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(54) Title: POLYPEPTIDES BINDING TO HUMAN COMPLEMENT C5

(57) **Abrégé/Abstract:**

The present invention relates to C5 binding polypeptides, comprising a C5 binding motif, BM, which motif consists of an amino acid sequence selected from i) EX₂X₃X₄A X₆X₇EID X₁₁LPNL X₁₆X₁₇X₁₈QW X₂₁AFIX₂₅X₂₆LX₂₈D, and ii) an amino acid sequence which has at least 86 % identity to the sequence defined in i), wherein the polypeptide binds to C5. The present invention moreover relates to C5 binding polypeptides for use in therapy, such as for use in treatment of a C5 related condition, and to methods of treatments.

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POLYPEPTIDES BINDING TO HUMAN COMPLEMENT C5

Technical field

The present disclosure relates to polypeptides that bind to human complement component 5
5 (C5) and to the use of such polypeptides in therapy.

Background

The complement protein C5 is a central component of the complement system; a key part of the innate immune system. The complement system is an intricate immune survival system with
10 numerous tasks in tightly controlled, diverse processes. One of its functions is as first line host defense against infection by other organisms by discriminating healthy host tissues from cellular debris and apoptotic and necrotic cells. Furthermore, it is involved in clearance of immune complexes, regulation of the adaptive immune response, promotion of tissue regeneration, angiogenesis, mobilization of stem cells and development of the central nervous system
15 (Woodruff *et al.* Mol Immunol 2011, 48 (14):1631-1642); Ricklin *et al.* Nat Immunol 2010, 11(9):785-795). Any trigger, for example erroneous or unrestricted activation or insufficient regulation, that disturbs the fine balance of complement activation and regulation may lead to pathologic conditions including self-attack of the host's cells leading to extensive tissue damage.

20 The complement system consists of about 30 proteins. There are three pathways to initiate complement immunity; the classical pathway that employs C1q to recognize immune complexes on the surface of cells, the lectin pathway that is initiated when mannose-binding lectin (MBL) recognizes certain sugars and the alternative pathway that is initiated spontaneously by hydrolysis of complement factor 3 (C3), a process suppressed by certain
25 mammalian cell surface molecules not present on invading pathogens. The alternative pathway also acts as an amplification loop for the complement system. All three pathways converge at the level of C3. Cleavage of C3 into C3a and C3b leads to the formation of a convertase that in turn cleaves complement factor 5 (C5) into C5a and C5b. C5a is a very potent attractant of various immune cells while C5b oligomerizes with C6-9 to form a pore known as the membrane
30 attack complex (MAC) or sometimes the terminal complement complex (TCC). Activation of the complement system leads to a number of mechanisms with the purpose of neutralizing the pathogen; formation of MAC on the surface of a cell such as an invading bacteria lead to lysis, deposition of C3 and C4 cleavage products C3b and C4b aids opsonization leading to phagocytosis of the pathogen by macrophages and anaphylatoxins such as C3a and C5a attracts
35 monocytes and neutrophils to the site of activation, upregulates surface markers leading to increased immunologic susceptibility and to the release of cytokines.

C5 is a 190-kDa glycoprotein comprised of 2 disulfide-linked polypeptide chains, alpha and beta, with a molecular mass of 115 and 75 kDa, respectively (Tack *et al.* Biochem 1979, 18:1490-1497). Haviland *et al.* (J Immun 1991, 146: 362-368) constructed the complete cDNA sequence of human complement pro-C5, which is predicted to encode a 1,676-amino acid pro-molecule that contains an 18-amino acid leader peptide and a 4-amino acid linker separating the beta and alpha chains. Blockade of C5 cleavage into C5a and C5b prevents MAC formation and formation of the pro-inflammatory C5a but leaves the upstream complement effector system intact allowing the C3/C4 mediated opsonization.

The complement system's key role in the defense against pathogens in general makes it an interesting target for pharmaceutical intervention. This is emphasized by the fact that many mutations or impaired regulation of complement is involved in various diseases and conditions. These include increased susceptibility to auto-immune diseases such as systemic lupus erythematosus (SLE) where deposition of immune complexes triggers the classical pathway (Manderson *et al.* Annu Rev Immunol 2004, 22:431-456). In addition, mutations of the complement proteins C1-C5 often result in SLE or SLE like symptoms. Other autoimmune diseases with a strong involvement of the complement system are rheumatoid arthritis (RA) where immune complexes may activate complement in the RA joint, Sjögren's syndrome, dermatomyositis and other autoantibody driven diseases such as Guillain-Barré syndrome (GBS), Fisher syndrome (Kaida *et al.* J. Neuroimmun 2010, 223:5-12) different types of vasculitis, systemic sclerosis, anti-glomerular basement membrane (anti-GBM) and anti-phospholipid syndrome (APS) (Chen *et al.* J Autoimmun 2010, 34:J276-J286).

The complement system is furthermore involved in neurodegenerative disorders such as Alzheimer's disease (AD) where A β plaques directly activate the complement system leading to C5a mediated recruitment of microglia. This was further confirmed when a C5aR antagonist was shown to be neuroprotective in a mouse model of AD (Fonseca *et al.* J Immunol 2009, 183:1375-1383). Auto-antibodies against the acetylcholine receptor and subsequent complement activation is the most common cause to myasthenia gravis, a disease that affects the neuromuscular junction (Toyka and Gold, Schweizer Archive Neurol Psych 2007, 158:309-321). MAC formation is involved in the pathophysiology of multiple sclerosis (MS) (Oh *et al.* Immunol Res 2008, 40:224-234). Also in Parkinson's disease, Huntington's disease and prion diseases such as Creutzfeld-Jacob disease, complement activation is a part of the pathology (Bonifati and Kishore, Mol Immunol 2007, 44:999-1010). In wound healing, inflammatory responses are a key component to restore tissue homeostasis and the complement system is involved in the early recognition of damaged tissue. However, in models of chronic wounds and severe burns, for example, inhibition of complement by e. g. C1 inhibitor resulted in improved healing and decreased tissue damage suggesting that complement. Furthermore, various complement deficiencies, such as exemplified by the C4 knockout mouse, have been found to

be protective against long-term tissue damage resulting from wounds (reviewed in Cazender *et al.* Clinical and Developmental Immunology 2012, on-line publication). Lately it has been shown that tumor growth and proliferation is facilitated by complement activation, in particular by C5a, and that blockade of the C5a receptor slows down this process. In addition, mice
5 lacking C3 display significantly slower tumor growth than wild-type littermates (Markiewski *et al.* Nat Immunol 2008, 9:1225-1235).

Dysfunctional complement regulation is the cause of several rare to ultra-rare conditions, such as paroxysmal nocturnal hemoglobinuria (PNH) and atypical hemolytic uremic syndrome (aHUS), where hemolysis is a key feature in the pathology. In PNH, a clone of
10 hematopoietic stem cells with mutated PIG-A gene encoding phosphatidylinositol N-acetylglucosaminyltransferase subunit A take over the pool of blood cells. This mutation leads to loss of GPI anchored proteins such as the complement regulators CD55 and CD59. Red blood cells lacking CD55 and CD59 on the surface are exposed for complement mediated lysis by MAC. Clinically, PNH is manifested by hemolysis leading to anemia, thrombosis and bone
15 marrow failure. Atypical HUS is caused by mutations in regulatory proteins of mainly the alternative pathway, such as by mutations in factor H.

The eye is strongly indicated as a site for complement driven pathology. The most common cause of visual loss is age-related macular degeneration (AMD) where, in its more severe form (exudative or wet AMD), pathologic choroidal neurovascular membranes develop
20 under the retina. In the US, about 10 % of the population aged 65-74 shows sign of macular degeneration and as many as 5 % have visual impairment as a result to AMD. These numbers increase dramatically with age, but there are also genetic factors. Among the genes strongest associated with AMD are complement factor H, factor B and C3 and the C1 inhibitor (Bradley *et al.* Eye 2011, 25:683-693). Furthermore, several studies and clinical trials using various
25 complement blocking molecules have proven beneficial, suggesting that a C5 blocking molecule could help these patient groups. However, the current treatments of advanced AMD aims at inhibition of vascular endothelial growth factor (VEGF) induced vascularization by intravitreal injections of e.g. Ranibizumab (a monoclonal antibody fragment) and Bevacizumab (monoclonal antibody). In animal models of uveitis, inflammation of the eye due to immune
30 responses to ocular antigens, blocking antibodies against alternative pathway factor B (Manickam *et al.* J Biol Chem 2011, 286:8472-8480) as well as against C5 (Copland *et al.* Clin Exp Immunol 2009, 159:303-314), improved the disease state.

In transplantation of solid organs, there are two major mechanistic pathways leading to rejection or delayed/impaired function of the graft: 1) the immunologic barriers between donor
35 and recipient with respect to blood group (ABO) and MHC classes as well as extent of pre-sensitization of the recipient against the donor, i.e. occurrence of donor specific antibodies (DSA) leading to acute antibody mediated rejection (AMR); and 2) the condition of the

transplanted organ as well as the period of time it has been kept without constant blood perfusion, i.e. the degree of ischemic damage or ischemia reperfusion injury (IRI) of the graft. In both AMR and IRI, the complement system is attacking the organ recognized as foreign and, therefore, an entity that should be rejected. In AMR, the pre-existing anti-donor antibodies rapidly form immune complexes on the surface of the foreign organ leading to recognition by C1q and subsequent activation of the complement system via the classical pathway. This process, known as hyper-acute rejection happens within minutes and, therefore modern transplantation of mismatched organs includes elimination of DSA prior to transplantation by plasmapheresis or plasma exchange and intravenous IgG combined with different immunosuppressants. Novel treatments also include B-cell depletion via usage of the anti-CD20 antibody Rituximab (Genberg *et al.* Transplant 2008, 85:1745-1754). These protocols have vastly eliminated the occurrence of hyper-acute rejection but still, in highly sensitized patients, the incidence of acute AMR (weeks-months) is as high as 40 % (Burns *et al.* Am J Transplant 2008, 6:2684-2694; Stegall *et al.* Am J Transplant 2011, early on-line publication). With respect to IRI, most evidence points at the terminal pathway with subsequent MAC formation and lysis as the main cause of tissue damage. Thus, a C5 blocking polypeptide would be protective against rejection regardless of the cause being AMR, IRI or, as often happens, a combination of both AMR and IRI. As expected, highly perfused organs, such as the liver (Qin *et al.* Cell Mol Immunol 2006, 3:333-340), the heart and the kidneys are particularly susceptible to complement mediated damage.

The central placement of the C5 protein; connecting the proximal and the terminal parts of the complement cascade, makes it an attractive target for pharmaceutical intervention. Since C5 is common to all pathways of complement activation, blocking of C5 will stop the progression of the cascade regardless of the stimuli and thereby prevent the deleterious properties of terminal complement activation while leaving the immunoprotective and immunoregulatory functions of the proximal complement cascade intact.

Antibodies targeted to human complement C5 are known from, e.g., WO 95/29697; WO 02/30985; and WO 2004/007553. Eculizumab (Soliris™) is a humanized monoclonal antibody directed against protein C5 and prevents cleavage of C5 into C5a and C5b. Eculizumab has been shown to be effective in treating PNH, a rare and sometimes life threatening disease of the blood characterized by intravascular hemolytic anemia, thrombophilia and bone marrow failure, and is approved for this indication. Eculizumab was also recently approved by the FDA for treatment of atypical hemolytic syndrome (aHUS), a rare but life threatening disease caused by loss of control of the alternative complement pathway leading to over-activation manifested as thrombotic microangiopathy (TMA) leading to constant risk of damage to vital organs such as kidney, heart and the brain. In aHUS, transplantation of the damaged organ only temporarily helps the patient as the liver continues to produce the mutated

form of controlling protein (most often complement factor H or other proteins of the alternative pathway). A related disease with a transient acute pathophysiology is HUS caused by infection of Shiga toxin positive *E. coli* (STEC-HUS) and there are promising clinical data suggesting efficacy also for this condition (Lapeyraque *et al.*, N Engl J Med 2011, 364:2561-2563). Finally, the C5 blocking antibody Eculizumab has proven efficacious in preventing AMR in recipients of highly mismatched kidneys (Stegall, M. D. *et al.*, Am J Transplant 2011, 11:2405-2413).

Apart from full length antibodies, single-chain variable fragments (scFV), minibodies and aptamers targeting C5 are described in literature. These C5 inhibitors may bind to different sites (epitopes) on the C5 molecule and may have different modes of action. For example, whereas Eculizumab interacts with C5 at some distance of the convertase cleavage site, the minibody Mubodina® interacts with the cleavage site of C5. The C5 inhibitory protein *Ornithodoros moubata* Complement Inhibitor (OmCI, Nunn, M. A. *et al.*, J Immunol 2005, 174:2084-2091) from soft tick *Ornithodoros moubata* has been hypothesized to bind to the distal end of the CUB-C5d-MG8 superdomain, which is close to the convertase cleavage site (Fredslund *et al.*, Nat Immunol 2008, 9 (7):753-760). In contrast to the three proteins mentioned above inhibiting cleavage of C5, the monoclonal antibody TNX-558 binds to a C5a epitope present both on intact C5 and released C5a without inhibiting the cleavage of C5. (Fung *et al.*, Clin Exp Immunol 2003, 133 (2):160-169).

Antibodies with their large, multidomain structure, 12 intra-chain and 4 inter-chain disulfide bridges and complex glycosylation patterns, have a number of intrinsic disadvantages related to their molecular structure. For example, the size of Eculizumab is about 148 kDa. The concentration of C5 in human blood is about 400 nM and in order to block C5 activity entirely, the concentration of the inhibitor must be at least equal or higher than that. Therefore, the standard life-long treatment regimen of PNH using Soliris™ is intravenous infusions of 900 mg protein every second week, a treatment that mainly take place in the clinic leading to great inconvenience to the patient and cost to the society. Soliris™ has also been reported to cause chest pain, fever, chills, itching, hives, flushing of the face, rash, dizziness, troubled breathing, or swelling of the face, tongue, and throat, although the reasons for these side effects are not clear. Furthermore, Eculizumab is not active in any tested animal model, including primates, making animal studies with the active drug impossible. As mentioned above, the current treatments of AMD are also antibody dependent and, thus, treatments based on injections or other routes of administration with molecules of lower molecular weight, are highly required.

In addition, antibody production is more difficult and more expensive than production of small proteins (Kenanova *et al.*, Expert Opin Drug Deliv 2006, 3 (1):53-70). Other drawbacks generally related to antibodies are listed by Reilly *et al.* (Clin Pharmacokinet 1995, 28:126-142), such as cross-reactivity and non-specific binding to normal tissues, increased metabolism

of injected antibodies and formation of human anti-human antibodies (HAMA) causing decreased or loss of the therapeutic effect.

Thus, continued provision of agents with comparable C5 blocking activity remains a matter of substantial interest within the field. In particular, there is a continued need for
 5 molecules that prevent the terminal complement cascade as well as the formation of the pro-inflammatory molecule C5a. Of great interest is also a provision of uses of such molecules in the treatment of disease.

Description

10 It is an object of the invention to provide new C5 binding agents. It is moreover an object of the invention to provide new C5 binding agents for use in therapeutic applications.

In one aspect, there is provided a C5 binding polypeptide, comprising a C5 binding motif, *BM*, which motif consists of the amino acid sequence selected from

15 i) $EX_2X_3X_4AX_6X_7EIDX_{11}LPNLX_{16}X_{17}X_{18}QWX_{21}AFIX_{25}X_{26}LX_{28}D$,

wherein, independently of each other,

X_2 is selected from H, Q, S, T and V;

X_3 is selected from I, L, M and V;

20 X_4 is selected from A, D, E, H, K, L, N, Q, R, S, T and Y;

X_6 is selected from N and W;

X_7 is selected from A, D, E, H, N, Q, R, S and T;

X_{11} is selected from A, E, G, H, K, L, Q, R, S, T and Y;

X_{16} is selected from N and T;

25 X_{17} is selected from I, L and V;

X_{18} is selected from A, D, E, H, K, N, Q, R, S and T;

X_{21} is selected from I, L and V;

X_{25} is selected from D, E, G, H, N, S and T;

X_{26} is selected from K and S;

30 X_{28} is selected from A, D, E, H, N, Q, S, T and Y;

and

ii) an amino acid sequence which has at least 86 % identity to the sequence defined in i), wherein the polypeptide binds to C5.

35 The above defined class of sequence related polypeptides having a binding affinity for C5 is derived from a common parent polypeptide sequence. More specifically, the definition of the class is based on an analysis of a large number of random polypeptide variants of the parent

polypeptide that were selected for their interaction with C5 in selection experiments. The identified C5 binding motif, or “*BM*”, corresponds to the target binding region of the parent scaffold, which region constitutes two alpha helices within a three-helical bundle protein domain. In the parent scaffold, the varied amino acid residues of the two *BM* helices constitute a binding surface for interaction with the constant Fc part of antibodies. By random variation of binding surface residues and subsequent selection of variants, the Fc interaction capacity of the binding surface has been replaced with a capacity for interaction with C5.

As accounted for in the following Examples, selection of C5 binding polypeptide variants may for example be achieved by phage display for selection of naïve variants of a protein scaffold optionally followed by affinity maturation and cell display for selection of affinity matured C5 binding variants. It is however understood that any selection system, whether phage-based, bacterial-based, cell-based or other, may be used for selection of C5 binding polypeptides.

The terms “C5 binding” and “binding affinity for C5” as used in this specification refers to a property of a polypeptide which may be tested for example by the use of surface plasmon resonance technology, such as in a Biacore instrument (GE Healthcare). C5 binding affinity may e.g. be tested in an experiment in which C5 is immobilized on a sensor chip of a Biacore instrument, and the sample containing the polypeptide to be tested is passed over the chip. Alternatively, the polypeptide to be tested is immobilized on a sensor chip of the instrument, and a sample containing C5, or fragment thereof, is passed over the chip. The skilled person may then interpret the results obtained by such experiments to establish at least a qualitative measure of the binding of the polypeptide to C5. If a quantitative measure is desired, for example to determine the apparent equilibrium dissociation constant K_D for the interaction, surface plasmon resonance methods may also be used. Binding values may for example be defined in a Biacore 2000 instrument (GE Healthcare). C5 is immobilized on a sensor chip of the measurement, and samples of the polypeptide whose affinity is to be determined are prepared by serial dilution and injected over the chip. K_D values may then be calculated from the results using for example the 1:1 Langmuir binding model of the BIAevaluation software provided by the instrument manufacturer. The C5 or fragment thereof used in the K_D determination may for example comprise the amino acid sequence represented by SEQ ID NO:760.

In one embodiment of the C5 binding polypeptide according to the present invention, the C5 binding polypeptide binds to C5 such that the K_D value of the interaction is at most 1×10^{-6} M, such as at most 1×10^{-7} M, 1×10^{-8} M, or 1×10^{-9} M.

A C5 binding polypeptide according to the present invention may be used as an alternative to conventional antibodies or low molecular weight substances in various medical, veterinary and diagnostic applications. In particular, the C5 binding polypeptide may be useful

in any method requiring affinity for C5 of a reagent. Accordingly, the C5 binding polypeptide may be used as a detection reagent, a capture reagent, a separation reagent, a diagnostic agent or a therapeutic agent in such methods.

As the skilled person will realize, the function of any polypeptide, such as the C5
5 binding capacity of the polypeptides as defined herein, is dependent on the tertiary structure of the polypeptide. It is therefore possible to make minor changes to the amino acid sequence of a polypeptide without largely affecting the tertiary structure and the function thereof. Thus, in one embodiment, the polypeptide comprises modified variants of the BM of i), which are such that the resulting sequence is at least 89% identical to a sequence belonging to the class defined by
10 i), such as at least 93 % identical, such as at least 96 % identical to a sequence belonging to the class defined by i). For example, it is possible that an amino acid residue belonging to a certain functional grouping of amino acid residues (e.g. hydrophobic, hydrophilic, polar etc) could be exchanged for another amino acid residue from the same functional group.

In another embodiment of the C5 binding polypeptide as defined above, the amino acid
15 sequence is selected from i) as defined above, and iii) an amino acid sequence which in the 13 variable positions as denoted by X_n , wherein n is 2-4, 6-7, 11, 16-18, 21, 25-26 and 28, has at least 84 % identity to the sequence defined in i), and which in positions 1, 5, 8-10, 12-15, 19-20, 22-24, 27 and 29 has at least 87 % identity to the sequence defined in i).

In one embodiment of the polypeptide according to the present invention, X_2 is selected
20 from H, T and V. In another embodiment, X_2 is selected from T and V. In yet another embodiment, X_2 is V.

In one embodiment of the polypeptide according to the present invention, X_3 is selected from I, L and V. In another embodiment, X_3 is selected from I and L. In yet another embodiment, X_3 is I. In an alternative embodiment, X_3 is L.

25 In one embodiment of the polypeptide according to the present invention, X_4 is selected from A, D, E, K, L, Q and R. In another embodiment, X_4 is selected from A, D, E, K and R. In yet another related embodiment, X_4 is selected from D and E.

In one embodiment of the polypeptide according to the present invention, X_6 is W.

In one embodiment of the polypeptide according to the present invention, X_7 is selected
30 from A, D, N and T. In another embodiment, X_7 is selected from D and N. In yet another related embodiment, X_7 is D. In an alternative embodiment, X_7 is N.

In one embodiment of the polypeptide according to the present invention, X_{11} is selected from A, H, K, Q, R and S. In another embodiment, X_{11} is selected from A, H, K and R. In yet another related embodiment, X_{11} is selected from A, K and R. In yet another related
35 embodiment, X_{11} is selected from K and R.

In one embodiment of the polypeptide according to the present invention, X_{16} is T.

In one embodiment of the polypeptide according to the present invention, X_{17} is selected from I and L. In another embodiment, X_{17} is I. In an alternative embodiment, X_{17} is L.

In one embodiment of the polypeptide according to the present invention, X_{18} is selected from A, D, E, N, Q, S and T. In another embodiment, X_{18} is selected from A, D, E, Q and S. In yet another related embodiment, X_{18} is selected from D, E and Q. In yet another related embodiment, X_{18} is selected from D and E. In yet another related embodiment, X_{18} is D. In an alternative embodiment, X_{18} is E.

In one embodiment of the polypeptide according to the present invention, X_{21} is selected from I and L. In another embodiment, X_{21} is I. In an alternative embodiment, X_{21} is L.

In one embodiment of the polypeptide according to the present invention, X_{25} is selected from E, H, N and T. In another embodiment, X_{25} is selected from E and N. In yet another related embodiment, X_{25} is N.

In one embodiment of the polypeptide according to the present invention, X_{26} is K.

In one embodiment of the polypeptide according to the present invention, X_{28} is selected from A, D, E, H, N, Q and S. In another embodiment of the above disclosed polypeptide, X_{28} is selected from A, D, E and S. In yet another related embodiment, X_{28} is selected from A, D and E. In yet another related embodiment, X_{28} is selected from D and E. In yet another related embodiment, X_{28} is D.

In one embodiment of the polypeptide according to the present invention, X_3X_4 is selected from LE and LD.

In one embodiment of the polypeptide according to the present invention, $X_{17}X_{18}$ is selected from IE and LD.

In the above embodiments of the first aspect, examples of C5 binding polypeptides falling within the class of polypeptides are identified. It is contemplated that the above individual embodiments may be combined in all conceivable ways and still fall within the scope of the present invention. Such combinations of individual embodiments define a restricted, in one or more of the positions X_2 - X_{28} , amino acid sequence as compared to the amino acid definition in i).

The above embodiments of a C5 binding polypeptide may for example be combined such that the amino acid i) fulfils at least four of the following eight conditions I-VIII:

- I. X_2 is V;
- II. X_3 is selected from I and L;
- III. X_6 is W;
- IV. X_7 is selected from D and N;
- V. X_{17} is selected from I and L;
- VI. X_{21} is L;
- VII. X_{25} is N;

VIII. X_{28} is D.

In some examples of a C5 binding polypeptide according to the first aspect, the amino acid sequence i) fulfils at least five of the eight conditions I-VIII. More specifically, the amino acid sequence i) may fulfill at least six of the eight conditions I-VIII, such at least seven of the eight conditions I-VIII, such as all of the eight conditions I-VIII.

As described in the following Examples, the selection of C5 binding variants has led to the identification of individual C5 binding motif (*BM*) sequences. These sequences constitute individual embodiments of C5 binding polypeptides according to this aspect. The sequences of individual C5 binding motifs are presented in Figure 1 and as SEQ ID NO:1-248. In some embodiments of this aspect, the *BM* sequence i) is selected from any one of SEQ ID NO:1-12, SEQ ID NO:20, SEQ ID NO:23-24, SEQ ID NO:26-28, SEQ ID NO:32-35, SEQ ID NO:38-39, SEQ ID NO:41, SEQ ID NO:46, SEQ ID NO:49, SEQ ID NO:56-57, SEQ ID NO:59, SEQ ID NO:66, SEQ ID NO:78-79, SEQ ID NO:87, SEQ ID NO:92, SEQ ID NO:106, SEQ ID NO:110, SEQ ID NO:119, SEQ ID NO:125, SEQ ID NO:141, SEQ ID NO:151, SEQ ID NO:161, SEQ ID NO:166, SEQ ID NO:187, SEQ ID NO:197, SEQ ID NO:203, SEQ ID NO:205, SEQ ID NO:215 and SEQ ID NO:243. More specifically, the *BM* sequence i) is selected from any one of SEQ ID NO:1-12, such as from SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4 and SEQ ID NO:5. In particular, the *BM* sequence i) may be selected from SEQ ID NO:1 and SEQ ID NO:4.

In particular embodiments, the C5 binding motif (*BM*) forms part of a three-helix bundle protein domain. For example, the *BM* may essentially constitute two alpha helices with an interconnecting loop, within said three-helix bundle protein domain.

The three-helix bundle protein domain is, in another embodiment, selected from domains of bacterial receptor proteins. Non-limiting examples of such domains are the five different three-helical domains of Protein A from *Staphylococcus aureus*, such as domain B, and derivatives thereof. In some embodiments, the three-helical bundle protein domain is a variant of protein Z, which is derived from said domain B of staphylococcal Protein A.

In embodiments where the C5 binding polypeptide of the invention forms part of a three-helix bundle protein domain, the C5 binding polypeptide may comprise an amino acid sequence selected from:

i) $K-[BM]-DPSQS X_a X_b LLX_c EAKKL NDX_d Q;$

wherein

[*BM*] is a C5 binding motif as defined above;

X_a is selected from A and S;

X_b is selected from N and E;

X_c is selected from A, S and C;

X_d is selected from A and S;

and

- ii) an amino acid sequence which has at least 79 % identity to any one of the sequences defined above. Said amino acid sequence may have at least 81 %, such as at least 83 %, such as at least 85 %, such as at least 87 %, such as at least 89 %, such as at least 91 %, such as at least 93 %, such as at least 95 %, such as at least 97 % identity to any one of the sequences defined above.

In one embodiment of the C5 binding polypeptide as defined above, X_a is A. In an alternative embodiment of the C5 binding polypeptide as defined above, X_a is S.

In one embodiment of the C5 binding polypeptide as defined above, X_b is N. In an alternative embodiment, X_b is E.

In one embodiment of the C5 binding polypeptide as defined above, X_c is A. In an alternative embodiment, X_c is S. In yet another alternative embodiment, X_c is C.

In one embodiment of the C5 binding polypeptide as defined above, X_d is A. In an alternative embodiment, X_d is S.

In one embodiment of the C5 binding polypeptide as defined above, X_a is A; X_b is N; X_c is A and X_d is A.

In a further embodiment of the C5 binding polypeptide as defined above, X_a is A; X_b is N; X_c is C and X_d is A.

In a further embodiment of the C5 binding polypeptide as defined above, X_a is S; X_b is E; X_c is S and X_d is S.

In a further embodiment of the C5 binding polypeptide as defined above, X_a is S; X_b is E; X_c is C and X_d is S.

In yet a further embodiment, the amino acid sequence of the C5 binding polypeptide as defined above is selected from SEQ ID NO:249-496, in particular from SEQ ID NO:249-260, SEQ ID NO:268, SEQ ID NO:271-272, SEQ ID NO:274-276, SEQ ID NO:280-283, SEQ ID NO:286-287, SEQ ID NO:289, SEQ ID NO:294, SEQ ID NO:297, SEQ ID NO:304-305, SEQ ID NO:307, SEQ ID NO:314, SEQ ID NO:326-327, SEQ ID NO:335, SEQ ID NO:340, SEQ ID NO:354, SEQ ID NO:358, SEQ ID NO:367, SEQ ID NO:373, SEQ ID NO:389, SEQ ID NO:399, SEQ ID NO:409, SEQ ID NO:414, SEQ ID NO:435, SEQ ID NO:445, SEQ ID NO:451, SEQ ID NO:453, SEQ ID NO:463 and SEQ ID NO:491, such as from SEQ ID NO:249-260. In a further embodiment, the amino acid sequence is selected from SEQ ID NO:249, SEQ ID NO:250, SEQ ID NO:251, SEQ ID NO:252 and SEQ ID NO:253, such as from SEQ ID NO:249 and SEQ ID NO:252.

Thus, in a further embodiment, there is provided a C5 binding polypeptide which comprises an amino acid sequence selected from:

i) YAK-[BM]-DPSQS SELLX_c EAKKL NDSQA P;

wherein [BM] is a C5 binding motif as defined above and

X_c is selected from S and C; and

ii) an amino acid sequence which has at least 81 % identity to any one of the sequences defined in i) above.

Alternatively, there is provided a C5 binding polypeptide which comprises an amino acid sequence selected from:

i) FNK-[BM]-DPSQS ANLLX_c EAKKL NDAQA P;

wherein [BM] is a C5 binding motif as defined above and

X_c is selected from A and C; and

ii) an amino acid sequence which has at least 81 % identity to any one of the sequences defined in i) above.

As discussed above, polypeptides comprising minor changes as compared to the above amino acid sequences without largely affecting the tertiary structure and the function thereof are also within the scope of the present application. Thus, in some embodiments, the C5 binding polypeptides as defined above may for example have a sequence which is at least 83 %, at least 84 %, at least 86 %, at least 88 %, at least 90 %, at least 92 %, at least 94 %, at least 96 % or at least 98 % identical to the sequence defined in i).

