists entirely of ribonucleotides and one strand consists entirely of deoxyribonucleotides, or one or both strands contain a mixture of ribonucleotides and deoxyribonucleotides. In certain embodiments, the regions of complementarity are at least 70, 80, 90, 95, 98, or 100% complementary to each other and to a target nucleic acid sequence. In certain embodiments, the region of the dsRNA that is present in a double-stranded conformation includes at least 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 50, 75, 100, 200, 500, 1000, 2000 or 5000 nucleotides or includes all of the nucleotides in a cDNA or other target nucleic acid sequence being represented in the dsRNA. In some embodiments, the dsRNA does not contain any single stranded regions, such as single stranded ends, or the dsRNA is a hairpin. In other embodiments, the dsRNA has one or more single stranded regions or overhangs. In certain embodiments, RNA/DNA hybrids include a DNA strand or region that is an antisense strand or region (e.g, has at least 70, 80, 90, 95, 98, or 100% complementarity to a target nucleic acid) and an RNA strand or region that is a sense strand or region (e.g, has at least 70, 80, 90, 95, 98, or 100% identity to a target nucleic acid), and vice versa.

In various embodiments, the RNA/DNA hybrid is made in vitro using enzymatic or chemical synthetic methods such as those described herein or those described in WO 00/63364, filed Apr. 19, 2000, or U.S. Ser. No. 60/130,377, filed Apr. 21, 1999. In other embodiments, a DNA strand synthesized in vitro is complexed with an RNA strand made in vivo or in vitro before, after, or concurrent with the transformation of the DNA strand into the cell. In yet other embodiments, the dsRNA is a single circular nucleic acid containing a sense and an antisense region, or the dsRNA includes a circular nucleic acid and either a second circular nucleic acid or a linear nucleic acid (see, for example, WO 00/63364, filed Apr. 19, 2000, or U.S. Ser. No. 60/130,377, filed Apr. 21, 1999.) Exemplary circular nucleic acids include lariat structures in which the free 5' phosphoryl group of a nucleotide becomes linked to the 2' hydroxyl group of another nucleotide in a loop back fashion.

In other embodiments, the dsRNA includes one or more modified nucleotides in which the 2' position in the sugar contains a halogen (such as fluorine group) or contains an alkoxy group (such as a methoxy group) which increases the half-life of the dsRNA in vitro or in vivo compared to the corresponding dsRNA in which the corresponding 2' position contains a hydrogen or an hydroxyl group. In yet other embodiments, the dsRNA includes one or more linkages between adjacent nucleotides other than a naturally-occurring phosphodiester linkage. Examples of such linkages include phosphoramidate, phosphorothioate, and phosphorodithioate linkages. The dsRNAs may also be chemically modified nucleic acid molecules as taught in U.S. Pat. No. 6,673,661. In other embodiments, the dsRNA contains one or two capped strands, as disclosed, for example, by WO 00/63364, filed Apr. 19, 2000, or U.S. Ser. No. 60/130,377, filed Apr. 21, 1999.

In other embodiments, the dsRNA can be any of the at least partially dsRNA molecules disclosed in WO 00/63364, as well as any of the dsRNA molecules described in U.S. Provisional Application 60/399,998; and U.S. Provisional Application 60/419,532, and PCT/US2003/033466, the teaching of which is hereby incorporated by reference. Any of the dsRNAs may be expressed in vitro or in vivo using the methods
 5 described herein or standard methods, such as those described in WO 00/63364.

Occupancy

In certain embodiments, antisense compounds are not expected to result in cleavage or the target nucleic acid via RNase H or to result in cleavage or sequestration through the RISC pathway. In certain such
 10 embodiments, antisense activity may result from occupancy, wherein the presence of the hybridized antisense compound disrupts the activity of the target nucleic acid. In certain such embodiments, the antisense compound may be uniformly modified or may comprise a mix of modifications and/or modified and unmodified nucleosides.

Target Nucleic Acids, Target Regions and Nucleotide Sequences

Nucleotide sequences that encode ANGPTL3 include, without limitation, the following: the human sequence as set forth in GenBank Accession No. NM_014495.2 (incorporated herein as SEQ ID NO: 1) or GenBank Accession No. NT_032977.9 nucleotides 33032001 to 33046000 (incorporated herein as SEQ ID NO: 2). It is understood that the sequence set forth in each SEQ ID NO in the Examples contained herein is independent of any modification to a sugar moiety, an internucleoside linkage, or a nucleobase. As such,
 15 antisense compounds defined by a SEQ ID NO can comprise, independently, one or more modifications to a sugar moiety, an internucleoside linkage, or a nucleobase. Antisense compounds described by Isis Number (Isis No) indicate a combination of nucleobase sequence and motif.

In certain embodiments, a target region is a structurally defined region of the target nucleic acid. For example, a target region can encompass a 3' UTR, a 5' UTR, an exon, an intron, an exon/intron junction, a
 25 coding region, a translation initiation region, translation termination region, or other defined nucleic acid region. The structurally defined regions for ANGPTL3 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 can 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.

In certain embodiments, a "target segment" is a smaller, sub-portion of a target region within a nucleic acid. For example, a target segment can be the sequence of nucleotides of a target nucleic acid to which one or more antisense compound is targeted. "5' target site" or "5' start site" refers to the 5'-most nucleotide of a target segment. "3' target site" or "3' stop site" refers to the 3'-most nucleotide of a target
 30 segment.

Targeting includes determination of at least one target segment to which an antisense compound hybridizes, such that a desired effect occurs. In certain embodiments, the desired effect is a reduction in mRNA target nucleic acid levels. In certain embodiments, the desired effect is reduction of levels of protein encoded by the target nucleic acid or a phenotypic change associated with the target nucleic acid.

5 A target region can contain one or more target segments. Multiple target segments within a target region can be overlapping. Alternatively, they can be non-overlapping. In certain embodiments, target segments within a target region are separated by no more than about 300 nucleotides. In certain
embodiments, target segments within a target region are separated by a number of nucleotides that is, is
about, is no more than, is no more than about, 250, 200, 150, 100, 90, 80, 70, 60, 50, 40, 30, 20, or 10
10 nucleotides on the target nucleic acid, or is a range defined by any two of the preceding values. In certain
embodiments, target segments within a target region are separated by no more than, or no more than about, 5
nucleotides on the target nucleic acid. In certain embodiments, target segments are contiguous.
Contemplated are target regions defined by a range having a starting nucleic acid that is any of the 5' target
sites or 3' target sites listed herein.

15 Suitable target segments can be found within a 5' UTR, a coding region, a 3' UTR, an intron, an
exon, or an exon/intron junction. Target segments containing a start codon or a stop codon are also suitable
target segments. A suitable target segment can specifically exclude a certain structurally defined region such
as the start codon or stop codon.

The determination of suitable target segments can include a comparison of the sequence of a target
20 nucleic acid to other sequences throughout the genome. For example, the BLAST algorithm can be used to
identify regions of similarity amongst different nucleic acids. This comparison can prevent the selection of
antisense compound sequences that can hybridize in a non-specific manner to sequences other than a selected
target nucleic acid (i.e., non-target or off-target sequences).

There can be variation in activity (e.g., as defined by percent reduction of target nucleic acid levels)
25 of the antisense compounds within an active target region. In certain embodiments, reductions in ANGPTL3
mRNA levels are indicative of inhibition of ANGPTL3 protein expression. Reductions in levels of an
ANGPTL3 protein are also indicative of inhibition of target mRNA expression. Further, phenotypic changes,
such as a reduction of the level of cholesterol, LDL, triglyceride, or glucose, can be indicative of inhibition of
ANGPTL3 mRNA and/or protein expression.

30 *Hybridization*

In some embodiments, hybridization occurs between an antisense compound disclosed herein and an
ANGPTL3 nucleic acid. The most common mechanism of hybridization involves hydrogen bonding (e.g.,
Watson-Crick, Hoogsteen or reversed Hoogsteen hydrogen bonding) between complementary nucleobases of
the nucleic acid molecules.

Hybridization can occur under varying conditions. Stringent conditions are sequence-dependent and are determined by the nature and composition of the nucleic acid molecules to be hybridized.

Methods of determining whether a sequence is specifically hybridizable to a target nucleic acid are well known in the art (Sambrook and Russell, *Molecular Cloning: A Laboratory Manual*, 3rd Ed., 2001). In certain embodiments, the antisense compounds provided herein are specifically hybridizable with an ANGPTL3 nucleic acid.

Complementarity

An antisense compound and a target nucleic acid are complementary to each other when a sufficient number of nucleobases of the antisense compound can hydrogen bond with the corresponding nucleobases of the target nucleic acid, such that a desired effect will occur (e.g., antisense inhibition of a target nucleic acid, such as an ANGPTL3 nucleic acid).

An antisense compound can hybridize over one or more segments of an ANGPTL3 nucleic acid such that intervening or adjacent segments are not involved in the hybridization event (e.g., a loop structure, mismatch or hairpin structure).

In certain embodiments, the antisense compounds provided herein, or a specified portion thereof, are, or are at least, 70%, 75%, 80%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% complementary to an ANGPTL3 nucleic acid, a target region, target segment, or specified portion thereof. In certain embodiments, the antisense compounds provided herein, or a specified portion thereof, are, or are at least, 70%, 75%, 80%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% complementary to the sequence of one or more of SEQ ID NOs: 1-2. Percent complementarity of an antisense compound with a target nucleic acid can be determined using routine methods.

For example, an antisense compound in which 18 of 20 nucleobases of the antisense compound are complementary to a target region, and would therefore specifically hybridize, would represent 90 percent complementarity. In this example, the remaining noncomplementary nucleobases can be clustered or interspersed with complementary nucleobases and need not be contiguous to each other or to complementary nucleobases. As such, an antisense compound which is 18 nucleobases in length having 4 (four) noncomplementary nucleobases which are flanked by two regions of complete complementarity with the target nucleic acid would have 77.8% overall complementarity with the target nucleic acid and would thus fall within the scope of the present invention. Percent complementarity of an antisense compound with a region of a target nucleic acid can be determined routinely using BLAST programs (basic local alignment search tools) and PowerBLAST programs known in the art (Altschul et al., *J. Mol. Biol.*, 1990, 215, 403 410; Zhang and Madden, *Genome Res.*, 1997, 7, 649 656). Percent homology, sequence identity or complementarity, can be determined by, for example, the Gap program (Wisconsin Sequence Analysis

Package, Version 8 for Unix, Genetics Computer Group, University Research Park, Madison Wis.), using default settings, which uses the algorithm of Smith and Waterman (Adv. Appl. Math., 1981, 2, 482-489).

In certain embodiments, the antisense compounds provided herein, or specified portions thereof, are fully complementary (i.e. 100% complementary) to a target nucleic acid, or specified portion thereof. For example, an antisense compound can be fully complementary to an ANGPTL3 nucleic acid, or a target region, or a target segment or target sequence thereof. As used herein, “fully complementary” means each nucleobase of an antisense compound is capable of precise base pairing with the corresponding nucleobases of a target nucleic acid. For example, a 20 nucleobase antisense compound is fully complementary to a target sequence that is 400 nucleobases long, so long as there is a corresponding 20 nucleobase portion of the target nucleic acid that is fully complementary to the antisense compound. Fully complementary can also be used in reference to a specified portion of the first and/or the second nucleic acid. For example, a 20 nucleobase portion of a 30 nucleobase antisense compound can be “fully complementary” to a target sequence that is 400 nucleobases long. The 20 nucleobase portion of the 30 nucleobase oligonucleotide is fully complementary to the target sequence if the target sequence has a corresponding 20 nucleobase portion wherein each nucleobase is complementary to the 20 nucleobase portion of the antisense compound. At the same time, the entire 30 nucleobase antisense compound can be fully complementary to the target sequence, depending on whether the remaining 10 nucleobases of the antisense compound are also complementary to the target sequence.

The location of a non-complementary nucleobase can be at the 5' end or 3' end of the antisense compound. Alternatively, the non-complementary nucleobase or nucleobases can be at an internal position of the antisense compound. When two or more non-complementary nucleobases are present, they can be either contiguous (i.e. linked) or non-contiguous. In one embodiment, a non-complementary nucleobase is located in the wing segment of a gapmer antisense oligonucleotide.

In certain embodiments, antisense compounds that are, or are up to 10, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleobases in length comprise no more than 4, no more than 3, no more than 2, or no more than 1 non-complementary nucleobase(s) relative to a target nucleic acid, such as an ANGPTL3 nucleic acid, or specified portion thereof.

In certain embodiments, antisense compounds that are, or are up to 10, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, or 30 nucleobases in length comprise no more than 6, no more than 5, no more than 4, no more than 3, no more than 2, or no more than 1 non-complementary nucleobase(s) relative to a target nucleic acid, such as an ANGPTL3 nucleic acid, or specified portion thereof.

The antisense compounds provided herein also include those which are complementary to a portion of a target nucleic acid. As used herein, “portion” refers to a defined number of contiguous (i.e. linked) nucleobases within a region or segment of a target nucleic acid. A “portion” can also refer to a defined number of contiguous nucleobases of an antisense compound. In certain embodiments, the antisense compounds, are complementary to at least an 8 nucleobase portion of a target segment. In certain

embodiments, the antisense compounds are complementary to at least a 10 nucleobase portion of a target segment. In certain embodiments, the antisense compounds are complementary to at least a 15 nucleobase portion of a target segment. Also contemplated are antisense compounds that are complementary to at least an 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, or more nucleobase portion of a target segment, or a range defined by any two of these values.

Identity

The antisense compounds provided herein can also have a defined percent identity to a particular nucleotide sequence, SEQ ID NO, or the sequence of a compound represented by a specific Isis number, or portion thereof. As used herein, an antisense compound is identical to the sequence disclosed herein if it has the same nucleobase pairing ability. For example, a RNA which contains uracil in place of thymidine in a disclosed DNA sequence would be considered identical to the DNA sequence since both uracil and thymidine pair with adenine. Shortened and lengthened versions of the antisense compounds described herein as well as compounds having non-identical bases relative to the antisense compounds provided herein also are contemplated. The non-identical bases can be adjacent to each other or dispersed throughout the antisense compound. Percent identity of an antisense compound is calculated according to the number of bases that have identical base pairing relative to the sequence to which it is being compared.

In certain embodiments, the antisense compounds, or portions thereof, are at least 70%, 75%, 80%, 85%, 90%, 95%, 96%, 97%, 98%, 99% or 100% identical to one or more of the antisense compounds or SEQ ID NOs, or a portion thereof, disclosed herein.

Modifications

A nucleoside is a base-sugar combination. The nucleobase (also known as base) portion of the nucleoside is normally a heterocyclic base moiety. Nucleotides are nucleosides that further include a phosphate group covalently linked to the sugar portion of the nucleoside. For those nucleosides that include a pentofuranosyl sugar, the phosphate group can be linked to the 2', 3' or 5' hydroxyl moiety of the sugar.

Oligonucleotides are formed through the covalent linkage of adjacent nucleosides to one another, to form a linear polymeric oligonucleotide. Within the oligonucleotide structure, the phosphate groups are commonly referred to as forming the internucleoside linkages of the oligonucleotide.

Modifications to antisense compounds encompass substitutions or changes to internucleoside linkages, sugar moieties, or nucleobases. Modified antisense compounds are often preferred over native forms because of desirable properties such as, for example, enhanced cellular uptake, enhanced affinity for nucleic acid target, increased stability in the presence of nucleases, or increased inhibitory activity.

Chemically modified nucleosides can also be employed to increase the binding affinity of a shortened or truncated antisense oligonucleotide for its target nucleic acid. Consequently, comparable results can often be obtained with shorter antisense compounds that have such chemically modified nucleosides.

Modified Internucleoside Linkages

The naturally occurring internucleoside linkage of RNA and DNA is a 3' to 5' phosphodiester linkage. Antisense compounds having one or more modified, i.e. non-naturally occurring, internucleoside linkages are often selected over antisense compounds having naturally occurring internucleoside linkages because of desirable properties such as, for example, enhanced cellular uptake, enhanced affinity for target nucleic acids, and increased stability in the presence of nucleases.

Oligonucleotides having modified internucleoside linkages include internucleoside linkages that retain a phosphorus atom as well as internucleoside linkages that do not have a phosphorus atom.

Representative phosphorus containing internucleoside linkages include, but are not limited to, phosphodiester, phosphotriester, methylphosphonate, phosphoramidate, and phosphorothioate. Methods of preparation of phosphorous-containing and non-phosphorous-containing linkages are well known.

In certain embodiments, antisense compounds targeted to an ANGPTL3 nucleic acid comprise one or more modified internucleoside linkages. In certain embodiments, the modified internucleoside linkages are phosphorothioate linkages. In certain embodiments, each internucleoside linkage of an antisense compound is a phosphorothioate internucleoside linkage.

Modified Sugar Moieties

Antisense compounds of the invention can optionally contain one or more nucleosides wherein the sugar group has been modified. Such sugar modified nucleosides may impart enhanced nuclease stability, increased binding affinity, or some other beneficial biological property to the antisense compounds. In certain embodiments, nucleosides comprise chemically modified ribofuranose ring moieties. Examples of chemically modified ribofuranose rings include without limitation, addition of substituent groups (including 5' and 2' substituent groups, bridging of non-geminal ring atoms to form bicyclic nucleic acids (BNA), replacement of the ribosyl ring oxygen atom with S, N(R), or C(R₁)(R₂) (R, R₁ and R₂ are each independently H, C₁-C₁₂ alkyl or a protecting group) and combinations thereof. Examples of chemically modified sugars include 2'-F-5'-methyl substituted nucleoside (see PCT International Application WO 2008/101157 Published on 8/21/08 for other disclosed 5',2'-bis substituted nucleosides) or replacement of the ribosyl ring oxygen atom with S with 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 BNA (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).

Examples of nucleosides having modified sugar moieties include without limitation nucleosides comprising 5'-vinyl, 5'-methyl (*R* or *S*), 4'-S, 2'-F, 2'-OCH₃, 2'-OCH₂CH₃, 2'-OCH₂CH₂F and 2'-O(CH₂)₂OCH₃ substituent groups. The substituent at the 2' position can also be selected from allyl, amino, azido, thio, O-allyl, O-C₁-C₁₀ alkyl, OCF₃, OCH₂F, O(CH₂)₂SCH₃, O(CH₂)₂-O-N(R_m)(R_n), O-CH₂-C(=O)-

$N(R_m)(R_n)$, and $O-CH_2-C(=O)-N(R_1)-(CH_2)_2-N(R_m)(R_n)$, where each R_1 , R_m and R_n is, independently, H or substituted or unsubstituted C_1-C_{10} alkyl.

As used herein, “bicyclic nucleosides” refer to modified nucleosides comprising a bicyclic sugar moiety. Examples of bicyclic nucleic acids (BNAs) include without limitation nucleosides comprising a bridge between the 4' and the 2' ribosyl ring atoms. In certain embodiments, antisense compounds provided herein include one or more BNA nucleosides wherein the bridge comprises one of the formulas: 4'-(CH_2)-O-2' (LNA); 4'-(CH_2)-S-2'; 4'-(CH_2)₂-O-2' (ENA); 4'-CH(CH_3)-O-2' and 4'-CH(CH_2OCH_3)-O-2' (and analogs thereof see U.S. Patent 7,399,845, issued on July 15, 2008); 4'-C(CH_3)(CH_3)-O-2' (and analogs thereof see PCT/US2008/068922 published as WO/2009/006478, published January 8, 2009); 4'- CH_2 -N(OCH_3)-2' (and analogs thereof see PCT/US2008/064591 published as WO/2008/150729, published December 11, 2008); 4'- CH_2 -O-N(CH_3)-2' (see published U.S. Patent Application US2004-0171570, published September 2, 2004); 4'- CH_2 -N(R)-O-2', wherein R is H, C_1-C_{12} alkyl, or a protecting group (see U.S. Patent 7,427,672, issued on September 23, 2008); 4'- CH_2 -C(H)(CH_3)-2' (see Chattopadhyaya *et al.*, *J. Org. Chem.*, 2009, 74, 118-134); and 4'- CH_2 -C(= CH_2)-2' (and analogs thereof see PCT/US2008/066154 published as WO 2008/154401, published on December 8, 2008).

Further bicyclic nucleosides have been reported in published literature (see for example: Srivastava *et al.*, *J. Am. Chem. Soc.*, 2007, 129(26) 8362-8379; Frieden *et al.*, *Nucleic Acids Research*, 2003, 31, 6365-6372; Elayadi *et al.*, *Curr. Opinion Inven. Drugs*, 2001, 2, 558-561; Braasch *et al.*, *Chem. Biol.*, 2001, 8, 1-7; Orum *et al.*, *Curr. Opinion Mol. Ther.*, 2001, 3, 239-243; Wahlestedt *et al.*, *Proc. Natl. Acad. Sci. U. S. A.*, 2000, 97, 5633-5638; Singh *et al.*, *Chem. Commun.*, 1998, 4, 455-456; Koshkin *et al.*, *Tetrahedron*, 1998, 54, 3607-3630; Kumar *et al.*, *Bioorg. Med. Chem. Lett.*, 1998, 8, 2219-2222; Singh *et al.*, *J. Org. Chem.*, 1998, 63, 10035-10039; U.S. Patents Nos.: 7,399,845; 7,053,207; 7,034,133; 6,794,499; 6,770,748; 6,670,461; 6,525,191; 6,268,490; U.S. Patent Publication Nos.: US2008-0039618; US2007-0287831; US2004-0171570; U.S. Patent Applications, Serial Nos.: 12/129,154; 61/099,844; 61/097,787; 61/086,231; 61/056,564; 61/026,998; 61/026,995; 60/989,574; International applications WO 2007/134181; WO 2005/021570; WO 2004/106356; WO 94/14226; and PCT International Applications Nos.: PCT/US2008/068922; PCT/US-2008/066154; and PCT/US2008/064591). Each of the foregoing bicyclic nucleosides can be prepared having one or more stereochemical sugar configurations including for example α -L-ribofuranose and β -D-ribofuranose (see PCT international application PCT/DK98/00393, published on March 25, 1999 as WO 99/14226).

As used herein, “monocyclic nucleosides” refer to nucleosides comprising modified sugar moieties that are not bicyclic sugar moieties. In certain embodiments, the sugar moiety, or sugar moiety analogue, of a nucleoside may be modified or substituted at any position.

As used herein, “4'-2' bicyclic nucleoside” or “4' to 2' bicyclic nucleoside” refers to a bicyclic nucleoside comprising a furanose ring comprising a bridge connecting two carbon atoms of the furanose ring connects the 2' carbon atom and the 4' carbon atom of the sugar ring.

In certain embodiments, bicyclic sugar moieties of BNA nucleosides include, but are not limited to, compounds having at least one bridge between the 4' and the 2' carbon atoms of the pentofuranosyl sugar moiety including without limitation, bridges comprising 1 or 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, C_1-C_{12} alkyl, substituted C_1-C_{12} alkyl, C_2-C_{12} alkenyl, substituted C_2-C_{12} alkenyl, C_2-C_{12} alkynyl, substituted C_2-C_{12} alkynyl, C_5-C_{20} aryl, substituted C_5-C_{20} aryl, heterocycle radical, substituted heterocycle radical, heteroaryl, substituted heteroaryl, C_5-C_7 alicyclic radical, substituted C_5-C_7 alicyclic radical, halogen, OJ_1 , NJ_1J_2 , SJ_1 , N_3 , $COOJ_1$, acyl ($C(=O)-H$), substituted acyl, CN, sulfonyl ($S(=O)_2-J_1$), or sulfoxyl ($S(=O)-J_1$); and

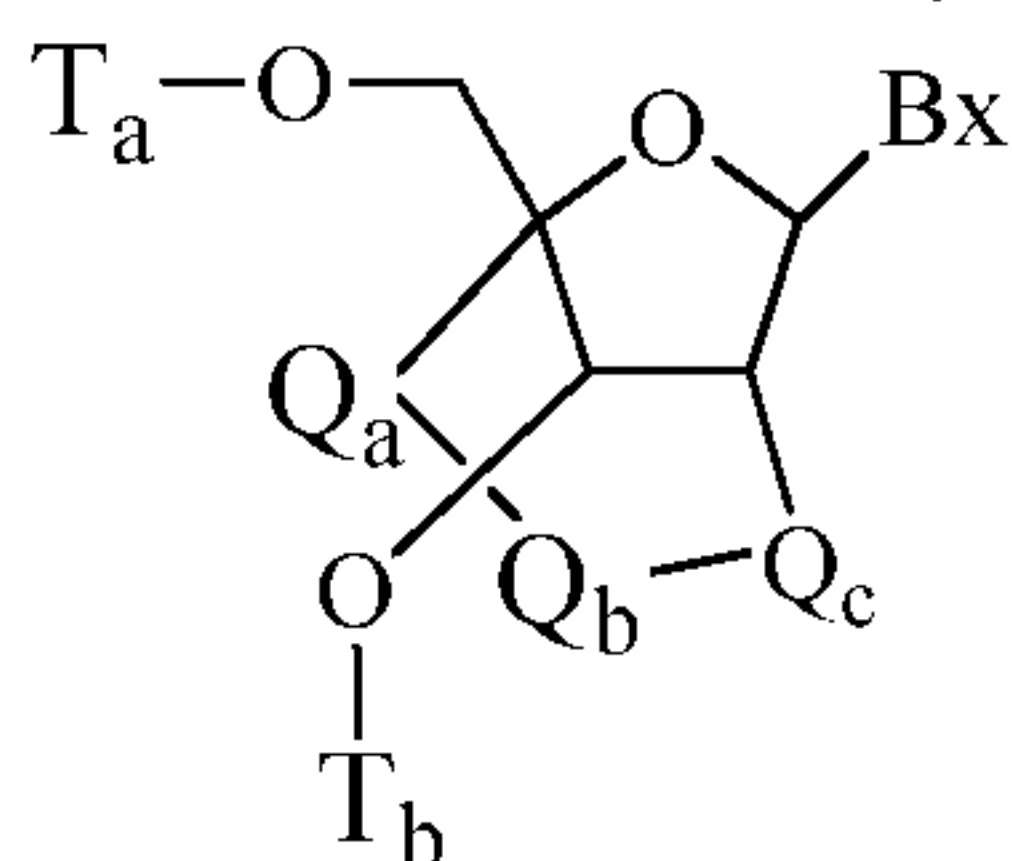
each J_1 and J_2 is, independently, H, C_1-C_{12} alkyl, substituted C_1-C_{12} alkyl, C_2-C_{12} alkenyl, substituted C_2-C_{12} alkenyl, C_2-C_{12} alkynyl, substituted C_2-C_{12} alkynyl, C_5-C_{20} aryl, substituted C_5-C_{20} aryl, acyl ($C(=O)-H$), substituted acyl, a heterocycle radical, a substituted heterocycle radical, C_1-C_{12} aminoalkyl, substituted C_1-C_{12} aminoalkyl or a protecting group.

In certain embodiments, the bridge of a bicyclic sugar moiety is $-[C(R_a)(R_b)]_n-$, $-[C(R_a)(R_b)]_n-O-$, $-C(R_aR_b)-N(R)-O-$ or $-C(R_aR_b)-O-N(R)-$. In certain embodiments, the bridge is 4'-CH₂-2', 4'-(CH₂)₂-2', 4'-(CH₂)₃-2', 4'-CH₂-O-2', 4'-(CH₂)₂-O-2', 4'-CH₂-O-N(R)-2' and 4'-CH₂-N(R)-O-2' wherein each R is, independently, H, a protecting group or C_1-C_{12} alkyl.

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

In certain embodiments, bicyclic nucleosides include those having a 4' to 2' bridge wherein such bridges include without limitation, α -L-4'-(CH₂)-O-2', β -D-4'-CH₂-O-2', 4'-(CH₂)₂-O-2', 4'-CH₂-O-N(R)-2', 4'-CH₂-N(R)-O-2', 4'-CH(CH₃)-O-2', 4'-CH₂-S-2', 4'-CH₂-N(R)-2', 4'-CH₂-CH(CH₃)-2', and 4'-(CH₂)₃-2', wherein R is H, a protecting group or C_1-C_{12} alkyl.

In certain embodiment, bicyclic nucleosides have the formula:



wherein:

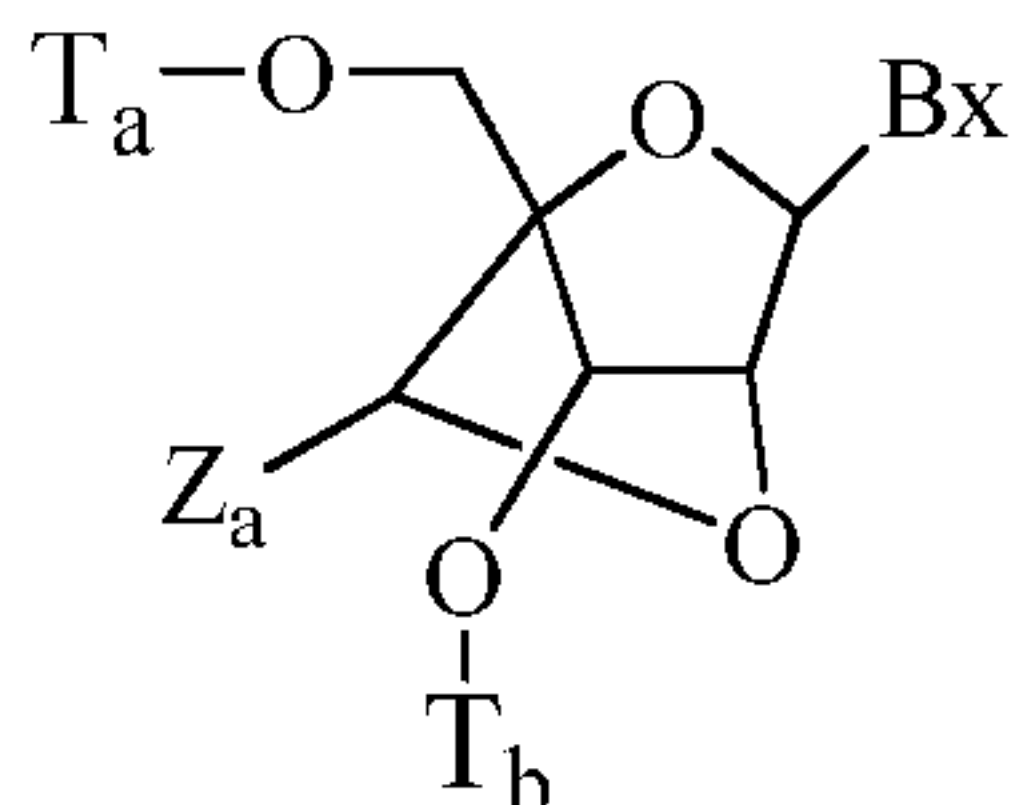
Bx is a heterocyclic base moiety;

$-Q_a-Q_b-Q_c-$ is $-CH_2-N(R_c)-CH_2-$, $-C(=O)-N(R_c)-CH_2-$, $-CH_2-O-N(R_c)-$, $-CH_2-N(R_c)-O-$ or $-N(R_c)-O-CH_2-$;

R_c is C_1 - C_{12} alkyl or an amino protecting group; and

T_a and T_b are each, independently H, a hydroxyl protecting group, a conjugate group, a reactive phosphorus group, a phosphorus moiety or a covalent attachment to a support medium.

In certain embodiments, bicyclic nucleosides have the formula:



5

wherein:

Bx is a heterocyclic base moiety;

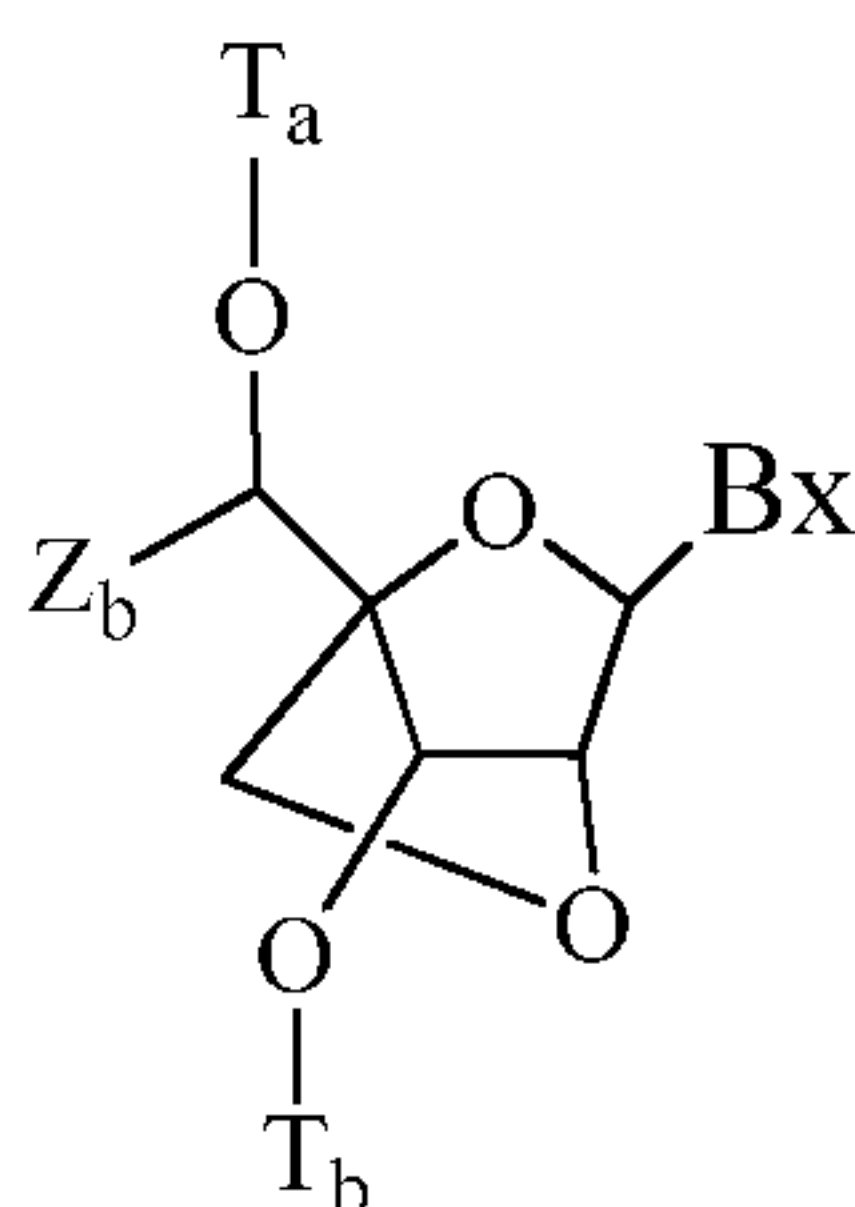
T_a and T_b are each, independently H, a hydroxyl protecting group, a conjugate group, a reactive phosphorus group, a phosphorus moiety or a covalent attachment to a support medium;

10 Z_a is C_1 - C_6 alkyl, C_2 - C_6 alkenyl, C_2 - C_6 alkynyl, substituted C_1 - C_6 alkyl, substituted C_2 - C_6 alkenyl, substituted C_2 - C_6 alkynyl, acyl, substituted acyl, substituted amide, thiol or substituted thiol.

In one embodiment, each of the substituted groups, is, independently, mono or poly substituted with substituent groups independently selected from halogen, oxo, hydroxyl, OJ_c , NJ_cJ_d , SJ_c , N_3 , $OC(=X)J_c$, and $NJ_eC(=X)NJ_cJ_d$, wherein each J_c , J_d and J_e is, independently, H, C_1 - C_6 alkyl, or substituted C_1 - C_6 alkyl and X is O or NJ_c .

15

In certain embodiments, bicyclic nucleosides have the formula:



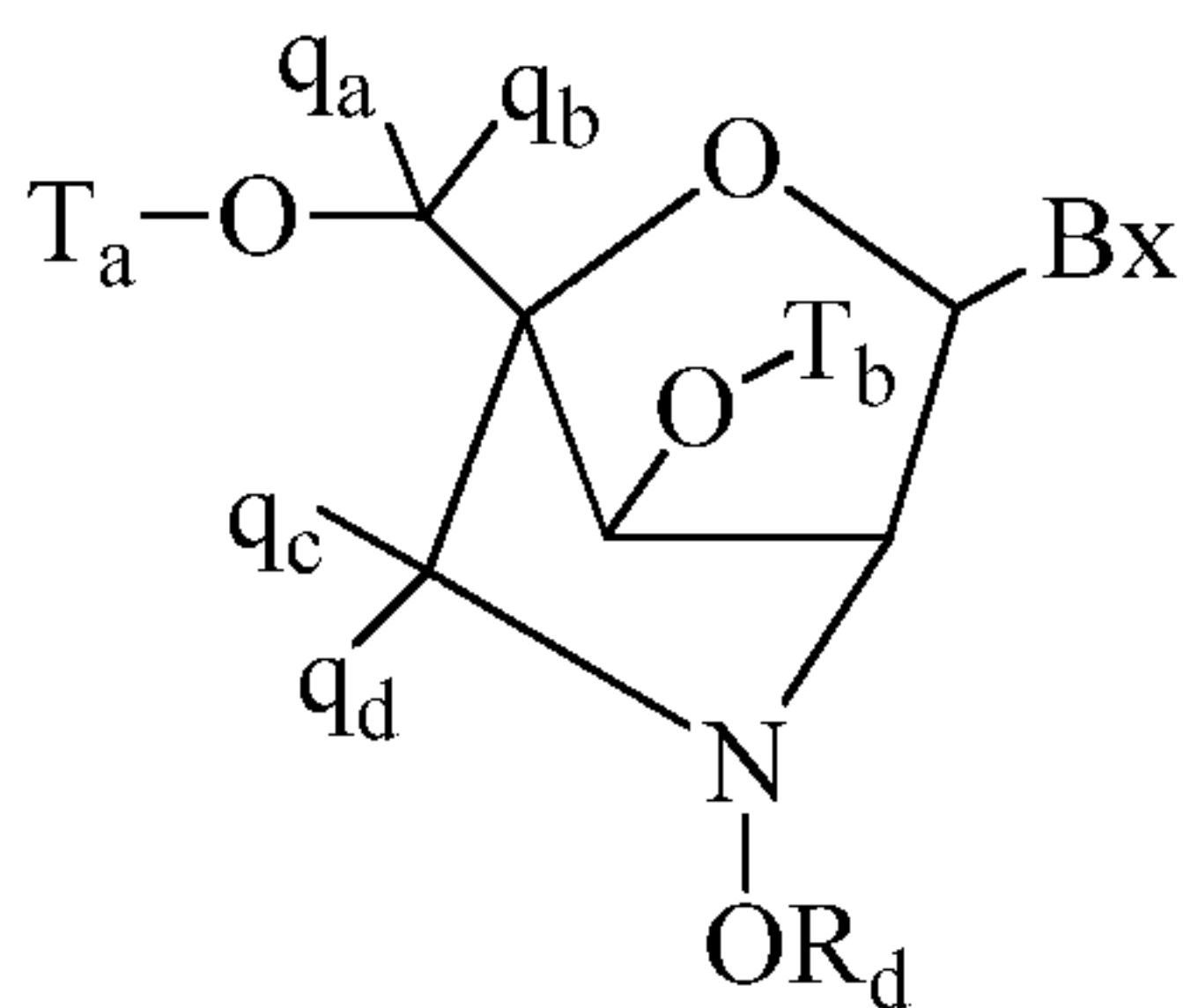
wherein:

Bx is a heterocyclic base moiety;

20 T_a and T_b are each, independently H, a hydroxyl protecting group, a conjugate group, a reactive phosphorus group, a phosphorus moiety or a covalent attachment to a support medium;

Z_b is C_1 - C_6 alkyl, C_2 - C_6 alkenyl, C_2 - C_6 alkynyl, substituted C_1 - C_6 alkyl, substituted C_2 - C_6 alkenyl, substituted C_2 - C_6 alkynyl or substituted acyl ($C(=O)-$).

In certain embodiments, bicyclic nucleosides have the formula:



wherein:

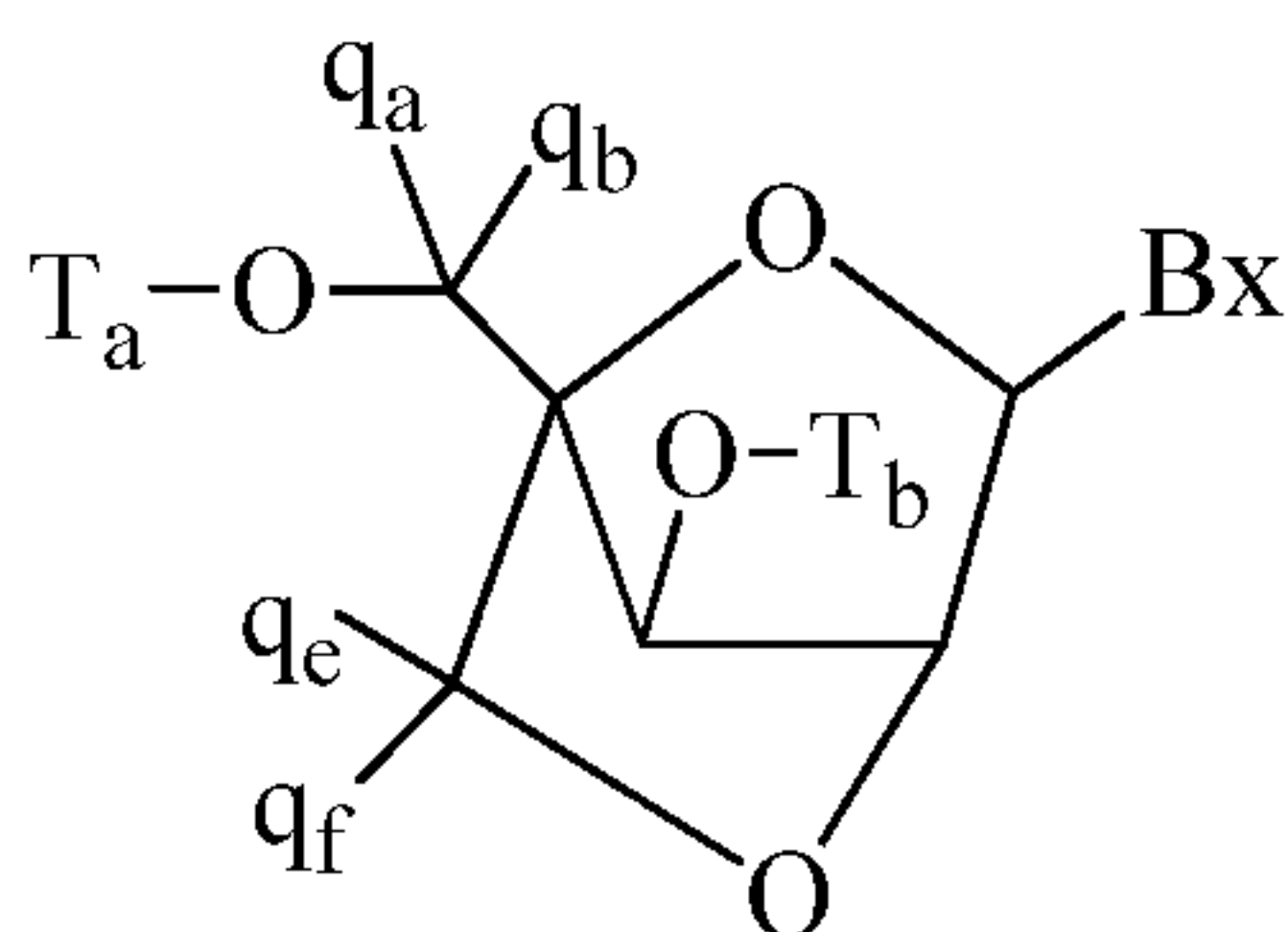
Bx is a heterocyclic base moiety;

T_a and T_b are each, independently H, a hydroxyl protecting group, a conjugate group, a reactive
5 phosphorus group, a phosphorus moiety or a covalent attachment to a support medium;

R_d is C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₂-C₆ alkenyl, substituted C₂-C₆ alkenyl, C₂-C₆ alkynyl or
substituted C₂-C₆ alkynyl;

each q_a, q_b, q_c and q_d is, independently, H, halogen, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₂-C₆
alkenyl, substituted C₂-C₆ alkenyl, C₂-C₆ alkynyl or substituted C₂-C₆ alkynyl, C₁-C₆ alkoxy, substituted C₁-
10 C₆ alkoxy, acyl, substituted acyl, C₁-C₆ aminoalkyl or substituted C₁-C₆ aminoalkyl;

In certain embodiments, bicyclic nucleosides have the formula:



wherein:

Bx is a heterocyclic base moiety;

15 T_a and T_b are each, independently H, a hydroxyl protecting group, a conjugate group, a reactive
phosphorus group, a phosphorus moiety or a covalent attachment to a support medium;

q_a, q_b, q_e and q_f are each, independently, hydrogen, halogen, C₁-C₁₂ alkyl, substituted C₁-C₁₂ alkyl, C₂-
C₁₂ alkenyl, substituted C₂-C₁₂ alkenyl, C₂-C₁₂ alkynyl, substituted C₂-C₁₂ alkynyl, C₁-C₁₂ alkoxy, substituted
C₁-C₁₂ alkoxy, OJ_j, SJ_j, SOJ_j, SO₂J_j, NJ_jJ_k, N₃, CN, C(=O)OJ_j, C(=O)NJ_jJ_k, C(=O)J_j, O-C(=O)NJ_jJ_k,
20 N(H)C(=NH)NJ_jJ_k, N(H)C(=O)NJ_jJ_k or N(H)C(=S)NJ_jJ_k;

or q_e and q_f together are =C(q_g)(q_h);

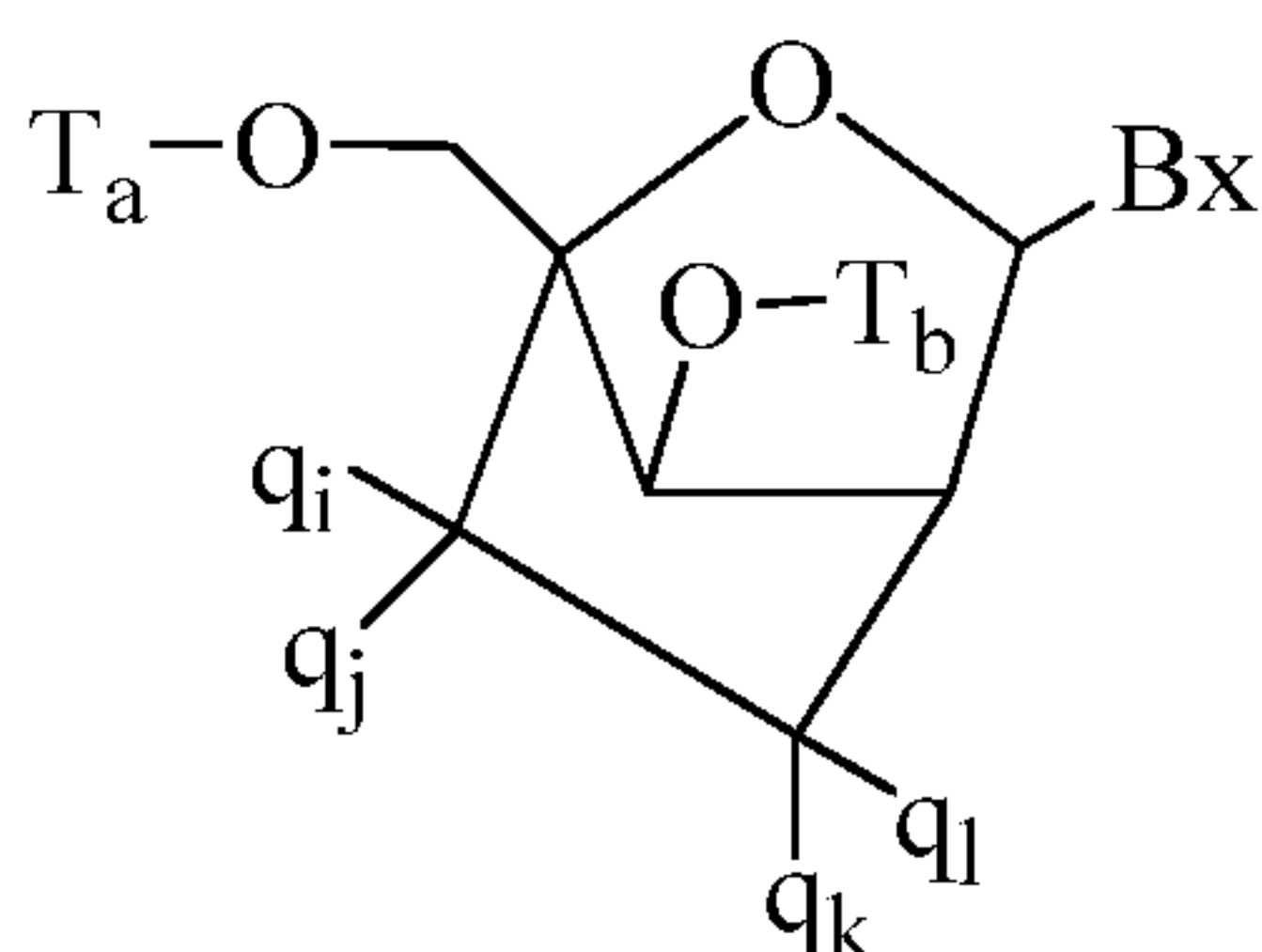
q_g and q_h are each, independently, H, halogen, C₁-C₁₂ alkyl or substituted C₁-C₁₂ alkyl.

The synthesis and preparation of adenine, cytosine, guanine, 5-methyl-cytosine, thymine and uracil
bicyclic nucleosides having a 4'-CH₂-O-2' bridge, along with their oligomerization, and nucleic acid
25 recognition properties have been described (Koshkin et al., *Tetrahedron*, 1998, 54, 3607-3630). The
synthesis of bicyclic nucleosides has also been described in WO 98/39352 and WO 99/14226.

Analogues of various bicyclic nucleosides that have 4' to 2' bridging groups such as 4'-CH₂-O-2' and 4'-CH₂-S-2', have also been prepared (Kumar *et al.*, *Bioorg. Med. Chem. Lett.*, 1998, 8, 2219-2222).

Preparation of oligodeoxyribonucleotide duplexes comprising bicyclic nucleosides for use as substrates for nucleic acid polymerases has also been described (Wengel *et al.*, WO 99/14226). Furthermore, synthesis of 2'-amino-BNA, a novel conformationally restricted high-affinity oligonucleotide analog has been described in the art (Singh *et al.*, *J. Org. Chem.*, 1998, 63, 10035-10039). In addition, 2'-amino- and 2'-methylamino-BNA's have been prepared and the thermal stability of their duplexes with complementary RNA and DNA strands has been previously reported.

In certain embodiments, bicyclic nucleosides have the formula:



wherein:

Bx is a heterocyclic base moiety;

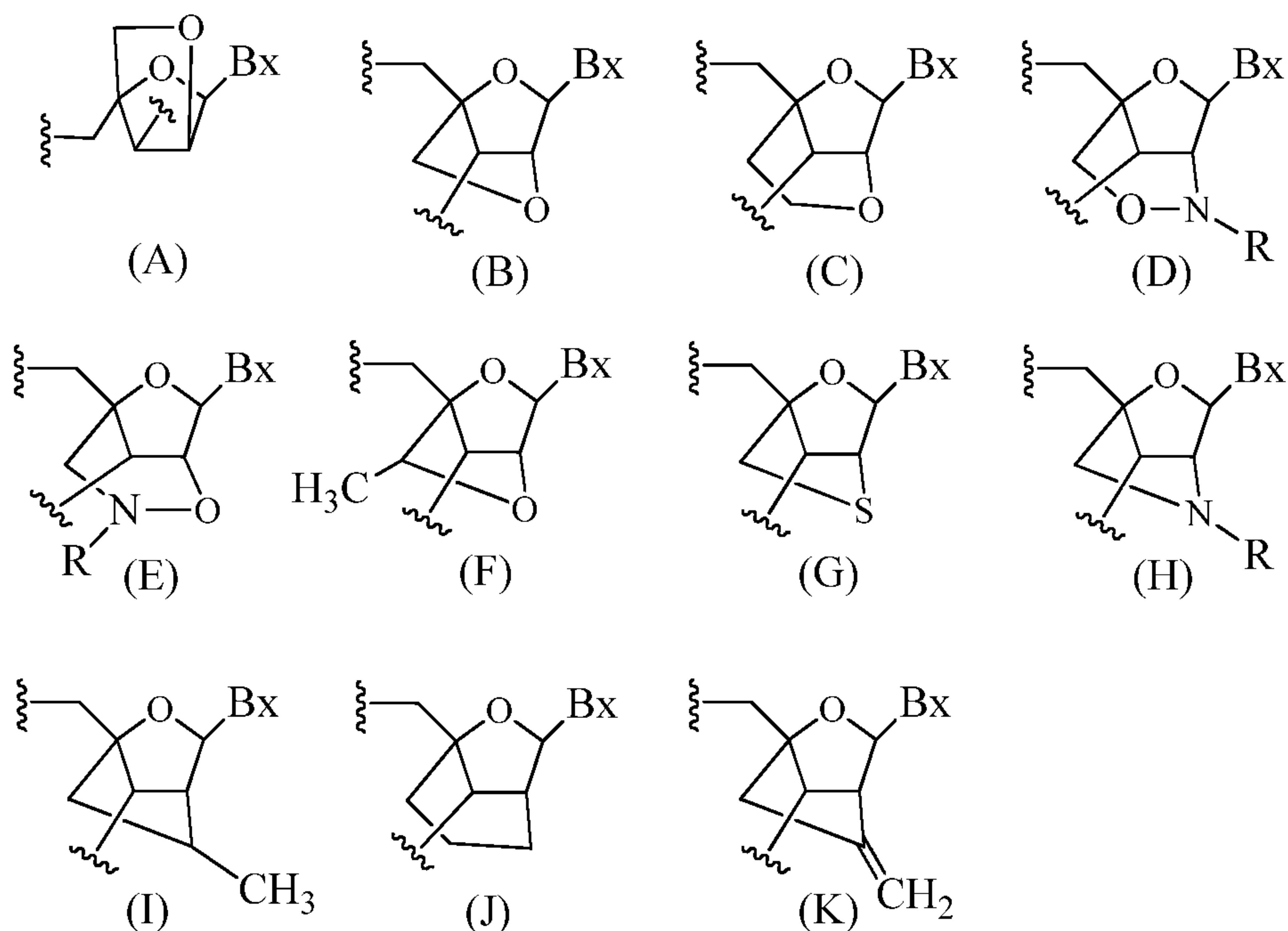
T_a and T_b are each, independently H, a hydroxyl protecting group, a conjugate group, a reactive phosphorus group, a phosphorus moiety or a covalent attachment to a support medium;

each q_i, q_j, q_k and q_l is, independently, H, halogen, C₁-C₁₂ alkyl, substituted C₁-C₁₂ alkyl, C₂-C₁₂ alkenyl, substituted C₂-C₁₂ alkenyl, C₂-C₁₂ alkynyl, substituted C₂-C₁₂ alkynyl, C₁-C₁₂ alkoxy, substituted C₁-C₁₂ alkoxy, OJ_j, SJ_j, SOJ_j, SO₂J_j, NJ_jJ_k, N₃, CN, C(=O)OJ_j, C(=O)NJ_jJ_k, C(=O)J_j, O-C(=O)NJ_jJ_k, N(H)C(=NH)NJ_jJ_k, N(H)C(=O)NJ_jJ_k or N(H)C(=S)NJ_jJ_k; and

q_i and q_j or q_l and q_k together are =C(q_g)(q_h), wherein q_g and q_h are each, independently, H, halogen, C₁-C₁₂ alkyl or substituted C₁-C₁₂ alkyl.

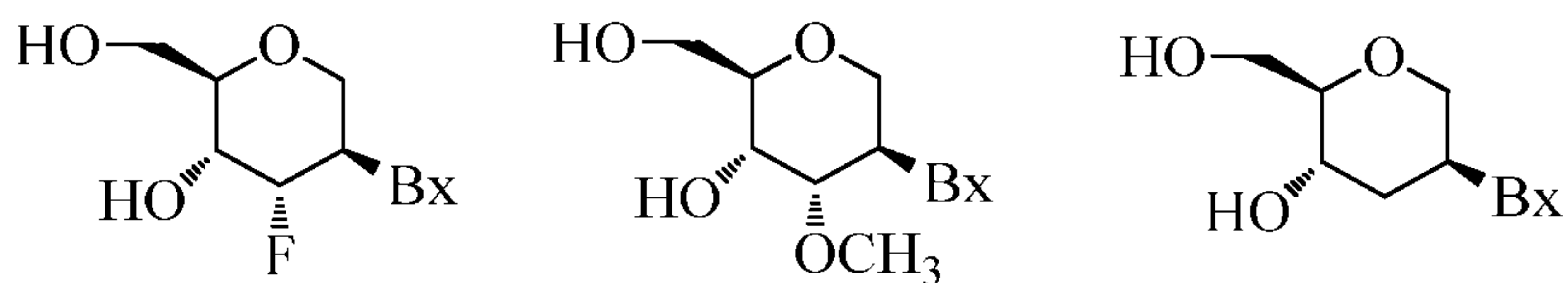
One carbocyclic bicyclic nucleoside having a 4'-(CH₂)₃-2' bridge and the alkenyl analog bridge 4'-CH=CH-CH₂-2' have been described (Frier *et al.*, *Nucleic Acids Research*, 1997, 25(22), 4429-4443 and Albaek *et al.*, *J. Org. Chem.*, 2006, 71, 7731-7740). The synthesis and preparation of carbocyclic bicyclic nucleosides along with their oligomerization and biochemical studies have also been described (Srivastava *et al.*, *J. Am. Chem. Soc.* 2007, 129(26), 8362-8379).

In certain embodiments, 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, (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, (J) propylene carbocyclic (4'-(CH₂)₃-2') BNA, and (K) vinyl BNA as depicted below.

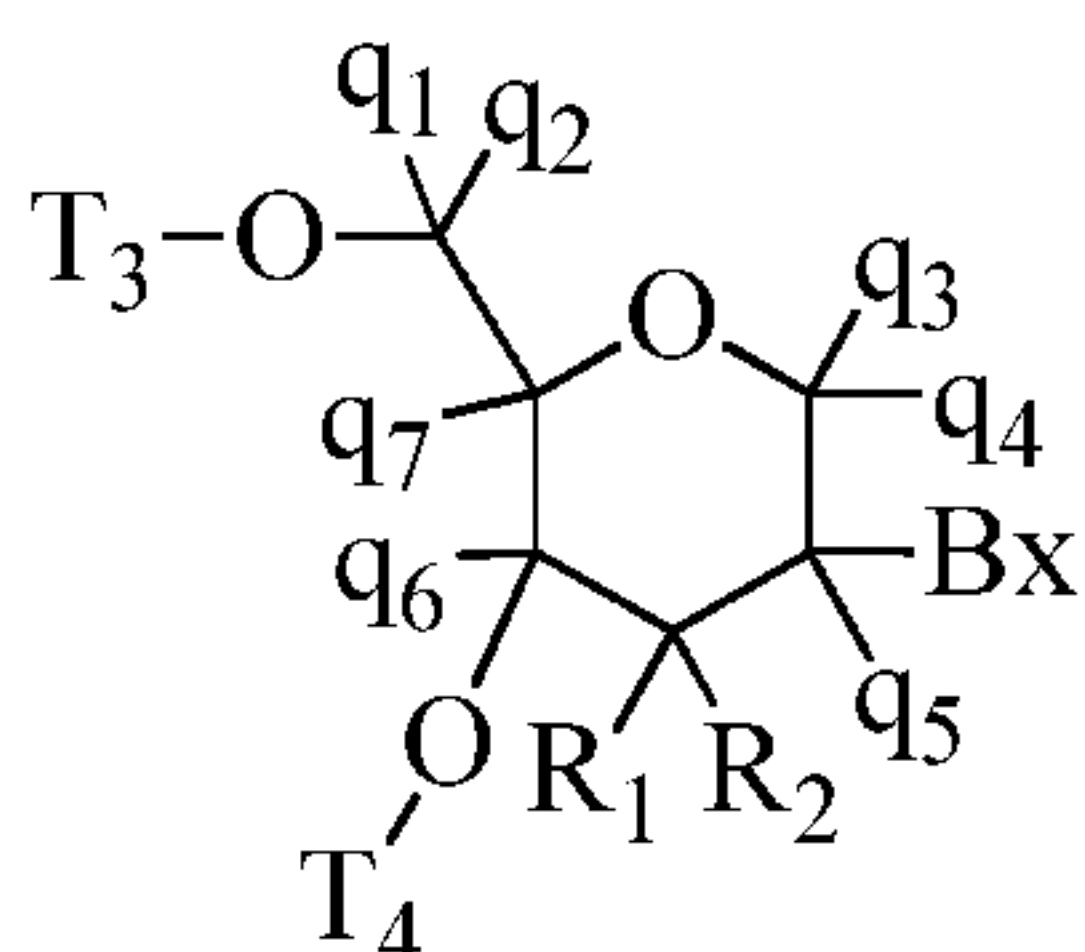


wherein Bx is the base moiety and R is, independently, H, a protecting group, C₁-C₆ alkyl or C₁-C₆ alkoxy.

As used herein, the term “modified tetrahydropyran nucleoside” or “modified THP nucleoside” means a nucleoside having a six-membered tetrahydropyran “sugar” substituted for the pentofuranosyl residue in normal nucleosides and can be referred to as a sugar surrogate. Modified THP nucleosides include, but are not limited to, what is referred to in the art as hexitol nucleic acid (HNA), anitol nucleic acid (ANA), manitol nucleic acid (MNA) (see Leumann, *Bioorg. Med. Chem.*, 2002, 10, 841-854) or fluoro HNA (F-HNA) having a tetrahydropyranyl ring system as illustrated below.



In certain embodiment, sugar surrogates are selected having the formula:



wherein:

Bx is a heterocyclic base moiety;

T₃ and T₄ are each, independently, an internucleoside linking group linking the tetrahydropyran

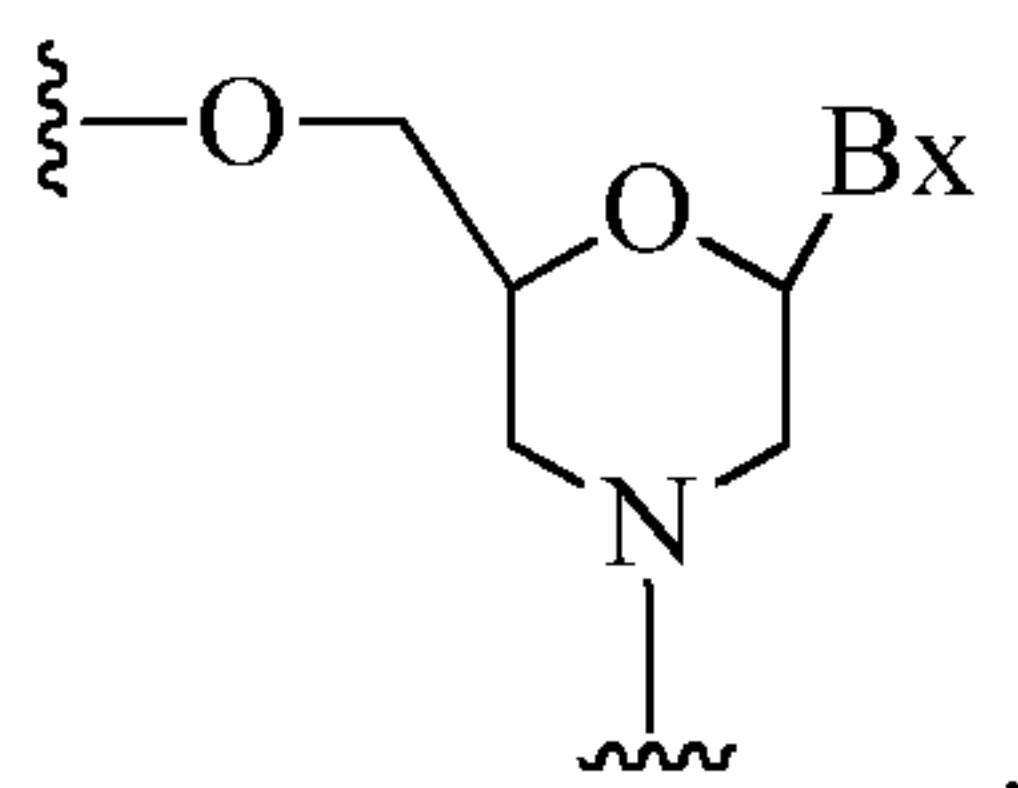
nucleoside analog to the oligomeric compound or one of T₃ and T₄ is an internucleoside linking group linking the tetrahydropyran nucleoside analog to an oligomeric compound or oligonucleotide and the other of T₃ and T₄ is H, a hydroxyl protecting group, a linked conjugate group or a 5' or 3'-terminal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

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

one of R₁ and R₂ is hydrogen and the other is selected from 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.

In certain embodiments, 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 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.

In certain embodiments, sugar surrogates comprise rings having more than 5 atoms and more than one heteroatom. For example nucleosides comprising morpholino sugar moieties and their use in oligomeric compounds has been reported (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 formula:

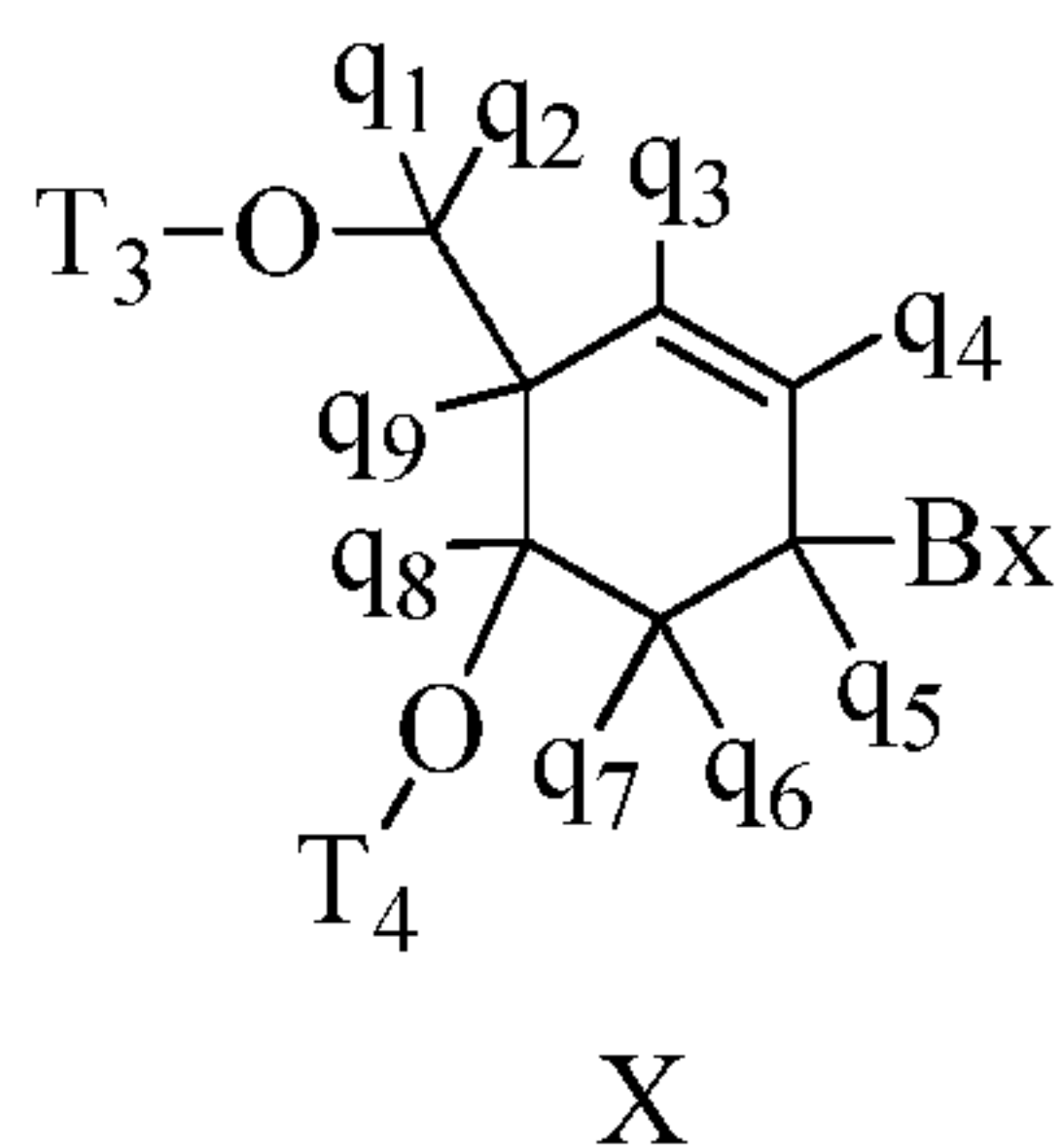


In certain embodiments, morpholinos may be modified, for example by adding or altering various substituent groups from the above morpholino structure. Such sugar surrogates are referred to herein as “modified morpholinos.”

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, antisense compounds comprise one or more modified cyclohexenyl nucleosides, which is a nucleoside having a six-membered cyclohexenyl in place of the pentofuranosyl

residue in naturally occurring nucleosides. Modified cyclohexenyl nucleosides include, but are not limited to those described in the art (see for example commonly owned, published PCT Application WO 2010/036696, published on April 10, 2010, Robeyns *et al.*, *J. Am. Chem. Soc.*, 2008, 130(6), 1979-1984; Horváth *et al.*, *Tetrahedron Letters*, 2007, 48, 3621-3623; Nauwelaerts *et al.*, *J. Am. Chem. Soc.*, 2007, 129(30), 9340-9348; Gu *et al.*, *Nucleosides, Nucleotides & Nucleic Acids*, 2005, 24(5-7), 993-998; Nauwelaerts *et al.*, *Nucleic Acids Research*, 2005, 33(8), 2452-2463; Robeyns *et al.*, *Acta Crystallographica, Section F: Structural Biology and Crystallization Communications*, 2005, F61(6), 585-586; Gu *et al.*, *Tetrahedron*, 2004, 60(9), 2111-2123; Gu *et al.*, *Oligonucleotides*, 2003, 13(6), 479-489; Wang *et al.*, *J. Org. Chem.*, 2003, 68, 4499-4505; Verbeure *et al.*, *Nucleic Acids Research*, 2001, 29(24), 4941-4947; Wang *et al.*, *J. Org. Chem.*, 2001, 66, 8478-82; Wang *et al.*, *Nucleosides, Nucleotides & Nucleic Acids*, 2001, 20(4-7), 785-788; Wang *et al.*, *J. Am. Chem.*, 2000, 122, 8595-8602; Published PCT application, WO 06/047842; and Published PCT Application WO 01/049687; the text of each is incorporated by reference herein, in their entirety). Certain modified cyclohexenyl nucleosides have Formula X.



wherein independently for each of said at least one cyclohexenyl nucleoside analog of Formula X:

Bx is a heterocyclic base moiety;

15 T₃ and T₄ are each, independently, an internucleoside linking group linking the cyclohexenyl nucleoside analog to an antisense compound or one of T₃ and T₄ is an internucleoside linking group linking the tetrahydropyran nucleoside analog to an 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; and

20 q₁, q₂, q₃, q₄, q₅, q₆, q₇, q₈ and q₉ are each, independently, H, C₁-C₆ alkyl, substituted C₁-C₆ alkyl, C₂-C₆ alkenyl, substituted C₂-C₆ alkenyl, C₂-C₆ alkynyl, substituted C₂-C₆ alkynyl or other sugar substituent group.

Many other monocyclic, bicyclic and tricyclic ring systems are known in the art and are suitable as sugar surrogates that can be used to modify nucleosides for incorporation into oligomeric compounds as provided herein (see for example review article: Leumann, Christian J. *Bioorg. & Med. Chem.*, 2002, 10, 841-854). Such ring systems can undergo various additional substitutions to further enhance their activity.

As used herein, "2'-modified sugar" means a furanosyl sugar modified at the 2' position. In certain embodiments, such modifications include substituents selected from: a halide, including, but not limited to substituted and unsubstituted alkoxy, substituted and unsubstituted thioalkyl, substituted and unsubstituted

amino alkyl, substituted and unsubstituted alkyl, substituted and unsubstituted allyl, and substituted and unsubstituted alkynyl. In certain embodiments, 2' modifications are selected from substituents including, but not limited to: $O[(CH_2)_nO]_mCH_3$, $O(CH_2)_nNH_2$, $O(CH_2)_nCH_3$, $O(CH_2)_nF$, $O(CH_2)_nONH_2$, $OCH_2C(=O)N(H)CH_3$, and $O(CH_2)_nON[(CH_2)_nCH_3]_2$, where n and m are from 1 to about 10. Other 2'-

5 substituent groups can also be selected from: C_1 - C_{12} alkyl, substituted alkyl, alkenyl, alkynyl, alkaryl, aralkyl, O-alkaryl or O-aralkyl, SH, SCH₃, OCN, Cl, Br, CN, F, CF₃, OCF₃, SOCH₃, SO₂CH₃, ONO₂, NO₂, N₃, NH₂, heterocycloalkyl, heterocycloalkaryl, aminoalkylamino, polyalkylamino, substituted silyl, an RNA cleaving group, a reporter group, an intercalator, a group for improving pharmacokinetic properties, or a group for improving the pharmacodynamic properties of an antisense compound, and other substituents having similar

10 properties. In certain embodiments, modified nucleosides comprise a 2'-MOE side chain (Baker *et al.*, *J. Biol. Chem.*, 1997, 272, 11944-12000). Such 2'-MOE substitution have been described as having improved binding affinity compared to unmodified nucleosides and to other modified nucleosides, such as 2'-O-methyl, O-propyl, and O-aminopropyl. Oligonucleotides having the 2'-MOE substituent also have been shown to be antisense inhibitors of gene expression with promising features for *in vivo* use (Martin, *Helv. Chim. Acta*, 1995, 78, 486-504; Altmann *et al.*, *Chimia*, 1996, 50, 168-176; Altmann *et al.*, *Biochem. Soc. Trans.*, 1996, 24, 630-637; and Altmann *et al.*, *Nucleosides Nucleotides*, 1997, 16, 917-926).

As used herein, "2'-modified" or "2'-substituted" refers to a nucleoside comprising a sugar comprising a substituent at the 2' position other than H or OH. 2'-modified nucleosides, include, but are not limited to, bicyclic nucleosides wherein the bridge connecting two carbon atoms of the sugar ring connects

20 the 2' carbon and another carbon of the sugar ring; and nucleosides with non-bridging 2' substituents, such as allyl, amino, azido, thio, O-allyl, O- C_1 - C_{10} alkyl, -OCF₃, O-(CH₂)₂-O-CH₃, 2'-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 or substituted or unsubstituted C_1 - C_{10} alkyl. 2'-modified nucleosides may further comprise other modifications, for example at other positions of the sugar and/or at the nucleobase.

25 As used herein, "2'-F" refers to a nucleoside comprising a sugar comprising a fluoro group at the 2' position of the sugar ring.

As used herein, "2'-OMe" or "2'-OCH₃", "2'-O-methyl" or "2'-methoxy" each refers to a nucleoside comprising a sugar comprising an -OCH₃ group at the 2' position of the sugar ring.

30 As used herein, "MOE" or "2'-MOE" or "2'-OCH₂CH₂OCH₃" or "2'-O-methoxyethyl" each refers to a nucleoside comprising a sugar comprising a -OCH₂CH₂OCH₃ group at the 2' position of the sugar ring.

Methods for the preparations of modified sugars are well known to those skilled in the art. Some representative U.S. patents that teach the preparation of such modified sugars include without limitation, U.S.: 4,981,957; 5,118,800; 5,319,080; 5,359,044; 5,393,878; 5,446,137; 5,466,786; 5,514,785; 5,519,134; 5,567,811; 5,576,427; 5,591,722; 5,597,909; 5,610,300; 5,627,053; 5,639,873; 5,646,265; 5,670,633;

35 5,700,920; 5,792,847 and 6,600,032 and International Application PCT/US2005/019219, filed June 2, 2005

and published as WO 2005/121371 on December 22, 2005, and each of which is herein incorporated by reference in its entirety.

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

In nucleotides having modified sugar moieties, the nucleobase moieties (natural, modified or a combination thereof) are maintained for hybridization with an appropriate nucleic acid target.

In certain embodiments, antisense compounds comprise one or more nucleosides having modified sugar moieties. In certain embodiments, the modified sugar moiety is 2'-MOE. In certain embodiments, the 2'-MOE modified nucleosides are arranged in a gapmer motif. In certain embodiments, the modified sugar moiety is a bicyclic nucleoside having a (4'-CH(CH₃)-O-2') bridging group. In certain embodiments, the (4'-CH(CH₃)-O-2') modified nucleosides are arranged throughout the wings of a gapmer motif.

Modified Nucleobases

Nucleobase (or base) modifications or substitutions are structurally distinguishable from, yet functionally interchangeable with, naturally occurring or synthetic unmodified nucleobases. Both natural and modified nucleobases are capable of participating in hydrogen bonding. Such nucleobase modifications can impart nuclease stability, binding affinity or some other beneficial biological property to antisense compounds. Modified nucleobases include synthetic and natural nucleobases such as, for example, 5-methylcytosine (5-me-C). Certain nucleobase substitutions, including 5-methylcytosine substitutions, are particularly useful for increasing the binding affinity of an antisense compound for a target nucleic acid. For example, 5-methylcytosine substitutions have been shown to increase nucleic acid duplex stability by 0.6-1.2°C (Sanghvi, Y.S., Crooke, S.T. and Lebleu, B., eds., *Antisense Research and Applications*, CRC Press, Boca Raton, 1993, pp. 276-278).

Additional modified nucleobases include 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≡C-CH₃) 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 and 3-deazaguanine and 3-deazaadenine.

Heterocyclic base moieties can 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.

Nucleobases that are particularly useful for increasing the binding affinity of antisense compounds include 5-

substituted pyrimidines, 6-azapyrimidines and N-2, N-6 and O-6 substituted purines, including 2 aminopropyladenine, 5-propynyluracil and 5-propynylcytosine.

In certain embodiments, antisense compounds targeted to an ANGPTL3 nucleic acid comprise one or more modified nucleobases. In certain embodiments, shortened or gap-widened antisense oligonucleotides
5 targeted to an ANGPTL3 nucleic acid comprise one or more modified nucleobases. In certain embodiments, the modified nucleobase is 5-methylcytosine. In certain embodiments, each cytosine is a 5-methylcytosine.

Compositions and Methods for Formulating Pharmaceutical Compositions

Antisense oligonucleotides can be admixed with pharmaceutically acceptable active or inert
10 substance for the preparation of pharmaceutical compositions or formulations. Compositions and methods for the formulation of pharmaceutical compositions are dependent upon a number of criteria, including, but not limited to, route of administration, extent of disease, or dose to be administered.

Antisense compound targeted to an ANGPTL3 nucleic acid can be utilized in pharmaceutical compositions by combining the antisense compound with a suitable pharmaceutically acceptable diluent or
15 carrier. A pharmaceutically acceptable diluent includes phosphate-buffered saline (PBS). PBS is a diluent suitable for use in compositions to be delivered parenterally. Accordingly, in one embodiment, employed in the methods described herein is a pharmaceutical composition comprising an antisense compound targeted to an ANGPTL3 nucleic acid and a pharmaceutically acceptable diluent. In certain embodiments, the pharmaceutically acceptable diluent is PBS. In certain embodiments, the antisense compound is an antisense
20 oligonucleotide.

Pharmaceutical compositions comprising antisense compounds encompass any pharmaceutically acceptable salts, esters, or salts of such esters, or any other 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
25 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 antisense compound which are cleaved by endogenous nucleases within the body, to form the active antisense
30 compound.

Conjugated Antisense Compounds

Antisense compounds can be covalently linked to one or more moieties or conjugates which enhance the activity, cellular distribution or cellular uptake of the resulting antisense oligonucleotides. Typical
35 conjugate groups include cholesterol moieties and lipid moieties. Additional conjugate groups include

carbohydrates, phospholipids, biotin, phenazine, folate, phenanthridine, anthraquinone, acridine, fluoresceins, rhodamines, coumarins, and dyes.

Antisense compounds can also be modified to have one or more stabilizing groups that are generally attached to one or both termini of antisense compounds to enhance properties such as, for example, nuclease stability. Included in stabilizing groups are cap structures. These terminal modifications protect the antisense compound having terminal nucleic acids from exonuclease degradation, and can help in delivery and/or localization within a cell. The cap can be present at the 5'-terminus (5'-cap), or at the 3'-terminus (3'-cap), or can be present on both termini. Cap structures are well known in the art and include, for example, inverted deoxy abasic caps. Further 3' and 5'-stabilizing groups that can be used to cap one or both ends of an antisense compound to impart nuclease stability include those disclosed in WO 03/004602 published on January 16, 2003.

Cell culture and antisense compounds treatment

The effects of antisense compounds on the level, activity or expression of ANGPTL3 nucleic acids can be tested in vitro in a variety of cell types. Cell types used for such analyses are available from commercial vendors (e.g. American Type Culture Collection, Manassus, VA; Zen-Bio, Inc., Research Triangle Park, NC; Clonetics Corporation, Walkersville, MD) and cells are cultured according to the vendor's instructions using commercially available reagents (e.g. Invitrogen Life Technologies, Carlsbad, CA). Illustrative cell types include, but are not limited to, HepG2 cells, Hep3B cells, Huh7 (hepatocellular carcinoma) cells, primary hepatocytes, A549 cells, GM04281 fibroblasts and LLC-MK2 cells.

In vitro testing of antisense oligonucleotides

Described herein are methods for treatment of cells with antisense oligonucleotides, which can be modified appropriately for treatment with other antisense compounds.

In general, cells are treated with antisense oligonucleotides when the cells reach approximately 60-80% confluence in culture.

One reagent commonly used to introduce antisense oligonucleotides into cultured cells includes the cationic lipid transfection reagent LIPOFECTIN® (Invitrogen, Carlsbad, CA). Antisense oligonucleotides are mixed with LIPOFECTIN® in OPTI-MEM® 1 (Invitrogen, Carlsbad, CA) to achieve the desired final concentration of antisense oligonucleotide and a LIPOFECTIN® concentration that typically ranges 2 to 12 ug/mL per 100 nM antisense oligonucleotide.

Another reagent used to introduce antisense oligonucleotides into cultured cells includes LIPOFECTAMINE 2000® (Invitrogen, Carlsbad, CA). Antisense oligonucleotide is mixed with LIPOFECTAMINE 2000® in OPTI-MEM® 1 reduced serum medium (Invitrogen, Carlsbad, CA) to achieve the desired concentration of antisense oligonucleotide and a LIPOFECTAMINE® concentration that typically ranges 2 to 12 ug/mL per 100 nM antisense oligonucleotide.

Another reagent used to introduce antisense oligonucleotides into cultured cells includes Cytofectin® (Invitrogen, Carlsbad, CA). Antisense oligonucleotide is mixed with Cytofectin® in OPTI-MEM® 1 reduced serum medium (Invitrogen, Carlsbad, CA) to achieve the desired concentration of antisense oligonucleotide and a Cytofectin® concentration that typically ranges 2 to 12 ug/mL per 100 nM antisense oligonucleotide.

Another reagent used to introduce antisense oligonucleotides into cultured cells includes Oligofectamine™ (Invitrogen Life Technologies, Carlsbad, CA). Antisense oligonucleotide is mixed with Oligofectamine™ in Opti-MEM™-1 reduced serum medium (Invitrogen Life Technologies, Carlsbad, CA) to achieve the desired concentration of oligonucleotide with an Oligofectamine™ to oligonucleotide ratio of approximately 0.2 to 0.8 µL per 100 nM.

Another reagent used to introduce antisense oligonucleotides into cultured cells includes FuGENE 6 (Roche Diagnostics Corp., Indianapolis, IN). Antisense oligomeric compound was mixed with FuGENE 6 in 1 mL of serum-free RPMI to achieve the desired concentration of oligonucleotide with a FuGENE 6 to oligomeric compound ratio of 1 to 4 µL of FuGENE 6 per 100 nM.

Another technique used to introduce antisense oligonucleotides into cultured cells includes electroporation (Sambrook and Russell, Molecular Cloning: A Laboratory Manual, 3rd Ed., 2001).

Cells are treated with antisense oligonucleotides by routine methods. Cells are typically harvested 16-24 hours after antisense oligonucleotide treatment, at which time RNA or protein levels of target nucleic acids are measured by methods known in the art and described herein. In general, when treatments are performed in multiple replicates, the data are presented as the average of the replicate treatments.

The concentration of antisense oligonucleotide used varies from cell line to cell line. Methods to determine the optimal antisense oligonucleotide concentration for a particular cell line are well known in the art. Antisense oligonucleotides are typically used at concentrations ranging from 1 nM to 300 nM when transfected with LIPOFECTAMINE2000®, Lipofectin or Cytofectin. Antisense oligonucleotides are used at higher concentrations ranging from 625 to 20,000 nM when transfected using electroporation.

RNA Isolation

RNA analysis can be performed on total cellular RNA or poly(A)+ mRNA. Methods of RNA isolation are well known in the art (Sambrook and Russell, Molecular Cloning: A Laboratory Manual, 3rd Ed., 2001). RNA is prepared using methods well known in the art, for example, using the TRIZOL® Reagent (Invitrogen, Carlsbad, CA) according to the manufacturer's recommended protocols.

Analysis of inhibition of target levels or expression

Inhibition of levels or expression of an ANGPTL3 nucleic acid can be assayed in a variety of ways known in the art (Sambrook and Russell, Molecular Cloning: A Laboratory Manual, 3rd Ed., 2001). For example, target nucleic acid levels can be quantitated by, e.g., Northern blot analysis, competitive polymerase chain reaction (PCR), or quantitative real-time PCR. RNA analysis can be performed on total cellular RNA

or poly(A)+ mRNA. Methods of RNA isolation are well known in the art. Northern blot analysis is also routine in the art. Quantitative real-time PCR can be conveniently accomplished using the commercially available ABI PRISM® 7600, 7700, or 7900 Sequence Detection System, available from PE-Applied Biosystems, Foster City, CA and used according to manufacturer's instructions.

5

Quantitative Real-Time PCR Analysis of Target RNA Levels

Quantitation of target RNA levels can be accomplished by quantitative real-time PCR using the ABI PRISM® 7600, 7700, or 7900 Sequence Detection System (PE-Applied Biosystems, Foster City, CA) according to manufacturer's instructions. Methods of quantitative real-time PCR are well known in the art.

10

Prior to real-time PCR, the isolated RNA is subjected to a reverse transcriptase (RT) reaction, which produces complementary DNA (cDNA) that is then used as the substrate for the real-time PCR amplification. The RT and real-time PCR reactions are performed sequentially in the same sample well. RT and real-time PCR reagents are obtained from Invitrogen (Carlsbad, CA). RT and real-time-PCR reactions are carried out by methods well known to those skilled in the art.

15

Gene (or RNA) target quantities obtained by real time PCR can be normalized using either the expression level of a gene whose expression is constant, such as cyclophilin A or GADPH or by quantifying total RNA using RIBOGREEN® (Life Technologies™, Inc. Carlsbad, CA). Cyclophilin A or GADPH expression can be quantified by real time PCR, by being run simultaneously with the target, multiplexing, or separately. Total RNA can be quantified using RIBOGREEN® RNA quantification reagent. Methods of RNA quantification by RIBOGREEN® are taught in Jones, L.J., et al, (Analytical Biochemistry, 1998, 265, 368-374). A CYTOFLUOR® 4000 instrument (PE Applied Biosystems) can be used to measure RIBOGREEN® fluorescence.

20

Methods for designing real-time PCR probes and primers are well known in the art, and can include the use of software such as PRIMER EXPRESS® Software (Applied Biosystems, Foster City, CA). Probes and primers used in real-time PCR were designed to hybridize to ANGPTL3 specific sequences and are disclosed in the Examples below. The target specific PCR probes can have FAM covalently linked to the 5' end and TAMRA or MGB covalently linked to the 3' end, where FAM is the fluorescent dye and TAMRA or MGB is the quencher dye.

25

30

Analysis of Protein Levels

Antisense inhibition of ANGPTL3 nucleic acids can be assessed by measuring ANGPTL3 protein levels. Protein levels of ANGPTL3 can be evaluated or quantitated in a variety of ways well known in the art, such as immunoprecipitation, Western blot analysis (immunoblotting), enzyme-linked immunosorbent assay (ELISA), quantitative protein assays, protein activity assays (for example, caspase activity assays), immunohistochemistry, immunocytochemistry or fluorescence-activated cell sorting (FACS) (Sambrook and Russell, Molecular Cloning: A Laboratory Manual, 3rd Ed., 2001). Antibodies directed to a target can be

35

identified and obtained from a variety of sources, such as the MSRS catalog of antibodies (Aerie Corporation, Birmingham, MI), or can be prepared via conventional monoclonal or polyclonal antibody generation methods well known in the art.

5 *In vivo testing of antisense compounds*

Antisense compounds, for example, antisense oligonucleotides, are tested in animals to assess their ability to inhibit expression of ANGPTL3 and produce phenotypic changes. Testing can be performed in normal animals, or in experimental disease models. For administration to animals, antisense oligonucleotides are formulated in a pharmaceutically acceptable diluent, such as phosphate-buffered saline. Administration includes parenteral routes of administration. Following a period of treatment with antisense oligonucleotides, RNA is isolated from tissue and changes in ANGPTL3 nucleic acid expression are measured. Changes in ANGPTL3 protein levels are also measured.

Certain Indications

15 In certain embodiments, provided herein are methods of treating an individual comprising administering one or more pharmaceutical compositions as described herein. In certain embodiments, the individual has a metabolic disease and/or cardiovascular disease. In certain embodiments, the individual has hypercholesterolemia (e.g., familial homozygous hypercholesterolemia (HoFH), familial heterozygous hypercholesterolemia (HeFH)), dyslipidemia, hypertriglyceridemia (e.g., heterozygous LPL deficiency, 20 homozygous LPL deficiency), coronary artery disease (CAD), familial chylomicronemia syndrome (FCS), hyperlipoproteinemia Type IV), lipodystrophy, hyperlipidemia (e.g., combined hyperlipidemia, familial combined hyperlipidemia (FCHL)), metabolic syndrome, non-alcoholic fatty liver disease (NAFLD), nonalcoholic steatohepatitis (NASH), diabetes (e.g., Type 2 diabetes), vascular wall thickening, high blood pressure (e.g., pulmonary arterial hypertension), sclerosis (e.g., atherosclerosis, systemic sclerosis, 25 progressive skin sclerosis and proliferative obliterative vasculopathy such as digital ulcers and pulmonary vascular involvement).

In certain embodiments, the compounds targeted to ANGPTL3 described herein modulate lipid and/or energy metabolism in an animal. In certain embodiments, the compounds targeted to ANGPTL3 described herein modulate physiological markers or phenotypes of hypercholesterolemia, dyslipidemia, 30 hypertriglyceridemia, metabolic syndrome, NAFLD, NASH and/or diabetes. For example, administration of the compounds to animals can modulate one or more of VLDL, non-esterified fatty acids (NEFA), LDL, cholesterol, triglyceride, glucose, insulin or ANGPTL3 levels. In certain embodiments, the modulation of the physiological markers or phenotypes can be associated with inhibition of ANGPTL3 by the compounds.

In certain embodiments, the compounds targeted to ANGPTL3 described herein reduce and/or 35 prevent one or more of hepatic TG accumulation (i.e. hepatic steatosis), atherosclerosis, vascular wall thickening (e.g., arterial intima-media thickening), hypercholesterolemia (e.g., familial homozygous

hypercholesterolemia (HoFH), familial heterozygous hypercholesterolemia (HeFH)), dyslipidemia, hypertriglyceridemia (e.g., heterozygous LPL deficiency, homozygous LPL deficiency, familial chylomicronemia syndrome (FCS), hyperlipoproteinemia Type IV), lipodystrophy, hyperlipidemia (e.g., combined hyperlipidemia, familial combined hyperlipidemia (FCHL)), metabolic syndrome, non-alcoholic fatty liver disease (NAFLD), nonalcoholic steatohepatitis (NASH), diabetes (e.g., Type 2 diabetes), high blood pressure and sclerosis. In certain embodiments, the compounds targeted to ANGPTL3 described herein improve insulin sensitivity.

In certain embodiments, administration of an antisense compound targeted to an ANGPTL3 nucleic acid as described herein results in reduction of ANGPTL3 expression by about at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95% or at least 99%, or a range defined by any two of these values.

