Title: METHOD FOR PREPARING SOLID DOSAGE FORM OF DESMOPRESSIN

Abstract: The present invention relates to a novel method for the preparation of a solid dosage form of perorally effective peptide, preferably desmopressin or a pharmaceutically acceptable salt thereof, comprising providing a said peptide containing granulate suitable for compression to a pharmaceutically acceptable tablet, as well as to solid dosage forms, preferably tablets, obtainable by said method.
METHOD FOR PREPARING SOLID DOSAGE FORM OF DESMOPRESSIN

Field of the Invention

The present invention relates to a novel method for the preparation of a solid dosage form of a perorally effective peptide, preferably desmopressin, or a pharmaceutically acceptable salt thereof, as well as to solid dosage forms, preferably tablets, obtainable by said method.

Background

Desmopressin, also known as dDAVP, is a nonapeptide and the therapeutically active ingredient (as the acetate salt) in the pharmaceutical product Minirin®, which is marketed inter alia as a nasal spray and a tablet formulation. Desmopressin is primarily used in the treatment of primary nocturnal enuresis, i.e. bedwetting, in children, but it is approved also for the treatment of nocturia and diabetes insipidus. The first market introduction of the tablet formulation was in Sweden in 1987. The composition of the marketed tablet form of desmopressin has remained the same to date.

The tablet form of desmopressin was first disclosed as set forth in the patent US 5,047,398. The subsequently issued marketing authorisations relate to a tablet where i.a. the mannitol, talc and cellulose components exemplified in US 5,047,398 are replaced with potato starch. In addition to desmopressin acetate and potato starch, the present tablet components are lactose, polyvinylpyrrolidone (PVP) and magnesium stearate that together form a homogeneous tablet compressed from a granulate. This composition is inter alia disclosed on page 28 in the publication WO 2003/094886 A1.

Since desmopressin is a nonapeptide containing a disulfide bond, its stability must always be considered. Representative publications addressing the problem of the

Desmopressin containing granulate has to date been prepared in a wet granulation process involving a sequence of several sieving and mixing steps performed at ambient temperature and humidity followed by drying (cf. example 1 herein). One of the objectives of that procedure is to keep shearing forces that desmopressin may be subjected to at a minimum level. The main disadvantages of said procedure is that it is rather time-consuming and labor intensive.

The publication WO 97/15297 A1 (examples 6 and 10) discloses a wet granulation method for the preparation of a buccal delivery system for desmopressin.

As a mixture of water and ethanol is used as the granulation liquid in the prior art granulate preparation, the resulting tablet inevitably contains solvent residues, typically 5-6% of water and 0.1% of ethanol (percentage by weight). Complete removal of solvent residues by drying is impractical, as conditions for complete drying of solid dosage forms tend to be either too costly in industrial scale or potentially thermally damaging to the desmopressin. The primary purpose of the added ethanol is to shorten the time of drying (via an azeotrope).

US 2003/0091637 A1 discloses the use of a specific copolymer as the coating agent for a pharmaceutical core. The pharmaceutically active principle may, for example, be a peptide hormone. The coating may be conducted using a fluidised bed apparatus.

The aforementioned WO 2003/094886 A1 discloses the preparation of tablets containing desmopressin using a conventional wet granulation process.

WO 2004/096181 A2 (published 11 November 2004) exemplifies a wet granulation process for desmopressin utilising a water/ethanol 1:3 mixture as granulation liquid. The process is characterised by a purposively selected excipient average particle size.
It is an objective of the present invention to overcome the aforementioned disadvantages.

Summary of the Drawing

Figure 1 illustrates the results of an accelerated stability study of various tablets compressed from a granulate prepared according to the present invention.

Disclosure of the Invention

The present invention relates to a method for the preparation of a solid dosage form of a perorally effective peptide, preferably desmopressin, or a pharmaceutically acceptable salt thereof, comprising granulating said peptide, and at least one excipient, carrier or diluent or mixture thereof in a fluid bed granulation apparatus, wherein the resulting peptide containing granulate is suitable for compression to a pharmaceutically acceptable tablet. More specifically, the processing comprises providing conditions to provide mixing and shearing action. Said granulation typically comprises adjusting fluidising air flow and processing temperature and time.

As used herein, the expression perorally effective peptide relates to a non-conjugated drug substance comprising from three to ten (oligopeptide) or at least eleven (polypeptide) amino acids of any origin linked via peptidic bonds and providing a desired therapeutic effect in a mammal when perorally administered in the absence of a protease inhibitor. Desmopressin, octreotide and leuprolide are examples of perorally effective peptides.