In some embodiments and as disclosed in the Examples below, the C5 binding motif may form part of a 58 or 60 amino acid polypeptide. Such a polypeptide may e.g. comprise a sequence selected from any one of SEQ ID NO:497-757, in particular a sequence selected from any one of SEQ ID NO:497-508, SEQ ID NO:516, SEQ ID NO:519-520, SEQ ID NO:522-524, SEQ ID NO:528-531, SEQ ID NO:534-535, SEQ ID NO:537, SEQ ID NO:542, SEQ ID NO:545, SEQ ID NO:552-553, SEQ ID NO:555, SEQ ID NO:562, SEQ ID NO:574-575, SEQ ID NO:583, SEQ ID NO:588, SEQ ID NO:602, SEQ ID NO:606, SEQ ID NO:615, SEQ ID NO:621, SEQ ID NO:637, SEQ ID NO:647, SEQ ID NO:657, SEQ ID NO:662, SEQ ID NO:683, SEQ ID NO:693, SEQ ID NO:699, SEQ ID NO:701, SEQ ID NO:711, SEQ ID NO:739 and SEQ ID NO:745-757, such as a sequence selected from SEQ ID NO:497-508 and SEQ ID NO:745-757. In another embodiment, the amino acid sequence is selected from SEQ ID NO:497, SEQ ID NO:498, SEQ ID NO:499, SEQ ID NO:500, SEQ ID NO:501, SEQ ID NO:746, SEQ ID NO:747, SEQ ID NO:748, SEQ ID NO:750 and SEQ ID NO:753, such as from any one of SEQ ID NO:497, SEQ ID NO:500, SEQ ID NO:748 and SEQ ID NO:753.

Binding of a molecule to C5 does not necessarily inhibit cleavage of C5. Inhibition is dependent on binding site, and since it is not entirely clear what effects interaction with specific regions of C5 have, some C5 binding molecules may interact with C5 without inhibiting its cleavage into C5a and C5b. In one embodiment of the present invention, the C5 binding polypeptide, when e.g. administered to a mammalian subject, inhibits cleavage of C5. The C5

binding polypeptide according to the invention may more specifically inhibit cleavage of human C5 when administered to a human subject.

The structure of C5 differs somewhat between species, and thus, a C5 binder found to bind C5 of one species may be inactive in another species. The humanized antibody

5 Eculizumab, for example, binds to a domain of C5 often referred to as MG7. This region is highly variable between species and, therefore, Eculizumab binding is restricted to human C5. The C5 binding polypeptide of the present invention, however, is not restricted to human C5 but exhibits activity in animal models as well, as demonstrated in the appended Examples.

The skilled person will understand that various modifications and/or additions can be
10 made to a C5 binding polypeptide according to any aspect disclosed herein in order to tailor the polypeptide to a specific application without departing from the scope of the present invention. For example, a C5 binding polypeptide according to any aspect may comprise further C terminal and/or N terminal amino acids. Such a polypeptide should be comprehended as a polypeptide having additional amino acids residues at the very first and/or the very last position
15 in the polypeptide chain, i.e. at the N- and/or C-terminus. Thus, a C5 binding polypeptide may comprise any suitable number of additional amino acid residues, for example at least one additional amino acid residue. Each additional amino acid residue may individually or collectively be added in order to, for example, improve production, purification, stabilization *in vivo* or *in vitro*, coupling, or detection of the polypeptide. Such additional amino acid residues
20 may comprise one or more amino acid residues added for the purpose of chemical coupling. One example of this is the addition of a cysteine residue. Such additional amino acid residues may also provide a "tag" for purification or detection of the polypeptide, such as a His₆ tag or a "myc" (c-myc) tag or a "FLAG" tag for interaction with antibodies specific to the tag or immobilized metal affinity chromatography (IMAC) in the case of the His₆-tag.

25 The further amino acids as discussed above may be coupled to the C5 binding polypeptide by means of chemical conjugation (using known organic chemistry methods) or by any other means, such as expression of the C5 binding polypeptide as a fusion protein.

The further amino acids as discussed above may for example comprise one or more polypeptide domain(s). A further polypeptide domain may provide the C5 binding polypeptide
30 with another function, such as for example another binding function, or an enzymatic function, or a toxic function (e.g. an immunotoxin), or a fluorescent signaling function, or combinations thereof.

A further polypeptide domain may moreover provide the C5 binding polypeptide with the same binding function. Thus, in a further embodiment, there is provided a C5 binding
35 polypeptide comprising at least two C5 binding polypeptide monomer units, the amino acid sequences of which may be the same or different. Multimeric forms of the polypeptides may comprise a suitable number of domains, each having a C5 binding motif, and each forming a

"monomer" within the multimer. These domains may all have the same amino acid sequence, but alternatively, they may have different amino acid sequences. In particular, the C5 binding polypeptide of the invention may form homo- or heterodimers.

5 The further polypeptide domain(s) as described above may be joined to the C5 binding polypeptide by covalent coupling using known organic chemistry methods. Alternatively, the C5 binding polypeptide comprising the further polypeptide domain(s) may be expressed as one or more fusion polypeptides, for example in a system for recombinant expression of polypeptides, or joined in any other fashion, either directly or via a linker, for example an amino acid linker.

10 In some embodiments, the further polypeptide domain(s) may comprise a half-life extending moiety which increases the half life of the C5 binding polypeptide *in vivo*. As understood by the skilled person, increased, or extended, half life means slower clearance of a particular molecule from blood. There are a number of known strategies for prolonging the half life of a particular polypeptide *in vivo*, such as coupling to the Fc domain of an antibody (Fc
15 conjugation) or coupling to albumin. Another example is coupling to a half life extending moiety, e.g. a peptide or protein, that will associate to serum albumin *in vivo*. In particular, the half life extending moiety may be an albumin binding moiety. An albumin binding moiety may e.g. consist of a naturally occurring polypeptide, or an albumin binding fragment thereof, or an engineered polypeptide. An engineered polypeptide may be derived from a naturally occurring
20 starting polypeptide through subjecting it to protein engineering techniques, such as mutations and alterations in a site-directed or randomized approach, with a view to create novel or enhanced properties, such as binding affinity for a molecule such as albumin. Such an engineered albumin binding polypeptide may for example be a variant of a protein scaffold, which variant has been selected for its specific binding affinity for albumin. In a specific
25 embodiment, the protein scaffold may be selected from domains of streptococcal Protein G or derivatives thereof, such as for example domain GA1, domain GA2 and domain GA3 of Protein G from *Streptococcus* strain G148, in particular domain GA3.

Accordingly, in one embodiment of the C5 binding polypeptide, the further amino acids improves stabilization *in vivo* or *in vitro* and comprise an albumin binding domain (ABD) of
30 streptococcal protein G, or a derivative thereof. One example of an albumin binding domain which may be comprised as a further polypeptide domain in the C5 binding polypeptide of the invention is set out in SEQ ID NO:759. Other examples of suitable albumin binding domains are disclosed in WO 2009/016043 and WO 2012/004384. Such an ABD-extended polypeptide binds to serum albumin *in vivo*, and benefits from its longer half life, which increases the net
35 half life of the polypeptide itself (see e.g. WO 91/01743). The pharmacokinetic profile of a C5 binding polypeptide comprising an albumin binding moiety as defined above thus resembles that of serum albumin when administered for example to a mammalian subject. ABD and

derivatives thereof bind very strongly to human serum albumin (HSA) as well as to serum albumin from other species, such as mouse and rat.

ABD of streptococcal protein G is 46 amino acid long, and thus when a C5 binding polypeptide according to the invention comprises an ABD moiety or a derivative thereof, the overall size of the C5 binding polypeptide is relatively small. When administered for example to a mammalian subject, such as a human subject, the albumin binding part of the C5 binding polypeptide will associate non-covalently with serum albumin and the polypeptide may thereby benefit from decreased renal clearance and increased recirculation in epithelial cells. Tissue penetration may however still be fast due to extravasating properties of serum albumin. Furthermore, a C5 binding polypeptide comprising a half life extending moiety may not only display an extended half life *in vivo*, but also a reduced immunologic response *in vivo*, as compared to a polypeptide lacking a corresponding half life extending moiety (see e.g. WO 2005/097202).

In a related aspect, there is provided a C5 binding compound, comprising at least one C5 binding polypeptide according to any preceding claim; at least one albumin binding domain of streptococcal protein G, or a derivative thereof, and at least one linking moiety for linking said at least one domain or derivative thereof to the C or N terminus of said at least one C5 binding polypeptide. Such a C5 binding compound has high affinity for C5 as well as for serum albumin *in vivo*, when administered e.g. to a mammalian subject, and binding to serum albumin does not interfere with the interaction with C5, as demonstrated in the following Examples.

In one embodiment, the C5 binding compound has a structure selected from

[CBP1]-[L1]-[ALBD];

[CBP1]-[CBP2]-[L1]-[ALBD];

[CBP1]-[L1]-[ALBD]-[L2]-[CBP2];

[ALBD]-[L1]-[CBP1];

[ALBD]-[L1]- [CBP1]-[CBP2];

[CBP1]-[L1]- [CBP2]-[L2]-[ALBD]; and

[ALBD]-[L1]- [CBP1]- [L2]- [CBP2]

wherein, independently of each other,

[CBP1] and [CBP2] are C5 binding polypeptides which may be the same or different;

[L1] and [L2] are linking moieties which may be the same or different; and

[ALBD] is an albumin binding domain of streptococcal protein G, or derivative thereof.

Preferred C5 binding compounds have a structure selected from

[CBP1]-[CBP2]-[L1]-[ALBD];

[CBP1]-[L1]-[ALBD]-[L2]-[CBP2]; and most preferably,

[CBP1]-[L1]-[ALBD].

Examples of linking moieties that may be used in such C5 binding compounds are selected from G, GS, $[G_2S]_n$, $[G_3S]_n$, $[G_4S]_n$, $GS[G_4S]_n$, wherein n is 0-7 (preferably, n is 0-2); $[S_2G]_m$, $[S_3G]_m$, $[S_4G]_m$; wherein m is 0-7, and VDGS. Preferred linkers are GS and $GS[G_4S]_2$.

5 Examples of albumin binding domains or derivatives thereof that may be comprised in a C5 binding compound are as described above. In particular, one example of an albumin binding domain is set out in SEQ ID NO:759.

Particularly preferred C5 binding compounds have the structure $[CBP1]-[LI]-[ALBD]$, wherein $[CBP1]$ is a polypeptide selected from SEQ ID NO:748 and SEQ ID NO:753, $[LI]$ is GS, and $[ALBD]$ is a polypeptide shown as SEQ ID NO:759.

10 The C5 binding polypeptide(s) comprised in a C5 binding polypeptide are, in one embodiment, independently selected from 58-mer or 60-mer C5 binding polypeptides as previously described. In particular, the C5 binding compound may comprise one or more C5 binding polypeptides independently selected from any one of SEQ ID NO:497-508, SEQ ID NO:516, SEQ ID NO:519-520, SEQ ID NO:522-524, SEQ ID NO:528-531, SEQ ID NO:534-
15 535, SEQ ID NO:537, SEQ ID NO:542, SEQ ID NO:545, SEQ ID NO:552-553, SEQ ID NO:555, SEQ ID NO:562, SEQ ID NO:574-575, SEQ ID NO:583, SEQ ID NO:588, SEQ ID NO:602, SEQ ID NO:606, SEQ ID NO:615, SEQ ID NO:621, SEQ ID NO:637, SEQ ID NO:647, SEQ ID NO:657, SEQ ID NO:662, SEQ ID NO:683, SEQ ID NO:693, SEQ ID NO:699, SEQ ID NO:701, SEQ ID NO:711, SEQ ID NO:739 and SEQ ID NO:746-757, such as
20 a sequence selected from SEQ ID NO:497-508 and SEQ ID NO:746-757. In another embodiment, the amino acid sequence is selected from SEQ ID NO:497, SEQ ID NO:498, SEQ ID NO:499, SEQ ID NO:500, SEQ ID NO:501, SEQ ID NO:746, SEQ ID NO:747, SEQ ID NO:748, SEQ ID NO:750 and SEQ ID NO:753, such as from any one of SEQ ID NO:497, SEQ ID NO:500, SEQ ID NO:748 and SEQ ID NO:753.

25 In a further aspect, there is provided a polynucleotide encoding a C5 binding polypeptide or a compound as described above. An expression vector comprising such a polynucleotide may enable production of a C5 binding polypeptide or a C5 binding compound, for example by expression in a host cell.

It should be understood that the C5 binding polypeptide according to the present
30 invention may be useful as a therapeutic or diagnostic agent in its own right or as a means for targeting other therapeutic or diagnostic agents, with e.g. direct or indirect effects on the complement protein C5. A direct therapeutic effect may for example be accomplished by inhibiting C5 cleavage. In one embodiment, there is thus provided a combination of a C5 binding polypeptide or a C5 binding compound according to the invention with a therapeutic
35 agent. Non-limiting examples of therapeutic agents that may prove useful in such a combination are immunostimulatory agents and radionuclides.

Thus, the C5 binding polypeptide as such, or as comprised in a C5 binding compound or a combination according to the invention, is in one embodiment provided for use in therapy, for example for the treatment of a C5 related condition, such as a C5 related condition in a mammal, such as a human subject. In one embodiment, said C5 related condition is selected

5 from inflammatory disease, such as antigen-induced arthritis, sepsis, synovial inflammation, vasculitis and asthma; autoimmune disease, such as systemic lupus erythematosus (SLE), cold agglutinin disease, rheumatoid arthritis, multiple sclerosis (MS), Sjögren's syndrome, dermatomyositis, myasthenia gravis and other autoantibody driven diseases such as Guillain-Barré syndrome (GBS), Fisher syndrome, systemic sclerosis, anti-glomerular basement

10 membrane (anti-GBM) and anti-phospholipid syndrome (APS); infectious disease, such as hemolytic-uremic syndrome (HUS), viral infections, bacterial infections and fungal infections; cardiovascular disease, such as (acute) myocardial infarction (undergoing revascularization either by fibrinolysis or percutaneous coronary intervention (PCI)); neurodegenerative disorders such as Alzheimer's disease (AD), Huntington's disease, Creutzfeld-Jacob disease and

15 Parkinson's disease; cancers; wounds; graft injury, such as ischemia reperfusion injury (IRI) and acute antibody mediated rejection (AMR); eye disease, such as age-related macular degeneration (AMD), uveitis, diabetic ocular diseases and disorders, and retinopathy of prematurity; kidney disease, such as membranous glomerulonephritis, membranous nephritis, immunoglobulin A nephropathy, Lupus nephritis, Goodpasture syndrome and post-

20 streptococcal glomerulonephritis; pulmonary diseases, such as adult respiratory distress syndrome, chronic obstructive pulmonary disease and cystic fibrosis; hematological diseases; such as hemolytic anaemia, paroxysmal cold hemoglobinuria, atypical hemolytic uremic syndrome (aHUS) and paroxysmal nocturnal hemoglobinuria (PNH); allergic diseases, such as anaphylactic shock, allergy and asthma; and dermatological diseases, such as pemphigus,

25 bullous pemphigoid, phototoxic reactions and psoriasis. In a more particular embodiment, the C5 binding polypeptide, compound or combination according to the invention is used for treatment of paroxysmal nocturnal hemoglobinuria (PNH).

As mentioned when discussing organ transplantation in the background section above, differences between donor and recipient (e.g. ABO and MCH classes) as well as the condition

30 of the transplanted organ may lead to delayed functioning or even rejection of the transplanted organ. Treatment may thus be necessary to eliminate anti-donor antibodies despite a positive donor-recipient crossmatch or to eliminate ABO antibodies when transplantation occurs against the ABO barrier. Such treatment typically includes immunoadsorption, e.g. by use of affinity chromatography techniques, prior to as well as after transplantation or plasmapheresis. Such

35 procedures however runs the risk of eliminating nearly all antibodies present in the circulation, thus including therapeutic antibodies. The C5 binding polypeptides or compounds of the

invention are however not affected by any antibody removing procedures, and may thus be exploited in these treatments.

In some C5 related conditions where a more local acute pathology in readily accessible tissues, such as lung and the blood stream, dominates rather than systemic pathologies, a drug with a very short half-life could be advantageous over one with a slow elimination. Thus, in such C5 related conditions, a C5 binding polypeptide without a half-life extending moiety may be beneficial. As previously accounted for, a C5 binding polypeptide according to the invention will, due to its relatively small size, exhibit a relatively rapid pharmacokinetic profile when administered to a mammal such as a human. The C5 binding polypeptide according to the invention may potentially be active in treatment of C5 related conditions such as asthma (Zhang *et al.* Expert Rev Clin Immunol 2010, 6:269-277), sepsis (Ward *et al.* The Sci World J 2010, 10:2395-2402), and hypersensitivity syndrome including the C activation-related pseudoallergy (CARPA, a reaction to certain therapeutic liposomes and lipid excipient-based drugs that in rare cases can lead to life threatening cardiopulmonary distress (Szebeni *et al.* Adv Drug Delivery Rev 2011, 63:1020-1030). In addition, a C5 binding polypeptide according to the invention may be used for complement inhibition when a recipient of blood transfusion has received blood of an incompatible type (a situation occurring in about 1:14000 transfusion units in the US which is associated with high mortality, Goodnough *et al.* Lancet 2003, 361:161-169).

In a related aspect, there is provided a method of treatment of a C5 related condition, comprising administering of a C5 binding polypeptide, or combination as described above to a mammalian subject in need thereof. Consequently, in the method of treatment, the subject is treated with a C5 binding polypeptide, a C5 binding compound or a combination according to the invention. In a more specific embodiment of said method, the binding of the C5 binding polypeptide or the combination, to a C5 expressed on a cell surface in the subject inhibits C5 cleavage. In one embodiment of the method of treatment, the C5 related condition is selected from inflammatory disease; autoimmune disease; infectious disease; cardiovascular disease; neurodegenerative disorders; cancers; wounds; graft injury; eye disease; kidney disease; pulmonary diseases; hematological diseases; allergic diseases and dermatological diseases. In particular the C5 related condition may be as defined above in relation to therapeutic use of a C5 binding polypeptide, compound or combination according to the invention. The C5 related condition may for example be paroxysmal nocturnal hemoglobinuria (PNH). In one embodiment of the method of treatment, the said C5 binding polypeptide is administered intravenously, subcutaneously, by inhalation, nasally, orally, intravitreally, or topically.

The invention as claimed relates to:

- C5 binding polypeptide, comprising a C5 binding motif, *BM*, which motif forms part of a three-helix bundle protein domain and consists of an amino acid sequence according to

EX₂X₃X₄A X₆X₇EID X₁₁LPNL X₁₆X₁₇X₁₈QW X₂₁AFIX₂₅ X₂₆LX₂₈D,

- 5 wherein, independently of each other, X₂ is selected from H, Q, S, T and V; X₃ is selected from I, L, M and V; X₄ is selected from A, D, E, H, K, L, N, Q, R, S, T and Y; X₆ is selected from N and W; X₇ is selected from A, D, E, H, N, Q, R, S and T; X₁₁ is selected from A, E, G, H, K, L, Q, R, S, T and Y; X₁₆ is selected from N and T; X₁₇ is selected from I, L and V; X₁₈ is selected from A, D, E, H, K, N, Q, R, S and T; X₂₁ is selected from I, L and V; X₂₅ is selected from D, E, G, H, N, S and T; X₂₆ is selected
10 from K and S; and X₂₈ is selected from A, D, E, H, N, Q, S, T and Y;

- C5 binding compound, comprising at least one C5 binding polypeptide as described herein; at least one albumin binding domain of streptococcal protein G, and at least one linking moiety for linking said at least one domain to the C or N terminal of said at least one C5 binding polypeptide;

- a fusion polypeptide comprising the C5 binding polypeptide as described herein and a further
15 polypeptide domain;

- polynucleotide encoding the C5 binding polypeptide as described herein or the C5 binding compound as described herein;

- combination of the C5 binding polypeptide as described herein or the C5 binding compound as described herein with a therapeutic agent;

- 20 - C5 binding polypeptide as described herein, C5 binding compound as described herein or combination as described herein for treatment of a C5 related condition selected from inflammatory disease; autoimmune disease; infectious disease; cardiovascular disease; neurodegenerative disorders; cancer; graft injury; wounds; eye disease; kidney disease; pulmonary diseases; hematological diseases; allergic diseases and dermatological diseases;

- 25 - C5 binding polypeptide as described herein, C5 binding compound as described herein or combination as described herein for treatment of paroxysmal nocturnal hemoglobinuria (PNH); and

- use of the C5 binding polypeptide as described herein, C5 binding compound as described herein or combination as described herein, for treatment of a C5 related condition in a mammalian subject in need thereof, wherein said C5 related condition is selected from inflammatory disease;
30 autoimmune disease; infectious disease; cardiovascular disease; neurodegenerative disorders; cancer; graft injury; wounds; eye disease; kidney disease; pulmonary diseases; hematological diseases; allergic diseases and dermatological diseases.

The invention will now be further illustrated by the following non-limiting Examples.

Brief description of the Figures

Figure 1 is a listing of the amino acid sequences of examples of C5 binding motifs comprised in C5 binding polypeptides of the invention (SEQ ID NO:1-248), examples of 49-mer C5 binding polypeptides according to the invention (SEQ ID NO:249-496), examples of 58-mer C5 binding polypeptides according to the invention (SEQ ID NO:497-744) and examples of 60-mer C5 binding polypeptides according to the invention (SEQ ID NO:745-757), as well as the sequences of protein Z (SEQ ID NO:758), an albumin binding domain (ABD094, SEQ ID NO:759), the Swiss-Prot entry P01031 of human C5 (amino acid residues 1-1676, SEQ ID NO:760; of which the α -chain corresponds to amino acid residues 678-1676 and the β -chain corresponds to amino acid residues 19-673), the sequence of the His₆-tagged tic protein OmCI used herein (SEQ ID NO:761) and cynomolgus C5 (SEQ ID NO:762) derived from genomic sequence (published on-line at www.ebi.ac.uk/ena; Ebeling et al. (2001) Genome Res. 21(10):1746-1756) using human C5 as template. The sequence contains two unknown amino acids "X" in positions 63 and 1346.

Figure 2 shows the result of a typical binding analysis performed in a Biacore instrument as described in Example 2. Sensorgrams were obtained by injection of human C5 (hC5; black solid curve), cynomolgus C5 (cC5; black short-dashed curve), rat C5 (rC5; black long-dashed curve), human MG7 domain (hMG7; gray dotted curve), and human immunoglobulin G (hIgG; gray solid curve), respectively, over an immobilized dimeric Z variant (Z05477, SEQ ID NO:509).

Figure 3 is a column chart showing the response in ELISA against hC5 and rC5, respectively, for selected matured Z variants. The black columns corresponds to the absorbance at 450 nm obtained using 0.05 μ g/ml hC5 (left column in each group) and to the absorbance at 450 nm obtained using 4 μ g/ml rC5 for each Z variant (right column in each group), as described in Example 4. The responses for the Z variant Z05363 (SEQ ID NO:510) are plotted as a positive control.

Figure 4 schematically shows different constructs encompassing one or several C5 binding Z variants selected from SEQ ID NO:745-757, optionally linked to ABD094 (SEQ ID NO:759).

Figure 5 shows SDS-PAGE analyses of purified C5 binding Z variants (reduced condition) visualized by Instant Blue, as described in Example 6. A) represents one example of dimeric Z-Z-ABD (lane 1 where Z is equal to SEQ ID NO:745 and ABD is equal to SEQ ID NO:759) compared with different Z-ABD fusion proteins (where Z is equal to SEQ ID NO:745 (lane 2), SEQ ID NO:748-757 (lanes 4-13) fused to ABD094 (SEQ ID NO:759) by a GS linker); B) represents one C5 binding Z variant (SEQ ID NO:753) in different constructs, and C) represents two different C5 binding Z variants (SEQ ID NO:748, lanes 2-3 and 6 and SEQ ID

NO:753, lanes 4-5 and 7), in monomeric form (lanes 6-7) and in fusion with ABD094 (SEQ ID NO:759) via a GS(G₄S)₂ linker (lanes 2-5).

Figures 6A and B are diagrams showing exemplary data of dose-response characterization of the potency of different C5 binding Z variants to inhibit complement activation as seen in a hemolytic assay, described in Example 6. C5 deficient serum was diluted 63-fold and supplemented with 0.1 nM hC5. A) shows effect of different Z-ABD fusion proteins (Z variants corresponding to SEQ ID NO:745, SEQ ID NO:748-753 and SEQ ID NO:756 fused to ABD094 (SEQ ID NO:759) by a GS linker) to hC5, whereas B) shows effect of different C5 binding constructs comprising the same C5 binding Z variant (Z06175a, SEQ ID NO:753) as monomer or dimer, in fusion with ABD094 (SEQ ID NO:759), or as provided with a His₆-tag (six histidine residues), compared to the C5 binding tick protein OmCI (SEQ ID NO:761).

Figures 7A and B are diagrams showing exemplary data of equilibrium binding based on the displacement ECL technique described in Example 6. Fig. 7A shows C5 binding of different Z variants (SEQ ID NO:745, SEQ ID NO:748-757) in fusion with ABD094 (SEQ ID NO:759) compared to C5 binding of the tick protein OmCI (SEQ ID NO:761). Fig. 7B shows binding of different C5 binding constructs comprising the same C5 binding Z variant (SEQ ID NO:753) as monomer or dimer, in fusion with ABD094 (SEQ ID NO:759) or as provided with a His₆-tag.

Figures 8A and 8B show interactions between Z-ABD variants and human serum albumin (HSA) studied as described in Example 7. A) Size exclusion chromatography (SEC) where Z-ABD (Z06175a (SEQ ID NO:753) fused to ABD094 (SEQ ID NO:759) by a GS linker) has been preincubated with equimolar amounts of HSA (1). As a comparison, the chromatograms for HSA alone (2) and Z-ABD alone (3) are also shown in the graph. B) Biacore sensorgrams of Z-ABD and Z-ABD-Z (Z06175a (SEQ ID NO:753) fused to ABD094 (SEQ ID NO:759) by linkers specified in Figure 4, in construct 2 and construct 5, respectively) injected over an HSA coated surface. Each of the two constructs was injected at a concentration of 25, 100 and 400 nM.

Figure 9 is a diagram showing the pharmacokinetic profiles for the C5 binding compounds Z-ABD and Z-ABD-Z (Z06175a, SEQ ID NO:753) fused to ABD094 (SEQ ID NO:759) by linkers specified in Figure 4, in construct 2 and construct 5, respectively) in Male Sprague Dawley rats over time after intravenous (i.v., 0.25 µmol/kg) and subcutaneous (s.c., 0.5 µmol/kg) administration, as described in Example 8. Each data point represents an average from three individual animals at a specific time point ranging from five minutes to two weeks after dosing for animals dosed i.v and from 15 minutes to two weeks for animals dosed s.c.

Figure 10 shows *ex vivo* hemolysis in sheep erythrocytes after exposure to serum diluted 1:5 from animal samples taken from Sprague Dawley rats after intravenous (i.v.; 0.25 µmol/kg)

administration of Z-ABD (Z06175a (SEQ ID NO:753) fused to ABD094 (SEQ ID NO:759) by a GS linker), as described in Example 8. Each dot represents one individual animal at a specific time point ranging from five minutes to two weeks after dosing.

Figure 11 shows *ex vivo* hemolysis in sheep erythrocytes after exposure to serum diluted 1:5 from animal samples taken from Sprague Dawley rats after subcutaneous (s.c.; 0.5 μ mol/kg) administration of Z-ABD (Z06175a (SEQ ID NO:753) fused to ABD094 (SEQ ID NO:759) by a GS linker), as described in Example 8. Each dot represents one individual animal at a specific time point ranging from 15 minutes to two weeks.

Figure 12 shows the hemolysis versus Z-ABD (Z06175a (SEQ ID NO:753) fused to ABD094 (SEQ ID NO:759) by a GS linker) serum concentration following i.v. and s.c. administration to male Sprague Dawley rats, as described in Example 8.

Figure 13 shows hemolysis in sheep erythrocytes versus Z-ABD-Z (Z06175a (SEQ ID NO:753) fused to ABD094 (SEQ ID NO:759) by linkers specified in Figure 4, construct 5) serum concentration following i.v. and s.c. administration to male Sprague Dawley rats, as described in Example 8.

Figure 14 shows the serum exposure of Z-ABD (Z06175a (SEQ ID NO:753) fused to ABD094 (SEQ ID NO:759) by a GS linker) following i.v. (415 nmol/kg) and s.c. (1250 nmol/kg) administration in male Cynomolgus monkey, as described in Example 9. Each data point represents the mean of three individual animals.

Figure 15 is a diagram showing the effect (C5a concentration in lavage) of the pro-inflammatory molecule zymosan (40 mg/kg i.p.) alone and in combination with a C5 binding Z-ABD fusion molecule (Z06175a (SEQ ID NO:753) fused to ABD094 (SEQ ID NO:759) by a GS linker) or OmCI (SEQ ID NO:761) analyzed as described in Example 10. Z-ABD was administered at 20 nmol/kg (LD), 100 nmol/kg (MD) and 500 nmol/kg (HD) s.c. 18 h before induction with zymosan. OmCI (30 nmol/kg) was administered i.p. 1 h before zymosan treatment and samples were taken 1 h after zymosan induction.

Figures 16A and 16B show the pharmacokinetic profile of Z-ABD (Z06175a (SEQ ID NO:753) fused to ABD094 (SEQ ID NO:759) by a GS linker) following intratracheal administration of 500 nmol/kg into female C57bl mice, as described in Example 11. A) serum concentration in each animal (n=3 for each time point, 27 animals totally) and B) hemolysis in sheep erythrocytes exposed to these serum samples diluted 1:5.

Examples

The following materials were used throughout this work except where otherwise noted.

- *Escherichia coli* strain RR1 Δ M15 (Rüther, Nucleic Acids Res 10:5765–5772, 1982).
- *Escherichia coli* strain XL1-Blue (Agilent Technologies, cat. no. 200268).

