The invention relates to a modified tandem LHRH-peptide vaccine preparation in which the amino acid glycine at position (6) of one or both LHRH decapptides that constitute the tandem unit is substituted by a dextrorotatory amino acid that contains a side chain to which a carrier compound can be coupled. In addition, the tandem LHRH-peptide can be brought into a tandem-dimer form which is also suitable for producing a vaccine that is effective against LHRH (luteinizing hormone releasing hormone), also referred to as GaRH (gonadotropin-releasing hormone), for immunological castration to inhibit or affect reproductive functions or to affect behaviour in vertebrates in general and in domesticated animals and man in particular.

15 Claims, No Drawings
PEPTIDE, IMMUNOGENIC COMPOSITION AND VACCINE OR MEDICAL PREPARATION, A METHOD TO IMMUNISE ANIMALS AGAINST THE HORMONE LH-RH, ANALOGS OF THE LH-RH TANDEM, REPEAT PEPTIDE AND THEIR USE AS VACCINE

Matter enclosed in heavy brackets [ ] appears in the original patent but forms no part of this reissue specification; matter printed in italics indicates the additions made by reissue.

CROSS-REFERENCE TO RELATED APPLICATIONS

This application is a national entry under 35 U.S.C. § 371 PCT International Application Number PCT/NI96/00223 filed on Jun. 6, 1996, designating the United States of America which itself is a continuation in part from both U.S. patent application Ser. No. 08/477,298 filed on Jun. 7, 1995, now abandoned; and U.S. patent application Ser. No. 08/476,013, filed on Jun. 7, 1995, now abandoned.

This invention relates to a peptide suitable for producing a vaccine effective against the Luteinising Hormone Releasing Hormone (LH-RH), also referred to as Gonadotrophin Releasing Hormone (GnRH). The invention further relates to immunogenic compositions and vaccines or medicinal preparations (vaccines and pharmaceuticals) based on such a peptide and the use of such a vaccine or medicinal preparation in a method of immunizing a mammal against LH-RH and thereby influencing reproductive or behavioral characteristics of that mammal and in a method of improving the meat quality of pigs.

LH-RH is a small 10 amino acid long peptide (decapeptide) from the hypothalamus. The amino acid sequence (with, as usual, the amino terminal amino acid to the left and the carboxyl terminal amino acid to the right) of LH-RH is according to the formula in which the amino acids are coded with the three-letter code: pGlu-Lys-Trp-Ser-Tyr-Gly-Leu-Arg-Pro-Gly-NH2. (SEQ ID NO:1), or in the one letter code according to the formula: #E H W S Y G L R P G@ (SEQ ID NO:1), #E is pyroglutamic acid and G@ is glycine amide.

LH-RH acts at the hypothysis to cause an increase in release of biologically active FSH (follicle-stimulating hormone) and LH (luteinising hormone) in the blood, which in turn stimulate the development of the testes in the growing male animal and the synthesis of testicular steroids. In the growing female animal the development of the ovaries is stimulated and therein follicle development, synthesis of ovarian steroids and ovulation.

It is known that LH-RH, if coupled to a carrier protein, can be used to vaccinate animals. Such a vaccination can be carried out for different reasons which are as connected with the natural function of the LH-RH. As is known, a drastic reduction of LH and FSH in the blood inhibits the production of testicular steroids or androgens and sperm in the testis of the male and the formation of ovarian steroids or progestagens and estrogens and follicle maturation in the ovary of the female. Such a reduction of the amounts of androgens, progestagens and estrogens in the blood to a level comparable to the level obtainable by removing the testes or ovaries via castration can be achieved by effective immunization of the animal against LH-RH. In male animals in many cases the testes then appear to develop slowly or not at all (no synthesis of androgens (male steroid hormones) and no formation of spermatozoa) and in female animals the activity of the ovaria appears to diminish (synthesis of estrogen and progestagens (female steroid hormones), ripening of follicles and ovulation are inhibited).

In veterinary medicine, 100% effective immunisation against LH-RH could be used for the sterilisation of, e.g., small domestic animals such as male and female cats and dogs, or for the treatment of aggressiveness to male dogs and bulls, simply by vaccination instead of by drastic surgery, such as castration or ovariectomy. Other conceivable reasons for immunisation against LH-RH are to prevent heat in female animals, such as dogs, cats and cows, and restlessness in male animals, being fattened for slaughter. In human health care, immunisation against LH-RH can be used in the treatment of prostate cancer and breast cancer and, if required in the treatment of some forms of pituitary carcinoma.

Another use of a vaccine against LH-RH is in the field of stock breeding, particularly the fattening of pigs for slaughter. The meat of male, sexually mature pigs (boars) has a typical odour, the so-called boar taint or boar odour. In the sexually mature pig in the testes many C19-A16 steroids are formed which are stored in the fat tissue of the animal (Patterson, J. Sci. Food Agric. 19, 31–38 (1968); Brooks et al., J. Anim. Sci. 62, 652–645 (1986); Claus, Z. Zeitschrift Tierzüchtung, Züchtungsbiol. 33, 38–47 (1976); Claus, Z. Endocrinol. (Copenh.) 91, Suppl. 125, 432–433 (1979)). These steroids are mainly responsible for the formation of the disagreeable urine-like odour when the meat is heated (Fuchs, Swedish J. Agric. Res. J. 233–237 (1971); Bonneau, Livest. Prod. Sci. 9, 687–705 (1982)). Owing to this unpleasant odour, meat of male sexually mature pigs is hardly, if at all, suitable for consumption and unfit for export. Because about 10% of the male slaughter pigs are already sexually mature before the slaughter time, this potentially entails a great loss for the pig farming industry.

