The present invention is related to feed granules comprising a feed enzyme and a copper ion donor.
FIELD OF THE INVENTION
The present invention relates to enzyme granules especially for animal feed. The present invention further relates to the manufacturing of said granules.

BACKGROUND OF THE INVENTION
In the art concerning animal feed it is a well known fact that pelleting of feed is a desideratum, as pelleting of feed increases the digestibility of especially the starch fraction of the feed. Furthermore, it is known that pelleting of animal feed reduces dust problems.

In the process of producing feed pellets it is considered necessary to steam treat mash feed in order to kill pathogenic micro-organisms if present and to partly gelatinize starch in order to improve the physical properties of the pellets, whereby a steam treatment of around 70-120°C is appropriate. Active compounds present in the feed pellets such as enzymes are not very stable at high temperature or humidity, and thus, a large surplus of enzymes has to be used, or enzyme free feed components are pelletized and steam treated, where after an enzyme containing slurry or solution is coated onto the steam treated pellets. However, this coating process is cumbersome and is often not compatible with feed mill equipment. An attempt to obtain improved enzyme granules for feed is found in WO 92/12645. WO 92/12645 describes T-granules, which are coated with a fat or a wax. Said T-granules are mixed with feed components steam treated and subsequently pelletized. By this invention it was possible to heat treat the granules comprising enzymes and avoid the cumbersome coating with enzymes after the heat treatment. The use of wax coated T-granules was a significant improvement in this field as it was possible to maintain an acceptable enzyme activity during steam pelleting. Another attempt to improve the pelleting stability of enzyme granules is described in WO 2006034710 where it has been found that coating enzyme granules with a salt coating improves the pelleting stability. However some feed mills are run under very aggressive conditions which are still very harsh for the improved enzyme granules, thus there is still a demand for improved pelleting stability. Furthermore the drawback with coating of the enzyme granules is an additional process step.

The present invention provides very good pelleting stability and makes the step of coating optional.
It is described in WO 99/32595 to use copper sulphate as barrier coating in enzyme granules to stabilize enzyme granules for detergents. It is also known in the art to use inorganic salts of zinc and magnesium to stabilize enzyme granules, see WO 97/05245.

SUMMARY OF THE INVENTION

One object of the present invention is to provide an enzyme granule with good pelleting stability. A second object of the present invention is to provide a simple process for obtaining enzyme granules with good pelleting stability.

It has surprisingly been found that copper ions have a positive effect on enzyme stability during pelleting if formulated together with the enzyme.

The present invention provides thus in a first aspect a granule suitable for use in animal feed compositions comprising a feed enzyme and a Cu ion donor.
The present invention further provides feed compositions comprising the enzyme granule of the invention.

DETAILED DESCRIPTION OF THE INVENTION

Introduction

We have surprisingly found it possible to increase the stability of enzymes comprised in granules during steam pelleting by adding a copper ion donor. We have furthermore identified that the amount of added copper ions has to be adjusted in a way that optimize the pelleting stability but without destroying per se stability of the enzyme, which apparently seems to decrease if too high amounts of Cu is present in the enzyme granule.

Definitions

By the term "Cu ion donor" or "copper ion donor" is meant a compound comprising copper, and which is able of providing copper ions to the composition of the invention.

By the term "solution" is meant a homogeneous mixture of two or more substances.

By the term "suspension" is meant fine particles suspended in a liquid.

By the term "particle size" is meant the mass mean diameter of the granules.

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention pertains. As used in the specification and claims, the singular "a", "an" and "the" include the plural references unless the context clearly dictates otherwise. For example, the term granule may include a plurality of granules.
The granule

When referring to the granule of the present invention it can either be a single granule or several granules.

The granule of the present invention which is particularly well suited for steam pelleting and as part of a steam treated pelletized feed composition, comprises an enzyme comprising region wherein a copper ion donor is present to stabilize the enzyme. In a particular embodiment of the present invention the enzyme and copper ion donor is present in a homogeneous matrix. The matrix comprising the enzyme and copper ion donor may comprise other auxiliary components.

Suitable particle sizes of the granule of the present invention is found to be 20-2000 µm, 50-2000 µm, such as 50-800 µm, or 75-700 µm, more particular 100-1000 µm. The granule of the present invention may in a particular embodiment have a particle size below 700 µm. In another particular embodiment of the present invention the particle size of the finished granule is 100-800 µm. In a more particular embodiment of the present invention the particle size of the finished granule is 300-600 µm. In a most particular embodiment of the present invention the particle size is 450-550 µm. In another particular embodiment of the present invention the particle size of the finished granule is below 400 µm. In another most particular embodiment the particle size of the granules of the present invention is above 250 µm and below 350 µm.

In a particular embodiment of the present invention the particle size of the granule of the present invention is between 210 and 390 µm.

The enzyme comprising matrix

The enzyme comprising matrix or region of the granule may be the core of the granule, a layer surrounding the core, or the granule as such if the structure of the granule is homogenous throughout the granule. There may be layers present between the enzyme layer and the core, or the enzyme layer may be next to the core. The core may be an inert particle.

The enzyme comprising region may be a homogeneous blend comprising enzyme and a copper ion donor, or an inert particle with a matrix layer comprising an enzyme and a copper ion donor applied onto it.

The core particle which either consist of a homogeneous blend comprising the enzyme and the copper ion donor or consist of an inert particle upon which a layer comprising the enzyme and copper ion donor is applied has in a particular embodiment a particle size of 20-800 µm. In a more particular embodiment of the present invention the core particle size is 50-500 µm. In an even more particular embodiment of the present invention the core particle size is 100-300 µm. In a most particular embodiment of the present invention the core particle size is 150-250 µm.

In another particular embodiment of the present invention the core particle size is 400-500 µm.

The core particle comprising the enzyme has in a particular embodiment a particle size of 20-800 µm. In a more particular embodiment of the present invention the core particle size is 50-500 µm. In an even more particular embodiment of the present invention the core particle size
is 100-300 µm. In a most particular embodiment of the present invention the core particle size is 150-250 µm. In another particular embodiment of the present invention the core particle size is 400-500 µm.

5 Inert particle:

Inert particles such as placebo particles, carrier particles, inactive nuclei, inactive particles, non-pareil particles, non active particles or seeds, are particles not comprising enzymes or only minor amount of enzymes upon which a coating mixture comprising the enzyme can be layered. They may be formulated with organic or inorganic materials such as inorganic salts, sugars, sugar alcohols, small organic molecules such as organic acids or salts, starch, flour, treated flour, cellulose, polysaccharides, minerals such as clays or silicates or a combination of two or more of these.

In a particular embodiment of the present invention the particles to be coated are inactive particles. In a more particular embodiment of the present invention the material of the core particles are selected from the group consisting of inorganic salts, flour, sugar alcohols, small organic molecules, starch, cellulose and minerals.

Inert particles can be produced by a variety of granulation techniques including: crystallisation, precipitation, pan-coating, fluid bed coating, fluid bed agglomeration, rotary atomization, extrusion, prilling, spheronization, size reduction methods, drum granulation, and/or high shear granulation.

Enzymes

The enzyme in the context of the present invention may be any enzyme or combination of different enzymes that are suitable to be given to an animal, meaning that it in one way or the other will be good for the animal nutritionally to eat the enzyme. Accordingly, when reference is made to "an enzyme" this will in general be understood to include one enzyme or a combination of enzymes. In a particular embodiment it is not construed as including enzymes which have a therapeutic function in medical/pharmaceutical sense.

The feed enzymes should be feed/food grade, thus meaning that they may not be harmful to the animal and be a feed/food grade meaning that it should comply with recommended purity specifications for food grade enzymes. In a particular embodiment this means that the enzyme complies with recommended purity specifications for food grade enzymes given by the Joint FAO/WHO Expert Committee on Food Additives (JECFA) and the Food Chemical Codex (FCC).

The enzyme shall in a particular embodiment comprise less than 30 coliform bacteria pr gram and comprise a viable count of less than 50000/g.
The granules of the invention include between about 0.0005 to about 20% on a dry weight basis of the enzyme component of the granule. For instance, the weight percent of enzyme in embodiments of the invention comprises at least 0.0005 to about 15%, at least 0.001 to about 15%, at least 0.01 to about 10%, at least 0.1 to about 10%, at least 1.0 to about 10%, at least 1.0 to about 8%, at least 1.0 to about 5%, and at least 2.0 to at least 5% in the granule. Typical doses of 25 to 400 grams of the stable, enzyme granules per ton of feed will deliver about 0.0001 to about 80 grams of active enzyme protein per ton of feed, and the enzyme granules may be dosed as high as 5000 grams per ton of feed.

It is to be understood that enzyme variants (produced, for example, by recombinant techniques) are included within the meaning of the term "enzyme". Examples of such enzyme variants are disclosed, e.g. in EP 251,446 (Genencor), WO 91/00345 (Novo Nordisk), EP 525,610 (Solvay) and WO 94/02618 (Gist-Brocades NV).

Enzymes can be classified on the basis of the handbook Enzyme Nomenclature from NC-IUBMB, 1992), see also the ENZYME site at the internet: http://www.expasy.ch/enzyme - ENZYME is a repository of information relative to the nomenclature of enzymes. It is primarily based on the recommendations of the Nomenclature Committee of the International Union of Biochemistry and Molecular Biology (IUB-MB), Academic Press, Inc., 1992, and it describes each type of characterized enzyme for which an EC (Enzyme Commission) number has been provided (Bairoch A. The ENZYME database, 2000, Nucleic Acids Res 28:304-305). This IUB-MB Enzyme nomenclature is based on their substrate specificity and occasionally on their molecular mechanism; such a classification does not reflect the structural features of these enzymes.


