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Patentanmeldung Nr.

Patent application No.

Demande de brevet n°

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Im Auftrag

For the President of the European Patent Office

Le Président de l'Office européen des brevets
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V. Joseph

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COMPSTATIN ANALOGUES AND THEIR MEDICAL USES

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COMPSTATIN ANALOGUES AND THEIR MEDICAL USES

Field of the Invention

5 The present invention relates to inhibiting activation of the complement cascade in the body, and more particularly to compstatin analogues that are capable of binding to C3 protein and inhibiting complement activation. The present invention also relates to the medical uses of the compstatin analogues, in particular for the treatment of conditions characterized by unwanted activation of the complement cascade, such as autoimmune and inflammatory
10 diseases.

Background of the Invention

The human complement system is a powerful player in the defense against pathogenic organisms and the mediation of immune responses. Complement can be activated through
15 three different pathways: the classical, lectin and alternative pathways. The major activation event that is shared by all three pathways is the proteolytic cleavage of the central protein of the complement system, C3, into its activation products C3a and C3b by C3 convertases. Generation of these fragments leads to the opsonization of pathogenic cells by C3b and iC3b, a process that renders them susceptible to phagocytosis or clearance, and to the activation of
20 immune cells through an interaction with complement receptors (Markiewski & Lambris, 2007, Am. J. Pathol., 171: 715-727). Deposition of C3b on target cells also induces the formation of new convertase complexes and thereby initiates a self-amplification loop. An ensemble of plasma and cell surface-bound proteins carefully regulates complement activation to prevent host cells from self-attack by the complement cascade. However, excessive activation or
25 inappropriate regulation of complement can lead to a number of pathologic conditions, ranging from autoimmune to inflammatory diseases (Holers, 2003, Clin. Immunol., 107: 140-51; Markiewski & Lambris, 2007, supra; Ricklin & Lambris, 2007, Nat. Biotechnol., 25: 1265-75; Sahu et al., 2000, J. Immunol., 165: 2491-9). The development of therapeutic complement inhibitors is therefore highly desirable. In this context, C3 and C3b have
30 emerged as promising targets because their central role in the cascade allows for the simultaneous inhibition of the initiation, amplification, and downstream activation of complement (Ricklin & Lambris, 2007, supra).

Human compstatin was first identified as a 27 amino acid peptide and was the first non-host-
35 derived complement inhibitor that was shown to be capable of blocking all three activation pathways (Sahu et al., 1996, J. Immunol., 157: 884-91; U.S. Patent No: 6,319,897). It has been shown that it is possible to truncate compstatin without loss of activity to a 13 amino

acid peptide. However, attempts to further truncate this peptide led to loss of activity. The sequence of the 13 amino acid truncated compstatin peptide is Ile¹-Cys²-Val³-Val⁴-Gln⁵-Asp⁶-Trp⁷-Gly⁸-His⁹-His¹⁰-Arg¹¹-Cys¹²-Thr¹³-NH₂, where Cys² and Cys¹² are disulfide bonded. This cyclic tridecapeptide binds to C3 (and fragments of C3), thereby inhibiting the activation of the downstream complement cascade and preventing the cleavage of native C3 by the C3 convertases. Its inhibitory efficacy was confirmed by a series of studies using experimental models that pointed to its potential as a therapeutic agent (Fiane et al, 1999a, Xenotransplantation, 6: 52-65; Fiane et al., 1999b, Transplant Proc., 31:934- 935; Nilsson et al., 1998, Blood, 92: 1661-1667; Ricklin & Lambris, 2008, Adv. Exp. Med..Biol., 632: 273-292; Schmidt et al., 2003, J. Biomed. Mater. Res., A66: 491-499; Soulika et al., 2000, Clin. Immunol., 96: 212-221).

Progressive optimization of the 13 amino acid compstatin peptide has led to analogues with improved biological activity (Ricklin & Lambris, 2008, supra; WO 2004/026328; WO 2007/062249, WO 2013/036778, WO 2014/100407).

Earlier structure-activity studies have identified the cyclic nature of the compstatin peptide and the presence of both a β -turn and hydrophobic cluster as key features of the molecule (Morikis et al., 1998, Protein Sci., 7: 619-627; WO 99/13899; Morikis et al., 2002, J. Biol. Chem., 277:14942-14953; Ricklin & Lambris, 2008, supra). Hydrophobic residues at positions 4 and 7 were found to be of particular importance, and their modification with unnatural amino acids generated an analogue with 264-fold improved activity over the original compstatin peptide (Katragadda et al., 2006, J. Med. Chem., 49: 4616-4622; WO 2007/062249). Further attempts to optimize compstatin for use in the treatment of eye disorders are described in WO 2007/044668.

While previous optimization steps have been based on combinatorial screening studies, solution structures, and computational models (Chiu et al., 2008, Chem. Biol. Drug Des., 72: 249-256; Mulakala et al., 2007, Bioorg. Med. Chem., 15: 1638-1644; Ricklin & Lambris, 2008, supra), the publication of a co-crystal structure of compstatin complexed with the complement fragment C3c (Janssen et al., 2007, J. Biol. Chem., 282: 29241-29247; WO 2008/153963) provided a basis for initiating rational optimization. The crystal structure revealed a shallow binding site at the interface of macroglobulin (MG) domains 4 and 5 of C3c and showed that 9 of the 13 amino acids were directly involved in the binding, either through hydrogen bonds or hydrophobic interactions. As compared to the structure of the compstatin peptide in solution (Morikis et al., 1998, supra), the bound form of compstatin experienced a conformational

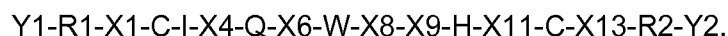
change, with a shift in the location of the β -turn from residues 5-8 to 8-11 (Janssen et al., 2007, *supra*; WO 2008/153963).

In view of its therapeutic potential in AMD, C3G, PNH and other diseases, it remains a problem in the art to further optimize compstatin analogues, for example to achieve an even greater activity and/or to modulate pharmacokinetic properties, such as increased half-life in vivo and/or physicochemical properties such as increased solubility.

Summary of the Invention

Broadly, the present invention is based on work to develop a new family of compstatin analogues having improved binding and complement-inhibiting activity as compared to the 13 amino acid compstatin peptide (ICVVQDWGHRCT (cyclic C2-C12)). In some cases, these compstatin analogues additionally possess useful physicochemical properties, such as increased solubility. In particular, the present inventors found that introducing an isoleucine residue at position 3 in place of the wild type valine residue led to compstatin peptides with improved binding and complement-inhibiting activity. The present inventors further discovered that the introduction of isoleucine at position 3 was compatible with other modifications, for example modifications that are capable of increasing solubility, such as the introduction of charged or polar amino acids at position 9 and/or the introduction of N-and/or C-terminal sequences. Example of such additional modifications include the replacement of Ile at position 1 with Tyr, Phe or Sar, replacement of Val at position 4 with Trp, a Trp analogue (as described herein); replacement of His at position 9 with Ala, Glu, Asp, Lys, Ser or Arg; replacement of Arg at position 11 with Ser; replacement of Thr at position 13 with Ser, Glu, Sar or Ile. Preferred compstatin peptides including one or more of these modifications have improved solubility, for example as compared to the 13 amino acid compstatin peptide (ICVVQDWGHRCT (cyclic C2-C12)). Further examples of these compstatin peptides combine modification at position 9 with extensions to the N-terminal and/or C-terminus of the peptide.

Accordingly, the present invention provides a compstatin analogue represented by the formula:



5

wherein:

Y1 is hydrogen, acetyl or a lipophilic group Φ ;

10 X1 is I, Y, F or Sar;

X4 is W, F, V, Y, 1MeTrp, D-Trp, N-Me-Trp, 1-For-Trp, 1-Nal, 2-Nal, 5MeTrp, Bpa or 2Igl;

X6 is E, K or D;

15

X8 is G or Sar;

X9 is H, A, E, D, K, R or S;

20 X11 is R, S or K;

X13 is T, S, E, F, H, K, Sar, G, I, D, N-Me-Ile or N-Me-Thr;

Y2 is NH₂, OH or a lipophilic group Φ ;

25

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof; and

30 R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof;

35 wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

and wherein the compstatin analogue optionally has a lipophilic group Φ covalently linked to the side chain of one or more amino acid residues;

or a pharmaceutically acceptable salt and/or solvate thereof.

5

In some embodiments, X11 is R or S.

In some embodiments, if a lipophilic group Φ is linked to the side chain of an amino acid residue, that residue is the residue at position X1, X11 or X13, or is a residue in R1 or R2. It may be a lysine residue. For example, it may be a lysine residue at position X11 or X13, or a lysine residue in R1 or R2.

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In some embodiments, Y1 is hydrogen or acetyl.

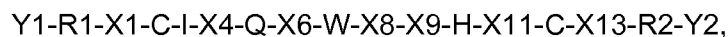
In some embodiments, the compstatin analogue comprises at least one lipophilic group Φ , e.g. exactly one lipophilic group Φ .

15

In some embodiments, the compstatin analogue does not comprise a lipophilic group Φ .

The present invention further provides a compstatin analogue represented by the formula:

20



wherein:

25

Y1 is hydrogen, acetyl, or a lipophilic group Φ ;

X1 is I, Y, F or Sar;

X4 is W, V, Y, 2-Nal, 1-Nal or 1MeTrp;

30

X6 is E or D;

X8 is G or Sar;

35

X9 is A, E, D, K or S;

X11 is R, S or K;

X13 is T, S, E, I, Sar, K, G or N-Me-Ile;

5 Y2 is NH₂, OH or a lipophilic group Φ ;

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof, or Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof; and

10

R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg3 or Peg4, or 8-aminooctanoyl, or derivatives thereof;

15 wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

and wherein the compstatin analogue optionally has a lipophilic group Φ covalently linked to the side chain of one or more amino acids;

20

or a pharmaceutically acceptable salt and/or solvate thereof.

In some embodiments, X11 is R or S.

25 In some embodiments, if a lipophilic group Φ is linked to the side chain of an amino acid residue, that residue is the residue at position X1, X11 or X13, or is a residue in R1 or R2. It may be a lysine residue. For example, it may be a lysine residue at position X13, or a lysine residue in R1 or R2.

30 In some embodiments, Y1 is hydrogen or acetyl.

In some embodiments, the compstatin analogue comprises at least one lipophilic group Φ , e.g. exactly one lipophilic group Φ .

35 In some embodiments of this formula, the compstatin analogue does not comprise a lipophilic group Φ .

The present invention further provides a compstatin analogue represented by the formula:



5 wherein:

Y1 is hydrogen, acetyl or a lipophilic group Φ ;

X1 is I, Y, F or Sar;

10

X4 is W, V, Y, 1-Nal, 2-Nal or 1MeTrp;

X6 is E or D;

15 X9 is A, E, D, K or S;

X11 is R, S or K;

X13 is T, I, S, E, K or Sar;

20

Y2 is NH_2 , OH or a lipophilic group Φ ;

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof, or

25 Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof; and

R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg3 or Peg4, or 8-aminooctanoyl, or derivatives thereof;

30

wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

and wherein the compstatin analogue optionally has a lipophilic group Φ covalently linked to the side chain of one or more amino acids;

35

or a pharmaceutically acceptable salt and/or solvate thereof.

In some embodiments, X11 is R or S.

5 In some embodiments, if a lipophilic group Φ is linked to the side chain of an amino acid residue, that residue is the residue at position X1, X11 or X13, or is a residue in R1 or R2. It may be a lysine residue. For example, it may be a lysine residue at position X11 or X13, or a lysine residue in R1 or R2.

10 In some embodiments, Y1 is hydrogen or acetyl.

In some embodiments, the compstatin analogue comprises at least one lipophilic group Φ , e.g. exactly one lipophilic group Φ .

15 In some embodiments of this formula, the compstatin analogue does not comprise a lipophilic group Φ .

The compstatin analogue may be represented by the formula:

Y1-R1-X1-C-I-X4-Q-X6-W-G-X9-H-R-C-X13-R2-Y2,

20 wherein:

Y1 is hydrogen, acetyl or a lipophilic group Φ ;

X1 is I, Y, F or Sar;

25

X4 is W, V, Y, 1-Nal, 2-Nal or 1MeTrp;

X6 is E or D;

30 X9 is A, E, D, K or S;

X13 is T, S, E or Sar;

Y2 is NH_2 , OH or a lipophilic group Φ ;

35

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof, or Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof; and

- 5 R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg3 or Peg4, or 8-aminooctanoyl, or derivatives thereof;

10 wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

and wherein the compstatin analogue optionally has a lipophilic group Φ covalently linked to the side chain of one or more amino acids;

- 15 or a pharmaceutically acceptable salt and/or solvate thereof.

20 In some embodiments, if a lipophilic group Φ is linked to the side chain of an amino acid residue, that residue is the residue at position X1, X11 or X13, or is a residue in R1 or R2. It may be a lysine residue. For example, it may be a lysine residue at position X13, or a lysine residue in R1 or R2.

In some embodiments, Y1 is hydrogen or acetyl.

25 In some embodiments, the compstatin analogue comprises at least one lipophilic group Φ , e.g. exactly one lipophilic group Φ .

In some embodiments of this formula, the compstatin analogue does not comprise a lipophilic group Φ .

30 In some embodiments of the formulae above, X6 is D.

In one aspect, compstatin analogues which do not possess a lipophilic group Φ may be represented by the formula:

35 Y1-R1-X1-C-I-X4-Q-D-W-G-X9-H-R-C-X13-R2-Y2,

wherein:

Y1 is hydrogen or acetyl;

X1 is Y or F ;

5

X4 is W, Y, 1MeTrp;

X9 is A, E or K;

10 X13 is T, E or Sar;

Y2 is NH₂ or OH;

15 R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γGlu, βAsp, or βAla, or a corresponding D form thereof, or Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof; and

20 R2 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γGlu, βAsp, or βAla, or a corresponding D form thereof; or Peg3 or Peg4, or 8-aminooctanoyl, or derivatives thereof;

wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

25 or a pharmaceutically acceptable salt and/or solvate thereof.

The compstatin analogue may be represented by the formula:

Y1-R1-X1-C-I-1MeTrp-Q-D-W-G-E-H-R-C-X13-R2-Y2,

30

wherein:

Y1 is hydrogen or acetyl;

X1 is Y or F;

35

X13 is T, E or Sar;

Y2 is NH₂ or OH;

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γGlu, βAsp, or βAla, or a corresponding D form thereof, or

5 Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof; and

R2 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γGlu, βAsp, or βAla, or a corresponding D form thereof; or Peg3 or Peg4, or 8-aminooctanoyl, or derivatives thereof;

10

wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

or a pharmaceutically acceptable salt and/or solvate thereof.

15

In some embodiments, the compstatin analogue has the formula:

Y1-R1-X1-C-I-X4-Q-X6-W-X8-X9-H-X11-C-X13-R2-Y2

20

wherein:

Y1 is hydrogen, acetyl or a lipophilic group Φ;

X1 is I, Y, F or Sar;

25

X4 is W, V, 1MeTrp, 1-Nal or 2-Nal;

X6 is E, K or D;

30

X8 is G or Sar;

X9 is H, A, E, D, K, R or S;

X11 is R, S, K or K*;

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X13 is T, S, E, Sar or N-Me-Ile;

Y2 is NH₂ or OH;

R1 and R2 may be as defined in any of the formulae above, or elsewhere in this specification.

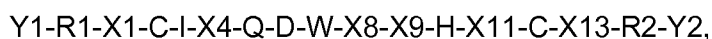
In some embodiments, R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, K, K*, S, Y, or a corresponding D form thereof; and/or R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, K, K*, P, S, Peg3, γGlu, 8-aminooctanoyl, or a corresponding D form thereof;

wherein * indicates that the amino acid residue bears a lipophilic group Φ covalently linked to its side chain.

10

It may be desirable that the compstatin analogue comprises at least one lipophilic group Φ, e.g. exactly one lipophilic group Φ. Alternatively, it may comprise no lipophilic group Φ.

In an alternative aspect, compstatin analogues which comprise a lipophilic group Φ may be represented by formula:



wherein:

20 Y1 is hydrogen, acetyl or a lipophilic group Φ;

X1 is I, Y, F or Sar;

X4 is W, V, Y, 2-Nal, 1-Nal or 1MeTrp;

25

X8 is G or Sar;

X9 is A, E, D, K or S;

30 X11 is R, S or K*;

X13 is T, S, E, I, Sar, K, G or N-Me-Ile;

Y2 is NH₂, OH or a lipophilic group Φ;

35

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, K*, F, P, S, T, W, Y, R, V or Sar, or a corresponding D form thereof;

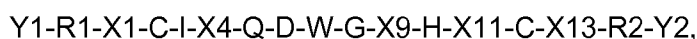
R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg 3 or Peg4, or 8-aminooctanoyl, or derivatives thereof; wherein * indicates that the amino acid residue bears a lipophilic group Φ covalently linked to its side chain;

wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

and wherein the compstatin analogue comprises at least one lipophilic group Φ , e.g. exactly one lipophilic group Φ ;

or a pharmaceutically acceptable salt and/or solvate thereof.

The compstatin analogue may be represented by the formula:



wherein:

Y1 is hydrogen, acetyl, or a lipophilic group Φ ;

X1 is I, Y, F or Sar;

X4 is W, V, Y, 1-Nal, 2-Nal or 1MeTrp;

X9 is A, E, D, K or S;

X11 is R, S or K*;

X13 is T, I, S, E, K or Sar;

Y2 is NH_2 , OH or a lipophilic group Φ ;

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, K*, F, P, S, T, W, Y, R, V or Sar, or a corresponding D form thereof;

R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, K* F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg 3 or Peg4, or 8-aminooctanoyl, or derivatives thereof; wherein * indicates that the amino acid residue bears a lipophilic group Φ covalently attached to its side chain;

5

wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

and wherein the compstatin analogue comprises at least one lipophilic group Φ , e.g. exactly one lipophilic group Φ ;

10

or a pharmaceutically acceptable salt and/or solvate thereof.

The compstatin analogue may be represented by the formula:

15

Y1-R1-X1-C-I-X4-Q-D-W-G-X9-H-R-C-X13-R2-Y2,

wherein:

Y1 is hydrogen, acetyl or a lipophilic group Φ ;

20

X1 is I, Y, F or Sar;

X4 is W, V, 1-Nal, 2-Nal or 1MeTrp;

X9 is A, E, D, K or S;

25

X13 is T, S, E or Sar;

Y2 is NH_2 , OH or a lipophilic group Φ ;

30

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, K*, F, P, S, T, W, Y, R, V or Sar, or a corresponding D form thereof;

R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, K*, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg3 or Peg4, or 8-aminooctanoyl, or derivatives thereof; wherein * indicates that the amino

35

acid residue bears a lipophilic group Φ covalently attached to its amino acid side chain; wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

and wherein the compstatin analogue comprises at least one lipophilic group Φ , e.g. exactly one lipophilic group Φ ;

- 5 or a pharmaceutically acceptable salt and/or solvate thereof.

In some embodiments, the 13-mer peptide portion (X1-X13) of the compstatin analogue has a sequence selected from:

- 10 [Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)[Sar];
 [Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)T;
 [Sar]C(1)I[1-Me-Trp]QEW[Sar]EHRC(1)T;
 [Sar]C(1)I[1-Me-Trp]QEWGEHRC(1)[Sar];
 [Sar]C(1)IWQDWGEHRC(1)T;
 15 FC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)[Sar];
 FC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)T;
 FC(1)I[1-Me-Trp]QDWGEHKC(1)[Sar];
 FC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar];
 FC(1)I[1-Me-Trp]QDWGEHRC(1)E;
 20 FC(1)I[1-Me-Trp]QDWGEHRC(1)S;
 FC(1)I[1-Me-Trp]QDWGEHRC(1)T;
 FC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar];
 FC(1)I[1Na]QDWGEHRC(1)T;
 FC(1)I[2Na]QDWGEHRC(1)T;
 25 FC(1)IWQDWGEHRC(1)[Sar];
 FC(1)IWQDWGEHRC(1)T;
 IC(1)I[1-Me-Trp]QDW[Sar]AHRC(1)[N-Me-Ile];
 IC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar];
 IC(1)I[1-Me-Trp]QDWGEHRC(1)T;
 30 IC(1)I[2Na]QDWGEHRC(1)[Sar];
 IC(1)IWQDWGAHRC(1)E;
 IC(1)IWQDWGAHRC(1)T;
 IC(1)IWQDWGAHSC(1)T;
 IC(1)IWQDWGDHRC(1)T;
 35 IC(1)IWQDWGEHRC(1)[Sar];
 IC(1)IWQDWGEHRC(1)E;
 IC(1)IWQDWGEHRC(1)S;

IC(1)IWQDWGEHRC(1)T;
 IC(1)IWQDWGEHSC(1)T;
 IC(1)IWQDWGKHRC(1)T;
 IC(1)IWQDWGRHRC(1)T;
 5 IC(1)IWQDWGSHRC(1)T;
 IC(1)IWQEWGEHRC(1)T;
 IC(1)IWQKWGAHRC(1)T;
 IC(1)IWQKWGEHRC(1)T;
 YC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar];
 10 YC(1)I[1-Me-Trp]QDWGEHRC(1)T;
 YC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar];
 YC(1)I[2NaI]QDWGEHRC(1)T;
 YC(1)IWQDWGEHRC(1)T;
 YC(1)I[1-Me-Trp]QDWGEH[K*]C(1)[Sar]; and
 15 YC(1)I[1-Me-Trp]QEW[Sar]EHRC(1)[Sar].

In some embodiments, R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, R, V or Sar, or a corresponding D form thereof, and/or R2 may be a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, R, V or Sar, or a corresponding D form thereof.

For example, R1 is selected from ESSA, AKGE, ASSE, ASES, GSAE, ESSE, ESGA, SEG, GES, ESS, EGSA, ESE, EGE, ESA, SAE, SGA, YLEA, GSA, KEK, EKG, ES, AS, SE, SA or E, and/or R2 is selected from GAES, EYGS, EGYA, EAGS, EAKS, EKSA, EGGS, EGGA, ESSG, ESAG, GEES, AEES, ESEG, AEGS, ESGS, SEGA, SEG, ESG, EAG, GAE, EGEA, EGE, EA, E, GE, EG, EKE or EKP.

In alternative embodiments, R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof, or Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof.

In some embodiments, R1 is absent or is a sequence of 1 to 6 amino acid residue selected from A, E, G, L, K, F, P, S, T, W, Y, R, V or Sar, or a corresponding D form thereof.

For example, R1 may be absent or a sequence of 1 to 6 amino acid residues selected from A, E, G, K, S and Y, or a corresponding D-form thereof.

A lipophilic group Φ may be covalently linked to the side chain of one or more of the residues in Y1, especially to the side chain of a lysine residue (which may be designated K*). It may be desirable that the residue bearing Φ is at the N-terminus of Y1.

5 Examples of sequences for the group R1 include:

{d}Y, EGSE, AGSE, SASE, EYSE, GSE, ASE, ESSA, KGSA, AKGE, ASGE, ASSE, ASES, GSAE, ESSE, ESGA, SEG, GES, ESS, EGSA, ESE, EGE, ESA, SAE, SGA, YLEA, GSA, KEK, EKG, ES, RS, SR, AE, TE, KE, GE, FE, YE, AS, SE, RS, SR, SA, GE, S, Y and E.

10 In some embodiments, R1 is two amino acid residues in length, for example, AE, TE, KE, GE, FE, YE, AS, SE, SA, or GE; preferably AE, TE, KE, GE, FE, YE, SE, or GE.

As mentioned above, a lipophilic group Φ may be covalently linked to the side chain of one or more of the residues in Y1, especially to the side chain of a lysine residue (which may be

15 designated K*), e.g. to yield a sequence K*GSA.

R2 may be absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof.

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For example, R2 may be absent or a sequence of 1 to 8 amino acid residues selected from A, E, G, K, S, γ Glu, Peg3 or 8-aminooctanoyl or selected from A, E, G, K and S.

When K is present in R2, it may be desirable that K is present at the C-terminus of R2.

25

A lipophilic group Φ may be covalently linked to the side chain of one or more of the residues in Y2, especially to the side chain of a lysine residue. It may be desirable that the residue bearing Φ is at the C-terminus of Y2.

30 Examples of sequences for the group R2 include:

EGASGSG, EGAGSG, EGASAG, EGAGAG, EGESGSG, EGE GSG, EGESAG, EGE GAG, EK[γ Glu]AK, EGE-Peg3-Peg3-K, EGE G, EGAGG, EGE S, GAESK, EGAK, EGEK, EGG, EGK, EGKK, EGS, EK, EGA, EK[γ Glu], EK[γ Glu]K, EGE-Peg3, EGE-Peg3-K, EGE-Peg3-Peg3, EGE-Peg3-Peg3-K, EGE-Peg3-Peg3-Peg3, GESESE, GAESSES, EGESES, EGESESK, EGE-Peg3-ES, EGE-Peg3-ESK, GESESE, EGE-[8-aminooctanoyl], EGE-[8-aminooctanoyl]-K, EGE-[8-aminooctanoyl]-EK, EGE GGG, EGE GGGK, EK[γ Glu]GGG, EK[γ Glu]GGGK, EGE-[8-aminooctanoyl]-E, GAES, EYGS, EGYA, EAGS, EAKS, EKSA,

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ESGA, EGGS, EGGA, ESSG, ESAG, GEES, AEES, ESEG, AEGS, ESGS, SEGA, SEG, EGK, EGA, ESG, EAG, GAE, EGEA, EGE, EA, E, S, GE, GEK, EG, EKE and EKP.

As mentioned above, a lipophilic group Φ may be covalently linked to the side chain of one or more of the residues in Y2, especially the side chain of a lysine residue, e.g. to yield a sequence EK[γ Glu]AK*, EGKK, EK[γ Glu]K*, EGE-Peg3-K*, EGE-Peg3-Peg3-K*, EGESESK*, EGE-Peg3-ESK*, EGE-[8-aminooctanoyl]-K*, EGE-[8-aminooctanoyl]-EK*, EGE GGGK*, EK[γ Glu]GGGK*, EGE-Peg3-Peg3-K*, GAESK*, EGAK*, EGEK*, EGK* EGE-Peg3-ESK*, GESESEK*, GEK* or EK*.

