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MARKUSSEN J ET AL: "SOLUBLE, PROLONGED-ACTING INSULIN DERIVATIVES. I. DEGREE OF PROTRACTION AND CRYSTALLIZABILITY OF INSULINS SUBSTITUTED IN THE TERMINI OF THE B-CHAIN", PROTEIN ENGINEERING, OXFORD UNIVERSITY PRESS, SURREY, GB, vol. 1, no. 3, 1 June 1987 (1987-06-01), pages 205-213, XP000000187, ISSN: 0269-2139

Fortsættes ...

Description

The invention relates to novel insulin analogues with basal time/action profile, their preparation and use.

5

The incidence of diabetes has increased in recent years to an almost epidemic extent. The disorder may result in a serious shortening of life expectancy. People with diabetes must frequently supply their body with insulin from outside. It is
10 sensible to optimize the treatment with insulin. Different insulins with specific pharmacological properties are now available. In practice, the different insulins are differentiated according to their duration of action into short-acting insulins, fast-acting insulins, long-acting
15 insulins and mixed insulins. Designations used synonymously for long-acting insulins are slow insulins, depot insulin or else basal insulin. The active ingredients in many of these insulin products are so-called insulin analogues which have been derived from human insulin by substitution, deletion
20 and/or addition of one or more amino acids. The terms "insulin analogues" and "insulins" are used synonymously herein.

The policy of intensified insulin therapy attempts to diminish the health risk by aiming at a stable control of the blood
25 glucose level by early administration of basal insulins. One example of a current basal insulin is the medicament Lantus® (active ingredient: insulin glargin = Gly (A21), Arg (B31), Arg (B32) human insulin). The general aim of developing novel, improved basal insulins is to minimize the number of
30 hypoglycemic events. An ideal basal insulin in this connection is one acting reliably for at least 24 hours in each patient. The insulin effect ideally has a delayed onset and a time/action profile which is as shallow as possible, so that the risk of brief hypoglycemia is distinctly minimized and
35 administration is even possible without previous intake of foodstuffs. There is a good supply of basal insulin when the insulin effect persists at the same level for as long as possible, i.e. the body is supplied with a constant amount of

insulin. The risk of hypoglycemic events is thus low and a patient- and a day-specific variability is minimized. The pharmacokinetic profile of an ideal basal insulin should thus be characterized by a delayed onset of action and by a
5 delayed, i.e. long-lasting and uniform, action.

However - despite the therapeutic advantages already achieved - none of the slow insulins described to date shows the pharmacokinetic properties of an ideal basal insulin.
10 Desirable insulins have such a shallow and long-lasting time/action profile that the risk of hypoglycemic events and of the day-dependent variations in the patient is further minimized and the duration of action is further delayed, so that it is no longer necessary in some circumstances to
15 administer insulin daily. This would make simplified treatment of diabetics possible, especially of elderly diabetics and those in need of care, who are no longer able to inject insulin themselves, and would thus also be of great economic benefit. Such basal insulins would additionally be beneficial
20 in the early phase of type 2 diabetes. Clinicians report that the injection phobia present in many people deters them from starting insulin therapy in good time. As a consequence, the control of blood glucose is poor, leading to the late sequelae of diabetes. A basal insulin which reduces the number of
25 insulin doses given by injection might have the effect of making insulin therapy more acceptable to patients.

US 6,100,376 discloses insulin derivatives having an isoelectric point between 5 and 8.5 with increased stability
30 in slightly acidic aqueous medium. US 6,100,376 describes the C-terminal modification Arg-Arg-OH at the B chain.

WO 2007/081824 discloses fibrillation-resistant insulin analogues having histidine residues at positions A4, A8 and
35 B1.

WO 89/10937 relates to the stabilization of insulin analogues against chemical modifications such as desamidation or di- or

polymerization. WO 89/10937 discloses modifications at positions A15, A18, A21, B3, B4 and B10.

Markussen et al. (Protein Engineering, 1987, 1: 205-213) disclose insulin analogues having prolonged action and describe substitutions at positions B-1, B1, B29, B30, B31 and B32. Substitution is with the amino acids Thr, Lys, Anti-sticking properties, Arg and Phe.

10 Kohn et al. (Peptides 28 (2007) 935-948) describe how it is possible to optimize the pharmacodynamics of insulin by preparing insulin analogues whose isoelectric point (pI) is shifted, by addition of lysine or arginine at the B chain end or at the N terminus of the A and B chain, in the direction of
15 the alkaline range compared with the isoelectric point of human insulin (pI = 5.6), so that the solubility under physiological conditions is reduced and a prolonged time/action profile results. Compound 18 from Kohn et al. (Arg (A0), Gly (A21), Arg (B31), Arg (B32) human insulin
20 (experimentally determined pI = 7.3; calculated pI = 7.58) is described in this connection as the best compound in the context of the idea. Kohn et al. therefore regard the main aim in designing novel insulin analogues as being the addition of positively charged amino acids to the amino acid sequence of
25 human insulin for the purpose of increasing the isoelectric point from pI = 5.6 into the neutral range.

This aim in the design of novel insulin analogues is the opposite of substitution of neutral amino acids in human
30 insulin by acidic amino acids and/or addition of acidic amino acids, because such a substitution and/or additions at least partly abolishes the effect of introducing positively charged amino acids. However, it has now surprisingly been found that the described desirable basal time/action profile is obtained
35 with insulin analogues which are characterized by the features that

- the B chain end consists of an amidated basic amino acid

residue such as lysine or argininamide, and

- the N-terminal amino acid residue of the insulin A chain is a lysine or arginine residue, and

5

- the A8 amino acid position is occupied by a histidine residue, and

- the A21 amino acid position is occupied by a glycine residue, i.e. in the amidated basic amino acid residue at the B chain end the carboxyl group of the terminal amino acid is present in its amidated form, and

- there have been two substitutions of neutral amino acids by acidic amino acids, two additions of negatively charged amino acid residues or one such substitution and one such addition respectively in the A5, A15, A18, B-1, B0, B1, B2, B3 and B4 positions.

20 Whereas the first three features mentioned tend, through introduction of positive charges or elimination of negative charges, to contribute to increasing the pI of a corresponding insulin analogue, the last-mentioned substitutions and/or additions of negatively charged amino acid residues have the

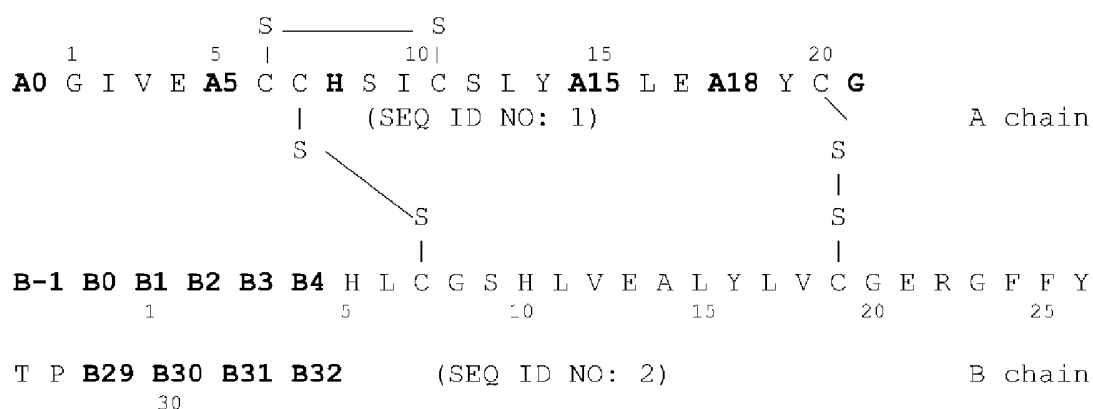
25 opposite effect and contribute to reducing the pI. Surprisingly, precisely the insulin analogues described have the desired advantageous time/action profiles. The pI values of these compounds are lower than that of compound 18 from Kohn et al. (Arg (A0), Gly (A21), Arg (B31), Arg (B32) human

30 insulin), but nevertheless moreover show a delayed onset of action and a longer duration of action, i.e. an extremely shallow and long-lasting, uniform action profile. The risk of hypoglycemic events is thus distinctly minimized. The delay is so marked that it is surprisingly possible to detect the

35 effect even in model experiments on rats, although the delayed action of insulin glargine cannot by contrast be unambiguously observed in rats. Figure 1 shows the hypoglycemic effect of the compound YKL205 of the invention compared with that of

insulin glargine. Similar results are obtained in dogs (see figure 2). Thus, novel basal insulins which need to be administered distinctly less frequently have been provided. Besides these pharmacokinetic advantages described, the analogues of the invention show distinctly better properties compared with insulin glargine in pharmacological respects such as, for example, receptor specificity and in vitro mitogenicity. The claimed insulins also show advantages in physicochemical respects.

The invention thus relates to an insulin analogue of the formula I



where

A0 corresponds to Arg;

A5 corresponds to Asp, Gln or Glu;

A15 corresponds to Asp, Glu or Gln;

A18 corresponds to Asp, Glu or Asn;

B-1 corresponds to Asp, Glu or an amino group;

B0 corresponds to Asp, Glu or a chemical bond;

B1 corresponds to Asp, Glu or Phe;

B2 corresponds to Asp, Glu or Val;

B3 corresponds to Asp, Glu or Asn;

5

B4 corresponds to Asp, Glu or Gln;

B29 corresponds to Lys;

10 B30 corresponds to Thr;

B31 corresponds to Arg or Lys;

B32 corresponds to Arg-NH₂ or Lys-NH₂,

15

where two amino acid residues of the group comprising A5, A15, A18, B-1, B0, B1, B2, B3 and B4 correspond simultaneously and independently of one another to Asp or Glu.

20 The invention relates in particular to insulin analogues as detailed above in which independently of one another A5 corresponds to Glu, or where A15 corresponds to Glu, or where A18 corresponds to Asp, or where B-1 corresponds to an amino group, or where B0 corresponds to Glu, or where B1 corresponds
25 to Asp, or where B2 corresponds to Val, or where B3 corresponds to Asp, or where B4 corresponds to Glu.