The types of enzymes which may be incorporated in granules of the invention include oxidoreductases (EC 1.-.-.-), transferases (EC 2.-.-.-), hydrolases (EC 3.-.-.-), lyases (EC 4.-.-.-), isomerases (EC 5.-.-.-) and ligases (EC 6.-.-.-).
Preferred oxidoreductases in the context of the invention are peroxidases (EC 1.11.1), laccases (EC 1.10.3.2) and glucose oxidases (EC 1.1.3.4). An Example of a commercially available oxidoreductase (EC 1.*) is Gluzyme™ (enzyme available from Novozymes A/S). Further oxidoreductases are available from other suppliers. Preferred transferases are transferases in any of the following sub-classes:

a) Transferases transferring one-carbon groups (EC 2.1);
b) transferases transferring aldehyde or ketone residues (EC 2.2); acyltransferases (EC 2.3);
c) glycosyltransferases (EC 2.4);
d) transferases transferring alkyl or aryl groups, other that methyl groups (EC 2.5); and

e) transferases transferring nitrogenous groups (EC 2.6).

A most preferred type of transferase in the context of the invention is a transglutaminase (protein-glutamine γ-glutamyltransferase; EC 2.3.2.13).

Further examples of suitable transglutaminases are described in WO 96/06931 (Novo Nordisk A/S).

Preferred hydrolases in the context of the invention are: carboxylic ester hydrolases (EC 3.1.1.-) such as lipases (EC 3.1.1.3); phytases (EC 3.1.3.-), e.g. 3-phytases (EC 3.1.3.8) and 6-phytases (EC 3.1.3.26); glycosidases (EC 3.2, which fall within a group denoted herein as "carbohydrases"), such as α-amylases (EC 3.2.1.1); peptidases (EC 3.4, also known as proteases); and other carbonyl hydrolases. Examples of commercially available phytases include Bio-Feed™ Phytase (Novozymes), Ronozyme™ P (DSM Nutritional Products), Natuphos™ (BASF), Finase™ (AB Enzymes), and the Phyzone™ product series (Danisco). Other preferred phytases include those described in WO 98/28408, WO 00/43503, and WO 03/066847.

In the present context, the term "carbohydrase" is used to denote not only enzymes capable of breaking down carbohydrate chains (e.g. starches or cellulose) of especially five- and six-membered ring structures (i.e. glycosidases, EC 3.2), but also enzymes capable of isomerizing carbohydrates, e.g. six-membered ring structures such as D-glucose to five-membered ring structures such as D-fructose.

Carbohydrases of relevance include the following (EC numbers in parentheses):

α-amylases (EC 3.2.1.1), β-amylases (EC 3.2.1.2), glucan 1,4-α-glucosidases (EC 3.2.1.3), endo-1,4-beta-glucanase (cellulases, EC 3.2.1.4), endo-1,3(4)-β-glucanases (EC 3.2.1.6), endo-1,4-β-xylanases (EC 3.2.1.8), dextranases (EC 3.2.1.11), chitinases (EC 3.2.1.14), polygalacturonases (EC 3.2.1.15), lysozymes (EC 3.2.1.17), β-glucosidases (EC 3.2.1.21), α-galactosidases (EC 3.2.1.22), β-galactosidases (EC 3.2.1.23), amylo-1,6-glucosidases (EC 3.2.1.33), xylan 1,4-β-xilosidases (EC 3.2.1.37), glucan endo-1,3-β-D-glucosidases (EC 3.2.1.39), α-dextrin endo-1,6-α-glucosidases (EC3.2.1.41), sucrose α-glucosidases (EC 3.2.1.48), glucan endo-1,3-α-glucosidases (EC 3.2.1.59), glucan 1,4-β-glucosidases (EC
3.2.1 .74), glucan endo-1,6-β-glucosidases (EC 3.2.1 .75), galactanases (EC 3.2.1 .89), arabinan endo-1,5-α-L-arabinosidases (EC 3.2.1.99), lactases (EC 3.2.1.108), chitosanases (EC 3.2.1.132) and xylose isomerases (EC 5.3.1.5).

In the present context a phytase is an enzyme which catalyzes the hydrolysis of phytate (myo-inositol hexakisphosphate) to (1) myo-inositol and/or (2) mono-, di-, tri-, tetra- and/or penta-phosphates thereof and (3) inorganic phosphate.

Three different types of phytases are known: A so-called 3-phytase (alternative name 1-phytase; a myo-inositol hexaphosphatase 3-phosphohydrolase, EC 3.1.3.8), a so-called 4-phytase (alternative name 6-phytase, name based on 1L-numbering system and not D-numbering, EC 3.1.3.26), and a so-called 5-phytase (EC 3.1.3.72). For the purposes of the present invention, all three types are included in the definition of phytase.

For the purposes of the present invention phytase activity may be, preferably is, determined in the unit of FYT, one FYT being the amount of enzyme that liberates 1 micro-mol inorganic ortho-phosphate per min. under the following conditions: pH 5.5; temperature 37°C; substrate: sodium phytate (C₆H₈O₂₄PNa₁₂) in a concentration of 0.0050 mol/l. Suitable phytase assays are described in Example 1 of WO 00/20569. FTU is for determining phytase activity in feed and premix.

Preferred examples of phytases are microbial phytases, such as fungal or bacterial phytases, e.g. derived from the following:


ii. Thermomyces or Humicola, such as the Thermomyces lanuginosus phytase disclosed in WO 97/35017;

iii. Basidiomycetes, such as Peniophora (WO 98/28408 and WO 98/28409);

iv. Other fungal phytases such as those disclosed in JP 11000164 (Penicillium phytase), or WO98/13480 (Monascus anka phytase);


vi. Escherichia coli (e.g. US 6110719);

vii. Citrobacter, such as Citrobacter freundii (disclosed in WO 2006/038062, WO 2006/038128, or with the sequence of UniProt Q676V7), Citrobacter braakii (disclosed in WO 2004/085638
(Geneseqp ADU50737), and WO 2006/037328), and Citrobacter amalonaticus or Citrobacter
gillenii (disclosed in WO 2006/037327);
viii. Other bacterial phytases such as the phytase from Buttiauxella (disclosed in WO
2006/043178);
ix. Yeast phytases, e.g. from Schwanniomyces occidentalis (e.g. disclosed in US 5830732); as well as
x. a phytase having an amino acid sequence of at least 75% identity to a mature amino acid
sequence of any one of the phytases of (i)-(ix);
xi. a variant of the phytase of (i)-(ix) comprising a substitution, deletion, and/or insertion of one
or more amino acids;
xii. an allelic variant of the phytase of (i)-(ix);
xiii. a fragment of the phytase of (i)-(ix) that retains phytase activity; or
xiv. a synthetic polypeptide designed on the basis of (i)-(ix) and having phytase activity.

Preferred examples of phytase variants are disclosed in, e.g., WO 99/49022, WO 99/48380,
WO 00/43503, EP 0897010, EP 0897985, WO 2003/66847, as well as in the above-mentioned

Examples of commercially available proteases (peptidases) include Kannase™, Everlase™,
Esperase™, Alcalase™, Neutrase™, Durazym™, Savinase™, Ovozyme™, Pyrase™,
Pancreatic Trypsin NOVO (PTN), Bio-Feed™ Pro and Clear-Lens™ Pro (all available from
Novozymes A/S, Bagsvaerd, Denmark). Other preferred proteases include those described in
WO 01/58275 and WO 01/58276.

Other commercially available proteases include Ronozyme™ Pro, Maxatase™, Maxacal™,
Maxapem™, Opticlean™, Propease™, Purafect™ and Purafect Ox™ (available from Genencor
International Inc., Gist-Brocades, BASF, or DSM Nutritional Products).

Examples of commercially available lipases include Lipex™, Lipoprime™, Lipopan™,
Lipolase™, Lipolase™ Ultra, Lipozyme™, Palatase™, Resinase™, Novozym™ 435 and
Lecitase™ (all available from Novozymes A/S).

Other commercially available lipases include Lumafast™ (Pseudomonas mendocina lipase
from Genencor International Inc.); Lipomax™ (Ps. pseudoalcaligenes lipase from Gist-
Brocades/Genencor Int. Inc.; and Bacillus sp. lipase from Solvay enzymes. Further lipases are
available from other suppliers.

Examples of commercially available carbohydrases include Alpha-Gal™, Bio-Feed™ Alpha,
Bio-Feed™ Beta, Bio-Feed™ Plus, Bio-Feed™ Wheat, Bio-Feed™ Z, Novozyme™ 188,
Carezyme™, Celluclast™, Cellusoft™, Celluzyme™, Ceremyl™, Citrozym™, Denimax™,
Dezyme™, Dextrozyme™, Duramyl™, Energex™, Finizym™, Fungamyl™, Gamanase™,
Glucanex™, Lactozym™, Liquezyme™, Maltogenase™, Natalase™, Pentopan™, Pectinex™,
Promozyme™, Pulpzyme™, Novamyl™, Termamyl™, AMG™ (Amyloglucosidase Novo),
Maltogenase™, Sweetzyme™ and Aquazym™ (all available from Novozymes A/S). Further carbohydrates are available from other suppliers, such as the Roxazyme™ and Ronozyme™ product series (DSM Nutritional Products), the Avizyme™, Porzyme™ and Grindazyme™ product series (Danisco, Finnfeeds), and Natugrain™ (BASF), Purastar™ and Purastar™ OxAm (Genencor).

Other commercially available enzymes include Mannaway™, Pectaway™, Stainzyme™ and Ronozyme™.

In a particular embodiment of the present invention the feed enzyme is selected from the group consisting of endoglucanases, endo-1,3(4)-beta-glucanases, proteases, phytases, galactanases, mannanases, dextranases and alpha-galactosidase, and reference is made to WO 2003/062409 which is hereby incorporated by reference.