In order to control and prevent these losses, nearly all male piglets are castrated when they are young, with a surgical procedure that is generally executed without any form of anaesthesia. Apart from the animal unfriendly aspect of such a castration, castration also leads to infections, growth inhibition and a final meat quality inferior to that of an intact animal (at least as long as that intact animal has not yet developed boar taint) (Walstra, Livest. Prod. Sci. 1, 187–96 (1974)).

An animal friendly alternative which in addition benefits meat quality consists in the reduction of the LH-RH concentration at the pituitary of the young animal by means of immunisation against LH-RH. This reduction to LHRH levels leads to a reduction in the concentrations of biologically active FSH and LH, which in turn will inhibit development of the testes in the growing animals and inhibit to synthesis of testicular steroids among which the C19-A16 steroids. Animal unfriendly castration becomes unnecessary; infections and growth inhibition are being prevented while the ultimate meat quality certainly is not less than after castration. In addition this method prevents the occurrence of boar taint in male pigs before slaughter time.

However, a strict requirement for a good vaccine against boar taint is that in all pigs development of the testes is being delayed to such an extent that in no case boar taint will occur before slaughter time, even not when tested in a very large population of pigs. The known vaccine preparation do not meet this requirement.

The existing literature and previous patent applications regarding the anti-fertility properties of vaccines against LH-RH, the results of vaccinations often appear to be variable, for instance, some vaccinated animals hardly, if at all, respond to the vaccination, or large doses in commercially unacceptable adjuvants are seeded for the desired effect (Chaffaux et al., Recueil de Médicine Vétérinaire 161 (2), 133–145 (1985); Canatly et al., C. R. Acad. Sc. Paris t. 303. Série III. No. 16, 673–676 (1986); Falvo et al., J. Anim. Sci 63, 986–994 (1986); Goubau et al., Domest. Anim.
More in particular, the tandem-LHRH vaccine invention provides a peptide which is characterised in that it comprises at least 2 LHRH sequences in tandem according to the general formula (with the amino terminal amino acid (left) and the carboxy terminal amino acid to the right) pGlu-His-Trp'-Ser-Tyr-Gly-Leu-Arg-Pro'-Gly-Glu-His-Trp'-Ser-Tyr-Gly-Leu-Arg-Pro' Gly-cys-NH₂ (SEQ 1D NO.3), in which amino acids are indicated according to the three-letter code, Trp¹ and Trp² are either Trp or N-formyl-Trp, and n is a number having a value of at least 1.

This tandem-LHRH vaccine appeared to block completely testes growth and the development of steroid, producing (Leydig) cells in the testis. However, this vaccine appeared to have a number of disadvantages which made it less practical for use. One disadvantage was the need for a high dose to be applied in vaccination to make immunocapsulation successful, at least 1 mg per pig was needed to elicit the wanted response, which makes it expensive to apply the vaccine on a large scale in the pig industry.

Another important disadvantage of this vaccine preparation is that this vaccine is so highly effective only in a composition with complete Freund’s adjuvant. Use of this harsh adjuvant elicits many unwanted side effects such as difficulties in preparation and application due to its viscose nature, furthermore, the application itself can be very painful for the animal, and finally, remains of the adjuvant and the possible development of chronic inflammatory reactions to theadjuvant such as adjuvant related abscesses in the muscle at the injection site may decrease the meat quality of the injected animal.

The present invention, however, provides solutions to the undesired side effects of the above described vaccination against LHRH with the tandem-LHRH peptide preparation, without losing the beneficial advantages of the effective vaccination with a tandem-LHRH peptide in comparison with other existing vaccines directed against LHRH.

It was shown that the monomeric form of the tandem-LHRH vaccine with complete Freund’s adjuvant but in total absence of the carrier protein KLH was fully effective in blocking testicle growth and boar odor in pigs and that dimeric forms of the tandem-LHRH vaccine applied without using the complete Freund’s adjuvant but with the milder incomplete Freund’s adjuvant instead resulted in fairly high efficacy. Surprisingly, it then also appeared that the tandem principle applied to a variant of the LHRH molecule, namely with a substitution of the sixth amino acid Gly of the decapetide by a dextrorotatory (D)-amino acid, D-Lys, after which the resulting peptide was coupled to a common carrier compound. (here ovalbumine was used), resulted in a vaccine that was very effective in several mild adjuvants, namely Specol and a double oil emulsion, and was furthermore also effective in low doses. Thus, whereas a vaccine using D-amino acid substitutions of Gly at position 6 of the original and single LHRH decapetide with a D-amino acid did decrease the immunogenicity as compared the original LHRH sequence, such substitutions with a D-amino acid applied to a tandem-LHRH vaccine were able to generate even more immunogenic LHRH vaccine preparations. It can further be expected that substitutions of Gly at position 6 of one or each the LHRH decapetides that constitute the tandem unit with other d-amino terminal amino acid may also result in improving the vaccine, in addition, when using for example D-Lys as substituting amino acid, this allows for dimerisation or multimulation of the tandem peptide without losing the possibility to conjugate the peptide composition to a carrier compound. Although the C-terminal cysteines can now be utilised for dimerisation via disulfide bond formation and thus would not be available for conjugation to a carrier compound any longer, the side chains of the amino acid substitutions still can be used for coupling to carrier
compounds. Of course, substitutions with other amino acids that contain suitable side chains (such as D-Glu, but other possible substituting amino acids carrying suitable side chains are also known to the average person) would also allow for additional coupling possibilities to a carrier compound.