In certain embodiments, administration of an antisense compound targeted to an ANGPTL3 nucleic acid as described herein results in reduction of one or more of triglycerides, LDL-cholesterol, non-HDL cholesterol, VLDL-cholesterol, total cholesterol, ApoB and ApoC-III by about at least 5%, at least 10%, at least 15%, at least 20%, at least 25%, at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95% or at least 99%, or a range defined by any two of these values.

In certain embodiments, pharmaceutical compositions comprising an antisense compound targeted to ANGPTL3 are used for the preparation of a medicament for treating a patient suffering from, or susceptible to, a metabolic disease or cardiovascular disease. In certain embodiments, pharmaceutical compositions comprising an antisense compound targeted to ANGPTL3 are used in the preparation of a medicament for treating a patient suffering from, or susceptible to, one or more of hypercholesterolemia (e.g., familial homozygous hypercholesterolemia (HoFH), familial heterozygous hypercholesterolemia (HeFH)), dyslipidemia, hypertriglyceridemia (e.g., familial chylomicronemia syndrome (FCS), hyperlipoproteinemia Type IV), lipodystrophy, hyperlipidemia (e.g., combined hyperlipidemia, familial combined hyperlipidemia (FCHL)), metabolic syndrome, non-alcoholic fatty liver disease (NAFLD), nonalcoholic steatohepatitis (NASH), diabetes (e.g., Type 2 diabetes) vascular wall thickening, high blood pressure and sclerosis.

Administration

In certain embodiments, the compounds and compositions as described herein are administered parenterally.

In certain embodiments, parenteral administration is by infusion. Infusion can be chronic or continuous or short or intermittent. In certain embodiments, infused pharmaceutical agents are delivered with a pump.

In certain embodiments, parenteral administration is by injection. The injection can be delivered with a syringe or a pump. In certain embodiments, the injection is a bolus injection. In certain embodiments, the injection is administered directly to a tissue or organ. In certain embodiments, the injection is subcutaneous.

5

Certain Combination Therapies

In certain embodiments, a first agent comprising the modified oligonucleotide disclosed herein is co-administered with one or more secondary agents. In certain embodiments, such second agents are designed to treat the same disease, disorder or condition as the first agent described herein. In certain embodiments, such second agents are designed to treat a different disease, disorder, or condition as the first agent described herein. In certain embodiments, such second agents are designed to treat an undesired side effect of one or more pharmaceutical compositions as described herein. In certain embodiments, second agents are co-administered with the first agent to treat an undesired effect of the first agent. In certain embodiments, second agents are co-administered with the first agent to produce a combinational effect. In certain
10
15
embodiments, second agents are co-administered with the first agent to produce a synergistic effect.

In certain embodiments, a first agent and one or more second agents are administered at the same time. In certain embodiments, the first agent and one or more second agents are administered at different times. In certain embodiments, the first agent and one or more second agents are prepared together in a single pharmaceutical formulation. In certain embodiments, the first agent and one or more second agents are
20
prepared separately.

In certain embodiments, second agents include, but are not limited to a glucose-lowering agent or a lipid-lowering agent. The glucose lowering agent can include, but is not limited to, a therapeutic lifestyle change, PPAR agonist, a dipeptidyl peptidase (IV) inhibitor, a GLP-1 analog, insulin or an insulin analog, an insulin secretagogue, a SGLT2 inhibitor, a human amylin analog, a biguanide, an alpha-glucosidase inhibitor, or a combination thereof. The glucose-lowering agent can include, but is not limited to metformin, sulfonylurea, rosiglitazone, meglitinide, thiazolidinedione, alpha-glucosidase inhibitor or a combination thereof. The sulfonylurea can be acetohexamide, chlorpropamide, tolbutamide, tolazamide, glimepiride, a glipizide, a glyburide, or a gliclazide. The meglitinide can be nateglinide or repaglinide. The thiazolidinedione can be pioglitazone or rosiglitazone. The alpha-glucosidase can be acarbose or miglitol. In
25
30
certain embodiments the lipid lowering therapy can include, but is not limited to, a therapeutic lifestyle change, niacin, HMG-CoA reductase inhibitor, cholesterol absorption inhibitor, MTP inhibitor (e.g., a small molecule, polypeptide, antibody or antisense compound targeted to MTP), fibrate, PCSK9 inhibitor (e.g., PCSK9 antibodies, polypeptides, small molecules nucleic acid compounds targeting PCSK9), CETP inhibitor (e.g., small molecules such as torcetrapib and anacetrapib, polypeptides, antibodies or nucleic acid
35
compounds targeted to CETP), apoC-III inhibitor (e.g., a small molecule, polypeptide, antibody or nucleic acid compounds targeted to apoC-III), apoB inhibitor (e.g., a small molecule, polypeptide, antibody or

nucleic acid compounds targeted to apoB), beneficial oils rich in omega-3 fatty acids, omega-3 fatty acids or any combination thereof. The HMG-CoA reductase inhibitor can be atorvastatin, rosuvastatin, fluvastatin, lovastatin, pravastatin, simvastatin and the like. The cholesterol absorption inhibitor can be ezetimibe. The fibrate can be fenofibrate, bezafibrate, ciprofibrate, clofibrate, gemfibrozil and the like. The beneficial oil rich in omega-3 fatty acids can be krill, fish (e.g., Vascepa^R), flaxseed oil and the like. The omega-3 fatty acid can be ALA, DHA, EPA and the like.

Certain Compounds

Antisense oligonucleotides targeting human ANGPTL3 were described in an earlier publication (see PCT Patent Publication No. WO 2011/085271 published July 14, 2011, incorporated by reference herein, in its entirety). Several oligonucleotides (233676, 233690, 233710, 233717, 233721, 233722, 337459, 337460, 337474, 337477, 337478, 337479, 337481, 337484, 337487, 337488, 337490, 337491, 337492, 337497, 337498, 337503, 337505, 337506, 337508, 337513, 337514, 337516, 337520, 337521, 337525, 337526 and 337528) described therein, including the top ten most potent antisense compounds *in vitro*, were used as benchmarks throughout select *in vitro* screens for new antisense compounds described hereinbelow. Of the most potent compounds described in WO 2011/085271, ISIS 233722 was found to be highly variable in its ability to inhibit ANGPTL3. According, although initially included in some *in vitro* studies, 233722 was not selected as a benchmark for further studies. Of the previously identified potent *in vitro* benchmark compounds, five (233710, 233717, 337477, 337478, 337479 and 337487) were selected for testing *in vivo*, as described hereinbelow, in huANGPTL3 transgenic mice to assess the most potent in reducing human mRNA transcript and protein expression (Example 11). The antisense oligonucleotide with the highest initial *in vivo* potency in reducing ANGPTL3 levels (233710) was used as a benchmark for *in vivo* assessment of the new antisense compounds described hereinbelow.

In certain embodiments, the antisense compounds described herein benefit from one or more improved properties relative to the antisense compounds described in WO 2011/085271. These improved properties are demonstrated in the examples herein, and a non-exhaustive summary of the examples is provided below for ease of reference.

In a first screen described herein, about 3000 newly designed 5-10-5 MOE gapmer antisense compounds targeting human ANGPTL3 were tested in Hep3B cells for their effect on human ANGPTL3 mRNA *in vitro* (Example 1). The mRNA inhibition levels of the new antisense compounds were assessed with some previously designed antisense compounds (233717, 337484, 337487, 337492 and 337516) used as benchmarks in select studies. Of the about 3000 newly designed antisense compounds from this first screen, about 85 antisense compounds were selected for *in vitro* dose-dependent inhibition studies to determine their half maximal inhibitory concentration (IC₅₀) (Examples 2-3). Of the about 85 new antisense compounds tested for their half maximal inhibitory concentration (IC₅₀), about 38 antisense compounds that demonstrated

potent dose-dependent reduction of ANGPTL3 were selected for *in vivo* potency and tolerability (ALT and AST) testing in mice (Examples 11-12) with antisense compound 233710 used as a benchmark.

In a second screen described herein, about 2000 newly designed antisense compounds targeting human ANGPTL3 with a MOE gapmer motif or a mixed motif (deoxy, 5-10-5 MOE and cET gapmers) were also tested in Hep3B cells for their effect on human ANGPTL3 mRNA *in vitro* (Examples 4-6). The inhibition levels of the new antisense compounds were assessed with some previously designed antisense compounds (233717, 337487, 337513, 337514 and 337516) used as benchmarks in select studies. Of the about 2000 newly designed antisense compounds from this second screen, about 147 antisense compounds were selected for *in vitro* dose-dependent inhibition studies to determine their half maximal inhibitory concentration (IC₅₀) (Examples 7-10). Of the about 147 new antisense compounds from tested for their half maximal inhibitory concentration (IC₅₀), about 73 antisense compounds that demonstrated potent dose-dependent reduction of ANGPTL3 were selected for *in vivo* potency and tolerability (e.g., ALT and AST) testing in mice (Examples 11-12) with antisense compound 233710 used as a benchmark.

Of the about 111 antisense compounds from screens one and two that were tested for potency and tolerability in mice, 24 were selected for more extensive tolerability testing in mice by assessing liver metabolic markers, such as alanine transaminase (ALT), aspartate transaminase (AST), albumin and bilirubin, as well as kidney metabolic markers BUN and creatinine and organ weight (Example 12).

In parallel with the *in vivo* murine studies seventeen antisense compounds were selected for viscosity testing (Example 13). Generally, antisense compounds that were not optimal for viscosity were not taken forward in further studies.

Based on the results of the mice tolerability study, twenty antisense compounds were selected for *in vivo* tolerability testing in rats (Example 14). In the rats, liver metabolic markers, such as ALT, AST, albumin and bilirubin, body and organ weights, as well as kidney metabolic markers, such as BUN, creatinine and total protein/creatinine ratio, were measured to determine the tolerability of a compound *in vivo*.

The twenty antisense compounds tested in the rats were also assessed for cross-reactivity to a rhesus monkey ANGPTL3 gene sequence (Example 15). Although the antisense compounds in this study were tested in cynomolgus monkeys, the cynomolgus monkey ANGPTL3 sequence was not available for comparison to the sequences of the full-length compounds, therefore the sequences of the antisense compounds were compared to that of the closely related rhesus monkey. The sequences of eight antisense compounds were found to have 0-2 mismatches with the rhesus ANGPTL3 gene sequence and were further studied in cynomolgus monkeys (Example 15). The eight antisense compounds (ISIS 563580, ISIS 560400, ISIS 567320, ISIS 567321, ISIS 544199, ISIS 567233, ISIS 561011 and ISIS 559277) were tested for inhibition of ANGPTL3 mRNA and protein expression as well as tolerability in the monkeys. In the tolerability studies, body weights, liver metabolic markers (ALT, AST and bilirubin), kidney metabolic markers (BUN and creatinine), hematology parameters (blood cell counts, hemoglobin and hematocrit), and pro-inflammatory markers (CRP and C3) were measured. Additionally, the full-length oligonucleotide

concentration present in liver and kidney was measured and the ratio of full-length oligonucleotide in the kidney/liver was calculated.

Accordingly, provided herein are antisense compounds with any one or more improved characteristics e.g., improved relative to the antisense compounds described in WO 2011/085271. In certain
 5 embodiments, provided herein are antisense compounds comprising a modified oligonucleotide as described herein targeted to, or specifically hybridizable with, a region of nucleotides of any one of SEQ ID NOs: 1-2.

In certain embodiments, certain antisense compounds as described herein are efficacious by virtue of their potency in inhibiting ANGPTL3 expression. In certain embodiments, the compounds or compositions inhibit ANGPTL3 by at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least
 10 70%, at least 75%, at least 80%, at least 85%, at least 90% or at least 95%.

In certain embodiments, certain antisense compounds as described herein are efficacious by virtue of an *in vitro* IC₅₀ of less than 20 μ M, less than 10 μ M, less than 8 μ M, less than 5 μ M, less than 2 μ M, less than 1 μ M, less than 0.9 μ M, less than 0.8 μ M, less than 0.7 μ M, less than 0.6 μ M, or less than 0.5 μ M when tested in human cells, for example, in the Hep3B cell line (as described in Examples 2-3 and 7-10). In certain
 15 embodiments, preferred antisense compounds having an IC₅₀ <1.0 μ M include SEQ ID NOs: 15, 20, 24, 34, 35, 36, 37, 42, 43, 44, 47, 50, 51, 57, 58, 60, 77, 79, 82, 87, 88, 90, 91, 93, 94, 100, 101, 104, 110, 111, 112, 113, 114, 115, 116, 117, 118, 119, 120, 122, 123, 124, 125, 126, 127, 128, 129, 130, 131, 132, 133, 134, 135, 136, 137, 138, 139, 140, 141, 142, 143, 144, 145, 146, 147, 148, 149, 150, 151, 152, 153, 154, 155, 156, 157, 158, 169, 170, 177, 188, 209, 210, 211, 212, 213, 214, 215, 216, 217, 218, 219, 220, 221, 222, 223, 224, 225,
 20 226, 227, 228, 229, 230, 231, and 232. In certain embodiments, preferred antisense compounds having an IC₅₀ <0.9 μ M include SEQ ID NOs: 15, 20, 35, 36, 42, 43, 44, 50, 57, 60, 77, 79, 87, 88, 90, 91, 93, 94, 101, 104, 110, 111, 112, 113, 114, 115, 116, 117, 118, 119, 122, 123, 124, 125, 126, 127, 128, 129, 130, 131, 132, 133, 134, 135, 136, 137, 138, 139, 140, 141, 142, 143, 144, 145, 146, 147, 148, 149, 150, 151, 152, 153, 154, 155, 156, 157, 158, 177, 209, 210, 211, 212, 213, 214, 215, 216, 217, 218, 219, 220, 221, 222, 223, 224, 225,
 25 226, 227, 228, 229, 230, 231, and 232. In certain embodiments, preferred antisense compounds having an IC₅₀ <0.8 μ M include SEQ ID NOs: 15, 20, 35, 36, 42, 43, 44, 50, 57, 60, 77, 79, 87, 88, 90, 91, 93, 94, 101, 104, 110, 111, 112, 113, 114, 115, 116, 117, 118, 122, 123, 124, 125, 126, 127, 128, 129, 130, 131, 132, 133, 134, 135, 136, 137, 138, 139, 140, 141, 142, 143, 144, 145, 146, 147, 148, 149, 150, 151, 152, 153, 154, 155, 156, 157, 158, 177, 209, 210, 211, 212, 213, 214, 215, 217, 218, 219, 220, 221, 222, 223, 224, 225, 228, 229,
 30 230, 231, and 232. In certain embodiments, preferred antisense compounds having an IC₅₀ <0.7 μ M include SEQ ID NOs: 15, 20, 36, 42, 43, 57, 60, 114, 117, 127, 131, 177, 209, 210, 211, 212, 213, 214, 215, 217, 218, 219, 220, 221, 222, 223, 224, 225, 228, 229, 230, 231, and 232. In certain embodiments, preferred antisense compounds having an IC₅₀ <0.6 μ M include SEQ ID NOs: 15, 20, 36, 42, 43, 57, 60, 114, 117, 127, 131, 177, 209, 210, 211, 212, 213, 215, 217, 218, 219, 220, 221, 222, 224, 225, 228, 229, 230, 231, and

232. In certain embodiments, preferred antisense compounds having an $IC_{50} < 0.5 \mu M$ include SEQ ID NOs: 43, 114, 117, 127, 131, 177, 209, 210, 211, 212, 215, 217, 218, 219, 220, 221, 222, 229, 230, and 232.

In certain embodiments, certain 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, less than 15 cP, or less than 10 cP when measured by an assay (as described in Example 13). Oligonucleotides having a viscosity greater than 40 cP would have less than optimal viscosity. In certain embodiments, preferred antisense compounds having a viscosity < 20 cP include SEQ ID NOs: 16, 18, 20, 34, 35, 36, 38, 49, 77, 90, 93, and 94. In certain embodiments, preferred antisense compounds having a viscosity < 15 cP include SEQ ID NOs: 16, 18, 20, 34, 35, 38, 49, 90, 93, and 94. In certain embodiments, preferred antisense compounds having a viscosity < 10 cP include SEQ ID NOs: 18, 34, 35, 49, 90, 93, and 94.

In certain embodiments, certain antisense compounds as described herein are highly tolerable, as demonstrated by the *in vivo* tolerability measurements described in the examples. In certain embodiments, the certain antisense compounds as described herein are highly tolerable, as demonstrated by having an increase in ALT and/or AST value of no more than 3 fold, 2 fold or 1.5 fold over saline treated animals.

In certain embodiments, certain antisense compounds as described herein are efficacious by virtue of having one or more of an inhibition potency of greater than 50%, an *in vitro* IC_{50} of less than $1 \mu M$, a viscosity of less than 20 cP, and no more than a 3 fold increase in ALT and/or AST.

In certain embodiments, ISIS 563580 (SEQ ID NO: 77) is preferred. This compound was found to be a potent inhibitor in ANGPTL3 transgenic mice and the most tolerable antisense compound. It had an acceptable viscosity of about 16.83 cP and an IC_{50} value of $< 0.8 \mu M$ *in vitro*. In mice it had no more than a 3 fold increase in ALT and/or AST levels over saline treated animals. Also, in monkeys, it was among the most tolerable and potent compounds in inhibiting ANGPTL3 and had the best ratio of full-length oligonucleotide concentration.

In certain embodiments, ISIS 544199 (SEQ ID NO: 20) is preferred. This compound was found to be a potent and tolerable antisense compound. It had an acceptable viscosity of 1.7 cP and an IC_{50} value of $< 0.5 \mu M$ *in vitro*. In mice it had no more than a 3 fold increase in ALT and/or AST levels over saline treated animals. Also, in monkeys, it was among the most potent compounds in inhibiting ANGPTL3 and had a good ratio of full-length oligonucleotide concentration.

In certain embodiments, ISIS 559277 (SEQ ID NO: 110) is preferred. This compound was found to be a potent and tolerable antisense compound. It had an IC_{50} value of $< 0.8 \mu M$ *in vitro*. In mice it had no more than a 3 fold increase in ALT and/or AST levels over saline treated animals. Also, in monkeys, it was among the most potent compounds in inhibiting ANGPTL3 and had a good ratio of full-length oligonucleotide concentration.

EXAMPLES***Non-limiting 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 recited in the present application is incorporated herein by reference in its entirety.

Example 1: Antisense inhibition of human Angiopoietin-like 3 in Hep3B cells by MOE gapmers

Antisense oligonucleotides were designed targeting an Angiopoietin-like 3 (ANGPTL3) nucleic acid and were tested for their effects on ANGPTL3 mRNA in vitro. The antisense oligonucleotides were tested in a series of experiments that had similar culture conditions. The results for each experiment are presented in separate tables shown below. Cultured Hep3B cells at a density of 20,000 cells per well were transfected using electroporation with 4,500 nM antisense oligonucleotide. After a treatment period of approximately 24 hours, RNA was isolated from the cells and ANGPTL3 mRNA levels were measured by quantitative real-time PCR. Human primer probe set RTS3492_MGB (forward sequence

CCGTGGAAGACCAATATAACAATT, designated herein as SEQ ID NO: 4; AGTCCTTCTGAGCTGATTTTCTATTTCT; reverse sequence, designated herein as SEQ ID NO: 5; probe sequence AACCAACAGCATAGTCAAATA, designated herein as SEQ ID NO: 6) was used to measure mRNA levels. ANGPTL3 mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN®. Results are presented as percent inhibition of ANGPTL3, relative to untreated control cells.

The newly designed chimeric antisense oligonucleotides in the Tables below were designed as 5-10-5 MOE gapmers. The 5-10-5 MOE 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. "Start site" indicates the 5'-most nucleoside to which the gapmer is targeted in the human gene sequence. "Stop site" indicates the 3'-most nucleoside to which the gapmer is targeted human gene sequence. Each gapmer listed in the Tables below is targeted to either the human ANGPTL3 mRNA, designated herein as SEQ ID NO: 1 (GENBANK Accession No. NM_014495.2) or the human ANGPTL3 genomic sequence, designated herein as SEQ ID NO: 2 (GENBANK Accession No. NT_032977.9 truncated from nucleotides 33032001 to 33046000). 'n/a' indicates that the antisense oligonucleotide does not target that particular gene sequence with 100% complementarity.

Table 1

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
544059	23	42	GATTTTCAATTTCAGCAAC	40	3127	3146	238
337459	49	68	AGCTTAATTGTGAACATTTT	47	3153	3172	239
544060	54	73	GAAGGAGCTTAATTGTGAAC	1	3158	3177	240
544061	63	82	CAATAAAAAGAAGGAGCTTA	37	3167	3186	241
544062	66	85	GAACAATAAAAAGAAGGAGC	38	3170	3189	242
544063	85	104	CTGGAGGAAATAACTAGAGG	30	3189	3208	243
337460	88	107	ATTCTGGAGGAAATAACTAG	39	3192	3211	244
544064	112	131	TCAAATGATGAATTGTCTTG	36	3216	3235	245
544065	138	157	TTGATTTTGGCTCTGGAGAT	26	3242	3261	246
544066	145	164	GCAAATCTTGATTTTGGCTC	56	3249	3268	247
233676	148	167	ATAGCAAATCTTGATTTTGG	69	3252	3271	248
544067	156	175	CGTCTAACATAGCAAATCTT	64	3260	3279	249
544068	174	193	TGGCTAAAATTTTACATCG	28	3278	3297	250
544069	178	197	CCATTGGCTAAAATTTTAC	0	3282	3301	251
544070	184	203	AGGAGGCCATTGGCTAAAAT	7	3288	3307	252
544071	187	206	TGAAGGAGGCCATTGGCTAA	32	3291	3310	253
544072	195	214	GTCCCAACTGAAGGAGGCCA	9	3299	3318	254
544073	199	218	CCATGTCCCAACTGAAGGAG	6	3303	3322	255
544074	202	221	AGACCATGTCCCAACTGAAG	18	3306	3325	256
544075	206	225	TTTAAGACCATGTCCCAACT	0	3310	3329	257
544076	209	228	GTCTTTAAGACCATGTCCCA	0	3313	3332	258
544077	216	235	GGACAAAGTCTTTAAGACCA	0	3320	3339	259
544078	222	241	TCTTATGGACAAAGTCTTTA	0	3326	3345	260
544079	245	264	TATGTCATTAATTTGGCCCT	0	3349	3368	261
544080	270	289	GATCAAATATGTTGAGTTTT	27	3374	3393	262
233690	274	293	GACTGATCAAATATGTTGAG	49	3378	3397	263
544081	316	335	TCTTCTTTGATTTCACTGGT	62	3420	3439	264
544082	334	353	CTTCTCAGTTCCTTTTCTTC	35	3438	3457	265
544083	337	356	GTTCTTCTCAGTTCCTTTTC	60	3441	3460	266
544084	341	360	TGTAGTTCTTCTCAGTTCCT	51	3445	3464	267
544431	345	364	TATATGTAGTTCTTCTCAGT	9	3449	3468	268
544086	348	367	GTTTATATGTAGTTCTTCTC	39	3452	3471	269
544087	352	371	TGTAGTTTATATGTAGTTCT	30	3456	3475	270
544088	356	375	GACTTGTAGTTTATATGTAG	12	3460	3479	271
544089	364	383	TCATTTTTGACTTGTAGTTT	31	3468	3487	272
544090	369	388	CCTCTTCATTTTTGACTTGT	61	3473	3492	273
544091	375	394	TCTTTACCTCTTCATTTTTG	48	3479	3498	274

544092	380	399	CATATTCTTTACCTCTTCAT	35	3484	3503	275
544093	384	403	GTGACATATTCTTTACCTCT	63	3488	3507	276
544094	392	411	GAGTTCAAGTGACATATTCT	53	3496	3515	277
544095	398	417	TGAGTTGAGTTCAAGTGACA	31	3502	3521	278
544096	403	422	AGTTTTGAGTTGAGTTCAAG	14	3507	3526	279
544097	406	425	TCAAGTTTTGAGTTGAGTTC	38	3510	3529	280
544098	414	433	GGAGGCTTTCAAGTTTTGAG	39	3518	3537	281
544099	423	442	TTTCTTCTAGGAGGCTTTCA	57	3527	3546	282
544100	427	446	ATTTTTTCTTCTAGGAGGCT	39	3531	3550	283
544101	432	451	GTAGAATTTTTTCTTCTAGG	28	3536	3555	284
544102	462	481	GCTCTTCTAAATATTTCACT	60	3566	3585	285
544103	474	493	AGTTAGTTAGTTGCTCTTCT	40	3578	3597	286
544104	492	511	CAGGTTGATTTTGAATTAAG	38	3596	3615	287
544105	495	514	TTTCAGGTTGATTTTGAATT	28	3599	3618	288
544106	499	518	GGAGTTTCAGGTTGATTTTG	38	3603	3622	289
544107	504	523	GTTCTGGAGTTTCAGGTTGA	50	3608	3627	290
544108	526	545	TTAAGTGAAGTTACTTCTGG	20	3630	3649	291
544109	555	574	TGCTATTATCTTGTTTTTCT	23	4293	4312	292
544110	564	583	GGTCTTTGATGCTATTATCT	67	4302	4321	293
544111	567	586	GAAGGTCTTTGATGCTATTA	49	4305	4324	294
544112	572	591	CTGGAGAAGGTCTTTGATGC	52	4310	4329	295
544113	643	662	CTGAGCTGATTTTCTATTTC	12	n/a	n/a	296
337477	664	683	GGTTCTTGAATACTAGTCCT	70	6677	6696	234
544114	673	692	ATTTCTGTGGGTTCTTGAAT	32	6686	6705	297
337478	675	694	AAATTTCTGTGGGTTCTTGA	51	6688	6707	235
544115	678	697	GAGAAATTTCTGTGGGTTCT	54	6691	6710	298
544116	682	701	GATAGAGAAATTTCTGTGGG	25	6695	6714	299
544117	689	708	CTTGGAAGATAGAGAAATTT	16	6702	6721	300
337479	692	711	TGGCTTGGAAGATAGAGAAA	34	6705	6724	236
544118	699	718	GTGCTCTTGGCTTGGAAGAT	64	6712	6731	301
544119	703	722	CTTGGTGCTCTTGGCTTGGA	70	6716	6735	302
544120	707	726	AGTTCTTGGTGCTCTTGGCT	82	6720	6739	15
233710	710	729	AGTAGTTCTTGGTGCTCTTG	63	6723	6742	233
544121	713	732	GGGAGTAGTTCTTGGTGCTC	64	6726	6745	303
544122	722	741	CTGAAGAAAGGGAGTAGTTC	24	6735	6754	304
544123	752	771	ATCATGTTTTACATTTCTTA	0	6765	6784	305
544124	755	774	GCCATCATGTTTTACATTTTC	35	n/a	n/a	306
544125	759	778	GAATGCCATCATGTTTTACA	8	n/a	n/a	307
544126	762	781	CAGGAATGCCATCATGTTTT	6	n/a	n/a	308
337487	804	823	CACTTGTATGTTACCTCTG	65	7389	7408	28
233717	889	908	TGAATTAATGTCCATGGACT	33	7876	7895	14

Table 2

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
544204	n/a	n/a	GACTTCTTAACTCTATATAT	0	3076	3095	309
544205	n/a	n/a	CTAGACTTCTTAACTCTATA	0	3079	3098	310
544206	n/a	n/a	GACCTAGACTTCTTAACTCT	0	3082	3101	311
544207	n/a	n/a	GGAAGCAGACCTAGACTTCT	21	3089	3108	312
544208	n/a	n/a	TCTGGAAGCAGACCTAGACT	23	3092	3111	313
544209	n/a	n/a	TCTTCTGGAAGCAGACCTAG	7	3095	3114	314
544210	n/a	n/a	CTAATCTTTAGGGATTTAGG	24	11433	11452	315
544211	n/a	n/a	TGTATCTAATCTTTAGGGAT	2	11438	11457	316
544213	n/a	n/a	TAACTTGGGCACTATATCCT	44	11553	11572	317
544214	n/a	n/a	ATTGACAAAGGTAGGTCACC	59	11576	11595	318
544215	n/a	n/a	ATATGACATGTATATTGGAT	41	11620	11639	319
544216	n/a	n/a	TTTTGTACTTTTCTGGAACA	34	11704	11723	320
544217	n/a	n/a	TAGTCTGTGGTCCTGAAAAT	32	11748	11767	321
544218	n/a	n/a	AGCTTAGTCTGTGGTCCTGA	20	11752	11771	322
544219	n/a	n/a	GACAGCTTAGTCTGTGGTCC	45	11755	11774	323
544220	n/a	n/a	GTATTCTGGCCCTAAAAAAA	2	11789	11808	324
544221	n/a	n/a	ATTTTGGTATTCTGGCCCTA	39	11795	11814	325
544223	n/a	n/a	TTTGCATTTGAAATTGTCCA	32	11837	11856	326
544224	n/a	n/a	GGAAGCAACTCATATATTAA	39	11869	11888	327
544225	n/a	n/a	TATCAGAAAAAGATACCTGA	0	9821	9840	328
544226	n/a	n/a	ATAATAGCTAATAATGTGGG	15	9875	9894	329
544227	n/a	n/a	TGCAGATAATAGCTAATAAT	31	9880	9899	330
544228	n/a	n/a	TGTCATTGCAGATAATAGCT	61	9886	9905	331
544229	n/a	n/a	TAAAAGTTGTCATTGCAGAT	38	9893	9912	332
544230	n/a	n/a	CGGATTTTAAAAGTTGTCA	45	9901	9920	333
544231	n/a	n/a	GGGATTCGGATTTTAAAAG	0	9907	9926	334
544232	n/a	n/a	TTTGGGATTCGGATTTTAA	24	9910	9929	335
544233	n/a	n/a	ACGCTTATTTGGGATTCGGA	53	9917	9936	336
544251	n/a	n/a	TTTAAGAGATTTACAAGTCA	11	2811	2830	337
544252	n/a	n/a	GACTACCTGTTTTTAAAAGC	6	2851	2870	338
544253	n/a	n/a	TATGGTGACTACCTGTTTTT	12	2857	2876	339
544254	n/a	n/a	ACTTTGCTGTATTATAAACT	12	2890	2909	340
544255	n/a	n/a	ATTGTATTTAACTTTGCTGT	0	2900	2919	341
544256	n/a	n/a	GAGCAACTAACTTAATAGGT	13	2928	2947	342
544257	n/a	n/a	GAAATGAGCAACTAACTTAA	25	2933	2952	343
544258	n/a	n/a	AATCAAAGAAATGAGCAACT	0	2940	2959	344
544259	n/a	n/a	ACCTTCTTCCACATTGAGTT	8	2977	2996	345

544260	n/a	n/a	CACGAATGTAACCTTCTTCC	0	2987	3006	346
544261	n/a	n/a	TTAACTTGCACGAATGTAAC	27	2995	3014	347
544262	n/a	n/a	TATATATACCAATATTTGCC	0	3063	3082	348
544263	n/a	n/a	TCTTAACTCTATATATACCA	0	3072	3091	349
544264	n/a	n/a	CTTTAAGTGAAGTTACTTCT	17	3632	3651	350
544265	n/a	n/a	TCTACTTACTTTAAGTGAAG	9	3640	3659	351
544266	n/a	n/a	GAACCCTCTTTATTTTCTAC	1	3655	3674	352
544267	n/a	n/a	ACATAAACATGAACCCTCTT	6	3665	3684	353
544268	n/a	n/a	CCACATTGAAAACATAAACA	25	3676	3695	354
544269	n/a	n/a	GCATGCCTTAGAAATATTTT	7	3707	3726	355
544270	n/a	n/a	CAATGCAACAAAGTATTTCA	0	3731	3750	356
544271	n/a	n/a	CTGGAGATTATTTTCTTGG	34	3768	3787	357
544272	n/a	n/a	TTCATATATAACATTAGGGA	0	3830	3849	358
544273	n/a	n/a	TCAGTGTTTTTCATATATAAC	18	3838	3857	359
544274	n/a	n/a	GACATAGTGTTCTAGATTGT	14	3900	3919	360
544275	n/a	n/a	CAATAGTGTAATGACATAGT	21	3912	3931	361
544276	n/a	n/a	TTACTTACCTTCAGTAATTT	0	3933	3952	362
544277	n/a	n/a	ATCTTTTCCATTTACTGTAT	8	4005	4024	363
544278	n/a	n/a	AGAAAAAGCCCAGCATATTT	11	4037	4056	364
544279	n/a	n/a	GTATGCTTCTTTCAAATAGC	36	4130	4149	365
544280	n/a	n/a	CCTTCCCCTTGTATGCTTCT	41	4140	4159	366
544281	n/a	n/a	CCTGTAACACTATCATAATC	1	4207	4226	367
544282	n/a	n/a	TGACTTACCTGATTTTCTAT	6	4384	4403	368
544283	n/a	n/a	GATGGGACATACCATTAAAA	0	4407	4426	369
544284	n/a	n/a	GTGAAAGATGGGACATACCA	20	4413	4432	370
544285	n/a	n/a	CCTGTGTGAAAGATGGGACA	6	4418	4437	371
544286	n/a	n/a	CATTGGCTGCTATGAATTAA	41	4681	4700	372
544287	n/a	n/a	GATGACATTGGCTGCTATGA	40	4686	4705	373
544288	n/a	n/a	GAGAAACATGATCTAATTTG	12	4717	4736	374
544289	n/a	n/a	ATGGAAAGCTATTGTGTGGT	0	4747	4766	375
544290	n/a	n/a	GTCTAAAGAGCCAATATGAG	22	4771	4790	376
544291	n/a	n/a	AATCTTGGTCTAAAGAGCCA	46	4778	4797	377
544433	n/a	n/a	GAGATTTACAAGTCAAAAAT	4	2806	2825	378
544434	n/a	n/a	ATTTAACTTTGCTGTATTAT	0	2895	2914	379
544435	n/a	n/a	ATCAATGCTAAATGAAATCA	0	2955	2974	380
544436	n/a	n/a	TATTTTCTGGAGATTATTTT	0	3774	3793	381
544437	n/a	n/a	AAAATGAATATTGGCAATTC	0	4159	4178	382
233717	889	908	TGAATTAATGTCCATGGACT	36	7876	7895	14
544202	2081	2100	AAAGTCAATGTGACTTAGTA	42	11053	11072	383
544203	2104	2123	AAGGTATAGTGATACCTCAT	56	11076	11095	384

Table 3

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
544127	765	784	CAGCAGGAATGCCATCATGT	4	N/A	N/A	385
544128	819	838	TGATGGCATAACATGCCACTT	0	7404	7423	386
544129	828	847	TGCTGGGTCTGATGGCATAAC	44	7413	7432	387
544130	832	851	GAGTTGCTGGGTCTGATGGC	16	7417	7436	388
544131	841	860	AAAACCTTGAGAGTTGCTGGG	0	7426	7445	389
544132	848	867	GACATGAAAAACTTGAGAGT	0	7433	7452	390
544133	859	878	ACATCACAGTAGACATGAAA	25	7444	7463	391
233717	889	908	TGAATTAATGTCCATGGACT	36	7876	7895	14
544134	915	934	AGTTTTGTGATCCATCTATT	46	7902	7921	392
544135	918	937	TGAAGTTTTGTGATCCATCT	42	7905	7924	393
544136	926	945	CGTTTCATTGAAGTTTTGTG	45	7913	7932	394
544137	946	965	CCATATTTGTAGTTCTCCCA	44	7933	7952	395
544138	949	968	AAACCATATTTGTAGTTCTC	25	7936	7955	396
544139	970	989	AATTCTCCATCAAGCCTCCC	35	N/A	N/A	397
233722	991	1010	ATCTTCTCTAGGCCCAACCA	65	9566	9585	398
544432	997	1016	GAGTATATCTTCTCTAGGCC	0	9572	9591	399
544140	1002	1021	CTATGGAGTATATCTTCTCT	6	9577	9596	400
544141	1008	1027	GCTTCACTATGGAGTATATC	63	9583	9602	401
544142	1013	1032	AGATTGCTTCACTATGGAGT	52	9588	9607	402
544143	1046	1065	CCAGTCTTCCAACCTCAATTC	35	9621	9640	403
544144	1052	1071	GTCTTTCCAGTCTTCCAACCT	64	9627	9646	404
544145	1055	1074	GTTGTCTTTCCAGTCTTCCA	80	9630	9649	16
544146	1059	1078	GTTTGTGTCTTTCCAGTCT	59	9634	9653	405
544147	1062	1081	AATGTTTGTGTCTTTCCAG	12	9637	9656	406
544148	1095	1114	CGTGATTTCCCAAGTAAAAA	56	9670	9689	407
544149	1160	1179	GTTTTCCGGGATTGCATTGG	33	9735	9754	408
544150	1165	1184	TCTTTGTTTTCCGGGATTGC	54	9740	9759	409
544151	1170	1189	CCAAATCTTTGTTTTCCGGG	64	9745	9764	410
544152	1173	1192	ACACCAAATCTTTGTTTTCC	37	9748	9767	411
544153	1178	1197	AGAAAACACCAAATCTTTGT	32	9753	9772	412
544154	1183	1202	CAAGTAGAAAACACCAAATC	13	9758	9777	413
544155	1188	1207	GATCCCAAGTAGAAAACACC	0	9763	9782	414
544156	1195	1214	GCTTTGTGATCCCAAGTAGA	74	9770	9789	17
544157	1198	1217	TTTGCTTTGTGATCCCAAGT	73	9773	9792	415
544158	1202	1221	TCCTTTTGCTTTGTGATCCC	62	9777	9796	416
544159	1208	1227	GAAGTGTCTTTTGCTTTGT	30	9783	9802	417
544160	1246	1265	TGCCACCACCAGCCTCCTGA	60	N/A	N/A	418

544161	1253	1272	CTCATCATGCCACCACCAGC	73	10225	10244	419
544162	1269	1288	GGTTGTTTTCTCCACACTCA	76	10241	10260	18
544163	1276	1295	CCATTTAGGTTGTTTTCTCC	25	10248	10267	420
544164	1283	1302	ATATTTACCATTTAGGTTGT	25	10255	10274	421
544165	1294	1313	CTTGGTTTGTTATATTTACC	63	10266	10285	422
544166	1353	1372	ACCTTCCATTTTGAGACTTC	75	10325	10344	19
544167	1363	1382	ATAGAGTATAACCTTCCATT	71	10335	10354	423
544168	1367	1386	TTTTATAGAGTATAACCTTC	37	10339	10358	424
544169	1374	1393	TGGTTGATTTTATAGAGTAT	37	10346	10365	425
544170	1378	1397	ATTTTGGTTGATTTTATAGA	3	10350	10369	426
544171	1383	1402	TCAACATTTTGGTTGATTTT	16	10355	10374	427
544172	1390	1409	GGATGGATCAACATTTTGGT	51	10362	10381	428
544173	1393	1412	GTTGGATGGATCAACATTTT	62	10365	10384	429
544174	1396	1415	TCTGTTGGATGGATCAACAT	5	10368	10387	430
544175	1401	1420	CTGAATCTGTTGGATGGATC	55	10373	10392	431
544176	1407	1426	AGCTTTCTGAATCTGTTGGA	65	10379	10398	432
544177	1414	1433	CATTCAAAGCTTTCTGAATC	21	10386	10405	433
544178	1417	1436	GTTCATTCAAAGCTTTCTGA	66	10389	10408	434
544179	1420	1439	TCAGTTCATTCAAAGCTTTC	6	10392	10411	435
544180	1423	1442	GCCTCAGTTCATTCAAAGCT	68	10395	10414	436
544181	1427	1446	ATTTGCCTCAGTTCATTCAA	53	10399	10418	437
544182	1431	1450	TTAAATTTGCCTCAGTTCAT	40	10403	10422	438
544183	1436	1455	GCCTTTTAAATTTGCCTCAG	70	10408	10427	439
544184	1498	1517	AGGATTTAATACCAGATTAT	38	10470	10489	440
544185	1502	1521	CTTAAGGATTTAATACCAGA	56	10474	10493	441
544186	1505	1524	TCTCTTAAGGATTTAATACC	33	10477	10496	442
544187	1546	1565	GACAGTGACTTTAAGATAAA	35	10518	10537	443
544188	1572	1591	TGTGATTGTATGTTTAATCT	48	10544	10563	444
544189	1578	1597	AGGTTATGTGATTGTATGTT	48	10550	10569	445
544190	1583	1602	CTTTAAGGTTATGTGATTGT	48	10555	10574	446
544191	1589	1608	GGTATTCTTTAAGGTTATGT	62	10561	10580	447
544192	1656	1675	ATTGATTCCCACATCACAAA	47	10628	10647	448
544193	1661	1680	CTAAAATTGATTCCCACATC	67	10633	10652	449
544194	1665	1684	CCATCTAAAATTGATTCCCA	63	10637	10656	450
544195	1771	1790	TTGTGATATTAGCTCATATG	59	10743	10762	451
544196	1794	1813	ACTAGTTTTTTTAAACTGGGA	28	10766	10785	452
544197	1820	1839	GTCAAGTTTAGAGTTTTAAC	44	10792	10811	453
544198	1826	1845	TATTTAGTCAAGTTTAGAGT	14	10798	10817	454
544199	1907	1926	TACACATACTCTGTGCTGAC	82	10879	10898	20
544200	1913	1932	GATTTTTACACATACTCTGT	57	10885	10904	455
544201	2008	2027	CTGCTTCATTAGGTTTCATA	61	10980	10999	456

Table 4

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
544127	765	784	CAGCAGGAATGCCATCATGT	0	N/A	N/A	457
544128	819	838	TGATGGCATACATGCCACTT	13	7404	7423	458
544129	828	847	TGCTGGGTCTGATGGCATAAC	49	7413	7432	459
544130	832	851	GAGTTGCTGGGTCTGATGGC	27	7417	7436	460
544131	841	860	AAAACCTTGAGAGTTGCTGGG	0	7426	7445	461
544132	848	867	GACATGAAAAACTTGAGAGT	0	7433	7452	462
544133	859	878	ACATCACAGTAGACATGAAA	18	7444	7463	463
233717	889	908	TGAATTAATGTCCATGGACT	55	7876	7895	14
544134	915	934	AGTTTTGTGATCCATCTATT	68	7902	7921	464
544135	918	937	TGAAGTTTTGTGATCCATCT	77	7905	7924	465
544136	926	945	CGTTTCATTGAAGTTTTGTG	60	7913	7932	466
544137	946	965	CCATATTTGTAGTTCTCCCA	64	7933	7952	467
544138	949	968	AAACCATATTTGTAGTTCTC	45	7936	7955	468
544139	970	989	AATTCTCCATCAAGCCTCCC	70	N/A	N/A	469
233722	991	1010	ATCTTCTCTAGGCCCAACCA	96	9566	9585	470
544432	997	1016	GAGTATATCTTCTCTAGGCC	69	9572	9591	471
544140	1002	1021	CTATGGAGTATATCTTCTCT	37	9577	9596	472
544141	1008	1027	GCTTCACTATGGAGTATATC	65	9583	9602	473
544142	1013	1032	AGATTGCTTCACTATGGAGT	55	9588	9607	474
544143	1046	1065	CCAGTCTTCCAACCTCAATTC	31	9621	9640	475
544144	1052	1071	GTCTTTCCAGTCTTCCAACCT	72	9627	9646	476
544145	1055	1074	GTTGTCTTTCCAGTCTTCCA	86	9630	9649	16
544146	1059	1078	GTTTGTGTCTTTCCAGTCT	66	9634	9653	477
544147	1062	1081	AATGTTTGTGTCTTTCCAG	21	9637	9656	478
544148	1095	1114	CGTGATTTCCCAAGTAAAAA	63	9670	9689	479
544149	1160	1179	GTTTTCCGGGATTGCATTGG	32	9735	9754	480
544150	1165	1184	TCTTTGTTTTCCGGGATTGC	48	9740	9759	481
544151	1170	1189	CCAAATCTTTGTTTTCCGGG	72	9745	9764	482
544152	1173	1192	ACACCAAATCTTTGTTTTCC	39	9748	9767	483
544153	1178	1197	AGAAAACACCAAATCTTTGT	39	9753	9772	484
544154	1183	1202	CAAGTAGAAAACACCAAATC	22	9758	9777	485
544155	1188	1207	GATCCCAAGTAGAAAACACC	5	9763	9782	486
544156	1195	1214	GCTTTGTGATCCCAAGTAGA	79	9770	9789	17
544157	1198	1217	TTTGCTTTGTGATCCCAAGT	80	9773	9792	487
544158	1202	1221	TCCTTTTGCTTTGTGATCCC	73	9777	9796	488
544159	1208	1227	GAAGTGTCCTTTTGCTTTGT	33	9783	9802	489

544160	1246	1265	TGCCACCACCAGCCTCCTGA	67	N/A	N/A	490
544161	1253	1272	CTCATCATGCCACCACCAGC	79	10225	10244	491
544162	1269	1288	GGTTGTTTTCTCCACACTCA	84	10241	10260	18
544163	1276	1295	CCATTTAGGTTGTTTTCTCC	34	10248	10267	492
544164	1283	1302	ATATTTACCATTTAGGTTGT	17	10255	10274	493
544165	1294	1313	CTTGGTTTGTTATATTTACC	76	10266	10285	494
544166	1353	1372	ACCTTCCATTTTGAGACTTC	79	10325	10344	19
544167	1363	1382	ATAGAGTATAACCTTCCATT	73	10335	10354	495
544168	1367	1386	TTTTATAGAGTATAACCTTC	41	10339	10358	496
544169	1374	1393	TGGTTGATTTTATAGAGTAT	53	10346	10365	497
544170	1378	1397	ATTTTGGTTGATTTTATAGA	28	10350	10369	498
544171	1383	1402	TCAACATTTTGGTTGATTTT	19	10355	10374	499
544172	1390	1409	GGATGGATCAACATTTTGGT	66	10362	10381	500
544173	1393	1412	GTTGGATGGATCAACATTTT	71	10365	10384	501
544174	1396	1415	TCTGTTGGATGGATCAACAT	35	10368	10387	502
544175	1401	1420	CTGAATCTGTTGGATGGATC	68	10373	10392	503
544176	1407	1426	AGCTTTCTGAATCTGTTGGA	70	10379	10398	504
544177	1414	1433	CATTCAAAGCTTTCTGAATC	35	10386	10405	505
544178	1417	1436	GTTCATTCAAAGCTTTCTGA	76	10389	10408	506
544179	1420	1439	TCAGTTCATTCAAAGCTTTC	15	10392	10411	507
544180	1423	1442	GCCTCAGTTCATTCAAAGCT	68	10395	10414	508
544181	1427	1446	ATTTGCCTCAGTTCATTCAA	67	10399	10418	509
544182	1431	1450	TTAAATTTGCCTCAGTTCAT	51	10403	10422	510
544183	1436	1455	GCCTTTTAAATTTGCCTCAG	80	10408	10427	511
544184	1498	1517	AGGATTTAATACCAGATTAT	54	10470	10489	512
544185	1502	1521	CTTAAGGATTTAATACCAGA	69	10474	10493	513
544186	1505	1524	TCTCTTAAGGATTTAATACC	58	10477	10496	514
544187	1546	1565	GACAGTGACTTTAAGATAAA	34	10518	10537	515
544188	1572	1591	TGTGATTGTATGTTTAATCT	47	10544	10563	516
544189	1578	1597	AGGTTATGTGATTGTATGTT	68	10550	10569	517
544190	1583	1602	CTTTAAGGTTATGTGATTGT	62	10555	10574	518
544191	1589	1608	GGTATTCTTTAAGGTTATGT	66	10561	10580	519
544192	1656	1675	ATTGATTCCCACATCACAAA	50	10628	10647	520
544193	1661	1680	CTAAAATTGATTCCCACATC	73	10633	10652	521
544194	1665	1684	CCATCTAAAATTGATTCCCA	73	10637	10656	522
544195	1771	1790	TTGTGATATTAGCTCATATG	57	10743	10762	523
544196	1794	1813	ACTAGTTTTTTAAACTGGGA	21	10766	10785	524
544197	1820	1839	GTCAAGTTTAGAGTTTTAAC	53	10792	10811	525
544198	1826	1845	TATTTAGTCAAGTTTAGAGT	11	10798	10817	526
544199	1907	1926	TACACATACTCTGTGCTGAC	84	10879	10898	20
544200	1913	1932	GATTTTTTACACATACTCTGT	53	10885	10904	527

544201	2008	2027	CTGCTTCATTAGGTTTCATA	67	10980	10999	528
--------	------	------	----------------------	----	-------	-------	-----

Table 5

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
544127	765	784	CAGCAGGAATGCCATCATGT	18	N/A	N/A	529
544128	819	838	TGATGGCATACATGCCACTT	0	7404	7423	530
544129	828	847	TGCTGGGTCTGATGGCATAAC	48	7413	7432	531
544130	832	851	GAGTTGCTGGGTCTGATGGC	14	7417	7436	532
544131	841	860	AAAACCTTGAGAGTTGCTGGG	5	7426	7445	533
544132	848	867	GACATGAAAAACTTGAGAGT	0	7433	7452	534
544133	859	878	ACATCACAGTAGACATGAAA	28	7444	7463	535
233717	889	908	TGAATTAATGTCCATGGACT	51	7876	7895	14
544134	915	934	AGTTTTGTGATCCATCTATT	36	7902	7921	536
544135	918	937	TGAAGTTTTGTGATCCATCT	61	7905	7924	537
544136	926	945	CGTTTCATTGAAGTTTTGTG	54	7913	7932	538
544137	946	965	CCATATTTGTAGTTCTCCCA	67	7933	7952	539
544138	949	968	AAACCATATTTGTAGTTCTC	39	7936	7955	540
544139	970	989	AATTCTCCATCAAGCCTCCC	77	N/A	N/A	541
233722	991	1010	ATCTTCTCTAGGCCCAACCA	95	9566	9585	542
544432	997	1016	GAGTATATCTTCTCTAGGCC	86	9572	9591	543
544140	1002	1021	CTATGGAGTATATCTTCTCT	57	9577	9596	544
544141	1008	1027	GCTTCACTATGGAGTATATC	52	9583	9602	545
544142	1013	1032	AGATTGCTTCACTATGGAGT	40	9588	9607	546
544143	1046	1065	CCAGTCTTCCAACCTCAATTC	32	9621	9640	547
544144	1052	1071	GTCTTTCCAGTCTTCCAACCT	53	9627	9646	548
544145	1055	1074	GTTGTCTTTCCAGTCTTCCA	80	9630	9649	16
544146	1059	1078	GTTTGTTGTCTTTCCAGTCT	59	9634	9653	549
544147	1062	1081	AATGTTTGTTGTCTTTCCAG	42	9637	9656	550
544148	1095	1114	CGTGATTTCCCAAGTAAAAA	76	9670	9689	551
544149	1160	1179	GTTTTCCGGGATTGCATTGG	29	9735	9754	552
544150	1165	1184	TCTTTGTTTTCCGGGATTGC	50	9740	9759	553
544151	1170	1189	CCAAATCTTTGTTTTCCGGG	56	9745	9764	554
544152	1173	1192	ACACCAAATCTTTGTTTTCC	26	9748	9767	555
544153	1178	1197	AGAAAACACCAAATCTTTGT	22	9753	9772	556
544154	1183	1202	CAAGTAGAAAACACCAAATC	29	9758	9777	557
544155	1188	1207	GATCCCAAGTAGAAAACACC	16	9763	9782	558
544156	1195	1214	GCTTTGTGATCCCAAGTAGA	71	9770	9789	17
544157	1198	1217	TTTGCTTTGTGATCCCAAGT	55	9773	9792	559
544158	1202	1221	TCCTTTTGCTTTGTGATCCC	51	9777	9796	560

544159	1208	1227	GAAGTGTCTTTTGCTTTGT	8	9783	9802	561
544160	1246	1265	TGCCACCACCAGCCTCCTGA	68	N/A	N/A	562
544161	1253	1272	CTCATCATGCCACCACCAGC	48	10225	10244	563
544162	1269	1288	GGTTGTTTTCTCCACACTCA	74	10241	10260	18
544163	1276	1295	CCATTTAGGTTGTTTTCTCC	33	10248	10267	564
544164	1283	1302	ATATTTACCATTTAGGTTGT	0	10255	10274	565
544165	1294	1313	CTTGGTTTGTTATATTTACC	52	10266	10285	566
544166	1353	1372	ACCTTCCATTTTGAGACTTC	69	10325	10344	19
544167	1363	1382	ATAGAGTATAACCTTCCATT	72	10335	10354	567
544168	1367	1386	TTTTATAGAGTATAACCTTC	27	10339	10358	568
544169	1374	1393	TGGTTGATTTTATAGAGTAT	39	10346	10365	569
544170	1378	1397	ATTTTGGTTGATTTTATAGA	7	10350	10369	570
544171	1383	1402	TCAACATTTTGTTGATTTT	0	10355	10374	571
544172	1390	1409	GGATGGATCAACATTTTGGT	48	10362	10381	572
544173	1393	1412	GTTGGATGGATCAACATTTT	51	10365	10384	573
544174	1396	1415	TCTGTTGGATGGATCAACAT	46	10368	10387	574
544175	1401	1420	CTGAATCTGTTGGATGGATC	58	10373	10392	575
544176	1407	1426	AGCTTTCTGAATCTGTTGGA	57	10379	10398	576
544177	1414	1433	CATTCAAAGCTTTCTGAATC	0	10386	10405	577
544178	1417	1436	GTTCATTCAAAGCTTTCTGA	62	10389	10408	578
544179	1420	1439	TCAGTTCATTCAAAGCTTTC	21	10392	10411	579
544180	1423	1442	GCCTCAGTTCATTCAAAGCT	73	10395	10414	580
544181	1427	1446	ATTTGCCTCAGTTCATTCAA	46	10399	10418	581
544182	1431	1450	TTAAATTTGCCTCAGTTCAT	52	10403	10422	582
544183	1436	1455	GCCTTTTAAATTTGCCTCAG	66	10408	10427	583
544184	1498	1517	AGGATTTAATACCAGATTAT	31	10470	10489	584
544185	1502	1521	CTTAAGGATTTAATACCAGA	49	10474	10493	585
544186	1505	1524	TCTCTTAAGGATTTAATACC	49	10477	10496	586
544187	1546	1565	GACAGTGACTTTAAGATAAA	27	10518	10537	587
544188	1572	1591	TGTGATTGTATGTTTAATCT	30	10544	10563	588
544189	1578	1597	AGGTTATGTGATTGTATGTT	35	10550	10569	589
544190	1583	1602	CTTTAAGGTTATGTGATTGT	50	10555	10574	590
544191	1589	1608	GGTATTCTTTAAGGTTATGT	54	10561	10580	591
544192	1656	1675	ATTGATTCCCACATCACAAA	47	10628	10647	592
544193	1661	1680	CTAAAATTGATTCCCACATC	69	10633	10652	593
544194	1665	1684	CCATCTAAAATTGATTCCCA	74	10637	10656	594
544195	1771	1790	TTGTGATATTAGCTCATATG	54	10743	10762	595
544196	1794	1813	ACTAGTTTTTTTAAACTGGGA	27	10766	10785	596
544197	1820	1839	GTCAAGTTTAGAGTTTTAAC	18	10792	10811	597
544198	1826	1845	TATTTAGTCAAGTTTAGAGT	12	10798	10817	598
544199	1907	1926	TACACATACTCTGTGCTGAC	83	10879	10898	20

544200	1913	1932	GATTTTTTACACATACTCTGT	58	10885	10904	599
544201	2008	2027	CTGCTTCATTAGGTTTCATA	62	10980	10999	600

Table 6

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
337520	N/A	N/A	CAGTGTTATTCAGATTGTAC	64	6517	6536	601
337521	N/A	N/A	AGTGTCTTACCATCATGTTT	40	6776	6795	602
337525	N/A	N/A	CACCAGCCTCCTAAAGGAGA	39	10212	10231	603
544292	N/A	N/A	GAGGAGGTGAAGTCAGTGAG	35	4815	4834	604
544293	N/A	N/A	TAGAGTAGAGGAGGTGAAGT	23	4822	4841	605
544294	N/A	N/A	TGTTTGATGTGTTTGAATAC	19	4863	4882	606
544295	N/A	N/A	GAAACAACAAGGGCAAAGGC	23	4898	4917	607
544296	N/A	N/A	TGTTTGATAACGACCCTAAG	43	4974	4993	608
544297	N/A	N/A	TTTTTGGTTAAGTGACCTTG	48	5016	5035	609
544298	N/A	N/A	GTAGAAGTTTTTCAGGGATGG	23	5052	5071	610
544299	N/A	N/A	AGGAAGTAGAAGTTTTTCAGG	5	5057	5076	611
544300	N/A	N/A	AGGTGAGTGTGCAGGAGAAA	11	5085	5104	612
544301	N/A	N/A	TTAAATAAAGGTGAGTGTGC	14	5093	5112	613
544302	N/A	N/A	AGTGCAGGAATAGAAGAGAT	35	5136	5155	614
544303	N/A	N/A	CATTTTAGTGCAGGAATAGA	21	5142	5161	615
544306	N/A	N/A	CTATATTCTGGAGTATATAC	39	5216	5235	616
544307	N/A	N/A	CAGTATTCTATATTCTGGAG	72	5223	5242	617
544308	N/A	N/A	GTGCCATACAGTATTCTATA	50	5231	5250	618
544309	N/A	N/A	CTGTGTGAATATGACATTAC	52	5281	5300	619
544310	N/A	N/A	TGAGGCACACTATTTCTAGT	47	5333	5352	620
544311	N/A	N/A	GACCTTTAATTATGAGGCAC	67	5345	5364	621
544312	N/A	N/A	GAATGTTGACCTTTAATTAT	23	5352	5371	622
544313	N/A	N/A	TTGTTGAATGTTGACCTTTA	69	5357	5376	623
544314	N/A	N/A	TCTACTAAGTAACTATGTGA	37	5915	5934	624
544315	N/A	N/A	CTCTTTTCTACTAAGTAACT	31	5921	5940	625
544316	N/A	N/A	AAGGATCTATTGTAAAGTTT	24	5956	5975	626
544317	N/A	N/A	CTAGGACCTTATTTAAAAGG	24	5972	5991	627
544318	N/A	N/A	ATTCCTAGGACCTTATTTA	8	5977	5996	628
544319	N/A	N/A	TTGACAGTAAGAAAAGCAGA	28	6051	6070	629
544320	N/A	N/A	TTCTCATTGACAGTAAGAAA	56	6057	6076	630
544321	N/A	N/A	AGTTTTTCTCATTGACAGTA	50	6062	6081	631
544322	N/A	N/A	ATTGAATGATAGTTTTTCTC	42	6072	6091	632
544323	N/A	N/A	TTGGGTTTGCAATTTATTGA	36	6087	6106	633
544324	N/A	N/A	AGTGTGTTGGGTTTGCAATT	25	6093	6112	634

544325	N/A	N/A	TATTTAAGTGTGTTGGGTTT	27	6099	6118	635
544326	N/A	N/A	ATATATTCAGTAGTTTATCG	25	6145	6164	636
544327	N/A	N/A	AGATGTTGGCAGGTTGGCAA	51	6184	6203	637
544328	N/A	N/A	TCTGTAGATGTTGGCAGGTT	48	6189	6208	638
544329	N/A	N/A	TTGATAATTTTTGACCTGTA	34	6215	6234	639
544330	N/A	N/A	GGCTTTCTTGATAATTTGAT	52	6230	6249	640
544331	N/A	N/A	GTCTTACTGATCTTCAGACC	27	6282	6301	641
544332	N/A	N/A	TTAGGTCTTACTGATCTTC	14	6287	6306	642
544333	N/A	N/A	TCAGTTTTAGGTCTTACTGA	28	6292	6311	643
544334	N/A	N/A	TGATATTCTGTTTCAGATTTT	44	6326	6345	644
544335	N/A	N/A	TAGAGACTGCTTTGCTTAGA	31	6388	6407	645
544336	N/A	N/A	AGGCCAAAAGTAGAGACTGC	29	6398	6417	646
544337	N/A	N/A	GGCAAAAAAGCAGACATTGG	38	6433	6452	647
544338	N/A	N/A	AATCAGGGACATTATTTAAT	13	6473	6492	648
544339	N/A	N/A	TATTTAATCAGGGACATTAT	28	6478	6497	649
544340	N/A	N/A	CTCAAAATATTTAATCAGGG	27	6485	6504	650
544341	N/A	N/A	TACCTGTTCTCAAAATATTT	18	6493	6512	651
544342	N/A	N/A	GTACAGATTACCTGTTCTCA	68	6501	6520	652
544343	N/A	N/A	GGTGTTTGATATTTAGATAA	25	6538	6557	653
544344	N/A	N/A	TTGTCTTTCAGTTCATAATG	29	6565	6584	654
544345	N/A	N/A	ACAGTTTGTCTTTCAGTTCA	23	6570	6589	655
544346	N/A	N/A	TCTGAGCTGATAAAAGAATA	15	6657	6676	656
544347	N/A	N/A	CCCACCAAAGTGTCTTACCA	49	6784	6803	657
544348	N/A	N/A	CTTCAAGAAGGAAACCCACC	39	6798	6817	658
544349	N/A	N/A	AATAGCTTCAAGAAGGAAAC	12	6803	6822	659
544350	N/A	N/A	ACAAGTCCTAAGAATAGGGA	25	6833	6852	660
544351	N/A	N/A	GTCTAGAACAAGTCCTAAGA	53	6840	6859	661
544352	N/A	N/A	TCTAATAATCAAGTCCATAT	33	6972	6991	662
544353	N/A	N/A	ACCTTCTATATTATCTAATA	19	6985	7004	663
544354	N/A	N/A	GCATGTATCTCTTAAACAGG	50	7060	7079	664
544355	N/A	N/A	TTTCAGCATGTATCTCTTAA	79	7065	7084	21
544356	N/A	N/A	GTCCAGTGACCTTTAACTCC	69	7092	7111	665
544357	N/A	N/A	TCTTACCAAACCTATTTTCTT	28	7166	7185	666
544358	N/A	N/A	GTAATGTTTATGTTAAAGCA	17	7226	7245	667
544359	N/A	N/A	TTGTGGCAAATGTAGCATTT	52	7251	7270	668
544360	N/A	N/A	GAGATTTCACTTGACATTTT	30	7277	7296	669
544361	N/A	N/A	GGAGCTTGAGATTTCACTTG	30	7284	7303	670
544362	N/A	N/A	CATCAGATTTAGTAATAGGA	0	7315	7334	671
544363	N/A	N/A	GTTATTACATCAGATTTAGT	6	7322	7341	672
544365	N/A	N/A	CAGCAGGAATGCCTAGAATC	32	7350	7369	673
544366	N/A	N/A	CTCCTTAGACAGGTTTTACC	31	7471	7490	674

544367	N/A	N/A	GTCTATTCTCCTTAGACAGG	23	7478	7497	675
544368	N/A	N/A	ACCAGGTTAATCTTCCTAAT	71	7526	7545	22
544369	N/A	N/A	ATGAATGATTGAATGTAGTC	26	7977	7996	676
544370	N/A	N/A	ATATGAAGGCTGAGACTGCT	58	8072	8091	677
544371	N/A	N/A	ATAAATTATATGAAGGCTGA	7	8079	8098	678
544372	N/A	N/A	ATATTTAAGAACAGACATGT	12	8175	8194	679
544373	N/A	N/A	AGTTATGATCATTGTAAGCC	60	8217	8236	23
544374	N/A	N/A	ATTTGTAACAGTTACTACTT	51	8276	8295	680
544375	N/A	N/A	CACAGCTTATTTGTAACAGT	70	8284	8303	681
544376	N/A	N/A	GGAGTGGTTCTTTTCACAGC	71	8298	8317	24
544377	N/A	N/A	GTGACTAATGCTAGGAGTGG	34	8311	8330	682
544378	N/A	N/A	GAATAGAGTGACTAATGCTA	45	8318	8337	683
544379	N/A	N/A	ATGAGAGAATAGAGTGACTA	58	8324	8343	684
544380	N/A	N/A	TGGTCCTTTTAACTTCCAAT	70	8365	8384	25
544381	N/A	N/A	TATACTGTATGTCTGAGTTT	66	8387	8406	685
544382	N/A	N/A	AACTAATTCATTATAAGCCA	67	8450	8469	686
544383	N/A	N/A	GCATTGAGTTAACTAATTCA	64	8460	8479	26
544385	N/A	N/A	TTTGGATTTTAAACATCTGT	61	8528	8547	687
544386	N/A	N/A	TGTATGTGCTTTTTTGGATTT	37	8539	8558	688
544387	N/A	N/A	CATGGATTTTTTGTATGTGCT	62	8549	8568	689
544388	N/A	N/A	TCATTCATGGATTTTTTGTAT	34	8554	8573	690
544389	N/A	N/A	ACTTAGACATCATTCATGGA	55	8563	8582	691
544390	N/A	N/A	GTGAGTACTTAGACATCATT	66	8569	8588	692
544391	N/A	N/A	TTTATAAGTGAGTACTTAGA	36	8576	8595	693
544392	N/A	N/A	GTCTTCTACTTTATAAGTGA	65	8585	8604	694
544393	N/A	N/A	ATGAATGTCTTCTACTTTAT	34	8591	8610	695
544394	N/A	N/A	CAAATAGTACTGAGCATTTA	30	8627	8646	696
544395	N/A	N/A	TTAGAAGATTTGGAGCTACA	54	8718	8737	697
544396	N/A	N/A	TCACTATTAGAAGATTTGGA	37	8724	8743	698
544397	N/A	N/A	GGGTTACACTCACTATTAGA	36	8733	8752	699
544398	N/A	N/A	ACTTACCTGTCAGCCTTTTA	54	8758	8777	700
544399	N/A	N/A	CTTACCAGAATTAAGTGAGT	26	8785	8804	701
544400	N/A	N/A	AATACAAGTACAAATGGGTT	22	8810	8829	702
544401	N/A	N/A	CTGGTAAATACAAGTACAAA	55	8816	8835	703
544402	N/A	N/A	GGATTGCTGGTAAATACAAG	40	8822	8841	704
544403	N/A	N/A	TCATTTTAAGGATTGCTGGT	62	8831	8850	705
544404	N/A	N/A	AGTTAGTAGGAAGCTTCATT	56	8846	8865	706
544405	N/A	N/A	GCTATTGAGTTAGTAGGAAG	67	8853	8872	707
544407	N/A	N/A	AGCATGGTTCTTAATAACTT	67	9012	9031	708
544408	N/A	N/A	CTTTGTAGAAAAAGACAGGA	27	9062	9081	709
544409	N/A	N/A	ACCTGGCCTTTGGTATTTGC	49	9096	9115	710

544410	N/A	N/A	CATCCATATACAGTCAAGAG	80	9174	9193	27
544411	N/A	N/A	AGTCTTTATATGGATAAACT	15	9215	9234	711
544412	N/A	N/A	CGTCATTGGTAGAGGAATAT	51	9240	9259	712
544413	N/A	N/A	GATTATCCTTTCTATAATGC	48	9321	9340	713
544414	N/A	N/A	GTCTTGAATCCCTTGATCAT	40	9436	9455	714
544415	N/A	N/A	GGTGCAACTAATTGAGTTGT	27	9459	9478	715
544416	N/A	N/A	GTGTTTTTTTATTGGTGCAAC	31	9471	9490	716
544417	N/A	N/A	ATTCTCCTGAAAAGAAAAGT	24	9544	9563	717
544418	N/A	N/A	ATGCCACCACCAGCCTCCTA	73	10219	10238	718
544419	N/A	N/A	ATATCCTTTAACAAATGGGT	62	11540	11559	719
544420	N/A	N/A	GCACTATATCCTTTAACAAA	50	11545	11564	720
544421	N/A	N/A	ACTTGGGCACTATATCCTTT	68	11551	11570	721
544422	N/A	N/A	GAAACATGTCCTATGAGAGT	32	11918	11937	722
544424	N/A	N/A	TTGAGCACTTTAAGCAAAGT	7	12070	12089	723
544425	N/A	N/A	GGAATTTGAGCACTTTAAGC	34	12075	12094	724
544426	N/A	N/A	TAGATTAGACAACGTGAGT	52	12101	12120	725
544427	N/A	N/A	AAAATGAAGGTCAAGTTTGA	17	12197	12216	726
544428	N/A	N/A	GTGAAAGCAAAATGAAGGTC	33	12205	12224	727
544429	N/A	N/A	GTATTGTGAAAGCAAAATGA	39	12210	12229	728
544430	N/A	N/A	TGGAGAGTATAGTATTGTGA	35	12221	12240	729
544438	N/A	N/A	AGGAATAGAAGAGATAAATA	10	5131	5150	730
544439	N/A	N/A	TGGAGTATATACAAATAATG	30	5208	5227	731
544440	N/A	N/A	TGTTTACATTGTAGATTAAT	15	5381	5400	732
544441	N/A	N/A	CAGAATATATAATATCTTGC	57	6035	6054	733
544442	N/A	N/A	TGCAATTTATTGAATGATAG	31	6080	6099	734
544443	N/A	N/A	CATAATACATAATTTGAACC	0	6251	6270	735
544444	N/A	N/A	ATAATTTTCAGTTTTAGGTC	0	6299	6318	736
544445	N/A	N/A	TTTCAGTAATGTTTATGTTA	9	7231	7250	737
544446	N/A	N/A	AATGCCTAGAATCAATAAAA	36	7343	7362	738
544447	N/A	N/A	GTAAATATTTGTAGATTAGC	49	8003	8022	739
544448	N/A	N/A	ACAAATGTGTAATTGTTTGA	25	8101	8120	740
544449	N/A	N/A	TACTAACAAATGTGTAATTG	35	8106	8125	741
544450	N/A	N/A	TGATAAGTATATTTAAGAAC	35	8183	8202	742
544451	N/A	N/A	TTAACTTCCAATTAATTGAT	29	8357	8376	743
544452	N/A	N/A	TCTGTTATTTTATCTTGCTT	67	8513	8532	744
544453	N/A	N/A	ATCACAATCCTTTTTATTAA	18	8921	8940	745
544454	N/A	N/A	AGAGACTTGAGTAATAATAA	25	9137	9156	746
544455	N/A	N/A	AACAAAATGAAACATGTCCT	59	11926	11945	747
544127	765	784	CAGCAGGAATGCCATCATGT	33	N/A	N/A	748
544128	819	838	TGATGGCATACATGCCACTT	13	7404	7423	749
544129	828	847	TGCTGGGTCTGATGGCATAC	53	7413	7432	750

544130	832	851	GAGTTGCTGGGTCTGATGGC	22	7417	7436	751
544131	841	860	AAAACCTTGAGAGTTGCTGGG	13	7426	7445	752
544132	848	867	GACATGAAAAACTTGAGAGT	0	7433	7452	753
544133	859	878	ACATCACAGTAGACATGAAA	27	7444	7463	754
233717	889	908	TGAATTAATGTCCATGGACT	58	7876	7895	14
544134	915	934	AGTTTTGTGATCCATCTATT	46	7902	7921	755
544135	918	937	TGAAGTTTTGTGATCCATCT	54	7905	7924	756
544136	926	945	CGTTTCATTGAAGTTTTGTG	40	7913	7932	757
544137	946	965	CCATATTTGTAGTTCTCCCA	45	7933	7952	758
544138	949	968	AAACCATATTTGTAGTTCTC	41	7936	7955	759
544139	970	989	AATTCTCCATCAAGCCTCCC	43	N/A	N/A	760
233722	991	1010	ATCTTCTCTAGGCCCAACCA	65	9566	9585	761
544432	997	1016	GAGTATATCTTCTCTAGGCC	40	9572	9591	762
544140	1002	1021	CTATGGAGTATATCTTCTCT	28	9577	9596	763
544141	1008	1027	GCTTCACTATGGAGTATATC	55	9583	9602	764
544142	1013	1032	AGATTGCTTCACTATGGAGT	47	9588	9607	765
544143	1046	1065	CCAGTCTTCCAACCTCAATTC	33	9621	9640	766
544144	1052	1071	GTCTTTCCAGTCTTCCAACCT	59	9627	9646	767
544145	1055	1074	GTTGTCTTTCCAGTCTTCCA	77	9630	9649	16
544146	1059	1078	GTTTGTTGTCTTTCCAGTCT	58	9634	9653	768
544147	1062	1081	AATGTTTGTTGTCTTTCCAG	43	9637	9656	769
544148	1095	1114	CGTGATTTCCCAAGTAAAAA	57	9670	9689	770
544149	1160	1179	GTTTTCCGGGATTGCATTGG	44	9735	9754	771
544150	1165	1184	TCTTTGTTTTCCGGGATTGC	53	9740	9759	772
544151	1170	1189	CCAAATCTTTGTTTTCCGGG	57	9745	9764	773
544152	1173	1192	ACACCAAATCTTTGTTTTCC	44	9748	9767	774
544153	1178	1197	AGAAAACACCAAATCTTTGT	36	9753	9772	775
544154	1183	1202	CAAGTAGAAAACACCAAATC	29	9758	9777	776
544155	1188	1207	GATCCCAAGTAGAAAACACC	29	9763	9782	777
544156	1195	1214	GCTTTGTGATCCCAAGTAGA	71	9770	9789	17
544157	1198	1217	TTTGCTTTGTGATCCCAAGT	66	9773	9792	778
544158	1202	1221	TCCTTTTGCTTTGTGATCCC	53	9777	9796	779
544159	1208	1227	GAAGTGTCCTTTTGCTTTGT	10	9783	9802	780
544160	1246	1265	TGCCACCACCAGCCTCCTGA	65	N/A	N/A	781
544161	1253	1272	CTCATCATGCCACCACCAGC	59	10225	10244	782
544162	1269	1288	GGTTGTTTTCTCCACACTCA	74	10241	10260	18
544163	1276	1295	CCATTTAGGTTGTTTTCTCC	38	10248	10267	783
544164	1283	1302	ATATTTACCATTTAGGTTGT	13	10255	10274	784
544165	1294	1313	CTTGGTTTGTTATATTTACC	53	10266	10285	785
544166	1353	1372	ACCTTCCATTTTGAGACTTC	70	10325	10344	19
544167	1363	1382	ATAGAGTATAACCTTCCATT	69	10335	10354	786

544168	1367	1386	TTTTATAGAGTATAACCTTC	34	10339	10358	787
544169	1374	1393	TGGTTGATTTTATAGAGTAT	38	10346	10365	788
544170	1378	1397	ATTTTGGTTGATTTTATAGA	0	10350	10369	789
544171	1383	1402	TCAACATTTTGGTTGATTTT	12	10355	10374	790
544172	1390	1409	GGATGGATCAACATTTTGGT	58	10362	10381	791
544173	1393	1412	GTTGGATGGATCAACATTTT	66	10365	10384	792
544174	1396	1415	TCTGTTGGATGGATCAACAT	49	10368	10387	793
544175	1401	1420	CTGAATCTGTTGGATGGATC	60	10373	10392	794
544176	1407	1426	AGCTTTCTGAATCTGTTGGA	64	10379	10398	795
544177	1414	1433	CATTCAAAGCTTTCTGAATC	21	10386	10405	796
544178	1417	1436	G TTCATTCAAAGCTTTCTGA	60	10389	10408	797
544179	1420	1439	TCAGTTCATTCAAAGCTTTC	18	10392	10411	798
544180	1423	1442	GCCTCAGTTCATTCAAAGCT	72	10395	10414	799
544181	1427	1446	ATTTGCCTCAGTTCATTCAA	51	10399	10418	800
544182	1431	1450	TTAAATTTGCCTCAGTTCAT	48	10403	10422	801
544183	1436	1455	GCCTTTTAAATTTGCCTCAG	70	10408	10427	802
544184	1498	1517	AGGATTTAATACCAGATTAT	44	10470	10489	803
544185	1502	1521	CTTAAGGATTTAATACCAGA	47	10474	10493	804
544186	1505	1524	TCTCTTAAGGATTTAATACC	44	10477	10496	805
544187	1546	1565	GACAGTGACTTTAAGATAAA	38	10518	10537	806
544188	1572	1591	TGTGATTGTATGTTTAATCT	47	10544	10563	807
544189	1578	1597	AGGTTATGTGATTGTATGTT	43	10550	10569	808
544190	1583	1602	CTTTAAGGTTATGTGATTGT	42	10555	10574	809
544191	1589	1608	GGTATTCTTTAAGGTTATGT	60	10561	10580	810
544192	1656	1675	ATTGATTCCCACATCACAAA	46	10628	10647	811
544193	1661	1680	CTAAAATTGATTCCCACATC	65	10633	10652	812
544194	1665	1684	CCATCTAAAATTGATTCCCA	70	10637	10656	813
544195	1771	1790	TTGTGATATTAGCTCATATG	56	10743	10762	814
544196	1794	1813	ACTAGTTTTTTTAAACTGGGA	33	10766	10785	815
544197	1820	1839	GTCAAGTTTAGAGTTTTAAC	39	10792	10811	816
544198	1826	1845	TATTTAGTCAAGTTTAGAGT	21	10798	10817	817
544199	1907	1926	TACACATACTCTGTGCTGAC	80	10879	10898	20
544200	1913	1932	GATTTTACACATACTCTGT	56	10885	10904	818
544201	2008	2027	CTGCTTCATTAGGTTTCATA	65	10980	10999	819

Table 7

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
337525	N/A	N/A	CACCAGCCTCCTAAAGGAGA	58	10212	10231	820
544204	N/A	N/A	GACTTCTTAACTCTATATAT	67	3076	3095	821

544205	N/A	N/A	CTAGACTTCTTAACTCTATA	61	3079	3098	822
544206	N/A	N/A	GACCTAGACTTCTTAACTCT	54	3082	3101	823
544207	N/A	N/A	GGAAGCAGACCTAGACTTCT	58	3089	3108	824
544208	N/A	N/A	TCTGGAAGCAGACCTAGACT	48	3092	3111	825
544209	N/A	N/A	TCTTCTGGAAGCAGACCTAG	54	3095	3114	826
544210	N/A	N/A	CTAATCTTTAGGGATTAGG	57	11433	11452	827
544211	N/A	N/A	TGTATCTAATCTTTAGGGAT	53	11438	11457	828
544213	N/A	N/A	TAACTTGGGCACTATATCCT	74	11553	11572	829
544214	N/A	N/A	ATTGACAAAGGTAGGTCACC	79	11576	11595	830
544215	N/A	N/A	ATATGACATGTATATTGGAT	66	11620	11639	831
544216	N/A	N/A	TTTTGTACTTTTCTGGAACA	61	11704	11723	832
544217	N/A	N/A	TAGTCTGTGGTCCTGAAAAT	56	11748	11767	833
544218	N/A	N/A	AGCTTAGTCTGTGGTCCTGA	72	11752	11771	834
544219	N/A	N/A	GACAGCTTAGTCTGTGGTCC	74	11755	11774	835
544220	N/A	N/A	GTATTCTGGCCCTAAAAAAA	52	11789	11808	836
544221	N/A	N/A	ATTTTGGTATTCTGGCCCTA	56	11795	11814	837
544222	N/A	N/A	GAAATTGTCCAATTTTGGG	56	N/A	N/A	838
544223	N/A	N/A	TTTGCAATTTGAAATTGTCCA	61	11837	11856	839
544224	N/A	N/A	GGAAGCAACTCATATATTAA	57	11869	11888	840
544225	N/A	N/A	TATCAGAAAAAGATACCTGA	56	9821	9840	841
544226	N/A	N/A	ATAATAGCTAATAATGTGGG	59	9875	9894	842
544227	N/A	N/A	TGCAGATAATAGCTAATAAT	60	9880	9899	843
544228	N/A	N/A	TGTCATTGCAGATAATAGCT	79	9886	9905	844
544229	N/A	N/A	TAAAAGTTGTCATTGCAGAT	59	9893	9912	845
544230	N/A	N/A	CGGATTTTTTAAAAGTTGTCA	61	9901	9920	846
544231	N/A	N/A	GGGATTCGGATTTTTTAAAAG	28	9907	9926	847
544232	N/A	N/A	TTTGGGATTCGGATTTTTTAA	44	9910	9929	848
544233	N/A	N/A	ACGCTTATTTGGGATTCGGA	72	9917	9936	849
544251	N/A	N/A	TTTAAGAGATTTACAAGTCA	52	2811	2830	850
544252	N/A	N/A	GACTACCTGTTTTTAAAAGC	48	2851	2870	851
544253	N/A	N/A	TATGGTGACTACCTGTTTTT	39	2857	2876	852
544254	N/A	N/A	ACTTTGCTGTATTATAAACT	35	2890	2909	853
544255	N/A	N/A	ATTGTATTTAACTTTGCTGT	35	2900	2919	854
544256	N/A	N/A	GAGCAACTAACTTAATAGGT	42	2928	2947	855
544257	N/A	N/A	GAAATGAGCAACTAACTTAA	32	2933	2952	856
544258	N/A	N/A	AATCAAAGAAATGAGCAACT	42	2940	2959	857
544259	N/A	N/A	ACCTTCTTCCACATTGAGTT	44	2977	2996	858
544260	N/A	N/A	CACGAATGTAACCTTCTTCC	52	2987	3006	859
544261	N/A	N/A	TTAACTTGCACGAATGTAAC	45	2995	3014	860
544262	N/A	N/A	TATATATACCAATTTTGCC	43	3063	3082	861
544263	N/A	N/A	TCTTAACTCTATATATACCA	49	3072	3091	862

544264	N/A	N/A	CTTTAAGTGAAGTTACTTCT	53	3632	3651	863
544265	N/A	N/A	TCTACTTACTTTAAGTGAAG	44	3640	3659	864
544266	N/A	N/A	GAACCCTCTTTATTTTCTAC	46	3655	3674	865
544267	N/A	N/A	ACATAAACATGAACCCTCTT	50	3665	3684	866
544268	N/A	N/A	CCACATTGAAAACATAAACA	57	3676	3695	867
544269	N/A	N/A	GCATGCCTTAGAAATATTTT	23	3707	3726	868
544270	N/A	N/A	CAATGCAACAAAGTATTTCA	37	3731	3750	869
544271	N/A	N/A	CTGGAGATTATTTTTCTTGG	61	3768	3787	870
544272	N/A	N/A	TTCATATATAACATTAGGGA	14	3830	3849	871
544273	N/A	N/A	TCAGTGTTTTTCATATATAAC	32	3838	3857	872
544274	N/A	N/A	GACATAGTGTTCTAGATTGT	47	3900	3919	873
544275	N/A	N/A	CAATAGTGTAATGACATAGT	39	3912	3931	874
544276	N/A	N/A	TTACTTACCTTCAGTAATTT	35	3933	3952	875
544277	N/A	N/A	ATCTTTTCCATTTACTGTAT	39	4005	4024	876
544278	N/A	N/A	AGAAAAAGCCCAGCATATTT	23	4037	4056	877
544279	N/A	N/A	GTATGCTTCTTTCAAATAGC	46	4130	4149	878
544280	N/A	N/A	CCTTCCCCTTGTATGCTTCT	47	4140	4159	879
544281	N/A	N/A	CCTGTAACACTATCATAATC	49	4207	4226	880
544282	N/A	N/A	TGACTTACCTGATTTTCTAT	24	4384	4403	881
544283	N/A	N/A	GATGGGACATACCATTAAAA	41	4407	4426	882
544284	N/A	N/A	GTGAAAGATGGGACATACCA	54	4413	4432	883
544285	N/A	N/A	CCTGTGTGAAAGATGGGACA	27	4418	4437	884
544286	N/A	N/A	CATTGGCTGCTATGAATTAA	45	4681	4700	885
544287	N/A	N/A	GATGACATTGGCTGCTATGA	49	4686	4705	886
544288	N/A	N/A	GAGAAACATGATCTAATTTG	33	4717	4736	887
544289	N/A	N/A	ATGGAAAGCTATTGTGTGGT	42	4747	4766	888
544290	N/A	N/A	GTCTAAAGAGCCAATATGAG	39	4771	4790	889
544291	N/A	N/A	AATCTTGGTCTAAAGAGCCA	65	4778	4797	890
544361	N/A	N/A	GGAGCTTGAGATTTCACTTG	66	7284	7303	891
544362	N/A	N/A	CATCAGATTTAGTAATAGGA	61	7315	7334	892
544363	N/A	N/A	GTTATTACATCAGATTTAGT	63	7322	7341	893
544365	N/A	N/A	CAGCAGGAATGCCTAGAATC	72	7350	7369	894
544366	N/A	N/A	CTCCTTAGACAGGTTTTACC	67	7471	7490	895
544367	N/A	N/A	GTCTATTCTCCTTAGACAGG	59	7478	7497	896
544368	N/A	N/A	ACCAGGTTAATCTTCCTAAT	79	7526	7545	22
544369	N/A	N/A	ATGAATGATTGAATGTAGTC	56	7977	7996	897
544370	N/A	N/A	ATATGAAGGCTGAGACTGCT	73	8072	8091	898
544371	N/A	N/A	ATAAATTATATGAAGGCTGA	51	8079	8098	899
544372	N/A	N/A	ATATTTAAGAACAGACATGT	54	8175	8194	900
544373	N/A	N/A	AGTTATGATCATTGTAAGCC	77	8217	8236	23
544374	N/A	N/A	ATTTGTAACAGTTACTACTT	69	8276	8295	901

544375	N/A	N/A	CACAGCTTATTTGTAACAGT	72	8284	8303	902
544376	N/A	N/A	GGAGTGGTTCTTTTCACAGC	82	8298	8317	24
544377	N/A	N/A	GTGACTAATGCTAGGAGTGG	54	8311	8330	903
544378	N/A	N/A	GAATAGAGTGACTAATGCTA	55	8318	8337	904
544379	N/A	N/A	ATGAGAGAATAGAGTGACTA	66	8324	8343	905
544380	N/A	N/A	TGGTCCTTTTAACTTCCAAT	79	8365	8384	25
544381	N/A	N/A	TATACTGTATGTCTGAGTTT	72	8387	8406	906
544382	N/A	N/A	AACTAATTCATTATAAGCCA	56	8450	8469	907
544383	N/A	N/A	GCATTGAGTTAACTAATTCA	78	8460	8479	26
544385	N/A	N/A	TTTGGATTTTAAACATCTGT	73	8528	8547	908
544386	N/A	N/A	TGTATGTGCTTTTTTGGATTT	57	8539	8558	909
544387	N/A	N/A	CATGGATTTTTTGTATGTGCT	64	8549	8568	910
544388	N/A	N/A	TCATTCATGGATTTTTTGTAT	53	8554	8573	911
544389	N/A	N/A	ACTTAGACATCATTCATGGA	66	8563	8582	912
544390	N/A	N/A	GTGAGTACTTAGACATCATT	74	8569	8588	913
544391	N/A	N/A	TTTATAAGTGAGTACTTAGA	32	8576	8595	914
544392	N/A	N/A	GTCTTCTACTTTATAAGTGA	63	8585	8604	915
544393	N/A	N/A	ATGAATGTCTTCTACTTTAT	68	8591	8610	916
544394	N/A	N/A	CAAATAGTACTGAGCATTTA	53	8627	8646	917
544395	N/A	N/A	TTAGAAGATTTGGAGCTACA	55	8718	8737	918
544396	N/A	N/A	TCACTATTAGAAGATTTGGA	60	8724	8743	919
544397	N/A	N/A	GGGTTACACTCACTATTAGA	52	8733	8752	920
544398	N/A	N/A	ACTTACCTGTCAGCCTTTTA	61	8758	8777	921
544399	N/A	N/A	CTTACCAGAATTAAGTGAGT	43	8785	8804	922
544400	N/A	N/A	AATACAAGTACAAATGGGTT	29	8810	8829	923
544401	N/A	N/A	CTGGTAAATACAAGTACAAA	76	8816	8835	924
544402	N/A	N/A	GGATTGCTGGTAAATACAAG	59	8822	8841	925
544403	N/A	N/A	TCATTTTAAGGATTGCTGGT	63	8831	8850	926
544404	N/A	N/A	AGTTAGTAGGAAGCTTCATT	54	8846	8865	927
544405	N/A	N/A	GCTATTGAGTTAGTAGGAAG	63	8853	8872	928
544407	N/A	N/A	AGCATGGTTCTTAATAACTT	69	9012	9031	929
544408	N/A	N/A	CTTTGTAGAAAAAGACAGGA	45	9062	9081	930
544409	N/A	N/A	ACCTGGCCTTTGGTATTTGC	66	9096	9115	931
544410	N/A	N/A	CATCCATATACAGTCAAGAG	78	9174	9193	27
544411	N/A	N/A	AGTCTTTATATGGATAAACT	46	9215	9234	932
544412	N/A	N/A	CGTCATTGGTAGAGGAATAT	45	9240	9259	933
544413	N/A	N/A	GATTATCCTTTCTATAATGC	45	9321	9340	934
544414	N/A	N/A	GTCTTGAATCCCTTGATCAT	61	9436	9455	935
544415	N/A	N/A	GGTGCAACTAATTGAGTTGT	49	9459	9478	936
544416	N/A	N/A	GTGTTTTTTTATTGGTGCAAC	46	9471	9490	937
544417	N/A	N/A	ATTCTCCTGAAAAGAAAAGT	50	9544	9563	938

544418	N/A	N/A	ATGCCACCACCAGCCTCCTA	73	10219	10238	939
544419	N/A	N/A	ATATCCTTTAACAAATGGGT	68	11540	11559	940
544420	N/A	N/A	GCACTATATCCTTTAACAAA	74	11545	11564	941
544421	N/A	N/A	ACTTGGGCACTATATCCTTT	68	11551	11570	942
544422	N/A	N/A	GAAACATGTCCTATGAGAGT	56	11918	11937	943
544424	N/A	N/A	TTGAGCACTTTAAGCAAAGT	15	12070	12089	944
544425	N/A	N/A	GGAATTTGAGCACTTTAAGC	35	12075	12094	945
544426	N/A	N/A	TAGATTAGACAACGTGTGAGT	54	12101	12120	946
544427	N/A	N/A	AAAATGAAGGTCAAGTTTGA	45	12197	12216	947
544428	N/A	N/A	GTGAAAGCAAAAATGAAGGTC	55	12205	12224	948
544429	N/A	N/A	GTATTGTGAAAGCAAAAATGA	54	12210	12229	949
544430	N/A	N/A	TGGAGAGTATAGTATTGTGA	53	12221	12240	950
544433	N/A	N/A	GAGATTTACAAGTCAAAAAT	41	2806	2825	951
544434	N/A	N/A	ATTTAACCTTTGCTGTATTAT	29	2895	2914	952
544435	N/A	N/A	ATCAATGCTAAATGAAATCA	34	2955	2974	953
544436	N/A	N/A	TATTTTCTGGAGATTATTTT	24	3774	3793	954
544437	N/A	N/A	AAAATGAATATTGGCAATTC	34	4159	4178	955
544446	N/A	N/A	AATGCCTAGAATCAATAAAA	50	7343	7362	956
544447	N/A	N/A	GTAAATATTTGTAGATTAGC	38	8003	8022	957
544448	N/A	N/A	ACAAATGTGTAATTGTTTGA	43	8101	8120	958
544449	N/A	N/A	TACTAACAAATGTGTAATTG	59	8106	8125	959
544450	N/A	N/A	TGATAAGTATATTTAAGAAC	45	8183	8202	960
544451	N/A	N/A	TTAACTTCCAATTAATTGAT	55	8357	8376	961
544452	N/A	N/A	TCTGTTATTTTATCTTGCTT	67	8513	8532	962
544453	N/A	N/A	ATCACAATCCTTTTTATTAA	39	8921	8940	963
544454	N/A	N/A	AGAGACTTGAGTAATAATAA	43	9137	9156	964
544455	N/A	N/A	AACAAAATGAAACATGTCCT	47	11926	11945	965
544059	23	42	GATTTTCAATTTCAAGCAAC	74	3127	3146	966
337459	49	68	AGCTTAATTGTGAACATTTT	77	3153	3172	967
544060	54	73	GAAGGAGCTTAATTGTGAAC	59	3158	3177	968
544061	63	82	CAATAAAAAGAAGGAGCTTA	64	3167	3186	969
544062	66	85	GAACAATAAAAAGAAGGAGC	67	3170	3189	970
544063	85	104	CTGGAGGAAATAACTAGAGG	49	3189	3208	971
337460	88	107	ATTCTGGAGGAAATAACTAG	65	3192	3211	972
544064	112	131	TCAAATGATGAATTGTCTTG	58	3216	3235	973
544065	138	157	TTGATTTTGGCTCTGGAGAT	67	3242	3261	974
544066	145	164	GCAAATCTTGATTTTGGCTC	82	3249	3268	975
233676	148	167	ATAGCAAATCTTGATTTTGG	81	3252	3271	976
544067	156	175	CGTCTAACATAGCAAATCTT	87	3260	3279	977
544068	174	193	TGGCTAAAATTTTACATCG	66	3278	3297	978
544069	178	197	CCATTGGCTAAAATTTTAC	41	3282	3301	979