Particular suitable feed enzymes include: amylases, phosphotases, such as phytases, and/or acid phosphatases; carbohydrates, such as amylolytic enzymes and/or plant cell wall degrading enzymes including cellulases such as β-glucanases and/or hemicellulases such as xylanases or galactanases; proteases or peptidases such as lysozyme; galatosidases, pectinases, esterases, lipases, in particular phospholipases such as the mammalian pancreatic phospholipases A2 and glucose oxidase. In particular the feed enzymes have a neutral and/or acidic pH optimum. In a particular embodiment of the present invention the feed enzyme is selected from the group consisting of amylases, phosphotases, phytases, cellulases, β-glucanases, hemicellulases, proteases, peptidases, galatosidases, pectinases, esterases, lipases and glucose oxidase.

In a particular embodiment of the present invention the enzyme is selected from the group consisting of amylases, proteases, beta-glucanases, phytases, xylanases, phospholipases and glucose oxidases.

The above enzyme lists are examples only and are not meant to be exclusive. Any enzyme may be used in the durable granules of the present invention, including wild type, recombinant and variant enzymes of bacterial, fungal, yeast, plant, insect and animal sources, and acid, neutral or alkaline enzymes.

Copper ion donor

The copper ion donor may be any compound which is capable of donating copper ions. The Copper ion donor is preferably water soluble. The copper ion donor may be organic or inorganic and may be selected but is not limited to the group consisting of copper salts of chloride, bromide, iodide, sulfate, sulfite, bisulfite, thiosulfate, phosphate, monobasic phosphate, dibasic phosphate, hypophosphite, dihydrogen pyrophosphate, tetraborate, borate, carbonate, bicarbonate, metasilicate, citrate, malate, maleate, malonate, methionate, succinate,
lactate, formate, acetate, butyrate, propionate, benzoate, tartrate, ascorbate, gluconate and copper chelates such as amino acids chelates.

In a particular embodiment of the present invention the copper ion donor is selected from the group consisting of Cu-acetate \( \times 1 \) \( H_2O \), CuCO\( _3 \), Cu(OH)\( _2 \) \( \times 1 \) \( H_2O \), CuCl\( _2 \) \( \times 2 \) \( H_2O \), Cu(C\( _5H_{10}NO_2S \)_2) (copper methionate), CuO, CuSO\( _4 \) \( \times 5 \) \( H_2O \), CuSO\( _4 \) \( \times 1 \) \( H_2O \), Cupric chelate of amino acids, Cu(C\( _6H_{13}N_2O_2 \)_2SO\( _4 \)\( \times 2 \), Cu-sulphate-lysine and copper methionine chelate.

In a particular embodiment of the present invention the copper ion donor is selected from the group consisting of copper chloride, copper bromide, copper iodide, copper sulfate, copper sulfite, copper bisulfite, copper thiosulfate, copper phosphate, copper monobasic phosphate, copper dibasic phosphate, copper hypophosphite, copper dihydrogen pyrophosphate, copper tetraborate, copper borate, copper carbonate, copper bicarbonate, copper metasilicate, copper citrate, copper malate, copper maleate, copper malonate, copper methionate, copper succinate, copper lactate, copper formate, copper acetate, copper butyrate, copper propionate, copper benzoate, copper tartrate, copper ascorbate, copper gluconate, copper chelates such as amino acids chelates, Cu-acetate \( \times 1 \) \( H_2O \), CuCO\( _3 \), Cu(OH)\( _2 \) \( \times 1 \) \( H_2O \), CuCl\( _2 \) \( \times 2 \) \( H_2O \), Cu(C\( _5H_{10}NO_2S \)_2) (copper methionate), CuO, CuSO\( _4 \) \( \times 5 \) \( H_2O \), CuSO\( _4 \) \( \times 1 \) \( H_2O \), Cupric chelate of amino acids, Cu(C\( _6H_{13}N_2O_2 \)_2SO\( _4 \)\( \times 2 \), Cu-sulphate-lysine, copper methionine chelate and combinations thereof. Said compounds are selected with the proviso that if any of the mentioned compounds are not healthy or in other ways not good for the animal they should be deselected.

In a more particular embodiment of the present invention the copper salt is copper sulphate pentahydrate.

The copper ion donor may be added in the form of a solution, dispersion, emulsion or in dry state. In one embodiment the copper ion donor is added as a powder. In a particular embodiment of the present invention the copper ion donor is added in form of a solution. The solution is in a particular embodiment an aqueous solution. In one embodiment the copper ion donor is added as a liquid.

We have surprisingly found that the copper ion donor may be added in a certain ratio to the enzyme. If the amount of copper ions is too low the pelleting stability of the enzyme is not significantly improved. If the amount of copper ions is too high we have found the per se stability of the enzyme decreases which is most unfortunate. Thus the amount of copper ions may not be too high compared to the amount of enzyme present.

The below described amounts are given in copper ions per enzyme protein molecule. The amount of copper ions is calculated on the basis of the total amount of copper comprising compound added. Thus the copper amount is a theoretical amount calculated as if all the copper present in the granule is on ion form. This means that practically the copper is not necessarily on ion form in the granule.
In a particular embodiment of the present invention the amount of copper ions is between 1 and 600 Cu\textsuperscript{2+} enzyme protein molecule. In a more particular embodiment the amount of Cu\textsuperscript{2+} is between 1 and 400 Cu\textsuperscript{2+} enzyme protein molecule. In an even more particular embodiment of the present invention the amount of Cu\textsuperscript{2+} is between 1 and 200 Cu\textsuperscript{2+} enzyme protein molecule.

In a most particular embodiment of the present invention the copper sulphate is added in an amount of Cu\textsuperscript{2+} is between 1 and 100 Cu\textsuperscript{2+} enzyme protein molecule.

In a particular embodiment of the present invention the amount of copper ions is between 1 and 600 Cu\textsuperscript{2+} phytase protein molecule. In a more particular embodiment the amount of Cu\textsuperscript{2+} is between 1 and 400 Cu\textsuperscript{2+} phytase protein molecule. In an even more particular embodiment of the present invention the amount of Cu\textsuperscript{2+} is between 1 and 200 Cu\textsuperscript{2+} phytase protein molecule.

In a most particular embodiment of the present invention the copper sulphate is added in an amount of Cu\textsuperscript{2+} is between 1 and 100 Cu\textsuperscript{2+} phytase protein molecule.

In a particular embodiment of the present invention the copper ion donor is selected from the group consisting of copper chloride, copper bromide, copper iodide, copper sulfate, copper sulfite, copper bisulfite, copper thiosulfate, copper phosphate, copper monobasic phosphate, copper dibasic phosphate, copper hypophosphite, copper dihydrogen pyrophosphate, copper tetraborate, copper borate, copper carbonate, copper bicarbonate, copper metasilicate, copper citrate, copper malate, copper maleate, copper malonate, cupr methionate, copper succinate, copper lactate, copper formate, copper acetate, copper butyrate, copper propionate, copper benzoate, copper tartrate, copper ascorbate, copper gluconate, copper chelates such as amino acids chelates, Cu-acetate \( \times \) 1 \( \text{H}_2\text{O} \), CuCO\textsubscript{3} \( \times \) 1 \( \text{H}_2\text{O} \), CuCl\textsubscript{2} \( \times \) 2 \( \text{H}_2\text{O} \), Cu(C\textsubscript{6}H\textsubscript{10}NO\textsubscript{2}S)\textsubscript{2} (cupric methionate), CuO, CuSO\textsubscript{4} \( \times \) 5 \( \text{H}_2\text{O} \), CuSO\textsubscript{4} \( \times \) 1 \( \text{H}_2\text{O} \), Cupric chelate of amino acids, Cu(C\textsubscript{6}H\textsubscript{13}N\textsubscript{2}O\textsubscript{2})\textsubscript{2}SO\textsubscript{4}, Cu-sulphate-lysine, copper methionine chelate and combinations thereof and wherein the amount of copper ions added is between 1 and 400 Cu\textsuperscript{2+} enzyme protein molecule, such as between 1 and 200 Cu\textsuperscript{2+} enzyme protein molecule, even between 1 and 100 Cu\textsuperscript{2+} enzyme protein molecule.

Additional granulation agents
The granule may comprise additional materials such as binders, fillers, fibre materials, stabilizing agents, solubilising agents, suspension agents, viscosity regulating agents, light spheres, plasticizers, salts, lubricants and fragrances.

Binders of the present invention can be synthetic polymers, waxes including fats, fermentation broth, carbohydrates, salts or polypeptides.

Synthetic polymers
By synthetic polymers is meant polymers which backbone has been polymerised synthetically.
Suitable synthetic polymers of the invention includes in particular polyvinyl pyrrolidone (PVP), polyvinyl alcohol (PVA), polyvinyl acetate, polyacrylate, polymethacrylate, polyacrylamide, polysulfonate, polycarboxylate, and copolymers thereof, in particular water soluble polymers or co-polymers.

In a particular embodiment of the present invention the synthetic polymer is a vinyl polymer.

Waxes
A "wax" in the context of the present invention is to be understood as a polymeric material having a melting point between 25 -150 °C, particularly 30 to100°C more particularly 35 to 85°C most particularly 40 to 75°C. The wax is preferably in a solid state at room temperature, 25°C.

The lower limit is preferred to set a reasonable distance between the temperature at which the wax starts to melt to the temperature at which the granules or compositions comprising the granules are usually stored, 20 to 30°C.

For some granules a preferable feature of the wax is that the wax should be water soluble or water dispersible, the wax should disintegrate and/or dissolve providing a quick release and dissolution of the active incorporated in the particles to the aqueous solution. Examples of water soluble waxes are poly ethylene glycols (PEG's). Amongst water insoluble waxes, which are dispersible in an aqueous solution are triglycerides and oils. For some granules it is preferable that the wax is insoluble.

In a particular embodiment of the present invention the wax composition is a hydrophilic composition. In a particular embodiment at least 25 % w/w of the constituents comprised in the wax composition is soluble in water, preferably at least 50% w/w, preferably at least 75% w/w, preferably at least 85% w/w, preferably at least 95% w/w, preferably at least 99% w/w.