The present invention relates to a peptide or a peptide composition consisting of at least two LHRH sequences in tandem wherein the sixth amino acid Gly of the original LHRH decapetide sequence is substituted with a D-amino Acid. The Gly at position 6 may be substituted by a functional amino acid which in addition contains a side chain by which the LHRH tandem unit can be coupled to a carrier compound. This peptide may or may not be C-terminally amidated, depending on the peptide synthesis techniques used. In addition this peptide may be dimerised or multimerised, and then at least one of the tandem-LHRH-peptide sequences in the dimer or multimer will contain a functional amino acid substitution at position 6 of the original LHRH decapetide.

The peptide or peptide composition according to the invention contains a consecutive sequence that can be described according to the following general formula (SEQ ID NO:4):

```
  1  6  16  21
#ENHGY*LRPGQHWSY*LRPGC
```

in which * indicates possible replacement of Gly by a dextrorotatory amino acid which in addition contains a side chain by which the LHRH Modern unit can be coupled to a carrier compound.

A first feature of this invention is that in this peptide or peptide composition a possible replacement amino acid or amino acids can be substituted in which the sixth amino acid per LHRH decapetide within the tandem unit, Gly, (thus in position 6 or/and 16) of the above general formula is for instance replaced by a D-amino acid to generate a peptide that is different enough from the normal LHRH sequence to be recognised by the immune system without losing the proper immunogenicity.

Furthermore, a second feature of this invention is that individual tandem units are dimerised to further enhance its immunogenicity without losing the possibility to couple the peptide or peptide composition to a carrier compound protein. In this peptide or peptide composition such a dimerisation of the tandem units can for example take place via the carboxyl-terminus or via the amino-terminus, two tandem units may for instance be dimerised by means of a disulfide or thioether bridge. To this purpose the Cys at position 21 can be used, or Cys can be synthesised before the glutamic acid at position 1, but other methods to dimerise or multimerise the LHRH-tandem units can also be found in the prior art. In case the dissociation or multimerisation results in the loss of accessible sites where a carrier compound can be conjugated, it is sufficient to restrict the choice of D-amino acids replacing Gly at position 6 or/and 16 to an amino acid with an appropriate side chain. Such a replacing amino acid can for example be D-Lys, D-Glu or another dextrorotatory amino acid containing a side chain that allows coupling to a carrier compound.

More in particular, a concrete example of such a preferred peptide according to the invention, a D-Lys₆-tandem-LHRH dimer according to the following formula:

```
  1  6  16  21
#ENHGY*LRPGQHWSY*LRPGC
```

Another concrete example of such a preferred peptide according to the invention is a D-Glu₆-tandem-LHRH dimer according to the following formula:

```
  1  6  16  21
#ENHGY*LRPGQHWSY*LRPGC
```

But other peptides or peptide in which monomerised, dimerised or multimerised LHRH tandem units that contain D-amino acid substitutions at positions 6 and/or 16 and/or (as in the last two examples regarding dimeric forms) at positions 27 and/or 37 are also part of the invention.

The invention further provided a composition which is characterized in the it comprises a peptide brought into an immunogenic form. As a skilled worker knows, there are different methods of bringing a substance which is in itself not immunogenic, into an immunogenic form. A first possibility is to couple a peptide according to the invention to a suitable carrier protein. In a tandem peptide, a cystiene at the N- or C-terminus an be suitably used for a chemical coupling. In the tandem-dimer peptide, coupling can also be performed using the plain or the modified side chain of D-lysin. D-glutamine, or any other modified amino acid replacing glycine at position 6 and/or 16 and/or 27 and/or 37. Those skilled in the art perfectly know what coupling methods and what carrier proteins are eligible. According to the invention there is preferred a composition which is characterized in that it comprises an immunogenic conjugate of a protein, such as ovalbumine, and a peptide or peptide composition according to the invention. Of course, the vaccine preparation according to the invention can be combined with at least one immunoadjuvant. Suitable immuno-adjuvants are known to those skilled in the art. A preferred adjuvant according to the invention can be Specol or a double oil emulsion, but other adjuvants that elicit no a only mild side-reactions an be used as well. The invention can be used in methods for immunising individuals selected from a wide range of vertebrates, but more in particular mammals, against LHRH. Immunisation against LHRH could for instance be used for the sterilisation of, e.g., small domestic animals such male and female cats and dogs, or for the treatment of aggressiveness in male dogs and bulls. Other conceivable reasons of immunisation against LHRH with the present invention are preventing heat in female animals, such as dogs, cats and cows, and preventing or treating restlessness in male animals, being fattened for slaughter. In human health care, immunisation against LHRH an be used in the treatment of prostate cancer and breast cancer and, if required, in the treatment of some forms of pituitary carcinoma.