544070	184	203	AGGAGGCCATTGGCTAAAAT	36	3288	3307	980
544071	187	206	TGAAGGAGGCCATTGGCTAA	44	3291	3310	981
544072	195	214	GTCCCAACTGAAGGAGGCCA	59	3299	3318	982
544073	199	218	CCATGTCCCAACTGAAGGAG	54	3303	3322	983
544074	202	221	AGACCATGTCCCAACTGAAG	68	3306	3325	984
544075	206	225	TTTAAGACCATGTCCCAACT	51	3310	3329	985
544076	209	228	GTCTTTAAGACCATGTCCCA	64	3313	3332	986
544077	216	235	GGACAAAGTCTTTAAGACCA	45	3320	3339	987
544078	222	241	TCTTATGGACAAAGTCTTTA	40	3326	3345	988
544079	245	264	TATGTCATTAATTTGGCCCT	30	3349	3368	989
544080	270	289	GATCAAATATGTTGAGTTTT	65	3374	3393	990
233690	274	293	GACTGATCAAATATGTTGAG	75	3378	3397	991
544081	316	335	TCTTCTTTGATTTCACCTGGT	86	3420	3439	992
544082	334	353	CTTCTCAGTTCCTTTTCTTC	69	3438	3457	993
544083	337	356	GTTCTTCTCAGTTCCTTTTC	77	3441	3460	994
544084	341	360	TGTAGTTCTTCTCAGTTCCT	75	3445	3464	995
544431	345	364	TATATGTAGTTCTTCTCAGT	15	3449	3468	996
544086	348	367	GTTTATATGTAGTTCTTCTC	65	3452	3471	997
544087	352	371	TGTAGTTTATATGTAGTTCT	49	3456	3475	998
544088	356	375	GACTTGTAGTTTATATGTAG	21	3460	3479	999
544089	364	383	TCATTTTTGACTTGTAGTTT	60	3468	3487	1000
544090	369	388	CCTCTTCATTTTTGACTTGT	83	3473	3492	1001
544091	375	394	TCTTTACCTCTTCATTTTTG	75	3479	3498	1002
544092	380	399	CATATTCTTTACCTCTTCAT	77	3484	3503	1003
544093	384	403	GTGACATATTCTTTACCTCT	76	3488	3507	1004
544094	392	411	GAGTTCAAGTGACATATTCT	71	3496	3515	1005
544095	398	417	TGAGTTGAGTTCAAGTGACA	44	3502	3521	1006
544096	403	422	AGTTTTGAGTTGAGTTCAAG	33	3507	3526	1007
544097	406	425	TCAAGTTTTGAGTTGAGTTC	69	3510	3529	1008
544098	414	433	GGAGGCTTTCAAGTTTTGAG	68	3518	3537	1009
544099	423	442	TTTCTTCTAGGAGGCTTTCA	79	3527	3546	1010
544100	427	446	ATTTTTTCTTCTAGGAGGCT	63	3531	3550	1011
544101	432	451	GTAGAATTTTTTCTTCTAGG	56	3536	3555	1012
544102	462	481	GCTCTTCTAAATATTTCACT	85	3566	3585	1013
544103	474	493	AGTTAGTTAGTTGCTCTTCT	71	3578	3597	1014
544104	492	511	CAGGTTGATTTTGAATTAAG	69	3596	3615	1015
544105	495	514	TTTCAGGTTGATTTTGAATT	53	3599	3618	1016
544106	499	518	GGAGTTTCAGGTTGATTTTG	64	3603	3622	1017
544107	504	523	GTTCTGGAGTTTCAGGTTGA	74	3608	3627	1018
544108	526	545	TTAAGTGAAGTTACTTCTGG	60	3630	3649	1019
544109	555	574	TGCTATTATCTTGTTTTTCT	63	4293	4312	1020

544110	564	583	GGTCTTTGATGCTATTATCT	65	4302	4321	1021
544111	567	586	GAAGGTCTTTGATGCTATTA	49	4305	4324	1022
544112	572	591	CTGGAGAAGGTCTTTGATGC	65	4310	4329	1023
544113	643	662	CTGAGCTGATTTTCTATTTC	64	N/A	N/A	1024
337477	664	683	GGTTCTTGAATACTAGTCCT	82	6677	6696	234
544114	673	692	ATTTCTGTGGGTTCTTGAAT	57	6686	6705	1025
337478	675	694	AAATTTCTGTGGGTTCTTGA	29	6688	6707	235
544115	678	697	GAGAAATTTCTGTGGGTTCT	68	6691	6710	1026
544116	682	701	GATAGAGAAATTTCTGTGGG	54	6695	6714	1027
544117	689	708	CTTGGAAGATAGAGAAATTT	36	6702	6721	1028
337479	692	711	TGGCTTGGAAGATAGAGAAA	54	6705	6724	236
544118	699	718	GTGCTCTTGGCTTGGAAGAT	64	6712	6731	1029
544119	703	722	CTTGGTGCTCTTGGCTTGGA	68	6716	6735	1030
544120	707	726	AGTTCTTGGTGCTCTTGGCT	91	6720	6739	15
233710	710	729	AGTAGTTCTTGGTGCTCTTG	80	6723	6742	233
544121	713	732	GGGAGTAGTTCTTGGTGCTC	76	6726	6745	1031
544122	722	741	CTGAAGAAAGGGAGTAGTTC	55	6735	6754	1032
544123	752	771	ATCATGTTTTACATTTCTTA	52	6765	6784	1033
544124	755	774	GCCATCATGTTTTACATTTC	61	N/A	N/A	1034
544125	759	778	GAATGCCATCATGTTTTACA	30	N/A	N/A	1035
544126	762	781	CAGGAATGCCATCATGTTTT	34	N/A	N/A	1036
337487	804	823	CACTTGTATGTTACCTCTG	83	7389	7408	28
233717	889	908	TGAATTAATGTCCATGGACT	75	7876	7895	14
544202	2081	2100	AAAGTCAATGTGACTTAGTA	70	11053	11072	1037
544203	2104	2123	AAGGTATAGTGATACCTCAT	84	11076	11095	1038

Table 8

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
560535	N/A	N/A	ACTGTTTTCTTCTGGAAGCA	0	3102	3121	1039
560536	N/A	N/A	AAATAAGGTATAGTGATACC	0	11080	11099	1040
560537	N/A	N/A	ACAAATAAGGTATAGTGATA	1	11082	11101	1041
560538	N/A	N/A	TAACAAATAAGGTATAGTGA	0	11084	11103	1042
560539	N/A	N/A	TTTAACAAATAAGGTATAGT	16	11086	11105	1043
560540	N/A	N/A	ATATATTTTAACAAATAAGG	0	11092	11111	1044
560541	N/A	N/A	CAGTATATATTTTAACAAAT	0	11096	11115	1045
560542	N/A	N/A	TACAGTATATATTTTAACAA	0	11098	11117	1046
560543	N/A	N/A	TATACAGTATATATTTTAAC	0	11100	11119	1047
560544	N/A	N/A	ATAGTATTAAGTGTTAAAAT	0	11130	11149	1048

560545	N/A	N/A	TCATAGTATTAAGTGTTAAA	0	11132	11151	1049
560546	N/A	N/A	GTTTTCATAGTATTAAGTGT	26	11136	11155	1050
560547	N/A	N/A	ATTATTTGTTTTTCATAGTAT	0	11143	11162	1051
560548	N/A	N/A	CTTTACAATTATTTGTTTTTC	0	11150	11169	1052
560549	N/A	N/A	ATTCCTTTACAATTATTTGT	21	11154	11173	1053
560550	N/A	N/A	AGATTCCTTTACAATTATTT	18	11156	11175	1054
560551	N/A	N/A	CAAGATTCCTTTACAATTAT	21	11158	11177	1055
560552	N/A	N/A	GACAAGATTCCTTTACAATT	55	11160	11179	1056
560553	N/A	N/A	CTGACAAGATTCCTTTACAA	47	11162	11181	1057
560554	N/A	N/A	AATCTGACAAGATTCCTTTA	52	11165	11184	1058
560555	N/A	N/A	GTAATCTGACAAGATTCCTT	56	11167	11186	1059
560556	N/A	N/A	CTGTAATCTGACAAGATTCC	51	11169	11188	1060
560557	N/A	N/A	TACTGTAATCTGACAAGATT	18	11171	11190	1061
560558	N/A	N/A	CTTACTGTAATCTGACAAGA	33	11173	11192	1062
560559	N/A	N/A	TTCTTACTGTAATCTGACAA	47	11175	11194	1063
560560	N/A	N/A	CATTCTTACTGTAATCTGAC	65	11177	11196	1064
560561	N/A	N/A	TTCATTCTTACTGTAATCTG	54	11179	11198	1065
560562	N/A	N/A	TGTTCACTTACTGTAATC	44	11181	11200	1066
560563	N/A	N/A	TATGTTCACTTACTGTAA	39	11183	11202	1067
560564	N/A	N/A	AATATGTTCACTTACTGT	0	11185	11204	1068
560565	N/A	N/A	ACAAATATGTTCACTTAC	3	11188	11207	1069
560566	N/A	N/A	CCACAAATATGTTCACTT	75	11190	11209	42
560567	N/A	N/A	TGCCACAAATATGTTCACTC	80	11192	11211	43
560568	N/A	N/A	CGATGCCACAAATATGTTCA	64	11195	11214	1070
560569	N/A	N/A	CTCGATGCCACAAATATGTT	65	11197	11216	1071
560570	N/A	N/A	AACTCGATGCCACAAATATG	46	11199	11218	1072
560571	N/A	N/A	TTAACTCGATGCCACAAATA	52	11201	11220	1073
560572	N/A	N/A	CTTTAACTCGATGCCACAAA	66	11203	11222	1074
560573	N/A	N/A	AACTTTAACTCGATGCCACA	53	11205	11224	1075
560574	N/A	N/A	TAAACTTTAACTCGATGCCA	72	11207	11226	44
560575	N/A	N/A	AATATAAACTTTAACTCGAT	6	11211	11230	1076
560576	N/A	N/A	GAAATATAAACTTTAACTCG	17	11213	11232	1077
560577	N/A	N/A	GGGAAATATAAACTTTAACT	0	11215	11234	1078
560578	N/A	N/A	GAATCACAGCATATTTAGGG	46	11233	11252	1079
560579	N/A	N/A	TAGAATCACAGCATATTTAG	32	11235	11254	1080
560580	N/A	N/A	GTATTAGAATCACAGCATAT	51	11239	11258	1081
560581	N/A	N/A	ATGTATTAGAATCACAGCAT	64	11241	11260	1082
560582	N/A	N/A	GAATGTATTAGAATCACAGC	44	11243	11262	1083
560583	N/A	N/A	ACGAATGTATTAGAATCACA	44	11245	11264	1084
560584	N/A	N/A	ACACGAATGTATTAGAATCA	41	11247	11266	1085
560585	N/A	N/A	CTACACGAATGTATTAGAAT	15	11249	11268	1086

560586	N/A	N/A	ACCTACACGAATGTATTAGA	37	11251	11270	1087
560587	N/A	N/A	AAACCTACACGAATGTATTA	3	11253	11272	1088
560588	N/A	N/A	GAAAACCTACACGAATGTAT	27	11255	11274	1089
560589	N/A	N/A	TTGAAAACCTACACGAATGT	19	11257	11276	1090
560590	N/A	N/A	ACTTGAAAACCTACACGAAT	21	11259	11278	1091
560591	N/A	N/A	CTACTTGAAAACCTACACGA	43	11261	11280	1092
560592	N/A	N/A	TATTTCTACTTGAAAACCTA	29	11266	11285	1093
560593	N/A	N/A	TTTATTTCTACTTGAAAACC	2	11268	11287	1094
560594	N/A	N/A	GGTTTATTTCTACTTGAAAA	27	11270	11289	1095
560595	N/A	N/A	GAGGTTTATTTCTACTTGAA	45	11272	11291	1096
560596	N/A	N/A	ACGAGGTTTATTTCTACTTG	75	11274	11293	45
560597	N/A	N/A	TTACGAGGTTTATTTCTACT	49	11276	11295	1097
560598	N/A	N/A	TGTTACGAGGTTTATTTCTA	39	11278	11297	1098
560599	N/A	N/A	CTTGTTACGAGGTTTATTTC	32	11280	11299	1099
560600	N/A	N/A	AACTTGTTACGAGGTTTATT	27	11282	11301	1100
560601	N/A	N/A	GTAACCTTGTTACGAGGTTTA	55	11284	11303	1101
560602	N/A	N/A	CAGTAACCTTGTTACGAGGTT	51	11286	11305	1102
560603	N/A	N/A	TTCAGTAACCTTGTTACGAGG	40	11288	11307	1103
560604	N/A	N/A	CGTTCAGTAACCTTGTTACGA	53	11290	11309	1104
560605	N/A	N/A	CTTGTCAGGCTGTTTAAACG	24	11308	11327	1105
560606	N/A	N/A	TGCTTGTCAGGCTGTTTAAA	46	11310	11329	1106
560607	N/A	N/A	CATGCTTGTCAGGCTGTTTA	72	11312	11331	46
560608	N/A	N/A	TACATGCTTGTCAGGCTGTT	72	11314	11333	47
560609	N/A	N/A	TATACATGCTTGTCAGGCTG	63	11316	11335	1107
560610	N/A	N/A	TATATACATGCTTGTCAGGC	55	11318	11337	1108
560611	N/A	N/A	CATATATACATGCTTGTCAG	47	11320	11339	1109
560235	2	21	TGGAACCTGTTTTCTTCTGGA	43	3106	3125	1110
337526	4	23	CGTGGAACCTGTTTTCTTCTG	54	3108	3127	1111
560236	25	44	TTGATTTTCAATTTCAAGCA	91	3129	3148	30
560237	27	46	TCTTGATTTTCAATTTCAAG	33	3131	3150	1112
560238	32	51	TTTTATCTTGATTTTCAATT	0	3136	3155	1113
560239	35	54	CATTTTTATCTTGATTTTCA	6	3139	3158	1114
560240	43	62	ATTGTGAACATTTTTATCTT	0	3147	3166	1115
560241	45	64	TAATTGTGAACATTTTTATC	20	3149	3168	1116
560242	56	75	AAGAAGGAGCTTAATTGTGA	39	3160	3179	1117
560243	58	77	AAAAGAAGGAGCTTAATTGT	17	3162	3181	1118
560244	60	79	TAAAAAGAAGGAGCTTAATT	0	3164	3183	1119
560245	75	94	TAAC TAGAGGAACAATAAAA	37	3179	3198	1120
560246	77	96	AATAACTAGAGGAACAATAA	3	3181	3200	1121
560247	79	98	GAAATAACTAGAGGAACAAT	13	3183	3202	1122
560248	81	100	AGGAAATAACTAGAGGAACA	28	3185	3204	1123

560249	83	102	GGAGGAAATAACTAGAGGAA	12	3187	3206	1124
560250	90	109	CAATTCTGGAGGAAATAACT	34	3194	3213	1125
560251	92	111	ATCAATTCTGGAGGAAATAA	32	3196	3215	1126
560252	96	115	CTTGATCAATTCTGGAGGAA	15	3200	3219	1127
560253	98	117	GTCTTGATCAATTCTGGAGG	53	3202	3221	1128
560254	100	119	TTGTCTTGATCAATTCTGGA	48	3204	3223	1129
560255	102	121	AATTGTCTTGATCAATTCTG	23	3206	3225	1130
560256	104	123	TGAATTGTCTTGATCAATTC	14	3208	3227	1131
560257	106	125	GATGAATTGTCTTGATCAAT	46	3210	3229	1132
560258	108	127	ATGATGAATTGTCTTGATCA	33	3212	3231	1133
560259	110	129	AAATGATGAATTGTCTTGAT	24	3214	3233	1134
560260	114	133	AATCAAATGATGAATTGTCT	25	3218	3237	1135
560261	116	135	AGAATCAAATGATGAATTGT	16	3220	3239	1136
560262	119	138	TAGAGAATCAAATGATGAAT	7	3223	3242	1137
560263	126	145	CTGGAGATAGAGAATCAAAT	40	3230	3249	1138
560264	128	147	CTCTGGAGATAGAGAATCAA	51	3232	3251	1139
560265	130	149	GGCTCTGGAGATAGAGAATC	63	3234	3253	31
560266	132	151	TTGGCTCTGGAGATAGAGAA	49	3236	3255	1140
560267	135	154	ATTTTGGCTCTGGAGATAGA	49	3239	3258	1141
560268	140	159	TCTTGATTTTGGCTCTGGAG	69	3244	3263	32
560269	142	161	AATCTTGATTTTGGCTCTGG	53	3246	3265	1142
560270	150	169	ACATAGCAAATCTTGATTTT	25	3254	3273	1143
560271	152	171	TAACATAGCAAATCTTGATT	0	3256	3275	1144
560272	154	173	TCTAACATAGCAAATCTTGA	53	3258	3277	1145
560273	176	195	ATTGGCTAAAATTTTACAT	12	3280	3299	1146
560274	180	199	GGCCATTGGCTAAAATTTT	34	3284	3303	1147
560275	182	201	GAGGCCATTGGCTAAAATTT	26	3286	3305	1148
560276	189	208	ACTGAAGGAGGCCATTGGCT	51	3293	3312	1149
560277	191	210	CAACTGAAGGAGGCCATTGG	28	3295	3314	1150
560278	193	212	CCCAACTGAAGGAGGCCATT	10	3297	3316	1151
560279	197	216	ATGTCCCAACTGAAGGAGGC	0	3301	3320	1152
560280	204	223	TAAGACCATGTCCCAACTGA	13	3308	3327	1153
560281	211	230	AAGTCTTTAAGACCATGTCC	4	3315	3334	1154
560282	213	232	CAAAGTCTTTAAGACCATGT	24	3317	3336	1155
560283	219	238	TATGGACAAAGTCTTTAAGA	8	3323	3342	1156
560284	224	243	CGTCTTATGGACAAAGTCTT	11	3328	3347	1157
560285	242	261	GTCATTAATTTGGCCCTTCG	57	3346	3365	33
560286	247	266	AATATGTCATTAATTTGGCC	0	3351	3370	1158
560287	249	268	GAAATATGTCATTAATTTGG	0	3353	3372	1159
560288	252	271	TTTGAAATATGTCATTAATT	4	3356	3375	1160
560289	256	275	AGTTTTTGAATATGTCATT	7	3360	3379	1161

560290	258	277	TGAGTTTTTTGAAATATGTCA	41	3362	3381	1162
560291	267	286	CAAATATGTTGAGTTTTTTGA	30	3371	3390	1163
560292	272	291	CTGATCAAATATGTTGAGTT	32	3376	3395	1164
560293	276	295	AAGACTGATCAAATATGTTG	37	3380	3399	1165
560294	280	299	TAAAAAGACTGATCAAATAT	0	3384	3403	1166
560295	282	301	CATAAAAAGACTGATCAAAT	6	3386	3405	1167
560296	284	303	ATCATAAAAAGACTGATCAA	10	3388	3407	1168
560297	287	306	TAGATCATAAAAAGACTGAT	0	3391	3410	1169
560298	289	308	GATAGATCATAAAAAGACTG	21	3393	3412	1170
560299	291	310	GCGATAGATCATAAAAAGAC	20	3395	3414	1171
560300	293	312	CAGCGATAGATCATAAAAAG	16	3397	3416	1172
560301	295	314	TGCAGCGATAGATCATAAAA	38	3399	3418	1173
560302	297	316	TTTGCAGCGATAGATCATAA	32	3401	3420	1174
560303	299	318	GGTTTGCAGCGATAGATCAT	34	3403	3422	1175
560304	301	320	CTGGTTTGCAGCGATAGATC	25	3405	3424	1176
560305	303	322	CACTGGTTTGCAGCGATAGA	28	3407	3426	1177
560306	305	324	TTCCTGGTTTGCAGCGATA	65	3409	3428	34
560307	307	326	ATTTCCTGGTTTGCAGCGA	23	3411	3430	1178
560308	310	329	TTGATTTCCTGGTTTGCAG	5	3414	3433	1179
560309	318	337	CTTCTTCTTTGATTTCCTG	25	3422	3441	1180
560310	327	346	GTTCCCTTTTCTTCTTCTTG	19	3431	3450	1181
544120	707	726	AGTTCTTGGTGCTCTTGGCT	77	6720	6739	15
560311	801	820	TTGTATGTTACCTCTGTTA	25	7386	7405	1182
560312	802	821	CTTGTATGTTACCTCTGTT	37	7387	7406	1183
337487	804	823	CACTTGTATGTTACCTCTG	83	7389	7408	28
560313	806	825	GCCACTTGTATGTTACCTC	40	7391	7410	1184
560314	807	826	TGCCACTTGTATGTTACCT	56	7392	7411	1185
560315	808	827	ATGCCACTTGTATGTTACC	39	7393	7412	1186
337488	809	828	CATGCCACTTGTATGTTAC	19	7394	7413	1187
560316	810	829	ACATGCCACTTGTATGTTCA	26	7395	7414	1188
560317	811	830	TACATGCCACTTGTATGTTT	20	7396	7415	1189
560318	814	833	GCATACATGCCACTTGTATG	2	7399	7418	1190
560319	815	834	GGCATAACATGCCACTTGTAT	24	7400	7419	1191
560320	816	835	TGGCATAACATGCCACTTGTA	7	7401	7420	1192
560321	817	836	ATGGCATAACATGCCACTTGT	0	7402	7421	1193
560322	821	840	TCTGATGGCATAACATGCCAC	26	7406	7425	1194
560323	822	841	GTCTGATGGCATAACATGCCA	39	7407	7426	1195
560324	824	843	GGGTCTGATGGCATAACATGC	15	7409	7428	1196
560325	825	844	TGGGTCTGATGGCATAACATG	23	7410	7429	1197
560326	826	845	CTGGGTCTGATGGCATAACAT	9	7411	7430	1198
560327	834	853	GAGAGTTGCTGGGTCTGATG	0	7419	7438	1199

560328	835	854	TGAGAGTTGCTGGGTCTGAT	2	7420	7439	1200
560329	836	855	TTGAGAGTTGCTGGGTCTGA	35	7421	7440	1201
560330	837	856	CTTGAGAGTTGCTGGGTCTG	17	7422	7441	1202
560331	838	857	ACTTGAGAGTTGCTGGGTCT	0	7423	7442	1203
560332	839	858	AACTTGAGAGTTGCTGGGTC	13	7424	7443	1204
560333	843	862	GAAAAAAGTTGAGAGTTGCTG	22	7428	7447	1205
560334	844	863	TGAAAAAAGTTGAGAGTTGCT	16	7429	7448	1206
560335	845	864	ATGAAAAAAGTTGAGAGTTGC	10	7430	7449	1207
560336	846	865	CATGAAAAAAGTTGAGAGTTG	2	7431	7450	1208
560337	851	870	GTAGACATGAAAAAAGTTGAG	13	7436	7455	1209
560338	853	872	CAGTAGACATGAAAAAAGTTG	3	7438	7457	1210
560339	861	880	TAACATCACAGTAGACATGA	30	7446	7465	1211
560340	862	881	ATAACATCACAGTAGACATG	34	7447	7466	1212
560341	863	882	TATAACATCACAGTAGACAT	0	7448	7467	1213
560342	864	883	ATATAACATCACAGTAGACA	10	7449	7468	1214
560343	865	884	GATATAACATCACAGTAGAC	9	7450	7469	1215
560344	866	885	TGATATAACATCACAGTAGA	20	7451	7470	1216
337490	867	886	CTGATATAACATCACAGTAG	24	7452	7471	1217
560345	868	887	CCTGATATAACATCACAGTA	36	7453	7472	1218
560346	869	888	ACCTGATATAACATCACAGT	35	7454	7473	1219
560347	870	889	TACCTGATATAACATCACAG	26	7455	7474	1220
560348	871	890	CTACCTGATATAACATCACA	38	N/A	N/A	1221
560349	872	891	ACTACCTGATATAACATCAC	12	N/A	N/A	1222
560350	873	892	GACTACCTGATATAACATCA	28	N/A	N/A	1223
560351	874	893	GGACTACCTGATATAACATC	15	N/A	N/A	1224
560352	875	894	TGGACTACCTGATATAACAT	0	N/A	N/A	1225
560353	876	895	ATGGACTACCTGATATAACA	11	N/A	N/A	1226
337491	877	896	CATGGACTACCTGATATAAC	3	N/A	N/A	1227
560354	878	897	CCATGGACTACCTGATATAA	0	N/A	N/A	1228
560355	879	898	TCCATGGACTACCTGATATA	13	N/A	N/A	1229
560356	880	899	GTCCATGGACTACCTGATAT	50	N/A	N/A	1230
560357	881	900	TGTCCATGGACTACCTGATA	12	N/A	N/A	1231
560358	882	901	ATGTCCATGGACTACCTGAT	20	N/A	N/A	1232
560359	883	902	AATGTCCATGGACTACCTGA	16	7870	7889	1233
560360	884	903	TAATGTCCATGGACTACCTG	26	7871	7890	1234
560361	885	904	TTAATGTCCATGGACTACCT	31	7872	7891	1235
560362	886	905	ATTAATGTCCATGGACTACC	42	7873	7892	1236
560363	887	906	AATTAATGTCCATGGACTAC	21	7874	7893	1237
560364	891	910	GTTGAATTAATGTCCATGGA	18	7878	7897	1238
560365	892	911	TGTTGAATTAATGTCCATGG	36	7879	7898	1239
560366	893	912	ATGTTGAATTAATGTCCATG	13	7880	7899	1240

560367	894	913	GATGTTGAATTAATGTCCAT	14	7881	7900	1241
560368	895	914	CGATGTTGAATTAATGTCCA	30	7882	7901	1242
560369	896	915	TCGATGTTGAATTAATGTCC	29	7883	7902	1243
560370	897	916	TTCGATGTTGAATTAATGTC	4	7884	7903	1244
560371	898	917	ATTCGATGTTGAATTAATGT	22	7885	7904	1245
560372	899	918	TATTCGATGTTGAATTAATG	0	7886	7905	1246
560373	900	919	CTATTCGATGTTGAATTAAT	0	7887	7906	1247
337492	901	920	TCTATTCGATGTTGAATTAA	59	7888	7907	29
560374	902	921	ATCTATTCGATGTTGAATTA	18	7889	7908	1248
560375	903	922	CATCTATTCGATGTTGAATT	27	7890	7909	1249
560376	904	923	CCATCTATTCGATGTTGAAT	40	7891	7910	1250
560377	905	924	TCCATCTATTCGATGTTGAA	23	7892	7911	1251
560378	906	925	ATCCATCTATTCGATGTTGA	47	7893	7912	1252
560379	907	926	GATCCATCTATTCGATGTTG	46	7894	7913	1253
560380	908	927	TGATCCATCTATTCGATGTT	16	7895	7914	1254
560381	909	928	GTGATCCATCTATTCGATGT	24	7896	7915	1255
560382	910	929	TGTGATCCATCTATTCGATG	21	7897	7916	1256
560383	911	930	TTGTGATCCATCTATTCGAT	19	7898	7917	1257
560384	1273	1292	TTTAGGTTGTTTTCTCCACA	35	10245	10264	1258
560385	1274	1293	ATTTAGGTTGTTTTCTCCAC	34	10246	10265	1259
560386	1278	1297	TACCATTTAGGTTGTTTTCT	15	10250	10269	1260
560387	1286	1305	GTTATATTTACCATTTAGGT	20	10258	10277	1261
560388	1287	1306	TGTTATATTTACCATTTAGG	17	10259	10278	1262
560389	1288	1307	TTGTTATATTTACCATTTAG	21	10260	10279	1263
560390	1289	1308	TTTGTTATATTTACCATTTA	4	10261	10280	1264
560391	1292	1311	TGGTTTGTTATATTTACCAT	23	10264	10283	1265
560392	1296	1315	CTCTTGTTTGTATATTTA	63	10268	10287	1266
560393	1297	1316	GCTCTTGTTTGTATATTT	61	10269	10288	1267
560394	1298	1317	TGCTCTTGTTTGTATATT	51	10270	10289	1268
560395	1301	1320	TTTTGCTCTTGTTTGTAT	2	10273	10292	1269
560396	1302	1321	ATTTTGCTCTTGTTTGTTA	0	10274	10293	1270
560397	1303	1322	GATTTTGCTCTTGTTTGT	0	10275	10294	1271
560398	1304	1323	AGATTTTGCTCTTGTTTGT	16	10276	10295	1272
560399	1305	1324	TAGATTTTGCTCTTGTTTG	28	10277	10296	1273
560400	1307	1326	CTTAGATTTTGCTCTTGTT	69	10279	10298	35
560401	1308	1327	GCTTAGATTTTGCTCTTGGT	77	10280	10299	36
560402	1309	1328	GGCTTAGATTTTGCTCTTGG	72	10281	10300	37
560403	1315	1334	CTCTCTGGCTTAGATTTTGC	38	10287	10306	1274
560404	1316	1335	CCTCTCTGGCTTAGATTTTG	49	10288	10307	1275
560405	1317	1336	TCCTCTCTGGCTTAGATTTT	46	10289	10308	1276
560406	1321	1340	CTTCTCCTCTCTGGCTTAGA	40	10293	10312	1277

560407	1322	1341	TCTTCTCCTCTCTGGCTTAG	57	10294	10313	1278
560408	1323	1342	CTCTTCTCCTCTCTGGCTTA	40	10295	10314	1279
337505	1328	1347	TAATCCTCTTCTCCTCTCTG	28	10300	10319	1280
560409	1329	1348	ATAATCCTCTTCTCCTCTCT	30	10301	10320	1281
560410	1330	1349	GATAATCCTCTTCTCCTCTC	9	10302	10321	1282
560411	1331	1350	AGATAATCCTCTTCTCCTCT	23	10303	10322	1283
560412	1332	1351	AAGATAATCCTCTTCTCCTC	12	10304	10323	1284
560413	1333	1352	CAAGATAATCCTCTTCTCCT	40	10305	10324	1285
560414	1334	1353	CCAAGATAATCCTCTTCTCC	52	10306	10325	1286
560415	1335	1354	TCCAAGATAATCCTCTTCTC	56	10307	10326	1287
560416	1336	1355	TTCCAAGATAATCCTCTTCT	60	10308	10327	1288
560417	1337	1356	CTTCCAAGATAATCCTCTTC	58	10309	10328	1289
560418	1338	1357	ACTTCCAAGATAATCCTCTT	31	10310	10329	1290
560419	1339	1358	GACTTCCAAGATAATCCTCT	52	10311	10330	1291
560420	1340	1359	AGACTTCCAAGATAATCCTC	49	10312	10331	1292
560421	1341	1360	GAGACTTCCAAGATAATCCT	56	10313	10332	1293
337506	1342	1361	TGAGACTTCCAAGATAATCC	49	10314	10333	1294
560422	1343	1362	TTGAGACTTCCAAGATAATC	34	10315	10334	1295
560423	1344	1363	TTTGAGACTTCCAAGATAAT	14	10316	10335	1296
560424	1345	1364	TTTTGAGACTTCCAAGATAA	27	10317	10336	1297
560425	1346	1365	ATTTTGAGACTTCCAAGATA	23	10318	10337	1298
560426	1348	1367	CCATTTTGAGACTTCCAAGA	40	10320	10339	1299
560427	1351	1370	CTTCCATTTTGAGACTTCCA	58	10323	10342	1300
560428	1355	1374	TAACCTTCCATTTTGAGACT	36	10327	10346	1301
560429	1356	1375	ATAACCTTCCATTTTGAGAC	51	10328	10347	1302
560430	1357	1376	TATAACCTTCCATTTTGAGA	33	10329	10348	1303
560431	1358	1377	GTATAACCTTCCATTTTGAG	53	10330	10349	1304
337508	1360	1379	GAGTATAACCTTCCATTTTG	28	10332	10351	1305
560432	1361	1380	AGAGTATAACCTTCCATTTT	50	10333	10352	1306
560433	1365	1384	TTATAGAGTATAACCTTCCA	63	10337	10356	1307
560434	1369	1388	GATTTTATAGAGTATAACCT	31	10341	10360	1308
560435	1370	1389	TGATTTTATAGAGTATAACC	6	10342	10361	1309
560436	1371	1390	TTGATTTTATAGAGTATAAC	14	10343	10362	1310
560437	1372	1391	GTTGATTTTATAGAGTATAA	2	10344	10363	1311
560438	1376	1395	TTTGGTTGATTTTATAGAGT	20	10348	10367	1312
560439	1386	1405	GGATCAACATTTTGGTTGAT	42	10358	10377	1313
560440	1387	1406	TGGATCAACATTTTGGTTGA	10	10359	10378	1314
560441	1388	1407	ATGGATCAACATTTTGGTTG	34	10360	10379	1315
560442	1398	1417	AATCTGTTGGATGGATCAAC	52	10370	10389	1316
560443	1399	1418	GAATCTGTTGGATGGATCAA	47	10371	10390	1317
560444	1403	1422	TTCTGAATCTGTTGGATGGA	30	10375	10394	1318

560445	1404	1423	TTTCTGAATCTGTTGGATGG	34	10376	10395	1319
560446	1405	1424	CTTTCTGAATCTGTTGGATG	50	10377	10396	1320
560447	1409	1428	AAAGCTTTCTGAATCTGTTG	29	10381	10400	1321
560448	1425	1444	TTGCCTCAGTTCATTCAAAG	38	10397	10416	1322
560449	1429	1448	AAATTTGCCTCAGTTCATTC	27	10401	10420	1323
560450	1434	1453	CTTTTAAATTTGCCTCAGTT	34	10406	10425	1324
560451	1440	1459	TATTGCCTTTTAAATTTGCC	21	10412	10431	1325
560452	1441	1460	TTATTGCCTTTTAAATTTGC	23	10413	10432	1326
560453	1446	1465	TTAAATTATTGCCTTTTAAA	1	10418	10437	1327
560454	1447	1466	TTTAAATTATTGCCTTTTAA	1	10419	10438	1328
560455	1448	1467	GTTTAAATTATTGCCTTTTA	48	10420	10439	1329
560456	1449	1468	TGTTTAAATTATTGCCTTTT	25	10421	10440	1330
560457	1450	1469	ATGTTTAAATTATTGCCTTT	0	10422	10441	1331
560458	1704	1723	TTTAATAAGTTCACCTATTG	26	10676	10695	1332
560459	1705	1724	ATTTAATAAGTTCACCTATT	26	10677	10696	1333
560460	1706	1725	TATTTAATAAGTTCACCTAT	16	10678	10697	1334
560461	1707	1726	TTATTTAATAAGTTCACCTA	4	10679	10698	1335
560462	1708	1727	GTTATTTAATAAGTTCACCT	36	10680	10699	1336
560463	1709	1728	AGTTATTTAATAAGTTCACC	0	10681	10700	1337
560464	1712	1731	AAAAGTTATTTAATAAGTTC	12	10684	10703	1338
560465	1719	1738	TATTTAGAAAAGTTATTTAA	0	10691	10710	1339
560466	1738	1757	TAAAAGTCTCTAAATTTTTT	0	10710	10729	1340
560467	1739	1758	ATAAAAGTCTCTAAATTTTT	0	10711	10730	1341
560468	1740	1759	AATAAAAGTCTCTAAATTTTT	25	10712	10731	1342
560469	1760	1779	GCTCATATGATGCCTTTTAA	77	10732	10751	38
560470	1761	1780	AGCTCATATGATGCCTTTTA	73	10733	10752	39
560471	1762	1781	TAGCTCATATGATGCCTTTT	67	10734	10753	40
560472	1763	1782	TTAGCTCATATGATGCCTTT	42	10735	10754	1343
560473	1764	1783	ATTAGCTCATATGATGCCTT	61	10736	10755	1344
560474	1765	1784	TATTAGCTCATATGATGCCT	55	10737	10756	41
560475	1766	1785	ATATTAGCTCATATGATGCC	42	10738	10757	1345
560476	1767	1786	GATATTAGCTCATATGATGC	36	10739	10758	1346
560477	1768	1787	TGATATTAGCTCATATGATG	21	10740	10759	1347
560478	1769	1788	GTGATATTAGCTCATATGAT	40	10741	10760	1348
560479	1776	1795	GAAAGTTGTGATATTAGCTC	43	10748	10767	1349
560480	1777	1796	GGAAAGTTGTGATATTAGCT	19	10749	10768	1350
560481	1778	1797	GGGAAAGTTGTGATATTAGC	17	10750	10769	1351
560482	1779	1798	TGGGAAAGTTGTGATATTAG	29	10751	10770	1352
560483	1780	1799	CTGGGAAAGTTGTGATATTA	35	10752	10771	1353
560484	1781	1800	ACTGGGAAAGTTGTGATATT	25	10753	10772	1354
560485	1782	1801	AACTGGGAAAGTTGTGATAT	12	10754	10773	1355

560486	1783	1802	AAACTGGGAAAGTTGTGATA	21	10755	10774	1356
560487	1784	1803	TAAACTGGGAAAGTTGTGAT	22	10756	10775	1357
560488	1785	1804	TTAAACTGGGAAAGTTGTGA	12	10757	10776	1358
560489	1786	1805	TTTAAACTGGGAAAGTTGTG	22	10758	10777	1359
560490	1787	1806	TTTTAAACTGGGAAAGTTGT	23	10759	10778	1360
560491	1790	1809	GTTTTTTTAAACTGGGAAAGT	1	10762	10781	1361
560492	1791	1810	AGTTTTTTTAAACTGGGAAAG	0	10763	10782	1362
560493	1792	1811	TAGTTTTTTTAAACTGGGAAA	0	10764	10783	1363
560494	1796	1815	GTACTAGTTTTTTTAAACTGG	23	10768	10787	1364
560495	1799	1818	AGAGTACTAGTTTTTTTAAAC	0	10771	10790	1365
560496	1801	1820	CAAGAGTACTAGTTTTTTTAA	0	10773	10792	1366
560497	1806	1825	TTTAACAAGAGTACTAGTTT	21	10778	10797	1367
560498	1807	1826	TTTTAACAAGAGTACTAGTT	19	10779	10798	1368
560499	1808	1827	GTTTTAACAAGAGTACTAGT	37	10780	10799	1369
560500	1809	1828	AGTTTTAACAAGAGTACTAG	20	10781	10800	1370
560501	1810	1829	GAGTTTTAACAAGAGTACTA	21	10782	10801	1371
560502	1811	1830	AGAGTTTTAACAAGAGTACT	0	10783	10802	1372
560503	1814	1833	TTTAGAGTTTTTAACAAGAGT	0	10786	10805	1373
560504	1815	1834	GTTTAGAGTTTTTAACAAGAG	18	10787	10806	1374
560505	1817	1836	AAGTTTAGAGTTTTTAACAAG	9	10789	10808	1375
560506	1818	1837	CAAGTTTAGAGTTTTTAACAA	1	10790	10809	1376
560507	1822	1841	TAGTCAAGTTTAGAGTTTTA	21	10794	10813	1377
560508	1823	1842	TTAGTCAAGTTTAGAGTTTT	10	10795	10814	1378
560509	1824	1843	TTTAGTCAAGTTTAGAGTTT	20	10796	10815	1379
560510	1828	1847	TGTATTTAGTCAAGTTTAGA	8	10800	10819	1380
560511	1829	1848	CTGTATTTAGTCAAGTTTAG	37	10801	10820	1381
560512	1830	1849	TCTGTATTTAGTCAAGTTTA	46	10802	10821	1382
560513	1834	1853	GTCCTCTGTATTTAGTCAAG	38	10806	10825	1383
560514	1835	1854	AGTCCTCTGTATTTAGTCAA	29	10807	10826	1384
560515	1836	1855	CAGTCCTCTGTATTTAGTCA	47	10808	10827	1385
560516	1837	1856	CCAGTCCTCTGTATTTAGTC	31	10809	10828	1386
560517	1838	1857	ACCAGTCCTCTGTATTTAGT	31	10810	10829	1387
560518	1839	1858	TACCAGTCCTCTGTATTTAG	35	10811	10830	1388
560519	1840	1859	TTACCAGTCCTCTGTATTTA	30	10812	10831	1389
560520	1841	1860	ATTACCAGTCCTCTGTATTT	37	10813	10832	1390
560521	1842	1861	AATTACCAGTCCTCTGTATT	12	10814	10833	1391
560522	1843	1862	CAATTACCAGTCCTCTGTAT	38	10815	10834	1392
560523	1844	1863	ACAATTACCAGTCCTCTGTA	35	10816	10835	1393
560524	1845	1864	TACAATTACCAGTCCTCTGT	51	10817	10836	1394
560525	1846	1865	GTACAATTACCAGTCCTCTG	52	10818	10837	1395
560526	1847	1866	TGTACAATTACCAGTCCTCT	38	10819	10838	1396

560527	1848	1867	CTGTACAATTACCAGTCCTC	19	10820	10839	1397
560528	1849	1868	ACTGTACAATTACCAGTCCT	13	10821	10840	1398
560529	1850	1869	AACTGTACAATTACCAGTCC	27	10822	10841	1399
560530	1851	1870	GAAGTGTACAATTACCAGTC	20	10823	10842	1400
560531	1852	1871	AGAACTGTACAATTACCAGT	24	10824	10843	1401
560532	1854	1873	TAAGAACTGTACAATTACCA	22	10826	10845	1402
560533	1855	1874	TTAAGAACTGTACAATTACC	20	10827	10846	1403
560534	1856	1875	TTTAAGAACTGTACAATTAC	1	10828	10847	1404

Table 9

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
544355	N/A	N/A	TTTCAGCATGTATCTCTTAA	69	7065	7084	21
544376	N/A	N/A	GGAGTGGTTCCTTTTCACAGC	64	8298	8317	24
544380	N/A	N/A	TGGTCCTTTTAACTTCCAAT	50	8365	8384	25
560612	N/A	N/A	ACTTGAAATTATAATAGGAA	0	3798	3817	1405
560613	N/A	N/A	AAAAAACTAACTTGAAATTA	0	3807	3826	1406
560614	N/A	N/A	GAAACAAAAAACTAACTTGA	21	3812	3831	1407
560615	N/A	N/A	GTGTTTTTCATATATAACATT	19	3835	3854	1408
560616	N/A	N/A	AATTTTCAGTGTTTTTCATAT	0	3843	3862	1409
560617	N/A	N/A	AAAATGCAAATTTTCAGTGT	0	3851	3870	1410
560618	N/A	N/A	GTAATTTTCATATAAAATGC	0	3864	3883	1411
560619	N/A	N/A	GATTTGTAATTTTCATATAA	0	3869	3888	1412
560620	N/A	N/A	TAACCGATTTGTAATTTTCA	16	3874	3893	1413
560621	N/A	N/A	TAATTTAACCGATTTGTAAT	5	3879	3898	1414
560622	N/A	N/A	TTGTATAATTTAACCGATTT	13	3884	3903	1415
560623	N/A	N/A	CTAGATTGTATAATTTAACC	8	3889	3908	1416
560624	N/A	N/A	GTGTTCTAGATTGTATAATT	24	3894	3913	1417
560625	N/A	N/A	AATGACATAGTGTTCTAGAT	0	3903	3922	1418
560626	N/A	N/A	AGTGTAATGACATAGTG TTC	10	3908	3927	1419
560627	N/A	N/A	TTACAATAGTGTAATGACAT	0	3915	3934	1420
560628	N/A	N/A	TTCAGTAATTTACAATAGTG	12	3924	3943	1421
560629	N/A	N/A	TTACCTTCAGTAATTTACAA	9	3929	3948	1422
560630	N/A	N/A	TTAACTTTTTACTTACCTTC	7	3941	3960	1423
560631	N/A	N/A	GAATAGTTTTTAAATTTTTTTT	0	3960	3979	1424
560632	N/A	N/A	ACACTGGAGAATAGTTTTTAA	10	3968	3987	1425
560633	N/A	N/A	TTTAAACACTGGAGAATAGT	0	3973	3992	1426
560634	N/A	N/A	TCTGTTTTTAAACACTGGAGA	25	3978	3997	1427
560635	N/A	N/A	GTATTATTTAATCTGTTTTTA	0	3989	4008	1428

560636	N/A	N/A	TTACTGTATTATTTAATCTG	5	3994	4013	1429
560637	N/A	N/A	TAAATCTTTTCCATTTACTG	18	4008	4027	1430
560638	N/A	N/A	ATGAATAAATCTTTTCCATT	12	4013	4032	1431
560639	N/A	N/A	GCATATTTTTCATATGAATAA	9	4025	4044	1432
560640	N/A	N/A	GCCCAGCATATTTTTCATATG	20	4030	4049	1433
560641	N/A	N/A	AAAAGAAAAAGCCCAGCATA	20	4040	4059	1434
560642	N/A	N/A	CTGAACTTCAATTAAAAGAA	5	4053	4072	1435
560643	N/A	N/A	GATTTTCTGAACTTCAATTA	9	4059	4078	1436
560644	N/A	N/A	TCTAAAATTTGATTTTCTGA	0	4069	4088	1437
560645	N/A	N/A	ACTATCTCTAAAATTTGATT	8	4075	4094	1438
560646	N/A	N/A	TTAAATTGTACTATCTCTAA	5	4084	4103	1439
560647	N/A	N/A	ACATTTTATTTAATTGTAC	17	4093	4112	1440
560648	N/A	N/A	GTCCTTAACATTTTATTTAA	0	4100	4119	1441
560649	N/A	N/A	CATATTTTGTCTTAACAT	0	4109	4128	1442
560650	N/A	N/A	TAGCACATATTTTGTCTT	25	4114	4133	1443
560651	N/A	N/A	TCAAATAGCACATATTTTG	0	4119	4138	1444
560652	N/A	N/A	CTTCTTTCAAATAGCACATA	41	4125	4144	1445
560653	N/A	N/A	CTTGTATGCTTCTTTCAAAT	19	4133	4152	1446
560654	N/A	N/A	ATTCCTTCCCCTTGTATGCT	12	4143	4162	1447
560655	N/A	N/A	TTGGCAATTCCTTCCCCTTG	36	4149	4168	1448
560656	N/A	N/A	GAATATTGGCAATTCCTTCC	38	4154	4173	1449
560657	N/A	N/A	TGAAAAATGAATATTGGCAA	0	4162	4181	1450
560658	N/A	N/A	TAATGGATTTGAAAAATGAA	0	4171	4190	1451
560659	N/A	N/A	ACTAATAATGGATTTGAAAA	1	4176	4195	1452
560660	N/A	N/A	CATAATCTAAATTTTAAAC	6	4194	4213	1453
560661	N/A	N/A	CACTATCATAATCTAAATTT	4	4200	4219	1454
560662	N/A	N/A	AATTTCCTGTAACACTATCA	2	4212	4231	1455
560663	N/A	N/A	CTATTAATTCCTGTAACAC	9	4217	4236	1456
560664	N/A	N/A	CTTTTCTATTAATTCCTGT	5	4222	4241	1457
560665	N/A	N/A	CTCTTTCTTTTCTATTAATT	0	4228	4247	1458
560666	N/A	N/A	AGTTGCTTTCCTCTTTCTTT	0	4238	4257	1459
560667	N/A	N/A	TTATAAGTTGCTTTCCTCTT	10	4243	4262	1460
560668	N/A	N/A	GTTGGTTATAAGTTGCTTTC	6	4248	4267	1461
560669	N/A	N/A	AGTAGGTTGGTTATAAGTTG	4	4253	4272	1462
560670	N/A	N/A	TAGAGAGTAGGTTGGTTATA	0	4258	4277	1463
560671	N/A	N/A	GGATATAGAGAGTAGGTTGG	0	4263	4282	1464
560672	N/A	N/A	AGTCTGGATATAGAGAGTAG	0	4268	4287	1465
560673	N/A	N/A	TACAAAAGTCTGGATATAGA	7	4274	4293	1466
560674	N/A	N/A	GTTTTTCTACAAAAGTCTGG	12	4281	4300	1467
560675	N/A	N/A	TTACCTGATTTTCTATTTCT	15	4380	4399	1468
560676	N/A	N/A	ATACTGACTTACCTGATTTT	15	4388	4407	1469

560677	N/A	N/A	TTAAAATACTGACTTACCTG	2	4393	4412	1470
560678	N/A	N/A	TACCATTAAAATACTGACTT	0	4398	4417	1471
560679	N/A	N/A	GGACATACCATTAAAATACT	7	4403	4422	1472
560680	N/A	N/A	AAAGATGGGACATACCATTA	0	4410	4429	1473
560681	N/A	N/A	AGACCTGTGTGAAAGATGGG	19	4421	4440	1474
560682	N/A	N/A	TTTACAGACCTGTGTGAAAG	22	4426	4445	1475
560683	N/A	N/A	GTGTTTTTACAGACCTGTGT	47	4431	4450	1476
560684	N/A	N/A	ATTCAGTGTTTTTACAGACC	44	4436	4455	1477
560685	N/A	N/A	TTAGGATTCAGTGTTTTTAC	46	4441	4460	1478
560686	N/A	N/A	ATAATTTTAGGATTCAGTGT	15	4447	4466	1479
560687	N/A	N/A	GCTTGTAATAAATTTTAGGA	0	4455	4474	1480
560688	N/A	N/A	GTAAAGCTTGTAATAAATT	0	4461	4480	1481
560689	N/A	N/A	TGTTTTATATCTCTTGAAAA	0	5571	5590	1482
560690	N/A	N/A	TTGGTAATAATTTTGTTTT	9	5585	5604	1483
560691	N/A	N/A	GGAAATTGGTAATAATATTT	0	5590	5609	1484
560692	N/A	N/A	TTAGTGGAAATTGGTAATAA	22	5595	5614	1485
560693	N/A	N/A	TTTGTTTAGTGGAAATTGGT	8	5600	5619	1486
560694	N/A	N/A	TTATGTTTGTTTAGTGGAAA	0	5605	5624	1487
560695	N/A	N/A	TAACATTATGTTTGTTTAGT	12	5610	5629	1488
560696	N/A	N/A	ACTACTAACATTATGTTTGT	4	5615	5634	1489
560697	N/A	N/A	GCAGCACTACTAACATTATG	38	5620	5639	1490
560698	N/A	N/A	TTTTAGCAGCACTACTAACA	15	5625	5644	1491
560699	N/A	N/A	AAACCTTTTAGCAGCACTAC	52	5630	5649	1492
560700	N/A	N/A	GATAAAAAACCTTTTAGCAG	0	5636	5655	1493
560701	N/A	N/A	TAGTTGATAAAAAACCTTTT	0	5641	5660	1494
560702	N/A	N/A	CAAAAGTAGTTGATAAAAAA	0	5647	5666	1495
560703	N/A	N/A	ATGGAAACCAAAGTAGTTG	13	5655	5674	1496
560704	N/A	N/A	AAAGTATGGAAACCAAAGT	20	5660	5679	1497
560705	N/A	N/A	GAAGGAAAGTATGGAAACCA	45	5665	5684	1498
560706	N/A	N/A	CATAAGAAGGAAAGTATGGA	10	5670	5689	1499
560707	N/A	N/A	TAACATCATAAGAAGGAAAG	0	5676	5695	1500
560708	N/A	N/A	GAATAATAACATCATAAGAA	0	5682	5701	1501
560709	N/A	N/A	GAATTTAGAATAATAACATC	1	5689	5708	1502
560710	N/A	N/A	TATAATTGAAAAGAATTTAG	8	5701	5720	1503
560711	N/A	N/A	TAGTAAAAGATATAATTGAA	0	5711	5730	1504
560712	N/A	N/A	AATCATAGTAAAAGATATAA	10	5716	5735	1505
560713	N/A	N/A	CAGGTTCATTTAATCATAGT	43	5727	5746	1506
560714	N/A	N/A	CTATAGTAACATTTTGCTTT	24	5753	5772	1507
560715	N/A	N/A	GTATATTACTATAGTAACAT	18	5761	5780	1508
560716	N/A	N/A	ACAATGTATATTACTATAGT	0	5766	5785	1509
560717	N/A	N/A	TAGACACAATGTATATTACT	46	5771	5790	1510

560718	N/A	N/A	TATTTTTTAGACACAATGTAT	29	5777	5796	1511
560719	N/A	N/A	ACACATTTTTTATTTTTAGAC	15	5786	5805	1512
560720	N/A	N/A	TTGGTTTCTTCACACATTTT	62	5797	5816	1513
560721	N/A	N/A	TTCATTGTTTTTGGTTTCTTC	55	5806	5825	1514
560722	N/A	N/A	CAGAAATTCATTGTTTTGGT	55	5812	5831	1515
560723	N/A	N/A	TCCAACCTCAGAAATTCATTG	65	5819	5838	48
560724	N/A	N/A	CTTCTTCCAACCTCAGAAATT	41	5824	5843	1516
560725	N/A	N/A	TGATCTAACTCTTCTTCCAA	24	5834	5853	1517
560726	N/A	N/A	TTAAATGATCTAACTCTTCT	23	5839	5858	1518
560727	N/A	N/A	TGAGAAAGTTAAATGATCTA	0	5847	5866	1519
560728	N/A	N/A	TACTTAAATTTTTAGAGTTT	10	5886	5905	1520
560729	N/A	N/A	AAAGTTACTTAAATTTTTAG	3	5891	5910	1521
560730	N/A	N/A	ATCTTAAAGTTACTTAAATT	0	5896	5915	1522
560731	N/A	N/A	ATGTGATCTTAAAGTTACTT	24	5901	5920	1523
560732	N/A	N/A	TAACATATGTGATCTTAAAGT	0	5906	5925	1524
560733	N/A	N/A	TTACTCTTTTCTACTAAGTA	39	5924	5943	1525
560734	N/A	N/A	GGGTATTACTCTTTTCTACT	48	5929	5948	1526
560735	N/A	N/A	TTGCTGGGTATTACTCTTTT	75	5934	5953	49
560736	N/A	N/A	TTTGCTTGCTGGGTATTACT	65	5939	5958	50
560737	N/A	N/A	TAAAGTTTGCTTGCTGGGTA	49	5944	5963	1527
560738	N/A	N/A	TATTGTAAAGTTTGCTTGCT	15	5949	5968	1528
560739	N/A	N/A	TAAAAGGATCTATTGTAAAG	0	5959	5978	1529
560740	N/A	N/A	TTATTTAAAAGGATCTATTG	9	5964	5983	1530
560741	N/A	N/A	GGACCTTATTTAAAAGGATC	17	5969	5988	1531
560742	N/A	N/A	GATATTTCCCTAGGACCTTAT	27	5980	5999	1532
560743	N/A	N/A	TGAATGATATTTCCCTAGGAC	0	5985	6004	1533
560744	N/A	N/A	TGGCATGAATGATATTTCCCT	74	5990	6009	51
560745	N/A	N/A	GATGCTGGCATGAATGATAT	40	5995	6014	1534
560746	N/A	N/A	TTTTTTGATGCTGGCATGAA	38	6001	6020	1535
560747	N/A	N/A	GTTAGTTTTTTGATGCTGGC	35	6006	6025	1536
560748	N/A	N/A	TTAGTGTTAGTTTTTTGATG	0	6011	6030	1537
560749	N/A	N/A	GCATTATTAGTGTTAGTTTT	50	6017	6036	1538
560750	N/A	N/A	ATCTTGCATTATTAGTGTTA	49	6022	6041	1539
560751	N/A	N/A	ATAATATCTTGCATTATTAG	17	6027	6046	1540
560752	N/A	N/A	CAGTAAGAAAAGCAGAATAT	15	6047	6066	1541
560753	N/A	N/A	TCATTGACAGTAAGAAAAGC	47	6054	6073	1542
560754	N/A	N/A	GATAGTTTTTCTCATTGACA	40	6065	6084	1543
560755	N/A	N/A	GTTTGCAATTTATTGAATGA	12	6083	6102	1544
560756	N/A	N/A	GTGTTGGGTTTGCAATTTAT	55	6090	6109	1545
560757	N/A	N/A	TTAAGTGTGTTGGGTTTGCA	50	6096	6115	1546
560758	N/A	N/A	TTTTATTTAAGTGTGTTGGG	5	6102	6121	1547

560759	N/A	N/A	TTTAGCAGTAACATTTTATT	19	6121	6140	1548
560760	N/A	N/A	GTTAGTTTAGCAGTAACATT	30	6126	6145	1549
560761	N/A	N/A	TCTATATATTCAGTAGTTTA	17	6148	6167	1550
560762	N/A	N/A	TTACTTTCTATATATTCAGT	14	6154	6173	1551
560763	N/A	N/A	GTTTGCTTACTTTCTATATA	20	6160	6179	1552
560764	N/A	N/A	AGTTTGTTTGCTTACTTTCT	36	6165	6184	1553
560765	N/A	N/A	TGGCAAGTTTGTTTGCTTAC	43	6170	6189	1554
560766	N/A	N/A	TTACTGTTACTGTATTTCCC	39	10155	10174	1555
560767	N/A	N/A	ATGTAGTTACTGTTACTGTA	18	10161	10180	1556
560768	N/A	N/A	ATTTAATGGGTACAGACTCG	47	10182	10201	61
560769	N/A	N/A	ATGCAATTTAATGGGTACAG	32	10187	10206	1557
560770	N/A	N/A	TAGATATGCAATTTAATGGG	4	10192	10211	1558
560771	N/A	N/A	AGGAGATAGATATGCAATTT	5	10198	10217	1559
560772	N/A	N/A	CCTAAAGGAGATAGATATGC	36	10203	10222	1560
560773	N/A	N/A	AGCCTCCTAAAGGAGATAGA	0	10208	10227	1561
560774	N/A	N/A	CACCACCAGCCTCCTAAAGG	35	10215	10234	1562
560775	N/A	N/A	ATCTAAGAAAATTAATAAAC	17	7003	7022	1563
560776	N/A	N/A	ATGATCACATCTAAGAAAAT	8	7011	7030	1564
560777	N/A	N/A	ATACCATGATCACATCTAAG	49	7016	7035	62
560778	N/A	N/A	GCAATACCATGATCACATCT	59	7019	7038	52
560779	N/A	N/A	AACTGCAATACCATGATCAC	35	7023	7042	1565
560780	N/A	N/A	TAAAACCTGCAATACCATGAT	43	7026	7045	1566
560781	N/A	N/A	CTTTAAACCTGCAATACCAT	13	7029	7048	1567
560782	N/A	N/A	TCTCCTTTAAACCTGCAATA	18	7033	7052	1568
560783	N/A	N/A	TGTTCTCCTTTAAACCTGCA	13	7036	7055	1569
560784	N/A	N/A	GATTGTTCTCCTTTAAACCT	23	7039	7058	1570
560785	N/A	N/A	AGGAGATTGTTCTCCTTTAA	14	7043	7062	1571
560786	N/A	N/A	AACAGGAGATTGTTCTCCTT	0	7046	7065	1572
560787	N/A	N/A	TTAAACAGGAGATTGTTCTC	7	7049	7068	1573
560788	N/A	N/A	CTCTTAAACAGGAGATTGTT	10	7052	7071	1574
560789	N/A	N/A	ACTCCGTAAATATTTTCAGCA	55	7077	7096	53
560790	N/A	N/A	CTTTAACTCCGTAAATATTT	22	7082	7101	1575
560791	N/A	N/A	GACCTTTAACTCCGTAAATA	54	7085	7104	63
560792	N/A	N/A	AGTGACCTTTAACTCCGTAA	35	7088	7107	1576
560793	N/A	N/A	GGAGTCCAGTGACCTTTAAC	15	7095	7114	1577
560794	N/A	N/A	TCTGGAGTCCAGTGACCTTT	46	7098	7117	64
560795	N/A	N/A	ACCAGTCTGGAGTCCAGTGA	8	7103	7122	1578
560796	N/A	N/A	TCATCTTACCAAACCTATTTT	22	7169	7188	1579
560797	N/A	N/A	GAATCATCTTACCAAACCTAT	39	7172	7191	1580
560798	N/A	N/A	TAAGAATCATCTTACCAAAC	35	7175	7194	1581
560799	N/A	N/A	ATGTAAGAATCATCTTACCA	52	7178	7197	65

560800	N/A	N/A	AAGAATGTAAGAATCATCTT	22	7182	7201	1582
560801	N/A	N/A	GTTATTTAAGAATGTAAGAA	0	7189	7208	1583
560802	N/A	N/A	CGTGTTATTTAAGAATGTAA	3	7192	7211	1584
560803	N/A	N/A	AGCATTTTTCTTAGATGGCG	48	7210	7229	66
560804	N/A	N/A	TAAAGCATTTTTCTTAGATG	0	7213	7232	1585
560805	N/A	N/A	TGTTAAAGCATTTTTCTTAG	0	7216	7235	1586
560806	N/A	N/A	TTTATGTTAAAGCATTTTTTC	20	7220	7239	1587
560807	N/A	N/A	ATGTTTATGTTAAAGCATTT	8	7223	7242	1588
560808	N/A	N/A	GCATTTTTTTCAGTAATGTTT	40	7237	7256	1589
560809	N/A	N/A	TGTAGCATTTTTTTCAGTAAT	24	7241	7260	1590
560810	N/A	N/A	CAAATGTAGCATTTTTTTCAG	0	7245	7264	1591
560811	N/A	N/A	TGGCAAATGTAGCATTTTTTT	60	7248	7267	54
560812	N/A	N/A	AAGTTGTGGCAAATGTAGCA	26	7254	7273	1592
560813	N/A	N/A	ATGAAGTTGTGGCAAATGTA	11	7257	7276	1593
560814	N/A	N/A	TTTATGAAGTTGTGGCAAAT	36	7260	7279	1594
560815	N/A	N/A	CATTTTATGAAGTTGTGGCA	45	7263	7282	67
560816	N/A	N/A	TGACATTTTATGAAGTTGTG	16	7266	7285	1595
560817	N/A	N/A	CACTTGACATTTTATGAAGT	47	7270	7289	68
560818	N/A	N/A	CTTGAGATTTCACTTGACAT	18	7280	7299	1596
560819	N/A	N/A	TTTGGAGCTTGAGATTTTAC	0	7287	7306	1597
560820	N/A	N/A	ATCTTTGGAGCTTGAGATTT	0	7290	7309	1598
560821	N/A	N/A	AATATCTTTGGAGCTTGAGA	6	7293	7312	1599
560822	N/A	N/A	AATAATATCTTTGGAGCTTG	24	7296	7315	1600
560823	N/A	N/A	AGGAATAATATCTTTGGAGC	1	7299	7318	1601
560824	N/A	N/A	AATAGGAATAATATCTTTGG	0	7302	7321	1602
560825	N/A	N/A	AGTAATAGGAATAATATCTT	0	7305	7324	1603
560826	N/A	N/A	TTACATCAGATTTAGTAATA	0	7318	7337	1604
560827	N/A	N/A	AAATGTTATTACATCAGATT	0	7326	7345	1605
560828	N/A	N/A	ATAAAATGTTATTACATCAG	12	7329	7348	1606
560829	N/A	N/A	CCTAGAATCAATAAAATGTT	13	7339	7358	1607
560830	N/A	N/A	AGGAATGCCTAGAATCAATA	9	7346	7365	1608
560831	N/A	N/A	ATTCAGCAGGAATGCCTAGA	26	7353	7372	1609
560832	N/A	N/A	TACATTCAGCAGGAATGCCT	23	7356	7375	1610
560833	N/A	N/A	TTACCTGATATAACATCACA	30	7456	7475	1611
560834	N/A	N/A	GTTTTACCTGATATAACATC	6	7459	7478	1612
560835	N/A	N/A	CAGGTTTTACCTGATATAAC	4	7462	7481	1613
560836	N/A	N/A	TTAGACAGGTTTTACCTGAT	6	7467	7486	1614
560837	N/A	N/A	ATTCTCCTTAGACAGGTTTT	6	7474	7493	1615
560838	N/A	N/A	ACTGTCTATTCTCCTTAGAC	0	7481	7500	1616
560839	N/A	N/A	ACTACTGTCTATTCTCCTTA	17	7484	7503	1617
560840	N/A	N/A	ACTAACTACTGTCTATTCTC	0	7488	7507	1618

560841	N/A	N/A	TGAACTAACTACTGTCTATT	0	7491	7510	1619
560842	N/A	N/A	AGTTGAACTAACTACTGTCT	0	7494	7513	1620
560844	N/A	N/A	ATTAATTGATATGTAAAACG	0	8347	8366	1621
560845	N/A	N/A	CCAATTAATTGATATGTAAA	15	8350	8369	1622
560846	N/A	N/A	TCCTTTTAACTTCCAATTAA	29	8362	8381	1623
560847	N/A	N/A	TCCTGGTCCTTTTAACTTCC	58	8368	8387	69
560848	N/A	N/A	GTTTCCTGGTCCTTTTAACT	0	8371	8390	1624
560849	N/A	N/A	TCTGAGTTTCCTGGTCCTTT	36	8376	8395	1625
560850	N/A	N/A	ATGTCTGAGTTTCCTGGTCC	31	8379	8398	1626
560851	N/A	N/A	TGTATGTCTGAGTTTCCTGG	0	8382	8401	1627
560852	N/A	N/A	ATGTATACTGTATGTCTGAG	19	8390	8409	1628
560853	N/A	N/A	AAAATGTATACTGTATGTCT	12	8393	8412	1629
560854	N/A	N/A	TTTTAAAATGTATACTGTAT	0	8397	8416	1630
560855	N/A	N/A	CATACATTCTATATATTATA	29	8432	8451	1631
560856	N/A	N/A	AAGCCATACATTCTATATAT	38	8436	8455	55
560857	N/A	N/A	ATTATAAGCCATACATTCTA	6	8441	8460	1632
560858	N/A	N/A	TTCATTATAAGCCATACATT	0	8444	8463	1633
560859	N/A	N/A	TAATTCATTATAAGCCATAC	19	8447	8466	1634
560860	N/A	N/A	TGAGTTAACTAATTCATTAT	0	8456	8475	1635
560861	N/A	N/A	TTTGCATTGAGTTAACTAAT	26	8463	8482	1636
560862	N/A	N/A	TAATTTGCATTGAGTTAACT	0	8466	8485	1637
560863	N/A	N/A	GAATAATTTGCATTGAGTTA	0	8469	8488	1638
560864	N/A	N/A	ATAGAATAATTTGCATTGAG	0	8472	8491	1639
560865	N/A	N/A	AAAATAGAATAATTTGCATT	0	8475	8494	1640
560866	N/A	N/A	TTGTAATCAAAATAGAATAA	0	8483	8502	1641
560867	N/A	N/A	TATTTGTAATCAAAATAGAA	16	8486	8505	1642
560868	N/A	N/A	TACTATTTGTAATCAAAATA	0	8489	8508	1643
560869	N/A	N/A	TTTTACTATTTGTAATCAAA	0	8492	8511	1644
560870	N/A	N/A	GCTTATTTTACTATTTGTAA	0	8497	8516	1645
560871	N/A	N/A	CTTGCTTATTTTACTATTTG	0	8500	8519	1646
560872	N/A	N/A	TTATCTTGCTTATTTTACTA	1	8504	8523	1647
560873	N/A	N/A	GTTATTTTATCTTGCTTATT	0	8510	8529	1648
560874	N/A	N/A	AAACATCTGTTATTTTATCT	0	8518	8537	1649
560875	N/A	N/A	GGATTTTAAACATCTGTTAT	0	8525	8544	1650
560876	N/A	N/A	CTTTTTGGATTTTAAACATC	24	8531	8550	1651
560877	N/A	N/A	GTGCTTTTTTGGATTTTAAAC	6	8534	8553	1652
560878	N/A	N/A	TTTTGTATGTGCTTTTTTGA	24	8542	8561	1653
560879	N/A	N/A	GACATCATTCATGGATTTTT	50	8558	8577	70
560880	N/A	N/A	AGTACTTAGACATCATTCAT	43	8566	8585	71
560881	N/A	N/A	TAAGTGAGTACTTAGACATC	17	8572	8591	1654
560882	N/A	N/A	TACTTTATAAGTGAGTACTT	0	8579	8598	1655

560883	N/A	N/A	TTCTACTTTATAAGTGAGTA	32	8582	8601	1656
560884	N/A	N/A	AATGTCTTCTACTTTATAAG	0	8588	8607	1657
560885	N/A	N/A	AATAATGAATGTCTTCTACT	9	8595	8614	1658
560886	N/A	N/A	TATAATAATGAATGTCTTCT	0	8598	8617	1659
560887	N/A	N/A	TGATATAATAATGAATGTCT	29	8601	8620	1660
560888	N/A	N/A	AAAATTTGATATAATAATGA	0	8607	8626	1661
560889	N/A	N/A	CATTTAAAAATTTGATATAA	0	8613	8632	1662
560890	N/A	N/A	GTA CTGAGCATTTAAAAATT	8	8621	8640	1663
560891	N/A	N/A	GGTCAAATAGTACTGAGCAT	40	8630	8649	72
560892	N/A	N/A	AATGGTCAAATAGTACTGAG	23	8633	8652	1664
560893	N/A	N/A	TTAAATGGTCAAATAGTACT	17	8636	8655	1665
560894	N/A	N/A	AGTTTGAATACAAAATTTTT	0	8654	8673	1666
560895	N/A	N/A	GGTAGTTTGAATACAAAATT	38	8657	8676	73
560896	N/A	N/A	ACTGGTAGTTTGAATACAAA	0	8660	8679	1667
560897	N/A	N/A	TTC ACTGGTAGTTTGAATAC	0	8663	8682	1668
560898	N/A	N/A	GCTTTC ACTGGTAGTTTGAA	25	8666	8685	1669
560899	N/A	N/A	AGGGCTTTC ACTGGTAGTTT	30	8669	8688	1670
560900	N/A	N/A	GGTAGGGCTTTC ACTGGTAG	9	8672	8691	1671
560901	N/A	N/A	CTAGGTAGGGCTTTC ACTGG	37	8675	8694	1672
560902	N/A	N/A	CTTCTAGGTAGGGCTTTCAC	32	8678	8697	1673
560903	N/A	N/A	TACCTTCTAGGTAGGGCTTT	26	8681	8700	1674
560904	N/A	N/A	GTATACCTTCTAGGTAGGGC	0	8684	8703	1675
560905	N/A	N/A	TGAGTATACCTTCTAGGTAG	15	8687	8706	1676
560906	N/A	N/A	CACTGAGTATACCTTCTAGG	36	8690	8709	1677
560907	N/A	N/A	TATCACTGAGTATACCTTCT	0	8693	8712	1678
560908	N/A	N/A	ACTTATCACTGAGTATACCT	28	8696	8715	1679
560909	N/A	N/A	ACAAA ACTTATCACTGAGTA	32	8701	8720	1680
560910	N/A	N/A	GCTACAAA ACTTATCACTGA	15	8704	8723	1681
560911	N/A	N/A	GGAGCTACAAA ACTTATCAC	21	8707	8726	1682
560912	N/A	N/A	GATTTGGAGCTACAAA ACTT	0	8712	8731	1683
560913	N/A	N/A	GAAGATTTGGAGCTACAAA	0	8715	8734	1684
560914	N/A	N/A	CTATTAGAAGATTTGGAGCT	0	8721	8740	1685
560915	N/A	N/A	CACTCACTATTAGAAGATTT	33	8727	8746	1686
560916	N/A	N/A	TGTCAGCCTTTTATTTTGGG	0	8751	8770	1687
560917	N/A	N/A	ACCTGTCAGCCTTTTATTTT	11	8754	8773	1688
560918	N/A	N/A	TCGACTTACCTGTCAGCCTT	0	8761	8780	1689
560919	N/A	N/A	TTCTCGACTTACCTGTCAGC	0	8764	8783	1690
560920	N/A	N/A	GTATTCTCGACTTACCTGTC	0	8767	8786	1691
560921	N/A	N/A	TAACATCCATATACAGTCAA	25	9177	9196	1692
560922	N/A	N/A	TATTAACATCCATATACAGT	20	9180	9199	1693
560923	N/A	N/A	ATTTATTAACATCCATATAC	20	9183	9202	1694

560924	N/A	N/A	GCTATTTATTAACATCCATA	47	9186	9205	1695
560925	N/A	N/A	TCAGCTATTTATTAACATCC	58	9189	9208	56
560926	N/A	N/A	CTGTCAGCTATTTATTAACA	30	9192	9211	1696
560927	N/A	N/A	TTACTGTCAGCTATTTATTA	22	9195	9214	1697
560928	N/A	N/A	ACTTTACTGTCAGCTATTTA	27	9198	9217	1698
560929	N/A	N/A	TAAACTTTACTGTCAGCTAT	41	9201	9220	1699
560930	N/A	N/A	GGATAAACTTTACTGTCAGC	45	9204	9223	1700
560931	N/A	N/A	TATGGATAAACTTTACTGTC	15	9207	9226	1701
560932	N/A	N/A	TTATATGGATAAACTTTACT	0	9210	9229	1702
560933	N/A	N/A	TTGCAAGTCTTTATATGGAT	47	9220	9239	1703
560934	N/A	N/A	TATTTGCAAGTCTTTATATG	26	9223	9242	1704
560935	N/A	N/A	GAATATTTGCAAGTCTTTAT	4	9226	9245	1705
560936	N/A	N/A	GAGGAATATTTGCAAGTCTT	58	9229	9248	57
560937	N/A	N/A	GTAGAGGAATATTTGCAAGT	47	9232	9251	1706
560938	N/A	N/A	TTGGTAGAGGAATATTTGCA	65	9235	9254	58
560939	N/A	N/A	GTTACATTATTATAGATATT	33	9269	9288	1707
560940	N/A	N/A	TGTGTTACATTATTATAGAT	20	9272	9291	1708
560941	N/A	N/A	GAAATGTGTTACATTATTAT	0	9276	9295	1709
560942	N/A	N/A	ACCAGTGAAATGTGTTACAT	56	9282	9301	59
560943	N/A	N/A	TTCACCAGTGAAATGTGTTA	19	9285	9304	1710
560944	N/A	N/A	TGTTTCACCAGTGAAATGTG	41	9288	9307	1711
560945	N/A	N/A	ACATGTTTCACCAGTGAAAT	0	9291	9310	1712
560946	N/A	N/A	AAGACATGTTTCACCAGTGA	48	9294	9313	1713
560947	N/A	N/A	GACAAGACATGTTTCACCAG	28	9297	9316	1714
560948	N/A	N/A	TATGACAAGACATGTTTCAC	13	9300	9319	1715
560949	N/A	N/A	GCATATGACAAGACATGTTT	12	9303	9322	1716
560950	N/A	N/A	TAATGCATATGACAAGACAT	4	9307	9326	1717
560951	N/A	N/A	CTATAATGCATATGACAAGA	22	9310	9329	1718
560952	N/A	N/A	TTTCTATAATGCATATGACA	23	9313	9332	1719
560953	N/A	N/A	TCCTTTCTATAATGCATATG	16	9316	9335	1720
560954	N/A	N/A	TCTGATTATCCTTTCTATAA	32	9324	9343	1721
560955	N/A	N/A	AAGTCTGATTATCCTTTCTA	42	9327	9346	1722
560956	N/A	N/A	TGAAAGTCTGATTATCCTTT	51	9330	9349	60
560957	N/A	N/A	AACTGAAAGTCTGATTATCC	31	9333	9352	1723
560958	N/A	N/A	TATAACTGAAAGTCTGATTA	6	9336	9355	1724
560959	N/A	N/A	GTAAAAAATATTAATATAAC	3	9350	9369	1725
560960	N/A	N/A	TGTGCACAAAAATGTTAAAA	0	9363	9382	1726
560961	N/A	N/A	CTATGTGCACAAAAATGTTA	9	9366	9385	1727
560962	N/A	N/A	TAGCTATGTGCACAAAAATG	29	9369	9388	1728
560963	N/A	N/A	AGATAGCTATGTGCACAAAA	41	9372	9391	1729
560964	N/A	N/A	TGAAGATAGCTATGTGCACA	23	9375	9394	1730

560965	N/A	N/A	TATTGAAGATAGCTATGTGC	13	9378	9397	1731
560966	N/A	N/A	TTTTATTGAAGATAGCTATG	4	9381	9400	1732
560967	N/A	N/A	CAATTTTATTGAAGATAGCT	17	9384	9403	1733
560968	N/A	N/A	AAACAATTTTATTGAAGATA	27	9387	9406	1734
560969	N/A	N/A	GTGTATCTTAAAATAATACC	7	9412	9431	1735
560970	N/A	N/A	TTAGTGTATCTTAAAATAAT	25	9415	9434	1736
560971	N/A	N/A	TGATCATTTTGTGTATCTT	34	9423	9442	1737
560972	N/A	N/A	CCCTTGATCATTTTAGTGTA	7	9427	9446	1738
560973	N/A	N/A	AATCCCTTGATCATTTTAGT	0	9430	9449	1739
560974	N/A	N/A	TTGAATCCCTTGATCATTTT	20	9433	9452	1740
560975	N/A	N/A	TTAGTCTTGAATCCCTTGAT	28	9439	9458	1741
560976	N/A	N/A	TTGTTTAGTCTTGAATCCCT	40	9443	9462	1742
560977	N/A	N/A	GAGTTGTTTAGTCTTGAATC	6	9446	9465	1743
560978	N/A	N/A	ATTGAGTTGTTTAGTCTTGA	14	9449	9468	1744
560979	N/A	N/A	CTAATTGAGTTGTTTAGTCT	0	9452	9471	1745
560980	N/A	N/A	CAACTAATTGAGTTGTTTAG	0	9455	9474	1746
560981	N/A	N/A	ATTGGTGCAACTAATTGAGT	0	9462	9481	1747
560982	N/A	N/A	TTTATTGGTGCAACTAATTG	9	9465	9484	1748
560983	N/A	N/A	TTTTTTATTGGTGCAACTAA	8	9468	9487	1749
560984	N/A	N/A	TAAGTGTTTTTTATTGGTGC	20	9474	9493	1750
560985	N/A	N/A	ACTGACAGTTTTTTTAAGTG	16	9488	9507	1751
560986	N/A	N/A	GACACTGACAGTTTTTTTAA	6	9491	9510	1752
560987	N/A	N/A	TTGGACACTGACAGTTTTTTT	0	9494	9513	1753
560988	N/A	N/A	AGGTTGGACACTGACAGTTT	6	9497	9516	1754
560989	N/A	N/A	TACAGGTTGGACACTGACAG	0	9500	9519	1755
544120	707	726	AGTTCTTGGTGCTCTTGGCT	72	6720	6739	15
337487	804	823	CACTTGTATGTTACCTCTG	80	7389	7408	28
544145	1055	1074	GTTGTCTTCCAGTCTTCCA	69	9630	9649	16
544156	1195	1214	GCTTTGTGATCCCAAGTAGA	61	9770	9789	17
544162	1269	1288	GGTTGTTTTCTCCACACTCA	71	10241	10260	18
544166	1353	1372	ACCTTCCATTTTGAGACTTC	65	10325	10344	19
544199	1907	1926	TACACATACTCTGTGCTGAC	69	10879	10898	20

Table 10

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
563720	N/A	N/A	TATATTGGATAATTTGAAAT	7	11610	11629	1756
563721	N/A	N/A	ATGTATATTGGATAATTTGA	17	11613	11632	1757
563722	N/A	N/A	GACATGTATATTGGATAATT	20	11616	11635	1758

563723	N/A	N/A	ATGACATGTATATTGGATAA	29	11618	11637	1759
563724	N/A	N/A	TATATATGACATGTATATTG	9	11623	11642	1760
563725	N/A	N/A	ATGTGACATATAAAAATATA	4	11639	11658	1761
563726	N/A	N/A	ATATGTGACATATAAAAATA	0	11641	11660	1762
563727	N/A	N/A	TTTATATATGTGACATATAA	0	11646	11665	1763
563728	N/A	N/A	CTTTTATATATGTGACATAT	16	11648	11667	1764
563729	N/A	N/A	ATCTTTTATATATGTGACAT	13	11650	11669	1765
563730	N/A	N/A	CATATCTTTTATATATGTGA	2	11653	11672	1766
563731	N/A	N/A	TCATACATATCTTTTATATA	2	11658	11677	1767
563732	N/A	N/A	TAGATCATACATATCTTTTA	31	11662	11681	1768
563733	N/A	N/A	CATAGATCATACATATCTTT	28	11664	11683	1769
563734	N/A	N/A	CACATAGATCATACATATCT	56	11666	11685	1770
563735	N/A	N/A	AGGATTCACATAGATCATAC	56	11672	11691	1771
563736	N/A	N/A	TTAGGATTCACATAGATCAT	24	11674	11693	1772
563737	N/A	N/A	ACTTAGGATTCACATAGATC	49	11676	11695	1773
563738	N/A	N/A	TTACTTAGGATTCACATAGA	15	11678	11697	1774
563739	N/A	N/A	TATTTACTTAGGATTCACAT	6	11681	11700	1775
563740	N/A	N/A	AATATTTACTTAGGATTCAC	28	11683	11702	1776
563741	N/A	N/A	TGTACTTTTCTGGAACAAAA	63	11701	11720	1777
563742	N/A	N/A	GATTATTTTTTACCTTTATTA	21	11724	11743	1778
563743	N/A	N/A	TAGATTATTTTTTACCTTTAT	5	11726	11745	1779
563744	N/A	N/A	ATTATAGATTATTTTTTACCT	12	11730	11749	1780
563745	N/A	N/A	GAAAATTATAGATTATTTTTT	15	11734	11753	1781
563746	N/A	N/A	GGTCCTGAAAATTATAGATT	7	11740	11759	1782
563747	N/A	N/A	GTGGTCCTGAAAATTATAGA	29	11742	11761	1783
563748	N/A	N/A	CTGTGGTCCTGAAAATTATA	37	11744	11763	1784
563749	N/A	N/A	GTCTGTGGTCCTGAAAATTA	47	11746	11765	1785
563750	N/A	N/A	TCGACAGCTTAGTCTGTGGT	66	11757	11776	1786
563751	N/A	N/A	TTTCGACAGCTTAGTCTGTG	41	11759	11778	1787
563752	N/A	N/A	AATTTTCGACAGCTTAGTCTG	40	11761	11780	1788
563753	N/A	N/A	TTAATTTTCGACAGCTTAGTC	35	11763	11782	1789
563754	N/A	N/A	CGTTAATTTTCGACAGCTTAG	50	11765	11784	1790
563755	N/A	N/A	TGGCCCTAAAAAAATCAGCG	7	11783	11802	1791
563756	N/A	N/A	TCTGGCCCTAAAAAAATCAG	0	11785	11804	1792
563757	N/A	N/A	TGGTATTCTGGCCCTAAAAA	37	11791	11810	1793
563758	N/A	N/A	TTTGGTATTCTGGCCCTAAA	29	11793	11812	1794
563759	N/A	N/A	CCATTTTGGTATTCTGGCCC	35	11797	11816	1795
563760	N/A	N/A	GAGGAGCCATTTTGGTATTC	34	11803	11822	1796
563761	N/A	N/A	GAGAGGAGCCATTTTGGTAT	18	11805	11824	1797
563762	N/A	N/A	AAGAGAGGAGCCATTTTGGT	17	11807	11826	1798
563763	N/A	N/A	TGAAATTGTCCAATTTTGGG	28	11829	11848	1799

563764	N/A	N/A	TTTGAAATTGTCCAATTTTG	10	11831	11850	1800
563765	N/A	N/A	CATTTGAAATTGTCCAATTT	22	11833	11852	1801
563766	N/A	N/A	TGCATTTGAAATTGTCCAAT	45	11835	11854	1802
563767	N/A	N/A	ATTTTGCATTTGAAATTGTC	35	11839	11858	1803
563768	N/A	N/A	ATAATGAATTATTTTGCATT	0	11849	11868	1804
563769	N/A	N/A	TAAATAATGAATTATTTTGC	17	11852	11871	1805
563770	N/A	N/A	CTCATATATTAAATAATGAA	0	11861	11880	1806
563771	N/A	N/A	AACTCATATATTAAATAATG	16	11863	11882	1807
563772	N/A	N/A	TAGAGGAAGCAACTCATATA	7	11873	11892	1808
563773	N/A	N/A	AATAGAGGAAGCAACTCATA	20	11875	11894	1809
563774	N/A	N/A	CAAATAGAGGAAGCAACTCA	29	11877	11896	1810
563775	N/A	N/A	ACCAAATAGAGGAAGCAACT	27	11879	11898	1811
563776	N/A	N/A	AAACCAAATAGAGGAAGCAA	22	11881	11900	1812
563777	N/A	N/A	GGAAACCAAATAGAGGAAGC	37	11883	11902	1813
563778	N/A	N/A	TAAGGAAACCAAATAGAGGA	0	11886	11905	1814
563779	N/A	N/A	TTTAAGGAAACCAAATAGAG	0	11888	11907	1815
563780	N/A	N/A	TGTTTTCTTCTGGAAGCAGA	5	3100	3119	1816
563781	N/A	N/A	CTTACTTTAAGTGAAGTTAC	0	3636	3655	1817
563782	N/A	N/A	TTTTCTACTTACTTTAAGTG	3	3643	3662	1818
563783	N/A	N/A	ACATGAACCCTCTTTATTTT	0	3659	3678	1819
563784	N/A	N/A	GAAACATAAACATGAACCC	0	3669	3688	1820
563785	N/A	N/A	AGATCCACATTGAAAACATA	8	3680	3699	1821
563786	N/A	N/A	TTAAAAGATCCACATTGAAA	8	3685	3704	1822
563787	N/A	N/A	GCCTTAGAAATATTTTTTTT	2	3703	3722	1823
563788	N/A	N/A	CAAATGGCATGCCTTAGAAA	29	3713	3732	1824
563789	N/A	N/A	TATTTCAAATGGCATGCCTT	24	3718	3737	1825
563790	N/A	N/A	CAAAGTATTTCAAATGGCAT	8	3723	3742	1826
563791	N/A	N/A	TGCAACAAAGTATTTCAAAT	0	3728	3747	1827
563792	N/A	N/A	TCAACAATGCAACAAAGTAT	3	3735	3754	1828
563793	N/A	N/A	GAAAAAAAAGTATTTCAACA	4	3749	3768	1829
563794	N/A	N/A	GATTATTTTTCTTGGA AAAA	11	3763	3782	1830
563795	N/A	N/A	GAAATTTTATTTTCTGGAGA	10	3781	3800	1831
563796	N/A	N/A	AAATTATAATAGGAAATTTT	14	3793	3812	1832
563797	N/A	N/A	CTGAATATAATGAATGAAAT	1	7854	7873	1833
563798	N/A	N/A	TACCTGAATATAATGAATGA	4	7857	7876	1834
563799	N/A	N/A	GACTACCTGAATATAATGAA	25	7860	7879	1835
563800	N/A	N/A	ATGGACTACCTGAATATAAT	15	7863	7882	1836
563801	N/A	N/A	TCCATGGACTACCTGAATAT	39	7866	7885	1837
563802	N/A	N/A	ACCATCAAGCCTCCCAAAAC	23	7952	7971	1838
563803	N/A	N/A	CCTTACCATCAAGCCTCCCA	29	7956	7975	1839
563804	N/A	N/A	AGTCCCCTTACCATCAAGCC	31	7961	7980	1840

563805	N/A	N/A	TGTAGTCCCCTTACCATCAA	18	7964	7983	1841
563806	N/A	N/A	GAATGTAGTCCCCTTACCAT	0	7967	7986	1842
563807	N/A	N/A	ATTGAATGTAGTCCCCTTAC	12	7970	7989	1843
563808	N/A	N/A	ATGATTGAATGTAGTCCCCT	14	7973	7992	1844
563809	N/A	N/A	GATTAGCAAGTGAATGAATG	13	7990	8009	1845
563810	N/A	N/A	GTAGATTAGCAAGTGAATGA	25	7993	8012	1846
563811	N/A	N/A	TTTGTAGATTAGCAAGTGAA	9	7996	8015	1847
563812	N/A	N/A	ATATTTGTAGATTAGCAAGT	0	7999	8018	1848
563813	N/A	N/A	CCATAAGAGGTTCTCAGTAA	44	8019	8038	1849
563814	N/A	N/A	GGTCCATAAGAGGTTCTCAG	37	8022	8041	1850
563815	N/A	N/A	CCTGGTCCATAAGAGGTTCT	25	8025	8044	1851
563816	N/A	N/A	TAATACCTGGTCCATAAGAG	9	8030	8049	1852
563817	N/A	N/A	TCCTAATACCTGGTCCATAA	39	8033	8052	1853
563818	N/A	N/A	TTTTCCTAATACCTGGTCCA	43	8036	8055	1854
563819	N/A	N/A	TACTTTTCCTAATACCTGGT	43	8039	8058	1855
563820	N/A	N/A	CGTTACTACTTTTCCTAATA	47	8045	8064	1856
563821	N/A	N/A	AAGGCTGAGACTGCTTCTCG	46	8067	8086	1857
563822	N/A	N/A	GATAATAAATTATATGAAGG	5	8083	8102	1858
563823	N/A	N/A	GTTTGATAATAAATTATATG	0	8087	8106	1859
563824	N/A	N/A	GTGTAATTGTTTGATAATAA	14	8095	8114	1860
563825	N/A	N/A	AATGTGTAATTGTTTGATAA	0	8098	8117	1861
563826	N/A	N/A	GTAATTTACTAACAAATGTG	18	8112	8131	1862
563827	N/A	N/A	AGTGTAATTTACTAACAAAT	0	8115	8134	1863
563828	N/A	N/A	ATAAGTGTAATTTACTAACA	0	8118	8137	1864
563829	N/A	N/A	GTAATAAGTGTAATTTACTA	0	8121	8140	1865
563830	N/A	N/A	GTTGTAATAAGTGTAATTTA	20	8124	8143	1866
563831	N/A	N/A	ACAGTTGTAATAAGTGTAAT	1	8127	8146	1867
563832	N/A	N/A	ATAACAGTTGTAATAAGTGT	4	8130	8149	1868
563833	N/A	N/A	TTCAAATAATAACAGTTGTA	0	8138	8157	1869
563834	N/A	N/A	ATAATTCAAATAATAACAGT	16	8142	8161	1870
563835	N/A	N/A	AATTGTGATAAATATAATTC	0	8155	8174	1871
563836	N/A	N/A	ATGTAATTGTGATAAATATA	0	8159	8178	1872
563837	N/A	N/A	GACATGTAATTGTGATAAAT	8	8162	8181	1873
563838	N/A	N/A	ACAGACATGTAATTGTGATA	33	8165	8184	1874
563839	N/A	N/A	AGAACAGACATGTAATTGTG	34	8168	8187	1875
563840	N/A	N/A	TTAAGAACAGACATGTAATT	0	8171	8190	1876
563841	N/A	N/A	AAGTATATTTAAGAACAGAC	0	8179	8198	1877
563842	N/A	N/A	TTAAATTGTGATAAGTATAT	1	8191	8210	1878
563843	N/A	N/A	GAATTAAATTGTGATAAGTA	0	8194	8213	1879
563844	N/A	N/A	GTGGAATTAAATTGTGATAA	0	8197	8216	1880
563845	N/A	N/A	GCCGTGGAATTAAATTGTGA	20	8200	8219	1881

563846	N/A	N/A	TAAGCCGTGGAATTAAATTG	16	8203	8222	1882
563847	N/A	N/A	TTGTAAGCCGTGGAATTAAG	28	8206	8225	1883
563848	N/A	N/A	TCATTGTAAGCCGTGGAATT	25	8209	8228	1884
563849	N/A	N/A	TGATCATTGTAAGCCGTGGA	49	8212	8231	1885
563850	N/A	N/A	TATAGTTATGATCATTGTAA	0	8220	8239	1886
563851	N/A	N/A	AATTATAGTTATGATCATTG	0	8223	8242	1887
563852	N/A	N/A	CTTTAATAATTATAGTTATG	0	8230	8249	1888
563853	N/A	N/A	TGTCTTTAATAATTATAGTT	4	8233	8252	1889
563854	N/A	N/A	AATTGTCTTTAATAATTATA	0	8236	8255	1890
563855	N/A	N/A	TCAAAATTGTCTTTAATAAT	7	8240	8259	1891
563856	N/A	N/A	ATTTAATCAAAATTGTCTTT	0	8246	8265	1892
563857	N/A	N/A	TAACATTTAATCAAAATTGT	0	8250	8269	1893
563858	N/A	N/A	ACATAACATTTAATCAAAAT	0	8253	8272	1894
563859	N/A	N/A	ATGACATAACATTTAATCAA	13	8256	8275	1895
563860	N/A	N/A	TACTTATGACATAACATTTA	0	8261	8280	1896
563861	N/A	N/A	TTACTACTTATGACATAACA	0	8265	8284	1897
563862	N/A	N/A	AACAGTTACTACTTATGACA	31	8270	8289	1898
563863	N/A	N/A	TGTAACAGTTACTACTTATG	29	8273	8292	1899
563864	N/A	N/A	CTTATTTGTAACAGTTACTA	0	8279	8298	1900
563865	N/A	N/A	TTTCACAGCTTATTTGTAAC	29	8287	8306	1901
563866	N/A	N/A	TCTTTTCACAGCTTATTTGT	22	8290	8309	1902
563867	N/A	N/A	GGTTCTTTTCACAGCTTATT	66	8293	8312	1903
563868	N/A	N/A	CTAGGAGTGGTTCTTTTCAC	37	8301	8320	1904
563869	N/A	N/A	ATGCTAGGAGTGGTTCTTTT	20	8304	8323	1905
563870	N/A	N/A	CTAATGCTAGGAGTGGTTCT	30	8307	8326	1906
563871	N/A	N/A	AGAGTGACTAATGCTAGGAG	41	8314	8333	1907
563872	N/A	N/A	AGAGAATAGAGTGACTAATG	28	8321	8340	1908
563873	N/A	N/A	TTAATGAGAGAATAGAGTGA	4	8327	8346	1909
563496	608	627	CTGTTGGTTTAATTGTTTAT	33	4346	4365	1910
563497	610	629	TGCTGTTGGTTTAATTGTTT	29	4348	4367	1911
563498	612	631	TATGCTGTTGGTTTAATTGT	27	4350	4369	1912
563499	614	633	ACTATGCTGTTGGTTTAATT	24	4352	4371	1913
563500	616	635	TGACTATGCTGTTGGTTTAA	68	4354	4373	1914
563501	619	638	ATTTGACTATGCTGTTGGTT	45	4357	4376	1915
563502	621	640	TTATTTGACTATGCTGTTGG	39	4359	4378	1916
563503	623	642	TTTTATTTGACTATGCTGTT	33	4361	4380	1917
563504	625	644	TCTTTTATTTGACTATGCTG	55	4363	4382	1918
563505	627	646	TTTCTTTTATTTGACTATGC	29	4365	4384	1919
563506	646	665	CTTCTGAGCTGATTTTCTAT	40	N/A	N/A	1920
563507	648	667	TCCTTCTGAGCTGATTTTCT	76	N/A	N/A	1921
563508	650	669	AGTCCTTCTGAGCTGATTTT	37	N/A	N/A	1922