The invention particularly preferably relates to an insulin analogue selected from a group comprising:

30

Arg (A0), His (A8), Glu (A5), Asp (A18), Gly (A21), Arg (B31), Arg (B32) - NH₂ human insulin,

Arg (A0), His (A8), Glu (A5), Asp (A18), Gly (A21), Arg (B31), Lys (B32) - NH₂ human insulin,

35 Arg (A0), His (A8), Glu (A15), Asp (A18), Gly (A21), Arg (B31), Arg (B32) - NH₂ human insulin,

Arg (A0), His (A8), Glu (A15), Asp (A18), Gly (A21), Arg (B31), Lys (B32) - NH₂ human insulin,

Arg (A0), His (A8), Glu(A5), Glu (A15), Gly (A21), Arg (B31),
Arg (B32) - NH₂ human insulin,
Arg (A0), His (A8), Glu (A5), Glu (A15), Gly (A21), Arg (B31),
Lys (B32) - NH₂ human insulin,
5 Arg (A0), His(A8), Glu (A5), Gly (A21), Asp (B3), Arg (B31),
Arg (B32) - NH₂ human insulin,
Arg (A0), His(A8), Glu (A5), Gly (A21), Asp (B3), Arg (B31),
Lys (B32) - NH₂ human insulin,
Arg (A0), His (A8), Glu (A15), Gly (A21), Asp (B3), Arg (B31),
10 Arg (B32) - NH₂ human insulin,
Arg (A0), His (A8), Glu (A15), Gly (A21), Asp (B3), Arg (B31),
Lys (B32) - NH₂ human insulin,
Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B3), Arg (B31),
Arg (B32) - NH₂ human insulin,
15 Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B3), Arg (B31),
Lys (B32) - NH₂ human insulin,
Arg (A0), His(A8), Gly (A21), Asp (B3), Glu (B4), Arg (B31),
Arg (B32) - NH₂ human insulin,
Arg (A0), His (A8), Gly (A21), Asp (B3), Glu (B4), Arg (B31),
20 Lys (B32) - NH₂ human insulin,
Arg (A0), His (A8), Glu (A5), Gly (A21), Glu (B4), Arg (B31),
Arg (B32) - NH₂ human insulin,
Arg (A0), His (A8), Glu (A5), Gly (A21), Glu (B4), Arg (B31),
Lys (B32) - NH₂ human insulin,
25 Arg (A0), His (A8), Glu (A15), Gly (A21), Glu (B4), Arg (B31),
Arg (B32) - NH₂ human insulin,
Arg (A0), His (A8), Glu (A15), Gly (A21), Glu (B4), Arg (B31),
Lys (B32) - NH₂ human insulin,
Arg (A0), His (A8), Asp (A18), Gly (A21), Glu (B4), Arg (B31),
30 Arg (B32) - NH₂ human insulin,
Arg (A0), His (A8), Asp (A18), Gly (A21), Glu (B4), Arg (B31),
Lys (B32) - NH₂ human insulin,
Arg (A0), His (A8), Glu (A5), Gly (A21), Glu (B0), Arg (B31),
Arg (B32) - NH₂ human insulin,
35 Arg (A0), His (A8), Glu (A5), Gly (A21), Glu (B0), Arg (B31),
Lys (B32) - NH₂ human insulin,
Arg (A0), His (A8), Glu (A15), Gly (A21), Glu (B0), Arg (B31),
Arg (B32) - NH₂ human insulin,

Arg (A0), His (A8), Glu (A15), Gly (A21), Glu (B0), Arg (B31),
Lys (B32) - NH₂ human insulin,
Arg (A0), His (A8), Asp (A18), Gly (A21), Glu (B0), Arg (B31),
Arg (B32) - NH₂ human insulin,
5 Arg (A0), His (A8), Asp (A18), Gly (A21), Glu (B0), Arg (B31),
Lys (B32) - NH₂ human insulin,
Arg (A0), His (A8), Glu (A5), Gly (A21), Asp (B1), Arg (B31),
Arg (B32) - NH₂ human insulin,
Arg (A0), His (A8), Glu (A5), Gly (A21), Asp (B1), Arg (B31),
10 Lys (B32) - NH₂ human insulin,
Arg (A0), His (A8), Glu (A15), Gly (A21), Asp (B1), Arg (B31),
Arg (B32) - NH₂ human insulin,
Arg (A0), His (A8), Glu (A15), Gly (A21), Asp (B1), Arg (B31),
Lys (B32) - NH₂ human insulin,
15 Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B1), Arg (B31),
Arg (B32) - NH₂ human insulin,
Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B1), Arg (B31),
Lys (B32) - NH₂ human insulin,
Arg (A0), His (A8), Gly (A21), Glu (B0), Asp (B1), Arg (B31),
20 Arg (B32) - NH₂ human insulin,
Arg (A0), His (A8), Gly (A21), Glu (B0), Asp (B1), Arg (B31),
Lys (B32) - NH₂ human insulin,
Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B3), Arg (B30),
Arg (B31) - NH₂ human insulin,
25 Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B3), Arg (B30),
Lys (B31) - NH₂ human insulin.

Specification of the term "human insulin" in the designations
of the insulin analogues mentioned makes reference to the
30 amino acid sequences of the A chain and B chain of human
insulin, and all deviations (additions, substitutions,
deletions) therefrom are indicated in a given designation of
an insulin analogue.

35 The invention further relates to a process for preparing an
insulin analogue as mentioned above, where a precursor of the
insulin analogue is prepared recombinantly, the precursor is
processed enzymatically to two-chain insulin, and a coupling

with argininamide is carried out in the presence of an enzyme having trypsin activity, and the insulin analogue is isolated.

5 The invention further relates to a use of an insulin analogue as described above for the manufacture of a medicament for the treatment of diabetes, in particular of diabetes of type I or type II. The invention likewise relates to a use of an insulin analogue as described above for the manufacture of a medicament for assisting beta cell regeneration.

10

The invention further relates to a pharmaceutical comprising an insulin analogue as described above and/or physiologically acceptable salts thereof.

15 The invention further relates to a formulation of the insulin analogue as described above, where the formulation is in aqueous form comprising the dissolved insulin analogue.

20 The invention further relates to a formulation of the insulin analogue as described above, where the formulation is in the form of powder.

The invention further relates to a formulation as described above, where the insulin analogue as described above is
25 present in crystalline and/or amorphous form.

The invention further relates to a formulation of the insulin analogue as described above, where the formulation is in the form of a suspension.

30

The invention further relates to a formulation of the insulin analogue as described above, where the formulation additionally comprises a chemical chaperone.

35 The invention further relates to a DNA coding for the A chain or B chain of an insulin analogue as described above.

The invention further relates to a vector comprising a DNA as

described above.

The invention further relates to a host organism comprising a DNA as described above or a vector as described above.

5

The invention further relates to a formulation as described above which additionally comprises also a glucagon-like peptide-1 (GLP1) or an analogue or derivative thereof, or exendin-3 or -4 or an analogue or derivative thereof,
10 preferably exendin-4.

The invention further relates to a formulation as described above wherein an analogue of exendin-4 is selected from a group comprising

15

H-desPro³⁶-exendin-4-Lys₆-NH₂,
H-des(Pro^{36,37})-exendin-4-Lys₄-NH₂ and
H-des(Pro^{36,37})-exendin-4-Lys₅-NH₂,
or a pharmacologically tolerable salt thereof.

20

The invention further relates to a formulation as described above wherein an analogue of exendin-4 is selected from a group comprising

25

desPro³⁶ [Asp²⁸]exendin-4 (1-39),
desPro³⁶ [IsoAsp²⁸]exendin-4 (1-39),
desPro³⁶ [Met(O)¹⁴, Asp²⁸]exendin-4 (1-39),
desPro³⁶ [Met(O)¹⁴, IsoAsp²⁸]exendin-4 (1-39),
desPro³⁶ [Trp(O₂)²⁵, Asp²⁸]exendin-2 (1-39),
30 desPro³⁶ [Trp(O₂)²⁵, IsoAsp²⁸]exendin-2 (1-39),
desPro³⁶ [Met(O)¹⁴Trp(O₂)²⁵, Asp²⁸]exendin-4 (1-39) and
desPro³⁶ [Met(O)¹⁴Trp(O₂)²⁵, IsoAsp²⁸]exendin-4 (1-39),
or a pharmacologically tolerable salt thereof.

35

The invention further relates to a formulation as described in the preceding paragraph, where the peptide -Lys₆-NH₂ is attached to the C termini of the analogues of exendin-4.

The invention further relates to a formulation as described above where an analogue of exendin-4 is selected from a group comprising

- 5 H-(Lys)₆- des Pro³⁶ [Asp²⁸]exendin-4(1-39)-Lys₆-NH₂
des Asp²⁸Pro³⁶, Pro³⁷, Pro³⁸ exendin-4(1-39) -NH₂,
H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Asp²⁸]exendin-4(1-39) -NH₂,
H-Asn-(Glu)₅ des Pro³⁶, Pro³⁷, Pro³⁸ [Asp²⁸]exendin-4(1-39) -NH₂,
des Pro³⁶, Pro³⁷, Pro³⁸ [Asp²⁸]exendin-4(1-39)-(Lys)₆-NH₂,
- 10 H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Asp²⁸]exendin-4(1-39)-(Lys)₆-
NH₂,
H-Asn-(Glu)₅- des Pro³⁶, Pro³⁷, Pro³⁸ [Asp²⁸]exendin-4(1-39)-
(Lys)₆-NH₂,
- 15 H-(Lys)₆- des Pro³⁶ [Trp(O₂)²⁵, Asp²⁸]exendin-4(1-39)-Lys₆-NH₂,
H- des Asp²⁸ Pro³⁶, Pro³⁷, Pro³⁸ [Trp(O₂)²⁵]exendin-4(1-39) -NH₂,
H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Trp(O₂)²⁵, Asp²⁸]exendin-4(1-
39) -NH₂,
H-Asn-(Glu)₅- des Pro³⁶, Pro³⁷, Pro³⁸ [Trp(O₂)²⁵, Asp²⁸]exendin-
20 4(1-39) -NH₂,
des Pro³⁶, Pro³⁷, Pro³⁸ [Trp(O₂)²⁵, Asp²⁸]exendin-4(1-39)-(Lys)₆-
NH₂,
H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Trp(O₂)²⁵, Asp²⁸]exendin-4(1-
39)-(Lys)₆-NH₂,
- 25 H-Asn-(Glu)₅- des Pro³⁶, Pro³⁷, Pro³⁸ [Trp(O₂)²⁵, Asp²⁸]exendin-
4(1-39)-(Lys)₆-NH₂,
H-(Lys)₆- des Pro³⁶ [Met(O)¹⁴, Asp²⁸]exendin-4(1-39)-Lys₆-NH₂,
des Met(O)¹⁴ Asp²⁸ Pro³⁶, Pro³⁷, Pro³⁸ exendin-4(1-39) -NH₂,
H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Asp²⁸]exendin-4(1-
30 39) -NH₂,
H-Asn-(Glu)₅- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Asp²⁸] exendin-
4(1-39) -NH₂,
des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Asp²⁸]exendin-4(1-39)-(Lys)₆-
NH₂,
- 35 H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Asp²⁸]exendin-4(1-
39)-Lys₆-NH₂,
H-Asn-(Glu)₅ des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Asp²⁸] exendin-
4(1-39)-(Lys)₆-NH₂,

H-(Lys)₆- des Pro³⁶ [Met(O)¹⁴, Trp(O₂)²⁵, Asp²⁸]exendin-4(1-39)-
 Lys₆-NH₂,
 des Asp²⁸ Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Trp(O₂)²⁵]exendin-4(1-39)
 5 -NH₂,
 H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Trp(O₂)²⁵,
 Asp²⁸]exendin-4(1-39) -NH₂,
 H-Asn-(Glu)₅- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Asp²⁸] exendin-
 4(1-39) -NH₂,
 10 des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Trp(O₂)²⁵, Asp²⁸]exendin-4(1-
 39)-(Lys)₆-NH₂,
 H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Trp(O₂)²⁵,
 Asp²⁸]exendin-4(1-39)-(Lys)₆-NH₂,
 H-Asn-(Glu)₅- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Trp(O₂)²⁵, Asp²⁸]
 15 exendin-4(1-39)-(Lys)₆-NH₂,
 or a pharmacologically tolerable salt thereof.

The invention further relates to a formulation as described
 above which additionally comprises Arg³⁴, Lys²⁶ (N^ε(γ-
 20 glutamyl(N^α-hexadecanoyl))) GLP-1 (7-37) [liraglutide] or a
 pharmacologically tolerable salt thereof.

