Note: Within nine months of the publication of the mention of the grant of the European patent in the European Patent Bulletin, any person may give notice to the European Patent Office of opposition to that patent, in accordance with the Implementing Regulations. Notice of opposition shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).
The present invention relates to acylated glucagon analogues and their medical use, for example in the treatment of obesity and diabetes.

Obesity and diabetes are globally increasing health problems and are associated with various diseases, particularly cardiovascular disease (CVD), obstructive sleep apnea, stroke, peripheral artery disease, microvascular complications and osteoarthritis.

There are 246 million people worldwide with diabetes, and by 2025 it is estimated that 380 million will have diabetes. Many have additional cardiovascular risk factors including high/aberrant LDL and triglycerides and low HDL.

Cardiovascular disease accounts for about 50% of the mortality in people with diabetes and the morbidity and mortality rates relating to obesity and diabetes underscore the medical need for efficacious treatment options.

Preproglucagon is a 158 amino acid precursor polypeptide that is differentially processed in the tissues to form a number of structurally related proglucagon-derived peptides, including glucagon (Glu), glucagon-like peptide-1 (GLP-1), glucagon-like peptide-2 (GLP-2), and oxyntomodulin (OXM). These molecules are involved in a wide variety of physiological functions, including glucose homeostasis, insulin secretion, gastric emptying and intestinal growth, as well as regulation of food intake.

Glucagon is a 29-amino acid peptide that corresponds to amino acids 53 to 81 of pre-proglucagon and has the sequence His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-Arg-Ala-Gln-Asp-Phe-Val-Gln-Trp-Leu-Met-Asn-Thr. Oxyntomodulin (OXM) is a 37 amino acid peptide which includes the complete 29 amino acid sequence of glucagon with an octapeptide carboxyterminal extension (amino acids 82 to 89 of pre-proglucagon, having the sequence Lys-Arg-Asn-Arg-Asn-Asn-Ile-Ala) and termed “intervening peptide 1” or IP-1; the full sequence of human oxyntomodulin is thus His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-Arg-Ala-Gln-Asp-Phe-Val-Gln-Trp-Leu-Met-Asn-Thr-Lys-Arg-Asn-Ala-Ile-Ala). The major biologically active fragment of GLP-1 is produced as a 30-amino acid, C-terminally amidated peptide that corresponds to amino acids 98 to 127 of pre-proglucagon.

Glucagon helps maintain the level of glucose in the blood by binding to glucagon receptors on hepatocytes, causing the liver to release glucose - stored in the form of glycogen - through glycogenolysis. As these stores become depleted, glucagon stimulates the liver to synthesize additional glucose by gluconeogenesis. This glucose is released into the bloodstream, preventing the development of hypoglycemia. Additionally, glucagon has been demonstrated to increase lipolysis and decrease body weight.

GLP-1 decreases elevated blood glucose levels by improving glucose-stimulated insulin secretion and promotes weight loss chiefly through decreasing food intake. Oxyntomodulin is released into the blood in response to food ingestion and in proportion to meal calorie content. The mechanism of action of oxyntomodulin is not well understood. In particular, it is not known whether the effects of the hormone are mediated exclusively through the glucagon receptor and the GLP-1 receptor, or through one or more as-yet unidentified receptors.


Stabilization of peptides has been shown to provide a better pharmacokinetic profile for several drugs. In particular addition of one or more polyethylene glycol (PEG) or acyl group has been shown to prolong half-life of peptides such as GLP-1 and other peptides with short plasma stability.

In WO 00/55184A1 and WO 00/55119 are disclosed methods for acylation of a range of peptides, in particular GLP-1. Madsen et al (J. Med. Chem. 2007, 50, 6126-6132) describe GLP-1 acylated at position 20 (Liraglutide) and provide data on its stability.

Stabilization of OXM by PEGylation and C-terminal acylation has also been shown to improve the pharmacokinetic profile of selected analogues in WO2007/100535, WO08/071972 and in Endocrinology 2009, 150(4), 1712-1721 by Druce, M R et al.

It has recently been shown that PEGylation of glucagon analogues has a significant effect on the pharmacokinetic profile of the tested compounds (WO2008/101017) but also interferes with the potency of these compounds.

The invention provides a compound having the formula:
R₁-Z-R₂

wherein R₁ is H, C₁₋₄ alkyl, acetyl, formyl, benzoyl or trifluoroacetyl;
R₂ is OH or NH₂;
and Z is a peptide having the formula I

His-X₂-Gtn-Giy-Thr-Thr-Asp-Tyr-Ser-X₁₂-Tyr-Leu-Asp-X₁₆-Ala-Ala-X₂₀-X₂₁-
Phe-Val-X₂₄-Trp-Leu-X₂₇-X₂₈-Ala-X₃₀; (I)

wherein

X₂ is selected from Aib or Ser;
X₁₂ is selected from Lys, Arg and Leu;
X₁₆ is selected from Arg and X;
X₁₇ is selected from Arg and X;
X₂₀ is selected from Arg, His and X;
X₂₁ is selected from Asp and Glu;
X₂₄ is selected from Ala and X;
X₂₇ is selected from Leu and X;
X₂₈ is selected from Arg and X;
X₃₀ is X or is absent;
wherein at least one of X₁₆, X₁₇, X₂₀, X₂₄, X₂₇, X₂₈, and X₃₀ is X;
and wherein each residue X is independently selected from the group consisting of Glu, Lys, Ser, Cys, Dbu, Dpr and Om;

wherein the side chain of at least one residue X is conjugated to a lipophilic substituent having the formula:

(i) Z¹, wherein Z¹ is a lipophilic moiety conjugated directly to the side chain of X; or
(ii) Z¹Z², wherein Z¹ is a lipophilic moiety, Z² is a spacer, and Z¹ is conjugated to the side chain of X via Z²;

with the proviso that Z is not HSQGTFTSDYSKYLDS-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLRA.

[0015] X₃₀ may be present or absent. In those embodiments when X₃₀ is present, it may be desirable for it to be Lys.

[0016] In certain embodiments, any residue X, and especially any residue X which is conjugated to a lipophilic substituent, is independently selected from Lys, Glu or Cys.

[0017] In certain embodiments,
X₁₆ is selected from Glu, Lys and Ser;
X₁₇ is selected from Lys and Cys;
X₂₀ is selected from His, Lys, Arg and Cys;
X₂₄ is selected from Lys, Glu and Ala;
X₂₇ is selected from Leu and Lys; and/or
X₂₈ is selected from Ser, Arg and Lys.

[0018] Specific combinations of residues which may be present in the peptide of formula I include the following:

X₂ is Aib and X₁₇ is Lys;
X₂ is Aib and X₁₇ is Cys;
X₂ is Aib and X₂₀ is Cys;
X₂ is Aib and X₂₈ is Lys;
X₁₂ is Arg and X₁₇ is Lys;
X₁₂ is Leu and X₁₇ is Lys;
X₁₂ is Lys and X₂₀ is Lys;
X₁₂ is Lys and X₁₇ is Lys;
X₁₆ is Lys and X₁₇ is Lys;
X₁₆ is Ser and X₁₇ is Lys;
X₁₇ is Lys and X₂₀ is Lys;
X₁₇ is Lys and X₂₁ is Asp;
X₁₇ is Lys and X₂₄ is Glu;
X17 is Lys and X27 is Leu;  
X17 is Lys and X27 is Lys;  
X17 is Lys and X28 is Ser;  
X17 is Lys and X28 is Arg;  
X20 is Lys and X27 is Leu;  
X21 is Asp and X27 is Leu;  
X2 is Aib, X12 is Lys and X16 is Ser;  
X12 is Lys, X17 is Lys and X16 is Ser;  
X12 is Arg, X17 is Lys and X16 is Glu;  
X16 is Glu, X17 is Lys and X20 is Lys;  
X16 is Ser, X21 is Asp and X24 is Glu;  
X17 is Lys, X24 is Glu and X28 is Arg;  
X17 is Lys, X24 is Glu and X28 is Lys;  
X17 is Lys, X27 is Leu and X28 is Ser;  
X17 is Lys, X27 is Leu and X28 is Arg;  
X20 is Lys, X24 is Glu and X27 is Leu;  
X24 is Glu and X28 is Ser;  
X17 is Lys, X20 is His, X24 is Glu and X27 is Leu;  
X17 is Lys, X20 is His, X24 is Glu and X28 is Ser;  
X17 is Lys, X20 is Lys, X24 is Glu and X27 is Leu; or  
X17 is Cys, X20 is Lys, X24 is Glu and X27 is Leu.

[0019] It may be desirable that the peptide of formula I contains only one amino acid of the type which is to be derivatised by addition of the lipophilic substituent. For example, the peptide may contain only one Lys residue, only one Cys residue or only one Glu residue for the lipophilic substituent to be conjugated to that residue.

[0020] The compounds of the invention may carry one or more intramolecular bridge within the peptide sequence of formula I. Each such bridge is formed between the side chains of two amino acid residues of formula I which are typically separated by three amino acids in the linear amino acid sequence (i.e. between amino acid A and amino acid A+4).

[0021] More particularly, the bridge may be formed between the side chains of residue pairs 16 and 20, 17 and 21, 20 and 24, or 24 and 28. The two side chains can be linked to one another through ionic interactions, or by covalent bonds. Thus these pairs of residues may comprise oppositely charged side chains in order to form a salt bridge by ionic interactions. For example, one of the residues may be Glu or Asp, while the other may be Lys or Arg. The pairings of Lys and Glu and Lys and Asp, may also be capable of reacting to form a lactam ring.

[0022] Examples of suitable pairs of residues at positions 16 and 20 include:

- X16 is Glu and X20 is Lys;  
- X16 is Glu and X20 is Arg;  
- X16 is Lys and X20 is Glu; and  
- X16 is Arg and X20 is Glu.

[0023] Examples of suitable pairs of residues at positions 17 and 21 include:

- X17 is Arg and X21 is Glu;  
- X17 is Lys and X21 is Glu;  
- X17 is Arg and X21 is Asp; and  
- X17 is Lys and X21 is Asp.

[0024] Examples of suitable pairs of residues at positions 20 and 24 include:

- X20 is Glu and X24 is Lys;  
- X20 is Glu and X24 is Arg;  
- X20 is Lys and X24 is Glu; and  
- X20 is Arg and X24 is Glu.

[0025] Examples of suitable pairs of residues at positions 24 and 28 include:

- X24 is Glu and X28 is Lys;
X24 is Glu and X28 is Arg;  
X24 is Lys and X28 is Glu; and  
X24 is Arg and X28 is Glu.

[0026] The pairing of Lys and Glu, e.g. to form a lactam ring, may be particularly desirable, especially between positions 24 and 28.

[0027] It will be apparent that a residue involved in an intramolecular bridge cannot also be derivatised with a lipophilic substituent. Thus, when a residue X is involved in an intramolecular bridge, at least one of the other residues X is conjugated to a lipophilic substituent.

[0028] Without wishing to be bound by any particular theory, it is believed that such intramolecular bridges stabilise the alpha helical structure of the molecule and so increase potency and/or selectivity at the GLP-1 receptor and possibly also the glucagon receptor.

[0029] The compound may have the formula:

R₁-Z-R²

wherein R₁ is H, C₁₋₄ alkyl, acetyl, formyl, benzoyl or trifluoroacetyl;  
R² is OH or NH₂;  
and Z is a peptide having the formula Ila


wherein

X₁₂ is selected from Lys, Arg and Leu;  
X₁₆ is selected from Ser and X;  
X₁₇ is X;  
X₂₀ is selected from His and X;  
X₂₁ is selected from Asp and Glu;  
X₂₄ is selected from Ala and Glu;  
X₂₈ is selected from Ser, Lys and Arg;  
and wherein each residue X is independently selected from the group consisting of Glu, Lys, and Cys;  
wherein the side chain of at least one residue X is conjugated to a lipophilic substituent having the formula:

(i) Z₁', wherein Z₁' is a lipophilic moiety conjugated directly to the side chain of X; or  
(ii) Z₁'Z₂, wherein Z₁' is a lipophilic moiety, Z₂ is a spacer, and Z₁' is conjugated to the side chain of X via Z₂.

[0030] Alternatively, the compound may have the formula:

R₁-Z-R²

wherein R₁ is H, C₁₋₄ alkyl, acetyl, formyl, benzoyl or trifluoroacetyl;  
R² is OH or NH₂;  
and Z is a peptide having the formula Iib

His-Ser-Gln-Gly-Thr-Asp-Ser-X₁₂-Tyr-Leu-Asp-X₁₆-X₁₇-Ala-Ala-X₂₀-X₂₁-Phe-Val-X₂₄-Trp-Leu-Leu-X₂₈-Ala; (IIb)

wherein

X₁₂ is selected from Lys, Arg and Leu;  
X₁₆ is selected from Ser and X;  
X₁₇ is X;  
X₂₀ is selected from His and X;
X21 is selected from Asp and Glu;
X24 is selected from Ala and Glu;
X28 is selected from Ser, Lys and Arg;
and wherein each residue X is independently selected from the group consisting of Glu, Lys, and Cys;

wherein the side chain of at least one residue X is conjugated to a lipophilic substituent having the formula:

(i) Z1, wherein Z1 is a lipophilic moiety conjugated directly to the side chain of X; or
(ii) Z1Z2, wherein Z1 is a lipophilic moiety, Z2 is a spacer, and Z1 is conjugated to the side chain of X via Z2;

with the proviso that Z is not HSQGFTSDYSKYLDS-K(Hexadecanoyl-γ-Glu))-AAHDFVEWLLRA.

[0031] The compound may have the formula:

\[ R^1-Z-R^2 \]

wherein \( R^1 \) is H, C1-4 alkyl, acetyl, formyl, benzoyl or trifluoroacetyl;
\( R^2 \) is OH or NH2;
and Z is a peptide having the formula IIIa

\[
\text{His-Aib-Gln-Gly-Thr-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X17-Ala-Ala-X20-X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala; (IIIa)}
\]

wherein

X12 is selected from Lys and Arg;
X17 is X;
X20 is selected from His and X;
X21 is selected from Asp and Glu;
X24 is selected from Ala and Glu;
X28 is selected from Ser, Lys and Arg;

and wherein each residue X is independently selected from Glu, Lys, and Cys;

wherein the side chain of at least one residue X is conjugated to a lipophilic substituent having the formula:

(i) Z1, wherein Z1 is a lipophilic moiety conjugated directly to the side chain of X; or
(ii) Z1Z2, wherein Z1 is a lipophilic moiety, Z2 is a spacer, and Z1 is conjugated to the side chain of X via Z2.

[0032] Alternatively the compound may have the formula:

\[ R^1-Z-R^2 \]

wherein \( R^1 \) is H, Alkyl, acetyl, formyl, benzoyl or trifluoroacetyl;
\( R^2 \) is OH or NH2;
and Z is a peptide having the formula IIIb

\[
\text{His-Ser-Gln-Gly-Thr-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X17-Ala-Ala-X20-X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala; (IIIb)}
\]

wherein

X12 is selected from Lys or Arg;
X17 is X;
X20 is selected from His and X;
X21 is selected from Asp and Glu;
X24 is selected from Ala and Glu;
X28 is selected from Ser, Lys and Arg;
and wherein each residue X is independently selected from Glu, Lys, and Cys;

wherein the side chain of at least one residue X is conjugated to a lipophilic substituent having the formula:

(i) $Z^1$, wherein $Z^1$ is a lipophilic moiety conjugated directly to the side chain of X; or
(ii) $Z^1Z^2$, wherein $Z^1$ is a lipophilic moiety, $Z^2$ is a spacer, and $Z^1$ is conjugated to the side chain of X via $Z^2$;

with the proviso that $Z$ is not HSQGTFTDSYLDAS-K(Hexadecanoyl-$\gamma$-Glu))-AAHDFVEWLLRA. [0033] The compound may have the formula:

$$R^1-Z-R^2$$

wherein $R^1$ is $H$, C$_1$-4 alkyl, acetyl, formyl, benzoyl or trifluoroacetyl;

$R^2$ is $OH$ or $NH_2$;

and $Z$ is a peptide having the formula IVa

His-Alb-Gln-Gly-Thr-Thr-Ser-Asp-Tyr-Ser-X$_{12}$-Tyr-Leu-Asp-Ser-X$_{17}$-Ala-Ala-His-X$_{21}$-Phe-Val-X$_{24}$-Trp-Leu-Leu-X$_{28}$-Ala; (IVa)

wherein

X$_{12}$ is selected from Lys and Arg;

X$_{17}$ is X;

X$_{21}$ is selected from Asp and Glu;

X$_{24}$ is selected from Ala and Glu;

X$_{28}$ is selected from Ser, Lys and Arg;

and wherein X is selected from the group consisting of Glu, Lys, and Cys;

and wherein the side chain of X is conjugated to a lipophilic substituent having the formula:

(i) $Z^1$, wherein $Z^1$ is a lipophilic moiety conjugated directly to the side chain of X; or
(ii) $Z^1Z^2$, wherein $Z^1$ is a lipophilic moiety, $Z^2$ is a spacer, and $Z^1$ is conjugated to the side chain of X via $Z^2$.

[0034] Alternatively the compound may have the formula:

$$R^1-Z-R^2$$

wherein $R^1$ is $H$, C$_1$-4 alkyl, acetyl, formyl, benzoyl or trifluoroacetyl;

$R^2$ is $OH$ or $NH_2$;

and $Z$ is a peptide having the formula IVb

His-Ser-Gln-Gly-Thr-Thr-Ser-Asp-Tyr-Ser-X$_{12}$-Tyr-Leu-Asp-Ser-X$_{17}$-Ala-Ala-His-X$_{21}$-Phe-Val-X$_{24}$-Trp-Leu-Leu-X$_{28}$-Ala; (IVb)

wherein

X$_{12}$ is selected from Lys and Arg;

X$_{17}$ is X;

X$_{21}$ is selected from Asp and Glu;

X$_{24}$ is selected from Ala and Glu;

X$_{28}$ is selected from Ser, Lys and Arg;

and wherein X is selected from the group consisting of Glu, Lys, and Cys;

and wherein the side chain of X is conjugated to a lipophilic substituent having the formula:

(i) $Z^1$, wherein $Z^1$ is a lipophilic moiety conjugated directly to the side chain of X; or
(ii) $Z_1Z_2$, wherein $Z_1$ is a lipophilic moiety, $Z_2$ is a spacer, and $Z_1$ is conjugated to the side chain of $X$ via $Z_2$;

with the proviso that $Z$ is not HSQGTFTSDYSKYLDS-K(Decanoyl-γ-Glu))-AAHDFVEWLLRA.

Alternatively the compound may have the formula:

$$R^1-Z-R^2$$

wherein $R^1$ is H, $C_{1-4}$ alkyl, acetyl, formyl, benzoyl or trifluoroacetyl;
$R^2$ is OH or NH$_2$;
and $Z$ is a peptide having the formula $V$

$$\text{His-Aib-Gln-Gly-Thr-Ser-Thr-Ser-Leu-Asp-Ser-Lys-Ala-Ala-His-Asp-Phe-Val-Glu-Trp-Leu-Leu-X} \_28$$

(V)

wherein

$X_{28}$ is Ser or absent;
$X_{17}$ is $X$

wherein $X$ is selected from the group consisting of Glu, Lys, and Cys;
and wherein the side chain of $X$ is conjugated to a lipophilic substituent having the formula:

(i) $Z_1$, wherein $Z_1$ is a lipophilic moiety conjugated directly to the side chain of $X$; or
(ii) $Z_1Z_2$, wherein $Z_1$ is a lipophilic moiety, $Z_2$ is a spacer, and $Z_1$ is conjugated to the side chain of $X$ via $Z_2$;

In certain embodiments of the invention, the peptide of formula I may have the sequence:

- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
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- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAHDFVEWLLRA;
In certain embodiments these peptides may carry a lipophilic substituent at the position marked "*" as follows:

- H-Aib-QGTFTSDYSKYLDSKAAK()DFVE()WLLRA;
- H-Aib-QGTFTSDYSKYLDSK()AAHE()FVEWLLKA; or
- H-Aib-QGTFTSDYSKYLDSKQAAKE()FVEWLLRA.

[0037] Residues marked "()" participate in an intramolecular bond, such as a lactam ring. The side chain(s) of one or more of the residues X are conjugated to a lipophilic substituent. For example, one side chain of a residue X may be conjugated to a lipophilic substituent. Alternatively, two, or even more than two, side chains of residues X may be conjugated to a lipophilic substituent.