- Human complement protein C5 (hC5). The full 1676 amino acid pro-protein has GenBank accession number: NP_001726 (SEQ ID NO:760) wherein amino acids 19-673 is the beta chain and amino acids 678-1676 is the alpha chain. Human C5 used herein was purchased from Quidel (cat. no. A403)
- 5 • Cynomolgus complement protein C5 (cC5; SEQ ID NO:762). The cC5 sequence was derived from the Cynomolgus (*Macaca fascicularis*) genomic sequence (www.ebi.ac.uk/ena; Ebeling et al. (2001) Genome Res. 21(10):1746-1756). The coding region of human C5 was retrieved from www.ensembl.org, and the C5 gene was localized to chromosome 15. The region containing the gene is approximately 110 000
10 bases long and is contained in contigs CAEC01154150 to CAEC01154178. The contigs were manually joined to a single file and used as a genomic context for the sim4 software to align the coding region of human C5 to the raw Cynomolgus genomic material. Cynomolgus C5 used herein was purified in-house from serum using a three-step procedure; PEG6000 precipitation, ion exchange and OmCI affinity
15 chromatography.
- Rat Complement protein C5 (rC5; GenBank accession number: XP_001079130) Rat C5 used herein was purified in-house from serum using a three-step procedure; PEG6000 precipitation, ion exchange and OmCI affinity chromatography.
- Human MG7 (hMG7) domain of complement protein C5, corresponding to amino acid
20 residues 822-931 of human C5 (SEQ ID NO:760; Fredslund et al. (2008) Nature Immunology 9: 753-760) produced in-house in Freestyle HEK293 cells.
- hMG7 binding protein.
- OmCI (AF2999, Nunn, M. A. *et al. supra*) from soft tick *Ornithodoros moubata* OmCI with a His₆ tag in the C-terminus (SEQ ID NO:761) was produced in-house in *E. coli*
25 strain Origami(DE3) and purified on a HisTrap1 column.

Example 1: Selection and screening of complement protein C5 binding polypeptides

Materials and methods

- 30 Biotinylation of target protein hC5: hC5 was biotinylated according to the manufacturer's recommendations at room temperature (RT) for 40 min using No-Weigh EZ-Link Sulfo-NHS-LC-Biotin (Pierce, cat. no. 21327) at a ten times (10x) molar excess. Subsequent buffer exchange to PBS (10 mM phosphate, 137 mM NaCl, 2.68 mM KCl, pH 7.4) was performed using Protein Desalting Spin Columns (Pierce, cat. no. 89849) according to the manufacturer's
35 instructions.

Phage display selection of C5-binding polypeptides: Libraries of random variants of protein Z displayed on bacteriophage, constructed in phagemid pAffi1/pAY00065/ pAY02947/pAY02592 essentially as described in Grönwall *et al.* J Biotechnol 2007, 128:162-183), were used to select C5 binding polypeptides. Three different library vectors were used. Two of these utilize an

5 albumin binding domain (ABD, GA3 of protein G from *Streptococcus* strain G148) as fusion partner to the Z variants generating the libraries Zlib003Naive.I and Zlib006Naive.II. The third library, Zlib004Naive.I utilizes the *Taq* DNA polymerase binding molecule Z03639 (denoted $Z_{TaqSI-1}$ in Gunneriusson *et al.* Protein Eng 1999, 12:873-878) as fusion partner. The libraries had the following actual sizes: 3×10^9 (Zlib003Naive.I); 1.5×10^{10} (Zlib006Naive.II); and $1.4 \times$

10 10^{10} (Zlib004Naive.I), the number referring to the amount of variants.

Phage stocks were prepared either in shake flasks (Zlib003Naive.I) as described in Grönwall *et al. supra* or in a 20 l fermenter (Zlib006Naive.II and Zlib004Naive.I). Cells from a glycerol stock containing the phagemid library Zlib004Naive.I were inoculated in 20 l of TSB-YE (Tryptic Soy Broth-Yeast Extract; 30 g/l TSB, 5 g/l yeast extract) supplemented with 2 %

15 glucose and 100 µg/ml ampicillin. Cells from a glycerol stock containing the phagemid library Zlib006Naive.II were inoculated in 20 l of a defined proline free medium [dipotassium hydrogenphosphate 7 g/l, trisodium citrate dihydrate 1 g/l, uracil 0.02 g/l, YNB (Difco™ Yeast Nitrogen Base w/o amino acids, Becton Dickinson) 6.7 g/l, glucose monohydrate 5.5 g/l, L-

20 alanine 0.3 g/l, L-arginine monohydrochloride 0.24 g/l, L-asparagine monohydrate 0.11 g/l, L-cysteine 0.1 g/l, L-glutamic acid 0.3 g/l, L-glutamine 0.1 g/l, glycine 0.2 g/l, L-histidine 0.05 g/l, L-isoleucine 0.1 g/l, L-leucine 0.1 g/l, L-lysine monohydrochloride 0.25 g/l, L-methionine 0.1 g/l, L-phenylalanine 0.2 g/l, L-serine 0.3 g/l, L-threonine 0.2 g/l, L-tryptophane 0.1 g/l, L-tyrosine 0.05 g/l, L-valine 0.1 g/l] supplemented with 100 µg/ml ampicillin. The cultivations were grown at 37°C in a fermenter (Belach Biotechnik, BR20). When the cells reached an optical

25 density (OD) of 0.7-0.8, approximately 2.6 l of the cultivation was infected using a 10x molar excess of M13K07 helper phage (New England Biolabs #N0315S). The cells were incubated for 30 minutes, whereupon the fermenter were filled up to 20 l with TSB-YE supplemented with 100 µM IPTG (isopropyl-β-D-1-thiogalactopyranoside, for induction of expression), 25 µg/ml kanamycin and 12.5 µg/ml carbenicillin and grown at 30°C for 22 h. The cells in the cultivation

30 were pelleted by centrifugation at 15,900 g and the phage particles remaining in the medium were thereafter precipitated twice in PEG/NaCl (polyethylene glycol/sodium chloride), filtered and dissolved in PBS and glycerol as described in Grönwall *et al. supra*. Phage stocks were stored at -80°C before use.

Selections were performed in four cycles against biotinylated hC5. Phage stock

35 preparation, selection procedure and amplification of phage between selection cycles were performed essentially as described in WO 2009/077175. PBS supplemented with 3 % bovine serum albumin (BSA) and 0.1 % Tween20 was used as selection buffer and the target-phage

complexes were directly captured by Dynabeads® M-280 Streptavidin (Dyna, cat. no. 112.06). 1 mg beads per 10 µg complement protein C5 was used. *E. coli* strain RR1ΔM15 was used for phage amplification. In cycle 1 of the selections, 100 nM hC5 was used and two washes with PBST 0.1 % (PBS supplemented with 0.1 % Tween-20) were performed. An increased stringency, using a lowered target concentration and an increased number of washes, was applied in the subsequent three cycles. In cycle 2, 3 and 4; 50 or 33 nM hC5, 25 or 11 nM hC5 and 12.5 or 3.7 nM hC5 were used. In cycle 2, 3 and 4; 4, 6 and 8 washes were performed, using PBST 0.1 % in all cycles or PBST 0.2 %, 0.3 % and 0.4 % in cycle 2, 3 and 4.

ELISA screening of Z variants: To test if the selected Z variant molecules could indeed interact with human complement protein C5, ELISA assays were performed. The Z variants were produced by inoculating single colonies from the selections into 1 ml TSB-YE medium supplemented with 100 µg/ml ampicillin and 0.1 mM IPTG in deep-well plates (Nunc, cat. no. 278752). The plates were incubated for 18-24 h at 37°C. Cells were pelleted by centrifugation, re-suspended in 300 µl PBST 0.05 % and frozen at -80°C to release the periplasmic fraction of the cells. Frozen samples were thawed in a water bath and cells were pelleted by centrifugation. The periplasmic supernatant contained the Z variants as fusions to an albumin binding domain (GA3 of protein G from *Streptococcus* strain G148), expressed as AQHDEALE-[Z#####]-VDYV-[ABD]-YVPG (Grönwall *et al, supra*), or to the *Taq* DNA polymerase binding molecule Z03639, expressed as AQHDEALE-[Z#####]-VDYV-[Z03639]-YVPG. Z##### refers to individual 58 amino acid residues Z variants.

Half-area 96-well ELISA plates (Costar, cat. no. 3690) were coated with 50 µl/well of coating buffer (50 mM sodium carbonate, pH 9.6) containing 4 µg/ml of an antibody specific for Z variants (Affibody, cat. no. 20.1000.01.0005) and incubated over-night at 4°C. The antibody solution was poured off and the wells were blocked with 100 µl of PBSC (PBS supplemented with 0.5 % casein (Sigma, cat. no. C8654) for 1-2 h at RT. The blocking solution was discarded and 50 µl periplasmic solution was added to the wells and incubated for 1 h at RT under slow shaking. The supernatants were poured off and the wells were washed 4 times with PBST 0.05 %. Then 50 µl of biotinylated complement protein hC5, at a concentration of 5 µg/ml in PBSC, was added to each well. The plates were incubated for 1.5 h at RT followed by washes as described above. Streptavidin-HRP (Horseradish peroxidase; Dako, cat. no. P0397) was diluted 1:10,000 in PBSC, added to the wells which were then incubated for 45 min. After washing as described above, 50 µl ImmunoPure TMB substrate (Thermo Scientific, cat. no. 34021) was added to the wells and the plates were treated according to the manufacturer's recommendations. Absorbance of the wells was measured at 450 nm using a multi-well plate reader, Victor³ (Perkin Elmer).

As positive control, a periplasmic fraction also containing the PSMA binding molecule Z03938 expressed as AQHDEALE-[Z03938]-VDYV-[Z03639]-YVPG was assayed against 5

µg/ml biotinylated PSMA protein. As negative control; the same periplasmic preparation was assayed against complement protein hC5. Sequencing was performed for the clones with positive absorbance values against hC5.

Sequencing: PCR fragments were amplified from single colonies using a standard PCR program and the primers AFFI-21 (5'-tgcttcggctcgatgtgtgtg) and AFFI-22 (5'-cggaaccagaccaccgg). Sequencing of amplified fragments was performed using the biotinylated oligonucleotide AFFI-72 (5'-biotin-cggaaccagaccaccgg) and a BigDye® Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems), used in accordance with the manufacturer's protocol. The sequencing reactions were purified by binding to magnetic streptavidin coated beads (Detach Streptavidin Beads, Nordiag, cat. no. 2012-01) using a Magnatrix 8000 (Magnetic Biosolution), and analyzed on ABI PRISM® 3100 Genetic Analyzer (PE Applied Biosystems).

Blocking ELISA: Clones found positive for hC5 in the ELISA screening were subjected to an ELISA blocking assay in order to elucidate if their target binding was affected by the presence of the hC5 binding proteins OmCI and/or hMG7 binding protein. The blocking ELISA was run using Z variants expressed in periplasmic fractions as described in the section for ELISA screening above, but setting up 5 ml cultures in 12 ml round-bottom tubes and using 2 ml PBST 0.05 % for pellet dissolution. The ELISA blocking assay was run as the ELISA screening assay, with a protocol modification introduced at the target step; OmCI or hMG7 binding protein were mixed with the target protein before addition to the assay plate. 5 µg/ml biotinylated hC5 was mixed with 5 times or 20 times molar excess of OmCI or hMG7 binding protein, respectively, then incubated 1 h at RT to allow complex formation before addition to the plate. For each clone, a reference (1), a negative control (2) and a background (3) response/signal, respectively, were obtained as follows: at the target step, solely hC5 was added to the Z variants (as in the screening ELISA) (1); the irrelevant protein PSMA (in house produced) was added to complement protein hC5, instead of OmCI or hMG7 binding protein (2); only buffer was added to the Z variants (3).

Results

Phage display selection of complement protein C5 binding polypeptides: Individual clones were obtained after two-four cycles of phage display selections against biotinylated hC5.

ELISA screening of Z variants: The clones obtained after four cycles of selection were produced in 96-well plates and screened for complement protein C5 binding activity in ELISA. In total, nearly 400 clones were screened. The absorbance measurements showed many clearly hC5 positive clones. The result from a selection of clones is displayed in Table 1; the Z05363 (SEQ ID NO:510) variant is tagged with ABD, whereas the other listed Z variants are tagged with the Taq binding molecule Z03639 as described in the methods section. The PSMA specific

molecule Z03938 used as a negative control gave a positive signal for PSMA, whereas no signal was obtained against hC5.

Blocking ELISA: Clones positive for hC5 were subjected to a blocking assay using the hC5 binding proteins OmCI and hMG7 binding protein. For five clones, the binding signal to complement protein C5 was completely extinguished by the presence of OmCI, reaching the same level as the background (Table 1). One of these clones, namely the Z05363 variant (SEQ ID NO:510), was also tested for its ability to bind hC5 in the presence of hMG7 binding protein. The hMG7 binding protein did not inhibit the binding of Z05363 to hC5.

Table 1. Response in ELISA to target, with or without blocking molecule for a number of Z variants.

Z variant	SEQ ID NO:#	hC5 (OD 450 nm)	OmCI-block
Z05363	SEQ ID NO:510	3.143	complete
Z05477	SEQ ID NO:509	2.872	complete
Z05483	SEQ ID NO:511	0.531	complete
Z05538	SEQ ID NO:512	0.099	complete
Z05692	SEQ ID NO:513	0.944	complete

Sequencing: Sequencing was performed for the clones with positive absorbance values against complement protein C5 in the ELISA screening. Each variant was given a unique identification number #####, and individual variants are referred to as Z#####. The amino acid sequences of the 58 amino acid residues long Z variants are listed in Figure 1 and in the sequence listing as SEQ ID NO:509-513. The deduced complement protein C5 binding motifs of these Z variants are listed in Figure 1 and in the sequence listing as SEQ ID NO:13-17. The amino acid sequences of the 49 amino acid residues long polypeptides predicted to constitute the complete three-helix bundle within each of these Z variants are listed in Figure 1 and in the sequence listing as SEQ ID NO:261-265.

Example 2: Production and characterization of Z variants

Materials and methods

Subcloning of Z variants, protein expression and purification:

Five complement protein C5 binding Z variants (Z05363 (SEQ ID NO:510); Z05477 (SEQ ID NO:509); Z05483 (SEQ ID NO:511); Z05538 (SEQ ID NO:512) and Z05692 (SEQ ID NO:513)) were amplified from pAffi1/pAY00065/pAY02947 library vectors. A subcloning strategy for construction of dimeric Affibody molecules with N-terminal His₆ tags was applied

using standard molecular biology techniques and as described in detail in WO 2009/077175. The Z gene fragments were subcloned into the expression vector pAY01448 resulting in the encoded sequence MGSSHHHHHHLQ-[Z#####][Z#####]-VD.

5 The subcloned Z variants were transformed into *E. coli* BL21(DE3) and expressed in the multifermenter system Greta (Belach Biotechnik). In brief, cultures were grown at 37°C in 800 ml TSB-YE-medium containing 50 µg/ml kanamycin. At an OD₆₀₀ of ~1, the cultures were induced through the automatic addition of IPTG to a final concentration of 0.05 mM. Cultures were cooled down to approximately 10°C after 5 h of induction, and harvested by centrifugation (20 min, 15,900 g). Supernatants were discarded and the cell pellets were collected and stored at
10 -20°C until further use. Expression levels and the degree of solubility were estimated by SDS-PAGE analysis on 4-12 % NuPAGE™ gels (Invitrogen) using Coomassie blue staining.

For Z variants expressed mainly as soluble protein, the cell pellets were resuspended in binding buffer (20 mM sodium phosphate, 0.5 M NaCl, 20 mM imidazole, pH 7.4) with an addition of 1000 U Benzonase® (Merck, cat. no. 1.01654.001) and disrupted by ultrasonication.
15 For each of the Z variants, the sonicated suspension was clarified by centrifugation (40 min, 25,000 g, 4 °C) and the supernatant was loaded onto a 1 ml His GraviTrap™ column (GE Healthcare). The column was washed with wash buffer (20 mM sodium phosphate, 0.5 M NaCl, 60 mM imidazole, pH 7.4), before eluting the Z variants with 3 ml elution buffer (20 mM sodium phosphate, 0.5 M NaCl, 0.5 M imidazole, pH 7.4). Z variants which expressed mainly
20 as insoluble protein were purified likewise, but 8 M urea was included in the binding and wash buffer. If required, the Z variants were further purified by reversed phase chromatography (RPC) on 1 ml Resource™ columns (GE Healthcare) using water including 0.1 % TFA (trifluoroacetic acid) as mobile phase and elution with an appropriate gradient (typically 0-50 % over 20 column volumes) of acetonitrile including 0.1 % TFA.

25 The buffer was exchanged to PBS using PD-10 columns (GE Healthcare).

Protein characterization: The concentration of the purified Z variants was determined by absorbance measurements at 280 nm using theoretical extinction coefficients. The purity was estimated by SDS-PAGE analysis on 4-12 % NuPAGE™ gels (Invitrogen) using Coomassie blue staining. To verify the identity and to determine the molecular weights of purified Z
30 variants, LC/MS-analyses were performed on an Agilent 1100 LC/MSD system (Agilent Technologies).

CD analysis: The purified Z variants were diluted to 0.5 mg/ml in PBS. For each diluted Z variant, a CD spectrum was recorded between 250-195 nm at a temperature of 20°C. In addition, a variable temperature measurement (VTM) was performed to determine the melting
35 temperature (T_m). In the VTM, the absorbance was measured at 221 nm while the temperature was raised from 20 to 90°C, with a temperature slope of 5°C/min. The ability of the Z variant to refold was assessed by collecting an additional CD spectrum at 250-195 nm after cooling to

20°C. The CD measurements were performed on a Jasco J-810 spectropolarimeter (Jasco Scandinavia AB) using a cell with an optical path length of 1 mm.

Biacore binding analysis: The interactions of the five subcloned His₆-tagged dimeric hC5-binding Z variants with hC5, cC5, rC5, hMG7 and hIgG (Sigma, cat. no. G4386) were analyzed in a Biacore instrument (GE Healthcare). The Z variants were immobilized in different flow cells on the carboxylated dextran layer of several CM5 chip surfaces (GE Healthcare). The immobilization was performed using amine coupling chemistry according to the manufacturer's protocol. One flow cell surface on each chip was activated and deactivated for use as blank during analyte injections. The analytes, diluted in HBS-EP running buffer (GE Healthcare) to a final concentration of 100 nM, were injected at a flow rate of 10 µl/min for 1 min. After 2 min of dissociation, the surfaces were regenerated with one injection of 10 mM HCl. The results were analyzed in BiaEvaluation software (GE Healthcare). Curves of the blank surface were subtracted from the curves of the ligand surfaces.

15 *Results*

Subcloning of Z variants: Five selected unique clones (Z05477 (SEQ ID NO:509), Z05363 (SEQ ID NO:510), Z05483 (SEQ ID NO:511), Z05538 (SEQ ID NO:512) and Z05692 (SEQ ID NO:513)) were chosen for subcloning as dimers in the expression vector pAY01448 and were subsequently verified by sequencing.

Protein production: The histidine-tagged dimeric Z variants yielded acceptable expression levels of soluble gene product. The purity of produced batches was estimated to exceed 90 % as assessed by SDS-PAGE analysis. LC/MS analysis verified the correct molecular weight for all Z variant molecules.

CD analysis: The melting temperatures (T_m) of the different Z variants were calculated by determining the midpoint of the transition in the CD signal vs. temperature plot. The results for a number of reversibly folding Z variants are summarized in Table 2 below.

Table 2. Melting temperatures for a number of Z variants.

Z variant	SEQ ID NO:# of monomeric Z variant	T _m (°C)
His ₆ -(Z05477) ₂	SEQ ID NO:509	45
His ₆ -(Z05363) ₂	SEQ ID NO:510	35
His ₆ -(Z05483) ₂	SEQ ID NO:511	44
His ₆ -(Z05538) ₂	SEQ ID NO:512	54
His ₆ -(Z05692) ₂	SEQ ID NO:513	52

Biacore binding analysis: The binding of the five subcloned dimeric Z variants to different species of C5 and MG7, a subdomain of hC5, as well as the background binding to IgG was tested in a Biacore instrument by injecting the different proteins over surfaces containing the Z variants. The ligand immobilization levels for the different Z variants on the surfaces were:

5 Z05363: 2080 RU, Z05477: 2180 RU, Z05483: 2010 RU, Z05538: 2570 RU and Z05692: 3270 RU. The different Z variants were tested for binding to different sets of proteins injected at concentrations of 100 nM, see Table 3. The result for the tested Z variants is displayed in the table as a +/- outcome for each protein. As an example of the Biacore binding analysis, Figure 2 shows the sensorgrams obtained from immobilized dimeric Z05477 assayed against hC5, cC5,

10 rC5, hMG7 and hIgG.

Table 3. Biacore response of different Z variants against C5 from various species and relevant selected background proteins.

Z variant	SEQ ID NO:# of monomeric Z variant	hC5	cC5	rC5	hMG7	hIgG
His ₆ -(Z05477) ₂	SEQ ID NO:509	+	+	+	-	-
His ₆ -(Z05363) ₂	SEQ ID NO:510	+	+	+	-	-
His ₆ -(Z05483) ₂	SEQ ID NO:511	+	+	+	-	-
His ₆ -(Z05538) ₂	SEQ ID NO:512	+	+	+	-	-
His ₆ -(Z05692) ₂	SEQ ID NO:513	+	+	-	-	-

15 Example 3: Design and construction of a matured library of complement protein C5 binding Z variants

In this Example, a matured library was constructed. The library was used for selections of hC5-binding polypeptides. Selections from matured libraries are usually expected to result in

20 binders with increased affinity (Orlova *et al.* Cancer Res 2006, 66(8):4339-48). In this study, randomized double stranded linkers were generated by the Slonomics® technology which enables incorporation of randomized sets of trinucleotide building blocks using ligations and restrictions of the subsequently built up double stranded DNA.

25 *Materials and methods*

Library design: The library was based on a selection of sequences of the hC5 binding Z variants described in Examples 1 and 2. In the new library, 13 variable positions in the Z molecule scaffold were biased towards certain amino acid residues, according to a strategy based on the Z variant sequences defined in SEQ ID NO:509-513 (Z05477, Z05363, Z05483, Z05538,

30 Z05692). A SlonoMax® library of double-stranded DNA, containing the 147 bp partially

randomized helix 1 and 2 of the amino acid sequence 5'-AA ATA AAT CTC GAG GTA GAT GCC AAA TAC GCC AAA GAA/GAG NNN NNN NNN GCA/GCC NNN NNN GAG/GAA ATC/ATT NNN NNN TTA/CTG CCT AAC TTA ACC/ACT NNN NNN CAA/CAG TGG NNN GCC/GCG TTC ATC/ATT NNN AAA/AAG TTA/CTG NNN GAT/GAC GAC CCA AGC CAG AGC TCA TTA TTT A-3' (randomized codons are illustrated as NNN) flanked with restriction sites *Xho*I and *Sac*I, was ordered from Sloning BioTechnology GmbH (Puchheim, Germany). The theoretical distributions of amino acid residues in the new library finally including 12 variable Z positions are given in Table 4.

10 **Table 4:** Library design.

Amino acid position in the Z variant molecule	Randomization (amino acid abbreviations)	No of amino acids	Proportion
9	H,Q,S,T,V	5	1/5
10	I,L,V,W	4	1/4
11	A,D,E,H,K,L,N,R,S,T,Y	12	1/12
13	N,Q,W,Y	4	1/4
14	A,D,E,H,I,K,L,N,Q,R,S,T,V,W,Y	15	1/14
17	D,E	2	1/2
18	A,D,E,G,H,I,K,L,Q,R,S,T,V,Y	14	1/14
24	I,L,V	3	1/3
25	A,D,E,H,K,N,Q,R,S,T,Y	11	1/11
28	I,L,V	3	1/3
32	A,D,E,F,G,H,K,L,N,Q,R,S,T,V	14	1/14
35	A,D,E,H,K,N,Q,R,S,T,W,Y	12	1/12

Library construction: The library was amplified using *AmpliTaq* Gold polymerase (Applied Biosystems, cat. no. 4311816) during 12 cycles of PCR and pooled products were purified with QIAquick PCR Purification Kit (QIAGEN, cat. no. 28106) according to the supplier's recommendations. The purified pool of randomized library fragments was digested with restriction enzymes *Xho*I and *Sac*I (New England Biolabs, cat. no. R01460L, and cat. no. R0156L) and purified once more with PCR Purification Kit. Subsequently, the product was purified using preparative gel electrophoresis on a 1 % agarose gel.

The phagemid vector pAY02592 (essentially as pAffi1 described in Grönwall *et al. supra*) was restricted with the same enzymes, purified using phenol/chloroform extraction and ethanol precipitation. The restricted fragments and the restricted vector were ligated in a molar ratio of 5:1 with T4 DNA ligase (New England Biolabs, cat. no. M0202S), for 2 hours at RT

followed by overnight incubation at 4°C. The ligated DNA was recovered by phenol/chloroform extraction and ethanol precipitation, followed by dissolution in 10 mM Tris-HCl, pH 8.5.

The ligation reactions (approximately 250 ng DNA/transformation) were electroporated into electrocompetent *E. coli* RR1ΔM15 cells (100 μl). Immediately after electroporation, approximately 1 ml of SOC medium (TSB-YE media, 1 % glucose, 50 μM MgCl₂, 50 μM MgSO₄, 50 μM NaCl and 12.5 μM KCl) was added. The transformed cells were incubated at 37°C for 50 min. Samples were taken for titration and for determination of the number of transformants. The cells were thereafter pooled and cultivated overnight at 37°C in 7 l of TSB-YE medium, supplemented with 2 % glucose and 100 μg/ml ampicillin. The cells were pelleted for 15 min at 4,000 g, resuspended in a PBS/glycerol solution (approximately 40 % glycerol). The cells were aliquoted and stored at -80°C. Clones from the library of Z variants were sequenced in order to verify the content and to evaluate the outcome of the constructed library vis-à-vis the library design. Sequencing was performed as described in Example 1 and the amino acid distribution was verified.

Preparation of phage stock: Cells from the glycerol stock containing the C5 phagemid library were inoculated in 20 l of a defined proline free medium (described in Example 1) supplemented with 100 μg/ml ampicillin, and grown at 37°C in a fermenter (Belach Biotechnik, BR20). All steps were performed as described in Example 1 for the library Zlib006Naive.II. After cultivation, the cells were pelleted by centrifugation at 15,900 g and the phage particles remaining in the medium were thereafter precipitated twice in PEG/NaCl, filtered and dissolved in PBS and glycerol as described in Example 1. Phage stocks were stored at -80°C until use in selection.

Results

Library construction: The new library was designed based on a set of OmCI-blocked C5 binding Z variants with verified binding properties (Example 1 and 2). The theoretical size of the designed library was 6.7×10^9 Z variants. The actual size of the library, determined by titration after transformation to *E. coli* RR1ΔM15 cells, was 1.4×10^9 transformants.

The library quality was tested by sequencing of 64 transformants and by comparing their actual sequences with the theoretical design. The contents of the actual library compared to the designed library were shown to be satisfying. The locked position in the designed amino acid sequence (W in position 27) was reflected in the actual sequence in that only the expected amino acid occurred in that position. A matured library of hC5 binding polypeptides was thus successfully constructed.

Example 4: Selection, screening and characterization of Z variants from a matured library*Materials and methods*Phage display selection of complement protein C5 binding polypeptides: The target protein hC5

5 was biotinylated as described in Example 1. Phage display selections were performed against hC5 essentially as described in Example 1 using the new library of Z variant molecules described in Example 3. *E. coli* XL1-Blue was used for phage amplification. Selection was initially performed in two parallel tracks. In one track, the time of selection was 2 h, while in the other track, shorter selection times were used: 20 min in the first cycle and 10 min for

10 subsequent cycles 2-4. These two tracks (1 and 2) were further divided in the second cycle, resulting in totally six tracks (1a-c and 2a-c, differing in target concentration and wash conditions). Selection was performed in a total of four cycles. In cycle 1 of the selections, 25 nM complement protein C5 was used and five washes with PBST 0.1 % were performed. An increased stringency, using a lowered target concentration and an increased number of washes,

15 was applied in the subsequent three cycles. In cycle 2, 3 and 4; 10, 5 or 2.5 nM complement protein C5, 4, 1 or 0.25 nM complement protein C5 and 1.6, 0.2 or 0.05 nM complement protein C5 were used. In cycle 2, 3 and 4; 10, 15 and 20 washes were performed using PBST 0.1 %. In addition, the second last wash was prolonged to 3 h with a 50x excess of non-biotinylated hC5 in the washing solution for two of the tracks (1c and 2c).

20 Sequencing of potential binders: Individual clones from the different selection tracks were picked for sequencing. All clones run in the ELISA screening were sequenced. Amplification of gene fragments and sequence analysis of gene fragments were performed as described in Example 1.

ELISA screening of Z variants: Single colonies containing Z variants were randomly picked

25 from the selected clones of the complement protein C5 matured library and grown in 1 ml cultivations as described in Example 1. Periplasmic proteins were released by 8 repeated freeze-thawing cycles. ELISA screenings were performed essentially as described in Example 1 with the following exceptions. Half-area 96-well ELISA plates were coated with 2 µg/ml of an ABD specific goat antibody (in house produced) diluted in coating buffer. Biotinylated hC5 was used

30 at a concentration of 0.15 µg/ml and incubation performed for 1.5-2 h. Streptavidin conjugated HRP was obtained from Thermo Scientific (cat. no. N100). The Z variant Z05363 (SEQ ID NO:510) originating from the primary selections (Example 1) was used as a positive control as well as a negative control omitting hC5.

Selected matured Z variants were subjected to a second screen against hC5 at a lower

35 concentration and compared to rC5. The assay was essentially performed as described above. hC5 and rC5 was used at a concentration of 0.05 µg/ml and 4 µg/ml, respectively. The Z variant Z05363 (SEQ ID NO:510) was used as a positive control in this experiment as well. As a

negative control, a Z variant binding to PDGF-R β (Z01977; described in WO 2009/077175) was assayed against biotinylated hC5 or rC5.

In deep sequence analysis of selected Z variants and correlation of amino acids in the 13 randomized positions with measured melting temperatures and IC₅₀ values for human C5 and mouse C5 in the hemolysis assay (described in Example 6) suggested a favorable Z variant not identified among the 558 sequenced clones. Based on the Z variant Z05998 (SEQ ID No:499), a single amino acid, Ile in position 10 was substituted with Leu using conventional technology for site directed mutagenesis. The new variant is referred to as Z08044 (SEQ ID NO:498). The deduced complement protein C5 binding motif of this Z variant is listed in Figure 1 and in the sequence listing as SEQ ID NO:2. The amino acid sequences of the 49 amino acid residues long polypeptide predicted to constitute the complete three-helix bundle within these Z variant is listed in Figure 1 and in the sequence listing as SEQ ID NO:250.