Standard literature (see "Pharmaceutical Dosage Forms; Tablets", vol.1, pages 297-298, Eds. H.A. Lieberman, L. Lachman and J.B. Schwartz, Marcel Dekker, Inc., New York and Basel, 1989) teaches that the conditions involved in fluid bed granulation may be harmful e.g. to enzymes. More specifically, the heat and moisture combined with the circulating air and particles in a fluid bed granulation process generate significant shearing and abrasion forces with the purpose of providing
a granulate having flow properties ideal for tablet compression at industrial scale and speed. Such flow properties are due to the resulting smooth surface structure of the granulate subjected to said shearing and abrasion forces.

It is a surprising observation that a molecule as sensitive as desmopressin can withstand the processing conditions of fluid bed granulation. The most significant advantages of the method of the present invention are the short processing time compared to conventional wet granulation, and the excellent flow properties for compression of the resulting granulate.

Fluid bed granulation per se is a conventional technology, and it is extensively disclosed in various standard literature, such as "Pharmaceutical Dosage Forms; Tablets", vol.3, pages 27-29, Eds. H.A. Lieberman, L. Lachman and J.B. Schwartz, Marcel Dekker, Inc., New York and Basel, 1990) and "Pharmaceutics - The science of dosage form design", pages 625-627; Ed. M.E. Aulton, Churchill Livingstone, Edinburgh, London, Melbourne and New York 1988. The proper selection of the general equipment set up & processing conditions is therefore within the capacity of a person skilled in the art of manufacturing pharmaceutical formulations. Examples of commercial providers of apparatus adapted for fluid bed granulation are Aeromatic-Fielder AG, CH (Strea series) and Glatt GmbH, DE.

In a preferred embodiment of the present method, said desmopressin containing granulate is prepared by a process comprising the steps of:

i) providing a powder comprising, or consisting of, at least one excipient, carrier or diluent, or mixture thereof;

ii) providing a granulation liquid containing a solvent and desmopressin, or a pharmaceutically acceptable salt thereof, and optionally a binder; and
iii) contacting said granulation liquid, preferably by spraying, with said powder within said apparatus, wherein the fluidising air flow and processing temperature and time are simultaneously, and optionally also after said contacting is completed, adjusted to provide said mixing and shearing action.

The processing conditions of fluid bed granulation usually also provide drying while the fluidisation is ongoing, i.e. also during said step iii). Continued processing conditions after said contacting thus provide further drying in addition to the mixing and shearing action. As an example, a spraying operation in fluid bed granulation is typically performed at a constant spraying rate over a time period of from 10 to 60 minutes. Optionally, the spraying is followed by continued processing conditions for 10 to 240 minutes if further drying, mixing and/or shearing action is desired.

Said solvent is preferably water, which is a particularly advantageous aspect of the present invention. It is noteworthy that the use of water as sole solvent nevertheless keeps the time of drying short, whereas explosion risks and organic solvent exposure are reduced while providing a granulate of required quality. In addition, the granulation process is simplified by removing a component.

Fluidising air flow refers to an air flow that is sufficient to accomplish fluidisation of the powder and resulting granulate within the fluid bed granulation apparatus. The required air flow depends upon several parameters, including particle size and density. As a non-limiting example, the air flow may be in the range of from 10 to 2,500 m³/h, preferably from 20 to 1,500 m³/h. Different operating scales will inherently require somewhat different fluidising air flows. Selecting an optimal flow for the operating scale in question is not an impractical burden for a person skilled in the art, as the
machinery per se required in the practising of the present invention is commercially available and thus of a conventional nature.

Said processing temperature is typically in the range of from 25 to 80°C, preferably from 30 to 60°C. Temperature ranges of from 35 to 55°C and from 40 to 50°C are also conceivably.

It is preferred that said processing time is in the range of from 10 to 240 minutes. For practical purposes, the process is typically regarded as complete when the formed granulate, which is also dried during the process, reaches a water content that is essentially equal to that of said powder comprising excipient, carrier or diluent.

In many cases the terms excipient, diluent and carrier can be used interchangeably, and they may even refer to one and the same substance, or to a mixture of similar such substances. The proper use and understanding of these terms is well known to a person skilled in the art.

In the present method it is preferred that said excipient, carrier or diluent is selected from cellulose, starch and lactose. As used herein, the term cellulose includes, taken alone or in mixture, neat cellulose, microcrystalline cellulose, carboxymethyl cellulose, hydroxypropyl methylcellulose as well as other variants thereof that may be employed in pharmaceutical formulations.

As used herein, the term starch includes, taken alone or in mixture, potato starch, wheat starch, corn starch, rice starch and sheared and/or acid-hydrolysed variants of the aforementioned starches as well as other variants of starch that are typical in pharmaceutical formulations. As the use of corn starch unexpectedly provided the most stable tablets (cf. example 3), corn starch is the most preferred starch.