[0038] For example, at least one of X16, X17, X20 and X28 may be conjugated to a lipophilic substituent. In such cases, X30 may be absent. When X30 is present, it is typically conjugated to a lipophilic substituent.

Thus the compound may have just one lipophilic substituent, at position 16, 17, 20, 24, 27, 28 or 30, preferably at position 16, 17 or 20, particularly at position 17.

Alternatively, the compound may have precisely two lipophilic substituents, each at one of positions 16, 17, 20, 24, 27, 28 or 30. Preferably one or both lipophilic substituents are present at one of positions 16, 17 or 20.

Thus, the compound may have lipophilic substituents at positions 16 and 17, 16 and 20, 16 and 24, 16 and 27, 16 and 28 or 16 and 30; at 17 and 20, 17 and 24, 17 and 27, 17 and 28 or 17 and 30; at 20 and 24, 20 and 27, 20 and 28 or 20 and 30; at 24 and 27, 24 and 28 or 24 and 30; at 27 and 28 or 27 and 30; or at 28 and 30.

In yet further embodiments, the compound may have one or more further lipophilic substituents (giving three or more in total) at further positions selected from positions 16, 17, 20, 24, 27, 28 or 30. However it may be desirable...
that a maximum of two positions are derivatised in this way.

**[0044]** Z1 may comprise a hydrocarbon chain having 10 to 24 C atoms, e.g. 10 to 22 C atoms, e.g. 10 to 20 C atoms. It may have at least 11 C atoms, and/or 18 C atoms or fewer. For example, the hydrocarbon chain may contain 12, 13, 14, 15, 16, 17 or 18 carbon atoms. Thus Z1 may be a dodecanoyl, 2-butyloctanoyl, tetradecanoyl, hexadecanoyl, heptadecanoyl, octadecanoyl or eicosanoyl moiety.

**[0045]** Independently, where present, Z2 may be or comprise one or more amino acid residues. For example, Z2 may be a γ-Glu, Glu, β-Ala or ε-Lys residue, or a 4-aminobutanoyl, 8-aminooctanoyl or 8-amino-3,6-dioxaoctanoyl moiety. Certain combinations of Z1 and Z2 are dodecanoyl-γ-Glu, hexadecanoyl-γ-Glu, hexadecanoyl-Glu, hexadecanoyl-[3-aminopropanoyl], hexadecanoyl-[8-aminooctanoyl], hexadecanoyl-ε-Lys, 2-butyloctanoyl-γ-Glu, octadecanoyl-γ-Glu and hexadecanoyl-[4-aminobutanoyl].

**[0046]** In particular embodiments, Z has the formula:

- HSQGTFTSDYSKYLDS-K(Hexadecanoyl-γ-Glu)-KAAHDFVEWLLRA;
- HSQGTFTSDYSKYLDSKAA-K(Hexadecanoyl-γ-Glu)-DFVWLLRA;
- HSQGTFTSDYSKYLDSKAAHDFVEWLL-K(Hexadecanoyl-γ-Glu)-RA;
- H-Aib-QGTFTSDYSKYLDS-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLRA;
- H-Aib-QGTFTSDYSKYLDS-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLA;
- H-Aib-QGTFTSDYSKYLDS-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLA;

Residues marked "()" participate in an intramolecular bond, such as a lactam ring.

**[0047]** In a further embodiment, Z has the formula:

- H-Aib-QGTFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-YLDSKAAHDFVEWLLSA;
- H-Aib-QGTFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-KAHDFVEWLLSA;
- H-Aib-QGTFTSDYSKYLDSKAA-K(Hexadecanoyl-isoGlu)-DFVWLLSA;
- H-Aib-QGTFTSDYSKYLDSKAAHDFVEWLL-K(Hexadecanoyl-isoGlu)-LLSA;
- H-Aib-QGTFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA;
- H-Aib-QGTFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLA;
- H-Aib-QGTFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLA;

**[0048]** In a further aspect, Z has the formula:

- H-Aib-QGTFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLL;

Residues marked "(" participate in an intramolecular bond, such as a lactam ring.
In still a further aspect, Z has the formula:

\[
\text{H-Aib-QGFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLL;}
\]

In a further aspect, the present invention provides a composition comprising a compound as defined herein, or a salt or derivative thereof, in admixture with a carrier. In preferred embodiments, the composition is a pharmaceutically acceptable composition and the carrier is a pharmaceutically acceptable carrier. The salt may be a pharmaceutically acceptable acid addition salt of the compound, e.g. an acetate or chloride salt.

The compounds described find use in preventing weight gain or promoting weight loss. By "preventing" is meant inhibiting or reducing weight gain when compared to the absence of treatment, and is not necessarily meant to imply complete cessation of weight gain. The peptides may cause a decrease in food intake and/or increased energy expenditure, resulting in the observed effect on body weight. Independently of their effect on body weight, the compounds of the invention may have a beneficial effect on circulating glucose levels, glucose tolerance, and/or on circulating cholesterol levels, being capable of lowering circulating LDL levels and increasing HDL/DLD ratio. Thus the compounds of the invention can be used for direct or indirect therapy of any condition caused or characterised by excess body weight, such as the treatment and/or prevention of obesity, morbidity obesity, obesity linked inflammation, obesity linked gallbladder disease, obesity induced sleep apnoea. They may also be used for the treatment of pre-diabetes, insulin resistance, glucose intolerance, type 2 diabetes, type 1 diabetes, hypertension or atherogenic dyslipidaemia (or a combination of two or more of these metabolic risk factors), atherosclerosis, arteriosclerosis, coronary heart disease, peripheral artery disease, stroke and microvascular disease. Their effects in these conditions may be as a result of or associated with their effect on body weight, or may be independent thereof.

The invention provides a compound of the invention for use in a method of medical treatment, particularly for use in a method of treatment of a condition as described above.

The invention also provides the use of a compound of the invention in the preparation of a medicament for the treatment of a condition as described above.

The compound of the invention may be administered as part of a combination therapy with an agent for treatment of diabetes, obesity, dyslipidaemia or hypertension.

Thus the compound of the invention (or the salt thereof) can be used in combination with an antidiabetic agent including but not limited to metformin, a sulfonylurea, a glinide, a DPP-IV inhibitor, a glitazone, or insulin. In a preferred embodiment the compound or salt thereof is used in combination with insulin, DPP-IV inhibitor, sulfonylurea or metformin, particularly sulfonylurea or metformin, for achieving adequate glycemic control. In an even more preferred embodiment the compound or salt thereof is used in combination with a metformin, a sulfonylurea, insulin or an insulin analogue for achieving adequate glycemic control. Examples of insulin analogues include but are not limited to Lantus, Novorapid, Humalog, Novomix, Actraphane HM, Levemir and Apidra.

The compound or salt thereof can further be used in combination with an anti-obesity agent including but not limited to a glucagon-like peptide receptor 1 agonist, peptide YY or analogue thereof, cannabinoid receptor 1 antagonist, lipase inhibitor, melanocortin receptor 4 agonist, or melanin concentrating hormone receptor 1 antagonist.

The compound or salt thereof can further be used in combination with an anti-hypertension agent including but not limited to an angiotensin-converting enzyme inhibitor, angiotensin II receptor blocker, diuretic, beta-blocker, or calcium channel blocker.

The compound or salt thereof can be used in combination with an anti-dyslipidemia agent including but not limited to a statin, a fibrate, a niacin or a cholesterol absorption inhibitor.

DESCRIPTION OF THE FIGURES

**Figure 1.** Pharmacokinetic profile of compound 13 after subcutaneous (s.c.) administration to mice at a dose of 100 nmol/kg.

**Figure 2.** Effect of 21 days s.c. administration of compound 11 (10 nmol/kg) on oral glucose tolerance in long term high fat fed C57BL/6J mice. Data are shown as mean ± SEM.

**Figure 3.** Diabetic (db/db) mice were treated with vehicle or compound 7 (12.7 nmol/kg) for 4 weeks and HbA1c
was determined (Cobas® application note: A1C-2) in whole blood samples (20 μl) collected from the treated mice. The $\Delta$HbA1c (%) was calculated for each mouse by subtracting its HbA1c (%) at start of treatment from HbA1c (%) at 4 weeks. $\Delta$HbA1c (%) of db/db mice treated for 4 weeks with vehicle = 100%. * (P = 0.03, Students t-test).

**Figure 4.** Effect of 21 days s.c. administration of compound 11 on body weight in long term high fat fed C57BL/6J mice. Data are shown as mean+SEM.

**Figure 5.** Diet Induced Obese (DIO) mice were treated with vehicle or compound 7 (12.7 nmol/kg) for 4 weeks and plasma prepared from the collected blood samples. Total cholesterol was determined in each plasma sample (Cobas®; application note CHOL2). *** (P < 0.0001, Students t-test). Data are shown as mean+SEM.

**Figure 6.** Diet Induced Obese (DIO) mice were treated with vehicle or compound 7 (12.7 nmol/kg) and plasma prepared from the collected blood samples. LDL and HDL cholesterol were determined in each plasma sample (Cobas®; application notes HDLC3 and LDL_C). *** (P < 0.0001, Students t-test). Data are shown as mean+SEM.

**Figure 7.** Effect of s.c. administration of GluGLP-1 agonists on body weight gain in high fat fed C57BL/6J mice. Data are mean±SEM. Black line: Vehicle (PBS), Grey line: Low dose (0.5 nmol/kg), Broken line: High dose (5 nmol/kg).

**Figure 8.** Effect of acute s.c. administration of Compound 7 on oral glucose tolerance 2, 4, 6, 8, 10 and 12 h after dosing in high fat fed C57BL/6J mice. Data are expressed as mean+SEM.

**Figure 9.** Effect of s.c. administration of Compound 7 and exendin-4 on food intake/body weight in young lean C57BL/6J mice. Data are mean+SEM. *p<0.05 versus young lean vehicle. Data are expressed as mean+SEM.

**Figure 10.** Effect of s.c. administration of Compound 7 and exendin-4 on cumulative food intake/body weight in old obese C57BL/6J mice. Data are mean+SEM. *p<0.05 versus old obese vehicle. Data are expressed as mean+SEM.

**Figure 11.** Effect of s.c. administration of Vehicle, exendin-4 (10 nmol/kg) and Compound 11 (10 nmol/kg) on plasma lipid concentration in old obese C57BL/6J mice. Data are mean+SEM.

**Figure 12.** Mice were treated twice daily s.c. with Compound. 1 and Compound. 11 (at two doses: 0.5 and 5 nmol/kg) or vehicle for 2 weeks. On the day of sacrifice, the liver was exposed, and weighed. Compound 1 significantly increased "liver weight/body weight ratio" at the high dose. Compound. 11 did not affect "liver weight/body weight ratio" at the two doses (0.5 and 5 nmol/kg). Compound 1 is a non-acylated dual GluGLP-1 agonists and Compound. 11 is a long-acting acylated dual GluGLP-1 agonists (Figure 12).

**Figure 13.** Diabetic (db/db) mice were treated with vehicle or compound 11 (12.7 nmol/kg) for 4 weeks and HbA1c was determined (Cobas® application note: A1C-2) in whole blood samples (20 μl) collected from the treated mice. The $\Delta$HbA1c (%) was calculated for each mouse by subtracting its HbA1c (%) at start of treatment from HbA1c (%) at 4 weeks. $\Delta$HbA1c (%) of db/db mice treated for 4 weeks with vehicle = 100%. * (P = 0.03, Students t-test).

**DETAILED DESCRIPTION OF THE INVENTION**

Throughout this specification, the conventional one letter and three letter codes for naturally occurring amino acids are used, as well as generally accepted three letter codes for other amino acids, including Aib (α-aminoisobutyric acid), Orn (ornithine), Dbu (2,4 diaminobutyric acid) and Dpr (2,3-diaminopropanoic acid).

The term "native glucagon" refers to native human glucagon having the sequence H-His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-Lys-Tyr-Leu-Asp-Ser-Arg-Arg-Ala-Gln-Asp-Phe-Val-Gln-Trp-Leu-Met-Asn-Thr-OH.

The peptide sequence of the compound of the invention differs from that of native glucagon at least at positions 18, 20, 24, 27, 28 and 29. In addition, it may differ from that of native glucagon at one or more of positions 12, 16 and 17. Native glucagon has Arg at position 18. The compound of the invention has the small hydrophobic residue Ala at position 18 which is believed to increase potency at both glucagon and GLP-1 receptors but particularly the GLP-1 receptor.

The residues at positions 27, 28 and 29 of native glucagon appear to provide significant selectivity for the glucagon receptor. The substitutions at these positions with respect to the native glucagon sequence, particularly the Ala at position 29, may increase potency at and/or selectivity for the GLP-1 receptor, potentially without significant reduction of potency at the glucagon receptor. Further examples which may be included in the compunds of the invention
include Leu at position 27 and Arg at position 28. Furthermore, Arg at position 28 may be particularly preferred when there is a Glu at position 24 with which it can form an intramolecular bridge, since this may increase its effect on potency at the GLP-1 receptor.

Substitution of the naturally-occurring Met residue at position 27 (e.g. with Leu, Lys or Glu) also reduces the potential for oxidation, thereby increasing the chemical stability of the compounds.

Substitution of the naturally-occurring Asn residue at position 28 (e.g. by Arg or Ser) also reduces the potential for deamination in acidic solution, thereby increasing the chemical stability of the compounds.

Potency and/or selectivity at the GLP-1 receptor, potentially without significant loss of potency at the glucagon receptor, may also be increased by introducing residues that are likely to stabilise an alpha-helical structure in the C-terminal portion of the peptide. It may be desirable, but is not believed essential, for this helical portion of the molecule to have an amphipathic character. Introduction of residues such as Leu at position 12 and/or Ala at position 24 may assist. Additionally or alternatively charged residues may be introduced at one or more of positions 16, 20, 24, and 28. Thus the residues of positions 24 and 28 may all be charged, the residues at positions 20, 24, and 28 may all be charged, or the residues at positions 16, 20, 24, and 28 may all be charged. For example, the residue at position 20 may be His or Arg, particularly His. The residue at position 24 may be Glu, Lys or Arg, particularly Glu. The residue at position 28 may be Arg. Introduction of an intramolecular bridge in this portion of the molecule, as discussed above, may also contribute to stabilising the helical character, e.g. between positions 24 and 28.

Substitution of one or both of the naturally-occurring Gln residues at positions 20 and 24 also reduces the potential for deamination in acidic solution, so increasing the chemical stability of the compounds.

A substitution relative to the native glucagon sequence at position 12 (i.e. of Arg or Leu) may increase potency at both receptors and/or selectivity at the GLP-1 receptor.

C-terminal truncation of the peptide does not reduce potency of both receptors and/or selectivity of the GLP-1 receptor. In particular, truncation of position 29 or truncation of both position 28 and 29 does not reduce the receptor potency to any of the two receptors.

The side chain of one or more of the residues designated X (i.e. positions 16, 17, 20, 24, 27 and 28, and/or 30 if present) is conjugated to a lipophilic substituent. It will be appreciated that conjugation of the lipophilic substituent to a particular side chain may affect (e.g. reduce) certain of the benefits which the unconjugated side chain may provide at that position. The inventors have found that compounds of the invention provide a balance between the benefits of acylation and the benefits of particular substitutions relative to the native glucagon sequence.

Compositions of the invention may further be compounded in, or attached to, for example through covalent, hydrophobic and electrostatic interactions, a drug carrier, drug delivery system and advanced drug delivery system in order to further enhance stability of the compound, increase bioavailability, increase solubility, decrease adverse effects, achieve chronotherapy well known to those skilled in the art, and increase patient compliance or any combination thereof. Examples of carriers, drug delivery systems and advanced drug delivery systems include, but are not limited to, polymers, for example cellulose and derivatives, polysaccharides, for example dextran and derivatives, starch and derivatives, copolymeric systems well known to those skilled in the art, micelles, liposomes, microspheres, nanoparticulates, liquid crystals and dispersions thereof, L2 phase and dispersions thereof, well known to those skilled in the art of phase behaviour in lipid-water systems, polymeric micelles, multiple emulsions, self-emulsifying, self-microemulsifying, cyclo-dextrins and derivatives thereof, and dendrimers.

Other groups have attempted to prolong the half life of GluGLP-1 dual agonist compounds by derivatisation with PEG (WO2008/101017). However such derivatisation appears to be most effective when applied to the C-terminus of the molecule rather than in the central core of the peptide backbone, and potency of these compounds is still decreased compared to the corresponding unmodified peptide.

By contrast, the compounds of the present invention retain high potency at both the glucagon and GLP-1 receptors while having significantly protracted pharmacokinetic profiles compared to the corresponding unmodified peptides.

Native glucagon has Ser at position 16. Substitution with Ala, Gly or Thr has been shown to reduce adenylyl cyclase activation at the glucagon receptor significantly (Unson et al. Proc. Natl. Acad. Sci. 1994, 91, 454-458). Hence, derivatisation with a lipophilic substituent at position 16 would not have been expected to yield compounds retaining potency at the glucagon receptor, as is surprisingly shown by the compounds described in this specification. In WO2008/101017 a negatively charged residue was found to be desirable at position 16 to minimise loss of potency. The presence of basic amino acids at positions 17 and 18 is generally believed to be necessary for full glucagon receptor activation (Unson et al. J. Biol. Chem. 1998, 273, 10308-10312). The present inventors have found that, when position 18 is alanine, substitution with a hydrophobic amino acid in position 17 can still yield a highly potent compound. Even compounds in which the amino acid in position 17 is derivatised with a lipophilic substituent retain almost full potency at both glucagon and GLP-1 receptors, as well as displaying a significantly protracted pharmacokinetic profile. This is
so even when a lysine at position 17 is derivatised, converting the basic amine side chain into a neutral amide group.

[0077] The present inventors have also found that compounds with acylation at position 20 are still highly active dual agonists, despite indications from other studies that substitution in position 20 should be a basic amino acid having a side chain of 4-6 atoms in length to enhance GLP-1 receptor activity compared to glucagon (WO2008/101017). The compounds described herein retain both GLP-1 and glucagon receptor activity when position 20 is substituted with lysine and acylated.

Peptide synthesis

[0078] The peptide component of the compounds of the invention may be manufactured by standard synthetic methods, recombinant expression systems, or any other suitable method. Thus the peptides may be synthesized in a number of ways including for example, a method which comprises:

(a) synthesizing the peptide by means of solid phase or liquid phase methodology either stepwise or by fragment assembling and isolation and purification of the final peptide product;

(b) expressing a nucleic acid construct that encodes the peptide in a host cell and recovering the expression product from the host cell culture; or

(c) effecting cell-free in vitro expression of a nucleic acid construct that encodes the peptide and recovering the expression product;

or any combination of methods of (a), (b), and (c) to obtain fragments of the peptide, subsequently ligating the fragments to obtain the peptide, and recovering the peptide.

[0079] It may be preferred to synthesize the analogues of the invention by means of solid phase or liquid phase peptide synthesis. In this context, reference is given to WO 98/11125 and, amongst many others, Fields, GB et al., 2002, "Principles and practice of solid-phase peptide synthesis". In: Synthetic Peptides (2nd Edition) and the examples herein.

Lipophilic substituent

[0080] One or more of the amino acid side chains in the compound of the invention is conjugated to a lipophilic substituent Z1. Without wishing to be bound by theory, it is thought that the lipophilic substituent binds albumin in the blood stream, thus shielding the compounds of the invention from enzymatic degradation which can enhance the half-life of the compounds. It may also modulate the potency of the compound, e.g. with respect to the glucagon receptor and/or the GLP-1 receptor.

[0081] 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.

[0082] The lipophilic substituent Z1 may be covalently bonded to an atom in the amino acid side chain, or alternatively may be conjugated to the amino acid side chain by a spacer Z2.

[0083] The term "conjugated" is used here to describe the physical 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.

[0084] The spacer Z2, when present, is used to provide a spacing between the compound and the lipophilic moiety. The spacer Z2 may be attached to the amino acid side chain or to the spacer via an ester, a sulphonyl ester, a thioester, an amide or a sulphonamide. Accordingly it will be understood that preferably the lipophilic substituent includes 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. Preferably, an acyl group in the lipophilic substituent forms part of an amide or ester with the amino acid side chain or the spacer.