A preferred embodiment is a method of improving the meat quality of pigs, wherein the pigs are vaccinated with such a vaccine preparation according the invention. The invention is illustrated in the following experimental part.
SUCCESSFUL VACCINATION OF BOARS IS DEFINED AS TESTIS WEIGHT AT SLAUGHTER OF LESS THAN 150 GRAM. THE AIM IS VISIBLY SMALL TESTES IN ALL ANIMALS WITHIN A TREATMENT GROUP. TESTIS WEIGHT APPEARS TO BE DIRECTLY CORRELATED TO THE PRODUCTION OF ANDROSTERONE AND OR BOAR TANN STEROIDS. WHEN TESTIS WEIGHT IS LESS THAN 60 GRAMS THE TESTES HISTOLOGICALLY WERE COMpletely INACTIVE (Meloen et al., Vaccine 12, 741–746 (1994)) AND NO TESTOSTERONE IN THE SERUM COULD BE DETECTED. WE HAVE DESCRIBED AN EXCELLENT CORRELATION EXISTS BETWEEN SIZE AND PARTICULARLY WEIGHT OF THE TESTES AND THE LEVEL OF ANDROSTEROINE IN BOAR (Onk et al., Livest. Prod. Sci. 42, 63–71 (1995)). IT APPEARS THAT TESTES OF IMMUNIZED ANIMALS WEIGHING LESS THAN 150 GRAM ARE A CLEAR INDICATION FOR THE ABSENCE OF BOAR TAIN. ANDROSTERONE IN BACKFAT WAS USUALLY PRESENT IN UNDETECTABLE LOW CONCENTRATIONS, BUT IF PRESENT WAS ALWAYS LOWER THAN 0.5 MU/g BACKFAT THIS VALUE IN THE LITERATURE IS REFERRED TO AS SAFE LOWER LIMIT FOR THE POSSIBLE PERCEPTION OF BOAR TAIN ALTHOUGH OTHERS REPORT 1 MU/g BACKFAT AS SUFICIENTLY LOW. WE TAKE THE LOWEST VALUE AS LOWER LIMIT, BELOW WHICH WE CONSIDER A PIG AS SUCCESSFULLY IMMUNOCATRACATED. THIS RELATIONSHIP IS BASED ON MEASUREMENTS IN MORE THAN 100 PIGS. TESTICLE WEIGHTS OF CONTROL ANIMALS IN OUR EXPERIMENTS APPEARED TO BE BETWEEN 200 AND 350 GRAMS. FINALLY FEED CONVERSION AND MEAT/FAT RATIO APPEARED TO BE IMPROVED IN IMMUNOCATRACATED COMPARED TO BARROWS (SURGICALLY CATRACATED AT YOUNG AGE).

METHODS

PEPTIDE SYNTHESSES WERE PERFORMED ON A ABI 430A PEPTIDE SYNTHESIZER USING FASTMOC CYCLES ON A 0.25 MMOL SCALE WITH CYCLE TIMES OF ABOUT 60 MIN (Fields C G, Lloyd D H, Macdonald R L, Ottesen K M, Noble R L HBTU ACTIVATION FOR AUTOMATED FMOC SOLID-PHASE PEPTIDE SYNTHESIS PEPTIDE RESEARCH 4, 95–101 (1991); User Bulletin #32. APPLIED BIOSYSTEMS (1990)).

PEPTIDE PURIFICATION: THE PURIFICATIONS WERE CARRIED OUT USING A WATERS PREP.C4000 SYSTEM, EQUIPPED WITH A WATERS PREP/P 25 MMX100 MM FOR DELTA-PACK C18 (15 μM, 100A) MATERIAL AND A GUARD COLUMN.

FOR ANALYTICAL HPLC WE USED TWO WATERS PUMPS MODEL 510, A WATERS GRADIENT CONTROLLER MODEL 680, A WATERS AUTOINJECTOR MODEL WISP 712, AND A WATERS PHOTODIODE ARRAY DETECTOR MODEL 991. THE PRODUCTS WERE ANALYZED IN A LINEAR GRADIENT FROM WATER WITH 0.1% TFA TO 60% ACETONITRIL IN WATER WITH 0.1% TFA IN 60 MIN ON A WATERS DELTA PACK C18-100A (3.9×150 MM, 5μM) COLUMN AT 1 ML/MIN AT 215 Nm. ALL PRODUCTS WERE >95% PURE ACCORDING TO THE PEAK SURFACE.

AMINO ACID ANALYSIS WAS DONE WITH THE WATERS PICO TAG SYSTEM. THE RESULTS WERE IN AGREEMENT WITH THE EXPECTED VALUES ACCORDING TO THE AMINO ACID SEQUENCES.

FORMATION PROCEDURE: THE PRODUCT WAS DETERMINED BY DISSOLVING THE PRODUCT IN WATER. THE pH SHOULD BE ADJUSTED TO 5–6 WITH 1% OR 2% NH4HCO3. THE SOLUTION SHOULD STAY CLEAR. TOO HIGH pH CAN BE CORRECTED WITH 1–10% ACETIC ACID. STIR AT ROOM TEMPERATURE FOR AT LEAST 5 H. THE PRODUCT WAS PURIFIED DIRECTLY USING HPLC.