It is clear to a skilled worker in this connection that the
 insulins of the invention may be item of a pharmaceutical
 25 formulation which has an advantageous effect after
 administration. Aqueous solutions are the starting point in
 this connection. Further components must accordingly be
 miscible. The risk of viral animal contamination is minimized
 in that the preparation ought not to comprise any components
 30 derived from animal sources. It is further advantageous to
 prevent microbial contamination by adding preservatives. It is
 possible by adding isotonic agents to compensate for a
 possible negative effect of the formulation on the physiology
 of the tissue cells at the administration site. The addition
 35 of protamine may have a stabilizing effect, so that
 substantially salt-free insulin preparation can be obtained by
 adding protamine to the formulation. Addition of a phenolic
 component may lead to stabilization of the structure of the

insulin analogue used and thus additionally bring about inter alia the delaying effect on the onset of action. It is also possible to add to the formulation substances which stabilize the spatial structure of the slow insulins of the invention and lead to better thermal stability. Such chemical chaperones may be for example short synthetic peptides, which may also comprise amino acid analogues or include for example peptide sequences derived from the C peptide of insulin.

The insulins of the invention can be incorporated into nanoparticles for developing depot forms. Also conceivable are so-called slow release formulations in which the slow insulin of the invention is present reversibly bound to a polymer carrier.

The insulins of the invention can be administered in parallel with fast-acting insulin such as Apidra®, NovoRapid®, Humalog® or insulin derivatives undergoing development or formulations with an appropriate time/action profile or inhalable insulin or nasally or orally administered insulins which are undergoing development. It will be clear to a skilled worker in this connection that appropriately formulated mixtures of fast-acting and slow insulin of the invention can also be used for this purpose. The insulin analogues of the invention can further be used in pharmaceutical preparations which comprise peptides which are described by an activity comparable to GLP-1 (glucagon like Peptide-1) or exendin-4 or exendin-3. GLP-1 (7-37), exenatide (Byetta®) or peptides whose preparation is described in the patent applications WO 2006/058620, WO 2001/04156, WO 2004/005342 and WO 98/08871 represent examples of such peptides. Formulations particularly advantageous in this connection are those comprising a depot formulation of these peptides. Types of therapy advantageous especially in the initial phase of type II diabetes are those which provide in parallel with the administration of the pharmaceuticals of the invention, which increase the effect of insulin, such as, for example, metformin. Combination therapies with dipeptidyl peptidase-4 inhibitors which increase the level of incretins

are, like combinations with sulfonylureas which increase insulin secretion in the pancreas, likewise possible. The slow insulins of the invention can be employed particularly advantageously when regeneration of pancreatic beta cells from appropriate stem cells is initiated by administration of differentiation factors. All these applications are mentioned by way of example for the therapy of diabetes, and the invention likewise relates thereto. The invention thus further relates to the use of the insulins of the invention in combination with other active ingredients for the treatment of diabetes, especially diabetes of type I or type II diabetes.

The invention further relates to a pharmaceutical which comprises an insulin analogue of the invention which represents in particular an aqueous formulation or a powder.

The pharmaceutical is a pharmaceutical preparation which is preferably a solution or suspension for injection purposes; it is characterized by a content of at least one insulin analogue of the invention, and/or at least one of the physiologically tolerated salts thereof in dissolved, amorphous and/or crystalline - preferably in dissolved - form.

The preparation preferably has a pH of between about 2.5 and 8.5, in particular between 4.0 and 8.5, preferably comprises a suitable tonicity agent, a suitable preservative and, where appropriate, a suitable buffer, and preferably also a particular zinc ion concentration, in sterile aqueous solution. The totality of the preparation ingredients apart from the active ingredient forms the preparation carrier. Suitable tonicity agents are for example glycerol, glucose, mannitol, NaCl, calcium or magnesium compounds such as CaCl_2 etc. The solubility of the insulins of the invention or the physiologically tolerated salts thereof at weakly acidic pH values is influenced by the choice of the tonicity agent and/or preservative.

Examples of suitable preservatives are phenol, m-cresol,

benzyl alcohol and/or p-hydroxybenzoic esters.

Buffer substances which can be used in particular for adjusting a pH between about 4.0 and 8.5 are for example sodium acetate, sodium citrate, sodium phosphate etc. Otherwise, physiologically acceptable dilute acids (typically HCl) or alkalis (typically NaOH) are also suitable for adjusting the pH.

10 If the preparation has a zinc content, preference is given to one of from 1 to 2 mg/ml, in particular from 1 µg/ml to 200 µg zinc/ml.

15 The action profile of the insulin analogues of the invention can surprisingly be influenced satisfactorily by adding Zn. This allows preparations which differ in relation to the total duration of action, the speed of onset of action and the profile of the effect curve and thus allow individual stabilization of the patient. Another possibility arises
20 through the use of a "two-chamber insulin device" which allows a formulation with a rapid onset of action and/or slow gradual onset of action to be administered depending on the life situation.

25 For the purpose of varying the active ingredient profile of the preparation of the invention it is also possible to admix unmodified insulin, preferably bovine, porcine or human insulin, especially human insulin, or insulin analogues and derivatives thereof. It is likewise possible to admix one or
30 more exendin-4 derivatives or peptides which are characterized by an activity comparable to GLP-1 (glucagon like peptide-1) or correspond directly to GLP-1. The invention likewise relates to such pharmaceuticals (preparations).

35 Preferred active ingredient concentrations are those corresponding to about 1-1500, more preferably about 5-1000 and in particular about 40-400 international units/ml.

The insulin analogues of the invention are initially prepared biotechnologically as precursor which does not yet include the amide. The skilled worker is familiar with a large number of possibilities for preparing insulins. Host cell systems used
5 in this connection are bacteria, yeasts and plants or plant cells for cultivation by fermentation. If cost considerations permit, expression systems which use animal cells as host system are also conceivable. However, the precondition therefor is reliable freedom from animal viruses. It is thus
10 clear that the expression systems described by way of example represent only a small segment of the host/vector systems developed for the recombinant preparation of proteins. For example, biotechnological processes based on yeast or plant systems such as mosses, algae or higher plants such as
15 tobacco, pea, safflower, barley, corn or oilseed rape are not described in the application. Nevertheless, the invention likewise includes host/vector systems and coding DNA sequences which allow the target peptides to be prepared in appropriate biotechnological expression systems. Host organisms can thus
20 be selected in particular from the plant kingdom from organisms of the first division Schizophyta comprising Schizomycetes, bacteria or blue algae, organisms of the 2nd division Phycophyta class V Chlorophyceae, organisms of the 2nd division Phycophyta class VII Rhodophyceae, organisms of the
25 3rd division Mycophyta, organisms of the 5th division Bryophyta and organisms of the 7th division Spermatophyta.

European patent application EP-A 1 222 207 describes a plasmid pINT358d which codes for a preproinsulin which includes a
30 modified C peptide. It is now possible with the aid of the polymerase chain reaction (PCR) to modify the proinsulin-encoding sequence specifically so that it is possible to express preproinsulins which can serve as precursors of the insulins of the invention. Corresponding fusion proteins need
35 not necessarily be prepared intracellularly. It is clear to the skilled worker that such proteins can also be prepared by bacterial expression with subsequent secretion into the periplasm and/or into the culture supernatant. European patent

application EP-A 1 364 029 describes this by way of example. The invention likewise relates to the proinsulin precursors which lead to the analogues of the invention.

- 5 The proinsulins prepared in this way can in principle be converted into an insulin analogue precursor which includes lysine or arginine in position A0 and carries lysine or arginine at the C-terminal end of the B chain.
- 10 If the proinsulins of the invention are in the form of inclusion bodies or soluble form after intracellular expression in bacteria, these precursors must be folded by in vitro folding into the correct conformation before the processing and biochemical modification can be undertaken. In
- 15 this connection, the described fusion protein allows direct folding after denaturation by means of urea or guanidinium hydrochloride, and the invention likewise relates to folding intermediates.
- 20 Biochemical methods are used to concentrate the individual intermediates, especially separation processes whose underlying principles are published and in fact the subject of textbooks. It is clear to the skilled worker that such principles can consequently be combined and thus may lead to
- 25 processes which have not previously been published in their sequence. The invention thus likewise relates to processes which lead to purification of the analogues of the invention.

The invention further relates to a process for preparing the

30 insulin analogues of the invention, where a precursor of the insulin analogue is prepared recombinantly and converted enzymatically into a two-chain insulin precursor which carries arginine or lysine N-terminally in relation to amino acid 1 of the A chain, and has at the C-terminal end of the B chain a

35 lysine or arginine residue which is converted with argininamide or lysinamide in the presence of an enzyme having trypsin activity into the amide and thus into the slow insulin of the invention, and is prepared with high purity by a

biochemical purification process.

Proteins which differ through substitution of at least one naturally occurring amino acid residue by other amino acid residues and/or addition and/or deletion of at least one amino acid residue from the corresponding, otherwise identical naturally occurring protein are referred to as "analogues" of proteins. It is also possible in this connection for the added and/or replaced amino acid residues to be ones which do not occur naturally.

Proteins which are obtained by chemical modification of certain amino acid residues of initial proteins are referred to as "derivatives" of proteins. The chemical modification may consist for example of addition of one or more particular chemical groups to one or more amino acids.

Key to figures:

Fig. 1: Blood glucose-lowering effect of novel insulin analogues in rats

Fig. 2: Blood glucose-lowering effect of novel insulin analogues in dogs

Fig. 3: Blood glucose-lowering effect of YKL205 in dogs

Fig. 4: Zinc dependence of the hypoglycemic effect of YKL205 in dogs

The following examples are intended to illustrate the concept of the invention without having a restrictive effect in this connection.

Example 1: Preparation of the vector derivative pINT3580 which codes for Gly (A21)-insulin and a modified C peptide which carries Arg Arg at the C/A chain boundary.

European patent application EP-A 1 222 207 describes the plasmids pINT358d, pINT91d and the primer sequence Tir. DNA of these products is used to construct the plasmid pINT3580. The plasmid pINT358d is moreover characterized by a gene sequence which codes for a modified C peptide having particular properties. Three primer sequences are synthesized:

pint3580_glya21rev

10 5'- CAAAGGTCGACTATTAGGCCGCAGTAGTTCTCCAGCTGG-3' (SEQ ID NO: 3)

This primer serves after working up to introduce glycine (bold print, underlined) instead of asparagin in position 21 of the A chain of the proinsulin sequence encoded by pINT358d.

15 arg_cjuncf

5'-GTCCCTGCAG**CGT**CGCGGCATCGTGGAGCAG-3' (SEQ ID NO: 4)

20 This primer serves like the primer arg_cjunc_rev for introducing arginine instead of lysine at the insulin A/B chain boundary.

arg_cjunc_rev

25 5'- CCACGATGCC GCG**ACG**CTGC AGGGACCCCT CCAGCG-3' (SEQ ID NO: 5)

The codon for the arginine to be introduced is in bold print in both primers.

30 A PCR is carried out in accordance with the European patent application EP-A 1 222 207 with each of the primer pairs Tir / arg_cjunc_rev and arg_cjuncf / pint3580_glya21rev and with DNA of the plasmid pINT358d as template. Aliquots of the products of the two reactions are combined and employed together with the primer pair Tir / pint3580_glya21rev in a third PCR. The product of this reaction is purified after fractionation of the reaction mixture by gel electrophoresis and is digested

with the restriction enzymes Sall / NcoI in accordance with the manufacturer's instructions in one and the same reaction, the reaction mixture is fractionated by gel electrophoresis, and the DNA fragment encoding the proinsulin sequence is isolated. The fragment is then inserted by a DNA ligase reaction into the NcoI / Sall-opened pINT91d vector DNA.

The ligation mixture is used to transform competent E. coli bacterial cells. The transformation mixture is taken out on selection plates which contain 25 mg/l ampicillin. Plasmid DNA is isolated from colonies and characterized by DNA sequence analysis. Correct plasmids are called pINT3580.