[0085] The lipophilic substituent may include a hydrocarbon chain having 10 to 24 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. From the discussion above it will be understood that the hydrocarbon chain is preferably 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 acyl, and accordingly the hydrocarbon chain may be part of an alkanoic group, for example a dodecanoyl, 2-butyloctanoyl, tetradecanoyl, hexadecanoyl, heptadecanoyl, octadecanoyl or eicosanoyl group.
As mentioned above, the lipophilic substituent Z1 may be conjugated to the amino acid side chain by a spacer Z2. When present, the spacer is attached to the lipophilic substituent and to the amino acid side chain. 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 C1,10 hydrocarbon chain or more preferably a linear C1,6 hydrocarbon chain. Furthermore the spacer can be substituted with one or more substituents selected from C1,6 alkyl, Alkyl amine, C1,6 alkyl hydroxy and Alkyl carboxy.

The spacer may be, for example, a residue of any naturally occurring or unnatural amino acid. For example, the spacer may be a residue of Gly, Pro, Ala, Val, Leu, Ile, Met, Cys, Phe, Tyr, Trp, His, Lys, Arg, Gin, Asn, ε-Glu, γ-Glu, ε-Lys, Asp, Ser, Thr, Gaba, Aib, β-Ala (i.e. 3-aminopropanoyl), 4-aminobutanoyl, 5-aminopentanoyl, 6-aminohexanoyl, 7-aminooctanoyl, 8-aminononanoyl, 9-aminodecanoyl or 8-amino-3,6-dioxaoctanoyl. In certain embodiments, the spacer is a residue of Glu, γ-Glu, ε-Lys, β-Ala (i.e. 3-aminopropanoyl), 4-aminobutanoyl, 8-aminooctanoyl or 8-amino-3,6-dioxaoctanoyl. In the present invention, γ-Glu and isoGlu are used interchangeably.

The amino acid side chain to which the lipophilic substituent is conjugated is a side chain of a Glu, Lys, Ser, Cys, Dbu, Dpr or Om 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 these 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.

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

Here, the side chain of a Lys residue from the peptide of formula I is covalently attached to an 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.

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

PEGylated and/or acylation have a short half-life (T½), which gives rise to burst increases of GluGLP-1 agonist concentrations. The glucagon receptor is thus being subjected to burst exposure to the glucagon agonism once (or twice) daily throughout the treatment period.

Without being bound to any theory repeated burst exposure of GluR to glucagon agonism seems to bring havoc to the lipid and free fatty acid trafficking between the liver and adipose tissue with the result that fat accumulates in the liver. Constant exposure of GluR to glucagon agonism blocks accumulation of fat in the liver It has thus been found, that repeated treatment with glucagon or short acting dual GluGLP-1 agonists give rise to enlarged liver due to fat and glycogen accumulation (Chan et al., 1984. Exp. Mol. Path. 40, 320-327).

Repeated treatment with long-acting acylated dual GluGLP-1 agonists do not give rise to change in liver size.
(enlarged or shrunken) in normal weight subjects, but normalize liver lipid content (Day et al., 2009; Nat.Chem.Biol. 5, 749 - 57).

**Efficacy**

[0096] Binding of the relevant compounds to GLP-1 or glucagon (Glu) receptors may be used as an indication of agonist activity, but in general it is preferred to use a biological assay which measures intracellular signalling caused by binding of the compound to the relevant receptor. For example, activation of the glucagon receptor by a glucagon agonist will stimulate cellular cyclic AMP (cAMP) formation. Similarly, activation of the GLP-1 receptor by a GLP-1 agonist will stimulate cellular cAMP formation. Thus, production of cAMP in suitable cells expressing one of these two receptors can be used to monitor the relevant receptor activity. Use of a suitable pair of cell types, each expressing one receptor but not the other, can hence be used to determine agonist activity towards both types of receptor.

[0097] The skilled person will be aware of suitable assay formats, and examples are provided below. The GLP-1 receptor and/or the glucagon receptor may have the sequence of the receptors as described in the examples. For example, the assays may make use the human glucagon receptor (Glucagon-R) having primary accession number GI: 4503947 (NMP_000151.1) and/or the human glucagon-like peptide 1 receptor (GLP-1R) having primary accession number GI:166795283 (NP_002053.3).

(Where sequences of precursor proteins are referred to, it should of course be understood that assays may make use of the mature protein, lacking the signal sequence).

[0098] EC50 values may be used as a numerical measure of agonist potency at a given receptor. An EC50 value is a measure of the concentration of a compound required to achieve half of that compound’s maximal activity in a particular assay. Thus, for example, a compound having EC50 [GLP-1 R] lower than the EC50 [GLP-1 R] of native glucagon in a particular assay may be considered to have higher potency at the GLP-1 R than glucagon.

[0099] The compounds described in this specification are typically Glu-GLP-1 dual agonists, i.e. they are capable of stimulating cAMP formation at both the glucagon receptor and the GLP-1R. The stimulation of each receptor can be measured in independent assays and afterwards compared to each other.

[0100] By comparing the EC50 value for the glucagon receptor (EC50 [Glucagon-R]) with the EC50 value for the GLP-1 receptor (EC50 [GLP-1R]) for a given compound the relative glucagon selectivity (%) of that compound can be found:

\[
\text{Relative Glucagon-R selectivity [Compound]} = (1/\text{EC50 [Glucagon-R]}) \times 100\% / (1/\text{EC50 [Glucagon-R]}) + 1/\text{EC50 [GLP-1R]})
\]

[0101] The relative GLP-1R selectivity can likewise be found:

\[
\text{Relative GLP-1R selectivity [Compound]} = (1/\text{EC50 [GLP-1R]}) \times 100\% / (1/\text{EC50 [Glucagon-R]}) + 1/\text{EC50 [GLP-1R]})
\]

[0102] A compound’s relative selectivity allows its effect on the GLP-1 or glucagon receptor to be compared directly to its effect on the other receptor. For example, the higher a compound’s relative GLP-1 selectivity is, the more effective that compound is on the GLP-1 receptor as compared to the glucagon receptor.

[0103] Using the assays described below, we have found the relative GLP-1 selectivity for human glucagon to be approximately 5%.

[0104] The compounds of the invention have a higher relative GLP-1 R selectivity than human glucagon. Thus, for a particular level of glucagon-R agonist activity, the compound will display a higher level of GLP-1 R agonist activity (i.e. greater potency at the GLP-1 receptor) than glucagon. It will be understood that the absolute potency of a particular compound at the glucagon and GLP-1 receptors may be higher, lower or approximately equal to that of native human glucagon, as long as the appropriate relative GLP-1 R selectivity is achieved.

[0105] Nevertheless, the compounds of this invention may have a lower EC50 [GLP-1 R] than human glucagon. The compounds may have a lower EC50 [GLP-1R] than glucagon while maintaining an EC50 [Glucagon-R] that is less than 10-fold higher than that of human glucagon, less than 5-fold higher than that of human glucagon, or less than 2-fold higher than that of human glucagon.

[0106] It may be desirable that EC50 of any given compound for both the Glucagon-R and GLP-1 R should be less than 1 nM.

[0107] The compounds of the invention may have an EC50 [Glucagon-R] that is less than two-fold that of human glucagon. The compounds may have an EC50 [Glucagon-R] that is less than two-fold that of human glucagon and have
an EC50 [GLP-1R] that is less than half that of human glucagon, less than a fifth of that of human glucagon, or less than a tenth of that of human glucagon.  

**0108** The relative GLP-1 selectivity of the compounds may be greater than 5% and less than 95%. For example, the compounds may have a relative selectivity of 5-20%, 10-30%, 20-50%, 30-70%, or 50-80%, or of 30-50%, 40-60%, 50-70% or 75-95%.

**Therapeutic uses**

**0109** The compounds of the invention may provide an attractive treatment option for metabolic diseases including obesity and diabetes mellitus (diabetes).

**0110** Diabetes comprises a group of metabolic diseases characterized by hyperglycemia resulting from defects in insulin secretion, insulin action, or both. Acute signs of diabetes include excessive urine production, resulting compensatory thirst and increased fluid intake, blurred vision, unexplained weight loss, lethargy, and changes in energy metabolism. The chronic hyperglycemia of diabetes is associated with long-term damage, dysfunction, and failure of various organs, notably the eyes, kidneys, nerves, heart and blood vessels. Diabetes is classified into type 1 diabetes, type 2 diabetes and gestational diabetes on the basis on pathogenetic characteristics.

**0111** Type 1 diabetes accounts for 5-10% of all diabetes cases and is caused by auto-immune destruction of insulin-secreting pancreatic β-cells.

**0112** Type 2 diabetes accounts for 90-95% of diabetes cases and is a result of a complex set of metabolic disorders. Type 2 diabetes is the consequence of endogenous insulin production becoming insufficient to maintain plasma glucose levels below the diagnostic thresholds.

**0113** Gestational diabetes refers to any degree of glucose intolerance identified during pregnancy.

**0114** Pre-diabetes includes impaired fasting glucose and impaired glucose tolerance and refers to those states that occur when blood glucose levels are elevated but below the levels that are established for the clinical diagnosis for diabetes.

**0115** A large proportion of people with type 2 diabetes and pre-diabetes are at increased risk of morbidity and mortality due to the high prevalence of additional metabolic risk factors including abdominal obesity (excessive fat tissue around the abdominal internal organs), atherogenic dyslipidemia (blood fat disorders including high triglycerides, low HDL cholesterol and/or high LDL cholesterol, which foster plaque buildup in artery walls), elevated blood pressure (hypertension) a prothrombotic state (e.g. high fibrinogen or plasminogen activator inhibitor-1 in the blood), and proinflammatory state (e.g., elevated C-reactive protein in the blood).

**0116** Conversely, obesity confers an increased risk of developing pre-diabetes, type 2 diabetes as well as e.g. certain types of cancer, obstructive sleep apnea and gall-bladder disease.

**0117** Dyslipidaemia is associated with increased risk of cardiovascular disease. High Density Lipoprotein (HDL) is of clinical importance since an inverse correlation exists between plasma HDL concentrations and risk of atherosclerotic disease. The majority of cholesterol stored in atherosclerotic plaques originates from LDL and hence elevated concentrations Low Density Lipoproteins (LDL) is closely associated with atherosclerosis. The HDULDL ratio is a clinical risk indicator for atherosclerosis and coronary atherosclerosis in particular.

**0118** Without wishing to be bound by any particular theory, it is believed that the compounds of the invention act as GluGLP-1 dual agonists. The dual agonist may combine the effect of glucagon e.g. on fat metabolism with the effect of GLP-1 e.g. on blood glucose levels and food intake. They might therefore act to accelerate elimination of excessive adipose tissue, induce sustainable weight loss, and improve glycaemic control. Dual GluGLP-1 agonists might also act to reduce cardiovascular risk factors such as high cholesterol and LDL-cholesterol.

**0119** The compounds of the present invention can therefore be used as pharmaceutical agents for preventing weight gain, promoting weight loss, reducing excess body weight or treating obesity (e.g. by control of appetite, feeding, food intake, calorie intake, and/or energy expenditure), including morbid obesity, as well as associated diseases and health conditions including but not limited to obesity linked inflammation, obesity linked gallbladder disease and obesity induced sleep apnea. The compounds of the invention may also be used for treatment of insulin resistance, glucose intolerance, pre-diabetes, increased fasting glucose, type 2 diabetes, hypertension, dyslipidemia (or a combination of these metabolic risk factors), atherosclerosis, arteriosclerosis, coronary heart disease, peripheral artery disease and stroke. These are all conditions which can be associated with obesity. However, the effects of the compounds of the invention on these conditions may be mediated in whole or in part via an effect on body weight, or may be independent thereof.

**Pharmaceutical compositions**

**0120** The compounds of the present invention, or salts thereof, may be formulated as pharmaceutical compositions prepared for storage or administration, which typically comprise a therapeutically effective amount of a compound of the invention, or a salt thereof, in a pharmaceutically acceptable carrier.
The therapeutically effective amount of a compound of the present invention will depend on the route of administration, the type of mammal being treated, and the physical characteristics of the specific mammal under consideration. These factors and their relationship to determining this amount are well known to skilled practitioners in the medical arts. This amount and the method of administration can be tailored to achieve optimal efficacy, and may depend on such factors as weight, diet, concurrent medication and other factors, well known to those skilled in the medical arts. The dosage sizes and dosing regimen most appropriate for 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 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. Such considerations are known to the skilled person.

The term "pharmaceutically acceptable carrier" includes any of the standard pharmaceutical carriers. Pharmaceutically acceptable carriers for therapeutic use are well known in the pharmaceutical art, and are described, for example, in Remington's Pharmaceutical Sciences, Mack Publishing Co. (A. R. Gennaro edit. 1985). For example, sterile saline and phosphate-buffered saline at slightly acidic or physiological pH may be used. pH buffering agents may be phosphate, citrate, acetate, tris(hydroxymethyl)aminomethane (TRIS), N-Tris(hydroxymethyl)methyl-3-aminopropanesulphonic acid (TAPS), ammonium bicarbonate, diethanolamine, histidine, which is a preferred buffer, arginine, lysine, or acetate or mixtures thereof. The term further encompasses any agents listed in the US Pharmacopeia for use in animals, including humans.

The term "pharmaceutically acceptable salt" refers to the salt of the compounds. Salts include pharmaceutically acceptable salts such as acid addition salts and basic salts. Examples of acid addition salts include hydrochloride salts, citrate salts and acetate salts. Examples of basic salts include salts where the cation is selected from alkali metals, such as sodium and potassium, alkaline earth metals such as calcium, and ammonium ions +N(R3)3(R4), where R3 and R4 independently designate optionally substituted C1-C6-alkyl, optionally substituted C2-C6-alkenyl, optionally substituted aryl, or optionally substituted heteroaryl. Other examples of pharmaceutically acceptable salts are described in "Remington's Pharmaceutical Sciences", 17th edition. Ed. Alfonso R. Gennaro (Ed.), Mark Publishing Company, Easton, PA, U.S.A., 1985 and more recent editions, and in the Encyclopaedia of Pharmaceutical Technology.

"Treatment" is an approach for obtaining beneficial or desired clinical results. For the purposes of this invention, beneficial or desired clinical results include, but are not limited to, alleviation of symptoms, diminishment of extent of disease, stabilized (i.e., not worsening) state of disease, delay or slowing of disease progression, amelioration or palliation of the disease state, and remission (whether partial or total), whether detectable or undetectable. "Treatment" can also mean prolonging survival as compared to expected survival if not receiving treatment. "Treatment" is an intervention performed with the intention of preventing the development of or altering the pathology of a disorder. Accordingly, "treatment" refers to both therapeutic treatment and prophylactic or preventative measures. Those in need of treatment include those already with the disorder as well as those in which the disorder is to be prevented. By treatment is meant inhibiting or reducing an increase in pathology or symptoms (e.g. weight gain, hyperglycaemia) when compared to the absence of treatment, and is not necessarily meant to imply complete cessation of the relevant condition.

The pharmaceutical compositions can be in unit dosage form. In such form, the composition is divided into unit doses containing appropriate quantities of the active component. The unit dosage form can be a packaged preparation, the package containing discrete quantities of the preparations, for example, packeted tablets, capsules, and powders in vials or ampoules. The unit dosage form can also be a capsule, cachet, or tablet itself, or it can be the appropriate number of any of these packaged forms. It may be provided in single dose injectable form, for example in the form of a pen. Compositions may be formulated for any suitable route and means of administration. Pharmaceutically acceptable carriers or diluents include those used in formulations suitable for oral, rectal, nasal or parenteral (including subcutaneous, intramuscular, intravenous, intradermal, and transdermal) administration. The formulations may conveniently be presented in unit dosage form and may be prepared by any of the methods well known in the art of pharmacy.

Subcutaneous or transdermal modes of administration may be particularly suitable for the compounds described herein.

Combination therapy

The compound of the invention may be administered as part of a combination therapy with an agent for treatment of diabetes, obesity, dyslipidaemia or hypertension.

In such cases, the two active agents may be given together or separately, and as part of the same pharmaceutical formulation or as separate formulations.

Thus the compound of the invention (or the salt thereof) can be used in combination with an antidiabetic agent including but not limited to metformin, a sulfonylurea, a glinide, a DPP-IV inhibitor, a glitazone, or insulin. In a preferred embodiment the compound or salt thereof is used in combination with insulin, DPP-IV inhibitor, sulfonylurea or metformin,
particularly sulfonylurea or metformin, for achieving adequate glycemic control. In an even more preferred embodiment the compound or salt thereof is used in combination with insulin or an insulin analogue for achieving adequate glycemic control. Examples of insulin analogues include but are not limited to Lantus, Novorapid, Humalog, Novomix, Actraphane HM, Levemir and Apidra.

[0131] The compound or salt thereof can further be used in combination with an anti-obesity agent including but not limited to a glucagon-like peptide receptor 1 agonist, peptide YY or analogue thereof, cannabinoid receptor 1 agonist, lipase inhibitor, melanocortin receptor 4 agonist, or melanin concentrating hormone receptor 1 antagonist.

[0132] The compound or salt thereof can be used in combination with an anti-hypertension agent including but not limited to an angiotensin-converting enzyme inhibitor, angiotensin II receptor blocker, diuretics, beta-blocker, or calcium channel blocker.

[0133] The compound or salt thereof can be used in combination with an anti-dyslipidaemia agent including but not limited to a statin, a fibrate, a niacin and/or a cholesterol absorption inhibitor.

METHODS

General synthesis of acylated glucagon analogues

[0134] Solid phase peptide synthesis was performed on a CEM Liberty Peptide Synthesizer using standard Fmoc chemistry. TentaGel S Ram resin (1 g; 0.25 mmol/g) was swelled in NMP (10 ml) prior to use and transferred between tube and reaction vessel using DCM and NMP.

Coupling:

[0135] An Fmoc-amino acid in NMP/DMF/DCM (1:1:1; 0.2 M; 5 ml) was added to the resin in a CEM Discover microwave unit together with HATU/NMP (0.5 M; 2 ml) and DIPEA/NMP (2.0 M; 1 ml). The coupling mixture was heated to 75°C for 5 min while nitrogen was bubbled through the mixture. The resin was then washed with NMP (4 x 10 ml).

Deprotection:

[0136] Piperidine/NMP (20%; 10 ml) was added to the resin for initial deprotection and the mixture was heated by microwaves (30 sec.; 40°C). The reaction vessel was drained and a second portion of piperidine/NMP (20%; 10 ml) was added and heated (75°C; 3 min.) again. The resin was then washed with NMP (6 x 10 ml).

Side chain acylation:

[0137] Fmoc-Lys(ivDde)-OH or alternatively another amino acid with an orthogonal side chain protective group was introduced at the position of the acylation. The N-terminal of the peptide backbone was then Boc-protected using Boc₂O or alternatively by using a Boc-protected amino acid in the last coupling. 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 first coupled with Fmoc-Glu-OtBu or another spacer amino acid, which was deprotected with piperidine and acylated with a lipophilic moiety using the peptide coupling methodology as described above.

Abbreviations employed are as follows:

ivDde: 1-(4,4-dimethyl-2,6-dioxocyclohexylidene)3-methyl-butyl
Dde: 1-(4,4-dimethyl-2,6-dioxocyclohexylidene)-ethyl
DCM: dichloromethane
DMF: N,N-dimethylformamide
DIPEA: diisopropylethylamine
EtOH: ethanol
Et₂O: diethyl ether
HATU: N-[(dimethylamino)-1H-1,2,3-triazol[4,5-b]pyridine-1-ylmethylene]-N-methylmethanaminium hexafluoro-phosphat N-oxide
MeCN: acetonitrile
NMP: N-methylpyrrolidone
TFA: trifluoroacetic acid
TIS: trisopropylsilane
Cleavage:

[0138] The 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 crude peptide was cleaved from the resin by treatment with TFA/TIS/water (95/2.5/2.5; 40 ml, 2 h; r.t.). Most of the TFA was removed at reduced pressure and the crude peptide was precipitated and washed three times with diethylether and dried to constant weight at room temperature.

HPLC purification of the crude peptide:

[0139] The crude peptide was purified to greater than 90% by preparative reverse phase HPLC using a PerSeptive Biosystems VISION Workstation equipped with a C-18 column (5 cm; 10 μm) and a fraction collortor 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.). Fractions were analysed by analytical HPLC and MS and relevant fractions were pooled and lyophilised. The final product was characterised by HPLC and MS.

Generation of cell lines expressing human glucagon- and GLP-1 receptors

[0140] The cDNA encoding either the human glucagon receptor (Glucagon-R) (primary accession number P47871) or the human glucagon-like peptide 1 receptor (GLP-1 R) (primary accession number P43220) were cloned from the cDNA clones BC104854 (MGC:132514/IMAGE:8143857) or BC112126 (MGC:138331/IMAGE:8327994), respectively. The DNA encoding the Glucagon-R or the GLP-1 R was amplified by PCR using primers encoding terminal restriction sites for subcloning. The 5'-end primers additionally encoded a near Kozak consensus sequence to ensure efficient translation. The fidelity of the DNA encoding the Glucagon-R and the GLP-1 R was confirmed by DNA sequencing. The PCR products encoding the Glucagon-R or the GLP-1 R were subcloned into a mammalian expression vector containing a neomycin (G418) resistance marker.