Where R1 or R2 is one amino acid in length, it may be a D amino acid, e.g. {d}Y.

R1 and R2 may independently be present or absent. It may be desirable that R2 is present. Without wishing to be bound by any particular theory, it is believed that the presence of R1 and/or R2 may improve the stability of the compounds.

Preferred classes of compstatin analogues and exemplified compounds are discussed further below.

In a further aspect, the present invention provides a composition comprising a compstatin analogue of the present invention, or a pharmaceutically acceptable salt or solvate thereof, in admixture with a carrier. In some instances, the composition is a pharmaceutical composition and the carrier is a pharmaceutically acceptable carrier.

In a further aspect, the present invention provides a pharmaceutical composition comprising a compstatin analogue of the present invention, or a pharmaceutically acceptable salt or solvate thereof, in admixture with a pharmaceutically acceptable carrier, excipient or vehicle.

In a further aspect, the present invention provides a compstatin analogue of the present invention for use in therapy.

In a further aspect, the present invention provides a compstatin analogue of the present invention for use in a method of inhibiting complement activation. By way of example, inhibiting complement activation includes one or more biological activities selected from (1) binding to C3 protein, (2) binding to C3b protein and/or (3) inhibiting the cleavage of native C3 by C3 convertases. Examples of disease or condition that may be treated using the compstatin analogues of the present invention are discussed below.

In a further aspect, the present invention provides a compstatin analogue of the present invention for use in a method of inhibiting complement activation that occurs during cell or organ transplantation.

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In a further aspect, the present invention provides a method of inhibiting complement activation for treating a subject in need thereof, the method comprising administering to the subject a compstatin analogue of the present invention thereby to inhibit complement activation in the subject. Examples of disease or condition that may be treated using the compstatin analogues of the present invention are discussed below.

10

In a further aspect, the present invention provides an *ex vivo* method of inhibiting complement activation during extracorporeal shunting of a physiological fluid, the method comprising contacting the physiological fluid with a compstatin of the present invention, thereby to inhibiting complement activation.

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In a further aspect, the present invention provides the use of a compstatin analogue of the present invention in the preparation of a medicament for inhibiting complement activation. Examples of disease or condition that may be treated using the compstatin analogues of the present invention are discussed below.

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Embodiments of the present invention will now be described by way of example and not limitation.

Description of the Figures

Figure 1 (a-d): Normalized “ex vivo” activity of the alternative complement pathway over time after administration of test compounds at time 0 to one or two non-human primates. Compounds were given subcutaneously at a dose of 1840 nmol/kg. Complement activity was measured using the Wieslab kit. Activity was normalized using the predose (0) sample and the negative control included in the kit and set to 100%.. Normalized activity or Average normalized activity for both animals and standard deviation is shown. (a) compound 61 (2 animals), (b) compound 123, compound 126 & comp 128, all with one animals per compound, (c) compound 107, compound 111, compound 118 & compound 119 all with 2 animals per compound, and (d) compound 94, compound 96, compound 104 & compound 106 all with 2 animals per compound.

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Detailed Description

“and/or” where used herein is to be taken as specific disclosure of each of the two specified features or components with or without the other. For example “A and/or B” is to be taken as specific disclosure of each of (i) A, (ii) B and (iii) A and B, just as if each is set out individually herein.

Unless context dictates otherwise, the descriptions and definitions of the features set out above are not limited to any particular aspect or embodiment of the invention and apply equally to all aspects and embodiments which are described.

Various publications, including patents, published applications, technical articles and scholarly articles are cited throughout the specification. Each of these cited publications is incorporated by reference herein, in its entirety.

Unless otherwise defined herein, scientific and technical terms used in this application shall have the meanings that are commonly understood by those of ordinary skill in the art. Generally, nomenclature used in connection with, and techniques of, chemistry, molecular biology, cell and cancer biology, immunology, microbiology, pharmacology, and protein and nucleic acid chemistry, described herein, are those well known and commonly used in the art.

Each embodiment of the invention described herein may be taken alone or in combination with one or more other embodiments of the invention.

Unless specified otherwise, the following definitions are provided for specific terms which are used in the present written description.

Definitions

Throughout this specification, the word “comprise”, and grammatical variants thereof, such as “comprises” or “comprising”, will be understood to imply the inclusion of a stated integer or component, or group of integers or components, but not the exclusion of any other integer or component, or group of integers or components.

The singular forms “a,” “an,” and “the” include the plurals unless the context clearly dictates otherwise.

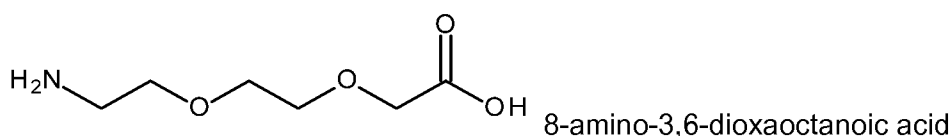
The term “including” is used to mean “including but not limited to”. “Including” and “including but not limited to” may be used interchangeably.

The terms “patient”, “subject” and “individual” may be used interchangeably. A subject may be a mammal, including a human or a non-human mammal, such as a non-human primate (e.g. ape, Old World monkey or New World monkey), livestock animal (e.g. bovine or porcine), companion animal (e.g. canine or feline) or laboratory animal such as a rodent (e.g. mouse or rat).

- Throughout the present description and claims the conventional three-letter and one-letter codes for naturally occurring amino acids are used, i.e. A (Ala), G (Gly), L (Leu), I (Ile), V (Val), F (Phe), W (Trp), S (Ser), T (Thr), Y (Tyr), N (Asn), Q (Gln), D (Asp), E (Glu), K (Lys), R (Arg), H (His), M (Met), C (Cys) and P (Pro); as well as generally accepted three-letter codes for other α -amino acids, such as norleucine (Nle), sarcosine (Sar), α -aminoisobutyric acid (Aib), 2,3-diaminopropanoic acid (Dap), 2,4-diaminobutanoic acid (Dab) and 2,5-diaminopentanoic acid (ornithine; Orn), 1-methyl-tryptophan (1MeTrp), 1-formyl-tryptophan (1-ForTrp), 1-naphthalin (1-Nal), 2-naphthalin (2-Nal), 5-methyl-tryptophan (5MeTrp), p-Benzoyl-phenylalanine (Bpa) 2-indanylglycine (2Igl). Other α -amino acids may be shown in square brackets “[]” (e.g. “[Nle]”) when used in a general formula or sequence in the present specification, especially when the rest of the formula or sequence is shown using the single letter code. The 20 “naturally occurring” amino acids listed above are those which are encoded by the standard genetic code, and may also be referred to as “proteinogenic” amino acids.
- Gamma-Glu and beta-Asp, also referred to as γ -Glu and β -Asp (or isoGlu and isoAsp), refers to glutamate or aspartate participating in peptide bonds via the γ - or β -carboxylic acid respectively (normally regarded as the side chain carboxyl groups), rather than the conventional configuration. Similarly, ϵ Lys or isoLys refers to lysine participating in a peptide bond via the epsilon amino group (normally regarded as the side chain amino group) rather than the alpha amino group.

Beta-Ala, also referred to as β -Ala, refers to 3-aminopropanoic acid.

- Peg3 refers to a residue of 8-amino-3,6-dioxaoctanoic acid (also known as {2-[2-aminoethoxy]ethoxy}acetic acid) and Peg4 refers to a residue of 11-amino-3,6,9-trioxaundecanoic acid. The residue may also be denoted [8-Amino-3,6-dioxaoctanoyl].



Unless otherwise specified, amino acid residues in peptides of the invention are of the L-configuration. However, in some instances, D-configuration amino acids may be

5 incorporated. In the present context, an amino acid code written with a small letter represents the D-configuration of said amino acid, e.g. "k" represents the D-configuration of lysine (K), or a D-configuration amino acid may be written as (d)X or {d}X, where X is the amino acid, e.g. (d)Y or {d}Y represents the D-configuration of tyrosine (Y).

10 Cysteine residues shown as "C(1)" indicate that their side-chains participate in a disulphide bond. Thus there will typically be two such residues in any given molecule.

The terminal groups present at the N- and C-termini of the peptide backbone are designated Y1 and Y2 respectively. Thus Y1 is bonded to the nitrogen atom of the N-terminal amino
15 group and Y2 is bonded to the C-terminal carbonyl carbon atom.

Y1 = hydrogen (also indicated as "H-" or "Hy-") indicates a hydrogen atom, corresponding to the presence of a free primary or secondary amino group at the N-terminus. Y1 = acetyl ("Ac") indicates the presence of an N-terminal secondary acetyl amide group.

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Y2 = "OH" or "NH₂" indicates the presence of a carboxy (COOH) group or an amido (CONH₂) group at the C-terminus of the molecule.

25 Either or both of Y1 and Y2 may alternatively be a lipophilic group Φ . Typically, only one of Y1 or Y2 will be a lipophilic group Φ .

In some embodiments, Y1 is hydrogen or acetyl, and Y2 is OH or NH₂.

30 Various terms relating to the methods and other aspects of the present invention are used throughout the specification and claims. Such terms are to be given their ordinary meaning in the art unless otherwise indicated. Other specifically defined terms are to be construed in a manner consistent with the definition provided herein. The term "about" as used herein when referring to a measurable value such as an amount, a temporal duration, and the like, is meant to encompass variations of $\pm 20\%$ or $\pm 10\%$, in some embodiments $\pm 5\%$, in some
35 embodiments $\pm 1\%$, and in some embodiments $\pm 0.1\%$ from the specified value, as such

variations are appropriate to make and used the disclosed compounds and compositions.

The term "full length compstatin" as used herein refers to a 27 amino acid peptide having the sequence IC(1)VVQDWGHHRC(1)TAGHMANLTSHASAI, wherein C(1) denotes the cysteine residue linked by a disulphide bond. As described above, a truncated form of full length compstatin, the tridecapeptide Ile¹-Cys²-Val³-Val⁴-Gln⁵-Asp⁶-Trp⁷-Gly⁸-His⁹-His¹⁰-Arg¹¹-Cys¹²-Thr¹³-NH₂ linked by a disulphide bond between the cysteine residues at positions 2 and 12 retains the activity of the full length peptide and is referred to herein as "Ac-compstatin"

The term "compstatin analogue" refers to a modified Ac-compstatin comprising substitutions of natural and unnatural amino acids, or amino acid analogs, as well as modifications within or between various amino acids, as described in greater detail herein. When referring to the position of amino acids or analogs within Ac-compstatin or compstatin analogs, the positions are numbered from 1 (Ile in compstatin) to 13 (Thr in compstatin). For example, the Gly residue occupies "position 8." As used to describe the compstatin analogue peptides of the present invention "C(1)" denotes a disulphide bond between the respective cysteine residues in the compstatin analogue.

The terms "pharmaceutically active" and "biologically active" refer to the ability of the compounds of the invention to bind C3 or fragments thereof and inhibit complement activation. The biological activities of compstatin analogs may be measured by one or more of several art-recognized assays, as described in greater detail herein.

As used herein, "L-amino acid" refers to any of the naturally occurring levorotatory alpha-amino acids normally present in proteins or the alkyl esters of those alpha-amino acids. The term "D-amino acid" refers to dextrorotatory alpha-amino acids. Unless specified otherwise, all amino acids referred to herein are L-amino acids.

"Hydrophobic" or "non-polar" are used synonymously herein, and refer to any inter- or intra-molecular interaction not characterized by a dipole.

As used herein, "pharmaceutically-acceptable salts" refers to derivatives of the disclosed compounds wherein the parent compound is modified by making acid or base salts thereof. Examples of pharmaceutically-acceptable salts include, but are not limited to, mineral or organic acid salts of basic residues such as amines; alkali or organic salts of acidic residues such as carboxylic acids; and the like. Thus, the term "acid addition salt" refers to the corresponding salt derivative of a parent compound that has been prepared by the addition of

an acid. The pharmaceutically-acceptable salts include the conventional salts or the quaternary ammonium salts of the parent compound formed, for example, from inorganic or organic acids. For example, such conventional salts include, but are not limited to, those derived from inorganic acids such as hydrochloric, hydrobromic, sulfuric, sulfamic, phosphoric, nitric and the like; and the salts prepared from organic acids such as acetic, propionic, succinic, glycolic, stearic, lactic, malic, tartaric, citric, ascorbic, pantoic, maleic, hydroxymaleic, phenylacetic, glutamic, benzoic, salicylic, sulfanilic, 2-acetoxybenzoic, fumaric, toluenesulfonic, methanesulfonic, ethane disulfonic, oxalic, isethionic, and the like. Certain acidic or basic compounds of the present invention may exist as zwitterions. All forms of the compounds, including free acid, free base, and zwitterions, are contemplated to be within the scope of the present invention.

Compstatin Analogues

Ac-Compstatin, a 13 amino acid peptide, is known to bind to C3 and prevent C3 convertase-mediated cleavage. Since its discovery by phage display, modification to the 13 amino acid Ac-Compstatin sequence has been carried out in an effort to find analogues with increased biological activity. However, in the core sequence between the two cysteines residues at positions 2 and 12, alanine scanning experiments have previously produced analogues showing only modest improvements in biological activity, with few modifications being tolerated. The modifications include changing the valine at position 4 to tryptophan, or a tryptophan analogue, that leads to an increase in biological activity and changing the histidine at position 9 to alanine or analogs thereof.

In particular, previous attempts to introduce modifications to the valine residue at position 3, replacing it with glycine, alanine, D-valine or leucine have been shown to lead to a decrease in biological activity. In contrast to these prior art findings, the present inventors surprisingly found that a change of valine to isoleucine is well tolerated and provides improvements in biological activity, as shown in the Examples below.

Without wishing to be bound by any specific theory, the present inventors reasoned that this modification might be combined with introduction of one or more polar or charged amino acids in the core sequence and may be used as an approach to increase the ability of the compstatin peptides to solubilize. Initially, glutamic acid or serine at position 9 were combined with valine 3 and led to a decrease in activity compared to the reference sequence 4W9A. However, when these changes were combined with the introduction of isoleucine at position 3, a surprising increase in biological activity was observed, in particular for the

combination of isoleucine at position 3 and glutamic acid at position 9. This observation correlates with improved binding to C3 as measured by surface plasmon resonance (SPR).

In a further series of experiments to validate these findings, compstatin peptides with glutamic acid at position 9 are combined with different substitutions in position 3 which would normally be considered “conservative” replacements for isoleucine, again showing that the peptides with isoleucine at position 3 are most active.

Taken together, these experiments show that replacing the valine residue at position 3 with isoleucine surprisingly provides compstatin peptides having increased biological activity and improved binding to C3. Furthermore, the experiments surprisingly demonstrate that these changes can be readily combined with other modifications in the core sequence of the compstatin analogues and with addition of N and C-terminal sequences, for example for improving the solubility of the compstatin peptides, e.g. at higher concentrations.

Introduction of isoleucine instead of valine at position 3 of a further prior art compound designated “Cp40” (Qu et al., Immunobiology 2013, 281(4): 496-505; also referred to in that paper as “peptide 14”) also increased the binding affinity to C3 as measured by SPR.

In any embodiment X1 may be Y, I or F. In any embodiment, X4 may be W, V, 1-Nal, 2-Nal or 1MeTrp. In any embodiment, X6 may be E or D. In any embodiment, X9 may be A, E, D, K or S. In any embodiment, X13 may be T, S, E, I, Sar, K, or G. In any embodiment, X13 may be T, I, S, E, K or Sar. In any embodiment, X13 may be T, S, E or Sar.

Lipophilic substituents

The compstatin analogues may bear a lipophilic group, designated Φ .

The lipophilic group may be covalently linked to the N-terminus and/or the C terminus of the molecule, i.e. Y1 may be Φ (in place of H or Ac) and/or Y2 may be Φ (in place of OH or NH₂).

Additionally or alternatively, the lipophilic group may be covalently linked to the side chain of an amino acid residue within the analogue. The residue may be part of R1, R2 or the compstatin analogue portion X1-X13 of the molecule.

The lipophilic group Φ is typically attached via an acyl group. The modification may therefore be termed acylation but can also be referred to as lipidation.

The lipophilic group includes a long chain alkylene group derived from a fatty acid, termed Z^1 herein and referred to as the lipophilic substituent. Without wishing to be bound by theory, it is believed that a lipophilic substituent binds plasma proteins (e.g. albumin) in the blood stream, thus shielding the compounds employed in the context of the invention from enzymatic degradation, and thereby enhancing the half-life of the compounds. The lipophilic substituent may also modulate the potency of the compound.

Z^1 may be attached directly to the amino acid sequence (including the $R1$ and $R2$ extensions, or as $Y1$) or via a spacer Z^2 as defined herein.

In other words, Φ may be Z^1 - or Z^1 - Z^2 -.

Where $Y1$ is Φ , Φ is preferably Z^1 -.

Where the lipophilic group Φ is linked to an amino acid side chain (i.e. where $Y1$ is hydrogen or Ac) Φ may preferably be Z^1 - Z^2 -.

In certain embodiments, only one amino acid side chain is conjugated to a lipophilic substituent. In other embodiments, two amino acid side chains are each conjugated to a lipophilic substituent. In yet further embodiments, three or even more amino acid side chains are each conjugated to a lipophilic substituent. When a compound contains two or more lipophilic substituents, they may be the same or different substituents.

In certain embodiments, only one lipophilic group Φ is present in the molecule.

The term "conjugated" is used here to describe the covalent attachment of one identifiable chemical moiety to another, and the structural relationship between such moieties. It should not be taken to imply any particular method of synthesis. The one or more spacers Z^2 , when present, are used to provide a spacing between the compound and the lipophilic substituent Z^1 .

A lipophilic substituent may be attached to an N-terminal nitrogen, or to an amino acid side chain or to a spacer via an ester, a sulphonyl ester, a thioester, an amide or a sulphonamide. Accordingly, it will be understood that a lipophilic substituent may include an acyl group, a sulphonyl group, an N atom, an O atom or an S atom which forms part of the ester, sulphonyl ester, thioester, amide or sulphonamide.

Suitably, an acyl group in the lipophilic substituent forms part of an amide or ester with the N-terminal nitrogen, or amino acid side chain, or the spacer. The lipophilic substituent may include a hydrocarbon chain having 10 to 24 carbon (C) atoms, e.g. 10 to 22 C atoms, e.g. 10 to 20 C atoms. Preferably, it has at least 11 C atoms, and preferably it has 18 C atoms or fewer. For example, the hydrocarbon chain may contain 12, 13, 14, 15, 16, 17 or 18 carbon atoms. The hydrocarbon chain may be linear or branched and may be saturated or unsaturated.

The hydrocarbon chain may incorporate a phenylene or piperazinylene moiety in its length as, for example, shown below (wherein --- represents the points of attachment within the chain). These groups should be "counted" as 4 carbon atoms in the chain length.



From the discussion above, it will be understood that the hydrocarbon chain may be substituted with a moiety which forms part of the attachment to the amino acid side chain or the spacer, for example an acyl group, a sulphonyl group, an N atom, an O atom or an S atom. Most preferably, the hydrocarbon chain is substituted with an acyl group, and accordingly the hydrocarbon chain may be part of an alkanoyl group, for example a dodecanoyl, 2-butyloctanoyl, tetradecanoyl, hexadecanoyl, heptadecanoyl, octadecanoyl or eicosanoyl group. Alternatively, Z^1 groups are derived from long-chain saturated α,ω -dicarboxylic acids of formula $\text{HOOC}-(\text{CH}_2)_{12-22}-\text{COOH}$, preferably from long-chain saturated α,ω -dicarboxylic acids having an even number of carbon atoms in the aliphatic chain.

In other words, Z^1 may be $\text{A}-\text{C}_{12-22}\text{alkylene}-(\text{CO})-$, where A is H or $-\text{COOH}$, and wherein the alkylene may be linear or branched and may be saturated or unsaturated, and may optionally incorporate a phenylene or piperazinylene moiety in its length.

For example, Z^1 may be:

- Tridecanoyl i.e. $\text{H}-(\text{CH}_2)_{12}-(\text{CO})-$;
- 13-carboxytridecanoyl, i.e. $\text{HOOC}-(\text{CH}_2)_{12}-(\text{CO})-$;
- 15-carboxypentadecanoyl, i.e. $\text{HOOC}-(\text{CH}_2)_{14}-(\text{CO})-$;
- 17-carboxyheptadecanoyl, i.e. $\text{HOOC}-(\text{CH}_2)_{16}-(\text{CO})-$;
- 19-carboxynonadecanoyl, i.e. $\text{HOOC}-(\text{CH}_2)_{18}-(\text{CO})-$; or
- 21-carboxyheneicosanoyl, i.e. $\text{HOOC}-(\text{CH}_2)_{20}-(\text{CO})-$

The carboxylic acid, if present, may be replaced by a bioisotere, phosphate or sulfonate.

Suitable bioisoterres for carboxylic acids are known in the art and include tetrazole,

5 acylsulfonamides, acylhydroxylamine, and squaric acid derivatives.

As mentioned above, the lipophilic substituent Z^1 may be conjugated to the amino acid side chain or N-terminal nitrogen by one or more spacers Z^2 .

10 When present, the spacer is attached to the lipophilic substituent and to the amino acid side chain or N-terminal nitrogen. The spacer may be attached to the lipophilic substituent and to the amino acid side chain independently by an ester, a sulphonyl ester, a thioester, an amide or a sulphonamide. Accordingly, it may include two moieties independently selected from acyl, sulphonyl, an N atom, an O atom or an S atom. The spacer may consist of a linear C_{1-10}
 15 hydrocarbon chain or more preferably a linear C_{1-5} hydrocarbon chain. Furthermore the spacer can be substituted with one or more substituents selected from C_{1-6} alkyl, C_{1-6} alkyl amine, C_{1-6} alkyl hydroxy and C_{1-6} alkyl carboxy.

The spacer may be, for example, a residue of any naturally occurring or unnatural amino acid.

20 For example, the spacer may be a residue of Gly, Pro, Ala, Val, Leu, Ile, Met, Cys, Phe, Tyr, Trp, His, Lys, Arg, Gln, Asn, α -Glu, γ -Glu, β -Asp, ϵ -Lys, Asp, Ser, Thr, Dapa, Gaba, Aib, β -Ala (i.e., 3-aminopropanoyl), 4-aminobutanoyl, 5-aminopentanoyl, 6-aminohexanoyl, 7-aminohexanoyl, 8-aminooctanoyl, 9-aminononanoyl, 10-aminodecanoyl, 8-amino-3,6-dioxaoctanoyl. In certain embodiments, the spacer is a residue of Glu, γ -Glu, ϵ -Lys, β -Ala
 25 (i.e., 3-aminopropanoyl), 4-aminobutanoyl, 8-aminooctanoyl or 8-amino-3,6-dioxaoctanoyl (Peg3), 11-amino-3,6,9-trioxaundecanoic acid (Peg4) or (piperazine-1-yl)-carboxylic acid. In the present invention, γ Glu and isoGlu are used interchangeably.

Z^2 is suitably a sequence of 1 to 6 residues of compounds selected from γ Glu, α -Glu E, K,

30 Orn, S, T, A, β Ala, G, P, V, L, I, Y, Q, N, Dapa, Gaba, or Aib, or a corresponding D form thereof, 5-aminopentanoyl, 6-aminohexanoyl, 7-aminohexanoyl, 8-aminooctanoyl, 9-aminononanoyl, and 10-aminodecanoyl. 8-amino-3,6-dioxaoctanoic acid (Peg3), 11-amino-3,6,9-trioxaundecanoic acid (Peg4) or (piperazine-1-yl)-carboxylic acid.

35 For example, Z^2 may be, or may comprise:

γ Glu,

γ Glu-Peg3-Peg3;

[(Piperazine-1-yl)-acetyl]-Peg3-Peg3;

(γ Glu)-G-(γ Glu);

(γ Glu)-K-(γ Glu);

(γ Glu)-K-G-(γ Glu); or

5 (γ Glu)-G-(Peg3)-(γ Glu)-(Peg3).

Z² is suitably bound at each side by amide linkage. Other suitable linkages may be used, with the commensurate atom replacement; for example sulfinamide, sulfonamide, or ester linkages or amino, ether, or thioether linkages are envisaged.

10

In other words, in some aspects the lipophilic group Φ is Z¹- or Z¹-Z²-; wherein

Z¹ is A-C₁₂₋₂₂alkylene-(CO)-;

where A is H or -COOH, and wherein the alkylene may be linear or branched and may be

15

saturated or unsaturated, and may optionally incorporate a phenylene or piperazinylene moiety in its length; and

Z² is a sequence of 1 to 6 of residues of compounds selected from γ -Glu, α -Glu E, K, Orn, S, T, A, β -Ala, G, P, V, L, I, Y, Q, N, Dapa, Gaba, or Aib, or a corresponding D form thereof, 5-aminopentanoyl, 6-aminohexanoyl, 7-aminoheptanoyl, 8-aminooctanoyl, 9-aminononanoyl, and 10-aminodecanoyl. 8-amino-3,6-dioxaoctanoic acid (Peg3), 11-amino-3,6,9-

20

trioxaundecanoic acid (Peg4) or (piperazine-1-yl)-carboxylic acid, e.g. a linker selected from γ Glu,

γ Glu-Peg3-Peg3-;

25

[(Piperazine-1-yl)-acetyl]-Peg3-Peg3;

(γ Glu)-G-(γ Glu);

(γ Glu)-K-(γ Glu);

(γ Glu)-K-G-(γ Glu); and

(γ Glu)-G-(Peg3)-(γ Glu)-(Peg3).