Results

Phage display selection of complement protein C5 binding polypeptides: Selection was performed in totally six parallel tracks containing four cycles each. The different selection tracks differed in target concentration and wash conditions as follows: 1a) 2 h selection time, high concentration, standard wash, 1b) 2 h selection time, low concentration, standard wash, 1c) 2 h selection time, medium concentration, long wash, 2a) 10 min selection time, high concentration, standard wash, 2b) 10 min selection time, low concentration, standard wash, and 2c) 10 min selection time, medium concentration, long wash. For each selection cycle, the target concentration was decreased and the washing conditions were more stringent. All tracks gave in each round sufficient amounts of phage particles in the eluate. Most phage particles were found in tracks 1a and 2a, representing the highest target concentration and mildest wash conditions.

Sequencing: Randomly picked clones (558) were sequenced. Each individual Z variant was given an identification number, Z#####, as described in Example 1. In total, 242 new unique Z variant molecules were identified. The amino acid sequences of the 58 amino acid residues long Z variants are listed in Figure 1 and in the sequence listing as SEQ ID NO:497, SEQ ID NO:499-508 and SEQ ID NO:514-744. The deduced complement protein C5 binding motifs of these Z variants are listed in Figure 1 and in the sequence listing as SEQ ID NO:1, SEQ ID NO:3-12 and SEQ ID NO:18-248. The amino acid sequences of the 49 amino acid residues long polypeptides predicted to constitute the complete three-helix bundle within each of these Z variants are listed in Figure 1 and in the sequence listing as SEQ ID NO:249, SEQ ID NO:251-260 and SEQ ID NO:266-496. Among the sequenced clones, 63 sequences occurred two or more times.

ELISA screening of Z variants: Clones obtained after four selection cycles were produced in 96-well plates and screened for hC5-binding activity using ELISA. All randomly picked clones

were analyzed. 229 of the 242 unique Z variants were found to give a higher response (0.3-3.1 AU) against hC5 at a concentration of 0.15 µg/ml compared to the positive control clone Z05363 (SEQ ID NO:510; an average absorbance signal of 0.3 AU), obtained from the primary selections (Example 1). Clones from all selection tracks showed positive signals. The negative controls had an absorbance of approximately 0.1 AU.

Z variants were selected based on their performance in the ELISA screen against hC5 and the occurrence frequency. 43 unique Z variants were assayed against a lower concentration of hC5 (0.05 µg/ml) as well as rC5 (4 µg/ml). A positive result against rC5 was obtained for 40 of the tested Z variants, defined as 2x the signal for the negative control (0.4 AU). The results for all the tested Z variants against the lower concentration of hC5 as well as against rC5 are shown in Figure 3.

Example 5: Subcloning, production and characterization of a subset of complement protein C5 binding Z variants

Materials and methods

Subcloning of Z variant molecules into expression vectors: Based on sequence analysis and the performance in the ELISA against human and rat complement protein C5, 45 clones were selected for subcloning into the expression vector pAY01448. Monomer Z variant fragments were amplified from the phagemid vector pAY02592 and the subcloning into pAY01448 was performed as described in Example 2, resulting in a vector encoding the protein sequence MGSSHHHHHLQ-[Z#####]-VD.

Protein expression and purification: The 45 Z variants in the His₆-(Z#####) format, were expressed in an automated multifermenter system as described in Example 2 or similarly in a small scale set-up of 100 ml cultures in shaker flasks induced manually with IPTG to a final concentration of 0.4 mM. Purification was performed using 1 ml HisGraviTrapTM columns essentially as described in Example 2 or in a smaller scale using 0.1 ml His SpinTrap (GE Healthcare, cat. no. 28-4013-53). Buffer was exchanged to PBS using PD-10 columns or PD SpinTrap G-25 (GE Healthcare, cat. no. 28-9180-04) according to the manufacturer's instructions. The concentration of purified Z variants was determined by absorbance measurements at 280 nm and the purity and identity was assessed by SDS-PAGE and LC/MS as described in Example 2. Samples were aliquoted and stored at -80°C until further use.

CD analysis: The CD analysis for determination of melting temperatures and folding reversibility was performed as described in Example 2.

Results

Protein expression and purification: All 45 subcloned Z variants could be expressed and the *in vitro* solubility for all purified variants was good. The purity was estimated by LC/MS to exceed 90 % for all variants. The correct molecular weights were verified by LC-MS.

- 5 CD analysis: CD spectrum measurements performed at 20°C confirmed the α -helical structure of the Z variants at this temperature. An overlay of the spectrums obtained after the variable temperature measurements (heating to 90°C followed by cooling to 20°C) on the spectrums obtained before the variable temperature measurement showed that all Z variants fold back completely, or nearly completely, to their α -helical structures after heating to 90°C (results not shown). The melting temperatures for a set of Z variants were determined from the variable temperature measurements and are shown in Table 5.

Table 5. Melting temperatures of matured Z variants with a histidine tag fused directly to the amino terminus of SEQ ID NO:497 and SEQ ID NO:499-508.

Z variant	SEQ ID NO:# of Z variant	T _m (°C)
His ₆ -Z06175	SEQ ID NO:497	44
His ₆ -Z05998	SEQ ID NO:499	45
His ₆ -Z06009	SEQ ID NO:500	45
His ₆ -Z06079	SEQ ID NO:501	46
His ₆ -Z06126	SEQ ID NO:502	44
His ₆ -Z06140	SEQ ID NO:503	42
His ₆ -Z06189	SEQ ID NO:504	47
His ₆ -Z06214	SEQ ID NO:505	44
His ₆ -Z06215	SEQ ID NO:506	41
His ₆ -Z06226	SEQ ID NO:507	44
His ₆ -Z06018	SEQ ID NO:508	46

Example 6: *In vitro* characterization of C5 binding Z variants

Materials and methods

Cloning and protein production: DNA encoding a subset of C5 binding Z variants (SEQ ID NO:745-757) where *E. coli* codon optimized and synthesized by GeneArt, GmbH. The synthetic genes representing the C5 binding Z variants were subcloned and expressed in *E. coli*. The expression vectors encoding constructs of monomers or dimers of Z variants optionally linked to an albumin binding domain (ABD094, SEQ ID NO:759) are schematically illustrated in Figure 4.

Intracellularly expressed Z variants were purified using conventional chromatography methods. Homogenization and clarification was performed by sonication followed by centrifugation and filtration. Anion exchange chromatography was used as capture step. Further purification was obtained by hydrophobic interaction chromatography. The purifications were executed at acidic conditions (pH 5.5). Polishing and buffer exchange was performed by size exclusion chromatography. Before concentration to final protein content, the endotoxin level was reduced by polymyxin B affinity chromatography. Produced proteins were analyzed by MALDI-TOF MS and on SDS-PAGE.

In addition, recombinantly expressed OmCI protein (SEQ ID NO:761) was used as a reference molecule in the *in vitro* studies.

Inhibition of hemolysis: For studies of classical complement pathway function and inhibition thereof by C5 binding polypeptides, sheep erythrocytes were prepared from fresh sheep whole blood in Alsever's solution (Swedish National Veterinary Institute) and thereafter treated with rabbit anti-sheep erythrocyte antiserum (Sigma) to become antibody sensitized sheep erythrocyte (EA). The whole process was conducted under aseptic conditions. All other reagents were from commercial sources.

The *in vitro* assay was run in 96-well U-form microtiter plate by consecutive additions of a test protein, a complement serum and EA suspension. The final concentrations of all reagents, in a total reaction volume of 50 μ l per well and at pH 7.3-7.4, were: 0.15 mM CaCl_2 ; 0.5 mM MgCl_2 ; 3 mM NaN_3 ; 138 mM NaCl ; 0.1% gelatin; 1.8 mM sodium barbital; 3.1 mM barbituric acid; 5 million EA; complement protein C5 serum at suitable dilution, and C5 binding Z variant at desired concentrations. Different species of complement sera were used in the assay to define cross-species potencies of the Z variants. For mouse serum, a C5 depleted human serum (C5D from Quidel cat. no. A501) had to be supplemented in an equal amount.

The Z variants were pre-incubated with the above described complement serum for 20 min on ice prior to starting the reaction by the addition of EA suspension. The hemolytic reaction was allowed to proceed at 37°C during agitation for 45 min and was then optionally ended by addition of 100 μ l ice-cold saline containing 0.02% Tween 20. The cells were centrifuged to the bottom and the upper portion, corresponding to 100 μ l supernatant, was transferred to a transparent microplate having half-area and flat-bottom wells. The reaction results were analyzed as optical density using a microtiter plate reader at a wavelength of 415 nm.

On all test occasions, controls, vehicle and OmCI (SEQ ID NO:761), were included in each plate to define the values of uninhibited and fully inhibited reactions, respectively. These values were used to calculate the % inhibition of the complement hemolysis at any given sample concentration. The inhibitory potencies (IC_{50} values) of tested Z variants were defined by applying the same assay in the presence of a controlled concentration of human C5 added to C5

depleted serum. For highly potent inhibitors (low nanomolar to sub-nanomolar), a final C5 concentration of the reaction mixture was controlled at 0.1 nM, which was optionally established by using C5 depleted or deficient sera.

In vitro kinetics and affinity of C5 binding Z variants to immobilized hC5: The binding affinity of a number of C5 binding Z variants (SEQ ID NO:748-757) to hC5 were analyzed using a Biacore T200 instrument (GE Healthcare). Human C5 (A403, Quidel Corporation) was coupled to a CM5 sensor chip (900 RU) using amine coupling chemistry according to the manufacturer's protocol. The coupling was performed by injecting hC5 at a concentration of 7.5 µg/ml in 10 mM Na-acetate buffer pH 5 (GE Healthcare). The reference cell was treated with the same reagents but without injecting human C5.

All experiments were performed in 10 mM HEPES pH 7.4, 150 mM NaCl, 3 mM EDTA, 0.005% Surfactant P20 (HBS-EP buffer, GE Healthcare). For kinetic analyses, the flow rate was 30 µl/min and data were collected at 25°C. Data from the reference cell were subtracted to compensate for bulk refractive index changes. In most cases, an injection of HBS-EP was also included as control so that the sensorgrams were double blanked. The surfaces were regenerated in HBS-EP buffer.

Binding of Z variants to immobilized hC5 was studied with the single cycle kinetics method, in which five concentrations of sample are injected one after the other in the same cycle without regeneration between injections. Kinetic constants were calculated from the sensorgrams using the Langmuir 1:1 or bivalent analyte model of Biacore T200 Evaluation Software version 1.0.

In vitro kinetics and affinity of C5 binding Z-ABD molecules to immobilized hC5: Binding of Z-ABD molecules (SEQ ID NO:748-757 fused to ABD094 (SEQ ID NO:759) by a GS linker), to immobilized hC5 was evaluated using a Biacore T200 instrument (GE Healthcare).

Z-ABD constructs where Z06175a (SEQ ID NO:753) as a monomer or dimer have been fused to ABD094 (SEQ ID NO:759) either in the N-terminus or the C-terminus via different linkers as specified in Figure 4 (constructs 2, 7, 5 and 4) were also pre-incubated with recombinant human albumin (Cell Prime rAlbumin AF-G, 9301, Novozymes), diluted and then injected over immobilized human C5 according to the single-cycle kinetics method as described above. As a comparison, the same constructs were injected in the absence of HSA. Two constructs, Z06175a-GS (Figure 4, construct 1) and Z06175a-GSGGGSGGGGS-ABD094 (Figure 4, construct 3) were only tested in the absence of HSA.

Steady state binding of C5 binding Z variants to C5 coated ECL plates: The affinity of a number of C5 binding constructs comprising Z variants (SEQ ID NO:745, SEQ ID NO:748-757 optionally fused to ABD094 (SEQ ID NO:759) in constructs as specified in Figure 4) to human C5 was measured by displacement of a ruthenium labeled C5 binding Z-ABD variant (SEQ ID NO:748 fused to SEQ ID NO:759 by a GS-linker).

The Z-ABD variant (SEQ ID NO:748 fused to SEQ ID NO:759 by a GS-linker) to be used as tracer was labeled at a molar ratio 1:12 to 1:20 (protein: SULFO-TAG NHS-Ester, Meso Scale Discovery, cat. no. R91AN-1). The labeling reaction was performed on ice for two hours. Unbound SULFO-TAG was removed using a ZebaTM spin desalting column (Thermo Scientific, cat no. 89889) and final protein concentration was measured by using Bradford reagent (Bradford, M.M., Anal. Biochem. 72: 248–254, 1976). The affinity (dissociation constant, K_D) of the SULFO-TAG labeled Z-ABD variant was determined by saturation binding analysis of increasing concentrations of the labeled Z-ABD variant to C5 coated electrochemoluminescence wells (ECL, Meso Scale Discovery). The labeled Z-ABD variant was further analyzed by LC/MS in order to determine the distribution of SULFO-TAG molecules on the Z-ABD variant.

Displacement was carried out by coating ECL, Multi-array 96-well high-bind, non-coated (Meso Scale Discovery, cat. no. L15XB) plates with 50 fmol/well hC5 over night at 4°C. Subsequently, non-specific sites were blocked with PBS with 1% Casein for two hours at RT. Different Z variants optionally fused with ABD094 (SEQ ID NO:759) (see Figure 4) were incubated at different concentrations along with approximately 100 pM of the SULFO-TAG labeled C5 binding Z-ABD variant in PBS with 1 % Casein. Incubation lasted three hours at RT while agitating the plate at 300 rpm. Finally, incubation was terminated by washing 3 times with 150 µl ice-cold PBS-Tween20. Immediately after the final wash, 150 µl 2x reading buffer (4x reading buffer T, Meso Scale Discovery cat. no. R92TC-3 diluted 1:1 in ultrapure H₂O) was added to each well and the signal was detected using a plate reader (SECTOR Imager 2400, Meso Scale Discovery). The naturally occurring C5 binding protein OmCI (Nunn *et al. supra*, SEQ ID NO:761) was included in the displacement assay as a positive control. Binding affinity of competing C5 binding constructs and controls to C5 was determined by non-linear regression analysis using Excel plugin XLfit5 and GraphPad Prism 4.

Selectivity of Z-ABD binding to C5 over C3, C4 and IgG: Binding of one Z-ABD variant (SEQ ID NO:748 fused to SEQ ID NO:759 by a GS-linker) to the closely related complement proteins C3 and C4 from human as well as binding to human IgG (since the origin of the Z-domain, Staphylococcal protein A, is an IgG binding protein) was addressed by surface plasmon resonance (SPR) using a Biacore 2000 instrument (GE Healthcare). The Z-ABD construct was immobilized on a CM5 chip (GE-Healthcare) using amine coupling (70 RU). 40 nM and 400 nM of each of human C3 (A401, Quidel), C4 (A402, Quidel) and IgG (I2511, Sigma) diluted in HBS-P buffer (GE Healthcare) were injected over the surface. Each injection was followed by a regeneration cycle with 20 mM NaOH injected for 30 s. Human C5 at the same concentrations was run in parallel as a positive control.

Results

Cloning and protein production: Produced protein variants as schematically described in Figure 4 where “Z” can be represented by SEQ ID NO:745 and SEQ ID NO:748-757 were analyzed by MALDI-TOF MS and on SDS-PAGE. (Figure 5)

- 5 Inhibition of hemolysis: A subset of C5 binding Z variants were assayed for C5 binding activity *in vitro* and inhibition of hemolysis in sheep erythrocytes. The concentration of Z variant resulting in 50 % inhibition of hemolysis (IC₅₀) or 50 % inhibition of tracer binding to human C5 was calculated. Representative concentration-response curves for Z variants shown as SEQ ID NO:745 and SEQ ID NO:748-757 inhibiting hemolysis as described in the methods section are shown in Figures 6A and 6B. The result for different Z variants fused to ABD094 (SEQ ID NO:759) via a short GS-linker are shown in Figure 6A.

The parental Z variant Z05477a (SEQ ID NO:745) fused to ABD094 (SEQ ID NO:759) separated by a short GS linker exhibited an IC₅₀ value of about 100 nM, whereas the tested second-generation C5 binding Z-ABD variants typically inhibited hemolysis with IC₅₀ values around or below 1 nM. This suggests a more than 100-fold increase in potency for the C5 binding Z variants identified in the maturation selection and subsequent screening.

In Figure 6B, C5 binding is shown for various combinations of one representative Z variant (Z06175a; SEQ ID NO:753) alone, as a dimer and in fusion with ABD094 (SEQ ID NO:759) either in the N-terminus or the C-terminus via different linkers as specified in the figure. The C5 binding combinations exhibited IC₅₀ values ranging from 86 pM to 12 nM with human serum as measured using the above described assay. The corresponding value for the tic protein OmCI was typically 300 to 500 pM.

In vitro kinetics: Kinetic studies of binding characteristics for a number of Z variants (SEQ ID NO:748-757) optionally fused to ABD094 (SEQ ID NO:759), to immobilized hC5, as well as to C5 in the presence of human albumin, were performed using the Biacore T200 instrument.

Data for ten different Z variants fused to ABD094 via a GS linker are presented in Table 6.

Table 6. Human C5-binding characteristics for different Z-ABD fusions

Construct	SEQ ID NO: # of Z variant	k _a (1/Ms)	k _d (1/s)	K _D (M)
Z-GS-ABD094	SEQ ID NO:748	6.93 x 10 ⁵	9.04 x 10 ⁻⁴	1.31 x 10 ⁻⁹
	SEQ ID NO:749	6.75 x 10 ⁵	1.23 x 10 ⁻³	1.83 x 10 ⁻⁹
	SEQ ID NO:750	7.65 x 10 ⁵	1.34 x 10 ⁻³	1.75 x 10 ⁻⁹
	SEQ ID NO:751	6.90 x 10 ⁵	1.29 x 10 ⁻³	1.87 x 10 ⁻⁹
	SEQ ID NO:752	7.02 x 10 ⁵	1.81 x 10 ⁻³	2.58 x 10 ⁻⁹

	SEQ ID NO:753	7.90×10^5	1.01×10^{-3}	1.18×10^{-9}
	SEQ ID NO:754	5.00×10^5	1.14×10^{-3}	2.28×10^{-9}
	SEQ ID NO:755	6.84×10^5	2.08×10^{-3}	3.05×10^{-9}
	SEQ ID NO:756	3.17×10^5	6.37×10^{-3}	2.01×10^{-9}
	SEQ ID NO:757	4.63×10^5	1.08×10^{-3}	2.34×10^{-9}

Binding of the same Z variant (SEQ ID NO:753) but in different constructs; i.e. with/without ABD and different linkers, were also analyzed using Biacore T200. In addition, the effect of albumin on some Z-ABD fusions was also assessed by running the same analysis in the absence and in the presence of human albumin. These data are presented below in Table 7.

Table 7. Human C5-binding characteristics for a Z-ABD fusion variant Z06175a (SEQ ID NO:753, abbreviated Z) comprised in different constructs.

Construct	Human albumin	k_a (1/Ms)	k_d (1/s)	K_D (M)
Z-GS-ABD094	-	7.37×10^5	1.06×10^{-3}	1.43×10^{-9}
Z-GS-ABD094	+	6.74×10^5	9.62×10^{-4}	1.43×10^{-9}
Z-Z-GS-ABD094	-	5.93×10^5	3.74×10^{-4}	6.30×10^{-10}
Z-Z-GS-ABD094	+	6.02×10^5	4.67×10^{-4}	7.76×10^{-10}
Z-GS-ABD094-GSGGGGSGGGGS-Z	-	8.69×10^5	5.75×10^{-4}	6.62×10^{-10}
Z-GS-ABD094-GSGGGGSGGGGS-Z	+	6.55×10^5	3.83×10^{-4}	5.86×10^{-10}
Z-Z-GSGGGGSGGGGS-ABD094	-	4.59×10^5	6.32×10^{-4}	1.38×10^{-9}
Z-Z-GSGGGGSGGGGS-ABD094	+	8.32×10^5	9.39×10^{-4}	1.13×10^{-9}
Z-GS	-	2.42×10^6	1.40×10^{-3}	5.79×10^{-10}
Z-GSGGGGSGGGGS-ABD094	-	3.64×10^5	1.37×10^{-3}	3.75×10^{-9}

Surprisingly small effects could be seen when comparing the affinities of the constructs for hC5 (SEQ ID NO:760) in the presence and absence of albumin. This suggests that simultaneous binding of albumin to the ABD moiety of the constructs does not interfere with C5 interaction.

Steady state binding of C5 binding Z variants to C5 coated ECL plates: Steady state binding of C5 binding constructs composed of different Z variants (SEQ ID NO:745 and 748-757), optionally fused to ABD094 (SEQ ID NO:759) in constructs as specified in Figure 4, to hC5 was assessed in a competition assay. By competing for binding to C5 coated on ECL plates with a SULFO-TAG labeled C5 binding Z variants (SEQ ID NO:748) fused to ABD (SEQ ID NO:759), steady state binding of the C5 constructs was evaluated. As a comparison the protein OmCI (SEQ ID NO:761) was also included. The labeled Z-ABD variant containing SEQ ID NO:748 had an affinity (K_d) of 0.9 nM for hC5. This labeled Z-ABD variant was further found to bind to an antibody specific for the constant region of Z variants in a concentration-dependent manner with a K_d of 0.34 nM.

The C5 binding Z-variants (SEQ ID NO:748-757) fused in the carboxy terminus to ABD094 (SEQ ID NO:759) by a GS linker were found to displace 200 pM SULFO-TAG labeled Z-ABD variant with IC_{50} values ranging from about 300 pM to 1 nM (Figure 7A), whereas the corresponding construct containing the parental Z variant Z05477a (SEQ ID NO:745) exhibited an affinity IC_{50} value of about 30 nM. In contrast, the naturally occurring C5 binding protein OmCI was found to bind hC5 with an IC_{50} of 1.5 nM (Figure 7A). Thus, all the tested second-generation Z variants (SEQ ID NO:748-757) exhibited a higher binding affinity for human C5 than the parental Z variant Z05477a (SEQ ID NO:745). In addition, the affinities were higher than that of OmCI binding to human C5 using the same method.

A number of different constructs containing the same C5 binding domain as a monomer, dimer, with or without ABD as well as a few different linkers between the different domains were also tested (Figure 7B). Monomeric variants of Z06175a (SEQ ID NO:753, optionally fused to a His₆-tag or a C-terminal ABD) and the dimeric variants with a C-terminal ABD linker were found to displace 200 pM SULFO-TAG labeled Z-ABD variant with IC_{50} values ranging from about 500 pM to 1.7 nM whereas the dimeric variant without an ABD and the monomeric variant with a N-terminal ABD displaced 200 pM SULFO-TAG labeled Z-ABD with IC_{50} values of 4 nM and 17 nM, respectively.

Selectivity: Selectivity was addressed using SPR analysis and the surface with the immobilized Z-ABD variant (SEQ ID NO:748 fused to SEQ ID NO:759 by a GS-linker) displayed no significant SPR signal when subjected to 40 and 400 nM of the C5 paralogs human C3 and C4 as well as human IgG. As a comparison, 400 nM human C5 elicited an SPR response of about 450 RU showing that the tested Z-ABD variant indeed is selective for C5 over C3, C4 and IgG.

Example 7: Interaction studies of Z-ABD variants with HSA, BSA and serum album from rat and mouse.

Materials and methods

5 Two different methods, size exclusion chromatography and Biacore, were used to study the interaction between the albumin binding domain ABD094 fused to a C5 binding Z variants.

Size exclusion chromatography (SEC) was employed to study the interaction between Z06175a-GS-ABD094 (SEQ ID NO:753 fused to SEQ ID NO:759 by a GS linker) and HSA. Briefly, equimolar amounts of Z06175a-GS-ABD094 and recombinant HSA (Novozymes) were
10 preincubated in PBS at room temperature for 60 minutes and subsequently run on a Superdex200 column (GE Healthcare) using the SMART system (GE Healthcare). Z06175a-GS-ABD094 and HSA were also run separately as controls.

Binding to immobilized albumin was studied using a Biacore 2000 instrument (GE Healthcare). Recombinant human albumin (Recombunin[®], Novozymes) was coupled to a CM5
15 sensor chip (385 RU) using amine coupling chemistry as described by the manufacturer. The coupling was performed by injecting human albumin in 10 mM Na-acetate buffer pH 4.5 (GE Healthcare). The reference cell was treated with the same reagents but without injecting human albumin. Injection of HBS-EP was also included as control so that the sensorgrams were double blanked. Experiments were performed in HBS-EP buffer, 10 mM glycine-HCl pH 2 (GE
20 Healthcare) was used for regeneration, the flow rate was 30 µl/min and data were collected at 25°C. Two different constructs were tested, Z-ABD (Z06175a-GS-ABD094) and Z-ABD-Z (Z06175a-GS-ABD094-GSGGGGSGGGGS-Z06175a) at three different concentrations; 25 nM, 100 nM and 400 nM. BIAevaluation version 4.1.1 was used for evaluation of sensorgram data. In a similar fashion, binding of Z-ABD (Z06175a-GS-ABD094) to surfaces immobilized with
25 serum albumin from rat (A4538, Sigma), mouse (A3559, Sigma), and cow (BSA, Sigma) was also investigated.

Results

On a SEC column, larger molecules elute faster than small. As seen in Figure 8A, the co-
30 injected HSA+ Z06175a-GS-ABD094 elute faster than when HSA is injected alone suggesting that the two molecules behave as a stable complex under these conditions. The smaller Z06175a-GS-ABD094 elute slower than either the complex or HSA alone showing that these proteins alone are smaller than the complex.

Biacore 2000 data for the analyzed Z-ABD and Z-ABD-Z variants show that the Z-
35 ABD has a faster on-rate than when ABD is flanked by Z-domains on either side (Figure 8B). Analysis of the binding affinity of ABD fused Z domains points at an affinity below 1 nM for Z-ABD whereas the Z-ABD-Z variant bind to immobilized HSA with a K_D above 1 nM.

Z06175a-GS-ABD094 bound to rat serum albumin with very high affinity ($K_D < 100$ pM) whereas the interaction with immobilized mouse serum albumin was weaker (K_D of about 4 nM) than both with human and rat serum albumin. Interaction with bovine serum albumin was not measureable.

5 These data agree well with published data on an earlier variant of ABD (Jonsson *et al.* Protein Engineering, Design & Selection 2008, 21: 515-527) and show that the tested Z-ABD variant is strongly bound to serum albumin in human at clinically relevant concentrations as well as in mouse and rat allowing comparisons of pharmacokinetic data between animals and humans.

10

Example 8: Pharmacokinetic studies of C5 binding Z variant in rats

Materials and methods

Rodent in-life phase: The pharmacokinetics of two C5 binding constructs Z-ABD (Z06175a-GS-ABD094; SEQ ID NO:753 fused to SEQ ID NO:759 by a GS linker, Figure 4, construct 2) and Z-ABD-Z (Z06175a-GS-ABD094-GSGGGSGGGGS-Z06175a; (SEQ ID NO:753 fused to SEQ ID NO:759 by a GS linker, followed by a GS(G₄S)₂ linker and a second SEQ ID NO:753 motif, Figure 4, construct 5) was studied in Male Sprague Dawley (SD) rats (250-300 g body weight). Each rat was given a single dose administration, i.v. (250 nmol/kg) or s.c. (500 nmol/kg), of Z-ABD or A-ABD-Z (n=3 per dose group). Blood samples (200 µL) were drawn at 5, 20, and 45 min, as well as 1.5, 4, 7, 24, 48, 72, 120, 168, 240, and 336 h following administration for the i.v. group and at 15 and 30 min, 1, 2, 4, 7, 24, 48, 72, 120, 168, 240, and 336 h following administration for the s.c. group. Blood was collected in tubes and placed in the fridge for 20 min to allow clotting. Serum was subsequently harvested following centrifugation at 4000 rpm for 10 minutes. Serum samples were kept at -70°C pending analysis.

Determination of C5 binding Z variant concentrations in serum samples from animals using LC/LC/MS/MS: Serum concentrations of the administrated C5 binding constructs Z-ABD and Z-ABD-Z, as described above, were determined by mass spectrometry (LC/LC/MS/MS).

Scrum or plasma samples (25 µl) were diluted with 150 µl of a pepsin agarose (7 mg/ml, Sigma, cat. no. P0609) suspended in 1 M ammonium formate buffer pH 3.0 in a 500 µl Eppendorf tube. The tubes were capped and agitated in an Eppendorf thermomixer compact at 37°C for 20 min. Following agitation, 25 µl of an internal standard solution I(¹³C₆,¹⁵N)NKLDDDPSSSEL (amino acids 31-44 of the SEQ ID NO:746-757) (Thermo Fisher Scientific GmbH), diluted to 0.5 µM in 0.1 % trifluoroacetic acid (TFA), was added. Following addition of internal standard, the samples were mixed and filtered through 0.45 µm cellulose spin filters (Grace).

Standard samples for calibration were prepared by weighing 20 µl of protein stock solution with known protein concentration (5-10 mg/ml) followed by dilution with blank plasma from the species to be analyzed. The first stock plasma standard (3 µM) was diluted further down to 0.1 µM.

5 40 µl of the samples were injected into a coupled column system followed by tandem mass spectrometry with multiple reaction monitoring (MRM). The first column was an Ascentis RP-Amide column packed with 5 µm particles (2.1 x 150 mm, Supelco). An enrichment column; a Brownlee newgard column (3.2 x 15 mm) packed with 7 µm C18 particles, was used to trap the analyte peptide fraction from the first column. The effluent from the first column was
10 diluted with 1 ml/min water pumped by Shimadzu pump into a whirl mixer (Lee Scientific). The last column was a mixed mode reversed phase and cation exchange column (2.1 x 100 mm) packed with 5 µm particles Primesep 100 (SIELC Inc).

 The mobile phases for the first column (RP-Amide) provided on a first liquid chromatograph (Acquity UPLC) were A: 2 % acetonitrile, 0.1 % acetic acid, 0.1 % TFA, and
15 97.8 % water, and B: acetonitrile with 0.1 % acetic acid and 0.02 % TFA. The flow was 0.5 ml/min and a linear gradient was used for elution. The sample was eluted at isocratic conditions with 100 % A for 1 min, followed by 80 % A at 7.9 min. At 8.1 min, the column was washed with 100 % B for one minute, followed by reconditioning with 100 % A. The effluent from the column was connected to a Valco six port valve controlled from the mass spectrometer
20 software.