[0141] The mammalian expression vectors encoding the Glucagon-R or the GLP-1 R were transfected into HEK293 cells by a standard calcium phosphate transfection method. 48 hr after transfection cells were seeded for limited dilution cloning and selected with 1 mg/ml G418 in the culture medium. Three weeks later 12 surviving colonies of Glucagon-R and GLP-1 R expressing cells were picked, propagated and tested in the Glucagon-R and GLP-1 R efficacy assays as described below. One Glucagon-R expressing clone and one GLP-1 R expressing clone were chosen for compound profiling.

Glucagon receptor and GLP-1 Receptor efficacy assays

[0142] HEK293 cells expressing the human Glucagon-R, or human GLP-1 R were seeded at 40,000 cells per well in 96-well microtiter plates coated with 0.01 % poly-L-lysine and grown for 1 day in culture in 100 μl growth medium. On the day of analysis, growth medium was removed and the cells washed once with 200 μl Tyrode buffer. Cells were incubated in 100 μl Tyrode buffer containing increasing concentrations of test peptides, 100 μM IBMX, and 6 mM glucose for 15 min at 37° C. The reaction was stopped by addition of 25 μl 0.5 M HCl and incubated on ice for 60 min. The cAMP content was estimated using the FlashPlate® cAMP kit from Perkin-Elmer. EC50 and relative efficacies compared to reference compounds (glucagon and GLP-1) were estimated by computer aided curve fitting.

Bioanalytical screening-method for quantification of peptides Glu-GLP1 agonists in mouse plasma after subcutaneous administration

[0143] Mice were dosed 100 nmol/kg subcutaneously (s.c.). The mice were sacrificed and the blood collected at the following time points; 0.5, 2, 4, 6, 16 and 24 h. Plasma samples were analyzed using protein precipitation, followed by solid phase extraction (SPE) and liquid chromatography mass spectrometry (LC-MS).

Oral Glucose Tolerance Test (OGTT), blood lipids and body weight in high fat fed C57BI/6J normal mice and HbA1c in db/db mice

[0144] Male mice (Long term high fat fed C57BI/6J, short term high fat fed C57BI/6J and db/db) were acclimatized with free access to food and water. They were housed in groups of 5-6 in a light-, temperature-, and humidity-controlled room (12-hour light:12-hour dark cycle, lights On/Off at 2000/0800 hour; 24°C; 50% relative humidity).

[0145] The animals were injected s.c. with 100 μl vehicle (once a day) for a period of three days to acclimatize the animals to handling and injections. Blood samples were taken from the eye or from the tip of the tail. The animals were randomized before treatment.
Mice were treated twice daily s.c. with GluGLP-1 agonist or vehicle (injection volume = 2.5 ml/kg). Throughout the study, body weights were recorded daily and used to administer the body weight-corrected doses of peptide. Peptide solutions were prepared fresh immediately before dosing.

Oral glucose tolerance tests (OGTT) were performed after subjecting the animals to a short fast. To prevent confounding food intake, the animals were kept fasted during the OGTTs. After peptide dosing an initial blood sample was taken. Thereafter an oral dose of glucose (1 g/kg), dissolved in phosphate buffer (pH = 7.4) was given (5 ml/kg), and the animals were returned to their home cages (t = 0). The whole blood glucose (BG) was measured at t=15 min, t=30 min, t=60 min, t=90 min and t=120 min.

The BG concentration was analyzed by the immobilized glucose oxidase method using a drop of blood (< 5 µl; Elite Autoanalyser, Bayer, Denmark) following the manufacturer’s instructions.

HbA1c determination

It is possible to assess the long term effect of a compound on a subject’s glucose level by determining the level of haemoglobin A1C (HbA1c). HbA1c is a glycated form of haemoglobin whose level in a cell reflects the average level of glucose to which the cell has been exposed during its lifetime. In mice, HbA1c is a relevant biomarker for the average blood glucose level during the preceding 4 weeks, because conversion to HbA1c is limited by the erythrocyte’s life span of approximately 47 days (Abbrecht & Littell, 1972; J. Appl. Physiol. 32, 443-445).

The HbA1c determination is based on Turbidimetric Inhibition ImmunoAssay (TINIA) in which HbA1c in the sample reacts with anti-HbA1c to form soluble antigen-antibody complexes. Additions of polyhapten reacts with excess anti-HbA1c antibodies to form an insoluble antibody-polyhapten complex, which can be measured turbidimetrically. Liberated hemoglobin in the hemolyzed sample is converted to a derivative having a characteristic absorption spectrum, which is measured bichromatically during the preincubation phases. The final result is expressed as percent HbA1c of total hemoglobin (Cobas®Application note A1C-2).

Cholesterol level determination

The assay is an enzymatic colorimetric method. In the presence of magnesium ions, dextran sulfate selectively forms water-soluble complexes with LDL, VLDLA and chylomicrons, which are resistant to PEG-modified enzymes. The HDL cholesterol is determined enzymatically by cholesterol esterase and cholesterol oxidase coupled with PEG to the amino groups. Cholesterol esters are broken down quantitatively to free cholesterol and fatty acids. HDL cholesterol is enzymatically oxidized to choles-4-en-3-one and H2O2, and the formed H2O2 is measured colorimetrically (Cobas®; Application note HDLC3).

The direct determination of LDL takes advantage of the selective micellary solubilization of LDL by a nonionic detergent and the interaction of a sugar compound and lipoproteins (VLDL and chylomicrons). The combination of a sugar compound with detergent enables the selective determination of LDL in plasma. The test principle is the same as that of cholesterol and HDL, but due to the sugar and detergent only LDL-cholesterol esters are broken down to free cholesterol and fatty acids. Free cholesterol is then oxidized and the formed H2O2 is measured colorimetrically (Application note LDL_C, Cobas®).

Body weight gain in high fat fed C57BL/6J mice.

C57Bl/6J male mice, 6 weeks old, were acclimatized in their new environment for 4 weeks with free access to high fat diet (HFD) (D12492, Research Diet Inc., New Brunswick, USA) and water. The animals were injected s.c. with 100 µl vehicle for a period of three days to acclimatize the animals to handling and injections, prior to initiation of peptide treatment. The mice were treated twice daily s.c. with exendin-4, Compound 3, Compound 6, Compound 7, Compound 8, Compound 11 and Compound 12 or vehicle. Throughout the study, body weights were recorded daily and used to administer the body weight-corrected doses of peptide. All animals were sacrificed on the same day by cervical dislocation.

Oral glucose tolerance 2, 4, 6, 8, 10 and 12 h after dosing in high fat fed C57BL/6J mice

C57Bl/6J male mice, 6 weeks old, were acclimatized to their new environment with free access to a high fat diet (D12492, Research Diet Inc., New Brunswick, USA) and water. The animals were injected s.c. with vehicle for a period of three days to acclimatize the animals to handling and injections. Blood samples were taken from the tip of the tail and blood glucose measured. The blood glucose (mM) concentration was analyzed by the immobilized glucose oxidase method using a drop of blood (< 5 µl; Contour Autoanalyser, Bayer, Denmark) following the manufacturer’s manual. After 4 weeks on the high fat diet the animals were weighed and the body weight was used to administer a body weight-corrected dose of peptide. An oral glucose tolerance test (OGTT) was performed after subjecting the animals to
4 hours of fasting. At 2, 4, 6, 8, 10 and 12 hours after single peptide or vehicle dosing an initial blood sample were taken (t=0 min). Immediately thereafter, an oral dose of glucose (1 g/kg) was given and the animals were returned to their home cages (t=0). BG levels were measured at t=15 min, t=30 min, t=60 min and t=90 min. Immediately following blood sampling, all animals were sacrificed by CO2 anesthesia followed by cervical dislocation.

Food intake in young lean and old obese C57BL/6J mice.

C57BL/6J mice were on a high fat diet for 11 days and C57BL/6J mice were on a high fat diet for 52 weeks. 3 days before study, the mice were transferred to individual cages and weighed. 4 days before study, they were acclimatized to handling and treatment by daily s.c. injections. On the day before the experiment food was removed at 20:00. On the day of the experiment, the mice were weighed and treated with s.c. injections of Exendin-4, Compound 7 or Vehicle at t=0 h (8:00) and t=12 h (20:00). Immediately after treatment (t=0), pre-weighted food were introduced to the mice and the cumulative food intake was measured by weighing the remaining food after t=1, 2, 4, 8, 12 and 24 hours. After weighing the food and the animals at t=24 h, the mice were sacrificed by cervical dislocation.

Hepatocyte cAMP formation.

Experimental procedure

Primary human hepatocytes provided by Lonza Walkersville, Inc. were carefully washed in TB buffer and incubated at 37°C with peptides dissolved in TB buffer supplemented with 100 μM IBMX and 0.1 % casein for 15 minutes. Prior to addition to the cells, the peptide dilutions were pre-warmed to 37°C. The reaction was stopped by addition of 25 μl of ice cold 0.5 M HCl, and the cells were incubated on ice for 60 min. The cAMP content in the wells was determined by adding 25 μl of the acid extracts from the wells to 75 μl sodium acetate buffer, pH 6.2, in 96-well microtiter "FlashPlates" coated with scintillant and anti-cAMP antibodies. Following addition of 100 μl of 10 μCi [125I]cAMP solution to each well, the plates were incubated overnight at 4°C, emptied, and the amount of [125I]cAMP bound to the FlashPlates was counted using the program "[125I]cAMP flashplate 10 min" on the TopCount NXT.

Peptides were tested at a concentration range of 0.1 - 1000 nM.

Data analysis and statistics

The amount of cAMP produced by the cells was calculated by extrapolation to a cAMP standard curve. EC50 values were estimated by fitting the cAMP data to the below formula using Sigma Plot:

\[
\text{cAMP response} = \frac{(cAMP_{\text{max}} - cAMP_{\text{min}}) \times c}{c + \text{EC}_{50}} + cAMP_{\text{min}},
\]

The invention is further illustrated by the following examples.

Liver Weight/Body weight of C57BL/6J Mice.

Mice were treated twice daily s.c. with Cpd. 1 and Cpd. 11 (at two doses: 0.5 and 5 nmol/kg) or vehicle for 2 weeks. Throughout the study, body weights were recorded daily and used to administer the body weight-corrected doses of peptide. On the day of sacrifice, the liver was exposed, and weighed.

EXAMPLES

Example 1: Synthesis of compounds and peptide properties

Synthesis example:

Compound 9 was synthesized on a CEM Liberty Peptide Synthesizer using TentaGel S Ram resin (1,17 g; 0.23 mmol/g) and Fmoc-chemistry as described above. Fmoc-Lys(ivDde)-OH was used in position 17 and pseudoprolines Fmoc-Phe-Thr(.Psi. Me, Me pro)-OH and Fmoc-Asp(OtBu)-Ser(.Psi., Me, Me pro)-OH were used in the peptide backbone. After completion of the peptide backbone on the resin the N-terminal Fmoc-group was cleaved manually followed by Boc-protection using Boc2O (226 mg) and DIEA (54 μl) in DCM. The ivDde-group was then cleaved with freshly prepared
hydrazine hydrate/NMP (4%; 2 x 15 min.). Back on the CEM Liberty Peptide Synthesizer the remaining two building blocks, Fmoc-Glu-OBu and hexadecanoic acid, were added to the unprotected lysine side chain.

The peptide was cleaved from the resin as described above, and the purification was performed on a Gemini-NX column (5 cm, 10 μm, C18) with a 35 ml/min flow of a mixture of buffer A (0.1% TFA, aq.) and buffer B (0.1% TFA, 90% MeCN, aq.). The product was eluted with a linear gradient from 25% to 65% buffer B over 47 min., and fractions (9 ml) were collected by a fraction collector. Relevant fractions were analysed by analytical HPLC and MS and fractions with purities above 95% were pooled and lyophilised to a white powder. The 72 mg yield had a purity of 97% determined by analytical HPLC and the mass was 3697.05 Da as determined by MS (Calc. 3696.97 Da).

Example 2: Efficacy on GLP-1 and Glucagon receptors

Efficacy of the GluGLP-1 agonists were estimated by exposing cells expressing hGlucagonR and hGLP-1R to the listed acylated compounds at increasing concentrations and measuring the formed cAMP as described in Methods. Results are shown in Table 1:

<table>
<thead>
<tr>
<th>Sequence</th>
<th>Compound</th>
</tr>
</thead>
<tbody>
<tr>
<td>GLP-1R</td>
<td>EC50 (nM)</td>
</tr>
<tr>
<td></td>
<td>EC50 (nM)</td>
</tr>
<tr>
<td>H-HSQGFTDSKYLDKAAHDFVEWLLRA-NH2</td>
<td>Compound 1</td>
</tr>
<tr>
<td>H-HSQGFTDSKYLD-K(Hexadecanoyl-γ-Glu)-KAAHDFVEWLLRA-NH2</td>
<td>Compound 2</td>
</tr>
<tr>
<td>H-HSQGFTDSKYLD-S-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLRA-NH2</td>
<td>Compound 3</td>
</tr>
<tr>
<td>H-HSQGFTDSKYLDKAAHDFVEWLL-K(Hexadecanoyl-γ-Glu)-RA-NH2</td>
<td>Compound 4</td>
</tr>
<tr>
<td>H-HSQGFTDSKYLDKAAHDFVEWLL-K(Hexadecanoyl-γ-Glu)-A-NH2</td>
<td>Compound 5</td>
</tr>
<tr>
<td>H-HSQGFTDSKYLDKAAHDFVEWLL-K(Hexadecanoyl-γ-Glu)-A-NH2</td>
<td>Compound 6</td>
</tr>
<tr>
<td>H-H-Aib-QGTFTDSKYLD-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLSA-NH2</td>
<td>Compound 7</td>
</tr>
<tr>
<td>H-H-Aib-QGTFTDSKYLD-K(Hexadecanoyl-γ-Glu)-AARDFVAWLLRA-NH2</td>
<td>Compound 9</td>
</tr>
<tr>
<td>H-H-Aib-QGTFTDSKYLDKAAHDFVEWLL-K(Hexadecanoyl-γ-Glu)-RA-NH2</td>
<td>Compound 10</td>
</tr>
<tr>
<td>H-H-Aib-QGTFTDSKYLDKAAHDFVEWLL-K(Hexadecanoyl-γ-Glu)-A-NH2</td>
<td>Compound 11</td>
</tr>
<tr>
<td>H-H-Aib-QGTFTDSKYLDKAAHDFVEWLL-K(Hexadecanoyl-γ-Glu)-A-NH2</td>
<td>Compound 12</td>
</tr>
<tr>
<td>H-H-Aib-QGTFTDSKYLDKAAHDFVEWLL-K(Hexadecanoyl-γ-Glu)-A-NH2</td>
<td>Compound 13</td>
</tr>
<tr>
<td>H-H-Aib-QGTFTDSKYLDKAAHDFVEWLL-K(Dodecanoyl-ε-Lys)-AAHDFVE()WLLK()A-NH2</td>
<td>Compound 14</td>
</tr>
<tr>
<td>H-H-Aib-QGTFTDSKYLDKAAHDFVEWLLS-K(Dodecanoyl-γ-Glu)-AAHDFVEWLLSA-NH2</td>
<td>Compound 15</td>
</tr>
<tr>
<td>H-H-Aib-QGTFTDSKYLDKAAHDFVEWLLS-K(Hexadecanoyl-[3-Aminopropanoyl]-AAHDFVEWLLSA-NH2</td>
<td>Compound 16</td>
</tr>
<tr>
<td>H-H-Aib-QGTFTDSKYLDKAAHDFVEWLLS-K(Hexadecanoyl-[8-Aminooctanoyl]-AAHDFVEWLLSA-NH2</td>
<td>Compound 17</td>
</tr>
</tbody>
</table>

The residues marked () form an intramolecular lactam ring.
### Table 1a EC_{50} values of additional acylated compounds according to the invention

<table>
<thead>
<tr>
<th>Sequence</th>
<th>Compound</th>
<th>EC_{50} (nM) GLP-1R</th>
<th>EC_{50} (nM) GluR</th>
</tr>
</thead>
<tbody>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA-OH</td>
<td>Compound 18</td>
<td>0.066</td>
<td>0.091</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLL-OH</td>
<td>Compound 19</td>
<td>0.048</td>
<td>0.483</td>
</tr>
<tr>
<td>H-H-Aib-EGFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA-OH</td>
<td>Compound 20</td>
<td>0.057</td>
<td>13.266</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA-OH</td>
<td>Compound 21</td>
<td>0.077</td>
<td>0.150</td>
</tr>
<tr>
<td>H-H-Aib-EGFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA-NH2</td>
<td>Compound 22</td>
<td>0.014</td>
<td>26.370</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K(Hexadecanoyl)-AAHDFVEWLLSA-NH2</td>
<td>Compound 23</td>
<td>0.140</td>
<td>0.124</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K([2-Butyloctanoyl]-isoGlu)-AAHDFVEWLLSA-NH2</td>
<td>Compound 24</td>
<td>0.161</td>
<td>0.133</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K(Octadecanoyl-isoGlu)-AAHDFVEWLLSA-NH2</td>
<td>Compound 25</td>
<td>0.069</td>
<td>0.103</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K(Octadecanoyl-isoGlu)-AAHDFVEWLLSA-NH2</td>
<td>Compound 26</td>
<td>0.097</td>
<td>0.116</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K(Dodecanoyl-isoGlu)-AAHDFVEWLLSA-NH2</td>
<td>Compound 27</td>
<td>0.152</td>
<td>0.147</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K([4-Aminobutanoyl])-AAHDFVEWLLSA-NH2</td>
<td>Compound 28</td>
<td>0.149</td>
<td>0.108</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K([4-Aminobutanoyl])-AAHDFVEWLLSA-NH2</td>
<td>Compound 29</td>
<td>0.199</td>
<td>0.123</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K([8-Aminooctanoyl])-AAHDFVEWLLSA-NH2</td>
<td>Compound 30</td>
<td>0.132</td>
<td>0.110</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K([4-Aminobutanoyl])-AAHDFVEWLLSA-NH2</td>
<td>Compound 31</td>
<td>0.103</td>
<td>0.151</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-Orn(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA-NH2</td>
<td>Compound 32</td>
<td>0.195</td>
<td>0.193</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYS-K(Hexadecanoyl-isoGlu)-YLDSKAHDFVEWLLSA-NH2</td>
<td>Compound 33</td>
<td>0.131</td>
<td>0.389</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLK-DK(Hexadecanoyl-isoGlu)-KAAHDFVEWLLSA-NH2</td>
<td>Compound 34</td>
<td>0.109</td>
<td>0.053</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDSA-K(Hexadecanoyl-isoGlu)-DFVWLLSA-NH2</td>
<td>Compound 35</td>
<td>0.202</td>
<td>0.180</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDSKAAHDFV-K(Hexadecanoyl-isoGlu)-WLLSA-NH2</td>
<td>Compound 36</td>
<td>0.191</td>
<td>0.213</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDSKAAHDFVEWLL-K(Hexadecanoyl-isoGlu)-A-NH2</td>
<td>Compound 37</td>
<td>0.207</td>
<td>0.147</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K(Hexadecanoyl-isoLys)-AADFVAWLLRA-NH2</td>
<td>Compound 38</td>
<td>0.132</td>
<td>0.183</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAKDFVEWLLSA-NH2</td>
<td>Compound 39</td>
<td>0.16</td>
<td>0.24</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDE-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA-NH2</td>
<td>Compound 40</td>
<td>0.20</td>
<td>0.18</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAHEFVEWLLSA-NH2</td>
<td>Compound 41</td>
<td>0.13</td>
<td>0.08</td>
</tr>
<tr>
<td>H-H-Aib-QGFTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-AAEDFVEWLLSA-NH2</td>
<td>Compound 42</td>
<td>0.03</td>
<td>0.27</td>
</tr>
<tr>
<td>Sequence</td>
<td>Compound 43</td>
<td>EC$_{50}$ (nM) GLP-1R</td>
<td>GLP-1R EC$_{50}$ (nM)</td>
</tr>
<tr>
<td>---------------------------</td>
<td>-------------</td>
<td>-----------------------</td>
<td>-----------------------</td>
</tr>
<tr>
<td>H-H-Ala-O-GFTS-DY-SK-L-D-S-K-Hexadecanoyl iso-Glu(4)AA-HDFWLEA-NH$_2$</td>
<td>Compound 43</td>
<td>0.082</td>
<td>0.12</td>
</tr>
</tbody>
</table>
For compound 28 H-H-Aib-QGTFTSDYSKYLDS-K(Hexadecanoyl-E)-AAHDFVEWLSSA-NH2 could also be written as H-H-Aib-QGTFTSDYSKYLDS-K(Hexadecanoyl-αGlu)-AAHDFVEWLSSA-NH2

Example 3: Pharmacokinetic screening:

Pharmacokinetic profiles were determined for various acylated compounds. Calculated $T_{1/2}$ values are shown in Table 2, compared to (non-acylated) compound 1.