In another embodiment the wax composition is hydrophilic and dispersible in an aqueous solution.

In a particular embodiment the wax composition comprises less than 75% w/w hydrophobic constituents, preferably less than 50% w/w, preferably less than 25% w/w, preferably less than 15% w/w, preferably less than 5% w/w, preferably less than 1% w/w.

Suitable waxes are organic compounds or salts of organic compounds having one or more of the above mentioned properties.

The wax composition of the invention may comprise any wax, which is chemically synthesized. It may also equally well comprise waxes isolated from a natural source or a derivative thereof.

Accordingly, the wax composition of the invention may comprise waxes selected from the following non limiting list of waxes.

- Poly ethylene glycols, PEG. Different PEG waxes are commercially available having different molecular sizes, wherein PEG'S with low molecular sizes also have low melting
points. Examples of suitable PEG’s are PEG 1500, PEG 2000, PEG 3000, PEG 4000, PEG 6000, PEG 8000, PEG 9000 etc. e.g. from BASF (Pluriol E series) or from Clariant or from Ineos. Derivatives of Poly ethylene glycols may also be used.

- polypropylene (e.g. polypropylene glycol Pluriol P series from BASF) or polyethylene or mixtures thereof. Derivatives of polypropylenes and polyethylene may also be used.

- Polymers of ethyleneoxide, propyleneoxide or copolymers thereof are useful, such as in block polymers, e.g. Pluronic PE 6800 from BASF. Derivatives of ethoxylated fatty alcohols.

- Waxes isolated from a natural source, such as Carnauba wax (melting point between 80-88 0C), Candelilla wax (melting point between 68-70 0C) and bees wax. Other natural waxes or derivatives thereof are waxes derived from animals or plants, e.g. of marine origin. Hydrogenated plant oil or animal tallow. Examples of such waxes are hydrogenated ox tallow, hydrogenated palm oil, hydrogenated cotton seeds and/or hydrogenated soy bean oil, wherein the term "hydrogenated" as used herein is to be construed as saturation of unsaturated carbohydrate chains, e.g. in triglycerides, wherein carbon-carbon double bonds are converted to carbon-carbon single bonds. Hydrogenated palm oil is commercially available e.g. from Hobum Oefe und Fette GmbH - Germany or Deutche Cargill GmbH - Germany.

- Fatty acid alcohols, such as the linear long chain fatty acid alcohol NAFOL 1822 (C18, 20, 22) from Condea Chemie GMBH - Germany, having a melting point between 55-60°C. Derivatives of fatty acid alcohols.

- Mono-glycerides and/or di-glycerides, such as glyceryl stearate, wherein stearate is a mixture of stearic and palmitic acid, are useful waxes. An example of this is Dimodan PM - from Danisco Ingredients, Denmark.

- Fatty acids, such as hydrogenated linear long chained fatty acids and derivatives of fatty acids.

- Paraffines, i.e. solid hydrocarbons.

- Micro-crystalline wax.

In further embodiments waxes which are useful in the invention can be found in CM. McTaggart et. al., Int. J. Pharm. 19, 139 (1984) or Flanders et. al., Drug Dev. Ind. Pharm. 13, 1001 (1987) both incorporated herein by reference.

In a particular embodiment of the present invention the wax of the present invention is a mixture of two or more different waxes.

In a particular embodiment of the present invention the wax or waxes is selected from the group consisting of PEG, fatty acids, fatty acid alcohols and glycerides.

In another particular embodiment of the present invention the waxes are chosen from synthetic waxes. In a more particular embodiment the waxes of the present invention are PEG. In a most
particular embodiment of the present invention the wax is selected from the group of beef tallow, PEG and palm oil.

Fermentation broth

A fermentation broth in accordance with the invention comprises microbial cells and/or cell debris thereof (biomass).

In a preferred embodiment the fermentation broth comprises at least 10% of the biomass, more preferably at least 50%, even more preferably at least 75% and most preferably at least 90% or at least 95% of the biomass originating from the fermentation. In another preferred embodiment the broth contains 0-31% w/w dry matter, preferably 0-20% w/w, more preferably 0-15% w/w such as 10-15% w/w dry matter, 0% dry matter being excluded from said ranges. The biomass may constitute up to 90% w/w of the dry matter, preferably up to 75% w/w, more preferably up to 50% w/w of the dry matter, while the enzyme may constitute up to 50% w/w of the dry matter, preferably up to 25% w/w, more preferably up to 10% w/w of the dry matter.

Polysaccharides

The polysaccharides of the present invention may be un-modified naturally occurring polysaccharides or modified naturally occurring polysaccharides.

Suitable polysaccharides include cellulose, pectin, dextrin and starch. The starches may be soluble or insoluble in water.

In a particular embodiment of the present invention the polysaccharide is a starch. In a particular embodiment of the present invention the polysaccharide is an insoluble starch.

Naturally occurring starches from a wide variety of plant sources are suitable in the context of the invention (either as starches per se, or as the starting point for modified starches), and relevant starches include starch from: rice, corn, wheat, potato, oat, cassava, sago-palm, yuca, barley, sweet potato, sorghum, yams, rye, millet, buckwheat, arrowroot, taro, tannia, and may for example be in the form of flour.

Cassava starch is among preferred starches in the context of the invention; in this connection it may be mentioned that cassava and cassava starch are known under various synonyms, including tapioca, manioc, mandioca and manihot.

As employed in the context of the present invention, the term "modified starch" denotes a naturally occurring starch, which has undergone some kind of at least partial chemical modification, enzymatic modification, and/or physical or physicochemical modification, and which - in general - exhibits altered properties relative to the "parent" starch.

In a particular embodiment of the present invention the granule comprise a polysaccharide.
Salts

The core may comprise additional salt. The salt may be an inorganic salt, e.g. salts of sulfate, sulfite, phosphate, phosphonate, nitrate, chloride or carbonate or salts of simple organic acids (less than 10 carbon atoms e.g. 6 or less carbon atoms) such as citrate, malonate or acetate.

Examples of cations in these salt are alkali or earth alkali metal ions, although the ammonium ion or metal ions of the first transition series, such as sodium, potassium, magnesium, calcium, zinc or aluminium. Examples of anions include chloride, iodide, sulfate, sulfite, bisulfite, thiosulfate, phosphate, monobasic phosphate, dibasic phosphate, hypophosphite, dihydrogen pyrophosphate, carbonate, bicarbonate, metasilicate, citrate, malate, maleate, malonate, succinate, lactate, formate, acetate, butyrate, propionate, benzoate, tartrate, ascorbate or gluconate. In particular alkali- or earth alkali metal salts of sulfate, sulfite, phosphate, phosphonate, nitrate, chloride or carbonate or salts of simple organic acids such as citrate, malonate or acetate may be used. Specific examples include NaH$_2$PO$_4$, Na$_2$HPO$_4$, Na$_3$PO$_4$, (NH$_4$)$_2$HPO$_4$, K$_2$HPO$_4$, KH$_2$PO$_4$, Na$_2$SO$_4$, K$_2$SO$_4$, KHSO$_4$, ZnSO$_4$, MgSO$_4$, Mg(NO$_3$)$_2$, (NH$_4$)$_2$SO$_4$, sodium borate, magnesium acetate and sodium citrate.

The salt may also be a hydrated salt, i.e. a crystalline salt hydrate with bound water(s) of crystallization, such as described in WO 99/32595. Examples of hydrated salts include magnesium sulfate heptahydrate (MgSO$_4$(7H$_2$O)), zinc sulfate heptahydrate (ZnSO$_4$(7H$_2$O)), sodium phosphate dibasic heptahydrate (Na$_2$HPO$_4$(7H$_2$O)), magnesium nitrate hexahydrate (Mg(NO$_3$)$_2$(6H$_2$O)), sodium borate decahydrate, sodium citrate dihydrate and magnesium acetate tetrahydrate.

In a particular embodiment of the present invention the binder is a polypeptide. The polypeptide may be selected from gelatin, collagen, casein, chitosan, poly aspartic acid and poly glutamic acid. In another particular embodiment the binder is a cellulose derivative such as hydroxypropyl cellulose, methyl cellulose or CMC. A suitable binder is a carbohydrate binder such as dextrin e.g Glucidex 21D or Avedex W80.

Fillers

Suitable fillers are water soluble and/or insoluble inorganic salts such as finely ground alkali sulphate, alkali carbonate and/or alkali chloride, clays such as kaolin (e.g. SPESWHITE™, English China Clay), bentonites, talcs, zeolites, chalk, calcium carbonate and/or silicates. Typical fillers are di-sodium sulphate and calcium-lignosulphonate. Other fillers are silica, gypsum, kaolin, talc, magnesium aluminium silicate and cellulose fibres.
Fibre materials

Pure or impure cellulose in fibrous form such as sawdust, pure fibrous cellulose, cotton, or other forms of pure or impure fibrous cellulose. Also, filter aids based on fibrous cellulose can be used. Several brands of cellulose in fibrous form are on the market, e.g. CEPO™ and ARBOCELL™.

Pertinent examples of fibrous cellulose filter aids are ARBOCELL BFC 200™ and ARBOCELL BC 200™. Also synthetic fibres may be used as described in EP 304331 B1.

Stabilizing agents

Stabilising or protective agents such as conventionally used in the field of granulation. Stabilising or protective agents may fall into several categories: alkaline or neutral materials, reducing agents, antioxidants and/or salts of first transition series metal ions. Each of these may be used in conjunction with other protective agents of the same or different categories. Examples of alkaline protective agents are alkali metal silicates, carbonates or bicarbonates. Examples of reducing protective agents are salts of sulfite, thiosulfite, thiosulfate or MnSO₄ while examples of antioxidants are methionine, butylated hydroxytoluene (BHT) or butylated hydroxyanisol (BHA).