563509	652	671	CTAGTCCTTCTGAGCTGATT	52	N/A	N/A	1923
563510	654	673	TACTAGTCCTTCTGAGCTGA	52	6667	6686	1924
563511	656	675	AATACTAGTCCTTCTGAGCT	41	6669	6688	1925
563512	658	677	TGAATACTAGTCCTTCTGAG	55	6671	6690	1926
563513	660	679	CTTGAATACTAGTCCTTCTG	43	6673	6692	1927
563514	662	681	TTCTTGAATACTAGTCCTTC	34	6675	6694	1928
563515	666	685	TGGGTTCTTGAATACTAGTC	52	6679	6698	1929
563516	668	687	TGTGGGTTCTTGAATACTAG	34	6681	6700	1930
563517	670	689	TCTGTGGGTTCTTGAATACT	43	6683	6702	1931
563518	680	699	TAGAGAAATTTCTGTGGGTT	0	6693	6712	1932
563519	684	703	AAGATAGAGAAATTTCTGTG	4	6697	6716	1933
563520	686	705	GGAAGATAGAGAAATTTCTG	0	6699	6718	1934
563521	694	713	CTTGGCTTGGAAGATAGAGA	29	6707	6726	1935
563522	696	715	CTCTTGGCTTGGAAGATAGA	51	6709	6728	1936
563523	705	724	TTCTTGGTGCTCTTGGCTTG	63	6718	6737	75
544120	707	726	AGTTCTTGGTGCTCTTGGCT	86	6720	6739	15
563524	715	734	AAGGGAGTAGTTCTTGGTGC	44	6728	6747	1937
563525	716	735	AAAGGGAGTAGTTCTTGGTG	14	6729	6748	1938
563526	717	736	GAAAGGGAGTAGTTCTTGGT	33	6730	6749	1939
563527	718	737	AGAAAGGGAGTAGTTCTTGG	0	6731	6750	1940
563528	719	738	AAGAAAGGGAGTAGTTCTTG	0	6732	6751	1941
563529	720	739	GAAGAAAGGGAGTAGTTCTT	0	6733	6752	1942
563530	726	745	TCAACTGAAGAAAGGGAGTA	0	6739	6758	1943
337481	728	747	ATTCAACTGAAGAAAGGGAG	23	6741	6760	1944
563531	729	748	CATTCAACTGAAGAAAGGGA	16	6742	6761	1945
563532	730	749	TCATTCAACTGAAGAAAGGG	23	6743	6762	1946
563533	732	751	TTTCATTCAACTGAAGAAAG	8	6745	6764	1947
563534	733	752	ATTTCAATTCAACTGAAGAAA	6	6746	6765	1948
563535	734	753	TATTTCAATTCAACTGAAGAA	0	6747	6766	1949
563536	735	754	TTATTTCAATTCAACTGAAGA	0	6748	6767	1950
563537	736	755	CTTATTTCAATTCAACTGAAG	11	6749	6768	1951
337482	737	756	TCTTATTTCAATTCAACTGAA	26	6750	6769	1952
563538	738	757	TTCTTATTTCAATTCAACTGA	17	6751	6770	1953
563539	740	759	ATTTCTTATTTCAATTCAACT	18	6753	6772	1954
563540	743	762	TACATTTCTTATTTCAATTCA	20	6756	6775	1955
563541	767	786	TTCAGCAGGAATGCCATCAT	34	N/A	N/A	1956
563542	768	787	ATTCAGCAGGAATGCCATCA	2	N/A	N/A	1957
563543	769	788	CATTCAGCAGGAATGCCATC	21	N/A	N/A	1958
563544	770	789	ACATTCAGCAGGAATGCCAT	5	N/A	N/A	1959
563545	771	790	TACATTCAGCAGGAATGCCA	37	N/A	N/A	1960
563546	772	791	GTACATTCAGCAGGAATGCC	50	7357	7376	1961

563547	773	792	GGTACATTCAGCAGGAATGC	64	7358	7377	76
563548	774	793	TGGTACATTCAGCAGGAATG	42	7359	7378	1962
563549	775	794	GTGGTACATTCAGCAGGAAT	51	7360	7379	1963
563550	776	795	GGTGGTACATTCAGCAGGAA	24	7361	7380	1964
563551	777	796	TGGTGGTACATTCAGCAGGA	47	7362	7381	1965
563552	778	797	ATGGTGGTACATTCAGCAGG	0	7363	7382	1966
563553	779	798	AATGGTGGTACATTCAGCAG	15	7364	7383	1967
563554	780	799	AAATGGTGGTACATTCAGCA	32	7365	7384	1968
563555	781	800	TAAATGGTGGTACATTCAGC	29	7366	7385	1969
563556	783	802	TATAAATGGTGGTACATTCA	33	7368	7387	1970
563557	784	803	TTATAAATGGTGGTACATTC	1	7369	7388	1971
563558	785	804	GTTATAAATGGTGGTACATT	4	7370	7389	1972
563559	786	805	TGTTATAAATGGTGGTACAT	0	7371	7390	1973
563560	787	806	CTGTTATAAATGGTGGTACA	4	7372	7391	1974
563561	788	807	TCTGTTATAAATGGTGGTAC	29	7373	7392	1975
337484	789	808	CTCTGTTATAAATGGTGGTA	62	7374	7393	74
563562	792	811	CACCTCTGTTATAAATGGTG	22	7377	7396	1976
563563	793	812	TCACCTCTGTTATAAATGGT	38	7378	7397	1977
337485	794	813	TTCACCTCTGTTATAAATGG	18	7379	7398	1978
563564	795	814	GTTACCTCTGTTATAAATG	52	7380	7399	1979
563565	797	816	ATGTTACCTCTGTTATAAA	24	7382	7401	1980
563566	798	817	TATGTTACCTCTGTTATAA	2	7383	7402	1981
337486	799	818	GTATGTTACCTCTGTTATA	32	7384	7403	1982
563567	800	819	TGTATGTTACCTCTGTTAT	38	7385	7404	1983
337487	804	823	CACTTGTATGTTACCTCTG	87	7389	7408	28
563568	1128	1147	TAATCGCAACTAGATGTAGC	39	9703	9722	1984
563569	1129	1148	GTAATCGCAACTAGATGTAG	26	9704	9723	1985
563570	1130	1149	AGTAATCGCAACTAGATGTA	17	9705	9724	1986
563571	1131	1150	CAGTAATCGCAACTAGATGT	43	9706	9725	1987
563572	1132	1151	CCAGTAATCGCAACTAGATG	39	9707	9726	1988
563573	1133	1152	GCCAGTAATCGCAACTAGAT	59	9708	9727	1989
563574	1134	1153	TGCCAGTAATCGCAACTAGA	57	9709	9728	1990
563575	1135	1154	TTGCCAGTAATCGCAACTAG	54	9710	9729	1991
563576	1136	1155	ATTGCCAGTAATCGCAACTA	43	9711	9730	1992
563577	1137	1156	CATTGCCAGTAATCGCAACT	49	9712	9731	1993
563578	1138	1157	ACATTGCCAGTAATCGCAAC	59	9713	9732	1994
563579	1139	1158	GACATTGCCAGTAATCGCAA	64	9714	9733	1995
563580	1140	1159	GGACATTGCCAGTAATCGCA	79	9715	9734	77
563581	1141	1160	GGGACATTGCCAGTAATCGC	47	9716	9735	1996
563582	1162	1181	TTGTTTTCCGGGATTGCATT	20	9737	9756	1997
563583	1163	1182	TTTGTTTTCCGGGATTGCAT	31	9738	9757	1998

563584	1167	1186	AATCTTTGTTTTCCGGGATT	14	9742	9761	1999
563585	1168	1187	AAATCTTTGTTTTCCGGGAT	54	9743	9762	2000
563586	1175	1194	AAACACCAAATCTTTGTTTT	32	9750	9769	2001
563587	1176	1195	AAAACACCAAATCTTTGTTTT	7	9751	9770	2002
563588	1180	1199	GTAGAAAACACCAAATCTTT	18	9755	9774	2003
563589	1181	1200	AGTAGAAAACACCAAATCTT	0	9756	9775	2004
563590	1185	1204	CCCAAGTAGAAAACACCAAA	26	9760	9779	2005
563591	1186	1205	TCCCAAGTAGAAAACACCAA	27	9761	9780	2006
563592	1190	1209	GTGATCCCAAGTAGAAAACA	26	9765	9784	2007
563593	1191	1210	TGTGATCCCAAGTAGAAAAC	28	9766	9785	2008
563594	1192	1211	TTGTGATCCCAAGTAGAAAA	12	9767	9786	2009
563595	1193	1212	TTTGTGATCCCAAGTAGAAA	14	9768	9787	2010
563596	1200	1219	CTTTTGCTTTGTGATCCCAA	64	9775	9794	2011
563597	1204	1223	TGTCCTTTTGCTTTGTGATC	24	9779	9798	2012
563598	1205	1224	GTGTCCTTTTGCTTTGTGAT	31	9780	9799	2013
563599	1206	1225	AGTGTCTTTTGCTTTGTGA	41	9781	9800	2014
563600	1210	1229	TTGAAGTGTCCTTTTGCTTT	21	9785	9804	2015
563601	1211	1230	GTTGAAGTGTCCTTTTGCTT	35	9786	9805	2016
563602	1212	1231	AGTTGAAGTGTCCTTTTGCT	27	9787	9806	2017
563603	1213	1232	CAGTTGAAGTGTCCTTTTGC	17	9788	9807	2018
563604	1214	1233	ACAGTTGAAGTGTCCTTTTG	0	9789	9808	2019
563605	1215	1234	GACAGTTGAAGTGTCCTTTT	19	9790	9809	2020
563606	1216	1235	GGACAGTTGAAGTGTCCTTT	34	9791	9810	2021
563607	1217	1236	TGGACAGTTGAAGTGTCCTT	12	9792	9811	2022
563608	1218	1237	CTGGACAGTTGAAGTGTCCT	39	9793	9812	2023
563609	1219	1238	TCTGGACAGTTGAAGTGTC	10	9794	9813	2024
563610	1220	1239	CTCTGGACAGTTGAAGTGTC	6	9795	9814	2025
563611	1221	1240	CCTCTGGACAGTTGAAGTGT	24	9796	9815	2026
563612	1222	1241	CCCTCTGGACAGTTGAAGTG	24	9797	9816	2027
563613	1223	1242	ACCCTCTGGACAGTTGAAGT	31	9798	9817	2028
563614	1224	1243	AACCCTCTGGACAGTTGAAG	34	9799	9818	2029
563615	1225	1244	TAACCCTCTGGACAGTTGAA	34	9800	9819	2030
563616	1226	1245	ATAACCCTCTGGACAGTTGA	31	9801	9820	2031
563617	1227	1246	AATAACCCTCTGGACAGTTG	22	9802	9821	2032
563618	1228	1247	GAATAACCCTCTGGACAGTT	25	9803	9822	2033
563619	1229	1248	TGAATAACCCTCTGGACAGT	18	9804	9823	2034
563620	1230	1249	CTGAATAACCCTCTGGACAG	24	9805	9824	2035
563621	1231	1250	CCTGAATAACCCTCTGGACA	39	9806	9825	2036
563622	1232	1251	TCCTGAATAACCCTCTGGAC	31	N/A	N/A	2037
563623	1233	1252	CTCCTGAATAACCCTCTGGA	15	N/A	N/A	2038
563624	1234	1253	CCTCCTGAATAACCCTCTGG	27	N/A	N/A	2039

563625	1235	1254	GCCTCCTGAATAACCCTCTG	25	N/A	N/A	2040
563626	1236	1255	AGCCTCCTGAATAACCCTCT	32	N/A	N/A	2041
563627	1237	1256	CAGCCTCCTGAATAACCCTC	44	N/A	N/A	2042
563628	1238	1257	CCAGCCTCCTGAATAACCCT	26	N/A	N/A	2043
563629	1239	1258	ACCAGCCTCCTGAATAACCC	23	N/A	N/A	2044
337503	1240	1259	CACCAGCCTCCTGAATAACC	25	N/A	N/A	2045
563630	1241	1260	CCACCAGCCTCCTGAATAAC	26	N/A	N/A	2046
563631	1242	1261	ACCACCAGCCTCCTGAATAA	25	N/A	N/A	2047
563632	1243	1262	CACCACCAGCCTCCTGAATA	33	N/A	N/A	2048
563633	1244	1263	CCACCACCAGCCTCCTGAAT	45	N/A	N/A	2049
563634	1248	1267	CATGCCACCACCAGCCTCCT	54	10220	10239	2050
563635	1250	1269	ATCATGCCACCACCAGCCTC	58	10222	10241	2051
563636	1251	1270	CATCATGCCACCACCAGCCT	61	10223	10242	2052
563637	1255	1274	CACTCATCATGCCACCACCA	68	10227	10246	78
563638	1256	1275	AACTCATCATGCCACCACC	65	10228	10247	2053
563639	1260	1279	CTCCACACTCATCATGCCAC	76	10232	10251	79
563640	1262	1281	TTCTCCACACTCATCATGCC	55	10234	10253	2054
563641	1263	1282	TTTCTCCACACTCATCATGC	63	10235	10254	80
563642	1264	1283	TTTTCTCCACACTCATCATG	24	10236	10255	2055
563643	1265	1284	GTTTTCTCCACACTCATCAT	53	10237	10256	2056
563644	1857	1876	ATTTAAGAACTGTACAATTA	7	10829	10848	2057
563645	1858	1877	CATTTAAGAACTGTACAATT	15	10830	10849	2058
563646	1859	1878	ACATTTAAGAACTGTACAAT	4	10831	10850	2059
563647	1860	1879	AACATTTAAGAACTGTACAA	4	10832	10851	2060
563648	1861	1880	CAACATTTAAGAACTGTACA	4	10833	10852	2061
563649	1862	1881	ACAACATTTAAGAACTGTAC	22	10834	10853	2062
563650	1863	1882	TACAACATTTAAGAACTGTA	21	10835	10854	2063
563651	1864	1883	CTACAACATTTAAGAACTGT	44	10836	10855	2064
563652	1865	1884	ACTACAACATTTAAGAACTG	20	10837	10856	2065
563653	1866	1885	TACTACAACATTTAAGAACT	15	10838	10857	2066
563654	1867	1886	ATACTACAACATTTAAGAAC	17	10839	10858	2067
563655	1868	1887	AATACTACAACATTTAAGAA	11	10840	10859	2068
563656	1869	1888	TAATACTACAACATTTAAGA	9	10841	10860	2069
563657	1870	1889	TTAATACTACAACATTTAAG	3	10842	10861	2070
563658	1874	1893	GAAATTAATACTACAACATT	0	10846	10865	2071
563659	1878	1897	TTTTGAAATTAATACTACAA	0	10850	10869	2072
563660	1879	1898	GTTTTGAAATTAATACTACA	15	10851	10870	2073
563661	1880	1899	AGTTTTGAAATTAATACTAC	2	10852	10871	2074
563662	1881	1900	TAGTTTTGAAATTAATACTA	14	10853	10872	2075
563663	1882	1901	TTAGTTTTGAAATTAATACT	8	10854	10873	2076
563664	1888	1907	CGATTTTTAGTTTTGAAATT	0	10860	10879	2077

563665	1889	1908	ACGATTTTTAGTTTTGAAAT	0	10861	10880	2078
563666	1890	1909	GACGATTTTTAGTTTTGAAA	20	10862	10881	2079
563667	1891	1910	TGACGATTTTTAGTTTTGAA	17	10863	10882	2080
563668	1892	1911	CTGACGATTTTTAGTTTTGA	64	10864	10883	2081
563669	1893	1912	GCTGACGATTTTTAGTTTTG	66	10865	10884	81
563670	1894	1913	TGCTGACGATTTTTAGTTTT	45	10866	10885	2082
563671	1895	1914	GTGCTGACGATTTTTAGTTT	42	10867	10886	2083
563672	1896	1915	TGTGCTGACGATTTTTAGTT	50	10868	10887	2084
563673	1897	1916	CTGTGCTGACGATTTTTAGT	55	10869	10888	2085
563674	1898	1917	TCTGTGCTGACGATTTTTAG	53	10870	10889	2086
563675	1899	1918	CTCTGTGCTGACGATTTTTA	49	10871	10890	2087
563676	1900	1919	ACTCTGTGCTGACGATTTTT	22	10872	10891	2088
563677	1901	1920	TACTCTGTGCTGACGATTTT	8	10873	10892	2089
563678	1902	1921	ATACTCTGTGCTGACGATTT	61	10874	10893	2090
563679	1903	1922	CATACTCTGTGCTGACGATT	68	10875	10894	2091
563680	1904	1923	ACATACTCTGTGCTGACGAT	4	10876	10895	2092
563681	1905	1924	CACATACTCTGTGCTGACGA	73	10877	10896	82
563682	1909	1928	TTTACACATACTCTGTGCTG	67	10881	10900	83
563683	1911	1930	TTTTTACACATACTCTGTGC	58	10883	10902	2093
563684	1915	1934	CAGATTTTTACACATACTCT	54	10887	10906	2094
563685	1916	1935	ACAGATTTTTACACATACTC	52	10888	10907	2095
563686	1917	1936	TACAGATTTTTACACATACT	40	10889	10908	2096
563687	1918	1937	TTACAGATTTTTACACATAC	22	10890	10909	2097
337528	1920	1939	TATTACAGATTTTTACACAT	4	6720	6739	2098
563688	1922	1941	TGTATTACAGATTTTTACAC	0	10894	10913	2099
563689	1935	1954	CAGTTTAAAAATTTGTATTA	8	10907	10926	2100
563690	1938	1957	CATCAGTTTAAAAATTTGTA	18	10910	10929	2101
563691	1941	1960	AAGCATCAGTTTAAAAATTT	16	10913	10932	2102
563692	1942	1961	GAAGCATCAGTTTAAAAATT	16	10914	10933	2103
563693	1951	1970	TAGCAAAATGAAGCATCAGT	40	10923	10942	2104
563694	1952	1971	GTAGCAAAATGAAGCATCAG	42	10924	10943	2105
563695	1953	1972	TGTAGCAAAATGAAGCATCA	44	10925	10944	2106
563696	1954	1973	TTGTAGCAAAATGAAGCATC	48	10926	10945	2107
563697	1955	1974	TTTGTAGCAAAATGAAGCAT	19	10927	10946	2108
563698	1974	1993	AACATTTACTCCAAATTATT	27	10946	10965	2109
563699	1976	1995	CAAACATTTACTCCAAATTA	23	10948	10967	2110
563700	1978	1997	ATCAAACATTTACTCCAAAT	24	10950	10969	2111
563701	1981	2000	CATATCAAACATTTACTCCA	61	10953	10972	2112
563702	1982	2001	TCATATCAAACATTTACTCC	50	10954	10973	2113
563703	1983	2002	ATCATATCAAACATTTACTC	31	10955	10974	2114
563704	1990	2009	TAAATAAATCATATCAAACA	10	10962	10981	2115

563705	1993	2012	TCATAAATAAATCATATCAA	20	10965	10984	2116
563706	1994	2013	TTCATAAATAAATCATATCA	11	10966	10985	2117
563707	1995	2014	TTTCATAAATAAATCATATC	5	10967	10986	2118
563708	1996	2015	GTTTCATAAATAAATCATAT	0	10968	10987	2119
563709	1997	2016	GGTTTCATAAATAAATCATA	8	10969	10988	2120
563710	1998	2017	AGGTTTCATAAATAAATCAT	15	10970	10989	2121
563711	1999	2018	TAGGTTTCATAAATAAATCA	19	10971	10990	2122
563712	2001	2020	ATTAGGTTTCATAAATAAAT	12	10973	10992	2123
563713	2002	2021	CATTAGGTTTCATAAATAAA	2	10974	10993	2124
563714	2003	2022	TCATTAGGTTTCATAAATAA	7	10975	10994	2125
563715	2004	2023	TTCATTAGGTTTCATAAATA	11	10976	10995	2126
563716	2005	2024	CTTCATTAGGTTTCATAAAT	15	10977	10996	2127
563717	2006	2025	GCTTCATTAGGTTTCATAAA	49	10978	10997	2128
563718	2010	2029	TTCTGCTTCATTAGGTTTCA	57	10982	11001	2129
563719	2013	2032	TAATTCTGCTTCATTAGGTT	43	10985	11004	2130

Table 11

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
566915	343	362	TATGTAGTTCTTCTCAGTTC	22	3447	3466	2131
566916	350	369	TAGTTTATATGTAGTTCTTC	21	3454	3473	2132
566917	354	373	CTTGTAGTTTATATGTAGTT	12	3458	3477	2133
566918	358	377	TTGACTTGTAGTTTATATGT	12	3462	3481	2134
566919	360	379	TTTTGACTTGTAGTTTATAT	0	3464	3483	2135
566920	362	381	ATTTTTGACTTGTAGTTTAT	7	3466	3485	2136
566921	367	386	TCTTCATTTTTGACTTGTAG	33	3471	3490	2137
566922	371	390	TACCTCTTCATTTTTGACTT	22	3475	3494	2138
566923	377	396	ATTCTTTACCTCTTCATTTT	12	3481	3500	2139
566924	387	406	CAAGTGACATATTCTTTACC	36	3491	3510	2140
566925	389	408	TTCAAGTGACATATTCTTTA	31	3493	3512	2141
566926	394	413	TTGAGTTCAAGTGACATATT	18	3498	3517	2142
566927	396	415	AGTTGAGTTCAAGTGACATA	6	3500	3519	2143
566928	400	419	TTTGAGTTGAGTTCAAGTGA	11	3504	3523	2144
566929	408	427	TTTCAAGTTTTGAGTTGAGT	15	3512	3531	2145
566930	410	429	GCTTTCAAGTTTTGAGTTGA	13	3514	3533	2146
566931	412	431	AGGCTTTCAAGTTTTGAGTT	22	3516	3535	2147
566932	416	435	TAGGAGGCTTTCAAGTTTGT	4	3520	3539	2148
566933	419	438	TTCTAGGAGGCTTTCAAGTT	35	3523	3542	2149
566934	421	440	TCTTCTAGGAGGCTTTCAAG	26	3525	3544	2150

566935	429	448	GAATTTTTTCTTCTAGGAGG	1	3533	3552	2151
566936	434	453	AAGTAGAATTTTTTCTTCTA	0	3538	3557	2152
566937	436	455	TGAAGTAGAATTTTTTCTTC	11	3540	3559	2153
566938	438	457	GTTGAAGTAGAATTTTTTCT	29	3542	3561	2154
566939	441	460	TTTGTTGAAGTAGAATTTTT	11	3545	3564	2155
566940	443	462	TTTTTGTTGAAGTAGAATTT	35	3547	3566	2156
566941	464	483	TTGCTCTTCTAAATATTTCA	35	3568	3587	2157
566942	466	485	AGTTGCTCTTCTAAATATTT	53	3570	3589	2158
566943	468	487	TTAGTTGCTCTTCTAAATAT	18	3572	3591	2159
566944	471	490	TAGTTAGTTGCTCTTCTAAA	38	3575	3594	2160
566945	476	495	TAAGTTAGTTAGTTGCTCTT	28	3580	3599	2161
566946	478	497	ATTAAGTTAGTTAGTTGCTC	28	3582	3601	2162
566947	480	499	GAATTAAGTTAGTTAGTTGC	27	3584	3603	2163
566948	482	501	TTGAATTAAGTTAGTTAGTT	21	3586	3605	2164
566949	484	503	TTTTGAATTAAGTTAGTTAG	2	3588	3607	2165
566950	487	506	TGATTTTGAATTAAGTTAGT	9	3591	3610	2166
566951	490	509	GGTTGATTTTGAATTAAGTT	52	3594	3613	2167
566952	497	516	AGTTTCAGGTTGATTTTGAA	13	3601	3620	2168
566953	501	520	CTGGAGTTTCAGGTTGATTT	50	3605	3624	2169
566954	507	526	GGTGTTCTGGAGTTTCAGGT	35	3611	3630	2170
566955	509	528	TGGGTGTTCTGGAGTTTCAG	18	3613	3632	2171
566956	511	530	TCTGGGTGTTCTGGAGTTTC	32	3615	3634	2172
566957	513	532	CTTCTGGGTGTTCTGGAGTT	28	3617	3636	2173
566958	515	534	TACTTCTGGGTGTTCTGGAG	23	3619	3638	2174
566959	517	536	GTTACTTCTGGGTGTTCTGG	12	3621	3640	2175
566960	519	538	AAGTTACTTCTGGGTGTTCT	1	3623	3642	2176
566961	522	541	GTGAAGTTACTTCTGGGTGT	0	3626	3645	2177
566962	528	547	TTTAAAGTGAAGTTACTTCT	6	N/A	N/A	2178
566963	530	549	AGTTTTAAAGTGAAGTTACTT	16	N/A	N/A	2179
566964	532	551	AAAGTTTTAAAGTGAAGTTAC	12	N/A	N/A	2180
566965	535	554	ACAAAAGTTTTAAAGTGAAGT	8	N/A	N/A	2181
337474	537	556	CTACAAAAGTTTTAAAGTGAA	10	N/A	N/A	2182
566966	539	558	TTCTACAAAAGTTTTAAAGTG	46	N/A	N/A	2183
566967	544	563	TGTTTTTCTACAAAAGTTTT	12	N/A	N/A	2184
566968	546	565	CTTGTTTTTCTACAAAAGTT	0	N/A	N/A	2185
566969	552	571	TATTATCTTGTTTTTCTACA	0	4290	4309	2186
566970	557	576	GATGCTATTATCTTGTTTTT	18	4295	4314	2187
566971	560	579	TTTGATGCTATTATCTTGTT	22	4298	4317	2188
566972	562	581	TCTTTGATGCTATTATCTTG	21	4300	4319	2189
566973	569	588	GAGAAGGTCTTTGATGCTAT	37	4307	4326	2190
566974	574	593	GTCTGGAGAAGGTCTTTGAT	26	4312	4331	2191

566975	576	595	CGGTCTGGAGAAGGTCTTTG	20	4314	4333	2192
566976	578	597	CACGGTCTGGAGAAGGTCTT	53	4316	4335	2193
566977	580	599	TCCACGGTCTGGAGAAGGTC	58	4318	4337	2194
566978	582	601	CTTCCACGGTCTGGAGAAGG	39	4320	4339	2195
566979	584	603	GTCTTCCACGGTCTGGAGAA	63	4322	4341	2196
566980	586	605	TGGTCTTCCACGGTCTGGAG	81	4324	4343	2197
566981	588	607	ATTGGTCTTCCACGGTCTGG	57	4326	4345	2198
566982	590	609	ATATTGGTCTTCCACGGTCT	60	4328	4347	2199
566983	592	611	TTATATTGGTCTTCCACGGT	49	4330	4349	2200
566984	594	613	GTTTATATTGGTCTTCCACG	54	4332	4351	2201
566985	596	615	TTGTTTATATTGGTCTTCCA	36	4334	4353	2202
566986	598	617	AATTGTTTATATTGGTCTTC	23	4336	4355	2203
566987	600	619	TTAATTGTTTATATTGGTCT	26	4338	4357	2204
566988	602	621	GTTTAATTGTTTATATTGGT	23	4340	4359	2205
566989	604	623	TGGTTTAATTGTTTATATTG	8	4342	4361	2206
566990	606	625	GTTGGTTTAATTGTTTATAT	1	4344	4363	2207
544120	707	726	AGTTCTTGGTGCTCTTGGCT	78	6720	6739	15
337487	804	823	CACTTGTATGTTACCTCTG	82	7389	7408	28
566991	912	931	TTTGTGATCCATCTATTCGA	25	7899	7918	2208
566992	913	932	TTTTGTGATCCATCTATTCG	12	7900	7919	2209
566993	920	939	ATTGAAGTTTTGTGATCCAT	32	7907	7926	2210
566994	921	940	CATTGAAGTTTTGTGATCCA	26	7908	7927	2211
566995	922	941	TCATTGAAGTTTTGTGATCC	0	7909	7928	2212
566996	923	942	TTCATTGAAGTTTTGTGATC	1	7910	7929	2213
566997	924	943	TTTCATTGAAGTTTTGTGAT	20	7911	7930	2214
566998	944	963	ATATTTGTAGTTCTCCCACG	35	7931	7950	2215
566999	952	971	CCAAAACCATATTTGTAGTT	13	7939	7958	2216
567000	953	972	CCCAAACCATATTTGTAGT	21	7940	7959	2217
567001	954	973	TCCCAAACCATATTTGTAG	0	7941	7960	2218
567002	955	974	CTCCCAAACCATATTTGTA	5	7942	7961	2219
567003	958	977	AGCCTCCCAAACCATATTT	0	7945	7964	2220
567004	960	979	CAAGCCTCCCAAACCATAT	14	7947	7966	2221
567005	961	980	TCAAGCCTCCCAAACCATATA	0	7948	7967	2222
567006	962	981	ATCAAGCCTCCCAAACCAT	17	7949	7968	2223
567007	963	982	CATCAAGCCTCCCAAACCA	31	7950	7969	2224
567008	964	983	CCATCAAGCCTCCCAAACC	11	7951	7970	2225
567009	965	984	TCCATCAAGCCTCCCAAAC	27	N/A	N/A	2226
567010	966	985	CTCCATCAAGCCTCCCAAAA	42	N/A	N/A	2227
567011	972	991	AAAATTCTCCATCAAGCCTC	48	N/A	N/A	2228
567012	974	993	CCAAAATTCTCCATCAAGCC	41	N/A	N/A	2229
567013	975	994	ACCAAATTCTCCATCAAGC	49	N/A	N/A	2230

567014	978	997	CCAACCAAAATTCTCCATCA	32	N/A	N/A	2231
567015	979	998	CCCAACCAAAATTCTCCATC	47	N/A	N/A	2232
337497	980	999	GCCCAACCAAAATTCTCCAT	46	N/A	N/A	2233
567016	981	1000	GGCCCAACCAAAATTCTCCA	48	N/A	N/A	2234
567017	982	1001	AGGCCCAACCAAAATTCTCC	30	9557	9576	2235
567018	983	1002	TAGGCCCAACCAAAATTCTC	0	9558	9577	2236
567019	984	1003	CTAGGCCCAACCAAAATTCT	31	9559	9578	2237
567020	985	1004	TCTAGGCCCAACCAAAATTC	39	9560	9579	2238
233721	986	1005	CTCTAGGCCCAACCAAAATT	15	9561	9580	2239
567021	987	1006	TCTCTAGGCCCAACCAAAAT	36	9562	9581	2240
567022	988	1007	TTCTCTAGGCCCAACCAAAA	26	9563	9582	2241
567023	989	1008	CTTCTCTAGGCCCAACCAAA	44	9564	9583	2242
567024	993	1012	ATATCTTCTCTAGGCCCAAC	29	9568	9587	2243
567025	994	1013	TATATCTTCTCTAGGCCCAA	41	9569	9588	2244
567026	995	1014	GTATATCTTCTCTAGGCCCA	53	9570	9589	2245
567027	1000	1019	ATGGAGTATATCTTCTCTAG	18	9575	9594	2246
567028	1004	1023	CACTATGGAGTATATCTTCT	35	9579	9598	2247
567029	1005	1024	TCACTATGGAGTATATCTTC	9	9580	9599	2248
567030	1006	1025	TTCACTATGGAGTATATCTT	11	9581	9600	2249
567031	1010	1029	TTGCTTCACTATGGAGTATA	43	9585	9604	2250
567032	1011	1030	ATTGCTTCACTATGGAGTAT	4	9586	9605	2251
567033	1015	1034	TTAGATTGCTTCACTATGGA	17	9590	9609	2252
567034	1016	1035	ATTAGATTGCTTCACTATGG	35	9591	9610	2253
567035	1017	1036	AATTAGATTGCTTCACTATG	18	9592	9611	2254
567036	1018	1037	TAATTAGATTGCTTCACTAT	17	9593	9612	2255
567037	1019	1038	ATAATTAGATTGCTTCACTA	19	9594	9613	2256
567038	1020	1039	CATAATTAGATTGCTTCACT	27	9595	9614	2257
567039	1021	1040	ACATAATTAGATTGCTTCAC	17	9596	9615	2258
337498	1022	1041	AACATAATTAGATTGCTTCA	9	9597	9616	2259
567040	1023	1042	AAACATAATTAGATTGCTTC	0	9598	9617	2260
567041	1024	1043	AAAACATAATTAGATTGCTT	0	9599	9618	2261
567042	1025	1044	TAAAACATAATTAGATTGCT	23	9600	9619	2262
567043	1026	1045	GTAAAACATAATTAGATTGC	25	9601	9620	2263
567044	1027	1046	CGTAAAACATAATTAGATTG	0	9602	9621	2264
567045	1048	1067	TTCCAGTCTTCCAACCTCAAT	9	9623	9642	2265
337500	1050	1069	CTTTCAGTCTTCCAACCTCA	30	9625	9644	2266
567046	1057	1076	TTGTTGTCTTTCCAGTCTTC	40	9632	9651	2267
567047	1064	1083	ATAATGTTTGTTGTCTTTCC	26	9639	9658	2268
567048	1065	1084	TATAATGTTTGTTGTCTTTC	6	9640	9659	2269
567049	1066	1085	ATATAATGTTTGTTGTCTTT	9	9641	9660	2270
567050	1069	1088	TCAATATAATGTTTGTTGTC	20	9644	9663	2271

567051	1073	1092	ATATTCAATATAATGTTTGT	15	9648	9667	2272
567052	1074	1093	AATATTCAATATAATGTTTG	16	9649	9668	2273
567053	1075	1094	GAATATTCAATATAATGTTT	7	9650	9669	2274
567054	1076	1095	AGAATATTCAATATAATGTT	3	9651	9670	2275
567055	1077	1096	AAGAATATTCAATATAATGT	7	9652	9671	2276
567056	1085	1104	CAAGTAAAAAGAATATTCAA	0	9660	9679	2277
567057	1086	1105	CCAAGTAAAAAGAATATTCA	0	9661	9680	2278
567058	1087	1106	CCCAAGTAAAAAGAATATTC	13	9662	9681	2279
567059	1090	1109	TTTCCCAAGTAAAAAGAATA	0	9665	9684	2280
567060	1091	1110	ATTTCCCAAGTAAAAAGAAT	2	9666	9685	2281
567061	1092	1111	GATTTCCCAAGTAAAAAGAA	14	9667	9686	2282
567062	1093	1112	TGATTTCCCAAGTAAAAAGA	14	9668	9687	2283
567063	1127	1146	AATCGCAACTAGATGTAGCG	15	9702	9721	2284
563874	1586	1605	ATTCTTTAAGGTTATGTGAT	13	10558	10577	2285
563875	1587	1606	TATTCTTTAAGGTTATGTGA	25	10559	10578	2286
563876	1591	1610	ACGGTATTCTTTAAGGTTAT	50	10563	10582	2287
563877	1592	1611	AACGGTATTCTTTAAGGTTA	48	10564	10583	2288
563878	1593	1612	AAACGGTATTCTTTAAGGTT	45	10565	10584	2289
563879	1594	1613	TAAACGGTATTCTTTAAGGT	16	10566	10585	2290
563880	1595	1614	GTAAACGGTATTCTTTAAGG	14	10567	10586	2291
563881	1596	1615	TGTAAACGGTATTCTTTAAG	0	10568	10587	2292
563882	1597	1616	ATGTAAACGGTATTCTTTAA	10	10569	10588	2293
563883	1598	1617	AATGTAAACGGTATTCTTTA	12	10570	10589	2294
563884	1599	1618	AAATGTAAACGGTATTCTTT	15	10571	10590	2295
563885	1600	1619	GAAATGTAAACGGTATTCTT	13	10572	10591	2296
563886	1601	1620	AGAAATGTAAACGGTATTCT	22	10573	10592	2297
563887	1602	1621	GAGAAATGTAAACGGTATTC	35	10574	10593	2298
563888	1603	1622	TGAGAAATGTAAACGGTATT	14	10575	10594	2299
563889	1604	1623	TTGAGAAATGTAAACGGTAT	0	10576	10595	2300
563890	1605	1624	ATTGAGAAATGTAAACGGTA	18	10577	10596	2301
563891	1606	1625	GATTGAGAAATGTAAACGGT	40	10578	10597	2302
563892	1607	1626	TGATTGAGAAATGTAAACGG	33	10579	10598	2303
563893	1608	1627	TTGATTGAGAAATGTAAACG	7	10580	10599	2304
563894	1609	1628	TTTGATTGAGAAATGTAAAC	0	10581	10600	2305
563895	1610	1629	TTTTGATTGAGAAATGTAAA	0	10582	10601	2306
563896	1611	1630	ATTTTGATTGAGAAATGTAA	0	10583	10602	2307
563897	1612	1631	AATTTTGATTGAGAAATGTA	0	10584	10603	2308
563898	1613	1632	GAATTTTGATTGAGAAATGT	4	10585	10604	2309
563899	1614	1633	AGAATTTTGATTGAGAAATG	4	10586	10605	2310
563900	1615	1634	AAGAATTTTGATTGAGAAAT	26	10587	10606	2311
563901	1617	1636	ATAAGAATTTTGATTGAGAA	4	10589	10608	2312

563902	1618	1637	TATAAGAATTTTGATTGAGA	0	10590	10609	2313
563903	1619	1638	TTATAAGAATTTTGATTGAG	0	10591	10610	2314
563904	1620	1639	ATTATAAGAATTTTGATTGA	0	10592	10611	2315
563905	1621	1640	TATTATAAGAATTTTGATTG	3	10593	10612	2316
563906	1622	1641	GTATTATAAGAATTTTGATT	1	10594	10613	2317
563907	1623	1642	AGTATTATAAGAATTTTGAT	44	10595	10614	2318
563908	1624	1643	TAGTATTATAAGAATTTTGA	29	10596	10615	2319
563909	1632	1651	AAAACAAATAGTATTATAAG	11	10604	10623	2320
563910	1633	1652	TAAAACAAATAGTATTATAA	16	10605	10624	2321
563911	1652	1671	ATTCCCACATCACAAAATTT	27	10624	10643	2322
563912	1653	1672	GATTCCCACATCACAAAATT	21	10625	10644	2323
563913	1654	1673	TGATTCCCACATCACAAAAT	49	10626	10645	2324
563914	1658	1677	AAATTGATTCCCACATCACA	47	10630	10649	2325
563915	1659	1678	AAAATTGATTCCCACATCAC	48	10631	10650	2326
563916	1663	1682	ATCTAAAATTGATTCCCACA	58	10635	10654	2327
563917	1667	1686	GACCATCTAAAATTGATTCC	41	10639	10658	2328
563918	1668	1687	TGACCATCTAAAATTGATTC	25	10640	10659	2329
563919	1669	1688	GTGACCATCTAAAATTGATT	33	10641	10660	2330
563920	1670	1689	TGTGACCATCTAAAATTGAT	34	10642	10661	2331
563921	1671	1690	TTGTGACCATCTAAAATTGA	20	10643	10662	2332
563922	1672	1691	ATTGTGACCATCTAAAATTG	2	10644	10663	2333
563923	1673	1692	GATTGTGACCATCTAAAATT	43	10645	10664	2334
563924	1674	1693	AGATTGTGACCATCTAAAAT	39	10646	10665	2335
563925	1675	1694	TAGATTGTGACCATCTAAAA	36	10647	10666	2336
563926	1676	1695	CTAGATTGTGACCATCTAAA	56	10648	10667	2337
563927	1677	1696	TCTAGATTGTGACCATCTAA	37	10649	10668	2338
563928	1678	1697	ATCTAGATTGTGACCATCTA	46	10650	10669	2339
563929	1679	1698	AATCTAGATTGTGACCATCT	56	10651	10670	2340
563930	1680	1699	TAATCTAGATTGTGACCATC	46	10652	10671	2341
563931	1681	1700	ATAATCTAGATTGTGACCAT	35	10653	10672	2342
563932	1682	1701	TATAATCTAGATTGTGACCA	45	10654	10673	2343
563933	1683	1702	TTATAATCTAGATTGTGACC	37	10655	10674	2344
563934	1686	1705	TGATTATAATCTAGATTGTG	28	10658	10677	2345
563935	1687	1706	TTGATTATAATCTAGATTGT	0	10659	10678	2346
563936	1688	1707	ATTGATTATAATCTAGATTG	0	10660	10679	2347
563937	1689	1708	TATTGATTATAATCTAGATT	0	10661	10680	2348
563938	1690	1709	CTATTGATTATAATCTAGAT	5	10662	10681	2349
563939	1691	1710	CCTATTGATTATAATCTAGA	0	10663	10682	2350
563940	1692	1711	ACCTATTGATTATAATCTAG	9	10664	10683	2351
563941	1693	1712	CACCTATTGATTATAATCTA	5	10665	10684	2352
563942	1694	1713	TCACCTATTGATTATAATCT	0	10666	10685	2353

563943	1695	1714	TTCACCTATTGATTATAATC	10	10667	10686	2354
563944	1696	1715	GTTCACCTATTGATTATAAT	31	10668	10687	2355
563945	1697	1716	AGTTCACCTATTGATTATAA	15	10669	10688	2356
563946	1698	1717	AAGTTCACCTATTGATTATA	31	10670	10689	2357
563947	1700	1719	ATAAGTTCACCTATTGATTA	9	10672	10691	2358
563948	1701	1720	AATAAGTTCACCTATTGATT	5	10673	10692	2359
563949	1702	1721	TAATAAGTTCACCTATTGAT	14	10674	10693	2360
563950	1703	1722	TTAATAAGTTCACCTATTGA	0	10675	10694	2361

Table 12

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
567064	N/A	N/A	TGAGTATTCTCGACTTACCT	26	8770	8789	2362
567065	N/A	N/A	AAGTGAGTATTCTCGACTTA	2	8773	8792	2363
567066	N/A	N/A	ATTAAGTGAGTATTCTCGAC	20	8776	8795	2364
567067	N/A	N/A	CCAGAATTAAGTGAGTATTC	36	8781	8800	2365
567068	N/A	N/A	GCTTTCTTACCAGAATTAAG	75	8790	8809	84
567069	N/A	N/A	GTTGCTTTCTTACCAGAATT	78	8793	8812	85
567070	N/A	N/A	TGGGTTGCTTTCTTACCAGA	26	8796	8815	2366
567071	N/A	N/A	AAATGGGTTGCTTTCTTACC	3	8799	8818	2367
567072	N/A	N/A	TACAAATGGGTTGCTTTCTT	24	8802	8821	2368
567073	N/A	N/A	AAGTACAAATGGGTTGCTTT	24	8805	8824	2369
567074	N/A	N/A	GTAAATACAAGTACAAATGG	7	8813	8832	2370
567075	N/A	N/A	TTGCTGGTAAATACAAGTAC	24	8819	8838	2371
567076	N/A	N/A	TAAGGATTGCTGGTAAATAC	6	8825	8844	2372
567077	N/A	N/A	TTTTAAGGATTGCTGGTAAA	4	8828	8847	2373
567078	N/A	N/A	GCTTCATTTTAAGGATTGCT	60	8834	8853	87
567079	N/A	N/A	GAAGCTTCATTTTAAGGATT	0	8837	8856	2374
567080	N/A	N/A	TAGGAAGCTTCATTTTAAGG	9	8840	8859	2375
567081	N/A	N/A	TAGTAGGAAGCTTCATTTTA	18	8843	8862	2376
567082	N/A	N/A	TTGAGTTAGTAGGAAGCTTC	30	8849	8868	2377
567083	N/A	N/A	ATTGCTATTGAGTTAGTAGG	21	8856	8875	2378
567084	N/A	N/A	CTTATTGCTATTGAGTTAGT	28	8859	8878	2379
567085	N/A	N/A	ATTGTCTTATTGCTATTGAG	16	8864	8883	2380
567086	N/A	N/A	ACTATTGTCTTATTGCTATT	10	8867	8886	2381
567087	N/A	N/A	TTCACCTATTGTCTTATTGCT	35	8870	8889	2382
567088	N/A	N/A	ACATTCACCTATTGTCTTATT	30	8873	8892	2383
567089	N/A	N/A	TAAACATTCACCTATTGTCTT	58	8876	8895	2384
567090	N/A	N/A	CATTAAACATTCACCTATTGT	28	8879	8898	2385

567091	N/A	N/A	GTTTTTCATTAAACATTCACT	54	8884	8903	2386
567092	N/A	N/A	AAATACTGTTTTTCATTAAAC	34	8891	8910	2387
567093	N/A	N/A	AAAGTATTTATAAAAATACTG	0	8903	8922	2388
567094	N/A	N/A	CCTTTTTTATTAAAGTATTTA	0	8913	8932	2389
567095	N/A	N/A	CAATCCTTTTTTATTAAAGTA	10	8917	8936	2390
567096	N/A	N/A	CTTCATCACAATCCTTTTTTA	52	8925	8944	2391
567097	N/A	N/A	GTTCTTCATCACAATCCTTT	57	8928	8947	2392
567098	N/A	N/A	ATTGTTCTTCATCACAATCC	37	8931	8950	2393
567099	N/A	N/A	TAGATTGTTCTTCATCACAA	31	8934	8953	2394
567100	N/A	N/A	AAATAGATTGTTCTTCATCA	11	8937	8956	2395
567101	N/A	N/A	AACAAATATAAATAGATTGT	0	8946	8965	2396
567102	N/A	N/A	CAAATAACAAATATAAATAG	3	8951	8970	2397
567103	N/A	N/A	TGGAATTAAAAACAAATAAC	3	8963	8982	2398
567104	N/A	N/A	TTATTGGAATTAAAAACAAA	12	8967	8986	2399
567105	N/A	N/A	TTTTTATTGGAATTAAAAAC	17	8970	8989	2400
567106	N/A	N/A	TAATAACTTTTTTCTGTAAT	6	9001	9020	2401
567107	N/A	N/A	GTTCTTAATAACTTTTTTCT	21	9006	9025	2402
567108	N/A	N/A	AAAAGCATGGTTCTTAATAA	0	9015	9034	2403
567109	N/A	N/A	AAATTTAAAAGCATGGTTCT	0	9021	9040	2404
567110	N/A	N/A	AGGAATAAATTTAAAAAATC	0	9046	9065	2405
567111	N/A	N/A	AGACAGGAATAAATTTAAAA	7	9050	9069	2406
567112	N/A	N/A	AAAAGACAGGAATAAATTTA	0	9053	9072	2407
567113	N/A	N/A	CTTTCTTTGTAGAAAAAGAC	29	9066	9085	2408
567114	N/A	N/A	ATGCTTTCTTTGTAGAAAAA	12	9069	9088	2409
567115	N/A	N/A	GCTTAATGTATGCTTTCTTT	67	9078	9097	88
567116	N/A	N/A	TTTGCTTAATGTATGCTTTC	21	9081	9100	2410
567117	N/A	N/A	GTATTTGCTTAATGTATGCT	0	9084	9103	2411
567118	N/A	N/A	TTGGTATTTGCTTAATGTAT	0	9087	9106	2412
567119	N/A	N/A	CCTTTGGTATTTGCTTAATG	35	9090	9109	2413
567120	N/A	N/A	TGGCCTTTGGTATTTGCTTA	0	9093	9112	2414
567121	N/A	N/A	TAAACCTGGCCTTTGGTATT	27	9099	9118	2415
567122	N/A	N/A	ATGTAAACCTGGCCTTTGGT	16	9102	9121	2416
567123	N/A	N/A	CAAATGTAAACCTGGCCTTT	0	9105	9124	2417
567124	N/A	N/A	CTTCAAATGTAAACCTGGCC	25	9108	9127	2418
567125	N/A	N/A	TTTCTTCAAATGTAAACCTG	2	9111	9130	2419
567126	N/A	N/A	TGTCACTTTCTTCAAATGTA	57	9117	9136	2420
567127	N/A	N/A	TAATGTCACTTTCTTCAAAT	6	9120	9139	2421
567128	N/A	N/A	AATAATAATGTCACTTTCTT	3	9125	9144	2422
567129	N/A	N/A	GAGTAATAATAATGTCACTT	18	9129	9148	2423
567130	N/A	N/A	GACTTGAGTAATAATAATGT	1	9134	9153	2424
567131	N/A	N/A	CCTAGAGACTTGAGTAATAA	32	9140	9159	2425

567132	N/A	N/A	ATTCCTAGAGACTTGAGTAA	8	9143	9162	2426
567133	N/A	N/A	AAGTATTCCTAGAGACTTGA	11	9147	9166	2427
567134	N/A	N/A	GTTAAGTATTCCTAGAGACT	61	9150	9169	89
567135	N/A	N/A	TGTGTTAAGTATTCCTAGAG	28	9153	9172	2428
567136	N/A	N/A	AGAGATGTGTTAAGTATTCC	31	9158	9177	2429
567137	N/A	N/A	GTCAAGAGATGTGTTAAGTA	52	9162	9181	2430
567138	N/A	N/A	ACAGTCAAGAGATGTGTAA	22	9165	9184	2431
567139	N/A	N/A	TATACAGTCAAGAGATGTGT	30	9168	9187	2432
567140	N/A	N/A	CCATATACAGTCAAGAGATG	45	9171	9190	2433
567141	N/A	N/A	GTAAGTTGAACTAACTACTG	9	7497	7516	2434
567142	N/A	N/A	TGAGTAAGTTGAACTAACTA	0	7500	7519	2435
567143	N/A	N/A	TAATGAGTAAGTTGAACTAA	2	7503	7522	2436
567144	N/A	N/A	AGGTTAATCTTCCTAATACG	18	7523	7542	2437
567145	N/A	N/A	ATAACCAGGTTAATCTTCCT	34	7529	7548	2438
567146	N/A	N/A	ATGATAACCAGGTTAATCTT	13	7532	7551	2439
567147	N/A	N/A	AACAATGATAACCAGGTAA	7	7536	7555	2440
567148	N/A	N/A	TAAAACAATGATAACCAGGT	45	7539	7558	2441
567149	N/A	N/A	GTATAAAACAATGATAACCA	26	7542	7561	2442
567150	N/A	N/A	CGAATACTCATATATATTTC	25	7572	7591	2443
567151	N/A	N/A	ATACGAATACTCATATATAT	30	7575	7594	2444
567152	N/A	N/A	TTTATACGAATACTCATATA	32	7578	7597	2445
567153	N/A	N/A	ATATTTATACGAATACTCAT	25	7581	7600	2446
567154	N/A	N/A	GTATTATATTTATACGAATA	0	7586	7605	2447
567155	N/A	N/A	AAAAGTATTATATTTATACG	0	7590	7609	2448
567156	N/A	N/A	GGTAAAAGTATTATATTTAT	0	7593	7612	2449
567157	N/A	N/A	ACAAGGTAAAAGTATTATAT	10	7597	7616	2450
567158	N/A	N/A	TAAACAAGGTAAAAGTATTA	11	7600	7619	2451
567159	N/A	N/A	ACATAAACAAGGTAAAAGTA	3	7603	7622	2452
567160	N/A	N/A	TTGAGTAAATACATAAACAA	12	7613	7632	2453
567161	N/A	N/A	GAGAATATTGAGTAAATACA	4	7620	7639	2454
567162	N/A	N/A	AAGGAGAATATTGAGTAAAT	8	7623	7642	2455
567163	N/A	N/A	GAAAAGGAGAATATTGAGTA	3	7626	7645	2456
567164	N/A	N/A	GAGGAAAAGGAGAATATTGA	19	7629	7648	2457
567165	N/A	N/A	TTAGAGGAAAAGGAGAATAT	41	7632	7651	2458
567166	N/A	N/A	ATTATTTTAGAGGAAAAGGA	30	7638	7657	2459
567167	N/A	N/A	CAGATTATTTTAGAGGAAAA	9	7641	7660	2460
567168	N/A	N/A	CTTCAGATTATTTTAGAGGA	24	7644	7663	2461
567169	N/A	N/A	TAGTCACTTCAGATTATTTT	38	7650	7669	2462
567170	N/A	N/A	TAATAGTCACTTCAGATTAT	13	7653	7672	2463
567171	N/A	N/A	TGATAATAGTCACTTCAGAT	39	7656	7675	2464
567172	N/A	N/A	TATTGATAATAGTCACTTCA	41	7659	7678	2465

567173	N/A	N/A	ACTTATTGATAATAGTCACT	29	7662	7681	2466
567174	N/A	N/A	TAAACTTATTGATAATAGTC	14	7665	7684	2467
567175	N/A	N/A	TAGTAAACTTATTGATAATA	31	7668	7687	2468
567176	N/A	N/A	GCATAGTAAACTTATTGATA	23	7671	7690	2469
567177	N/A	N/A	TTGGCATAGTAAACTTATTG	21	7674	7693	2470
567178	N/A	N/A	ATTTTGGCATAGTAAACTTA	8	7677	7696	2471
567179	N/A	N/A	TGAATTTTGGCATAGTAAAC	5	7680	7699	2472
567180	N/A	N/A	TTAATGAATTTTGGCATAGT	0	7684	7703	2473
567181	N/A	N/A	CAATTAATGAATTTTGGCAT	39	7687	7706	2474
567182	N/A	N/A	AAAGGCAATTAATGAATTTT	12	7692	7711	2475
567183	N/A	N/A	GTGAAAGGCAATTAATGAAT	28	7695	7714	2476
567184	N/A	N/A	TTAAGTGAAAGGCAATTAAT	7	7699	7718	2477
567185	N/A	N/A	AAGTTAAGTGAAAGGCAATT	25	7702	7721	2478
567186	N/A	N/A	CCAAAAGTTAAGTGAAAGGC	50	7706	7725	2479
567187	N/A	N/A	GTCCCAAAAAGTTAAGTGAAA	30	7709	7728	2480
567188	N/A	N/A	ATGGTCCCAAAAAGTTAAGTG	39	7712	7731	2481
567189	N/A	N/A	ATTATGGTCCCAAAAAGTTAA	19	7715	7734	2482
567190	N/A	N/A	TTTATTATGGTCCCAAAAAGT	33	7718	7737	2483
567191	N/A	N/A	TTATTATTTATTATGGTCCC	50	7724	7743	2484
567192	N/A	N/A	ATGGCAATACATTTTATTAT	13	7737	7756	2485
567193	N/A	N/A	GTTATGGCAATACATTTTAT	39	7740	7759	2486
567194	N/A	N/A	TAATGTTATGGCAATACATT	0	7744	7763	2487
567195	N/A	N/A	TATTAATGTTATGGCAATAC	22	7747	7766	2488
567196	N/A	N/A	GTTTATTAATGTTATGGCAA	28	7750	7769	2489
567197	N/A	N/A	GTAGTTTATTAATGTTATGG	20	7753	7772	2490
567198	N/A	N/A	AAGGTAGTTTATTAATGTTA	27	7756	7775	2491
567199	N/A	N/A	TGTAAGGTAGTTTATTAATG	0	7759	7778	2492
567200	N/A	N/A	TTTTGTAAGGTAGTTTATTA	0	7762	7781	2493
567201	N/A	N/A	TGGTTTTGTAAGGTAGTTTA	18	7765	7784	2494
567202	N/A	N/A	TGGTGGTTTTTGTAAGGTAGT	0	7768	7787	2495
567203	N/A	N/A	AATTGGTGGTTTTGTAAGGT	11	7771	7790	2496
567204	N/A	N/A	TTTAATTGGTGGTTTTGTAA	0	7774	7793	2497
567205	N/A	N/A	TTGATTTTAATTGGTGGTTT	19	7779	7798	2498
567206	N/A	N/A	TGTTTGATTTTAATTGGTGG	26	7782	7801	2499
567207	N/A	N/A	ATGTAAATAACACTTTTTTG	1	7804	7823	2500
567208	N/A	N/A	CAGATGTAAATAACACTTTT	1	7807	7826	2501
567209	N/A	N/A	TGACAGATGTAAATAACACT	21	7810	7829	2502
567210	N/A	N/A	ATGTTGACAGATGTAAATAA	0	7814	7833	2503
567211	N/A	N/A	TTTATGTTGACAGATGTAAA	0	7817	7836	2504
567212	N/A	N/A	AGATTTATGTTGACAGATGT	0	7820	7839	2505
567213	N/A	N/A	AGTAGATTTATGTTGACAGA	19	7823	7842	2506

567214	N/A	N/A	TTTAGTAGATTTATGTTGAC	4	7826	7845	2507
567215	N/A	N/A	ATTTT TAGTAGATTTATGTT	0	7829	7848	2508
567216	N/A	N/A	CATGTATTTT TAGTAGATTT	5	7834	7853	2509
567217	N/A	N/A	GAAATCATGTATTTT TAGTA	0	7839	7858	2510
567218	N/A	N/A	ATTGTATTTGATGGATATCT	43	6875	6894	2511
567219	N/A	N/A	GATACATTGTATTTGATGGA	20	6880	6899	2512
567220	N/A	N/A	TAGGTTGATACATTGTATTT	18	6886	6905	2513
567221	N/A	N/A	CAGTTTAGGTTGATACATTG	18	6891	6910	2514
567222	N/A	N/A	GCATCCAGTTTAGGTTGATA	31	6896	6915	2515
567223	N/A	N/A	CCCCAGCATCCAGTTTAGGT	14	6901	6920	2516
567224	N/A	N/A	AAGAACCCCAGCATCCAGTT	41	6906	6925	2517
567225	N/A	N/A	GTGTAAAAAGAACCCCAGCA	0	6913	6932	2518
567226	N/A	N/A	ATAGGGTGTA AAAAGAACCC	13	6918	6937	2519
567227	N/A	N/A	CTTTTATAGGGTGTA AAAAG	0	6923	6942	2520
567228	N/A	N/A	TATGTCTTTTATAGGGTGTA	26	6928	6947	2521
567229	N/A	N/A	TTAGGTATGTCTTTTATAGG	0	6933	6952	2522
567230	N/A	N/A	TTGTCTTAGGTATGTCTTTT	30	6938	6957	2523
567231	N/A	N/A	CTCTGATTGTCTTAGGTATG	27	6944	6963	2524
567232	N/A	N/A	TATTTCTCTGATTGTCTTAG	21	6949	6968	2525
567233	N/A	N/A	TCCATATTTGTATTTCTCTG	61	6959	6978	90
567234	N/A	N/A	TCAAGTCCATATTTGTATTT	20	6964	6983	2526
567235	N/A	N/A	AATAATCAAGTCCATATTTG	0	6969	6988	2527
567236	N/A	N/A	TTATCTAATAATCAAGTCCA	0	6975	6994	2528
567237	N/A	N/A	CTATATTATCTAATAATCAA	12	6980	6999	2529
567238	N/A	N/A	TAAACCTTCTATATTATCTA	12	6988	7007	2530
567239	N/A	N/A	AATTAATAAACCTTCTATAT	0	6994	7013	2531
567240	N/A	N/A	TAAGTACAGGTTGGACACTG	0	9504	9523	2532
567241	N/A	N/A	GTTATTAAGTACAGGTTGGA	2	9509	9528	2533
567242	N/A	N/A	TGTGAGTTATTAAGTACAGG	0	9514	9533	2534
567243	N/A	N/A	AAATCTGTGAGTTATTAAGT	0	9519	9538	2535
567244	N/A	N/A	GTTTTAAAAATCTGTGAGTT	19	9526	9545	2536
567245	N/A	N/A	CAAAATTCTCCTGAAAAGAA	20	9548	9567	2537
567246	N/A	N/A	CCCAACCAAAATTCTCCTGA	48	9554	9573	2538
567247	N/A	N/A	ACCTGAATAACCCTCTGGAC	21	9807	9826	2539
567248	N/A	N/A	AAGATACCTGAATAACCCTC	30	9812	9831	2540
567249	N/A	N/A	AGAAAAAGATACCTGAATAA	0	9817	9836	2541
567250	N/A	N/A	TGGTATCAGAAAAAGATACC	0	9824	9843	2542
567251	N/A	N/A	AGTATTGGTATCAGAAAAAG	0	9829	9848	2543
567252	N/A	N/A	AATAAAGTATTGGTATCAGA	10	9834	9853	2544
567253	N/A	N/A	ATGAAAATAAAGTATTGGTA	3	9839	9858	2545
567254	N/A	N/A	AGATACTTTGAAGATATGAA	0	9854	9873	2546

567255	N/A	N/A	TGGGAAGATACTTTGAAGAT	0	9859	9878	2547
567256	N/A	N/A	CTAATAATGTGGGAAGATAC	0	9868	9887	2548
567257	N/A	N/A	CATTGCAGATAATAGCTAAT	0	9883	9902	2549
567258	N/A	N/A	AAGTTGTCATTGCAGATAAT	0	9890	9909	2550
567259	N/A	N/A	TTTAAAAGTTGTCATTGCA	7	9896	9915	2551
567260	N/A	N/A	ATTCGGATTTTAAAAGTTG	5	9904	9923	2552
567261	N/A	N/A	TTATTTGGGATTCGGATTTT	15	9913	9932	2553
567262	N/A	N/A	TTATAGTTAAGAGGTTTTTCG	27	9949	9968	2554
567263	N/A	N/A	TTTCATTATAGTTAAGAGGT	12	9954	9973	2555
567264	N/A	N/A	GAACACTTTCATTATAGTTA	13	9960	9979	2556
567265	N/A	N/A	GAACTAGAATGAACACTTTC	28	9970	9989	2557
567266	N/A	N/A	TGATTGAACTAGAATGAACA	23	9975	9994	2558
567267	N/A	N/A	ATACCTGATTGAACTAGAAT	9	9980	9999	2559
567268	N/A	N/A	GTAAAATACCTGATTGAACT	6	9985	10004	2560
567269	N/A	N/A	TAGAGGTAAAATACCTGATT	16	9990	10009	2561
567270	N/A	N/A	AAGATTAGAGGTAAAATACC	0	9995	10014	2562
567271	N/A	N/A	TGAGGAAGATTAGAGGTAAA	6	10000	10019	2563
567272	N/A	N/A	GAAAATCTGAGGAAGATTAG	0	10007	10026	2564
567273	N/A	N/A	AAATAGAAAATCTGAGGAAG	0	10012	10031	2565
567274	N/A	N/A	ATCTATACACTACCAAAAAA	0	10029	10048	2566
567275	N/A	N/A	AAATAATCTATACACTACCA	19	10034	10053	2567
567276	N/A	N/A	AAATAATCTGTATAAATAAT	3	10047	10066	2568
567277	N/A	N/A	CCCAATTTTAAATAATCTGT	24	10056	10075	2569
567278	N/A	N/A	TAAGTCCCAATTTTAAATAA	0	10061	10080	2570
567279	N/A	N/A	TCTGTATAAGTCCCAATTTT	15	10067	10086	2571
567280	N/A	N/A	AATAATCTGTATAAGTCCCA	47	10072	10091	2572
567281	N/A	N/A	AGTTTTAAATAATCTGTATA	0	10079	10098	2573
567282	N/A	N/A	ATCCCAGTTTTTAAATAATCT	6	10084	10103	2574
567283	N/A	N/A	CATGTATCCCAGTTTTTAAAT	6	10089	10108	2575
567284	N/A	N/A	TAGATGCATGTATCCCAGTT	41	10095	10114	2576
567285	N/A	N/A	TGTTTTAGATGCATGTATCC	4	10100	10119	2577
567286	N/A	N/A	TACAGTGTTTTAGATGCATG	25	10105	10124	2578
567287	N/A	N/A	AATATTACAGTGTTTTAGAT	0	10110	10129	2579
567288	N/A	N/A	CTTATAAATATTACAGTGTT	2	10116	10135	2580
567289	N/A	N/A	CTTCCTTTCTTATAAATATT	12	10124	10143	2581
567290	N/A	N/A	TTTATCTTCCTTTCTTATAA	0	10129	10148	2582
567291	N/A	N/A	CGTAAGTTTATCTTCCTTTC	61	10135	10154	91
567292	N/A	N/A	TTCCCCGTAAGTTTATCTTC	22	10140	10159	2583
567293	N/A	N/A	TGTATTTCCCCGTAAGTTTA	0	10145	10164	2584
567294	N/A	N/A	GTTACTGTATTTCCCCGTAA	43	10150	10169	2585
544120	707	726	AGTTCTTGGTGCTCTTGGCT	80	6720	6739	15

337487	804	823	CACTTGTATGTTACCTCTG	80	7389	7408	28
--------	-----	-----	---------------------	----	------	------	----

Table 13

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
563780	N/A	N/A	TGTTTTCTTCTGGAAGCAGA	10	3100	3119	2586
568085	N/A	N/A	CAGACCTAGACTTCTTAAC	8	3084	3103	2587
568086	N/A	N/A	AGCAGACCTAGACTTCTTAA	6	3086	3105	2588
568087	N/A	N/A	TTTTCTTCTGGAAGCAGACC	0	3098	3117	2589
568088	N/A	N/A	AAACATATATACATGCTTGT	52	11323	11342	2590
568089	N/A	N/A	TTAAACATATATACATGCTT	39	11325	11344	2591
568090	N/A	N/A	GTTTATTGAATTTTAAACAT	0	11337	11356	2592
568091	N/A	N/A	TTGTTTATTGAATTTTAAAC	9	11339	11358	2593
568092	N/A	N/A	CTTTGTTTATTGAATTTTAA	0	11341	11360	2594
568093	N/A	N/A	GTCTTTGTTTATTGAATTTT	28	11343	11362	2595
568094	N/A	N/A	GGGTCTTTGTTTATTGAATT	0	11345	11364	2596
568095	N/A	N/A	CTGGGTCTTTGTTTATTGAA	11	11347	11366	2597
568096	N/A	N/A	GACTGGGTCTTTGTTTATTG	35	11349	11368	2598
568097	N/A	N/A	TTTCTATAATTTAGGGACTG	12	11364	11383	2599
568098	N/A	N/A	AATTTCTATAATTTAGGGAC	0	11366	11385	2600
568099	N/A	N/A	TAAATTTCTATAATTTAGGG	5	11368	11387	2601
568100	N/A	N/A	CAAGAATAATTTAAATTTCT	38	11379	11398	2602
568101	N/A	N/A	GATAAACATGCAAGAATAAT	1	11389	11408	2603
568102	N/A	N/A	TCGATAAACATGCAAGAATA	51	11391	11410	2604
568103	N/A	N/A	TGTCGATAAACATGCAAGAA	37	11393	11412	2605
568104	N/A	N/A	GATGTCGATAAACATGCAAG	57	11395	11414	2606
568105	N/A	N/A	GTGATGTCGATAAACATGCA	61	11397	11416	2607
568106	N/A	N/A	TTGTGATGTCGATAAACATG	57	11399	11418	2608
568107	N/A	N/A	TGTTGTGATGTCGATAAACA	47	11401	11420	2609
568108	N/A	N/A	TCTGTTGTGATGTCGATAAA	53	11403	11422	2610
568109	N/A	N/A	GATCTGTTGTGATGTCGATA	36	11405	11424	2611
568110	N/A	N/A	GGGATCTGTTGTGATGTCGA	41	11407	11426	2612
568111	N/A	N/A	TAGGGATCTGTTGTGATGTC	43	11409	11428	2613
568112	N/A	N/A	TTAGGGATCTGTTGTGATG	18	11411	11430	2614
568113	N/A	N/A	GATTTAGGGATCTGTTGTGA	41	11413	11432	2615
568114	N/A	N/A	ATCTAATCTTTAGGGATTTA	37	11435	11454	2616
568115	N/A	N/A	TTTGTATCTAATCTTTAGGG	28	11440	11459	2617
568116	N/A	N/A	AATTTGTATCTAATCTTTAG	0	11442	11461	2618
568117	N/A	N/A	GTGGTAAAAAATTTGTATCT	13	11451	11470	2619

568118	N/A	N/A	CTGTGGTAAAAAATTTGTAT	5	11453	11472	2620
568119	N/A	N/A	TACTGTGGTAAAAAATTTGT	10	11455	11474	2621
568120	N/A	N/A	GATACTGTGGTAAAAAATTT	17	11457	11476	2622
568121	N/A	N/A	AGTGATACTGTGGTAAAAAA	38	11460	11479	2623
568122	N/A	N/A	CAAGTGATACTGTGGTAAAA	58	11462	11481	2624
568123	N/A	N/A	GACAAGTGATACTGTGGTAA	52	11464	11483	2625
568124	N/A	N/A	CTGACAAGTGATACTGTGGT	62	11466	11485	2626
568125	N/A	N/A	TTCTGACAAGTGATACTGTG	27	11468	11487	2627
568126	N/A	N/A	AATTCTGACAAGTGATACTG	33	11470	11489	2628
568127	N/A	N/A	ATAAATTCTGACAAGTGATA	38	11473	11492	2629
568128	N/A	N/A	CTGGCAGTTTTTAAAAAATCA	28	11502	11521	2630
568129	N/A	N/A	TTCTTACTGGCAGTTTTTAAA	56	11508	11527	2631
568130	N/A	N/A	ATTTCTTACTGGCAGTTTTTA	47	11510	11529	2632
568131	N/A	N/A	AAATTTCTTACTGGCAGTTT	53	11512	11531	2633
568132	N/A	N/A	TTTAAAATTTCTTACTGGCA	46	11516	11535	2634
568133	N/A	N/A	TTAATTTAAAATTTCTTACT	9	11520	11539	2635
568134	N/A	N/A	CAAATGGGTTTAATTTAAAA	1	11529	11548	2636
568135	N/A	N/A	AACAAATGGGTTTAATTTAA	11	11531	11550	2637
568136	N/A	N/A	TTAACAAATGGGTTTAATTT	12	11533	11552	2638
568137	N/A	N/A	CTTTAACAAATGGGTTTAAT	27	11535	11554	2639
568138	N/A	N/A	TCCTTTAACAAATGGGTTTA	52	11537	11556	2640
568139	N/A	N/A	CTATATCCTTTAACAAATGG	24	11542	11561	2641
568140	N/A	N/A	GGGCACTATATCCTTTAACA	45	11547	11566	2642
568141	N/A	N/A	TTGGGCACTATATCCTTTAA	20	11549	11568	2643
568142	N/A	N/A	TATAACTTGGGCACTATATC	27	11555	11574	2644
568143	N/A	N/A	CATATAACTTGGGCACTATA	40	11557	11576	2645
568144	N/A	N/A	ACCATATAACTTGGGCACTA	69	11559	11578	103
568145	N/A	N/A	TCACCATATAACTTGGGCAC	60	11561	11580	2646
568146	N/A	N/A	GGTCACCATATAACTTGGGC	73	11563	11582	104
568147	N/A	N/A	TAGGTCACCATATAACTTGG	51	11565	11584	2647
568148	N/A	N/A	GGTAGGTCACCATATAACTT	57	11567	11586	2648
568149	N/A	N/A	AAGGTAGGTCACCATATAAC	52	11569	11588	2649
568150	N/A	N/A	CAAAGGTAGGTCACCATATA	28	11571	11590	2650
568151	N/A	N/A	GACAAAGGTAGGTCACCATA	67	11573	11592	105
568152	N/A	N/A	GTATTGACAAAGGTAGGTCA	55	11578	11597	2651
568153	N/A	N/A	AAGTATTGACAAAGGTAGGT	36	11580	11599	2652
568154	N/A	N/A	CTAAGTATTGACAAAGGTAG	24	11582	11601	2653
568155	N/A	N/A	TGCTAAGTATTGACAAAGGT	49	11584	11603	2654
568156	N/A	N/A	AATGCTAAGTATTGACAAAG	10	11586	11605	2655
568157	N/A	N/A	CATAATGCTAAGTATTGACA	19	11589	11608	2656
568158	N/A	N/A	TACATAATGCTAAGTATTGA	4	11591	11610	2657

568159	N/A	N/A	AATACATAATGCTAAGTATT	1	11593	11612	2658
568160	N/A	N/A	GAAATACATAATGCTAAGTA	23	11595	11614	2659
568161	N/A	N/A	TTTGAAATACATAATGCTAA	8	11598	11617	2660
568162	N/A	N/A	GGATAATTTGAAATACATAA	16	11604	11623	2661
568163	N/A	N/A	TTGGATAATTTGAAATACAT	0	11606	11625	2662
568164	N/A	N/A	TATTGGATAATTTGAAATAC	0	11608	11627	2663
568165	N/A	N/A	ATCCAGTTAAAGCTTGTAAG	46	4466	4485	2664
568166	N/A	N/A	TCATGATCCAGTTAAAGCTT	32	4471	4490	2665
568167	N/A	N/A	TTTACTCATGATCCAGTTAA	24	4476	4495	2666
568168	N/A	N/A	GATAATTTTACTCATGATCC	53	4482	4501	2667
568169	N/A	N/A	GATGTGATAATTTTACTCAT	27	4487	4506	2668
568170	N/A	N/A	ATGCTGATGTGATAATTTTA	42	4492	4511	2669
568171	N/A	N/A	CAGTTATGCTGATGTGATAA	0	4497	4516	2670
568172	N/A	N/A	TTTAACAGTTATGCTGATGT	17	4502	4521	2671
568173	N/A	N/A	GCAATTTTAACAGTTATGCT	11	4507	4526	2672
568174	N/A	N/A	AGAGCCTGCAATTTTAACAG	25	4514	4533	2673
568175	N/A	N/A	GCTTCAGAGCCTGCAATTTT	47	4519	4538	2674
568176	N/A	N/A	TATTAGCTTCAGAGCCTGCA	48	4524	4543	2675
568177	N/A	N/A	TAGTTTATTAGCTTCAGAGC	20	4529	4548	2676
568178	N/A	N/A	GCAGGTAGTTTATTAGCTTC	39	4534	4553	2677
568179	N/A	N/A	TAAATGCAGGTAGTTTATTA	0	4539	4558	2678
568180	N/A	N/A	ATGGTTTAAATGCAGGTAGT	20	4545	4564	2679
568181	N/A	N/A	GAGCCATGGTTTAAATGCAG	33	4550	4569	2680
568182	N/A	N/A	TTTTAGAGCCATGGTTTAAA	40	4555	4574	2681
568183	N/A	N/A	CAAAGTTTTAGAGCCATGGT	54	4560	4579	2682
568184	N/A	N/A	TCACACAAAGTTTTAGAGCC	61	4565	4584	2683
568185	N/A	N/A	CAAGGTCACACAAAGTTTTA	17	4570	4589	2684
568186	N/A	N/A	GGGTGAAGTAATTTATTCAA	0	4587	4606	2685
568187	N/A	N/A	GTGAGGAAACTGAGAGATAA	12	4609	4628	2686
568188	N/A	N/A	TGTAGTATATGTGAGGAAAC	38	4619	4638	2687
568189	N/A	N/A	ATCTTTGTAGTATATGTGAG	30	4624	4643	2688
568190	N/A	N/A	TTATTATCTTTGTAGTATAT	19	4629	4648	2689
568191	N/A	N/A	TTCTGTTATTATCTTTGTAG	48	4634	4653	2690
568192	N/A	N/A	ATAAGTTCTGTTATTATCTT	16	4639	4658	2691
568193	N/A	N/A	ATCCTATAAGTTCTGTTATT	22	4644	4663	2692
568194	N/A	N/A	CAATAATCCTATAAGTTCTG	0	4649	4668	2693
568195	N/A	N/A	TAAGATGACATTGGCTGCTA	49	4689	4708	2694
568196	N/A	N/A	TTTAGTAAGATGACATTGGC	32	4694	4713	2695
568197	N/A	N/A	TTGAATTTTAGTAAGATGAC	19	4700	4719	2696
568198	N/A	N/A	CTAATTTGAATTTTAGTAAG	34	4705	4724	2697
568199	N/A	N/A	CATGATCTAATTTGAATTTT	29	4711	4730	2698

568200	N/A	N/A	CAAAGAGAAACATGATCTAA	27	4721	4740	2699
568201	N/A	N/A	GTTTTGAGCAAAGAGAAACA	36	4729	4748	2700
568202	N/A	N/A	GTGTGGTTTTGAGCAAAGAG	3	4734	4753	2701
568203	N/A	N/A	AGCTATTGTGTGGTTTTGAG	13	4741	4760	2702
568204	N/A	N/A	TGAAATGGAAAGCTATTGTG	15	4751	4770	2703
568205	N/A	N/A	TATGAGTGAAATGGAAAGCT	27	4757	4776	2704
568206	N/A	N/A	GCCAATATGAGTGAAATGGA	62	4762	4781	106
568207	N/A	N/A	AAAGAGCCAATATGAGTGAA	25	4767	4786	2705
568208	N/A	N/A	TTGGTCTAAAGAGCCAATAT	42	4774	4793	2706
568209	N/A	N/A	GGTAATCTTGGTCTAAAGAG	29	4781	4800	2707
568210	N/A	N/A	GTGAGATGACGAAGGGTTGG	0	4800	4819	2708
568211	N/A	N/A	AGTCAGTGAGATGACGAAGG	5	4805	4824	2709
568212	N/A	N/A	GGTGAAGTCAGTGAGATGAC	12	4810	4829	2710
568213	N/A	N/A	GTAGAGGAGGTGAAGTCAGT	13	4818	4837	2711
568214	N/A	N/A	AACTAGAGTAGAGGAGGTGA	20	4825	4844	2712
568215	N/A	N/A	AGAATAACTAGAGTAGAGGA	33	4830	4849	2713
568216	N/A	N/A	CGGTCAGAATAACTAGAGTA	39	4835	4854	2714
568217	N/A	N/A	TAAAGCGGTCAGAATAACTA	29	4840	4859	2715
568218	N/A	N/A	ACTGGTAAAGCGGTCAGAAT	17	4845	4864	2716
568219	N/A	N/A	TGAATACTGGTAAAGCGGTC	37	4850	4869	2717
568220	N/A	N/A	TGTGTTTGAATACTGGTAAA	21	4856	4875	2718
568221	N/A	N/A	AGTATGTTTGATGTGTTTGA	25	4867	4886	2719
568222	N/A	N/A	GTGGCAGTATGTTTGATGTG	15	4872	4891	2720
568223	N/A	N/A	TTGAGGTGGCAGTATGTTTG	14	4877	4896	2721
568224	N/A	N/A	AGGCTTTGAGGTGGCAGTAT	33	4882	4901	2722
568225	N/A	N/A	GGCAAAGGCTTTGAGGTGGC	27	4887	4906	2723
568226	N/A	N/A	AACAAGGGCAAAGGCTTTGA	24	4893	4912	2724
568227	N/A	N/A	TAGAGGAAACAACAAGGGCA	24	4903	4922	2725
568228	N/A	N/A	CCAGTTAGAGGAAACAACAA	4	4908	4927	2726
568229	N/A	N/A	GATACCAGGGCAGAAGAGCG	24	4930	4949	2727
568230	N/A	N/A	AAATCAGAGAGTGGGCCACG	24	4952	4971	2728
568231	N/A	N/A	CCTAAGGGAAATCAGAGAGT	19	4960	4979	2729
568232	N/A	N/A	ACGACCCTAAGGGAAATCAG	30	4965	4984	2730
568233	N/A	N/A	TGATAACGACCCTAAGGGAA	0	4970	4989	2731
568234	N/A	N/A	TTTTGTTTGATAACGACCCT	22	4977	4996	2732
568235	N/A	N/A	GTCTTCATTGGGAATTTTTT	37	4993	5012	2733
568236	N/A	N/A	TGTAAGTCTTCATTGGGAAT	23	4998	5017	2734
568237	N/A	N/A	GACCTTGTAAGTCTTCATTG	52	5003	5022	2735
568238	N/A	N/A	TAAGTGACCTTGTAAGTCTT	36	5008	5027	2736
568239	N/A	N/A	TTGGTTAAGTGACCTTGTA	11	5013	5032	2737
568240	N/A	N/A	TGATTTTTGGTTAAGTGACC	12	5019	5038	2738

568241	N/A	N/A	GGTTGTGATTTTTGGTTAAG	11	5024	5043	2739
568242	N/A	N/A	CAGGCGGTTGTGATTTTTGG	41	5029	5048	2740
568243	N/A	N/A	GGGACCAGGCGGTTGTGATT	22	5034	5053	2741
568244	N/A	N/A	CTAAGGAAGTAGAAGTTTTC	42	5060	5079	2742
568245	N/A	N/A	AGTAGCTAAGGAAGTAGAAG	11	5065	5084	2743
568246	N/A	N/A	CAGGAGAAAAGTAGCTAAGG	36	5074	5093	2744
568247	N/A	N/A	GTGTGCAGGAGAAAAGTAGC	14	5079	5098	2745
568248	N/A	N/A	TAAAGGTGAGTGTGCAGGAG	7	5088	5107	2746
568249	N/A	N/A	ATGTTAAATAAAGGTGAGTG	8	5096	5115	2747
568250	N/A	N/A	ATGTTATGTTAAATAAAGGT	27	5101	5120	2748
568251	N/A	N/A	AATTTATGTTATGTTAAATA	27	5106	5125	2749
568252	N/A	N/A	TAACTAAAATTTATGTTATG	28	5113	5132	2750
568253	N/A	N/A	GATAAATAACTAAAATTTAT	32	5119	5138	2751
568254	N/A	N/A	TTTAGTGCAGGAATAGAAGA	33	5139	5158	2752
568255	N/A	N/A	AATCCCTGTATTCACAGAGC	68	5165	5184	2753
568256	N/A	N/A	GAAAAAATCCCTGTATTCAC	0	5170	5189	2754
568257	N/A	N/A	TAATGGAAAAAATCCCTGTA	8	5175	5194	2755
568258	N/A	N/A	AAATATGAAGATAATGGAAA	26	5186	5205	2756
568259	N/A	N/A	ATAATGGAAAATATGAAGAT	18	5194	5213	2757
568260	N/A	N/A	TATACAAATAATGGAAAATA	30	5201	5220	2758
568261	N/A	N/A	TTCTGGAGTATATACAAATA	45	5211	5230	2759
568262	N/A	N/A	ATTCTATATTCTGGAGTATA	40	5219	5238	2760
568263	N/A	N/A	CCATACAGTATTCTATATTC	57	5228	5247	2761
568264	N/A	N/A	CTGTGTGCCATACAGTATTC	28	5235	5254	2762
568265	N/A	N/A	GCCTACTGTGTGCCATACAG	60	5240	5259	2763
568266	N/A	N/A	AGAAATGCCTACTGTGTGCC	42	5246	5265	2764
568267	N/A	N/A	TCAACAGAAATGCCTACTGT	52	5251	5270	2765
568268	N/A	N/A	ATTAATTCAACAGAAATGCC	46	5257	5276	2766
568269	N/A	N/A	GACATTACATTTATTAATTC	32	5269	5288	2767
568270	N/A	N/A	GTGAATATGACATTACATTT	32	5277	5296	2768
568271	N/A	N/A	CTTCTGTGTGAATATGACAT	50	5284	5303	2769
568272	N/A	N/A	ACACGCTTCTGTGTGAATAT	43	5289	5308	2770
568273	N/A	N/A	ATAGCACACGCTTCTGTGTG	31	5294	5313	2771
568274	N/A	N/A	TAATCATAGCACACGCTTCT	40	5299	5318	2772
568275	N/A	N/A	AATAATAATCATAGCACACG	20	5304	5323	2773
568276	N/A	N/A	CCAAGTAATAATAATCATAG	35	5310	5329	2774
568277	N/A	N/A	CTAGTAATCCAAGTAATAAT	38	5318	5337	2775
568278	N/A	N/A	TATTTCTAGTAATCCAAGTA	39	5323	5342	2776
568279	N/A	N/A	CACACTATTTCTAGTAATCC	51	5328	5347	2777
568280	N/A	N/A	TTATGAGGCACACTATTTCT	25	5336	5355	2778
568281	N/A	N/A	TTTAATTATGAGGCACACTA	35	5341	5360	2779

568282	N/A	N/A	GTTGACCTTTAATTATGAGG	63	5348	5367	2780
568283	N/A	N/A	TTACATTGTTGAATGTTGAC	45	5362	5381	2781
568284	N/A	N/A	ATTAATTACATTGTTGAATG	31	5367	5386	2782
568285	N/A	N/A	TGTAGATTAATTACATTGTT	49	5372	5391	2783
568286	N/A	N/A	TACATTGTAGATTAATTACA	43	5377	5396	2784
568287	N/A	N/A	AGATGTTTACATTGTAGATT	28	5384	5403	2785
568288	N/A	N/A	TTCACCAGATGTTTACATTG	36	5390	5409	2786
568289	N/A	N/A	GTCACCTCACCAGATGTTTA	65	5395	5414	2787
568290	N/A	N/A	CCTCTGTCACCTCACCAGAT	67	5400	5419	2788
568291	N/A	N/A	GCTTCCCTCTGTCACCTCAC	70	5405	5424	2789
568292	N/A	N/A	CAAGTGCTTCCCTCTGTCAC	33	5410	5429	2790
568293	N/A	N/A	TTTCTAAACAAGTGCTTCCC	70	5418	5437	107
568294	N/A	N/A	GCTTTTTTCTAAACAAGTGC	45	5423	5442	2791
568295	N/A	N/A	ACATAGCTTTTTTCTAAACA	9	5428	5447	2792
568296	N/A	N/A	TTCTGACATAGCTTTTTTCT	23	5433	5452	2793
568297	N/A	N/A	ATGGATTCTGACATAGCTTT	46	5438	5457	2794
568298	N/A	N/A	AATACATGGATTCTGACATA	37	5443	5462	2795
568299	N/A	N/A	ATTAGAATACATGGATTCTG	57	5448	5467	2796
568300	N/A	N/A	CTGCATATTAGAATACATGG	75	5454	5473	108
568301	N/A	N/A	TTGTACTGCATATTAGAATA	53	5459	5478	2797
568302	N/A	N/A	AACTATTGTACTGCATATTA	25	5464	5483	2798
568303	N/A	N/A	TTTTAACTATTGTACTGCA	25	5469	5488	2799
568304	N/A	N/A	TGAGAGTATTATTAATATTT	8	5487	5506	2800
568305	N/A	N/A	GCTGTTTGAGAGTATTATTA	50	5493	5512	2801
568306	N/A	N/A	GAATAGCTGTTTGAGAGTAT	38	5498	5517	2802
568307	N/A	N/A	CCTCTTGAATAGCTGTTTGA	55	5504	5523	2803
568308	N/A	N/A	TGAATCCTCTTGAATAGCTG	55	5509	5528	2804
568309	N/A	N/A	TTTTTTGAATCCTCTTGAAT	46	5514	5533	2805
568310	N/A	N/A	TTATGTTTTTTGAATCCTCT	36	5519	5538	2806
568311	N/A	N/A	GTTTATATTATGTTTTTTGA	6	5526	5545	2807
568312	N/A	N/A	TCTGAGTTTATATTATGTTT	29	5531	5550	2808
568313	N/A	N/A	CAGTTTCTCTGAGTTTATAT	28	5538	5557	2809
568314	N/A	N/A	GTTTACCAGTTTCTCTGAGT	44	5544	5563	2810
568315	N/A	N/A	ATTTTGTTTACCAGTTTCTC	58	5549	5568	2811
568316	N/A	N/A	AAATGATTTTGTTTACCAGT	29	5554	5573	2812
568317	N/A	N/A	CTCTTGAAAATGATTTTGTT	22	5561	5580	2813
568318	N/A	N/A	TATATCTCTTGAAAATGATT	5	5566	5585	2814
568319	N/A	N/A	CAGGTTGGCAAGTTTGTTTG	27	6175	6194	2815
568320	N/A	N/A	GTTGGCAGGTTGGCAAGTTT	44	6180	6199	2816
568321	N/A	N/A	ATATCTGTAGATGTTGGCAG	59	6192	6211	2817
568322	N/A	N/A	TAAACATATCTGTAGATGTT	18	6197	6216	2818

568323	N/A	N/A	ACCTGTAAACATATCTGTAG	57	6202	6221	2819
568324	N/A	N/A	TTTTGACCTGTAAACATATC	23	6207	6226	2820
568325	N/A	N/A	ATAATTTTTTGACCTGTAAAC	7	6212	6231	2821
568326	N/A	N/A	TAATTTGATAATTTTTTGACC	7	6219	6238	2822
568327	N/A	N/A	TTCTTGATAATTTGATAATT	8	6226	6245	2823
568328	N/A	N/A	ACCAGGCTTTCTTGATAATT	55	6234	6253	2824
568329	N/A	N/A	TTTGAACCAGGCTTTCTTGA	49	6239	6258	2825
568330	N/A	N/A	CATAATTTGAACCAGGCTTT	68	6244	6263	109
568331	N/A	N/A	AGACATAATACATAATTTGA	8	6254	6273	2826
568332	N/A	N/A	CTGTGATAAAGACATAATAC	40	6263	6282	2827
568333	N/A	N/A	CAGACCTGTGATAAAGACAT	16	6268	6287	2828
568334	N/A	N/A	ATCTTCAGACCTGTGATAAA	7	6273	6292	2829
568335	N/A	N/A	TACTGATCTTCAGACCTGTG	47	6278	6297	2830
568336	N/A	N/A	TTAATAATTTTCAGTTTTAG	35	6302	6321	2831
568337	N/A	N/A	TAAGTTTAATAATTTTCAGT	23	6307	6326	2832
568338	N/A	N/A	TTCAGATTTTAAGTTTAATA	10	6316	6335	2833
568339	N/A	N/A	TATATTTGATATTCTGTTCA	42	6332	6351	2834
568340	N/A	N/A	ATATTGTAATGTATTCTTTT	0	6368	6387	2835
568341	N/A	N/A	TTAGAATATTGTAATGTATT	19	6373	6392	2836
568342	N/A	N/A	TTTGCTTAGAATATTGTAAT	9	6378	6397	2837
568343	N/A	N/A	ACTGCTTTGCTTAGAATATT	36	6383	6402	2838
568344	N/A	N/A	AAGTAGAGACTGCTTTGCTT	60	6391	6410	2839
568345	N/A	N/A	GCAAGGCCAAAAGTAGAGAC	59	6401	6420	2840
568346	N/A	N/A	ACAGAGCAAGGCCAAAAGTA	45	6406	6425	2841
568347	N/A	N/A	GGAAAACAGAGCAAGGCCAA	49	6411	6430	2842
568348	N/A	N/A	TGGTCGGAAAACAGAGCAAG	38	6416	6435	2843
568349	N/A	N/A	GACATTGGTCGGAAAACAGA	26	6421	6440	2844
568350	N/A	N/A	AAGCAGACATTGGTCGGAAA	50	6426	6445	2845
568351	N/A	N/A	CAAGGCCAAAAAAGCAGACAT	39	6436	6455	2846
568352	N/A	N/A	ATAAAGCAAGGCCAAAAAAGC	20	6442	6461	2847
568353	N/A	N/A	CATTATTTAATAAGATAAAA	29	6464	6483	2848
568354	N/A	N/A	AAATATTTAATCAGGGACAT	35	6481	6500	2849
568355	N/A	N/A	TGTTCTCAAAATATTTAATC	32	6489	6508	2850
568356	N/A	N/A	GATTACCTGTTCTCAAAATA	40	6496	6515	2851
568357	N/A	N/A	GATTGTACAGATTACCTGTT	12	6505	6524	2852
568358	N/A	N/A	ATTCAGATTGTACAGATTAC	34	6510	6529	2853
568359	N/A	N/A	AAACAGTGTTATTCAGATTG	32	6520	6539	2854
568360	N/A	N/A	TAGATAAACAGTGTTATTCA	25	6525	6544	2855
568361	N/A	N/A	ATATTTAGATAAACAGTGTT	14	6530	6549	2856
568362	N/A	N/A	GTTTGATATTTAGATAAACA	27	6535	6554	2857
568363	N/A	N/A	AACGGTGTTTGATATTTAGA	33	6541	6560	2858