 The trap column (3.2 x 15 mm) was connected to the six port valve in back flush mode. The mobile phases for the second column, provided on a second liquid chromatograph (Agilent 1100), were A: 80 % acetonitrile, 19.9 % water, and 0.1 % formic acid, and B: 80 %
25 acetonitrile, 19 % water, 0.5 % acetic acid and 0.5 % TFA pumped by an Agilent 1100 liquid chromatograph at 0.5 ml/min and eluted with the following gradient: 100% A during the first 5 minutes followed by B gradually being raised from 0 to 40 % from 5 to 10 minutes followed by a raise to 100 % B during the next 6 seconds (10 to 10.1 minutes). B was kept at 100 % until 11.5 minutes followed by a drop to 0 % (100 % A) during the next 6 seconds (11.5 to 11.6 minute) and kept at 0 % B throughout the cycle until stopped at 13 minutes.

30 The effluent from the last column was connected to a triple quadrupole mass spectrometer (Sciex API 4000) equipped with an electrospray ion source operated in positive ion mode. The MRM transitions were 780.9>814.4 for the analyte and 784.5>821.4 for the internal standard. The declustering potential was optimized at 55 V and the collision energy to 35 V. The effective collision energy was 70 eV since the precursor ion was doubly charged
35 giving a singly charged fragment ion. The peak area ratios between the analyte and internal standard were used for quantification. Linear calibration curves were obtained with a recovery of 85 % and a limit of quantification of about 40 nM.

Ex vivo hemolysis: An *ex vivo* hemolytic assay for complement activation was performed in order to optimally assemble *in vivo* conditions for the serum samples from the above described *in vivo* studies. The serum samples were 5x diluted in a total reaction volume of 25 μ l/well comprising 5 million antibody sensitized sheep erythrocytes (EA). In general, a portion of 20 μ l EA suspension containing all other components (see Example 6) was mixed (agitation 10 minutes) with 5 μ l serum sample to initiate the hemolytic activation at 37°C. For mouse serum samples, such as in example 11, however, 1 μ l C5D had to be included in the 20 μ l EA suspension. The *ex vivo* assay was performed essentially as described for the *in vitro* assay of Example 6.

Calculations: Evaluation of the pharmacokinetic parameters was based on individual serum concentration data, the mean (\pm stdev) is reported for each dose group. Levels below lower limit of quantitation (LLOQ) appearing at terminal sampling points were omitted from the pharmacokinetic analysis. Maximum serum concentration, C_{max} , and time to observed maximum serum concentration, t_{max} , were obtained directly from the serum concentration data. The pharmacokinetic parameters; area under curve (AUC, $AUC_{0-\infty}$ and AUC_{0-last} calculated by the linear trapezoidal method), subcutaneous bioavailability (F, calculated as $(AUC_{sc}/AUC_{iv}) \cdot (Dose_{iv}/Dose_{sc})$), terminal serum half-life ($T_{1/2z}$, calculated as $\ln 2/\lambda_z$ where estimation of terminal slope, λ_z , was based on at least 4 $C=f(t)$ observations), mean residence time (MRT, calculated as $AUMC/AUC$), serum clearance (CL, calculated as $Dose/AUC_{0-\infty}$), volume of distribution at steady state (V_{ss} , calculated as $CL \cdot MRT$) and volume of distribution at the terminal phase (V_z , calculated as CL/λ_z) were calculated using WinNonlin software version 5.2.1 (Pharsight Corp., USA), Non-Compartmental-Analysis.

Results

The pharmacokinetic data for Z-ABD and Z-ABD-Z following i.v. (250 nmol/kg) and s.c. (500 nmol/kg) administration are summarized in Table 8. Z-ABD was quantifiable in serum up to 10-14 days post dose in the i.v. group and 14 days in the s.c. group whereas Z-ABD-Z was quantifiable in serum up to 10 days post dose in both dose groups (Figure 9). 14 days was the final sampling time point.

Correlating the serum concentration of C5 binding polypeptide with the amount of hemolysis in sheep erythrocytes, it was found that full inhibition of hemolysis under the conditions described (e.g. serum dilution 1:5) was obtained by Z-ABD at serum concentrations above 1 μ M (Figure 12) whereas Z-ABD-Z reached full inhibition at serum concentrations around 0.5 μ M (Figure 13). Surprisingly, as seen in Figure 9 and Table 8, Z-ABD has a lower serum clearance, a longer terminal serum half-life and a higher bioavailability than Z-ABD-Z. In terms of time this lead to full inhibition of hemolysis for about three days after administration of 250 nmol/kg Z-ABD (Figure 10) i.v. or 500 nmol/kg s.c (Figure 11) to S.D.rats.

Table 8. Mean (\pm stdev) pharmacokinetics of Z-ABD and Z-ABD-Z following i.v. and s.c. administration in male Sprague Dawley rats.

		Z-ABD		Z-ABD-Z	
Administration route		i.v.	s.c.	i.v.	s.c.
Dose	nmol/kg	250	500	250	500
C _{max}	μ M		2.8 (0.2)		0.90 (0.10)
T _{max}	h		18 (9.8)		17 (12)
AUC _{0-∞}	μ M*h	233 (34)	252 (11)	79 (7.5)	64 (1.2)
AUC _{0-last}	μ M*h	226 (37)	247 (11)	79 (6.9)	63 (1.0)
F	%		55 (3.1)		41 (2.6)
T _{1/2,z}	h	58 (4.6)	57 (4.2)	36 (0.6)	46 (1.2)
MRT	h	69 (2.6)	80 (4.6)	27 (1.5)	63 (2.6)
CL	mL/h*kg	1.1 (0.2)		3.2 (0.2)	
V _{ss}	mL/kg	73 (12)		83 (10)	
V _z	mL/kg	90 (18)		159 (12)	

5 Example 9: Pharmacokinetic studies of C5 binding Z variants in monkey

Materials and methods

The study in life phase was performed at Charles River, Nevada (www.criver.com), formulation of administered drug and analysis of serums samples were performed *in house*. The pharmacokinetics of a Z-ABD variant (Z06175a (SEQ ID NO:753) fused to ABD094 (SEQ ID NO:759) by a GS linker) was investigated in the male Cynomolgus monkey (n=3) following i.v. (intravenous) and s.c. (subcutaneous) administration. Evaluation of the pharmacokinetic parameters was performed according to Example 8, however following i.v. administration the initial serum half-life ($T_{1/2\alpha}$) corresponding to the initial slope of the log-linear serum concentration-time curve, intermediate serum half-life ($T_{1/2\beta}$) corresponding to the slope of the log-linear serum concentration-time curve associated with the secondary (intermediate) phase and terminal serum half-life ($T_{1/2\gamma}$) corresponding to the terminal slope of the log-linear serum concentration-time curve was determined. $T_{1/2}$ was calculated as $\ln 2/\lambda$ where estimation of the slope, λ , was based on at least 4 $C=f(t)$ observations. The pharmacokinetic data presented for sc administration are compensated for pre-dose levels of Z-ABD while the graph displaying serum concentration versus time after sc administration show the actual serum concentrations determined. The monkeys were 2-4 years old with a body weight of 2.3-3 kg. Each monkey received a single i.v. dose (540 nmol/kg) followed by a single s.c. dose (1635 nmol/kg) three

weeks after the i.v. administration. Blood samples were taken at 10 and 30 minutes and 1, 2, 4, 8, 24, 48, 72, 120, 168, 240, 336 and 504 hours post dose following both administrations. The blood samples were allowed to clot for 20-40 minutes in room temperature and then centrifuged at 1500 to 2200 RCF at 2-8°C for 10-15 minutes before the serum was harvested and frozen.

5 The serum samples were stored at a temperature below -20°C until analysis.

Serum concentrations of Z-ABD were analyzed by LC/LC/MS/MS as described in Example 8. Serum concentrations determined by LC/LC/MS/MS were also confirmed by a quantitative sandwich enzyme immunoassay technique. A polyclonal antibody specific for the Z compartment of Z-ABD was coated on to a microplate. Unbound polyclonal antibody was
 10 washed away and casein was added as blocking agent to reduce unspecific binding to the plastic surface. Samples and standards were diluted in PBS containing 0.5% casein and between 1-5% monkey normal serum. After washing away unbound casein, standards and samples were pipetted to the wells allowing any Z-ABD, presumed mainly to be associated with serum albumin, present in the sample to bind to the immobilized antibody. After washing away any
 15 unbound Z-ABD, an HRP labeled polyclonal antibody specific for albumin was added to detect the immobilized Z-ABD-albumin complex by colorimetric methods. Unbound polyclonal antibody was washed away and a substrate solution was added to the wells and color develops in proportion to the amount of Z-ABD bound. Evaluation and calculation of pharmacokinetic parameters were performed as described in Example 8.

20 *Ex vivo* hemolysis in serum from cynomolgus monkeys dosed with above described Z-ABD variant was monitored using the method described in Examples 6 and 8 with the modification that the monkey serum was diluted only two-fold compared to five-fold for rodent serum.

25 *Results*

Data on the mean (\pm stdev) pharmacokinetics of each dose group are presented. Serum concentrations of Z-ABD were quantifiable at all time points following both i.v. and s.c. administration by LC/LC/MS/MS (Figure 14). ELISA data and LC/LC/MS/MS data correlated linearly by a coefficient of 0.986 but LC/LC/MS/MS data were used for the calculations.

30 Following i.v. administration of Z-ABD the initial serum half-life was 9.1 (0.8) hours, intermediate serum half-life was 84 (4) hours and the terminal serum half-life was 198 (51) hours. The mean residence time was 246 (62) hours. The volume of distribution, V_{ss} and V_z was calculated to 110 (23) ml/kg and 127 (27) ml/kg respectively and clearance was estimated to 0.45 (0.02) mL/h*kg.

35 Following s.c. administration, , and corrected for pre-dose serum levels remaining from the i.v. administration, maximum serum concentrations (mean C_{max} 21(3) μ M) were reached at

8-24 h after dose. The terminal serum half-life was 206 (40) hours and the mean residence time was 250 (68) hours. The subcutaneous bioavailability was estimated to be above 70%.

The pharmacodynamic effect of the injected Z-ABD variant (Z06175a (SEQ ID NO:753) fused to ABD094 (SEQ ID NO:759) by a GS linker) was monitored by hemolysis. The hemolytic effect in cynomolgus monkey was completely suppressed (≤ 20 % of pre-dose) for at least seven days after administration of 5 mg/kg Z-ABD i.v. and 15 mg/kg Z-ABD s.c.

Example 10: *In vivo* studies using Zymosan induced peritonitis

10 *Materials and methods*

Administration to mice: C57BL/6 female mice received different concentrations of a Z-ABD fusion molecule (Z06175a-GS-ABD094, SEQ ID NO:753 fused to SEQ ID NO:759 by a GS linker) or the positive control OmCI intraperitoneally (i.p.) 1 hour before induction with zymosan, or subcutaneously (s.c.) 18 hours before induction with zymosan.

15 0.8 mg/mouse zymosan was administered i.p. 1 hour later orbital blood samples (in serum vials with coagulation activator) were taken under isoflurane anaesthesia. The animals were killed by cervical dislocation. A skin incision was made, and the abdominal muscular wall was visualized. PBS solution (including 2 mM EDTA) was gently injected into the abdominal cavity. The abdomen was massaged and a sample of fluid (1-2 ml) was withdrawn. The samples were transferred to test tubes and stored on wet ice before centrifugation at 600 g for 10 min. Total protein and C5a concentrations in the supernatant were analyzed.

Blood samples were kept in a refrigerator for at least 30 min and centrifugation was thereafter performed at 2000 g. Serum samples were stored in freezer (-70°C) for later analysis of hemolytic activity and levels of Z06175a-GS-ABD094.

25 Analysis of hemolysis activity in serum samples from animals: Analysis of hemolysis activity was performed according to the hemolysis assay described in Examples 6 and 7.

Analysis of C5a concentration in lavage from mice dosed with zymosan and C5 binding Z-ABD fusion molecules: For detection of C5a in mouse peritoneal lavage samples, microtiter plates (MaxiSorp, Nunc) were coated overnight at 4°C with 100 μ l/well of anti-C5a antibody (cat. no.

30 MAB21501, R&D Systems) at a concentration of 1 μ g/ml in 0.05 M sodium carbonate-bicarbonate buffer, pH 9.6 (cat. no. C-3041, Sigma). The plates were washed three times with PBS containing 0.05 % Tween 20 (PBST, cat. no. 09-9410-100, Medicago) and blocked with 200 μ l/well of 1% BSA (cat. no. A7030, Sigma) in PBST for 1-1.5 h at RT during agitation at 450 rpm. The plate was again washed three times with PBST and then incubated with 100 μ l/well of recombinant mouse C5a standard (cat. no. 2150-C5, R&D Systems) at various concentrations in PBST with 0.1% BSA or samples for 2 h at RT during agitation at 450 rpm. High concentration samples were also diluted in PBST with 0.1% BSA. The plate was once

again washed three times with PBST and then incubated with 100 µl/well of biotinylated anti-C5a antibody (cat. no. BAF2150, R&D Systems) at a concentration of 0.1 µg/ml for 1.5 h at RT while shaking the plate at 450 rpm. Following 3x washing with PBST, the plate was incubated with 100 µl/well of streptavidin-HRP (cat. no. DY998, R&D Systems) at a 200 fold dilution in blocking buffer for 20 min at RT during agitation at 450 rpm. After three final washes, the plate was developed with 100 µl/well TMB substrate (cat. no. T0440, Sigma) and read after 20-30 min at 650 nm using a Spectramax Plus plate reader (Molecular Devices).

A standard curve was constructed by plotting the absorbance at 650 nm for each standard against its concentration (range 0-4000 pg/ml).

- 10 Determination of Z variant concentration in serum samples from animals using ECL: Serum concentrations of administrated C5 binding Z06175a-GS-ABD094 (SEQ ID NO:753 fused to SEQ ID NO:759 by a GS linker) and Z06175a-GS-ABD094-GSGGGGSGGGGS-Z06175a (SEQ ID NO:753 fused to SEQ ID NO:759 by a GS linker, followed by a GS(G₄S)₂ linker and a second SEQ ID NO:753 motif, see Figure 4 for construct description) were determined by ECL.
- 15 Multi-array 96-well high-bind, non-coated (Meso Scale Discovery cat. no. L15XB) plates were coated with a goat anti-Affibody molecule Ig (Affibody AB, cat. no. 20.1000.01.0005).

- In similarity with Example 6, a Z-ABD variant (Z06009a, SEQ ID NO:748 fused to ABD094, SEQ ID NO:759 Multi-array plates were coated with the goat anti-Affibody molecule IgG (Affibody AB) overnight at 4°C, and subsequently non-specific sites were blocked with PBS with 1 % Casein for two hours at RT.

- Meanwhile, serum samples were thawed from -70°C and diluted in PBS with casein in serum from the same animal strain. Standards and controls were diluted in the corresponding buffer. Samples and standards were incubated for three hours at RT while shaking the plate at 300 rpm. Incubation was terminated by washing 3x 150 µL ice-cold PBS-Tween20.
- 25 Immediately after the final wash, 150 µl 2x reading buffer (4x reading buffer T, Meso Scale Discovery cat. no. R92TC-3 diluted 1:1 in ultrapure H₂O) was added to each well and the signal was detected using a plate reader (SECTOR Imager 2400, Meso Scale Discovery).

- In an alternative experiment, plates were coated with human C5 (SEQ ID NO:760, 1 pmol/well). Prior to addition to the coated plate, serum samples and standards, diluted in serum or in serum and PBS with casein (all samples and standards were matched to the same serum concentration), were heated to 60°C for 30 min in order to denature endogenous C5. This alternative experiment provided a method for exclusive detection of C5 binding proteins, whereas the antibody dependent strategy described above can be applied to all proteins binding to that particular antibody.

Results

Analysis of serum concentrations of Z-ABD and hemolysis activity in serum samples from

animals: The serum concentrations as well as the ability to affect hemolysis in sheep

- 5 erythrocytes of the Z-ABD fusion molecule (Z06175a-GS-ABD094, SEQ ID NO:753 fused to SEQ ID NO:759 by a GS linker)) was assessed after administration of a low (20 nmol/kg), medium (100 nmol/kg) and high dose (500 nmol/kg). The serum concentrations were relatively linear with dose, and inhibition of hemolysis confirmed that the molecules in serum were active and that the inhibition of hemolysis indeed also was concentration dependent.
- 10 Analysis of C5a concentration in lavage from mice dosed with zymosan and C5 binding Z-ABD fusion molecules: The pro-inflammatory molecule zymosan was administered i.p. and in Figure 15 the effect on the highly inflammatory C5 cleavage product C5a in lavage as a function of zymosan dosing alone and zymosan dosed after a dosing of a C5 binding Z variant at 20, 100 and 500 nmol/kg administered s.c. 18 h before zymosan treatment or OmCI administered i.p. 1
- 15 h before zymosan treatment, is shown. Zymosan administration alone leads to a potent elevation of C5a in the lavage. This effect is blocked in a dose dependent manner by the presented C5 binding Z-ABD fusion molecule.

Example 11: Pharmacokinetic studies of C5 binding protein in mice following intratracheal

- 20 administration

Materials and methods

The pharmacokinetic profile of the C5 binding construct Z06175a-GS-ABD094 (SEQ ID NO: 753 fused to SEQ ID NO:759 by a GS linker) following intratracheal administration to female

25 C57bl mice was studied. Temperature, relative humidity and lighting was set to maintain $22 \pm 1^\circ\text{C}$, $55 \pm 5\%$ and a 12 h light – 12 h dark cycle and diet and water was provided ad libitum.

Animals were anesthetized with isoflurane and dosed directly into the lungs using a microspray with 500 nmol/kg Z06175a-GS-ABD094. As much blood as possible was drawn, under anesthesia by isoflurane, from vena cava at 5 min, 30 min, 1 h, 3 h, 7 h, 16 h, 24 h, 48 h and 72

30 h (three animals/time point) for preparation of serum samples. Serum samples were prepared by collecting blood in tubes and placing the tubes in the fridge for 20 min. Subsequently, the tubes were centrifuged at 4000 rpm for 10 minutes. A minimum of 100 μl serum was prepared from each blood sample. Serum samples were kept at -70°C prior to analysis. Serum concentrations of Z06175a-GS-ABD094 in each sample was determined by ECL as described in Example 10

35 and the ability of serum samples to affect hemolysis in sheep erythrocytes was determined as described in Examples 6 and 8.

Results

The serum concentration in each sample and the corresponding ability to affect hemolysis in sheep erythrocytes are described in Figure 16A and Figure 16B, respectively. Within 30 minutes, a plateau is reached with serum concentrations ranging from 300 to 1000 nM where hemolysis is nearly completely blocked. In serum sampled at time points later than 7 h post-administration, hemolysis is gradually reoccurring. At the final time point three days after dosing, hemolysis was about 70 % of control (Figure 16B). These data clearly demonstrate absorption of Z06175a-GS-ABD094 into the systemic circulation following intratracheal administration and that the molecule functionally inhibits hemolysis.

Example 12: Pharmacokinetic studies of C5 binding Z variant in rabbit eye following topical and intravitreal administration

Materials and methods

Rabbit in-life phase: The pharmacokinetics of a Z variant (Z06175a, SEQ ID NO:753 followed by GS (Figure 4, construct 1)) was studied in rabbit eye following intra-vitreous administration.

The study in-life phase and dissection of eyes from dosed animals (pigmented rabbits, 2 - 2.5 kg) was performed at Iris Pharma, La Gaude, France (www.iris-pharma.com). Animals were housed individually at 20 ± 2 °C at 55 ± 10 % relative humidity with access to food and water *ad lib*.

Animals were divided in three groups: 1) intravitreal administration (50 µl in each eye, n=3, six eyes totally) followed by dissection and serum sampling after one day, 2) intravitreal administration (50 µl in each eye, n=3) followed by dissection and serum sampling after four days and 3) untreated animals (n=5).

Four distinct eye compartments were dissected (aqueous humor, vitreous, neuro-retina and RPE-choroid) and immediately frozen at -80 °C. Formulation of administered drug (20.2 mg/ml in 10 mM phosphate buffer, 145 mM NaCl, pH 7.4) and analysis of drug in various eye compartments were performed *in house*.

Analysis of Z-variant in dissected eye compartments: Dissected eye compartments were shipped on dry ice and stored at -80 °C until analysis. The retina and choroid samples were thawed in 10 times (volume/weight) PBS containing 1 % human serum albumin in Lysing Matrix D tubes (MP Biomedical) containing ceramic beads and agitated at speed 4 for 2 x 20 s in a Savant Bio 101 homogenizer. The homogenate was removed from the beads using a pipette and transferred to a 1.5 ml Eppendorf tube and centrifuged at 900 rpm for ten minutes. The aqueous humor and vitreous samples were treated the same way as retina and choroid with the exception that no homogenization was needed. The vitreous samples from groups one and two were diluted 10 times further in the same buffer as above. Five standards were prepared in PBS

with HSA (35.8 μ M, 3.58 μ M, 358 nM, 35.8 nM and 17.9 nM). Subsequently, standards and samples were subjected to pepsin digestion and analysis of the concentration of Z variant in tissue extracts was determined using the LC/LC/MS/MS method described in Example 8.

5 *Results*

The concentrations of Z variant after intravitreal administration were high in all compartments after one day (6 – 200 μ M) and, surprisingly, remained high 4 days post-administration (1.5 – 78 μ M). In particular, the concentration of the Z molecule in the vitreous ranged from 118 to 201 μ M (average 161 μ M, n = 6 eyes) one day after injection and remained at 26 to 78 μ M (average 46 μ M, n = 6) four days post-injection, pointing at a $T_{1/2}$ of several days. There appears to be an inverse relationship between size and elimination of drugs after intravitreal injection in rabbit eye described by the following examples; Moxifloxacin (MW < 0.35 kDa, $T_{1/2}$ = 1.72 h, Mohan *et al.* Trans Am Ophthalmol Soc 2005, 103:76-83), ESBA105 (MW = 26 kDa, $T_{1/2}$ = 25 h, Ottiger *et al.* Investigative Ophthalmology & Visual Science 2009, 50: 779-786) and

10 Ranibizumab (MW = 48 kDa, $T_{1/2}$ = 2.88 days, Bakri *et al.* American Academy of Ophthalmology 2007, 114:2179-2182). The Z variant tested here had a molecular weight of 7.0 kDa, suggesting that the elimination of the Z molecule was slower than what would be expected for such a small molecule in vitreous.

15

SEQUENCE LISTING IN ELECTRONIC FORM

In accordance with Section 111(1) of the Patent Rules, this description contains a sequence listing in electronic form in ASCII text format (file: 22819-654 Seq 09-07-14 v1.txt).

A copy of the sequence listing in electronic form is available from the Canadian Intellectual Property Office.

CLAIMS:

1. C5 binding polypeptide, comprising a C5 binding motif, *BM*, which motif forms part of a three-helix bundle protein domain and consists of an amino acid sequence according to

5

EX₂X₃X₄A X₆X₇EID X₁₁LPNL X₁₆X₁₇X₁₈QW X₂₁AFIX₂₅ X₂₆LX₂₈D,

wherein, independently of each other,

X₂ is selected from H, Q, S, T and V;

10 X₃ is selected from I, L, M and V;

X₄ is selected from A, D, E, H, K, L, N, Q, R, S, T and Y;

X₆ is selected from N and W;

X₇ is selected from A, D, E, H, N, Q, R, S and T;

X₁₁ is selected from A, E, G, H, K, L, Q, R, S, T and Y;

15 X₁₆ is selected from N and T;

X₁₇ is selected from I, L and V;

X₁₈ is selected from A, D, E, H, K, N, Q, R, S and T;

X₂₁ is selected from I, L and V;

X₂₅ is selected from D, E, G, H, N, S and T;

20 X₂₆ is selected from K and S; and

X₂₈ is selected from A, D, E, H, N, Q, S, T and Y.

2. C5 binding polypeptide according to claim 1, wherein the amino acid sequence fulfills at least four of the following eight conditions I-VIII:

25

I. X₂ is V;

II. X₃ is selected from I and L;

III. X₆ is W;

IV. X₇ is selected from D and N;

- V. X_{17} is selected from I and L;
- VI. X_{21} is L;
- VII. X_{25} is N;
- VIII. X_{28} is D.

5

3. C5 binding polypeptide according to claim 1 or 2, wherein the amino acid sequence is selected from any one of SEQ ID NOs:1-248.

4. C5 binding polypeptide according to claim 3, wherein the amino acid sequence is selected from any one of SEQ ID NOs:1-12, SEQ ID NO:20, SEQ ID NOs:23-24, SEQ ID NOs:26-28, SEQ ID NOs:32-35, SEQ ID NOs:38-39, SEQ ID NO:41, SEQ ID NO:46, SEQ ID NO:49, SEQ ID NOs:56-57, SEQ ID NO:59, SEQ ID NO:66, SEQ ID NOs:78-79, SEQ ID NO:87, SEQ ID NO:92, SEQ ID NO:106, SEQ ID NO:110, SEQ ID NO:119, SEQ ID NO:125, SEQ ID NO:141, SEQ ID NO:151, SEQ ID NO:161, SEQ ID NO:166, SEQ ID NO:187, SEQ ID NO:197, SEQ ID NO:203, SEQ ID NO:205, SEQ ID NO:215 and SEQ ID NO:243.

20

5. C5 binding polypeptide according to claim 4, wherein the amino acid sequence is selected from any one of SEQ ID NOs:1-12.

6. C5 binding polypeptide according to any one of claims 1-5, which comprises an amino acid sequence according to:

$K-[BM]-DPSQS X_a X_b LLX_c EAKKL NDX_d Q;$

25 wherein

$[BM]$ is a C5 binding motif as defined in any one of claims 1-5;

X_a is selected from A and S;

X_b is selected from N and E;

X_c is selected from A, S and C; and

30 X_d is selected from A and S.

7. C5 binding polypeptide according to claim 6, wherein the amino acid sequence is selected from any one of SEQ ID NOs:249-496.

8. C5 binding polypeptide according to claim 7, wherein the amino acid sequence is selected from any one of SEQ ID NOs:249-260, SEQ ID NO:268, SEQ ID NOs:271-272, SEQ ID NOs:274-276, SEQ ID NOs:280-283, SEQ ID NOs:286-287, SEQ ID NO:289, SEQ ID NO:294, SEQ ID NO:297, SEQ ID NOs:304-305, SEQ ID NO:307, SEQ ID NO:314, SEQ ID NOs:326-327, SEQ ID NO:335, SEQ ID NO:340, SEQ ID NO:354, SEQ ID NO:358, SEQ ID NO:367, SEQ ID NO:373, SEQ ID NO:389, SEQ ID NO:399, SEQ ID NO:409, SEQ ID NO:414, SEQ ID NO:435, SEQ ID NO:445, SEQ ID NO:451, SEQ ID NO:453, SEQ ID NO:463 and SEQ ID NO:491.

9. C5 binding polypeptide according to claim 8, wherein the amino acid sequence is selected from any one of SEQ ID NOs:249-260.

15

10. C5 binding polypeptide according to any one of claims 1-9, wherein the amino acid sequence is selected from any one of SEQ ID NOs:497-757.

11. C5 binding polypeptide according to claim 10, wherein the amino acid sequence is selected from any one of SEQ ID NOs:497-508, SEQ ID NO:516, SEQ ID NOs:519-520, SEQ ID NOs:522-524, SEQ ID NOs:528-531, SEQ ID NOs:534-535, SEQ ID NO:537, SEQ ID NO:542, SEQ ID NO:545, SEQ ID NOs:552-553, SEQ ID NO:555, SEQ ID NO:562, SEQ ID NOs:574-575, SEQ ID NO:583, SEQ ID NO:588, SEQ ID NO:602, SEQ ID NO:606, SEQ ID NO:615, SEQ ID NO:621, SEQ ID NO:637, SEQ ID NO:647, SEQ ID NO:657, SEQ ID NO:662, SEQ ID NO:683, SEQ ID NO:693, SEQ ID NO:699, SEQ ID NO:701, SEQ ID NO:711, SEQ ID NO:739 and SEQ ID NOs:746-757.

25

12. C5 binding polypeptide according to claim 11, wherein the amino acid sequence is selected from any one of SEQ ID NOs:497-508 and SEQ ID NOs:746-757.

13. C5 binding polypeptide according to claim 12, wherein the amino acid sequence is selected from any one of SEQ ID NO:497, SEQ ID NO:498, SEQ ID NO:499, SEQ ID NO:500, SEQ ID NO:501, SEQ ID NO:746, SEQ ID NO:747, SEQ ID NO:748, SEQ ID NO:750 and SEQ ID NO:753.

5

14. C5 binding polypeptide according to any one of claims 1-13, which inhibits cleavage of C5.

15. C5 binding polypeptide according to any one of claims 1-14, wherein the C5
10 binding polypeptide binds to C5 such that the K_D value is at most 1×10^{-6} .

16. C5 binding polypeptide according to claim 15, wherein the K_D value is at most 1×10^{-7} M.

15 17. C5 binding polypeptide according to claim 15, wherein the K_D value is at most 1×10^{-8} M.

18. C5 binding polypeptide according to claim 15, wherein the K_D value is at most 1×10^{-9} M.

20

19. C5 binding polypeptide according to any one of claims 1-18, comprising further C terminal and/or N terminal amino acids that improves production, purification, stabilization *in vivo* or *in vitro*, coupling, or detection of the polypeptide.

25 20. C5 binding polypeptide according to any one of claims 1-19 in multimeric form, comprising at least two C5 binding polypeptide monomer units, the amino acid sequences of which may be the same or different.

21. C5 binding compound, comprising at least one C5 binding polypeptide
30 according to any one of claims 1-20; at least one albumin binding domain of streptococcal

protein G, and at least one linking moiety for linking said at least one domain to the C or N terminal of said at least one C5 binding polypeptide.

22. C5 binding compound according to claim 21, having a structure selected from

- 5 [CBP1]-[LI]-[ALBD];
 [CBP1]-[CBP2]-[LI]-[ALBD];
 [CBP1]-[LI]-[ALBD]-[L2]-[CBP2];
 [ALBD]-[LI]-[CBP1];
 [ALBD]-[LI]- [CBP1]-[CBP2];
 10 [CBP1]-[LI]- [CBP2]-[L2]-[ALBD]; and
 [ALBD]-[LI]- [CBP1]- [L2]- [CBP2]

wherein, independently of each other,

[CBP1] and [CBP2] are C5 binding polypeptides which may be the same or different;
 [LI] and [L2] are linking moieties which may be the same or different; and

- 15 [ALBD] is an albumin binding domain of streptococcal protein G.