In particular stabilising agents may be salts of thiosulfates, e.g. sodium thiosulfate or methionine. Still other examples of useful stabilizers are gelatine, urea, sorbitol, glycerol, casein, Poly vinyl pyrrolidone (PVP), hydroxypropylmethylcellulose (HPMC), carboxymethyl cellulose (CMC), hydroxyethylcellulose (HEC), powder of skimmed milk and/or edible oils, such as soy oil or canola oil. Particular stabilizing agents in feed granules are a lactic acid source or starch.

In a particular embodiment of the present invention the granule comprise a lactic acid source according to patent application no. EP 1,117,771 which is hereby incorporated as reference. A preferred lactic acid source is corn steep liquor. It is also well known in the art that enzyme substrates such as starch, lipids, proteins etc can act as stabilizers for enzymes.

Solubilising agents

As is known by the person skilled in the art, many agents, through a variety of methods, serve to increase the solubility of formulations, and typical agents known to the art can be found in National Pharmacopeia's.

Light spheres:

Light spheres are small particles with low true density. Typically, they are hollow spherical particles with air or gas inside. Such materials are usually prepared by expanding a solid material. These light spheres may be inorganic of nature or organic of nature. Polysaccharides are preferred, such as starch or derivatives thereof. Biodac® is an example of non-hollow lightweight material made from cellulose (waste from papermaking), available from GranTek Inc. These materials may be included in the granules of the invention either alone or as a mixture of different light materials.
Suspension agents:  
Suspension agents, mediators and/or solvents may be incorporated.

Viscosity regulating agents:  
Viscosity regulating agents may be present.

Plasticizers:  
Plasticizers of the present invention include, for example: polyols such as sugars, sugar alcohols, glycerine, glycerol trimethylol propane, neopentyl glycol, triethanolamine, mono-, di- and triethylene glycol or polyethylene glycols (PEGs) having a molecular weight less than 1000; urea and water.

Lubricants:  
As used in the present context, the term "lubricant" refers to any agent, which reduces surface friction, lubricates the surface of the granule, decreases tendency to build-up of static electricity, and/or reduces friability of the granules. Lubricants can serve as anti-agglomeration agents and wetting agents. Examples of suitable lubricants are lower polyethylene glycols (PEGs) and mineral oils. The lubricant is particularly a mineral oil or a nonionic surfactant, and more particularly the lubricant is not miscible with the other materials.

Coatings:  
The granules of the present invention may comprise one, two or more additional coating layers.

Coatings may be applied to the granule to provide additional characteristics or properties. Thus, for example, an additional coating may achieve one or more of the following effects:

(i) reduction of the dust-formation tendency of a granule;
(ii) protection of the active compound in the granule against hostile compounds in the surroundings.
(iii) dissolution at a desired rate upon introduction of the granule into a liquid medium (such as an acid medium);
(iv) provide a better physical strength of the granule.

The coatings of the present invention generally are applied as one or more layers surrounding the core. Embodiments include one, two, three or four protective coating layers. Suitable coating materials are polymers, carbohydrates, proteins, lipids, fats and oils, fatty acids, inorganic salts, and gums and mixtures thereof.

The coatings include moisture barrier coatings and moisture hydrating coatings. The moisture barrier coatings function by excluding moisture, for instance by forming a shell layer that typically does not absorb moisture and prevents or retards the rate of moisture migration into the granule. Moisture hydrating coatings on the granule absorb or bind moisture as either free
water or water of hydration, thereby acting to impede or retard the extent or rate of transport of external moisture into the granule. The moisture hydrating coatings typically constitute at least about 35% w/w of the granule. The moisture hydrating materials in the coatings thermally insulate the enzymes and will absorb a certain amount of moisture and retain it within the hydrating material without allowing it to pass through into the portion of the granule having the enzyme. For moisture hydrating coatings on stable, granules that do not contain appreciable amounts of water prior to steam treatment, such coatings may constitute about 25% w/w of the granule. Moisture barrier coatings typically comprise hydrophobic materials, such as hydrophobic polymers, for example PVA, HPMC, acid-thinned hydroxypropyl starches and oxidized starch; proteins, for example whey and whey protein concentrates; lipids, for example, lecithin; fats and oils, fatty acids, latex and gums, for example, gum arabic. Certain moisture barrier coatings, such as PVA and gum arabic, are not readily oxidized and find particularly applicability in providing chemical stability when the granules of the invention are stored in unpelleted or untableted mixtures, for instance, in premixes that contain choline chloride.

Moisture hydrating coating materials typically are hydrophilic materials, such as carbohydrates and inorganic salts, including hydrated salts. Examples of moisture hydrating materials are magnesium sulfate, sodium sulfate, maltodextrin, ammonium sulfate, sugars, for example, sucrose, and native cornstarch. Polymers used for the protective coatings are polyvinyl alcohol (PVA), polyethylene glycol, polyvinyl pyrrolidone, polyacrylates, polyethylene oxides (PEO), polyactic acid, polyvinylchloride, polyvinylacetate, polyvinyl pyrrolidones (PVP), cellulose ethers, alginates, gelatin, modified starches and substituted derivatives, hydrolysates and copolymers thereof, such as acid-thinned hydroxypropyl starch, such as, Pure cote™ hydroxypropyl methyl cellulose (HPMC), methyl cellulose (MC), carboxymethyl cellulose (CMC), and ethyl cellulose. Most preferred polymers for the protective coatings are PVA, modified PVA, as described in U.S. Pat. NO. 6,872,696, and modified cellulose, such as methyl cellulose and hydroxylpropylmethyl cellulose, as described in PCT Publication No. WO 99/51210, both of which are incorporated by reference herein. Carbohydrates used for the protective coatings are maltodextrin hydroxymethyl cellulose, modified or native starches made from corn, sorghum, arrowroot, rice, wheat, rye, barley, oat, potato, yam, tapioca, cassava, sago, and sugars including sucrose, corn syrup solids, molasses, glucose, fructose, and lactose. Proteins used for the protective coatings are whey powder, whey protein concentrate, whey protein isolate, caseinates, soy protein concentrate and isolate, zein, albumin and gelatin.

Simple, compound and derived lipids that may be used in the protective coatings are waxes (for example, vegetable, mineral and synthetic, such as candelilla, bees wax, cerumen, carnuba (carnauba), shellac, paraffin, and microcrystalline waxes); lecithin (for example mono-and diglycerides); fatty acids (for example stearic, palmitic, linoleic, oleic, butyric, and arachidonic fatty acids and their salts of sodium, potassium, calcium and zinc); and fats and oils (for
example, hydrogenated or partially hydrogenated fats and oils, such as soy, corn, cottonseed, tallow, canola, and linseed oil. A preferred lipid for the protective coatings is lecithin. Inorganic salts used for the protective coatings include salts of sulfate, citrate, chloride, carbonate, sulfite, phosphate, phosphonate, and bicarbonate salts of sodium, ammonium, potassium, calcium, magnesium and zinc. Preferred salts are magnesium, sodium and ammonium sulfates.

Gums that may be used in the protective coatings include gum arabic, guar gum, agar, gum tragacanth, karya gum, locust bean gum, carageenan, xanthan gum, and alginites. The protective coatings of the present invention may further include plasticizers, lubricants, pigments and powders, such as talc, bentonite, kaolin, corn starch, magnesium silicate, calcium carbonate, and chitosan.

Certain embodiments of the present invention typically have a single layer of a moisture hydrating material that is approximately at least 55% w/w of the granule. Because the capacity of moisture hydrating coatings to take up and sequester water has a limit, relatively high levels of single layer coatings are applied. Alternatively, moisture hydrating material(s) may be applied in two layers. Other embodiments of the present invention have protective coatings utilizing both moisture hydrating materials and moisture barrier materials. In these embodiments, the amount of moisture hydrating material may be lower, at least about 25% w/w of the granule and the moisture barrier material is about 2% to 25% w/w of the granule. Using both moisture hydrating materials and moisture barrier materials combines protective mechanisms and typically reduces cost, particularly of the moisture barrier materials. Moisture barrier materials, particularly film-forming materials may be subject to mechanical damage which, if these materials are used alone as a thin coating, may lead to loss of protection for the enzyme. The combination allows for the use of less of both materials than would be required if the materials were used alone. The combination allows for some damage to the moisture layer in view of the presence of the moisture hydrating material.


In a particular embodiment of the present invention the additional coating is a wax coating, according to US 4,106,991 or EP 0,569,468 which is hereby incorporated by reference. For suitable waxes see the section "Waxes" above. In a particular embodiment of the present invention an additional coating may comprise PVA, PEG and/or palm oil.
Additional Coating materials

The coating may comprise additional coating materials such as binders, fillers, fibre materials, enzyme stabilizing agents, salts, solubilising agents, suspension agents, viscosity regulating agents, light spheres, plasticizers, salts, lubricants and fragrances as mentioned in the section "additional granulation agents" above. Further coating ingredients may be pigments.

Pigments

Suitable pigments include, but are not limited to, finely divided whiteners, such as titanium dioxide or kaolin, coloured pigments, water soluble colorants, as well as combinations of one or more pigments and water soluble colorants.

Optionally, the granules can be coated with a coating mixture. Such mixtures may comprise but are not limited to coating agents, preferably hydrophobic coating agents, such as hydrogenated palm oil and beef tallow, and if desired other additives, such as calcium carbonate or kaolin.

In a particular embodiment of the present invention the granule of the present invention further comprises a wax coating.

In a particular embodiment of the present invention the granule of the present invention further comprise a lactic acid source.

In a particular embodiment of the present invention the granule of the present invention further comprise dry matter of corn steep liquor.

In a particular embodiment of the present invention the granule is coated with a salt coating.

Preparation of the granule

The raw granule comprises an enzyme and a copper ion donor.