30

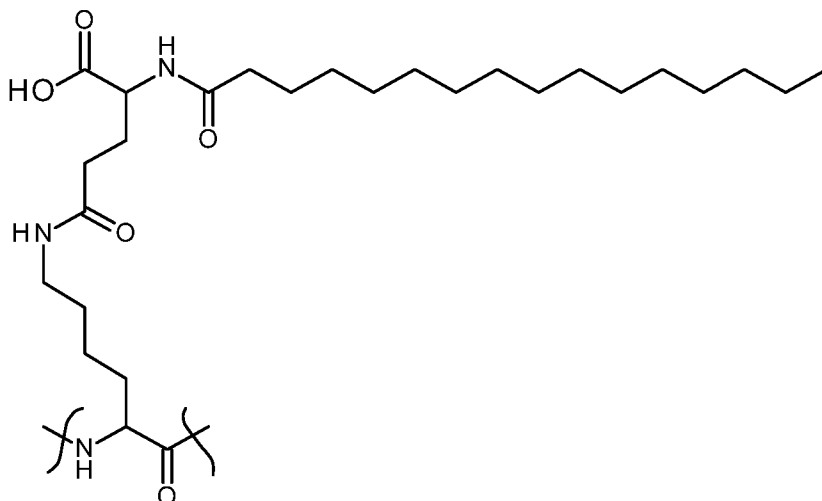
The amino acid side chain to which the lipophilic substituent is conjugated may be a side chain of a Glu, Lys, Ser, Cys, Dbu, Dpr or Orn residue. For example, it may be a side chain of a Lys, Glu or Cys residue. Where two or more side chains carry a lipophilic substituent, they may be independently selected from those residues. Thus, the amino acid side chain includes an carboxy, hydroxyl, thiol, amide or amine group, for forming an ester, a sulphonyl ester, a thioester, an amide, or a sulphonamide with the spacer or lipophilic substituent.

35

Typically, the amino acid side chain is a side chain of a Lys residue.

An example of a lipophilic substituent comprising a lipophilic moiety Z¹ and spacer Z² is shown in the formula below:

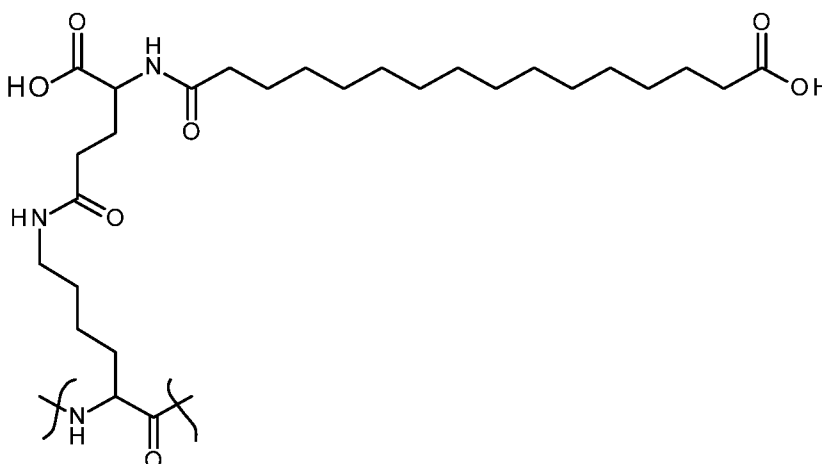
5



Here, the side chain of a Lys residue is covalently attached to a γ Glu spacer (Z²) via an amide linkage. A hexadecanoyl group (Z¹) is covalently attached to the γ Glu spacer via an amide linkage. This combination of lipophilic moiety and spacer, conjugated to a Lys residue, may be referred to by the short-hand notation K(Hexadecanoyl- γ Glu), *e.g.*, when shown in formulae of specific compounds. γ Glu can also be referred to as isoGlu, and a hexadecanoyl group as a palmitoyl group. Thus it will be apparent that the notation (Hexadecanoyl- γ Glu) is equivalent to the notations (isoGlu(Palm)) or (isoGlu(Palmitoyl)) as used for example in PCT/GB2008/004121.

15

Alternative Z¹ groups are derived from long-chain saturated α,ω -dicarboxylic acids of formula HOOC-(CH₂)₁₂₋₂₂-COOH as exemplified below



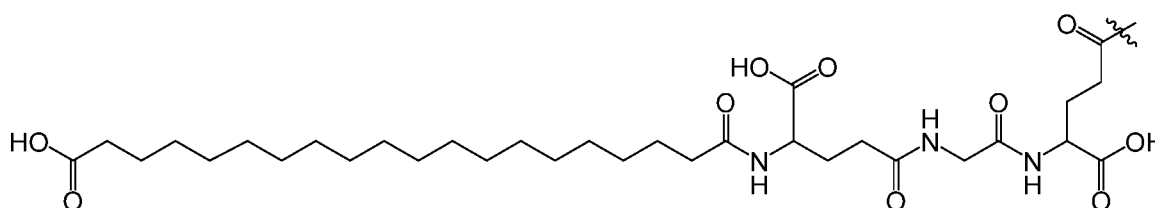
Here, the side chain of a Lys residue is covalently attached to a γ -Glu spacer (Z^2) via an amide linkage. A 15-carboxypentadecanoyl group (Z^1) is covalently attached to the γ -Glu spacer via an amide linkage. This combination of lipophilic moiety and spacer, conjugated to a Lys residue, may be referred to by the short-hand notation K(15-carboxypentadecanoyl- γ -Glu), *e.g.*, when shown in formulae of specific compounds. γ Glu can also be referred to as isoGlu.

Certain preferred Φ groups (Z^1 - and Z^1 - Z^2 -) include:

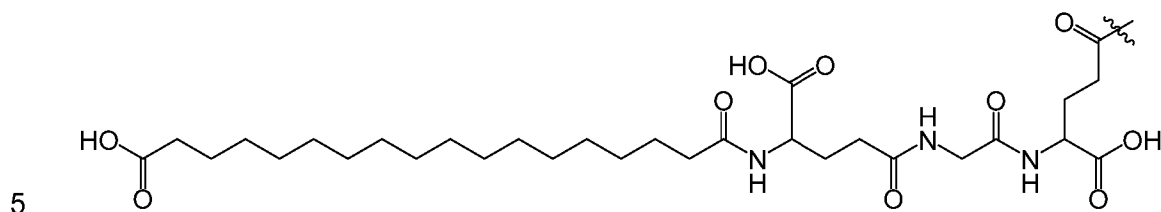
- 15-Carboxy-pentadecanoyl;
- 10 15-carboxy-pentadecanoyl- γ Glu-,
- 15-carboxy-pentadecanoyl- γ Glu-Peg3-Peg3-;
- 17-carboxy-heptadecanoyl- γ Glu-Peg3-Peg3-;
- 19-carboxy-nonadecanoyl- γ Glu-Peg3-Peg3-;
- 15-carboxy-pentadecanoyl-[(Piperazine-1-yl)-acetyl]-Peg3-Peg3-;
- 15 17-carboxy-heptadecanoyl-(γ Glu)-G-(γ Glu);
- 17-carboxy-heptadecanoyl-(γ Glu)-K-(γ Glu);
- 17-carboxy-heptadecanoyl-(γ Glu)-K-G-(γ Glu);
- 17-carboxy-heptadecanoyl-(γ Glu)-G-(Peg3)-(γ Glu)-(Peg3);
- 15-carboxy-hexadecanoyl- γ Glu-G- γ Glu;
- 20 15-carboxy-pentadecanoyl;
- 15-carboxy-pentadecanoyl- γ Glu-G- γ Glu;
- 17-carboxy-heptadecanoyl;
- 17-carboxy-heptadecanoyl- γ Glu]
- 19-carboxy-nonadecanoyl- γ Glu-G- γ Glu;
- 25 15-carboxy-pentadecanoyl- γ Glu;
- 17-carboxy-heptadecanoyl- γ Glu-Peg3-Peg3.

Illustrative structures of Φ groups (Z^1 - and Z^1 - Z^2 -) are shown below, where the wavy line indicates the linkage to the peptide (to an amino acid side chain, N-terminal nitrogen, or C-terminal carbon):

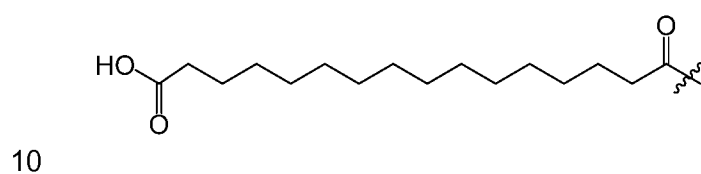
[(19-carboxy-nonadecanoyl)- γ Glu-G- γ Glu]:



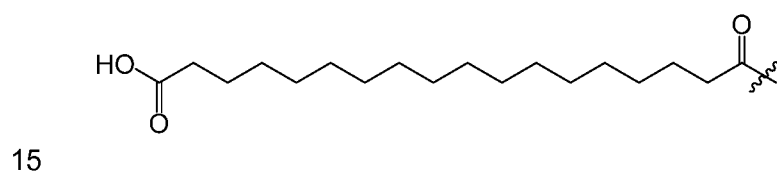
(17-carboxy-heptadecanoyl)-γGlu-G-γGlu]:



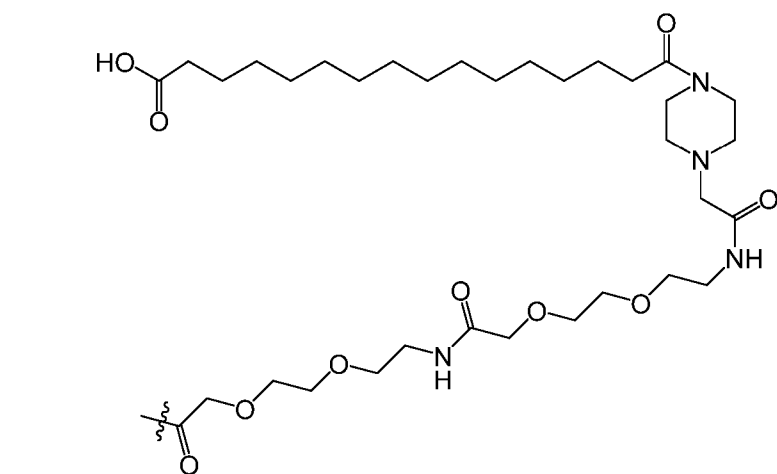
(15-carboxy-pentadecanoyl)- :



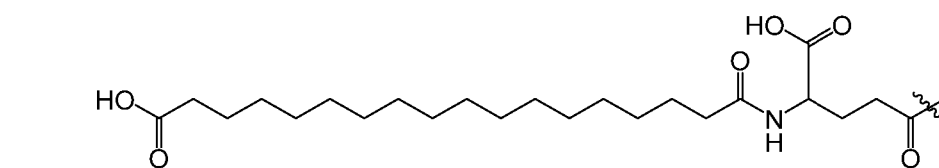
[(17-carboxy-heptadecanoyl)]- :



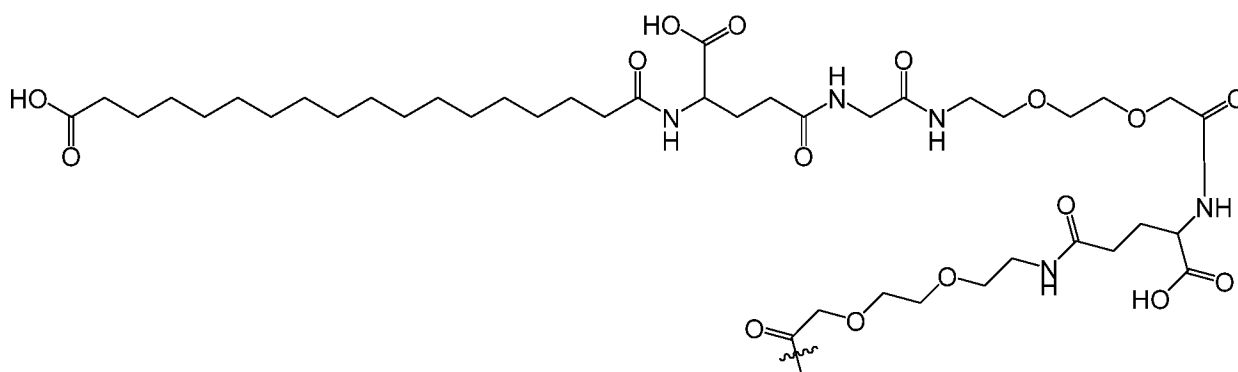
[(15-carboxy-pentadecanoyl)-[(Piperazine-1-yl)-acetyl]-Peg3-Peg3]:



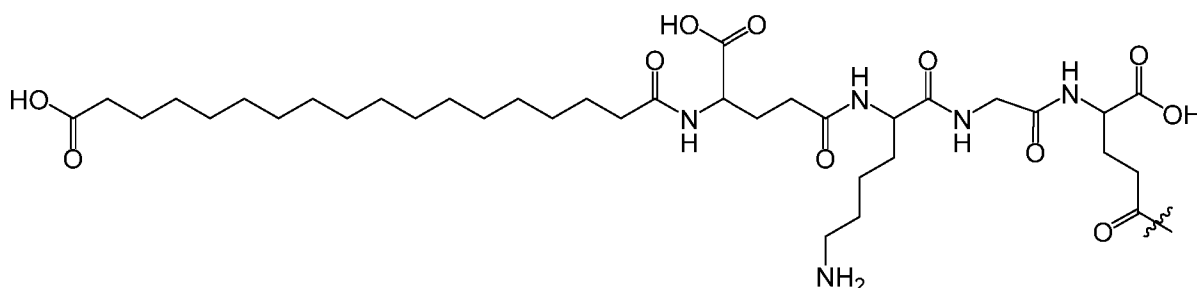
[(17-carboxy-heptadecanoyl)-γGlu]:



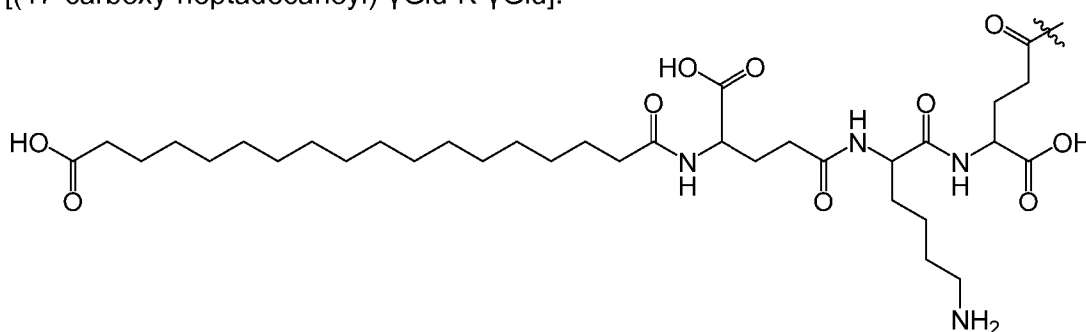
[(17-carboxy-heptadecanoyl)-γGlu-G-Peg3-γGlu-Peg3]:



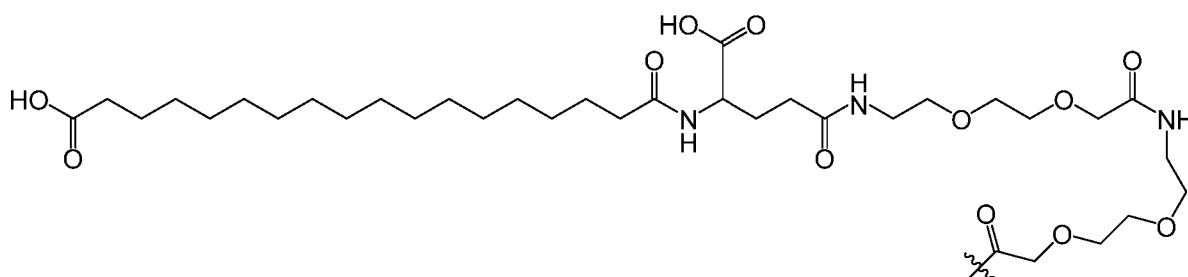
5 [(17-carboxy-heptadecanoyl)- γ Glu-KG- γ Glu]:



10 [(17-carboxy-heptadecanoyl)- γ Glu-K- γ Glu]:



15 [(17-carboxy-heptadecanoyl)- γ Glu-Peg3-Peg3]:



The skilled person will be well aware of suitable techniques for preparing the compounds employed in the context of the invention. For examples of suitable chemistry, see WO98/08871, WO00/55184, WO00/55119, Madsen et al., J. Med. Chem. 50:6126-32 (2007), and Knudsen et al., J. Med Chem. 43:1664-1669 (2000), incorporated herein by reference.

In some embodiments, the compstatin analogue has a lipophilic group Φ as described above conjugated to an amino acid at one or more of positions corresponding to positions 1, 3, 4, 5, 6, 7, 8, 9, 10, 11 or 13 of the compstatin-like sequence, i.e. positions X1-X13.

5

In certain embodiments, the compstatin analogue has a lipophilic substituent as described above conjugated to an amino acid at one or more of positions corresponding to positions X1, X11 or X13, or to an amino acid within R1 or R2, or at the N-terminus as group Y1.

10 The compstatin analogue may comprise one of the following sequences:

IC(1)IWQDWGAHRC(1)T

IC(1)IWQDWGEHRC(1)T

ESSAIC(1)IWQDWGEHRC(1)T

IC(1)I[1MeTrp]QDWGEHRC(1)T

15 IC(1)IWQDWGKHRC(1)T

IC(1)IWQDWGSHRC(1)T

IC(1)IWQKWGEHRC(1)T

IC(1)IWQKWGAHRC(1)TGAES

YC(1)IWQDWGEHRC(1)T

20 ESSAYC(1)IWQDWGEHRC(1)T

[Sar]C(1)IWQDWGEHRC(1)T

IC(1)IWQDWGAHRC(1)E

IC(1)IWQDWGEHRC(1)[Sar]

ESSAIC(1)IWQDWGEHRC(1)TGAES

25 IC(1)IWQDWGEHRC(1)TGAES

IC(1)IWQEWGEHRC(1)T

IC(1)IWQDWGDHRC(1)T

IC(1)IWQDWGRHRC(1)T

IC(1)IWQDWGAHSC(1)T

30 IC(1)IWQDWGEHSC(1)T

IC(1)IWQDWGEHRC(1)S

IC(1)IWQDWGEHRC(1)E

FC(1)IWQDWGEHRC(1)T

IC(1)IWQDWGEHRC(1)TEGE

35 IC(1)IWQDWGEHRC(1)TEA

IC(1)IWQDWGEHRC(1)TE

IC(1)IWQDWGEHRC(1)EGE

EGSAIC(1)IWQDWGEHRC(1)[Sar]E
 EGSAIC(1)IWQDWGEHRC(1)T
 EGEIC(1)IWQDWGEHRC(1)T
 ESEIC(1)IWQDWGEHRC(1)T
 5 SEIC(1)IWQDWGEHRC(1)TEA
 EIC(1)IWQDWGEHRC(1)TE
 EIC(1)IWQDWGEHRC(1)TEGE
 EGEIC(1)IWQDWGEHRC(1)EGE
 ESEIC(1)IWQDWGEHRC(1)EGE
 10 KEKIC(1)IWQDWGEHRC(1)TEKE
 EKGIC(1)IWQDWGEHRC(1)TEKP
 IC(1)IWQDWGEHRC(1)TEGK
 GSAIC(1)IWQDWGEHRC(1)[Sar]E
 SAIC(1)IWQDWGEHRC(1)[Sar]E
 15 SAIC(1)IWQDWGEHRC(1)TEG
 FC(1)IWQDWGEHRC(1)TGAE
 EGSAIC(1)IWQDWGEHRC(1)[Sar]EGE
 EGSAFC(1)IWQDWGEHRC(1)[Sar]E
 ESSAIC(1)IWQDWGAHRC(1)T
 20 IC(1)IWQDWGAHRC(1)TGAES
 {d}YIC(1)I[1-Me-Trp]QDW[Sar]AHRC(1)-[N-Me-Ile]
 EGSAIC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E
 EGSAIC(1)I-2NaI-QDWGEHRC(1)[Sar]E
 IC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES
 25 IC(1)I[2NaI]QDWGEHRC(1)TGAES
 EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E
 EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E
 EGSAIC(1)IWQDWGEHRC(1)TE
 EGSAFC(1)I[1NaI]QDWGEHRC(1)TE
 30 EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)TE
 EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)EGE
 EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)TE
 EGSAFC(1)I[2NaI]QDWGEHRC(1)TE
 FC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES
 35 YC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES
 FC(1)I[1NaI]QDWGEHRC(1)TGAES
 FC(1)I[2NaI]QDWGEHRC(1)TGAES

YC(1)I[2Na]QDWGEHRC(1)TGAES
 YC(1)IWQDWGEHRC(1)TGAES
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES
 YC(1)I[1-Me-Trp]QDWGEHRC(1)TEAGS
 5 YC(1)I[1-Me-Trp]QDWGEHRC(1)TESGA
 EGSAYC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]E
 SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA
 FC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TGAES
 {d}YFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TGAES
 10 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]GAES
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA
 SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)[Sar]EA
 SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TEA
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E
 15 SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)[Sar]E
 EFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA
 SE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA
 SE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA
 SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EA
 20 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)SEA
 EFC(1)I[1-Me-Trp]QDWGEHRC(1)ES
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA
 GEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA
 GE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA
 25 SE[Sar]C(1)I[1-Me-Trp]QEW[Sar]EHRC(1)TEA
 SE[Sar]C(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EA
 {d}Y[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA

For example, the compstatin analogue may be:

30 Ac-IC(1)IWQDWGAHRC(1)T-NH₂ (Compound 1)
 Ac-IC(1)IWQDWGEHRC(1)T-NH₂ (Compound 2)
 Ac-ESSAIC(1)IWQDWGEHRC(1)T-NH₂ (Compound 3)
 Ac-IC(1)I[1-Me-Trp]QDWGEHRC(1)T-NH₂ (Compound 4)
 Ac-IC(1)IWQDWGKHRC(1)T-NH₂ (Compound 5)
 35 Ac-IC(1)IWQDWGSHRC(1)T-NH₂ (Compound 6)
 Ac-IC(1)IWQKWGEHRC(1)T-NH₂ (Compound 7)
 Ac-IC(1)IWQKWGAHRC(1)TGAES-NH₂ (Compound 8)

- Ac-YC(1)IWQDWGEHRC(1)T-NH₂ (Compound 9)
 Ac-ESSAYC(1)IWQDWGEHRC(1)T-NH₂ (Compound 10)
 Ac-[Sar]C(1)IWQDWGEHRC(1)T-NH₂ (Compound 11)
 Ac-IC(1)IWQDWGAHRC(1)E-NH₂ (Compound 12)
 5 Ac-IC(1)IWQDWGEHRC(1)[Sar]-NH₂ (Compound 13)
 Ac-ESSAIC(1)IWQDWGEHRC(1)TGAES-NH₂ (Compound 14)
 Ac-IC(1)IWQDWGEHRC(1)TGAES-NH₂ (Compound 15)
 Ac-IC(1)IWQEWGEHRC(1)T-NH₂ (Compound 16)
 Ac-IC(1)IWQDWGDHRC(1)T-NH₂ (Compound 17)
 10 Ac-IC(1)IWQDWGRHRC(1)T-NH₂ (Compound 18)
 Ac-IC(1)IWQDWGAHSC(1)T-NH₂ (Compound 19)
 Ac-IC(1)IWQDWGEHSC(1)T-NH₂ (Compound 20)
 Ac-IC(1)IWQDWGEHRC(1)S-NH₂ (Compound 21)
 Ac-IC(1)IWQDWGEHRC(1)E-NH₂ (Compound 22)
 15 Ac-FC(1)IWQDWGEHRC(1)T-NH₂ (Compound 23)
 Ac-IC(1)IWQDWGEHRC(1)TEGE-NH₂ (Compound 24)
 Ac-IC(1)IWQDWGEHRC(1)TEA-NH₂ (Compound 25)
 Ac-IC(1)IWQDWGEHRC(1)TE-NH₂ (Compound 26)
 Ac-IC(1)IWQDWGEHRC(1)EGE-NH₂ (Compound 27)
 20 Ac-EGSAIC(1)IWQDWGEHRC(1)[Sar]E-NH₂ (Compound 28)
 Ac-EGSAIC(1)IWQDWGEHRC(1)T-NH₂ (Compound 29)
 Ac-EGEIC(1)IWQDWGEHRC(1)T-NH₂ (Compound 30)
 Ac-ESEIC(1)IWQDWGEHRC(1)T-NH₂ (Compound 31)
 Ac-SEIC(1)IWQDWGEHRC(1)TEA-NH₂ (Compound 32)
 25 Ac-EIC(1)IWQDWGEHRC(1)TE-NH₂ (Compound 33)
 Ac-EIC(1)IWQDWGEHRC(1)TEGE-NH₂ (Compound 34)
 Ac-EGEIC(1)IWQDWGEHRC(1)EGE-NH₂ (Compound 35)
 Ac-ESEIC(1)IWQDWGEHRC(1)EGE-NH₂ (Compound 36)
 Ac-KEKIC(1)IWQDWGEHRC(1)TEKE-NH₂ (Compound 37)
 30 Ac-EKGIC(1)IWQDWGEHRC(1)TEKP-NH₂ (Compound 38)
 Ac-IC(1)IWQDWGEHRC(1)TEGK-NH₂ (Compound 39)
 Ac-GSAIC(1)IWQDWGEHRC(1)[Sar]E-NH₂ (Compound 40)
 Ac-SAIC(1)IWQDWGEHRC(1)[Sar]E-NH₂ (Compound 41)
 Ac-SAIC(1)IWQDWGEHRC(1)TEG-NH₂ (Compound 42)
 35 Ac-FC(1)IWQDWGEHRC(1)TGAE-NH₂ (Compound 43)
 Ac-EGSAIC(1)IWQDWGEHRC(1)[Sar]EGE-NH₂ (Compound 44)
 Ac-EGSAFC(1)IWQDWGEHRC(1)[Sar]E-NH₂ (Compound 45)