568364	N/A	N/A	GTTATAACGGTGTTTGATAT	29	6546	6565	2859
568365	N/A	N/A	ATAATGTTATAACGGTGTTT	21	6551	6570	2860
568366	N/A	N/A	AGTTCATAATGTTATAACGG	37	6556	6575	2861
568367	N/A	N/A	CTTTCAGTTCATAATGTTAT	46	6561	6580	2862
568368	N/A	N/A	AGTACAGTTTGTCTTTCAGT	48	6573	6592	2863
568369	N/A	N/A	TCAGAAGTACAGTTTGTCTT	47	6578	6597	2864
568370	N/A	N/A	GGATGTCAGAAGTACAGTTT	46	6583	6602	2865
568371	N/A	N/A	GAGTAAGGATGTCAGAAGTA	45	6589	6608	2866
568372	N/A	N/A	GAAATCTGAGTAAGGATGTC	31	6596	6615	2867
568373	N/A	N/A	TACTGAATATACAATTAGGG	5	6616	6635	2868
568374	N/A	N/A	AATGATACTGAATATACAAT	21	6621	6640	2869
568375	N/A	N/A	GAATATAAATCTGTTTTTTA	19	6642	6661	2870
568376	N/A	N/A	TAAAAGAATATAAATCTGTT	32	6647	6666	2871
568377	N/A	N/A	GCTGATAAAAGAATATAAAT	50	6652	6671	2872
568378	N/A	N/A	CCTTCTGAGCTGATAAAAGA	37	6660	6679	2873
568379	N/A	N/A	CTAGTCCTTCTGAGCTGATA	45	6665	6684	2874
568380	N/A	N/A	TTACCATCATGTTTTACATT	30	6770	6789	2875
568381	N/A	N/A	CAAAGTGTCTTACCATCATG	24	6779	6798	2876
568382	N/A	N/A	AAACCCACCAAAGTGTCTTA	15	6787	6806	2877
568383	N/A	N/A	AGAAGGAAACCCACCAAAGT	22	6793	6812	2878
568384	N/A	N/A	AATAATAGCTTCAAGAAGGA	25	6806	6825	2879
568385	N/A	N/A	AATTTGATAATAATAGCTTC	24	6814	6833	2880
568386	N/A	N/A	TAGGGAATTTGATAATAATA	20	6819	6838	2881
568387	N/A	N/A	AAGAATAGGGAATTTGATAA	0	6824	6843	2882
568388	N/A	N/A	GTCCTAAGAATAGGGAATTT	45	6829	6848	2883
568389	N/A	N/A	TAGAACAAGTCCTAAGAATA	21	6837	6856	2884
568390	N/A	N/A	TTAGTCTAGAACAAGTCCTA	28	6843	6862	2885
568391	N/A	N/A	ATCTTTTAGTCTAGAACAAG	21	6848	6867	2886
568392	N/A	N/A	TAACTATCTTTTAGTCTAGA	13	6853	6872	2887
568393	N/A	N/A	ATCTCTTA ACTATCTTTTAG	28	6859	6878	2888
568394	N/A	N/A	TGGATATCTCTTA ACTATCT	48	6864	6883	2889
568395	N/A	N/A	TTTGATGGATATCTCTTAAC	35	6869	6888	2890
544120	707	726	AGTTCTTGGTGCTCTTGGCT	80	6720	6739	15
337487	804	823	CACTTGTATGTTACCTCTG	76	7389	7408	28
568006	2014	2033	TTAATTCTGCTTCATTAGGT	53	10986	11005	2891
568007	2015	2034	TTTAATTCTGCTTCATTAGG	38	10987	11006	2892
568008	2020	2039	CAGTATTTAATTCTGCTTCA	56	10992	11011	2893
568009	2021	2040	ACAGTATTTAATTCTGCTTC	63	10993	11012	2894
568010	2022	2041	TACAGTATTTAATTCTGCTT	56	10994	11013	2895
568011	2023	2042	ATACAGTATTTAATTCTGCT	39	10995	11014	2896
568012	2024	2043	AATACAGTATTTAATTCTGC	21	10996	11015	2897

WO 2015/100394

PCT/US2014/072303

568013	2025	2044	TAATACAGTATTTAATTCTG	12	10997	11016	2898
568014	2027	2046	TTTAATACAGTATTTAATTC	0	10999	11018	2899
568015	2028	2047	TTTTAATACAGTATTTAATT	15	11000	11019	2900
568016	2031	2050	TTATTTTAATACAGTATTTA	0	11003	11022	2901
568017	2034	2053	AACTTATTTTAATACAGTAT	24	11006	11025	2902
568018	2035	2054	GAACTTATTTTAATACAGTA	21	11007	11026	2903
568019	2036	2055	CGAACTTATTTTAATACAGT	2	11008	11027	2904
568020	2037	2056	GCGAACTTATTTTAATACAG	54	11009	11028	2905
568021	2038	2057	AGCGAACTTATTTTAATACA	35	11010	11029	2906
568022	2039	2058	CAGCGAACTTATTTTAATAC	50	11011	11030	2907
568023	2040	2059	ACAGCGAACTTATTTTAATA	34	11012	11031	2908
568024	2041	2060	GACAGCGAACTTATTTTAAT	52	11013	11032	2909
568025	2042	2061	AGACAGCGAACTTATTTTAA	58	11014	11033	2910
568026	2044	2063	AAAGACAGCGAACTTATTTT	32	11016	11035	2911
568027	2045	2064	TAAAGACAGCGAACTTATTT	26	11017	11036	2912
568028	2048	2067	GTTTAAAGACAGCGAACTTA	62	11020	11039	2913
568029	2049	2068	TGTTTAAAGACAGCGAACTT	58	11021	11040	2914
568030	2050	2069	TTGTTTAAAGACAGCGAACT	52	11022	11041	2915
568031	2051	2070	TTTGTTTAAAGACAGCGAAC	61	11023	11042	2916
568032	2052	2071	ATTTGTTTAAAGACAGCGAA	41	11024	11043	2917
568033	2053	2072	CATTTGTTTAAAGACAGCGA	60	11025	11044	2918
568034	2054	2073	CCATTTGTTTAAAGACAGCG	88	11026	11045	98
568035	2055	2074	TCCATTTGTTTAAAGACAGC	57	11027	11046	2919
568036	2056	2075	CTCCATTTGTTTAAAGACAG	58	11028	11047	2920
568037	2058	2077	ATCTCCATTTGTTTAAAGAC	56	11030	11049	2921
568038	2059	2078	CATCTCCATTTGTTTAAAGA	54	11031	11050	2922
568039	2060	2079	TCATCTCCATTTGTTTAAAG	62	11032	11051	2923
568040	2061	2080	GTCATCTCCATTTGTTTAAA	53	11033	11052	2924
568041	2063	2082	TAGTCATCTCCATTTGTTTA	48	11035	11054	2925
568042	2064	2083	GTAGTCATCTCCATTTGTTT	44	11036	11055	2926
568043	2065	2084	AGTAGTCATCTCCATTTGTT	48	11037	11056	2927
568044	2066	2085	TAGTAGTCATCTCCATTTGT	45	11038	11057	2928
568045	2067	2086	TTAGTAGTCATCTCCATTTG	66	11039	11058	2929
568046	2068	2087	CTTAGTAGTCATCTCCATTT	66	11040	11059	2930
568047	2069	2088	ACTTAGTAGTCATCTCCATT	68	11041	11060	99
568048	2070	2089	GACTTAGTAGTCATCTCCAT	77	11042	11061	100
568049	2071	2090	TGACTTAGTAGTCATCTCCA	70	11043	11062	101
568050	2072	2091	GTGACTTAGTAGTCATCTCC	65	11044	11063	2931
568051	2073	2092	TGTGACTTAGTAGTCATCTC	49	11045	11064	2932
568052	2074	2093	ATGTGACTTAGTAGTCATCT	47	11046	11065	2933
568053	2075	2094	AATGTGACTTAGTAGTCATC	48	11047	11066	2934

568054	2076	2095	CAATGTGACTTAGTAGTCAT	60	11048	11067	2935
568055	2077	2096	TCAATGTGACTTAGTAGTCA	54	11049	11068	2936
568056	2078	2097	GTCAATGTGACTTAGTAGTC	72	11050	11069	102
568057	2079	2098	AGTCAATGTGACTTAGTAGT	62	11051	11070	2937
568058	2083	2102	TTAAAGTCAATGTGACTTAG	15	11055	11074	2938
568059	2084	2103	GTAAAGTCAATGTGACTTA	28	11056	11075	2939
568060	2085	2104	TGTTAAAGTCAATGTGACTT	35	11057	11076	2940
568061	2086	2105	ATGTTAAAGTCAATGTGACT	17	11058	11077	2941
568062	2087	2106	CATGTTAAAGTCAATGTGAC	27	11059	11078	2942
568063	2089	2108	CTCATGTTAAAGTCAATGTG	28	11061	11080	2943
568064	2090	2109	CCTCATGTTAAAGTCAATGT	50	11062	11081	2944
568066	2091	2110	ACCTCATGTTAAAGTCAATG	48	11063	11082	2945
568068	2092	2111	TACCTCATGTTAAAGTCAAT	13	11064	11083	2946
568069	2093	2112	ATACCTCATGTTAAAGTCAA	43	11065	11084	2947
568072	2094	2113	GATACCTCATGTTAAAGTCA	40	11066	11085	2948
568073	2095	2114	TGATACCTCATGTTAAAGTC	40	11067	11086	2949
568075	2096	2115	GTGATACCTCATGTTAAAGT	37	11068	11087	2950
568077	2097	2116	AGTGATACCTCATGTTAAAG	6	11069	11088	2951
568078	2098	2117	TAGTGATACCTCATGTTAAA	12	11070	11089	2952
568079	2099	2118	ATAGTGATACCTCATGTTAA	8	11071	11090	2953
568080	2100	2119	TATAGTGATACCTCATGTTA	13	11072	11091	2954
568081	2101	2120	GTATAGTGATACCTCATGTT	41	11073	11092	2955
568082	2102	2121	GGTATAGTGATACCTCATGT	53	11074	11093	2956
568083	2106	2125	ATAAGGTATAGTGATACCTC	54	11078	11097	2957
568084	2107	2126	AATAAGGTATAGTGATACCT	38	11079	11098	2958

Table 14

Inhibition of ANGPTL3 mRNA by 5-10-5 MOE gapmers targeting SEQ ID NO: 1 and 2

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
544120	707	726	AGTTCTTGGTGCTCTTGGCT	83	6720	6739	15
337487	804	823	CACTTGTATGTTACCTCTG	81	7389	7408	28
567295	1452	1471	TAATGTTTAAATTATTGCCT	43	10424	10443	2959
567296	1455	1474	GGTTAATGTTTAAATTATTG	22	10427	10446	2960
567297	1456	1475	AGGTTAATGTTTAAATTATT	0	10428	10447	2961
567298	1457	1476	GAGGTTAATGTTTAAATTAT	0	10429	10448	2962
567299	1458	1477	TGAGGTTAATGTTTAAATTA	6	10430	10449	2963
567300	1460	1479	AATGAGGTTAATGTTTAAAT	14	10432	10451	2964
567301	1461	1480	GAATGAGGTTAATGTTTAAA	5	10433	10452	2965
567302	1462	1481	GGAATGAGGTTAATGTTTAA	27	10434	10453	2966

567303	1463	1482	TGGAATGAGGTTAATGTTTA	32	10435	10454	2967
567304	1464	1483	TTGGAATGAGGTTAATGTTT	37	10436	10455	2968
567305	1465	1484	CTTGGAATGAGGTTAATGTT	25	10437	10456	2969
567306	1468	1487	TAACTTGGAATGAGGTTAAT	29	10440	10459	2970
567307	1469	1488	TTAACTTGGAATGAGGTTAA	44	10441	10460	2971
337513	1470	1489	ATTA ACTTGGAATGAGGTTA	52	10442	10461	2972
567308	1471	1490	CATTA ACTTGGAATGAGGTT	62	10443	10462	2973
567309	1472	1491	ACATTA ACTTGGAATGAGGT	58	10444	10463	2974
567310	1473	1492	CACATTA ACTTGGAATGAGG	78	10445	10464	92
567311	1475	1494	ACCACATTA ACTTGGAATGA	59	10447	10466	2975
567312	1476	1495	GACCACATTA ACTTGGAATG	57	10448	10467	2976
337514	1477	1496	AGACCACATTA ACTTGGAAT	71	10449	10468	2977
567313	1478	1497	TAGACCACATTA ACTTGGA	43	10450	10469	2978
567314	1479	1498	TTAGACCACATTA ACTTGGA	59	10451	10470	2979
567315	1480	1499	ATTAGACCACATTA ACTTGG	70	10452	10471	2980
567316	1481	1500	TATTAGACCACATTA ACTTG	53	10453	10472	2981
567317	1482	1501	TTATTAGACCACATTA ACTT	49	10454	10473	2982
567318	1483	1502	ATTATTAGACCACATTA ACT	41	10455	10474	2983
567319	1484	1503	GATTATTAGACCACATTA AC	47	10456	10475	2984
567320	1487	1506	CCAGATTATTAGACCACATT	86	10459	10478	93
567321	1489	1508	TACCAGATTATTAGACCACA	85	10461	10480	94
337516	1490	1509	ATACCAGATTATTAGACCAC	77	10462	10481	86
567322	1491	1510	AATACCAGATTATTAGACCA	50	10463	10482	2985
567323	1492	1511	TAATACCAGATTATTAGACC	56	10464	10483	2986
567324	1494	1513	TTTAATACCAGATTATTAGA	9	10466	10485	2987
567325	1495	1514	ATTTAATACCAGATTATTAG	24	10467	10486	2988
567326	1496	1515	GATTTAATACCAGATTATTA	37	10468	10487	2989
567327	1500	1519	TAAGGATTTAATACCAGATT	60	10472	10491	2990
567328	1507	1526	TTTCTCTTAAGGATTTAATA	34	10479	10498	2991
567329	1508	1527	CTTTCTCTTAAGGATTTAAT	46	10480	10499	2992
567330	1509	1528	GCTTTCTCTTAAGGATTTAA	75	10481	10500	95
567331	1510	1529	AGCTTTCTCTTAAGGATTTA	59	10482	10501	2993
567332	1511	1530	AAGCTTTCTCTTAAGGATTT	30	10483	10502	2994
567333	1513	1532	TCAAGCTTTCTCTTAAGGAT	65	10485	10504	2995
567334	1514	1533	CTCAAGCTTTCTCTTAAGGA	77	10486	10505	96
567335	1515	1534	TCTCAAGCTTTCTCTTAAGG	75	10487	10506	97
567336	1516	1535	TTCTCAAGCTTTCTCTTAAG	59	10488	10507	2996
567337	1517	1536	TTTCTCAAGCTTTCTCTTAA	52	10489	10508	2997
567338	1521	1540	TCTATTTCTCAAGCTTTCTC	68	10493	10512	2998
567339	1522	1541	ATCTATTTCTCAAGCTTTCT	71	10494	10513	2999
567340	1523	1542	AATCTATTTCTCAAGCTTTC	74	10495	10514	3000

567341	1524	1543	AAATCTATTTCTCAAGCTTT	63	10496	10515	3001
567342	1525	1544	AAAATCTATTTCTCAAGCTT	54	10497	10516	3002
567343	1532	1551	GATAAAAAAAAAATCTATTTCT	30	10504	10523	3003
567344	1548	1567	TAGACAGTGACTTTAAGATA	37	10520	10539	3004
567345	1549	1568	ATAGACAGTGACTTTAAGAT	29	10521	10540	3005
567346	1550	1569	AATAGACAGTGACTTTAAGA	48	10522	10541	3006
567347	1551	1570	AAATAGACAGTGACTTTAAG	26	10523	10542	3007
567348	1552	1571	TAAATAGACAGTGACTTTAA	26	10524	10543	3008
567349	1553	1572	TTAAATAGACAGTGACTTTA	50	10525	10544	3009
567350	1554	1573	CTTAAATAGACAGTGACTTT	63	10526	10545	3010
567351	1555	1574	TCTTAAATAGACAGTGACTT	57	10527	10546	3011
567352	1556	1575	ATCTTAAATAGACAGTGACT	69	10528	10547	3012
567353	1557	1576	AATCTTAAATAGACAGTGAC	40	10529	10548	3013
567354	1558	1577	TAATCTTAAATAGACAGTGA	30	10530	10549	3014
567355	1559	1578	TTAATCTTAAATAGACAGTG	25	10531	10550	3015
567356	1560	1579	TTTAATCTTAAATAGACAGT	0	10532	10551	3016
567357	1561	1580	GTTTAATCTTAAATAGACAG	34	10533	10552	3017
567358	1562	1581	TGTTTAATCTTAAATAGACA	5	10534	10553	3018
567359	1563	1582	ATGTTTAATCTTAAATAGAC	0	10535	10554	3019
567360	1567	1586	TTGTATGTTTAATCTTAAAT	0	10539	10558	3020
567361	1568	1587	ATTGTATGTTTAATCTTAAA	8	10540	10559	3021
567362	1569	1588	GATTGTATGTTTAATCTTAA	20	10541	10560	3022
567363	1570	1589	TGATTGTATGTTTAATCTTA	29	10542	10561	3023
567364	1574	1593	TATGTGATTGTATGTTTAAT	7	10546	10565	3024
567365	1576	1595	GTTATGTGATTGTATGTTTA	43	10548	10567	3025
567366	1580	1599	TAAGGTTATGTGATTGTATG	28	10552	10571	3026
567367	1581	1600	TTAAGGTTATGTGATTGTAT	31	10553	10572	3027
567368	1585	1604	TTCTTTAAGGTTATGTGATT	12	10557	10576	3028

Example 2: Dose-dependent antisense inhibition of human ANGPTL3 in Hep3B cells by MOE gapmers

5-10-5 MOE gapmers from the studies described above exhibiting significant *in vitro* inhibition of ANGPTL3 mRNA were selected and tested at various doses in Hep3B cells. Cells were plated at a density of 20,000 cells per well and transfected using electroporation with 0.75 μ M, 1.50 μ M, 3.00 μ M, 6.00 μ M and 12.00 μ M 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 ANGPTL3 mRNA levels were measured by quantitative real-time PCR. Human primer probe set RTS3492_MGB was used to measure mRNA levels. ANGPTL3 mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN[®]. Results are presented as percent inhibition of ANGPTL3, relative to untreated control cells.

The half maximal inhibitory concentration (IC₅₀) of each oligonucleotide is also presented. ANGPTL3 mRNA levels were significantly reduced in a dose-dependent manner in antisense oligonucleotide treated cells.

Table 15

ISIS No	0.75 μM	1.50 μM	3.00 μM	6.00 μM	12.00 μM	IC ₅₀ (μM)	SEQ ID NO
233717	23	45	13	33	40	>12	14
544120	45	65	76	88	91	0.7	15
544145	38	42	61	82	84	1.6	16
544156	31	42	63	78	84	1.8	17
544162	35	43	71	76	82	1.6	18
544166	30	47	60	76	84	1.8	19
544199	54	61	73	83	84	0.5	20
544355	45	46	69	77	83	1.2	21
544368	12	37	63	74	81	2.6	22
544373	1	27	40	29	28	>12	23
544376	26	53	61	63	59	2.4	24
544380	16	33	41	64	39	8.4	25
544383	14	33	46	61	63	4.4	26
544410	10	41	48	62	69	3.6	27

5

Example 3: Dose-dependent antisense inhibition of human ANGPTL3 in Hep3B cells by MOE gapmers

5-10-5 MOE gapmers from the studies described above exhibiting significant *in vitro* inhibition of ANGPTL3 mRNA were selected and tested at various doses in Hep3B cells. Cells were plated at a density of 20,000 cells per well and transfected using electroporation with 0.813 μM, 1.625 μM, 3.25 μM, 6.50 μM and 13.00 μM 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 ANGPTL3 mRNA levels were measured by quantitative real-time PCR. Human primer probe set RTS3492_MGB was used to measure mRNA levels. ANGPTL3 mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN[®]. Results are presented as percent inhibition of ANGPTL3, relative to untreated control cells.

15

The half maximal inhibitory concentration (IC₅₀) of each oligonucleotide is also presented. ANGPTL3 mRNA levels were significantly reduced in a dose-dependent manner in antisense oligonucleotide treated cells.

Table 16

ISIS No	0.813 μM	1.625 μM	3.25 μM	6.50 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
337487	17	37	58	72	92	2.7	28
337492	0	0	0	5	58	>13	29
544120	23	40	65	81	91	2.2	15
560236	39	22	46	9	60	>13	30
560265	38	48	58	64	73	2.0	31
560268	37	57	60	71	83	1.5	32
560285	5	29	48	68	78	3.8	33
560306	45	64	67	81	86	0.9	34
560400	48	63	75	87	88	0.7	35
560401	49	75	79	89	88	0.5	36
560402	42	67	70	85	90	0.9	37
560469	43	55	70	74	83	1.2	38
560470	31	54	64	73	81	1.8	39
560471	26	43	59	62	77	2.7	40
560474	42	50	60	54	72	1.8	41

Table 17

ISIS No	0.813 μM	1.625 μM	3.25 μM	6.50 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
337487	20	35	51	78	89	1.8	28
544120	31	46	62	84	90	0.5	15
544145	4	36	60	58	89	3.8	16
544156	22	35	46	66	73	1.8	17
544162	2	21	54	69	87	>13	18
544166	15	0	25	59	89	>13	19
544199	61	37	57	53	81	0.9	20
544355	0	47	50	73	84	>13	21
544376	4	14	38	66	88	0.9	24
560566	53	68	70	76	85	>13	42
560567	55	70	75	78	89	2.7	43
560574	49	63	68	74	84	2.0	44
560596	28	40	41	52	75	1.5	45
560607	35	53	65	70	85	3.8	46
560608	40	50	62	68	83	0.9	47
560723	36	51	59	65	75	2.2	48
560735	36	44	59	72	85	>13	49
560736	26	34	50	64	80	0.7	50
560744	28	49	59	75	83	0.9	51
560778	24	46	60	67	85	1.8	52

560789	14	23	36	49	71	2.7	53
560811	32	50	65	73	87	1.2	54
560856	0	20	17	32	69	3.8	55
560925	2	16	38	52	82	2.7	56
560936	0	0	24	41	65	0.5	57
560938	0	26	30	43	50	0.9	58
560942	0	0	12	36	74	1.8	59
560956	0	16	16	68	81	0.5	60

Table 18

ISIS No	0.813 μM	1.625 μM	3.25 μM	6.50 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
337487	20	35	51	78	89	2.7	28
544120	31	46	62	84	90	1.9	15
560566	53	68	70	76	85	0.5	42
560567	55	70	75	78	89	0.4	43
560574	49	63	68	74	84	0.7	44
560596	28	40	41	52	75	3.9	45
560607	35	53	65	70	85	1.6	46
560608	40	50	62	68	83	1.6	47
560723	36	51	59	65	75	1.9	48
560735	36	44	59	72	85	2.0	49
560736	26	34	50	64	80	3.2	50
560744	28	49	59	75	83	2.1	51
560778	24	46	60	67	85	2.4	52
560789	14	23	36	49	71	5.7	53
560811	32	50	65	73	87	1.8	54

Table 19

ISIS No	0.813 μM	1.625 μM	3.25 μM	6.50 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
337487	10	21	49	73	90	3.4	28
544120	19	38	62	77	88	2.5	15
560768	1	14	14	28	51	>13	61
560777	13	35	37	56	80	4.2	62
560791	13	28	28	24	11	>13	63
560794	8	31	42	57	76	4.4	64
560799	0	14	21	43	72	7.2	65
560803	26	44	52	55	69	3.4	66
560815	16	26	26	52	60	7.6	67

560817	0	0	11	18	37	>13	68
560847	37	52	56	68	87	1.8	69
560879	15	18	38	53	72	5.4	70
560880	0	8	21	38	71	8.0	71
560891	7	25	32	35	62	8.9	72
560895	11	10	0	5	48	>13	73

Table 20

ISIS No	0.813 μM	1.625 μM	3.25 μM	6.50 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
337487	20	14	38	65	88	3.9	28
544120	22	34	51	71	86	2.9	15
544145	21	39	62	63	90	2.6	16
544156	31	41	55	72	78	2.4	17
544162	0	37	59	75	87	2.7	18
544166	8	43	45	55	75	4.0	19
544199	53	46	64	62	81	1.1	20
544355	0	0	52	72	84	2.9	21
544376	2	22	39	51	76	5.2	24
560856	10	29	36	41	69	6.4	55
560925	0	35	46	59	81	3.5	56
560936	18	9	35	55	69	5.9	57
560938	14	34	42	49	58	6.5	58
560942	8	13	27	47	77	6.1	59
560956	16	31	0	69	81	3.9	60

Table 21

ISIS No	0.813 μM	1.625 μM	3.25 μM	6.50 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
233717	11	0	33	58	75	5.0	14
337484	39	54	55	66	79	1.7	74
337487	35	42	67	82	92	1.8	28
544120	53	47	78	84	92	<0.8	15
563523	12	44	59	63	79	3.0	75
563547	33	51	55	43	58	4.6	76
563580	61	73	71	82	91	<0.8	77
563637	36	55	69	77	88	1.4	78
563639	56	71	79	88	93	<0.8	79
563641	30	42	56	77	84	2.2	80

563669	28	61	66	79	85	1.6	81
563681	35	67	74	75	70	0.9	82
563682	41	45	68	76	85	1.5	83
567068	32	37	50	66	81	2.8	84
567069	23	28	48	56	62	5.0	85

Table 22

ISIS No	0.813 μM	1.625 μM	3.25 μM	6.50 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
233717	9	0	25	62	74	5.5	14
337487	22	40	71	84	92	2.1	28
337516	36	54	78	81	92	1.3	86
544120	25	50	72	86	92	1.8	15
567078	54	64	70	78	78	<0.8	87
567115	55	65	72	80	81	<0.8	88
567134	33	58	53	57	69	2.2	89
567233	54	74	83	87	91	<0.8	90
567291	54	67	71	80	89	<0.8	91
567310	36	61	73	80	89	1.2	92
567320	63	77	88	88	92	<0.8	93
567321	55	75	89	89	93	<0.8	94
567330	31	68	76	85	93	1.2	95
567334	36	54	76	82	87	1.3	96
567335	31	49	72	80	92	1.7	97

Table 23

ISIS No	0.813 μM	1.625 μM	3.25 μM	6.50 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
233717	0	0	23	66	64	6.6	14
337487	13	44	60	74	85	2.6	28
544120	24	47	53	78	83	2.3	15
568034	35	54	51	59	46	4.2	98
568047	36	55	70	69	72	1.4	99
568048	41	64	63	66	66	0.9	100
568049	50	70	70	74	73	<0.8	101
568056	33	56	68	63	64	1.7	102
568144	27	57	63	63	76	2.0	103
568146	50	61	61	63	77	<0.8	104
568151	23	46	59	68	66	2.8	105

568206	24	40	56	61	75	3.0	106
568293	0	39	46	59	78	4.1	107
568300	22	36	61	68	73	3.0	108
568330	16	48	54	73	82	2.7	109

Example 4: Antisense inhibition of human ANGPTL3 in Hep3B cells by deoxy, MOE and (S)-cEt gapmers

Additional antisense oligonucleotides were designed targeting an ANGPTL3 nucleic acid and were tested for their effects on ANGPTL3 mRNA in vitro. Cultured Hep3B cells at a density of 20,000 cells per well were transfected using electroporation with 4,500 nM of antisense oligonucleotide. After a treatment period of approximately 24 hours, RNA was isolated from the cells and ANGPTL3 mRNA levels were measured by quantitative real-time PCR. Human primer probe set RTS3492_MGB was used to measure mRNA levels. ANGPTL3 mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN®. Results are presented as percent inhibition of ANGPTL3, relative to untreated control cells.

The newly designed chimeric antisense oligonucleotides in the Tables below were designed as deoxy, MOE, and (S)-cEt oligonucleotides. The deoxy, MOE and (S)-cEt oligonucleotides are 16 nucleosides in length wherein the nucleoside have either a MOE sugar modification, a (S)-cEt sugar modification, or a deoxy sugar residue. The sugar modifications of each antisense oligonucleotide is described as ‘eek-d10-kke’, where ‘k’ indicates a (S)-cEt sugar modification; ‘d’ indicates deoxyribose; the number indicates the number of deoxyribose sugars residues; and ‘e’ indicates a MOE sugar modification. The internucleoside linkages throughout each oligonucleotide are phosphorothioate (P=S) linkages. All cytosine residues throughout each oligonucleotide are 5-methylcytosines. “Start site” indicates the 5’-most nucleoside to which the oligonucleotide is targeted in the human gene sequence. “Stop site” indicates the 3’-most nucleoside to which the oligonucleotide is targeted human gene sequence. Each oligonucleotide listed in the Tables below is targeted to either the human ANGPTL3 mRNA, designated herein as SEQ ID NO: 1 (GENBANK Accession No. NM_014495.2) or the human ANGPTL3 genomic sequence, designated herein as SEQ ID NO: 2 (GENBANK Accession No. NT_032977.9 truncated from nucleotides 33032001 to 33046000). ‘n/a’ indicates that the antisense oligonucleotide does not target that particular gene sequence with 100% complementarity.

Table 24

Inhibition of ANGPTL3 mRNA by deoxy, MOE and cEt oligonucleotides targeting SEQ ID NO: 1 and 2

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
561681	N/A	N/A	TCTGGAAGCAGACCTA	37	3096	3111	3029
561682	N/A	N/A	CTTCTGGAAGCAGACC	27	3098	3113	3030
561683	N/A	N/A	AAATAAGGTATAGTGA	2	11084	11099	3031
561684	N/A	N/A	TAGTATTAAGTGTTAA	14	11133	11148	3032
561685	N/A	N/A	TCATAGTATTAAGTGT	0	11136	11151	3033
561686	N/A	N/A	AGATTCCTTTACAATT	21	11160	11175	3034
561687	N/A	N/A	ACAAGATTCCTTTACA	21	11163	11178	3035
561688	N/A	N/A	CTGACAAGATTCCTTT	70	11166	11181	3036
561689	N/A	N/A	AATCTGACAAGATTCC	83	11169	11184	180
561690	N/A	N/A	TGTAATCTGACAAGAT	46	11172	11187	3037
561691	N/A	N/A	TACTGTAATCTGACAA	47	11175	11190	3038
561692	N/A	N/A	TCTTACTGTAATCTGA	50	11178	11193	3039
561693	N/A	N/A	CATTCTTACTGTAATC	40	11181	11196	3040
561694	N/A	N/A	GTTCATTCTTACTGTA	71	11184	11199	3041
561695	N/A	N/A	ATATGTTTCATTCTTAC	2	11188	11203	3042
561696	N/A	N/A	GCCACAAATATGTTCA	80	11195	11210	3043
561697	N/A	N/A	GATGCCACAAATATGT	70	11198	11213	3044
561698	N/A	N/A	CTCGATGCCACAAATA	80	11201	11216	181
561699	N/A	N/A	TAAC TCGATGCCACAA	86	11204	11219	182
561700	N/A	N/A	CTTTAACTCGATGCCA	77	11207	11222	3045
561701	N/A	N/A	AAACTTTAACTCGATG	39	11210	11225	3046
561702	N/A	N/A	TATAAACTTTAACTCG	13	11213	11228	3047
561703	N/A	N/A	CACAGCATATTTAGGG	71	11233	11248	3048
561704	N/A	N/A	TAGAATCACAGCATAT	68	11239	11254	3049
561705	N/A	N/A	TATTAGAATCACAGCA	73	11242	11257	3050
561706	N/A	N/A	AATGTATTAGAATCAC	40	11246	11261	3051
561707	N/A	N/A	ACGAATGTATTAGAAT	22	11249	11264	3052
561708	N/A	N/A	TACACGAATGTATTAG	33	11252	11267	3053
561709	N/A	N/A	ACCTACACGAATGTAT	42	11255	11270	3054
561710	N/A	N/A	AAAACCTACACGAATG	24	11258	11273	3055
561711	N/A	N/A	TTGAAAACCTACACGA	34	11261	11276	3056
561712	N/A	N/A	TACTTGAAAACCTACA	33	11264	11279	3057
561713	N/A	N/A	GTTTATTTCTACTTGA	53	11273	11288	3058
561714	N/A	N/A	GAGGTTTATTTCTACT	69	11276	11291	3059
561715	N/A	N/A	TACGAGGTTTATTTCT	21	11279	11294	3060

561716	N/A	N/A	TGTTACGAGGTTTATT	47	11282	11297	3061
561717	N/A	N/A	ACTTGTTACGAGGTTT	70	11285	11300	3062
561718	N/A	N/A	CAGTAACTTGTTACGA	60	11290	11305	3063
561719	N/A	N/A	GTTCAGTAACTTGTTA	40	11293	11308	3064
561720	N/A	N/A	TCAGGCTGTTTAAACG	59	11308	11323	3065
561721	N/A	N/A	TTGTCAGGCTGTTTAA	74	11311	11326	3066
561722	N/A	N/A	TGCTTGTCAGGCTGTT	82	11314	11329	183
561723	N/A	N/A	ACATGCTTGTCAGGCT	84	11317	11332	184
561724	N/A	N/A	TATACATGCTTGTCAG	75	11320	11335	3067
561725	N/A	N/A	GTCTTTGTTTATTGAA	49	11347	11362	3068
561726	N/A	N/A	TGGGTCTTTGTTTATT	27	11350	11365	3069
561727	N/A	N/A	GACTGGGTCTTTGTTT	20	11353	11368	3070
561728	N/A	N/A	ATAATTTAGGGACTGG	20	11363	11378	3071
561729	N/A	N/A	TCTATAATTTAGGGAC	39	11366	11381	3072
561730	N/A	N/A	CGATAAACATGCAAGA	68	11394	11409	3073
561731	N/A	N/A	TGTCGATAAACATGCA	80	11397	11412	3074
561732	N/A	N/A	TGATGTCGATAAACAT	68	11400	11415	3075
561733	N/A	N/A	TTGTGATGTCGATAAA	28	11403	11418	3076
561734	N/A	N/A	CTGTTGTGATGTCGAT	74	11406	11421	3077
561735	N/A	N/A	GATCTGTTGTGATGTC	59	11409	11424	3078
561736	N/A	N/A	AGGGATCTGTTGTGAT	24	11412	11427	3079
561737	N/A	N/A	TTAGGGATCTGTTGT	19	11415	11430	3080
561738	N/A	N/A	GGATTTAGGGATCTGT	27	11418	11433	3081
561739	N/A	N/A	GATTTAGGGATTTAGG	44	11425	11440	3082
561740	N/A	N/A	TCTTTAGGGATTTAGG	38	11433	11448	3083
561741	N/A	N/A	TAATCTTTAGGGATTT	0	11436	11451	3084
561742	N/A	N/A	ATCTAATCTTTAGGGA	0	11439	11454	3085
561743	N/A	N/A	TGTATCTAATCTTTAG	15	11442	11457	3086
561744	N/A	N/A	AAATTTGTATCTAATC	21	11447	11462	3087
561745	N/A	N/A	GTAAAAAATTTGTATC	23	11452	11467	3088
561746	N/A	N/A	GTGGTAAAAAATTTGT	32	11455	11470	3089
561747	N/A	N/A	GATACTGTGGTAAAAA	45	11461	11476	3090
561748	N/A	N/A	AGTGATACTGTGGTAA	60	11464	11479	3091
561749	N/A	N/A	ACAAGTGATACTGTGG	75	11467	11482	3092
561750	N/A	N/A	CTGACAAGTGATACTG	59	11470	11485	3093
561751	N/A	N/A	ATTCTGACAAGTGATA	48	11473	11488	3094
561752	N/A	N/A	TAAATTCTGACAAGTG	59	11476	11491	3095
561753	N/A	N/A	TACTGGCAGTTTTAAA	42	11508	11523	3096
561754	N/A	N/A	TCTTACTGGCAGTTTT	51	11511	11526	3097
561755	N/A	N/A	ATTTCTTACTGGCAGT	69	11514	11529	3098
561756	N/A	N/A	AAAATTTCTTACTGGC	57	11517	11532	3099

561757	N/A	N/A	AACAAATGGGTTTAAT	0	11535	11550	3100
562374	N/A	N/A	GAATATTTGCAAGTCT	68	9230	9245	3101
562375	N/A	N/A	GTAGAGGAATATTTGC	83	9236	9251	151
562376	N/A	N/A	TCATTGGTAGAGGAAT	23	9242	9257	3102
562377	N/A	N/A	ATATTTTAAAGTCTCG	17	9258	9273	3103
562378	N/A	N/A	GTTACATTATTATAGA	29	9273	9288	3104
562379	N/A	N/A	GTGAAATGTGTTACAT	54	9282	9297	3105
562380	N/A	N/A	TCACCAGTGAAATGTG	64	9288	9303	3106
562381	N/A	N/A	CATGTTTCACCAGTGA	78	9294	9309	3107
562382	N/A	N/A	ACAAGACATGTTTCAC	36	9300	9315	3108
562383	N/A	N/A	CATATGACAAGACATG	42	9306	9321	3109
562384	N/A	N/A	CTATAATGCATATGAC	5	9314	9329	3110
562385	N/A	N/A	TCCTTTCTATAATGCA	65	9320	9335	3111
562386	N/A	N/A	TGATTATCCTTTCTAT	27	9326	9341	3112
562387	N/A	N/A	AAAGTCTGATTATCCT	90	9332	9347	152
562388	N/A	N/A	TAAGTCAAAGTCTGAT	59	9338	9353	3113
562389	N/A	N/A	GTGCACAAAAATGTTA	42	9366	9381	3114
562390	N/A	N/A	AGCTATGTGCACAAAA	77	9372	9387	3115
562391	N/A	N/A	GAAGATAGCTATGTGC	64	9378	9393	3116
562392	N/A	N/A	TTTATTGAAGATAGCT	33	9384	9399	3117
562393	N/A	N/A	TCATTTTAGTGTATCT	40	9424	9439	3118
562394	N/A	N/A	CCTTGATCATTTTAGT	15	9430	9445	3119
562395	N/A	N/A	TGAATCCCTTGATCAT	59	9436	9451	3120
562396	N/A	N/A	TAGTCTTGAATCCCTT	83	9442	9457	153
562397	N/A	N/A	GTTGTTTAGTCTTGAA	65	9448	9463	3121
562398	N/A	N/A	AATTGAGTTGTTTAGT	21	9454	9469	3122
562399	N/A	N/A	GCAACTAATTGAGTTG	15	9460	9475	3123
562400	N/A	N/A	ATTGGTGCAACTAATT	25	9466	9481	3124
562401	N/A	N/A	GTTTTTTATTGGTGCA	53	9473	9488	3125
562402	N/A	N/A	GGACACTGACAGTTTT	43	9496	9511	3126
562403	N/A	N/A	CAGGTTGGACACTGAC	23	9502	9517	3127
562404	N/A	N/A	TAAGTACAGGTTGGAC	33	9508	9523	3128
562405	N/A	N/A	AGTTATTAAGTACAGG	34	9514	9529	3129
562406	N/A	N/A	TCTGTGAGTTATTAAG	10	9520	9535	3130
562407	N/A	N/A	ACCAAAATTCTCCTGA	1	9554	9569	3131
562408	N/A	N/A	ACCTGAATAACCCTCT	73	9811	9826	3132
562409	N/A	N/A	GGTATCAGAAAAAGAT	14	9827	9842	3133
562410	N/A	N/A	AGTATTGGTATCAGAA	13	9833	9848	3134
562411	N/A	N/A	GGAAGATACTTTGAAG	25	9861	9876	3135
562412	N/A	N/A	AATGTGGGAAGATACT	23	9867	9882	3136
562413	N/A	N/A	CAGATAATAGCTAATA	29	9882	9897	3137

562414	N/A	N/A	TCATTGCAGATAATAG	45	9888	9903	3138
562415	N/A	N/A	AAGTTGTCATTGCAGA	86	9894	9909	154
562416	N/A	N/A	GATTCGGATTTTTTAAA	19	9909	9924	3139
562417	N/A	N/A	ATTTGGGATTCGGATT	34	9915	9930	3140
562418	N/A	N/A	ACGCTTATTTGGGATT	64	9921	9936	3141
562419	N/A	N/A	TCTAGAGAGAAAACGC	64	9933	9948	3142
562420	N/A	N/A	AGTTAAGAGGTTTTTCG	34	9949	9964	3143
562421	N/A	N/A	CATTATAGTTAAGAGG	24	9955	9970	3144
562422	N/A	N/A	CACTTTCATTATAGTT	13	9961	9976	3145
562423	N/A	N/A	TAGAATGAACACTTTC	63	9970	9985	3146
562424	N/A	N/A	TTGAACTAGAATGAAC	16	9976	9991	3147
562425	N/A	N/A	ACCTGATTGAACTAGA	51	9982	9997	3148
562426	N/A	N/A	TAAAATACCTGATTGA	19	9988	10003	3149
562427	N/A	N/A	TAGAGGTAAAATACCT	12	9994	10009	3150
562428	N/A	N/A	GAAGATTAGAGGTAAA	1	10000	10015	3151
562429	N/A	N/A	TCTGAGGAAGATTAGA	31	10006	10021	3152
562430	N/A	N/A	TATACACTACCAAAAA	0	10030	10045	3153
562431	N/A	N/A	ATAATCTATACACTAC	0	10036	10051	3154
562432	N/A	N/A	TAAGTCCCAATTTTAA	33	10065	10080	3155
562433	N/A	N/A	TCTGTATAAGTCCCAA	89	10071	10086	155
562434	N/A	N/A	CCAGTTTTAAATAATC	20	10085	10100	3156
562435	N/A	N/A	TGTATCCCAGTTTTAA	44	10091	10106	3157
562436	N/A	N/A	GATGCATGTATCCCAG	91	10097	10112	156
562437	N/A	N/A	GTTTTAGATGCATGTA	69	10103	10118	3158
562438	N/A	N/A	TACAGTGTTTTAGATG	28	10109	10124	3159
562439	N/A	N/A	GTAAGTTTATCTTCCT	78	10138	10153	157
562440	N/A	N/A	TTCCCCGTAAGTTTAT	33	10144	10159	3160
562441	N/A	N/A	CTGTATTTCCCCGTAA	55	10150	10165	3161
562442	N/A	N/A	CTGTTACTGTATTTCC	79	10156	10171	158
562443	N/A	N/A	TAGTTACTGTTACTGT	70	10162	10177	3162
562444	N/A	N/A	CGTATGTAGTTACTGT	66	10168	10183	3163
562445	N/A	N/A	AATGGGTACAGACTCG	72	10182	10197	3164
562446	N/A	N/A	GCAATTTAATGGGTAC	59	10189	10204	3165
562447	N/A	N/A	GATAGATATGCAATTT	20	10198	10213	3166
562448	N/A	N/A	AAAGGAGATAGATATG	22	10204	10219	3167
562449	N/A	N/A	CCTCCTAAAGGAGATA	42	10210	10225	3168
562450	N/A	N/A	CACCAGCCTCCTAAAG	37	10216	10231	3169
560990	709	724	TTCTTGGTGCTCTTGG	89	6722	6737	111
561373	1197	1212	TTTGTGATCCCAAGTA	40	9772	9787	3170
561374	1199	1214	GCTTTGTGATCCCAAG	76	9774	9789	3171
561375	1201	1216	TTGCTTTGTGATCCCA	82	9776	9791	3172

561376	1203	1218	TTTTGCTTTGTGATCC	40	9778	9793	3173
561377	1205	1220	CCTTTTGCTTTGTGAT	38	9780	9795	3174
561378	1207	1222	GTCCTTTTGCTTTGTG	75	9782	9797	3175
561379	1209	1224	GTGTCCTTTTGCTTTG	40	9784	9799	3176
561527	1604	1619	GAAATGTAAACGGTAT	47	10576	10591	3177
561528	1606	1621	GAGAAATGTAAACGGT	89	10578	10593	174
561529	1608	1623	TTGAGAAATGTAAACG	55	10580	10595	3178
561530	1611	1626	TGATTGAGAAATGTAA	18	10583	10598	3179
561531	1613	1628	TTTGATTGAGAAATGT	30	10585	10600	3180
561532	1619	1634	AAGAATTTTGATTGAG	53	10591	10606	3181
561533	1621	1636	ATAAGAATTTTGATTG	29	10593	10608	3182
561534	1632	1647	CAAATAGTATTATAAG	6	10604	10619	3183
561535	1653	1668	CCCACATCACAAAATT	70	10625	10640	3184
561536	1657	1672	GATTCCCACATCACAA	77	10629	10644	3185
561537	1659	1674	TTGATTCCCACATCAC	78	10631	10646	3186
561538	1661	1676	AATTGATTCCCACATC	68	10633	10648	3187
561539	1663	1678	AAAATTGATTCCCACA	72	10635	10650	3188
561540	1665	1680	CTAAAATTGATTCCCA	54	10637	10652	3189
561541	1668	1683	CATCTAAAATTGATTC	0	10640	10655	3190
561542	1670	1685	ACCATCTAAAATTGAT	35	10642	10657	3191
561543	1672	1687	TGACCATCTAAAATTG	55	10644	10659	3192
561544	1674	1689	TGTGACCATCTAAAAT	56	10646	10661	3193
561545	1676	1691	ATTGTGACCATCTAAA	73	10648	10663	3194
561546	1678	1693	AGATTGTGACCATCTA	67	10650	10665	3195
561547	1680	1695	CTAGATTGTGACCATC	50	10652	10667	3196
561548	1682	1697	ATCTAGATTGTGACCA	77	10654	10669	3197
561549	1684	1699	TAATCTAGATTGTGAC	55	10656	10671	3198
561550	1686	1701	TATAATCTAGATTGTG	28	10658	10673	3199
561551	1688	1703	ATTATAATCTAGATTG	52	10660	10675	3200
561552	1690	1705	TGATTATAATCTAGAT	43	10662	10677	3201
561553	1692	1707	ATTGATTATAATCTAG	53	10664	10679	3202
561554	1694	1709	CTATTGATTATAATCT	54	10666	10681	3203
561555	1696	1711	ACCTATTGATTATAAT	44	10668	10683	3204
561556	1698	1713	TCACCTATTGATTATA	52	10670	10685	3205
561557	1700	1715	G TTCACCTATTGATTA	50	10672	10687	3206
561558	1702	1717	AAGTTCACCTATTGAT	58	10674	10689	3207
561559	1704	1719	ATAAGTTCACCTATTG	66	10676	10691	3208
561560	1706	1721	TAATAAGTTCACCTAT	38	10678	10693	3209
561561	1708	1723	TTTAATAAGTTCACCT	50	10680	10695	3210
561562	1710	1725	TATTTAATAAGTTCAC	32	10682	10697	3211
561563	1712	1727	GTTATTTAATAAGTTC	47	10684	10699	3212

561564	1761	1776	CATATGATGCCTTTTA	63	10733	10748	3213
561565	1763	1778	CTCATATGATGCCTTT	81	10735	10750	175
561566	1765	1780	AGCTCATATGATGCCT	81	10737	10752	176
561567	1767	1782	TTAGCTCATATGATGC	84	10739	10754	177
561568	1769	1784	TATTAGCTCATATGAT	46	10741	10756	3214
561569	1771	1786	GATATTAGCTCATATG	49	10743	10758	3215
561570	1773	1788	GTGATATTAGCTCATA	81	10745	10760	3216
561571	1775	1790	TTGTGATATTAGCTCA	85	10747	10762	178
561572	1777	1792	AGTTGTGATATTAGCT	68	10749	10764	3217
561573	1779	1794	AAAGTTGTGATATTAG	45	10751	10766	3218
561574	1781	1796	GGAAAGTTGTGATATT	27	10753	10768	3219
561575	1783	1798	TGGGAAAGTTGTGATA	36	10755	10770	3220
561576	1785	1800	ACTGGGAAAGTTGTGA	83	10757	10772	179
561577	1787	1802	AAACTGGGAAAGTTGT	56	10759	10774	3221
561578	1789	1804	TTAAACTGGGAAAGTT	44	10761	10776	3222
561579	1794	1809	GTTTTTTTAAACTGGGA	58	10766	10781	3223
561580	1796	1811	TAGTTTTTTTAAACTGG	0	10768	10783	3224
561581	1802	1817	GAGTACTAGTTTTTTTA	18	10774	10789	3225
561582	1804	1819	AAGAGTACTAGTTTTT	55	10776	10791	3226
561583	1806	1821	ACAAGAGTACTAGTTT	51	10778	10793	3227
561584	1808	1823	TAACAAGAGTACTAGT	53	10780	10795	3228
561585	1810	1825	TTTAACAAGAGTACTA	48	10782	10797	3229
561586	1812	1827	GTTTTTAACAAGAGTAC	49	10784	10799	3230
561587	1814	1829	GAGTTTTTAACAAGAGT	54	10786	10801	3231
561588	1816	1831	TAGAGTTTTTAACAAGA	9	10788	10803	3232
561589	1819	1834	GTTTAGAGTTTTTAACA	24	10791	10806	3233
561590	1822	1837	CAAGTTTAGAGTTTTTA	30	10794	10809	3234
561591	1824	1839	GTCAAGTTTAGAGTTT	60	10796	10811	3235
561592	1826	1841	TAGTCAAGTTTAGAGT	56	10798	10813	3236
561593	1828	1843	TTTAGTCAAGTTTAGA	41	10800	10815	3237
561594	1830	1845	TATTTAGTCAAGTTTA	14	10802	10817	3238
561595	1832	1847	TGTATTTAGTCAAGTT	39	10804	10819	3239
561596	1834	1849	TCTGTATTTAGTCAAG	51	10806	10821	3240
561597	1836	1851	CCTCTGTATTTAGTCA	72	10808	10823	3241
561598	1838	1853	GTCCTCTGTATTTAGT	55	10810	10825	3242
561599	1840	1855	CAGTCCTCTGTATTTA	63	10812	10827	3243
561600	1842	1857	ACCAGTCCTCTGTATT	66	10814	10829	3244
561601	1844	1859	TTACCAGTCCTCTGTA	57	10816	10831	3245
561602	1846	1861	AATTACCAGTCCTCTG	43	10818	10833	3246
561603	1848	1863	ACAATTACCAGTCCTC	67	10820	10835	3247

Table 25

Inhibition of ANGPTL3 mRNA by deoxy, MOE and (S)-cEt gapmers targeting SEQ ID NO: 1 and 2

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
561770	N/A	N/A	ACAAAGGTAGGTCACC	77	11576	11591	143
586719	N/A	N/A	TCTGACAAGATTCCTT	76	11167	11182	3248
586720	N/A	N/A	ATCTGACAAGATTCCT	79	11168	11183	3249
586721	N/A	N/A	TAATCTGACAAGATTC	50	11170	11185	3250
586722	N/A	N/A	GTAATCTGACAAGATT	41	11171	11186	3251
586723	N/A	N/A	CTTGTCAGGCTGTTTA	50	11312	11327	3252
586724	N/A	N/A	GCTTGTCAGGCTGTTT	81	11313	11328	3253
586725	N/A	N/A	ATGCTTGTCAGGCTGT	78	11315	11330	3254
586726	N/A	N/A	TACATGCTTGTCAGGC	78	11318	11333	3255
586727	N/A	N/A	ATACATGCTTGTCAGG	76	11319	11334	3256
586728	N/A	N/A	AAAGGTAGGTCACCAT	72	11574	11589	3257
586729	N/A	N/A	CAAAGGTAGGTCACCA	69	11575	11590	3258
586730	N/A	N/A	GACAAAGGTAGGTCAC	55	11577	11592	3259
586731	N/A	N/A	TGACAAAGGTAGGTCA	32	11578	11593	3260
586732	N/A	N/A	TCTGACATAGCTTTTT	63	5436	5451	3261
586733	N/A	N/A	ATTCTGACATAGCTTT	76	5438	5453	3262
586734	N/A	N/A	GATTCTGACATAGCTT	73	5439	5454	3263
586735	N/A	N/A	GGATTCTGACATAGCT	81	5440	5455	3264
586736	N/A	N/A	ATGGATTCTGACATAG	74	5442	5457	3265
586737	N/A	N/A	CATGGATTCTGACATA	72	5443	5458	3266
586738	N/A	N/A	ACATGGATTCTGACAT	59	5444	5459	3267
586739	N/A	N/A	TACATGGATTCTGACA	71	5445	5460	3268
586740	N/A	N/A	ATACATGGATTCTGAC	60	5446	5461	3269
586741	N/A	N/A	TTTAGCAGCACTACTA	65	5628	5643	3270
586742	N/A	N/A	TTTTAGCAGCACTACT	51	5629	5644	3271
586743	N/A	N/A	CTTTTAGCAGCACTAC	74	5630	5645	3272
586744	N/A	N/A	CCTTTTAGCAGCACTA	83	5631	5646	223
586745	N/A	N/A	ACCTTTTAGCAGCACT	84	5632	5647	224
586746	N/A	N/A	AAACCTTTTAGCAGCA	87	5634	5649	225
586747	N/A	N/A	AAAACCTTTTAGCAGC	80	5635	5650	3273
586748	N/A	N/A	GATAAAAAACCTTTTA	16	5640	5655	3274
586749	N/A	N/A	TGATAAAAAACCTTTT	25	5641	5656	3275
586750	N/A	N/A	AGATGTTGGCAGGTTG	72	6188	6203	3276
586751	N/A	N/A	TAGATGTTGGCAGGTT	76	6189	6204	3277
586752	N/A	N/A	GTAGATGTTGGCAGGT	73	6190	6205	3278

586753	N/A	N/A	TGTAGATGTTGGCAGG	65	6191	6206	3279
586754	N/A	N/A	CTGTAGATGTTGGCAG	61	6192	6207	3280
586755	N/A	N/A	ATCTGTAGATGTTGGC	84	6194	6209	226
586756	N/A	N/A	TATCTGTAGATGTTGG	71	6195	6210	3281
586757	N/A	N/A	ATATCTGTAGATGTTG	61	6196	6211	3282
586758	N/A	N/A	CATATCTGTAGATGTT	63	6197	6212	3283
586759	N/A	N/A	TTTGAACCAGGCTTTC	47	6243	6258	3284
586760	N/A	N/A	AATTTGAACCAGGCTT	78	6245	6260	3285
586761	N/A	N/A	TAATTTGAACCAGGCT	83	6246	6261	227
586762	N/A	N/A	CATAATTTGAACCAGG	81	6248	6263	3286
586763	N/A	N/A	ACATAATTTGAACCAG	36	6249	6264	3287
586764	N/A	N/A	TACATAATTTGAACCA	38	6250	6265	3288
586765	N/A	N/A	ATACATAATTTGAACC	15	6251	6266	3289
586766	N/A	N/A	ACATTGGTCGGAAAAC	43	6424	6439	3290
586767	N/A	N/A	GACATTGGTCGGAAAA	49	6425	6440	3291
586768	N/A	N/A	AGACATTGGTCGGAAA	59	6426	6441	3292
586769	N/A	N/A	CAGACATTGGTCGGAA	66	6427	6442	3293
586770	N/A	N/A	GCAGACATTGGTCGGA	80	6428	6443	3294
586771	N/A	N/A	AAGCAGACATTGGTCG	65	6430	6445	3295
586772	N/A	N/A	TGTACAGATTACCTGT	51	6506	6521	3296
586773	N/A	N/A	TTGTACAGATTACCTG	34	6507	6522	3297
586774	N/A	N/A	ATTGTACAGATTACCT	62	6508	6523	3298
586775	N/A	N/A	GATTGTACAGATTACC	59	6509	6524	3299
586776	N/A	N/A	AGATTGTACAGATTAC	46	6510	6525	3300
586777	N/A	N/A	TCAGATTGTACAGATT	63	6512	6527	3301
586778	N/A	N/A	TTCAGATTGTACAGAT	63	6513	6528	3302
586779	N/A	N/A	ATTCAGATTGTACAGA	71	6514	6529	3303
586780	N/A	N/A	TATTCAGATTGTACAG	55	6515	6530	3304
586781	N/A	N/A	TTATTCAGATTGTACA	52	6516	6531	3305
586782	N/A	N/A	TAGGTATGTCTTTTAT	52	6936	6951	3306
586783	N/A	N/A	TGTCTTAGGTATGTCT	76	6941	6956	3307
586784	N/A	N/A	ATTGTCTTAGGTATGT	73	6943	6958	3308
586785	N/A	N/A	GATTGTCTTAGGTATG	60	6944	6959	3309
586786	N/A	N/A	TTCTTAGATGGCGTGT	74	7207	7222	3310
586787	N/A	N/A	TTTTCTTAGATGGCGT	86	7209	7224	228
586788	N/A	N/A	ATTTTCTTAGATGGC	75	7211	7226	3311
586789	N/A	N/A	CATTTTCTTAGATGG	49	7212	7227	3312
586790	N/A	N/A	GCATTTTCTTAGATG	47	7213	7228	3313
586791	N/A	N/A	ATAAGTCCCAATTTTA	27	10066	10081	3314
586792	N/A	N/A	TATAAGTCCCAATTTT	27	10067	10082	3315
586793	N/A	N/A	GTATAAGTCCCAATTT	28	10068	10083	3316

586794	N/A	N/A	TGTATAAGTCCCAATT	38	10069	10084	3317
586795	N/A	N/A	CTGTATAAGTCCCAAT	69	10070	10085	3318
586796	N/A	N/A	ATCTGTATAAGTCCCA	88	10072	10087	229
586797	N/A	N/A	AATCTGTATAAGTCCC	84	10073	10088	230
586798	N/A	N/A	TAATCTGTATAAGTCC	58	10074	10089	3319
586799	N/A	N/A	ATAATCTGTATAAGTC	21	10075	10090	3320
586800	N/A	N/A	AATAATCTGTATAAGT	12	10076	10091	3321
586801	N/A	N/A	TGCATGTATCCCAGTT	80	10095	10110	3322
586802	N/A	N/A	ATGCATGTATCCCAGT	83	10096	10111	231
586803	N/A	N/A	AGATGCATGTATCCCA	79	10098	10113	232
586804	N/A	N/A	TAGATGCATGTATCCC	87	10099	10114	3323
586805	N/A	N/A	TTAGATGCATGTATCC	78	10100	10115	3324
586806	N/A	N/A	TTTAGATGCATGTATC	50	10101	10116	3325
586653	7	22	GTGGAAGTGTCTTCTT	63	3111	3126	3326
586656	9	24	ACGTGGAAGTGTCTTC	72	3113	3128	3327
586658	99	114	TTGATCAATTCTGGAG	74	3203	3218	3328
586660	101	116	TCTTGATCAATTCTGG	71	3205	3220	3329
561011	102	117	GTCTTGATCAATTCTG	91	3206	3221	114
586661	103	118	TGTCTTGATCAATTCT	85	3207	3222	209
586663	134	149	GGCTCTGGAGATAGAG	63	3238	3253	3330
586665	136	151	TTGGCTCTGGAGATAG	63	3240	3255	3331
586668	140	155	GATTTTGGCTCTGGAG	64	3244	3259	3332
586669	142	157	TTGATTTTGGCTCTGG	89	3246	3261	210
561026	143	158	CTTGATTTTGGCTCTG	84	3247	3262	117
586670	144	159	TCTTGATTTTGGCTCT	71	3248	3263	3333
586671	146	161	AATCTTGATTTTGGCT	70	3250	3265	3334
586672	148	163	CAAATCTTGATTTTGG	81	3252	3267	3335
586673	298	313	GCAGCGATAGATCATA	76	3402	3417	3336
586674	300	315	TTGCAGCGATAGATCA	76	3404	3419	3337
586675	304	319	TGGTTTGCAGCGATAG	82	3408	3423	3338
586676	306	321	ACTGGTTTGCAGCGAT	89	3410	3425	211
586677	315	330	TTTGATTTCACTGGTT	62	3419	3434	3339
586678	317	332	TCTTTGATTTCACTGG	66	3421	3436	3340
586679	342	357	AGTTCTTCTCAGTTCC	77	3446	3461	3341
586680	476	491	TTAGTTAGTTGCTCTT	65	3580	3595	3342
586681	478	493	AGTTAGTTAGTTGCTC	69	3582	3597	3343
586682	703	718	GTGCTCTTGGCTTGGA	78	6716	6731	3344
586683	705	720	TGGTGCTCTTGGCTTG	77	6718	6733	3345
586684	802	817	TATGTTACCTCTGTT	55	7387	7402	3346
586685	804	819	TGTATGTTACCTCTG	79	7389	7404	3347
586686	1260	1275	ACACTCATCATGCCAC	72	10232	10247	3348

586687	1262	1277	CCACACTCATCATGCC	82	10234	10249	3349
586688	1308	1323	AGATTTTGCTCTTGGT	87	10280	10295	212
586689	1310	1325	TTAGATTTTGCTCTTG	78	10282	10297	3350
586690	1351	1366	CATTTTGAGACTTCCA	91	10323	10338	213
586691	1353	1368	TCCATTTTGAGACTTC	86	10325	10340	214
586692	1365	1380	AGAGTATAACCTTCCA	88	10337	10352	220
586693	1367	1382	ATAGAGTATAACCTTC	69	10339	10354	3351
586694	1402	1417	AATCTGTTGGATGGAT	59	10374	10389	3352
586695	1404	1419	TGAATCTGTTGGATGG	79	10376	10391	3353
586696	1420	1435	TTCATTCAAAGCTTTC	82	10392	10407	3354
586697	1422	1437	AGTTCATTCAAAGCTT	73	10394	10409	3355
561463	1423	1438	CAGTTCATTCAAAGCT	88	10395	10410	127
586698	1424	1439	TCAGTTCATTCAAAGC	69	10396	10411	3356
586699	1488	1503	GATTATTAGACCACAT	63	10460	10475	3357
586700	1490	1505	CAGATTATTAGACCAC	90	10462	10477	221
561487	1491	1506	CCAGATTATTAGACCA	95	10463	10478	131
586701	1492	1507	ACCAGATTATTAGACC	85	10464	10479	215
586702	1552	1567	TAGACAGTGACTTTAA	83	10524	10539	216
586703	1554	1569	AATAGACAGTGACTTT	70	10526	10541	3358
586704	1605	1620	AGAAATGTAAACGGTA	76	10577	10592	3359
586705	1607	1622	TGAGAAATGTAAACGG	83	10579	10594	217
586706	1762	1777	TCATATGATGCCTTTT	69	10734	10749	3360
586707	1764	1779	GCTCATATGATGCCTT	84	10736	10751	218
586708	1766	1781	TAGCTCATATGATGCC	83	10738	10753	222
561567	1767	1782	TTAGCTCATATGATGC	81	10739	10754	177
586709	1768	1783	ATTAGCTCATATGATG	40	10740	10755	3361
586710	1774	1789	TGTGATATTAGCTCAT	73	10746	10761	3362
586711	1776	1791	GTTGTGATATTAGCTC	80	10748	10763	3363
586712	1905	1920	TACTCTGTGCTGACGA	81	10877	10892	3364
586713	1907	1922	CATACTCTGTGCTGAC	81	10879	10894	3365
586714	2052	2067	GTTTAAAGACAGCGAA	72	11024	11039	3366
586715	2054	2069	TTGTTTAAAGACAGCG	81	11026	11041	3367
586716	2068	2083	GTAGTCATCTCCATTT	63	11040	11055	3368
586717	2070	2085	TAGTAGTCATCTCCAT	74	11042	11057	3369
561650	2071	2086	TTAGTAGTCATCTCCA	79	11043	11058	142
586718	2072	2087	CTTAGTAGTCATCTCC	84	11044	11059	219

Example 5: Antisense inhibition of human ANGPTL3 in Hep3B cells by deoxy, MOE and (S)-cEt gapmers

Additional antisense oligonucleotides were designed targeting an ANGPTL3 nucleic acid and were tested for their effects on ANGPTL3 mRNA in vitro. ISIS 337487 and ISIS 233717, which are 5-10-5 MOE gapmers, were also included in the assay as benchmark oligonucleotides. Cultured Hep3B cells at a density of 20,000 cells per well were transfected using electroporation with 4,500 nM antisense oligonucleotide. After a treatment period of approximately 24 hours, RNA was isolated from the cells and ANGPTL3 mRNA levels were measured by quantitative real-time PCR. Human primer probe set RTS3492_MGB was used to measure mRNA levels. ANGPTL3 mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN®. Results are presented as percent inhibition of ANGPTL3, relative to untreated control cells.

The newly designed chimeric antisense oligonucleotides in the Tables below were designed as deoxy, MOE, and (S)-cEt oligonucleotides or 5-10-5 MOE gapmers. The deoxy, MOE and (S)-cEt oligonucleotides are 16 nucleosides in length wherein the nucleoside have either a MOE sugar modification, an (S)-cEt sugar modification, or a deoxy sugar residue. The sugar modifications of each antisense oligonucleotide is described as ‘eek-d10-kke’, where ‘k’ indicates an (S)-cEt sugar modification; ‘d’ indicates deoxyribose; the number indicates the number of deoxyribose sugars residues; and ‘e’ indicates a MOE modification. The 5-10-5 MOE 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. The internucleoside linkages throughout each oligonucleotide are phosphorothioate (P=S) linkages. All cytosine residues throughout each oligonucleotide are 5-methylcytosines. “Start site” indicates the 5’-most nucleoside to which the oligonucleotide is targeted in the human gene sequence. “Stop site” indicates the 3’-most nucleoside to which the oligonucleotide is targeted human gene sequence. Each oligonucleotide listed in the Tables below is targeted to either the human ANGPTL3 mRNA, designated herein as SEQ ID NO: 1 (GENBANK Accession No. NM_014495.2) or the human ANGPTL3 genomic sequence, designated herein as SEQ ID NO: 2 (GENBANK Accession No. NT_032977.9 truncated from nucleotides 33032001 to 33046000). ‘n/a’ indicates that the antisense oligonucleotide does not target that particular gene sequence with 100% complementarity.

Table 26
Inhibition of ANGPTL3 mRNA by oligonucleotides targeting SEQ ID NO: 1 and 2

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	Chemistry	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
561671	N/A	N/A	TCTTAACTCTATATAT	Deoxy, MOE, and cEt	12	3076	3091	3370
561672	N/A	N/A	CTTCTTAACTCTATAT	Deoxy, MOE, and cEt	12	3078	3093	3371
561673	N/A	N/A	GACTTCTTAACTCTAT	Deoxy, MOE, and cEt	18	3080	3095	3372
561674	N/A	N/A	TAGACTTCTTAACTCT	Deoxy, MOE, and cEt	20	3082	3097	3373
561675	N/A	N/A	CCTAGACTTCTTAACT	Deoxy, MOE, and cEt	9	3084	3099	3374

561676	N/A	N/A	GACCTAGACTTCTTAA	Deoxy, MOE, and cEt	0	3086	3101	3375
561677	N/A	N/A	CAGACCTAGACTTCTT	Deoxy, MOE, and cEt	18	3088	3103	3376
561678	N/A	N/A	AGCAGACCTAGACTTC	Deoxy, MOE, and cEt	26	3090	3105	3377
561679	N/A	N/A	GAAGCAGACCTAGACT	Deoxy, MOE, and cEt	24	3092	3107	3378
561680	N/A	N/A	TGGAAGCAGACCTAGA	Deoxy, MOE, and cEt	30	3094	3109	3379
561758	N/A	N/A	CTTTAACAAATGGGTT	Deoxy, MOE, and cEt	25	11539	11554	3380
561759	N/A	N/A	ATCCTTTAACAAATGG	Deoxy, MOE, and cEt	31	11542	11557	3381
561760	N/A	N/A	CTATATCCTTTAACAA	Deoxy, MOE, and cEt	28	11546	11561	3382
561761	N/A	N/A	GCACTATATCCTTTAA	Deoxy, MOE, and cEt	59	11549	11564	3383
561762	N/A	N/A	TGGGCACTATATCCTT	Deoxy, MOE, and cEt	34	11552	11567	3384
561763	N/A	N/A	ACTTGGGCACTATATC	Deoxy, MOE, and cEt	30	11555	11570	3385
561764	N/A	N/A	ATAACTTGGGCACTAT	Deoxy, MOE, and cEt	51	11558	11573	3386
561765	N/A	N/A	CATATAACTTGGGCAC	Deoxy, MOE, and cEt	47	11561	11576	3387
561766	N/A	N/A	CACCATATAACTTGGG	Deoxy, MOE, and cEt	47	11564	11579	3388
561767	N/A	N/A	GGTCACCATATAACTT	Deoxy, MOE, and cEt	58	11567	11582	3389
561768	N/A	N/A	GTAGGTCACCATATAA	Deoxy, MOE, and cEt	62	11570	11585	3390
561769	N/A	N/A	AAGGTAGGTCACCATA	Deoxy, MOE, and cEt	65	11573	11588	3391
561770	N/A	N/A	ACAAAGGTAGGTCACC	Deoxy, MOE, and cEt	73	11576	11591	143
561771	N/A	N/A	TTGACAAAGGTAGGTC	Deoxy, MOE, and cEt	70	11579	11594	3392
561772	N/A	N/A	GTATTGACAAAGGTAG	Deoxy, MOE, and cEt	58	11582	11597	3393
561773	N/A	N/A	TAAGTATTGACAAAGG	Deoxy, MOE, and cEt	42	11585	11600	3394
561774	N/A	N/A	TGCTAAGTATTGACAA	Deoxy, MOE, and cEt	51	11588	11603	3395
561775	N/A	N/A	TAATGCTAAGTATTGA	Deoxy, MOE, and cEt	42	11591	11606	3396
561776	N/A	N/A	TACATAATGCTAAGTA	Deoxy, MOE, and cEt	36	11595	11610	3397
561777	N/A	N/A	GGATAATTTGAAATAC	Deoxy, MOE, and cEt	24	11608	11623	3398
561778	N/A	N/A	TATTGGATAATTTGAA	Deoxy, MOE, and cEt	35	11612	11627	3399
561779	N/A	N/A	GTATATTGGATAATTT	Deoxy, MOE, and cEt	0	11615	11630	3400
561780	N/A	N/A	CATGTATATTGGATAA	Deoxy, MOE, and cEt	20	11618	11633	3401
561781	N/A	N/A	TGACATGTATATTGGA	Deoxy, MOE, and cEt	73	11621	11636	144
561782	N/A	N/A	CTTTTATATATGTGAC	Deoxy, MOE, and cEt	37	11652	11667	3402
561783	N/A	N/A	GATCATACATATCTTT	Deoxy, MOE, and cEt	51	11664	11679	3403
561784	N/A	N/A	ATAGATCATACATATC	Deoxy, MOE, and cEt	46	11667	11682	3404
561785	N/A	N/A	CACATAGATCATACAT	Deoxy, MOE, and cEt	65	11670	11685	3405
561786	N/A	N/A	ATTCACATAGATCATA	Deoxy, MOE, and cEt	48	11673	11688	3406
561787	N/A	N/A	AGGATTCACATAGATC	Deoxy, MOE, and cEt	48	11676	11691	3407
561788	N/A	N/A	CTTAGGATTCACATAG	Deoxy, MOE, and cEt	42	11679	11694	3408
561789	N/A	N/A	TTACTTAGGATTCACA	Deoxy, MOE, and cEt	58	11682	11697	3409
561790	N/A	N/A	TATTTACTTAGGATTC	Deoxy, MOE, and cEt	45	11685	11700	3410
561791	N/A	N/A	GTACTTTTCTGGAACA	Deoxy, MOE, and cEt	77	11704	11719	145
561792	N/A	N/A	CCTGAAAATTATAGAT	Deoxy, MOE, and cEt	35	11741	11756	3411
561793	N/A	N/A	GGTCCTGAAAATTATA	Deoxy, MOE, and cEt	32	11744	11759	3412

561794	N/A	N/A	TGTGGTCCTGAAAATT	Deoxy, MOE, and cEt	45	11747	11762	3413
561795	N/A	N/A	GTCTGTGGTCCTGAAA	Deoxy, MOE, and cEt	47	11750	11765	3414
561796	N/A	N/A	TTAGTCTGTGGTCCTG	Deoxy, MOE, and cEt	67	11753	11768	3415
561797	N/A	N/A	AGCTTAGTCTGTGGTC	Deoxy, MOE, and cEt	55	11756	11771	3416
561798	N/A	N/A	GACAGCTTAGTCTGTG	Deoxy, MOE, and cEt	47	11759	11774	3417
561799	N/A	N/A	TTCGACAGCTTAGTCT	Deoxy, MOE, and cEt	68	11762	11777	3418
561800	N/A	N/A	AATTTTCGACAGCTTAG	Deoxy, MOE, and cEt	61	11765	11780	3419
561801	N/A	N/A	GTTAATTTTCGACAGCT	Deoxy, MOE, and cEt	70	11768	11783	3420
561802	N/A	N/A	CCTAAAAAAATCAGCG	Deoxy, MOE, and cEt	19	11783	11798	3421
561803	N/A	N/A	GGCCCTAAAAAAATCA	Deoxy, MOE, and cEt	0	11786	11801	3422
561804	N/A	N/A	TTCTGGCCCTAAAAAA	Deoxy, MOE, and cEt	10	11790	11805	3423
561805	N/A	N/A	GTATTCTGGCCCTAAA	Deoxy, MOE, and cEt	44	11793	11808	3424
561806	N/A	N/A	TTGGTATTCTGGCCCT	Deoxy, MOE, and cEt	45	11796	11811	3425
561807	N/A	N/A	ATTTTGGTATTCTGGC	Deoxy, MOE, and cEt	59	11799	11814	3426
561808	N/A	N/A	GCCATTTTGGTATTCT	Deoxy, MOE, and cEt	58	11802	11817	3427
561809	N/A	N/A	GGAGCCATTTTGGTAT	Deoxy, MOE, and cEt	33	11805	11820	3428
561810	N/A	N/A	AGAGGAGCCATTTTGG	Deoxy, MOE, and cEt	36	11808	11823	3429
561811	N/A	N/A	AAGAGAGGAGCCATTT	Deoxy, MOE, and cEt	14	11811	11826	3430
561812	N/A	N/A	ATTGTCCAATTTTGGG	Deoxy, MOE, and cEt	25	11829	11844	3431
561813	N/A	N/A	GAAATTGTCCAATTTT	Deoxy, MOE, and cEt	38	11832	11847	3432
561814	N/A	N/A	TTTGAAATTGTCCAAT	Deoxy, MOE, and cEt	36	11835	11850	3433
561815	N/A	N/A	GCATTTGAAATTGTCC	Deoxy, MOE, and cEt	67	11838	11853	3434
561816	N/A	N/A	GCAACTCATATATTAA	Deoxy, MOE, and cEt	57	11869	11884	3435
561817	N/A	N/A	GAAGCAACTCATATAT	Deoxy, MOE, and cEt	46	11872	11887	3436
561818	N/A	N/A	GAGGAAGCAACTCATA	Deoxy, MOE, and cEt	14	11875	11890	3437
561819	N/A	N/A	ATAGAGGAAGCAACTC	Deoxy, MOE, and cEt	60	11878	11893	3438
561820	N/A	N/A	CAAATAGAGGAAGCAA	Deoxy, MOE, and cEt	36	11881	11896	3439
561821	N/A	N/A	AACCAAATAGAGGAAG	Deoxy, MOE, and cEt	38	11884	11899	3440
561822	N/A	N/A	GGAAACCAAATAGAGG	Deoxy, MOE, and cEt	51	11887	11902	3441
561823	N/A	N/A	CTTTAAGTGAAGTTAC	Deoxy, MOE, and cEt	30	3636	3651	3442
561824	N/A	N/A	TACTTACTTTAAGTGA	Deoxy, MOE, and cEt	27	3642	3657	3443
561825	N/A	N/A	GAACCCTCTTTATTTT	Deoxy, MOE, and cEt	25	3659	3674	3444
561826	N/A	N/A	AAACATGAACCCTCTT	Deoxy, MOE, and cEt	14	3665	3680	3445
561827	N/A	N/A	GATCCACATTGAAAAC	Deoxy, MOE, and cEt	0	3683	3698	3446
561828	N/A	N/A	CATGCCTTAGAAATAT	Deoxy, MOE, and cEt	33	3710	3725	3447
561829	N/A	N/A	AAATGGCATGCCTTAG	Deoxy, MOE, and cEt	46	3716	3731	3448
561830	N/A	N/A	GTATTTCAAATGGCAT	Deoxy, MOE, and cEt	54	3723	3738	3449
561831	N/A	N/A	GCAACAAAGTATTTCA	Deoxy, MOE, and cEt	60	3731	3746	3450
561832	N/A	N/A	GTATTTCAACAATGCA	Deoxy, MOE, and cEt	28	3744	3759	3451
561833	N/A	N/A	ATAACATTAGGGAAAC	Deoxy, MOE, and cEt	18	3827	3842	3452
561834	N/A	N/A	TCATATATAACATTAG	Deoxy, MOE, and cEt	18	3833	3848	3453

561912	N/A	N/A	GTGGTTTTGAGCAAAG	Deoxy, MOE, and cEt	5	4736	4751	3454
561913	N/A	N/A	CTATTGTGTGGTTTTG	Deoxy, MOE, and cEt	36	4743	4758	3455
561914	N/A	N/A	GGAAAGCTATTGTGTG	Deoxy, MOE, and cEt	18	4749	4764	3456
561915	N/A	N/A	TATGAGTGAAATGGAA	Deoxy, MOE, and cEt	13	4761	4776	3457
561916	N/A	N/A	AGCCAATATGAGTGAA	Deoxy, MOE, and cEt	57	4767	4782	3458
561917	N/A	N/A	CTAAAGAGCCAATATG	Deoxy, MOE, and cEt	33	4773	4788	3459
561918	N/A	N/A	CTTGGTCTAAAGAGCC	Deoxy, MOE, and cEt	70	4779	4794	146
561919	N/A	N/A	GGTAATCTTGGTCTAA	Deoxy, MOE, and cEt	46	4785	4800	3460
561920	N/A	N/A	GATGACGAAGGGTTGG	Deoxy, MOE, and cEt	28	4800	4815	3461
561921	N/A	N/A	CAGTGAGATGACGAAG	Deoxy, MOE, and cEt	39	4806	4821	3462
561922	N/A	N/A	TGAAGTCAGTGAGATG	Deoxy, MOE, and cEt	49	4812	4827	3463
561923	N/A	N/A	AGGAGGTGAAGTCAGT	Deoxy, MOE, and cEt	35	4818	4833	3464
561924	N/A	N/A	GAGTAGAGGAGGTGAA	Deoxy, MOE, and cEt	33	4824	4839	3465
561925	N/A	N/A	TAAGTAGAGTAGAGGA	Deoxy, MOE, and cEt	35	4830	4845	3466
561926	N/A	N/A	TCAGAATAACTAGAGT	Deoxy, MOE, and cEt	24	4836	4851	3467
561927	N/A	N/A	AAGCGGTCAGAATAAC	Deoxy, MOE, and cEt	39	4842	4857	3468
561928	N/A	N/A	CTGGTAAAGCGGTCAG	Deoxy, MOE, and cEt	51	4848	4863	3469
561929	N/A	N/A	TGAATACTGGTAAAGC	Deoxy, MOE, and cEt	63	4854	4869	3470
561930	N/A	N/A	TGTGTTTGAATACTGG	Deoxy, MOE, and cEt	65	4860	4875	3471
561931	N/A	N/A	GTTTGATGTGTTTGAA	Deoxy, MOE, and cEt	49	4866	4881	3472
561932	N/A	N/A	CAGTATGTTTGATGTG	Deoxy, MOE, and cEt	48	4872	4887	3473
561933	N/A	N/A	AGGTGGCAGTATGTTT	Deoxy, MOE, and cEt	0	4878	4893	3474
561934	N/A	N/A	GCTTTGAGGTGGCAGT	Deoxy, MOE, and cEt	48	4884	4899	3475
561935	N/A	N/A	GGGCAAAGGCTTTGAG	Deoxy, MOE, and cEt	28	4892	4907	3476
561936	N/A	N/A	CAACAAGGGCAAAGGC	Deoxy, MOE, and cEt	65	4898	4913	3477
561937	N/A	N/A	GAGGAAACAACAAGGG	Deoxy, MOE, and cEt	42	4905	4920	3478
561938	N/A	N/A	CCAGTTAGAGGAAACA	Deoxy, MOE, and cEt	52	4912	4927	3479
561939	N/A	N/A	CCAGGGCAGAAGAGCG	Deoxy, MOE, and cEt	61	4930	4945	3480
561940	N/A	N/A	TAGATACCAGGGCAGA	Deoxy, MOE, and cEt	68	4936	4951	3481
561941	N/A	N/A	CAGAGAGTGGGCCACG	Deoxy, MOE, and cEt	46	4952	4967	3482
561942	N/A	N/A	GGAAATCAGAGAGTGG	Deoxy, MOE, and cEt	42	4958	4973	3483
561943	N/A	N/A	CCTAAGGGGAAATCAGA	Deoxy, MOE, and cEt	26	4964	4979	3484
561944	N/A	N/A	AACGACCCTAAGGGAA	Deoxy, MOE, and cEt	45	4970	4985	3485
561945	N/A	N/A	TTTGATAACGACCCTA	Deoxy, MOE, and cEt	57	4976	4991	3486
561946	N/A	N/A	TTTTTGTTTGATAACG	Deoxy, MOE, and cEt	21	4982	4997	3487
561947	N/A	N/A	CATTGGGAATTTTTTG	Deoxy, MOE, and cEt	35	4992	5007	3488
561948	N/A	N/A	AGTCTTCATTGGGAAT	Deoxy, MOE, and cEt	69	4998	5013	3489
561949	N/A	N/A	CTTGTAAGTCTTCATT	Deoxy, MOE, and cEt	35	5004	5019	3490
561950	N/A	N/A	AGTGACCTTGTAAGTC	Deoxy, MOE, and cEt	56	5010	5025	3491
561951	N/A	N/A	TGGTTAAGTGACCTTG	Deoxy, MOE, and cEt	67	5016	5031	3492
561952	N/A	N/A	GATTTTTGGTTAAGTG	Deoxy, MOE, and cEt	43	5022	5037	3493

561953	N/A	N/A	GGTTGTGATTTTTGGT	Deoxy, MOE, and cEt	58	5028	5043	3494
561954	N/A	N/A	CCAGGCGGTTGTGATT	Deoxy, MOE, and cEt	49	5034	5049	3495
561955	N/A	N/A	ATGGGACCAGGCGGTT	Deoxy, MOE, and cEt	52	5040	5055	3496
561956	N/A	N/A	AAGTTTTTCAGGGATGG	Deoxy, MOE, and cEt	49	5052	5067	3497
561957	N/A	N/A	AAGTAGAAGTTTTTCAG	Deoxy, MOE, and cEt	16	5058	5073	3498
561958	N/A	N/A	CTAAGGAAGTAGAAGT	Deoxy, MOE, and cEt	33	5064	5079	3499
561959	N/A	N/A	AAGTAGCTAAGGAAGT	Deoxy, MOE, and cEt	35	5070	5085	3500
561960	N/A	N/A	GGAGAAAAGTAGCTAA	Deoxy, MOE, and cEt	36	5076	5091	3501
561961	N/A	N/A	TGTGCAGGAGAAAAGT	Deoxy, MOE, and cEt	53	5082	5097	3502
561962	N/A	N/A	GGTGAGTGTGCAGGAG	Deoxy, MOE, and cEt	44	5088	5103	3503
561963	N/A	N/A	AATAAAGGTGAGTGTG	Deoxy, MOE, and cEt	38	5094	5109	3504
561964	N/A	N/A	TGCAGGAATAGAAGAG	Deoxy, MOE, and cEt	58	5138	5153	3505
561965	N/A	N/A	TTTGTAGTGCAGGAATA	Deoxy, MOE, and cEt	20	5144	5159	3506
561966	N/A	N/A	TATTCACAGAGCTTAC	Deoxy, MOE, and cEt	63	5161	5176	3507
561967	N/A	N/A	TCCCTGTATTCACAGA	Deoxy, MOE, and cEt	61	5167	5182	3508
561968	N/A	N/A	GAAAAAATCCCTGTAT	Deoxy, MOE, and cEt	22	5174	5189	3509
561969	N/A	N/A	TATGAAGATAATGGAA	Deoxy, MOE, and cEt	34	5187	5202	3510
561970	N/A	N/A	GGAGTATATACAAATA	Deoxy, MOE, and cEt	46	5211	5226	3511
561971	N/A	N/A	TATTCTGGAGTATATA	Deoxy, MOE, and cEt	29	5217	5232	3512
561972	N/A	N/A	ATTCTATATTCTGGAG	Deoxy, MOE, and cEt	58	5223	5238	3513
561973	N/A	N/A	CATACAGTATTCTATA	Deoxy, MOE, and cEt	39	5231	5246	3514
561974	N/A	N/A	GTGTGCCATACAGTAT	Deoxy, MOE, and cEt	48	5237	5252	3515
561975	N/A	N/A	AGAAATGCCTACTGTG	Deoxy, MOE, and cEt	34	5250	5265	3516
561976	N/A	N/A	ATTCAACAGAAATGCC	Deoxy, MOE, and cEt	52	5257	5272	3517
561977	N/A	N/A	GAATATGACATTACAT	Deoxy, MOE, and cEt	33	5279	5294	3518
561978	N/A	N/A	CTGTGTGAATATGACA	Deoxy, MOE, and cEt	63	5285	5300	3519
561979	N/A	N/A	ACGCTTCTGTGTGAAT	Deoxy, MOE, and cEt	59	5291	5306	3520
561980	N/A	N/A	TAGCACACGCTTCTGT	Deoxy, MOE, and cEt	29	5297	5312	3521
561981	N/A	N/A	TAATCATAGCACACGC	Deoxy, MOE, and cEt	64	5303	5318	3522
561982	N/A	N/A	CCAAGTAATAATAATC	Deoxy, MOE, and cEt	26	5314	5329	3523
561983	N/A	N/A	AGTAATCCAAGTAATA	Deoxy, MOE, and cEt	33	5320	5335	3524
561984	N/A	N/A	ATTTCTAGTAATCCAA	Deoxy, MOE, and cEt	42	5326	5341	3525
561985	N/A	N/A	CACACTATTTCTAGTA	Deoxy, MOE, and cEt	40	5332	5347	3526
561986	N/A	N/A	ATGAGGCACACTATTT	Deoxy, MOE, and cEt	47	5338	5353	3527
561987	N/A	N/A	TTAATTATGAGGCACA	Deoxy, MOE, and cEt	58	5344	5359	3528
561988	N/A	N/A	TGACCTTTAATTATGA	Deoxy, MOE, and cEt	38	5350	5365	3529
562066	N/A	N/A	GCAATTTATTGAATGA	Deoxy, MOE, and cEt	27	6083	6098	3530
562067	N/A	N/A	GGGTTTGCAATTTATT	Deoxy, MOE, and cEt	38	6089	6104	3531
562068	N/A	N/A	TGTGTTGGGTTTGCAA	Deoxy, MOE, and cEt	43	6095	6110	3532
562069	N/A	N/A	TTTAAGTGTGTTGGGT	Deoxy, MOE, and cEt	71	6101	6116	3533
562070	N/A	N/A	GTTTAGCAGTAACATT	Deoxy, MOE, and cEt	38	6126	6141	3534

562071	N/A	N/A	ATTCAGTAGTTTATCG	Deoxy, MOE, and cEt	17	6145	6160	3535
562072	N/A	N/A	CTATATATTCAGTAGT	Deoxy, MOE, and cEt	0	6151	6166	3536
562073	N/A	N/A	GCTTACTTTCTATATA	Deoxy, MOE, and cEt	21	6160	6175	3537
562074	N/A	N/A	AGTTTGTTTGCTTACT	Deoxy, MOE, and cEt	63	6169	6184	3538
562075	N/A	N/A	TTGGCAAGTTTGTTTG	Deoxy, MOE, and cEt	55	6175	6190	3539
562076	N/A	N/A	GGCAGGTTGGCAAGTT	Deoxy, MOE, and cEt	68	6181	6196	3540
562077	N/A	N/A	GATGTTGGCAGGTTGG	Deoxy, MOE, and cEt	54	6187	6202	3541
562078	N/A	N/A	TCTGTAGATGTTGGCA	Deoxy, MOE, and cEt	81	6193	6208	147
562079	N/A	N/A	AACATATCTGTAGATG	Deoxy, MOE, and cEt	32	6199	6214	3542
562080	N/A	N/A	CCTGTAAACATATCTG	Deoxy, MOE, and cEt	51	6205	6220	3543
562081	N/A	N/A	TTTTGACCTGTAAACA	Deoxy, MOE, and cEt	14	6211	6226	3544
562082	N/A	N/A	GATAATTTTGGACCTG	Deoxy, MOE, and cEt	49	6217	6232	3545
562083	N/A	N/A	TCTTGATAATTTGATA	Deoxy, MOE, and cEt	13	6229	6244	3546
562084	N/A	N/A	AGGCTTTCTTGATAAT	Deoxy, MOE, and cEt	55	6235	6250	3547
562085	N/A	N/A	TGAACCAGGCTTTCTT	Deoxy, MOE, and cEt	74	6241	6256	3548
562086	N/A	N/A	ATAATTTGAACCAGGC	Deoxy, MOE, and cEt	82	6247	6262	148
562087	N/A	N/A	GATAAAGACATAATAC	Deoxy, MOE, and cEt	21	6263	6278	3549
562088	N/A	N/A	ACCTGTGATAAAGACA	Deoxy, MOE, and cEt	27	6269	6284	3550
562089	N/A	N/A	CTTCAGACCTGTGATA	Deoxy, MOE, and cEt	23	6275	6290	3551
562090	N/A	N/A	ACTGATCTTCAGACCT	Deoxy, MOE, and cEt	48	6281	6296	3552
562091	N/A	N/A	GGTCTTACTGATCTTC	Deoxy, MOE, and cEt	59	6287	6302	3553
562092	N/A	N/A	GTTTTAGGTCTTACTG	Deoxy, MOE, and cEt	21	6293	6308	3554
562093	N/A	N/A	GTTCAGATTTTAAGTT	Deoxy, MOE, and cEt	31	6321	6336	3555
562094	N/A	N/A	ATATTCTGTTCAGATT	Deoxy, MOE, and cEt	36	6328	6343	3556
562095	N/A	N/A	ATATTGTAATGTATTC	Deoxy, MOE, and cEt	52	6372	6387	3557
562096	N/A	N/A	CTTAGAATATTGTAAT	Deoxy, MOE, and cEt	13	6378	6393	3558
562097	N/A	N/A	GCTTTGCTTAGAATAT	Deoxy, MOE, and cEt	47	6384	6399	3559
562098	N/A	N/A	GAGACTGCTTTGCTTA	Deoxy, MOE, and cEt	48	6390	6405	3560
562099	N/A	N/A	AAAGTAGAGACTGCTT	Deoxy, MOE, and cEt	44	6396	6411	3561
562100	N/A	N/A	AGGCCAAAAGTAGAGA	Deoxy, MOE, and cEt	59	6402	6417	3562
562101	N/A	N/A	TCGGAAAACAGAGCAA	Deoxy, MOE, and cEt	63	6417	6432	3563
562102	N/A	N/A	CATTGGTCGGAAAACA	Deoxy, MOE, and cEt	53	6423	6438	3564
562103	N/A	N/A	AGCAGACATTGGTCGG	Deoxy, MOE, and cEt	83	6429	6444	149
562104	N/A	N/A	AGCAAGGCAAAAAAGC	Deoxy, MOE, and cEt	22	6442	6457	3565
562105	N/A	N/A	GACATTATTTAATAAG	Deoxy, MOE, and cEt	21	6470	6485	3566
562106	N/A	N/A	ATCAGGGACATTATTT	Deoxy, MOE, and cEt	34	6476	6491	3567
562107	N/A	N/A	TATTTAATCAGGGACA	Deoxy, MOE, and cEt	47	6482	6497	3568
562108	N/A	N/A	ATTACCTGTTCTCAAA	Deoxy, MOE, and cEt	30	6499	6514	3569
562109	N/A	N/A	GTACAGATTACCTGTT	Deoxy, MOE, and cEt	38	6505	6520	3570
562110	N/A	N/A	CAGATTGTACAGATTA	Deoxy, MOE, and cEt	76	6511	6526	150
562111	N/A	N/A	GTTATTCAGATTGTAC	Deoxy, MOE, and cEt	32	6517	6532	3571