CONJUGATION OF D-LYS-TANDER-DIMER TO OVALBUMIN: [WEIGHT EQUIVALENTS TO USE: 1 MG OF OVALBUMIN AND 1 MG OF D-LYS-TANDER-DIMER IS COUPLED USING 10 MG OF ECDI IN MILIQ WATER. FIRST BOTH THE PePTIDE AND THE OVALBUMIN ARE DISSOlvED IN MILIQ WATER (A–PEPTIDE SOLUTION; B–OVALBUMIN SOLUTION). A AND B ARE MIXED WELL. NEXT A 10X EXCESS, BASED ON WEIGHT EQUIVALENTS, OF CARBOBIOBRINE (ECDI) IS DISSOLVED IN MILIQ WATER (C=ECDI SOLUTION). SUBSEQUENTLY C IS SLOWLY ADDED TO THE A+B SOLUTION UNDER CONTINUOUS STIRRING. AFTER 6 H THE PRODUCT IS DIALYZED (MW cutoff 10,000) AGAINST WATER.

DETERMINATION OF THE LOADING: THE LOADING IS CALCULATED FROM COMPARATIVE AMINO ACID ANALYSIS OF THE CONJUGATE AND THE SEPARATE PEPTIDE AND CARRIER PROTEIN. ACCORDING TO THE AMINO ACID ANALYSES THE CONJUGATES CONTAIN APPROXIMATELY 0.5 MG PEPTIDE PER MG OVALBUMIN.

VACCINE PREPARATION: VACCINES WERE PREPARED BY MIXING PEPTIDE WITH ADJUVANT (SEE DETAILS AT SPECOL).


CFA A IFA WAX MARRIED 1:1 WITH THE PEPTIDE SOLUTION TO A STABLE EMULSION.


SPECOL (SPECIAL OFF PHASE) IS A PRODUCT SUITABLE FOR PRODUCTION OF WATER-IN-OIL EMULSIONS FOR RESEARCH PURPOSES. COMPOSITION: PER 10 ML: 0.453 g TWEEN 85 (ICI). 0.532 g SPAN 85 (ICI). 9 ML MARCOL 52 (ESSO BELGIUM).


VACCINATION PROTOCOL: FOR EACH VACCINE, AND PER ANIMAL, 1 MG OF PEPTIDE (AS CALCULATED FROM THE LOADING), OR A LOWER AMOUNT AS INDICATED, AS PREVIOUSLY SHOWN, WAS INJECTED INTO THE VAIN OR INGUINAL TROMG. THE VACCINE WAS EMULSIFIED WITH THE INDICATED ADJUVANT. INTESTINAL INJURIES WERE APPROXIMATELY 10 WEEKS AFTER THE FIRST VACCINATION WHEN THEY RECEIVED THE FIRST VACCINATION. THE BOOSTER ADMINISTRATION 8 WEEKS AFTER THE FIRST VACCINATION HAD THE SAME COMPOSITION.

EVALUATION: THE SIZE OF THE PIG’S TESTICLES WAS MEASURED EXTERNALLY USING CALIPERS. FROM APPROXIMATELY 12 WEEKS AFTER THE FIRST VACCINATION THE SIZE OF THE TESTICLES DOES NOT INCREASE FURTHER (OR EVEN DIMinishes) IN PIGS WITH A LOWERED CONCENTRATION OF LHRH. SERUM SAMPLES WERE TAKEN FOR DETECTION OF ANTI-LHRH-TITERS AND TESTOSTERONE. ANTIBODIES TO GNRH WERE DETERMINED BY BINDING OF SERIAL DILUTIONS OF THE PIG ANTI SERA TO 125I-LHRH. ANIMALS WERE ALLOWED 6 WEEKS AFTER THE FIRST VACCINATION. AFTER SLAUGHTER TESTICLES WERE WEIGHED AND A BACKFAT SAMPLE WAS TAKEN FOR DETERMINATION OF THE BOAR TAIL STEROL ANDROSTEROINE USING AN ELISA (RIDASCREEN). FURTHERMORE MEAT QUALITY WAS JUDGED.

EXPERIMENT 1

IN THESE EXPERIMENTS DIFFERENT PEPTIDE CONSTRUCTS HAVE BEEN TESTED IN UNCONJUGATED FORM. EACH PEPTIDE WAS ADMINISTERED TWICE AT THE AGE OF 10 AND 18 WEEKS IN A QUANTITY OF 1 MG USING IFA AS ADJUVANT. TESTED CONSTRUCTS WERE CARBOXY-TERMINAL DIMERISED LHRH-MONOMER, CARBOXY-TERMINAL DIMERISED LHRH-TANDER, AMINO-TERMINAL DIMERISED LHRH-TANDER, CARBOXY-TERMINAL DIMERISED [D-NAF(2)2]-LHRH (NAFARELIN®-MONOMER, CARBOXY-TERMINAL DIMERISED [D-NAF(2)2]-LHRH (NAFARELIN®)-TANDER CARBOXY-TERMINAL DIMERISED [D-LYS]3 PHLHRH-MONOMER AND CARBOXY-TERMINAL DIMERISED [D-LYS]3 PHLHRH-TANDER.
US RE39,048 E

1) Peptide formula of C-monomer-LHRH-dimer:

1  11
pEHWSYGLRPC  (SEQ ID NO:5 where residue 6 is Gly)

pEHWSYGLRPC  (SEQ ID NO:5 where residue 6 is Gly)

2) Peptide formula of C-tandem-LHRH-dimer:

1  21
pEHWSYGLRPQKHWSYGLRPC  (SEQ ID NO:4 where residues 6 and 16 are Gly)

pEHWSYGLRPQKHWSYGLRPC  (SEQ ID NO:4 where residues 6 and 16 are Gly)