The lactose type used is preferably lactose-α-monohydrate.
As indicated above the present solid dosage form may optionally comprise at least one further additive typically selected from a disintegrating agent, binder, lubricant, flavoring agent, preservative, colorant and any suitable mixture thereof. Examples of additives that may be considered in practising the present invention are found in "Handbook of Pharmaceutical Excipients"; Ed. A.H. Kibbe, 3rd Ed., American Pharmaceutical Association, USA and Pharmaceutical Press UK, 2000.

In a preferred embodiment of the present method, said desmopressin containing granulate is compressed to a tablet, preferably in a process where a lubricant is added to said granulate before compression thereof.

Said lubricant is typically selected from a group consisting of stearic acid, salts or esters of stearic acid, hydrogenated vegetable oils, magnesium oxide, polyethylene glycol, sodium lauryl sulphate and talc, and mixtures thereof. Preferably said lubricant is selected from magnesium stearate, calcium stearate, zinc stearate, glyceryl palmitostearate and sodium stearyl fumarate, and mixtures thereof. Magnesium stearate is most preferred. The content of lubricant is typically from 0.05 to 1.0, preferably from 0.25 to 0.50, percent by weight of each unit of solid dosage form.

The practising of the present method preferably includes a binder, e.g. PVP. Typically an amount of binder of from 1 to 6 percent by weight of each unit of solid dosage form is employed.

In the most preferred embodiment said solid dosage form lacks an enteric coating. By avoiding an enteric coating the preparation of the solid dosage form of the present invention is further simplified.

The solid dosage form as eventually prepared preferably lacks an agent that exerts buffering capacity at a pH of from 2 to 6.
The method of the present invention most preferably provides an amount of desmopressin acetate of from 20 to 600 µg per unit of said solid dosage form.

Said solid dosage form is preferably selected from a group consisting of tablets, granulate powder, lozenge, cachet, and wafer sheet. A tablet is most preferred.

The present pharmaceutical composition in a solid dosage form is typically a perorally available tablet. A tablet may be manufactured by compression of a granulate by procedures well established in the art. Examples of suitable tablet compressing equipment are rotary presses provided by Elizabeth-Hata International, USA, and Courtoy NV, BE. For a comprehensive overview of pharmaceutical tablet manufacturing, see "Tableting" (by N.A. Armstrong) in the aforementioned "Pharmaceutics - The science of dosage form design", pages 647-668.

Accordingly, a further aspect of the present invention relates to a solid dosage form, preferably a tablet, that is obtainable by a method as defined above, both in general and as outlined in the specific embodiments.

The following illustrates the present invention in more detail. It shall not be construed as a limitation of how the invention may be practised.

Experimental

Example 1 (prior art): Preparation of a tablet containing desmopressin acetate via wet granulation

Lactose (900 g, Pharmatose 150M; provided by DMV, NL) and potato starch (550 g, AmylSolVat; provided by Lyckeby Stärkelse AB, SE) are mixed in a planetary mixer for 15 minutes at room temperature and sieved through a 1 mm sieve. A granulation liquid consisting of water (75 ml) and PVP (13.8 g, Kollidon® 25; provided by BASF GmbH, DE) is prepared, to which desmopressin acetate (0.75 g; provided by PolyPeptide Laboratories AB, SE) and ethanol (225 g) are added. The granulation liquid is then gradually added to the lactose/starch mixture during
mixing for 20 minutes, followed by further mixing for 10
minutes at room temperature. After sieving (1.4 mm),
drying for about 20 hours at 40°C and further sieving (1.4
mm), the obtained granulate is admixed with magnesium
stearate (11.3 g, 1.0 mm sieved; provided by Peter Greven
NV, NL) and subsequently compressed to 7500 tablets using
a single punch tablet compression machine (Fette Exacta
1). A typical prepared tablet for commercial use contains
0.1 mg of desmopressin acetate and is white, convex and
oval (6.8 x 9.6 mm) with a thickness of 3-4 mm and a
target weight of 192 mg. It has a smooth surface without
scratches or chipped edges, and shows no tendencies to
lamination (so-called capping).