562112	N/A	N/A	AACAGTGTTATTCAGA	Deoxy, MOE, and cEt	58	6523	6538	3572
562113	N/A	N/A	TAGATAAACAGTGTTA	Deoxy, MOE, and cEt	33	6529	6544	3573
562114	N/A	N/A	TGATATTTAGATAAAC	Deoxy, MOE, and cEt	26	6536	6551	3574
562115	N/A	N/A	GGTGTTTGATATTTAG	Deoxy, MOE, and cEt	60	6542	6557	3575
562116	N/A	N/A	TATAACGGTGTTTGAT	Deoxy, MOE, and cEt	42	6548	6563	3576
562117	N/A	N/A	TAATGTTATAACGGTG	Deoxy, MOE, and cEt	62	6554	6569	3577
562118	N/A	N/A	AGTTCATAATGTTATA	Deoxy, MOE, and cEt	21	6560	6575	3578
562119	N/A	N/A	GTCTTTCAGTTCATAA	Deoxy, MOE, and cEt	57	6567	6582	3579
562120	N/A	N/A	ACAGTTTGTCTTTCAG	Deoxy, MOE, and cEt	59	6574	6589	3580
562121	N/A	N/A	AGAAGTACAGTTTGTC	Deoxy, MOE, and cEt	3	6580	6595	3581
562122	N/A	N/A	GATGTCAGAAGTACAG	Deoxy, MOE, and cEt	45	6586	6601	3582
562123	N/A	N/A	AGTAAGGATGTCAGAA	Deoxy, MOE, and cEt	44	6592	6607	3583
562124	N/A	N/A	AATCTGAGTAAGGATG	Deoxy, MOE, and cEt	45	6598	6613	3584
562125	N/A	N/A	GAATATACAATTAGGG	Deoxy, MOE, and cEt	13	6616	6631	3585
562126	N/A	N/A	TGATACTGAATATACA	Deoxy, MOE, and cEt	13	6623	6638	3586
562127	N/A	N/A	CTGAGCTGATAAAAGA	Deoxy, MOE, and cEt	1	6660	6675	3587
562128	N/A	N/A	ACCATCATGTTTTACA	Deoxy, MOE, and cEt	44	6772	6787	3588
562129	N/A	N/A	TGTCTTACCATCATGT	Deoxy, MOE, and cEt	29	6778	6793	3589
562130	N/A	N/A	CCAAAGTGTCTTACCA	Deoxy, MOE, and cEt	42	6784	6799	3590
562131	N/A	N/A	AACCCACCAAAGTGTC	Deoxy, MOE, and cEt	33	6790	6805	3591
562132	N/A	N/A	GAAGGAAACCCACCAA	Deoxy, MOE, and cEt	24	6796	6811	3592
562133	N/A	N/A	CTTCAAGAAGGAAACC	Deoxy, MOE, and cEt	28	6802	6817	3593
562134	N/A	N/A	TAATAGCTTCAAGAAG	Deoxy, MOE, and cEt	1	6808	6823	3594
562135	N/A	N/A	GGGAATTTGATAATAA	Deoxy, MOE, and cEt	0	6821	6836	3595
562136	N/A	N/A	AGAATAGGGAATTTGA	Deoxy, MOE, and cEt	18	6827	6842	3596
562137	N/A	N/A	GTCCTAAGAATAGGGA	Deoxy, MOE, and cEt	9	6833	6848	3597
562138	N/A	N/A	GAACAAGTCCTAAGAA	Deoxy, MOE, and cEt	7	6839	6854	3598
562139	N/A	N/A	AGTCTAGAACAAGTCC	Deoxy, MOE, and cEt	70	6845	6860	3599
562140	N/A	N/A	TCTTTTAGTCTAGAAC	Deoxy, MOE, and cEt	22	6851	6866	3600
562141	N/A	N/A	TAACTATCTTTTAGTC	Deoxy, MOE, and cEt	15	6857	6872	3601
562142	N/A	N/A	ATCTCTTA ACTATCTT	Deoxy, MOE, and cEt	35	6863	6878	3602
560991	3	18	AAGTGTCTTCTCTGG	Deoxy, MOE, and cEt	37	3107	3122	3603
560992	8	23	CGTGGAAGTCTTCT	Deoxy, MOE, and cEt	74	3112	3127	112
560993	22	37	TCAATTTCAAGCAACG	Deoxy, MOE, and cEt	68	3126	3141	3604
560994	51	66	CTTAATTGTGAACATT	Deoxy, MOE, and cEt	21	3155	3170	3605
560995	53	68	AGCTTAATTGTGAACA	Deoxy, MOE, and cEt	59	3157	3172	3606
560996	55	70	GGAGCTTAATTGTGAA	Deoxy, MOE, and cEt	0	3159	3174	3607
560997	57	72	AAGGAGCTTAATTGTG	Deoxy, MOE, and cEt	36	3161	3176	3608
560998	59	74	AGAAGGAGCTTAATTG	Deoxy, MOE, and cEt	47	3163	3178	3609
560999	61	76	AAAGAAGGAGCTTAAT	Deoxy, MOE, and cEt	20	3165	3180	3610
561000	76	91	CTAGAGGAACAATAAA	Deoxy, MOE, and cEt	23	3180	3195	3611

561001	79	94	TAAC TAGAGGAACAAT	Deoxy, MOE, and cEt	19	3183	3198	3612
561002	81	96	AATAACTAGAGGAACA	Deoxy, MOE, and cEt	38	3185	3200	3613
561003	84	99	GGAAATAACTAGAGGA	Deoxy, MOE, and cEt	48	3188	3203	3614
561004	86	101	GAGGAAATAACTAGAG	Deoxy, MOE, and cEt	37	3190	3205	3615
561005	88	103	TGGAGGAAATAACTAG	Deoxy, MOE, and cEt	68	3192	3207	3616
561006	90	105	TCTGGAGGAAATAACT	Deoxy, MOE, and cEt	49	3194	3209	3617
561007	94	109	CAATTCTGGAGGAAAT	Deoxy, MOE, and cEt	43	3198	3213	3618
561008	96	111	ATCAATTCTGGAGGAA	Deoxy, MOE, and cEt	73	3200	3215	3619
561009	98	113	TGATCAATTCTGGAGG	Deoxy, MOE, and cEt	72	3202	3217	3620
561010	100	115	CTTGATCAATTCTGGA	Deoxy, MOE, and cEt	82	3204	3219	113
561011	102	117	GTCTTGATCAATTCTG	Deoxy, MOE, and cEt	85	3206	3221	114
561012	104	119	TTGTCTTGATCAATTC	Deoxy, MOE, and cEt	64	3208	3223	3621
561013	106	121	AATTGTCTTGATCAAT	Deoxy, MOE, and cEt	21	3210	3225	3622
561014	108	123	TGAATTGTCTTGATCA	Deoxy, MOE, and cEt	66	3212	3227	3623
561015	110	125	GATGAATTGTCTTGAT	Deoxy, MOE, and cEt	51	3214	3229	3624
561016	112	127	ATGATGAATTGTCTTG	Deoxy, MOE, and cEt	71	3216	3231	3625
561017	115	130	CAAATGATGAATTGTC	Deoxy, MOE, and cEt	36	3219	3234	3626
561018	117	132	ATCAAATGATGAATTG	Deoxy, MOE, and cEt	27	3221	3236	3627
561019	125	140	GATAGAGAATCAAATG	Deoxy, MOE, and cEt	11	3229	3244	3628
561020	129	144	TGGAGATAGAGAATCA	Deoxy, MOE, and cEt	73	3233	3248	3629
561021	131	146	TCTGGAGATAGAGAAT	Deoxy, MOE, and cEt	51	3235	3250	3630
561022	135	150	TGGCTCTGGAGATAGA	Deoxy, MOE, and cEt	76	3239	3254	115
561023	137	152	TTTGGCTCTGGAGATA	Deoxy, MOE, and cEt	73	3241	3256	3631
561024	139	154	ATTTTGGCTCTGGAGA	Deoxy, MOE, and cEt	61	3243	3258	3632
561025	141	156	TGATTTTGGCTCTGGA	Deoxy, MOE, and cEt	83	3245	3260	116
561026	143	158	CTTGATTTTGGCTCTG	Deoxy, MOE, and cEt	83	3247	3262	117
561027	145	160	ATCTTGATTTTGGCTC	Deoxy, MOE, and cEt	67	3249	3264	3633
559277	147	162	AAATCTTGATTTTGGC	Deoxy, MOE, and cEt	75	3251	3266	110
561028	149	164	GCAAATCTTGATTTTG	Deoxy, MOE, and cEt	53	3253	3268	3634
561029	151	166	TAGCAAATCTTGATTT	Deoxy, MOE, and cEt	27	3255	3270	3635
561030	153	168	CATAGCAAATCTTGAT	Deoxy, MOE, and cEt	63	3257	3272	3636
561031	155	170	AACATAGCAAATCTTG	Deoxy, MOE, and cEt	56	3259	3274	3637
561032	157	172	CTAACATAGCAAATCT	Deoxy, MOE, and cEt	67	3261	3276	3638
561033	159	174	GTCTAACATAGCAAAT	Deoxy, MOE, and cEt	51	3263	3278	3639
561034	174	189	TAAAATTTTACATCG	Deoxy, MOE, and cEt	4	3278	3293	3640
561035	177	192	GGCTAAAATTTTACA	Deoxy, MOE, and cEt	0	3281	3296	3641
561036	182	197	CCATTGGCTAAAATTT	Deoxy, MOE, and cEt	3	3286	3301	3642
561037	184	199	GGCCATTGGCTAAAAT	Deoxy, MOE, and cEt	16	3288	3303	3643
561038	186	201	GAGGCCATTGGCTAAA	Deoxy, MOE, and cEt	42	3290	3305	3644
561039	188	203	AGGAGGCCATTGGCTA	Deoxy, MOE, and cEt	61	3292	3307	3645
561040	190	205	GAAGGAGGCCATTGGC	Deoxy, MOE, and cEt	35	3294	3309	3646

561041	192	207	CTGAAGGAGGCCATTG	Deoxy, MOE, and cEt	37	3296	3311	3647
561042	194	209	AACTGAAGGAGGCCAT	Deoxy, MOE, and cEt	22	3298	3313	3648
561043	196	211	CCAACCTGAAGGAGGCC	Deoxy, MOE, and cEt	33	3300	3315	3649
561044	198	213	TCCCAACTGAAGGAGG	Deoxy, MOE, and cEt	19	3302	3317	3650
561045	200	215	TGTCCCAACTGAAGGA	Deoxy, MOE, and cEt	33	3304	3319	3651
561046	202	217	CATGTCCCAACTGAAG	Deoxy, MOE, and cEt	19	3306	3321	3652
561047	204	219	ACCATGTCCCAACTGA	Deoxy, MOE, and cEt	19	3308	3323	3653
561048	206	221	AGACCATGTCCCAACT	Deoxy, MOE, and cEt	19	3310	3325	3654
561049	208	223	TAAGACCATGTCCCAA	Deoxy, MOE, and cEt	0	3312	3327	3655
561050	210	225	TTTAAGACCATGTCCC	Deoxy, MOE, and cEt	5	3314	3329	3656
561051	212	227	TCTTTAAGACCATGTC	Deoxy, MOE, and cEt	10	3316	3331	3657
561052	214	229	AGTCTTTAAGACCATG	Deoxy, MOE, and cEt	10	3318	3333	3658
561053	216	231	AAAGTCTTTAAGACCA	Deoxy, MOE, and cEt	29	3320	3335	3659
561054	218	233	ACAAAGTCTTTAAGAC	Deoxy, MOE, and cEt	19	3322	3337	3660
561055	220	235	GGACAAAGTCTTTAAG	Deoxy, MOE, and cEt	21	3324	3339	3661
561056	222	237	ATGGACAAAGTCTTTA	Deoxy, MOE, and cEt	12	3326	3341	3662
561057	224	239	TTATGGACAAAGTCTT	Deoxy, MOE, and cEt	10	3328	3343	3663
561058	226	241	TCTTATGGACAAAGTC	Deoxy, MOE, and cEt	9	3330	3345	3664
561059	228	243	CGTCTTATGGACAAAG	Deoxy, MOE, and cEt	0	3332	3347	3665
561060	242	257	TTAATTTGGCCCTTCG	Deoxy, MOE, and cEt	28	3346	3361	3666
561061	244	259	CATTAATTTGGCCCTT	Deoxy, MOE, and cEt	13	3348	3363	3667
561062	246	261	GTCATTAATTTGGCCC	Deoxy, MOE, and cEt	63	3350	3365	3668
561063	248	263	ATGTCATTAATTTGGC	Deoxy, MOE, and cEt	37	3352	3367	3669
561064	267	282	TATGTTGAGTTTTTGA	Deoxy, MOE, and cEt	16	3371	3386	3670
561065	272	287	TCAAATATGTTGAGTT	Deoxy, MOE, and cEt	21	3376	3391	3671
561066	274	289	GATCAAATATGTTGAG	Deoxy, MOE, and cEt	36	3378	3393	3672
560990	709	724	TTCTTGCTGCTCTTGG	Deoxy, MOE, and cEt	73	6722	6737	111
337487	804	823	CACTTGTATGTTACCTCTG	5-10-5 MOE	76	7389	7408	28
561604	1850	1865	GTACAATTACCAGTCC	Deoxy, MOE, and cEt	59	10822	10837	3673
561605	1852	1867	CTGTACAATTACCAGT	Deoxy, MOE, and cEt	54	10824	10839	3674
561606	1854	1869	AACTGTACAATTACCA	Deoxy, MOE, and cEt	57	10826	10841	3675
561607	1856	1871	AGAACTGTACAATTAC	Deoxy, MOE, and cEt	36	10828	10843	3676
561608	1858	1873	TAAGAACTGTACAATT	Deoxy, MOE, and cEt	29	10830	10845	3677
561609	1862	1877	CATTTAAGAACTGTAC	Deoxy, MOE, and cEt	24	10834	10849	3678
561610	1870	1885	TACTACAACATTTAAG	Deoxy, MOE, and cEt	1	10842	10857	3679
561611	1874	1889	TTAATACTACAACATT	Deoxy, MOE, and cEt	0	10846	10861	3680
561612	1880	1895	TTGAAATTAATACTAC	Deoxy, MOE, and cEt	6	10852	10867	3681
561613	1883	1898	GTTTTGAAATTAATAC	Deoxy, MOE, and cEt	34	10855	10870	3682
561614	1892	1907	CGATTTTTAGTTTTGA	Deoxy, MOE, and cEt	22	10864	10879	3683
561615	1894	1909	GACGATTTTTAGTTTT	Deoxy, MOE, and cEt	29	10866	10881	3684
561616	1896	1911	CTGACGATTTTTAGTT	Deoxy, MOE, and cEt	50	10868	10883	3685

561617	1898	1913	TGCTGACGATTTT TAG	Deoxy, MOE, and cEt	54	10870	10885	3686
561618	1900	1915	TGTGCTGACGATTTT	Deoxy, MOE, and cEt	70	10872	10887	3687
561619	1902	1917	TCTGTGCTGACGATT	Deoxy, MOE, and cEt	69	10874	10889	3688
561620	1904	1919	ACTCTGTGCTGACGAT	Deoxy, MOE, and cEt	78	10876	10891	135
561621	1906	1921	ATACTCTGTGCTGACG	Deoxy, MOE, and cEt	87	10878	10893	134
561622	1908	1923	ACATACTCTGTGCTGA	Deoxy, MOE, and cEt	80	10880	10895	136
561623	1911	1926	TACACATACTCTGTGC	Deoxy, MOE, and cEt	61	10883	10898	3689
561624	1913	1928	TTACACATACTCTGT	Deoxy, MOE, and cEt	68	10885	10900	3690
561625	1917	1932	GATTTTACACATACT	Deoxy, MOE, and cEt	17	10889	10904	3691
561626	1946	1961	GAAGCATCAGTTTAAA	Deoxy, MOE, and cEt	27	10918	10933	3692
561627	1948	1963	ATGAAGCATCAGTTTA	Deoxy, MOE, and cEt	5	10920	10935	3693
561628	1956	1971	GTAGCAAAATGAAGCA	Deoxy, MOE, and cEt	73	10928	10943	137
561629	1958	1973	TTGTAGCAAAATGAAG	Deoxy, MOE, and cEt	42	10930	10945	3694
561630	1976	1991	CATTTACTCCAAATTA	Deoxy, MOE, and cEt	43	10948	10963	3695
561631	1981	1996	TCAAACATTTACTCCA	Deoxy, MOE, and cEt	82	10953	10968	138
561632	2006	2021	CATTAGGTTTCATAAA	Deoxy, MOE, and cEt	19	10978	10993	3696
561633	2008	2023	TTCATTAGGTTTCATA	Deoxy, MOE, and cEt	15	10980	10995	3697
561634	2010	2025	GCTTCATTAGGTTTCA	Deoxy, MOE, and cEt	57	10982	10997	3698
561635	2012	2027	CTGCTTCATTAGGTTT	Deoxy, MOE, and cEt	0	10984	10999	3699
561636	2014	2029	TTCTGCTTCATTAGGT	Deoxy, MOE, and cEt	65	10986	11001	3700
561637	2016	2031	AATTCTGCTTCATTAG	Deoxy, MOE, and cEt	48	10988	11003	3701
561638	2024	2039	CAGTATTTAATTCTGC	Deoxy, MOE, and cEt	38	10996	11011	3702
561639	2039	2054	GAAC TTATTTTAATAC	Deoxy, MOE, and cEt	29	11011	11026	3703
561640	2041	2056	GCGAACTTATTTTAAT	Deoxy, MOE, and cEt	38	11013	11028	3704
561641	2043	2058	CAGCGAACTTATTTTA	Deoxy, MOE, and cEt	46	11015	11030	3705
561642	2045	2060	GACAGCGAACTTATTT	Deoxy, MOE, and cEt	64	11017	11032	3706
561643	2047	2062	AAGACAGCGAACTTAT	Deoxy, MOE, and cEt	19	11019	11034	3707
561644	2049	2064	TAAAGACAGCGAACTT	Deoxy, MOE, and cEt	76	11021	11036	139
561645	2051	2066	TTTAAAGACAGCGAAC	Deoxy, MOE, and cEt	49	11023	11038	3708
561646	2053	2068	TGTTTAAAGACAGCGA	Deoxy, MOE, and cEt	81	11025	11040	140
561647	2065	2080	GTCATCTCCATTTGTT	Deoxy, MOE, and cEt	60	11037	11052	3709
561648	2067	2082	TAGTCATCTCCATTTG	Deoxy, MOE, and cEt	69	11039	11054	3710
561649	2069	2084	AGTAGTCATCTCCATT	Deoxy, MOE, and cEt	82	11041	11056	141
561650	2071	2086	TTAGTAGTCATCTCCA	Deoxy, MOE, and cEt	79	11043	11058	142
561651	2073	2088	ACTTAGTAGTCATCTC	Deoxy, MOE, and cEt	66	11045	11060	3711
561652	2075	2090	TGACTTAGTAGTCATC	Deoxy, MOE, and cEt	62	11047	11062	3712
561653	2077	2092	TGTGACTTAGTAGTCA	Deoxy, MOE, and cEt	52	11049	11064	3713
561654	2079	2094	AATGTGACTTAGTAGT	Deoxy, MOE, and cEt	44	11051	11066	3714
561655	2081	2096	TCAATGTGACTTAGTA	Deoxy, MOE, and cEt	65	11053	11068	3715
561656	2083	2098	AGTCAATGTGACTTAG	Deoxy, MOE, and cEt	70	11055	11070	3716
561657	2085	2100	AAAGTCAATGTGACTT	Deoxy, MOE, and cEt	2	11057	11072	3717

561658	2087	2102	TTAAAGTCAATGTGAC	Deoxy, MOE, and cEt	15	11059	11074	3718
561659	2089	2104	TGTTAAAGTCAATGTG	Deoxy, MOE, and cEt	27	11061	11076	3719
561660	2091	2106	CATGTTAAAGTCAATG	Deoxy, MOE, and cEt	51	11063	11078	3720
561661	2093	2108	CTCATGTTAAAGTCAA	Deoxy, MOE, and cEt	53	11065	11080	3721
561662	2095	2110	ACCTCATGTTAAAGTC	Deoxy, MOE, and cEt	55	11067	11082	3722
561663	2097	2112	ATACCTCATGTTAAAG	Deoxy, MOE, and cEt	25	11069	11084	3723
561664	2099	2114	TGATACCTCATGTAA	Deoxy, MOE, and cEt	0	11071	11086	3724
561665	2101	2116	AGTGATACCTCATGTT	Deoxy, MOE, and cEt	38	11073	11088	3725
561666	2103	2118	ATAGTGATACCTCATG	Deoxy, MOE, and cEt	61	11075	11090	3726
561667	2105	2120	GTATAGTGATACCTCA	Deoxy, MOE, and cEt	63	11077	11092	3727
561668	2107	2122	AGGTATAGTGATACCT	Deoxy, MOE, and cEt	27	11079	11094	3728
561669	2109	2124	TAAGGTATAGTGATAC	Deoxy, MOE, and cEt	34	11081	11096	3729
561670	2111	2126	AATAAGGTATAGTGAT	Deoxy, MOE, and cEt	22	11083	11098	3730

Table 27

Inhibition of ANGPTL3 mRNA by oligonucleotides targeting SEQ ID NO: 1 and 2

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	Chemistry	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
562220	N/A	N/A	GTAAACTTATTGATAA	Deoxy, MOE, and cEt	0	7670	7685	3731
562221	N/A	N/A	GGCATAGTAAACTTAT	Deoxy, MOE, and cEt	22	7676	7691	3732
562222	N/A	N/A	AATTTTGGCATAGTAA	Deoxy, MOE, and cEt	0	7682	7697	3733
562223	N/A	N/A	GGCAATTAATGAATTT	Deoxy, MOE, and cEt	15	7693	7708	3734
562224	N/A	N/A	GTGAAAGGCAATTAAT	Deoxy, MOE, and cEt	7	7699	7714	3735
562225	N/A	N/A	AGTTAAGTGAAAGGCA	Deoxy, MOE, and cEt	0	7705	7720	3736
562226	N/A	N/A	CCCAAAAGTTAAGTGA	Deoxy, MOE, and cEt	27	7711	7726	3737
562227	N/A	N/A	TATGGTCCCAAAAGTT	Deoxy, MOE, and cEt	35	7717	7732	3738
562228	N/A	N/A	ATTTATTATGGTCCCA	Deoxy, MOE, and cEt	67	7723	7738	3739
562229	N/A	N/A	GTTATGGCAATACATT	Deoxy, MOE, and cEt	37	7744	7759	3740
562230	N/A	N/A	ATTAATGTTATGGCAA	Deoxy, MOE, and cEt	33	7750	7765	3741
562231	N/A	N/A	GTAGTTTATTAATGTT	Deoxy, MOE, and cEt	15	7757	7772	3742
562232	N/A	N/A	TGTAAGGTAGTTTATT	Deoxy, MOE, and cEt	23	7763	7778	3743
562233	N/A	N/A	TGGTTTTGTAAGGTAG	Deoxy, MOE, and cEt	43	7769	7784	3744
562234	N/A	N/A	AATTGGTGGTTTTGTA	Deoxy, MOE, and cEt	18	7775	7790	3745
562235	N/A	N/A	GATTTTAATTGGTGGT	Deoxy, MOE, and cEt	21	7781	7796	3746
562236	N/A	N/A	GATGTAAATAACACTT	Deoxy, MOE, and cEt	9	7809	7824	3747
562237	N/A	N/A	TTGACAGATGTAAATA	Deoxy, MOE, and cEt	11	7815	7830	3748
562238	N/A	N/A	TTTATGTTGACAGATG	Deoxy, MOE, and cEt	20	7821	7836	3749
562239	N/A	N/A	AGTAGATTTATGTTGA	Deoxy, MOE, and cEt	9	7827	7842	3750
562240	N/A	N/A	CCTGAATATAATGAAT	Deoxy, MOE, and cEt	29	7859	7874	3751
562241	N/A	N/A	GGACTACCTGAATATA	Deoxy, MOE, and cEt	17	7865	7880	3752

562242	N/A	N/A	ACCATCAAGCCTCCCA	Deoxy, MOE, and cEt	45	7956	7971	3753
562243	N/A	N/A	CCCCTTACCATCAAGC	Deoxy, MOE, and cEt	31	7962	7977	3754
562244	N/A	N/A	TGTAGTCCCCTTACCA	Deoxy, MOE, and cEt	16	7968	7983	3755
562245	N/A	N/A	ATTGAATGTAGTCCCC	Deoxy, MOE, and cEt	19	7974	7989	3756
562246	N/A	N/A	GATTAGCAAGTGAATG	Deoxy, MOE, and cEt	6	7994	8009	3757
562247	N/A	N/A	TTTGTAGATTAGCAAG	Deoxy, MOE, and cEt	24	8000	8015	3758
562248	N/A	N/A	AAGAGGTTCTCAGTAA	Deoxy, MOE, and cEt	28	8019	8034	3759
562249	N/A	N/A	GTCCATAAGAGGTTCT	Deoxy, MOE, and cEt	34	8025	8040	3760
562250	N/A	N/A	TACCTGGTCCATAAGA	Deoxy, MOE, and cEt	10	8031	8046	3761
562251	N/A	N/A	TCCTAATACCTGGTCC	Deoxy, MOE, and cEt	32	8037	8052	3762
562252	N/A	N/A	TACTTTTCCTAATACC	Deoxy, MOE, and cEt	20	8043	8058	3763
562253	N/A	N/A	CGTTACTACTTTTCCT	Deoxy, MOE, and cEt	29	8049	8064	3764
562254	N/A	N/A	CTGAGACTGCTTCTCG	Deoxy, MOE, and cEt	36	8067	8082	3765
562255	N/A	N/A	TGAAGGCTGAGACTGC	Deoxy, MOE, and cEt	40	8073	8088	3766
562256	N/A	N/A	TAAATTATATGAAGGC	Deoxy, MOE, and cEt	9	8082	8097	3767
562257	N/A	N/A	GTAATTGTTTGATAAT	Deoxy, MOE, and cEt	0	8097	8112	3768
562258	N/A	N/A	TACTAACAAATGTGTA	Deoxy, MOE, and cEt	0	8110	8125	3769
562259	N/A	N/A	GTAATTTACTAACAAA	Deoxy, MOE, and cEt	0	8116	8131	3770
562260	N/A	N/A	ATAAGTGTAATTTACT	Deoxy, MOE, and cEt	0	8122	8137	3771
562261	N/A	N/A	GTTGTAATAAGTGTA	Deoxy, MOE, and cEt	0	8128	8143	3772
562262	N/A	N/A	GTGATAAATATAATTC	Deoxy, MOE, and cEt	0	8155	8170	3773
562263	N/A	N/A	CATGTAATTGTGATAA	Deoxy, MOE, and cEt	20	8164	8179	3774
562264	N/A	N/A	GTATATTTAAGAACAG	Deoxy, MOE, and cEt	13	8181	8196	3775
562265	N/A	N/A	TTGTGATAAGTATATT	Deoxy, MOE, and cEt	3	8190	8205	3776
562266	N/A	N/A	TGGAATTAAATTGTGA	Deoxy, MOE, and cEt	0	8200	8215	3777
562267	N/A	N/A	AAGCCGTGGAATTAAA	Deoxy, MOE, and cEt	10	8206	8221	3778
562268	N/A	N/A	CATTGTAAGCCGTGGA	Deoxy, MOE, and cEt	54	8212	8227	3779
562269	N/A	N/A	TATGATCATTGTAAGC	Deoxy, MOE, and cEt	0	8218	8233	3780
562270	N/A	N/A	TATAGTTATGATCATT	Deoxy, MOE, and cEt	0	8224	8239	3781
562271	N/A	N/A	GACATAACATTTAATC	Deoxy, MOE, and cEt	21	8258	8273	3782
562272	N/A	N/A	ACTTATGACATAACAT	Deoxy, MOE, and cEt	14	8264	8279	3783
562273	N/A	N/A	GTTACTACTTATGACA	Deoxy, MOE, and cEt	30	8270	8285	3784
562274	N/A	N/A	GTAACAGTTACTACTT	Deoxy, MOE, and cEt	24	8276	8291	3785
562275	N/A	N/A	GCTTATTTGTAACAGT	Deoxy, MOE, and cEt	17	8284	8299	3786
562276	N/A	N/A	TTCACAGCTTATTTGT	Deoxy, MOE, and cEt	20	8290	8305	3787
562277	N/A	N/A	GTTCTTTTCACAGCTT	Deoxy, MOE, and cEt	46	8296	8311	3788
562278	N/A	N/A	GGAGTG GTTCTTTTCA	Deoxy, MOE, and cEt	35	8302	8317	3789
562279	N/A	N/A	ATGCTAGGAGTGGTTC	Deoxy, MOE, and cEt	29	8308	8323	3790
562280	N/A	N/A	TGACTAATGCTAGGAG	Deoxy, MOE, and cEt	4	8314	8329	3791
562281	N/A	N/A	ATAGAGTGACTAATGC	Deoxy, MOE, and cEt	23	8320	8335	3792
562282	N/A	N/A	GAGAGAATAGAGTGAC	Deoxy, MOE, and cEt	15	8326	8341	3793

562284	N/A	N/A	ATTGATATGTAAAACG	Deoxy, MOE, and cEt	7	8347	8362	3794
562285	N/A	N/A	CAATTAATTGATATGT	Deoxy, MOE, and cEt	14	8353	8368	3795
562286	N/A	N/A	CCTTTTAACTTCCAAT	Deoxy, MOE, and cEt	40	8365	8380	3796
562287	N/A	N/A	CCTGGTCCTTTTAACT	Deoxy, MOE, and cEt	29	8371	8386	3797
562288	N/A	N/A	GAGTTTCCTGGTCCTT	Deoxy, MOE, and cEt	49	8377	8392	3798
562289	N/A	N/A	ATGTCTGAGTTTCCTG	Deoxy, MOE, and cEt	16	8383	8398	3799
562290	N/A	N/A	TACTGTATGTCTGAGT	Deoxy, MOE, and cEt	33	8389	8404	3800
562291	N/A	N/A	CCATACATTCTATATA	Deoxy, MOE, and cEt	10	8437	8452	3801
562292	N/A	N/A	TATAAGCCATACATTC	Deoxy, MOE, and cEt	24	8443	8458	3802
562293	N/A	N/A	ATTCATTATAAGCCAT	Deoxy, MOE, and cEt	38	8449	8464	3803
562295	N/A	N/A	CATTGAGTTAACTAAT	Deoxy, MOE, and cEt	7	8463	8478	3804
562296	N/A	N/A	AATTTGCATTGAGTTA	Deoxy, MOE, and cEt	18	8469	8484	3805
561144	525	540	TGAAGTTACTTCTGGG	Deoxy, MOE, and cEt	39	3629	3644	3806
561145	527	542	AGTGAAGTTACTTCTG	Deoxy, MOE, and cEt	51	3631	3646	3807
561146	529	544	TAAGTGAAGTTACTTC	Deoxy, MOE, and cEt	40	3633	3648	3808
561147	533	548	GTTTTAAGTGAAGTTA	Deoxy, MOE, and cEt	29	N/A	N/A	3809
561148	535	550	AAGTTTTAAGTGAAGT	Deoxy, MOE, and cEt	19	N/A	N/A	3810
561149	547	562	GTTTTTCTACAAAAGT	Deoxy, MOE, and cEt	38	4285	4300	3811
561150	560	575	ATGCTATTATCTTGTT	Deoxy, MOE, and cEt	30	4298	4313	3812
561151	562	577	TGATGCTATTATCTTG	Deoxy, MOE, and cEt	36	4300	4315	3813
561152	564	579	TTTGATGCTATTATCT	Deoxy, MOE, and cEt	23	4302	4317	3814
561153	567	582	GTCTTTGATGCTATTA	Deoxy, MOE, and cEt	51	4305	4320	3815
561154	569	584	AGGTCTTTGATGCTAT	Deoxy, MOE, and cEt	60	4307	4322	3816
561155	571	586	GAAGGTCTTTGATGCT	Deoxy, MOE, and cEt	61	4309	4324	3817
561156	573	588	GAGAAGGTCTTTGATG	Deoxy, MOE, and cEt	30	4311	4326	3818
561157	575	590	TGGAGAAGGTCTTTGA	Deoxy, MOE, and cEt	40	4313	4328	3819
561158	577	592	TCTGGAGAAGGTCTTT	Deoxy, MOE, and cEt	46	4315	4330	3820
561159	579	594	GGTCTGGAGAAGGTCT	Deoxy, MOE, and cEt	57	4317	4332	3821
561160	581	596	ACGGTCTGGAGAAGGT	Deoxy, MOE, and cEt	57	4319	4334	3822
561161	583	598	CCACGGTCTGGAGAAG	Deoxy, MOE, and cEt	56	4321	4336	3823
561162	585	600	TTCCACGGTCTGGAGA	Deoxy, MOE, and cEt	50	4323	4338	3824
561163	587	602	TCTTCCACGGTCTGGA	Deoxy, MOE, and cEt	77	4325	4340	3825
561164	589	604	GGTCTTCCACGGTCTG	Deoxy, MOE, and cEt	89	4327	4342	3826
561165	591	606	TTGGTCTTCCACGGTC	Deoxy, MOE, and cEt	79	4329	4344	3827
561166	593	608	TATTGGTCTTCCACGG	Deoxy, MOE, and cEt	39	4331	4346	3828
561167	595	610	TATATTGGTCTTCCAC	Deoxy, MOE, and cEt	22	4333	4348	3829
561168	597	612	TTTATATTGGTCTTCC	Deoxy, MOE, and cEt	43	4335	4350	3830
561169	599	614	TGTTTATATTGGTCTT	Deoxy, MOE, and cEt	50	4337	4352	3831
561170	601	616	ATTGTTTATATTGGTC	Deoxy, MOE, and cEt	27	4339	4354	3832
561171	603	618	TAATTGTTTATATTGG	Deoxy, MOE, and cEt	21	4341	4356	3833
561172	607	622	GGTTTAATTGTTTATA	Deoxy, MOE, and cEt	22	4345	4360	3834

561173	610	625	GTTGGTTTAATTGTTT	Deoxy, MOE, and cEt	33	4348	4363	3835
561174	612	627	CTGTTGGTTTAATTGT	Deoxy, MOE, and cEt	13	4350	4365	3836
561175	614	629	TGCTGTTGGTTTAATT	Deoxy, MOE, and cEt	26	4352	4367	3837
561176	616	631	TATGCTGTTGGTTTAA	Deoxy, MOE, and cEt	40	4354	4369	3838
561177	618	633	ACTATGCTGTTGGTTT	Deoxy, MOE, and cEt	68	4356	4371	3839
561178	620	635	TGACTATGCTGTTGGT	Deoxy, MOE, and cEt	64	4358	4373	3840
561179	622	637	TTTGACTATGCTGTTG	Deoxy, MOE, and cEt	42	4360	4375	3841
561180	624	639	TATTTGACTATGCTGT	Deoxy, MOE, and cEt	16	4362	4377	3842
561181	626	641	TTTATTTGACTATGCT	Deoxy, MOE, and cEt	17	4364	4379	3843
561182	628	643	CTTTTATTTGACTATG	Deoxy, MOE, and cEt	7	4366	4381	3844
561183	645	660	GAGCTGATTTTCTATT	Deoxy, MOE, and cEt	18	N/A	N/A	3845
561184	647	662	CTGAGCTGATTTTCTA	Deoxy, MOE, and cEt	42	N/A	N/A	3846
561185	649	664	TTCTGAGCTGATTTTC	Deoxy, MOE, and cEt	32	N/A	N/A	3847
561186	651	666	CCTTCTGAGCTGATTT	Deoxy, MOE, and cEt	14	N/A	N/A	3848
561187	653	668	GTCCTTCTGAGCTGAT	Deoxy, MOE, and cEt	39	6666	6681	3849
561188	655	670	TAGTCCTTCTGAGCTG	Deoxy, MOE, and cEt	7	6668	6683	3850
561189	657	672	ACTAGTCCTTCTGAGC	Deoxy, MOE, and cEt	32	6670	6685	3851
561190	659	674	ATACTAGTCCTTCTGA	Deoxy, MOE, and cEt	19	6672	6687	3852
561191	661	676	GAATACTAGTCCTTCT	Deoxy, MOE, and cEt	37	6674	6689	3853
561192	663	678	TTGAATACTAGTCCTT	Deoxy, MOE, and cEt	50	6676	6691	3854
561193	665	680	TCTTGAATACTAGTCC	Deoxy, MOE, and cEt	28	6678	6693	3855
561194	667	682	GTTCTTGAATACTAGT	Deoxy, MOE, and cEt	34	6680	6695	3856
561195	669	684	GGGTTCTTGAATACTA	Deoxy, MOE, and cEt	61	6682	6697	3857
561196	671	686	GTGGGTTCTTGAATAC	Deoxy, MOE, and cEt	21	6684	6699	3858
561197	673	688	CTGTGGGTTCTTGAAT	Deoxy, MOE, and cEt	45	6686	6701	3859
561198	675	690	TTCTGTGGGTTCTTGA	Deoxy, MOE, and cEt	0	6688	6703	3860
561199	679	694	AAATTTCTGTGGGTTC	Deoxy, MOE, and cEt	31	6692	6707	3861
561200	681	696	AGAAATTTCTGTGGGT	Deoxy, MOE, and cEt	60	6694	6709	3862
561201	684	699	TAGAGAAATTTCTGTG	Deoxy, MOE, and cEt	35	6697	6712	3863
561202	686	701	GATAGAGAAATTTCTG	Deoxy, MOE, and cEt	36	6699	6714	3864
561203	694	709	GCTTGGAAGATAGAGA	Deoxy, MOE, and cEt	39	6707	6722	3865
561204	696	711	TGGCTTGGAAGATAGA	Deoxy, MOE, and cEt	32	6709	6724	3866
561205	698	713	CTTGGCTTGGAAGATA	Deoxy, MOE, and cEt	23	6711	6726	3867
561206	700	715	CTCTTGGCTTGGAAGA	Deoxy, MOE, and cEt	21	6713	6728	3868
561207	702	717	TGCTCTTGGCTTGGA	Deoxy, MOE, and cEt	34	6715	6730	3869
561208	704	719	GGTGCTCTTGGCTTGG	Deoxy, MOE, and cEt	71	6717	6732	118
561209	706	721	TTGGTGCTCTTGGCTT	Deoxy, MOE, and cEt	59	6719	6734	3870
561210	708	723	TCTTGGTGCTCTTGGC	Deoxy, MOE, and cEt	65	6721	6736	3871
560990	709	724	TTCTTGGTGCTCTTGG	Deoxy, MOE, and cEt	54	6722	6737	111
561211	710	725	GTTCTTGGTGCTCTTG	Deoxy, MOE, and cEt	60	6723	6738	3872
561212	712	727	TAGTTCTTGGTGCTCT	Deoxy, MOE, and cEt	53	6725	6740	3873

561213	714	729	AGTAGTTCTTGGTGCT	Deoxy, MOE, and cEt	50	6727	6742	3874
561214	716	731	GGAGTAGTTCTTGGTG	Deoxy, MOE, and cEt	31	6729	6744	3875
561215	718	733	AGGGAGTAGTTCTTGG	Deoxy, MOE, and cEt	0	6731	6746	3876
561216	720	735	AAAGGGAGTAGTTCTT	Deoxy, MOE, and cEt	25	6733	6748	3877
561217	722	737	AGAAAGGGAGTAGTTC	Deoxy, MOE, and cEt	28	6735	6750	3878
561218	724	739	GAAGAAAGGGAGTAGT	Deoxy, MOE, and cEt	10	6737	6752	3879
561219	726	741	CTGAAGAAAGGGAGTA	Deoxy, MOE, and cEt	47	6739	6754	3880
561220	730	745	TCAACTGAAGAAAGGG	Deoxy, MOE, and cEt	50	6743	6758	3881
337487	804	823	CACTTGTATGTTACCTCTG	5-10-5 MOE	52	7389	7408	28
561297	926	941	TCATTGAAGTTTTGTG	Deoxy, MOE, and cEt	28	7913	7928	3882
561298	930	945	CGTTTCATTGAAGTTT	Deoxy, MOE, and cEt	35	7917	7932	3883
561299	944	959	TTGTAGTTCTCCCACG	Deoxy, MOE, and cEt	30	7931	7946	3884
561300	946	961	ATTTGTAGTTCTCCCA	Deoxy, MOE, and cEt	32	7933	7948	3885
561301	948	963	ATATTTGTAGTTCTCC	Deoxy, MOE, and cEt	24	7935	7950	3886
561302	950	965	CCATATTTGTAGTTCT	Deoxy, MOE, and cEt	5	7937	7952	3887
561303	952	967	AACCATATTTGTAGTT	Deoxy, MOE, and cEt	3	7939	7954	3888
561304	956	971	CCAAAACCATATTTGT	Deoxy, MOE, and cEt	19	7943	7958	3889
561305	959	974	CTCCCAAACCATATT	Deoxy, MOE, and cEt	23	7946	7961	3890
561306	961	976	GCCTCCCAAACCATATA	Deoxy, MOE, and cEt	25	7948	7963	3891
561307	963	978	AAGCCTCCCAAACCA	Deoxy, MOE, and cEt	30	7950	7965	3892
561308	965	980	TCAAGCCTCCCAAAC	Deoxy, MOE, and cEt	16	7952	7967	3893
561309	969	984	TCCATCAAGCCTCCCA	Deoxy, MOE, and cEt	46	N/A	N/A	3894
561310	971	986	TCTCCATCAAGCCTCC	Deoxy, MOE, and cEt	13	N/A	N/A	3895
561311	973	988	ATTCTCCATCAAGCCT	Deoxy, MOE, and cEt	16	N/A	N/A	3896
561312	975	990	AAATTCTCCATCAAGC	Deoxy, MOE, and cEt	20	N/A	N/A	3897
561313	979	994	ACCAAATTCTCCATC	Deoxy, MOE, and cEt	18	N/A	N/A	3898
561314	981	996	CAACCAAAATTCTCCA	Deoxy, MOE, and cEt	26	N/A	N/A	3899
561315	983	998	CCCAACCAAATTCTC	Deoxy, MOE, and cEt	38	9558	9573	3900
559316	985	1000	GGCCCAACCAAATTTC	Deoxy, MOE, and cEt	14	9560	9575	3901
561316	987	1002	TAGGCCCAACCAAAT	Deoxy, MOE, and cEt	38	9562	9577	3902
561317	989	1004	TCTAGGCCCAACCAAA	Deoxy, MOE, and cEt	51	9564	9579	3903
561318	991	1006	TCTCTAGGCCCAACCA	Deoxy, MOE, and cEt	35	9566	9581	3904
561319	993	1008	CTTCTCTAGGCCCAAC	Deoxy, MOE, and cEt	31	9568	9583	3905
561320	995	1010	ATCTTCTCTAGGCCCA	Deoxy, MOE, and cEt	68	9570	9585	119
561321	997	1012	ATATCTTCTCTAGGCC	Deoxy, MOE, and cEt	30	9572	9587	3906
561322	999	1014	GTATATCTTCTCTAGG	Deoxy, MOE, and cEt	25	9574	9589	3907
561323	1001	1016	GAGTATATCTTCTCTA	Deoxy, MOE, and cEt	26	9576	9591	3908
561324	1003	1018	TGGAGTATATCTTCTC	Deoxy, MOE, and cEt	46	9578	9593	3909
561325	1005	1020	TATGGAGTATATCTTC	Deoxy, MOE, and cEt	20	9580	9595	3910
561326	1007	1022	ACTATGGAGTATATCT	Deoxy, MOE, and cEt	20	9582	9597	3911
561327	1009	1024	TCACTATGGAGTATAT	Deoxy, MOE, and cEt	22	9584	9599	3912

561328	1011	1026	CTTCACTATGGAGTAT	Deoxy, MOE, and cEt	33	9586	9601	3913
561329	1013	1028	TGCTTCACTATGGAGT	Deoxy, MOE, and cEt	50	9588	9603	3914
561330	1015	1030	ATTGCTTCACTATGGA	Deoxy, MOE, and cEt	43	9590	9605	3915
561331	1017	1032	AGATTGCTTCACTATG	Deoxy, MOE, and cEt	31	9592	9607	3916
561332	1019	1034	TTAGATTGCTTCACTA	Deoxy, MOE, and cEt	36	9594	9609	3917
561333	1021	1036	AATTAGATTGCTTCAC	Deoxy, MOE, and cEt	17	9596	9611	3918
561334	1023	1038	ATAATTAGATTGCTTC	Deoxy, MOE, and cEt	23	9598	9613	3919
561335	1025	1040	ACATAATTAGATTGCT	Deoxy, MOE, and cEt	13	9600	9615	3920
561336	1031	1046	CGTAAAACATAATTAG	Deoxy, MOE, and cEt	25	9606	9621	3921
561337	1045	1060	CTTCCAACCTCAATTCTG	Deoxy, MOE, and cEt	0	9620	9635	3922
561338	1047	1062	GTCTTCCAACCTCAATT	Deoxy, MOE, and cEt	0	9622	9637	3923
561339	1049	1064	CAGTCTTCCAACCTCAA	Deoxy, MOE, and cEt	15	9624	9639	3924
561340	1051	1066	TCCAGTCTTCCAACCTC	Deoxy, MOE, and cEt	22	9626	9641	3925
561341	1053	1068	TTTCCAGTCTTCCAAC	Deoxy, MOE, and cEt	2	9628	9643	3926
561342	1056	1071	GTCTTTCCAGTCTTCC	Deoxy, MOE, and cEt	45	9631	9646	3927
561343	1059	1074	GTTGTCTTTCCAGTCT	Deoxy, MOE, and cEt	67	9634	9649	120
561344	1061	1076	TTGTTGTCTTTCCAGT	Deoxy, MOE, and cEt	43	9636	9651	3928
561345	1063	1078	GTTTGTGTGTCTTTCCA	Deoxy, MOE, and cEt	57	9638	9653	121
561346	1068	1083	ATAATGTTTGTGTGTCT	Deoxy, MOE, and cEt	6	9643	9658	3929
561347	1098	1113	GTGATTTCCCAAGTAA	Deoxy, MOE, and cEt	66	9673	9688	122
561348	1113	1128	CGTATAGTTGGTTTCG	Deoxy, MOE, and cEt	54	9688	9703	3930
561349	1127	1142	GCAACTAGATGTAGCG	Deoxy, MOE, and cEt	50	9702	9717	3931
561350	1129	1144	TCGCAACTAGATGTAG	Deoxy, MOE, and cEt	9	9704	9719	3932
561351	1131	1146	AATCGCAACTAGATGT	Deoxy, MOE, and cEt	9	9706	9721	3933
561352	1133	1148	GTAATCGCAACTAGAT	Deoxy, MOE, and cEt	15	9708	9723	3934
561353	1135	1150	CAGTAATCGCAACTAG	Deoxy, MOE, and cEt	41	9710	9725	3935
561354	1137	1152	GCCAGTAATCGCAACT	Deoxy, MOE, and cEt	38	9712	9727	3936
561355	1139	1154	TTGCCAGTAATCGCAA	Deoxy, MOE, and cEt	32	9714	9729	3937
561356	1141	1156	CATTGCCAGTAATCGC	Deoxy, MOE, and cEt	54	9716	9731	3938
561357	1143	1158	GACATTGCCAGTAATC	Deoxy, MOE, and cEt	20	9718	9733	3939
561358	1145	1160	GGGACATTGCCAGTAA	Deoxy, MOE, and cEt	0	9720	9735	3940
561359	1160	1175	TCCGGGATTGCATTGG	Deoxy, MOE, and cEt	43	9735	9750	3941
561360	1162	1177	TTTCCGGGATTGCATT	Deoxy, MOE, and cEt	31	9737	9752	3942
561361	1164	1179	GTTTTCCGGGATTGCA	Deoxy, MOE, and cEt	31	9739	9754	3943
561362	1166	1181	TTGTTTTCCGGGATTG	Deoxy, MOE, and cEt	36	9741	9756	3944
561363	1168	1183	CTTTGTTTTCCGGGAT	Deoxy, MOE, and cEt	22	9743	9758	3945
561364	1170	1185	ATCTTTGTTTTCCGGG	Deoxy, MOE, and cEt	13	9745	9760	3946
561365	1172	1187	AAATCTTTGTTTTCCG	Deoxy, MOE, and cEt	7	9747	9762	3947
561366	1177	1192	ACACCAAATCTTTGTT	Deoxy, MOE, and cEt	8	9752	9767	3948
561367	1179	1194	AAACACCAAATCTTTG	Deoxy, MOE, and cEt	11	9754	9769	3949
561368	1187	1202	CAAGTAGAAAACACCA	Deoxy, MOE, and cEt	16	9762	9777	3950

561369	1189	1204	CCCAAGTAGAAAACAC	Deoxy, MOE, and cEt	23	9764	9779	3951
561370	1191	1206	ATCCCAAGTAGAAAAC	Deoxy, MOE, and cEt	27	9766	9781	3952
561371	1193	1208	TGATCCCAAGTAGAAA	Deoxy, MOE, and cEt	25	9768	9783	3953
561372	1195	1210	TGTGATCCCAAGTAGA	Deoxy, MOE, and cEt	45	9770	9785	3954

Table 28

Inhibition of ANGPTL3 mRNA by oligonucleotides targeting SEQ ID NO: 1 and 2

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	Chemistry	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
561067	276	291	CTGATCAAATATGTTG	Deoxy, MOE, and cEt	54	3380	3395	3955
561068	278	293	GACTGATCAAATATGT	Deoxy, MOE, and cEt	19	3382	3397	3956
561069	280	295	AAGACTGATCAAATAT	Deoxy, MOE, and cEt	17	3384	3399	3957
561070	286	301	CATAAAAAGACTGATC	Deoxy, MOE, and cEt	18	3390	3405	3958
561071	289	304	GATCATAAAAAGACTG	Deoxy, MOE, and cEt	11	3393	3408	3959
561072	291	306	TAGATCATAAAAAGAC	Deoxy, MOE, and cEt	0	3395	3410	3960
561073	293	308	GATAGATCATAAAAAG	Deoxy, MOE, and cEt	15	3397	3412	3961
561074	295	310	GCGATAGATCATAAAA	Deoxy, MOE, and cEt	39	3399	3414	3962
561075	297	312	CAGCGATAGATCATAA	Deoxy, MOE, and cEt	53	3401	3416	3963
561076	299	314	TGCAGCGATAGATCAT	Deoxy, MOE, and cEt	70	3403	3418	159
561077	301	316	TTTGCAGCGATAGATC	Deoxy, MOE, and cEt	60	3405	3420	3964
561078	303	318	GGTTTGCAGCGATAGA	Deoxy, MOE, and cEt	63	3407	3422	3965
561079	305	320	CTGGTTTGCAGCGATA	Deoxy, MOE, and cEt	76	3409	3424	160
561080	307	322	CACTGGTTTGCAGCGA	Deoxy, MOE, and cEt	65	3411	3426	3966
561081	309	324	TTCAGTTTGCAGC	Deoxy, MOE, and cEt	45	3413	3428	3967
561082	311	326	ATTTCACTGGTTTGCA	Deoxy, MOE, and cEt	56	3415	3430	3968
561083	313	328	TGATTTCACTGGTTTG	Deoxy, MOE, and cEt	65	3417	3432	3969
561084	316	331	CTTTGATTTCACTGGT	Deoxy, MOE, and cEt	73	3420	3435	161
561085	341	356	GTTCTTCTCAGTTCCT	Deoxy, MOE, and cEt	79	3445	3460	162
561086	343	358	TAGTTCTTCTCAGTTC	Deoxy, MOE, and cEt	50	3447	3462	3970
561087	345	360	TGTAGTTCTTCTCAGT	Deoxy, MOE, and cEt	42	3449	3464	3971
561088	347	362	TATGTAGTTCTTCTCA	Deoxy, MOE, and cEt	27	3451	3466	3972
561089	349	364	TATATGTAGTTCTTCT	Deoxy, MOE, and cEt	37	3453	3468	3973
561090	352	367	GTTTATATGTAGTTCT	Deoxy, MOE, and cEt	39	3456	3471	3974
561091	355	370	GTAGTTTATATGTAGT	Deoxy, MOE, and cEt	55	3459	3474	3975
561092	358	373	CTTGTAGTTTATATGT	Deoxy, MOE, and cEt	48	3462	3477	3976
561093	360	375	GACTTGAGTTTATAT	Deoxy, MOE, and cEt	43	3464	3479	3977
561094	362	377	TTGACTTGAGTTTAT	Deoxy, MOE, and cEt	35	3466	3481	3978
561095	365	380	TTTTTGACTTGAGTT	Deoxy, MOE, and cEt	37	3469	3484	3979
561096	367	382	CATTTTGTAGTTG	Deoxy, MOE, and cEt	34	3471	3486	3980
561097	373	388	CCTCTTCATTTTGTAC	Deoxy, MOE, and cEt	48	3477	3492	3981

561098	386	401	GACATATTCTTTACCT	Deoxy, MOE, and cEt	40	3490	3505	3982
561099	388	403	GTGACATATTCTTTAC	Deoxy, MOE, and cEt	43	3492	3507	3983
561100	393	408	TTCAAGTGACATATTC	Deoxy, MOE, and cEt	51	3497	3512	3984
561101	395	410	AGTTCAAGTGACATAT	Deoxy, MOE, and cEt	27	3499	3514	3985
561102	397	412	TGAGTTCAAGTGACAT	Deoxy, MOE, and cEt	63	3501	3516	3986
561103	399	414	GTTGAGTTCAAGTGAC	Deoxy, MOE, and cEt	48	3503	3518	3987
561104	401	416	GAGTTGAGTTCAAGTG	Deoxy, MOE, and cEt	57	3505	3520	3988
561105	403	418	TTGAGTTGAGTTCAAG	Deoxy, MOE, and cEt	32	3507	3522	3989
561106	405	420	TTTTGAGTTGAGTTCA	Deoxy, MOE, and cEt	47	3509	3524	3990
561107	407	422	AGTTTTGAGTTGAGTT	Deoxy, MOE, and cEt	46	3511	3526	3991
561108	409	424	CAAGTTTTGAGTTGAG	Deoxy, MOE, and cEt	48	3513	3528	3992
561109	411	426	TTCAAGTTTTGAGTTG	Deoxy, MOE, and cEt	17	3515	3530	3993
561110	413	428	CTTCAAGTTTTGAGT	Deoxy, MOE, and cEt	48	3517	3532	3994
561111	415	430	GGCTTTCAAGTTTTGA	Deoxy, MOE, and cEt	56	3519	3534	3995
561112	417	432	GAGGCTTTCAAGTTTT	Deoxy, MOE, and cEt	39	3521	3536	3996
561113	419	434	AGGAGGCTTTCAAGTT	Deoxy, MOE, and cEt	49	3523	3538	3997
561114	421	436	CTAGGAGGCTTTCAAG	Deoxy, MOE, and cEt	49	3525	3540	3998
561115	423	438	TTCTAGGAGGCTTTCA	Deoxy, MOE, and cEt	40	3527	3542	3999
561116	425	440	TCTTCTAGGAGGCTTT	Deoxy, MOE, and cEt	66	3529	3544	4000
561117	427	442	TTTCTTCTAGGAGGCT	Deoxy, MOE, and cEt	74	3531	3546	4001
561118	442	457	GTTGAAGTAGAATTTT	Deoxy, MOE, and cEt	40	3546	3561	4002
561119	469	484	GTTGCTCTTCTAAATA	Deoxy, MOE, and cEt	44	3573	3588	4003
561120	471	486	TAGTTGCTCTTCTAAA	Deoxy, MOE, and cEt	19	3575	3590	4004
561121	473	488	GTTAGTTGCTCTTCTA	Deoxy, MOE, and cEt	67	3577	3592	4005
561122	475	490	TAGTTAGTTGCTCTTC	Deoxy, MOE, and cEt	51	3579	3594	4006
561123	477	492	GTTAGTTAGTTGCTCT	Deoxy, MOE, and cEt	73	3581	3596	163
561124	479	494	AAGTTAGTTAGTTGCT	Deoxy, MOE, and cEt	51	3583	3598	4007
561125	481	496	TTAAGTTAGTTAGTTG	Deoxy, MOE, and cEt	33	3585	3600	4008
561126	483	498	AATTAAGTTAGTTAGT	Deoxy, MOE, and cEt	0	3587	3602	4009
561127	485	500	TGAATTAAGTTAGTTA	Deoxy, MOE, and cEt	5	3589	3604	4010
561128	487	502	TTTGAATTAAGTTAGT	Deoxy, MOE, and cEt	18	3591	3606	4011
561129	494	509	GGTTGATTTTGAATTA	Deoxy, MOE, and cEt	20	3598	3613	4012
561130	496	511	CAGGTTGATTTTGAAT	Deoxy, MOE, and cEt	27	3600	3615	4013
561131	498	513	TTCAGGTTGATTTTGA	Deoxy, MOE, and cEt	33	3602	3617	4014
561132	500	515	GTTTCAGGTTGATTTT	Deoxy, MOE, and cEt	38	3604	3619	4015
561133	502	517	GAGTTTCAGGTTGATT	Deoxy, MOE, and cEt	33	3606	3621	4016
561134	504	519	TGGAGTTTCAGGTTGA	Deoxy, MOE, and cEt	67	3608	3623	4017
561135	507	522	TTCTGGAGTTTCAGGT	Deoxy, MOE, and cEt	32	3611	3626	4018
561136	509	524	TGTTCTGGAGTTTCAG	Deoxy, MOE, and cEt	14	3613	3628	4019
561137	511	526	GGTGTTCTGGAGTTTC	Deoxy, MOE, and cEt	23	3615	3630	4020
561138	513	528	TGGGTGTTCTGGAGTT	Deoxy, MOE, and cEt	30	3617	3632	4021

561139	515	530	TCTGGGTGTTCTGGAG	Deoxy, MOE, and cEt	24	3619	3634	4022
561140	517	532	CTTCTGGGTGTTCTGG	Deoxy, MOE, and cEt	17	3621	3636	4023
561141	519	534	TACTTCTGGGTGTTCT	Deoxy, MOE, and cEt	10	3623	3638	4024
561142	521	536	GTTACTTCTGGGTGTT	Deoxy, MOE, and cEt	11	3625	3640	4025
561143	523	538	AAGTTACTTCTGGGTG	Deoxy, MOE, and cEt	15	3627	3642	4026
560990	709	724	TTCTTGGTGCTCTTGG	Deoxy, MOE, and cEt	79	6722	6737	111
561221	758	773	CCATCATGTTTTACAT	Deoxy, MOE, and cEt	17	6771	6786	4027
561222	760	775	TGCCATCATGTTTTAC	Deoxy, MOE, and cEt	22	N/A	N/A	4028
561223	763	778	GAATGCCATCATGTTT	Deoxy, MOE, and cEt	12	N/A	N/A	4029
561224	765	780	AGGAATGCCATCATGT	Deoxy, MOE, and cEt	26	N/A	N/A	4030
561225	767	782	GCAGGAATGCCATCAT	Deoxy, MOE, and cEt	32	N/A	N/A	4031
561226	769	784	CAGCAGGAATGCCATC	Deoxy, MOE, and cEt	29	N/A	N/A	4032
561227	771	786	TTCAGCAGGAATGCCA	Deoxy, MOE, and cEt	22	N/A	N/A	4033
561228	773	788	CATTCAGCAGGAATGC	Deoxy, MOE, and cEt	23	7358	7373	4034
561229	775	790	TACATTCAGCAGGAAT	Deoxy, MOE, and cEt	28	7360	7375	4035
561230	777	792	GGTACATTCAGCAGGA	Deoxy, MOE, and cEt	61	7362	7377	4036
561231	779	794	GTGGTACATTCAGCAG	Deoxy, MOE, and cEt	57	7364	7379	4037
561232	781	796	TGGTGGTACATTCAGC	Deoxy, MOE, and cEt	59	7366	7381	4038
561233	787	802	TATAAATGGTGGTACA	Deoxy, MOE, and cEt	51	7372	7387	4039
561234	789	804	GTTATAAATGGTGGTA	Deoxy, MOE, and cEt	50	7374	7389	4040
561235	791	806	CTGTTATAAATGGTGG	Deoxy, MOE, and cEt	49	7376	7391	4041
561236	793	808	CTCTGTTATAAATGGT	Deoxy, MOE, and cEt	39	7378	7393	4042
561237	795	810	ACCTCTGTTATAAATG	Deoxy, MOE, and cEt	47	7380	7395	4043
561238	797	812	TCACCTCTGTTATAAA	Deoxy, MOE, and cEt	44	7382	7397	4044
561239	799	814	GTTACCTCTGTTATA	Deoxy, MOE, and cEt	43	7384	7399	4045
561240	801	816	ATGTTACCTCTGTTA	Deoxy, MOE, and cEt	59	7386	7401	4046
561241	803	818	GTATGTTACCTCTGT	Deoxy, MOE, and cEt	69	7388	7403	164
337487	804	823	CACTTGTATGTTACCTCTG	5-10-5 MOE	74	7389	7408	28
561242	805	820	TTGTATGTTACCTCT	Deoxy, MOE, and cEt	63	7390	7405	4047
561243	807	822	ACTTGTATGTTACCT	Deoxy, MOE, and cEt	63	7392	7407	4048
561244	809	824	CCACTTGTATGTTAC	Deoxy, MOE, and cEt	57	7394	7409	4049
561245	811	826	TGCCACTTGTATGTTC	Deoxy, MOE, and cEt	36	7396	7411	4050
561246	813	828	CATGCCACTTGTATGT	Deoxy, MOE, and cEt	33	7398	7413	4051
561247	815	830	TACATGCCACTTGTAT	Deoxy, MOE, and cEt	37	7400	7415	4052
561248	817	832	CATACATGCCACTTGT	Deoxy, MOE, and cEt	36	7402	7417	4053
561249	819	834	GGCATACATGCCACTT	Deoxy, MOE, and cEt	20	7404	7419	4054
561250	821	836	ATGGCATACATGCCAC	Deoxy, MOE, and cEt	0	7406	7421	4055
561251	823	838	TGATGGCATACATGCC	Deoxy, MOE, and cEt	22	7408	7423	4056
561252	825	840	TCTGATGGCATACATG	Deoxy, MOE, and cEt	34	7410	7425	4057
561253	827	842	GGTCTGATGGCATAACA	Deoxy, MOE, and cEt	46	7412	7427	4058
561254	829	844	TGGGTCTGATGGCATA	Deoxy, MOE, and cEt	51	7414	7429	4059

561255	834	849	GTTGCTGGGTCTGATG	Deoxy, MOE, and cEt	45	7419	7434	4060
561256	836	851	GAGTTGCTGGGTCTGA	Deoxy, MOE, and cEt	70	7421	7436	165
561257	838	853	GAGAGTTGCTGGGTCT	Deoxy, MOE, and cEt	57	7423	7438	4061
561258	840	855	TTGAGAGTTGCTGGGT	Deoxy, MOE, and cEt	47	7425	7440	4062
561259	842	857	ACTTGAGAGTTGCTGG	Deoxy, MOE, and cEt	53	7427	7442	4063
561260	844	859	AAACTTGAGAGTTGCT	Deoxy, MOE, and cEt	71	7429	7444	166
561261	846	861	AAAAACTTGAGAGTTG	Deoxy, MOE, and cEt	23	7431	7446	4064
561262	848	863	TGAAAAACTTGAGAGT	Deoxy, MOE, and cEt	11	7433	7448	4065
561263	850	865	CATGAAAAACTTGAGA	Deoxy, MOE, and cEt	34	7435	7450	4066
561264	852	867	GACATGAAAAACTTGA	Deoxy, MOE, and cEt	25	7437	7452	4067
561265	860	875	TCACAGTAGACATGAA	Deoxy, MOE, and cEt	16	7445	7460	4068
561266	862	877	CATCACAGTAGACATG	Deoxy, MOE, and cEt	37	7447	7462	4069
561267	864	879	AACATCACAGTAGACA	Deoxy, MOE, and cEt	57	7449	7464	4070
561268	866	881	ATAACATCACAGTAGA	Deoxy, MOE, and cEt	40	7451	7466	4071
561269	868	883	ATATAACATCACAGTA	Deoxy, MOE, and cEt	26	7453	7468	4072
561270	870	885	TGATATAACATCACAG	Deoxy, MOE, and cEt	35	7455	7470	4073
561271	872	887	CCTGATATAACATCAC	Deoxy, MOE, and cEt	60	7457	7472	4074
561272	874	889	TACCTGATATAACATC	Deoxy, MOE, and cEt	37	7459	7474	4075
561273	876	891	ACTACCTGATATAACA	Deoxy, MOE, and cEt	24	N/A	N/A	4076
561274	878	893	GGACTACCTGATATAA	Deoxy, MOE, and cEt	7	N/A	N/A	4077
561275	880	895	ATGGACTACCTGATAT	Deoxy, MOE, and cEt	33	N/A	N/A	4078
561276	882	897	CCATGGACTACCTGAT	Deoxy, MOE, and cEt	52	N/A	N/A	4079
561277	884	899	GTCCATGGACTACCTG	Deoxy, MOE, and cEt	71	7871	7886	167
561278	886	901	ATGTCCATGGACTACC	Deoxy, MOE, and cEt	67	7873	7888	4080
561279	888	903	TAATGTCCATGGACTA	Deoxy, MOE, and cEt	44	7875	7890	4081
559390	890	905	ATTAATGTCCATGGAC	Deoxy, MOE, and cEt	28	7877	7892	4082
561280	892	907	GAATTAATGTCCATGG	Deoxy, MOE, and cEt	51	7879	7894	4083
561281	894	909	TTGAATTAATGTCCAT	Deoxy, MOE, and cEt	30	7881	7896	4084
561282	896	911	TGTTGAATTAATGTCC	Deoxy, MOE, and cEt	38	7883	7898	4085
561283	898	913	GATGTTGAATTAATGT	Deoxy, MOE, and cEt	11	7885	7900	4086
561284	900	915	TCGATGTTGAATTAAT	Deoxy, MOE, and cEt	20	7887	7902	4087
561285	902	917	ATTCGATGTTGAATTA	Deoxy, MOE, and cEt	12	7889	7904	4088
561286	904	919	CTATTCGATGTTGAAT	Deoxy, MOE, and cEt	17	7891	7906	4089
561287	906	921	ATCTATTCGATGTTGA	Deoxy, MOE, and cEt	32	7893	7908	4090
561288	908	923	CCATCTATTCGATGTT	Deoxy, MOE, and cEt	69	7895	7910	168
561289	910	925	ATCCATCTATTCGATG	Deoxy, MOE, and cEt	32	7897	7912	4091
561290	912	927	TGATCCATCTATTCGA	Deoxy, MOE, and cEt	41	7899	7914	4092
561291	914	929	TGTGATCCATCTATTC	Deoxy, MOE, and cEt	50	7901	7916	4093
561292	916	931	TTTGTGATCCATCTAT	Deoxy, MOE, and cEt	50	7903	7918	4094
561293	918	933	GTTTTGTGATCCATCT	Deoxy, MOE, and cEt	41	7905	7920	4095
561294	920	935	AAGTTTTGTGATCCAT	Deoxy, MOE, and cEt	56	7907	7922	4096

561295	922	937	TGAAGTTTTGTGATCC	Deoxy, MOE, and cEt	57	7909	7924	4097
561296	924	939	ATTGAAGTTTTGTGAT	Deoxy, MOE, and cEt	0	7911	7926	4098
561450	1386	1401	CAACATTTTGGTTGAT	Deoxy, MOE, and cEt	45	10358	10373	4099
561451	1389	1404	GATCAACATTTTGGTT	Deoxy, MOE, and cEt	33	10361	10376	4100
561452	1391	1406	TGGATCAACATTTTGG	Deoxy, MOE, and cEt	81	10363	10378	123
561453	1393	1408	GATGGATCAACATTTT	Deoxy, MOE, and cEt	59	10365	10380	4101
561455	1397	1412	GTTGGATGGATCAACA	Deoxy, MOE, and cEt	53	10369	10384	4102
561456	1399	1414	CTGTTGGATGGATCAA	Deoxy, MOE, and cEt	71	10371	10386	4103
561457	1401	1416	ATCTGTTGGATGGATC	Deoxy, MOE, and cEt	71	10373	10388	4104
561458	1403	1418	GAATCTGTTGGATGGA	Deoxy, MOE, and cEt	84	10375	10390	124
561459	1405	1420	CTGAATCTGTTGGATG	Deoxy, MOE, and cEt	72	10377	10392	4105
561460	1407	1422	TTCTGAATCTGTTGGA	Deoxy, MOE, and cEt	78	10379	10394	125
561461	1414	1429	CAAAGCTTTCTGAATC	Deoxy, MOE, and cEt	45	10386	10401	4106
561462	1421	1436	GTTCATTCAAAGCTTT	Deoxy, MOE, and cEt	87	10393	10408	126
561463	1423	1438	CAGTTCATTCAAAGCT	Deoxy, MOE, and cEt	85	10395	10410	127
561464	1425	1440	CTCAGTTCATTCAAAG	Deoxy, MOE, and cEt	47	10397	10412	4107
561465	1427	1442	GCCTCAGTTCATTCAA	Deoxy, MOE, and cEt	60	10399	10414	4108
561466	1429	1444	TTGCCTCAGTTCATTC	Deoxy, MOE, and cEt	68	10401	10416	4109
561467	1431	1446	ATTTGCCTCAGTTCAT	Deoxy, MOE, and cEt	61	10403	10418	4110
561468	1433	1448	AAATTTGCCTCAGTTC	Deoxy, MOE, and cEt	48	10405	10420	4111
561469	1436	1451	TTTAAATTTGCCTCAG	Deoxy, MOE, and cEt	59	10408	10423	4112
561470	1438	1453	CTTTTAAATTTGCCTC	Deoxy, MOE, and cEt	50	10410	10425	4113
561471	1440	1455	GCCTTTTAAATTTGCC	Deoxy, MOE, and cEt	73	10412	10427	4114
561472	1452	1467	GTTTAAATTATTGCCT	Deoxy, MOE, and cEt	48	10424	10439	4115
561473	1463	1478	ATGAGGTTAATGTTTA	Deoxy, MOE, and cEt	33	10435	10450	4116
561474	1465	1480	GAATGAGGTTAATGTT	Deoxy, MOE, and cEt	29	10437	10452	4117
561475	1467	1482	TGGAATGAGGTTAATG	Deoxy, MOE, and cEt	66	10439	10454	4118
561476	1469	1484	CTTGGAATGAGGTAA	Deoxy, MOE, and cEt	72	10441	10456	4119
561477	1471	1486	AACTTGGAATGAGGTT	Deoxy, MOE, and cEt	69	10443	10458	4120
561478	1473	1488	TTAACTTGGAATGAGG	Deoxy, MOE, and cEt	74	10445	10460	128
561479	1475	1490	CATTA ACTTGGAATGA	Deoxy, MOE, and cEt	5	10447	10462	4121
561480	1477	1492	CACATTA ACTTGGAAT	Deoxy, MOE, and cEt	26	10449	10464	4122
561481	1479	1494	ACCACATTA ACTTGGA	Deoxy, MOE, and cEt	59	10451	10466	4123
561482	1481	1496	AGACCACATTA ACTTG	Deoxy, MOE, and cEt	76	10453	10468	129
561483	1483	1498	TTAGACCACATTA ACT	Deoxy, MOE, and cEt	47	10455	10470	4124
561484	1485	1500	TATTAGACCACATTAA	Deoxy, MOE, and cEt	38	10457	10472	4125
561485	1487	1502	ATTATTAGACCACATT	Deoxy, MOE, and cEt	59	10459	10474	4126
561486	1489	1504	AGATTATTAGACCACA	Deoxy, MOE, and cEt	84	10461	10476	130
561487	1491	1506	CCAGATTATTAGACCA	Deoxy, MOE, and cEt	93	10463	10478	131
561488	1493	1508	TACCAGATTATTAGAC	Deoxy, MOE, and cEt	22	10465	10480	4127
561489	1495	1510	AATACCAGATTATTAG	Deoxy, MOE, and cEt	48	10467	10482	4128

561490	1497	1512	TTAATACCAGATTATT	Deoxy, MOE, and cEt	22	10469	10484	4129
561491	1499	1514	ATTTAATACCAGATTA	Deoxy, MOE, and cEt	14	10471	10486	4130
561492	1501	1516	GGATTTAATACCAGAT	Deoxy, MOE, and cEt	74	10473	10488	4131
561493	1503	1518	AAGGATTTAATACCAG	Deoxy, MOE, and cEt	70	10475	10490	4132
561494	1505	1520	TTAAGGATTTAATACC	Deoxy, MOE, and cEt	14	10477	10492	4133
561495	1508	1523	CTCTTAAGGATTTAAT	Deoxy, MOE, and cEt	12	10480	10495	4134
561496	1510	1525	TTCTCTTAAGGATTTA	Deoxy, MOE, and cEt	47	10482	10497	4135
561497	1513	1528	GCTTTCTCTTAAGGAT	Deoxy, MOE, and cEt	73	10485	10500	4136
561498	1515	1530	AAGCTTTCTCTTAAGG	Deoxy, MOE, and cEt	59	10487	10502	4137
561499	1517	1532	TCAAGCTTTCTCTTAA	Deoxy, MOE, and cEt	62	10489	10504	4138
561500	1526	1541	ATCTATTTCTCAAGCT	Deoxy, MOE, and cEt	76	10498	10513	132
561501	1547	1562	AGTGACTTTAAGATAA	Deoxy, MOE, and cEt	23	10519	10534	4139
561502	1549	1564	ACAGTGACTTTAAGAT	Deoxy, MOE, and cEt	62	10521	10536	4140
561503	1551	1566	AGACAGTGACTTTAAG	Deoxy, MOE, and cEt	55	10523	10538	4141
561504	1553	1568	ATAGACAGTGACTTTA	Deoxy, MOE, and cEt	74	10525	10540	133
561505	1555	1570	AAATAGACAGTGACTT	Deoxy, MOE, and cEt	59	10527	10542	4142
561506	1557	1572	TTAAATAGACAGTGAC	Deoxy, MOE, and cEt	38	10529	10544	4143
561507	1559	1574	TCTTAAATAGACAGTG	Deoxy, MOE, and cEt	54	10531	10546	4144
561508	1561	1576	AATCTTAAATAGACAG	Deoxy, MOE, and cEt	22	10533	10548	4145
561509	1563	1578	TTAATCTTAAATAGAC	Deoxy, MOE, and cEt	0	10535	10550	4146
561510	1565	1580	GTTTAATCTTAAATAG	Deoxy, MOE, and cEt	0	10537	10552	4147
561511	1569	1584	GTATGTTTAATCTTAA	Deoxy, MOE, and cEt	13	10541	10556	4148
561512	1572	1587	ATTGTATGTTTAATCT	Deoxy, MOE, and cEt	40	10544	10559	4149
561513	1575	1590	GTGATTGTATGTTTAA	Deoxy, MOE, and cEt	71	10547	10562	4150
561514	1578	1593	TATGTGATTGTATGTT	Deoxy, MOE, and cEt	58	10550	10565	4151
561515	1580	1595	GTTATGTGATTGTATG	Deoxy, MOE, and cEt	68	10552	10567	4152
561516	1582	1597	AGGTTATGTGATTGTA	Deoxy, MOE, and cEt	73	10554	10569	4153
561517	1584	1599	TAAGGTTATGTGATTG	Deoxy, MOE, and cEt	64	10556	10571	4154
561518	1586	1601	TTTAAGGTTATGTGAT	Deoxy, MOE, and cEt	0	10558	10573	4155
561519	1588	1603	TCTTTAAGGTTATGTG	Deoxy, MOE, and cEt	53	10560	10575	4156
561520	1590	1605	ATTCTTTAAGGTTATG	Deoxy, MOE, and cEt	29	10562	10577	4157
561521	1592	1607	GTATTCTTTAAGGTTA	Deoxy, MOE, and cEt	24	10564	10579	4158
561522	1594	1609	CGGTATTCTTTAAGGT	Deoxy, MOE, and cEt	70	10566	10581	4159
561523	1596	1611	AACGGTATTCTTTAAG	Deoxy, MOE, and cEt	42	10568	10583	4160
561524	1598	1613	TAAACGGTATTCTTTA	Deoxy, MOE, and cEt	26	10570	10585	4161
561525	1600	1615	TGTAAACGGTATTCTT	Deoxy, MOE, and cEt	59	10572	10587	4162
561526	1602	1617	AATGTAAACGGTATTC	Deoxy, MOE, and cEt	57	10574	10589	4142

Table 29

Inhibition of ANGPTL3 mRNA by oligonucleotides targeting SEQ ID NO: 1 and 2

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	Chemistry	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
561681	N/A	N/A	TCTGGAAGCAGACCTA	Deoxy, MOE, and cEt	37	3096	3111	4164
561682	N/A	N/A	CTTCTGGAAGCAGACC	Deoxy, MOE, and cEt	27	3098	3113	4165
561683	N/A	N/A	AAATAAGGTATAGTGA	Deoxy, MOE, and cEt	2	11084	11099	4166
561684	N/A	N/A	TAGTATTAAGTGTTAA	Deoxy, MOE, and cEt	14	11133	11148	4167
561685	N/A	N/A	TCATAGTATTAAGTGT	Deoxy, MOE, and cEt	0	11136	11151	4168
561686	N/A	N/A	AGATTCCTTTACAATT	Deoxy, MOE, and cEt	21	11160	11175	4169
561687	N/A	N/A	ACAAGATTCCTTTACA	Deoxy, MOE, and cEt	21	11163	11178	4170
561688	N/A	N/A	CTGACAAGATTCCTTT	Deoxy, MOE, and cEt	70	11166	11181	4171
561689	N/A	N/A	AATCTGACAAGATTCC	Deoxy, MOE, and cEt	83	11169	11184	180
561690	N/A	N/A	TGTAATCTGACAAGAT	Deoxy, MOE, and cEt	46	11172	11187	4172
561691	N/A	N/A	TACTGTAATCTGACAA	Deoxy, MOE, and cEt	47	11175	11190	4173
561692	N/A	N/A	TCTTACTGTAATCTGA	Deoxy, MOE, and cEt	50	11178	11193	4174
561693	N/A	N/A	CATTCTTACTGTAATC	Deoxy, MOE, and cEt	40	11181	11196	4175
561694	N/A	N/A	G TTCATTCTTACTGTA	Deoxy, MOE, and cEt	71	11184	11199	4176
561695	N/A	N/A	ATATGTTTCATTCTTAC	Deoxy, MOE, and cEt	2	11188	11203	4177
561696	N/A	N/A	GCCACAAATATGTTCA	Deoxy, MOE, and cEt	80	11195	11210	4178
561697	N/A	N/A	GATGCCACAAATATGT	Deoxy, MOE, and cEt	70	11198	11213	4179
561698	N/A	N/A	CTCGATGCCACAAATA	Deoxy, MOE, and cEt	80	11201	11216	181
561699	N/A	N/A	TAAC TCGATGCCACAA	Deoxy, MOE, and cEt	86	11204	11219	182
561700	N/A	N/A	CTTTAACTCGATGCCA	Deoxy, MOE, and cEt	77	11207	11222	4180
561701	N/A	N/A	AAACTTTAACTCGATG	Deoxy, MOE, and cEt	39	11210	11225	4181
561702	N/A	N/A	TATAAACTTTAACTCG	Deoxy, MOE, and cEt	13	11213	11228	4182
561703	N/A	N/A	CACAGCATATTTAGGG	Deoxy, MOE, and cEt	71	11233	11248	4183
561704	N/A	N/A	TAGAATCACAGCATAT	Deoxy, MOE, and cEt	68	11239	11254	4184
561705	N/A	N/A	TATTAGAATCACAGCA	Deoxy, MOE, and cEt	73	11242	11257	4185
561706	N/A	N/A	AATGTATTAGAATCAC	Deoxy, MOE, and cEt	40	11246	11261	4186
561707	N/A	N/A	ACGAATGTATTAGAAT	Deoxy, MOE, and cEt	22	11249	11264	4187
561708	N/A	N/A	TACACGAATGTATTAG	Deoxy, MOE, and cEt	33	11252	11267	4188
561709	N/A	N/A	ACCTACACGAATGTAT	Deoxy, MOE, and cEt	42	11255	11270	4189
561710	N/A	N/A	AAAACCTACACGAATG	Deoxy, MOE, and cEt	24	11258	11273	4190
561711	N/A	N/A	TTGAAAACCTACACGA	Deoxy, MOE, and cEt	34	11261	11276	4191
561712	N/A	N/A	TACTTGAAAACCTACA	Deoxy, MOE, and cEt	33	11264	11279	4192
561713	N/A	N/A	GTTTATTTCTACTTGA	Deoxy, MOE, and cEt	53	11273	11288	4193
561714	N/A	N/A	GAGGTTTATTTCTACT	Deoxy, MOE, and cEt	69	11276	11291	4194
561715	N/A	N/A	TACGAGGTTTATTTCT	Deoxy, MOE, and cEt	21	11279	11294	4195
561716	N/A	N/A	TGTTACGAGGTTTATT	Deoxy, MOE, and cEt	47	11282	11297	4196

561717	N/A	N/A	ACTTGTTACGAGGTTT	Deoxy, MOE, and cEt	70	11285	11300	4197
561718	N/A	N/A	CAGTAACTTGTTACGA	Deoxy, MOE, and cEt	60	11290	11305	4198
561719	N/A	N/A	GTTCAGTAACTTGTTA	Deoxy, MOE, and cEt	40	11293	11308	4199
561720	N/A	N/A	TCAGGCTGTTTAAACG	Deoxy, MOE, and cEt	59	11308	11323	4200
561721	N/A	N/A	TTGTCAGGCTGTTTAA	Deoxy, MOE, and cEt	74	11311	11326	4201
561722	N/A	N/A	TGCTTGTCAGGCTGTT	Deoxy, MOE, and cEt	82	11314	11329	183
561723	N/A	N/A	ACATGCTTGTCAGGCT	Deoxy, MOE, and cEt	84	11317	11332	184
561724	N/A	N/A	TATACATGCTTGTCAG	Deoxy, MOE, and cEt	75	11320	11335	4202
561725	N/A	N/A	GTCTTTGTTTATTGAA	Deoxy, MOE, and cEt	49	11347	11362	4203
561726	N/A	N/A	TGGGTCTTTGTTTATT	Deoxy, MOE, and cEt	27	11350	11365	4204
561727	N/A	N/A	GACTGGGTCTTTGTTT	Deoxy, MOE, and cEt	20	11353	11368	4205
561728	N/A	N/A	ATAATTTAGGGACTGG	Deoxy, MOE, and cEt	20	11363	11378	4206
561729	N/A	N/A	TCTATAATTTAGGGAC	Deoxy, MOE, and cEt	39	11366	11381	4207
561730	N/A	N/A	CGATAAACATGCAAGA	Deoxy, MOE, and cEt	68	11394	11409	4208
561731	N/A	N/A	TGTCGATAAACATGCA	Deoxy, MOE, and cEt	80	11397	11412	4209
561732	N/A	N/A	TGATGTCGATAAACAT	Deoxy, MOE, and cEt	68	11400	11415	4210
561733	N/A	N/A	TTGTGATGTCGATAAA	Deoxy, MOE, and cEt	28	11403	11418	4211
561734	N/A	N/A	CTGTTGTGATGTCGAT	Deoxy, MOE, and cEt	74	11406	11421	4212
561735	N/A	N/A	GATCTGTTGTGATGTC	Deoxy, MOE, and cEt	59	11409	11424	4213
561736	N/A	N/A	AGGGATCTGTTGTGAT	Deoxy, MOE, and cEt	24	11412	11427	4214
561737	N/A	N/A	TTTAGGGATCTGTTGT	Deoxy, MOE, and cEt	19	11415	11430	4215
561738	N/A	N/A	GGATTTAGGGATCTGT	Deoxy, MOE, and cEt	27	11418	11433	4216
561739	N/A	N/A	GATTTAGGGATTTAGG	Deoxy, MOE, and cEt	44	11425	11440	4217
561740	N/A	N/A	TCTTTAGGGATTTAGG	Deoxy, MOE, and cEt	38	11433	11448	4218
561741	N/A	N/A	TAATCTTTAGGGATTT	Deoxy, MOE, and cEt	0	11436	11451	4219
561742	N/A	N/A	ATCTAATCTTTAGGGA	Deoxy, MOE, and cEt	0	11439	11454	4220
561743	N/A	N/A	TGTATCTAATCTTTAG	Deoxy, MOE, and cEt	15	11442	11457	4221
561744	N/A	N/A	AAATTTGTATCTAATC	Deoxy, MOE, and cEt	21	11447	11462	4222
561745	N/A	N/A	GTAAAAAATTTGTATC	Deoxy, MOE, and cEt	23	11452	11467	4223
561746	N/A	N/A	GTGGTAAAAAATTTGT	Deoxy, MOE, and cEt	32	11455	11470	4224
561747	N/A	N/A	GATACTGTGGTAAAAA	Deoxy, MOE, and cEt	45	11461	11476	4225
561748	N/A	N/A	AGTGATACTGTGGTAA	Deoxy, MOE, and cEt	60	11464	11479	4226
561749	N/A	N/A	ACAAGTGATACTGTGG	Deoxy, MOE, and cEt	75	11467	11482	4227
561750	N/A	N/A	CTGACAAGTGATACTG	Deoxy, MOE, and cEt	59	11470	11485	4228
561751	N/A	N/A	ATTCTGACAAGTGATA	Deoxy, MOE, and cEt	48	11473	11488	4229
561752	N/A	N/A	TAAATTCTGACAAGTG	Deoxy, MOE, and cEt	59	11476	11491	4230
561753	N/A	N/A	TACTGGCAGTTTTAAA	Deoxy, MOE, and cEt	42	11508	11523	4231
561754	N/A	N/A	TCTTACTGGCAGTTTT	Deoxy, MOE, and cEt	51	11511	11526	4232
561755	N/A	N/A	ATTTCTTACTGGCAGT	Deoxy, MOE, and cEt	69	11514	11529	4233
561756	N/A	N/A	AAAATTTCTTACTGGC	Deoxy, MOE, and cEt	57	11517	11532	4234
561757	N/A	N/A	AACAAATGGGTTTAAT	Deoxy, MOE, and cEt	0	11535	11550	4235

562374	N/A	N/A	GAATATTTGCAAGTCT	Deoxy, MOE, and cEt	68	9230	9245	4236
562375	N/A	N/A	GTAGAGGAATATTTGC	Deoxy, MOE, and cEt	83	9236	9251	151
562376	N/A	N/A	TCATTGGTAGAGGAAT	Deoxy, MOE, and cEt	23	9242	9257	4237
562377	N/A	N/A	ATATTTTAAAGTCTCG	Deoxy, MOE, and cEt	17	9258	9273	4238
562378	N/A	N/A	GTTACATTATTATAGA	Deoxy, MOE, and cEt	29	9273	9288	4239
562379	N/A	N/A	GTGAAATGTGTTACAT	Deoxy, MOE, and cEt	54	9282	9297	4240
562380	N/A	N/A	TCACCAGTGAAATGTG	Deoxy, MOE, and cEt	64	9288	9303	4241
562381	N/A	N/A	CATGTTTCACCAGTGA	Deoxy, MOE, and cEt	78	9294	9309	4242
562382	N/A	N/A	ACAAGACATGTTTCAC	Deoxy, MOE, and cEt	36	9300	9315	4243
562383	N/A	N/A	CATATGACAAGACATG	Deoxy, MOE, and cEt	42	9306	9321	4244
562384	N/A	N/A	CTATAATGCATATGAC	Deoxy, MOE, and cEt	5	9314	9329	4245
562385	N/A	N/A	TCCTTTCTATAATGCA	Deoxy, MOE, and cEt	65	9320	9335	4246
562386	N/A	N/A	TGATTATCCTTTCTAT	Deoxy, MOE, and cEt	27	9326	9341	4247
562387	N/A	N/A	AAAGTCTGATTATCCT	Deoxy, MOE, and cEt	90	9332	9347	152
562388	N/A	N/A	TAAGTCAAAGTCTGAT	Deoxy, MOE, and cEt	59	9338	9353	4248
562389	N/A	N/A	GTGCACAAAAATGTTA	Deoxy, MOE, and cEt	42	9366	9381	4249
562390	N/A	N/A	AGCTATGTGCACAAAA	Deoxy, MOE, and cEt	77	9372	9387	4250
562391	N/A	N/A	GAAGATAGCTATGTGC	Deoxy, MOE, and cEt	64	9378	9393	4251
562392	N/A	N/A	TTTATTGAAGATAGCT	Deoxy, MOE, and cEt	33	9384	9399	4252
562393	N/A	N/A	TCATTTTAGTGTATCT	Deoxy, MOE, and cEt	40	9424	9439	4253
562394	N/A	N/A	CCTTGATCATTTTAGT	Deoxy, MOE, and cEt	15	9430	9445	4254
562395	N/A	N/A	TGAATCCCTTGATCAT	Deoxy, MOE, and cEt	59	9436	9451	4255
562396	N/A	N/A	TAGTCTTGAATCCCTT	Deoxy, MOE, and cEt	83	9442	9457	153
562397	N/A	N/A	GTTGTTTAGTCTTGAA	Deoxy, MOE, and cEt	65	9448	9463	4256
562398	N/A	N/A	AATTGAGTTGTTTAGT	Deoxy, MOE, and cEt	21	9454	9469	4257
562399	N/A	N/A	GCAACTAATTGAGTTG	Deoxy, MOE, and cEt	15	9460	9475	4258
562400	N/A	N/A	ATTGGTGCAACTAATT	Deoxy, MOE, and cEt	25	9466	9481	4259
562401	N/A	N/A	GTTTTTTATTGGTGCA	Deoxy, MOE, and cEt	53	9473	9488	4260
562402	N/A	N/A	GGACACTGACAGTTTT	Deoxy, MOE, and cEt	43	9496	9511	4261
562403	N/A	N/A	CAGGTTGGACACTGAC	Deoxy, MOE, and cEt	23	9502	9517	4262
562404	N/A	N/A	TAAGTACAGGTTGGAC	Deoxy, MOE, and cEt	33	9508	9523	4263
562405	N/A	N/A	AGTTATTAAGTACAGG	Deoxy, MOE, and cEt	34	9514	9529	4264
562406	N/A	N/A	TCTGTGAGTTATTAAG	Deoxy, MOE, and cEt	10	9520	9535	4265
562407	N/A	N/A	ACCAAAATTCTCCTGA	Deoxy, MOE, and cEt	1	9554	9569	4266
562408	N/A	N/A	ACCTGAATAACCCTCT	Deoxy, MOE, and cEt	73	9811	9826	4267
562409	N/A	N/A	GGTATCAGAAAAAGAT	Deoxy, MOE, and cEt	14	9827	9842	4268
562410	N/A	N/A	AGTATTGGTATCAGAA	Deoxy, MOE, and cEt	13	9833	9848	4269
562411	N/A	N/A	GGAAGATACTTTGAAG	Deoxy, MOE, and cEt	25	9861	9876	4270
562412	N/A	N/A	AATGTGGGAAGATACT	Deoxy, MOE, and cEt	23	9867	9882	4271
562413	N/A	N/A	CAGATAATAGCTAATA	Deoxy, MOE, and cEt	29	9882	9897	4272
562414	N/A	N/A	TCATTGCAGATAATAG	Deoxy, MOE, and cEt	45	9888	9903	4273