<table>
<thead>
<tr>
<th>Compound</th>
<th>$T_{1/2}(h)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.23</td>
</tr>
<tr>
<td>2</td>
<td>5.8</td>
</tr>
<tr>
<td>5</td>
<td>5.3</td>
</tr>
<tr>
<td>4</td>
<td>2.0*</td>
</tr>
<tr>
<td>6</td>
<td>4.8</td>
</tr>
<tr>
<td>7</td>
<td>3.4</td>
</tr>
<tr>
<td>9</td>
<td>2.4*</td>
</tr>
<tr>
<td>11</td>
<td>4.9</td>
</tr>
<tr>
<td>12</td>
<td>6.0</td>
</tr>
<tr>
<td>13</td>
<td>6.4</td>
</tr>
</tbody>
</table>

*: Only two time points were used for calculation of $T_{1/2}$.

All of the acylated compounds have improved $T_{1/2}$ compared to compound 1.

A sample pharmacokinetic profile, for compound 13, is shown in Figure 1.

Example 4: Oral glucose tolerance test in DIO mice

Effect of 21 days s.c. administration of compound 11 (10 nmol/kg) on oral glucose tolerance in long term high fat-fed C57BL/6J mice. High fat-fed mice were fasted and an initial blood sample taken to determine fasting blood glucose level (t=0). An oral dose of glucose (1 g/kg in 5 ml/kg) was then given and blood glucose levels were measured at t=30 min, t=60 min, t=90 min and t=120 min. Compound 11 significantly improved glucose tolerance (two way ANOVA). Data are shown as mean±SEM.

Example 5: HbA1c in db/db mice after 28 days

Diabetic (db/db) mice were treated with vehicle or compound 7 for 4 weeks, and HbA1c was determined (Cobas® application note: A1 C-2) in whole blood samples (20 μl) collected from the treated mice. Results are shown in Figure 3. The $Δ$HbA1c (%) was calculated for each mice by subtracting its HbA1c (%) at start of treatment from HbA1c (%) at 4 weeks. Treatment with compound 7 decreased $Δ$HbA1c (%) significantly. (P = 0.03; Students t-test) compared to vehicle.

Example 6: Reduced Body weight

Effect of 21 days s.c. administration of compound 11 on body weight was determined in long term high fat-fed C57BL/6J mice. C57BL/6J male mice on high fat diet (HFD) were treated (b.i.d.; s.c.) with compound 11 (10 nmol/kg) or vehicle. Body weights were recorded daily and used to administer the body weight-corrected doses of peptide throughout the study. Data are shown as mean±SEM in Figure 4. Compound 11 significantly decreased body weight (p<0.05).

Example 7: Total Cholesterol and HDL/LDL ratio

Diet Induced Obese (DIO) mice were treated with vehicle or compound 7 for 4 weeks and plasma prepared from the collected blood samples. The total cholesterol, LDL and HDL were determined in each plasma sample (Cobas® application notes: CHOL2, HDLC3 and LDL_C) and results are shown in Figures 5 and 6. Treatment with compound 7 significantly (P <
0.0001, Students t-test) decreased total cholesterol concentrations (Figure 5) and significantly (P < 0.0001, Students t-test) increased the HDL/LDL-ratio (Figure 6).

**Example 8: Body weight gain in high fat fed C57BL/6J mice.**

**[0173]** Effect of 10 days s.c. administration of Exendin-4, Compound 8, Compound 3, Compound 7, Compound 11, Compound 12 and Compound 6 short term high fat-fed C57BL/6J mice. C57Bl/6J male mice on high fat diet (HFD) were treated (b.i.d.; s.c.) (0.5 and 5 nmol/kg) or vehicle. Body weights were recorded daily and used to administer the body weight-corrected doses of peptide throughout the study. Data are shown as mean±SEM in Figure 7.

**[0174]** The control peptide (exendin-4) as well as Compound 8, significantly decreased body weight gain at both doses (0.5 and 5 nmol/kg). Compound 3, Compound 7, Compound 11 and Compound 12 significantly decreased body weight gain at the high dose (5 nmol/kg) but not at the low dose (0.5 nmol/kg) (Fig. 7). Compound 6 significantly decreased body weight gain only at the low dose (0.5 nmol/kg).

**Example 9: Oral glucose tolerance 2, 4, 6, 8, 10 and 12 h after dosing in high fat fed C57BL/6J**

**[0175]** An oral glucose tolerance test (OGTT) was performed after subjecting the animals to 4 hours of fasting. At 2, 4, 6, 8, 10 and 12 hours after Compound 7 or vehicle dosing an initial blood sample was taken (t=0 min). Immediately thereafter, an oral dose of glucose (1 g/kg) was given. BG levels were measured at t=15 min, t=30 min, t=60 min and t=90 min. Immediately following blood sampling, all animals were sacrificed by CO2 anesthesia followed by cervical dislocation.

The study shows that subcutaneous administration with Compound 7 (10 nmol/kg) significantly improves glucose tolerance (measured as decreased AUC during an oral glucose tolerance test) 2, 4, 6, 8, 10 and 12 hours after dosing in high fat fed C57BL/6J mice.

**Example 10: Food intake in young lean and old obese C57BL/6J mice.**

**[0176]** C57BL/6J mice were on a high fat diet for 11 days and C57BL/6J mice were on a high fat diet for 52 weeks. On the day of the experiment, the mice were weighed and treated with s.c. injections of Exendin-4, Compound 7 or Vehicle at t=0 h (8:00) and t=12 h (20:00). Immediately after treatment (t=0), preweighed food were introduced to the mice and the cumulative food intake was measured by weighing the remaining food after t=1, 2, 4, 8, 12 and 24 hours.

In the young lean mice, Compound 7 statistically significantly (p<0.05) reduced food intake during the 0-4, 0-8, 0-12 and 0-24 time periods. Exendin-4 statistically significantly (p<0.05) reduced food intake during the 0-2, 0-4, 0-8, 0-12 and 0-24 time periods.

In the old obese mice, Compound 7 statistically significantly (p<0.05) reduced food intake during the 0-2, 0-4, 0-8, 0-12 and 0-24 time periods. Exendin-4 statistically significantly (p<0.05) reduced food intake in all time periods.

**Example 11: Effect of 3 weeks subcutaneous administration of GluGLP-1 agonist Compound 11 on lipids in 30 weeks High Fat Diet fed mice.**

**[0177]** Effect of 3 weeks treatment of mice that have been on 30 weeks High Fat Diet for 30 weeks prior treatment (s.c.) with vehicle (PBS), 10 nmol/kg exendin-4 or 10 nmol/kg Compound 11 twice daily for 3 weeks on lipids (Figure 11). The effect was measured on LDL, HDL and triglycerids (CHO: Total Cholesterol; HDL: High Density Cholesterol; LDL: Low Density Cholesterol; TRIG: Triglycerides; HDL/LDL: Ratio between HDL and LDL).

**[0178]** Compound 11 significantly decreased cholesterol, HDL, LDL (P < 0.001) and triglycerides (P < 0.05) significantly, while the ratio HDL/LDL was increased significantly (p < 0.001) (Fig. 11). The HDL/LDL ratio is considered a risk indicator for heart disease. The higher the ratio, the lower the risk of heart attack or other cardiovascular problems.

**Example 12: Effect of Compound 11 on Hepatocyt cAMP formation.**

**[0179]** All tested peptides behaved as full agonist with respect to GluR stimulated cAMP formation except of the pure GLP-1 agonists exendin-4 and liraglutide. From the table it can observed that the rank order of potency is: Compound 1 > glucagon > Compound 11 > oxyntomodulin >>> exendin-4 and liraglutide (Table 9).

**[0180]** Finally, no down regulation was observed of the E\textsubscript{MAX} cAMP response at the high concentrations, which is in contrast to what is observed in the hGluR HEK293 cells.
Example 13: Liver Weight of C57 Healthy Control Mice Treated for 2 Weeks

Repeated treatment with long-acting acylated dual GluGLP-1 agonists such as Compound 11 do not give rise to change in liver size (enlarged or shrunken) compared with the non-acylated dual GluGLP-1 agonists compound 1 (Figure 12).

Example 14: HbA1c in db/db mice after 28 days

Diabetic (db/db) mice were treated with vehicle or compound 11 for 4 weeks, and HbA1c was determined (Cobas® application note: A1C-2) in whole blood samples (20 μl) collected from the treated mice. Results are shown in Figure 13. The ΔHbA1c (%) was calculated for each mice by subtracting its HbA1c (%) at start of treatment from HbA1c (%) at 4 weeks. Treatment with compound 11 decreased ΔHbA1c (%) significantly. (P = 0.03; Students t-test) compared to vehicle.

Claims

1. A compound having the formula:

   \[ R_1^2 \cdot Z \cdot R_2 \]

   wherein \( R_1 \) is H, C1-4 alkyl, acetyl, formyl, benzoyl or trifluoroacetyl;
   \( R_2 \) is OH or NH2;
   and \( Z \) is a peptide having the formula (I)

   \[
   \text{His-X2-Gln-Gly-Thr-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-X16-X17-Ala-Ala-X20-X21-Phe-Val-X24-Trp-Leu-X27-X28-Ala-X30; (I)}
   \]

   wherein

   X2 is selected from Aib and Ser;
   X12 is selected from Lys, Arg or Leu;
   X16 is selected from Arg and X;
   X17 is selected from Arg and X;
   X20 is selected from Arg, His and X;
   X21 is selected from Asp and Glu;
   X24 is selected from Ala and X;
   X27 is selected from Leu and X;
X28 is selected from Arg and X;
X30 is X or is absent;
wherein at least one of X16, X17, X20, X24, X27, X28, and X30 is X;

and wherein each residue X is independently selected from the group consisting of Glu, Lys, Ser, Cys, Dbu, Dpr and Orn;
wherein the side chain of at least one residue X is conjugated to a lipophilic substituent having the formula:

(i) Z1, wherein Z1 is a lipophilic moiety conjugated directly to the side chain of X; or
(ii) Z1Z2, wherein Z1 is a lipophilic moiety, Z2 is a spacer, and Z1 is conjugated to the side chain of X via Z2;

with the proviso that Z is not HSQGTFTSDYSKLYDS-K(Hexadecanoyl-\textgamma-Glu)-AAHDFVEWLLRA.

2. A compound according to claim 1 wherein:

(a) one or more of said residues X is independently selected from Lys, Glu and Cys; and/or
(b) X16 is selected from Glu, Lys and Ser;
X17 is selected from Lys and Cys;
X20 is selected from His, Lys, Arg and Cys;
X24 is selected from Lys, Glu and Ala;
X27 is selected from Leu and Lys; and/or
X28 is selected from Ser, Arg and Lys; and/or
(c) the peptide of formula I includes one or more of the following combinations of residues:

X2 is Aib and X17 is Lys;
X2 is Aib and X20 is Cys;
X2 is Aib and X28 is Lys;
X12 is Arg and X17 is Lys;
X12 is Leu and X17 is Lys;
X12 is Lys and X20 is Lys;
X12 is Lys and X17 is Lys;
X16 is Lys and X17 is Lys;
X16 is Ser and X17 is Lys;
X16 is Lys and X20 is Lys;
X17 is Lys and X21 is Asp;
X17 is Lys and X24 is Glu;
X17 is Lys and X27 is Leu;
X17 is Lys and X27 is Lys;
X17 is Lys and X28 is Ser;
X17 is Lys and X28 is Arg;
X20 is Lys and X27 is Leu;
X21 is Asp and X27 is Leu;
X2 is Aib, X12 is Lys and X16 is Ser;
X12 is Lys, X17 is Lys and X16 is Ser;
X12 is Arg, X17 is Lys and X16 is Glu;
X16 is Glu, X17 is Lys and X20 is Lys;
X16 is Ser, X21 is Asp and X24 is Glu;
X17 is Lys, X24 is Glu and X28 is Ser;
X17 is Lys, X24 is Glu and X28 is Arg;
X17 is Lys, X27 is Leu and X28 is Ser;
X17 is Lys, X27 is Leu and X28 is Arg;
X20 is Lys, X24 is Glu and X27 is Leu;
X20 is Lys, X27 is Leu and X28 is Ser;
X20 is Lys, X27 is Leu and X28 is Arg;
X16 is Ser, X20 is His, X24 is Glu and X27 is Leu;
X17 is Lys, X20 is His, X24 is Glu and X28 is Ser;
X17 is Lys, X20 is Lys, X24 is Glu and X27 is Leu; or
X17 is Cys, X20 is Lys, X24 is Glu and X27 is Leu.

3. A compound according to claim 1 or claim 2 wherein the peptide of formula I contains only one amino acid of the type conjugated to the lipophilic substituent, e.g. wherein the peptide contains only one Lys residue, only one Cys residue or only one Glu residue, and wherein the lipophilic substituent is conjugated to that residue.

4. A compound according to any one of the preceding claims wherein the peptide sequence of formula I comprises one or more intramolecular bridges, and wherein optionally

(a) said intramolecular bridge is formed between the side chains of two amino acid residues which are separated by three amino acids in the linear amino acid sequence of formula I;
(b) the intramolecular bridge is formed between the side chains of residue pairs 16 and 20, 17 and 21, 20 and 24, or 24 and 28 (c) the intramolecular bridge is a salt bridge or a lactam ring; and/or
(d) the intramolecular bridge involves a pair of residues wherein:

X16 is Glu and X20 is Lys;
X16 is Glu and X20 is Arg;
X16 is Lys and X20 is Glu; or
X16 is Arg and X20 is Glu;
X17 is Arg and X21 is Glu;
X17 is Lys and X21 is Glu;
X17 is Arg and X21 is Asp; or
X17 is Lys and X21 is Asp;
X20 is Glu and X24 is Lys;
X20 is Glu and X24 is Arg;
X20 is Lys and X24 is Glu; or
X20 is Arg and X24 is Glu;
X24 is Glu and X28 is Lys;
X24 is Glu and X28 is Arg;
X24 is Lys and X28 is Glu; or
X24 is Arg and X28 is Glu.

5. A compound according to any one of the preceding claims wherein at least one of X16, X17, X20 and X28 is conjugated to a lipophilic substituent.

6. A compound according to any one of the preceding claims wherein:

(a) X30 is absent; or
(b) X30 is present and is conjugated to a lipophilic substituent.

7. A compound according to any one of the preceding claims wherein:

(a) the compound has just one lipophilic substituent, at position 16, 17, 20, 24, 27, 28 or 30, preferably at position 16, 17 or 20, particularly at position 17;
(b) the compound has precisely two lipophilic substituents, each at one of positions 16, 17, 20, 24, 27, 28 or 30; or
(c) the compound has lipophilic substituents at positions 16 and 17, 16 and 20, 16 and 24, 16 and 27, 16 and 28 or 16 and 30; at 17 and 20, 17 and 24, 17 and 27, 17 and 28 or 17 and 30; at 20 and 24, 20 and 27, 20 and 28 or 20 and 30; at 24 and 27, 24 and 28 or 24 and 30; at 27 and 28 or 27 and 30; or at 28 and 30.

8. A compound according to claim 1, having the formula:

\[ R^1-Z-R^2 \]

wherein \( R^1 \) is H, C1-4 alkyl, acetyl, formyl, benzoyl or trifluoroacetyl;
\( R^2 \) is OH or NH2;
and Z is a peptide having:
(a) the formula IIa:

\[
\text{His-Aib-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X}^{12}\text{-Tyr-Leu-Asp-X}^{16}\text{-X}^{17}\text{-Ala-Ala-X}^{20}\text{-X}^{21}\text{-Phe-Val-X}^{24}\text{-Trp-Leu-Leu-X}^{28}\text{-Ala};
\]  

\(\text{(IIa)}\)

wherein

- \(X^{12}\) is selected from Lys, Arg and Leu;
- \(X^{16}\) is selected from Ser and X;
- \(X^{17}\) is X;
- \(X^{20}\) is selected from His and X;
- \(X^{21}\) is selected from Asp and Glu;
- \(X^{24}\) is selected from Ala and Glu;
- \(X^{28}\) is selected from Ser, Lys and Arg;

and wherein each residue X is independently selected from the group consisting of Glu, Lys, and Cys;

(b) the formula IIIa:

\[
\text{His-Aib-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X}^{12}\text{-Tyr-Leu-Asp-Ser-X}^{17}\text{-Ala-Ala-X}^{20}\text{-X}^{21}\text{-Phe-Val-X}^{24}\text{-Trp-Leu-Leu-X}^{28}\text{-Ala};
\]  

\(\text{(IIIa)}\)

wherein

- \(X^{12}\) is selected from Lys and Arg;
- \(X^{17}\) is X;
- \(X^{20}\) is selected from His and X;
- \(X^{21}\) is selected from Asp and Glu;
- \(X^{24}\) is selected from Ala and Glu;
- \(X^{28}\) is selected from Ser, Lys and Arg;

and wherein each residue X is independently selected from Glu, Lys, and Cys; or

(c) the formula IVa:

\[
\text{His-Aib-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X}^{12}\text{-Tyr-Leu-Asp-Ser-X}^{17}\text{-Ala-Ala-His-X}^{21}\text{-Phe-Val-X}^{24}\text{-Trp-Leu-Leu-X}^{28}\text{-Ala};
\]  

\(\text{(IVa)}\)

wherein

- \(X^{12}\) is selected from Lys and Arg;
- \(X^{17}\) is X;
- \(X^{21}\) is selected from Asp and Glu;
- \(X^{24}\) is selected from Ala and Glu;
- \(X^{28}\) is selected from Ser, Lys and Arg;

wherein X is selected from the group consisting of Glu, Lys and Cys;

wherein the side chain of at least one residue X is conjugated to a lipophilic substituent having the formula:

- (i) \(Z^{1}\), wherein \(Z^{1}\) is a lipophilic moiety conjugated directly to the side chain of X; or
- (ii) \(Z^{1}\)\(Z^{2}\), wherein \(Z^{1}\) is a lipophilic moiety, \(Z^{2}\) is a spacer, and \(Z^{1}\) is conjugated to the side chain of X via \(Z^{2}\).

9. A compound according to claim 1 having the formula:

\[R^{1}\cdot Z\cdot R^{2}\]

wherein \(R^{1}\) is H, C\(_{1-4}\) alkyl, acetyl, formyl, benzoyl or trifluoroacetyl;
\(R^{2}\) is OH or NH\(_{2}\);
and Z is a peptide having:
(a) the formula Iib:

\[
\text{His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-X16-X17-Ala-Ala-X20-X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala; (IIb)}
\]

wherein

X12 is selected from Lys, Arg and Leu;
X16 is selected from Ser and X;
X17 is X;
X20 is selected from His and X;
X21 is selected from Asp and Glu;
X24 is selected from Ala and Glu;
X28 is selected from Ser, Lys and Arg;
and wherein each residue X is independently selected from the group consisting of Glu, Lys, and Cys;

(b) the formula IIIb:

\[
\text{His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X17-Ala-Ala-X20-X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala; (IIIb)}
\]

wherein

X12 is selected from Lys and Arg;
X17 is X;
X20 is selected from His and X;
X21 is selected from Asp and Glu;
X24 is selected from Ala and Glu;
X28 is selected from Ser, Lys and Arg;
and wherein each residue X is independently selected from Glu, Lys or Cys; or

(c) the formula IVb:

\[
\text{His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X17-Ala-Ala-His-X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala; (IVb)}
\]

wherein

X12 is selected from Lys and Arg;
X17 is X;
X20 is selected from His and X;
X21 is selected from Asp and Glu;
X24 is selected from Ala and Glu;
X28 is selected from Ser, Lys and Arg;
wherein X is selected from the group consisting of Glu, Lys and Cys;

wherein the side chain of at least one residue X is conjugated to a lipophilic substituent having the formula:

(i) Z1, wherein Z1 is a lipophilic moiety conjugated directly to the side chain of X; or
(ii) Z1Z2, wherein Z1 is a lipophilic moiety, Z2 is a spacer, and Z1 is conjugated to the side chain of X via Z2;

with the proviso that Z is not HSQGTFTSDYSKYLDS-K(Hexadecanoyl-γ-Glu))-AAHDFVEWLLRA.