Example 2: Construction of the plasmid pINT3581 coding for His (A8), Gly (A21) - preproinsulin

The construction takes place as described in example 1 by 3 polymerase chain reactions. The product of the third reaction is inserted after NcoI / Sall cleavage into the NcoI / Sall-opened pINT91d vector DNA. The primers Tir and pint3580_glya21rev are used. Two further primers are synthesized:

pint3580_Ha8f

5'-AGCAGTGCTGCC**CAC**AGCATCTGCTCCCTCTAC-3' (SEQ ID NO: 6)

pint3580_Ha8rev

5'-GAG CAGATGCT **GTG** GCAGCACTG CTCCACGATG-3' (SEQ ID NO: 7)

The codon which codes for histidine in position 8 of the A chain is emphasized by emboldening in each case. The construction is carried out as described in example 1. Template for PCR1 and 2 is DNA of the plasmid pINT3580. PCR1 is carried out with the primer pair Tir/ pint3580_Ha8rev and PCR2 is carried out with the primer pair pint3580_Ha8f/ pint3580_glya21rev. The primer pair Tir / pint3580_glya21rev

is employed in PCR 3. Template in this case is a mixture of the reaction products of PCR1 and PCR2. Correct plasmids are called pINT3581.

- 5 Example 3: Construction of the plasmid pINT3582 coding for His (A8), Glu (A5), Gly (A21) - preproinsulin

The construction takes place as described in example 1 and 2 by 3 polymerase chain reactions. The product of the third
10 reaction is inserted after Nco1/ Sall cleavage into the Nco1 / Sall-opened pINT91d vector DNA. The primers Tir and pint3580_glya21rev are used. Two further primers are synthesized.

15 pint3581_Ea5f

5'GCATCGTGGAG**GAG**TGCTGCCACAGCATCTG 3' (SEQ ID NO: 8)

pint3581_Ea5rev

20

5'-CTGT GGCAGCA**CTC** CTCCACGATG CCGCGACG-3' (SEQ ID NO: 9)

The codon which codes for glutamic acid in position 5 of the A chain is emphasized by emboldening in each case. The
25 construction is carried out as described in example 1. Template is DNA of the plasmid pINT3581. Correct plasmids are called pINT3582.

- 30 Example 4: Construction of the plasmid pINT3583 coding for His (A8), Asp (A18), Gly(A21) - preproinsulin

The construction differs from example 1 by taking place by only one polymerase chain reaction. The product of this reaction is inserted after Nco1 / Sall cleavage into the Nco1
35 / Sall-opened pINT91d vector DNA. The primer Tir is used. One further primer is synthesized:

pint3580_Da18rev

5' CAAAGGTCGACTATTAGCCGCAGTAG**CTC**CTCCAGCTGGTAGAGGGAG 3' (SEQ ID NO: 10)

5 The codon which codes for aspartic acid in position 18 of the A chain is emphasized by emboldening. Template is DNA of the plasmd pINT3581. Correct plasmids are called pINT3583.

Example 5: Construction of the plasmid pINT3584 coding for His
10 (A8), Glu (A5) Asp (A18), Gly (A21) - preproinsulin

The construction differs from example 1 by taking place by only one polymerase chain reaction. The product of this reaction is inserted after Nco1 / Sal1 cleavage into the Nco1
15 / Sal1-opened pINT91d vector DNA. The primer Tir . pint3580_Dal8rev (ex. 4) is used. Template is DNA of the plasmid pINT3582. Correct plasmids are called pINT3584. The preproinsulin encoded by the plasmid is precursor for the compound YKL205-1 which results after amidation with
20 argininamide and describes the following structure:

Arg (A0), Glu (A5), His (A8), Asp(A18), Gly(A21), Arg(B31), Arg(B32)-NH₂ - human insulin

25 Corresponding amidation with lysinamide leads to the compound YKL205-1b:

Arg (A0), Glu (A5), His (A8), Asp (A18), Gly(A21), Arg (B31), Lys (B32)-NH₂ - human insulin

30

Example 6: Construction of the plasmid pINT3585 coding for His (A8), Glu (A15), Gly (A21) - preproinsulin

The construction differs from example 1 by taking place by
35 only one polymerase chain reaction. The product of this reaction is inserted after Nco1/ Sal1 cleavage into the Nco1 / Sal1-opened pINT91d vector DNA. The primer Tir is used. One further primer is synthesized:

pint3580_Ea15rev

5'- CAAAGGTCGA CTATTAGCCG CAGTAGTTCTCCAG**CTCGTA** GAGGGAGCAG
 5 ATGCTG -3' (SEQ ID NO: 11)

The codon which codes for glutamic acid in position 15 of the A chain is emphasized by emboldening. Template is DNA of the plasmid pINT3581. Correct plasmids are called pINT3585.

10

Example 7: Construction of the plasmid pINT3586 coding for His (A8), Glu (A15), Asp (A18), Gly (A21) - preproinsulin

The construction differs from example 1 by taking place by
 15 only one polymerase chain reaction. The product of this reaction is inserted after Nco1/ Sall cleavage into the Nco1 / Sall-opened pINT91d vector DNA. The primer Tir is used. One further primer is synthesized:

20 pint3585_Ea15_Da18rev

5'- CAAAGGTCGACTATTAGCCGCAGTAG**GTCCTCCAGCTCGT**AGAGGGAGCAG ATGCTG
 -3' (SEQ ID NO: 12)

25 The codon for glutamic acid in position 15 of the A chain and aspartic acid in position A18 of the A chain is emphasized by emboldening in each case. Template is DNA of the plasmid pINT3581. Correct plasmids are called pINT3586. The preproinsulin encoded by the plasmid is precursor for the
 30 compound YKL205 which results after amidation with argininamide and describes the following structure:

Arg (A0), His (A8), Glu (A15), Asp (A18), Gly (A21), Arg (B31), Arg (B32) - NH₂ - human insulin

35

The preproinsulin encoded by the plasmid is precursor for the compound YKL205b which results after amidation with lysinamide and describes the following structure:

Arg (A0), His (A8), Glu (A15), Asp (A18), Gly (A21), Arg (B31),
Lys (B32) - NH₂ - human insulin

- 5 Example 8: Construction of the plasmid pINT3587 coding for
Glu (A5), His (A8), Glu (A15), Gly (A21) - preproinsulin

The construction differs from example 1 by taking place by
only one polymerase chain reaction. The product of this
10 reaction is inserted after Nco1/ Sal1 cleavage into the Nco1 /
Sal1-opened pINT91d vector DNA. The primer Tir and
pint3580_Ea15rev shown in example 6 is used. Template is DNA
of the plasmid pINT3582. Correct plasmids are called pINT3587.
The preproinsulin encoded by the plasmid is precursor for the
15 compound YKL205 -2 which results after amidation with
argininamide and describes the following structure:

Arg (A0), Glu (A5), His (A8), Glu (A15), Gly (A21), Arg (B31),
Arg (B32) - NH₂ - human insulin

20

The preproinsulin encoded by the plasmid is precursor for the
compound YKL205 -2b which results after amidation with
lysinamide and describes the following structure:

- 25 Arg (A0), Glu (A5), His (A8), Glu (A15), Gly (A21), Arg (B31),
Lys (B32) - NH₂ - human insulin

Example 9: Construction of the plasmid pINT3588 coding for His
(A8), Gly (A21), Asp (B3)- preproinsulin

30

Construction takes place as described in example 1 and 2 by 3
polymerase chain reactions. The product of the third reaction
is inserted after Nco1/ Sal1 cleavage into the Nco1 / Sal1-
opened pINT91d vector DNA. The primers Tir and
35 pint3580_glya21rev are used. Two further primers are
synthesized:

pint3581_Db3f

5'- GCACGATTTGTG**GACC**AGCACCTGTGCGGC -3' (SEQ ID NO: 13)

pint3581_Db3rev

5

5'- CACAGG TGCTG**GTCCA** CAAATCGTGC CGAATTTC -3' (SEQ ID NO: 14)

The codon which codes for aspartic acid in position 3 of the insulin B chain is emphasized by emboldening in each case.

10 Construction is carried out as described in example 1. Template is DNA of the plasmid pINT3581. Correct plasmids are called pINT3588.

Example 10: Construction of the plasmid pINT3589 coding for
15 Glu (A5), His (A8), Gly (A21), Asp (B3)- preproinsulin

Carrying out the reactions as described in example 9 but using DNA of the plasmid pINT3582 as template in PCR1 and PCR2 results in plasmid pINT3589.

20

The preproinsulin encoded by the plasmid is precursor for the compound YKL205-3 which results after amidation with argininamide and describes the following structure:

25 Arg (A0), Glu (A5), His (A8), Gly (A21), Asp (B3), Arg (B31), Arg (B32) - NH₂ - human insulin

The preproinsulin encoded by the plasmid is precursor for the compound YKL205-3b which results after amidation with
30 lysinamide and describes the following structure:

Arg (A0), Glu (A5), His (A8), Gly (A21), Asp (B3), Arg (B31), Lys (B32) - NH₂ - human insulin

35 Example 11: Construction of the plasmid pINT3590 coding for His (A8), Glu (A15), Gly (A21), Asp (B3) - preproinsulin

Carrying out the reactions as described in example 9 but using

DNA of the plasmid pINT3585 as template in PCR1 and PCR2 results in plasmid pINT3590. The preproinsulin encoded by the plasmid is precursor for the compound YKL205-4 which results after amidation with argininamide and describes the following structure:

Arg (A0), His (A8), Glu (A15), Gly (A21), Asp (B3), Arg (B31), Arg (B32) - NH₂ - human insulin

10 The preproinsulin encoded by the plasmid is precursor for the compound YKL205-4b which results after amidation with lysinamide and describes the following structure:

Arg (A0), His (A8), Glu (A15), Gly (A21), Asp (B3), Arg (B31),
15 Lys (B32) - NH₂ - human insulin

Example 12: Construction of the plasmid pINT3591 coding for His (A8), Asp (A18), Gly (A21), Asp (B3)- preproinsulin

20 Carrying out the reactions as described in example 9 but using DNA of the plasmid pINT3586 as template in PCR1 and PCR2 results in plasmid pINT3591. The preproinsulin encoded by the plasmid is precursor for the compound YKL205-5 which results after amidation with argininamide and describes the following structure:

Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B3), Arg (B31), Arg (B32) - NH₂ - human insulin

30 The preproinsulin encoded by the plasmid is precursor for the compound YKL205-5b which results after amidation with lysinamide and describes the following structure:

Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B3), Arg (B31),
35 Lys (B32) - NH₂ - human insulin

Example 13: Construction of the plasmid pINT3592 coding for His (A8), Gly (A21), Asp (B3)- Glu (B4) - preproinsulin

Construction takes place as described in example 1 and 2 by 3
polymerase chain reactions. The product of the third reaction
is inserted after Nco1/ Sal1 cleavage into the Nco1 / Sal1-
5 opened pINT91d vector DNA. The primers Tir and
pint3580_glya21rev are used. Two further primers are
synthesized:

pint3581_Db3_Eb4f

10

5'- GCACGATTTGTG**GACGAG**CACCTGTGCGGCTC -3' (SEQ ID NO: 15)

pint3581_Db3_Eb4rev

15 5'- CGCACAGG TG**CTCGTCCA** CAAATCGTGC CGAATTTC -3' (SEQ ID NO:
16)

The codon which codes for aspartic acid in position 3 and
glutamic acid in position 4 of the insulin B chain is
20 emphasized by emboldening in each case. The construction is
carried out as described in example 1. Template is DNA of the
plasmid pINT3581. Correct plasmids are called pINT3592. The
preproinsulin encoded by the plasmid is precursor for the
compound YKL205-6 which results after amidation with
25 argininamide and describes the following structure:

Arg (A0), His (A8), Gly (A21), Asp (B3), Glu (B4), Arg (B31),
Arg (B32) - NH₂ - human insulin

30 The preproinsulin encoded by the plasmid is precursor for the
compound YKL205-6b which results after amidation with
lysineamide and describes the following structure:

35 Arg (A0), His (A8), Gly (A21), Asp (B3), Glu (B4), Arg (B31),
Lys (B32) - NH₂ - human insulin

Example 14: Construction of the plasmid pINT3593 coding for
His (A8), Gly (A21), Glu (B4) - preproinsulin

Construction takes place as described in example 1 and 2 by 3
polymerase chain reactions. The product of the third reaction
is inserted after Nco1/ Sal1 cleavage into the Nco1 / Sal1-
5 opened pINT91d vector DNA. The primers Tir and
pint3580_glya21rev are used. Two further primers are
synthesized:

pint3581_Eb4f

10

5'- ACGATTTGTGAAC**GAG**CACCTGTGCGGCTC -3' (SEQ ID NO: 17)

pint3581_Eb4rev

15

5'- CGCACAGG TG**CTC**GTTCA CAAATCGTGC CGAATTTC -3' (SEQ ID NO:
18)

The codon which codes for glutamic acid in position 4 of the
insulin B chain is emphasized by emboldening. The construction
20 is carried out as described in example 1. Template is DNA of
the plasmid pINT3581. Correct plasmids are called pINT3593.