562415	N/A	N/A	AAGTTGTCATTGCAGA	Deoxy, MOE, and cEt	86	9894	9909	154
562416	N/A	N/A	GATTCGGATTTTAAA	Deoxy, MOE, and cEt	19	9909	9924	4274
562417	N/A	N/A	ATTTGGGATTCGGATT	Deoxy, MOE, and cEt	34	9915	9930	4275
562418	N/A	N/A	ACGCTTATTTGGGATT	Deoxy, MOE, and cEt	64	9921	9936	4276
562419	N/A	N/A	TCTAGAGAGAAAACGC	Deoxy, MOE, and cEt	64	9933	9948	4277
562420	N/A	N/A	AGTTAAGAGGTTTTCG	Deoxy, MOE, and cEt	34	9949	9964	4278
562421	N/A	N/A	CATTATAGTTAAGAGG	Deoxy, MOE, and cEt	24	9955	9970	4279
562422	N/A	N/A	CACTTTCATTATAGTT	Deoxy, MOE, and cEt	13	9961	9976	4280
562423	N/A	N/A	TAGAATGAACACTTTC	Deoxy, MOE, and cEt	63	9970	9985	4281
562424	N/A	N/A	TTGAACTAGAATGAAC	Deoxy, MOE, and cEt	16	9976	9991	4282
562425	N/A	N/A	ACCTGATTGAACTAGA	Deoxy, MOE, and cEt	51	9982	9997	4283
562426	N/A	N/A	TAAAATACCTGATTGA	Deoxy, MOE, and cEt	19	9988	10003	4284
562427	N/A	N/A	TAGAGGTAAAATACCT	Deoxy, MOE, and cEt	12	9994	10009	4285
562428	N/A	N/A	GAAGATTAGAGGTAAA	Deoxy, MOE, and cEt	1	10000	10015	4286
562429	N/A	N/A	TCTGAGGAAGATTAGA	Deoxy, MOE, and cEt	31	10006	10021	4287
562430	N/A	N/A	TATACACTACCAAAAA	Deoxy, MOE, and cEt	0	10030	10045	4288
562431	N/A	N/A	ATAATCTATACACTAC	Deoxy, MOE, and cEt	0	10036	10051	4289
562432	N/A	N/A	TAAGTCCCAATTTTAA	Deoxy, MOE, and cEt	33	10065	10080	4290
562433	N/A	N/A	TCTGTATAAGTCCCAA	Deoxy, MOE, and cEt	89	10071	10086	155
562434	N/A	N/A	CCAGTTTTAAATAATC	Deoxy, MOE, and cEt	20	10085	10100	4291
562435	N/A	N/A	TGTATCCCAGTTTTAA	Deoxy, MOE, and cEt	44	10091	10106	4292
562436	N/A	N/A	GATGCATGTATCCCAG	Deoxy, MOE, and cEt	91	10097	10112	156
562437	N/A	N/A	GTTTTAGATGCATGTA	Deoxy, MOE, and cEt	69	10103	10118	4293
562438	N/A	N/A	TACAGTGTTTTAGATG	Deoxy, MOE, and cEt	28	10109	10124	4294
562439	N/A	N/A	GTAAGTTTATCTTCCT	Deoxy, MOE, and cEt	78	10138	10153	157
562440	N/A	N/A	TTCCCCGTAAGTTTAT	Deoxy, MOE, and cEt	33	10144	10159	4295
562441	N/A	N/A	CTGTATTTCCCCGTAA	Deoxy, MOE, and cEt	55	10150	10165	4296
562442	N/A	N/A	CTGTACTGTATTTCC	Deoxy, MOE, and cEt	79	10156	10171	158
562443	N/A	N/A	TAGTTACTGTTACTGT	Deoxy, MOE, and cEt	70	10162	10177	4297
562444	N/A	N/A	CGTATGTAGTTACTGT	Deoxy, MOE, and cEt	66	10168	10183	4298
562445	N/A	N/A	AATGGGTACAGACTCG	Deoxy, MOE, and cEt	72	10182	10197	4299
562446	N/A	N/A	GCAATTTAATGGGTAC	Deoxy, MOE, and cEt	59	10189	10204	4300
562447	N/A	N/A	GATAGATATGCAATTT	Deoxy, MOE, and cEt	20	10198	10213	4301
562448	N/A	N/A	AAAGGAGATAGATATG	Deoxy, MOE, and cEt	22	10204	10219	4302
562449	N/A	N/A	CCTCCTAAAGGAGATA	Deoxy, MOE, and cEt	42	10210	10225	4303
562450	N/A	N/A	CACCAGCCTCCTAAAG	Deoxy, MOE, and cEt	37	10216	10231	4304
544120	707	726	AGTTCTTGGTGCTCTTGGCT	5-10-5 MOE	83	6720	6739	15
560990	709	724	TTCTTGGTGCTCTTGG	Deoxy, MOE, and cEt	89	6722	6737	111
337487	804	823	CACTTGTATGTTACCTCTG	5-10-5 MOE	81	7389	7408	28
561373	1197	1212	TTTGTGATCCCAAGTA	Deoxy, MOE, and cEt	40	9772	9787	4305
561374	1199	1214	GCTTTGTGATCCCAAG	Deoxy, MOE, and cEt	76	9774	9789	4306

561375	1201	1216	TTGCTTTGTGATCCCA	Deoxy, MOE, and cEt	82	9776	9791	4307
561376	1203	1218	TTTTGCTTTGTGATCC	Deoxy, MOE, and cEt	40	9778	9793	4308
561377	1205	1220	CCTTTTGCTTTGTGAT	Deoxy, MOE, and cEt	38	9780	9795	4309
561378	1207	1222	GTCCTTTTGCTTTGTG	Deoxy, MOE, and cEt	75	9782	9797	4310
561379	1209	1224	GTGTCCTTTTGCTTTG	Deoxy, MOE, and cEt	40	9784	9799	4311
561380	1212	1227	GAAGTGTCTTTTGCT	Deoxy, MOE, and cEt	23	9787	9802	4312
561381	1214	1229	TTGAAGTGTCTTTTG	Deoxy, MOE, and cEt	26	9789	9804	4313
561382	1216	1231	AGTTGAAGTGTCTTT	Deoxy, MOE, and cEt	34	9791	9806	4314
561383	1218	1233	ACAGTTGAAGTGTCT	Deoxy, MOE, and cEt	27	9793	9808	4315
561384	1220	1235	GGACAGTTGAAGTGTC	Deoxy, MOE, and cEt	19	9795	9810	4316
561385	1222	1237	CTGGACAGTTGAAGTG	Deoxy, MOE, and cEt	34	9797	9812	4317
561386	1224	1239	CTCTGGACAGTTGAAG	Deoxy, MOE, and cEt	19	9799	9814	4318
561387	1226	1241	CCCTCTGGACAGTTGA	Deoxy, MOE, and cEt	54	9801	9816	4319
561388	1228	1243	AACCCTCTGGACAGTT	Deoxy, MOE, and cEt	50	9803	9818	4320
561389	1230	1245	ATAACCCTCTGGACAG	Deoxy, MOE, and cEt	35	9805	9820	4321
561390	1232	1247	GAATAACCCTCTGGAC	Deoxy, MOE, and cEt	34	9807	9822	4322
561391	1234	1249	CTGAATAACCCTCTGG	Deoxy, MOE, and cEt	62	9809	9824	4323
561392	1236	1251	TCCTGAATAACCCTCT	Deoxy, MOE, and cEt	57	N/A	N/A	4324
561393	1238	1253	CCTCCTGAATAACCCT	Deoxy, MOE, and cEt	30	N/A	N/A	4325
561394	1246	1261	ACCACCAGCCTCCTGA	Deoxy, MOE, and cEt	70	N/A	N/A	4326
561395	1251	1266	ATGCCACCACCAGCCT	Deoxy, MOE, and cEt	68	10223	10238	4327
561396	1253	1268	TCATGCCACCACCAGC	Deoxy, MOE, and cEt	72	10225	10240	4328
561397	1255	1270	CATCATGCCACCACCA	Deoxy, MOE, and cEt	67	10227	10242	4329
561398	1257	1272	CTCATCATGCCACCAC	Deoxy, MOE, and cEt	77	10229	10244	172
561399	1259	1274	CACTCATCATGCCACC	Deoxy, MOE, and cEt	74	10231	10246	2330
561400	1261	1276	CACACTCATCATGCCA	Deoxy, MOE, and cEt	80	10233	10248	173
561401	1263	1278	TCCACACTCATCATGC	Deoxy, MOE, and cEt	64	10235	10250	4331
561402	1265	1280	TCTCCACACTCATCAT	Deoxy, MOE, and cEt	42	10237	10252	4332
561403	1267	1282	TTTCTCCACACTCATC	Deoxy, MOE, and cEt	47	10239	10254	4333
561404	1269	1284	GTTTTCTCCACACTCA	Deoxy, MOE, and cEt	77	10241	10256	4334
561405	1272	1287	GTTGTTTTCTCCACAC	Deoxy, MOE, and cEt	53	10244	10259	4335
561406	1274	1289	AGGTTGTTTTCTCCAC	Deoxy, MOE, and cEt	67	10246	10261	4336
561407	1276	1291	TTAGGTTGTTTTCTCC	Deoxy, MOE, and cEt	73	10248	10263	4337
561408	1282	1297	TACCATTTAGGTTGTT	Deoxy, MOE, and cEt	30	10254	10269	4338
561409	1284	1299	TTTACCATTTAGGTTG	Deoxy, MOE, and cEt	22	10256	10271	4339
561410	1286	1301	TATTTACCATTTAGGT	Deoxy, MOE, and cEt	24	10258	10273	4340
561411	1292	1307	TTGTTATATTTACCAT	Deoxy, MOE, and cEt	41	10264	10279	4341
561412	1294	1309	GTTTGTTATATTTACC	Deoxy, MOE, and cEt	37	10266	10281	4342
561413	1298	1313	CTTGGTTTGTTATATT	Deoxy, MOE, and cEt	45	10270	10285	4343
561414	1300	1315	CTCTTGGTTTGTTATA	Deoxy, MOE, and cEt	73	10272	10287	4344
561415	1302	1317	TGCTCTTGGTTTGTTA	Deoxy, MOE, and cEt	45	10274	10289	4345

561416	1304	1319	TTTGCTCTTGGTTTGT	Deoxy, MOE, and cEt	67	10276	10291	4346
561417	1307	1322	GATTTTGCTCTTGGTT	Deoxy, MOE, and cEt	75	10279	10294	4347
561418	1309	1324	TAGATTTTGCTCTTGG	Deoxy, MOE, and cEt	87	10281	10296	169
561419	1311	1326	CTTAGATTTTGCTCTT	Deoxy, MOE, and cEt	64	10283	10298	4348
561420	1313	1328	GGCTTAGATTTTGCTC	Deoxy, MOE, and cEt	58	10285	10300	4349
561421	1315	1330	CTGGCTTAGATTTTGC	Deoxy, MOE, and cEt	70	10287	10302	4350
561422	1317	1332	CTCTGGCTTAGATTTT	Deoxy, MOE, and cEt	38	10289	10304	4351
561423	1319	1334	CTCTCTGGCTTAGATT	Deoxy, MOE, and cEt	63	10291	10306	4352
561424	1321	1336	TCCTCTCTGGCTTAGA	Deoxy, MOE, and cEt	76	10293	10308	4353
561425	1323	1338	TCTCCTCTCTGGCTTA	Deoxy, MOE, and cEt	67	10295	10310	4354
561426	1332	1347	TAATCCTCTTCTCCTC	Deoxy, MOE, and cEt	50	10304	10319	4355
561427	1334	1349	GATAATCCTCTTCTCC	Deoxy, MOE, and cEt	36	10306	10321	4356
561428	1336	1351	AAGATAATCCTCTTCT	Deoxy, MOE, and cEt	43	10308	10323	4357
561429	1338	1353	CCAAGATAATCCTCTT	Deoxy, MOE, and cEt	59	10310	10325	4358
561430	1340	1355	TTCCAAGATAATCCTC	Deoxy, MOE, and cEt	65	10312	10327	4359
561431	1342	1357	ACTTCCAAGATAATCC	Deoxy, MOE, and cEt	74	10314	10329	4360
561432	1344	1359	AGACTTCCAAGATAAT	Deoxy, MOE, and cEt	52	10316	10331	4361
561433	1346	1361	TGAGACTTCCAAGATA	Deoxy, MOE, and cEt	49	10318	10333	4362
561434	1348	1363	TTTGAGACTTCCAAGA	Deoxy, MOE, and cEt	47	10320	10335	4363
561435	1350	1365	ATTTTGAGACTTCCAA	Deoxy, MOE, and cEt	64	10322	10337	4364
561436	1352	1367	CCATTTTGAGACTTCC	Deoxy, MOE, and cEt	84	10324	10339	170
561437	1354	1369	TTCCATTTTGAGACTT	Deoxy, MOE, and cEt	67	10326	10341	4365
561438	1356	1371	CCTTCCATTTTGAGAC	Deoxy, MOE, and cEt	53	10328	10343	4366
561439	1358	1373	AACCTTCCATTTTGAG	Deoxy, MOE, and cEt	37	10330	10345	4367
561440	1360	1375	ATAACCTTCCATTTTG	Deoxy, MOE, and cEt	50	10332	10347	4368
561441	1362	1377	GTATAACCTTCCATTT	Deoxy, MOE, and cEt	27	10334	10349	4369
561442	1364	1379	GAGTATAACCTTCCAT	Deoxy, MOE, and cEt	65	10336	10351	4370
561443	1366	1381	TAGAGTATAACCTTCC	Deoxy, MOE, and cEt	84	10338	10353	171
561444	1368	1383	TATAGAGTATAACCTT	Deoxy, MOE, and cEt	17	10340	10355	4371
561445	1370	1385	TTTATAGAGTATAACC	Deoxy, MOE, and cEt	37	10342	10357	4372
561446	1373	1388	GATTTTATAGAGTATA	Deoxy, MOE, and cEt	28	10345	10360	4373
561447	1375	1390	TTGATTTTATAGAGTA	Deoxy, MOE, and cEt	21	10347	10362	4374
561448	1377	1392	GGTTGATTTTATAGAG	Deoxy, MOE, and cEt	28	10349	10364	4375
561449	1379	1394	TTGGTTGATTTTATAG	Deoxy, MOE, and cEt	22	10351	10366	4376
567295	1452	1471	TAATGTTTAAATTATTGCCT	5-10-5 MOE	43	10424	10443	4377
567296	1455	1474	GGTTAATGTTTAAATTATTG	5-10-5 MOE	22	10427	10446	4378
567297	1456	1475	AGGTTAATGTTTAAATTATT	5-10-5 MOE	0	10428	10447	4379
567298	1457	1476	GAGGTTAATGTTTAAATTAT	5-10-5 MOE	0	10429	10448	4380
567299	1458	1477	TGAGGTTAATGTTTAAATTA	5-10-5 MOE	6	10430	10449	4381
567300	1460	1479	AATGAGGTTAATGTTTAAAT	5-10-5 MOE	14	10432	10451	4382
567301	1461	1480	GAATGAGGTTAATGTTTAAA	5-10-5 MOE	5	10433	10452	4383

WO 2015/100394

PCT/US2014/072303

567302	1462	1481	GGAATGAGGTTAATGTTTAA	5-10-5 MOE	27	10434	10453	4384
567303	1463	1482	TGGAATGAGGTTAATGTTTA	5-10-5 MOE	32	10435	10454	4385
567304	1464	1483	TTGGAATGAGGTTAATGTTT	5-10-5 MOE	37	10436	10455	4386
567305	1465	1484	CTTGGAATGAGGTTAATGTT	5-10-5 MOE	25	10437	10456	4387
567306	1468	1487	TAACTTGGAATGAGGTTAAT	5-10-5 MOE	29	10440	10459	4388
567307	1469	1488	TTAACTTGGAATGAGGTTAA	5-10-5 MOE	44	10441	10460	4389
337513	1470	1489	ATTA ACTTGGAATGAGGTTA	5-10-5 MOE	52	10442	10461	4390
567308	1471	1490	CATTA ACTTGGAATGAGGTT	5-10-5 MOE	62	10443	10462	4391
567309	1472	1491	ACATTA ACTTGGAATGAGGT	5-10-5 MOE	58	10444	10463	4392
567310	1473	1492	CACATTA ACTTGGAATGAGG	5-10-5 MOE	78	10445	10464	92
567311	1475	1494	ACCACATTA ACTTGGAATGA	5-10-5 MOE	59	10447	10466	4393
567312	1476	1495	GACCACATTA ACTTGGAATG	5-10-5 MOE	57	10448	10467	4394
337514	1477	1496	AGACCACATTA ACTTGGAAT	5-10-5 MOE	71	10449	10468	4395
567313	1478	1497	TAGACCACATTA ACTTGGA	5-10-5 MOE	43	10450	10469	4396
567314	1479	1498	TTAGACCACATTA ACTTGGA	5-10-5 MOE	59	10451	10470	4397
567315	1480	1499	ATTAGACCACATTA ACTTGG	5-10-5 MOE	70	10452	10471	4398
567316	1481	1500	TATTAGACCACATTA ACTTG	5-10-5 MOE	53	10453	10472	4399
567317	1482	1501	TTATTAGACCACATTA ACTT	5-10-5 MOE	49	10454	10473	4400
567318	1483	1502	ATTATTAGACCACATTA ACT	5-10-5 MOE	41	10455	10474	4401
567319	1484	1503	GATTATTAGACCACATTA AC	5-10-5 MOE	47	10456	10475	4402
567320	1487	1506	CCAGATTATTAGACCACATT	5-10-5 MOE	86	10459	10478	93
567321	1489	1508	TACCAGATTATTAGACCACA	5-10-5 MOE	85	10461	10480	94
337516	1490	1509	ATACCAGATTATTAGACCAC	5-10-5 MOE	77	10462	10481	86
567322	1491	1510	AATACCAGATTATTAGACCA	5-10-5 MOE	50	10463	10482	4403
567323	1492	1511	TAATACCAGATTATTAGACC	5-10-5 MOE	56	10464	10483	4404
567324	1494	1513	TTTAATACCAGATTATTAGA	5-10-5 MOE	9	10466	10485	4405
567325	1495	1514	ATTTAATACCAGATTATTAG	5-10-5 MOE	24	10467	10486	4406
567326	1496	1515	GATTTAATACCAGATTATTA	5-10-5 MOE	37	10468	10487	4407
567327	1500	1519	TAAGGATTTAATACCAGATT	5-10-5 MOE	60	10472	10491	4408
567328	1507	1526	TTTCTCTTAAGGATTTAATA	5-10-5 MOE	34	10479	10498	4409
567329	1508	1527	CTTTCTCTTAAGGATTTAAT	5-10-5 MOE	46	10480	10499	4410
567330	1509	1528	GCTTTCTCTTAAGGATTTAA	5-10-5 MOE	75	10481	10500	95
567331	1510	1529	AGCTTTCTCTTAAGGATTTA	5-10-5 MOE	59	10482	10501	4411
567332	1511	1530	AAGCTTTCTCTTAAGGATTT	5-10-5 MOE	30	10483	10502	4412
567333	1513	1532	TCAAGCTTTCTCTTAAGGAT	5-10-5 MOE	65	10485	10504	4413
567334	1514	1533	CTCAAGCTTTCTCTTAAGGA	5-10-5 MOE	77	10486	10505	96
567335	1515	1534	TCTCAAGCTTTCTCTTAAGG	5-10-5 MOE	75	10487	10506	97
567336	1516	1535	TTCTCAAGCTTTCTCTTAAG	5-10-5 MOE	59	10488	10507	4414
567337	1517	1536	TTTCTCAAGCTTTCTCTTAA	5-10-5 MOE	52	10489	10508	4415
567338	1521	1540	TCTATTTCTCAAGCTTTCTC	5-10-5 MOE	68	10493	10512	4416
567339	1522	1541	ATCTATTTCTCAAGCTTTCT	5-10-5 MOE	71	10494	10513	4417

567340	1523	1542	AATCTATTTCTCAAGCTTTC	5-10-5 MOE	74	10495	10514	4418
567341	1524	1543	AAATCTATTTCTCAAGCTTT	5-10-5 MOE	63	10496	10515	4419
567342	1525	1544	AAAATCTATTTCTCAAGCTT	5-10-5 MOE	54	10497	10516	4420
567343	1532	1551	GATAAAAAAAAAATCTATTTCT	5-10-5 MOE	30	10504	10523	4421
567344	1548	1567	TAGACAGTGACTTTAAGATA	5-10-5 MOE	37	10520	10539	4422
567345	1549	1568	ATAGACAGTGACTTTAAGAT	5-10-5 MOE	29	10521	10540	4423
567346	1550	1569	AATAGACAGTGACTTTAAGA	5-10-5 MOE	48	10522	10541	4424
567347	1551	1570	AAATAGACAGTGACTTTAAG	5-10-5 MOE	26	10523	10542	4425
567348	1552	1571	TAAATAGACAGTGACTTTAA	5-10-5 MOE	26	10524	10543	4426
567349	1553	1572	TTAAATAGACAGTGACTTTA	5-10-5 MOE	50	10525	10544	4427
567350	1554	1573	CTTAAATAGACAGTGACTTT	5-10-5 MOE	63	10526	10545	4428
567351	1555	1574	TCTTAAATAGACAGTGACTT	5-10-5 MOE	57	10527	10546	4429
567352	1556	1575	ATCTTAAATAGACAGTGACT	5-10-5 MOE	69	10528	10547	4430
567353	1557	1576	AATCTTAAATAGACAGTGAC	5-10-5 MOE	40	10529	10548	4431
567354	1558	1577	TAATCTTAAATAGACAGTGA	5-10-5 MOE	30	10530	10549	4432
567355	1559	1578	TTAATCTTAAATAGACAGTG	5-10-5 MOE	25	10531	10550	4433
567356	1560	1579	TTTAATCTTAAATAGACAGT	5-10-5 MOE	0	10532	10551	4434
567357	1561	1580	GTTTAATCTTAAATAGACAG	5-10-5 MOE	34	10533	10552	4435
567358	1562	1581	TGTTTAATCTTAAATAGACA	5-10-5 MOE	5	10534	10553	4436
567359	1563	1582	ATGTTTAATCTTAAATAGAC	5-10-5 MOE	0	10535	10554	4437
567360	1567	1586	TTGTATGTTTAATCTTAAAT	5-10-5 MOE	0	10539	10558	4438
567361	1568	1587	ATTGTATGTTTAATCTTAAA	5-10-5 MOE	8	10540	10559	4439
567362	1569	1588	GATTGTATGTTTAATCTTAA	5-10-5 MOE	20	10541	10560	4440
567363	1570	1589	TGATTGTATGTTTAATCTTA	5-10-5 MOE	29	10542	10561	4441
567364	1574	1593	TATGTGATTGTATGTTTAAT	5-10-5 MOE	7	10546	10565	4442
567365	1576	1595	GTTATGTGATTGTATGTTTA	5-10-5 MOE	43	10548	10567	4443
567366	1580	1599	TAAGGTTATGTGATTGTATG	5-10-5 MOE	28	10552	10571	4444
567367	1581	1600	TTAAGGTTATGTGATTGTAT	5-10-5 MOE	31	10553	10572	4445
567368	1585	1604	TTCTTTAAGGTTATGTGATT	5-10-5 MOE	12	10557	10576	4446
561527	1604	1619	GAAATGTAAACGGTAT	Deoxy, MOE, and cEt	47	10576	10591	4447
561528	1606	1621	GAGAAATGTAAACGGT	Deoxy, MOE, and cEt	89	10578	10593	174
561529	1608	1623	TTGAGAAATGTAAACG	Deoxy, MOE, and cEt	55	10580	10595	4448
561530	1611	1626	TGATTGAGAAATGTAA	Deoxy, MOE, and cEt	18	10583	10598	4449
561531	1613	1628	TTTGATTGAGAAATGT	Deoxy, MOE, and cEt	30	10585	10600	4450
561532	1619	1634	AAGAATTTTGATTGAG	Deoxy, MOE, and cEt	53	10591	10606	4451
561533	1621	1636	ATAAGAATTTTGATTG	Deoxy, MOE, and cEt	29	10593	10608	4452
561534	1632	1647	CAAATAGTATTATAAG	Deoxy, MOE, and cEt	6	10604	10619	4453
561535	1653	1668	CCCACATCACAAAATT	Deoxy, MOE, and cEt	70	10625	10640	4454
561536	1657	1672	GATTCCCACATCACAA	Deoxy, MOE, and cEt	77	10629	10644	4455
561537	1659	1674	TTGATTCCCACATCAC	Deoxy, MOE, and cEt	78	10631	10646	4456
561538	1661	1676	AATTGATTCCCACATC	Deoxy, MOE, and cEt	68	10633	10648	4457

561539	1663	1678	AAAATTGATTCCCACA	Deoxy, MOE, and cEt	72	10635	10650	4458
561540	1665	1680	CTAAAATTGATTCCCA	Deoxy, MOE, and cEt	54	10637	10652	4459
561541	1668	1683	CATCTAAAATTGATTC	Deoxy, MOE, and cEt	0	10640	10655	4460
561542	1670	1685	ACCATCTAAAATTGAT	Deoxy, MOE, and cEt	35	10642	10657	4461
561543	1672	1687	TGACCATCTAAAATTG	Deoxy, MOE, and cEt	55	10644	10659	4462
561544	1674	1689	TGTGACCATCTAAAAT	Deoxy, MOE, and cEt	56	10646	10661	4463
561545	1676	1691	ATTGTGACCATCTAAA	Deoxy, MOE, and cEt	73	10648	10663	4464
561546	1678	1693	AGATTGTGACCATCTA	Deoxy, MOE, and cEt	67	10650	10665	4465
561547	1680	1695	CTAGATTGTGACCATC	Deoxy, MOE, and cEt	50	10652	10667	4466
561548	1682	1697	ATCTAGATTGTGACCA	Deoxy, MOE, and cEt	77	10654	10669	4467
561549	1684	1699	TAATCTAGATTGTGAC	Deoxy, MOE, and cEt	55	10656	10671	4468
561550	1686	1701	TATAATCTAGATTGTG	Deoxy, MOE, and cEt	28	10658	10673	4469
561551	1688	1703	ATTATAATCTAGATTG	Deoxy, MOE, and cEt	52	10660	10675	4470
561552	1690	1705	TGATTATAATCTAGAT	Deoxy, MOE, and cEt	43	10662	10677	4471
561553	1692	1707	ATTGATTATAATCTAG	Deoxy, MOE, and cEt	53	10664	10679	4472
561554	1694	1709	CTATTGATTATAATCT	Deoxy, MOE, and cEt	54	10666	10681	4473
561555	1696	1711	ACCTATTGATTATAAT	Deoxy, MOE, and cEt	44	10668	10683	4474
561556	1698	1713	TCACCTATTGATTATA	Deoxy, MOE, and cEt	52	10670	10685	4475
561557	1700	1715	G TTCACCTATTGATTA	Deoxy, MOE, and cEt	50	10672	10687	4476
561558	1702	1717	AAGTTCACCTATTGAT	Deoxy, MOE, and cEt	58	10674	10689	4477
561559	1704	1719	ATAAGTTCACCTATTG	Deoxy, MOE, and cEt	66	10676	10691	4478
561560	1706	1721	TAATAAGTTCACCTAT	Deoxy, MOE, and cEt	38	10678	10693	4479
561561	1708	1723	TTTAATAAGTTCACCT	Deoxy, MOE, and cEt	50	10680	10695	4480
561562	1710	1725	TATTTAATAAGTTCAC	Deoxy, MOE, and cEt	32	10682	10697	4481
561563	1712	1727	GTTATTTAATAAGTTC	Deoxy, MOE, and cEt	47	10684	10699	4482
561564	1761	1776	CATATGATGCCTTTTA	Deoxy, MOE, and cEt	63	10733	10748	4483
561565	1763	1778	CTCATATGATGCCTTT	Deoxy, MOE, and cEt	81	10735	10750	175
561566	1765	1780	AGCTCATATGATGCCT	Deoxy, MOE, and cEt	81	10737	10752	176
561567	1767	1782	TTAGCTCATATGATGC	Deoxy, MOE, and cEt	84	10739	10754	177
561568	1769	1784	TATTAGCTCATATGAT	Deoxy, MOE, and cEt	46	10741	10756	4484
561569	1771	1786	GATATTAGCTCATATG	Deoxy, MOE, and cEt	49	10743	10758	4485
561570	1773	1788	GTGATATTAGCTCATA	Deoxy, MOE, and cEt	81	10745	10760	4486
561571	1775	1790	TTGTGATATTAGCTCA	Deoxy, MOE, and cEt	85	10747	10762	178
561572	1777	1792	AGTTGTGATATTAGCT	Deoxy, MOE, and cEt	68	10749	10764	4487
561573	1779	1794	AAAGTTGTGATATTAG	Deoxy, MOE, and cEt	45	10751	10766	4488
561574	1781	1796	GGAAAGTTGTGATATT	Deoxy, MOE, and cEt	27	10753	10768	4489
561575	1783	1798	TGGGAAAGTTGTGATA	Deoxy, MOE, and cEt	36	10755	10770	4490
561576	1785	1800	ACTGGGAAAGTTGTGA	Deoxy, MOE, and cEt	83	10757	10772	179
561577	1787	1802	AAACTGGGAAAGTTGT	Deoxy, MOE, and cEt	56	10759	10774	4491
561578	1789	1804	TTAAACTGGGAAAGTT	Deoxy, MOE, and cEt	44	10761	10776	4492
561579	1794	1809	GTTTTTTAAACTGGGA	Deoxy, MOE, and cEt	58	10766	10781	4493

561580	1796	1811	TAGTTTTTTTAAACTGG	Deoxy, MOE, and cEt	0	10768	10783	4494
561581	1802	1817	GAGTACTAGTTTTTTA	Deoxy, MOE, and cEt	18	10774	10789	4495
561582	1804	1819	AAGAGTACTAGTTTTT	Deoxy, MOE, and cEt	55	10776	10791	4496
561583	1806	1821	ACAAGAGTACTAGTTT	Deoxy, MOE, and cEt	51	10778	10793	4497
561584	1808	1823	TAACAAGAGTACTAGT	Deoxy, MOE, and cEt	53	10780	10795	4498
561585	1810	1825	TTTAACAAGAGTACTA	Deoxy, MOE, and cEt	48	10782	10797	4499
561586	1812	1827	GTTTTAACAAGAGTAC	Deoxy, MOE, and cEt	49	10784	10799	4500
561587	1814	1829	GAGTTTTTAACAAGAGT	Deoxy, MOE, and cEt	54	10786	10801	4501
561588	1816	1831	TAGAGTTTTTAACAAGA	Deoxy, MOE, and cEt	9	10788	10803	4502
561589	1819	1834	GTTTAGAGTTTTTAACA	Deoxy, MOE, and cEt	24	10791	10806	4503
561590	1822	1837	CAAGTTTAGAGTTTTA	Deoxy, MOE, and cEt	30	10794	10809	4504
561591	1824	1839	GTCAAGTTTAGAGTTT	Deoxy, MOE, and cEt	60	10796	10811	4505
561592	1826	1841	TAGTCAAGTTTAGAGT	Deoxy, MOE, and cEt	56	10798	10813	4506
561593	1828	1843	TTTAGTCAAGTTTAGA	Deoxy, MOE, and cEt	41	10800	10815	4507
561594	1830	1845	TATTTAGTCAAGTTTA	Deoxy, MOE, and cEt	14	10802	10817	4508
561595	1832	1847	TGTATTTAGTCAAGTT	Deoxy, MOE, and cEt	39	10804	10819	4509
561596	1834	1849	TCTGTATTTAGTCAAG	Deoxy, MOE, and cEt	51	10806	10821	4510
561597	1836	1851	CCTCTGTATTTAGTCA	Deoxy, MOE, and cEt	72	10808	10823	4511
561598	1838	1853	GTCCTCTGTATTTAGT	Deoxy, MOE, and cEt	55	10810	10825	4512
561599	1840	1855	CAGTCCTCTGTATTTA	Deoxy, MOE, and cEt	63	10812	10827	4513
561600	1842	1857	ACCAGTCCTCTGTATT	Deoxy, MOE, and cEt	66	10814	10829	4514
561601	1844	1859	TTACCAGTCCTCTGTA	Deoxy, MOE, and cEt	57	10816	10831	4515
561602	1846	1861	AATTACCAGTCCTCTG	Deoxy, MOE, and cEt	43	10818	10833	4516
561603	1848	1863	ACAATTACCAGTCCTC	Deoxy, MOE, and cEt	67	10820	10835	4517

Table 30
Inhibition of ANGPTL3 mRNA by oligonucleotides targeting SEQ ID NO: 1 and 2

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	Chemistry	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
561835	N/A	N/A	GCAAATTTTCAGTGTT	Deoxy, MOE, and cEt	49	3850	3865	4518
561836	N/A	N/A	CGATTTGTAATTTTCA	Deoxy, MOE, and cEt	20	3874	3889	4519
561837	N/A	N/A	TTTAACCGATTTGTAA	Deoxy, MOE, and cEt	42	3880	3895	4520
561838	N/A	N/A	GTATAATTTAACCGAT	Deoxy, MOE, and cEt	15	3886	3901	4521
561839	N/A	N/A	CTAGATTGTATAATTT	Deoxy, MOE, and cEt	15	3893	3908	4522
561840	N/A	N/A	AGTGTTCTAGATTGTA	Deoxy, MOE, and cEt	45	3899	3914	4523
561841	N/A	N/A	TGACATAGTGTTCTAG	Deoxy, MOE, and cEt	51	3905	3920	4524
561842	N/A	N/A	GTGTAATGACATAGTG	Deoxy, MOE, and cEt	58	3911	3926	4525
561843	N/A	N/A	ACAATAGTGTAATGAC	Deoxy, MOE, and cEt	12	3917	3932	4526
561844	N/A	N/A	GTAATTTACAATAGTG	Deoxy, MOE, and cEt	18	3924	3939	4527

561845	N/A	N/A	CCTTCAGTAATTTACA	Deoxy, MOE, and cEt	0	3930	3945	4528
561846	N/A	N/A	TACTTACCTTCAGTAA	Deoxy, MOE, and cEt	2	3936	3951	4529
561847	N/A	N/A	CTGGAGAATAGTTTTA	Deoxy, MOE, and cEt	19	3969	3984	4530
561848	N/A	N/A	TTAAACACTGGAGAAT	Deoxy, MOE, and cEt	14	3976	3991	4531
561849	N/A	N/A	GCCCAGCATATTTTCA	Deoxy, MOE, and cEt	22	4034	4049	4532
561850	N/A	N/A	GAAAAAGCCCAGCATA	Deoxy, MOE, and cEt	15	4040	4055	4533
561851	N/A	N/A	GATTTTCTGAACTTCA	Deoxy, MOE, and cEt	52	4063	4078	4534
561852	N/A	N/A	GTACTATCTCTAAAAT	Deoxy, MOE, and cEt	6	4081	4096	4535
561853	N/A	N/A	TAAATTGTACTATCTC	Deoxy, MOE, and cEt	13	4087	4102	4536
561854	N/A	N/A	CACATATTTTTGTCCT	Deoxy, MOE, and cEt	47	4115	4130	4537
561855	N/A	N/A	CTTTCAAATAGCACAT	Deoxy, MOE, and cEt	31	4126	4141	4538
561856	N/A	N/A	GTATGCTTCTTTCAAA	Deoxy, MOE, and cEt	22	4134	4149	4539
561857	N/A	N/A	CCCCTTGTATGCTTCT	Deoxy, MOE, and cEt	55	4140	4155	4540
561858	N/A	N/A	TTCCTTCCCCTTGTAT	Deoxy, MOE, and cEt	32	4146	4161	4541
561859	N/A	N/A	TGGCAATTCCTTCCCC	Deoxy, MOE, and cEt	43	4152	4167	4542
561860	N/A	N/A	GAATATTGGCAATTCC	Deoxy, MOE, and cEt	52	4158	4173	4543
561861	N/A	N/A	CTAATAATGGATTTGA	Deoxy, MOE, and cEt	0	4179	4194	4544
561862	N/A	N/A	CTATCATAATCTAAAT	Deoxy, MOE, and cEt	0	4202	4217	4545
561863	N/A	N/A	GTAACACTATCATAAT	Deoxy, MOE, and cEt	7	4208	4223	4546
561864	N/A	N/A	AATTCCTGTAACACT	Deoxy, MOE, and cEt	17	4216	4231	4547
561865	N/A	N/A	AAGTTGCTTTCCTCTT	Deoxy, MOE, and cEt	12	4243	4258	4548
561866	N/A	N/A	GGTTATAAGTTGCTTT	Deoxy, MOE, and cEt	6	4249	4264	4549
561867	N/A	N/A	TAGGTTGGTTATAAGT	Deoxy, MOE, and cEt	10	4255	4270	4550
561868	N/A	N/A	AGAGAGTAGGTTGGTT	Deoxy, MOE, and cEt	10	4261	4276	4551
561869	N/A	N/A	GGATATAGAGAGTAGG	Deoxy, MOE, and cEt	23	4267	4282	4552
561870	N/A	N/A	AAGTCTGGATATAGAG	Deoxy, MOE, and cEt	13	4273	4288	4553
561871	N/A	N/A	CTACAAAAGTCTGGAT	Deoxy, MOE, and cEt	1	4279	4294	4554
561872	N/A	N/A	CTTACCTGATTTTCTA	Deoxy, MOE, and cEt	0	4385	4400	4555
561873	N/A	N/A	TACTGACTTACCTGAT	Deoxy, MOE, and cEt	2	4391	4406	4556
561874	N/A	N/A	CCATTAAAATACTGAC	Deoxy, MOE, and cEt	1	4400	4415	4557
561875	N/A	N/A	GGACATAACCATTAAAA	Deoxy, MOE, and cEt	11	4407	4422	4558
561876	N/A	N/A	AAGATGGGACATACCA	Deoxy, MOE, and cEt	38	4413	4428	4559
561877	N/A	N/A	GTGTGAAAGATGGGAC	Deoxy, MOE, and cEt	25	4419	4434	4560
561878	N/A	N/A	AGACCTGTGTGAAAGA	Deoxy, MOE, and cEt	33	4425	4440	4561
561879	N/A	N/A	TTTTACAGACCTGTGT	Deoxy, MOE, and cEt	29	4431	4446	4562
561880	N/A	N/A	CAGTGTTTTTACAGAC	Deoxy, MOE, and cEt	40	4437	4452	4563
561881	N/A	N/A	TAGGATTCAGTGTTTT	Deoxy, MOE, and cEt	62	4444	4459	4564
561882	N/A	N/A	GTTAAAGCTTGTAAT	Deoxy, MOE, and cEt	16	4465	4480	4565
561883	N/A	N/A	GATCCAGTTAAAGCTT	Deoxy, MOE, and cEt	39	4471	4486	4566
561884	N/A	N/A	ACTCATGATCCAGTTA	Deoxy, MOE, and cEt	60	4477	4492	4567
561885	N/A	N/A	AATTTTACTCATGATC	Deoxy, MOE, and cEt	36	4483	4498	4568

561886	N/A	N/A	TGTGATAATTTTACTC	Deoxy, MOE, and cEt	30	4489	4504	4569
561887	N/A	N/A	TGCTGATGTGATAATT	Deoxy, MOE, and cEt	41	4495	4510	4570
561888	N/A	N/A	CAGTTATGCTGATGTG	Deoxy, MOE, and cEt	86	4501	4516	185
561889	N/A	N/A	GCAATTTTAAACAGTTA	Deoxy, MOE, and cEt	13	4511	4526	4571
561890	N/A	N/A	GAGCCTGCAATTTTAA	Deoxy, MOE, and cEt	14	4517	4532	4572
561891	N/A	N/A	TAGCTTCAGAGCCTGC	Deoxy, MOE, and cEt	61	4525	4540	4573
561892	N/A	N/A	GTTTATTAGCTTCAGA	Deoxy, MOE, and cEt	45	4531	4546	4574
561893	N/A	N/A	CAGGTAGTTTATTAGC	Deoxy, MOE, and cEt	37	4537	4552	4575
561894	N/A	N/A	TAAATGCAGGTAGTTT	Deoxy, MOE, and cEt	11	4543	4558	4576
561895	N/A	N/A	ATGGTTTAAATGCAGG	Deoxy, MOE, and cEt	53	4549	4564	4577
561896	N/A	N/A	TAGAGCCATGGTTTAA	Deoxy, MOE, and cEt	58	4556	4571	4578
561897	N/A	N/A	AAGTTTTAGAGCCATG	Deoxy, MOE, and cEt	81	4562	4577	186
561898	N/A	N/A	TCACACAAAGTTTTAG	Deoxy, MOE, and cEt	17	4569	4584	4579
561899	N/A	N/A	GTGAAGTAATTTATTC	Deoxy, MOE, and cEt	8	4589	4604	4580
561900	N/A	N/A	ACTGAGAGATAAAGGG	Deoxy, MOE, and cEt	34	4605	4620	4581
561901	N/A	N/A	GTATATGTGAGGAAAC	Deoxy, MOE, and cEt	18	4619	4634	4582
561902	N/A	N/A	TTTGTAGTATATGTGA	Deoxy, MOE, and cEt	3	4625	4640	4583
561903	N/A	N/A	ATTATCTTTGTAGTAT	Deoxy, MOE, and cEt	8	4631	4646	4584
561904	N/A	N/A	ATAAGTTCTGTTATTA	Deoxy, MOE, and cEt	18	4643	4658	4585
561905	N/A	N/A	AATCCTATAAGTTCTG	Deoxy, MOE, and cEt	55	4649	4664	4586
561906	N/A	N/A	CTGCTATGAATTAATT	Deoxy, MOE, and cEt	16	4679	4694	4587
561907	N/A	N/A	CATTGGCTGCTATGAA	Deoxy, MOE, and cEt	48	4685	4700	4588
561908	N/A	N/A	AGATGACATTGGCTGC	Deoxy, MOE, and cEt	71	4691	4706	4589
561909	N/A	N/A	TTAGTAAGATGACATT	Deoxy, MOE, and cEt	0	4697	4712	4590
561910	N/A	N/A	GATCTAATTTGAATTT	Deoxy, MOE, and cEt	7	4712	4727	4591
561911	N/A	N/A	TTGAGCAAAGAGAAAC	Deoxy, MOE, and cEt	6	4730	4745	4592
561989	N/A	N/A	GAATGTTGACCTTTAA	Deoxy, MOE, and cEt	49	5356	5371	4593
561990	N/A	N/A	ATTGTTGAATGTTGAC	Deoxy, MOE, and cEt	57	5362	5377	4594
561991	N/A	N/A	TTAATTACATTGTTGA	Deoxy, MOE, and cEt	0	5370	5385	4595
561992	N/A	N/A	TTGTAGATTAATTACA	Deoxy, MOE, and cEt	18	5377	5392	4596
561993	N/A	N/A	TTTACATTGTAGATTA	Deoxy, MOE, and cEt	3	5383	5398	4597
561994	N/A	N/A	CAGATGTTTACATTGT	Deoxy, MOE, and cEt	71	5389	5404	4598
561995	N/A	N/A	CTTCACCAGATGTTTA	Deoxy, MOE, and cEt	19	5395	5410	4599
561996	N/A	N/A	CTGTCACTTCACCAGA	Deoxy, MOE, and cEt	77	5401	5416	187
561997	N/A	N/A	AGTGCTTCCCTCTGTC	Deoxy, MOE, and cEt	66	5412	5427	4600
561998	N/A	N/A	TAAACAAGTGCTTCCC	Deoxy, MOE, and cEt	62	5418	5433	4601
561999	N/A	N/A	TAGCTTTTTTCTAAAC	Deoxy, MOE, and cEt	0	5429	5444	4602
562000	N/A	N/A	CTGACATAGCTTTTTT	Deoxy, MOE, and cEt	66	5435	5450	4603
562001	N/A	N/A	TGGATTCTGACATAGC	Deoxy, MOE, and cEt	85	5441	5456	188
562002	N/A	N/A	AATACATGGATTCTGA	Deoxy, MOE, and cEt	35	5447	5462	4604
562003	N/A	N/A	TATTAGAATACATGGA	Deoxy, MOE, and cEt	7	5453	5468	4605

562004	N/A	N/A	GTACTGCATATTAGAA	Deoxy, MOE, and cEt	48	5461	5476	4606
562005	N/A	N/A	ACTATTGTACTGCATA	Deoxy, MOE, and cEt	53	5467	5482	4607
562006	N/A	N/A	TTTTAAACTATTGTAC	Deoxy, MOE, and cEt	0	5473	5488	4608
562007	N/A	N/A	GAGAGTATTATTAATA	Deoxy, MOE, and cEt	8	5490	5505	4609
562008	N/A	N/A	CTGTTTGAGAGTATTA	Deoxy, MOE, and cEt	0	5496	5511	4610
562009	N/A	N/A	GAATAGCTGTTTGAGA	Deoxy, MOE, and cEt	34	5502	5517	4611
562010	N/A	N/A	AATCCTCTTGAATAGC	Deoxy, MOE, and cEt	62	5511	5526	4612
562011	N/A	N/A	TTTTTGAATCCTCTTG	Deoxy, MOE, and cEt	50	5517	5532	4613
562012	N/A	N/A	GAGTTTATATTATGTT	Deoxy, MOE, and cEt	5	5532	5547	4614
562013	N/A	N/A	GTTTCTCTGAGTTTAT	Deoxy, MOE, and cEt	58	5540	5555	4615
562014	N/A	N/A	TTACCAGTTTCTCTGA	Deoxy, MOE, and cEt	64	5546	5561	4616
562015	N/A	N/A	GATTTTGTTTACCAGT	Deoxy, MOE, and cEt	68	5554	5569	4617
562016	N/A	N/A	GTTTTATATCTCTTGA	Deoxy, MOE, and cEt	33	5574	5589	4618
562017	N/A	N/A	TTGGTAATAATATTTG	Deoxy, MOE, and cEt	13	5589	5604	4619
562018	N/A	N/A	TGGAAATTGGTAATAA	Deoxy, MOE, and cEt	1	5595	5610	4620
562019	N/A	N/A	GTTTAGTGGAAATTGG	Deoxy, MOE, and cEt	44	5601	5616	4621
562020	N/A	N/A	ATGTTTGTTTAGTGGA	Deoxy, MOE, and cEt	47	5607	5622	4622
562021	N/A	N/A	CTAACATTATGTTTGT	Deoxy, MOE, and cEt	0	5615	5630	4623
562022	N/A	N/A	GCACTACTAACATTAT	Deoxy, MOE, and cEt	42	5621	5636	4624
562023	N/A	N/A	TTAGCAGCACTACTAA	Deoxy, MOE, and cEt	35	5627	5642	4625
562024	N/A	N/A	AACCTTTTAGCAGCAC	Deoxy, MOE, and cEt	76	5633	5648	189
562025	N/A	N/A	TTGATAAAAAACCTTT	Deoxy, MOE, and cEt	0	5642	5657	4626
562026	N/A	N/A	CAAAAGTAGTTGATAA	Deoxy, MOE, and cEt	0	5651	5666	4627
562027	N/A	N/A	GGAAACCAAAAGTAGT	Deoxy, MOE, and cEt	28	5657	5672	4628
562028	N/A	N/A	GAAAGTATGGAAACCA	Deoxy, MOE, and cEt	52	5665	5680	4629
562029	N/A	N/A	ACATCATAAGAAGGAA	Deoxy, MOE, and cEt	8	5678	5693	4630
562030	N/A	N/A	TCATAGTAAAAGATAT	Deoxy, MOE, and cEt	0	5718	5733	4631
562031	N/A	N/A	TCATTTAATCATAGTA	Deoxy, MOE, and cEt	7	5726	5741	4632
562032	N/A	N/A	GCAGGTTCATTTAATC	Deoxy, MOE, and cEt	56	5732	5747	4633
562033	N/A	N/A	GTAACATTTTGCTTTG	Deoxy, MOE, and cEt	44	5752	5767	4634
562034	N/A	N/A	ATATTACTATAGTAAC	Deoxy, MOE, and cEt	4	5763	5778	4635
562035	N/A	N/A	CAATGTATATTACTAT	Deoxy, MOE, and cEt	19	5769	5784	4636
562036	N/A	N/A	TAGACACAATGTATAT	Deoxy, MOE, and cEt	17	5775	5790	4637
562037	N/A	N/A	GGTTTCTTCACACATT	Deoxy, MOE, and cEt	63	5799	5814	4638
562038	N/A	N/A	CTCAGAAATTCATTGT	Deoxy, MOE, and cEt	36	5818	5833	4639
562039	N/A	N/A	CTTCTTCCAACTCAGA	Deoxy, MOE, and cEt	56	5828	5843	4640
562040	N/A	N/A	CTAACTCTTCTTCCAA	Deoxy, MOE, and cEt	39	5834	5849	4641
562041	N/A	N/A	AATGATCTAACTCTTC	Deoxy, MOE, and cEt	33	5840	5855	4642
562042	N/A	N/A	GAAAGTTAAATGATCT	Deoxy, MOE, and cEt	3	5848	5863	4643
562043	N/A	N/A	ATCTTAAAGTTACTTA	Deoxy, MOE, and cEt	56	5900	5915	4644
562044	N/A	N/A	TATGTGATCTTAAAGT	Deoxy, MOE, and cEt	5	5906	5921	4645

562045	N/A	N/A	AGTAACTATGTGATCT	Deoxy, MOE, and cEt	60	5912	5927	4646
562046	N/A	N/A	CTACTAAGTAACTATG	Deoxy, MOE, and cEt	0	5918	5933	4647
562047	N/A	N/A	TCTTTTCTACTAAGTA	Deoxy, MOE, and cEt	18	5924	5939	4648
562048	N/A	N/A	TATTACTCTTTTCTAC	Deoxy, MOE, and cEt	3	5930	5945	4649
562049	N/A	N/A	GCTGGGTATTACTCTT	Deoxy, MOE, and cEt	76	5936	5951	4650
562050	N/A	N/A	TTGCTTGCTGGGTATT	Deoxy, MOE, and cEt	77	5942	5957	190
562051	N/A	N/A	TAAAGTTTGCTTGCTG	Deoxy, MOE, and cEt	58	5948	5963	4651
562052	N/A	N/A	CTATTGTAAAGTTTGC	Deoxy, MOE, and cEt	16	5954	5969	4652
562053	N/A	N/A	AAGGATCTATTGTAAA	Deoxy, MOE, and cEt	5	5960	5975	4653
562054	N/A	N/A	CTTATTTAAAAGGATC	Deoxy, MOE, and cEt	0	5969	5984	4654
562055	N/A	N/A	TAGGACCTTATTTAAA	Deoxy, MOE, and cEt	0	5975	5990	4655
562056	N/A	N/A	ATTTCCTAGGACCTTA	Deoxy, MOE, and cEt	10	5981	5996	4656
562057	N/A	N/A	CATGAATGATATTTCC	Deoxy, MOE, and cEt	39	5991	6006	4657
562058	N/A	N/A	TGCTGGCATGAATGAT	Deoxy, MOE, and cEt	62	5997	6012	4658
562059	N/A	N/A	TTTTGATGCTGGCATG	Deoxy, MOE, and cEt	74	6003	6018	4659
562060	N/A	N/A	TTAGTTTTTTTGATGCT	Deoxy, MOE, and cEt	25	6009	6024	4660
562061	N/A	N/A	GCATTATTAGTGTTAG	Deoxy, MOE, and cEt	44	6021	6036	4661
562062	N/A	N/A	TATCTTGCATTATTAG	Deoxy, MOE, and cEt	35	6027	6042	4662
562063	N/A	N/A	ATATAATATCTTGCAT	Deoxy, MOE, and cEt	0	6033	6048	4663
562064	N/A	N/A	CATTGACAGTAAGAAA	Deoxy, MOE, and cEt	0	6057	6072	4664
562065	N/A	N/A	AGTTTTTCTCATTGAC	Deoxy, MOE, and cEt	62	6066	6081	4665
562143	N/A	N/A	ATGGATATCTCTTAAC	Deoxy, MOE, and cEt	18	6869	6884	4666
562144	N/A	N/A	TATTTGATGGATATCT	Deoxy, MOE, and cEt	35	6875	6890	4667
562145	N/A	N/A	ACATTGTATTTGATGG	Deoxy, MOE, and cEt	41	6881	6896	4668
562146	N/A	N/A	GTTGATACATTGTATT	Deoxy, MOE, and cEt	8	6887	6902	4669
562147	N/A	N/A	GTTTAGGTTGATACAT	Deoxy, MOE, and cEt	35	6893	6908	4670
562148	N/A	N/A	CATCCAGTTTAGGTTG	Deoxy, MOE, and cEt	59	6899	6914	4671
562149	N/A	N/A	CCCCAGCATCCAGTTT	Deoxy, MOE, and cEt	37	6905	6920	4672
562150	N/A	N/A	AAAGAACCCCAGCATC	Deoxy, MOE, and cEt	35	6911	6926	4673
562151	N/A	N/A	GTGTAAAAAGAACCCC	Deoxy, MOE, and cEt	33	6917	6932	4674
562152	N/A	N/A	TATAGGGTGTA AAAAG	Deoxy, MOE, and cEt	0	6923	6938	4675
562153	N/A	N/A	GTCTTTTATAGGGTGT	Deoxy, MOE, and cEt	75	6929	6944	191
562154	N/A	N/A	AGGTATGTCTTTTATA	Deoxy, MOE, and cEt	21	6935	6950	4676
562155	N/A	N/A	TTGTCTTAGGTATGTC	Deoxy, MOE, and cEt	84	6942	6957	192
562156	N/A	N/A	CTCTGATTGTCTTAGG	Deoxy, MOE, and cEt	77	6948	6963	193
562157	N/A	N/A	GTATTTCTCTGATTGT	Deoxy, MOE, and cEt	77	6954	6969	194
562158	N/A	N/A	AGTCCATATTTGTATT	Deoxy, MOE, and cEt	49	6965	6980	4677
562159	N/A	N/A	TAATCAAGTCCATATT	Deoxy, MOE, and cEt	19	6971	6986	4678
562160	N/A	N/A	ATCTAATAATCAAGTC	Deoxy, MOE, and cEt	5	6977	6992	4679
562161	N/A	N/A	CCTTCTATATTATCTA	Deoxy, MOE, and cEt	38	6988	7003	4680
562162	N/A	N/A	TAATAAACCTTCTATA	Deoxy, MOE, and cEt	8	6995	7010	4681

562163	N/A	N/A	GATCACATCTAAGAAA	Deoxy, MOE, and cEt	25	7013	7028	4682
562164	N/A	N/A	TACCATGATCACATCT	Deoxy, MOE, and cEt	66	7019	7034	4683
562165	N/A	N/A	CTGCAATACCATGATC	Deoxy, MOE, and cEt	54	7025	7040	4684
562166	N/A	N/A	GTTCTCCTTTAAAACT	Deoxy, MOE, and cEt	0	7039	7054	4685
562167	N/A	N/A	GAGATTGTTCTCCTTT	Deoxy, MOE, and cEt	7	7045	7060	4686
562168	N/A	N/A	AAACAGGAGATTGTTC	Deoxy, MOE, and cEt	6	7051	7066	4687
562169	N/A	N/A	TCTCTTAAACAGGAGA	Deoxy, MOE, and cEt	1	7057	7072	4688
562170	N/A	N/A	CATGTATCTCTTAAAC	Deoxy, MOE, and cEt	40	7063	7078	4689
562171	N/A	N/A	CGTAAATATTTTCAGCA	Deoxy, MOE, and cEt	30	7077	7092	4690
562172	N/A	N/A	TAACTCCGTAAATATT	Deoxy, MOE, and cEt	0	7083	7098	4691
562173	N/A	N/A	GACCTTTAACTCCGTA	Deoxy, MOE, and cEt	68	7089	7104	4692
562174	N/A	N/A	TCCAGTGACCTTTAAC	Deoxy, MOE, and cEt	6	7095	7110	4693
562175	N/A	N/A	CACCAGTCTGGAGTCC	Deoxy, MOE, and cEt	52	7108	7123	4694
562176	N/A	N/A	TTCTATCACCAGTCTG	Deoxy, MOE, and cEt	67	7114	7129	4695
562177	N/A	N/A	ATCTTACCAAACCTATT	Deoxy, MOE, and cEt	23	7171	7186	4696
562178	N/A	N/A	AGAATCATCTTACCAA	Deoxy, MOE, and cEt	55	7177	7192	4697
562179	N/A	N/A	GAATGTAAGAATCATC	Deoxy, MOE, and cEt	0	7184	7199	4698
562180	N/A	N/A	GTGTTATTTAAGAATG	Deoxy, MOE, and cEt	0	7195	7210	4699
562181	N/A	N/A	TTTTTCTTAGATGGCG	Deoxy, MOE, and cEt	82	7210	7225	195
562182	N/A	N/A	GTTTATGTTAAAGCAT	Deoxy, MOE, and cEt	8	7225	7240	4700
562183	N/A	N/A	AGTAATGTTTATGTTA	Deoxy, MOE, and cEt	4	7231	7246	4701
562184	N/A	N/A	GTAGCATTTTTTTCAGT	Deoxy, MOE, and cEt	58	7244	7259	4702
562185	N/A	N/A	GCAAATGTAGCATTTT	Deoxy, MOE, and cEt	61	7250	7265	4703
562186	N/A	N/A	GTTGTGGCAAATGTAG	Deoxy, MOE, and cEt	32	7256	7271	4704
562187	N/A	N/A	TATGAAGTTGTGGCAA	Deoxy, MOE, and cEt	54	7262	7277	4705
562188	N/A	N/A	GATTTCACTTGACATT	Deoxy, MOE, and cEt	19	7279	7294	4706
562189	N/A	N/A	GCTTGAGATTTCACTT	Deoxy, MOE, and cEt	42	7285	7300	4707
562190	N/A	N/A	TTTGGAGCTTGAGATT	Deoxy, MOE, and cEt	22	7291	7306	4708
562191	N/A	N/A	AATATCTTTGGAGCTT	Deoxy, MOE, and cEt	36	7297	7312	4709
562192	N/A	N/A	AGGAATAATATCTTTG	Deoxy, MOE, and cEt	5	7303	7318	4710
562193	N/A	N/A	ATTAGTAATAGGAAT	Deoxy, MOE, and cEt	5	7313	7328	4711
562194	N/A	N/A	CATCAGATTTAGTAAT	Deoxy, MOE, and cEt	0	7319	7334	4712
562195	N/A	N/A	GTTATTACATCAGATT	Deoxy, MOE, and cEt	23	7326	7341	4713
562196	N/A	N/A	GCCTAGAATCAATAAA	Deoxy, MOE, and cEt	8	7344	7359	4714
562197	N/A	N/A	AGGAATGCCTAGAATC	Deoxy, MOE, and cEt	2	7350	7365	4715
562198	N/A	N/A	TTCAGCAGGAATGCCT	Deoxy, MOE, and cEt	46	7356	7371	4716
562199	N/A	N/A	TTACCTGATATAACAT	Deoxy, MOE, and cEt	41	7460	7475	4717
562200	N/A	N/A	CAGGTTTTACCTGATA	Deoxy, MOE, and cEt	31	7466	7481	4718
562201	N/A	N/A	CTTAGACAGGTTTTAC	Deoxy, MOE, and cEt	41	7472	7487	4719
562202	N/A	N/A	ATTCTCCTTAGACAGG	Deoxy, MOE, and cEt	37	7478	7493	4720
562203	N/A	N/A	CTGTCTATTCTCCTTA	Deoxy, MOE, and cEt	53	7484	7499	4721

562204	N/A	N/A	TAACTACTGTCTATTC	Deoxy, MOE, and cEt	5	7490	7505	4722
562205	N/A	N/A	TTGAACTAACTACTGT	Deoxy, MOE, and cEt	3	7496	7511	4723
562206	N/A	N/A	AGTAAGTTGAACTAAC	Deoxy, MOE, and cEt	11	7502	7517	4724
562207	N/A	N/A	GTAATGAGTAAGTTGA	Deoxy, MOE, and cEt	37	7508	7523	4725
562208	N/A	N/A	TAATCTTCCTAATACG	Deoxy, MOE, and cEt	5	7523	7538	4726
562209	N/A	N/A	ACCAGGTTAATCTTCC	Deoxy, MOE, and cEt	71	7530	7545	4727
562210	N/A	N/A	ATGATAACCAGGTAA	Deoxy, MOE, and cEt	42	7536	7551	4728
562211	N/A	N/A	CGAATACTCATATATA	Deoxy, MOE, and cEt	20	7576	7591	4729
562212	N/A	N/A	TTTATACGAATACTCA	Deoxy, MOE, and cEt	17	7582	7597	4730
562213	N/A	N/A	ATTATATTTATACGAA	Deoxy, MOE, and cEt	0	7588	7603	4731
562214	N/A	N/A	GGTAAAAGTATTATAT	Deoxy, MOE, and cEt	0	7597	7612	4732
562215	N/A	N/A	GAGAATATTGAGTAAA	Deoxy, MOE, and cEt	9	7624	7639	4733
562216	N/A	N/A	CAGATTATTTTAGAGG	Deoxy, MOE, and cEt	16	7645	7660	4734
562217	N/A	N/A	TCACTTCAGATTATTT	Deoxy, MOE, and cEt	34	7651	7666	4735
562218	N/A	N/A	TAATAGTCACTTCAGA	Deoxy, MOE, and cEt	33	7657	7672	4736
562219	N/A	N/A	TATTGATAATAGTCAC	Deoxy, MOE, and cEt	1	7663	7678	4737
562297	N/A	N/A	TACTATTTGTAATCAA	Deoxy, MOE, and cEt	0	8493	8508	4738
562298	N/A	N/A	CTTGCTTATTTTACTA	Deoxy, MOE, and cEt	24	8504	8519	4739
562299	N/A	N/A	CATCTGTTATTTTATC	Deoxy, MOE, and cEt	0	8519	8534	4740
562300	N/A	N/A	ATGTGCTTTTTGGATT	Deoxy, MOE, and cEt	20	8540	8555	4741
562301	N/A	N/A	GGATTTTTGTATGTGC	Deoxy, MOE, and cEt	64	8550	8565	4742
562302	N/A	N/A	CATCATTGATGGATTT	Deoxy, MOE, and cEt	55	8560	8575	4743
562303	N/A	N/A	CTTAGACATCATTCAT	Deoxy, MOE, and cEt	32	8566	8581	4744
562304	N/A	N/A	TGAGTACTTAGACATC	Deoxy, MOE, and cEt	58	8572	8587	4745
562305	N/A	N/A	TATAAGTGAGTACTTA	Deoxy, MOE, and cEt	3	8578	8593	4746
562306	N/A	N/A	CTACTTTATAAGTGAG	Deoxy, MOE, and cEt	0	8584	8599	4747
562307	N/A	N/A	TGAATGTCTTCTACTT	Deoxy, MOE, and cEt	42	8594	8609	4748
562308	N/A	N/A	TATAATAATGAATGTC	Deoxy, MOE, and cEt	2	8602	8617	4749
562309	N/A	N/A	GTACTGAGCATTTAAA	Deoxy, MOE, and cEt	24	8625	8640	4750
562310	N/A	N/A	CAAATAGTACTGAGCA	Deoxy, MOE, and cEt	48	8631	8646	4751
562311	N/A	N/A	AATGGTCAAATAGTAC	Deoxy, MOE, and cEt	0	8637	8652	4752
562312	N/A	N/A	GTAGTTTGAATACAAA	Deoxy, MOE, and cEt	9	8660	8675	4753
562313	N/A	N/A	TCACTGGTAGTTTGAA	Deoxy, MOE, and cEt	56	8666	8681	4754
562314	N/A	N/A	GGGCTTTCAGTGGTAG	Deoxy, MOE, and cEt	70	8672	8687	196
562315	N/A	N/A	TAGGTAGGGCTTTCAC	Deoxy, MOE, and cEt	50	8678	8693	4755
562316	N/A	N/A	ACCTTCTAGGTAGGGC	Deoxy, MOE, and cEt	47	8684	8699	4756
562317	N/A	N/A	GAGTATACCTTCTAGG	Deoxy, MOE, and cEt	38	8690	8705	4757
562318	N/A	N/A	ATCACTGAGTATACCT	Deoxy, MOE, and cEt	61	8696	8711	4758
562319	N/A	N/A	AAACTTATCACTGAGT	Deoxy, MOE, and cEt	0	8702	8717	4759
562320	N/A	N/A	GCTACAAAACCTTATCA	Deoxy, MOE, and cEt	8	8708	8723	4760
562321	N/A	N/A	TTTGAGCTACAAAAC	Deoxy, MOE, and cEt	0	8714	8729	4761

562322	N/A	N/A	AGAAGATTTGGAGCTA	Deoxy, MOE, and cEt	24	8720	8735	4762
562323	N/A	N/A	ACTATTAGAAGATTTG	Deoxy, MOE, and cEt	0	8726	8741	4763
562324	N/A	N/A	ACACTCACTATTAGAA	Deoxy, MOE, and cEt	0	8732	8747	4764
562325	N/A	N/A	AGCCTTTTATTTTGGG	Deoxy, MOE, and cEt	37	8751	8766	4765
562326	N/A	N/A	CCTGTCAGCCTTTTAT	Deoxy, MOE, and cEt	0	8757	8772	4766
562327	N/A	N/A	GACTTACCTGTCAGCC	Deoxy, MOE, and cEt	47	8763	8778	4767
562328	N/A	N/A	ATTCTCGACTTACCTG	Deoxy, MOE, and cEt	12	8769	8784	4768
562329	N/A	N/A	GTGAGTATTCTCGACT	Deoxy, MOE, and cEt	25	8775	8790	4769
562330	N/A	N/A	AATTAAGTGAGTATTC	Deoxy, MOE, and cEt	0	8781	8796	4770
562331	N/A	N/A	TACCAGAATTAAGTGA	Deoxy, MOE, and cEt	0	8787	8802	4771
562332	N/A	N/A	GCTTTCTTACCAGAAT	Deoxy, MOE, and cEt	23	8794	8809	4772
562333	N/A	N/A	TGGGTTGCTTTCTTAC	Deoxy, MOE, and cEt	0	8800	8815	4773
562334	N/A	N/A	TACAAGTACAAATGGG	Deoxy, MOE, and cEt	36	8812	8827	4774
562335	N/A	N/A	GGTAAATACAAGTACA	Deoxy, MOE, and cEt	19	8818	8833	4775
562336	N/A	N/A	ATTGCTGGTAAATACA	Deoxy, MOE, and cEt	13	8824	8839	4776
562337	N/A	N/A	TTAAGGATTGCTGGTA	Deoxy, MOE, and cEt	43	8830	8845	4777
562338	N/A	N/A	GCTTCATTTTAAGGAT	Deoxy, MOE, and cEt	12	8838	8853	4778
562339	N/A	N/A	GTAGGAAGCTTCATTT	Deoxy, MOE, and cEt	23	8845	8860	4779
562340	N/A	N/A	GAGTTAGTAGGAAGCT	Deoxy, MOE, and cEt	58	8851	8866	4780
562341	N/A	N/A	GCTATTGAGTTAGTAG	Deoxy, MOE, and cEt	21	8857	8872	4781
562342	N/A	N/A	CTTATTGCTATTGAGT	Deoxy, MOE, and cEt	34	8863	8878	4782
562343	N/A	N/A	TATTGTCTTATTGCTA	Deoxy, MOE, and cEt	17	8869	8884	4783
562344	N/A	N/A	ATTCACTATTGTCTTA	Deoxy, MOE, and cEt	22	8875	8890	4784
562345	N/A	N/A	ATCACAATCCTTTTTA	Deoxy, MOE, and cEt	18	8925	8940	4785
562346	N/A	N/A	TTCTTCATCACAATCC	Deoxy, MOE, and cEt	43	8931	8946	4786
562347	N/A	N/A	AGATTGTTCTTCATCA	Deoxy, MOE, and cEt	35	8937	8952	4787
562348	N/A	N/A	TATAAATAGATTGTTC	Deoxy, MOE, and cEt	10	8944	8959	4788
562349	N/A	N/A	GGTTCTTAATAACTTT	Deoxy, MOE, and cEt	31	9011	9026	4789
562350	N/A	N/A	AAGCATGGTTCTTAAT	Deoxy, MOE, and cEt	12	9017	9032	4790
562351	N/A	N/A	CTTTGTAGAAAAAGAC	Deoxy, MOE, and cEt	0	9066	9081	4791
562352	N/A	N/A	TATGCTTTCTTTGTAG	Deoxy, MOE, and cEt	26	9074	9089	4792
562353	N/A	N/A	CTTAATGTATGCTTTC	Deoxy, MOE, and cEt	55	9081	9096	4793
562354	N/A	N/A	GTATTTGCTTAATGTA	Deoxy, MOE, and cEt	0	9088	9103	4794
562355	N/A	N/A	CCTTTGGTATTTGCTT	Deoxy, MOE, and cEt	54	9094	9109	4795
562356	N/A	N/A	ACCTGGCCTTTGGTAT	Deoxy, MOE, and cEt	0	9100	9115	4796
562357	N/A	N/A	ATGTAAACCTGGCCTT	Deoxy, MOE, and cEt	1	9106	9121	4797
562358	N/A	N/A	CTTCAAATGTAAACCT	Deoxy, MOE, and cEt	0	9112	9127	4798
562359	N/A	N/A	GTAATAATAATGTCAC	Deoxy, MOE, and cEt	0	9131	9146	4799
562360	N/A	N/A	AGACTTGAGTAATAAT	Deoxy, MOE, and cEt	0	9139	9154	4800
562361	N/A	N/A	TCCTAGAGACTTGAGT	Deoxy, MOE, and cEt	25	9145	9160	4801
562362	N/A	N/A	AAGTATTCCTAGAGAC	Deoxy, MOE, and cEt	28	9151	9166	4802

562363	N/A	N/A	TGTGTTAAGTATTCCT	Deoxy, MOE, and cEt	50	9157	9172	4803
562364	N/A	N/A	AAGAGATGTGTTAAGT	Deoxy, MOE, and cEt	21	9163	9178	4804
562365	N/A	N/A	ACAGTCAAGAGATGTG	Deoxy, MOE, and cEt	74	9169	9184	197
562366	N/A	N/A	CCATATACAGTCAAGA	Deoxy, MOE, and cEt	49	9175	9190	4805
562367	N/A	N/A	TAACATCCATATACAG	Deoxy, MOE, and cEt	16	9181	9196	4806
562368	N/A	N/A	CTATTTATTAACATCC	Deoxy, MOE, and cEt	2	9189	9204	4807
562369	N/A	N/A	TGTCAGCTATTTATTA	Deoxy, MOE, and cEt	22	9195	9210	4808
562370	N/A	N/A	CTTTACTGTCAGCTAT	Deoxy, MOE, and cEt	56	9201	9216	4809
562371	N/A	N/A	GATAAACTTTACTGTC	Deoxy, MOE, and cEt	37	9207	9222	4810
562372	N/A	N/A	CTTTATATGGATAAAC	Deoxy, MOE, and cEt	31	9216	9231	4811
562373	N/A	N/A	GCAAGTCTTTATATGG	Deoxy, MOE, and cEt	62	9222	9237	4812
560990	709	724	TTCTTGGTGCTCTTGG	Deoxy, MOE, and cEt	74	6722	6737	111
337487	804	823	CACTTGATGTTACCTCTG	5-10-5 MOE	30	7389	7408	28
233717	889	908	TGAATTAATGTCCATGGACT	5-10-5 MOE	38	7876	7895	14

Example 6: Antisense inhibition of human ANGPTL3 in Hep3B cells by MOE gapmers

Additional antisense oligonucleotides were designed targeting an ANGPTL3 nucleic acid and were tested for their effects on ANGPTL3 mRNA in vitro. Cultured Hep3B cells at a density of 20,000 cells per well were transfected using electroporation with 4,500 nM antisense oligonucleotide. After a treatment period of approximately 24 hours, RNA was isolated from the cells and ANGPTL3mRNA levels were measured by quantitative real-time PCR. Human primer probe set RTS3492_MGB was used to measure mRNA levels. ANGPTL3 mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN®. Results are presented as percent inhibition of ANGPTL3, relative to untreated control cells.

The newly designed chimeric antisense oligonucleotides in the Tables below were designed as 5-10-5 MOE or 3-10-4 MOE gapmers. The 5-10-5 MOE 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. The 3-10-4 MOE 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 and four nucleosides respectively. Each nucleoside in the 5' wing segment and each nucleoside in the 3' wing segment has a 2'-MOE modification. 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. "Start site" indicates the 5'-most nucleoside to which the gapmer is targeted in the human gene sequence. "Stop site" indicates the 3'-most nucleoside to which the gapmer is targeted human gene sequence. Each gapmer listed in the Tables below is targeted to either the human ANGPTL3 mRNA, designated herein as SEQ ID NO: 1 (GENBANK Accession No. NM_014495.2) or the human ANGPTL3 genomic sequence, designated herein as SEQ ID NO: 2 (GENBANK Accession No. NT_032977.9 truncated from nucleotides 33032001 to 33046000). 'n/a'

indicates that the antisense oligonucleotide does not target that particular gene sequence with 100% complementarity.

Table 31
Inhibition of ANGPTL3 mRNA by MOE gapmers targeting SEQ ID NO: 1 and 2

ISIS NO	SEQ ID NO: 1 Start Site	SEQ ID NO: 1 Stop Site	Sequence	Motif	% inhibition	SEQ ID NO: 2 Start Site	SEQ ID NO: 2 Stop Site	SEQ ID NO
582715	N/A	N/A	CTGGGTATTACTCTTTTCTA	5-10-5	60	5931	5950	4813
582716	N/A	N/A	CTTGCTGGGTATTACTCTTT	5-10-5	59	5935	5954	4814
582717	N/A	N/A	TGCTTGCTGGGTATTACTCT	5-10-5	59	5937	5956	4815
582718	N/A	N/A	CATGAATGATATTCCTAGG	5-10-5	39	5987	6006	4816
582719	N/A	N/A	GGCATGAATGATATTCCTA	5-10-5	60	5989	6008	4817
582720	N/A	N/A	CTGGCATGAATGATATTTCC	5-10-5	46	5991	6010	4818
582721	N/A	N/A	TGCTGGCATGAATGATATTT	5-10-5	32	5993	6012	4819
582722	N/A	N/A	AAGTCCATATTTGTATTTCT	5-10-5	50	6962	6981	4820
582723	N/A	N/A	GCAAATGTAGCATTTTTTCA	5-10-5	32	7246	7265	4821
582724	N/A	N/A	GGCAAATGTAGCATTTTTTC	5-10-5	55	7247	7266	4822
582725	N/A	N/A	GTGGCAAATGTAGCATTTTTT	5-10-5	62	7249	7268	203
582726	N/A	N/A	CTGGTCCTTTTAACTTCCAA	5-10-5	40	8366	8385	4823
582727	N/A	N/A	CCTGGTCCTTTTAACTTCCA	5-10-5	58	8367	8386	4824
582728	N/A	N/A	TTCCTGGTCCTTTTAACTTC	5-10-5	32	8369	8388	4825
582729	N/A	N/A	TGCTTAATGTATGCTTTCTT	5-10-5	51	9079	9098	4826
582730	N/A	N/A	CCGTAAGTTTATCTTCCTTT	5-10-5	58	10136	10155	4827
582731	N/A	N/A	CCCCGTAAGTTTATCTTCCT	5-10-5	51	10138	10157	4828
582732	N/A	N/A	CACAAATATGTTCAATTCTTA	5-10-5	22	11189	11208	4829
582733	N/A	N/A	GCCACAAATATGTTCAATTCT	5-10-5	71	11191	11210	204
582734	N/A	N/A	AAACTTTAACTCGATGCCAC	5-10-5	51	11206	11225	4830
582735	N/A	N/A	ATAAACTTTAACTCGATGCC	5-10-5	57	11208	11227	4831
582736	N/A	N/A	ATGCTTGTCAGGCTGTTTAA	5-10-5	56	11311	11330	4832
582737	N/A	N/A	GTCACCATATAACTTGGGCA	5-10-5	48	11562	11581	4833
582738	N/A	N/A	AGGTCACCATATAACTTGGG	5-10-5	44	11564	11583	4834
582766	N/A	N/A	GCTGGGTATTACTCTTT	3-10-4	55	5935	5951	4835
582767	N/A	N/A	GCATGAATGATATTTCC	3-10-4	4	5991	6007	4836
582768	N/A	N/A	GGCAAATGTAGCATTTT	3-10-4	33	7250	7266	4837
582769	N/A	N/A	CTGGTCCTTTTAACTTC	3-10-4	29	8369	8385	4838
582770	N/A	N/A	GTAAGTTTATCTTCCTT	3-10-4	26	10137	10153	4839
582771	N/A	N/A	ACTTTAACTCGATGCCA	3-10-4	42	11207	11223	4840
582772	N/A	N/A	AACTTTAACTCGATGCC	3-10-4	55	11208	11224	4841
582773	N/A	N/A	AACTTTAACTCGATGC	3-10-4	1	11209	11225	4842
582774	N/A	N/A	GCTTGTCAGGCTGTTTA	3-10-4	65	11312	11328	208
582775	N/A	N/A	CACCATATAACTTGGGC	3-10-4	38	11563	11579	4843

WO 2015/100394

PCT/US2014/072303

582776	N/A	N/A	TCACCATATAACTTGGG	3-10-4	37	11564	11580	4844
582777	N/A	N/A	GTCACCATATAACTTGG	3-10-4	31	11565	11581	4845
582702	139	158	CTTGATTTTGGCTCTGGAGA	5-10-5	53	3243	3262	4846
582739	140	156	TGATTTTGGCTCTGGAG	3-10-4	41	3244	3260	4847
582703	141	160	ATCTTGATTTTGGCTCTGGA	5-10-5	64	3245	3264	198
582740	305	321	ACTGGTTTGCAGCGATA	3-10-4	58	3409	3425	4848
582704	306	325	TTTCACTGGTTTGCAGCGAT	5-10-5	60	3410	3429	4849
582741	306	322	CACTGGTTTGCAGCGAT	3-10-4	57	3410	3426	4850
582742	307	323	TCACTGGTTTGCAGCGA	3-10-4	60	3411	3427	4851
582705	706	725	GTTCTTGGTGCTCTTGGCTT	5-10-5	78	6719	6738	199
544120	707	726	AGTTCTTGGTGCTCTTGGCT	5-10-5	75	6720	6739	15
582743	707	723	TCTTGGTGCTCTTGGCT	3-10-4	63	6720	6736	205
582706	708	727	TAGTTCTTGGTGCTCTTGGC	5-10-5	69	6721	6740	200
582744	708	724	TTCTTGGTGCTCTTGGC	3-10-4	51	6721	6737	4852
582745	709	725	GTTCTTGGTGCTCTTGG	3-10-4	50	6722	6738	4853
337487	804	823	CACTTGTATGTTACCTCTG	5-10-5	25	7389	7408	28
233717	889	908	TGAATTAATGTCCATGGACT	5-10-5	22	7876	7895	14
582707	1054	1073	TTGTCTTTCCAGTCTTCCAA	5-10-5	42	9629	9648	4854
582708	1056	1075	TGTTGTCTTTCCAGTCTTCC	5-10-5	52	9631	9650	4855
582746	1140	1156	CATTGCCAGTAATCGCA	3-10-4	53	9715	9731	4856
582747	1141	1157	ACATTGCCAGTAATCGC	3-10-4	61	9716	9732	4857
582748	1142	1158	GACATTGCCAGTAATCG	3-10-4	34	9717	9733	4858
582709	1194	1213	CTTTGTGATCCCAAGTAGAA	5-10-5	28	9769	9788	4859
582749	1195	1211	TTGTGATCCCAAGTAGA	3-10-4	16	9770	9786	4860
582710	1196	1215	TGCTTTGTGATCCCAAGTAG	5-10-5	54	9771	9790	4861
582750	1196	1212	TTTGTGATCCCAAGTAG	3-10-4	19	9771	9787	4862
582751	1197	1213	CTTTGTGATCCCAAGTA	3-10-4	32	9772	9788	4863
582752	1260	1276	CACACTCATCATGCCAC	3-10-4	42	10232	10248	4864
582711	1268	1287	GTTGTTTTCTCCACACTCAT	5-10-5	51	10240	10259	4865
582712	1270	1289	AGGTTGTTTTCTCCACACTC	5-10-5	63	10242	10261	201
582753	1307	1323	AGATTTTGCTCTTGGTT	3-10-4	54	10279	10295	4866
582754	1308	1324	TAGATTTTGCTCTTGGT	3-10-4	52	10280	10296	4867
582755	1309	1325	TTAGATTTTGCTCTTGG	3-10-4	44	10281	10297	4868
582756	1310	1326	CTTAGATTTTGCTCTTG	3-10-4	34	10282	10298	4869
567320	1487	1506	CCAGATTATTAGACCACATT	5-10-5	77	10459	10478	93
582757	1488	1504	AGATTATTAGACCACAT	3-10-4	0	10460	10476	4870
582758	1489	1505	CAGATTATTAGACCACA	3-10-4	39	10461	10477	4871
582759	1490	1506	CCAGATTATTAGACCAC	3-10-4	63	10462	10478	206
582760	1491	1507	ACCAGATTATTAGACCA	3-10-4	31	10463	10479	4872
582761	1763	1779	GCTCATATGATGCCTTT	3-10-4	71	10735	10751	207
582713	1906	1925	ACACATACTCTGTGCTGACG	5-10-5	68	10878	10897	202

582762	1907	1923	ACATACTCTGTGCTGAC	3-10-4	57	10879	10895	4873
582714	1908	1927	TTACACATACTCTGTGCTGA	5-10-5	49	10880	10899	4874
582763	2071	2087	CTTAGTAGTCATCTCCA	3-10-4	49	11043	11059	4875
582764	2072	2088	ACTTAGTAGTCATCTCC	3-10-4	53	11044	11060	4876
582765	2073	2089	GACTTAGTAGTCATCTC	3-10-4	36	11045	11061	4877

Example 7: Dose-dependent antisense inhibition of human ANGPTL3 in Hep3B cells

Deoxy, MOE, and cEt oligonucleotides from the studies described above exhibiting significant *in vitro* inhibition of ANGPTL3 mRNA were selected and tested at various doses in Hep3B cells. ISIS 233717 and ISIS 337847, both 5-10-5 MOE gapmers, were also included in the studies. The antisense oligonucleotides were tested in a series of experiments that had similar culture conditions. The results of each experiment are presented in separate tables below.

Cells were plated at a density of 20,000 cells per well and transfected using electroporation with 0.813 μ M, 1.625 μ M, 3.25 μ M, 6.500 μ M and 13.00 μ M 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 ANGPTL3 mRNA levels were measured by quantitative real-time PCR. Human primer probe set RTS3492_MGB was used to measure mRNA levels. ANGPTL3 mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN[®]. Results are presented as percent inhibition of ANGPTL3, relative to untreated control cells.

The half maximal inhibitory concentration (IC₅₀) of each oligonucleotide is also presented. ANGPTL3 mRNA levels were significantly reduced in a dose-dependent manner in antisense oligonucleotide treated cells.

Table 32

ISIS No	0.813 μ M	1.625 μ M	3.25 μ M	6.50 μ M	13.00 μ M	IC ₅₀ (μ M)	SEQ ID NO
233717	0	27	43	66	79	4.4	14
337487	26	49	63	85	94	2.0	28
559277	54	68	70	82	91	<0.8	110
560990	36	61	74	90	96	1.2	111
560992	60	67	76	81	93	<0.8	112
561010	71	77	82	86	94	<0.8	113
561011	80	87	91	95	97	<0.8	114
561022	75	79	84	89	93	<0.8	115
561025	68	82	81	91	96	<0.8	116
561026	72	85	85	89	90	<0.8	117
561208	63	80	87	92	93	<0.8	118
561320	47	60	86	92	96	0.8	119

561343	45	59	79	86	93	0.9	120
561345	38	59	80	88	95	1.1	121
561347	53	63	84	88	97	<0.8	122

Table 33

ISIS No	0.813 μM	1.625 μM	3.25 μM	6.50 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
233717	7	19	55	60	77	4.2	14
337487	33	44	69	83	88	2.0	28
560990	36	64	81	87	95	1.1	111
561452	58	69	75	85	88	<0.8	123
561458	69	77	84	91	94	<0.8	124
561460	54	50	72	79	85	<0.8	125
561462	49	72	80	90	92	<0.8	126
561463	63	79	84	92	93	<0.8	127
561478	56	53	80	86	91	<0.8	128
561482	46	69	80	86	91	<0.8	129
561486	56	73	80	91	92	<0.8	130
561487	82	87	88	90	93	<0.8	131
561500	52	60	71	80	91	<0.8	132
561504	49	72	85	91	93	<0.8	133
561621	68	76	85	91	94	<0.8	134

Table 34

ISIS No	0.813 μM	1.625 μM	3.25 μM	6.50 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
233717	28	35	48	56	60	4.7	14
337487	43	58	72	82	89	1.0	28
560990	57	73	82	86	96	<0.8	111
561620	51	74	80	85	88	<0.8	135
561622	63	73	85	88	87	<0.8	136
561628	48	69	77	79	80	<0.8	137
561631	60	75	84	86	90	<0.8	138
561644	59	69	77	85	83	<0.8	139
561646	67	81	84	91	92	<0.8	140
561649	70	76	85	89	89	<0.8	141
561650	78	85	88	90	91	<0.8	142
561770	66	81	79	88	91	<0.8	143
561781	65	67	80	81	91	<0.8	144
561791	68	73	83	82	85	<0.8	145

561918	63	71	81	86	92	<0.8	146
--------	----	----	----	----	----	------	-----

Table 35

ISIS No	0.813 μM	1.625 μM	3.25 μM	6.50 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
233717	21	26	47	62	69	4.2	14
337487	35	54	73	82	92	1.0	28
560990	42	76	81	88	96	<0.8	111
562078	55	85	86	91	93	<0.8	147
562086	64	83	87	92	93	<0.8	148
562103	72	83	90	90	94	<0.8	149
562110	66	80	83	89	92	<0.8	150
562375	56	61	63	84	90	<0.8	151
562387	67	75	81	90	88	<0.8	152
562396	60	71	80	80	85	<0.8	153
562415	66	73	77	77	81	<0.8	154
562433	68	84	86	90	91	<0.8	155
562436	78	87	87	91	94	<0.8	156
562439	55	66	78	82	93	<0.8	157
562442	55	57	60	76	86	<0.8	158

Example 8: Dose-dependent antisense inhibition of human ANGPTL3 in Hep3B cells

- 5 Deoxy, MOE, and cEt oligonucleotides from the studies described above exhibiting significant *in vitro* inhibition of ANGPTL3 mRNA were selected and tested at various doses in Hep3B cells. ISIS 337847, a 5-10-5 MOE gapmer, was also included in the studies. The antisense oligonucleotides were tested in a series of experiments that had similar culture conditions. The results of each experiment are presented in separate tables below.
- 10 Cells were plated at a density of 20,000 cells per well and transfected using electroporation with 0.160 μM, 0.481 μM, 1.444 μM, 4.333 μM and 13.00 μM 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 ANGPTL3 mRNA levels were measured by quantitative real-time PCR. Human primer probe set RTS3492_MGB was used to measure mRNA levels. ANGPTL3 mRNA levels were adjusted according to
- 15 total RNA content, as measured by RIBOGREEN[®]. Results are presented as percent inhibition of ANGPTL3, relative to untreated control cells.

The half maximal inhibitory concentration (IC₅₀) of each oligonucleotide is also presented. ANGPTL3 mRNA levels were significantly reduced in a dose-dependent manner in antisense oligonucleotide treated cells.

Table 36

ISIS No	0.160 μM	0.481 μM	1.444 μM	4.333 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
337487	0	18	24	49	73	4.1	28
560990	2	27	39	59	80	2.0	111
561076	20	33	59	73	89	1.1	159
561079	24	39	51	72	84	1.0	160
561084	7	17	46	66	87	1.9	161
561085	21	35	55	69	86	1.2	162
561123	20	39	52	72	87	1.1	163
561241	13	22	41	68	86	2.0	164
561256	12	22	35	54	82	2.6	165
561260	22	16	34	54	82	2.6	166
561277	21	21	37	59	69	2.9	167
561288	6	8	23	36	68	6.9	168
561418	25	36	61	79	86	0.9	169
561436	21	40	61	77	88	0.9	170
561443	18	32	52	82	88	1.1	171

Table 37

ISIS No	0.160 μM	0.481 μM	1.444 μM	4.333 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
337487	0	8	21	52	81	3.7	28
560990	6	14	40	61	74	3.0	111
561398	3	9	22	64	79	3.0	172
561400	11	28	50	65	83	1.7	173
561528	2	39	59	74	84	1.3	174
561565	18	43	58	75	83	1.0	175
561566	21	29	54	71	79	1.4	176
561567	16	35	56	67	78	1.4	177
561571	18	32	60	80	86	1.1	178
561576	11	12	42	65	77	2.4	179
561689	16	27	52	76	80	1.4	180
561698	1	24	31	61	74	2.9	181
561699	2	19	48	65	81	2.0	182
561722	14	34	59	72	85	1.2	183
561723	7	31	69	71	75	1.4	184

Table 38

ISIS No	0.160 μM	0.481 μM	1.444 μM	4.333 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
337487	14	9	9	47	72	5.9	28
560990	13	26	39	58	81	2.0	111
561888	16	19	46	72	84	1.7	185
561897	6	31	50	67	82	2.0	186
561996	19	31	49	59	83	1.6	187
562001	22	46	57	67	89	0.9	188
562024	17	29	59	71	83	1.3	189
562050	21	38	46	62	74	1.6	190
562153	22	35	42	61	71	2.0	191
562155	29	29	50	72	84	1.2	192
562156	15	17	39	60	82	2.3	193
562157	14	15	43	54	75	3.0	194
562181	24	34	58	73	80	1.1	195
562314	22	30	42	54	64	3.1	196
562365	25	27	46	64	77	1.7	197

Example 9: Dose-dependent antisense inhibition of human ANGPTL3 in Hep3B cells by MOE gapmers

5 MOE gapmers from the Examples above exhibiting significant *in vitro* inhibition of ANGPTL3 mRNA were selected and tested at various doses in Hep3B cells. Cells were plated at a density of 20,000 cells per well and transfected using electroporation with 0.160 μM, 0.481 μM, 1.444 μM, 4.333 μM and 13.00 μM 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 ANGPTL3 mRNA levels were

10 measured by quantitative real-time PCR. Human primer probe set RTS3492_MGB was used to measure mRNA levels. ANGPTL3 mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN[®]. Results are presented as percent inhibition of ANGPTL3, relative to untreated control cells.

The half maximal inhibitory concentration (IC₅₀) of each oligonucleotide is also presented. ANGPTL3 mRNA levels were significantly reduced in a dose-dependent manner in antisense oligonucleotide

15 treated cells.

Table 39

ISIS No	Motif	0.16 μM	0.48 μM	1.44 μM	4.33 μM	13.00 μM	IC ₅₀ (μM)	SEQ ID NO
233717	5-10-5	0	3	12	38	64	8.0	14
337487	5-10-5	0	0	15	30	66	8.0	28
544120	5-10-5	10	37	62	81	94	1.0	15

567320	5-10-5	0	30	67	84	95	1.1	93
582703	5-10-5	0	18	47	71	83	2.0	198
582705	5-10-5	22	18	46	82	93	1.0	199
582706	5-10-5	2	0	32	67	85	2.6	200
582712	5-10-5	0	0	54	71	89	2.2	201
582713	5-10-5	25	25	52	75	85	1.2	202
582725	5-10-5	0	3	43	62	84	2.7	203
582733	5-10-5	0	30	66	77	87	1.3	204
582743	3-10-4	0	6	37	51	87	2.9	205
582759	3-10-4	0	2	51	76	93	2.0	206
582761	3-10-4	4	38	58	72	87	1.3	207
582774	3-10-4	5	29	46	72	86	1.6	208

Example 10: Dose-dependent antisense inhibition of human ANGPTL3 in Hep3B cells by deoxy, MOE and cEt oligonucleotides

Deoxy, MOE, and cEt oligonucleotides from the studies described above exhibiting significant *in vitro* inhibition of ANGPTL3 mRNA were selected and tested at various doses in Hep3B cells. Cells were plated at a density of 20,000 cells per well and transfected using electroporation with 0.111 μ M, 0.333 μ M, 1.00 μ M, 3.00 μ M and 9.00 μ M 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 ANGPTL3 mRNA levels were measured by quantitative real-time PCR. Human primer probe set RTS3492_MGB was used to measure mRNA levels. ANGPTL3 mRNA levels were adjusted according to total RNA content, as measured by RIBOGREEN[®]. Results are presented as percent inhibition of ANGPTL3, relative to untreated control cells.

The half maximal inhibitory concentration (IC₅₀) of each oligonucleotide is also presented. ANGPTL3 mRNA levels were significantly reduced in a dose-dependent manner in antisense oligonucleotide treated cells.