- Ac-ESSAIC(1)IWQDWGAHRC(1)T-NH₂ (Compound 46)
- Ac-IC(1)IWQDWGAHRC(1)TGAES-NH₂ (Compound 47)
- H-{d}YIC(1)I[1-Me-Trp]QDW[Sar]AHRC(1)[N-Me-Ile]-NH₂ (Compound 48)
- Ac-EGSAIC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 49)
- 5 Ac-EGSAIC(1)I[2NaI]QDWGEHRC(1)[Sar]E-NH₂ (Compound 50)
- Ac-IC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-NH₂ (Compound 51)
- Ac-IC(1)I[2NaI]QDWGEHRC(1)TGAES-NH₂ (Compound 52)
- Ac-EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 53)
- Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 54)
- 10 Ac-EGSAIC(1)IWQDWGEHRC(1)TE-NH₂ (Compound 55)
- Ac-EGSAFC(1)I[1NaI]QDWGEHRC(1)TE-NH₂ (Compound 56)
- Ac-EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)TE-NH₂ (Compound 57)
- Ac-EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)EGE-NH₂ (Compound 58)
- Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)TE-NH₂ (Compound 59)
- 15 Ac-EGSAFC(1)I[2NaI]QDWGEHRC(1)TE-NH₂ (Compound 60)
- Ac-FC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-NH₂ (Compound 61)
- Ac-YC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-NH₂ (Compound 62)
- Ac-FC(1)I[1NaI]QDWGEHRC(1)TGAES-NH₂ (Compound 63)
- Ac-FC(1)I[2NaI]QDWGEHRC(1)TGAES-NH₂ (Compound 64)
- 20 Ac-YC(1)I[2NaI]QDWGEHRC(1)TGAES-NH₂ (Compound 65)
- Ac-YC(1)IWQDWGEHRC(1)TGAES-NH₂ (Compound 66)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-NH₂ (Compound 67)
- Ac-YC(1)I[1-Me-Trp]QDWGEHRC(1)TEAGS-NH₂ (Compound 68)
- Ac-YC(1)I[1-Me-Trp]QDWGEHRC(1)TESGA-NH₂ (Compound 69)
- 25 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 70)
- Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH₂ (Compound 71)
- Ac-FC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TGAES-NH₂ (Compound 72)
- H-{d}YFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TGAES-NH₂ (Compound 73)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]GAES-NH₂ (Compound 74)
- 30 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH₂ (Compound 75)
- Ac-SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)[Sar]EA-NH₂ (Compound 76)
- Ac-SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TEA-NH₂ (Compound 77)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 78)
- Ac-SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)[Sar]E-NH₂ (Compound 79)
- 35 Ac-EFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH₂ (Compound 80)
- Ac-SE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH₂ (Compound 81)
- Ac-SE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA-NH₂ (Compound 82)

- Ac-SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EA-NH₂ (Compound 83)
 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)SEA-NH₂ (Compound 84)
 Ac-EFC(1)I[1-Me-Trp]QDWGEHRC(1)ES-NH₂ (Compound 85)
 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH₂ (Compound 86)
 5 Ac-GEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH₂ (Compound 87)
 Ac-GE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA-NH₂ (Compound 88)
 Ac-SE[Sar]C(1)I[1-Me-Trp]QEW[Sar]EHRC(1)TEA-NH₂ (Compound 89)
 Ac-SE[Sar]C(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EA-NH₂ (Compound 90)
 H-{d}Y[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA-NH₂ (Compound 91)

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Alternatively, the compstatin analogue may comprise one of the following sequences:

- [K*]GSAIC(1)IWQDWGEHRC(1)TEGE (Compound 100)
 15 ASGEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[K*] (Compound 113)
 EFC(1)I[1-Me-Trp]QDWGEHRC(1)EGE-[K*] (Compound 134)
 EGSAIC(1)IWQDWGEHRC(1)TEG[K*] (Compound 101)
 EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E (Compound 103)
 EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EG[K*] (Compound 104)
 20 EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[K*] (Compound 109)
 EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGK[K*] (Compound 110)
 EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EK[ΓGlu][K*] (Compound 111)
 FC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES[K*] (Compound 102)
 IC(1)IWQDWGEHRC(1)TEG[K*] (Compound 92)
 25 IC(1)IWQDWGEHRC(1)TEGE[K*] (Compound 94)
 SAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E[K*] (Compound 105)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGA[K*] (Compound 119)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[Peg3][Peg3][K*] (Compound 123)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGEGGG-[K*] (Compound 129)
 30 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-Peg3-[K*] (Compound 138)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-Peg3-ES-[K*] (Compound 140)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-Peg3-Peg3-[K*] (Compound 127)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGESES-[K*] (Compound 139)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EK-γGlu-GGG-[K*] (Compound 132)
 35 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE-[8-aminooctanoyl]-[K*] (Compound 136)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE-[8-aminooctanoyl]-E-[K*] (Compound 137)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGEGGG-[K*] (Compound 130)

- SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE-Peg3-ES-[K*] (Compound 142)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE-Peg3-Peg3-[K*] (Compound 126)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEK-γGlu-GGG-[K*] (Compound 133)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-[K*] (Compound 135)
 5 SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGA[K*] (Compound 120)
 SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[Peg3][Peg3][K*] (Compound 124)
 SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGA[K*] (Compound 112)
 SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[Peg3][Peg3][K*] (Compound 117)
 SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[K*] (Compound 114)
 10 SEYC(1)I[1-Me-Trp]QEW[Sar]EHRC(1)[Sar]EK[γGlu]A[K*] (Compound 121)
 SEYC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGA[K*] (Compound 122)
 SEYC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[Peg3][Peg3][K*] (Compound 125)
 EGSEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E (Compound 107)
 ESSAIC(1)IWQDWGEHRC(1)TEGE (Compound 99)

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For example, the compstatin analogue may comprise one of the following sequences:

- Ac-[K*]GSAIC(1)IWQDWGEHRC(1)TEGE-NH₂ (Compound 100)
 Ac-ASGEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[K*]-NH₂ (Compound 113)
 20 Ac-EFC(1)I[1-Me-Trp]QDWGEHRC(1)EGE-[K*]-NH₂ (Compound 134)
 Ac-EGSAIC(1)IWQDWGEHRC(1)TEG[K*]-NH₂ (Compound 101)
 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEH[K*]C(1)[Sar]E-NH₂ (Compound 103)
 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EG[K*]-NH₂ (Compound 104)
 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[K*]-NH₂ (Compound 109)
 25 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGK[K*]-NH₂ (Compound 110)
 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EK[γGlu][K*]-NH₂ (Compound 111)
 Ac-FC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES[K*]-NH₂ (Compound 102)
 Ac-IC(1)IWQDWGEHRC(1)TEG[K*]-NH₂ (Compound 92, 93, 95, 96, 98)
 Ac-IC(1)IWQDWGEHRC(1)TEGE[K*]-NH₂ (Compound 94, 97)
 30 Ac-SAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E[K*]-NH₂ (Compound 105, 106)
 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGA[K*]-NH₂ (Compound 119)
 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[Peg3][Peg3][K*]-NH₂ (Compound 123)
 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGEGGG-[K*]-NH₂ (Compound 129)
 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-Peg3-[K*]-NH₂ (Compound 138)
 35 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-Peg3-ES-[K*]-NH₂ (Compound 140)
 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-Peg3-Peg3-[K*]-NH₂ (Compound 127,
 128)

- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGESES-[K*]-NH₂ (Compound 139, 141)
 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EK[γGlu]GGG-[K*]-NH₂ (Compound 132)
 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEGE-[8-aminooctanoyl]-[K*]-NH₂ (Compound 136)
 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEGE-[8-aminooctanoyl]-E-[K*]-NH₂ (Compound
 5 137)
 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEGEGGG-[K*]-NH₂ (Compound 130, 131)
 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEGE-Peg3-ES-[K*]-NH₂ (Compound 142)
 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEGE-Peg3-Peg3-[K*]-NH₂ (Compound 126)
 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEK-γGlu-GGG-[K*]-NH₂ (Compound 133)
 10 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TGAES-[K*]-NH₂ (Compound 135)
 Ac-SEFC(1)|[1-Me-Trp]QEWGEHRC(1)[Sar]EGA[K*]-NH₂ (Compound 120)
 Ac-SEFC(1)|[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[Peg3][Peg3][K*]-NH₂ (Compound 124)
 Ac-SEYC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGA[K*]-NH₂ (Compound 112, 118)
 Ac-SEYC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[Peg3][Peg3][K*]-NH₂ (Compound 117)
 15 Ac-SEYC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[K*]-NH₂ (Compound 114, 115, 116)
 Ac-SEYC(1)|[1-Me-Trp]QEW[Sar]EHRC(1)[Sar]EK[ΓGlu]A[K*]-NH₂ (Compound 121)
 Ac-SEYC(1)|[1-Me-Trp]QEWGEHRC(1)[Sar]EGA[K*]-NH₂ (Compound 122)
 Ac-SEYC(1)|[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[Peg3][Peg3][K*]-NH₂ (Compound 125)
 Φ-EGSEYC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 107, 108)
 20 Φ-ESSAIC(1)|IWQDWGEHRC(1)TEGE-NH₂ (Compound 99)

For example, the compstatin analogue may be:

- Ac-IC(1)|IWQDWGEHRC(1)TEG-K([15-carboxy-pentadecanoyl]-γGlu)-NH₂ (Compound 92)
 25 Ac-IC(1)|IWQDWGEHRC(1)TEG-K([15-carboxy-pentadecanoyl]-γGlu-Peg3-Peg3)-NH₂
 (Compound 93)
 Ac-IC(1)|IWQDWGEHRC(1)TEGE-K([15-carboxy-pentadecanoyl]-γGlu-Peg3-Peg3)-NH₂
 (Compound 94)
 Ac-IC(1)|IWQDWGEHRC(1)TEG-K((15-carboxy-pentadecanoyl)-[(Piperazine-1-yl)-acetyl]-
 30 Peg3-Peg3)-NH₂ (Compound 95)
 Ac-IC(1)|IWQDWGEHRC(1)TEG-K([17-carboxy-heptadecanoyl]-γGlu-Peg3-Peg3)-NH₂
 (Compound 96)
 Ac-IC(1)|IWQDWGEHRC(1)TEGE-K([17-carboxy-heptadecanoyl]-γGlu-Peg3-Peg3)-NH₂
 (Compound 97)
 35 Ac-IC(1)|IWQDWGEHRC(1)TEG-K([19-carboxy-nonadecanoyl]-γGlu-Peg3-Peg3)-NH₂
 (Compound 98)
 [15-Carboxy-pentadecanoyl]-ESSAIC(1)|IWQDWGEHRC(1)TEGE-NH₂ (Compound 99)

- Ac-[K([15-carboxy-pentadecanoyl]-γGlu-Peg3-Peg3)]-GSAIC(1)IWQDWGEHRC(1)TEGE-NH₂ (Compound 100)
- Ac-EGSAIC(1)IWQDWGEHRC(1)TEG-K([15-carboxy-pentadecanoyl]-γGlu)-NH₂ (Compound 101)
- 5 Ac-FC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-K([15-carboxy-pentadecanoyl]-γGlu-Peg3-Peg3)-NH₂ (Compound 102)
- Ac-EGSAYC(1)I[1-Me-Trp]QDWGEH-K([15-carboxy-pentadecanoyl]-γGlu-Peg3-Peg3)-C(1)[Sar]E-NH₂ (Compound 103)
- Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EG-K([15-carboxy-pentadecanoyl]-γGlu-Peg3-Peg3)-NH₂ (Compound 104)
- 10 Ac-SAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-K([17-carboxy-heptadecanoyl]-γGlu-KG-γGlu)-NH₂ (Compound 105)
- Ac-SAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 106)
- 15 [15-Carboxy-pentadecanoyl]-EGSEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 107)
- [17-Carboxy-heptadecanoyl]-EGSEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 108)
- Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 109)
- 20 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGK-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 110)
- Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EK[γGlu]-K([17-carboxy-heptadecanoyl]-γGlu-Peg3-Peg3)-NH₂ (Compound 111)
- 25 Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGA-K[17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 112)
- Ac-ASGEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 113)
- Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 114)
- 30 Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGK-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 115)
- Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-K([17-carboxy-heptadecanoyl]-γGlu-K-γGlu)-NH₂ (Compound 116)
- 35 Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[Peg3][Peg3]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 117)

- Ac-SEYC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-(Peg3)-γGlu-(Peg3))-NH2 (Compound 118)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-(Peg3)-γGlu-(Peg3))-NH2 (Compound 119)
- 5 Ac-SEFC(1)|[1-Me-Trp]QEWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-(Peg3)-γGlu-(Peg3))-NH2 (Compound 120)
- Ac-SEYC(1)|[1-Me-Trp]QEW[Sar]EHRC(1)[Sar]EK[γGlu]A-K([17-carboxy-heptadecanoyl]-γGlu-G-(Peg3)-γGlu-(Peg3))-NH2 (Compound 121)
- Ac-SEYC(1)|[1-Me-Trp]QEWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-(8-amino-3,6-dioxaoctanoyl)-γGlu-(8-amino-3,6-dioxaoctanoyl))-NH2 (Compound 122)
- 10 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2 (Compound 123)
- Ac-SEFC(1)|[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2 (Compound 124)
- 15 Ac-SEYC(1)|[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2 (Compound 125)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2 (Compound 126)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]-EGE-[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([15-carboxy-pentadecanoyl]-γGlu-G-γGlu)-NH2 (Compound 127)
- 20 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]-EGE-[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([19-carboxy-nonadecanoyl]-γGlu-G-γGlu)-NH2 (Compound 128)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGEGGG-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2 (Compound 129)
- 25 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEGEGGG-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2 (Compound 130)
- Ac-SEFC(1)|[1-Me-Trp]-QDWGEHRC(1)TEGEGGG-K([15-carboxy-pentadecanoyl]-γGlu-G-γGlu)-NH2 (Compound 131)
- Ac-SEFC(1)|[1-Me-Trp]-QDWGEHRC(1)[Sar]EK[γGlu]GGG-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2 (Compound 132)
- 30 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEK[γGlu]GGG-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2 (Compound 133)
- Ac-EFC(1)|[1-Me-Trp]QDWGEHRC(1)EGE-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2 (Compound 134)
- 35 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TGAES-K([15-carboxy-hexadecanoyl]-γGlu-G-γGlu))-NH2 (Compound 135)

Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE[8-aminooctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 136)

Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE[8-aminooctanoyl]-E-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 137)

- 5 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 138)

Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGESES-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 139)

- 10 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl]ES-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 140)

Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGESES-K([17-carboxy-heptadecanoyl]-γGlu)-NH₂ (Compound 141)

Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE-[8-Amino-3,6-dioxaoctanoyl]ES-K([17-carboxy-heptadecanoyl]-γGlu)-NH₂ (Compound 142)

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Compstatin analogues made in the prior art have been shown to possess improved activity as compared with the parent peptide, i.e., up to about 99-fold (Mallik, B. et al, 2005, supra; WO 2004/026328), and up to about 264-fold (Katragadda et al., 2006, supra; WO2007/062249).

- 20 In accordance with the present invention, information about the biological and physico-chemical characteristics of Ac-compstatin binding to C3 have been employed to design compstatin analogues with significantly improved activity compared to the parent compstatin analogues.

- 25 Preferably, the compstatin analogs have greater activity than Ac-compstatin, e.g. at least 10-fold greater activity, at least 20-fold greater activity, at least 30-fold greater activity than Ac-compstatin. In other embodiments, the analogs have at least 40-, 50-, 60-, 70-, 80-, 90-, 100-, 110-, 120-, 130-, 140-, 150-fold or greater activity than Ac-compstatin, as compared utilizing the assays described in the examples.

30

A compound of the invention typically has greater activity than an otherwise identical compound having valine instead of isoleucine at the position corresponding to Val3 of compstatin.

- 35 The compstatin analogues are capable of binding to C3 and/or C3b, and of inhibiting activation of the complement cascade, particularly downstream of C3, e.g. by inhibiting cleavage of C3 by C3 convertases.

The compstatin analogues are also typically capable of inhibiting complement-driven haemolysis. Complement-driven haemolysis is typically assessed (in a “haemolysis assay”) by contacting serum from a first mammalian species (e.g. human serum) with erythrocytes (red blood cells; RBC) from a second mammalian species (e.g. sheep or any other suitable species), typically in the presence of mammalian immunoglobulin capable of binding to the erythrocytes. Complement in the serum is activated by the cell-bound immunoglobulin, leading to lysis of the erythrocytes, i.e. haemolysis. The immunoglobulin may be from the first species, or may be from a third mammalian species as long as it is capable of activating complement from the first species.

In such an assay, a test compound will typically be pre-incubated with the serum before the serum is contacted with the erythrocytes. The erythrocytes may also be pre-incubated with the immunoglobulin before contacting with the serum.

In the examples below, human serum is pre-incubated with a test compound, and sheep erythrocytes are pre-incubated with rabbit anti-serum against sheep erythrocytes, before the serum and erythrocytes are combined.

Thus, the activity of the compstatin analogues may be determined with reference to one or more biological activities selected from (1) binding to C3 protein, (2) binding to C3b protein, (3) inhibiting the cleavage of native C3 by C3 convertases, and (4) inhibiting the activation of the complement system.

Thus a compstatin analogue of the invention may bind C3 or C3b with a higher affinity than that of compstatin. For example, they may have a K_d at least 10-fold lower, at least 20-fold lower, or at least 30-fold lower than Ac-compstatin, e.g. at least 40-, 50-, 60-, 70-, 80-, 90-, 100-, 110-, 120-, 130-, 140-, or 150-fold lower than Ac-compstatin. The K_d may be determined by surface plasmon resonance (SPR), e.g. using an assay as described in Example 4.

A compstatin analogue of the invention typically binds C3 or C3b with a greater affinity (i.e. a lower K_d) than that of an otherwise identical compound having valine instead of isoleucine at the position corresponding to Val3 of compstatin.

A compstatin analogue of the invention may have a greater ability to inhibit haemolysis than Ac-compstatin. For example, it may inhibit haemolysis with an IC_{50} at least 10-fold, at least 20-fold, or at least 30-fold lower than Ac-compstatin, e.g. at least 40-, 50-, 60-, 70-, 80-, 90-,

100-, 110-, 120-, 130-, 140-, 150-, 200-, 250-, 300- 350-, 400-, 450-, 500-fold lower than Ac-compstatin.

A compstatin analogue of the invention typically has a greater ability to inhibit haemolysis (i.e. a lower IC_{50}) than an otherwise identical compound having valine instead of isoleucine at the position corresponding to Val3 of compstatin.

Preferably, the *in vitro* effect of the compounds of the present invention are assessed by measuring their inhibitory effect on the classical complement pathway in a haemolysis assay, e.g. using the assay described in Example 2.

Compstatin analogues having acylation may have a lower absolute activity than an otherwise identical compound lacking acylation, but have additional benefits including prolonged *in vivo* half life which may offset any apparent reduction of absolute activity.

Synthesis of Compstatin Analogues

It is preferred to synthesize compstatin analogues of the present invention by means of solid-phase or liquid-phase peptide synthesis methodology. In this context, reference may be made to WO 98/11125 and, among many others, Fields, G.B. et al., 2002, "Principles and practice of solid-phase peptide synthesis". In: Synthetic Peptides (2nd Edition), and the Examples herein.

In accordance with the present invention, a compstatin analogue of the invention may be synthesized or produced in a number of ways, including for example, a method which comprises:

(a) synthesizing the compstatin analogues by means of solid-phase or liquid-phase peptide synthesis methodology and recovering the synthesized compstatin analogues thus obtained; or

(b) expressing a precursor peptide sequence from a nucleic acid construct that encodes the precursor peptide, recovering the expression product, and modifying the precursor peptide to yield a compound of the invention.

The precursor peptide may be modified by introduction of one or more non-proteinogenic amino acids, e.g. Aib, Orn, Dap, 1MeTrp, 1Nal, 2Nal, Sar, γ Glu or Dab, or by the introduction of an appropriate terminal groups Y1 and/or Y2.

Expression is typically performed from a nucleic acid encoding the precursor peptide, which may be performed in a cell or a cell-free expression system comprising such a nucleic acid.

It is preferred to synthesize the analogues of the invention by means of solid-phase or liquid-phase peptide synthesis. In this context, reference is made to WO 98/11125 and, among many others, Fields, GB et al., 2002, "Principles and practice of solid-phase peptide synthesis". In: Synthetic Peptides (2nd Edition), and the Examples herein.

For recombinant expression, the nucleic acid fragments encoding the precursor peptide will normally be inserted in suitable vectors to form cloning or expression vectors. The vectors can, depending on purpose and type of application, be in the form of plasmids, phages, cosmids, mini-chromosomes, or virus, but also naked DNA which is only expressed transiently in certain cells is an important vector. Preferred cloning and expression vectors (plasmid vectors) are capable of autonomous replication, thereby enabling high copy-numbers for the purposes of high-level expression or high-level replication for subsequent cloning.

In general outline, an expression vector comprises the following features in the 5'→3' direction and in operable linkage: a promoter for driving expression of the nucleic acid fragment, optionally a nucleic acid sequence encoding a leader peptide enabling secretion (to the extracellular phase or, where applicable, into the periplasma), the nucleic acid fragment encoding the precursor peptide, and optionally a nucleic acid sequence encoding a terminator. They may comprise additional features such as selectable markers and origins of replication. When operating with expression vectors in producer strains or cell lines it may be preferred that the vector is capable of integrating into the host cell genome. The skilled person is very familiar with suitable vectors and is able to design one according to their specific requirements.

The vectors of the invention are used to transform host cells to produce the precursor peptide. Such transformed cells can be cultured cells or cell lines used for propagation of the nucleic acid fragments and vectors, and/or used for recombinant production of the precursor peptides.

Preferred transformed cells are micro-organisms such as bacteria [such as the species *Escherichia* (e.g. *E. coli*), *Bacillus* (e.g. *Bacillus subtilis*), *Salmonella*, or *Mycobacterium* (preferably non-pathogenic, e.g. *M. bovis* BCG), yeasts (e.g., *Saccharomyces cerevisiae* and *Pichia pastoris*), and protozoans. Alternatively, the transformed cells may be derived from a

multicellular organism, i.e. it may be fungal cell, an insect cell, an algal cell, a plant cell, or an animal cell such as a mammalian cell. For the purposes of cloning and/or optimised expression it is preferred that the transformed cell is capable of replicating the nucleic acid fragment of the invention. Cells expressing the nucleic fragment can be used for small-scale or large-scale preparation of the peptides of the invention.

When producing the precursor peptide by means of transformed cells, it is convenient, although far from essential, that the expression product is secreted into the culture medium.

10 Medical Conditions

In a broad aspect, the present invention provides compstatin analogues of the present invention for use as a medicament or for use in therapy.

The compstatin analogues described herein have biological activities of binding to C3 protein and/or inhibiting complement activation. Generally, the compstatin analogues of the present invention may be used for the treatment or prevention conditions associated with excessive or unwanted activation of the complement system. Complement can be activated through three different pathways: the classical, lectin and alternative pathways. The major activation event that is shared by all three pathways is the proteolytic cleavage of the central protein of the complement system, C3, into its activation products C3a and C3b by C3 convertases. Generation of these fragments leads to the opsonization of pathogenic cells by C3b and iC3b, a process that renders them susceptible to phagocytosis or clearance, and to the activation of immune cells through an interaction with complement receptors (Markiewski & Lambris, 2007, Am. J. Pathol., 171: 715-727). Deposition of C3b on target cells also induces the formation of new convertase complexes and thereby initiates a self-amplification loop. An ensemble of plasma and cell surface-bound proteins carefully regulates complement activation to prevent host cells from self-attack by the complement cascade. The 13 amino acid cyclic tridecapeptide used as a reference point for the design of the compstatin analogues of the present invention inhibits complement activation by binding to C3 and/or C3b, preventing the cleavage of native C3 by the C3 convertases. Without wishing to be bound by any particular theory, the present inventors believe that the compstatin analogues of the present invention also function in this way and may share one or more biological activities selected from (1) binding to C3 protein, (2) binding to C3b protein, (3) inhibiting the cleavage of native C3 by C3 convertases, and/or (4) inhibiting the activation of the complement system. The biological activity of the compstatin analogues of the present invention may be determined *in vitro* by

measuring their inhibitory effect of the classical complement pathway in a haemolysis assay, for example using a protocol set out in the examples below.

Excessive activation or inappropriate regulation of complement can lead to a number of pathologic conditions, ranging from autoimmune diseases to inflammatory diseases (Holers, 2003, Clin. Immunol., 107: 140-51; Markiewski & Lambris, 2007, supra; Ricklin & Lambris, 2007, Nat. Biotechnol., 25: 1265-75; Sahu et al., 2000, J. Immunol., 165: 2491-9). These conditions include: (1) inhibiting complement activation to facilitate treatment of diseases or conditions including age-related macular degeneration, periodontitis, diabetic retinopathy, glaucoma, uveitis, rheumatoid arthritis, spinal cord injury, stroke, multiple sclerosis, Parkinson's disease, Alzheimer's disease, cancer, and respiratory disorders such as asthma, chronic obstructive pulmonary disease (COPD), allergic inflammation, emphysema, bronchitis, bronchiectasis, cystic fibrosis, tuberculosis, pneumonia, respiratory distress syndrome (RDS - neonatal and adult), rhinitis and sinusitis; bacterial infections such as sepsis, ischemia-reperfusion injury in various tissues, myocardial infarction, anaphylaxis, paroxysmal nocturnal hemoglobinuria, autoimmune hemolytic anemia, psoriasis, hidradenitis suppurativa, myasthenia gravis, systemic lupus erythematosus, CHAPLE syndrome, C3 glomeropathy, IgA nephropathy, atypical hemolytic uremic syndrome, Crohn's disease, antiphospholipid syndrome, or (2) inhibiting complement activation that occurs during cell or organ transplantation, or in the use of artificial organs or implants (e.g., by coating or otherwise treating the cells, organs, artificial organs or implants with a peptide of the invention); or (3) inhibiting complement activation that occurs during extracorporeal shunting of physiological fluids (blood, urine) (e.g., by coating the tubing through which the fluids are shunted with a compstatin analogue of the present invention).

Pharmaceutical Compositions and Administration

In a further aspect, the present invention relates to a composition comprising a compstatin analogue according to the invention, or a pharmaceutically acceptable salt or solvate thereof, together with a carrier. In one embodiment of the invention, the composition is a pharmaceutical composition and the carrier is a pharmaceutically acceptable carrier. The present invention also relates to a pharmaceutical composition comprising a compstatin analogue according to the invention, or a salt and/or solvate thereof, together with a carrier, excipient or vehicle. Accordingly, the compstatin analogue of the present invention, or salts or solvates thereof, especially pharmaceutically acceptable salts and/or solvates thereof, may be formulated as compositions or pharmaceutical compositions prepared for storage or administration, and which comprise a therapeutically effective amount of a compstatin analogue of the present invention, or a salt or solvate thereof.