3) Peptide formula of N-tandem-LHRH-dimer (SEQ ID NO:6 where residues 7 and 17 are Gly and residue 21 is glycine amide):

1  21
COHWSYGLRPGQHWSYGLRPG8  COHWSYGLRPGQHWSYGLRPG8

4) Peptide formula of [D-Nal(2)]-monomer-LHRH-dimer:

1  6  11
pEHWSY*LRPGC  (SEQ ID NO:5)

pEHWSY*LRPGC  (SEQ ID NO:5)

* = [3-(2-naphthalenyl)-D-alanine]

5) Peptide formula of [D-Nal(2)]-tandem-LHRH-dimer:

1  6  16  21
pEHWSY*LRPGQKHWSY*LRPGC  (SEQ ID NO:4)

pEHWSY*LRPGQKHWSY*LRPGC  (SEQ ID NO:4)

* = [3-(2-naphthalenyl)-D-alanine]

6) Peptide formula of [D-Lys]-(amine)monomer LHRH-dimer:

1  6  11
pEHWSY*LRPGC  (SEQ ID NO:5)

pEHWSY*LRPGC  (SEQ ID NO:5)

* = [D-lysine]

7) Peptide formula of [D-Lys]-(amine)monomer LHRH-dimer:

1  6  16  21
pEHWSY*LRPGQKHWSY*LRPGC  (SEQ ID NO:4)

pEHWSY*LRPGQKHWSY*LRPGC  (SEQ ID NO:4)

* = [D-lysine]

In group 2, 3, and 7 70-80% of the animals appeared to be successfully immunocastrated. In these responders the highest measured testis weight was 86 g., whereas the non-responders had testis weights of 193 g. or higher. From these results it appears that replacement of Complete Freund’s Adjuvant by Incomplete Freund’s Adjuvant in a formula with an unconjugated peptide leads to loss of effectiveness. Furthermore, it particularly appears firstly that the tandem principle is essential for the activity of the vaccine, secondly that dimerisation can take place via the carboxy- or via the amino terminus and thirdly that not every amino acid substitution is allowed. In view of the inefficiency of the Nafarelin®-analog as a vaccine.

EXPERIMENT 2

The already known successful vaccine based on the tandem-LHRH peptide coupled to keyhole limpet hemocyanine (KLH) in CFA far the first and IFA for the second vaccination was tested without conjugation to a carrier protein. In single and dimerised form, in CFA/IFA and in 2xIFA, and in a dose of 1 mg and 100 µg. Pigs were vaccinated 2x at 10 and 18 weeks of age.

1) Peptide formula of Tandem-LHRH (SEQ ID NO:4 where residues 6 and 16 are Gly):

1  21
pEHWSYGLRPQKHWSYGLRPC

2) Peptide formula or Tandem-LHRH-dimer:

1  21
pEHWSYGLRPQKHWSYGLRPC  (SEQ ID NO:4 where residues 6 and 16 are Gly)

pEHWSYGLRPQKHWSYGLRPC  (SEQ ID NO:4 where residues 6 and 16 are Gly)
The Tandem-LHRH vaccine in single or dimerised form appears to be virtually completely effective even in complete absence of the carrier protein KLH in blotting testis growth and boar taint in pigs. However, an important disadvantage of this vaccine is that the very high effectiveness is achieved only in a composition with complete Freund's adjuvant and in a high dose. In addition, reasonable anti-LHRH Ab titers (expressed as % binding of sera taken at 8 weeks postvaccination), were achieved only with tandem-LHRH or tandem-LHRH dimer in CFA/IFA at 1 mg dose.

**EXPERIMENT 3**

The effect of conjugation to a carrier protein was tested in combination with the replacement of IFA by the milder adjuvants Specol and double oil emulsion (d.o.e.). The dimerised tandem-LHRH peptide with a replacement of Glycine by D-Lysine at positions 6, 16, 27 and 37 was conjugated to ovalbumin and emulsified in two different adjuvants. Specol and double oil emulsion. Pigs were vaccinated 2x, 10 and 18 weeks old, with 1 mg peptide-conjugate in adjuvant. The coupling efficiency is 50%, therefore the amount of antigen administered in reality is 500 μg.

The peptide formula of [D-Lys(5)]-tandem-LHRH-dimer:

\[
p_{\text{LHRH}}^{\text{D-Lys}} \cdot p_{\text{LHRH}}^{\text{D-Lys}} \quad (SEQ \ ID \ NO14) \]

\[
p_{\text{LHRH}}^{\text{D-Lys}} \cdot p_{\text{LHRH}}^{\text{D-Lys}} \quad (SEQ \ ID \ NO14) \]

* = [D-LysLyn]

The number of pigs with testis weight <150 g per group size and testis weight (mean ± s.d.) at 8 w.p.v.