10. A compound according to any one of the preceding claims wherein the peptide of formula I, Ila, IIa, IVa, Iib, IIib or IVb has the sequence:

HSQGTFTSDYSKYLDSKAAHDFVEWLLRA;
HSQGTFTSDYSKYLDSKAKAHEDFVEWLLRA;
HSQGTFTSDYSKYLDSKAAKDFVEWLLRA;
HSQGTFTSDYKLDSDKAAHDFVEWLLRA;
HSQGTFTSDYSKLDSDKAAHDFVEWLLLA;
HSQGTFTSDYSRLYLDSKAAHDFVEWLLRA;
HSQGTFTSDYSLYLDSDKAAHDFVEWLLRA;
5
HSQGTFTSDYKLDSDKAAHDFVEWLLRAK;
HSQGTFTSDYSKLDSDKAAHDFVEWLLSAK;
HSQGTFTSDYSKLDSDKAAHDFVEWLLSA;
HSQGTFTSDYSKLDSDKAAHDFVKKWLLRA;
10
HSQGTFTSDYSKLDSDKAAHDFVEWLLLA;
HSQGTFTSDYSKLDSDKAAHDFVEWLLSA;
HSQGTFTSDYKLDSDKAAHDFVEWLLRA;
HSQGTFTSDYSKLDSDKAAHDFVEWLLSA;
15
H-Aib-QGTFTSDYKLDSDKAAHDFVEWLLLA;
H-Aib-QGTFTSDYKLDSDKAAHDFVEWLLSA;
H-Aib-QGTFTSDYKLDSDKAAHDFVEWLLRA;
H-Aib-QGTFTSDYKLDSDKAAHDFVEWLLKA;
20
H-Aib-QGTFTSDYKLDSDKAAHDFVEWLLSA;
H-Aib-QGTFTSDYKLDSDKAAHDFVEWLLRA;
H-Aib-QGTFTSDYKLDSDKAAHDFVEWLLKA;
H-Aib-QGTFTSDYKLDSDKAAHDFVEWLLLA;
25
H-Aib-QGTFTSDYKLDSDKAAHDFVEWLLSA;
H-Aib-QGTFTSDYKLDSDKAAHDFVEWLLLA;
H-Aib-QGTFTSDYKLDSDKAAHDFVEWLLRA;
H-Aib-QGTFTSDYKLDSDKAAHDFVEWLLKA;
30
11. A compound according to any one of the preceding claims wherein:

(a) $Z_1$ comprises a hydrocarbon chain having 10 to 24 C atoms, 10 to 22 C atoms, or 10 to 20 C atoms, e.g. wherein $Z_1$ is a dodecanoyl, 2-butyloctanoyl, tetradecanoyl, hexadecanoyl, heptadecanoyl, octadecanoyl or eicosanoyl moiety;

(b) $Z_2$ is or comprises one or more amino acid residues, e.g. wherein $Z_2$ is a $\gamma$-Glu, Glu, $\beta$-Ala or $\varepsilon$-Lys residue, or a 3-aminopropanoyl, 4-aminobutanoyl, 8-aminooctanoyl or 8-amino-3,6-dioxo-octanoyl moiety; and/or

(c) wherein the lipophilic substituent is selected from the group consisting of dodecanoyl-$\gamma$-Glu, hexadecanoyl-$\gamma$-Glu, hexadecanoyl-$[3$-aminopropanoyl], hexadecanoyl-$[8$-aminooctanoyl], hexadecanoyl-$\varepsilon$-Lys, 2-butyloctanoyl-$\gamma$-Glu, octadecanoyl-$\gamma$-Glu and hexadecanoyl-$[4$-aminobutanoyl].

12. A compound according to claim 11 wherein $Z$ has the formula:

$$\text{HSQGTFTSDYS}	ext{K}(\text{Hexadecanoyl-$\gamma$-Glu})\text{-KAAHDFVEWLLSA};$$

$$\text{HSQGTFTSDYSKYLDS}	ext{K}(\text{Hexadecanoyl-$\gamma$-Glu})\text{-KAAHDFVEWLLSA};$$

$$\text{HSQGTFTSDYSKYLDSKAA}	ext{-C$^*_*$DFVEWLLRA};$$

$$\text{H-Aib-QGTFTSDYSKYLDSKAA-C$^*_*$DFVEWLLRA};$$

$$\text{H-Aib-QGTFTSDYSKYLDS}-2\text{AAHDFVEWLLSA};$$

$$\text{H-Aib-QGTFTSDYSKYLDS}-2\text{AAHDFVEWLLSA};$$

wherein "**" indicates the position of a lipophilic substituent.

wherein $Z$ has the formula:

$$\text{H-Aib-QGTFTSDYS-(KHexadecanoyl-isoGlu)-YLDSDKAAHDFVEWLLSA};$$

$$\text{H-Aib-QGTFTSDYS KYLD-(KHexadecanoyl-isoGlu)-KAAHDFVEWLLSA};$$
13. A compound according to claim 1 which is

H-Alb-QGFTTSDYSKYLDS-K(Hexadecanoyl-isoGlu)-DFVEWLLSA.

14. A compound having the formula:

R¹-Z-R²

wherein R¹ is H, C₁₋₄ alkyl, acetyl, formyl, benzyol or trifluoroacetyl;
R² is OH or NH₂;
and Z is a peptide having:

(a) the formula V:

His-Alb-Gln-Gly-Thr-Phe-Thr-Ser-Tyr-Ser-Lys-Tyr-Leu-Asp-Ser-X₁⁷-Ala-Ala-His-Asp-Phe-Val-Glu-Trp-Leu-Leu-X₂⁸; (V)

wherein

X₁⁷ is X
X₂⁸ is Ser or absent; or

(b) the formula VI:

His-Alb-Glu-Gly-Thr-Phe-Thr-Ser-Tyr-Ser-Lys-Tyr-Leu-Asp-Ser-X₁⁷-Ala-Ala-His-Asp-Phe-Val-Glu-Trp-Leu-Leu-Ser-Ala; (VI)

wherein

X₁⁷ is X;

wherein X is selected from the group consisting of Glu, Lys, and Cys;
and wherein the side chain of X is conjugated to a lipophilic substituent having the formula:

(i) Z¹, wherein Z¹ is a lipophilic moiety conjugated directly to the side chain of X; or
(ii) Z¹Z², wherein Z¹ is a lipophilic moiety, Z² is a spacer, and Z¹ is conjugated to the side chain of X via Z²;

15. A compound according to claim 14 wherein Z has the formula:

H-Alb-QGFTTSDYKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA;
H-Alb-QGFTTSDYKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLS; or
H-Alb-EGFTTSDYKYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA.

16. A composition comprising a compound according to any one of claims 1 to 15, or a salt or derivative thereof, in admixture with a carrier, e.g. wherein the composition is a pharmaceutically acceptable composition and the carrier is a pharmaceutically acceptable carrier.

17. A compound according to any one of claims 1 to 15 for use in a method of medical treatment, e.g.

(a) for use in preventing weight gain or promoting weight loss;
(b) for use in a method of improving circulating glucose levels, glucose tolerance and/or circulating cholesterol levels, lowering circulating LDL levels, and/or increasing HDL/LDL ratio; or
(c) for use in a method of treatment of a condition caused or characterised by excess body weight, e.g. the treatment and/or prevention of obesity, morbid obesity, obesity linked inflammation, obesity linked gallbladder disease, obesity induced sleep apnea, metabolic syndrome, pre-diabetes, insulin resistance, glucose intolerance, type 2 diabetes, type I diabetes, hypertension, atherogenic dyslipidaemia, atherosclerosis, arteriosclerosis, coronary heart disease, peripheral artery disease, stroke or microvascular disease.

18. A compound for use according to claim 17 wherein the compound is for administration as part of a combination therapy with an agent for treatment of diabetes, obesity, dyslipidaemia, or hypertension.

19. A compound for use according to claim 18 wherein the agent for treatment of diabetes is metformin, a sulphonylurea, a glinide, a DPP-IV inhibitor, a glitazone, insulin or an insulin analogue.

20. A compound for use according to claim 18 wherein the agent for treatment of obesity is a glucagon-like peptide receptor 1 agonist, peptide YY or analogue thereof, cannabinoid receptor 1 antagonist, lipase inhibitor, melanocortin receptor 4 agonist, or melanin concentrating hormone receptor 1 antagonist.

21. A compound for use according to claim 18 wherein the agent for treatment of hypertension is an angiotensin-converting enzyme inhibitor, angiotensin II receptor blocker, diuretic, beta-blocker, or calcium channel blocker.

22. A compound for use according to claim 18 wherein the agent for treatment of dyslipidaemia is a statin, a fibrate, a niacin and/or a cholesterol absorption inhibitor.

23. Use of a compound according to any one of claims 1 to 15 in the preparation of a medicament for:

(a) preventing weight gain or promoting weight loss in an individual in need thereof;
(b) improving circulating glucose levels, glucose tolerance and/or circulating cholesterol levels, lowering circulating LDL levels, and/or increasing HDULDL ratio in an individual in need thereof; or
(c) treatment of a condition caused or characterised by excess body weight, e.g. the treatment and/or prevention of obesity, morbid obesity, obesity linked inflammation, obesity linked gallbladder disease, obesity induced sleep apnea, pre-diabetes, insulin resistance, glucose intolerance, type 2 diabetes, type I diabetes, hypertension, atherogenic dyslipidimia, atherosclerosis, arteriosclerosis, coronary heart disease, peripheral artery disease, stroke or microvascular disease in an individual in need thereof.

24. Use according to claim 23 wherein the compound is for administration as part of a combination therapy with an agent for treatment of diabetes, obesity, dyslipidaemia, or hypertension, e.g.

(a) wherein the agent for treatment of diabetes is metformin, a sulphonylurea, a glinide, a DPP-IV inhibitor, a glitazone, insulin or an insulin analogue;
(b) wherein the agent for treatment of obesity is a glucagon-like peptide receptor 1 agonist, peptide YY or analogue thereof, cannabinoid receptor 1 antagonist, lipase inhibitor, melanocortin receptor 4 agonist, or melanin concentrating hormone receptor 1 antagonist;
(c) wherein the agent for treatment of hypertension is an angiotensin-converting enzyme inhibitor, angiotensin II receptor blocker, diuretic, beta-blocker, or calcium channel blocker; or
(d) wherein the agent for treatment of dyslipidaemia is a statin, a fibrate, a niacin and/or a cholesterol absorption inhibitor.

Patentansprüche

1. Verbindung der Formel:

\[ R^1-Z-R^2, \]

worin \( R^1 \) H, C_{1-4}-Alkyl, Acetyl, Formyl, Benzoyl oder Trifluoracetyl ist; \( R^2 \) OH oder \( NH_2 \) ist; und Z ein Peptid der Formel I ist:
His-X2-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-X16-X17-Ala-Ala-X20-X21-Phe-Val-X24-Trp-Leu-X27-X28-Ala-X30; (I)

worin

- X2 aus Aib und Ser ausgewählt ist;
- X12 aus Lys, Arg und Leu ausgewählt ist;
- X16 aus Arg und X ausgewählt ist;
- X17 aus Arg und X ausgewählt ist;
- X20 aus Arg, His und X ausgewählt ist;
- X21 aus Asp und Glu ausgewählt ist;
- X24 aus Ala und X ausgewählt ist;
- X27 aus Leu und X ausgewählt ist;
- X28 aus Arg und X ausgewählt ist;
- X30 X ist oder fehlt;

worin zumindest eines von X16, X17, X20, X24, X27, X28 und X30 X ist;

und worin jeder Rest X jeweils unabhängig aus der aus Glu, Lys, Ser, Cys, Dbu, Dpr und Orn bestehenden Gruppe ausgewählt ist;

worin die Seitenkette von zumindest einem Rest X an einen lipophilen Substituenten der folgenden Formel konjugiert ist:

(i) Z1, worin Z1 eine direkt an die Seitenkette von X konjugierte lipophile Gruppierung ist; oder
(ii) Z1Z2, worin Z1 eine lipophile Gruppierung ist, Z2 ein Spacer ist und Z1 über Z2 an die Seitenkette von X konjugiert ist;

mit der Maßgabe, dass Z nicht HSQGTFSTSDFYSKYLDS-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLRA ist.

2. Verbindung nach Anspruch 1, worin:

(a) einer oder mehrere der Reste X jeweils unabhängig aus Lys, Glu und Cys ausgewählt sind; und/oder
(b) X16 aus Glu, Lys und Ser ausgewählt ist;

- X17 aus Lys und Cys ausgewählt ist;
- X20 aus His, Lys, Arg und Cys ausgewählt ist;
- X24 aus Lys, Glu und Ala ausgewählt ist;
- X27 aus Leu und Lys ausgewählt ist; und/oder
- X28 aus Ser, Arg und Lys ausgewählt ist; und/oder

(c) das Peptid der Formel I eine oder mehrere der folgenden Kombinationen von Resten umfasst:

- X2 ist Aib und X17 ist Lys;
- X2 ist Aib und X17 ist Cys;
- X2 ist Aib und X20 ist Cys;
- X2 ist Aib und X28 ist Lys;
- X12 ist Arg und X17 ist Lys;
- X12 ist Leu und X17 ist Lys;
- X12 ist Lys und X20 ist Lys;
- X12 ist Lys und X17 ist Lys;
- X16 ist Lys und X17 ist Lys;
- X16 ist Ser und X17 ist Lys;
- X17 ist Lys und X20 ist Lys;
- X17 ist Lys und X21 ist Asp;
- X17 ist Lys und X24 ist Glu;
- X17 ist Lys und X27 ist Leu;
- X17 ist Lys und X27 ist Lys;
- X17 ist Lys und X28 ist Ser;
- X17 ist Lys und X28 ist Arg;
X20 ist Lys und X27 ist Leu;
X21 ist Asp und X27 ist Leu;
X2 ist Alb, X12 ist Lys und X16 ist Ser;
X12 ist Lys, X17 ist Lys und X16 ist Ser;
X12 ist Arg, X17 ist Lys und X16 ist Glu;
X16 ist Glu, X17 ist Lys und X20 ist Lys;
X16 ist Ser, X21 ist Asp und X24 ist Glu;
X17 ist Lys, X24 ist Glu und X28 ist Arg;
X17 ist Lys, X24 ist Glu und X28 ist Lys;
X17 ist Lys, X27 ist Leu und X28 ist Ser;
X17 ist Lys, X27 ist Leu und X28 ist Arg;
X20 ist Lys, X24 ist Glu und X27 ist Leu;
X20 ist Lys, X27 ist Leu und X28 ist Ser;
X16 ist Ser, X20 ist His, X24 ist Glu und X27 ist Leu;
X17 ist Lys, X20 ist His, X24 ist Glu und X28 ist Ser;
X17 ist Lys, X20 ist His, X24 ist Glu und X27 ist Leu; oder
X17 ist Cys, X20 ist His, X24 ist Glu und X27 ist Leu.

3. Verbindung nach Anspruch 1 oder Anspruch 2, worin das Peptid der Formel I nur eine Aminosäure der an den lipophilen Substituenten konjugierten Art enthält, z.B. worin das Peptid nur einen Lys-Rest, nur einen Cys-Rest oder nur einen Glu-Rest enthält, und worin der lipophile Substituent an diesen Rest konjugiert ist.

4. Verbindung nach einem der vorangegangenen Ansprüche, worin die Peptidsequenz der Formel I eine oder mehrere intramolekulare Brücken umfasst und worin gegebenenfalls

(a) die intramolekulare Brücke zwischen den Seitenketten zweier Aminosäurereste ausgebildet ist, die in der linearen Aminosäuresequenz der Formel I durch drei Aminosäuren getrennt sind;
(b) die intramolekulare Brücke zwischen den Seitenketten der Restepaare 16 und 20, 17 und 21, 20 und 24 oder 24 und 28 ausgebildet ist,
(c) die intramolekulare Brücke eine Salzbrücke oder ein Lactamring ist; und/oder
(d) an der intramolekularen Brücke ein Paar von Resten beteiligt ist, worin:

X16 Glu ist und X20 Lys ist;
X16 Glu ist und X20 Arg ist;
X16 Lys ist und X20 Glu ist; oder
X16 Arg ist und X20 Glu ist;
X17 Arg ist und X21 Glu ist;
X17 lys ist und X21 Glu ist;
X17 Arg ist und X21 Asp ist; oder
X17 Lys ist und X21 Asp ist;
X20 Glu ist und X24 Lys ist;
X20 Glu ist und X24 Arg ist;
X20 Lys ist und X24 Glu ist; oder
X20 Arg ist und X24 Glu ist;
X24 Glu ist und X28 Lys ist;
X24 Glu ist und X28 Arg ist;
X24 Lys ist und X28 Glu ist; oder
X24 Arg ist und X28 Glu ist.

5. Verbindung nach einem der vorangegangenen Ansprüche, worin zumindest einer von X16, X17, X20 und X28 an einen lipophilen Substituenten konjugiert ist.

6. Verbindung nach einem der vorangegangenen Ansprüche, worin:

(a) X30 fehlt; oder
(b) X30 vorhanden und an einen lipophilen Substituenten konjugiert ist.
7. Verbindung nach einem der vorangegangenen Ansprüche, worin:

(a) die Verbindung nur einen lipophilen Substituenten an Position 16, 17, 20, 24, 27, 28 oder 30, vorzugsweise an Position 16, 17 oder 20, insbesondere an Position 17, aufweist;
(b) die Verbindung genau zwei lipophile Substituenten, je einen an den Positionen 16, 17, 20, 24, 27, 28 oder 30 aufweist; oder
(c) die Verbindung lipophile Substituenten an den Positionen 16 und 17, 16 und 20, 16 und 24, 16 und 27, 16 und 28 oder 16 und 30; an 17 und 20, 17 und 24, 17 und 27, 17 und 28 oder 17 und 30; an 20 und 24, 20 und 27, 20 und 28 oder 20 und 30; an 24 und 27, 24 und 28 oder 24 und 30; an 27 und 28 oder 27 und 30; oder an 28 und 30 aufweist.

8. Verbindung nach Anspruch 1 der Formel:

\[ R^1 \cdot Z \cdot R^2, \]

worin \( R^1 \) H, C\(_{1-4}\)-Alkyl, Acetyl, Formyl, Benzoyl oder Trifluoracetyl ist;
\( R^2 \) OH oder NH\(_2\) ist;
und \( Z \) ein Peptid ist, das:

(a) der Formel Ilia entspricht:

\[ \text{His-Alb-Gln-Gly-Thr-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-X16-X17-Ala-Ala-X20-X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala:} \quad (\text{Ilia}) \]

worin

- X\(_{12}\) aus Lys, Arg und Leu ausgewählt ist;
- X\(_{16}\) aus Ser und X ausgewählt ist;
- X\(_{17}\) X ist;
- X\(_{20}\) aus His und X ausgewählt ist;
- X\(_{21}\) aus Asp und Glu ausgewählt ist;
- X\(_{24}\) aus Ala und Glu ausgewählt ist;
- X\(_{28}\) aus Ser, Lys und Arg ausgewählt ist;
und worin jeder Rest X jeweils unabhängig aus der aus Glu, Lys und Cys bestehenden Gruppe ausgewählt ist;

(b) der Formel IIIa entspricht:

\[ \text{His-Alb-Gln-Gly-Thr-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X17-Ala-Ala-X20-X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala}; \quad (\text{IIIa}) \]

worin

- X\(_{12}\) aus Lys und Arg ausgewählt ist;
- X\(_{16}\) aus Ser und X ausgewählt ist;
- X\(_{17}\) X ist;
- X\(_{20}\) aus His und X ausgewählt ist;
- X\(_{21}\) aus Asp und Glu ausgewählt ist;
- X\(_{24}\) aus Ala und Glu ausgewählt ist;
- X\(_{28}\) aus Ser, Lys und Arg ausgewählt ist;
und worin jeder Rest X jeweils unabhängig aus Glu, Lys und Cys ausgewählt ist; oder

(c) der Formel IVa entspricht:

\[ \text{His-Alb-Gln-Gly-Thr-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X17-Ala-Ala-His-X20-X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala}; \quad (\text{IVa}) \]

worin

- X\(_{12}\) aus Lys und Arg ausgewählt ist;
- X\(_{17}\) X ist;
- X\(_{20}\) aus His und X ausgewählt ist;
- X\(_{21}\) aus Asp und Glu ausgewählt ist;
- X\(_{24}\) aus Ala und Glu ausgewähl isst;
- X\(_{28}\) aus Ser, Lys und Arg ausgewählt ist;
und worin jeder Rest X jeweils unabhängig aus Glu, Lys und Cys ausgewählt ist; oder
X12 aus Lys und Arg ausgewählt ist;
X17 X ist;
X21 aus Asp und Glu ausgewählt ist;
X24 aus Ala und Glu ausgewählt ist;
X28 aus Ser, Lys und Arg ausgewählt ist;
worin X aus der aus Glu, Lys und Cys bestehenden Gruppe ausgewählt ist;
worin die Seitenkette von zumindest einem Rest X an einen lipophilen Substituenten der folgenden Formel konjugiert ist:

(i) $Z^1$, worin $Z^1$ eine direkt an die Seitenkette von X konjugierte lipophile Gruppierung ist; oder
(ii) $Z^1Z^2$, worin $Z^1$ eine lipophile Gruppierung ist, $Z^2$ ein Spacer ist und $Z^1$ über $Z^2$ an die Seitenkette von X konjugiert ist.