Suitable salts formed with bases include metal salts, such as alkali metal or alkaline earth metal salts.

- 5 In one embodiment, a pharmaceutical composition of the invention is one wherein the compstatin analogue is in the form of a pharmaceutically acceptable acid addition salt.

As will be apparent to one skilled in the medical art, a “therapeutically effective amount” of a compstatin analogue compound or pharmaceutical composition thereof of the present
10 invention will vary depending upon, inter alia, the age, weight and/or gender of the subject (patient) to be treated. Other factors that may be of relevance include the physical characteristics of the specific patient under consideration, the patient’s diet, the nature of any concurrent medication, the particular compound(s) employed, the particular mode of administration, the desired pharmacological effect(s) and the particular therapeutic indication.
15 Because these factors and their relationship in determining this amount are well known in the medical arts, the determination of therapeutically effective dosage levels, the amount necessary to achieve the desired result of treating and/or preventing and/or remedying malabsorption and/or low-grade inflammation described herein, as well as other medical indications disclosed herein, will be within the ambit of the skilled person.

20

As used herein, the term “a therapeutically effective amount” refers to an amount which reduces symptoms of a given condition or pathology, and preferably which normalizes physiological responses in an individual with that condition or pathology. Reduction of symptoms or normalization of physiological responses can be determined using methods
25 routine in the art and may vary with a given condition or pathology. In one aspect, a therapeutically effective amount of one or more compstatin analogues, or pharmaceutical compositions thereof, is an amount which restores a measurable physiological parameter to substantially the same value (preferably to within 30%, more preferably to within 20%, and still more preferably to within 10% of the value) of the parameter in an individual without the
30 condition or pathology in question.

In one embodiment of the invention, administration of a compound or pharmaceutical composition of the present invention is commenced at lower dosage levels, with dosage levels being increased until the desired effect of preventing/treating the relevant medical
35 indication is achieved. This would define a therapeutically effective amount. For the compstatin analogues of the present invention, alone or as part of a pharmaceutical composition, such human doses of the active compstatin analogue may be between about

0.01 pmol/kg and 500 μ mol/kg body weight, between about 0.01 pmol/kg and 300 μ mol/kg body weight, between 0.01 pmol/kg and 100 μ mol/kg body weight, between 0.1 pmol/kg and 50 μ mol/kg body weight, between 1 pmol/kg and 10 μ mol/kg body weight, between 5 pmol/kg and 5 μ mol/kg body weight, between 10 pmol/kg and 1 μ mol/kg body weight, between 50
 5 pmol/kg and 0.1 μ mol/kg body weight, between 100 pmol/kg and 0.01 μ mol/kg body weight, between 0.001 μ mol/kg and 0.5 μ mol/kg body weight, between 0.05 μ mol/kg and 0.1 μ mol/kg body weight.

The therapeutic dosing and regimen most appropriate for patient treatment will of course vary
 10 with the disease or condition to be treated, and according to the patient's weight and other parameters. Without wishing to be bound by any particular theory, it is expected that doses, in the mg/kg range, and shorter or longer duration or frequency of treatment may produce therapeutically useful results, such as a statistically significant inhibition of the alternative and classical complement pathways. The dosage sizes and dosing regimen most appropriate for
 15 human use may be guided by the results obtained by the present invention, and may be confirmed in properly designed clinical trials.

An effective dosage and treatment protocol may be determined by conventional means, starting with a low dose in laboratory animals and then increasing the dosage while
 20 monitoring the effects, and systematically varying the dosage regimen as well. Numerous factors may be taken into consideration by a clinician when determining an optimal dosage for a given subject.

For local delivery to the eye, the pharmaceutically acceptable compositions may be
 25 formulated in isotonic, pH adjusted sterile saline or water, either with or without a preservative such as benzylalkonium chloride. Alternatively, for ophthalmic uses, the pharmaceutically acceptable compositions may be formulated in an ointment such as petrolatum or as eyedrops. Methods of local administration to the eye include, e.g., choroidal injection, transscleral injection or placing a scleral patch, selective arterial catheterization, eyedrops or
 30 eye ointments, intraocular administration including transretinal, subconjunctival bulbar, intravitreal injection, suprachoroidal injection, subtenon injection, scleral pocket and scleral cutdown injection, by osmotic pump, etc. The agent can also be alternatively administered intravascularly, such as intravenously (IV) or intraarterially. In choroidal injection and scleral
 35 patching, the clinician uses a local approach to the eye after initiation of appropriate anesthesia, including painkillers and ophthalmoplegics. A needle containing the therapeutic compound is directed into the subject's choroid or sclera and inserted under sterile conditions. When the needle is properly positioned the compound is injected into either or

both of the choroid or sclera. When using either of these methods, the clinician can choose a sustained release or longer acting formulation. Thus, the procedure can be repeated only every several months or several years, depending on the subject's tolerance of the treatment and response.

5

The following examples are provided to describe the invention in greater detail. They are intended to illustrate, not to limit, the invention.

Example 1: Synthesis of Compstatin Analogues

10 **General Peptide Synthesis**

List of abbreviations and suppliers			
	Abbreviation	Name	Brand / Supplier
Resins			
		TentaGel™ PHB AA(Proct)-Fmoc	Rapp Polymere
		TentaGel™ SRAM	Rapp Polymere
Amino acids			
		Pseudoprolines (E.g. YS, FS, FT)	Jupiter Bioscience Ltd.
		Fmoc-L-AA-OH	Senn Chemicals AG
Coupling reagents			
	Oxyma Pure	Ethyl cyanoglyoxylate-2-oxime	Chem Impex international
	DIC	Diisopropylcarbodiimide	Fluka / Sigma Aldrich Co.
	HATU	N-[(dimethylamino)-1H-1,2,3-triazol[4,5-b]pyridine-1-ylmethylene]-N-methylmethanaminium hexafluorophosphate N-oxide	ChemPep Inc.
	HOBt	Hydroxybenzotriazole	Sigma-Aldrich Co.
Solvents and reagents			
	Boc ₂ O	Di-tert-butyl pyrocarbonate	Advanced ChemTech
	DCM	Dichloromethane	Prolabo (VWR)
	DIPEA	Diisopropylethylamine	Fluka / Sigma Aldrich Co.
	DMF	N,N-dimethylformamide	Taminco
	Et ₂ O	Diethyl ether	Prolabo (VWR)

	EtOH	Ethanol	CCS Healthcare AB
	HCOOH	Formic acid (HPLC grade)	Sigma-Aldrich Co.
	H ₂ O	Water, Milli-Q water	Millipore
	MeCN	Acetonitrile (HPLC)	Sigma-Aldrich Co.
	NMP	N-methylpyrrolidone	Sigma-Aldrich Co.
		Piperidine	Jubliant Life Sciences Ltd.
	TFA	Trifluoroacetic acid (HPLC)	Chemicals Raw Materials Ltd.
	TIS	Triisopropylsilane	Sigma-Aldrich Co.
	DODT	2,2'-(ethylenedioxy)diethanethiol	Sigma-Aldrich Co.
Other	MeOH	Methanol	Sigma-Aldrich Co.
		Ascorbic acid	Sigma-Aldrich Co.
	I ₂	Iodine	Sigma-Aldrich Co.

Apparatus and synthetic strategy

Peptides were synthesized batchwise on a peptide synthesiser, such as a CEM Liberty Peptide Synthesizer or a Symphony X Synthesizer, according to solid phase peptide synthetic procedures using 9-fluorenylmethyloxycarbonyl (Fmoc) as N- α -amino protecting group and suitable common protection groups for side-chain functionalities.

As polymeric support based resins, such as e.g. TentaGel™, was used. The synthesizer was loaded with resin that prior to usage was swelled in DMF.

Coupling

CEM Liberty Peptide Synthesizer

A solution of Fmoc-protected amino acid (4 equiv.) was added to the resin together with a coupling reagent solution (4 equiv.) and a solution of base (8 equiv.). The mixture was either heated by the microwave unit to 70-75°C and coupled for 5 minutes or coupled with no heat for 60 minutes. During the coupling nitrogen was bubbled through the mixture.

Symphony X Synthesizer

The coupling solutions were transferred to the reaction vessels in the following order: amino acid (4 equiv.), HATU (4 equiv.) and DIPEA (8 equiv.). The coupling time was 10 min at room temperature (RT) unless otherwise stated. The resin was washed with DMF (5 x 0,5 min). In case of repeated couplings the coupling time was in all cases 45 min at RT.

Deprotection*CEM Liberty Peptide Synthesizer*

The Fmoc group was deprotected using piperidine in DMF or other suitable solvents. The deprotection solution was added to the reaction vessel and the mixture was heated for 30 sec. reaching approx. 40°C. The reaction vessel was drained and fresh deprotection solution was added and subsequently heated to 70-75°C for 3 min. After draining the reaction vessel the resin was washed with DMF or other suitable solvents.

10 *Symphony X Synthesizer*

Fmoc deprotection was performed for 2,5 minutes using 40% piperidine in DMF and repeated using the same conditions. The resin was washed with DMF (5 x 0,5 min).

Side chain acylation

15 Fmoc-Lys(Dde)-OH or alternatively another amino acid with an orthogonal side chain protective group was introduced at the position of the acylation (side-chain lipidation). The N-terminal of the linear peptide was protected with Ac or Boc. While the peptide was still attached to the resin, the orthogonal side chain protective group was selectively cleaved using freshly prepared hydrazine hydrate (2-4%) in NMP for 2 x 15 min. The unprotected lysine side chain was then elongated using standard coupling conditions and Fmoc-deprotections with the desired building block. The lipidation moiety was coupled as the last step.

Cleavage

25 The dried peptide resin was treated with TFA and suitable scavengers for approximately 2 hours. The volume of the filtrate was reduced and the crude peptide was precipitated after addition of diethylether. The crude peptide precipitate was washed several times with diethylether and finally dried.

30 *HPLC purification of the crude peptide*

The crude peptide was purified by preparative reverse phase HPLC using a conventional HPLC apparatus, such as a Gilson GX-281 with 331/332 pump combination, for binary gradient application equipped with a column, such as 5 x 25 cm Gemini NX 5u C18 110A column, and a fraction collector using a flow 20-40 ml/min with a suitable gradient of buffer A (0.1% Formic acid, aq.) or A (0.1% TFA, aq.) and buffer B (0.1% Formic acid, 90% MeCN, aq.) or B (0.1% TFA, 90% MeCN, aq.). Fractions were analyzed by analytical HPLC and MS and

selected fractions were pooled and lyophilized. The final product was characterized by HPLC and MS.

Oxidation

- 5 Following purification and lyophilisation of the crude linear peptide, the peptide was redissolved in 0.1% TFA in water, acetonitrile and acetic acids until a clear solution. The concentration of the peptide solution was kept at approx. 1-2 mg/ml depending on the peptides ability to solubilize. The peptide solution was stirred, while a solution of iodine in methanol (approx. 1.5 equiv.) was added drop-wise until the peptide solution obtain an orange colour. After 10-15 minutes, the oxidation was finished and excess iodine was reduced with a solution of ascorbic acid in water (1 equiv.) until a colourless peptide solution. The peptide solution was diluted with water before preparative HPLC purification.

Analytical HPLC

- 15 Final purities were determined by analytic HPLC (Agilent 1100/1200 series) equipped with auto sampler, degasser, 20 µl flow cell and Chromeleon software. The HPLC was operated with a flow of 1.2 ml/min at 40°C using an analytical column, such as Kinetex 2.6 µm XB-C18 100A 100X8,6 mm column. The compound was detected and quantified at 215 nm. Buffers A (0.1% TFA, aq.) and buffer B (0.1% TFA, 90% MeCN, aq.).

20

Mass spectroscopy

- Final MS analysis were determined on a conventional mass spectroscopy, e.g. Waters Xevo G2 TOF, equipped with electrospray detector with lock-mass calibration and MassLynx software. It was operated in positive mode using direct injection and a cone voltage of 15V (1 TOF), 30 V (2 TOF) or 45 V (3 TOF) as specified on the chromatogram. Precision was 5 ppm with a typical resolution of 15,000-20,000.

25

Synthesis of compound No 24:

Ac-IC(1)IWQDWGEHRC(1)TEGE-NH₂

30

Solid phase peptide synthesis was performed on a Symphony X Synthesizer using standard Fmoc chemistry. TentaGel S RAM (2,51 g; 0.23 mmol/g) was swelled in DMF (20 ml) prior to use and the Fmoc-group was deprotected according to the procedure described above.

Coupling

- 35 Suitable protected Fmoc-amino acids according to the sequence were coupled as described above using HATU as coupling reagent. All couplings were performed at R.T.

Deprotection

Fmoc deprotection was performed according to the procedure described above.

5 Cleavage of the peptide from the solid support

The peptide-resin was washed with EtOH (3 x 10 ml) and Et₂O (3 x 10 ml) and dried to constant weight at room temperature (r.t.). The peptide was cleaved from the resin by treatment with TFA/DODT (95/5; 60 ml, 2 h; r.t.). The volume of the filtrate was reduced and the crude peptide was precipitated after addition of diethylether. The crude peptide

10 precipitate was washed several times with diethylether and finally dried to constant weight at room temperature yield 760 mg crude peptide product (purity ~30%).

HPLC purification of the crude linear peptide

The crude peptide was purified by preparative reverse phase HPLC using a Gilson GX-

15 281with 331/332 pump combination for binary gradient application equipped with a 5 x 25 cm Gemini NX 5u C18 110A, column and a fraction collector and run at 35 ml/min with a gradient of buffer A (0.1% TFA, aq.) and buffer B (0.1% TFA, 90% MeCN, aq.) gradient from 20%B to 45%B in 47 min. Fractions were analyzed by analytical HPLC and MS and relevant fractions were pooled and lyophilized to yield 190 mg, with a purity of 85% as characterized by HPLC

20 and MS as described above. Calculated monoisotopic MW = 2001.58 found 2001.81.

Oxidation of the crude linear peptide

The 190 mg purified linear peptide was dissolved in 220 ml 0.1% TFA in water (65%) and acetonitrile (35%) until a clear solution. The peptide solution was stirred, while a solution of

25 iodine in methanol (2.2 mL, approx. 1.5 equiv. iodine) was added drop-wise until the peptide solution obtain an orange colour. The reaction was followed by analytic HPLC but already after 10-15 minutes, the oxidation was finished. Excess iodine was reduced with a solution of ascorbic acid in water (220 µL, approx.1 equiv.) until a colourless peptide solution. The peptide solution was reduced slightly by rota evaporation before purification on preparative

30 HPLC.

HPLC purification of the oxidized peptide

The crude peptide was purified by preparative reverse phase HPLC using a Gilson GX-

35 281with 331/332 pump combination for binary gradient application equipped with a 5 x 25 cm Gemini NX 5u C18 110A, column and a fraction collector and run at 35 ml/min with a gradient of buffer A (0.1% TFA, aq.) and buffer B (0.1% TFA, 90% MeCN, aq.) gradient from 20%B to 45%B in 47 min. Fractions were analyzed by analytical HPLC and MS and relevant fractions

were pooled and lyophilized to yield 138 mg, with a purity of 92% as characterized by HPLC and MS as described above. Calculated monoisotopic MW = 1999.83 found 1999.54.

5 Synthesis of compound No 119

Ac-SEFC(1)-I[1-Me-Trp]QDWGEHRC(1)-[Sar]-EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-(8-amino-3,6-dioxaoctanoyl)-γGlu-(8-amino-3,6-dioxaoctanoyl))-NH₂

10 Solid phase peptide synthesis was performed on a Symphony X Synthesizer using standard Fmoc chemistry. TentaGel S RAM (3 x ~1.3 g; 0.22 mmol/g) was swelled in DMF (3 x 10 ml) prior to use and the Fmoc-group was deprotected according to the procedure described above.

Coupling

15 Suitable protected Fmoc-amino acids according to the sequence were coupled as described above using HATU as coupling reagent. All couplings were performed at R.T. The lysine used for the incorporation of the branched moiety was incorporated as Fmoc-Lys(Dde)-OH for orthogonal coupling

20 Deprotection

Fmoc deprotection was performed according to the procedure described above.

Side chain acylation

25 While the peptide was still attached to the resin, the orthogonal side-chain protective group (Dde) was selectively cleaved using freshly prepared hydrazine hydrate (2-4%) in NMP for 2 x 15 min. The unprotected lysine side-chain was doubled coupled with Fmoc-Peg3-OH followed by single couplings with Fmoc-Glu-OtBu, Fmoc-8-Amino-3,6-dioxaoctanoyl-OH, Fmoc-Gly-OH, Fmoc-Glu-OtBu and lastly the fatty acid moiety 17-carboxy-heptadecanoic acid mono tert-butyl ester using standard coupling conditions.

30 Cleavage of the peptide from the solid support

The peptide-resin was washed with EtOH (3 x 15 ml) and Et₂O (3 x 150 ml) and dried to constant weight at room temperature (r.t.). The peptide was cleaved from the resin by treatment with TFA/DODT (95/5; 120 ml, 2 h; r.t.). The volume of the filtrate was reduced and the crude peptide was precipitated after addition of diethylether. The crude peptide
35 precipitate was washed several times with diethylether and finally dried to constant weight at room temperature yield 2.36 g crude peptide product (purity ~41-48%).

HPLC purification of the crude linear peptide

The crude peptide was purified by preparative reverse phase HPLC using a Gilson GX-281 with 331/332 pump combination for binary gradient application equipped with a 5 x 25 cm Gemini NX 5u C18 110A, column and a fraction collector and run at 35 ml/min with a gradient of buffer A (0.1% TFA, aq.) and buffer B (0.1% TFA, 90% MeCN, aq.) gradient from 30%B to 60%B in 47 min. Fractions were analyzed by analytical HPLC and MS and relevant fractions were pooled and lyophilized to yield 744 mg, with a purity of 84% as characterized by HPLC and MS as described above. Calculated monoisotopic MW = 3207.47 found 3207.32.

Oxidation of the crude linear peptide

The 744 mg purified linear peptide was dissolved in 350 ml 0.1% TFA in water, 150 ml acetonitrile and 100 ml acetic acid until a clear solution (total volume 600 ml). The peptide solution was stirred, while a solution of iodine in methanol (4.7 mL, approx. 1.5 equiv. iodine) was added drop-wise until the peptide solution obtain an orange colour. The reaction was followed by analytic HPLC but already after 10-15 minutes, the oxidation was finished. Excess iodine was reduced with a solution of ascorbic acid in water (150 µL, approx. 1 equiv.) until a colourless peptide solution. The peptide solution was reduced slightly by rota evaporation before purification on preparative HPLC.

HPLC purification of the oxidized peptide

The crude peptide was purified by preparative reverse phase HPLC using a Gilson GX-281 with 331/332 pump combination for binary gradient application equipped with a 5 x 25 cm Gemini NX 5u C18 110A, column and a fraction collector and run at 35 ml/min with a gradient of buffer A (0.1% TFA, aq.) and buffer B (0.1% TFA, 90% MeCN, aq.) gradient from 30%B to 60%B in 47 min. Fractions were analysed by analytical HPLC and MS and relevant fractions were pooled and lyophilized to yield 510 mg, with a purity of 91% as characterized by HPLC and MS as described above. Calculated monoisotopic MW = 3205.47 found 3205.23.

Table 1: Synthesized compounds:

Compound	Sequence
Compstatin 1-13	H-IC(1)VVQDWGHHRC(1)T-NH ₂
Ac-compstatin	Ac-IC(1)VVQDWGHHRC(1)T-NH ₂
4W9A*	Ac-IC(1)VWQDWGAHRC(1)T-NH ₂
Cp40*	H-{d}YIC(1)V[1-Me-Trp]QDW[Sar]AHRC(1)[N-Me-Ile]-NH ₂
A	Ac-IC(1)VWQDWGEHRC(1)T-NH ₂
B	Ac-IC(1)VWQDWGSHRC(1)T-NH ₂

C	Ac-ESSAIC(1)VWQDWGEHRC(1)T-NH2
D	Ac-IC(1)VWQDWGEHRC(1)TGAES-NH2
E	Ac-IC(1)VWQDWGAHSC(1)T-NH2
F	Ac-IC(1)VWQDWGEHSC(1)T-NH2
G	Ac-IC(1)VWQDWGEHRC(1)S-NH2
H	Ac-EGSAIC(1)VWQDWGEHRC(1)[Sar]E-NH2
J	Ac-IC(1)VWQDWGEHRC(1)TEGE-NH2
1	Ac-IC(1)IWQDWGAHRC(1)T-NH2
2	Ac-IC(1)IWQDWGEHRC(1)T-NH2
3	Ac-ESSAIC(1)IWQDWGEHRC(1)T-NH2
4	Ac-IC(1)I[1-Me-Trp]QDWGEHRC(1)T-NH2
5	Ac-IC(1)IWQDWGKHRC(1)T-NH2
6	Ac-IC(1)IWQDWGSHRC(1)T-NH2
7	Ac-IC(1)IWQKWGEHRC(1)T-NH2
8	Ac-IC(1)IWQKWGAHRC(1)TGAES-NH2
9	Ac-YC(1)IWQDWGEHRC(1)T-NH2
10	Ac-ESSAYC(1)IWQDWGEHRC(1)T-NH2
11	Ac-[Sar]C(1)IWQDWGEHRC(1)T-NH2
12	Ac-IC(1)IWQDWGAHRC(1)E-NH2
13	Ac-IC(1)IWQDWGEHRC(1)[Sar]-NH2
14	Ac-ESSAIC(1)IWQDWGEHRC(1)TGAES-NH2
15	Ac-IC(1)IWQDWGEHRC(1)TGAES-NH2
16	Ac-IC(1)IWQEWGEHRC(1)T-NH2
17	Ac-IC(1)IWQDWGDHRC(1)T-NH2
18	Ac-IC(1)IWQDWGRHRC(1)T-NH2
19	Ac-IC(1)IWQDWGAHSC(1)T-NH2
20	Ac-IC(1)IWQDWGEHSC(1)T-NH2
21	Ac-IC(1)IWQDWGEHRC(1)S-NH2
22	Ac-IC(1)IWQDWGEHRC(1)E-NH2
23	Ac-FC(1)IWQDWGEHRC(1)T-NH2
24	Ac-IC(1)IWQDWGEHRC(1)TEGE-NH2
25	Ac-IC(1)IWQDWGEHRC(1)TEA-NH2
26	Ac-IC(1)IWQDWGEHRC(1)TE-NH2
27	Ac-IC(1)IWQDWGEHRC(1)EGE-NH2
28	Ac-EGSAIC(1)IWQDWGEHRC(1)[Sar]E-NH2
29	Ac-EGSAIC(1)IWQDWGEHRC(1)T-NH2

30	Ac-EGEIC(1)IWQDWGEHRC(1)T-NH2
31	Ac-ESEIC(1)IWQDWGEHRC(1)T-NH2
32	Ac-SEIC(1)IWQDWGEHRC(1)TEA-NH2
33	Ac-EIC(1)IWQDWGEHRC(1)TE-NH2
34	Ac-EIC(1)IWQDWGEHRC(1)TEGE-NH2
35	Ac-EGEIC(1)IWQDWGEHRC(1)EGE-NH2
36	Ac-ESEIC(1)IWQDWGEHRC(1)EGE-NH2
37	Ac-KEKIC(1)IWQDWGEHRC(1)TEKE-NH2
38	Ac-EKGIC(1)IWQDWGEHRC(1)TEKP-NH2
39	Ac-IC(1)IWQDWGEHRC(1)TEGK-NH2
40	Ac-GSAIC(1)IWQDWGEHRC(1)[Sar]E-NH2
41	Ac-SAIC(1)IWQDWGEHRC(1)[Sar]E-NH2
42	Ac-SAIC(1)IWQDWGEHRC(1)TEG-NH2
43	Ac-FC(1)IWQDWGEHRC(1)TGAE-NH2
44	Ac-EGSAIC(1)IWQDWGEHRC(1)[Sar]EGE-NH2
45	Ac-EGSAFC(1)IWQDWGEHRC(1)[Sar]E-NH2
46	Ac-ESSAIC(1)IWQDWGAHRC(1)T-NH2
47	Ac-IC(1)IWQDWGAHRC(1)TGAES-NH2
48	H-{d}YIC(1)I[1-Me-Trp]QDW[Sar]AHRC(1)[N-Me-Ile]-NH2
49	Ac-EGSAIC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH2
50	Ac-EGSAIC(1)I[2NaI]QDWGEHRC(1)[Sar]E-NH2
51	Ac-IC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-NH2
52	Ac-IC(1)I[2NaI]QDWGEHRC(1)TGAES-NH2
53	Ac-EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH2
54	Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH2
55	Ac-EGSAIC(1)IWQDWGEHRC(1)TE-NH2
56	Ac-EGSAFC(1)I[1NaI]QDWGEHRC(1)TE-NH2
57	Ac-EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)TE-NH2
58	Ac-EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)EGE-NH2
59	Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)TE-NH2
60	Ac-EGSAFC(1)I[2NaI]QDWGEHRC(1)TE-NH2
61	Ac-FC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-NH2
62	Ac-YC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-NH2
63	Ac-FC(1)I[1NaI]QDWGEHRC(1)TGAES-NH2
64	Ac-FC(1)I[2NaI]QDWGEHRC(1)TGAES-NH2
65	Ac-YC(1)I[2NaI]QDWGEHRC(1)TGAES-NH2