Example 2: Preparation of a tablet containing
desmopressin acetate via fluid bed granulation
Lactose (476.6 g, Granulac 140; provided by Meggle
AG, DE) and potato starch (294.6 g, M14; provided by KMC,
DK) are fed to a fluid bed granulation apparatus (Strea 1;
provided by Aeromatic Fielder AG, DE) and mixed for 2
minutes in an upwards directed fluidising air flow of 25
m³/h at a set temperature of 45°C. A granulation liquid is
prepared by dissolving PVP (24 g, Povidone; provided by
BASF, DE) and desmopressin acetate (0.80 g; provided by
PolyPeptide Laboratories AB, SE) in water (80 g). The
granulation liquid is then sprayed downwards at a constant
rate during 15 minutes onto the lactose/starch mixture
while the latter is simultaneously subjected to an upwards
directed fluidising air flow of 25 m³/h at a temperature
of 45°C. When all the granulation liquid is added the same
air flow and temperature is maintained for a further 20
minutes. The obtained dry granulate is then sieved (1.0
mm) and mixed with powdered magnesium stearate (4 g, 1.0
mm sieved; provided by Peter Greven NV, NL) for 2 minutes
in a conventional mixer (AR400E; provided by EWREKA GmbH,
DE), and subsequently compressed to 4000 tablets in a
rotary punch (Ø 8 mm) compression machine (Korsch XL 100;
provided by Korsch, DE) with a target weight of 200 mg. Tablets with a hardness of 5 kp (1 kp = 9.81 N) and each containing 0.2 mg of desmopressin acetate were prepared in this manner. The tablets had a smooth surface without scratches or chipped edges, and no capping was observed.

Example 3: Stability of desmopressin acetate tablets compressed by granulate from fluid bed granulation

Tablets were prepared in accordance with example 2, including variants with a varying content of PVP (% w/w of total) and where the potato starch (KMC M14) was replaced with either corn starch (White maize starch, C(star)PharmGel.03302; Cerestar Benelux B.V.) or microcrystalline cellulose (Avicel® PH 102; FMC BioPolymer). The five tablet types compared are depicted in Figure 1 as follows:

- Potato 1% PVP (potato starch)
- Microcryst 1% PVP (Avicel®)
- Corn 1% PVP (corn starch)
- Microcryst 0% PVP (Avicel®)
- Corn 0% PVP (corn starch)

Tablets were stored at 40°C at a relative humidity (RH) of 75% in climate chambers. The content of intact desmopressin (start content 100% at 0 months) was monitored over time, and the results are summarised in Figure 1. 20 tablets were investigated of each tablet type, and the mean value is shown. The content of desmopressin was monitored with conventional liquid chromatography and UV spectroscopy (220 nm). As a whole, corn starch unexpectedly provided the most stable desmopressin tablets stemming from fluid bed granulation.

All references listed are to be regarded as an integral part of the present writ.
1. Method for the preparation of a solid dosage form of a perorally effective peptide, or a pharmaceutically acceptable salt thereof, comprising granulating said peptide and at least one excipient, carrier or diluent or mixture thereof in a fluid bed granulation apparatus, wherein the resulting said peptide containing granulate is suitable for compression to a pharmaceutically acceptable tablet.

2. Method according to claim 1, wherein said perorally effective peptide is desmopressin.

3. Method according to any one of claims 1-2, wherein a granulation liquid containing water as sole solvent is utilised.

4. Method according to any one of claims 1-2, wherein the preparation of said granulate comprises adjusting fluidising air flow and processing temperature and time.

5. Method according to claim 4, wherein said desmopressin containing granulate is prepared by a process comprising the steps of:
   i) providing a powder comprising at least one excipient, carrier or diluent, or mixture thereof;
   ii) providing a granulation liquid containing a solvent and desmopressin, or a pharmaceutically acceptable salt thereof, and optionally a binder; and
   iii) contacting said granulation liquid with said powder within said apparatus, wherein the fluidising air flow and processing temperature and time are simultaneously adjusted to provide mixing and shearing action.
6. Method according to claim 5, wherein said solvent is water.

7. Method according to any one of claims 4-6, wherein said fluidising air flow is in the range of from 10 to 2500 m³/h, preferably from 20 to 1500 m³/h.

8. Method according to any one of claims 4-7, wherein said processing temperature is in the range of from 25 to 80°C, preferably from 30 to 60°C.

9. Method according to any one of claims 4-8, wherein said processing time is in the range of from 10 to 240 minutes.

10. Method according to any one of claims 1 and 5-9, wherein said excipient, carrier or diluent is selected from cellulose, starch and lactose.

11. Method according to claim 10, wherein said starch is corn starch.

12. Method according to any one of claims 1-11, wherein said desmopressin containing granulate is compressed to a tablet, preferably in a process where a lubricant is added to said granulate before compression thereof.

13. Method according to claim 12, wherein said lubricant is magnesium stearate.

14. Method according to any one of claims 5-13, wherein a binder, preferably PVP, is present.

15. Method according to any one of claims 1-14, wherein said solid dosage form lacks enteric coating.
16. Method according to any one of claims 1-15, where said solid dosage form lacks an agent that exerts buffering capacity at a pH of from 2 to 6.

17. Method according to any one of claims 1-16, wherein the solid dosage form contains desmopressin acetate in an amount of from 20 to 600 µg per unit of said solid dosage form.

18. Solid dosage form, preferably a tablet, obtainable by a method as defined in any one of the preceding claims.
Desmopressin tablet accelerated stability study

Fig. 1