9. Verbindung nach Anspruch 1 der Formel:

$$R^1-Z-R^2,$$
worin $R^1$ H, C$_{1-4}$-Alkyl, Acetyl, Formyl, Benzoyl oder Trifluoracetyl ist;
$R^2$ OH oder NH$_2$ ist;
und Z ein Peptid ist, das:

(a) der Formel IIb entspricht:

$$\text{His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-X16-X17-Ala-Ala-X20-X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala;}$$ (IIb)

worin

X12 aus Lys, Arg und Leu ausgewählt ist;
X16 aus Ser und X ausgewählt ist;
X17 X ist;
X20 aus His und X ausgewählt ist;
X21 aus Asp und Glu ausgewählt ist;
X24 aus Ala und Glu ausgewählt ist;
X28 aus Ser, Lys und Arg ausgewählt ist;
und worin jeder Rest X jeweils unabhängig aus der aus Glu, Lys und Cys bestehenden Gruppe ausgewählt ist;

(b) der Formel IIIb entspricht:

$$\text{His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X17-Ala-Ala-X20-X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala;}$$ (IIIb)

worin

X12 aus Lys und Arg ausgewählt ist;
X16 aus Ser und X ausgewählt ist;
X17 X ist;
X20 aus His und X ausgewählt ist;
X21 aus Asp und Glu ausgewählt ist;
X24 aus Ala und Glu ausgewählt ist;
X28 aus Ser, Lys und Arg ausgewählt ist;
und worin jeder Rest X jeweils unabhängig aus Glu, Lys und Cys ausgewählt ist; oder

(c) der Formel IVb entspricht:

$$\text{His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X17-Ala-Ala-X20-X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala;}$$ (IVb)

worin

X12 aus Lys und Arg ausgewählt ist;
X17 X ist;
X20 aus His und X ausgewählt ist;
X21 aus Asp und Glu ausgewählt ist;
X24 aus Ala und Glu ausgewählt ist;
X28 aus Ser, Lys und Arg ausgewählt ist;
und worin jeder Rest X jeweils unabhängig aus Glu, Lys und Cys ausgewählt ist; oder
worin X12 aus Lys und Arg ausgewählt ist;
X17 X ist;
X21 aus Asp und Glu ausgewählt ist;
X24 aus Ala und Glu ausgewählt ist;
X28 aus Ser, Lys und Arg ausgewählt ist;
worin X aus der aus Glu, Lys und Cys bestehenden Gruppe ausgewählt ist;
worin die Seitenkette von zumindest einem Rest X an einen lipophilen Substituenten der folgenden Formel konjugiert ist:

(i) Z1, worin Z1 eine direkt an die Seitenkette von X konjugierte lipophile Gruppierung ist; oder
(ii) Z1Z2, worin Z1 eine lipophile Gruppierung ist, Z2 ein Spacer ist und Z1 über Z2 an die Seitenkette von X konjugiert ist.

mit der Maßgabe, dass Z nicht HSQGFTSDYSKYLDS-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLRA ist.

10. Verbindung nach einem der vorangegangenen Ansprüche, worin das Peptid der Formel I, IIa, IIIa, IVa, IIb, IIIb oder IVb die folgende Sequenz aufweist:

```
HSQGFTSDYSKYLDSKAHDFVEWLLRA;
HSQGFTSDYSKYLDSKAHDFVEWLLRA;
HSQGFTSDYSKYLDSKAHDFVEWLLRA;
HSQGFTSDYSKYLDSKAHDFVEWLKRA;
HSQGFTSDYSKYLDSKAHDFVEWLLKA;
HSQGFTSDYSKYLDSKAHDFVEWLLRA;
HSQGFTSDYSKYLDSKAHDFVEWLLRA;
HSQGFTSDYSKYLDSKAHDFVEWLLRA;
HSQGFTSDYSKYLDSKAHDFVEWLLRA;
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oder