Table 40

ISIS No	0.111 μ M	0.333 μ M	1.00 μ M	3.00 μ M	9.00 μ M	IC ₅₀ (μ M)	SEQ ID NO
561011	20	39	65	81	94	0.5	114
561026	23	43	65	84	94	0.5	117
561463	26	25	59	76	91	0.7	127
561487	42	61	81	89	95	0.1	131
586661	24	36	46	76	92	0.7	209
586669	31	50	68	85	95	0.3	210
586676	26	50	73	83	95	0.3	211
586688	4	24	51	82	91	0.9	212

586690	19	39	64	84	95	0.5	213
586691	6	37	60	81	93	0.7	214
586701	10	32	55	76	90	0.8	215
586702	16	25	55	69	86	0.9	216
586705	10	30	54	80	89	0.8	217
586707	33	42	71	83	89	0.3	218
586718	38	54	72	78	85	0.2	219

Table 41

ISIS No	0.111 μM	0.333 μM	1.00 μM	3.00 μM	9.00 μM	IC ₅₀ (μM)	SEQ ID NO
561011	13	29	41	76	89	1.0	114
561567	20	46	57	75	78	0.7	177
586692	32	30	71	85	95	0.4	220
586700	3	46	70	82	95	1.0	221
586708	36	46	62	77	86	0.4	222
586744	0	19	54	81	92	1.0	223
586745	35	22	66	78	92	0.5	224
586746	14	30	59	82	92	0.7	225
586755	18	22	53	74	90	0.9	226
586761	26	26	54	73	90	0.8	227
586787	0	38	64	79	90	0.8	228
586796	12	13	56	83	93	0.9	229
586797	4	26	58	82	90	0.9	230
586802	12	28	56	76	81	0.9	231
586804	17	40	65	86	93	0.5	232

Table 42

ISIS No	0.111 μM	0.333 μM	1.00 μM	3.00 μM	9.00 μM	IC ₅₀ (μM)	SEQ ID NO
561011	20	48	75	84	94	0.4	114
561026	31	48	70	88	95	0.3	117
561463	27	40	67	85	94	0.4	127
561487	41	66	84	91	95	0.1	131
586661	36	45	64	82	91	0.3	209
586669	21	55	73	90	96	0.3	210
586676	23	59	77	87	94	0.3	211
586688	25	41	70	82	93	0.4	212
586690	16	45	74	86	92	0.5	213

586691	13	40	65	86	92	0.6	214
586701	22	49	70	82	93	0.4	215
586702	11	31	58	76	92	0.8	216
586705	26	45	66	82	89	0.4	217
586707	28	58	75	85	88	0.3	218
586718	33	59	73	80	88	0.2	219

Table 43

ISIS No	0.111 μM	0.333 μM	1.00 μM	3.00 μM	9.00 μM	IC ₅₀ (μM)	SEQ ID NO
561011	23	41	63	82	92	0.5	114
561567	31	44	65	75	83	0.4	177
586692	16	58	74	89	93	0.4	220
586700	25	62	75	91	94	0.3	221
586708	36	53	72	81	90	0.3	222
586744	30	29	64	75	94	0.6	223
586745	21	44	59	81	89	0.5	224
586746	19	48	57	85	87	0.5	225
586755	6	30	59	78	89	0.8	226
586761	12	29	59	72	87	0.9	227
586787	27	35	64	84	97	0.5	228
586796	31	40	72	91	95	0.3	229
586797	36	47	67	82	88	0.3	230
586802	35	32	61	76	90	0.5	231
586804	35	50	75	91	91	0.2	232

Example 11: Antisense Inhibition of human ANGPTL3 in huANGPTL3 transgenic mice

5 Antisense oligonucleotides described in the studies above were further evaluated for their ability to reduce human ANGPTL3 mRNA transcript in C57Bl/6 mice with the human ANGPTL3 transgene (Tg mice).

Study 1

10 Female Tg mice were maintained on a 12-hour light/dark cycle. Animals were acclimated for at least 7 days in the research facility before initiation of the experiment. Antisense oligonucleotides (ASOs) were prepared in buffered saline (PBS) and sterilized by filtering through a 0.2 micron filter. Oligonucleotides were dissolved in 0.9% PBS for injection.

Groups of mice received intraperitoneal injections of 5-10-5 MOE gapmers at a dose of 50 mg/kg once per week for 2 weeks. One group of mice received subcutaneous injections of PBS once weekly for 2 weeks. The PBS-injected group served as the control group to which the corresponding oligonucleotide-treated groups were compared.

5 *RNA analysis*

At the end of the treatment period, RNA was extracted from liver for real-time PCR analysis of measurement of mRNA expression of ANGPTL3 with RTS3492_MGB. mRNA levels were also measured with human primer probe set RTS1984 (forward sequence CTTCAATGAAACGTGGGAGAACT, designated herein as SEQ ID NO: 7; reverse sequence TCTCTAGGCCCAACCAAAATTC, designated
10 herein as SEQ ID NO: 8; probe sequence AAATATGGTTTTGGGAGGCTTGAT, designated herein as SEQ ID NO: 9). Results are presented as percent change of mRNA, relative to PBS control, normalized with RIBOGREEN[®]. As shown in the Table below, treatment with ISIS antisense oligonucleotides resulted in significant reduction of ANGPTL3 mRNA in comparison to the PBS control.

Table 44

15 Percent inhibition of ANGPTL3 mRNA in transgenic mouse liver relative to the PBS control

ISIS No	RTS3492_MGB	RTS1984	SEQ ID NO
233710	91	94	233
233717	49	58	14
337477	76	82	234
337478	52	65	235
337479	53	76	236
337487	80	92	28

Protein analysis

Human ANGPTL3 protein levels were quantified using a commercially available ELISA kit (Catalog #DANL30 by R&D Systems, Minneapolis, MN) with transgenic plasma samples diluted 1:20,000 using the
20 manufacturer described protocol. The results are presented in the Table below. The results indicate that treatment with ISIS oligonucleotides resulted in reduced ANGPTL3 protein levels.

Table 45

Percent inhibition of plasma protein levels in the transgenic mouse

ISIS No	%	SEQ ID NO
233710	92	233
233717	47	14
337477	68	234
337478	36	235

337479	48	236
337487	78	28

Plasma chemistry markers

To evaluate the effect of ISIS oligonucleotides on day 10, plasma levels of transaminases (ALT and AST) 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 these liver function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 46

Plasma transaminase levels (IU/L) in transgenic mice on day 10

	ALT	AST	SEQ ID NO
PBS	27	36	
ISIS 233710	19	37	233
ISIS 233717	16	32	14
ISIS 337477	22	35	234
ISIS 337478	23	49	235
ISIS 337479	21	29	236
ISIS 337487	19	35	28

Study 2

Male Tg mice were maintained on a 12-hour light/dark cycle. Animals were acclimated for at least 7 days in the research facility before initiation of the experiment. Antisense oligonucleotides (ASOs) were prepared in buffered saline (PBS) and sterilized by filtering through a 0.2 micron filter. Oligonucleotides were dissolved in 0.9% PBS for injection.

Groups of mice received intraperitoneal injections of 5-10-5 MOE gapmers at a dose of 50 mg/kg once per week for 2 weeks. One group of mice received subcutaneous injections of PBS once weekly for 2 weeks. The PBS-injected groups served as the control groups to which the corresponding oligonucleotide-treated groups were compared.

RNA analysis

At the end of the treatment period, RNA was extracted from liver for real-time PCR analysis of measurement of mRNA expression of ANGPTL3 with RTS1984. Results are presented as percent change of mRNA, relative to PBS control, normalized with RIBOGREEN[®]. As shown in the Table below, treatment

with ISIS antisense oligonucleotides resulted in significant reduction of ANGPTL3 mRNA in comparison to the PBS control.

Table 47

Percent inhibition of ANGPTL3 mRNA in transgenic mouse liver relative to the PBS control

ISIS No	%	SEQ ID NO
233710	81	233
337487	92	28
544145	98	16
544162	75	18
544199	97	20
560306	90	34
560400	97	35
560401	95	36
560402	98	37
560469	98	38
560735	87	49
567320	95	93
567321	93	94

5

Protein analysis

Human ANGPTL3 protein levels were quantified using a commercially available ELISA kit (Catalog #DANL30 by R&D Systems, Minneapolis, MN) with transgenic plasma samples diluted 1:20,000 using the manufacturer described protocol. The results are presented in the Table below. The results indicate that treatment with ISIS oligonucleotides resulted in reduced ANGPTL3 protein levels.

10

Table 48

Percent inhibition of plasma protein levels in the transgenic mouse

ISIS No	%	SEQ ID NO
233710	96	233
337487	78	28
544145	96	16
544162	97	18
544199	98	20
560306	97	34
560400	98	35
560401	97	36
560402	94	37
560469	96	38
560735	91	49

567320	98	93
567321	96	94

Plasma chemistry markers

To evaluate the effect of ISIS oligonucleotides on day 8, plasma levels of transaminases (ALT and AST) 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 these liver function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 49

Plasma transaminase levels (IU/L) in transgenic mice on day 8

	ALT	AST	SEQ ID NO
PBS	29	44	
ISIS 233710	29	47	233
ISIS 337487	22	36	28
ISIS 544145	29	45	16
ISIS 544162	31	62	18
ISIS 544199	29	51	20
ISIS 560306	23	42	34
ISIS 560400	24	52	35
ISIS 560401	20	38	36
ISIS 560402	29	49	37
ISIS 560469	22	50	38
ISIS 560735	20	38	49
ISIS 567320	49	71	93
ISIS 567321	20	44	94

Study 3

Male and female Tg mice were maintained on a 12-hour light/dark cycle. Animals were acclimated for at least 7 days in the research facility before initiation of the experiment. Antisense oligonucleotides (ASOs) were prepared in buffered saline (PBS) and sterilized by filtering through a 0.2 micron filter.

Oligonucleotides were dissolved in 0.9% PBS for injection.

Groups of mice received intraperitoneal injections of 5-10-5 MOE gapmers at a dose of 2.5 mg/kg, 12.5 mg/kg, or 25 mg/kg once per week for 3 weeks. One group of mice received subcutaneous injections of PBS once weekly for 2 weeks. The PBS-injected groups served as the control groups to which the corresponding oligonucleotide-treated groups were compared.

RNA analysis

At the end of the treatment period, RNA was extracted from liver for real-time PCR analysis of measurement of mRNA expression of ANGPTL3 with hANGPTL3_LTS01022 (forward sequence
5 AAATTTTAGCCAATGGCCTCC, designated herein as SEQ ID NO: 10; reverse sequence
TGTCATTAATTTGGCCCTTCG, designated herein as SEQ ID NO: 11; probe sequence
TCAGTTGGGACATGGTCTTAAAGACTTTGTCC, designated herein as SEQ ID NO: 12). Results are
presented as percent change of mRNA, relative to PBS control, normalized with RIBOGREEN[®]. As shown
in the Table below, treatment with ISIS antisense oligonucleotides resulted in significant reduction of
10 ANGPTL3 mRNA in comparison to the PBS control. The ED₅₀ of each gapmer is also presented in the Table
below. 'n.d.' indicates that the ED₅₀ could not be determined.

Table 50

Percent inhibition of ANGPTL3 mRNA in transgenic mouse liver relative to the PBS control

ISIS No	Dose (mg/kg)	%	ED ₅₀	SEQ ID NO
233710	25	88	8	233
	12.5	79		
	2.5	0		
544145	25	90	4	16
	12.5	74		
	2.5	39		
544162	25	53	9	18
	12.5	63		
	2.5	39		
544199	25	81	7	20
	12.5	82		
	2.5	7		
560306	25	0	n.d.	34
	12.5	0		
	2.5	0		
560400	25	87	5	35
	12.5	76		
	2.5	24		
560401	25	89	8	36
	12.5	62		
	2.5	5		
560469	25	73	3	38
	12.5	78		
	2.5	50		

560735	25	26	31	49
	12.5	37		
	2.5	51		
567320	25	74	12	93
	12.5	37		
	2.5	32		
567321	25	75	11	94
	12.5	61		
	2.5	0		

Protein analysis

Human ANGPTL3 protein levels were quantified using a commercially available ELISA kit (Catalog #DANL30 by R&D Systems, Minneapolis, MN) with transgenic plasma samples diluted 1:20,000 using the manufacturer described protocol. The results are presented in the Table below. The results indicate that treatment with ISIS oligonucleotides resulted in reduced ANGPTL3 protein levels. ‘n.d.’ indicates that the ED₅₀ could not be determined.

Table 51

Percent inhibition of plasma protein levels in the transgenic mouse

ISIS No	Dose (mg/kg)	%	ED ₅₀	SEQ ID NO
233710	25	80	11	233
	12.5	56		
	2.5	0		
544145	25	88	9	16
	12.5	64		
	2.5	0		
544162	25	56	15	18
	12.5	46		
	2.5	24		
544199	25	73	6	20
	12.5	73		
	2.5	31		
560306	25	63	n.d.	34
	12.5	55		
	2.5	53		
560400	25	88	6	35
	12.5	73		
	2.5	20		
560401	25	88	10	36
	12.5	61		
	2.5	0		

560469	25	75	4	38
	12.5	70		
	2.5	52		
560735	25	27	34	49
	12.5	37		
	2.5	34		
567320	25	69	10	93
	12.5	44		
	2.5	39		
567321	25	68	12	94
	12.5	62		
	2.5	1		

Plasma chemistry markers

To evaluate the effect of ISIS oligonucleotides on day 17, plasma levels of transaminases (ALT and AST) 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 these liver function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 52

Plasma transaminase levels (IU/L) in transgenic mice on day 17

	Dose (mg/kg)	ALT	AST	SEQ ID NO
PBS	-	25	38	
ISIS 233710	25	27	40	233
	12.5	24	45	
	2.5	23	36	
ISIS 544145	25	30	56	16
	12.5	25	52	
	2.5	28	43	
ISIS 544162	25	28	52	18
	12.5	36	53	
	2.5	28	50	
ISIS 544199	25	24	47	20
	12.5	23	60	
	2.5	24	44	
ISIS 560306	25	21	45	34
	12.5	24	49	
	2.5	24	47	

ISIS 560400	25	22	38	35
	12.5	21	53	
	2.5	23	52	
ISIS 560401	25	36	80	36
	12.5	27	75	
	2.5	22	49	
ISIS 560469	25	24	121	38
	12.5	23	53	
	2.5	21	88	
ISIS 560735	25	20	48	49
	12.5	22	138	
	2.5	24	78	
ISIS 567320	25	21	65	93
	12.5	20	58	
	2.5	23	46	
ISIS 567321	25	23	62	94
	12.5	21	49	
	2.5	24	67	

Study 4

Male and female Tg mice were maintained on a 12-hour light/dark cycle. Animals were acclimated for at least 7 days in the research facility before initiation of the experiment. Antisense oligonucleotides (ASOs) were prepared in buffered saline (PBS) and sterilized by filtering through a 0.2 micron filter. Oligonucleotides were dissolved in 0.9% PBS for injection.

Groups of mice received intraperitoneal injections of 5-10-5 MOE gapmers at a dose of 25 mg/kg once per week for 2 weeks. One group of mice received subcutaneous injections of PBS once weekly for 2 weeks. The PBS-injected group served as the control group to which the corresponding oligonucleotide-treated groups were compared.

RNA analysis

At the end of the treatment period, RNA was extracted from liver for real-time PCR analysis of measurement of mRNA expression of ANGPTL3 with hANGPTL3_LTS01022. Results are presented as percent change of mRNA, relative to PBS control, normalized with RIBOGREEN[®]. As shown in the Table below, treatment with ISIS antisense oligonucleotides resulted in significant reduction of ANGPTL3 mRNA in comparison to the PBS control.

Table 53

Percent inhibition of ANGPTL3 mRNA in transgenic mouse liver relative to the PBS control

ISIS No	%	SEQ ID NO
233710	68	233
544120	63	15
544199	82	20
544355	0	21
560268	36	32
560470	47	39
560471	67	40
560474	57	41
560566	45	42
560567	68	43
560607	37	46
560608	15	47
560744	25	51
560778	32	52
560811	27	54
560925	0	56
563639	5	79
567291	8	91
567330	30	95
568049	48	101
568146	26	104

Plasma chemistry markers

5 To evaluate the effect of ISIS oligonucleotides on day 10, plasma levels of transaminases (ALT and AST) 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 these liver function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 54

Plasma transaminase levels (IU/L) in transgenic mice on day 10

	ALT	AST	SEQ ID NO
PBS	29	41	
ISIS 233710	29	48	233
ISIS 544120	24	35	15
ISIS 544199	27	57	20

ISIS 544355	23	44	21
ISIS 560268	23	42	32
ISIS 560470	26	42	39
ISIS 560471	21	50	40
ISIS 560474	20	33	41
ISIS 560566	27	102	42
ISIS 560567	20	37	43
ISIS 560607	25	47	46
ISIS 560608	20	49	47
ISIS 560744	26	66	51
ISIS 560778	24	87	52
ISIS 560811	21	63	54
ISIS 560925	25	115	56
ISIS 563639	20	43	79
ISIS 567291	20	67	91
ISIS 567330	29	78	95
ISIS 568049	25	63	101
ISIS 568146	28	140	104

Study 5

Male and female Tg mice were maintained on a 12-hour light/dark cycle. Animals were acclimated for at least 7 days in the research facility before initiation of the experiment. Antisense oligonucleotides (ASOs) were prepared in buffered saline (PBS) and sterilized by filtering through a 0.2 micron filter. Oligonucleotides were dissolved in 0.9% PBS for injection.

Groups of mice received intraperitoneal injections of 5-10-5 MOE gapmers or deoxy, MOE, and cEt gapmers at a dose of 25 mg/kg once per week for 2 weeks. One group of mice received subcutaneous injections of PBS once weekly for 2 weeks. The PBS-injected group served as the control group to which the corresponding oligonucleotide-treated groups were compared.

RNA analysis

At the end of the treatment period, RNA was extracted from liver for real-time PCR analysis of measurement of mRNA expression of ANGPTL3 with RTS1984. Results are presented as percent change of mRNA, relative to PBS control, normalized with RIBOGREEN[®]. As shown in the Table below, treatment with ISIS antisense oligonucleotides resulted in significant reduction of ANGPTL3 mRNA in comparison to the PBS control.

Table 55

Percent inhibition of ANGPTL3 mRNA in transgenic mouse liver relative to the PBS control

ISIS No	Chemistry	%	SEQ ID NO
233710	5-10-5 MOE	79	233
544156	5-10-5 MOE	92	17
559277	Deoxy, MOE and cEt	75	110
560265	5-10-5 MOE	52	31
560285	5-10-5 MOE	42	33
560574	5-10-5 MOE	93	44
560847	5-10-5 MOE	61	69
560992	Deoxy, MOE and cEt	80	112
561010	Deoxy, MOE and cEt	66	113
561011	Deoxy, MOE and cEt	96	114
561022	Deoxy, MOE and cEt	79	115
561025	Deoxy, MOE and cEt	57	116
563580	5-10-5 MOE	80	77
567115	5-10-5 MOE	78	88
567233	5-10-5 MOE	91	90

Plasma chemistry markers

To evaluate the effect of ISIS oligonucleotides on day 9, plasma levels of transaminases (ALT and AST) 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 these liver function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 56

Plasma transaminase levels (IU/L) in transgenic mice on day 9

	Chemistry	ALT	AST	SEQ ID NO
PBS	-	48	65	
ISIS 233710	5-10-5 MOE	24	43	233
ISIS 544156	5-10-5 MOE	29	44	17
ISIS 559277	Deoxy, MOE and cEt	22	38	110
ISIS 560265	5-10-5 MOE	28	83	31
ISIS 560285	5-10-5 MOE	29	44	33
ISIS 560574	5-10-5 MOE	24	54	44
ISIS 560847	5-10-5 MOE	25	45	69
ISIS 560992	Deoxy, MOE and cEt	32	128	112
ISIS 561010	Deoxy, MOE and cEt	22	51	113

ISIS 561011	Deoxy, MOE and cEt	28	43	114
ISIS 561022	Deoxy, MOE and cEt	51	85	115
ISIS 561025	Deoxy, MOE and cEt	22	48	116
ISIS 563580	5-10-5 MOE	28	109	77
ISIS 567115	5-10-5 MOE	21	42	88
ISIS 567233	5-10-5 MOE	22	73	90

Study 6

Male and female Tg mice were maintained on a 12-hour light/dark cycle. Animals were acclimated for at least 7 days in the research facility before initiation of the experiment. Antisense oligonucleotides (ASOs) were prepared in buffered saline (PBS) and sterilized by filtering through a 0.2 micron filter. Oligonucleotides were dissolved in 0.9% PBS for injection.

Groups of mice received intraperitoneal injections of deoxy, MOE, and cEt oligonucleotides at a dose of 25 mg/kg once per week for 2 weeks. ISIS 233710, a 5-10-5 MOE gapmer, was also included as a benchmark. One group of mice received subcutaneous injections of PBS once weekly for 2 weeks. The PBS-injected group served as the control group to which the corresponding oligonucleotide-treated groups were compared.

RNA analysis

At the end of the treatment period, RNA was extracted from liver for real-time PCR analysis of measurement of mRNA expression of ANGPTL3 with hANGPTL3_LTS01022. Results are presented as percent change of mRNA, relative to PBS control, normalized with RIBOGREEN[®]. As shown in the Table below, treatment with several of the ISIS antisense oligonucleotides resulted in significant reduction of ANGPTL3 mRNA in comparison to the PBS control.

Table 57

Percent inhibition of ANGPTL3 mRNA in transgenic mouse liver relative to the PBS control

ISIS No	Chemistry	%	SEQ ID NO
233710	5-10-5 MOE	68	233
561026	Deoxy, MOE and cEt	94	117
561079	Deoxy, MOE and cEt	51	160
561084	Deoxy, MOE and cEt	56	161
561123	Deoxy, MOE and cEt	47	163
561208	Deoxy, MOE and cEt	42	118
561241	Deoxy, MOE and cEt	13	164
561400	Deoxy, MOE and cEt	31	173
561418	Deoxy, MOE and cEt	32	169

561436	Deoxy, MOE and cEt	67	170
561443	Deoxy, MOE and cEt	12	171
561458	Deoxy, MOE and cEt	57	124

Protein analysis

Human ANGPTL3 protein levels were quantified using a commercially available ELISA kit (Catalog #DANL30 by R&D Systems, Minneapolis, MN) with transgenic plasma samples diluted 1:20,000 using the manufacturer described protocol. The results are presented in the Table below. The results indicate that treatment with several of the ISIS oligonucleotides resulted in reduced ANGPTL3 protein levels.

Table 58

Percent inhibition of plasma protein levels in the transgenic mouse

ISIS No	Chemistry	%	SEQ ID NO
233710	5-10-5 MOE	82	233
561026	Deoxy, MOE and cEt	92	117
561079	Deoxy, MOE and cEt	80	160
561084	Deoxy, MOE and cEt	89	161
561123	Deoxy, MOE and cEt	62	163
561208	Deoxy, MOE and cEt	0	118
561241	Deoxy, MOE and cEt	36	164
561400	Deoxy, MOE and cEt	60	173
561418	Deoxy, MOE and cEt	42	169
561436	Deoxy, MOE and cEt	46	170
561443	Deoxy, MOE and cEt	27	171
561458	Deoxy, MOE and cEt	71	124

10 *Plasma chemistry markers*

To evaluate the effect of ISIS oligonucleotides on day 10, plasma levels of transaminases (ALT and AST) 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 these liver function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 59

Plasma transaminase levels (IU/L) in transgenic mice on day 10

	Chemistry	ALT	AST	SAE ID NO
PBS	-	41	64	
ISIS 233710	5-10-5 MOE	25	74	233
ISIS 561026	Deoxy, MOE and cEt	30	67	117
ISIS 561079	Deoxy, MOE and cEt	42	62	160
ISIS 561084	Deoxy, MOE and cEt	70	101	161
ISIS 561123	Deoxy, MOE and cEt	24	41	163
ISIS 561208	Deoxy, MOE and cEt	203	168	118
ISIS 561241	Deoxy, MOE and cEt	26	47	164
ISIS 561400	Deoxy, MOE and cEt	27	83	173
ISIS 561418	Deoxy, MOE and cEt	58	164	169
ISIS 561436	Deoxy, MOE and cEt	24	42	170
ISIS 561443	Deoxy, MOE and cEt	27	91	171
ISIS 561458	Deoxy, MOE and cEt	30	144	124

Study 7

5 Male and female Tg mice were maintained on a 12-hour light/dark cycle. Animals were acclimated for at least 7 days in the research facility before initiation of the experiment. Antisense oligonucleotides (ASOs) were prepared in buffered saline (PBS) and sterilized by filtering through a 0.2 micron filter. Oligonucleotides were dissolved in 0.9% PBS for injection.

10 Groups of mice received intraperitoneal injections of deoxy, MOE, and cEt oligonucleotides at a dose of 25 mg/kg once per week for 2 weeks. ISIS 233710, a 5-10-5 MOE gapmer, was also included as a benchmark. One group of mice received subcutaneous injections of PBS once weekly for 2 weeks. The PBS-injected group served as the control group to which the corresponding oligonucleotide-treated groups were compared.

RNA analysis

15 At the end of the treatment period, RNA was extracted from liver for real-time PCR analysis of measurement of mRNA expression of ANGPTL3 with hANGPTL3_LTS01022. Results are presented as percent change of mRNA, relative to PBS control, normalized with RIBOGREEN[®]. As shown in the Table below, treatment with ISIS antisense oligonucleotides resulted in significant reduction of ANGPTL3 mRNA in comparison to the PBS control.

Table 60

Percent inhibition of ANGPTL3 mRNA in transgenic mouse liver relative to the PBS control

ISIS No	Chemistry	%	SEQ ID NO
233710	5-10-5 MOE	80	233
561462	Deoxy, MOE and cEt	84	126
561463	Deoxy, MOE and cEt	84	127
561486	Deoxy, MOE and cEt	74	130
561487	Deoxy, MOE and cEt	82	131
561504	Deoxy, MOE and cEt	51	133
561528	Deoxy, MOE and cEt	87	174
561565	Deoxy, MOE and cEt	94	175
561566	Deoxy, MOE and cEt	76	176
561571	Deoxy, MOE and cEt	51	178
561621	Deoxy, MOE and cEt	93	134
561646	Deoxy, MOE and cEt	39	140
561649	Deoxy, MOE and cEt	93	141
561650	Deoxy, MOE and cEt	82	142
561689	Deoxy, MOE and cEt	51	180
561722	Deoxy, MOE and cEt	88	183
561723	Deoxy, MOE and cEt	85	184
561770	Deoxy, MOE and cEt	70	143
562024	Deoxy, MOE and cEt	82	189

Protein analysis

Human ANGPTL3 protein levels were quantified using a commercially available ELISA kit (Catalog #DANL30 by R&D Systems, Minneapolis, MN) with transgenic plasma samples diluted 1:20,000 using the manufacturer described protocol. The results are presented in the Table below. The results indicate that treatment with some of the ISIS oligonucleotides resulted in reduced ANGPTL3 levels. In this case, '0' value implies that treatment with the ISIS oligonucleotide did not inhibit expression; in some instances, increased levels of expression may have been recorded.

Table 61

Percent inhibition of plasma protein levels in the transgenic mouse

ISIS No	Chemistry	%	SEQ ID NO
233710	5-10-5 MOE	60	233
561462	Deoxy, MOE and cEt	62	126
561463	Deoxy, MOE and cEt	59	127
561486	Deoxy, MOE and cEt	0	130
561487	Deoxy, MOE and cEt	0	131

561504	Deoxy, MOE and cEt	0	133
561528	Deoxy, MOE and cEt	0	174
561565	Deoxy, MOE and cEt	71	175
561566	Deoxy, MOE and cEt	0	176
561571	Deoxy, MOE and cEt	0	178
561621	Deoxy, MOE and cEt	72	134
561646	Deoxy, MOE and cEt	0	140
561649	Deoxy, MOE and cEt	63	141
561650	Deoxy, MOE and cEt	0	142
561689	Deoxy, MOE and cEt	0	180
561722	Deoxy, MOE and cEt	0	183
561723	Deoxy, MOE and cEt	0	184
561770	Deoxy, MOE and cEt	0	143
562024	Deoxy, MOE and cEt	0	189

Plasma chemistry markers

To evaluate the effect of ISIS oligonucleotides on day 9, plasma levels of transaminases (ALT and AST) 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 these liver function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 62

Plasma transaminase levels (IU/L) in transgenic mice on day 9

	Chemistry	ALT	AST	SEQ ID NO
PBS	-	35	72	
ISIS 233710	5-10-5 MOE	23	39	233
ISIS 561462	Deoxy, MOE and cEt	26	56	126
ISIS 561463	Deoxy, MOE and cEt	34	61	127
ISIS 561486	Deoxy, MOE and cEt	23	61	130
ISIS 561487	Deoxy, MOE and cEt	21	64	131
ISIS 561504	Deoxy, MOE and cEt	26	66	133
ISIS 561528	Deoxy, MOE and cEt	26	86	174
ISIS 561565	Deoxy, MOE and cEt	24	43	175
ISIS 561566	Deoxy, MOE and cEt	23	62	176
ISIS 561571	Deoxy, MOE and cEt	26	68	178
ISIS 561621	Deoxy, MOE and cEt	26	96	134
ISIS 561646	Deoxy, MOE and cEt	24	77	140
ISIS 561649	Deoxy, MOE and cEt	22	94	141

ISIS 561650	Deoxy, MOE and cEt	34	121	142
ISIS 561689	Deoxy, MOE and cEt	24	73	180
ISIS 561722	Deoxy, MOE and cEt	34	89	183
ISIS 561723	Deoxy, MOE and cEt	24	65	184
ISIS 561770	Deoxy, MOE and cEt	22	69	143
ISIS 562024	Deoxy, MOE and cEt	32	162	189

Study 8

Male and female Tg mice were maintained on a 12-hour light/dark cycle. Animals were acclimated for at least 7 days in the research facility before initiation of the experiment. Antisense oligonucleotides (ASOs) were prepared in buffered saline (PBS) and sterilized by filtering through a 0.2 micron filter. Oligonucleotides were dissolved in 0.9% PBS for injection.

Groups of mice received intraperitoneal injections of deoxy, MOE, and cEt oligonucleotides at a dose of 25 mg/kg once per week for 2 weeks. ISIS 233710, a 5-10-5 MOE gapmer, was also included as a benchmark. One group of mice received subcutaneous injections of PBS once weekly for 2 weeks. The PBS-injected group served as the control group to which the corresponding oligonucleotide-treated groups were compared.

RNA analysis

At the end of the treatment period, RNA was extracted from liver for real-time PCR analysis of measurement of mRNA expression of ANGPTL3 with hANGPTL3_LTS01022. Results are presented as percent change of mRNA, relative to PBS control, normalized with RIBOGREEN[®]. As shown in the Table below, treatment with ISIS antisense oligonucleotides resulted in significant reduction of ANGPTL3 mRNA in comparison to the PBS control.

Table 63

Percent inhibition of ANGPTL3 mRNA in transgenic mouse liver relative to the PBS control

ISIS No	Chemistry	%	SEQ ID NO
233710	5-10-5 MOE	99	233
562078	Deoxy, MOE and cEt	73	147
562086	Deoxy, MOE and cEt	85	148
562103	Deoxy, MOE and cEt	58	149
562110	Deoxy, MOE and cEt	94	150
562155	Deoxy, MOE and cEt	85	192
562181	Deoxy, MOE and cEt	79	195
562433	Deoxy, MOE and cEt	59	155
562436	Deoxy, MOE and cEt	99	156

586669	Deoxy, MOE and cEt	95	210
586676	Deoxy, MOE and cEt	80	211

Protein analysis

Human ANGPTL3 protein levels were quantified using a commercially available ELISA kit (Catalog #DANL30 by R&D Systems, Minneapolis, MN) with transgenic plasma samples diluted 1:20,000 using the manufacturer described protocol. The results are presented in the Table below. The results indicate that treatment with the ISIS oligonucleotides resulted in reduced ANGPTL3 levels.

Table 64

Percent inhibition of plasma protein levels in the transgenic mouse

ISIS No	Chemistry	%	SEQ ID NO
233710	5-10-5 MOE	69	233
562078	Deoxy, MOE and cEt	44	147
562086	Deoxy, MOE and cEt	91	148
562103	Deoxy, MOE and cEt	26	149
562110	Deoxy, MOE and cEt	68	150
562155	Deoxy, MOE and cEt	75	192
562181	Deoxy, MOE and cEt	86	195
562433	Deoxy, MOE and cEt	80	155
562436	Deoxy, MOE and cEt	98	156
586669	Deoxy, MOE and cEt	98	210
586676	Deoxy, MOE and cEt	95	211

Plasma chemistry markers

To evaluate the effect of ISIS oligonucleotides on day 8, plasma levels of transaminases (ALT and AST) 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 these liver function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 65

Plasma transaminase levels (IU/L) in transgenic mice on day 8

	Chemistry	ALT	AST	SEQ ID NO
PBS	-	44	248	
ISIS 233710	5-10-5 MOE	27	52	233
ISIS 562078	Deoxy, MOE and cEt	41	130	147

ISIS 562086	Deoxy, MOE and cEt	30	62	148
ISIS 562103	Deoxy, MOE and cEt	35	99	149
ISIS 562110	Deoxy, MOE and cEt	30	161	150
ISIS 562155	Deoxy, MOE and cEt	68	622	192
ISIS 562181	Deoxy, MOE and cEt	37	168	195
ISIS 562433	Deoxy, MOE and cEt	33	209	155
ISIS 562436	Deoxy, MOE and cEt	30	93	156
ISIS 586669	Deoxy, MOE and cEt	27	141	210
ISIS 586676	Deoxy, MOE and cEt	22	60	211

Study 9

Male and female Tg mice were maintained on a 12-hour light/dark cycle. Animals were acclimated for at least 7 days in the research facility before initiation of the experiment. Antisense oligonucleotides (ASOs) were prepared in buffered saline (PBS) and sterilized by filtering through a 0.2 micron filter. Oligonucleotides were dissolved in 0.9% PBS for injection.

Groups of mice received intraperitoneal injections of deoxy, MOE, and cEt oligonucleotides at a dose of 25 mg/kg once per week for 2 weeks. ISIS 233710, a 5-10-5 MOE gapmer, was also included as a benchmark. One group of mice received subcutaneous injections of PBS once weekly for 2 weeks. The PBS-injected group served as the control group to which the corresponding oligonucleotide-treated groups were compared.

RNA analysis

At the end of the treatment period, RNA was extracted from liver for real-time PCR analysis of measurement of mRNA expression of ANGPTL3 with hANGPTL3_LTS01022. Results are presented as percent change of mRNA, relative to PBS control, normalized with RIBOGREEN[®]. As shown in the Table below, treatment with some of the ISIS antisense oligonucleotides resulted in significant reduction of ANGPTL3 mRNA in comparison to the PBS control. In this case, '0' value implies that treatment with the ISIS oligonucleotide did not inhibit expression; in some instances, increased levels of expression may have been recorded.

Table 66

Percent inhibition of ANGPTL3 mRNA in transgenic mouse liver relative to the PBS control

ISIS No	Chemistry	%	SEQ ID NO
233710	5-10-5 MOE	84	233
586690	Deoxy, MOE and cEt	45	213
586692	Deoxy, MOE and cEt	45	220

586700	Deoxy, MOE and cEt	46	221
586707	Deoxy, MOE and cEt	88	218
586708	Deoxy, MOE and cEt	73	222
586718	Deoxy, MOE and cEt	20	219
586744	Deoxy, MOE and cEt	0	223
586745	Deoxy, MOE and cEt	0	224
586755	Deoxy, MOE and cEt	75	226
586761	Deoxy, MOE and cEt	66	227
586787	Deoxy, MOE and cEt	47	228
586796	Deoxy, MOE and cEt	88	229
586797	Deoxy, MOE and cEt	81	230
586802	Deoxy, MOE and cEt	33	231
586804	Deoxy, MOE and cEt	60	232

Protein analysis

Human ANGPTL3 protein levels were quantified using a commercially available ELISA kit (Catalog #DANL30 by R&D Systems, Minneapolis, MN) with transgenic plasma samples diluted 1:20,000 using the manufacturer described protocol. The results are presented in the Table below. The results indicate that treatment with some of the ISIS oligonucleotides resulted in reduced ANGPTL3 levels. In this case, '0' value implies that treatment with the ISIS oligonucleotide did not inhibit expression; in some instances, increased levels of expression may have been recorded.

Table 67

Percent inhibition of plasma protein levels in the transgenic mouse

ISIS No	Chemistry	%	SEQ ID NO
233710	5-10-5 MOE	80	233
586690	Deoxy, MOE and cEt	21	213
586692	Deoxy, MOE and cEt	46	220
586700	Deoxy, MOE and cEt	0	221
586707	Deoxy, MOE and cEt	84	218
586708	Deoxy, MOE and cEt	32	222
586718	Deoxy, MOE and cEt	0	219
586744	Deoxy, MOE and cEt	0	223
586745	Deoxy, MOE and cEt	0	224
586755	Deoxy, MOE and cEt	0	226
586761	Deoxy, MOE and cEt	0	227
586787	Deoxy, MOE and cEt	0	228
586796	Deoxy, MOE and cEt	40	229
586797	Deoxy, MOE and cEt	50	230
586802	Deoxy, MOE and cEt	0	231
586804	Deoxy, MOE and cEt	0	232

Plasma chemistry markers

To evaluate the effect of ISIS oligonucleotides on day 9, plasma levels of transaminases (ALT and AST) 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 these liver function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 68

Plasma transaminase levels (IU/L) in transgenic mice on day 9

	Chemistry	ALT	AST	SEQ ID NO
PBS	-	28	73	
ISIS 233710	5-10-5 MOE	22	86	233
ISIS 586690	Deoxy, MOE and cEt	42	120	213
ISIS 586692	Deoxy, MOE and cEt	22	45	220
ISIS 586700	Deoxy, MOE and cEt	24	84	221
ISIS 586707	Deoxy, MOE and cEt	26	44	218
ISIS 586708	Deoxy, MOE and cEt	22	48	222
ISIS 586718	Deoxy, MOE and cEt	22	39	219
ISIS 586744	Deoxy, MOE and cEt	26	83	223
ISIS 586745	Deoxy, MOE and cEt	25	56	224
ISIS 586746	Deoxy, MOE and cEt	77	77	225
ISIS 586755	Deoxy, MOE and cEt	28	148	226
ISIS 586761	Deoxy, MOE and cEt	36	126	227
ISIS 586787	Deoxy, MOE and cEt	23	88	228
ISIS 586796	Deoxy, MOE and cEt	32	148	229
ISIS 586797	Deoxy, MOE and cEt	29	151	230
ISIS 586802	Deoxy, MOE and cEt	35	200	231
ISIS 586804	Deoxy, MOE and cEt	24	87	232

Study 10

Male and female Tg mice were maintained on a 12-hour light/dark cycle. Animals were acclimated for at least 7 days in the research facility before initiation of the experiment. Antisense oligonucleotides (ASOs) were prepared in buffered saline (PBS) and sterilized by filtering through a 0.2 micron filter.

Oligonucleotides were dissolved in 0.9% PBS for injection.

Groups of mice received intraperitoneal injections of 5-10-5 MOE gapmers or deoxy, MOE and cEt oligonucleotides at a dose of 5 mg/kg, 12.5 mg/kg, or 25 mg/kg once per week for 2 weeks. One group of

mice received subcutaneous injections of PBS once weekly for 2 weeks. The PBS-injected group served as the control group to which the corresponding oligonucleotide-treated groups were compared.

RNA analysis

- 5 At the end of the treatment period, RNA was extracted from liver for real-time PCR analysis of measurement of mRNA expression of ANGPTL3 with hANGPTL3_LTS01022, and also with RTS3492_MGB. Results are presented as percent change of mRNA, relative to PBS control, normalized with RIBOGREEN[®]. As shown in the Table below, treatment with some of the ISIS antisense oligonucleotides resulted in reduction of ANGPTL3 mRNA in comparison to the PBS control.

10

Table 69

Percent inhibition of ANGPTL3 mRNA in transgenic mouse liver relative to the PBS control

ISIS No	Chemistry	Dose (mg/kg)	RTS3492_MGB	hANGPTL3_LTS01022	SEQ ID NO
233710	5-10-5 MOE	25	0	8	233
		12.5	24	22	
		5	12	22	
544199	5-10-5 MOE	25	63	59	20
		12.5	43	43	
		5	17	24	
559277	Deoxy, MOE and cEt	25	37	46	110
		12.5	0	0	
		5	0	0	
560400	5-10-5 MOE	25	45	48	35
		12.5	36	50	
		5	0	0	
561010	Deoxy, MOE and cEt	25	5	37	113
		12.5	0	6	
		5	0	0	
563580	5-10-5 MOE	25	56	59	77
		12.5	43	44	
		5	5	9	
567320	5-10-5 MOE	25	47	50	93
		12.5	0	0	
		5	0	0	
567321	5-10-5 MOE	25	46	32	94
		12.5	0	0	
		5	0	0	

Plasma chemistry markers

To evaluate the effect of ISIS oligonucleotides on day 8, plasma levels of transaminases (ALT and AST) 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 these liver function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 70

Plasma transaminase levels (IU/L) in transgenic mice on day 8

	Chemistry	Dose (mg/kg)	ALT	AST	SEQ ID NO
PBS	-	-	22	82	
ISIS 233710	5-10-5 MOE	25	21	41	233
		12.5	23	66	
		5	22	118	
ISIS 544199	5-10-5 MOE	25	25	47	20
		12.5	20	40	
		5	27	43	
ISIS 559277	Deoxy, MOE and cEt	25	21	34	110
		12.5	21	37	
		5	22	39	
ISIS 560400	5-10-5 MOE	25	21	37	35
		12.5	20	44	
		5	24	35	
ISIS 561010	Deoxy, MOE and cEt	25	22	48	113
		12.5	33	64	
		5	24	41	
ISIS 563580	5-10-5 MOE	25	21	36	77
		12.5	29	81	
		5	21	59	
ISIS 567320	5-10-5 MOE	25	22	47	93
		12.5	29	58	
		5	21	70	
ISIS 567321	5-10-5 MOE	25	20	50	94
		12.5	24	102	
		5	19	53	

Example 12: Tolerability of antisense oligonucleotides targeting human ANGPTL3 in CD1 mice

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.

5

Study 1

Male CD1 mice (one animal per treatment group) were injected intraperitoneally with a single dose of 200 mg/kg of deoxy, MOE, and cEt oligonucleotide. One male CD1 mouse was injected subcutaneously with a single dose of PBS. Mice were euthanized 48 hours after the last dose, and organs and plasma were
10 harvested for further analysis.

Plasma chemistry markers

To evaluate the effect of ISIS oligonucleotides on day 4 plasma levels of transaminases (ALT and AST) were measured using an automated clinical chemistry analyzer (Hitachi Olympus AU400e, Melville,
15 NY). The results are presented in the Table below. ISIS oligonucleotides that caused changes in the levels of any of these liver function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 71

Plasma transaminase levels in CD1 mice plasma on day 4

	ALT (IU/L)	AST (IU/L)	SEQ ID NO
ISIS 559277	29	43	110
ISIS 560990	19	43	111
ISIS 560992	21	36	112
ISIS 561010	31	40	113
ISIS 561011	27	32	114
ISIS 561022	35	48	115
ISIS 561025	17	28	116
ISIS 561026	31	43	117
ISIS 561208	32	47	118
ISIS 561320	25	37	119
ISIS 561343	41	90	120
ISIS 561345	30	45	121
ISIS 561347	31	41	122
ISIS 561458	18	38	124
ISIS 561460	42	59	125
ISIS 561463	21	33	127
ISIS 561486	17	39	130

ISIS 561487	18	39	131
ISIS 561504	24	41	133
ISIS 561621	31	56	134

Body weights

Body weights were measured one day after the single dose of ISIS oligonucleotide, 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 72

Body weights (g) of CD1 mice after antisense oligonucleotide treatment

	Body weight	SEQ ID NO
ISIS 559277	27	110
ISIS 560990	28	111
ISIS 560992	29	112
ISIS 561010	30	113
ISIS 561011	27	114
ISIS 561022	24	115
ISIS 561025	28	116
ISIS 561026	27	117
ISIS 561208	29	118
ISIS 561320	27	119
ISIS 561343	24	120
ISIS 561345	25	121
ISIS 561347	28	122
ISIS 561458	25	124
ISIS 561460	26	125
ISIS 561463	26	127
ISIS 561486	26	130
ISIS 561487	27	131
ISIS 561504	26	133
ISIS 561621	27	134

10 Study 2

Male CD1 mice (one animal per treatment group) were injected intraperitoneally with a single dose of 200 mg/kg of deoxy, MOE and cEt oligonucleotides. One male CD1 mouse was injected subcutaneously with a single dose of 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 day 5 plasma levels of transaminases (ALT and AST) 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 these liver function markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 73

Plasma transaminase levels in CD1 mice plasma on day 5

	ALT (IU/L)	AST (IU/L)	SEQ ID NO
ISIS 561622	29	64	136
ISIS 561628	17	24	137
ISIS 561646	16	34	140
ISIS 561650	32	51	142
ISIS 561079	19	32	160
ISIS 561084	24	56	161
ISIS 561241	60	70	164
ISIS 561462	22	54	126
ISIS 561649	56	53	141
ISIS 561770	23	39	143
ISIS 561781	20	41	144
ISIS 561918	31	112	146
ISIS 562078	15	33	147
ISIS 562086	19	32	148
ISIS 562110	20	41	150
ISIS 562415	13	30	154
ISIS 562433	19	35	155
ISIS 562436	21	37	156
ISIS 562442	19	34	158

Body weights

Body weights were measured on day 5 after the single dose of ISIS oligonucleotide, 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 74

Body weights (g) of CD1 mice after antisense oligonucleotide treatment

	Body weights	SEQ ID NO
ISIS 561622	27	136
ISIS 561628	28	137
ISIS 561646	29	140
ISIS 561650	30	142
ISIS 561079	27	160
ISIS 561084	24	161
ISIS 561241	28	164
ISIS 561462	27	126
ISIS 561649	29	141
ISIS 561770	27	143
ISIS 561781	24	144
ISIS 561918	25	146
ISIS 562078	28	147
ISIS 562086	25	148
ISIS 562110	26	150
ISIS 562415	26	154
ISIS 562433	26	155
ISIS 562436	27	156
ISIS 562442	26	158

5 Study 3

Male CD1 mice (four animals per treatment group) were injected intraperitoneally with 100 mg/kg of 5-10-5 MOE gapmers given once a week for 6 weeks. One group of 4 male CD1 mice was injected intraperitoneally with PBS given once a week for 6 weeks. Mice were euthanized 48 hours after the last dose, and organs and plasma were harvested for further analysis.

10 *Plasma chemistry markers*

To evaluate the effect of ISIS oligonucleotides, plasma levels of various liver and kidney function markers were measured on day 45 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 these markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 75

Plasma chemistry marker levels in CD1 mice plasma on day 45

	ALT (IU/L)	AST (IU/L)	Albumin (g/dL)	BUN (mg/dL)	Creatinine (mg/dL)	Bilirubin (mg/dL)	SEQ ID NO
PBS	30	55	2.7	26	0.15	0.17	
ISIS 544145	1146	1081	2.5	29	0.14	0.24	16
ISIS 544199	244	213	2.6	25	0.13	0.15	20
ISIS 560400	211	244	2.5	28	0.14	0.14	35
ISIS 560401	212	269	2.4	31	0.14	0.12	36
ISIS 560469	165	160	2.4	24	0.11	0.14	38
ISIS 567320	141	146	2.7	25	0.14	0.15	93
ISIS 567321	106	122	2.5	24	0.11	0.13	94

Body weights

5 Body weights were measured on day 43, and are presented in the Table below. Kidney, liver and spleen weights were measured at the end of the study on day 45. ISIS oligonucleotides that caused any changes in organ weights outside the expected range for antisense oligonucleotides were excluded from further studies.

Table 76

Weights (g) of CD1 mice after antisense oligonucleotide treatment

	Body	Kidney	Liver	Spleen	SEQ ID NO
PBS	39	0.6	2.1	0.1	
ISIS 544145	30	0.5	1.9	0.1	16
ISIS 544199	42	0.6	2.9	0.3	20
ISIS 560400	40	0.6	2.8	0.3	35
ISIS 560401	38	0.6	2.7	0.2	36
ISIS 560469	40	0.6	2.7	0.2	38
ISIS 567320	39	0.6	2.3	0.3	93
ISIS 567321	42	0.6	2.6	0.3	94

Study 4

15 Male CD1 mice (four animals per treatment group) were injected intraperitoneally with 50 mg/kg or 100 mg/kg of 5-10-5 MOE gapmers or deoxy, MOE and cEt oligonucleotides given once a week for 6 weeks. One group of 4 male CD1 mice was injected intraperitoneally with PBS given once a week for 6 weeks. 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, plasma levels of various liver and kidney function markers were measured on day 46 using an automated clinical chemistry analyzer (Hitachi Olympus AU400e, Melville, NY). The results are presented in the Table below. ISIS oligonucleotides that caused

5 changes in the levels of any of these markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 77

Plasma chemistry marker levels in CD1 mice plasma on day 45

	Chemistry	Dose (mg/kg)	ALT (IU/L)	AST (IU/L)	Albumin (g/dL)	BUN (mg/dL)	Creatinine (mg/dL)	Bilirubin (mg/dL)	SEQ ID NO
PBS		-	28	46	2.7	28	0.13	0.13	
ISIS 544156	5-10-5 MOE	100	80	145	2.2	26	0.12	0.10	17
ISIS 560574	5-10-5 MOE	100	182	184	2.5	25	0.14	0.15	44
ISIS 561010	Deoxy, MOE and cEt	50	32	53	2.4	31	0.15	0.12	113
ISIS 561011	Deoxy, MOE and cEt	50	93	152	1.8	27	0.15	0.08	114
ISIS 560580	5-10-5 MOE	100	50	76	2.5	25	0.12	0.13	237
ISIS 567115	5-10-5 MOE	100	202	304	2.5	19	0.14	0.12	88
ISIS 567233	5-10-5 MOE	100	123	145	2.5	24	0.12	0.12	90

10 *Body weights*

Body weights were measured on day 44, and are presented in the Table below. Kidney, liver and spleen weights were measured at the end of the study on day 46. ISIS oligonucleotides that caused any changes in organ weights outside the expected range for antisense oligonucleotides were excluded from further studies.

15

Table 78

Weights (g) of CD1 mice after antisense oligonucleotide treatment

	Chemistry	Dose (mg/kg)	Body	Kidney	Liver	Spleen	SEQ ID NO
PBS		-	38	0.6	2.1	0.2	
ISIS 544156	5-10-5 MOE	100	36	0.5	2.2	0.2	17
ISIS 560574	5-10-5 MOE	100	40	0.6	2.6	0.4	44
ISIS 561010	Deoxy, MOE and cEt	50	39	0.5	2.2	0.2	113
ISIS 561011	Deoxy, MOE and cEt	50	39	0.6	2.9	0.3	114
ISIS 560580	5-10-5 MOE	100	39	0.5	2.4	0.2	237
ISIS 567115	5-10-5 MOE	100	36	0.5	2.2	0.2	88
ISIS 567233	5-10-5 MOE	100	39	0.6	2.2	0.3	90

Study 5

Male CD1 mice (four animals per treatment group) were injected intraperitoneally with 50 mg/kg of deoxy, MOE and cEt oligonucleotides given once a week for 6 weeks. One group of 4 male CD1 mice was injected intraperitoneally with PBS given once a week for 6 weeks. 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, plasma levels of various liver and kidney function markers were measured on day 43 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 these markers outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 79

Plasma chemistry marker levels in CD1 mice plasma on day 43

	ALT (IU/L)	AST (IU/L)	Albumin (g/dL)	BUN (mg/dL)	Creatinine (mg/dL)	Bilirubin (mg/dL)	SEQ ID NO
PBS	35	166	2.6	29	0.12	0.32	
ISIS 559277	45	77	2.5	29	0.13	0.16	110
ISIS 561022	826	802	2.9	29	0.13	0.99	115
ISIS 561025	146	183	2.3	28	0.14	0.13	116
ISIS 561026	93	154	2.6	26	0.11	0.16	117
ISIS 561079	1943	1511	2.9	28	0.15	0.94	160
ISIS 561084	153	227	2.6	27	0.12	0.16	161
ISIS 561123	49	90	2.5	31	0.13	0.13	163
ISIS 561436	29	57	2.6	25	0.12	0.12	170

Body weights

Body weights were measured on day 41, and are presented in the Table below. Kidney, liver and spleen weights were measured at the end of the study on day 43. ISIS oligonucleotides that caused any changes in organ weights outside the expected range for antisense oligonucleotides were excluded from further studies.

Table 80

Weights (g) of CD1 mice after antisense oligonucleotide treatment

	Body	Kidney	Liver	Spleen	SEQ ID NO
PBS	37	0.5	2.0	0.1	
ISIS 559277	38	0.6	2.5	0.3	110

ISIS 561022	31	0.4	3.2	0.1	115
ISIS 561025	37	0.5	2.6	0.2	116
ISIS 561026	39	0.6	2.1	0.2	117
ISIS 561079	42	0.6	4.0	0.2	160
ISIS 561084	37	0.6	2.4	0.2	161
ISIS 561123	36	0.6	2.2	0.2	163
ISIS 561436	41	0.6	2.4	0.2	170

Example 13: Measurement of viscosity of ISIS antisense oligonucleotides targeting human ANGPTL3

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 of 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, where the concentration of each antisense oligonucleotide was 350 mg/ml, and indicate that most of the antisense oligonucleotides solutions are optimal in their viscosity under the criterion stated above.

Table 81

Viscosity of ISIS antisense oligonucleotides targeting human ANGPTL3

ISIS No.	Viscosity (cP)	SEQ ID NO
233710	14.65	233
337478	13.34	235
544145	11.97	16
544162	8.50	18
544199	11.70	20
560306	5.67	34
560400	9.26	35
560401	18.11	36
560402	90.67	37
560469	12.04	38
560735	7.49	49
567320	9.05	93
567321	9.62	94
567233	6.72	90
563580	16.83	77

561010	26.32	113
561011	43.15	114

Example 14: Tolerability of antisense oligonucleotides targeting human ANGPTL3 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 from the studies described in the Examples above and evaluated for changes in the levels of various plasma chemistry markers.

Study 1

Male Sprague-Dawley rats were maintained on a 12-hour light/dark cycle and fed ad libitum with Purina normal rat chow, diet 5001. Groups of 4 Sprague-Dawley rats each were injected subcutaneously once a week for 6 weeks with PBS or with 100 mg/kg of 5-10-5 MOE gapmers. Forty eight hours after the last dose, rats were euthanized and organs and plasma were harvested for further analysis.

Liver function

To evaluate the effect of ISIS oligonucleotides on hepatic function, plasma levels of transaminases were measured using an automated clinical chemistry analyzer (Hitachi Olympus AU400e, Melville, NY). Plasma levels of ALT (alanine transaminase) and AST (aspartate transaminase) were measured on day 45 and the results are presented in the Table below expressed in IU/L. Plasma levels of bilirubin were also measured using the same clinical chemistry analyzer and the results are also presented in the Table below expressed in mg/dL. ISIS oligonucleotides that caused changes in the levels of any markers of liver function outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 82

Liver function markers in Sprague-Dawley rats

	ALT (IU/L)	AST (IU/L)	Bilirubin (mg/dL)	SEQ ID NO
PBS	25	65	0.11	
ISIS 544145	225	407	0.30	16
ISIS 544199	56	102	0.11	20
ISIS 560400	55	175	0.12	35
ISIS 560401	89	206	0.13	36
ISIS 560469	227	290	0.15	38
ISIS 567320	55	172	0.11	93
ISIS 567321	39	109	0.10	94

Kidney function

To evaluate the effect of ISIS oligonucleotides on kidney function, plasma levels of blood urea nitrogen (BUN) 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. ISIS oligonucleotides that caused changes in the levels of any of the kidney function markers outside the expected range for antisense oligonucleotides were excluded in further studies. Total urine protein and urine creatinine levels were measured, and the ratio of total urine protein to creatinine was evaluated. The results are presented in the Table below.

Table 83

Kidney function plasma markers (mg/dL) in Sprague-Dawley rats

	BUN	Creatinine	SEQ ID NO
PBS	16	0.27	
ISIS 544145	53	0.26	16
ISIS 544199	24	0.34	20
ISIS 560400	28	0.31	35
ISIS 560401	29	0.28	36
ISIS 560469	23	0.32	38
ISIS 567320	26	0.35	93
ISIS 567321	24	0.37	94

Table 84

Kidney function urine markers in Sprague-Dawley rats

	Creatinine (mg/dL)	Total protein (mg/dL)	Protein: Creatinine ratio	SEQ ID NO
PBS	59	90	1.5	
ISIS 544145	27	2131	84.8	16
ISIS 544199	24	199	8.6	20
ISIS 560400	32	176	5.4	35
ISIS 560401	29	521	17.3	36
ISIS 560469	43	351	8.2	38
ISIS 567320	34	177	5.2	93
ISIS 567321	54	269	5.3	94

Organ weights

Body weights were measured on day 42 and presented in the Table below. Liver, spleen and kidney weights were measured at the end of the study on day 45, 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 85

Body and organ weights (g) of Sprague Dawley rats

	Body	Kidney	Liver	Spleen	SEQ ID NO
PBS	441	3.3	11.8	0.8	
ISIS 544145	240	3.0	11.2	1.7	16
ISIS 544199	307	2.6	10.3	2.0	20
ISIS 560400	294	2.8	12.3	2.0	35
ISIS 560401	281	3.4	11.6	2.3	36
ISIS 560469	316	3.0	11.8	2.0	38
ISIS 567320	312	3.1	12.4	2.5	93
ISIS 567321	332	3.3	11.6	2.3	94

5

Study2

Male Sprague-Dawley rats were maintained on a 12-hour light/dark cycle and fed ad libitum with Purina normal rat chow, diet 5001. Groups of 4 Sprague-Dawley rats each were injected subcutaneously once a week for 6 weeks with PBS or with 50 mg/kg or 100 mg/kg of 5-10-5 MOE gapmers or deoxy, MOE and cEt oligonucleotides. Forty eight hours after the last dose, rats were euthanized and organs and plasma were harvested for further analysis.

10

Liver function

To evaluate the effect of ISIS oligonucleotides on hepatic function, plasma levels of transaminases were measured using an automated clinical chemistry analyzer (Hitachi Olympus AU400e, Melville, NY). Plasma levels of ALT (alanine transaminase) and AST (aspartate transaminase) were measured on day 44 and the results are presented in the Table below expressed in IU/L. Plasma levels of bilirubin were also measured using the same clinical chemistry analyzer and the results are also presented in the Table below expressed in mg/dL. ISIS oligonucleotides that caused changes in the levels of any markers of liver function outside the expected range for antisense oligonucleotides were excluded in further studies.

15

Table 86

Liver function markers in Sprague-Dawley rats

	Chemistry	Dose (mg/kg)	ALT (IU/L)	AST (IU/L)	Bilirubin (mg/dL)	SEQ ID NO
PBS	-	-	22	63	0.09	
ISIS 544156	5-10-5 MOE	100	153	221	0.19	17
ISIS 560574	5-10-5 MOE	100	62	128	0.24	44

20

ISIS 561010	Deoxy, MOE and cEt	50	32	99	0.12	113
ISIS 561011	Deoxy, MOE and cEt	50	56	100	0.11	114
ISIS 563580	5-10-5 MOE	100	74	89	0.09	77
ISIS 567233	5-10-5 MOE	100	41	136	0.08	90

Kidney function

To evaluate the effect of ISIS oligonucleotides on kidney function, plasma levels of blood urea nitrogen (BUN) and creatinine were measured on day 44 using an automated clinical chemistry analyzer (Hitachi Olympus AU400e, Melville, NY). Results are presented in the Table below, expressed in mg/dL. ISIS oligonucleotides that caused changes in the levels of any of the kidney function markers outside the expected range for antisense oligonucleotides were excluded in further studies. Total urine protein and urine creatinine levels were measured, and the ratio of total urine protein to creatinine was evaluated. The results are presented in the Table below.

Table 87

Kidney function plasma markers (mg/dL) in Sprague-Dawley rats

	Chemistry	Dose (mg/kg)	BUN	Creatinine	SEQ ID NO
PBS	-	-	18	0.31	
ISIS 544156	5-10-5 MOE	100	27	0.27	17
ISIS 560574	5-10-5 MOE	100	32	0.24	44
ISIS 561010	Deoxy, MOE and cEt	50	24	0.31	113
ISIS 561011	Deoxy, MOE and cEt	50	33	0.32	114
ISIS 563580	5-10-5 MOE	100	25	0.20	77
ISIS 567233	5-10-5 MOE	100	37	0.23	90

Table 88

Kidney function urine markers in Sprague-Dawley rats

	Chemistry	Dose (mg/kg)	Creatinine (mg/dL)	Total protein (mg/dL)	Protein: Creatinine ratio	SEQ ID NO
PBS	-	-	55	66	1.2	
ISIS 544156	5-10-5 MOE	100	26	166	6.2	17
ISIS 560574	5-10-5 MOE	100	39	276	6.8	44
ISIS 561010	Deoxy, MOE and cEt	50	54	299	5.6	113
ISIS 561011	Deoxy, MOE and cEt	50	41	525	11.7	114
ISIS 563580	5-10-5 MOE	100	44	338	8.1	77
ISIS 567233	5-10-5 MOE	100	46	307	6.4	90

Organ weights

Body weights were measured on day 42 and presented in the Table below. Liver, spleen and kidney weights were measured at the end of the study on day 44, 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 89

Body and organ weights (g) of Sprague Dawley rats

	Chemistry	Dose (mg/kg)	Body	Kidney	Liver	Spleen	SEQ ID NO
PBS	-	-	433	3.1	10.8	0.6	
ISIS 544156	5-10-5 MOE	100	291	2.4	10.6	1.6	17
ISIS 560574	5-10-5 MOE	100	315	3.1	10.7	2.1	44
ISIS 561010	Deoxy, MOE and cEt	50	386	3.0	11.9	2.1	113
ISIS 561011	Deoxy, MOE and cEt	50	324	4.1	12.5	2.4	114
ISIS 563580	5-10-5 MOE	100	358	3.0	12.8	1.5	77
ISIS 567233	5-10-5 MOE	100	286	2.9	13.0	2.9	90

Study3

Male Sprague-Dawley rats were maintained on a 12-hour light/dark cycle and fed ad libitum with Purina normal rat chow, diet 5001. Groups of 4 Sprague-Dawley rats each were injected subcutaneously once a week for 6 weeks with PBS or with 50 mg/kg of deoxy, MOE and cEt oligonucleotides. Forty eight hours after the last dose, rats were euthanized and organs and plasma were harvested for further analysis.

Liver function

To evaluate the effect of ISIS oligonucleotides on hepatic function, plasma levels of transaminases were measured using an automated clinical chemistry analyzer (Hitachi Olympus AU400e, Melville, NY). Plasma levels of ALT (alanine transaminase) and AST (aspartate transaminase) were measured on day 44 and the results are presented in the Table below expressed in IU/L. Plasma levels of bilirubin were also measured using the same clinical chemistry analyzer and the results are also presented in the Table below expressed in mg/dL. ISIS oligonucleotides that caused changes in the levels of any markers of liver function outside the expected range for antisense oligonucleotides were excluded in further studies.

Table 90

Liver function markers in Sprague-Dawley rats

	ALT (IU/L)	AST (IU/L)	Bilirubin (mg/dL)	SEQ ID NO
PBS	27	87	0.08	

ISIS 559277	36	108	0.10	110
ISIS 561025	150	260	0.15	116
ISIS 561026	53	105	0.08	117
ISIS 561079	87	196	0.09	160
ISIS 561084	62	177	0.11	161
ISIS 561123	39	94	0.07	163
ISIS 561436	64	225	0.13	170

Kidney function

To evaluate the effect of ISIS oligonucleotides on kidney function, plasma levels of blood urea nitrogen (BUN) and creatinine were measured on day 44 using an automated clinical chemistry analyzer (Hitachi Olympus AU400e, Melville, NY). Results are presented in the Table below, expressed in mg/dL. ISIS oligonucleotides that caused changes in the levels of any of the kidney function markers outside the expected range for antisense oligonucleotides were excluded in further studies. Total urine protein and urine creatinine levels were measured, and the ratio of total urine protein to creatinine was evaluated. The results are presented in the Table below.

10

Table 91

Kidney function plasma markers (mg/dL) in Sprague-Dawley rats

	BUN	Creatinine	SEQ ID NO
PBS	12	0.26	
ISIS 559277	16	0.30	110
ISIS 561025	24	0.34	116
ISIS 561026	61	0.38	117
ISIS 561079	87	0.67	160
ISIS 561084	24	0.35	161
ISIS 561123	16	0.31	163
ISIS 561436	39	0.37	170

Table 92

Kidney function urine markers in Sprague-Dawley rats

	Creatinine (mg/dL)	Total protein (mg/dL)	Protein: Creatinine ratio	SEQ ID NO
PBS	42	77	1.9	
ISIS 559277	35	253	7.2	110
ISIS 561025	47	583	14.3	116
ISIS 561026	22	1993	111.4	117

ISIS 561079	17	1313	75.5	160
ISIS 561084	73	571	7.9	161
ISIS 561123	33	925	29.5	163
ISIS 561436	25	789	36.6	170

Organ weights

Body weights were measured on day 42 and presented in the table below. Liver, spleen and kidney weights were measured at the end of the study on day 44, 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 93

Body and organ weights (g) of Sprague Dawley rats

	Body	Kidney	Liver	Spleen	SEQ ID NO
PBS	419	3.2	10.7	0.7	
ISIS 559277	365	3.5	11.2	1.6	110
ISIS 561025	335	3.2	12.8	2.7	116
ISIS 561026	334	4.9	13.9	2.3	117
ISIS 561079	302	3.9	9.9	0.9	160
ISIS 561084	317	3.5	12.2	1.9	161
ISIS 561123	367	3.3	13.5	1.5	163
ISIS 561436	272	3.1	9.8	2.9	170

Example 15: Effect of ISIS antisense oligonucleotides targeting human ANGPTL3 in cynomolgus monkeys

Cynomolgus monkeys were treated with ISIS antisense oligonucleotides selected from studies described in the Examples above. Antisense oligonucleotide efficacy and tolerability, as well as their pharmacokinetic profile in the liver and kidney, were evaluated.

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 cross-reactive with the rhesus genomic sequence (GENBANK Accession No. NW_001108682.1 truncated from nucleotides 3049001 to 3062000, designated herein as SEQ ID NO: 3). 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: 3 is presented in the Table below. “Start site” indicates the 5’-most nucleotide to which the gapmer is targeted in the rhesus monkey gene sequence. ‘Mismatches’ indicates the number of nucleobases in the human oligonucleotide that are mismatched with the rhesus genomic sequence.

Table 94

Antisense oligonucleotides complementary to the rhesus ANGPTL3 genomic sequence (SEQ ID NO: 3)

ISIS No	Target Start Site	Mismatches	Chemistry	SEQ ID NO
563580	9315	2	5-10-5 MOE	77
560400	10052	1	5-10-5 MOE	35
567320	10232	1	5-10-5 MOE	93
567321	10234	1	5-10-5 MOE	94
544199	10653	0	5-10-5 MOE	20
567233	6834	2	5-10-5 MOE	90
561011	3220	1	Deoxy, MOE and (S)-cEt	114
559277	3265	0	Deoxy, MOE and (S)-cEt	110

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. Nine groups of 5 randomly assigned male cynomolgus monkeys each were injected subcutaneously with ISIS oligonucleotide or PBS at four sites on the back in a clockwise rotation (i.e. left, top, right, and bottom), one site per dose. The monkeys were given loading doses of PBS or 40 mg/kg of ISIS oligonucleotide every two days for the first week (days 1, 3, 5, and 7) and were subsequently dosed once a week for 12 weeks (days 14, 21, 28, 35, 42, 49, 56, 63, 70, 77, and 84) with PBS or 40 mg/kg of ISIS oligonucleotide.

During the study period, the monkeys were observed twice 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 example, one animal in the ISIS 567321 treatment group was found moribund on day 45 and was terminated. Scheduled euthanasia of the animals was conducted on day 86 (approximately 48 hours after the final dose) by exsanguination after ketamine/xylazine-induced anesthesia and administration of sodium pentobarbital. The protocols described in the Example were approved by the Institutional Animal Care and Use Committee (IACUC).

Hepatic Target Reduction*RNA analysis*

On day 86, RNA was extracted from liver for real-time PCR analysis of measurement of mRNA expression of ANGPTL3. Results are presented as percent change of mRNA, relative to PBS control, normalized with RIBOGREEN[®]. As shown in the Table below, treatment with ISIS antisense oligonucleotides resulted in significant reduction of ANGPTL3 mRNA in comparison to the PBS control. Analysis of ANGPTL3 mRNA levels revealed that ISIS 544199 and ISIS 559277, which are both fully cross-reactive with the rhesus sequence, significantly reduced expression levels. Other ISIS oligonucleotides, which targeted the monkey sequence with mismatches, were also able to reduce ANGPTL3 mRNA levels.

Table 95

Percent inhibition of ANGPTL3 mRNA in the cynomolgus monkey liver relative to the PBS control

ISIS No	%	SEQ ID NO
563580	62	77
560400	59	35
567320	67	93
567321	34	94
544199	88	20
561011	47	114
559277	85	110

Protein analysis

Approximately 1 mL of blood was collected from all available animals at day 85 and placed in tubes containing the potassium salt of EDTA. The blood samples were placed in ice and centrifuged (3000 rpm for 10 min at 4°C) to obtain plasma.

Human ANGPTL3 protein levels were quantified using a commercially available ELISA kit (Catalog #DANL30 by R&D Systems, Minneapolis, MN) with transgenic plasma samples diluted 1:20,000 using the manufacturer described protocol. The results are presented in the Table below. Analysis of plasman ANGPTL3 revealed that ISIS 563580, 544199 and ISIS 559277 reduced protein levels in a sustained manner. Other ISIS oligonucleotides were also able to reduce ANGPTL3 levels.

Table 96

Plasma protein levels (ng/mL) in the cynomolgus monkey

	Day 1	Day 3	Day 16	Day 30	Day 44	Day 58	Day 72	Day 86	SEQ ID NO
PBS	142	113	122	75	147	170	130	158	
ISIS 563580	113	99	102	46	109	93	82	81	77
ISIS 560400	92	107	145	63	170	182	157	178	35

ISIS 567320	87	72	94	56	176	181	134	166	93
ISIS 567321	80	84	98	62	156	116	122	112	94
ISIS 544199	114	84	50	34	66	56	81	71	20
ISIS 567233	115	111	174	134	162	125	122	109	90
ISIS 561011	89	92	111	106	104	100	140	129	114
ISIS 559277	86	62	63	54	77	64	68	70	110

Tolerability studies

Body weight measurements

To evaluate the effect of ISIS oligonucleotides on the overall health of the animals, body and weights were measured and are presented in the Table below. The results indicate that effect of treatment with antisense oligonucleotides on body weights was within the expected range for antisense oligonucleotides. Specifically, treatment with ISIS 563580 was well tolerated in terms of the body weights of the monkeys.

Table 97

Final body weights (g) in cynomolgus monkey

	Day 1	Day 14	Day 28	Day 35	Day 56	Day 70	Day 84	SEQ ID NO
PBS	2713	2709	2721	2712	2761	2754	2779	
ISIS 563580	2678	2669	2724	2699	2797	2798	2817	77
ISIS 560400	2713	2738	2808	2767	2867	2920	2976	35
ISIS 567320	2682	2707	2741	2731	2804	2830	2853	93
ISIS 567321	2672	2745	2849	2845	2995	2965	3002	94
ISIS 544199	2760	2813	2851	2897	2905	2888	2871	20
ISIS 567233	2657	2668	2650	2677	2907	2963	2903	90
ISIS 561011	2753	2797	2801	2811	2921	2967	2941	114
ISIS 559277	2681	2688	2701	2755	2826	2831	2965	110

Liver function

To evaluate the effect of ISIS oligonucleotides on hepatic function, blood samples were collected from all the study groups. The blood samples were collected from the cephalic, saphenous, or femoral veins, 48 hours post-dosing. The monkeys were fasted overnight prior to blood collection. Blood was collected in tubes without anticoagulant for serum separation. The tubes were kept at room temperature for a minimum of 90 minutes and then centrifuged (approximately 3,000 rpm for 10 min) 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 most of the antisense oligonucleotides had no effect on liver

function outside the expected range for antisense oligonucleotides. Specifically, treatment with ISIS 563580 was well tolerated in terms of the liver function in monkeys.

Table 98

ALT levels (IU/L) in cynomolgus monkey plasma

	Day 1	Day 30	Day 58	Day 86	SEQ ID NO
PBS	47	35	32	46	
ISIS 563580	56	55	55	83	77
ISIS 560400	50	35	47	68	35
ISIS 567320	72	44	51	106	93
ISIS 567321	53	39	44	75	94
ISIS 544199	58	49	51	51	20
ISIS 567233	42	38	47	64	90
ISIS 561011	48	35	34	43	114
ISIS 559277	49	45	53	60	110

5

Table 99

AST levels (IU/L) in cynomolgus monkey plasma

	Day 1	Day 30	Day 58	Day 86	SEQ ID NO
PBS	76	42	39	60	
ISIS 563580	75	56	42	81	77
ISIS 560400	85	63	59	99	35
ISIS 567320	104	64	55	153	93
ISIS 567321	83	47	45	66	94
ISIS 544199	68	68	70	91	20
ISIS 567233	46	80	66	86	90
ISIS 561011	48	39	41	51	114
ISIS 559277	50	56	55	77	110

Table 100

Bilirubin levels (mg/dL) in cynomolgus monkey plasma

	Day 1	Day 30	Day 58	Day 86	SEQ ID NO
PBS	0.31	0.24	0.20	0.19	
ISIS 563580	0.34	0.23	0.17	0.18	77
ISIS 560400	0.29	0.19	0.14	0.13	35
ISIS 567320	0.38	0.24	0.16	0.19	93
ISIS 567321	0.35	0.20	0.16	0.17	94

10

ISIS 544199	0.23	0.16	0.17	0.15	20
ISIS 567233	0.26	0.17	0.15	0.12	90
ISIS 561011	0.20	0.13	0.16	0.13	114
ISIS 559277	0.22	0.15	0.16	0.15	110

Kidney function

To evaluate the effect of ISIS oligonucleotides on kidney function, blood samples were collected from all the study groups. The blood samples were collected from the cephalic, saphenous, or femoral veins, 48 hours post-dosing. The monkeys were fasted overnight prior to blood collection. Blood was collected in tubes without anticoagulant for serum separation. The tubes were kept at room temperature for a minimum of 90 minutes and then centrifuged (approximately 3,000 rpm for 10 min) to obtain serum. Levels of BUN and creatinine were measured using a Toshiba 200FR NEO chemistry analyzer (Toshiba Co., Japan). Results are presented in the Table below, expressed in mg/dL.

The plasma chemistry data indicate that most of the ISIS oligonucleotides did not have any effect on the kidney function outside the expected range for antisense oligonucleotides. Specifically, treatment with ISIS 563580 was well tolerated in terms of the kidney function of the monkeys.

Table 101

Plasma BUN levels (mg/dL) in cynomolgus monkeys

	Day 1	Day 30	Day 58	Day 86	SEQ ID NO
PBS	28	28	27	29	
ISIS 563580	27	27	25	27	77
ISIS 560400	25	24	21	27	35
ISIS 567320	27	28	26	32	93
ISIS 567321	25	24	23	24	94
ISIS 544199	23	25	24	23	20
ISIS 567233	23	32	30	29	90
ISIS 561011	25	24	23	24	114
ISIS 559277	26	28	24	26	110

Table 102

Plasma creatinine levels (mg/dL) in cynomolgus monkeys

	Day 1	Day 30	Day 58	Day 86	SEQ ID NO
PBS	0.96	0.95	0.89	0.88	
ISIS 563580	0.97	1.04	0.88	0.85	77
ISIS 560400	0.99	1.00	0.93	0.91	35

ISIS 567320	0.95	0.94	0.89	0.87	93
ISIS 567321	0.97	0.94	0.89	0.87	94
ISIS 544199	0.86	0.87	0.88	0.87	20
ISIS 567233	0.89	1.08	1.06	1.00	90
ISIS 561011	0.93	0.93	0.91	0.90	114
ISIS 559277	0.86	0.91	0.87	0.91	110

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 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, individual white blood cell counts, such as that of monocytes, neutrophils, lymphocytes, as well as for platelet count, hemoglobin content and hematocrit, using an ADVIA120 hematology analyzer (Bayer, USA). The data is presented in the Tables below.

The data indicate the oligonucleotides did not cause any changes in hematologic parameters outside the expected range for antisense oligonucleotides at this dose. Specifically, treatment with ISIS 563580 was well tolerated in terms of the hematologic parameters of the monkeys.

Table 103

Blood cell counts in cynomolgus monkeys

	RBC (x 10 ⁶ /μL)	Platelets (x 10 ³ /μL)	WBC (x 10 ³ /μL)	Neutrophils (% WBC)	Lymphocytes (% total)	Monocytes (% total)	SEQ ID NO
PBS	5.6	462	12.2	58	39	2	
ISIS 563580	5.5	394	10.7	52	44	2	77
ISIS 560400	5.7	269	10.2	44	50	3	35
ISIS 567320	5.1	329	9.1	51	44	3	93
ISIS 567321	5.3	363	8.9	60	36	2	94
ISIS 544199	5.6	316	9.7	34	61	3	20
ISIS 567233	5.0	298	12.1	40	53	4	90
ISIS 561011	5.5	356	10.2	33	62	3	114
ISIS 559277	5.1	343	8.3	45	49	3	110

15

Table 104

Hematologic parameters in cynomolgus monkeys

	Hemoglobin (g/dL)	HCT (%)	SEQ ID NO
PBS	13	43	
ISIS 563580	12	40	77

ISIS 560400	12	41	35
ISIS 567320	11	38	93
ISIS 567321	12	41	94
ISIS 544199	13	44	20
ISIS 567233	11	38	90
ISIS 561011	13	42	114
ISIS 559277	12	40	110

Effect on pro-inflammatory molecules

To evaluate any inflammatory effect of ISIS oligonucleotides in cynomolgus monkeys, blood samples were taken for analysis of C-reactive protein and C3 levels on day 84 pre-dose. 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) and complement C3, which serve as markers of inflammation, were measured using a Toshiba 200FR NEO chemistry analyzer (Toshiba Co., Japan). The results indicate that treatment with ISIS 563580 was tolerable in monkeys.

10

Table 105

C-reactive protein levels (mg/L) in cynomolgus monkey plasma

	Day 1	Day 30	Day 58	Day 86	SEQ ID NO
PBS	3.1	5.5	2.7	4.1	
ISIS 563580	2.4	2.4	4.5	3.9	77
ISIS 560400	3.4	7.5	9.2	14.4	35
ISIS 567320	2.5	1.7	2.5	4.3	93
ISIS 567321	3.7	3.1	5.5	7.0	94
ISIS 544199	1.2	1.5	8.8	8.1	20
ISIS 567233	1.9	12.0	6.8	6.6	90
ISIS 561011	1.7	1.2	2.1	3.7	114
ISIS 559277	1.8	2.1	10.9	5.2	110

Table 106

C3 levels (mg/dL) in cynomolgus monkey plasma

	Pre-dose	Day 84	SEQ ID NO
PBS	122	117	
ISIS 563580	116	84	77
ISIS 560400	120	105	35
ISIS 567320	114	100	93

ISIS 567321	106	93	94
ISIS 544199	113	66	20
ISIS 567233	113	63	90
ISIS 561011	115	79	114
ISIS 559277	119	87	110

Measurement of oligonucleotide concentration

The concentration of the full-length oligonucleotide 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, GCGTTTGCTCTTCTTCTTGCGTTTTTT, designated herein as SEQ ID NO: 13) was added prior to extraction. Tissue sample concentrations were calculated using calibration curves, with a lower limit of quantitation (LLOQ) of approximately 1.14 µg/g. The results are presented in the Table below, expressed as µg/g liver or kidney tissue. The ratio of full-length oligonucleotide concentrations in the kidney versus the liver was calculated. The ratio of full-length oligonucleotide concentrations in the kidney versus the liver after treatment with ISIS 563580 was found to be most optimal compared to other compounds assessed.

Table 107

Oligonucleotide full length concentration

ISIS No	Kidney	Liver	Kidney/Liver ratio	SEQ ID NO
563580	1822	1039	1.8	77
560400	3807	1375	2.8	35
567320	2547	569	4.5	93
567321	2113	463	4.6	94
544199	1547	561	2.8	20
561011	2027	477	4.3	114
559277	2201	508	4.3	110

Example 16: ISIS 563580 Clinical Trial

To assess the effect of Study Drug ISIS 563580 a Phase 1, blinded, randomized, placebo-controlled, dose-escalation study was performed on healthy volunteers with a normal lipid profile. The study evaluated the safety, tolerability, pharmacokinetics, effect on plasma levels of ANGPTL3 and the lipoprotein profile of the study subjects after single and multiple doses of the Study Drug ISIS 563580.

The study population was healthy males or females aged 18-65 inclusive. Exclusion criteria included clinically significant abnormalities in the medical history and in the screening laboratory values of any of the

subjects. Subjects were randomized 3:1 to receive ISIS 563580 or placebo within each single-dose and multiple-dose cohort. Subjects were administered the study drug or placebo subcutaneously (SC).

Blood and urine samples were collected regularly throughout the study for safety, pharmacokinetic, and pharmacodynamics analyses. The safety and tolerability of ISIS 563580 was assessed by determining the incidence, severity, and dose-relationship of adverse events, vital signs, and clinical laboratory parameters. Safety results in subjects dosed with ISIS 563580 were compared with those in subjects dosed with placebo.

The most frequent adverse events were mild, local reactions at the injection site. Treatment with ISIS 563580 was generally well-tolerated and demonstrated an acceptable safety profile.

10 *Study Drug and Treatment for single dose study*

A solution of the Study Drug ISIS 563580 (200 mg/mL, 1.0 mL) contained in 2-mL stoppered glass vials was provided. Vials were for single-use only. ISIS 563580 solution and placebo are prepared by a pharmacist (or qualified delegate). A trained professional administered a single dose of the Study Drug in the abdomen, thigh, or outer area of the upper arm on each dosing day. A total of 16 subjects were enrolled in this study. The study design is presented in the Table below:

Table 108: Study design for single dose study with ISIS 563580

Cohort	Dose (mg)
A	50
B	100
C	200
D	400

Subcutaneous injection volumes were 0.25, 0.5, 1.0, and 2.0 mL for cohorts A, B, C, and D, respectively, with the 2.0 mL volume given as 2 non-contiguous 1.0 mL injections. For cohort A, a period of at least 24 hours was required between administering the Study Drug to the first 2 subjects and the remaining 2 subjects in the cohort. Dose escalation proceeded when the subjects in the preceding single-dose cohort had completed dosing and Day 4 safety evaluations demonstrated an acceptable safety profile.

The length of each subject's participation was approximately 8 weeks, including a 4-week screening period, a single dose, and a 4-week post-treatment evaluation period. Subjects had follow-up visits at the

Study Center on Days 2, 4, 8, 15, and a telephone contact on Day 30. The effect of the study drug is being assessed.

Study Drug and Treatment for multiple dose study

A solution of the Study Drug ISIS 563580 (200 mg/mL, 1.0 mL) contained in stoppered glass vials was provided. Vials were for single-use only. ISIS 563580 solution and placebo are prepared by a pharmacist (or qualified delegate). A trained professional administered the Study Drug in the abdomen, thigh, or outer area of the upper arm on each dosing day. A total of 32 subjects were enrolled in this study. The study design is presented in the Table below

Table 109: Study design for multiple dose study with ISIS 563580

Cohort	Dose amount	Amount of drug per dose	Total Dose (mg)
AA	8	100	800
BB	8	200	1600
CC	8	300	2400
DD	8	400	3200

Subcutaneous injection volumes were 0.5, 1.0, 1.5, and 2.0 mL for cohorts AA, BB, CC, and DD, respectively, with the 2.0 mL volume given as 2 non-contiguous 1.0 mL injections. Dosing of the first cohort (Cohort AA) began after at least 4 subjects in the single-dose cohort (Cohort D) had completed dosing and Day 4 safety evaluations demonstrated an acceptable safety profile. Subjects received 3 subcutaneous doses of the Study Drug during week 1 on alternative days (Days 1, 3, and 5) followed by once-weekly SC administrations during the next 5 weeks (Days 8, 15, 22, 29, and 36) for a total of 8 doses.

The length of each subject's participation was approximately 5.5 months, including a 4-week screening period, a 6-week treatment period, and a 13-week post-treatment evaluation period. Subjects had follow-up visits at the Study Center on Days 37, 43, 50, 64, 78, 92, 106, and 127. The results of the lipid profiles of the subjects on Day 36 are presented in the Table below. The asterisks indicate statistically significant changes ($p < 0.05$).

Treatment with ISIS 563580 generally produced dose-dependent reductions in plasma ANGPTL3, triglycerides, LDL-cholesterol, non-HDL cholesterol, VLDL-cholesterol, total cholesterol, ApoB and ApoC-III at day 36. In general, the magnitude of the reductions was associated with baseline lipid levels, with larger reductions observed in subjects with higher baselines.

Table 110: Mean % change compared the baseline at Day 36

Parameter	Placebo (N=8)	100 mg ISIS 563580 (N=6)	200 mg ISIS 563580 (N=6)	300 mg ISIS 563580 (N=6)	400 mg ISIS 563580 (N=6)
ANGPTL3	-8.1	-17.3	-51.6**	-62.4**	-81.9**
LDL-cholesterol	-0.9	-5.3	-12.8	-20.4*	-22.4**
Triglycerides	-15.3	-23.8	-21.0	-49.1**	-44.3**
Non-HDL- cholesterol	-3.2	-9.0	-15.4*	-25.4**	-25.6**
VLDL-cholesterol	-15.2	-25.1	-21.5	-50.0**	-43.5*
Total cholesterol	0.4	-2.1	-12.4*	-22.4**	-28.1**
ApoB	-5.1	-8.2	-8.8	-27.1*	-12.5
ApoC-III	20.6	8.2	-20.3	-51.3**	-65.0**

* P<0.05; ** p<0.01

CLAIMS

1. A compound comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and comprising a nucleobase sequence comprising a portion of at least 8 contiguous nucleobases complementary to an equal length portion of nucleobases 1140 to 1159 of SEQ ID NO: 1, wherein the nucleobase sequence of the modified oligonucleotide is at least 80% complementary to SEQ ID NO: 1.
2. The compound of claim 1, wherein the modified oligonucleotide consists of 15 to 30, 18 to 24, 19 to 22, 13 to 25, 14 to 25, 15 to 25, 16 or 20 linked nucleosides.
3. The compound of claim 1, wherein the modified oligonucleotide comprises a nucleobase sequence comprising a portion of at least 10, at least 12, at least 14, at least 16, at least 18, at least 19, or at least 20 contiguous nucleobases complementary to an equal length portion of SEQ ID NO: 1
4. A compound comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and comprising a nucleobase sequence comprising a portion of 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 complementary to an equal length portion of nucleobases 1140 to 1159 of SEQ ID NO: 1, wherein the nucleobase sequence of the modified oligonucleotide is at least 80% complementary to SEQ ID NO: 1.
5. The compound of any preceding claim, wherein 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.
6. A compound comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and having 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: 77.
7. A compound comprising a modified oligonucleotide consisting of 12 to 30 linked nucleosides and having a nucleobase sequence comprising at least 8, least 9, least 10, least 11, at least 12, least 13, at least 14, at least 15, or at least 16 contiguous nucleobases of any of the nucleobase sequences of SEQ ID NO: 77, 20, 110.
8. The compound of any preceding claim, wherein the modified oligonucleotide is single-stranded.

9. The compound of any preceding claim, wherein at least one internucleoside linkage is a modified internucleoside linkage.
10. The compound of claim 9, wherein each internucleoside linkage is a phosphorothioate internucleoside linkage.
11. The compound of any preceding claim, wherein the modified oligonucleotide comprises at least one modified sugar.
12. The compound of claim 11, wherein at least one modified sugar is a bicyclic sugar.
13. The compound of claim 11, wherein 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.
14. The compound of any preceding claim, wherein at least one nucleoside comprises a modified nucleobase.
15. The compound of claim 14, wherein the modified nucleobase is a 5-methylcytosine.
16. The compound of any preceding claim, wherein the modified oligonucleotide consists of 12 to 30 linked nucleosides and comprises:
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.
17. The compound of claim 16, wherein the modified oligonucleotide consists of 20 linked nucleosides and 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 is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

18. The compound of claim 16, wherein the modified oligonucleotide consists of 20 linked nucleosides.

19. A compound comprising a modified oligonucleotide consisting of 20 linked nucleosides and having a nucleobase sequence comprising at least 8 contiguous nucleobases complementary to an equal length portion of any of SEQ ID NO: 77, 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 is a phosphorothioate linkage and wherein each cytosine residue is a 5-methylcytosine.

20. A modified oligonucleotide according to the following formula: Ges Ges Aes mCes Aes Tds Tds Gds mCds mCds Ads Gds Tds Ads Ads Tes mCes Ges mCes Ae; wherein,

A = an adenine,

mC = a 5'-methylcytosine

G = a guanine,

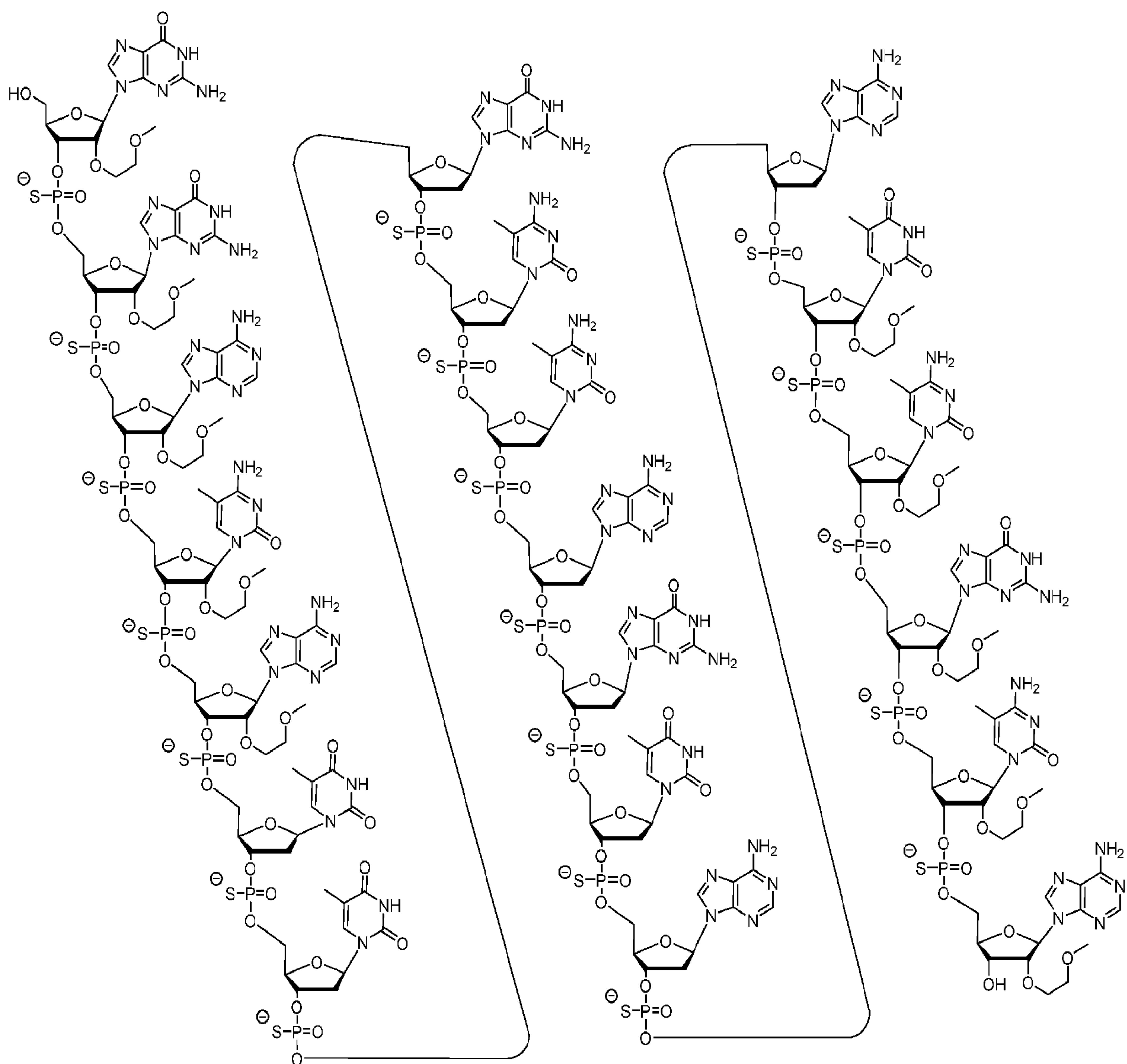
T = a thymine,

e = a 2'-O-methoxyethyl modified nucleoside,

d = a 2'-deoxynucleoside, and

s = a phosphorothioate internucleoside linkage.

21. A modified oligonucleotide according to the following formula:



22. A composition comprising a compound or modified oligonucleotide according to any preceding claim, or a salt thereof, and a pharmaceutically acceptable carrier or diluent.
23. A composition comprising a compound or modified oligonucleotide according to any preceding claim, for use in therapy.
24. The compound or modified oligonucleotide of claim 19, 20 or 21, for use in treating, preventing, or slowing progression of a disease related to elevated ANGPTL3.
25. The compound or modified oligonucleotide of claim 19, 20 or 21, wherein the disease is a cardiovascular and/or metabolic disease, disorder or condition.