Methods for preparing the a raw granule may be found in Handbook of Powder Technology; Particle size enlargement by C. E. Capes; Volume 1; 1980; Elsevier. Preparation methods include known feed and granule formulation technologies, i.e.:

a) Spray dried products, wherein a liquid enzyme-containing solution is atomized in a spray drying tower to form small droplets which during their way down the drying tower dry to form an enzyme-containing particulate material. Very small particles can be produced this way (Michael S. Showell (editor); Powdered detergents; Surfactant Science Series; 1998; vol. 71; page 140-142; Marcel Dekker).

b) Layered products, wherein the active compound is coated as a layer around a pre-formed inert core particle, wherein an enzyme-containing solution is atomized, typically in a fluid bed apparatus wherein the pre-formed core particles are fluidized, and the enzyme-containing solution adheres to the core particles and dries up to leave a layer of dry enzyme on the
surface of the core particle. Particles of a desired size can be obtained this way if a useful core particle of the desired size can be found. This type of product is described in e.g. WO 97/23606 c) Absorbed particles, wherein rather than coating the enzyme as a layer around the core, the active compound is absorbed onto and/or into the surface of the core. Such a process is described in WO 97/391 16.

d) Extrusion or pelletized products, wherein an enzyme-containing paste is pressed to pellets or under pressure is extruded through a small opening and cut into particles which are subsequently dried. Such particles usually have a considerable size because of the material in which the extrusion opening is made (usually a plate with bore holes) sets a limit on the allowable pressure drop over the extrusion opening. Also, very high extrusion pressures when using a small opening increase heat generation in the active compound paste, which is harmful to the enzyme. (Michael S. Showell (editor); *Powdered detergents*; Surfactant Science Series; 1998; vol. 71; page 140-142; Marcel Dekker)

e) Prilled products, wherein an enzyme powder is suspended in molten wax and the suspension is sprayed, e.g. through a rotating disk atomiser, into a cooling chamber where the droplets quickly solidify (Michael S. Showell (editor); *Powdered detergents*; Surfactant Science Series; 1998; vol. 71; page 140-142; Marcel Dekker). The product obtained is one wherein the enzyme is uniformly distributed throughout an inert material instead of being concentrated on its surface. Also US 4,016,040 and US 4,713,245 are documents relating to this technique

f) Mixer granulation products, wherein an enzyme-containing liquid is added to a dry powder composition of conventional granulating components. The liquid and the powder in a suitable proportion are mixed and as the moisture of the liquid is absorbed in the dry powder, the components of the dry powder will start to adhere and agglomerate and particles will build up, forming granulates comprising the enzyme. Such a process is described in US 4,106,991 (NOVO NORDISK) and related documents EP 170360 B1 (NOVO NORDISK), EP 304332 B1 (NOVO NORDISK), EP 304331 (NOVO NORDISK), WO 90/09440 (NOVO NORDISK) and WO 90/09428 (NOVO NORDISK). In a particular product of this process wherein various high-shear mixers can be used as granulators, granulates consisting of enzyme as active compound, fillers and binders etc. are mixed with cellulose fibres to reinforce the particles to give the so-called T-granulate. Reinforced particles, being more robust, release less enzymatic dust.

g) Size reduction, wherein the granules are produced by milling or crushing of larger particles, pellets, tablets, briquettes etc. containing the enzyme. The wanted core particle fraction is obtained by sieving the milled or crushed product. Over and undersized particles can be recycled. Size reduction is described in (Martin Rhodes (editor); *Principles of Powder Technology*; 1990; Chapter 10; John Wiley & Sons).
h) Fluid bed granulation. Fluid bed granulation involves suspending particulates in an air stream and spraying a liquid onto the fluidized particles via nozzles. Particles hit by spray droplets get wetted and become tacky. The tacky particles collide with other particles and adhere to them and form a granule.

i) The raw granules may be subjected to drying, such as in a fluid bed drier. Other known methods for drying granules in the feed or enzyme industry can be used by the skilled person. The drying preferably takes place at a product temperature of from 25 to 90°C. For some enzymes it is important the cores comprising the enzyme contain a low amount of water before coating with the salt. If water sensitive enzymes are coated with a salt before excessive water is removed, it will be trapped within the core and it may affect the activity of the enzyme negatively. After drying, the cores preferably contain 0.1-10 % w/w water.

Coating of the granule


The coating may be prepared by the same methods as mentioned above in the section "Preparation of the granule".

The granules obtained can be subjected to rounding off (e.g. spheronisation), such as in a Marumeriser™, or compaction.

The granules can be dried, such as in a fluid bed drier. Other known methods for drying granules in the feed or enzyme industry can be used by the skilled person. The drying preferably takes place at a product temperature of from 25 to 90°C.

Manufacturing of feed pellets

"Pellets" and "Pelleting" refer to solid rounded, spherical and cylindrical tablets or pellets and the processes for forming such solid shapes, particularly feed pellets and solid, extruded animal feed. Known feed pelleting manufacturing processes generally include mixing together the feed ingredients for about 1 to about 5 minutes, transferring the mixture to a surge bin, conveying the mixture to a steam conditioner, optionally transferring the steam conditioned mixture to an expander, transferring the mixture to the pellet mill or extruder, and finally transferring the pellets into a pellet cooler. Fairfield, D. 1994. Chapter 10, Pelleting Cost Center. In Feed Manufac-
In the manufacturing of feed pellets of the present invention it is preferred to involve steam treatment prior to pelleting, a process called conditioning. In the subsequent pelleting step the feed is forced through a die and the resulting strands are cut into suitable pellets of variable length. During this conditioning step the process temperature may rise to 60-100 °C or even up to 120 °C. The granule of the above mentioned embodiments may retain at least 50%, at least 60%, at least 70%, at least 80%, at least 90%, and at least 95% enzyme activity under steam treatment.

In a particular embodiment of the present invention the granule of the invention retain at least 70% enzyme activity after being exposed to 100 °C.

In the present invention the feed mixture (mash feed) may be prepared by mixing the granules comprising the enzyme and copper ion donor with desired feed components. The mixture is led to a conditioner e.g. a cascade mixer with steam injection. In a particular embodiment the feed mixture comprising granules comprising the enzyme and the copper ion donor and other feed ingredients are steam treated. Optionally the steam treated feed mixture is pelleted.

The steam conditioner treats the mixture for about 20 to about 90 seconds, and up to several minutes, at about 85 °C to about 95 °C. The feed is in the conditioner heated up to a specified temperature, 60-100 °C, e.g. 60 °C, 70 °C, 80 °C, 90 °C or 100 °C by injecting steam, measured at the outlet of the conditioner. The residence time can vary from seconds to minutes and even hours. Such as 5 seconds, 10 seconds, 15 seconds, 30 seconds, 1 minute, 2 minutes, 5 minutes, 10 minutes, 15 minutes, 30 minutes and 1 hour. In a particular embodiment of the present invention the temperature is 100 °C and the residence time is 60 seconds.

In a particular embodiment of the present invention the process temperature during steam treatment is at least 60 °C. In a more particular embodiment of the present invention the process temperature during steam treatment is at least 70 °C. In an even more particular embodiment of the present invention the process temperature during steam treatment is at least 80 °C. In a most particular embodiment of the present invention the process temperature during steam treatment is at least 90 °C.

The amount of steam may vary in accordance with the amount of moisture and the initial temperature of the feed mix. About 4% to about 6% added steam has been reported in pelleting processes, and the amount is selected to produce less than about 18% moisture in the mash prior to pelleting, or up to about 28% moisture in mash intended for extrusion.

An optional expander process occurs for about 4 to about 10 seconds at a temperature range of about 100 °C to about 140 °C. The pellet mill portion of the manufacturing process typically operates for about 3 to about 5 seconds at a 15 temperature of about 85 °C to about 95 °C.
After conditioning the feed may be led to a press e.g. a Simon Heesen press, and pressed to pellets with variable length e.g. 15 mm. After the press the pellets are placed in an air cooler and cooled for a specified time e.g. 15 minutes.

A particular embodiment of the present invention is a method for manufacturing a feed composition comprising the steps of:

i. mixing feed components with granules of the present invention,

ii. steam treating said composition (i), and

iii. pelleting said composition (ii).

In a particular embodiment of the present invention the granule is comprising an enzyme and a copper ion donor, wherein the enzyme is selected from amylases, phosphotases, phytases, cellulases, β-glucanases, hemicellulases, proteases, peptidases, galatosidases, pectinases, esterases, lipases and glucose oxidase, and wherein the amount of copper added to the granule during production of the granule is between 1 and 600 Cu²⁺ enzyme protein molecule and wherein the particle size of the granule is between 50-2000 µm, and wherein the granule has been steam treated at a temperature of at least 60°C.

Animal Feed

The granule of the present invention is suitable for use in animal feed compositions. The granule is mixed with feed substances, such feed composition is called mash feed. The characteristics of the granule allows its use as a component of a composition which is well suited as an animal feed, which is steam treated and optionally subsequently pelletized.

The term animal includes all animals. Examples of animals are non-ruminants, and ruminants, such as cows, sheep and horses. In a particular embodiment, the animal is a non-ruminant animal. Non-ruminant animals include mono-gastric animals, e.g. pigs or swine (including, but not limited to, piglets, growing pigs, and sows); poultry such as turkeys and chicken (including but not limited to broiler chickens, layers); young calves; and fish (including but not limited to salmon).

The feed of the present invention may comprise vegetable proteins. The term vegetable proteins as used herein refers to any compound, composition, preparation or mixture that includes at least one protein derived from or originating from a vegetable, including modified proteins and protein-derivatives. In particular embodiments, the protein content of the vegetable proteins is at least 10, 20, 30, 40, 50, or 60% (w/w).