```
HSQGFTSDYSKYLDS-K*-AAHDFVEWLLRA;
```
11. Verbindung nach einem der vorangegangenen Ansprüche, worin:

(a) \( Z_1 \) eine Kohlenwasserstoffkette mit 10 bis 24 C-Atomen, 10 bis 22 C-Atomen oder 10 bis 20 C-Atomen umfasst, z.B. worin \( Z_1 \) eine Dodecanoyl-, 2-Butyloctanoyl-, Tetradecanoyl-, Hexadecanoyl-, Heptadecanoyl-, Octadecanoyl- oder Eicosanoylgruppierung ist;

(b) \( Z_2 \) ein oder mehrere Aminosäurereste ist oder diese(n) umfasst, z.B. worin \( Z_2 \) ein \( \gamma \)-Glu-, Glu-, \( \beta \)-Ala- oder \( \varepsilon \)-Lys-Rest oder eine 3-Aminopropanoyl-, 4-Aminobutanoyl-, 8-Aminooctanoyl- oder 8-Amino-3,6-dioxaoctanoylgruppierung ist; und/oder

(c) worin der lipophile Substituent aus der aus Dodecanoyl-\( \gamma \)-Glu, Hexadecanoyl-\( \gamma \)-Glu, Hexadecanoyl-E-[3-aminopropanoyl], Hexadecanoyl-E-[8-aminooctanoyl], Hexadecanoyl-E-[\( \varepsilon \)-Lys, 2-Butyloctanoyl-\( \gamma \)-Glu, Octadecanoyl-\( \gamma \)-Glu und Hexadecanoyl-[2-aminobutanoyl] bestehenden Gruppe ausgewählt ist.

12. Verbindung nach Anspruch 11, worin \( Z \) die folgende Formel aufweist:

\[
\text{HSQGTFTSDYKLD-K(Hexadecanoyl-\( \gamma \)-Glu)-KAADFVEWLLRA;}
\text{HSQGTFTSDYKLDKAA-K(Hexadecanoyl-\( \gamma \)-Glu)-RA;}
\text{HSQGTFTSDYKLDKAHHDFVEWLL-K(Hexadecanoyl-\( \gamma \)-Glu)-RA;}
\text{H-Alb-QGTFTSDYKLDK(Hexadecanoyl-\( \gamma \)-Glu)-AAADFVEWLLRA;}
\text{H-Alb-QGTFTSDYKLDK(Hexadecanoyl-\( \gamma \)-Glu)-AARDFAWLLRA;}
\]
H-Aib-QGTFTSDYKS-YLD-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLSA;
H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLKA;
H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLA;
10  HSQGTFTSDYKSYLDS-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLRA;
H-Aib-QGTFTSDYKSYLDSAA-K(Hexadecanoyl-y-Glu)-DFVAWLLRA;
H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLSA;
H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-γ-Glu)-AAHDFVEWLLA;
H-Aib-QGTFTSDYKSYLDS-K(Dodecanoyl-γ-Glu)-AAHDFVEWLLSA;
H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-ε-Lys)-AAHDFVEWLLSA;
15  HSQGTFTSDYKSYLDS-K(Hexadecanoyl)-AAHDFVEWLLSA;
HSQGTFTSDYKSYLDS-K(Octadecanoyl-γ-Glu)-AAHDFVEWLLSA;
HSQGTFTSDYKSYLDS-K(Octadecanoyl-γ-Glu)-AAHDFVEWLLSA;
20  HSQGTFTSDYKSYLDS-K([2-Butyloctanoyl]-γ-Glu)-AAHDFVEWLLSA;
H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-ε-Lys)-AAHDFVEWLLSA;
H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-[4-Aminobutanoyl])-AAHDFVEWLLSA;
25  H-Aib-QGTFTSDYKSYLDS-K(Octadecanoyl-γ-Glu)-AAHDFVEWLLSA;
H-Aib-QGTFTSDYKSYLDS-K([2-Butyloctanoyl]-γ-Glu)-AAHDFVEWLLSA;
H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-ε-Lys)-AAHDFVEWLLSA;
H-Aib-QGTFTSDYKSYLDS-K(Octadecanoyl-γ-Glu)-AAHDFVEWLLSA;
30  HSQGTFTSDYKSYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA;
H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA;
H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA;
H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA;
35  H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA;
H-Aib-QGTFTSDYKSYLDS-K(Hexadecanoyl-isoGlu)-AAHDFVEWLLSA;


14. Verbindung der Formel:

\[ R^1 \cdot Z \cdot R^2, \]

worin \( R^1 \) H, C₁₋₄-Alkyl, Acetyl, Formyl, Benzoyl oder Trifluoracetyl ist;
\( R^2 \) OH oder NH₂ ist;
und Z ein Peptid ist, das:

(a) der Formel V entspricht:

\[ \text{His-} \text{Aib-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-Lys-Tyr-Leu-Asp-Ser-X17-Ala-Ata-His-Asp-Phe-Val-Glu-Trp-Leu-Leu-X28; } (V) \]

worin

X₁₇ X ist,
X₂₈ Ser ist oder fehlt; oder

40
(b) der Formel VI entspricht:

\[
\text{His-Aib-Glu-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-Leu-Asp-Ser-X17-Ala-Ala-His-}
\text{Asp-Phe-Val-Glu-Trp-Leu-Leu-Ser-Ala; (VI)}
\]

worin

\[
X_{17} \text{ X ist;}
\]

worin X aus der aus Glu, Lys und Cys bestehenden Gruppe ausgewählt ist;

und worin die Seitenkette von zumindest einem Rest X an einen lipophilen Substituenten der folgenden Formel konjugiert ist:

(i) \(Z^1\), worin \(Z^1\) eine direkt an die Seitenkette von X konjugierte lipophile Gruppierung ist; oder

(ii) \(Z^1Z^2\), worin \(Z^1\) eine lipophile Gruppierung ist, \(Z^2\) ein Spacer ist und \(Z^1\) über \(Z^2\) an die Seitenkette von X konjugiert ist.

15. Verbindung nach Anspruch 14, worin \(Z\) der folgenden Formel entspricht:

\[
\text{H-Aib-QGFTSDSKYLD-K[Hexadecanoyl-isoGlu]-AAHDFVEWLLS;}
\]

\[
\text{H-Aib-QGFTSDSKYLD-K[Hexadecanoyl-isoGlu]-AAHDFVEWLL;} \text{ oder}
\]

\[
\text{H-Aib-EGFTSDSKYLD-K[Hexadecanoyl-isoGlu]-AAHDFVEWLLSA.}
\]

16. Zusammensetzung, die eine Verbindung nach einem der Ansprüche 1 bis 15 oder ein Salz oder Derivat davon im Gemisch mit einem Träger umfasst, z.B. worin die Zusammensetzung eine pharmazeutisch annehmbare Zusammensetzung ist und der Träger ein pharmazeutisch annehmbarer Träger ist.

17. Verbindung nach einem der Ansprüche 1 bis 15 zur Verwendung in einem Verfahren zur medizinischen Behandlung, z.B.

(a) zur Verwendung zur Prävention von Gewichtszunahme oder der Förderung von Gewichtsverlust;

(b) zur Verwendung in einem Verfahren zur Verbesserung der zirkulierenden Glucosespiegel, Glucosetoleranz und/oder zirkulierenden Cholesterinspiegeln, zum Senken der zirkulierenden LDL-Spiegel und/oder zur Erhöhung des HDL/LDL-Verhältnisses; oder

(c) zur Verwendung in einem Verfahren zur Behandlung eines durch überschüssiges Körpergewicht verursachten oder dadurch gekennzeichneten Leidens, z.B. zur Behandlung und/oder Prävention von Adipositas, morbid Adipositas, adipositasbedingter Entzündung, adipositasbedingter Gallenblasenerkrankung, durch Adipositas ausgelöster Schlafapnoe, metabolischem Syndrom, Prädiabetes, Insulinresistenz, Glucosetolereznanz, Typ-2-Diabetes, Typ-1-Diabetes, Bluthochdruck, athergener Dyslipidämie, Atherosklerose, Arteriosklerose, koronarer Herzkrankheit, peripherer Arterienkrankheit, Schlaganfall oder mikrovaskulärer Krankheit.


19. Verbindung zur Verwendung nach Anspruch 18, worin das Mittel zur Behandlung von Diabetes Metformin, ein Sulfonylharnstoff, ein Glinit, ein DPP-IV-Inhibitor, ein Gltazon, Insulin oder ein Insulinanalagon ist.


22. Verbindung zur Verwendung nach Anspruch 18, worin das Mittel zur Behandlung von Dyslipidämie ein Statin, ein Fibrat, ein Niacin und/oder ein Cholesterinabsorptionsinhibitor ist.

23. Verwendung einer Verbindung nach einem der Ansprüche 1 bis 15 zur Herstellung eines Medikaments zur:
(a) Prävention von Gewichtszunahme oder Förderung von Gewichtsverlust bei einem Individuum mit Bedarf daran;
(b) Verbesserung von zirkulierenden Glucosespiegeln, Glucosetoleranz und/oder zirkulierenden Cholesterinspiegeln, zum Senken des HDL/LDL-Verhältnisses bei einem Individuum mit Bedarf daran; oder

24. Verwendung nach Anspruch 23, worin die Verbindung der Verabreichung als Teil einer Kombinationstherapie mit einem Mittel zur Behandlung von Diabetes, Adipositas, Dyslipidämie oder Bluthochdruck dient, z.B.
(a) worin das Mittel zur Behandlung von Diabetes Metformin, ein Sulfonylharnstoff, ein Glinid, ein DPP-IV-Inhibitor, ein Glitazon, Insulin oder ein Insulinanalagon ist;
(b) worin das Mittel zur Behandlung von Adipositas ein glucagonartiger Peptidrezeptor-1-Agonist, Peptid YY oder ein Analogon davon, ein Cannabinoidezeptor-1-Antagonist, Lipaseinhibitor, Melanocortinrezeptor-4-Antagonist oder Antagonist von Melanin konzentrierendem Hormonrezeptor 1 ist;
(c) worin das Mittel zur Behandlung von Bluthochdruck ein Inhibitor des Angiotensin konvertierenden Enyzms, ein Angiotensin-II-Rezeptor-Blocker, ein Diuretikum, ein Beta-blocker oder ein Calciumkanalblocker ist; oder
(d) worin das Mittel zur Behandlung von Dyslipidämie ein Statin, ein Fibrat, ein Niacin und/oder ein Cholesterinabsorptionsinhibitor ist.

Revendications

1. Composé ayant la formule :

\[ \text{R}^1-Z\text{-R}^2 \]

\[ \text{dans laquelle R}^1 \text{ est H, alkyle en C}_{1-4}, \text{acétyle, formyle, benzoyle ou trifluoroacétyle ;} \]
\[ \text{R}^2 \text{ est OH ou NH}_{2}; \]
\[ \text{et Z est un peptide ayant la formule I} \]


\( (I) \)

dans laquelle

\[ X2 \text{ est sélectionné parmi Alb et Ser ;} \]
\[ X12 \text{ est sélectionné parmi Lys, Arg ou Leu ;} \]
\[ X16 \text{ est sélectionné parmi Arg et X ;} \]
\[ X17 \text{ est sélectionné parmi Arg et X ;} \]
\[ X20 \text{ est sélectionné parmi Arg, His et X ;} \]
\[ X21 \text{ est sélectionné parmi Asp et Glu ;} \]
\[ X24 \text{ est sélectionné parmi Ala et X ;} \]
\[ X27 \text{ est sélectionné parmi Leu et X ;} \]
\[ X28 \text{ est sélectionné parmi Arg et X ;} \]
\[ X30 \text{ est X ou est absent ;} \]
\[ \text{où au moins l’un de X16, X17, X20, X24, X27, X28 et X30 est X ;} \]
\[ \text{et où chaque résidu X est indépendamment sélectionné dans le groupe consistant en Glu, Lys, Ser, Cys, Dbu, Dpr et Orn ;} \]
\[ \text{où la chaîne latérale d’au moins un résidu X est conjuguée à un substituant lipophile ayant la formule :} \]

(i) \( Z^1, \text{ où } Z^1 \text{ est un fragment lipophile conjugué directement à la chaîne latérale de } X ; \text{ ou} \)

(ii) \( Z^1Z^2, \text{ où } Z^1 \text{ est un fragment lipophile, } Z^2 \text{ est un espaceur, et } Z^1 \text{ est conjugué à la chaîne latérale de } X \)
EP 2 454 282 B1

via Z² ;

à condition que Z ne soit pas HSQGFTSDYSKYLDS-K(hexadécanoyl-γ-Glu)-AAHDFVEWLLRA.

2. Composé selon la revendication 1, dans lequel :

(a) un ou plusieurs desdits résidus X est indépendamment sélectionné parmi Lys, Glu et Cys ; et/ou
(b) X16 est sélectionné parmi Glu, Lys et Ser ;

X17 est sélectionné parmi Lys et Cys ;
X20 est sélectionné parmi His, Lys, Arg et Cys ;
X24 est sélectionné parmi Lys, Glu et Ala ;
X27 est sélectionné parmi Leu et Lys ; et/ou
X28 est sélectionné parmi Ser, Arg et Lys ; et/ou

(c) le peptide de formule I comprend une ou plusieurs des combinaisons de résidus suivantes :

X2 est Aib et X17 est Lys ;
X2 est Aib et X17 est Cys ;
X2 est Aib et X20 est Cys ;
X2 est Aib et X28 est Lys ;
X12 est Arg et X17 est Lys ;
X12 est Leu et X17 est Lys ;
X12 est Lys et X20 est Lys ;
X12 est Lys et X17 est Lys ;
X16 est Lys et X17 est Lys ;
X16 est Ser et X17 est Lys ;
X17 est Lys et X20 est Lys ;
X17 est Lys et X24 est Glu ;
X17 est Lys et X27 est Leu ;
X17 est Lys et X27 est Lys ;
X17 est Lys et X28 est Ser ;
X20 est Lys et X27 est Leu ;
X21 est Asp et X27 est Leu ;
X2 est Aib, X12 est Lys et X16 est Ser ;
X12 est Lys, X17 est Lys et X16 est Ser ;
X12 est Arg, X17 est Lys et X16 est Glu ;
X16 est Glu, X17 est Lys et X20 est Lys ;
X16 est Ser, X21 est Asp et X24 est Glu ;
X17 est Lys, X24 est Glu et X28 est Arg ;
X17 est Lys, X24 est Glu et X28 est Lys ;
X17 est Lys, X27 est Leu et X28 est Ser ;
X17 est Lys, X27 est Leu et X28 est Arg ;
X17 est Lys, X27 est Leu et X28 est Ser ;
X17 est Lys, X27 est Leu et X28 est Arg ;
X20 est Lys, X24 est Glu et X27 est Leu ;
X20 est Lys, X27 est Leu et X28 est Ser ;
X20 est Lys, X27 est Leu et X28 est Arg ;
X16 est Ser, X20 est His, X24 est Glu et X27 est Leu ;
X17 est Lys, X20 est His, X24 est Glu et X28 est Ser ;
X17 est Lys, X20 est Lys, X24 est Glu et X27 est Leu ; ou
X17 est Cys, X20 est Lys, X24 est Glu et X27 est Leu.

3. Composé selon la revendication 1 ou la revendication 2, dans lequel le peptide de formule I contient un seul acide améné du type conjugué au substituant lipophile, par exemple où le peptide contient un seul résidu Lys, un seul résidu Cys ou un seul résidu Glu, et où le substituant lipophile est conjugué à ce résidu.

4. Composé selon l’une quelconque des revendications précédentes, dans lequel la séquence peptidique de formule
I comprend un ou plusieurs ponts intramoléculaires, et où facultativement

(a) ledit pont intramoléculaire est formé entre les chaînes latérales de deux résidus d’acides aminés qui sont séparés par trois acides aminés dans la séquence d’acides aminés linéaire de formule I ;
(b) le pont intramoléculaire est formé entre les chaînes latérales des paires de résidus 16 et 20, 17 et 21, 20 et 24, ou 24 et 28 ;
(c) le pont intramoléculaire est un pont saîn ou un cycle lactame ; et/ou
(d) le pont intramoléculaire implique une paire de résidus dans laquelle :

<table>
<thead>
<tr>
<th></th>
<th>X16</th>
<th>X20</th>
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<tbody>
<tr>
<td>16</td>
<td>Glu</td>
<td>Lys</td>
</tr>
<tr>
<td>17</td>
<td>Glu</td>
<td>Lys</td>
</tr>
<tr>
<td>20</td>
<td>Glu</td>
<td>Lys</td>
</tr>
<tr>
<td>24</td>
<td>Glu</td>
<td>Lys</td>
</tr>
</tbody>
</table>

5. Composé selon l’une quelconque des revendications précédentes, dans lequel au moins l’un de X16, X17, X20 et X28 est conjugué à un substituant lipophile.

6. Composé selon l’une quelconque des revendications précédentes, dans lequel :

(a) X30 est absent ; ou
(b) X30 est présent et est conjugué à un substituant lipophile.

7. Composé selon l’une quelconque des revendications précédentes, dans lequel :

(a) le composé possède un seul substituant lipophile, à la position 16, 17, 20, 24, 27, 28 ou 30, de préférence à la position 16, 17 ou 20, particulièrement à la position 17 ;
(b) le composé possède précisément deux substituants lipophiles, chacun à l’une des positions 16, 17, 20, 24, 27, 28 ou 30 ; ou
(c) le composé possède des substituants lipophiles aux positions 16 et 17, 16 et 20, 16 et 24, 16 et 27, 16 et 28 ou 16 et 30 ; aux positions 17 et 20, 17 et 24, 17 et 27, 17 et 28 ou 17 et 30 ; aux positions 20 et 24, 20 et 27, 20 et 28 ou 20 et 30 ; aux positions 24 et 27, 24 et 28 ou 24 et 30 ; aux positions 27 et 28 ou 27 et 30 ; ou aux positions 28 et 30.

8. Composé selon la revendication 1, ayant la formule :

\[ R^1 - Z - R^2 \]

dans laquelle \( R^1 \) est H, alkyle en C\(_{1-4}\), acétyle, formyle, benzoyle ou trifluoroacétyle ;
\( R^2 \) est OH ou NH\(_2\) ;
et Z est un peptide ayant :

(a) la formule IIa :

\[ \text{His-Alb-Gln-Gly-Thr-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-X16-X17-Ala-Ala-X20-X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala} \] (IIa)
dans laquelle

X12 est sélectionné parmi Lys, Arg et Leu ;
X16 est sélectionné parmi Ser et X ;
X17 est X ;
X20 est sélectionné parmi His et X ;
X21 est sélectionné parmi Asp et Glu ;
X24 est sélectionné parmi Ala et Glu ;
X28 est sélectionné parmi Ser, Lys et Arg ;
et où chaque résidu X est indépendamment sélectionné dans le groupe consistant en Glu, Lys et Cys ;

(b) la formule IIIa :

His-Aib-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X17-Ala-Ala-X20-
X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala ; (IIIa)

dans laquelle

X12 est sélectionné parmi Lys et Arg ;
X17 est X ;
X20 est sélectionné parmi His et X ;
X21 est sélectionné parmi Asp et Glu ;
X24 est sélectionné parmi Ala et Glu ;
X28 est sélectionné parmi Ser, Lys et Arg ;
et où chaque résidu X est indépendamment sélectionné dans le groupe consistant en Glu, Lys et Cys ; ou

(c) la formule IVa .

His-Aib-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X17-Ala-Ala-His-X21-
Phe-Val-X24-Trp-Leu-Leu-X28-Ala ; (IVa)

dans laquelle

X12 est sélectionné parmi Lys et Arg ;
X17 est X ;
X21 est sélectionné parmi Asp et Glu ;
X24 est sélectionné parmi Ala et Glu ;
X28 est sélectionné parmi Ser, Lys et Arg ;
et X est sélectionné dans le groupe consistant en Glu, Lys et Cys ; où la chaîne latérale d’au moins un résidu X est conjuguée à un substituant lipophile ayant la formule :

(i) $Z^1$, où $Z^1$ est un fragment lipophile conjugué directement à la chaîne latérale de X ; ou
(ii) $Z^1Z^2$, où $Z^1$ est un fragment lipophile, $Z^2$ est un espaceur, et $Z^1$ est conjugué à la chaîne latérale de X via $Z^2$.

9. Composé selon la revendication 1, ayant la formule :

$$R^1-Z-R^2$$

dans laquelle $R^1$ est H, alkyle en $C_{1-4}$, acétylome, formyle, benzoyle ou trifluoroacétylme ;
$R^2$ est OH ou NH$_2$ ;
et Z est un peptide ayant :

(a) la formule IIb :

His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X16-X17-Ala-Ala-X20-
X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala ; (IIb)
dans laquelle

X12 est sélectionné parmi Lys, Arg et Leu ;
X16 est sélectionné parmi Ser et X ;
X17 est X ;
X20 est sélectionné parmi His et X ;
X21 est sélectionné parmi Asp et Glu ;
X24 est sélectionné parmi Ala et Glu ;
X28 est sélectionné parmi Ser, Lys et Arg ;
et où chaque résidu X est indépendamment sélectionné dans le groupe consistant en Glu, Lys et Cys ;

(b) la formule IIIb :

His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X17-Ala-Ala-X20-
X21-Phe-Val-X24-Trp-Leu-Leu-X28-Ala ; (IIIb)
dans laquelle

X12 est sélectionné parmi Lys et Arg ;
X17 est X ;
X20 est sélectionné parmi His et X ;
X21 est sélectionné parmi Asp et Glu ;
X24 est sélectionné parmi Ala et Glu ;
X28 est sélectionné parmi Ser, Lys et Arg ;
et où chaque résidu X est indépendamment sélectionné parmi Glu, Lys ou Cys ; ou

(c) la formule IVb :

His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-X12-Tyr-Leu-Asp-Ser-X17-Ala-Ala-His-X21-
Phe-Val-X24-Trp-Leu-Leu-X28-Ala ; (IVb)
dans laquelle

X12 est sélectionné parmi Lys et Arg ;
X17 est X ;
X20 est sélectionné parmi Asp et Glu ;
X24 est sélectionné parmi Ala et Glu ;
X28 est sélectionné parmi Ser, Lys et Arg ;
où X est sélectionné dans le groupe consistant en Glu, Lys et Cys ;
ou la chaîne latérale d’au moins un résidu X est conjuguée à un substituant lipophile ayant la formule :

(i) Z1, où Z1 est un fragment lipophile conjugué directement à la chaîne latérale de X ; ou
(ii) Z1Z2, où Z1 est un fragment lipophile, Z2 est un espaceur, et Z1 est conjugué à la chaîne latérale
do X via Z2 ;

à condition que Z ne soit pas HSQGTFTSDYSKYLDS-K(hexadécanoyl-γ-Glu)-AAHDFVEWLLRA.

10. Composé selon l’une quelconque des revendications précédentes, dans lequel le peptide de formule I, Ila, IIIa, IVa,
llb, IIIb ou IVb possède la séquence :

HSQGTFTSDYSKYLDSKAAHDFVEWLLRA;
HSQGTFTSDYSKYLDSKAAHDFVEWLLRA;
HSQGTFTSDYSKYLDSKAAKDFVEWLLRA;
HSQGTFTSDYSKYLDSKAAHDFVEWLLRA;
HSQGTFTSDYSKYLDSKAAHDFVEWLLRA;
HSQGTFTSDYSKYLDSKAAHDFVEWLLRA;
HSQGTFTSDYSKYLDSKAAHDFVEWLLRA;
HSQGTFTSDYSKYLDSKAAHDFVEWLLRA;
HSQGTFTSDYSKYLDSKAAHDFVEWLLRA;
HSQGTFTSDYSKYLDSKAAHDFVEWLLRA;
HSQGTFTSDYSKYLDKAAHDFVEWLLSAK;
HSQGTFTSDYSKYLDKAAHDFVEWLKSA;
HSQGTFTSDYSKYLDKAAHDFVWLARRA;
HSQGTFTSDYSKYLDSCAAHDFVEWLLRA;
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HSQGTFTSDYLDSCAAHDFVEWLLSA;
HSQGTFTSDYLDSCAAHDFVEWLLRA;
HSQGTFTSDYLDSCAAHDFVEWLRLA;
HSQGTFTSDYLDSCAAHDFVEWLLSA;
HSQGTFTSDYLDSCAAHDFVEWLLRA;
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HSQGTFTSDYLDSCAAHDFWEWLLRA;
H-Aib-QGTFTSDYLDKAAHDFVEWLLSA;
H-Aib-QGTFTSDYLDKAAHDFVEWLLSAK;
H-Aib-QGTFTSDYLDKAAHDFVWLARRA;
H-Aib-QGTFTSDYLDKAAHDFVWLARRA;
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H-Aib-QGTFTSDYLDKAAHDFVWLARRA;
H-Aib-QGTFTSDYLDKAAHDFVWLARRA;
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ou
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HSQGTFTSDYSKYLDL-K*-AAHDFVEWLLRA;
HSQGTFTSDYSKYLDL-K*-AAHDFVEWLLRA;
HSQGTFTSDYSKYLDL-K*-AAHDFVEWLLRA;
HSQGTFTSDYSKYLDL-K*-AAHDFVEWLLRA;
HSQGTFTSDYSKYLDL-K*-AAHDFVEWLLRA;
11. Composé selon l'une quelconque des revendications précédentes, dans lequel :

(a) Z₁ comprend une chaîne hydrocarbonée ayant 10 à 24 atomes de C, 10 à 22 atomes de C ou 10 à 20 atomes de C, par exemple où Z₁ est un fragment dodécanoyle, 2-butylloctanoyle, tétradécanoyle, hexadécanoyle, heptadécanoyle, octadécanoyle ou éicosanoyle ;
(b) Z₂ est ou comprend un ou plusieurs résidus d'acides aminés, par exemple où Z₂ est un résidu γ-Glu, Glu, β-Ala ou ε-Lys, ou un fragment 3-aminopropanoyle, 4-aminobutanoyl, 8-aminooctanoyle ou 8-amino-3,6-dioxoctanoyle ; et/ou
(c) où le substituant lipophile est sélectionné dans le groupe consistant en dodécanoyl-γ-Glu, hexadécanoyl-γ-Glu, hexadécanoyl-Glu, hexadécanoyl-[3-aminopropanoyl], hexadécanoyl-[8-aminooctanoyle], hexadécanoyle-ε-Lys, 2-butylloctanoyle-γ-Glu, octadécanoyle-γ-Glu et hexadécanoyle-[4-aminobutanoyl].

12. Composé selon la revendication 11, dans lequel Z possède la formule :

H-Aib-QGTFTSDYSKYLK(\text{Hexadecanoyl-\(\gamma\)-Glu})-KAAHDFVEWLLRA;
H-Aib-QGTFTSDYSKYLK(\text{Hexadecanoyl-\(\gamma\)-Glu})-DFVEWLLRA;
H-Aib-QGTFTSDYSKYLK(\text{Hexadecanoyl-\(\gamma\)-Glu})-DFVEWLLRA;
H-Aib-QGTFTSDYSKYLK(\text{Hexadecanoyl-\(\gamma\)-Glu})-DFVEWLLRA;

ou les résidus marqués « ( ) » participent à une liaison intramoléculaire ; ou

H-Aib-QGTFTSDYSKYLK(\text{Hexadecanoyl-\(\gamma\)-Glu})-DFVEWLLRA;
13. Composé selon la revendication 1, qui est H-Aib-QGTFTSDYSKYLDS-K(hexadécanoyl-\(\gamma\)-Glu)-AAHDFVEWLLEA-NH\(_2\).

14. Composé ayant la formule :

\[ R^1\cdot Z\cdot R^2 \]

dans laquelle 
\( R^1 \) est H, alkyle en C\(_{1-4}\), acétyle, formyle, benzoyle ou trifluoroacétyle ;
\( R^2 \) est OH ou NH\(_2\) ;
et \( Z \) est un peptide ayant :

(a) la formule V :

\[ \text{His-Aib-Gln-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-Lys-Tyr-Leu-Asp-Ser-X\(_{17}\)-Ala-Ala-His-Asp-Phe-Val-Glu-Trp-Leu-Leu-X\(_{28}\)} \]

(V)

dans laquelle

\( X\(_{17}\) \) est X
\( X\(_{28}\) \) est Ser ou absent ; ou

(b) la formule VI :

\[ \text{His-Aib-Glu-Gly-Thr-Phe-Thr-Ser-Asp-Tyr-Ser-Lys-Tyr-Leu-Asp-Ser-X\(_{17}\)-Ala-Ala-His-Asp-Phe-Val-Glu-Trp-Leu-Leu-Ser-Ala} \]

(VI)

dans laquelle

\( X\(_{17}\) \) est X ;
où X est sélectionné dans le groupe consistant en Glu, Lys et Cys ;
et où la chaîne latérale de X est conjuguée à un substituant lipophile ayant la formule :

(i) \( Z^1 \)
\( Z^1 \) est un fragment lipophile conjugué directement à la chaîne latérale de X ; ou

(ii) \( Z^1Z^2 \)
\( Z^1 \) est un fragment lipophile, \( Z^2 \) est un espaceur, et \( Z^1 \) est conjugué à la chaîne latérale de X via \( Z^2 \).

15. Composé selon la revendication 14, dans lequel \( Z \) possède la formule :

H-Aib-QGTFTSDYSKYLDS-K(hexadécanoyl-isoGlu)-AAHDFVEWLLSA ;
H-Aib-QGTFTSDYSKYLDS-K(hexadécanoyl-isoGlu)-AAHDFVEWLLS ;
H-Aib-EGTFTSDYSKYLDS-K(hexadécanoyl-isoGlu)-AAHDFVEWLLSA.

16. Composition comprenant un composé selon l’une quelconque des revendications 1 à 15, ou un sel ou dérivé de celui-ci, dans un mélange avec un véhicule, par exemple où la composition est une composition pharmaceutiquement acceptable et le véhicule est un véhicule pharmaceutiquement acceptable.

17. Composé selon l’une quelconque des revendications 1 à 15 destiné à être utilisé dans un procédé de traitement médical, par exemple :

(a) destiné à être utilisé pour prévenir la prise de poids ou promouvoir la perte de poids ;
(b) destiné à être utilisé dans un procédé pour améliorer les taux de glucose circulant, la tolérance au glucose et/ou les taux de cholestérol circulant, réduire les taux de LDL circulant, et/ou augmenter le rapport HDL/LDL ; ou
18. Composé destiné à être utilisé selon la revendication 17, où le composé est destiné à une administration en tant qu’élément d’une thérapie de combinaison avec un agent pour le traitement du diabète, de l’obésité, de la dyslipidémie ou de l’hypertension.

19. Composé destiné à être utilisé selon la revendication 18, dans lequel l’agent pour le traitement du diabète est la metformine, une sulfonylurée, un glinide, un inhibiteur de la DPP-IV, une glitazone, l’insuline ou un analogue de l’insuline.

20. Composé destiné à être utilisé selon la revendication 18, dans lequel l’agent pour le traitement de l’obésité est un agoniste du récepteur du peptide analogue au glucagon 1, un peptide YY ou un analogue de celui-ci, un antagoniste du récepteur cannabinoïde 1, un inhibiteur de la lipase, un agoniste du récepteur 4 de la mélanocortine, ou un antagoniste du récepteur 1 de l’hormone concentrant la mélanine.


22. Composé destiné à être utilisé selon la revendication 18, dans lequel l’agent pour le traitement de la dyslipidémie est une statine, un fibrate, une niacine et/ou un inhibiteur de l’absorption du cholestérol.

23. Utilisation d’un composé selon 1’une quelconque des revendications 1 à 15 dans la préparation d’un médicament pour :

(a) prévenir la prise de poids ou promouvoir la perte de poids chez un individu en ayant besoin ;
(b) améliorer les taux de glucose circulant, la tolérance au glucose et/ou les taux de cholestérol circulant, réduire les taux de LDL circulant, et/ou augmenter le rapport HDL/LDL chez un individu en ayant besoin ; ou
(c) le traitement d’un état pathologique provoqué ou caractérisé par un poids corporel excessif, par exemple le traitement ou la prévention de l’obésité, l’obésité morbide, une inflammation liée à l’obésité, l’apnée du sommeil induite par l’obésité, le pré-diabète, l’insulinorésistance, l’intolérance au glucose, le diabète de type 2, le diabète de type I, l’hypertension, la dyslipidémie athérogène, l’athérosclérose, l’artériosclérose, une coronaropathie, une maladie artérielle périphérique, un AVC ou une maladie microvasculaire.

24. Utilisation selon la revendication 23, dans laquelle le composé est destiné à une administration en tant qu’élément d’une thérapie de combinaison avec un agent pour le traitement du diabète, de l’obésité, de la dyslipidémie ou de l’hypertension, par exemple

(a) où l’agent pour le traitement du diabète est la metformine, une sulfonylurée, un glinide, un inhibiteur de la DPP-IV, une glitazone, l’insuline ou un analogue de l’insuline ;
(b) où l’agent pour le traitement de l’obésité est un agoniste du récepteur du peptide analogue au glucagon 1, un peptide YY ou un analogue de celui-ci, un antagoniste du récepteur cannabinoïde 1, un inhibiteur de la lipase, un agoniste du récepteur 4 de la mélanocortine, ou un antagoniste du récepteur 1 de l’hormone concentrant la mélanine ;
(c) où l’agent pour le traitement de l’hypertension est un inhibiteur de l’enzyme de conversion de l’angiotensine, un antagoniste des récepteurs de l’angiotensine II, un diurétique, un bêtabloquant, ou un inhibiteur des canaux calciques ; ou
(d) où l’agent pour le traitement de la dyslipidémie est une statine, un fibrate, une niacine et/ou un inhibiteur de l’absorption du cholestérol.
Figure 1.

Cpd. 13. dosed 100 nmol/kg to mice SC
Figure 2.

**Diagram:**
- **Vehicle**
- **Cpd. 11 (10 nmol/kg)**

**Y-axis (Plasma Glucose):**
- Values range from 4 to 10 mmol/l.

**X-axis (Time):**
- Time in minutes: 0, 30, 60, 90, 120.

The diagram illustrates the time course of plasma glucose levels after administering Vehicle and Cpd. 11 (10 nmol/kg).
Figure 3.
Figure 4.

![Bar chart showing body weight (g) for different treatments: Vehicle before treatment, Cpd 11 before treatment, Vehicle after treatment, Cpd 11 after treatment.](chart.png)
Figure 5.

![Bar chart showing total cholesterol levels for Vehicle and cpd. 7 treatments. The chart indicates a significant difference with **P < 0.0001.](https://example.com)
Figure 6.

![Graph showing HDL/LDL ratio for Vehicle and cpd. 7 treatments.]

- **Vehicle**
- **cpd. 7 12.7 nmol/kg**

*** P < 0.0001
Figure 7.
Figure 8.

Blood glucose after p.o. glucose at t=0. (n=7-8)

- Vehicle 2h
- Cpd 7 2h

Blood glucose after p.o. glucose at t=0. (n=7-8)

- Vehicle 4h
- Cpd 7 4h

Blood glucose after p.o. glucose at t=0. (n=7-8)

- Vehicle 6h
- Cpd 7 6h

Blood glucose after p.o. glucose at t=0. (n=7-8)

- Vehicle 8h
- Cpd 7 8h

Blood glucose after p.o. glucose at t=0. (n=7-8)

- Vehicle 10h
- Cpd 7 10h

Blood glucose after p.o. glucose at t=0. (n=7-8)

- Vehicle 12h
- Cpd 7 12h
Figure 9.

Cumulated Food Intake/Body Weight

Young Lean (Vehicle)
Young Lean (Exendin-4)
Young Lean (Cpd. 7)

Time (Hours)
Figure 10.
Figure 12.
Figure 13.

- **Vehicle**
- **Cpd. 11 12.7 nmol/kg**

ΔHbA1c (%)

- **P = 0.03**

Treatment: Cpd. 11
REFERENCES CITED IN THE DESCRIPTION

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