23. C5 binding compound according to claim 22, wherein the linking moiety is selected from G, GS; [G₂S]_n; [G₃S]_n; [G₄S]_n; GS[G₄S]_n, wherein n is 0-7; [S₂G]_m; [S₃G]_m; [S₄G]_m; wherein m is 0-7, and VDGS.

20

24. C5 binding compound according to any one of claims 21-23, wherein said albumin-binding domain is as set out in SEQ ID NO:759.

25. C5 binding compound according to any one of claims 21-23, wherein each of
 25 said C5 binding polypeptides is independently selected from the C5 binding polypeptide as defined in any one of claims 10-13.

26. A fusion polypeptide comprising the C5 binding polypeptide according to any one of the claims 1 to 20 and a further polypeptide domain.

27. The fusion polypeptide according to claim 26, wherein said further polypeptide domain comprises a half-life extending moiety which increases the half-life of the C5 binding polypeptide in vivo.

28. The fusion polypeptide according to claim 26, wherein the further polypeptide domain comprises an Fc domain.

29. Polynucleotide encoding the C5 binding polypeptide according to any one of claims 1-20 or the C5 binding compound according to any one of claims 21-25.

30. Combination of the C5 binding polypeptide according to any one of claims 1-20 or the C5 binding compound according to any one of claims 21-24 with a therapeutic agent.

31. C5 binding polypeptide according to any one of claims 1-20, C5 binding compound according to any one of claims 21-25 or combination according to claim 30 for treatment of a C5 related condition selected from inflammatory disease; autoimmune disease; infectious disease; cardiovascular disease; neurodegenerative disorders; cancer; graft injury; wounds; eye disease; kidney disease; pulmonary diseases; hematological diseases; allergic diseases and dermatological diseases.

32. C5 binding polypeptide according to any one of claims 1-20, C5 binding compound according to any one of claims 21-25 or combination according to claim 30 for treatment of paroxysmal nocturnal hemoglobinuria (PNH).

33. Use of the C5 binding polypeptide according to any one of claims 1-20, C5 binding compound according to any one of claims 21-25 or combination according to

claim 30, for treatment of a C5 related condition in a mammalian subject in need thereof, wherein said C5 related condition is selected from inflammatory disease; autoimmune disease; infectious disease; cardiovascular disease; neurodegenerative disorders; cancer; graft injury; wounds; eye disease; kidney disease; pulmonary diseases; hematological diseases; allergic diseases and dermatological diseases.

34. The use according to claim 33, in which binding of the C5 binding polypeptide, the C5 binding compound or the combination to C5 inhibits cleavage of C5.

35. The use according to claim 33, wherein said C5 related condition is paroxysmal nocturnal hemoglobinuria (PNH).

36. The use according to any one of claims 33 to 35, wherein the C5 binding polypeptide is for intravenous, subcutaneous, nasal, oral, intravitreal, or topical administration, or for administration by inhalation.

37. The C5 binding polypeptide according to any one of claims 1-20 or the C5 binding compound according to any one of claims 21-25, comprising the amino acid sequence of SEQ ID NO:1.

38. The C5 binding polypeptide according to any one of claims 1-20 or the C5 binding compound according to any one of claims 21-25, comprising the amino acid sequence of SEQ ID NO:4.

39. The fusion polypeptide according to claim 26, wherein the further polypeptide domain comprises a cysteine residue, a His₆ tag, a myc tag, or a FLAG tag.

40. The fusion polypeptide according to claim 26, wherein the further polypeptide domain comprises an albumin binding domain.

41. The fusion polypeptide according to claim 26, wherein the further polypeptide domain comprises an albumin binding domain of streptococcal protein G.

42. The fusion polypeptide according to claim 26, wherein the further polypeptide domain comprises an albumin binding domain comprising the amino acid sequence of SEQ ID NO: 759.

43. The fusion polypeptide according to any one of claims 26-28 or 39-42, wherein the fusion polypeptide binds to C5 such that the K_D value is at most 1×10^{-6} .

44. The fusion polypeptide according to claim 43, wherein the K_D value is at most 1×10^{-7} M.

45. The fusion polypeptide according to claim 43, wherein the K_D value is at most 1×10^{-8} M.

46. The fusion polypeptide according to claim 43, wherein the K_D value is at most 1×10^{-9} M.

47. The fusion polypeptide according to any one of claims 26-28 or 39-46, wherein the C5 binding polypeptide comprises SEQ ID NO:1.

48. The fusion polypeptide according to any one of claims 26-28 or 39-46, wherein the C5 binding polypeptide comprises SEQ ID NO:4.

49. A polynucleotide encoding the C5 binding polypeptide according to claim 37 or claim 38.

50. A polynucleotide encoding the fusion polypeptide according to any one of claims 26-28 or 39-48.

51. An expression vector comprising the polynucleotide according to any one of claims 29, 49 or 50.

5 52. The C5 binding polypeptide according to claim 37 or 38, or the fusion polypeptide according to any one of claims 26-28 or 39-48, for treatment of a C5 related condition selected from inflammatory disease; autoimmune disease; infectious disease; cardiovascular disease; neurodegenerative disorders; cancer; graft injury; wounds; eye disease; kidney disease; pulmonary diseases; hematological diseases; allergic diseases and
10 dermatological diseases.

53. The C5 binding polypeptide according to claim 37 or 38, or the fusion polypeptide according to any one of claims 26-28 or 39-48, for treatment of paroxysmal nocturnal hemoglobinuria (PNH).

15 54. Use of the C5 binding polypeptide according to claim 37 or 38, or the fusion polypeptide according to any one of claims 26-28 or 39-48, for treatment of a C5 related condition in a mammalian subject in need thereof, wherein said C5 related condition is selected from inflammatory disease; autoimmune disease; infectious disease; cardiovascular
20 disease; neurodegenerative disorders; cancer; graft injury; wounds; eye disease; kidney disease; pulmonary diseases; hematological diseases; allergic diseases and dermatological diseases.

55. The use according to claim 54, in which binding of the C5 binding polypeptide
25 or fusion polypeptide to C5 inhibits cleavage of C5.

56. The use according to claim 54, wherein said C5 related condition is paroxysmal nocturnal hemoglobinuria (PNH).

57. The use according to any one of claims 54 to 56, wherein the C5 binding polypeptide or fusion polypeptide is for intravenous, subcutaneous, nasal, oral, intravitreal, or topical administration, or for administration by inhalation.

Fig. 1

Polypeptide	Amino Acid Sequence	SEQ ID NO :
CEM06175	EVLEAWDEIDRLPNLTIEQWLAFINKLDD	1
CEM08044	EVLEAWNEIDRLPNLTIEQWLAFINKLDD	2
CEM05998	EVIEAWNEIDRLPNLTIEQWLAFINKLDD	3
CEM06009	EVLEAWDEIDRLPNLTLDQWLAFINKLDD	4
CEM06079	EVLDAWDEIDALPNLTIEQWLAFINKLDD	5
CEM06126	EVIDAWDEIDRLPNLTLDQWLAFINKLDD	6
CEM06140	ETLEAWDEIDRLPNLTIEQWLAFINKLDD	7
CEM06189	EVIDAWNEIDALPNLTLDQWLAFINKLDD	8
CEM06214	EVLDAWDEIDKLPNLTIDQWLAFINKLDD	9
CEM06215	EVLEAWDEIDHLPNLTLDQWLAFINKLDD	10
CEM06226	EVLEAWDEIDALPNLTIEQWLAFINKLDD	11
CEM06018	EVLDAWDEIDKLPNLTIEQWLAFINKLDD	12
CEM05477	ETITAWDEIDKLPNLTIEQWLAFIGKLED	13
CEM05363	ESMKAWDEIDRLPNLTININQWVAFIDSLYD	14
CEM05483	ESIEAWTEIDHLPNLTIEQWLAFINKLTD	15
CEM05538	EVLDAWHEIDTLPNLTIVRQWLAFISKLED	16
CEM05692	EHIQANEEIDRLPNLTIKQWLAFINKLHD	17
CEM05994	EVLHAWAEIDALPNLTIEQWLAFINKLDD	18
CEM05995	EVLAAWDEIDSLPNLTILQQWLAFINKLDD	19
CEM05996	EVIDAWNEIDALPNLTIEQWLAFINKLDD	20
CEM05997	EVLDAWNEIDALPNLTIDQWLAFINKLSD	21
CEM05999	EVIEAWDEIDGLPNLTIEQWLAFINKLDD	22
CEM06000	EVLEAWDEIDHLPNLTILQQWLAFINKLDD	23

CBM06001	EVIEAWNEIDALPNLTIEQWLAFINKLDD	24
CBM06002	EVIAAWNEIDRLPNLTTLTQWLAFINKLDD	25
CBM06003	EVIEAWDEIDALPNLTLOQWLAFINKLDD	26
CBM06004	EVIAAWDEIDKLPNLTIEQWLAFINKLDD	27
CBM06005	EVIAAWDEIDKLPNLTLOQWLAFINKLDD	28
CBM06006	ETIAAWDEIDKLPNLTIEQWLAFINKLDD	29
CBM06007	ETIEAWNEIDRLPNLTIEQWLAFINKLDD	30
CBM06008	EVLEAWREIDALPNLTIOQWLAFINKLDD	31
CBM06010	EVIEAWDEIDQLPNLTIEQWLAFINKLDD	32
CBM06011	EVLRAWDEIDHLPNLTLEQWLAFINKLDD	33
CBM06012	EVLEAWDEIDRLPNLTTLNQWLAFINKLDD	34
CBM06013	EVLDAWNEIDHLPNLTIEQWLAFINKLDD	35
CBM06014	EVIDAWNEIDKLPNLTIEQWLAFINKLDD	36
CBM06015	ETLEAWDEIDQLPNLTLOQWLAFINKLDD	37
CBM06016	EVIEAWNEIDALPNLTLDQWLAFINKLDD	38
CBM06017	EVIDAWNEIDRLPNLTLOQWLAFINKLDD	39
CBM06019	EVIDAWNEIDQLPNLTLEQWLAFINKLDD	40
CBM06020	ETIAAWDEIDHLPNLTLEQWLAFINKLDD	41
CBM06024	EVLQAWDEIDHLPNLTIOQWLAFINKLSD	42
CBM06025	ETLHAWAEIDRLPNLTIEQWLAFINKLDD	43
CBM06026	EVLEAWNEIDHLPNLTIAQWLAFINKLDD	44
CBM06027	EVIEAWDEIDKLPNLTIAQWLAFINKLDD	45
CBM06028	EVLDAWDEIDHLPNLTLOQWLAFINKLDD	46
CBM06029	ETIEAWNEIDKLPNLTTLTQWLAFINKLDD	47
CBM06030	EVLEAWNEIDLPLPNLTIEQWLAFINKLDD	48
CBM06031	EVIEAWDEIDHLPNLTIDQWLAFINKLDD	49

CBM06032	EVI SAWNEIDALPNLTLOQWLAFINKLDD	50
CBM06033	EVIAAWNEIDKLPNLTLEQWLAFINKLDD	51
CBM06034	ETIEAWNEIDSLPNLTLDQWLAFINKLDD	52
CBM06035	EVLDAWNEIDQLPNLTLOQWLAFINKLDD	53
CBM06037	EVIAAWNEIDHLPNLTIEQWLAFINKLDD	54
CBM06038	EVLEAWDEIDHLPNLTITQWLAFINKLDD	55
CBM06039	ETIDAWNEIDHLPNLTIEQWLAFINKLDD	56
CBM06040	EVIEAWNEIDHLPNLTIOQWLAFINKLDD	57
CBM06041	EVIQAWNEIDALPNLTI SQWLAFINKLDD	58
CBM06043	EVIAAWDEIDSLPNLTIEQWLAFINKLDD	59
CBM06044	EHIEAWNEIDALPNLTIEQWLAFINKLQD	60
CBM06045	EVLEAWNEIDKLPNLTLDQWLAFINKLDD	61
CBM06047	EVIDAWNEIDHLPNLTIEQWLAFINKLAD	62
CBM06048	ETIDAWDEIDKLPNLTIEQWLAFINKLDD	63
CBM06049	EVIAAWDEIDLLPNLTLOQWLAFINKLAD	64
CBM06050	EVIHAWDEIDKLPNLTIEQWLAFINKLDD	65
CBM06051	EVIAAWNEIDHLPNLTIEQWLAFINKLDD	66
CBM06052	ETLDAWNEIDKLPNLTLSQWLAFINKLDD	67
CBM06053	EVLEAWNEIDALPNLTLEQWLAFINKLDD	68
CBM06054	EVIQAWDEIDHLPNLTISQWLAFINKLDD	69
CBM06055	EVLQAWDEIDSLPNLTIEQWLAFINKLDD	70
CBM06056	ETLEAWDEIDHLPNLTIAQWLAFINKLDD	71
CBM06057	ETIDAWNEIDRLPNLTI SQWLAFINKLDD	72
CBM06058	EVLDAWHEIDHLPNLTIOQWLAFINKLDD	73
CBM06059	EQIRAWDEIDKLPNLTIEQWLAFINKLAD	74
CBM06060	ETLYAWNEIDKLPNLTIEQWLAFIEKLQD	75

CBM06061	EVIEAWNEIDALPNLTIDQWLAFINKLDD	76
CBM06062	EVLEAWNEIDHLPNLTIQQWLAFINKLDD	77
CBM06063	ETIEAWDEIDALPNLTIEQWLAFINKLDD	78
CBM06065	EVIEAWNEIDHLPNLTQQWLAFINKLDD	79
CBM06066	EVIEAWNEIDKLPNLTIQQWLAFINKLDD	80
CBM06068	ETLDAWAEIDHLPNLTLDQWLAFINKLDD	81
CBM06069	EHIDAWNEIDALPNLTLSQWLAFINKLDD	82
CBM06070	EVLDAWNEIDKLPNLTIAQWLAFINKLDD	83
CBM06071	EVIEAWTEIDYLPNLTQQWLAFINKLDD	84
CBM06072	ETIEAWNEIDHLPNLTIAQWLAFINKLDD	85
CBM06073	EVIQAWNEIDKLPNLTLEQWLAFINKLDD	86
CBM06074	EVIEAWDEIDHLPNLTIEQWLAFINKLDD	87
CBM06075	ETIDAWNEIDLLPNLTIEQWLAFINKLDD	88
CBM06076	EHIDAWNEIDKLPNLTLDQWLAFINKLDD	89
CBM06077	EVVAWNEIDALPNLTIEQWLAFINKLND	90
CBM06080	EVIEAWNEIDALPNLTIAQWLAFINKLDD	91
CBM06081	EVLQAWDEIDRLPNLTLDQWLAFINKLDD	92
CBM06082	EVIDAWDEIDHLPNLTIEQWLAFINKLSD	93
CBM06083	EVVEAWNEIDQLPNLTIEQWLAFINKLDD	94
CBM06084	EVIQAWNEIDALPNLTIEQWLAFINKLDD	95
CBM06085	EVIQAWDEIDKLPNLTIDQWLAFINKLAD	96
CBM06086	EVVAWDEIDALPNLTLTQWLAFINKLDD	97
CBM06087	EVIQAWNEIDGLPNLTLSQWLAFINKLDD	98
CBM06088	ETIEAWDEIDALPNLTITQWLAFINKLDD	99
CBM06089	EVIDAWNEIDHLPNLTIQQWLAFINKLAD	100
CBM06090	ETIEAWNEIDALPNLTLDQWLAFINKLED	101

CBM06091	EHIAWNEIDELPNLTIEQWLAFINKLAD	102
CBM06092	EVIDAWDEIDHLPNLTIDQWLAFINKLSD	103
CBM06093	EVIDANDEIDALPNLTIAQWLAFINKLHD	104
CBM06095	ETIEAWDEIDKLPNLTIEQWLAFINKLDD	105
CBM06097	EVLAWDEIDHLPNLTIEQWLAFINKLDD	106
CBM06098	EHIDAWNEIDGLPNLTIEQWLAFINKLDD	107
CBM06099	EVIEAWSEIDALPNLTIDQWLAFINKLAD	108
CBM06100	EQLNAWAEIDALPNLTIEQWLAFINKLDD	109
CBM06101	EVIDAWNEIDALPNLTIAQWLAFINKLDD	110
CBM06103	ETIDAWNEIDQLPNLTIEQWLAFINKLDD	111
CBM06104	EVIEAWDEIDKLPNLTIAQWLAFINKLDD	112
CBM06105	EVLAWAEIDHLPNLTIEQWLAFINKLDD	113
CBM06107	EQIDAWNEIDRLPNLTIQQWLAFINKLDD	114
CBM06108	EVLAAWDEIDRLPNLTIEQWLAFINKLDD	115
CBM06109	EVIEAWDEIDHLPNLTILHQWLAFINKLDD	116
CBM06110	EVIEAWNEIDKLPNLTILQQWLAFINKLDD	117
CBM06111	EVIDANDEIDALPNLTIEQWLAFINKLHD	118
CBM06112	EVIAAWDEIDALPNLTIEQWLAFINKLDD	119
CBM06113	EVIEAWTEIDQLPNLTIDQWLAFINKLDD	120
CBM06114	EVINAWNEIDALPNLTIQQWLAFINKLDD	121
CBM06115	EHIEAWDEIDHLPNLTIDQWLAFINKLAD	122
CBM06116	EHLEAWREIDALPNLTIEQWLAFINKLDD	123
CBM06117	EVLDAWNEIDKLPNLTILQQWLAFINKLDD	124
CBM06118	EVIAAWDEIDHLPNLTIIQQWLAFINKLDD	125
CBM06119	EVIQAWNEIDALPNLTIEQWLAFINKLDD	126
CBM06121	EVIDAWNEIDHLPNLTIAQWLAFINKLDD	127

CBM06122	EQLDAWDEIDHLPNLTIDQWLAFINKLSD	128
CBM06123	EVLNAWDEIDKLPNLTIEQWLAFINKLDD	129
CBM06124	EVLEAWNEIDHLPNLTIDQWLAFINKLDD	130
CBM06125	EVLAWDEIDRLPNLTIDQWLAFINKLAD	131
CBM06127	EVIAAWNEIDQLPNLTIDQWLAFINKLDD	132
CBM06128	ETLLAWDEIDALPNLTIEQWLAFINKLDD	133
CBM06129	EVIDAWNEIDTLPNLTIEQWLAFINKLDD	134
CBM06131	EVLHAWNEIDHLPNLTINQWLAFINKLQD	135
CBM06132	EVIQAWNEIDALPNLTIAQWLAFINKLDD	136
CBM06133	ETVDAWNEIDALPNLTIEQWLAFINKLDD	137
CBM06134	EVIQAWDEIDHLPNLTIDQWLAFINKLDD	138
CBM06135	EVLDAWNEIDQLPNLTIQQWLAFINKLDD	139
CBM06136	ETIEAWNEIDALPNLTIDQWLAFINKLDD	140
CBM06137	EVIEAWDEIDALPNLTIDQWLAFINKLDD	141
CBM06138	EVIEAWNEIDQLPNLTIQQWLAFINKLDD	142
CBM06139	EVIEAWTEIDHLPNLTIEQWLAFINKLDD	143
CBM06141	EVIQAWNEIDHLPNLTILQQWLAFINKLED	144
CBM06142	EVIQANNEIDQLPNLTIEQWLAFINKLHD	145
CBM06143	EVLHAWSEIDKLPNLTIEQWLAFINKLDD	146
CBM06144	ETIQAWDEIDKLPNLTIDQWLAFINKLSD	147
CBM06145	ETLRAWDEIDKLPNLTILQQWLAFINKLAD	148
CBM06146	EVIDAWNEIDHLPNLTIEQWLAFINKLED	149
CBM06147	EVIDAWNEIDHLPNLTILQQWLAFINKLAD	150
CBM06148	ETIDAWNEIDALPNLTIDQWLAFINKLDD	151
CBM06149	EVIEAWNEIDQLPNLTIEQWLAFINKLDD	152
CBM06150	EVIRAWDEIDQLPNLTLTSLQWLAFINKLDD	153

CBM06151	EVIEAWNEIDRLPNLTIHQWLAFAINKLDD	154
CBM06152	ETIEAWNEIDQLPNLTIEQWLAFAINKLDD	155
CBM06153	EVLTAWAEIDALPNLTLSQWLAFAINKLDD	156
CBM06154	EVIEAWDEIDKLPNLTVDQWLAFAINKLDD	157
CBM06155	EVIDAWNEIDHLPNLTTLTQWLAFAINKLDD	158
CBM06156	EVIEAWNEIDQLPNLTLDQWLAFAINKLDD	159
CBM06157	ETLQAWDEIDHLPNLTINQWLAFAINKLDD	160
CBM06158	EVIDAWNEIDHLPNLTIEQWLAFAINKLDD	161
CBM06159	EVIEAWNEIDLLPNLTLSQWLAFAINKLDD	162
CBM06160	EVIDAWDEIDRLPNLTIKQWLAFAINKLDD	163
CBM06161	ETLHAWDEIDKLPNLTIEQWLAFAINKLDD	164
CBM06162	EVIKAWDEIDHLPNLTINQWLAFAINKLDD	165
CBM06163	EVIEAWNEIDHLPNLTIAQWLAFAINKLDD	166
CBM06164	EVIQAWNEIDHLPNLTIDQWLAFAITKLED	167
CBM06165	EVIEAWNEIDRLPNLTIKQWLAFAINKLDD	168
CBM06167	EVIEAWNEIDSLPNLTLOQWLAFAINKLDD	169
CBM06168	ETIDAWNEIDKLPNLTIEQWLAFAINKLDD	170
CBM06169	EVLEAWAEIDALPNLTIAQWLAFAINKLDD	171
CBM06170	ETIDAWNEIDRLPNLTIEQWLAFAINKLDD	172
CBM06171	ETLKAWDEIDRLPNLTLEQWLAFAINKLDD	173
CBM06172	ETIAAWNEIDALPNLTLOQWLAFAINKLDD	174
CBM06173	EVLQAWNEIDHLPNLTIQWLAFAINKLDD	175
CBM06174	EVIEAWSEIDHLPNLTLOQWLAFAINKLDD	176
CBM06176	EVIDAWNEIDGLPNLTIEQWLAFAINKLDD	177
CBM06178	EVIHAWNEIDHLPNLTINQWLAFAINKLED	178
CBM06179	EVLDAWNEIDSLPNLTLDQWLAFAINKLDD	179

CBM06180	EQIEAWNEIDRLPNLTLEQWLAFINKLDD	180
CBM06181	EVVDAWNEIDALPNLTTLQQWLAFINKLDD	181
CBM06182	EVIEAWNEIDKLPNLTIEQWLAFINKLDD	182
CBM06183	EVIEANDEIDRLPNLTIEQWLAFINKLHD	183
CBM06184	ETLQAWDEIDKLPNLTIEQWLAFINKLDD	184
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OmCI - His6	MDSESDCTGSEPVDAFQAFSEGKEAYLVVRSTDPKARDCLKGEPAGEKQDNTLPMMTFKNGTDWASTDWTFTLLDGAQVT ATLGNLTQNRVVYDSQSHHCHVDKVEKEVPDYEMMMLDAGGLEVEVECCRQKLEELASGRNQMPHLKDCGGGGSENLY FQGSHHHHH	761
Cynomolgus C5	MGLLGILCFLLFLGKTWQEQTYYVISA PKIFRVGASENIVIQVYGYTEAFDATISIKSYDPDKXFSYSSGHVHLSENKFO NSAVLTIQPKQLPGGQNVSYVYLEVSVKHSKSKKIPITYDNGFLFIHTDKPVYTPDQSVKVRVYSLNDDDLKPAKRETV LTFIDPEGSEIDMVEEIDHIGIISFPDFKIPSNPRYGMWTIOAKYKEDFSTTGTAFFEVKEYVLPHFVSVEPESENFIGY KNFKNFEITIKARIFYNKVVTEADVYITFGIREDLKDDQKEMMQTAMQNTMLINGIAEVTFDSETAVKELSYYSLEDLNN KYLIAVTVIESTGGFSEAEIPGIKYVLSPYKLNLVATPLFLKPGIPYSIKVQKDALDQLVGGVPVTLNAQTIDVNQE TSDLEPRKSVTRVDDGVASFVNLP SGVTVLEFNVKTDAPDLDPENQAREGYRAIAYSSLSQSYLYIDWTDNHHKALLVGE YLNIIIVTPKSPYIDKIITHYNYLILSKGKIIHFGTREKLSASYQ SINIPVTQNMVPSRLLVYIIVTGEQTAELVSDSVW LNIEKCGNQLQVHLSPDADTYSPGQTVSLNMVTGMDSWALTAVDSAVYGVQRRRAKPLERVFOFLEKSDLGCGAGGGL NNANVFHLA GLTFLTNANADDQENDEPCKEIIIRPRMLQEKIEIEIAAKYKHLVVKCCYDGVRRINHDETCEQRAARI SV GPRCVKAFTTECCVVASQLRANNSHKDLQGLRLHMKTL LLPVSKPEIRSYFPE SWLWEVHLVPRRKQLQFALPD SVTTTWEIQ GVGINSNGICVADTIKAKVFKDVFLEMNIPYSVVRGEQVQLKGTVYNRTSGMQFCVKMSAVEGICTSESPVIDHQGTKS SKCVRQKVEGSSNHLVTFVLPLEIGLQININFSLETSFGKEIILVKSIRVVPEGVKRESYSGITLDPRGYGTISRRKEFP YRIPDLVPKTEIKRILSVKGLLVGEILSAVLSREGINILTHLPKGSAAEILMSVVPVYFVHYLETGNHWNIFHSDPLI EKRNLKKEGMSVMSYRNADYSYSVWKGGASITWLTAFALRVLGQVHKYVEQNQNSICNSLLWLVENYQLDNGSFKE NSQYQPIKLQGLTPVEARENSLYLTAFTVIGIRKAFD ICPLVKINTALIKADTFLENTLPAQSTFTTLAISAYALSLGDK THPQFRSIVSALKREALVKGNPPIYRFWKDSLQHKDSSVPNTGTARVETTAYALLTSLNKDINYVNP I IKWLSEEQRY GGGYSTQDTINAIEGLTEYSLLVKQLRLNMDIDVAYKHKGPLHNYKMTDKNFLGRPVLEVLLNDDXVVSTGFGSGLATVH VTTVVHKTSTSEVCSFYLKIDTQDVEASHYRGYGNSDYKRVACASYKPSKEESSGSHAVMDISLPTGINANEEDLK ALVEGDQLFTDYQIKDGHVILQINSIPSSDFLCVRFRI FELFEVGF LSPATFTVYEHPRPKQCTMFYSTSNIKI QKVC EGATCKCIEADCGQMOKELDLTISAETRQQTACNPEIAYAYKVIITSIITTENVFVKYKATLLDIYKTGEAVAEKDSEITF IKKVTCITNAELVKGROYLIMGKEALQIKYNFTFRYIYPLDSLITWIEYWPRDTTCSSCQAF LANLDEFAEDI FLNGC	762

Fig. 2

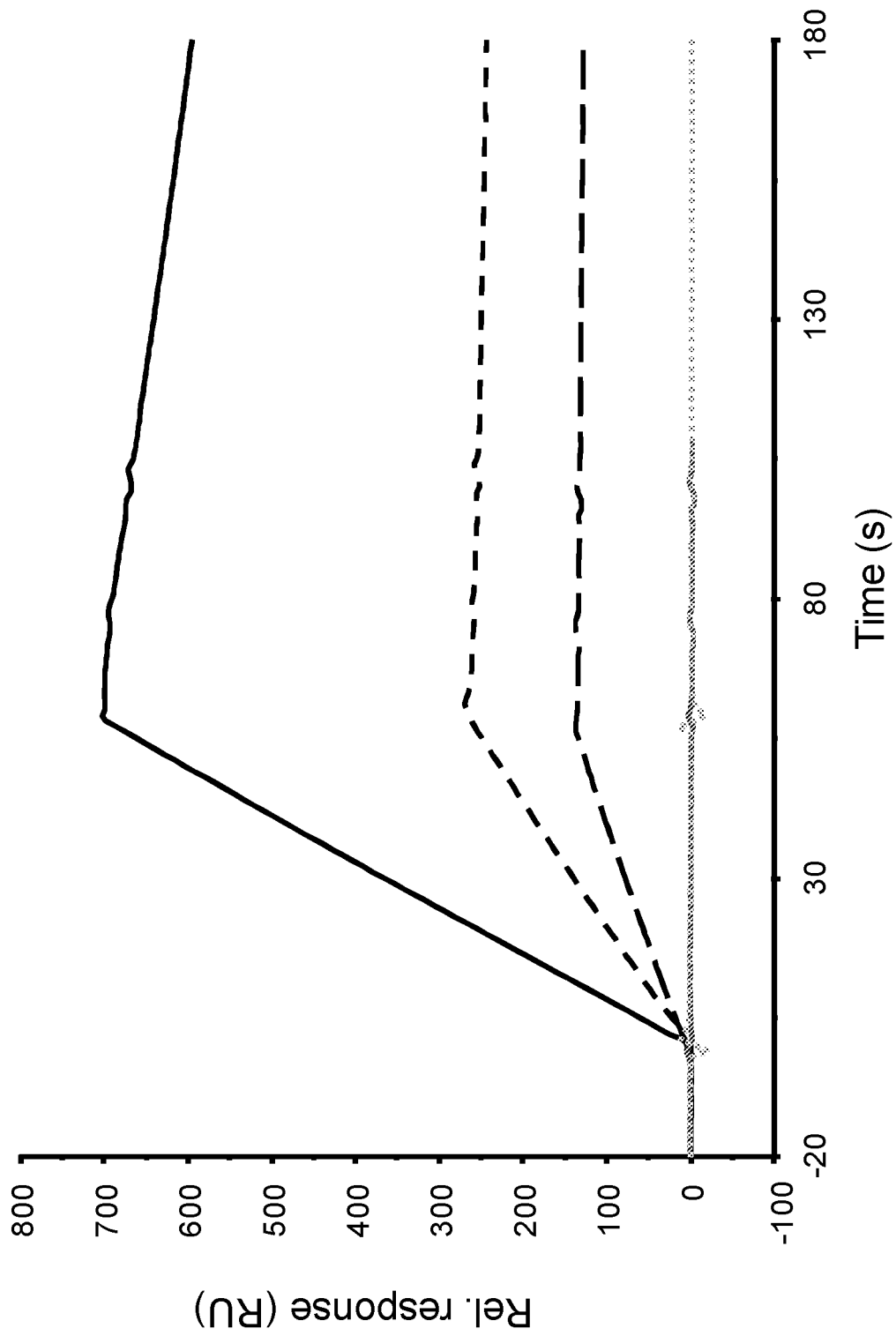


Fig. 3

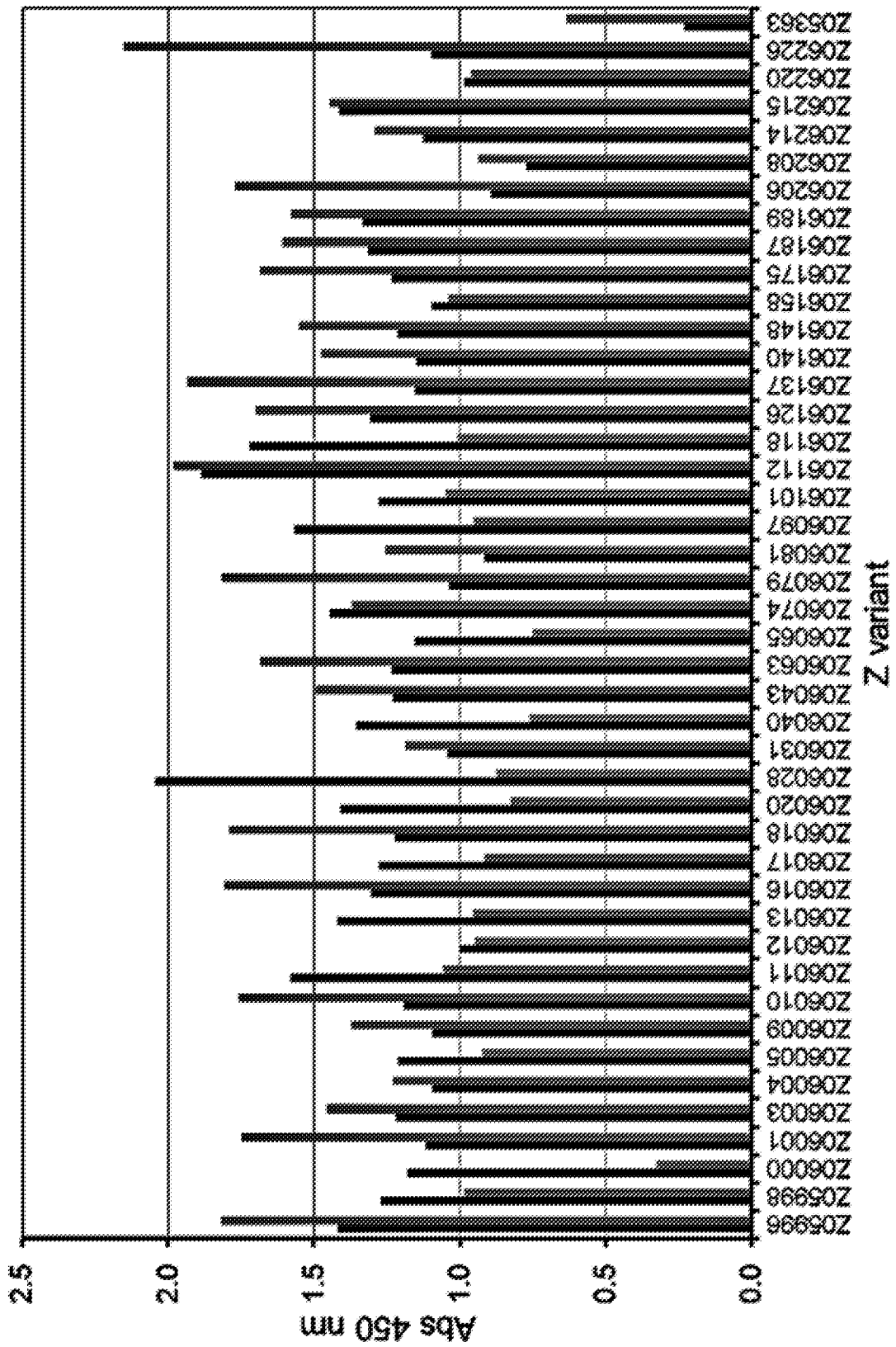


Fig. 4

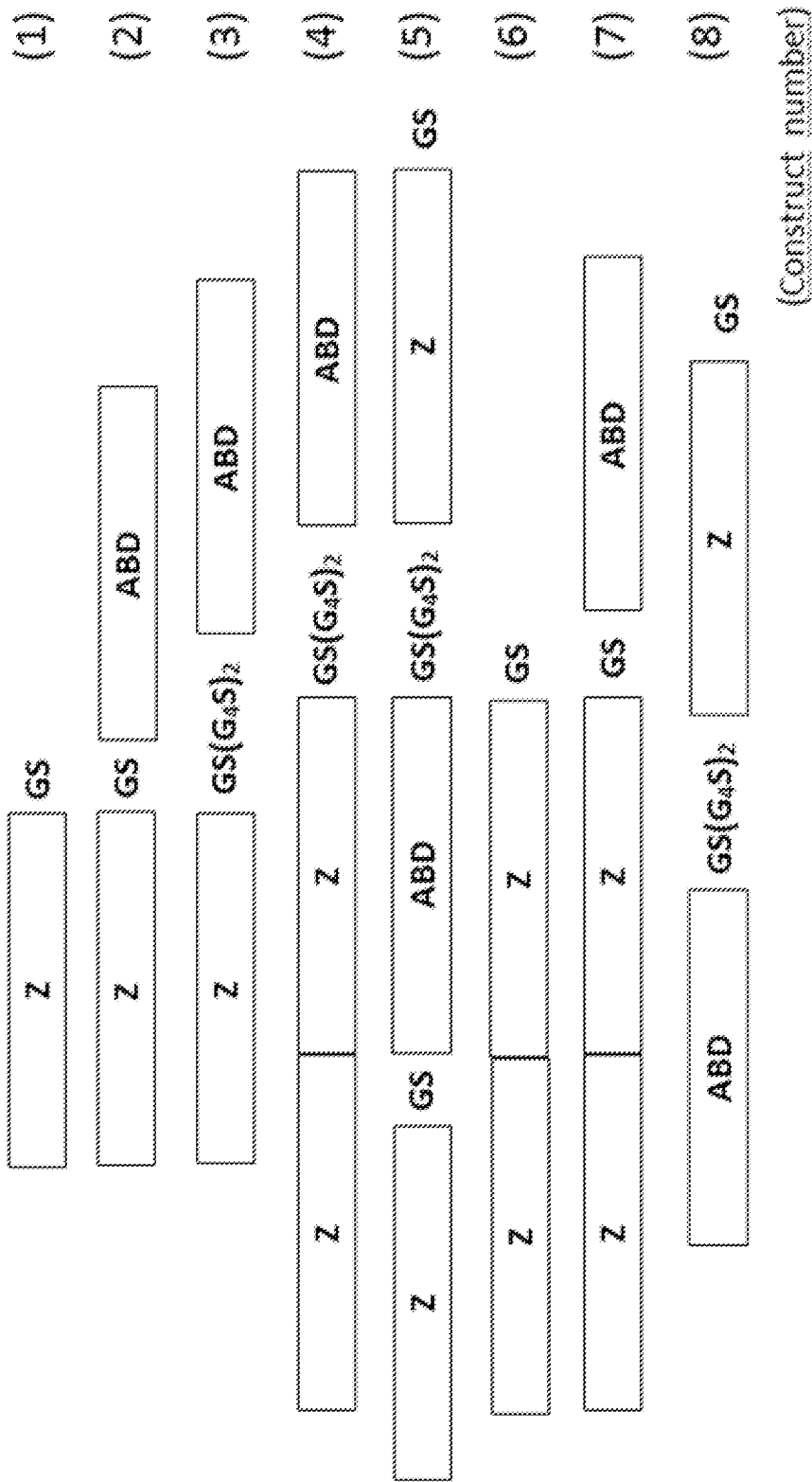


Fig. 5

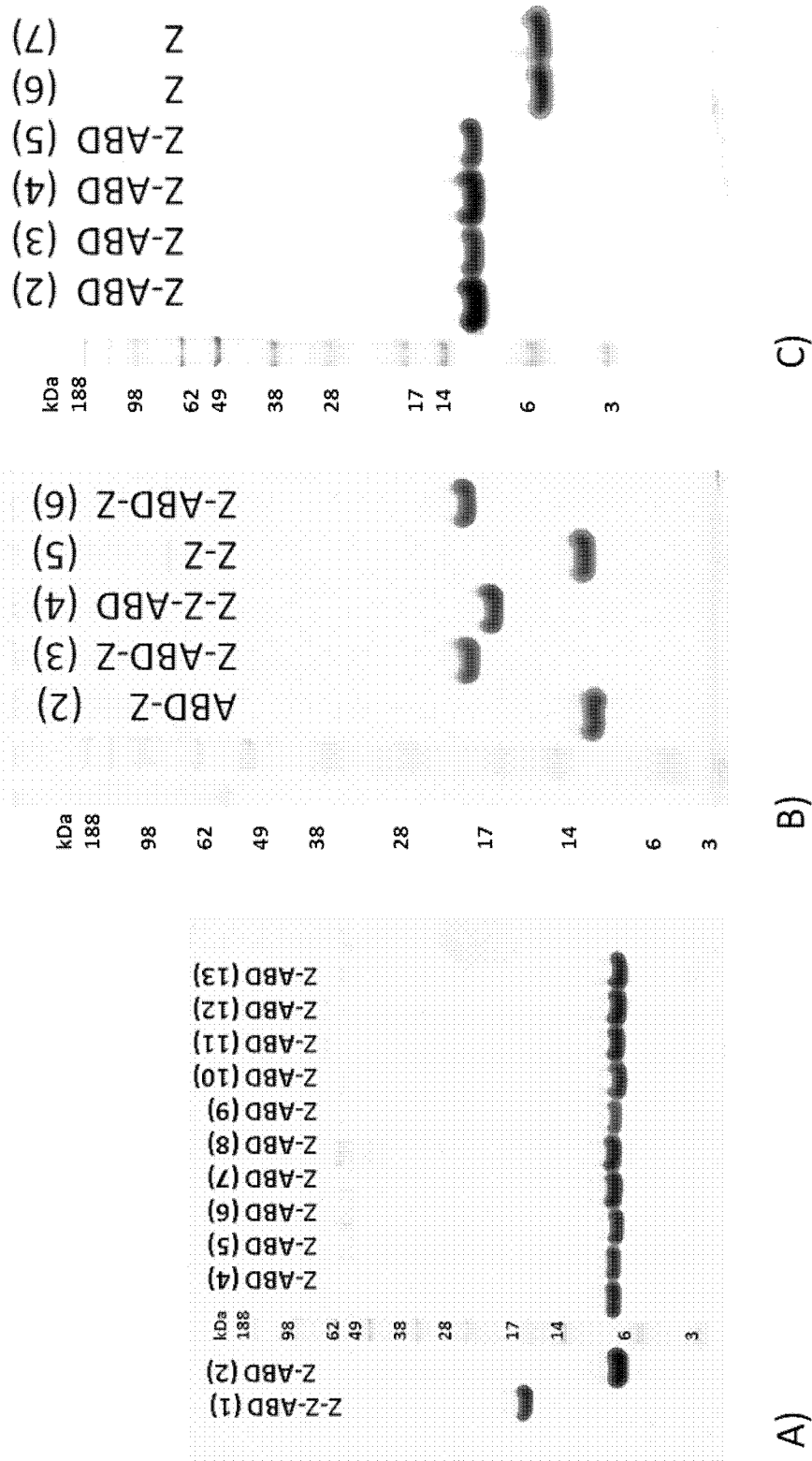


Fig. 6A

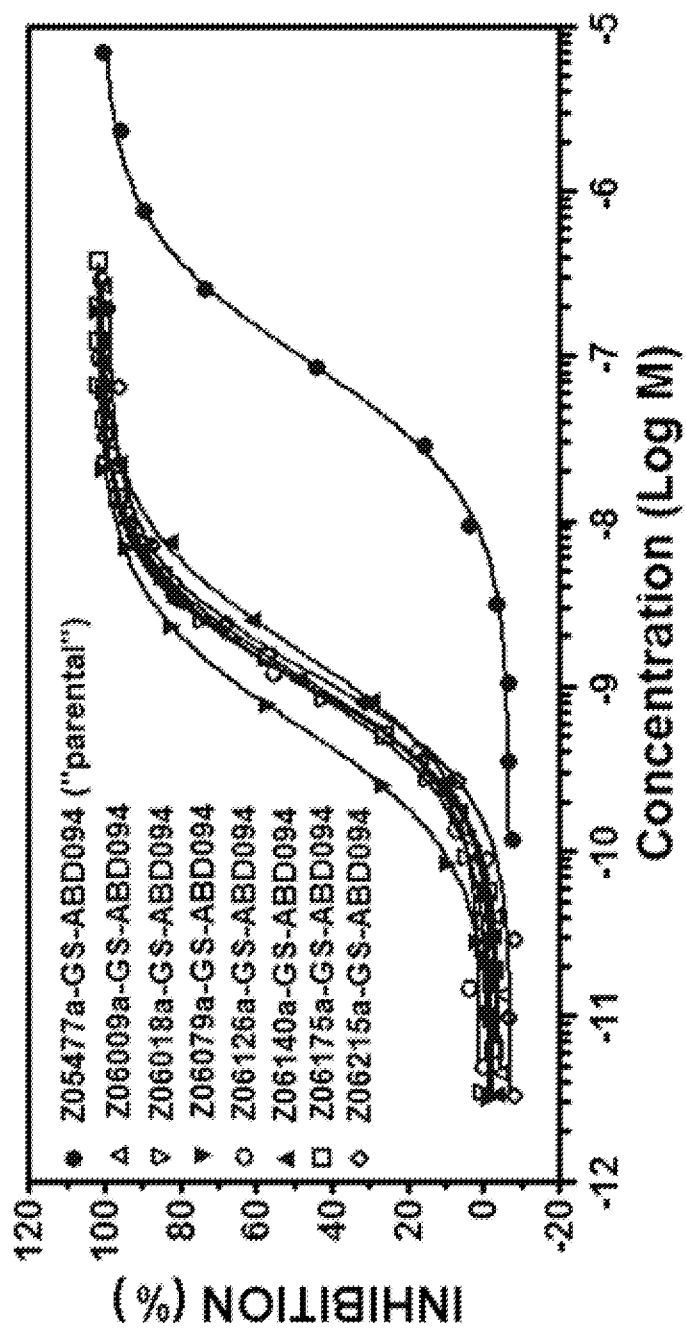


Fig. 6B

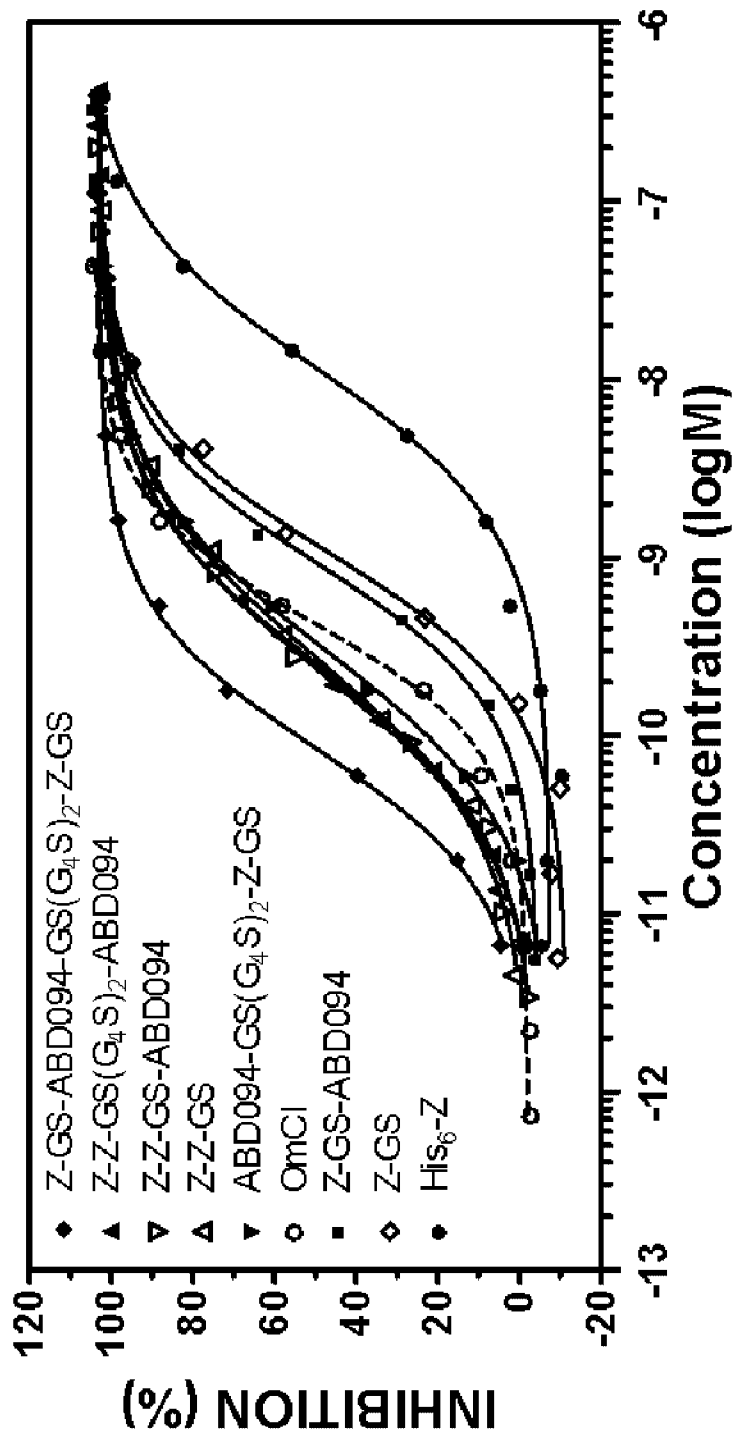


Fig. 7A

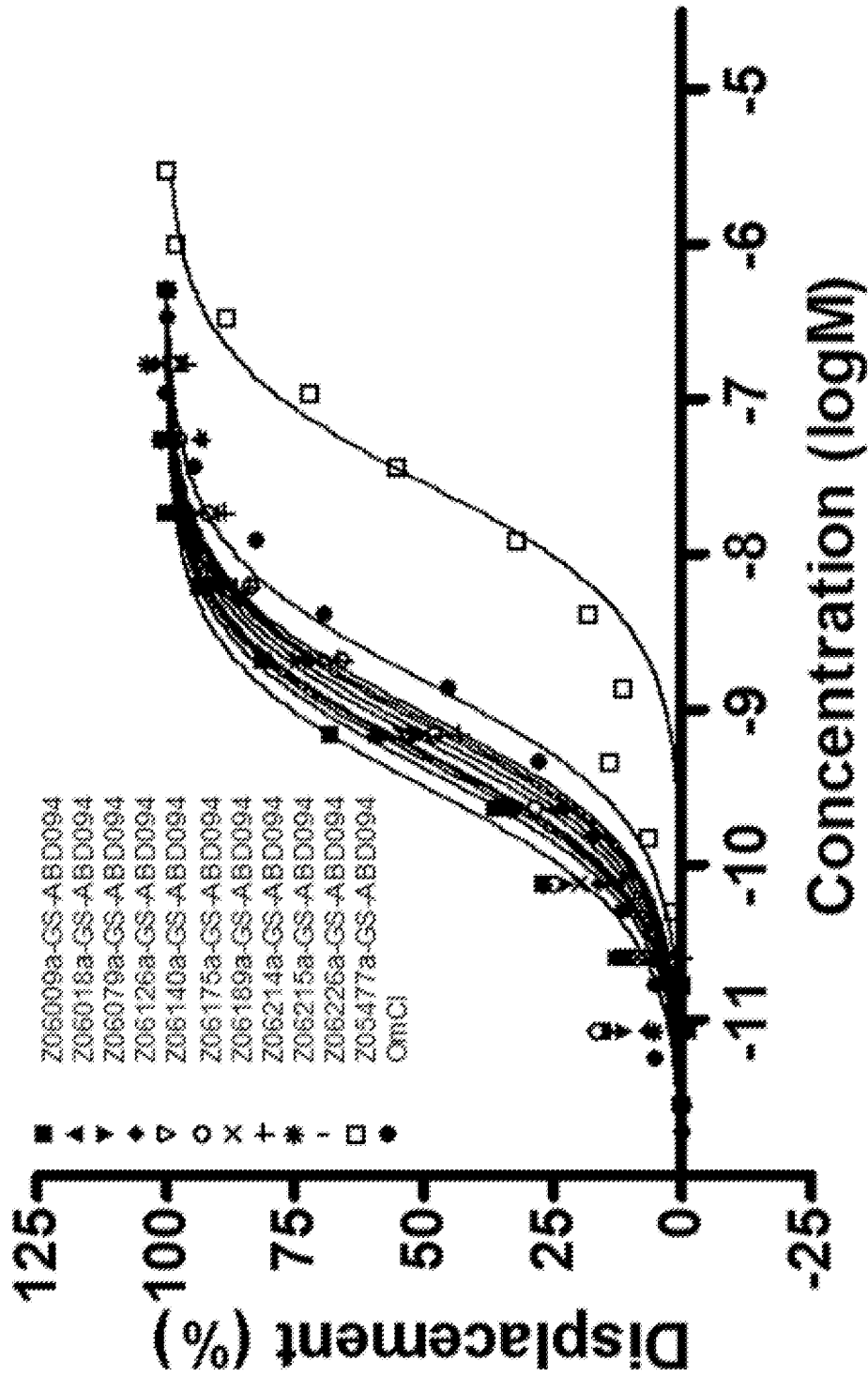


Fig. 7B

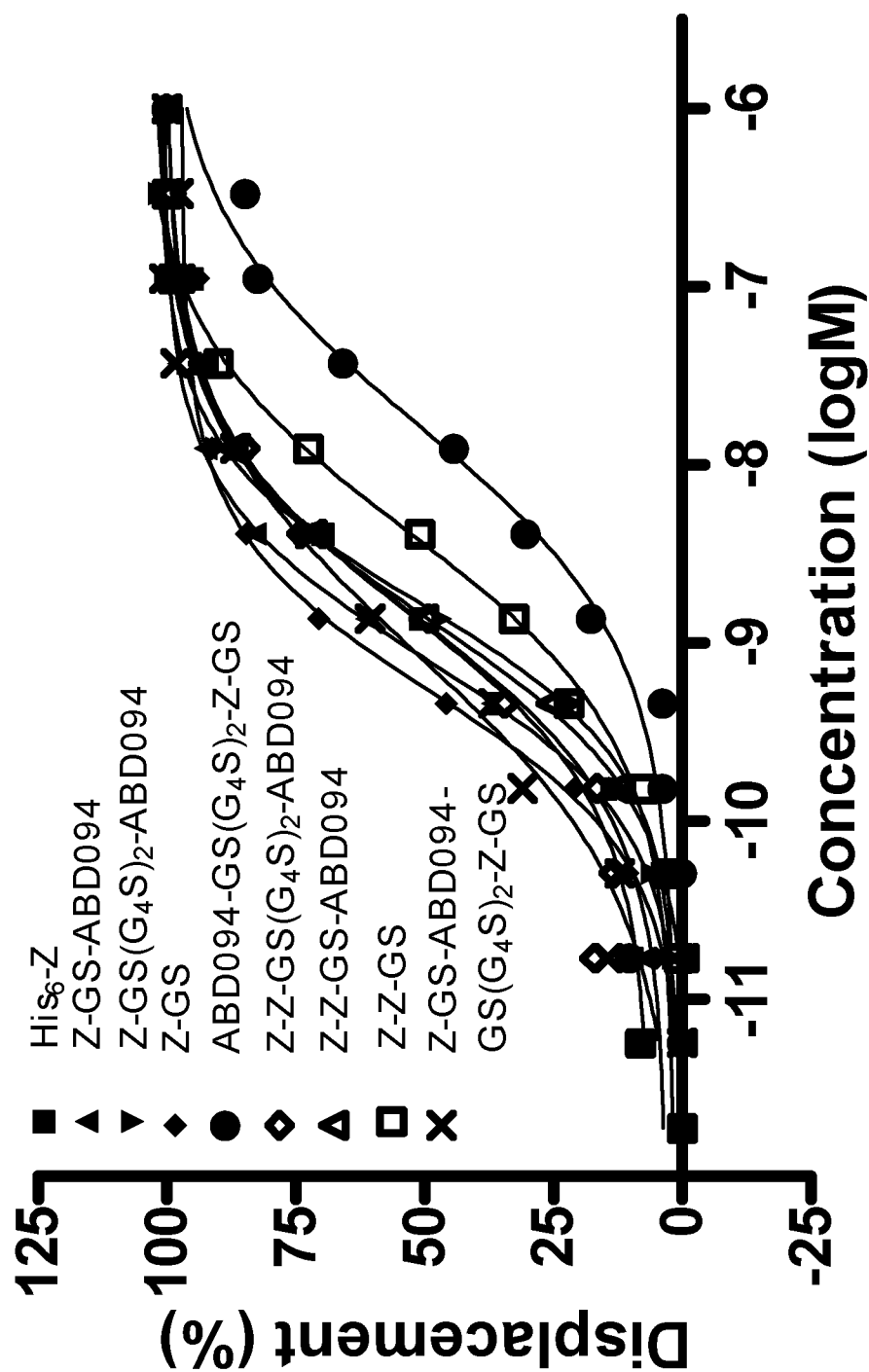


Fig. 8A

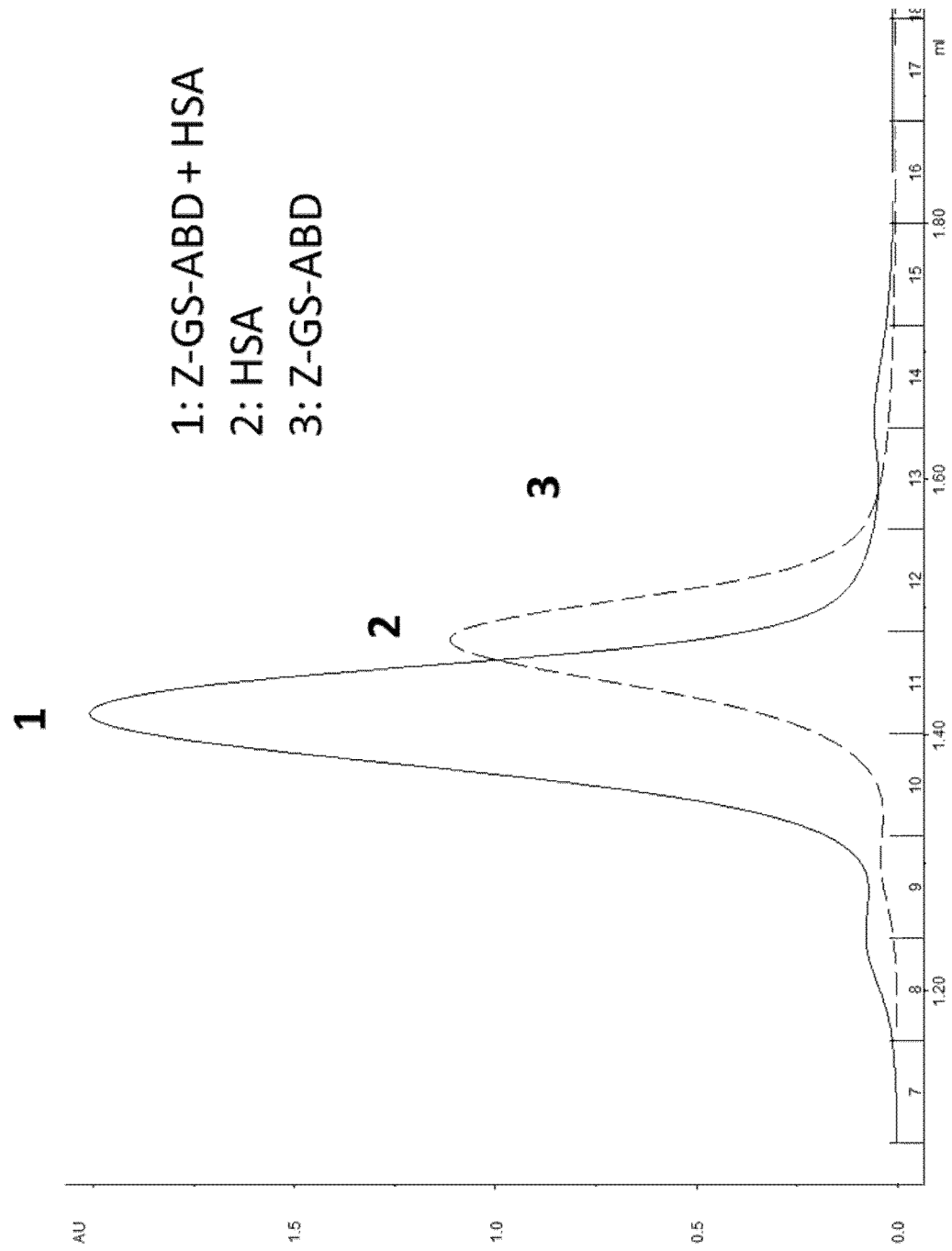