Vegetable proteins may be derived from vegetable protein sources, such as legumes and cereals, for example materials from plants of the families Fabaceae (Leguminosae), Cruciferaceae, Chenopodiaceae, and Poaceae, such as soy bean meal, lupin meal and rapeseed meal.
In a particular embodiment, the vegetable protein source is material from one or more plants of the family *Fabaceae*, e.g. soybean, lupine, pea, or bean.

In another particular embodiment, the vegetable protein source is material from one or more plants of the family *Chenopodiaceae*, e.g. beet, sugar beet, spinach or quinoa.

Other examples of vegetable protein sources are rapeseed, and cabbage.

Soybean is a preferred vegetable protein source.

Other examples of vegetable protein sources are cereals such as barley, wheat, rye, oat, maize (corn), rice, and sorghum.

Suitable animal feed additives are enzyme inhibitors, fat-soluble vitamins, water soluble vitamins, trace minerals and macro minerals.

Further, optional, feed-additive ingredients are colouring agents, aroma compounds, stabilisers, antimicrobial peptides, and/or at least one other enzyme selected from amongst phytases EC 3.1.3.8 or 3.1.3.26; xylanases EC 3.2.1.8; galactanases EC 3.2.1.89; and/or beta-glucanases EC 3.2.1.4.

Examples of anti microbial peptides (AMP's) are CAP18, Leucocin A, Tritripticin, Protegrin-1, Thanatin, Defensin, Ovispirin such as Novispirin (Robert Lehrer, 2000), and variants, or fragments thereof which retain antimicrobial activity.

Examples of anti fungal polypeptides (AFP's) are the *Aspergillus giganteus*, and *Aspergillus niger* peptides, as well as variants and fragments thereof which retain antifungal activity, as disclosed in WO 94/01459 and PCT/DK02/00289.

Usually fat- and water-soluble vitamins, as well as trace minerals form part of a so-called pre-mix intended for addition to the feed, whereas macro minerals are usually separately added to the feed.

The following are non-exclusive lists of examples of these components:

Examples of fat-soluble vitamins are vitamin A, vitamin D3, vitamin E, and vitamin K, e.g. vitamin K3.

Examples of water-soluble vitamins are vitamin B12, biotin and choline, vitamin B1, vitamin B2, vitamin B6, niacin, folic acid and panthothenate, e.g. Ca-D-panthothenate.

Examples of trace minerals are manganese, zinc, iron, copper, iodine, selenium, and cobalt.

Examples of macro minerals are calcium, phosphorus and sodium.

In still further particular embodiments, the animal feed composition of the invention contains 0-80% maize; and/or 0-80% sorghum; and/or 0-70% wheat; and/or 0-70% Barley; and/or 0-30% oats; and/or 0-40% soybean meal; and/or 0-10% fish meal; and/or 0-20% whey.

The present invention is further described by the following examples which should not be construed as limiting the scope of the invention.
Methods and Materials

Steam test

A laboratory scale steam treatment test has been developed to simulate the residual activities that are found after steam pelleting in large scale.

The laboratory scale steam treatment test comprises a 5 litre Lodige mixer equipped with a jacket. The mixer also has an opening in the bottom, which may be used for a thermocouple or introduction of steam through a pipe. Steam flow for treating the granulates are controlled by letting the steam at 1.5 bar pass a 2.0 mm hole. The mixer is preheated by steam through the mixer jacket to reach a temperature of 100°C to avoid condensate in steam-connections and pipe to be inserted into the mixer, a steam flush is allowed prior to the introduction of the granulate. The tests conditions comprise a series of actions precisely controlled time wise for good reproducibility. At time zero, 1 kg of enzyme granulates is added to the mixer while rotating the mixing tools at full speed. Immediately after the steam-pipe is inserted into the mixer for the granulates to reach 100°C, and after precisely 30 seconds steam is turned off. The mixture is maintained at 100°C for another 30 seconds after which approx 200 gram is poured on to a 75 micron sieving net. To immediately cool the warm granulate a net is positioned on top of the sample net and the sample is cooled during 60 second by use of a lid connected to air ventilation. A reference sample is taken prior and after steam treatment and send to FYT activity measurement for residual activity to be calculated.

Measurements of pelleting stability

Experimental set-up in example 5, 8 and 11:

Approximately 50 g enzyme granulate was pre-mixed with 10 kg feed for 10 minutes in a small horizontal mixer. This premix was mixed with 90 kg feed for 10 minutes in a larger horizontal mixer. From the mixer the feed was led to the conditioner (a cascade mixer with steam injection) at a rate of approximately 300 kg/hour. The conditioner heated up the feed to 100°C (measured at the outlet) by injecting steam. The residence time in the conditioner was 60 seconds. From the conditioner the feed was led to a Simon Heesen press equipped with 3.0x35 mm horizontal die and pressed to pellets with a length of around 15 mm. After the press the pellets were placed in an air cooler and cooled for 15 minutes.

Feed formulation:

- 74.0% Grind corn
- 20.7% Toasted soy grits
- 5.0% soy oil
- 0.3% Solivit Mikro 106 premix of minerals and vitamins
- 12% water content
Phytase activity analysis

Method: Phytase splits phytic acid into phosphate, released phosphate is reacted with vanadu-
dium and molydenium oxides into a colored (yellow) complex. Absorbance is measured at 415 nm.

Unit: 1 FTU = amount of enzyme which at standard conditions (as given below) releases phos-
phate equivalent to 1 µM phosphate per minute.

Buffers:
Extraction buffer: 0.01% Tween 20 (polyoxyethylene sorbitan monolaurate)
Substrate: 5 mM phytic acid, 0.22M acetate (sodium acetate/acetic acid), pH 5.5.

Reagent: 5 mM ammonium vanadate, 20 mM ammonium heptamolybdate tetrahydrate, 40 mM
ammonia, 2.4M nitric acid

Procedure:
Extraction of feed: 50 g feed is extracted in 500 ml extraction buffer for 1 hour. Eventual further
dilution in extraction buffer if the activity is higher than 2.5 FTU/g feed. (Detection level is 0.1
FTU/g feed). The sample is centrifuged (15 minutes at 4000 rpm). 300 µl supernatant is mixed
with 3 ml substrate and reacted for 60 minutes at 37 degree C. 2 ml reagent is added. Samples
are centrifuged (10 minutes at 4000 rpm.). Absorbance at 415 nm is measured. Activity is de-
termined relative to a standard curve prepared with KH$_2$PO$_4$.

Reference is made to WO 2003/66847.

EXAMPLES

Example 1

A powder consisting of:

- 1.5 kg fibrous cellulose, Arbocel BC200
- 0.75 kg carbohydrate binder, Avedex W80
- 11.362 kg finely ground sodium sulphate

was granulated in a Lodige mixer FM 50 with a granulation liquid consisting of:

- 0.75 kg carbohydrate binder, Avedex W80
- 0.3 kg wheat starch
- 1.7 kg phytase concentrate
- 1.275 kg water

The granulation was performed in a manner as described in US No. patent 4,106,991, Exam-
pie 1.
The obtained granulate was dried in a fluid bed to a water content below 1% and sifted to obtain a product with the particle range 250 µm to 850 µm. Finally, the product was coated with 11.5% palm oil and 22% calcium carbonate in a manner as described in US patent No. 4,106,991, Example 22.

Example 2

A powder consisting of:

- 1.5 kg fibrous cellulose, Arbocel BC200
- 0.75 kg carbohydrate binder, Avedex W80
- 11.272 kg finely ground sodium sulphate

was granulated in a Lodige mixer FM 50 with a granulation liquid consisting of:

- 0.75 kg carbohydrate binder, Avedex W80
- 0.3 kg wheat starch
- 0.09 kg CuSO4 x 5 H2O
- 1.7 kg phytase concentrate
- 0.016 kg 26% NaOH
- 1.2 kg water

The granulation was performed in a manner as described in US No. patent 4,106,991, Example 1.

The granulate was dried in a fluid bed to a water content below 1% and sifted to obtain a product with the particle range 250 µm to 850 µm. Finally, the product was coated with 11.5% palm oil and 22% calcium carbonate in a manner as described in US patent No. 4,106,991, Example 22.

Example 3

A powder consisting of:

- 1.5 kg fibrous cellulose, Arbocel BC200
- 0.75 kg carbohydrate binder, Avedex W80
- 11.107 kg finely ground sodium sulphate

Was granulated in a Lodige mixer FM 50 with a granulation liquid consisting of:

- 0.75 kg carbohydrate binder, Avedex W80
- 0.3 kg wheat starch
- 0.255 kg CuSO4 x 5 H2O
- 1.7 kg phytase concentrate
- 0.035 kg 26% NaOH
- 1.25 kg water
The granulation was performed in a manner as described in US No. patent 4,106,991, Example 1.

The granulate was dried in a fluid bed to a water content below 1% and sifted to obtain a product with the particle range 250 µm to 850 µm. Finally, the product was coated with 12.0% palm oil and 22% calcium carbonate in a manner as described in US patent No. 4,106,991, Example 22.

Example 4

A powder consisting of:
- 1.5 kg fibrous cellulose, Arbocel BC200
- 0.75 kg carbohydrate binder, Avedex W80
- 10.912 kg finely ground sodium sulphate

was granulated in a Lodige mixer FM 50 with a granulation liquid consisting of:
- 0.75 kg carbohydrate binder, Avedex W80
- 0.3 kg wheat starch
- 0.450 kg CuSO₄·5H₂O
- 1.7 kg phytase concentrate
- 0.035 kg 26% NaOH
- 1.25 kg water

The granulation was performed in a manner as described in US No. patent 4,106,991, Example 1.

The granulate was dried in a fluid bed to a water content below 1% and sifted to obtain a product with the particle range 250 µm to 850 µm. Finally, the product was coated with 11.5% palm oil and 22% calcium carbonate in a manner as described in US patent No. 4,106,991, Example 22.