<table>
<thead>
<tr>
<th>Antigen</th>
<th>Adjuvant</th>
<th>Dose</th>
<th>Group size</th>
<th>Number of pigs</th>
<th>Testis weight (mean ± s.d.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) Tandem LHRH</td>
<td>CEA/IFA</td>
<td>1 mg</td>
<td>6/6</td>
<td>38 ± 23</td>
<td>100 ± 13</td>
</tr>
<tr>
<td>2) Tandem LHRH</td>
<td>CEA/IFA</td>
<td>1 mg</td>
<td>11/11</td>
<td>39 ± 14</td>
<td>178 ± 11.5</td>
</tr>
<tr>
<td>1) Tandem LHRH</td>
<td>2 x IFA</td>
<td>1 mg</td>
<td>5/13</td>
<td>159 ± 108</td>
<td>10.8 ± 8.4</td>
</tr>
<tr>
<td>2) Tandem LHRH-dimer</td>
<td>2 x IFA</td>
<td>1 mg</td>
<td>6/13</td>
<td>154 ± 95</td>
<td>8.7 ± 6.8</td>
</tr>
<tr>
<td>1) Tandem LHRH</td>
<td>CEA/IFA</td>
<td>100 μg</td>
<td>7/12</td>
<td>121 ± 92</td>
<td>12.6 ± 5.9</td>
</tr>
<tr>
<td>2) Tandem LHRH-dimer</td>
<td>CEA/IFA</td>
<td>100 μg</td>
<td>6/12</td>
<td>132 ± 85</td>
<td>9.8 ± 5.2</td>
</tr>
<tr>
<td>1) Tandem LHRH</td>
<td>2 x IFA</td>
<td>100 μg</td>
<td>5/13</td>
<td>172 ± 92</td>
<td>3.5 ± 4.6</td>
</tr>
<tr>
<td>2) Tandem LHRH-dimer</td>
<td>2 x IFA</td>
<td>100 μg</td>
<td>1/13</td>
<td>220 ± 55</td>
<td>4.6 ± 5.4</td>
</tr>
</tbody>
</table>

Using the [D-Lys(5)]-tandem-LHRH-dimer-ovalbumin conjugate in Specol a marked reduction of vaccine dose needed can be achieved. In 43 pigs vaccinated with 125 μg conjugate (≈62.5 μg peptide) the effectivity was 100%. In the backfat of pigs as testes smaller than 150 g. In no occasion more than 0.5 μg androstenone/g backfat has been measured (Oonk et al., Livest. Prod. Sci. 42, 63–71 (1995)); in most cases the androstenone concentrations even are below the detection level in the ELISA (0.111 μg/g backfat). With doses of 10–100 μg conjugate there is an increased risk for non-responders, and doses lower than 10 μg are not effective. Anti-LHRH antibody titers are comparable only widen an experiment. Results clearly show that high doses of vaccine are able to elicit similar titers as high doses, in contrast to the results obtained using the tandem-LHRH or tandem-LHRH-dimer. Within an experiment, lower average titers and larger standard deviations result from the contribution from pigs, not fully responsive or unresponsive to the vaccine.

**EXPERIMENT 5**

In control groups of intact boars testis weights and androstenone have been determined.

<table>
<thead>
<tr>
<th>Antigen</th>
<th>Number of pigs with testis weight &lt;150 g</th>
<th>Testis weight (mean ± s.d.)</th>
<th>Androstenone (μg/g backfat)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) [D-Lys(5)]-tandem-LHRH-dimer-ovalbumin in Specol</td>
<td>9/9</td>
<td>19 ± 12</td>
<td>0.68 ± 0.42</td>
</tr>
<tr>
<td>2) [D-Lys(5)]-tandem-LHRH-dimer-ovalbumin in d.o.e.</td>
<td>8/8</td>
<td>43 ± 46</td>
<td>2.04 ± 1.55</td>
</tr>
</tbody>
</table>

As reported before in Oonk et al., Livest. Prod. Sci. 42, 63–71 (1995) the androstenone concentration varies strongly in individual intact adult male pigs.
SEQUENCE LISTING

<160> NUMBER OF SEQ ID NOS: 7

<210> SEQ ID NO 1
<211> LENGTH: 10
<212> TYPE: PRT
<213> ORGANISM: Unknown
<220> FEATURE:
<223> OTHER INFORMATION: Luteinizing Hormone Releasing Hormone (LHRH) from the hypothalamus of an undisclosed mammal.
<220> FEATURE:
<221> NAME/KEY: misc_feature
<222> LOCATION: (1) (1)
<223> OTHER INFORMATION: X at position 1 = pyroglutamic acid
<220> FEATURE:
<221> NAME/KEY: misc_feature
<222> LOCATION: (10) (10)
<223> OTHER INFORMATION: X at position 10 = glycine amide

<400> SEQUENCE: 1
Xaa His Trp Ser Tyr Gly Leu Arg Pro Xaa
 1  5  10

<210> SEQ ID NO 2
<211> LENGTH: 20
<212> TYPE: PRT
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Vaccine against LHRH from the hypothalamus of an undisclosed mammal.
<220> FEATURE:
<221> NAME/KEY: misc_feature
<222> LOCATION: (1) (1)
<223> OTHER INFORMATION: X at position 1 = preferably pyroglutamic acid, but can also be glutamine having attached thereto a tail comprising one or more additional amino acids.
<220> FEATURE:
<221> NAME/KEY: misc_feature
<222> LOCATION: (3) (3)
<223> OTHER INFORMATION: X at position 3 = tryptophan or formylated tryptophan
<220> FEATURE:
<221> NAME/KEY: misc_feature
<222> LOCATION: (10) (10)
<223> OTHER INFORMATION: The bond between amino acids 10 and 11 could comprise a direct peptide bond between 10 and 11 or a spacer consisting of one or more amino acids, a shorter or longer hydrocarbon chain, or compound groups or molecules.
<220> FEATURE:
<221> NAME/KEY: misc_feature
<222> LOCATION: (13) (13)
<223> OTHER INFORMATION: X at position 13 = tryptophan or formylated tryptophan
<220> FEATURE:
<221> NAME/KEY: misc_feature
<222> LOCATION: (19) (19)
<223> OTHER INFORMATION: The sequence comprising residues 10-19 may be repeated.
<220> FEATURE:
<221> NAME/KEY: misc_feature
<222> LOCATION: (20) (20)
<223> OTHER INFORMATION: X at position 20 = either nothing or a tail comprising an additional amino acid; preferably Cys, the C terminal cysteine being added in connection with a possible coupling of the peptide to a carrier protein.