Example 15: Construction of the plasmid pINT3594 coding for
25 Glu (A5), His (A8), Gly (A21), Glu (B4) - preproinsulin.

Carrying out the reactions as described in example 9 but using
DNA of the plasmid pINT3582 as template in PCR1 and PCR2
results in plasmid pINT3594.

30

The proinsulin is precursor for the compound YKL205-7 which
results after amidation with argininamide and describes the
following structure:

Arg (A0), Glu (A5), His (A8), Gly (A21), Glu (B4), Arg (B31),
35 Arg (B32) - NH₂ - human insulin.

The proinsulin is precursor for the compound YKL205-7b which
results after amidation with lysinamide and describes the

following structure:

Arg (A0), Glu (A5), His (A8), Gly (A21), Glu (B4), Arg (B31),
Lys (B32) - NH₂ - human insulin

- 5 Example 16: Construction of the plasmid pINT3595 coding for
His (A8), Glu (A15), Gly (A21), Glu (B4) - preproinsulin.

Carrying out the reactions as described in example 9 but using
DNA of the plasmid pINT3585 as template in PCR1 and PCR2
10 results in plasmid pINT3595. The preproinsulin encoded by the
plasmid is precursor for the compound YKL205-8 which results
after amidation with argininamide and describes the following
structure:

- 15 Arg (A0), His (A8), Glu (A15), Gly (A21), Glu (B4), Arg (B31),
Arg (B32) - NH₂ - human insulin

The preproinsulin encoded by the plasmid is precursor for the
compound YKL205-8b which results after amidation with
20 lysinamide and describes the following structure:

Arg (A0), His (A8), Glu (A15), Gly (A21), Glu (B4), Arg (B31),
Lys (B32) - NH₂ - human insulin

- 25 Example 17: Construction of the plasmid pINT3596 coding for
His (A8), Asp (A18), Gly (A21), Glu (B4)- preproinsulin

Carrying out the reactions as described in example 9 but using
DNA of the plasmid pINT3586 as template in PCR1 and PCR2
30 results in plasmid pINT3596. The preproinsulin encoded by the
plasmid is precursor for the compound YKL205-9 which results
after amidation with argininamide and describes the following
structure:

- 35 Arg (A0), His (A8), Asp (A18), Gly (A21), Glu (B4), Arg (B31),
Arg (B32) - NH₂ - human insulin

The preproinsulin encoded by the plasmid is precursor for the

compound YKL205-9b which results after amidation with lysinamide and describes the following structure:

Arg (A0), His (A8), Asp (A18), Gly (A21), Glu (B4), Arg (B31),
 5 Lys (B32) - NH₂ - human insulin

Example 18: Construction of the plasmid pINT3597 coding for His (A8), Gly (A21), Glu (B0) - preproinsulin

10 Construction takes place by 2 polymerase chain reactions. The primer pint3580_glya21rev is used. Two further primers are synthesized:

pint3581_Eb0f1

15

5'- CAAC**C**AGGAA ATTCGGCACG **AGAG**TTTGTG AACCAGCACC TGTG-3' (SEQ ID NO: 19)

pint3581_Eb01f2

20

5'- TATCGA CCAT GG CAACAACA TCAAC**C**AGGAA ATTCGGCACG **AGAG**-3' (SEQ ID NO: 20)

There is partial overlap of the two primers in this case.

25 Pint3581_Eb0f2 contains an NcoI recognition sequence. This is depicted underlined. The codon which codes for glutamic acid in position 0 at the start of the B chain is emphasized by boldening in each case. Template for PCR1 is DNA of the plasmid pINT3581.

30

PCR1 is carried out with the primer pair pint3581_Eb-1f2 / pint3580_glya21rev. Template for PCR2 is the product from PCR1. PCR2 is carried out with the primer pair pint3581_Eb-1f2 / pint3580_glya21rev. The product from PCR2 covers the complete
 35 preproinsulin sequence. The product of the second reaction is inserted after NcoI/ SalI cleavage into the NcoI / SalI-opened pINT91d vector DNA. Correct plasmids are called pINT3597. Replacement of the codon for glutamic acid in position B0 by

the codon of aspartic acid and following the example results in plasmids which have aspartic acid instead of glutamic acid in position B0.

- 5 Example 19: Construction of the plasmid pINT3598 coding for Glu (A5), His (A8), Gly (A21), Glu (B0) - preproinsulin

Carrying out the reactions as described in example 18 but using DNA of the plasmid pINT3582 as template in PCR1 results
10 in plasmid pINT3598. The preproinsulin encoded by the plasmid is precursor for the compound YKL205-10 which results after amidation with argininamide and describes the following structure:

- 15 Arg (A0), Glu (A5), His (A8), Gly (A21), Glu (B0), Arg (B31), Arg (B32) - NH₂ - human insulin

The preproinsulin encoded by the plasmid is precursor for the compound YKL205-10b which results after amidation with
20 lysinamide and describes the following structure:

Arg (A0), Glu (A5), His (A8), Gly (A21), Glu (B0), Arg (B31), Lys (B32) - NH₂ - human insulin

- 25 Example 20: Construction of the plasmid pINT3599 coding for His (A8), Glu (A15), Gly (A21), Glu (B0) - preproinsulin

Carrying out the reactions as described in example 18 but using DNA of the plasmid pINT3585 as template in PCR1 results
30 in plasmid pINT3599. The preproinsulin encoded by the plasmid is precursor for the compound YKL205-11 which results after amidation with argininamide and describes the following structure:

- 35 Arg (A0), His (A8), Glu (A15), Gly (A21), Glu (B0), Arg (B31), Arg (B32) - NH₂ - human insulin

The preproinsulin encoded by the plasmid is precursor for the

compound YKL205-11b which results after amidation with lysinamide and describes the following structure:

Arg (A0), His (A8), Glu (A15), Gly (A21), Glu (B0), Arg (B31),
5 Lys (B32) - NH₂ - human insulin

Example 21: Construction of the plasmid pINT3600 coding for His (A8), Asp (A18), Gly (A21), Glu (B0) - preproinsulin

10 Carrying out the reactions as described in example 18 but using DNA of the plasmid pINT3586 as template in PCR1 results in plasmid pINT3600. The preproinsulin encoded by the plasmid is precursor for the compound YKL205-12 which results after amidation with argininamide and describes the following
15 structure:

Arg (A0), His (A8), Asp (A18), Gly (A21), Glu (B0), Arg (B31),
Arg (B32) - NH₂ - human insulin

20 The preproinsulin encoded by the plasmid is precursor for the compound YKL205-12b which results after amidation with lysinamide and describes the following structure:

Arg (A0), His (A8), Asp (A18), Gly (A21), Glu (B0), Arg (B31),
25 Lys (B32) - NH₂ - human insulin

Example 22: Construction of the plasmid pINT3601 coding for His (A8), Gly (A21), Asp (B1) - preproinsulin

30 Construction takes place by 2 polymerase chain reactions. The primer pint3580_glya21rev is used. Two further primers are synthesized:

pint3581_Db1f1

35

5'-CAACAGGAA ATTCGGCACG AGACGTG AACCAGCACC TGTGCG-3' (SEQ ID NO: 21)

pint3581_Db1f2

5'-TATCGA CCAT GG CAACAACA TCAAC**C**AGGAA ATTCGGCACG **AGAC**-3' (SEQ
ID NO: 22)

5

There is partial overlap of the two primers in this case. Pint3581_Db-1f2 contains an NcoI recognition sequence. This is depicted underlined. The codon which codes for aspartic acid in position 1 of the B chain is emphasized by boldening in each case. Template for PCR1 is DNA of the plasmid pINT3581. PCR1 is carried out with the primer pint3581_Db1f1 / pint3580_glya21rev. Template for PCR2 is the product from PCR1. PCR2 is carried out with the primer pair pint3581_Db1f2 / pint3580_glya21rev. The product from PCR2 covers the complete preproinsulin sequence. The product of the second reaction is inserted after NcoI/ SalI cleavage into the NcoI / SalI-opened pINT91d vector DNA. Correct plasmids are called pINT3601.

Example 23: Construction of the plasmid pINT3602 coding for Glu (A5), His (A8), Gly (A21), Asp (B1) - preproinsulin

Carrying out the reactions as described in example 22 by using DNA of the plasmid pINT3582 as template in PCR1 results in plasmid pINT3602. The preproinsulin encoded by the plasmid is precursor for the compound YKL205-13 which results after amidation with argininamide and describes the following structure:

Arg (A0), Glu (A5), His (A8), Gly (A21), Asp (B1), Arg (B31), Arg (B32) - NH₂ - human insulin

The preproinsulin encoded by the plasmid is precursor for the compound YKL205-13b which results after amidation with lysinamide and describes the following structure:

35

Arg (A0), Glu (A5), His (A8), Gly (A21), Asp (B1), Arg (B31), Lys (B32) - NH₂ - human insulin

Example 24: Construction of the plasmid pINT3603 coding for His (A8), Glu (A15), Gly (A21), Asp (B1) - preproinsulin

5 Carrying out the reactions as described in example 22 by using DNA of the plasmid pINT3585 as template in PCR1 results in plasmid pINT3603. The preproinsulin encoded by the plasmid is precursor for the compound YKL205-14 which results after amidation with argininamide and describes the following
10 structure:

Arg (A0), His (A8), Glu (A15), Gly (A21), Asp (B1), Arg (B31), Arg (B32) - NH₂ - human insulin

15 The preproinsulin coded by the plasmid is precursor for the compound YKL205-14b which results after amidation with lysinamide and describes the following structure:

Arg (A0), His (A8), Glu (A15), Gly (A21), Asp (B1), Arg (B31),
20 Lys (B32) - NH₂ - human insulin

Example 25: Construction of the plasmid pINT3604 coding for His (A8), Asp (A18), Gly (A21), Asp (B1) - preproinsulin

25 Carrying out the reactions as described in example 22 but using DNA of the plasmid pINT3586 as template in PCR1 results in plasmid pINT3604. The preproinsulin encoded by the plasmid is precursor for the compound YKL205-15 which results after amidation with argininamide and describes the following
30 structure:

Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B1), Arg (B31), Arg (B32) - NH₂ - human insulin

35 The preproinsulin encoded by the plasmid is precursor for the compound YKL205-15b which results after amidation with lysinamide and describes the following structure:

Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B1), Arg (B31),
Lys (B32) - NH₂ - human insulin

5 Example 26: Construction of the plasmid pINT3605 coding for
His (A8), Gly (A21), Glu (B0), Asp (B1) - preproinsulin

Construction takes place by 2 polymerase chain reactions. The
primer pint3580_glya21rev and the primer pint3581_Eb01f2
described in example 18 is used. The primer pint3597_Db1f is
10 synthesized:

5'-CAACAGGAA ATTCGGCACG **AGAGGAC**GTG AACCAGCACC TGTGC-3' (SEQ ID
NO: 23)

15 The codon which codes for glutamic acid in position 0 and
which codes for aspartic acid in each case at the start of the
B chain is emphasized by boldening in each case. Template
for PCR1 is DNA of the plasmid pINT3597. PCR1 is carried out
with the primer pair pint3597_Db1f/ pint3580_glya21rev.
20 Template for PCR2 is the product from PCR1. PCR2 is carried
out with the primer pair pint3581_Eb1f2 / pint3580_glya21rev.
The product from PCR2 covers the complete preproinsulin
sequence. The product of the second reaction is inserted after
Nco1/ Sal1 cleavage into the Nco1 / Sal1-opened pINT91d vector
25 DNA. Correct plasmids are called pINT3605. The preproinsulin
encoded by the plasmid is precursor for the compound YKL205-16
which results after amidation with argininamide and describes
the following structure:

30 Arg (A0), His (A8), Gly (A21), Glu (B0), Asp (B1), Arg (B31),
Arg (B32) - NH₂ - human insulin

The preproinsulin encoded by the plasmid is precursor for the
compound YKL205-16a which results after amidation with
35 lysinamide and describes the following structure:

Arg (A0), His (A8), Gly (A21), Glu (B0), Asp (B1), Arg (B31),
Lys (B32) - NH₂ - human insulin

Example 27: Construction of the plasmid pINT3606 coding for His (A8), Glu (A15), Asp (A18), Gly (A21), desThr (B30) - preproinsulin

5

Construction takes place as described in example 1 and 2 by 3 polymerase chain reactions. The primers Tir and pint3580_glya21rev are used. Two further primers are synthesized:

10

desB30f

5'-TTCTACACACCC**AAG**CGCGATGTTTCCTCAGGTGG-3' (SEQ ID NO: 24)

15

desB30rev

5'-AGG AACATCGCGC TTGGGTGTGT AGAAGAAGC-3' (SEQ ID NO: 25)

Template for PCR1 and PCR2 is DNA of the plasmid pINT3586.

20

PCR1 is carried out with the primer pair desB30f / pint3580_glya21rev and PCR2 is carried out with the primer pair Tir / desB30rev template. The template used for PCR3 is an equimolar mixture of the products from PCR1 and PCR2. The reaction is carried out with the primer pair Tir/ pint3580_glya21rev. The product from PCR3 covers the complete preproinsulin sequence. The product of the third reaction is inserted after Nco1/ Sal1 cleavage into the Nco1 / Sal1-opened pINT91d vector DNA. The preproinsulin encoded by the plasmid is precursor for the compound YKL205-17 which results after amidation with argininamide and describes the following structure:

25

30

Arg (A0), His (A8), Glu (A15), Asp (A18), Gly (A21), Arg (B30), Arg (B31) - NH₂ - human insulin

35

The preproinsulin encoded by the plasmid is precursor for the compound YKL205-17b which results after amidation with lysinamide and describes the following structure:

Arg (A0), His (A8), Glu (A15), Asp (A18), Gly (A21), Arg (B30),
Lys (B31) - NH₂ - human insulin

5 Example 28: Expression of the proinsulin derivatives

The expression is carried out in accordance with example 1 of
European patent application EP-A 1 222 207.

10 Example 29: Folding of the proinsulin derivatives

The folding takes place in principle by the method described
in EP-A 0 668 282

15 Example 30: Enzymatic processing of the folded preproinsulin
to give the 2-chain Arg(A0) - insulin precursor whose C-
terminal B chain end is characterized by lysine or arginine.

The enzymatic processing of the folded preproinsulin precursor
20 takes place as described for example in example 4 of
W091/03550. It proves to be particularly advantageous in this
case to employ the trypsin variant described in WO 2007/031187
A1.

25 Example 31: Preparation of an Arg (A0), His (A8), Gly (A21),
Arg (B31), Arg (B32) - NH₂- human insulin

Irrespective of the positioning of the additional acidic amino
acids, a standard reaction is carried out as follows: 100 mg
30 of Arg (A0), Gly (A21), Arg (B31) - insulin analogue are
dissolved in 0.95 ml of argininamide solution (446 g/L), and
0.13 mL of M Na acetate buffer (pH 5.8) and 2 ml of DMF are
added. The reaction mixture is cooled to 12°C and started by
adding 0.094 ml of trypsin (0.075 mg, Roche Diagnostics). The
35 reaction is stopped after 8 h by adding TFA to pH 2.5 and
analyzed by HPLC. There is formation of >60% Arg (A0), Gly
(A21), Arg (B31), Arg (B32) - NH₂- human insulin. Addition of
trypsin inhibitor solution is followed by purification of the

amidated analogue in analogy to US 5,656,722.

Preparation of the corresponding lysinamide compound takes place analogously. However, an aqueous lysinamide stock solution containing 366g/L lysinamide in solution forms the starting material.

Example 32: Formulation of the amidated insulin derivatives

In order to test the insulin derivatives of the invention for their biopharmacological and physicochemical properties, a solution of the compounds was prepared as follows: the insulin derivative of the invention was dissolved with a target concentration of $240 \pm 5 \mu\text{M}$ in 1 mM hydrochloric acid with 80 $\mu\text{g/mL}$ zinc (as zinc chloride).

The following compositions were used as dissolving medium:

- a) 1 mM hydrochloric acid
- b) 1 mM hydrochloric acid, 5 $\mu\text{g/mL}$ zinc (added as zinc chloride or hydrochloric acid)
- c) 1 mM hydrochloric acid, 10 $\mu\text{g/mL}$ zinc (added as zinc chloride or hydrochloric acid)
- d) 1 mM hydrochloric acid, 15 $\mu\text{g/mL}$ zinc (added as zinc chloride or hydrochloric acid)
- e) 1 mM hydrochloric acid, 30 $\mu\text{g/mL}$ zinc (added as zinc chloride or hydrochloric acid)
- f) 1 mM hydrochloric acid, 80 $\mu\text{g/mL}$ zinc (added as zinc chloride or hydrochloric acid)
- g) 1 mM hydrochloric acid, 120 $\mu\text{g/mL}$ zinc (added as zinc chloride or hydrochloric acid)

For this purpose, initially an amount of the freeze-dried material which is about 30% higher than required on the basis of the molecular weight and the desired concentration was weighed out. The concentration present was then determined by analytical HPLC and the solution was subsequently made up to the volume necessary to achieve the target concentration with

5 mM hydrochloric acid with 80 µg/mL zinc. If necessary, the pH was readjusted to 3.5 ± 0.1 . After the final analysis by HPLC to verify the target concentration of 240 ± 5 µM, the finished solution was transferred by means of a syringe with a 0.2 µm filter attachment into a sterile vial which was closed with a septum and a crimped cap. No optimization of the formulations, e.g. in relation to addition of isotonic agents, preservatives or buffer substances, was carried out for the short-term single testing of the insulin derivatives of the invention.

Example 33: Evaluation of the blood glucose-lowering effect of novel insulin analogues in rats

The blood glucose-lowering effect of selected novel insulin analogues is tested in healthy male normoglycemic Wistar rats. Male rats receive subcutaneous injection of a dose of 9 nmol/kg of an insulin analogue. Blood samples are taken from the animals immediately before the injection of the insulin analogue and at regular intervals up to eight hours after the injection, and the blood glucose content therein is determined. The experiment shows clearly (cf. fig. 1) that the employed insulin analogue of the invention leads to a distinctly delayed onset of action and a longer, uniform duration of action.

Example 34: Evaluation of the blood glucose-lowering effect of novel insulin analogues in dogs

The blood glucose-lowering effect of selected novel insulin analogues is tested in healthy male normoglycemic beagle dogs. Male animals receive subcutaneous injection of a dose of 6 nmol/kg of an insulin analogue. Blood samples are taken from the animals immediately before the injection of the insulin analogue and at regular intervals up to 48 hours after the injection, and the blood glucose content therein is determined. The experiment shows clearly (cf. fig. 2) that the employed insulin analogue of the invention leads to a

distinctly delayed onset of action and a longer, uniform duration of action.

Example 35: Evaluation of the blood glucose-lowering effect in dogs with a dose increased two-fold

The blood glucose-lowering effect of selected novel insulin analogues is tested in healthy male normoglycemic beagle dogs. Male animals receive subcutaneous injection of a dose of 6 nmol/kg and 12 nmol/kg of an insulin analogue. Blood samples are taken from the animals immediately before the injection of the insulin analogue and at regular intervals up to 48 hours after the injection, and the blood glucose content therein is determined. The experiment shows clearly (cf. fig. 3) that the employed insulin analogue of the invention has a dose-dependent effect but that, despite the dose being increased two-fold, the action profile has a shallow profile, i.e. no pronounced low point (nadir) is observed. It can be deduced from this that the insulins of the invention lead to distinctly fewer hypoglycemic events by comparison with known slow insulins.

Example 36: Evaluation of the blood glucose-lowering effect in dogs with different zinc concentrations in the formulation

The experiments were carried out as described in Example 35. Figure 4 shows the result. According to this, the time-effect curve of the insulin analogue of the invention can be influenced through the content of zinc ions in the formulation with the same insulin concentration in such a way that a rapid onset of action is observed with a zero or low zinc content, and the effect is maintained for 24 hours, whereas a gradual onset of action is observed with a higher zinc content, and the insulin effect is maintained for distinctly longer than 24 hours.

SEQUENCE LISTING

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<140> 102008003569.8-41

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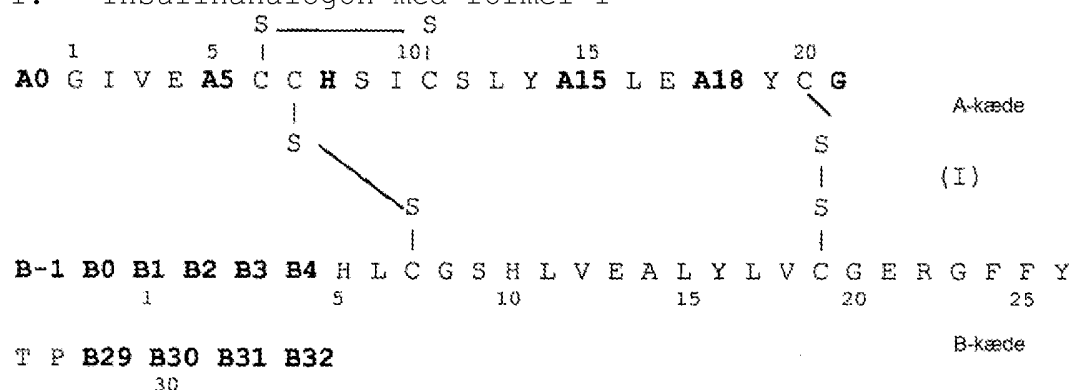
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Patentkrav

1. Insulinalanalogon med formel I



- 5 idet
- A0 repræsenterer Arg;
- A5 repræsenterer Asp, Gln eller Glu;
- A15 repræsenterer Asp, Glu eller Gln;
- A18 repræsenterer Asp, Glu eller Asn;
- 10 B-1 repræsenterer Asp, Glu eller en aminogruppe;
- B0 repræsenterer Asp, Glu eller en kemisk binding;
- B1 repræsenterer Asp, Glu eller Phe;
- B2 repræsenterer Asp, Glu eller Val;
- B3 repræsenterer Asp, Glu eller Asn;
- 15 B4 repræsenterer Asp, Glu eller Gln;
- B29 repræsenterer Lys;
- B30 repræsenterer Thr;
- B31 repræsenterer Arg eller Lys;
- B32 repræsenterer Arg-NH₂ eller Lys-NH₂
- 20 idet to aminosyrerester fra gruppen indeholdende A5, A15, A18, B-1, B0, B1, B2, B3 og B4 samtidigt og uafhængigt af hinanden repræsenterer Asp eller Glu.
- 25 2. Insulinalanalogon ifølge et eller flere af de foregående krav, idet A5 repræsenterer Glu.
3. Insulinalanalogon ifølge et eller flere af de foregående krav, idet A15 repræsenterer Glu.
- 30 4. Insulinalanalogon ifølge et eller flere af de foregående krav, idet A18 repræsenterer Asp.