Example 5

The samples produced in Example 1 to Example 4 were tested in a pelleting trial at 100°C in the outlet of the conditioner. The phytase content was measured using Analytical method EB-SM 0559.02 version 0.1 (available from Novozymes upon request) in a mash feed prior to pelleting and in the feed pellets after pelleting. The following residual activities of the phytase were found:
The conclusion is that CuSO₄ significantly improves the pelleting stability.

**Example 6**

The per se stability was tested. Residual activity of the phytase in [%]:

<table>
<thead>
<tr>
<th>Batch</th>
<th>Amount of Cu²⁺ enzyme protein molecule</th>
<th>After 4 weeks at 4°C</th>
<th>After 4 weeks at 50°C</th>
<th>After 4 weeks at 40°C/60% RH</th>
</tr>
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<tr>
<td>Example 2</td>
<td>135</td>
<td>91.3</td>
<td>68.2</td>
<td>26.3</td>
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<tr>
<td>Example 3</td>
<td>380</td>
<td>87.7</td>
<td>61.9</td>
<td>17.7</td>
</tr>
<tr>
<td>Example 4</td>
<td>675</td>
<td>91.6</td>
<td>68.8</td>
<td>15.0</td>
</tr>
</tbody>
</table>

It can be concluded that Copper ions have a negative influence on the per se stability. The per se stability decreases with an increasing amount of Cu²⁺ ions.

**Example 7**

A powder consisting of:

- 1.5 kg fibrous cellulose, Arboce BC200
- 0.75 kg carbohydrate binder, Avedex W80
- 11.312 kg finely ground sodium sulphate

was granulated in a Lodige mixer FM 50 with a granulation liquid consisting of:

- 0.75 kg carbohydrate binder, Avedex W80
- 0.3 kg wheat starch
- 1.595 kg phytase concentrate
- 1.450 kg water

The granulation was performed in a manner as described in US No. patent 4,106,991, Example 1.
The obtained granulate was dried in a fluid bed to a water content below 1% and sifted to obtain a product with the particle range 250 µm to 850 µm. Finally, the product was coated with 10% palm oil and 22% calcium carbonate in a manner as described in US patent No. 4,106,991, Example 22.

Example 8
A powder consisting of:
1.5 kg fibrous cellulose, Arbocel BC200
0.75 kg carbohydrate binder, Avedex W80
11.267 kg finely ground sodium sulphate
was granulated in a Lodige mixer FM 50 with a granulation liquid consisting of:
0.75 kg carbohydrate binder, Avedex W80
0.3 kg wheat starch
0.045 kg CuSO₄ x 5 H₂O
1.595 kg phytase concentrate
1.45 kg water

The granulation was performed in a manner as described in US No. patent 4,106,991, Example 1.

The granulate was dried in a fluid bed to a water content below 1% and sifted to obtain a product with the particle range 250 µm to 850 µm. Finally, the product was coated with 10.1% palm oil and 22% calcium carbonate in a manner as described in US patent No. 4,106,991, Example 22.

Example 9
A powder consisting of:
1.5 kg fibrous cellulose, Arbocel BC200
0.75 kg carbohydrate binder, Avedex W80
11.222 kg finely ground sodium sulphate
was granulated in a Lodige mixer FM 50 with a granulation liquid consisting of:
0.75 kg carbohydrate binder, Avedex W80
0.3 kg wheat starch
0.090 kg CuSO₄ x 5 H₂O
1.595 kg phytase concentrate
1.4 kg water

The granulation was performed in a manner as described in US No. patent 4,106,991, Example 1.
The granulate was dried in a fluid bed to a water content below 1% and sifted to obtain a product with the particle range 250 µm to 850 µm. Finally, the product was coated with 10.2% palm oil and 22% calcium carbonate in a manner as described in US patent No. 4,106,991, Example 22.

**Example 10**

The samples produced in Example 7 to Example 9 were tested in a pelleting trial at 100°C in the outlet of the conditioner. The phytase content was measured using analytical method EB-SM 0559.02 version 01 (available from Novozymes upon request) prior to pelleting and in the feed pellets after pelleting. The following residual activities of the phytase were found:

<table>
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<tr>
<th>Batch</th>
<th>Residual activity of the Phytase in [%]</th>
</tr>
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<tbody>
<tr>
<td>Example 7</td>
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</tr>
<tr>
<td>Example 8</td>
<td>76.2</td>
</tr>
<tr>
<td>Example 9</td>
<td>81.4</td>
</tr>
</tbody>
</table>

The conclusion is that CuSO₄ significantly improves the pelleting stability.
1. A granule suitable for feed comprising an enzyme and a copper ion donor, wherein the granule has a particle size of 50-2000 µm.

2. The granule of claim 1, wherein the granule has been exposed to a temperature of at least 50°C.

3. The granule of claim 1 wherein the granule has been exposed to a temperature of at least 80°C.

4. The granule of claims 1 to 3, wherein the enzyme and the copper ion donor are present as a mixture in the granule.

5. The granule of claims 1 to 4, wherein the granule comprises a core and a layer surrounding the core.

6. The granule of claim 5, wherein the enzyme and the copper ion donor is present in the core of the granule.

7. The granule of claim 5, wherein the enzyme and the copper ion donor is present in the layer surrounding the core.

8. The granule of claim 7, wherein the core of claim 6 is an inactive particle.

9. The granule according to any of the claims 1 to 8, wherein the amount of copper added is between 1 and 600 Cu\(^{2+}\)enzyme protein molecule.

10. The granule of claims 1 to 8, wherein the amount of copper added is between 1 and 400 Cu\(^{2+}\)enzyme protein molecule.

11. The granule according to any of the preceding claims, wherein the copper ion donor is selected from the group consisting of copper salts of chloride, bromide, iodide, sulfate, sulfite, bisulfite, thiosulfate, phosphate, monobasic phosphate, dibasic phosphate, hypophosphite, dihydrogen pyrophosphate, tetraborate, borate, carbonate, bicarbonate, metasilicate, citrate, malate, maleate, malonate, methionate, succinate, lactate, formate, acetate, butyrate, propionate, benzoate, tartrate, ascorbate, gluconate and combinations thereof.
12. The granule according to any of the claims 1 to 10, wherein the copper ion donor is selected from the group consisting of copper sulfate, copper acetate, copper citrate and copper methionate.

13. The granule according to any of the preceding claims, wherein the enzyme is selected from the group consisting of amylases, phosphotases, phytases, cellulases, β-glucanases, hemicellulases, proteases, peptidases, galatosidases, pectinases, esterases, lipases, glucose oxidase and mixtures thereof.

14. The granule according to any of the claims 1 to 12, wherein the enzyme is selected from the group consisting of amylases, proteases, beta-glucanases, phytases, xylanases, phospholipases, glucose oxidases and mixtures thereof.

15. The granule according to any of the claims 1 to 14, wherein the enzyme is food grade.

16. The granule according to any of the claims 1 to 15, wherein the copper ion donor is food grade.

17. The granule according to any of the preceding claims, wherein the granules have a particle size below 700 µm.

18. The granule according to any of the preceding claims, wherein the granules have a particle size between 210 and 390 µm.

19. The granule according to any of the preceding claims, wherein the active compound is thermo labile.

20. The granule according to any of the preceding claims, wherein the granule comprises a salt coating.

21. The granule according to any of the preceding claims, wherein the granule comprises a polymer coating.

22. The granule according to any of the preceding claims, wherein the granule comprises a wax coating.

23. The granule according to any of the preceding claims, wherein the granule comprises a lactic acid source.
24. A process of preparing the feed granules of claims 1-23, comprising the steps of:
   
   a) preparing a core comprising a copper ion donor and a feed enzyme
   
   b) coating the core with a coating material.

25. The process according to claim 24, wherein the granule is prepared in a mixer, a fluid bed, a fluidized spray dryer, a spray fluidizer, a spray dryer or an extruder.

26. The process according to claims 24 to 25, wherein the copper ion donor is added to the process as a liquid.

27. The process according to claims 24 to 25, wherein the copper ion donor is added as a powder.

28. The process according to claims 24 to 25, wherein the core comprises an inert particle.

29. A feed composition comprising feed components and the granule of any of claims 1 to 23.

30. The feed composition of claim 29, wherein the feed components are selected from the group consisting of vegetable protein, fat-soluble vitamins, water soluble vitamins, trace minerals and macro minerals.

31. A pelletized feed composition comprising any of the granules of claim 1 to 23.

32. A steam treated pelletized feed composition comprising any of the granules of claim 1 to 23.

33. A method for feeding animals comprising administering the feed composition of claim 29-32 to an animal.

34. A method for manufacturing a feed composition comprising the steps of:
   
   i. mixing feed components with any of the granules of claim 1 to 23,
   
   ii. steam treating said composition (i), and
   
   iii. pelleting said composition (ii).
35. The use of any of the granules of claim 1 to 23 for steam treated pelletized feed compositions.
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

INV. A23K1/165 A23K1/00 A23K1/175

According to International Patent Classification (IPC) or to both national classification and IPC.

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

A23K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched.

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal , WPI Data, FSTA, BIOSIS, CHEMABS Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No.


Further documents are listed in the continuation of Box C.

See patent family annex.

* Special categories of cited documents:

"A" document defining the general state of the art which is not considered to be of particular relevance

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"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)

"O" document referring to an oral disclosure, use, exhibition or other means

"P" document published prior to the international filing date but later than the priority date claimed

"R" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"Z" document member of the same patent family

Date of the actual completion of the international search 2 November 2007

Date of mailing of the International search report 12/11/2007

Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentilaan 2 NL-2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl Fax. (+31-70) 340-3016

Authorized officer

Rooney, Kevin
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<td>A</td>
<td>WO 2004/067739 A (NOVOZYMES A/S; SIMONSEN, OLE; MARKUSSEN, ERIK, KJÆR; NIELSEN, HANNE,) 12 August 2004 (2004-08-12) the whole document</td>
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