Fig. 8B

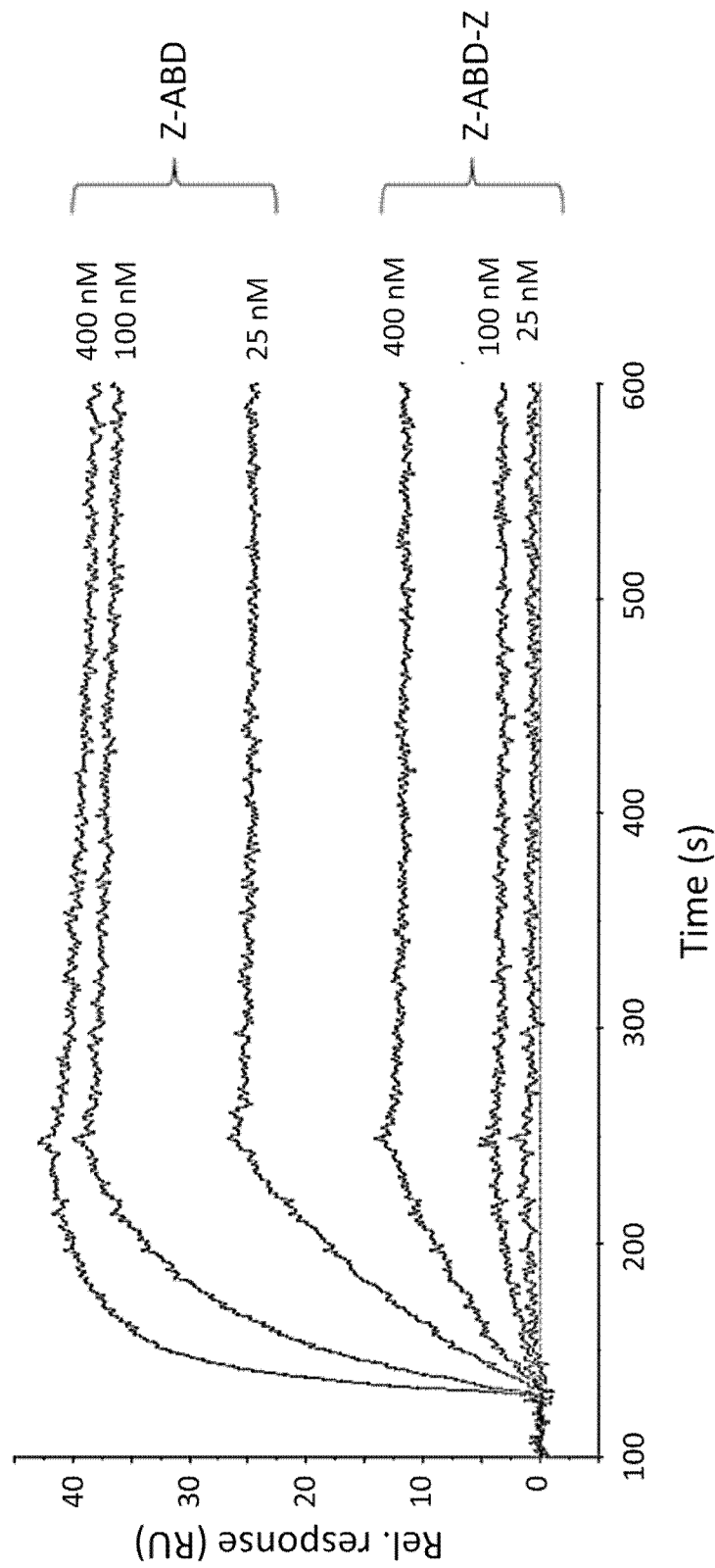


Fig. 9

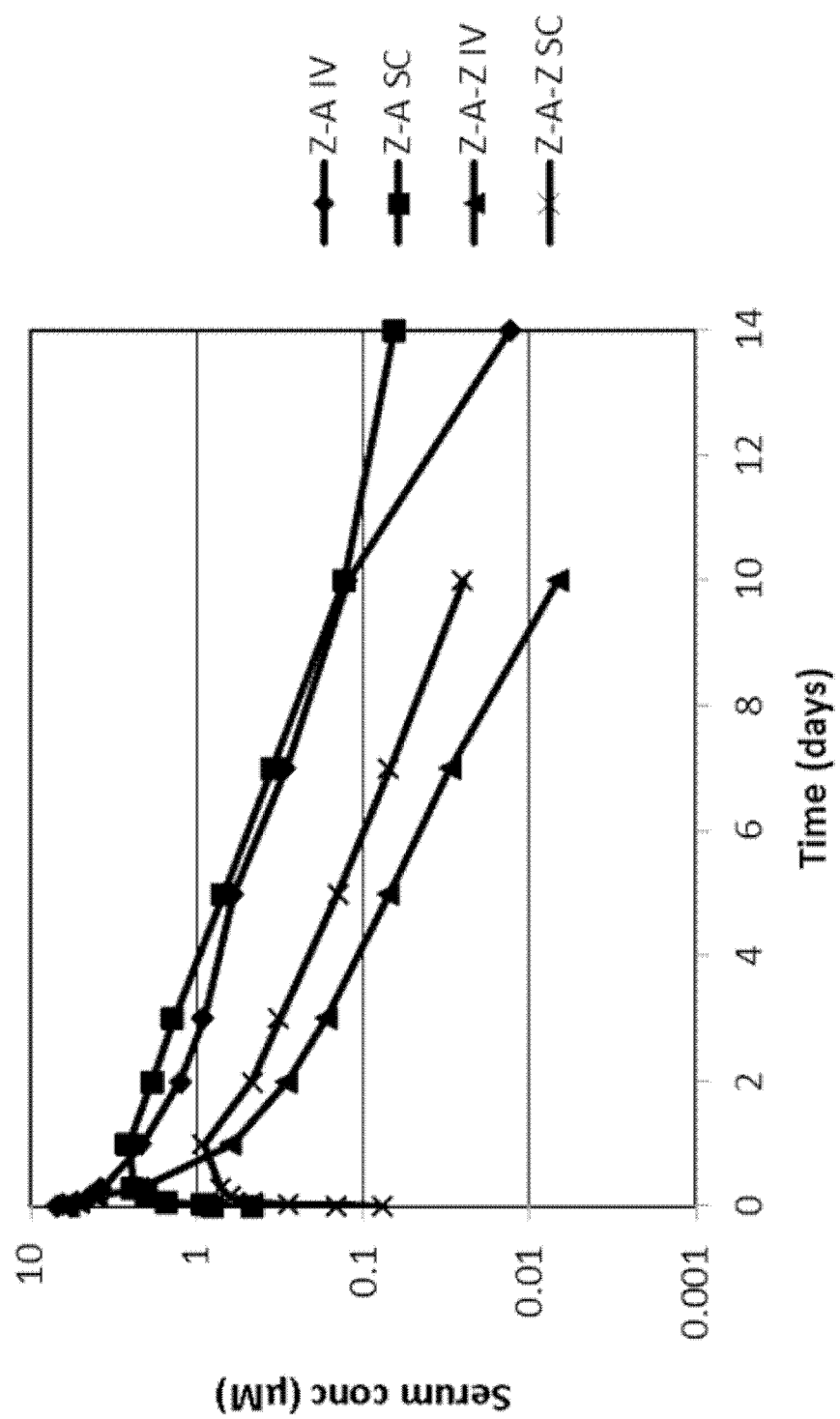


Fig. 10

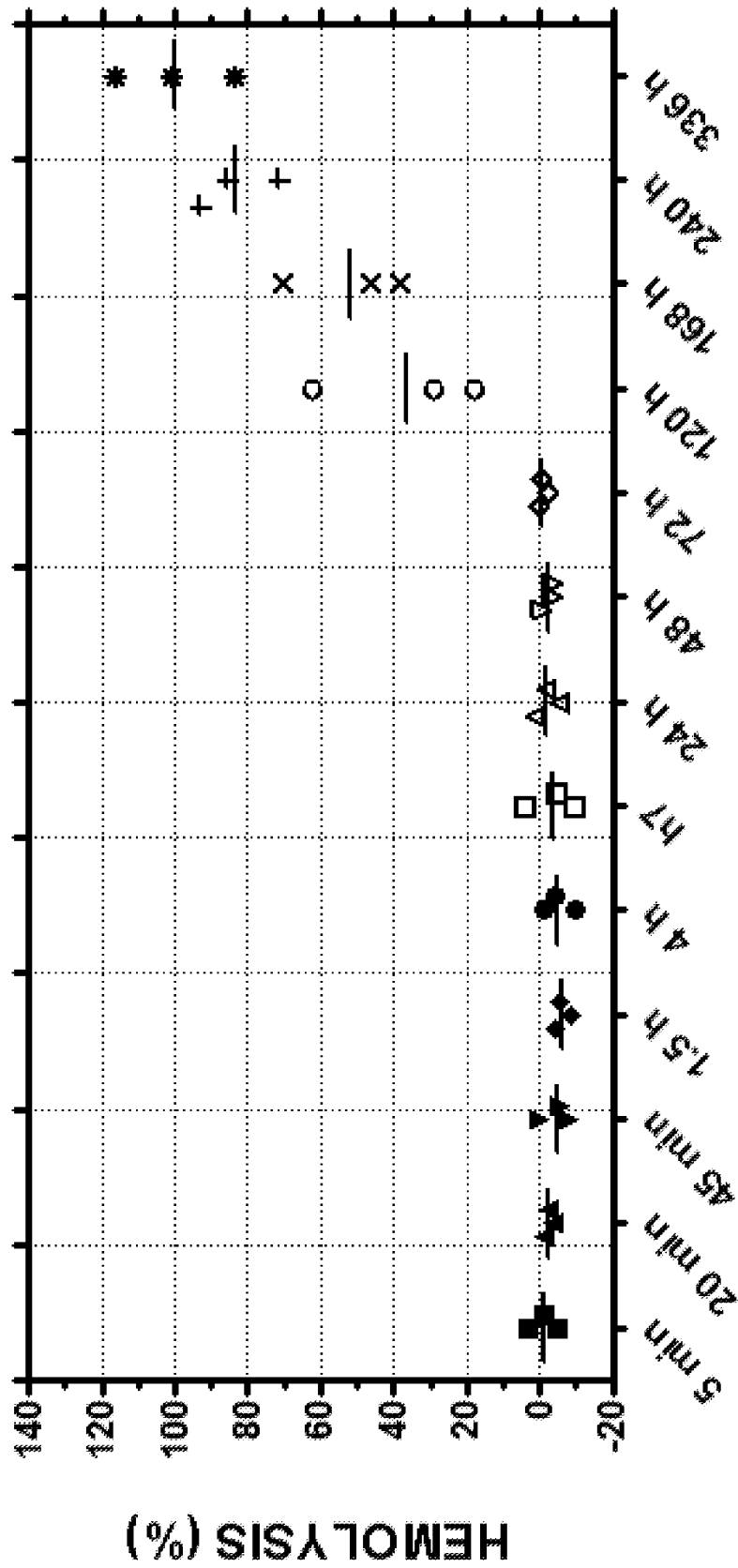


Fig. 11

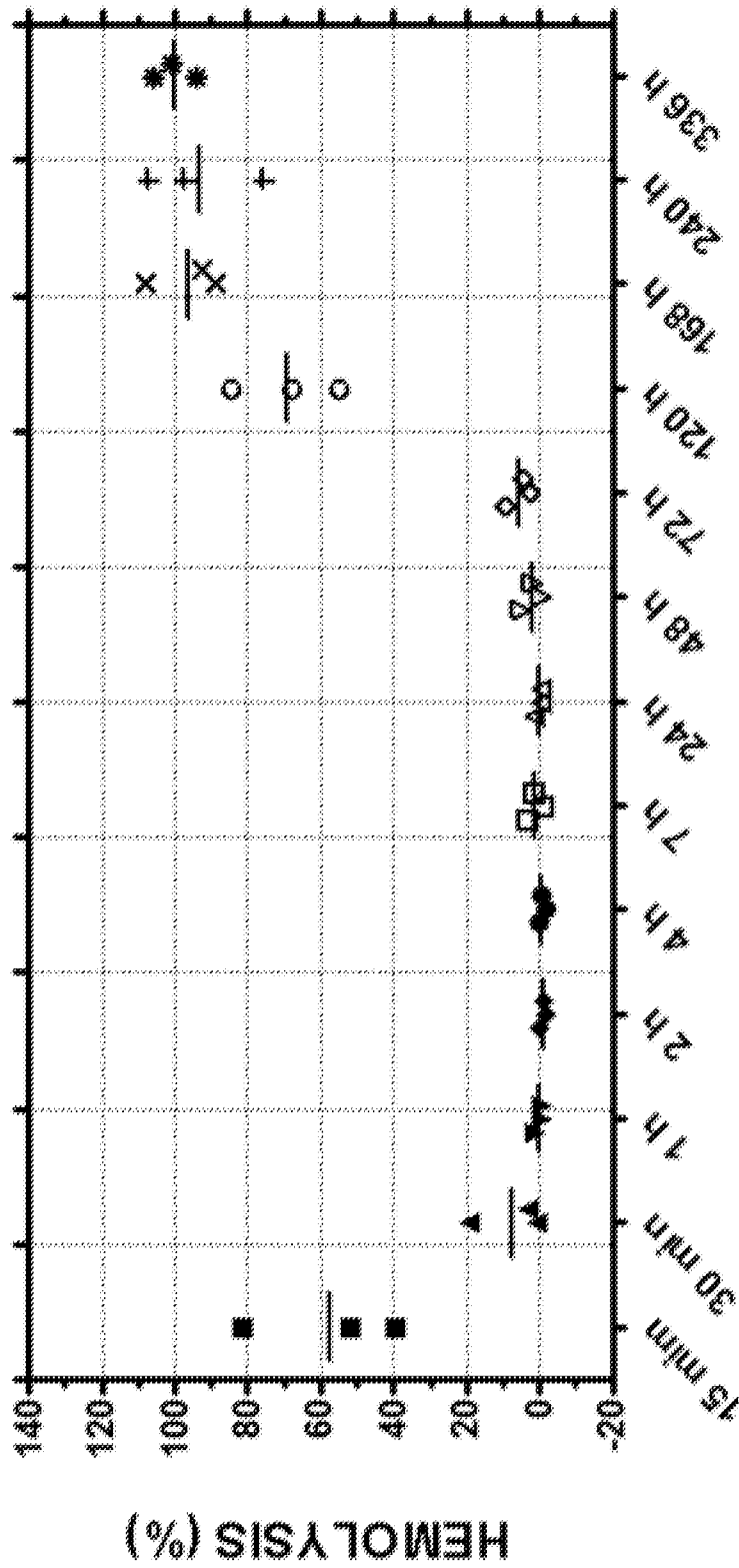


Fig. 12

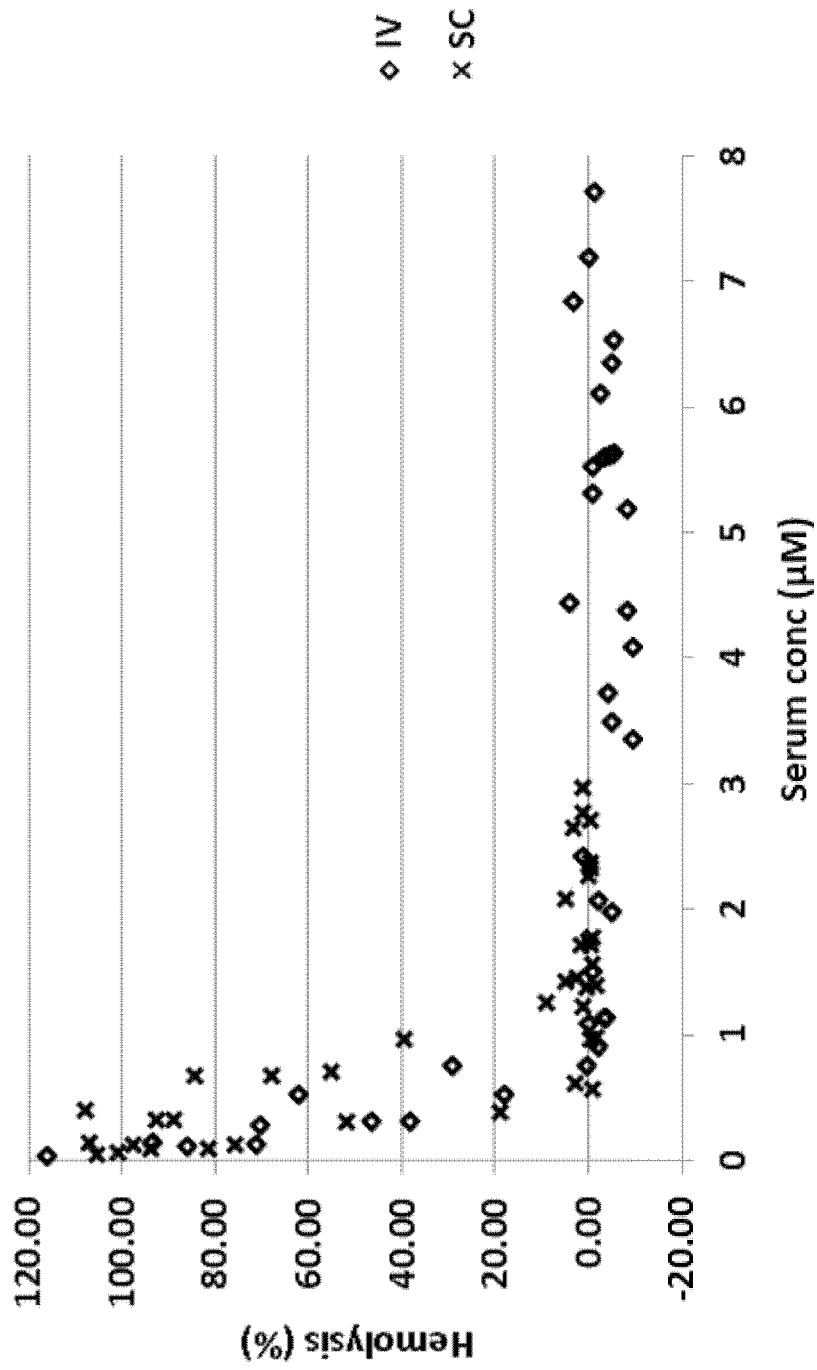


Fig. 13

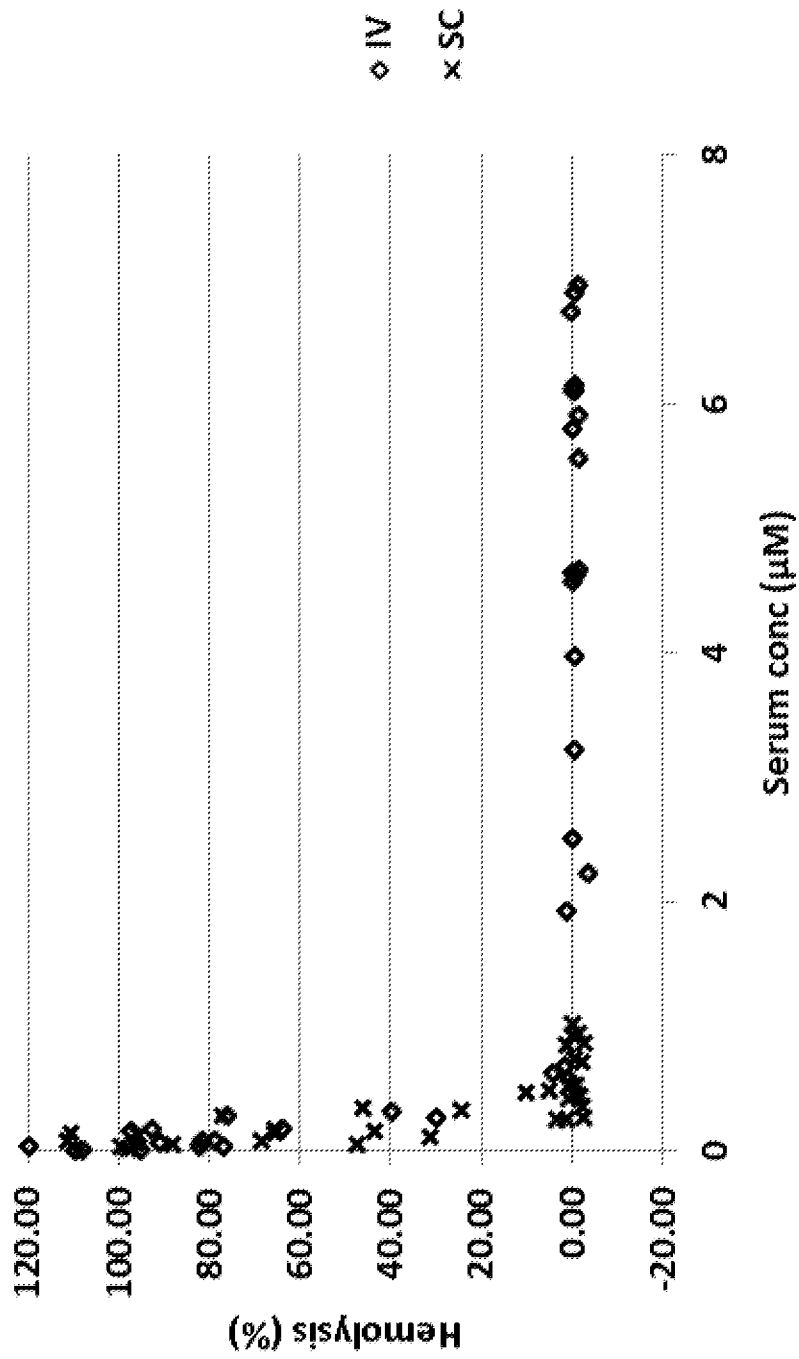
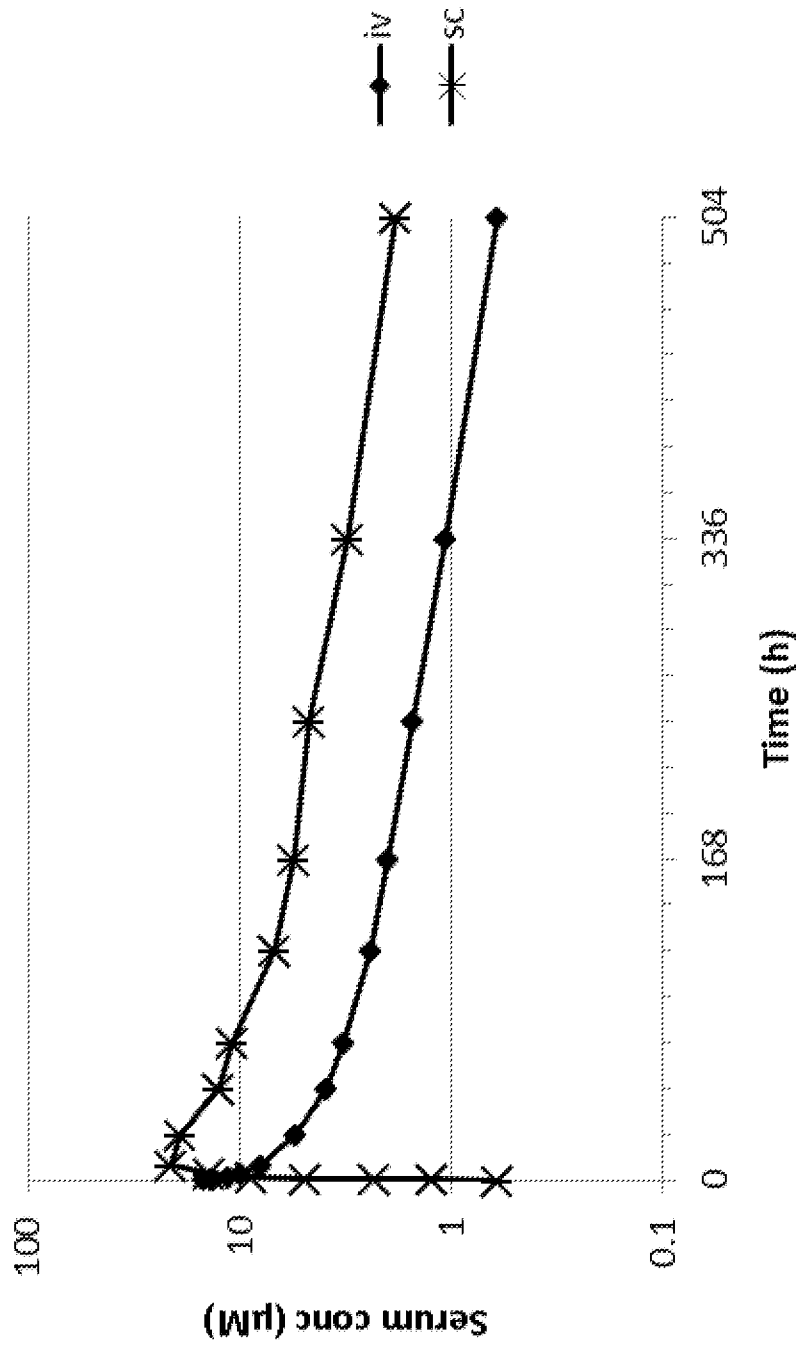


Fig. 14



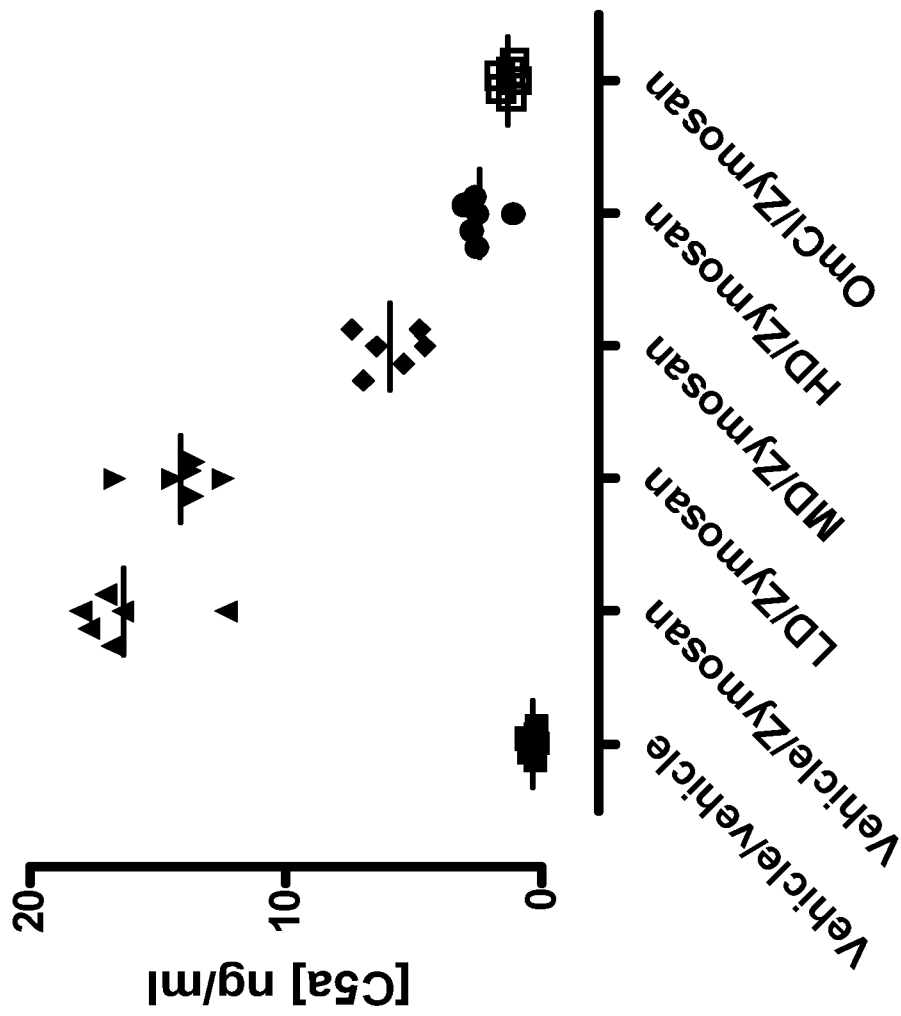


Fig. 16A

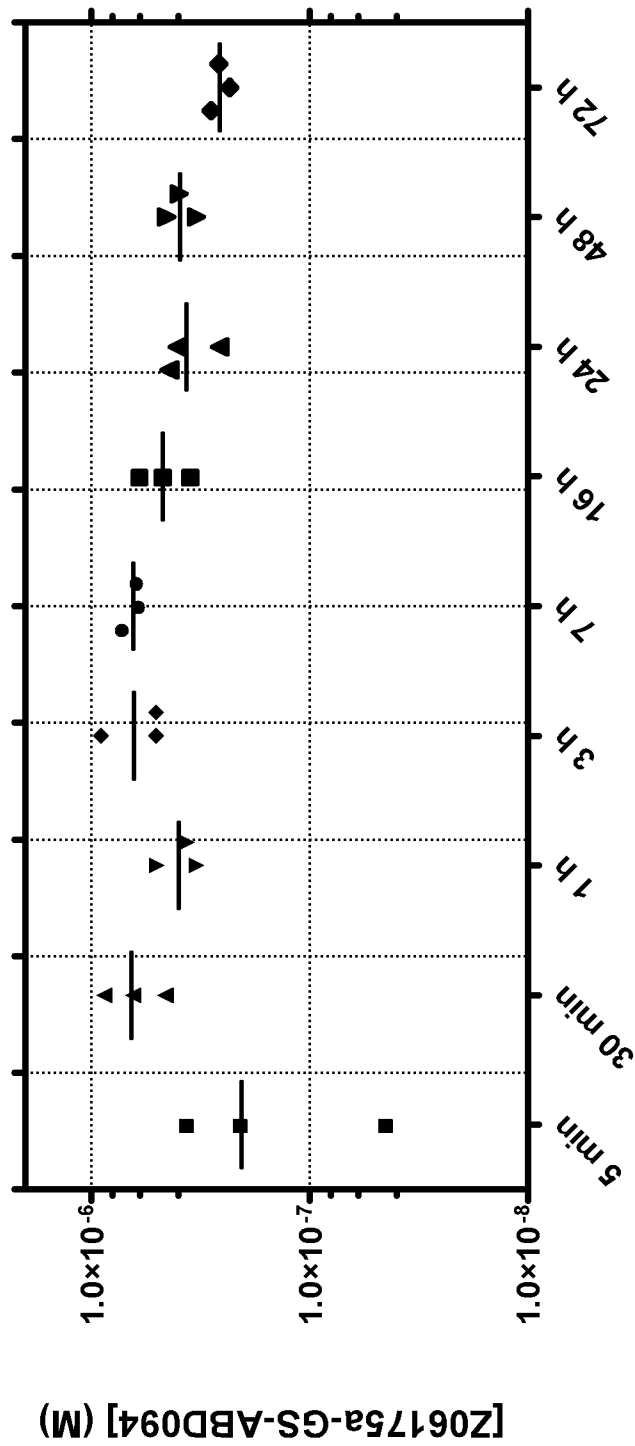


Fig. 16B