66	Ac-YC(1)IWQDWGEHRC(1)TGAES-NH2
67	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-NH2
68	Ac-YC(1)I[1-Me-Trp]QDWGEHRC(1)TEAGS-NH2
69	Ac-YC(1)I[1-Me-Trp]QDWGEHRC(1)TESGA-NH2
70	Ac-EGSAYC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]E-NH2
71	Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH2
72	Ac-FC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TGAES-NH2
73	H-{d}YFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TGAES-NH2
74	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]GAES-NH2
75	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH2
76	Ac-SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)[Sar]EA-NH2
77	Ac-SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TEA-NH2
78	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH2
79	Ac-SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)[Sar]E-NH2
80	Ac-EFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH2
81	Ac-SE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH2
82	Ac-SE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA-NH2
83	Ac-SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EA-NH2
84	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)SEA-NH2
85	Ac-EFC(1)I[1-Me-Trp]QDWGEHRC(1)ES-NH2
86	Ac-SEFC(1)I[1-Me-Trp]QDWGEHKC(1)[Sar]EA-NH2
87	Ac-GEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH2
88	Ac-GE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA-NH2
89	Ac-SE[Sar]C(1)I[1-Me-Trp]QEW[Sar]EHRC(1)TEA-NH2
90	Ac-SE[Sar]C(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EA-NH2
91	H-{d}Y[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA-NH2

Table 1 (contd.):

Compound	Sequence
92	Ac-IC(1)IWQDWGEHRC(1)TEG-K([15-carboxy-pentadecanoyl]-γGlu)-NH2
93	Ac-IC(1)IWQDWGEHRC(1)TEG-K([15-carboxy-pentadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH2
94	Ac-IC(1)IWQDWGEHRC(1)TEGE-K([15-carboxy-pentadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH2
95	Ac-IC(1)IWQDWGEHRC(1)TEG-K((15-carboxy-pentadecanoyl)-[(Piperazine-1-yl)-acetyl]-8-Amino-3,6-

	dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH2
96	Ac-IC(1)IWQDWGEHRC(1)TEG-K([17-carboxy-heptadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH2
97	Ac-IC(1)IWQDWGEHRC(1)TEGE-K([17-carboxy-heptadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH2
98	Ac-IC(1)IWQDWGEHRC(1)TEG-K([19-carboxy-nonadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH2
99	[15-Carboxy-pentadecanoyl]-ESSAIC(1)IWQDWGEHRC(1)TEGE-NH2
100	Ac-[K([15-carboxy-pentadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)]GSAIC(1)IWQDWGEHRC(1)TEGE-NH2
101	Ac-EGSAIC(1)IWQDWGEHRC(1)TEG-K([15-carboxy-pentadecanoyl]-γGlu)-NH2
102	Ac-FC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-K([15-carboxy-pentadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH2
103	Ac-EGSAYC(1)I[1-Me-Trp]QDWGEH-K([15-carboxy-pentadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-C(1)[Sar]E-NH2
104	Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EG-K([15-carboxy-pentadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH2
105	Ac-SAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-K([17-carboxy-heptadecanoyl]-γGlu-KG-γGlu)-NH2
106	Ac-SAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2
107	[15-Carboxy-pentadecanoyl]-EGSEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH2
108	[17-Carboxy-heptadecanoyl]-EGSEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH2
109	Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2
110	Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGK-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2
111	Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EK[γGlu]-K([17-carboxy-heptadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH2
112	Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2
113	Ac-ASGEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2
114	Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2
115	Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGK-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH2
116	Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-K([17-carboxy-heptadecanoyl]-γGlu-K-γGlu)-NH2
117	Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([17-

	carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
118	Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-(8-Amino-3,6-dioxaoctanoyl)-γGlu-(8-Amino-3,6-dioxaoctanoyl))-NH ₂
119	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-(8-Amino-3,6-dioxaoctanoyl)-γGlu-(8-Amino-3,6-dioxaoctanoyl))-NH ₂
120	Ac-SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-(8-Amino-3,6-dioxaoctanoyl)-γGlu-(8-Amino-3,6-dioxaoctanoyl))-NH ₂
121	Ac-SEYC(1)I[1-Me-Trp]QEW[Sar]EHRC(1)[Sar]EK[γGlu]A-K([17-carboxy-heptadecanoyl]-γGlu-G-(8-Amino-3,6-dioxaoctanoyl)-γGlu-(8-Amino-3,6-dioxaoctanoyl))-NH ₂
122	Ac-SEYC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-(8-Amino-3,6-dioxaoctanoyl)-γGlu-(8-Amino-3,6-dioxaoctanoyl))-NH ₂
123	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
124	Ac-SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu))-NH ₂
125	Ac-SEYC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
126	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
127	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)-[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([15-carboxy-pentadecanoyl]-γGlu-G-γGlu)-NH ₂
128	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([19-carboxy-nonadecanoyl]-γGlu-G-γGlu)-NH ₂
129	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGEGGG-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
130	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGEGGG-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
131	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGEGGG-K([15-carboxy-pentadecanoyl]-γGlu-G-γGlu)-NH ₂
132	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EK[γGlu]GGG-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
133	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEK[γGlu]GGG-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
134	Ac-EFC(1)I[1-Me-Trp]QDWGEHRC(1)EGE-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
135	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-K([15-carboxy-hexadecanoyl]-γGlu-G-γGlu)-NH ₂
136	Ac-SEFC(1)I[1-Me-Trp]-QDWGEHRC(1)TEGE-[8-aminooctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
137	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE-[8-aminooctanoyl]-E-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
138	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-[8-Amino-

	3,6-dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
139	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGESES-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
140	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl]ES-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH ₂
141	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGESES-K([17-carboxy-heptadecanoyl]-γGlu)-NH ₂
142	Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE[8-Amino-3,6-dioxaoctanoyl]ES-K([17-carboxy-heptadecanoyl]-γGlu)-NH ₂

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4W9A – described by Mallik et al., J. Med. Chem. 2005, 48, 274-286 (“V4W/H9A”).

- 5 Cp40- described by Qu et al., Immunobiology 2013, 281(4): 496-505 (also referred to in that paper as “peptide 14).

Example 2: *In vitro* haemolysis assay

Method

- 10 The *in vitro* effect of the compounds of the present invention was assessed by measuring their inhibitory effect of the classical complement pathway in a haemolysis assay.

Briefly, compounds of the present invention and reference compounds were dissolved in DMSO and diluted in Tris/Casein Assay Buffer (10 mM Tris, 145 mM NaCl, 0.5 mM MgCl₂,
 15 0.15 mM CaCl₂, and 0.1 % W/V Casein, adjusted to pH 7.4) as 9-point serial dilutions in a 96 well plate. Sensitized sheep red blood cells (RBC) coated with rabbit anti-sheep erythrocyte antiserum (Complement Technology, Inc., TX, USA) were washed in Tris/Casein Assay Buffer. 50 µl from each well of diluted compound was added to a 96-well plate containing 50 µl diluted human serum (Complement Technology, Inc., TX, USA) and incubated for 15
 20 minutes at room temperature. The serum dilution factor was optimized for every serum batch to obtain 70-90% of maximal haemolysis using the protocol. Then 50 µl sensitized sheep red blood cells were added to all wells (10⁷ per well).

After 30 minutes of incubation at 37 °C with gentle agitation, the reaction was stopped by addition of 50 µl Tris STOP Buffer per well (10 mM EDTA, 10 mM Tris, 145 mM NaCl
 25 adjusted to pH 7.4). The RBCs were then removed by centrifugation and the resulting supernatant measured for hemolysis by absorbance at 405 nm.

The response was normalized relative to a positive and negative control (vehicle) to calculate the IC₅₀ from the concentration response curve using the 4-parameter logistic (4PL)
 30 nonlinear model for curve fitting. All values are based on n=>2 independent determinations.

Table 2: Effect of exchange from valine to isoleucine. Compound 1 differs from the prior art compound 4W9A only by the presence of Ile instead of Val at position 3.

Comp	CP															
no	hemolysis	1	2	3	4	5	6	7	8	9	10	11	12	13		
	IC50 (nM)															
1	150	Ac	I	C(1)	I	W	Q	D	W	G	A	H	R	C(1)	T	NH ₂
4W9A	250	V														

5

Further compounds were tested as shown below.

Table 3: *in vitro* analysis of inhibition of hemolysis

Compound	IC50[nM]
Compstatin	>5μM
Ac-compstatin	>5μM
4W9A	<500
1	<250
2	<100
3	<100
4	<100
5	<250
6	<250
7	<1000
8	<500
9	<100
10	<100
11	<100
12	<100
13	<100
14	<100
15	<100
16	<100
17	<100
18	<100
19	<250
20	<100
21	<100
22	<100
23	<100
24	<100
25	<100
26	<100
27	<100
28	<100
29	<100
30	<100
31	<100

32	<100
33	<100
34	<250
35	<500
36	<250
37	<250
38	<250
39	<100
40	<250
41	<250
42	<250
43	<100
44	<250
45	<100
46	<100
47	<100
48	<100
49	<100
50	<100
51	<100
52	<100
53	<100
54	<100
55	<250
56	<100
57	<100
58	<100
59	<100
60	<100
61	<100
62	<100
63	<100
64	<100
65	<100
66	<100
67	<100
68	<100
69	<100
70	<100
71	<100
72	<100
73	<100
74	<100
75	<100
76	<100
77	<100
78	<100
79	<100
80	<100
81	<100
82	<100
83	<100
84	<100

85	<100
86	<100
87	<100
88	<100
89	<100
90	<250
91	<100

Table 3 (contd.)

Compound	IC50[nM]
92	<1000
93	<500
94	<500
95	<500
96	<1000
97	<250
98	<500
99	<250
100	<500
101	<500
102	<100
103	<100
104	<100
105	<100
106	<250
107	<100
108	<500
109	<250
110	<250
111	<100
112	<500
113	<500
114	<500
115	<250
116	<500
117	<250
118	<100
119	<100
120	<250
121	<250
122	<500
123	<100
124	<100
125	<500

126	<100
127	<100
128	<100
129	<100
130	<100
131	<100
132	<100
133	<100
134	<100
135	<100
136	<100
137	<100
138	<100
139	<100
140	<100
141	<100
142	<100

The following pairs of compounds, each of which differ only at position 3, show that the effects of replacing valine by isoleucine are seen in compounds having a variety of peptide backbone sequences.

5

Table 4 : Direct comparison of valine 3 to isoleucine 3 in combination with modification at position 9, 11 and/or 13.

Compound	CP hemolysis IC50 (nM)		1	2	3	4	5	6	7	8	9	10	11	12	13	
2	94	Ac	I	C(1)	I	W	Q	D	W	G	E	H	R	C(1)	T	NH ₂
A	350				V											
6	140	Ac	I	C(1)	I	W	Q	D	W	G	S	H	R	C(1)	T	NH ₂
B	360				V											
3	69	Ac ESSA	I	C(1)	I	W	Q	D	W	G	E	H	R	C(1)	T	NH ₂
C	300				V											
15	47	Ac	I	C(1)	I	W	Q	D	W	G	E	H	R	C(1)	T	GAES NH ₂
D	210				V											
19	140	Ac	I	C(1)	I	W	Q	D	W	G	A	H	S	C(1)	T	NH ₂
E	>1000				V											
20	59	Ac	I	C(1)	I	W	Q	D	W	G	E	H	S	C(1)	T	NH ₂
F	540				V											
21	77	Ac	I	C(1)	I	W	Q	D	W	G	E	H	R	C(1)	S	NH ₂

G	180	V												
28	88	Ac	EGSA	I	C(1)	I	W	Q	D	W	G	E	H	R C(1) Sar E NH ₂
H	330	V												
24	90	Ac		I	C(1)	I	W	Q	D	W	G	E	H	R C(1) T EGE NH ₂
J	240	V												

Isoleucine at position 3 was also demonstrated to be superior compared to other residues often considered to be “conservative” replacements for isoleucine.

5 Table 5: Effect on hemolysis of different residues at position 3

Ac-IC(1)XWQDWGEHRC(1)T-NH₂

Compound	Position 3 (X)	IC ₅₀ , CP hemolysis (nM)
A	Valine	350
2	Isoleucine	94
-	Leucine	500
-	Norvaline	>1000
-	Norleucine	480
-	Phenylalanine	>10000
-	Beta-Homo-Isoleucine	>10000

- Due to the high concentration of C3 found in serum, it may be difficult to use the hemolysis assay to differentiate between compounds having very high affinity for C3.
- In such circumstances, it may be possible to determine a more accurate hierarchy of binding affinity to C3 by SPR measurements using immobilized C3, as described below.

Example 3: Solubility test

15 **Materials and method**

Compound solubility at 10 mg/mL

The solubility of compounds was assessed by measuring light scattering over a pH interval from pH 4 to pH 7.5.

Compounds were dissolved in a stock solution of 20 mg/mL in H₂O at pH 2.5 or pH 10.

- 20 These stock solutions were diluted 1:1 with 200 mM buffered solution to reach a final solution of 10 mg/mL compound in 100 mM buffer. The 5 investigated conditions were (1) acetate pH 4.0, (2) acetate pH 5.0, (3) phosphate pH 6.0, (4) phosphate pH 7 and (5) phosphate pH 7.5.

These samples were equilibrated for 15 minutes at ambient temperature, before evaluating solubility by visual inspection and absorbance measurements in a SpectraMax 190 microplate reader (Molecular Devices).

5 Visual inspection

Visual inspection included manually checking the 96 well plate for wells that are clear or non-clear. In addition to this a picture of the 96 well plate is taken.

Microplate reader and light scattering

10 Absorbance was measured at four wavelengths: 280 nm, 325 nm, 340 nm and 360 nm in an UV transparent 96 well microplate in a SpectraMax 190 microplate reader (Molecular Devices). The compounds do not absorb at 325-360 nm and signal at these wavelengths are therefore an expression of light scattering, which reflects the presence of visible or sub-visible particles that are detected as increased signal.

15 The light scattering was normalized to the signal from pure buffer solutions (100 mM) and compound solubility was evaluated as good (+) or poor (-). The criteria for this was a combination of visual inspection and light scattering not exceeding 0.1 AU, where values below 0.1 AU are good in visually clear samples.

20

Solubility of Comp No 24: (Sequence Ac-IC(1)IWQDWGEHRC(1)TEGE-NH₂)

Stock solution

Comp No 24 was carefully weighed out and dissolved in pH 2.5 H₂O-Cl. The stock solution was equilibrated 15 minutes at ambient temperature, at which point no visible particles were present. 200 mM buffer stock solutions were prepared for each pH condition.

25

Solubility assay:

The formulations for solubility testing were made by mixing 50 µL Comp No 24 stock solution and 50 µL buffer stock solution with gentle mixing by pipetting the solution a couple of times.

30 This was done for each buffer/pH condition in a UV transparent 96 well microplate (Corning 96 well REF 3635). Reference samples without Comp No 24 were made by mixing 50 µL pH 2.5 H₂O-Cl and 50 µL buffer stock solution. The plate was covered with a lid and left 15 minutes at ambient temperature before assessing solubility.

Measuring solubility:

Solubility was assessed by visual inspection of each formulation and a picture taken in a photo box. Light scattering was measured at 280 nm, 325 nm, 340 nm and 360 nm in a SpectraMax 190 microplate reader (Molecular Devices).

5

The visual inspection revealed that condition 1, 2 and 3 were cloudy and condition 2 additionally contained visible precipitates. The absorbance measurement confirmed the visual evaluation with condition 1, 2 and 3 all exceeding 0.1 AU threshold. Condition 4 and 5 were thus deemed good conditions for solubility of 10 mg/mL Comp No 24.

10

Similarly, additional compounds were tested for solubility (Table 6).

Table 6: Table of most soluble compounds, as tested at 10 mg/mL. “+” denotes solubility at the given condition, as determined by UV absorbance being less than 0.1 AU at 340 nm and the sample being clear when manually inspected. “-” denotes lack of solubility at the given condition, as UV absorbance at 340 nm exceeds 0.1 AU and/or it is visibly turbid or contains particles.

Comp No	Buffer & pH				
	Condition 1	Condition 2	Condition 3	Condition 4	Condition 5
	Acetate pH 4	Acetate pH 5	Phosphate pH 6	Phosphate pH 7	Phosphate pH 7.5
1	+	-	-	-	-
3	+	-	-	+	+
14	+	-	-	+	+
15	+	-	-	+	+
22	+	-	-	+	+
24	-	-	-	+	+
25	+	-	-	+	+
27	-	-	-	+	+
28	+	+	+	+	+
30	-	-	-	+	+
31	+	+	+	+	+
32	-	-	-	+	+
33	-	-	+	+	+
36	-	-	-	+	+

40	-	-	+	+	+
41	-	-	+	+	+
44	-	-	+	+	+
45	-	+	+	+	+
49	-	-	+	+	+
50	-	-	+	+	+
51	-	-	+	+	+
52	-	-	+	+	+
53	-	-	+	+	+
54	-	-	+	+	+
55	-	-	+	+	+
56	-	-	+	+	+
57	-	-	+	+	+
60	-	-	+	+	+
61	-	-	+	+	+
62	-	-	+	+	+
63	-	-	+	+	+
65	-	-	+	+	+
66	-	-	+	+	+
67	-	-	+	+	+
68	-	-	+	+	+
72	-	-	+	+	+
73	+	-	-	+	+
74	-	+	+	+	+
76	-	-	+	+	+
77	-	-	+	+	+
78	-	-	+	+	+
79	-	-	+	+	+
80	-	-	+	+	+
81	-	-	+	+	+

Table 6 (contd.)

Comp No	Buffer & pH				
	Condition 1	Condition 2	Condition 3	Condition 4	Condition 5
	Acetate pH 4	Acetate pH 5	Phosphate pH 6	Phosphate pH 7	Phosphate pH 7.5
102	-	+	+	+	+
103	-	+	+	+	+
104	-	+	+	+	+
105	-	-	+	+	+
107	-	-	+	+	+
108	-	-	+	+	+
109	-	+	+	+	+
111	-	-	+	+	+
114	+	+	+	+	+
115	-	-	+	+	+
116	-	-	+	+	+
118	-	+	+	+	+

5 **Example 4:** Affinity measurements by surface plasmon resonance (SPR)

Method

Surface plasmon resonance (SPR) was used to characterize peptides with respect to their binding affinity (K_d) for C3. Human C3 (Complement tech cat #A113c) was immobilised on individual flow cells of CM5 sensor chips (GE Healthcare) using standard amine coupling to a density of approximately 3000 resonance units (RU) in a buffer consisting of 10 mM phosphate pH 7.4, 150 mM NaCl, 0.05% Tween20.

For interaction experiments a multi-cycle experiment approach was used and performed using a BiacoreT200™ instrument (GE Healthcare) at 25°C. Peptides were injected in increasing concentration series (6-8 different concentrations) for 60-120 s at a flow rate of 30 µL/min in a buffer consisting of 10 mM Tris buffer at pH 7.4, with 150 mM NaCl and 0.05% Tween20. This was followed by a dissociation period for up to 10 min. The C3 surface was regenerated between runs by a 45 s injection of 3 M MgCl₂.

Sensorgrams were double-referenced (reference surface, blanks) prior to analysis of the kinetic profiles by globally fitting data to a 1:1 Langmuir binding model to obtain association and dissociation rates for calculation of the equilibrium dissociation constant K_d . Each peptide was tested at in a least 3 independent experiments.

5

Table 7: Compstatin analogues binding affinities for C3 as determined by a surface plasmon resonance assay with immobilized C3.

Comp. no.	K_d [nM]	N
2	16	3
4	1.5	3
15	14	3
20	37	3
21	16	3
23	2.9	3
24	28	5
28	44	3
29	21	3
43	3.3	3
48	0.12	3
49	3.2	3
50	13	3
53	1.4	3
54	3.0	3
61	0.33	3
63	4.3	3
67	0.69	3
73	0.30	3
75	1.5	3
81	10	3
82	5.4	3
86	1.3	3
87	2.6	3

10

Comp. no.	Kd [nM]	N
		3
102	1.7	3
104	33.9	3
106	5.4	3
107	6	3
111	8.1	3
117	23.7	3
118	11.5	3
119	9.8	3
120	27.6	3
121	29.6	3
122	62.9	3
123	11.1	3
124	30.7	3
125	71	3
126	5.2	3
127	8.5	3
128	6.5	3
130	4.4	3
139	7.4	3
140	7.6	3
141	6.6	3
142	4.8	3

5 The following pairs of compounds, which differ only at position 3, show the effects of replacing valine by isoleucine in different peptide backbones.

Table 3: Binding affinity of compstatin analogues to immobilized C3 determined by a surface plasmon resonance (SPR) assay.

Comp no	SPR Kd (nM)		1	2	3	4	5	6	7	8	9	10	11	12	13	
2	16	Ac	I	C(1)	I	W	Q	D	W	G	E	H	R	C(1)	T	NH ₂
A	130				V											
15	14	Ac	I	C(1)	I	W	Q	D	W	G	E	H	R	C(1)	T	GAES NH ₂
D	230				V											
21	16	Ac	I	C(1)	I	W	Q	D	W	G	E	H	R	C(1)	S	NH ₂
G	160				V											
48	0.12	H dTyr	I	C(1)	I	1MeTrp	Q	D	W	Sar	A	H	R	C(1)	NMelle	NH ₂
Cp40	0.31				V											

5

Example 5: Profiling of test compounds in Non-Human Primates (NHP)

Healthy male Cynomolgus monkeys (*Macaca fascicularis*) received single subcutaneous administrations of each test substance. Compounds were formulated in 20 mM phosphate adjusted with NaOH to pH 7.5 and mannitol for isotonicity and dosed at 1850 nmol/kg. Blood was collected from a femoral vein from each animal at the following times: Pre-dose, 1, 2, 4, 8, 24, 48, 72, 96 and 120 h (10 sampling times). Blood was collected into serum separation tubes and allowed to clot at room temperature. The tubes were centrifuged and resulting serum was aliquoted and snap-frozen over dry-ice and stored at nominally -80°C until analysis. All NHP studies were performed in accordance with animal welfare laws and regulations, including approval of the study by a local ethical review process.

Serum isolated from non-human primates at specific time points after dosing were analyzed for alternative pathway complement activity using the Complement system Alternative Pathway WIESLAB® kit from Svar Life Science (previously Euro diagnostic AB, Sweden) following the manufacturer's protocol. Briefly, serum samples or controls were diluted in buffer and incubated in microtitre strips coated with specific activators of the alternative pathway. The wells were washed and formed C5b-9 was detected using included colorimetric reagents. Absorbance at 405 nm was measured. The percent activity of the alternative complement pathway was calculated for each animal and timepoint relative to the predose activity (0 hours) of the individual animal with subtraction of the negative control. This provides the pharmacological activity of the compounds. Results are shown in figure 1a-d.

In Fig 1a, the non-acylated compound 61 had a relatively short duration of action despite high affinity for C3. By contrast, the acylated compounds in Fig 1b, 1c and 1d in general possessed a longer-lasting pharmacological activity in vivo when compared to the non-acylated compounds in Fig 1a despite lower affinity. Although acylation of peptides is
5 generally known to increase the in vivo half-life, it was surprisingly found that the in vivo duration of the pharmacological efficacy was prolonged to this extent.

Claims:

1. A compstatin analogue represented by the formula:

5

Y1-R1-X1-C-I-X4-Q-X6-W-X8-X9-H-X11-C-X13-R2-Y2,

wherein:

10 Y1 is hydrogen, acetyl or a lipophilic group Φ ;

X1 is I, Y, F or Sar;

X4 is W, F, V, Y, 1MeTrp, D-Trp, N-Me-Trp, 1-For-Trp, 1-Nal, 2-Nal, 5MeTrp, Bpa or 2Igl;

15

X6 is E, K or D;

X8 is G or Sar;

20 X9 is H, A, E, D, K, R or S;

X11 is R, S or K;

X13 is T, S, E, F, H, K, Sar, G, I, D, N-Me-Ile or N-Me-Thr;

25

Y2 is NH₂, OH or a lipophilic group Φ ;

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or 8-

30

Amino-3,6-dioxaoctanoyl, Peg4, or 8-aminooctanoyl, or derivatives thereof; and

R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or 8-Amino-3,6-dioxaoctanoyl, Peg4, or 8-aminooctanoyl, or derivatives thereof;

35

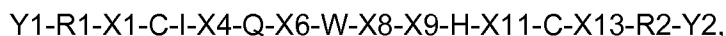
wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

and wherein the compstatin analogue optionally has a lipophilic group Φ covalently linked to the side chain of one or more amino acid residues;

or a pharmaceutically acceptable salt and/or solvate thereof.

5

2. The present invention further provides a compstatin analogue represented by the formula:



10

wherein:

Y1 is hydrogen, acetyl, or a lipophilic group Φ ;

15 X1 is I, Y, F or Sar;

X4 is W, V, Y, 2-Nal, 1-Nal or 1MeTrp;

X6 is E or D;

20

X8 is G or Sar;

X9 is A, E, D, K or S;

25 X11 is R, S or K;

X13 is T, S, E, I, Sar, K, G or N-Me-Ile;

Y2 is NH_2 , OH or a lipophilic group Φ ;

30

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof, or Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof; and

35 R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg3 or Peg4, or 8-aminooctanoyl, or derivatives thereof;

wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

- 5 and wherein the compstatin analogue optionally has a lipophilic group Φ covalently linked to the side chain of one or more amino acids;

or a pharmaceutically acceptable salt and/or solvate thereof.

- 10 3. The present invention further provides a compstatin analogue represented by the formula:



- 15 wherein:

Y1 is hydrogen, acetyl or a lipophilic group Φ ;

X1 is I, Y, F or Sar;

20

X4 is W, V, Y, 1-Nal, 2-Nal or 1MeTrp;

X6 is E or D;

- 25 X9 is A, E, D, K or S;

X11 is R, S or K;

X13 is T, I, S, E, K or Sar;

30

Y2 is NH_2 , OH or a lipophilic group Φ ;

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof, or

- 35 Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof; and

R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg3 or Peg4, or 8-aminooctanoyl, or derivatives thereof;

- 5 wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

and wherein the compstatin analogue optionally has a lipophilic group Φ covalently linked to the side chain of one or more amino acids;

10

or a pharmaceutically acceptable salt and/or solvate thereof.

4. The compstatin analogue may be represented by the formula:

15 Y1-R1-X1-C-I-X4-Q-X6-W-G-X9-H-R-C-X13-R2-Y2,

wherein:

Y1 is hydrogen, acetyl or a lipophilic group Φ ;

20 X1 is I, Y, F or Sar;

X4 is W, V, Y, 1-Nal, 2-Nal or 1MeTrp;

X6 is E or D;

25

X9 is A, E, D, K or S;

X13 is T, S, E or Sar;

30 Y2 is NH_2 , OH or a lipophilic group Φ ;

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof, or Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof; and

35

R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg3 or Peg4, or 8-aminooctanoyl, or derivatives thereof;

- 5 wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

and wherein the compstatin analogue optionally has a lipophilic group Φ covalently linked to the side chain of one or more amino acids;

10

or a pharmaceutically acceptable salt and/or solvate thereof.