5. Insulinanalogon ifølge et eller flere af de foregående krav, idet B-1 repræsenterer en aminogruppe.

5 6. Insulinanalogon ifølge et eller flere af de foregående krav, idet B0 repræsenterer Glu.

7. Insulinanalogon ifølge et eller flere af de foregående krav, idet B1 repræsenterer Asp.

10

8. Insulinanalogon ifølge et eller flere af de foregående krav, idet B2 repræsenterer Val.

9. Insulinanalogon ifølge et eller flere af de foregående
15 krav, idet B3 repræsenterer Asp.

10. Insulinanalogon ifølge et eller flere af de foregående krav, idet B4 repræsenterer Glu.

20 11. Insulinanalogon ifølge et eller flere af de foregående krav udvalgt af en gruppe indeholdende:

Arg (A0), His (A8), Glu (A5), Asp (A18), Gly (A21), Arg (B31), Arg (B32) - NH₂ humaninsulin,

Arg (A0), His (A8), Glu (A5), Asp (A18), Gly (A21), Arg (B31),
25 Lys (B32) - NH₂ humaninsulin,

Arg (A0), His (A8), Glu (A15), Asp (A18), Gly (A21), Arg (B31), Arg (B32) - NH₂ humaninsulin,

Arg (A0), His (A8), Glu (A15), Asp (A18), Gly (A21), Arg (B31), Lys (B32) - NH₂ humaninsulin,

30 Arg (A0), His (A8), Glu(A5), Glu (A15), Gly (A21), Arg (B31), Arg (B32) - NH₂ humaninsulin,

Arg (A0), His (A8), Glu (A5), Glu (A15), Gly (A21), Arg (B31), Lys (B32) - NH₂ humaninsulin,

Arg (A0), His(A8), Glu (A5), Gly (A21), Asp (B3), Arg (B31),
35 Arg (B32) - NH₂ humaninsulin,

Arg (A0), His(A8), Glu (A5), Gly (A21), Asp (B3), Arg (B31), Lys (B32) - NH₂ humaninsulin,

Arg (A0), His (A8), Glu (A15), Gly (A21), Asp (B3), Arg (B31),

Arg (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Glu (A15), Gly (A21), Asp (B3), Arg (B31),
 Lys (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B3), Arg (B31),
 5 Arg (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B3), Arg (B31),
 Lys (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Gly (A21), Asp (B3), Glu (B4), Arg (B31),
 Arg (B32) - NH₂ humaninsulin,
 10 Arg (A0), His (A8), Gly (A21), Asp (B3), Glu (B4), Arg (B31),
 Lys (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Glu (A5), Gly (A21), Glu (B4), Arg (B31),
 Arg (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Glu (A5), Gly (A21), Glu (B4), Arg (B31),
 15 Lys (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Glu (A15), Gly (A21), Glu (B4), Arg (B31),
 Arg (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Glu (A15), Gly (A21), Glu (B4), Arg (B31),
 Lys (B32) - NH₂ humaninsulin,
 20 Arg (A0), His (A8), Asp (A18), Gly (A21), Glu (B4), Arg (B31),
 Arg (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Asp (A18), Gly (A21), Glu (B4), Arg (B31),
 Lys (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Glu (A5), Gly (A21), Glu (B0), Arg (B31),
 25 Arg (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Glu (A5), Gly (A21), Glu (B0), Arg (B31),
 Lys (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Glu (A15), Gly (A21), Glu (B0), Arg (B31),
 Arg (B32) - NH₂ humaninsulin,
 30 Arg (A0), His (A8), Glu (A15), Gly (A21), Glu (B0), Arg (B31),
 Lys (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Asp (A18), Gly (A21), Glu (B0), Arg (B31),
 Arg (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Asp (A18), Gly (A21), Glu (B0), Arg (B31),
 35 Lys (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Glu (A5), Gly (A21), Asp (B1), Arg (B31),
 Arg (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Glu (A5), Gly (A21), Asp (B1), Arg (B31),

Lys (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Glu (A15), Gly (A21), Asp (B1), Arg (B31),
 Arg(B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Glu (A15), Gly (A21), Asp (B1), Arg (B31),
 5 Lys (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B1), Arg (B31),
 Arg (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B1), Arg (B31),
 Lys (B32) - NH₂ humaninsulin,
 10 Arg (A0), His (A8), Gly (A21), Glu (B0), Asp (B1), Arg (B31),
 Arg (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Gly (A21), Glu (B0), Asp (B1), Arg (B31),
 Lys (B32) - NH₂ humaninsulin,
 Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B3), Arg (B30),
 15 Arg (B31) - NH₂ humaninsulin,
 Arg (A0), His (A8), Asp (A18), Gly (A21), Asp (B3), Arg (B30),
 Lys (B31) - NH₂ humaninsulin.

12. Fremgangsmåde til fremstilling af et insulinanalogon
 20 ifølge et af kravene 1 til 11, idet der rekombinant
 fremstilles en forløber af insulinanalogonet, forløberen
 enzymatisk processeres til dobbeltkædet insulin, og der
 gennemføres en kobling med argininamid i nærvær af et enzym
 med trypsinaktivitet, og insulinanalogonet isoleres.

25

13. Anvendelse af et insulinanalogon ifølge et af kravene 1
 til 11 til fremstilling af et medikament til behandling af
 diabetes mellitus.

30 14. Anvendelse af et insulinanalogon ifølge et af kravene 1
 til 11 i en fremgangsmåde til fremstilling af et medikament
 til behandling af diabetes mellitus type I eller type II eller
 til terapeutisk understøttelse af beta-celleregenerationen.

35 15. Lægemiddel indeholdende et insulinanalog ifølge et af
 kravene 1 til 11 og/eller fysiologisk forligelige salte heraf.

16. Formulering af insulinanalogonet ifølge et af kravene 1

til 11, idet formuleringen foreligger i vandig form indeholdende det opløste insulinanalogon.

17. Formulering af insulinanalogonet ifølge et af kravene 1 til 11, idet formuleringen foreligger i form af pulver.

18. Formulering ifølge krav 17, idet insulinanalogonet ifølge et af kravene 1 til 11 er indeholdt i krystallin og/eller amorf form.

10

19. Formulering af insulinanalogonet ifølge et af kravene 1 til 11, idet formuleringen foreligger i form af en suspension.

15

20. Formulering af insulinanalogonet ifølge et af kravene 1 til 11, idet formuleringen desuden indeholder en kemisk chaperon.

20

21. DNA kodende for en præ-proinsulin af et insulinanalogon ifølge et af kravene 1 til 11.

22. Vektor indeholdende et DNA ifølge krav 21.

25

23. Ikke-human værtsorganisme indeholdende et DNA ifølge krav 21 eller en vektor ifølge krav 22.

24. Formulering ifølge et eller flere af kravene 16 til 20, hvori der endvidere også er indeholdt et glucagon-lignende peptid-1 (GLP1) eller et analogon eller derivat heraf eller exendin-3 eller -4 eller et analogon eller derivat heraf.

30

25. Formulering ifølge krav 24, hvori der endvidere er indeholdt exendin-4.

35

26. Formulering ifølge krav 24, hvori et analogon af exendin-4 udvælges af en gruppe indeholdende
H-desPro³⁶-exendin-4-Lys₆-NH₂,
H-des(Pro^{36,37})-exendin-4-Lys₄-NH₂ og
H-des(Pro^{36,37})-exendin-4-Lys₅-NH₂, eller et farmakologisk

acceptabelt salt heraf.

27. Formulering ifølge krav 24, hvori et analogon af exendin-4 udvælges af en gruppe indeholdende

- 5 desPro³⁶ [Asp²⁸]exendin-4 (1-39),
 desPro³⁶ [IsoAsp²⁸]exendin-4 (1-39),
 desPro³⁶ [Met(O)¹⁴, Asp²⁸]exendin-4 (1-39),
 desPro³⁶ [Met(O)¹⁴, IsoAsp²⁸]exendin-4 (1-39),
 desPro³⁶ [Trp(O₂)²⁵ Asp²⁸]exendin-2 (1-39),
 10 desPro³⁶ [Trp(O₂)²⁵, IsoAsp²⁸]exendin-2 (1-39),
 desPro³⁶ [Met(O)¹⁴Trp(O₂)²⁵, Asp²⁸]exendin-4 (1-39) og
 desPro³⁶ [Met(O)¹⁴Trp(O₂)²⁵, IsoAsp²⁸]exendin-4 (1-39), eller et
 farmakologisk acceptabelt salt heraf.

- 15 28. Formulering ifølge krav 27, hvori der på C-terminiene af
 analoger af exendin-4 er tilføjet peptidet -Lys₆-NH₂.

29. Formulering ifølge krav 24, hvori der udvælges et
 analogon af exendin-4 af en gruppe indeholdende

- 20 H-(Lys)₆- des Pro³⁶ [Asp²⁸]exendin-4(1-39)-Lys₆-NH₂
 des Asp²⁸Pro³⁶, Pro³⁷, Pro³⁸ exendin-4(1-39) -NH₂,
 H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Asp²⁸]exendin-4(1-39)-NH₂,
 H-Asn-(Glu)₅ des Pro³⁶, Pro³⁷, Pro³⁸ [Asp²⁸]exendin-4(1-39)-NH₂,
 des Pro³⁶, Pro³⁷, Pro³⁸ [Asp²⁸]exendin-4(1-39)-(Lys)₆-NH₂,
 25 H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Asp²⁸]exendin-4(1-39)-(Lys)₆-
 NH₂,
 H-Asn-(Glu)₅- des Pro³⁶, Pro³⁷, Pro³⁸ [Asp²⁸]exendin-4(1-39)-
 (Lys)₆-NH₂,
 H-(Lys)₆- des Pro³⁶ [Trp(O₂)²⁵, Asp²⁸]exendin-4(1-39)-Lys₆-NH₂,
 30 H- des Asp²⁸ Pro³⁶, Pro³⁷, Pro³⁸ [Trp(O₂)²⁵]exendin-4(1-39)-NH₂,
 H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Trp(O₂)²⁵, Asp²⁸]exendin-4(1-
 39) -NH₂,
 H-Asn-(Glu)₅- des Pro³⁶, Pro³⁷, Pro³⁸ [Trp(O₂)²⁵, Asp²⁸]exendin-
 4(1-39) -NH₂,
 35 des Pro³⁶, Pro³⁷, Pro³⁸ [Trp(O₂)²⁵, Asp²⁸]exendin-4(1-39)-(Lys)₆-
 NH₂,
 H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Trp(O₂)²⁵, Asp²⁸]exendin-4(1-
 39)-(Lys)₆-NH₂,

H-Asn-(Glu)₅- des Pro³⁶, Pro³⁷, Pro³⁸ [Trp(O₂)²⁵, Asp²⁸]exendin-4(1-39)-(Lys)₆-NH₂,

H-(Lys)₆- des Pro³⁶ [Met(O)¹⁴, Asp²⁸]exendin-4(1-39)-Lys₆-NH₂,
des Met(O)¹⁴ Asp²⁸ Pro³⁶, Pro³⁷, Pro³⁸ exendin-4(1-39) -NH₂,

5 H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Asp²⁸]exendin-4(1-39) -NH₂,

H-Asn-(Glu)₅- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Asp²⁸] exendin-4(1-39) -NH₂,

des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Asp²⁸]exendin-4(1-39)-(Lys)₆-
10 NH₂,

H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Asp²⁸]exendin-4(1-39)-Lys₆-NH₂,

H-Asn-(Glu)₅ des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Asp²⁸] exendin-4(1-39)-(Lys)₆-NH₂,

15 H-(Lys)₆- des Pro³⁶ [Met(O)¹⁴, Trp(O₂)²⁵, Asp²⁸]exendin-4(1-39)-Lys₆-NH₂,

des Asp²⁸ Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Trp(O₂)²⁵]exendin-4(1-39)-NH₂,

H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Trp(O₂)²⁵,
20 Asp²⁸]exendin-4(1-39) -NH₂,

H-Asn-(Glu)₅- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Asp²⁸] exendin-4(1-39) -NH₂,

des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Trp(O₂)²⁵, Asp²⁸]exendin-4(1-39)-(Lys)₆-NH₂,

25 H-(Lys)₆- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Trp(O₂)²⁵, Asp²⁸]exendin-4(1-39)-(Lys)₆-NH₂,

H-Asn-(Glu)₅- des Pro³⁶, Pro³⁷, Pro³⁸ [Met(O)¹⁴, Trp(O₂)²⁵, Asp²⁸]exendin-4(1-39)-(Lys)₆-NH₂, eller et farmakologisk acceptabelt salt heraf.

30

30. Formulering ifølge krav 24, hvori der endvidere er indeholdt Arg³⁴, Lys²⁶ (N^ε(γ-glutamyl(N^α-hexadecanoyl))) GLP-1 (7-37) [liraglutider] eller et farmakologisk acceptabelt salt heraf.

35

31. Vandig formulering af insulinanalogonet ifølge et af kravene 1 til 11, som indeholder ingen eller mindre end 15 µg/ml zink.

32. Vandig formulering af insulinanalogonet ifølge et af kravene 1 til 11, som indeholder ingen eller mindre end 15 µg/ml til 2 mg/ml zink.

5

33. Formulering ifølge krav 32, idet der er indeholdt 200 µg/ml zink.

Fig. 1

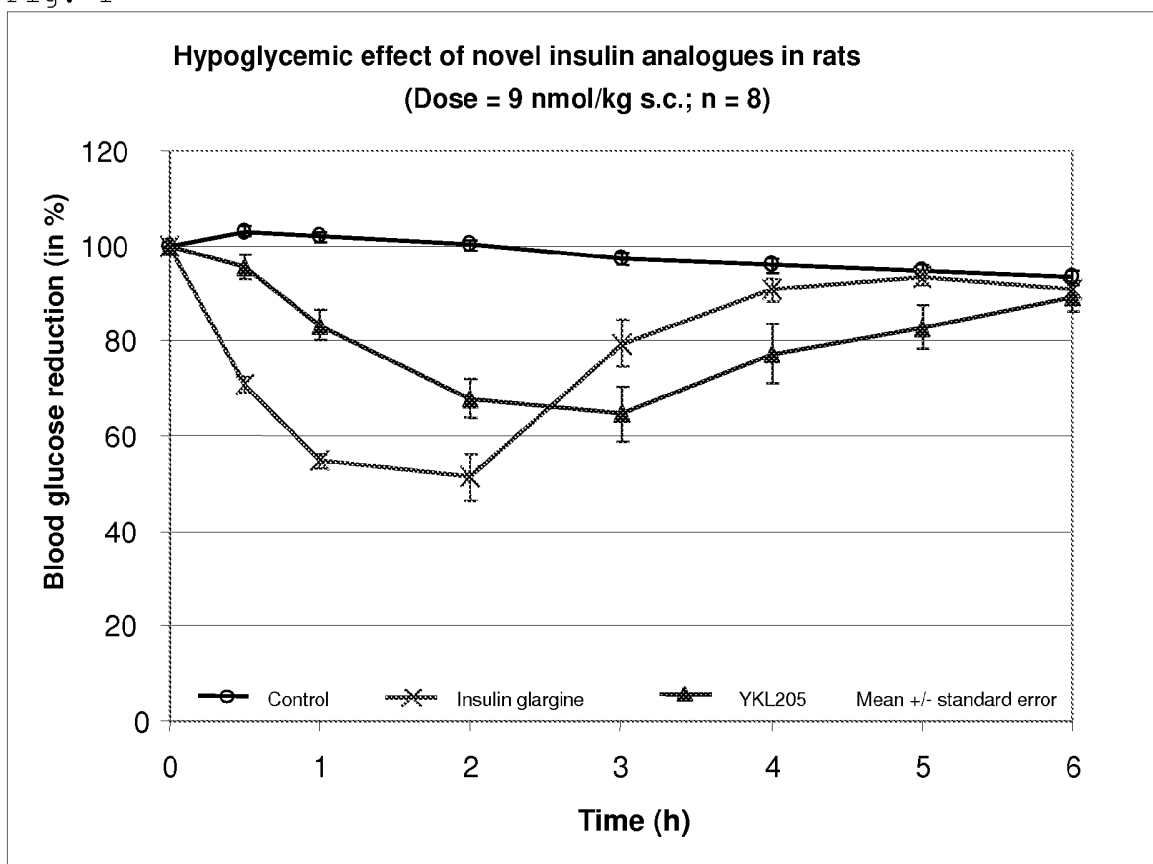


Fig. 2

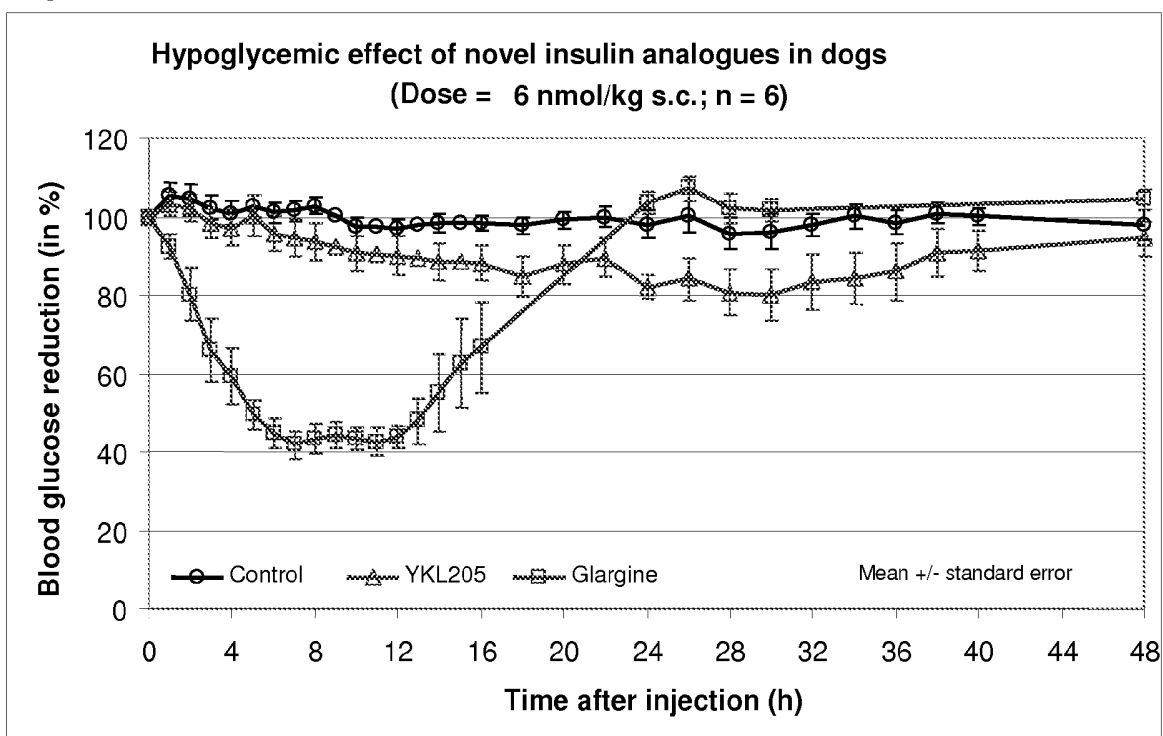


Fig. 3

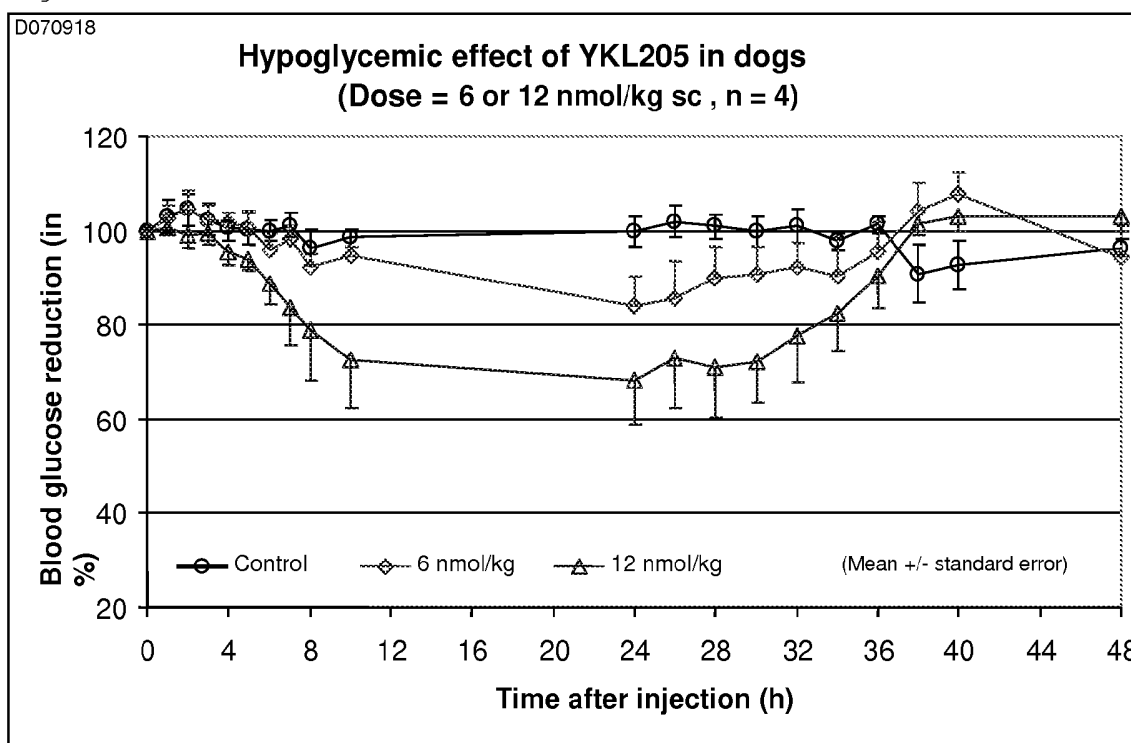


Fig. 4