<400> SEQUENCE: 2
Xaa His Xaa Ser Tyr Gly Leu Arg Pro Gly Gln His Xaa Ser Tyr Gly
 1  5  10  15

Leu Arg Pro Xaa
 20
pyroglutamic acid tryptophan or N-formyl-Trp

Xaa His Xaa Ser Tyr Gly Leu Arg Pro Gly Gln His Xaa Ser Tyr Gly
1 5 10 15

Leu Arg Pro Gly Cys
20

pyroglutamic acid tryptophan or N-formyl-Trp

Xaa His Xaa Ser Tyr Gly Leu Arg Pro Gly Gln His Xaa Ser Tyr Gly
1 5 10 15

Leu Arg Pro Gly Cys
20
FEATURE: Misc feature

OTHER INFORMATION: X at position 1 = pyroglutamic acid

OTHER INFORMATION: X at position 6 = Gly or a dextrorotatory amino acid containing a side chain that allows coupling to a carrier compound.

FEATURE: Misc feature

OTHER INFORMATION: X at position 21 = glycine amide

OTHER INFORMATION: The initial cysteine of the peptide comprising residues 1-21 is joined to the initial cysteine of an identical peptide (residues 22-42) to form a dimer.

FEATURE: Misc feature

OTHER INFORMATION: X at position 7 = a possible replacement of glycine by a dextrorotatory amino acid which in addition contains a side chain by which the LHRH tandem unit can be coupled to a carrier compound.

FEATURE: Misc feature

OTHER INFORMATION: X at position 17 = a possible replacement of glycine by a dextrorotatory amino acid which in addition contains a side chain by which the LHRH tandem unit can be coupled to a carrier compound.

FEATURE: Misc feature

OTHER INFORMATION: The initial cysteine of the peptide comprising residues 1-22 is joined to the initial cysteine of an identical peptide (residues 1-44) to form a dimer.
We claim:

1. A peptide that comprises consisting of at least two contiguous LHRH decapetide sequences wherein the amino acid glycine at position 6 of at least one of the constituting LHRH decapetides is replaced by a dextrorotatory amino acid with a side chain that can be coupled to a carrier compound wherein said contiguous LHRH decapetide sequences are joined with a terminus linkage.

2. A peptide according to claim 1 characterised in that it comprises an amino acid sequence that comprises the structure (SEQ ID NO:4):

wherein the amino acid * at position 6 or 16 is a dextrorotatory amino acid with a side chain that can be coupled to a carrier compound and the other amino acid * is either glycine or a dextrorotatory amino acid with a side chain that can be coupled to a carrier compound.

3. Peptides according to claim 1 that A peptide that comprises consisting of at least two contiguous LHRH decapetide sequences wherein the amino acid glycine at position 6 of at least one of the constituting LHRH decapetides is replaced by a dextrorotatory amino acid with a side chain that can be coupled to a carrier compound wherein said decapetides are joined with an N-terminus to N-terminus linkage or C-terminus to C-terminus linkage and are dimersed or multimerised.

4. A peptide according to claim 3 and comprising the structure:

wherein the amino acid * at position 6 or 16 or 27 or 37 is D-lysine or D-glutamine or another dextrorotatory amino acid with a side chain that can be coupled to a carrier compound and the other amino acid * is either glycine or D-lysine or D-glutamine or another dextrorotatory amino acid with a side chain that can be coupled to a carrier compound.

5. A peptide according to claim 3 and having the structure (SEQ ID NO:7 where residue 22 is Cys):

wherein the amino acid * at position 6 or 16 or 27 or 37 or 17 or 29 or 39 is D-lysine a D-glutamine or another dextrorotatory amino acid with a side chain that can be coupled to a carrier compound and the other amino acid * is either glycine or D-lysine a D-glutamine or another dextrorotatory amino acid with a side chain that can be coupled to a carrier compound.

6. A composition in which a peptide in accordance with claim 1 is coupled to a carrier compound.

7. A composition in accordance with claim 6 wherein the carrier compound is a protein.

8. A composition in accordance with claim 7 wherein the carrier compound is KLH a ovalbumin.

9. A composition peptide in accordance with claims 1 additionally comprising combined with a mild adjuvant.

10. A composition in accordance with claim 9 wherein the mild adjuvant is an oil phase of a water-in-oil emulsion or a double oil emulsion.

11. A vaccine comprising a composition peptide in accordance to claim 1.

12. A method comprising inoculating an animal with a vaccine according to claim 11.

13. A method comprising inoculating an animal with a vaccine according to claim 11 wherein the amount is less than about 1 mg.

14. A method to effect one or more reproductive or behavioural characteristics of an animal, characterized in that said animal is vaccinated in accordance with claim 12.

15. A method to immunocastrate a pig, characterized in that said pig is vaccinated in accordance with claim 12.