5. A compstatin analogue according to any one of claims 1 to 4 comprising at least one lipophilic group Φ .

15

6. A compstatin analogue according to claim 5 wherein Y1 or Y2 is a lipophilic group Φ .

7. A compstatin analogue according to claim 5 or claim 6 comprising a lipophilic group Φ linked to the side chain of an amino acid residue at position X1, X11 or X13, or an amino acid residue in R1 or R2.

20

8. A compstatin analogue according to claim 7 wherein said amino acid residue is a lysine residue.

- 25 9. A compstatin analogue according to any one of claims 1 to 4 which does not comprise a lipophilic group Φ .

10. A compstatin analogue according to claim 1, represented by the formula:

30

Y1-R1-X1-C-I-X4-Q-D-W-G-X9-H-R-C-X13-R2-Y2,

wherein:

Y1 is hydrogen or acetyl;

35

X1 is Y or F ;

X4 is W, Y, 1MeTrp;

X9 is A, E or K;

5 X13 is T, E or Sar;

Y2 is NH₂ or OH;

10 R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γGlu, βAsp, or βAla, or a corresponding D form thereof, or Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof; and

15 R2 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γGlu, βAsp, or βAla, or a corresponding D form thereof; or Peg3 or Peg4, or 8-aminooctanoyl, or derivatives thereof;

wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

20 or a pharmaceutically acceptable salt and/or solvate thereof.

11. A compstatin analogue according to claim 10, represented by the formula:

Y1-R1-X1-C-I-1MeTrp-Q-D-W-G-E-H-R-C-X13-R2-Y2,

25

wherein:

Y1 is hydrogen or acetyl;

X1 is Y or F;

30

X13 is T, E or Sar;

Y2 is NH₂ or OH;

35 R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γGlu, βAsp, or βAla, or a corresponding D form thereof, or Peg3, Peg4, or 8-aminooctanoyl, or derivatives thereof; and

R2 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg3 or Peg4, or 8-aminooctanoyl, or derivatives thereof;

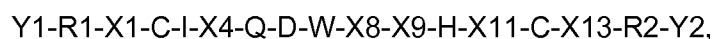
5

wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

or a pharmaceutically acceptable salt and/or solvate thereof.

10

12. A compstatin analogue according to claim 1, represented by the formula:



15 wherein:

Y1 is hydrogen, acetyl or a lipophilic group Φ ;

X1 is I, Y, F or Sar;

20

X4 is W, V, Y, 2-Nal, 1-Nal or 1MeTrp;

X8 is G or Sar;

25 X9 is A, E, D, K or S;

X11 is R, S or K*;

X13 is T, S, E, I, Sar, K, G or N-Me-Ile;

30

Y2 is NH₂, OH or a lipophilic group Φ ;

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, K*, F, P, S, T, W, Y, R, V or Sar, or a corresponding D form thereof;

35

R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, K*, F, P, S, T, W, Y, R, V Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or

Peg 3 or Peg4, or 8-aminooctanoyl, or derivatives thereof; wherein * indicates that the amino acid residue bears a lipophilic group Φ covalently linked to its side chain;

wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

and wherein the compstatin analogue comprises at least one lipophilic group Φ ;

or a pharmaceutically acceptable salt and/or solvate thereof.

13. A compstatin analogue according to claim 11, represented by the formula:



wherein:

Y1 is hydrogen, acetyl, or a lipophilic group Φ ;

X1 is I, Y, F or Sar;

X4 is W, V, Y, 1-Nal, 2-Nal or 1MeTrp;

X9 is A, E, D, K or S;

X11 is R, S or K*;

X13 is T, I, S, E, K or Sar;

Y2 is NH₂, OH or a lipophilic group Φ ;

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, K*, F, P, S, T, W, Y, R, V or Sar, or a corresponding D form thereof; and

R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, K*, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or Peg 3 or Peg4, or 8-aminooctanoyl, or derivatives thereof; wherein * indicates that the amino acid residue bears a lipophilic group Φ covalently attached to its side chain;

wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

5 and wherein the compstatin analogue comprises at least one lipophilic group Φ ;

or a pharmaceutically acceptable salt and/or solvate thereof.

14. A compstatin analogue according to claim 12 or claim 13, represented by the formula:

10



wherein:

Y1 is hydrogen, acetyl or a lipophilic group Φ ;

15

X1 is I, Y, F or Sar;

X4 is W, V, 1-Nal, 2-Nal or 1MeTrp;

20 X9 is A, E, D, K or S;

X13 is T, S, E or Sar;

Y2 is NH_2 , OH or a lipophilic group Φ ;

25

R1 is absent or is a sequence of 1 to 6 amino acid residues selected from A, E, G, L, K, K*, F, P, S, T, W, Y, R, V or Sar, or a corresponding D form thereof;

R2 is absent or is a sequence of 1 to 8 amino acid residues selected from A, E, G, L, K, K*, F, P, S, T, W, Y, R, V, Sar, isoLys, γ Glu, β Asp, or β Ala, or a corresponding D form thereof; or

30 Peg3 or Peg4, or 8-aminooctanoyl, or derivatives thereof; wherein * indicates that the amino acid residue bears a lipophilic group Φ covalently attached to its amino acid side chain;

wherein the compstatin analogue has a disulphide bond between the cysteine residues at positions 2 and 12;

35

and wherein the compstatin analogue comprises at least one lipophilic group Φ , e.g. exactly one lipophilic group Φ ;

or a pharmaceutically acceptable salt and/or solvate thereof.

15. A compstatin analogue according to any one of the preceding claims wherein the 13-mer peptide portion (X1-X13) of the compstatin analogue has a sequence selected from:

[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)[Sar];
 [Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)T;
 [Sar]C(1)I[1-Me-Trp]QEW[Sar]EHRC(1)T;
 10 [Sar]C(1)I[1-Me-Trp]QEWGEHRC(1)[Sar];
 [Sar]C(1)IWQDWGEHRC(1)T;
 FC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)[Sar];
 FC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)T;
 FC(1)I[1-Me-Trp]QDWGEHKC(1)[Sar];
 15 FC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar];
 FC(1)I[1-Me-Trp]QDWGEHRC(1)E;
 FC(1)I[1-Me-Trp]QDWGEHRC(1)S;
 FC(1)I[1-Me-Trp]QDWGEHRC(1)T;
 FC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar];
 20 FC(1)I[1Na]QDWGEHRC(1)T;
 FC(1)I[2Na]QDWGEHRC(1)T;
 FC(1)IWQDWGEHRC(1)[Sar];
 FC(1)IWQDWGEHRC(1)T;
 IC(1)I[1-Me-Trp]QDW[Sar]AHRC(1)[N-Me-Ile];
 25 IC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar];
 IC(1)I[1-Me-Trp]QDWGEHRC(1)T;
 IC(1)I[2Na]QDWGEHRC(1)[Sar];
 IC(1)IWQDWGAHRC(1)E;
 IC(1)IWQDWGAHRC(1)T;
 30 IC(1)IWQDWGAHSC(1)T;
 IC(1)IWQDWGDHRC(1)T;
 IC(1)IWQDWGEHRC(1)[Sar];
 IC(1)IWQDWGEHRC(1)E;
 IC(1)IWQDWGEHRC(1)S;
 35 IC(1)IWQDWGEHRC(1)T;
 IC(1)IWQDWGEHSC(1)T;
 IC(1)IWQDWGKHRC(1)T;

IC(1)IWQDWGRHRC(1)T;

IC(1)IWQDWGSHRC(1)T;

IC(1)IWQEWGEHRC(1)T;

IC(1)IWQKWGAHRC(1)T;

5 IC(1)IWQKWGEHRC(1)T;

YC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar];

YC(1)I[1-Me-Trp]QDWGEHRC(1)T;

YC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar];

YC(1)I[2NaI]QDWGEHRC(1)T;

10 YC(1)IWQDWGEHRC(1)T;

YC(1)I[1-Me-Trp]QDWGEH[K*]C(1)[Sar]; and

YC(1)I[1-Me-Trp]QEW[Sar]EHRC(1)[Sar].

16. A compstatin analogue according to any one of the preceding claims wherein R1 has
15 a sequence selected from:

{d}Y, EGSE, AGSE, SASE, EYSE, GSE, ASE, ESSA, KGSA, AKGE, ASGE, ASSE, ASES, GSAE, ESSE, ESGA, SEG, GES, ESS, EGSA, ESE, EGE, ESA, SAE, SGA, YLEA, GSA, KEK, EKG, ES, AE, TE, KE, GE, FE, YE, AS, SE, RS, SR, SA, GE, Y, S and E.

20 17. A compstatin analogue according to claim 16 comprising a lipophilic group Φ covalently linked to an amino acid side chain of R1.

18. A compstatin analogue according to claim 17 wherein R1 has the sequence K*GSA.

25 19. A compstatin analogue according to any one of the preceding claims wherein R2 has a sequence selected from:

EGASGSG, EGAGSG, EGASAG, EGAGAG, EGESGSG, EGE GSG, EGESAG, EGE GAG, EK[γ Glu]AK, EGE-Peg3-Peg3-K, EGE G G, EGAGG, EGE S S, GAESK, EGAK, EGEK, EGG, EGK, EGKK, EGS, EK, EGA, EK[γ Glu], EK[γ Glu]K, EGE-Peg3, EGE-Peg3-K, EGE-Peg3-Peg3, EGE-Peg3-Peg3-K, EGE-Peg3-Peg3-Peg3, GESESE, GAESSES, EGESES, EGESESK, EGE-Peg3-ES, EGE-Peg3-ESK, GESESE, EGE-[8-aminooctanoyl], EGE-[8-aminooctanoyl]-K, EGE-[8-aminooctanoyl]-EK, EGE G G G, EGE G G G K, EK[γ Glu]G G G, EK[γ Glu]G G G K, EGE-[8-aminooctanoyl]-E, GAES, EYGS, EGYA, EAGS, EAKS, EKSA, ESGA, EGGS, EGGA, ESSG, ESAG, GEES, AEES, ESEG, AEGS, ESGS, SEGA, SEG, EGK, EGA, ESG, EAG, GAE, EGEA, EGE, EA, E, S, GE, GEK, EG, EKE and EKP.

30
35

20. A compstatin analogue according to claim 19 comprising a lipophilic group Φ covalently linked to an amino acid side chain of R2.

21. A compstatin analogue according to claim 20 wherein R2 has the sequence

5 EK[γ Glu]AK*, EGKK, EK[γ Glu]K*, EGE-Peg3-K*, EGE-Peg3-Peg3-K*, EGESESK*, EGE-Peg3-ESK*, EGE-[8-aminooctanoyl]-K*, EGE-[8-aminooctanoyl]-EK*, EGE³GGK*, EK[γ Glu]GGGK*, EGE-Peg3-Peg3-K*, GAESK*, EGAK*, EGEK*, EGK* EGE-Peg3-ESK*, GESESEK*, GEK* or EK.

10 22. A compstatin analogue according to claim 1, comprising a sequence selected from:

IC(1)IWQDWGAHRC(1)T

IC(1)IWQDWGEHRC(1)T

ESSAIC(1)IWQDWGEHRC(1)T

IC(1)I[1MeTrp]QDWGEHRC(1)T

15 IC(1)IWQDWGKHRC(1)T

IC(1)IWQDWGSHRC(1)T

IC(1)IWQKWGEHRC(1)T

IC(1)IWQKWGAHRC(1)TGAES

YC(1)IWQDWGEHRC(1)T

20 ESSAYC(1)IWQDWGEHRC(1)T

[Sar]C(1)IWQDWGEHRC(1)T

IC(1)IWQDWGAHRC(1)E

IC(1)IWQDWGEHRC(1)[Sar]

ESSAIC(1)IWQDWGEHRC(1)TGAES

25 IC(1)IWQDWGEHRC(1)TGAES

IC(1)IWQEWGEHRC(1)T

IC(1)IWQDWGDHRC(1)T

IC(1)IWQDWGRHRC(1)T

IC(1)IWQDWGAHSC(1)T

30 IC(1)IWQDWGEHSC(1)T

IC(1)IWQDWGEHRC(1)S

IC(1)IWQDWGEHRC(1)E

FC(1)IWQDWGEHRC(1)T

IC(1)IWQDWGEHRC(1)TEGE

35 IC(1)IWQDWGEHRC(1)TEA

IC(1)IWQDWGEHRC(1)TE

IC(1)IWQDWGEHRC(1)EGE

EGSAIC(1)IWQDWGEHRC(1)[Sar]E
 EGSAIC(1)IWQDWGEHRC(1)T
 EGEIC(1)IWQDWGEHRC(1)T
 ESEIC(1)IWQDWGEHRC(1)T
 5 SEIC(1)IWQDWGEHRC(1)TEA
 EIC(1)IWQDWGEHRC(1)TE
 EIC(1)IWQDWGEHRC(1)TEGE
 EGEIC(1)IWQDWGEHRC(1)EGE
 ESEIC(1)IWQDWGEHRC(1)EGE
 10 KEKIC(1)IWQDWGEHRC(1)TEKE
 EKGIC(1)IWQDWGEHRC(1)TEKP
 IC(1)IWQDWGEHRC(1)TEGK
 GSAIC(1)IWQDWGEHRC(1)[Sar]E
 SAIC(1)IWQDWGEHRC(1)[Sar]E
 15 SAIC(1)IWQDWGEHRC(1)TEG
 FC(1)IWQDWGEHRC(1)TGAE
 EGSAIC(1)IWQDWGEHRC(1)[Sar]EGE
 EGSAFC(1)IWQDWGEHRC(1)[Sar]E
 ESSAIC(1)IWQDWGAHRC(1)T
 20 IC(1)IWQDWGAHRC(1)TGAES
 {d}YIC(1)I[1-Me-Trp]QDW[Sar]AHRC(1)-[N-Me-Ile]
 EGSAIC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E
 EGSAIC(1)I-2NaI-QDWGEHRC(1)[Sar]E
 IC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES
 25 IC(1)I[2NaI]QDWGEHRC(1)TGAES
 EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E
 EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E
 EGSAIC(1)IWQDWGEHRC(1)TE
 EGSAFC(1)I[1NaI]QDWGEHRC(1)TE
 30 EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)TE
 EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)EGE
 EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)TE
 EGSAFC(1)I[2NaI]QDWGEHRC(1)TE
 FC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES
 35 YC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES
 FC(1)I[1NaI]QDWGEHRC(1)TGAES
 FC(1)I[2NaI]QDWGEHRC(1)TGAES

YC(1)I[2Na]QDWGEHRC(1)TGAES
 YC(1)IWQDWGEHRC(1)TGAES
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES
 YC(1)I[1-Me-Trp]QDWGEHRC(1)TEAGS
 5 YC(1)I[1-Me-Trp]QDWGEHRC(1)TESGA
 EGSAYC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]E
 SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA
 FC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TGAES
 {d}YFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TGAES
 10 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]GAES
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA
 SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)[Sar]EA
 SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TEA
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E
 15 SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)[Sar]E
 EFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA
 SE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA
 SE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA
 SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EA
 20 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)SEA
 EFC(1)I[1-Me-Trp]QDWGEHRC(1)ES
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA
 GEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA
 GE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA
 25 SE[Sar]C(1)I[1-Me-Trp]QEW[Sar]EHRC(1)TEA
 SE[Sar]C(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EA
 {d}Y[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA

23. A compstatin analogue according to claim 1 which is:

30 Ac-IC(1)IWQDWGAHRC(1)T-NH₂ (Compound 1)
 Ac-IC(1)IWQDWGEHRC(1)T-NH₂ (Compound 2)
 Ac-ESSAIC(1)IWQDWGEHRC(1)T-NH₂ (Compound 3)
 Ac-IC(1)I[1-Me-Trp]QDWGEHRC(1)T-NH₂ (Compound 4)
 Ac-IC(1)IWQDWGKHRC(1)T-NH₂ (Compound 5)
 35 Ac-IC(1)IWQDWGSHRC(1)T-NH₂ (Compound 6)
 Ac-IC(1)IWQKWGEHRC(1)T-NH₂ (Compound 7)
 Ac-IC(1)IWQKWGAHRC(1)TGAES-NH₂ (Compound 8)

- Ac-YC(1)IWQDWGEHRC(1)T-NH2 (Compound 9)
 Ac-ESSAYC(1)IWQDWGEHRC(1)T-NH2 (Compound 10)
 Ac-[Sar]C(1)IWQDWGEHRC(1)T-NH2 (Compound 11)
 Ac-IC(1)IWQDWGAHRC(1)E-NH2 (Compound 12)
 5 Ac-IC(1)IWQDWGEHRC(1)[Sar]-NH2 (Compound 13)
 Ac-ESSAIC(1)IWQDWGEHRC(1)TGAES-NH2 (Compound 14)
 Ac-IC(1)IWQDWGEHRC(1)TGAES-NH2 (Compound 15)
 Ac-IC(1)IWQEWGEHRC(1)T-NH2 (Compound 16)
 Ac-IC(1)IWQDWGDHRC(1)T-NH2 (Compound 17)
 10 Ac-IC(1)IWQDWGRHRC(1)T-NH2 (Compound 18)
 Ac-IC(1)IWQDWGAHSC(1)T-NH2 (Compound 19)
 Ac-IC(1)IWQDWGEHSC(1)T-NH2 (Compound 20)
 Ac-IC(1)IWQDWGEHRC(1)S-NH2 (Compound 21)
 Ac-IC(1)IWQDWGEHRC(1)E-NH2 (Compound 22)
 15 Ac-FC(1)IWQDWGEHRC(1)T-NH2 (Compound 23)
 Ac-IC(1)IWQDWGEHRC(1)TEGE-NH2 (Compound 24)
 Ac-IC(1)IWQDWGEHRC(1)TEA-NH2 (Compound 25)
 Ac-IC(1)IWQDWGEHRC(1)TE-NH2 (Compound 26)
 Ac-IC(1)IWQDWGEHRC(1)EGE-NH2 (Compound 27)
 20 Ac-EGSAIC(1)IWQDWGEHRC(1)[Sar]E-NH2 (Compound 28)
 Ac-EGSAIC(1)IWQDWGEHRC(1)T-NH2 (Compound 29)
 Ac-EGEIC(1)IWQDWGEHRC(1)T-NH2 (Compound 30)
 Ac-ESEIC(1)IWQDWGEHRC(1)T-NH2 (Compound 31)
 Ac-SEIC(1)IWQDWGEHRC(1)TEA-NH2 (Compound 32)
 25 Ac-EIC(1)IWQDWGEHRC(1)TE-NH2 (Compound 33)
 Ac-EIC(1)IWQDWGEHRC(1)TEGE-NH2 (Compound 34)
 Ac-EGEIC(1)IWQDWGEHRC(1)EGE-NH2 (Compound 35)
 Ac-ESEIC(1)IWQDWGEHRC(1)EGE-NH2 (Compound 36)
 Ac-KEKIC(1)IWQDWGEHRC(1)TEKE-NH2 (Compound 37)
 30 Ac-EKGIC(1)IWQDWGEHRC(1)TEKP-NH2 (Compound 38)
 Ac-IC(1)IWQDWGEHRC(1)TEGK-NH2 (Compound 39)
 Ac-GSAIC(1)IWQDWGEHRC(1)[Sar]E-NH2 (Compound 40)
 Ac-SAIC(1)IWQDWGEHRC(1)[Sar]E-NH2 (Compound 41)
 Ac-SAIC(1)IWQDWGEHRC(1)TEG-NH2 (Compound 42)
 35 Ac-FC(1)IWQDWGEHRC(1)TGAE-NH2 (Compound 43)
 Ac-EGSAIC(1)IWQDWGEHRC(1)[Sar]EGE-NH2 (Compound 44)
 Ac-EGSAFC(1)IWQDWGEHRC(1)[Sar]E-NH2 (Compound 45)

- Ac-ESSAIC(1)IWQDWGAHRC(1)T-NH₂ (Compound 46)
 Ac-IC(1)IWQDWGAHRC(1)TGAES-NH₂ (Compound 47)
 H-{d}YIC(1)I[1-Me-Trp]QDW[Sar]AHRC(1)[N-Me-Ile]-NH₂ (Compound 48)
 Ac-EGSAIC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 49)
 5 Ac-EGSAIC(1)I[2NaI]QDWGEHRC(1)[Sar]E-NH₂ (Compound 50)
 Ac-IC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-NH₂ (Compound 51)
 Ac-IC(1)I[2NaI]QDWGEHRC(1)TGAES-NH₂ (Compound 52)
 Ac-EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 53)
 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 54)
 10 Ac-EGSAIC(1)IWQDWGEHRC(1)TE-NH₂ (Compound 55)
 Ac-EGSAFC(1)I[1NaI]QDWGEHRC(1)TE-NH₂ (Compound 56)
 Ac-EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)TE-NH₂ (Compound 57)
 Ac-EGSAFC(1)I[1-Me-Trp]QDWGEHRC(1)EGE-NH₂ (Compound 58)
 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)TE-NH₂ (Compound 59)
 15 Ac-EGSAFC(1)I[2NaI]QDWGEHRC(1)TE-NH₂ (Compound 60)
 Ac-FC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-NH₂ (Compound 61)
 Ac-YC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-NH₂ (Compound 62)
 Ac-FC(1)I[1NaI]QDWGEHRC(1)TGAES-NH₂ (Compound 63)
 Ac-FC(1)I[2NaI]QDWGEHRC(1)TGAES-NH₂ (Compound 64)
 20 Ac-YC(1)I[2NaI]QDWGEHRC(1)TGAES-NH₂ (Compound 65)
 Ac-YC(1)IWQDWGEHRC(1)TGAES-NH₂ (Compound 66)
 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-NH₂ (Compound 67)
 Ac-YC(1)I[1-Me-Trp]QDWGEHRC(1)TEAGS-NH₂ (Compound 68)
 Ac-YC(1)I[1-Me-Trp]QDWGEHRC(1)TESGA-NH₂ (Compound 69)
 25 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 70)
 Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH₂ (Compound 71)
 Ac-FC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TGAES-NH₂ (Compound 72)
 H-{d}YFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TGAES-NH₂ (Compound 73)
 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]GAES-NH₂ (Compound 74)
 30 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH₂ (Compound 75)
 Ac-SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)[Sar]EA-NH₂ (Compound 76)
 Ac-SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)TEA-NH₂ (Compound 77)
 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 78)
 Ac-SEFC(1)I[1-Me-Trp]QDW[Sar]EHRC(1)[Sar]E-NH₂ (Compound 79)
 35 Ac-EFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH₂ (Compound 80)
 Ac-SE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH₂ (Compound 81)
 Ac-SE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA-NH₂ (Compound 82)

Ac-SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EA-NH₂ (Compound 83)
 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)SEA-NH₂ (Compound 84)
 Ac-EFC(1)I[1-Me-Trp]QDWGEHRC(1)ES-NH₂ (Compound 85)
 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH₂ (Compound 86)
 5 Ac-GEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EA-NH₂ (Compound 87)
 Ac-GE[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA-NH₂ (Compound 88)
 Ac-SE[Sar]C(1)I[1-Me-Trp]QEW[Sar]EHRC(1)TEA-NH₂ (Compound 89)
 Ac-SE[Sar]C(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EA-NH₂ (Compound 90)
 H-{d}Y[Sar]C(1)I[1-Me-Trp]QDWGEHRC(1)TEA-NH₂ (Compound 91)

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24. A compstatin analogue according to claim 1 comprising a sequence selected from:

[K*]GSAIC(1)IWQDWGEHRC(1)TEGE (Compound 100)
 ASGEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[K*] (Compound 113)
 15 EFC(1)I[1-Me-Trp]QDWGEHRC(1)EGE-[K*] (Compound 134)
 EGSAIC(1)IWQDWGEHRC(1)TEG[K*] (Compound 101)
 EGSAYC(1)I[1-Me-Trp]QDWGEH[K*]C(1)[Sar]E (Compound 103)
 EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EG[K*] (Compound 104)
 EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[K*] (Compound 109)
 20 EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGK[K*] (Compound 110)
 EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EK[Glu][K*] (Compound 111)
 FC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES[K*] (Compound 102)
 IC(1)IWQDWGEHRC(1)TEG[K*] (Compound 92)
 IC(1)IWQDWGEHRC(1)TEGE[K*] (Compound 94)
 25 SAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E[K*] (Compound 105)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGA[K*] (Compound 119)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-
 dioxaoctanoyl][K*] (Compound 123)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGEGGG-[K*] (Compound 129)
 30 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][K*] (Compound
 138)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl]ES-[K*]
 (Compound 140)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-
 35 dioxaoctanoyl]-[K*] (Compound 127)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGESES-[K*] (Compound 139)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EK[yGlu]GGG-[K*] (Compound 132)

- SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE-[8-aminooctanoyl]-[K*] (Compound 136)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE-[8-aminooctanoyl]-E-[K*] (Compound 137)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGEGGG-[K*] (Compound 130)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE[8-Amino-3,6-dioxaoctanoyl]ES-[K*] (Compound
 5 142)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-
 dioxaoctanoyl]-[K*] (Compound 126)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEK[γGlu]GGG-[K*] (Compound 133)
 SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-[K*] (Compound 135)
 10 SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGA[K*] (Compound 120)
 SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-
 dioxaoctanoyl][K*] (Compound 124)
 SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGA[K*] (Compound 112)
 SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-
 15 dioxaoctanoyl][K*] (Compound 117)
 SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[K*] (Compound 114)
 SEYC(1)I[1-Me-Trp]QEW[Sar]EHRC(1)[Sar]EK[γGlu]A[K*] (Compound 121)
 SEYC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGA[K*] (Compound 122)
 SEYC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-
 20 dioxaoctanoyl][K*] (Compound 125)
 EGSEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E (Compound 107)
 ESSAIC(1)IWQDWGEHRC(1)TEGE (Compound 99)
25. A compstatin analogue according to claim 24 comprising a sequence selected from:
- 25 Ac-[K*]GSAIC(1)IWQDWGEHRC(1)TEGE-NH₂ (Compound 100)
 Ac-ASGEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[K*]-[NH₂] (Compound 113)
 Ac-EFC(1)I[1-Me-Trp]QDWGEHRC(1)EGE[NH₂] (Compound 134)
 Ac-EGSAIC(1)IWQDWGEHRC(1)TEG[K*]-[NH₂] (Compound 101)
 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEH[K*]C(1)[Sar]E-[NH₂] (Compound 103)
 30 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EG[K*]-NH₂ (Compound 104)
 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[K*]-NH₂ (Compound 109)
 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGK[K*]-NH₂ (Compound 110)
 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EK[γGlu][K*]-NH₂ (Compound 111)
 Ac-FC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES[K*]-NH₂ (Compound 102)
 35 Ac-IC(1)IWQDWGEHRC(1)TEG[K*]-NH₂ (Compound 92,93, 95. 96, 98)
 Ac-IC(1)IWQDWGEHRC(1)TEGE[K*]-NH₂ (Compound 94, 97)
 Ac-SAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E[K*]-NH₂ (Compound 105, 106)

- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGA[K*]-NH2 (Compound 119)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl][K*]-NH2 (Compound 123)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGEGGG-[K*]-NH2 (Compound 129)
- 5 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl]-[K*]-NH2 (Compound 138)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl]ES-[K*]-NH2 (Compound 140)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-[K*]-NH2 (Compound 127, 128)
- 10 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGESES-[K*]-NH2 (Compound 139, 141)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EK[γGlu]GGG-[K*]-NH2 (Compound 132)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEGE-[8-aminoctanoyl]-[K*]-NH2 (Compound 136)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEGE-[8-aminoctanoyl]-E-[K*]-NH2 (Compound 137)
- 15 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEGEGGG-[K*]-NH2 (Compound 130, 131)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEGE[8-Amino-3,6-dioxaoctanoyl]ES-[K*]-NH2 (Compound 142)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-[K*]-NH2 (Compound 126)
- 20 Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TEK[γGlu]GGG-[K*]-NH2 (Compound 133)
- Ac-SEFC(1)|[1-Me-Trp]QDWGEHRC(1)TGAES-[K*]-NH2 (Compound 135)
- Ac-SEFC(1)|[1-Me-Trp]QEWGEHRC(1)[Sar]EGA[K*]-NH2 (Compound 120)
- Ac-SEFC(1)|[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl][K*]-NH2 (Compound 124)
- 25 Ac-SEYC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGA[K*]-NH2 (Compound 112, 118)
- Ac-SEYC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl][K*]-NH2 (Compound 117)
- Ac-SEYC(1)|[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[K*]-NH2 (Compound 114, 115, 116)
- 30 Ac-SEYC(1)|[1-Me-Trp]QEW[Sal]EHRC(1)[Sal]EK[γGlu]A[K*]-NH2 (Compound 121)
- Ac-SEYC(1)|[1-Me-Trp]QEWGEHRC(1)[Sal]EGA[K*]-NH2 (Compound 122)
- Ac-SEYC(1)|[1-Me-Trp]QEWGEHRC(1)[Sal]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl][K*]-NH2 (Compound 125)
- Φ-EGSEYC(1)|[1-Me-Trp]QDWGEHRC(1)[Sal]E-NH2 (Compound 107, 108)
- 35 Φ-ESSAIC(1)|IWQDWGEHRC(1)TEGE-NH2 (Compound 99)

26. A compstatin analogue according to any one of claims 1 to 8, 12 to 22, 24 or 25 which

comprises a lipophilic group Φ , and wherein the lipophilic group Φ is Z^1 - or Z^1 - Z^2 -; wherein

Z^1 is $A-C_{12-22}alkylene-(CO)-$;

where A is H or $-COOH$, and wherein the alkylene may be linear or branched and may be saturated or unsaturated, and may optionally incorporate a phenylene or piperazinylene moiety in its length; and

Z^2 is a sequence of 1 to 6 residues of compounds selected from γ -Glu, α -Glu, E, K, Orn, S, T, A, β -Ala, G, P, V, L, I, Y, Q, N, Dapa, Gaba, or Aib, or a corresponding D form thereof, 5-aminopentanoyl, 6-aminohexanoyl, 7-aminoheptanoyl, 8-aminooctanoyl, 9-aminononanoyl, and 10-aminodecanoyl. 8-amino-3,6-dioxaoctanoic acid (8-Amino-3,6-dioxaoctanoyl), 11-amino-3,6,9-trioxaundecanoic acid (Peg4) and (piperazine-1-yl)-carboxylic acid.

27. A compstatin analogue according to claim 26 wherein Z^1 is selected from:

Tridecanoyl i.e. $H-(CH_2)_{12}-(CO)-$;

13-carboxytridecanoyl, i.e. $HOOC-(CH_2)_{12}-(CO)-$;

15-carboxypentadecanoyl, i.e. $HOOC-(CH_2)_{14}-(CO)-$;

17-carboxyheptadecanoyl, i.e. $HOOC-(CH_2)_{16}-(CO)-$;

19-carboxynonadecanoyl, i.e. $HOOC-(CH_2)_{18}-(CO)-$; or

21-carboxyheneicosanoyl, i.e. $HOOC-(CH_2)_{20}-(CO)-$.

28. A compstatin analogue according to claim 26 or claim 27 wherein Z^2 is selected from:
 γ Glu,

γ Glu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl-;

[(Piperazine-1-yl)-acetyl]-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl-;

(γ Glu)-G-(γ Glu);

(γ Glu)-K-(γ Glu);

(γ Glu)-K-G-(γ Glu); and

(γ Glu)-G-(8-Amino-3,6-dioxaoctanoyl)-(γ Glu)-(8-Amino-3,6-dioxaoctanoyl).

29. A compstatin analogue according to any one of claims 26 to 28 wherein Z^1 - or Z^1 - Z^2 - is selected from:

15-carboxy-pentadecanoyl;

15-carboxy-pentadecanoyl- γ Glu-,

15-carboxy-pentadecanoyl- γ Glu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl-;

17-carboxy-heptadecanoyl- γ Glu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl-;

19-carboxy-nonadecanoyl- γ Glu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl-;

- 15-carboxy-pentadecanoyl-[(Piperazine-1-yl)-acetyl]-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl);
- 17-carboxy-heptadecanoyl-(γGlu)-G-(γGlu);
- 17-carboxy-heptadecanoyl-(γGlu)-K-(γGlu);
- 5 17-carboxy-heptadecanoyl-(γGlu)-K-G-(γGlu);
- 17-carboxy-heptadecanoyl-(γGlu)-G-(8-Amino-3,6-dioxaoctanoyl)-(γGlu)-(8-Amino-3,6-dioxaoctanoyl);
- 15-carboxy-hexadecanoyl-γGlu-G-γGlu;
- 15-carboxy-pentadecanoyl;
- 10 15-carboxy-pentadecanoyl-γGlu-G-γGlu;
- 17-carboxy-heptadecanoyl;
- 17-carboxy-heptadecanoyl-γGlu]
- 19-carboxy-nonadecanoyl-γGlu-G-γGlu;
- 15-carboxy-pentadecanoyl-γGlu; and
- 15 17-carboxy-heptadecanoyl-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl.
30. A compstatin analogue according to claim 1 which is:
- Ac-IC(1)IWQDWGEHRC(1)TEG-K([15-carboxy-pentadecanoyl]-γGlu)-NH₂ (Compound 92)
- Ac-IC(1)IWQDWGEHRC(1)TEG-K([15-carboxy-pentadecanoyl]-γGlu-8-Amino-3,6-
- 20 dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 93)
- Ac-IC(1)IWQDWGEHRC(1)TEGE-K([15-carboxy-pentadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 94)
- Ac-IC(1)IWQDWGEHRC(1)TEG-K((15-carboxy-pentadecanoyl)-[(Piperazine-1-yl)-acetyl]-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 95)
- 25 Ac-IC(1)IWQDWGEHRC(1)TEG-K([17-carboxy-heptadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 96)
- Ac-IC(1)IWQDWGEHRC(1)TEGE-K([17-carboxy-heptadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 97)
- Ac-IC(1)IWQDWGEHRC(1)TEG-K([19-carboxy-nonadecanoyl]-γGlu-8-Amino-3,6-
- 30 dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 98)
- [15-Carboxy-pentadecanoyl]-ESSAIC(1)IWQDWGEHRC(1)TEGE-NH₂ (Compound 99)
- Ac-[K([15-carboxy-pentadecanoyl]-isoGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)]GSAIC(1)IWQDWGEHRC(1)TEGE-NH₂ (Compound 100)
- Ac-EGSAIC(1)IWQDWGEHRC(1)TEG-K([15-carboxy-pentadecanoyl]-γGlu)-NH₂ (Compound
- 35 101)
- Ac-FC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-K([15-carboxy-pentadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 102)

- Ac-EGSAYC(1)I[1-Me-Trp]QDWGEH-K([15-carboxy-pentadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-C(1)[Sar]E-NH₂ (Compound 103)
- Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EG-K([15-carboxy-pentadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 104)
- 5 Ac-SAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-K([17-carboxy-heptadecanoyl]-γGlu-KG-γGlu)-NH₂ (Compound 105)
- Ac-SAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 106)
- [15-Carboxy-pentadecanoyl]-EGSEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 107)
- 10 [17-Carboxy-heptadecanoyl]-EGSEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]E-NH₂ (Compound 108)
- Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 109)
- 15 Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGK-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 110)
- Ac-EGSAYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EK[γGlu]-K([17-carboxy-heptadecanoyl]-γGlu-8-Amino-3,6-dioxaoctanoyl-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 111)
- Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 112)
- 20 Ac-ASGEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-K([17-carboxy-heptadecanoyl]-γGI)-G-γGlu)-NH₂ (Compound 113)
- Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-K([17-carboxy-heptadecanoyl](γGI)-G-γGlu)-NH₂ (Compound 114)
- 25 Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGK-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 115)
- Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE-K([17-carboxy-heptadecanoyl]-γGlu-K-γGlu)-NH₂ (Compound 116)
- Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 117)
- 30 Ac-SEYC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-8-Amino-3,6-dioxaoctanoyl-γGlu-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 118)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-8-Amino-3,6-dioxaoctanoyl-γGlu-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 119)
- 35 Ac-SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-8-Amino-3,6-dioxaoctanoyl-γGlu-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 120)

- Ac-SEYC(1)I[1-Me-Trp]QEW[Sar]EHRC(1)[Sar]EK[ΓGlu]A-K([17-carboxy-heptadecanoyl]-
γGlu-G-8-Amino-3,6-dioxaoctanoyl-γGlu-(8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 121)
- Ac-SEYC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGA-K([17-carboxy-heptadecanoyl]-γGlu-G-8-
Amino-3,6-dioxaoctanoyl-γGlu-8-Amino-3,6-dioxaoctanoyl)-NH₂ (Compound 122)
- 5 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-
dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 123)
- Ac-SEFC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-
dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 124)
- Ac-SEYC(1)I[1-Me-Trp]QEWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-
10 dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 125)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-
dioxaoctanoyl]-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 126)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-
dioxaoctanoyl]-K([15-carboxy-pentadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 127)
- 15 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl][8-Amino-3,6-
dioxaoctanoyl]-K([19-carboxy-nonadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 128)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGEGGG-K([17-carboxy-heptadecanoyl]-γGlu-
G-γGlu)-NH₂ (Compound 129)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGEGGG-K([17-carboxy-heptadecanoyl]-γGlu-G-
20 γGlu)-NH₂ (Compound 130)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGEGGG-K([15-carboxy-pentadecanoyl]-γGlu-G-
γGlu)-NH₂ (Compound 131)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EK[γGlu]GGG-K([17-carboxy-heptadecanoyl]-
γGlu-G-γGlu)-NH₂ (Compound 132)
- 25 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEK[γGlu]GGG-K([17-carboxy-heptadecanoyl]-γGlu-
G-γGlu)-NH₂ (Compound 133)
- Ac-EFC(1)I[1-Me-Trp]QDWGEHRC(1)EGE-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂
(Compound 134)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TGAES-K([15-carboxy-hexadecanoyl]-γGlu-G-γGlu)-
30 NH₂ (Compound 135)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE[8-aminoctanoyl]-K([17-carboxy-
heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 136)
- Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE[8-aminoctanoyl]E-K([17-carboxy-
heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 137)
- 35 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl]-K([17-carboxy-
heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 138)

Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGESES-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 139)

Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGE[8-Amino-3,6-dioxaoctanoyl]ES-K([17-carboxy-heptadecanoyl]-γGlu-G-γGlu)-NH₂ (Compound 140)

5 Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)[Sar]EGESES-K([17-carboxy-heptadecanoyl]-γGlu)-NH₂ (Compound 141)

Ac-SEFC(1)I[1-Me-Trp]QDWGEHRC(1)TEGE[8-Amino-3,6-dioxaoctanoyl]ES-K([17-carboxy-heptadecanoyl]-γGlu)-NH₂ (Compound 142)

10 31. A composition comprising a compstatin analogue according to any one of claims 1 to 30, or a pharmaceutically acceptable salt or solvate thereof, in admixture with a carrier.

32. A composition according to claim 31, wherein the composition is a pharmaceutical composition and the carrier is a pharmaceutically acceptable carrier.

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33. A pharmaceutical composition comprising a compstatin analogue according to any one of claims 1 to 32, or a pharmaceutically acceptable salt or solvate thereof, in admixture with a pharmaceutically acceptable carrier, excipient or vehicle.

20 34. A compstatin analogue, or a pharmaceutically acceptable salt or solvate thereof, according to any one of claims 1 to 30 for use in therapy.

35. A compstatin analogue, or a pharmaceutically acceptable salt or solvate thereof, according to any one of claims 1 to 30 for use in a method of inhibiting complement
25 activation.

36. The compstatin analogue, or a pharmaceutically acceptable salt or solvate thereof, for use according to claim 35, wherein inhibiting complement activation comprises one or more biological activities selected from (1) binding to C3 protein, (2) binding to C3b protein and/or
30 (3) inhibiting the cleavage of native C3 by C3 convertases.

37. A compstatin analogue, or a pharmaceutically acceptable salt or solvate thereof, according to any one of claims 1 to 30 for use in a method of prophylaxis or treatment of age-related macular degeneration, diabetic retinopathy, glaucoma, uveitis, rheumatoid arthritis,
35 spinal cord injury, stroke, multiple sclerosis, Parkinson's disease, Alzheimer's disease, cancer, and respiratory disorders such as asthma, chronic obstructive pulmonary disease (COPD), allergic inflammation, emphysema, bronchitis, bronchiectasis, cystic fibrosis,

tuberculosis, pneumonia, respiratory distress syndrome, rhinitis and sinusitis; bacterial infections such as sepsis, ischemia-reperfusion injury, myocardial infarction, anaphylaxis, paroxysmal nocturnal haemoglobinuria, autoimmune haemolytic anemia, psoriasis, hidradentitis suppurativa, myasthenia gravis, systemic lupus erythematosus, CHAPLE syndrome, C3 glomeropathy, atypical haemolytic uremic syndrome, Crohn's disease or antiphospholipid syndrome.

38. A compstatin analogue, or a pharmaceutically acceptable salt or solvate thereof, according to any one of claims 1 to 30 for use in a method of inhibiting complement activation that occurs during cell or organ transplantation.

39. A method of inhibiting complement activation for treating a subject in need thereof, the method comprising administering to the subject a compstatin analogue, or a pharmaceutically acceptable salt or solvate thereof, according to any one of claims 1 to 30 thereby to inhibit complement activation in the subject.

40. The method of claim 39, wherein the subject has age-related macular degeneration, diabetic retinopathy, glaucoma, uveitis, rheumatoid arthritis, spinal cord injury, stroke, multiple sclerosis, Parkinson's disease, Alzheimer's disease, cancer, and respiratory disorders such as asthma, chronic obstructive pulmonary disease (COPD), allergic inflammation, emphysema, bronchitis, bronchiectasis, cystic fibrosis, tuberculosis, pneumonia, respiratory distress syndrome, rhinitis and sinusitis; bacterial infections such as sepsis, ischemia-reperfusion injury, myocardial infarction, anaphylaxis, paroxysmal nocturnal haemoglobinuria, autoimmune haemolytic anemia, psoriasis, hidradentitis suppurativa, myasthenia gravis, systemic lupus erythematosus, CHAPLE syndrome, IgA nephropathy, C3 glomeropathy, atypical haemolytic uremic syndrome, Crohn's disease or antiphospholipid syndrome, the method comprising administering to the subject a compstatin analogue according to any one of claims 1 to 10.

41. An *ex vivo* method of inhibiting complement activation during extracorporeal shunting of a physiological fluid, the method comprising contacting the physiological fluid with a compstatin analogue, or a pharmaceutically acceptable salt or solvate thereof, according to any one of claims 1 to 30, thereby inhibiting complement activation.

42. Use of a compstatin analogue, or a pharmaceutically acceptable salt or solvate thereof, according to any one of claims 1 to 30, in the preparation of a medicament for inhibiting complement activation.

43. Use of a compstatin analogue, or a pharmaceutically acceptable salt or solvate thereof, according to any one of claims 1 to 30 in the preparation of a medicament for the treatment of age-related macular degeneration, diabetic retinopathy, glaucoma, peridontitis, uveitis, rheumatoid arthritis, spinal cord injury, stroke, multiple sclerosis, Parkinson's disease, Alzheimer's disease, cancer, and respiratory disorders such as asthma, chronic obstructive pulmonary disease (COPD), allergic inflammation, emphysema, bronchitis, bronchiecstasis, cystic fibrosis, tuberculosis, pneumonia, respiratory distress syndrome, rhinitis and sinusitis; bacterial infections such as sepsis, ischemia-reperfusion injury, myocardial infarction, anaphylaxis, paroxysmal nocturnal haemoglobinuria, autoimmune haemolytic anemia, psoriasis, hidradentitis suppurativa, myasthenia gravis, systemic lupus erythematosus, CHAPLE syndrome, C3 glomeropathy, IgA nephropathy, atypical haemolytic uremic syndrome, Crohn's disease or antiphospholipid syndrome.

COMPSTATIN ANALOGUES AND THEIR MEDICAL USES**ABSTRACT**

5

Compstatin analogues having improved binding and complement-inhibiting activity as compared to the 13 amino acid compstatin peptide (ICVVQDWGHRCT (cyclic C2-C12)) are described, in particular compstatin analogues that additionally possess useful

10 physicochemical properties, such as increased solubility. These analogues include variants with an isoleucine residue at position 3 in place of the wild type valine residue led to compstatin peptides with improved binding and complement-inhibiting activity that was found to be compatible with other modifications, for example modifications that are capable of

15 increasing solubility, such as the introduction of charged or polar amino acids at position 9 and/or the introduction of N-and/or C-terminal sequences.

Figure 1

Figure 1a:

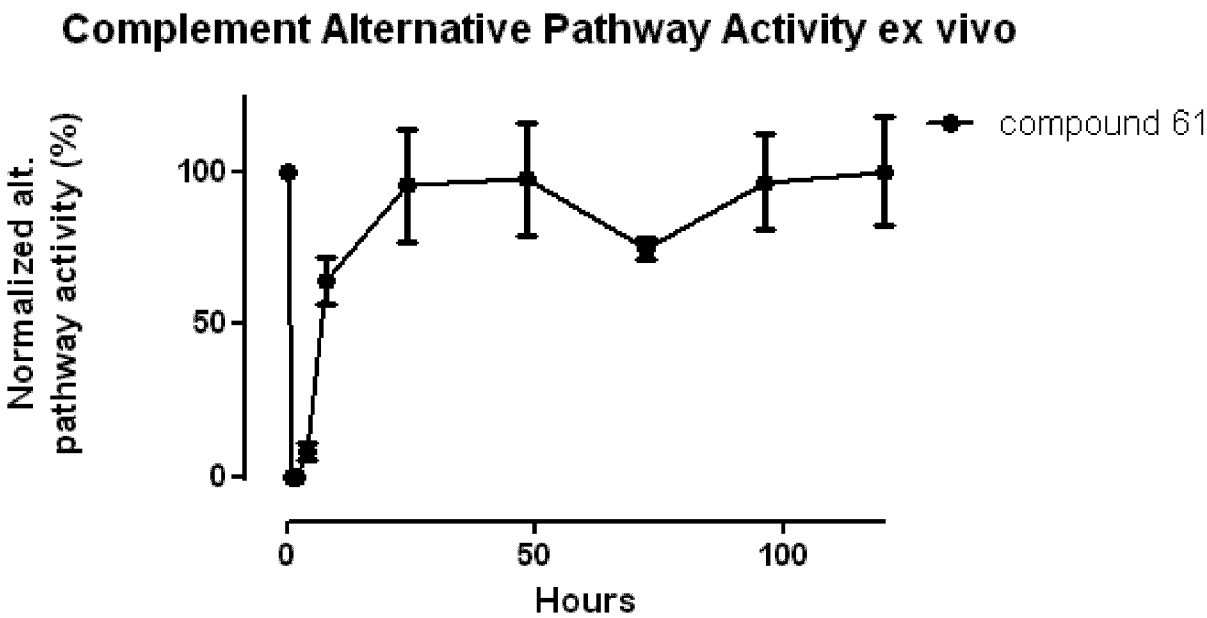


Figure 1b:

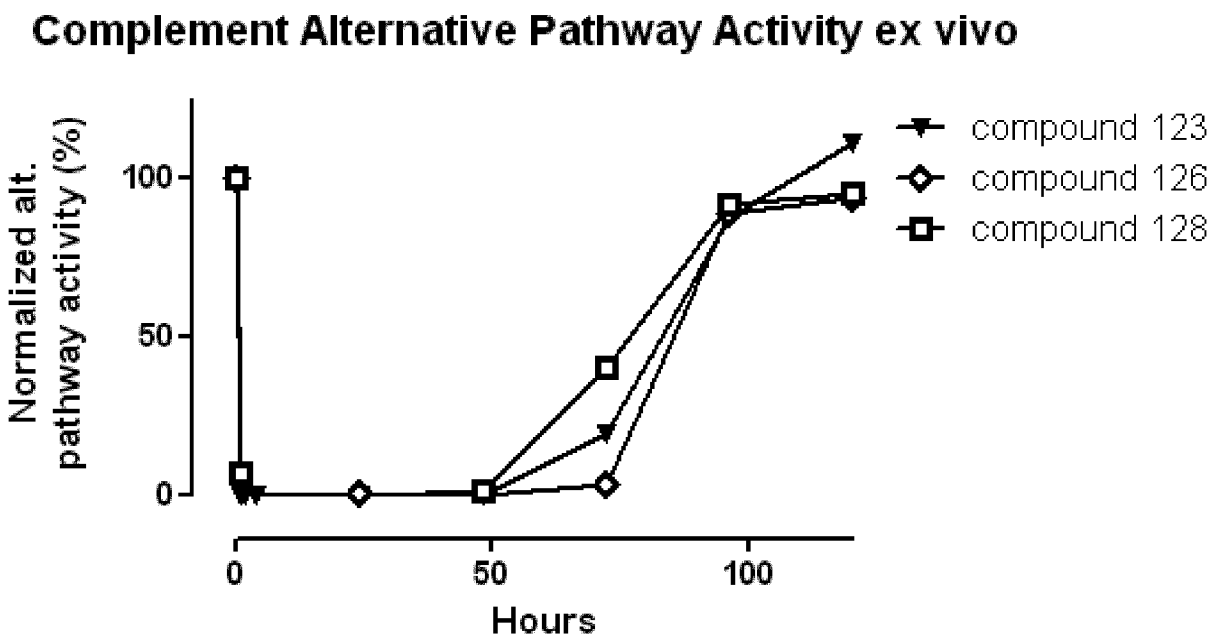


Figure 1c:

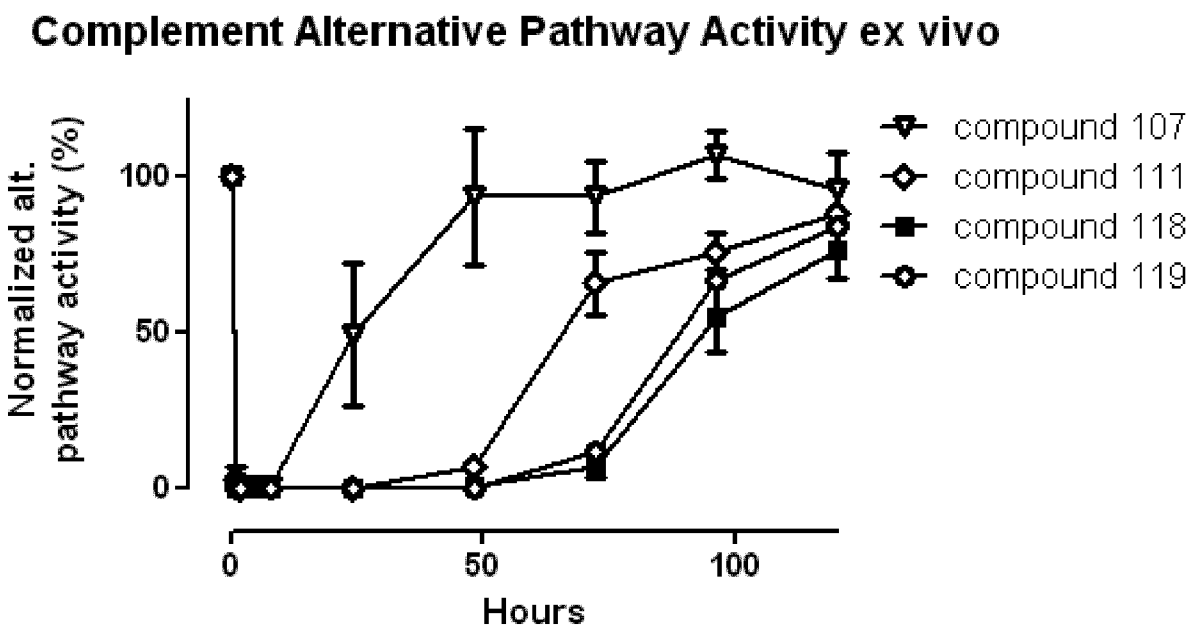


Figure 1d:

