Title: USE OF POLYPEPTIDES HAVING PROTEASE ACTIVITY IN ANIMAL FEED AND DETERGENTS

Abstract: The present invention relates to the use of isolated polypeptides having protease activity in animal feed and detergents. It also relates to the use of isolated nucleic acid sequences encoding the proteases in the recombinant production of isolated polypeptides having protease activity and isolated nucleic acid sequences encoding the proteases. The invention also relates to nucleic acid constructs, vectors, and host cells, including plant and animal cells, comprising the nucleic acid sequences, as well as methods for producing and using the proteases, particularly using the proteases in animal feed and detergents.
USE OF POLYPEPTIDES HAVING PROTEASE ACTIVITY
IN ANIMAL FEED AND DETERGENTS

REFERENCE TO A SEQUENCE LISTING

This application contains a Sequence Listing in computer readable form, which is
incorporated herein by reference.

BACKGROUND OF THE INVENTION

Field of the Invention

The present invention relates to the use of isolated polypeptides having protease activity
in animal feed and detergents. It also relates to the use of isolated nucleic acid sequences
encoding the proteases in the recombinant production of isolated polypeptides having protease
activity and isolated nucleic acid sequences encoding the proteases. The invention also relates
to nucleic acid constructs, vectors, and host cells, including plant and animal cells, comprising
the nucleic acid sequences, as well as methods for producing and using the proteases,
particularly using the proteases in animal feed and detergents.

Description of the Related Art

Proteases of the S1 group isolated from Saccharomonospora are known in the art. Pati
et al. have disclosed a serine protease from Saccharomonospora viridis in "Complete genome
sequence of Saccharomonospora viridis type strain (P101)", 2009, Stand. Genomic Sci. 1:141-
149, which has been submitted to the EMBL/GenBank under accession number CP001683
(SEQ ID NO: 1 herein). The amino acid sequence is registered with Uniprot number C7MV18
(SEQ ID NO: 2 herein) and the mature amino acid sequence is disclosed in SEQ ID NO: 3. The
strain was isolated from peat bog in Ireland.

Lucas et al have submitted a protease from Saccharomonospora cyanea NA-134
(Uniprot: H5XEH4, SEQ ID NO: 7) having 91.3% sequence identity to SEQ ID NO: 3. Csepregi
et al. have submitted a trypsin protease proenzyme from Saccharomonospora azurea SZMC
14600 (Uniprot: H0K7C9, SEQ ID NO: 8) having 89.4% identity to SEQ ID NO: 3. Lucas et al.
have submitted an endopeptidase from Saccharomonospora glauca K62 to the
EMBL/GenBank/DDBJ databases (Uniprot: H1JPF3, SEQ ID NO: 9) having 86.9% sequence
identity to SEQ ID NO: 3.

Lucas et al. have also submitted two endopeptidases from Saccharomonospora
paurometabolica to the EMBL/GenBank/DDBJ databases (Uniprot: G4J6Q2 and G4XC2, SEQ
ID NO: 10 and 11 respectively) having 81.8% and 80.0% sequence identity to SEQ ID NO: 3.
respectively. Oliynyk et al. have disclosed a serine protease from *Saccharopolyspora erythraea*
having 81.0% sequence identity with SEQ ID NO: 3 (Uniprot: A4FNQ0, SEQ ID NO: 12) in
"Complete genome sequence of the erythromycin-producing bacterium *Saccharopolyspora

Lucas et al. have submitted an alpha-lytic protease from *Saccharomonospora xinjiangensis* XJ-54 to the EMBL/GenBank/DDBJ databases (Uniprot: I0V8H8, SEQ ID NO: 13) having 91.3% sequence identity to SEQ ID NO: 3 and a membrane protein from *Saccharomonospora azurea* NA-128 to the EMBL/GenBank/DDBJ databases (Uniprot: H8GAL4, SEQ ID NO: 14) having 89.4% sequence identity to SEQ ID NO: 3. Other known proteases have sequence identities that are lower than 80%.

WO 05/052146 and WO 05/052161 describes a serine protease used for animal feed
having an identity to the protease of SEQ ID NO: 3 of 71.3%. US 2008/0004186 describes the
use of a protease, cellulase etc having 70.0% identity to the protease of SEQ ID NO: 3 as a
protease from *Streptomyces* 1AG3 for animal feed and for dishwashing having 69.4% identity to
SEQ ID NO: 3. The use of a serine protease having an identity of 69.4% to SEQ ID NO: 3 for
cleaning is disclosed in WO 08/048392. WO 08/153925 and WO 2008/153934 describes using
a protease having 69.4% identity to SEQ ID NO: 3 as a detergent.

WO 95/28850 discloses the combination of a phytase and one or more microbial
proteolytic enzymes to improve the solubility of vegetable proteins. WO 01/58275 discloses the
use of acid stable proteases of the subtilisin family in animal feed. WO 01/58276 discloses the
use of acid-stable proteases derived from *Nocardiopsis* sp. NRRL 18262 (a 10R protease), as
well as a protease derived from *Nocardiopsis alba* DSM 14010 in animal feed. WO 04/072221 ,
WO 04/11220, WO 04/11223, WO 05/035747, and WO 05/12391 disclose proteases related
to the 10R protease and their use in animal feed. WO 04/072279 discloses the use of other
proteases in animal feed. WO 04/034776 discloses the use of a subtilisin/keratinase, PWD-1
from *B. Licheniformis*, in the feed of poultry. WO 04/077960 discloses a method for increasing
the digestibility of forage or grain in ruminants by applying a bacterial or fungal protease.

Commercial products comprising a protease and marketed for use in animal feed include

**RONOZYME® ProAct** (DSM NP/Novozymes), **Axtra®** (Danisco), **Avizyme®** (Danisco),
**Porzyme®** (Danisco), **Allzyme™** (Alltech), **Versazyme®** (BioResources, Int.), **Poultrygrow™**
(Jefo) and **Cibenza® DP100** (Novus).
SUMMARY OF THE INVENTION

Background of the Invention

In the use of proteases in animal feed (in vivo), and/or the use of such proteases for treating vegetable proteins (in vitro) it is noted that proteins are essential nutritional factors for animals and humans. Most livestock and many human beings get the necessary proteins from vegetable protein sources. Important vegetable protein sources are e.g. oilseed crops, legumes and cereals.

When e.g. soybean meal is included in the feed of mono-gastric animals such as pigs and poultry, a significant proportion of the soybean meal is not digested efficiently (the apparent ileal protein digestibility in piglets, growing pigs and poultry such as broilers, laying hens and roosters is only around 80%).

The gastrointestinal tract of animals consists of a series of segments each representing different pH environments. In mono-gastric animals such as pigs and poultry and many types of fish, the stomach is strongly acidic with a pH potentially as low as 1-2, while the intestine has a more neutral pH of around 6-7.5. Apart from the stomach and intestine, poultry also have a crop preceding the stomach. The pH in the crop is mostly determined by the feed ingested and hence typically lies in the range of pH 4-6. Protein digestion by a protease may occur along the entire digestive tract, provided that the protease is active and survives the conditions in the digestive tract. Hence, proteases which are highly acid stable and so can survive in the gastric environment and at the same time are efficiently active at the broad range of physiological pH of the digestive tract in the target animal are especially desirable.

Since animal feed is often formulated in pelleted form, in which steam is applied in the pelleting process, it is also desirable that proteases used in animal feed are capable of remaining active after exposure to said steam treatment.

Proteases have also for many years been used in detergent compositions for hydrolysing proteinaceous materials on textiles, hard surfaces and other surfaces, such as the skin, etc. Such detergent compositions can be used for the cleaning of textiles, in hand washing, in automatic washing machines using powders, tablets or soap bars, and in dish washing by hand or machine using powders, liquids or tablets. The novel S1 proteases of the invention are also useful for these purposes.

In order to produce a protease for industrial use, it is important that the protease is produced in high yields making the product available in sufficient quantities in order to be able to provide the protease at a favourable price.

The present invention relates to the use of an isolated polypeptide having protease activity, selected from the group consisting of:
(a) a polypeptide having at least 80% sequence identity to the polypeptide of SEQ ID NO: 3; 
(b) a polypeptide encoded by a polynucleotide that hybridizes under medium stringency conditions, medium-high stringency conditions, high stringency conditions or very-high stringency conditions with:
   (i) the mature polypeptide coding sequence of SEQ ID NO: 1, and/or
   (ii) the full-length complementary strand of (i);
(c) a polypeptide encoded by a polynucleotide having at least 80% sequence identity to the mature polypeptide coding sequence of SEQ ID NO: 1;
(d) a variant of the polypeptide of SEQ ID NO: 3 comprising a substitution, deletion, and/or insertion of one or more (e.g. several) positions; and
(e) a fragment of the polypeptide of (a), (b), (c), or (d) that has protease activity, in animal feed and detergent compositions.

The present invention also relates to variant polypeptides having protease activity and having at least 85%, e.g. at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% sequence identity to SEQ ID NO: 3 comprising at least one substitution, deletion, and/or insertion of at least one or more (several) amino acids of SEQ ID NO: 3.

The present invention further relates to compositions comprising an isolated polypeptide having protease activity, selected from the group consisting of:
(a) a polypeptide having at least 80%, e.g., at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 3;
(b) a polypeptide encoded by a polynucleotide that hybridizes under medium stringency conditions, medium-high stringency conditions, high stringency conditions or very-high stringency conditions with:
   (i) the mature polypeptide coding sequence of SEQ ID NO: 1; and/or
   (ii) the full-length complementary strand of (i);
(c) a polypeptide encoded by a polynucleotide having at least 80%, e.g., at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to the mature polypeptide coding sequence of SEQ ID NO: 3;
(d) a variant comprising a substitution, deletion, and/or insertion of one or more (several) amino acids of SEQ ID NO: 3; and
(e) a fragment of a polypeptide of (a), (b), (c) or (d), that has protease activity.

The compositions can be detergent compositions or animal feed compositions. The present invention also relates to isolated polynucleotides encoding the polypeptides of the present invention, nucleic acid constructs, recombinant expression vectors, recombinant host cells comprising the polynucleotides, and to methods of recombinantly producing the polypeptides. The present invention also relates to methods for preparing a composition for use in animal feed, methods for improving the nutritional value of an animal feed, and methods of treating proteins to be used in animal feed compositions.

OVERVIEW OF SEQUENCE LISTING

SEQ ID NO: 1 is the DNA sequence of S1 protease 1 as isolated from *Saccharomonospora viridis*.

SEQ ID NO: 2 is the amino acid sequence as deduced from SEQ ID NO: 1 (Uniprot: C7MV18).

SEQ ID NO: 3 is the amino acid sequence of the mature *Saccharomonospora viridis* protease.

SEQ ID NO: 4 is a *Bacillus clausii* C360 secretion signal.

SEQ ID NO: 5 is the DNA sequence of the 10R protease (WO 05/035747, SEQ ID NO: 1).

SEQ ID NO: 6 is the amino acid sequence of the 10R protease (WO 05/035747, SEQ ID NO: 2).

SEQ ID NO: 7 is the amino acid sequence of a protease from *Saccharomonospora cyanea* NA-134 (Uniprot: H5XEH4).

SEQ ID NO: 8 is the amino acid sequence of a trypsin protease proenzyme from *Saccharomonospora azurea* SZMC 14600 (Uniprot: H0K7C9).

SEQ ID NO: 9 is the amino acid sequence of an endopeptidase from *Saccharomonospora glauca* K62 (Uniprot: H1JPF3).

SEQ ID NO: 10 is the amino acid sequence of an endopeptidase from *Saccharomonospora paurometabolica* (Uniprot: G4J6Q2).

SEQ ID NO: 11 is the amino acid sequence of an endopeptidase from *Saccharomonospora paurometabolica* (Uniprot: G41XC2).

SEQ ID NO: 12 is the amino acid sequence of a serine protease from *Saccharopolyspora erythraea* (Uniprot: A4FNQ0).

SEQ ID NO: 13 is the amino acid sequence of an alpha-lytic protease from *Saccharomonospora xinjiangensis* XJ-54 (Uniprot: I0V8H8).

SEQ ID NO: 14 is the amino acid sequence of a membrane protein from *Saccharomonospora azurea* NA-128 (Uniprot: H8GAL4).
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BRIEF DESCRIPTION OF THE FIGURES

Figure 1 shows the pH-activity profile of S1 protease 1 from *Saccharomonospora viridis* and 10R protease on the Suc-AAPF-pNA substrate at 25°C.

Figure 2 shows the pH-stability profile of *S1 protease 1 from Saccharomonospora viridis* and 10R protease (residual activity after 2 hours at 37°C).

Figure 3 shows the temperature activity profile of *S1 protease 1 from Saccharomonospora viridis* and 10R protease on Protazyme AK at pH 7.0.

Figure 4 shows the P1-specificity of *S1 protease 1 from Saccharomonospora viridis* and 10R protease on 10 Suc-AAPX-pNA substrates at pH 9.0, 25°C.

Figure 5 shows the pH-activity profile of *S1 protease 1 from Saccharomonospora viridis* and 10R protease on soybean-maize meal at 40°C.

DEFINITIONS

Allelic variant: The term "allelic variant" means any of two or more alternative forms of a gene occupying the same chromosomal locus. Allelic variation arises naturally through
mutation, and may result in polymorphism within populations. Gene mutations can be silent (no change in the encoded polypeptide) or may encode polypeptides having altered amino acid sequences. An allelic variant of a polypeptide is a polypeptide encoded by an allelic variant of a gene.

cDNA: The term "cDNA" means a DNA molecule that can be prepared by reverse transcription from a mature, spliced, mRNA molecule obtained from a eukaryotic cell. cDNA lacks intron sequences that may be present in the corresponding genomic DNA. The initial, primary RNA transcript is a precursor to mRNA that is processed through a series of steps, including splicing, before appearing as mature spliced mRNA.

Coding sequence: The term "coding sequence" means a polynucleotide, which directly specifies the amino acid sequence of a polypeptide. The boundaries of the coding sequence are generally determined by an open reading frame, which usually begins with the ATG start codon or alternative start codons such as GTG and TTG and ends with a stop codon such as TAA, TAG, and TGA. The coding sequence may be a DNA, cDNA, synthetic, or recombinant polynucleotide.

Colour clarification: During washing and wearing loose or broken fibers can accumulate on the surface of the fabrics. One consequence can be that the colours of the fabric appear less bright or less intense because of the surface contaminations. Removal of the loose or broken fibers from the textile will partly restore the original colours and looks of the textile. By the term "colour clarification", as used herein, is meant the partial restoration of the initial colours of textile.

Control sequences: The term "control sequences" means all components necessary for the expression of a polynucleotide encoding a polypeptide of the present invention. Each control sequence may be native or foreign to the polynucleotide encoding the polypeptide or native or foreign to each other. Such control sequences include, but are not limited to, a leader, polyadenylation sequence, propeptide sequence, promoter, signal peptide sequence, and transcription terminator. At a minimum, the control sequences include a promoter, and transcriptional and translational stop signals. The control sequences may be provided with linkers for the purpose of introducing specific restriction sites facilitating ligation of the control sequences with the coding region of the polynucleotide encoding a polypeptide.

Detergent component: the term "detergent component" is defined herein to mean the types of chemicals which can be used in detergent compositions. Examples of detergent components are surfactants, hydrolytes, builders, co-builders, chelators or chelating agents, bleaching system or bleach components, polymers, fabric hueing agents, fabric conditioners, foam boosters, suds suppressors, dispersants, dye transfer inhibitors, fluorescent whitening agents, perfume, optical brighteners, bactericides, fungicides, soil suspending agents, soil release polymers, anti-redeposition agents, enzyme inhibitors or stabilizers, enzyme activators,
antioxidants, and solubilizers. The detergent composition may comprise one or more of any type of detergent component.

**Detergent Composition:** the term "detergent composition" refers to compositions that find use in the removal of undesired compounds from items to be cleaned, such as textiles, dishes, and hard surfaces. The detergent composition may be used to e.g. clean textiles, dishes and hard surfaces for both household cleaning and industrial cleaning. The terms encompass any materials/compounds selected for the particular type of cleaning composition desired and the form of the product (e.g., liquid, gel, powder, granulate, paste, or spray compositions) and includes, but is not limited to, detergent compositions (e.g., liquid and/or solid laundry detergents and fine fabric detergents; hard surface cleaning formulations, such as for glass, wood, ceramic and metal counter tops and windows; carpet cleaners; oven cleaners; fabric fresheners; fabric softeners; and textile and laundry pre-spotters, as well as dish wash detergents). In addition to containing a protease of the invention, the detergent formulation may contain one or more additional enzymes (such as other proteases, amylases, lipases, cutinases, cellulases, endoglucanases, xyloglucanases, pectinases, pectin lyases, xanthanases, peroxidaes, haloperoxigenases, catalases and mannanases, or any mixture thereof), and/or components such as surfactants, builders, chelators or chelating agents, bleach system or bleach components, polymers, fabric conditioners, foam boosters, suds suppressors, dyes, perfume, tannish inhibitors, optical brighteners, bactericides, fungicides, soil suspending agents, anti corrosion agents, enzyme inhibitors or stabilizers, enzyme activators, transferase(s), hydrolytic enzymes, oxido reductases, bluing agents and fluorescent dyes, antioxidants, and solubilizers.

**Dish wash:** The term "dish wash" refers to all forms of washing dishes, e.g. by hand or automatic dish wash. Washing dishes includes, but is not limited to, the cleaning of all forms of crockery such as plates, cups, glasses, bowls, all forms of cutlery such as spoons, knives, forks and serving utensils as well as ceramics, plastics, metals, china, glass and acrylics.

**Dish washing composition:** The term "dish washing composition" refers to all forms of compositions for cleaning hard surfaces. The present invention is not restricted to any particular type of dish wash composition or any particular detergent.

**Enzyme Detergency benefit:** The term "enzyme detergency benefit" is defined herein as the advantageous effect an enzyme may add to a detergent compared to the same detergent without the enzyme. Important detergency benefits which can be provided by enzymes are stain removal with no or very little visible soils after washing and or cleaning, prevention or reduction of redeposition of soils released in the washing process an effect that also is termed anti-redeposition, restoring fully or partly the whiteness of textiles, which originally were white but after repeated use and wash have obtained a greyish or yellowish appearance an effect that also is termed whitening. Textile care benefits, which are not directly related to catalytic stain removal or prevention of redeposition of soils are also important for enzyme detergency.
benefits. Examples of such textile care benefits are prevention or reduction of dye transfer from one fabric to another fabric or another part of the same fabric an effect that is also termed dye transfer inhibition or anti-backstaining, removal of protruding or broken fibers from a fabric surface to decrease pilling tendencies or remove already existing pills or fuzz an effect that also is termed anti-pilling, improvement of the fabric-softness, colour clarification of the fabric and removal of particulate soils which are trapped in the fibers of the fabric or garment. Enzymatic bleaching is a further enzyme detergency benefit where the catalytic activity generally is used to catalyze the formation of bleaching component such as hydrogen peroxide or other peroxides.

Expression: The term "expression" includes any step involved in the production of the polypeptide including, but not limited to, transcription, post-transcriptional modification, translation, post-translational modification, and secretion.

Expression vector: The term "expression vector" means a linear or circular DNA molecule that comprises a polynucleotide encoding a polypeptide and is operably linked to additional nucleotides that provide for its expression.

Fragment: The term "fragment" means a polypeptide having one or more (several) amino acids deleted from the amino and/or carboxyl terminus of a mature polypeptide; wherein the fragment has protease activity. In one aspect, a fragment contains at least 130 amino acid residues (e.g., amino acids 15 to 144 of SEQ ID NO: 2); in another aspect a fragment contains at least 140 amino acid residues (e.g., amino acids 10 to 149 of SEQ ID NO: 2); in a further aspect a fragment contains at least 150 amino acid residues (e.g., amino acids 5 to 154 of SEQ ID NO: 2). In another aspect, a fragment contains at least 130 amino acid residues (e.g., amino acids 15 to 144 of SEQ ID NO: 3); in another aspect a fragment contains at least 140 amino acid residues (e.g., amino acids 10 to 149 of SEQ ID NO: 3); in a further aspect a fragment contains at least 150 amino acid residues (e.g., amino acids 5 to 154 of SEQ ID NO: 3).

Hard surface cleaning: The term "Hard surface cleaning" is defined herein as cleaning of hard surfaces wherein hard surfaces may include floors, tables, walls, roofs etc. as well as surfaces of hard objects such as cars (car wash) and dishes (dish wash). Dish washing includes but are not limited to cleaning of plates, cups, glasses, bowls, and cutlery such as spoons, knives, forks, serving utensils, ceramics, plastics, metals, china, glass and acrylics.

Host cell: The term "host cell" means any cell type that is susceptible to transformation, transfection, transduction, and the like with a nucleic acid construct or expression vector comprising a polynucleotide of the present invention. The term "host cell" encompasses any progeny of a parent cell that is not identical to the parent cell due to mutations that occur during replication.

Improved wash performance: The term "improved wash performance" is defined herein as a (variant) enzyme (also a blend of enzymes, not necessarily only variants but also backbones, and in combination with certain cleaning composition etc.) displaying an alteration
of the wash performance of a protease variant relative to the wash performance of the parent protease variant e.g. by increased stain removal. The term "wash performance" includes wash performance in laundry but also e.g. in dish wash.

**Isolated polynucleotide:** The term "isolated polynucleotide" means a polynucleotide that is in a form or environment that does not occur in nature, such as (1) any non-naturally occurring polynucleotide, (2) any polynucleotide that is at least partially removed from one or more or all of the naturally occurring constituents with which it is associated in nature; (3) any polynucleotide that is modified by the hand of man relative to that polynucleotide as found in nature or (4) any polynucleotide modified by increasing the amount of the polynucleotide relative to other components with which it is naturally associated (e.g., recombinant production in a host cell; multiple copies of a gene encoding the substance; and use of a stronger promoter than the promoter naturally associated with the gene encoding the substance). In one aspect, the isolated polynucleotide is at least 1% pure, e.g., at least 5% pure, more at least 10% pure, at least 20% pure, at least 40% pure, at least 60% pure, at least 80% pure, at least 90% pure, and at least 95% pure, as determined by agarose electrophoresis. The polynucleotides may be of genomic, cDNA, RNA, semisynthetic, synthetic origin, or any combinations thereof.

**Isolated polypeptide:** The term "isolated polypeptide" means a polypeptide that is in a form or environment that does not occur in nature, such as (1) any non-naturally occurring polypeptide, (2) any polypeptide that is at least partially removed from one or more or all of the naturally occurring constituents with which it is associated in nature; (3) any polypeptide that is modified by the hand of man relative to that polypeptide as found in nature in admixture with other components, such as other polypeptides, secondary metabolites, salts, *et alia* or (4) any polypeptide modified by increasing the amount of the polypeptide relative to other components with which it is naturally associated. In one aspect, the polypeptide is at least 1% pure, e.g., at least 5% pure, at least 10% pure, at least 20% pure, at least 40% pure, at least 60% pure, at least 80% pure, and at least 90% pure, as determined by SDS-PAGE.

**Laundering:** The term "laundering" relates to both household laundering and industrial laundering and means the process of treating textiles with a solution containing a cleaning or detergent composition of the present invention. The laundering process can for example be carried out using e.g. a household or an industrial washing machine or can be carried out by hand.

**Mature polypeptide:** The term "mature polypeptide" means a polypeptide in its final form following translation and any post-translational modifications, such as N-terminal processing, C-terminal truncation, glycosylation, phosphorylation, etc. In one aspect, the mature polypeptide is amino acids 1 to 160 in the numbering of SEQ ID NO: 2; amino acids -198 to -167 in the numbering of SEQ ID NO: 2 is a signal peptide.
Mature polypeptide coding sequence: The term "mature polypeptide coding sequence" means a polynucleotide that encodes a mature polypeptide having protease activity. In one aspect, the mature polypeptide coding sequence is nucleotides 595-1074 in the numbering of SEQ ID NO: 1. Further nucleotides 1 to 96 in the numbering of SEQ ID NO: 1 encode a signal peptide.

Nucleic acid construct: The term "nucleic acid construct" means a nucleic acid molecule, either single- or double-stranded, which is isolated from a naturally occurring gene or is modified to contain segments of nucleic acids in a manner that would not otherwise exist in nature or which is synthetic. The term nucleic acid construct is synonymous with the term "expression cassette" when the nucleic acid construct contains the control sequences required for expression of a coding sequence of the present invention.

Operably linked: The term "operably linked" means a configuration in which a control sequence is placed at an appropriate position relative to the coding sequence of a polynucleotide such that the control sequence directs the expression of the coding sequence.

Polypeptides Having Protease Activity: Polypeptides having protease activity, or proteases, are sometimes also designated peptidases, proteinases, peptide hydrolases, or proteolytic enzymes. Proteases may be of the exo-type that hydrolyse peptides starting at either end thereof, or of the endo-type that act internally in polypeptide chains (endopeptidases). Endopeptidases show activity on N- and C-terminally blocked peptide substrates that are relevant for the specificity of the protease in question.


The present invention provides for the use of polypeptides having protease activity in animal feed and detergent compositions. It also provides polynucleotides encoding the polypeptides. The proteases of the invention are serine proteases of the peptidase family S1. The proteases of the invention exhibit surprising pH properties, which makes them interesting candidates for use in animal feed. The proteases of the invention thus are active on Suc-Ala-Ala-Pro-Phe-pNA within a broad pH range of 5-11, exhibit especially high activity in the pH range of 7-11, are active on a feed relevant soybean meal-maize meal substrate within a broad
physiological pH range of pH 3-7 and retain 100% activity after being subjected for 2 hours to a pH as low as 3 and more than 40% after being subjected for 2 hours to a pH as low as 2.

The proteases of the invention and for use according to the invention are selected from the group consisting of:

(a) proteases belonging to the EC 3.4.21. enzyme group; and/or
(b) Serine proteases of the peptidase family S1;


Proteases of the invention are endopeptidases (EC 3.4.21). There are several protease activity types: The three main activity types are: trypsin-like where there is cleavage of amide substrates following Arg or Lys at P1, chymotrypsin-like where cleavage occurs following one of the hydrophobic amino acids at P1, and elastase-like with cleavage following an Ala at P1.

The polypeptides of the present invention have at least 20%, e.g., at least 40%, at least 50%, at least 60%, at least 70%, at least 80%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, and at least 100% of the protease activity of the mature polypeptide of SEQ ID NO: 2.

More specifically the proteases used in the invention are those that prefer a hydrophobic aromatic amino acid residue in the P1 position.

For determining whether a given protease is a serine protease, and a family S1 protease, reference is made to the above Handbook and the principles indicated therein. Such determination can be carried out for all types of proteases, be it naturally occurring or wild-type proteases; or genetically engineered or synthetic proteases.

The peptidases of family S1 contain the catalytic triad His, Asp and Ser in that order. Mutation of any of the amino acids of the catalytic triad will result in loss of enzyme activity. The amino acids of the catalytic triad of the S1 protease 1 from Saccharomonospora viridis (SEQ ID NO: 3) are probably positions His-32, Asp-56 and Ser-137.

Protease activity can be measured using any assay, in which a substrate is employed, that includes peptide bonds relevant for the specificity of the protease in question. Assay-pH and assay-temperature are likewise to be adapted to the protease in question. Examples of assay-pH-values are pH 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Examples of assay-temperatures are 15, 20, 25, 30, 35, 37, 40, 45, 50, 55, 60, 65, 70, 80, 90, or 95°C. Examples of general protease substrates are casein, bovine serum albumin and haemoglobin. In the classical Anson and Mirsky method, denatured haemoglobin is used as substrate and after the assay incubation with the protease in question, the amount of trichloroacetic acid soluble haemoglobin is determined

For the purpose of the present invention, protease activity was determined using assays which are described in "Materials and Methods", such as the Suc-AAPF-pNA assay, Protazyme AK assay, Suc-AAPX-pNA assay and o-Phthalaldehyde (OPA). For the Protazyme AK assay, insoluble Protazyme AK (Azurine-Crosslinked Casein) substrate liberates a blue colour when incubated with the protease and the colour is determined as a measurement of protease activity. For the Suc-AAPF-pNA assay, the colourless Suc-AAPF-pNA substrate liberates yellow paranitroaniline when incubated with the protease and the yellow colour is determined as a measurement of protease activity.

Sequence Identity: The relatedness between two amino acid sequences or between two nucleotide sequences is described by the parameter "sequence identity".

For purposes of the present invention, the degree of sequence identity between two amino acid sequences is determined using the Needleman-Wunsch algorithm (Needleman and Wunsch, 1970, J. Mol. Biol. 48: 443-453) as implemented in the Needle program of the EMBOSS package (EMBOSS: The European Molecular Biology Open Software Suite, Rice et al., 2000, Trends Genet. 16: 276-277), preferably version 3.0.0 or later. Version 6.1.0 was used. The optional parameters used are gap open penalty of 10, gap extension penalty of 0.5, and the EBLOSUM62 (EMBOSS version of BLOSUM62) substitution matrix. The output of Needle labelled "longest identity" (obtained using the -nobrief option) is used as the percent identity and is calculated as follows:

\[
\frac{\text{Identical Residues} \times 100}{\text{Length of Alignment} - \text{Total Number of Gaps in Alignment}}
\]

For purposes of the present invention, the degree of sequence identity between two deoxyribonucleotide sequences is determined using the Needleman-Wunsch algorithm (Needleman and Wunsch, 1970, supra) as implemented in the Needle program of the EMBOSS package (EMBOSS: The European Molecular Biology Open Software Suite, Rice et al., 2000, supra), preferably version 3.0.0 or later. Version 6.1.0 was used. The optional parameters used are gap open penalty of 10, gap extension penalty of 0.5, and the EDNAFULL (EMBOSS version of NCBI NUC4.4) substitution matrix. The output of Needle labelled "longest identity" (obtained using the -nobrief option) is used as the percent identity and is calculated as follows:

\[
\frac{\text{Identical Deoxyribonucleotides} \times 100}{\text{Length of Alignment} - \text{Total Number of Gaps in Alignment}}
\]

Stringency conditions: The different stringency conditions are defined as follows.

The term "very low stringency conditions" means for probes of at least 100 nucleotides in length, prehybridization and hybridization at 42°C in 5X SSPE, 0.3% SDS, 200 micrograms/ml sheared and denatured salmon sperm DNA, and 35% formamide, following standard Southern
blotting procedures for 12 to 24 hours. The carrier material is finally washed three times each for 15 minutes using 1.5X SSC, 0.2% SDS at 65°C.

The term "low stringency conditions" means for probes of at least 100 nucleotides in length, prehybridization and hybridization at 42°C in 5X SSPE, 0.3% SDS, 200 micrograms/ml sheared and denatured salmon sperm DNA, and 35% formamide, following standard Southern blotting procedures for 12 to 24 hours. The carrier material is finally washed three times each for 15 minutes using 0.8X SSC, 0.2% SDS at 65°C.

The term "medium stringency conditions" means for probes of at least 100 nucleotides in length, prehybridization and hybridization at 42°C in 5X SSPE, 0.3% SDS, 200 micrograms/ml sheared and denatured salmon sperm DNA, and 35% formamide, following standard Southern blotting procedures for 12 to 24 hours. The carrier material is finally washed three times each for 15 minutes using 0.4x SSC, 0.2% SDS at 65°C.

The term "medium-high stringency conditions" means for probes of at least 100 nucleotides in length, prehybridization and hybridization at 42°C in 5X SSPE, 0.3% SDS, 200 micrograms/ml sheared and denatured salmon sperm DNA, and 35% formamide, following standard Southern blotting procedures for 12 to 24 hours. The carrier material is finally washed three times each for 15 minutes using 0.2X SSC, 0.2% SDS at 65°C.

The term "high stringency conditions" means for probes of at least 100 nucleotides in length, prehybridization and hybridization at 42°C in 5X SSPE, 0.3% SDS, 200 micrograms/ml sheared and denatured salmon sperm DNA, and 35% formamide, following standard Southern blotting procedures for 12 to 24 hours. The carrier material is finally washed three times each for 15 minutes using 0.2X SSC, 0.2% SDS at 70°C.

The term "very high stringency conditions" means for probes of at least 100 nucleotides in length, prehybridization and hybridization at 42°C in 5X SSPE, 0.3% SDS, 200 micrograms/ml sheared and denatured salmon sperm DNA, and 35% formamide, following standard Southern blotting procedures for 12 to 24 hours. The carrier material is finally washed three times each for 15 minutes using 0.1X SSC, 0.2% SDS at 70°C.

Subsequence: The term "subsequence" means a polynucleotide having one or more (several) nucleotides deleted from the 5' and/or 3' end of a mature polypeptide coding sequence; wherein the subsequence encodes a fragment having protease activity. In one aspect, a subsequence contains at least 390 nucleotides (e.g., nucleotides 637 to 1026 of SEQ ID NO: 1), e.g., and at least 420 nucleotides (e.g., nucleotides 622 to 1041 of SEQ ID NO: 1); e.g., and at least 450 nucleotides (e.g., nucleotides 607 to 1056 of SEQ ID NO: 1).

Substantially pure polynucleotide: The term "substantially pure polynucleotide" means a polynucleotide preparation free of other extraneous or unwanted nucleotides and in a form suitable for use within genetically engineered polypeptide production systems. Thus, a substantially pure polynucleotide contains at most 10%, at most 8%, at most 6%, at most 5%, at
most 4%, at most 3%, at most 2%, at most 1%, and at most 0.5% by weight of other polynucleotide material with which it is natively or recombinantly associated. A substantially pure polynucleotide may, however, include naturally occurring 5' and 3' untranslated regions, such as promoters and terminators. Preferably, the polynucleotide is at least 90% pure, e.g., at least 92% pure, at least 94% pure, at least 95% pure, at least 96% pure, at least 97% pure, at least 98% pure, at least 99% pure, and at least 99.5% pure, and 100% pure by weight. The polynucleotides of the present invention are preferably in a substantially pure form.

Substantially pure polypeptide: The term "substantially pure polypeptide" means a preparation that contains at most 10%, at most 8%, at most 6%, at most 5%, at most 4%, at most 3%, at most 2%, at most 1%, and at most 0.5% by weight of other polypeptide material with which it is natively or recombinantly associated. Preferably, the polypeptide is at least 92% pure, e.g., at least 94% pure, at least 95% pure, at least 96% pure, at least 97% pure, at least 98% pure, at least 99%, at least 99.5% pure, and 100% pure by weight of the total polypeptide material present in the preparation. The polypeptides of the present invention are preferably in a substantially pure form. This can be accomplished, for example, by preparing the polypeptide by well known recombinant methods or by classical purification methods.

Textile: The term "textile" means any textile material including yarns, yarn intermediates, fibers, non-woven materials, natural materials, synthetic materials, and any other textile material, fabrics made of these materials and products made from fabrics (e.g., garments and other articles). The textile or fabric may be in the form of knits, wovens, denims, non-wovens, felts, yarns, and towelling. The textile may be cellulose based such as natural cellulosics, including cotton, flax/linen, jute, ramie, sisal or coir or manmade cellulosics (e.g. originating from wood pulp) including viscose/rayon, ramie, cellulose acetate fibers (tricell), lyocell or blends thereof. The textile or fabric may also be non-cellulose based such as natural polyamides including wool, camel, cashmere, mohair, rabbit and silk or synthetic polymer such as nylon, aramid, polyester, acrylic, polypropylene and spandex/elastane, or blends thereof as well as blend of cellulose based and non-cellulose based fibers. Examples of blends are blends of cotton and/or rayon/viscose with one or more companion material such as wool, synthetic fibers (e.g. polyamide fibers, acrylic fibers, polyester fibers, polyvinyl alcohol fibers, polyvinyl chloride fibers, polyurethane fibers, polyurea fibers, aramid fibers), and cellulose-containing fibers (e.g. rayon/viscose, ramie, flax/linen, jute, cellulose acetate fibers, lyocell). Fabric may be conventional washable laundry, for example stained household laundry. When the term fabric or garment is used it is intended to include the broader term textiles as well.

Textile care benefit: "Textile care benefits", which are not directly related to catalytic stain removal or prevention of redeposition of soils, are also important for enzyme detergency benefits. Examples of such textile care benefits are prevention or reduction of dye transfer from one textile to another textile or another part of the same textile an effect that is also termed dye
transfer inhibition or anti-backstaining, removal of protruding or broken fibers from a textile surface to decrease pilling tendencies or remove already existing pills or fuzz; an effect that also is termed anti-pilling, improvement of the textile-softness, colour clarification of the textile and removal of particulate soils which are trapped in the fibers of the textile. Enzymatic bleaching is a further enzyme detergency benefit where the catalytic activity generally is used to catalyze the formation of bleaching component such as hydrogen peroxide or other peroxides or other bleaching species.

Variant: The term "variant" means a polypeptide having protease activity comprising an alteration, \textit{i.e.}, a substitution, insertion, and/or deletion of one or more (several) amino acid residues at one or more (several) positions. A substitution means a replacement of an amino acid occupying a position with a different amino acid; a deletion means removal of an amino acid occupying a position; and an insertion means adding 1, 2, or 3 amino acids adjacent to an amino acid occupying a position.

Wash performance: The term "wash performance" is used as an enzyme's ability to remove stains present on the object to be cleaned during e.g. wash or hard surface cleaning.

Whiteness: The term "Whiteness" is defined herein as a broad term with different meanings in different regions and for different customers. Loss of whiteness can e.g. be due to greying, yellowing, or removal of optical brighteners/hueing agents. Greying and yellowing can be due to soil redeposition, body soils, colouring from e.g. iron and copper ions or dye transfer. Whiteness might include one or several issues from the list below: Colorant or dye effects; Incomplete stain removal (e.g. body soils, sebum etc); Re-deposition (greying, yellowing or other discolorations of the object) (removed soils re-associates with other part of textile, soiled or unsoiled); Chemical changes in textile during application; and Clarification or brightening of colours.

DETAILED DESCRIPTION OF THE INVENTION

Polypeptides Having Protease Activity

The present invention relates to the use in animal feed or detergents of isolated polypeptides having protease activity selected from the group consisting of:

(a) a polypeptide having at least 80% sequence identity to the polypeptide of SEQ ID NO: 3;

(b) a polypeptide encoded by a polynucleotide that hybridizes under medium stringency conditions, medium-high stringency conditions, high stringency conditions or very-high stringency conditions with:

(i) the mature polypeptide coding sequence of SEQ ID NO: 1, and/or

(ii) the full-length complementary strand of (i);
(c) a polypeptide encoded by a polynucleotide having at least 80% sequence identity to the mature polypeptide coding sequence of SEQ ID NO: 1;
(d) a variant of the polypeptide of SEQ ID NO: 3 comprising a substitution, deletion, and/or insertion of one or more (e.g. several) positions; and
(e) a fragment of the polypeptide of (a), (b), (c), or (d) that has protease activity.

The present invention relates to the use in animal feed or detergents of isolated polypeptides having a sequence identity to the polypeptide of SEQ ID NO: 3 of at least 80%, e.g. at least 85%, e.g., at least 87%, at least 89%, at least 90%, at least 93%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100%, which have protease activity. In one aspect, the polypeptides differ by no more than thirty two amino acids, e.g., by thirty amino acids, by twenty five amino acids, by twenty amino acids, by fifteen amino acids, by ten amino acids, by eight amino acids, by seven amino acids, by six amino acids, by five amino acids, by four amino acids, by three amino acids, by two amino acids, and by one amino acid from the polypeptide of SEQ ID NO: 3. Specifically the isolated polypeptides having protease activity for the use in animal feed or detergents should be selected from the group consisting of:

(a) a polypeptide having at least 85% sequence identity to the polypeptide of SEQ ID NO: 3;
(b) a polypeptide encoded by a polynucleotide that hybridizes under medium stringency conditions, medium-high stringency conditions, high stringency conditions or very-high stringency conditions with:
   (i) the mature polypeptide coding sequence of SEQ ID NO: 1, and/or
   (ii) the full-length complementary strand of (i);
(c) a polypeptide encoded by a polynucleotide having at least 85% sequence identity to the mature polypeptide coding sequence of SEQ ID NO: 1;
(d) a variant of the polypeptide of SEQ ID NO: 3 comprising a substitution, deletion, and/or insertion of one or more (e.g. several) positions; and
(e) a fragment of the polypeptide of (a), (b), (c), or (d) that has protease activity.

Further isolated polypeptides having protease activity and for the use in animal feed or detergents should be selected from the group consisting of:

(a) a polypeptide having at least 90% sequence identity to the polypeptide of SEQ ID NO: 3;
(b) a polypeptide encoded by a polynucleotide that hybridizes under medium stringency conditions, medium-high stringency conditions, high stringency conditions or very-high stringency conditions with:
(i) the mature polypeptide coding sequence of SEQ ID NO: 1, and/or
(ii) the full-length complementary strand of (i);
(c) a polypeptide encoded by a polynucleotide having at least 90% sequence identity
to the mature polypeptide coding sequence of SEQ ID NO: 1;
(d) a variant of the polypeptide of SEQ ID NO: 3 comprising a substitution, deletion,
and/or insertion of one or more (e.g. several) positions; and
(e) a fragment of the polypeptide of (a), (b), (c), or (d) that has protease activity.

Specifically the isolated polypeptides having protease activity for the use in animal feed
or detergents should be selected from the group consisting of:
(a) a polypeptide having at least 95% sequence identity to the polypeptide of SEQ ID
NO: 3;
(b) a polypeptide encoded by a polynucleotide that hybridizes under medium
stringency conditions, medium-high stringency conditions, high stringency conditions or very-
high stringency conditions with:
(i) the mature polypeptide coding sequence of SEQ ID NO: 1, and/or
(ii) the full-length complementary strand of (i);
(c) a polypeptide encoded by a polynucleotide having at least 95% sequence identity
to the mature polypeptide coding sequence of SEQ ID NO: 1;
(d) a variant of the polypeptide of SEQ ID NO: 3 comprising a substitution, deletion,
and/or insertion of one or more (e.g. several) positions; and
(e) a fragment of the polypeptide of (a), (b), (c), or (d) that has protease activity.

Further isolated polypeptides having protease activity and for the use in animal feed or
detergents should be selected from the group consisting of:
(a) a polypeptide having at least 97% sequence identity to the polypeptide of SEQ ID
NO: 3;
(b) a polypeptide encoded by a polynucleotide that hybridizes under medium
stringency conditions, medium-high stringency conditions, high stringency conditions or very-
high stringency conditions with:
(i) the mature polypeptide coding sequence of SEQ ID NO: 1, and/or
(ii) the full-length complementary strand of (i);
(c) a polypeptide encoded by a polynucleotide having at least 97% sequence identity
to the mature polypeptide coding sequence of SEQ ID NO: 1;
(d) a variant of the polypeptide of SEQ ID NO: 3 comprising a substitution, deletion,
and/or insertion of one or more (e.g. several) positions; and
(e) a fragment of the polypeptide of (a), (b), (c), or (d) that has protease activity.
Specifically the isolated polypeptides having protease activity for the use in animal feed or detergents should be selected from the group consisting of:

(a) a polypeptide having at least 98% sequence identity to the polypeptide of SEQ ID NO: 3; 

(b) a polypeptide encoded by a polynucleotide that hybridizes under medium stringency conditions, medium-high stringency conditions, high stringency conditions or very-high stringency conditions with:

   (i) the mature polypeptide coding sequence of SEQ ID NO: 1, and/or

   (ii) the full-length complementary strand of (i);

(c) a polypeptide encoded by a polynucleotide having at least 98% sequence identity to the mature polypeptide coding sequence of SEQ ID NO: 1;

(d) a variant of the polypeptide of SEQ ID NO: 3 comprising a substitution, deletion, and/or insertion of one or more (e.g. several) positions; and

(e) a fragment of the polypeptide of (a), (b), (c), or (d) that has protease activity.

Further isolated polypeptides having protease activity and for the use in animal feed or detergents should be selected from the group consisting of:

(a) a polypeptide having at least 99% sequence identity to the polypeptide of SEQ ID NO: 3; 

(b) a polypeptide encoded by a polynucleotide that hybridizes under medium stringency conditions, medium-high stringency conditions, high stringency conditions or very-high stringency conditions with:

   (i) the mature polypeptide coding sequence of SEQ ID NO: 1, and/or

   (ii) the full-length complementary strand of (i);

(c) a polypeptide encoded by a polynucleotide having at least 99% sequence identity to the mature polypeptide coding sequence of SEQ ID NO: 1;

(d) a variant of the polypeptide of SEQ ID NO: 3 comprising a substitution, deletion, and/or insertion of one or more (e.g. several) positions; and

(e) a fragment of the polypeptide of (a), (b), (c), or (d) that has protease activity.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 85% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 86% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 87% sequence identity to the polypeptide of SEQ ID NO: 3.
An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 88% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 89% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 90% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 91% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 92% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 93% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 94% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 95% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 96% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 97% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 98% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having at least 99% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is an isolated polypeptide for use in animal feed or detergents having 100% sequence identity to the polypeptide of SEQ ID NO: 3.

A polypeptide to be used in the present invention preferably comprises or consists of the amino acid sequence of SEQ ID NO: 3 or an allelic variant thereof; or is a fragment thereof having protease activity. In another aspect, the polypeptide comprises or consists of the mature polypeptide of SEQ ID NO: 2. In a further aspect, the polypeptide comprises or consists of the polypeptide of SEQ ID NO: 3. In another aspect, the polypeptide comprises or consists of amino acids 1 to 160 of SEQ ID NO: 2, amino acids 5 to 154 of SEQ ID NO: 2, or amino acids 10 to 149 of SEQ ID NO: 2. In another aspect, the polypeptide comprises or consists of amino acids 1 to 160 of SEQ ID NO: 3, amino acids 5 to 154 of SEQ ID NO: 3, or amino acids 10 to 149 of SEQ ID NO: 3.
The present invention also relates to isolated polypeptides having protease activity that are encoded by polynucleotides that hybridize under medium stringency conditions, medium-high stringency conditions, high stringency conditions or very-high stringency conditions with (i) the mature polypeptide coding sequence of SEQ ID NO: 1, and/or (ii) the full-length complementary strand of (i) (J. Sambrook, E.F. Fritsch, and T. Maniatis, 1989, Molecular Cloning, A Laboratory Manual, 2d edition, Cold Spring Harbor, New York).

The polynucleotide of SEQ ID NO: 1 or a subsequence thereof, as well as the amino acid sequence of SEQ ID NO: 2 or SEQ ID NO: 3, or a fragment thereof, may be used to design nucleic acid probes to identify and clone DNA encoding polypeptides having protease activity from strains of different genera or species according to methods well known in the art. In particular, such probes can be used for hybridization with the genomic or cDNA of the genus or species of interest, following standard Southern blotting procedures, in order to identify and isolate the corresponding gene therein. Such probes can be considerably shorter than the entire sequence, but should be at least 14, e.g., at least 25, at least 35, or at least 70 nucleotides in length. Preferably, the nucleic acid probe is at least 100 nucleotides in length, e.g., at least 200 nucleotides, at least 300 nucleotides, at least 400 nucleotides, at least 500 nucleotides, at least 600 nucleotides, at least 700 nucleotides, at least 800 nucleotides, or at least 900 nucleotides in length. Both DNA and RNA probes can be used. The probes are typically labelled for detecting the corresponding gene (for example, with $^{32}$P, $^3$H, $^{35}$S, biotin, or avidin). Such probes are encompassed by the present invention.

A genomic DNA or cDNA library prepared from such other strains may be screened for DNA that hybridizes with the probes described above and encodes a polypeptide having protease activity. Genomic or other DNA from such other strains may be separated by agarose or polyacrylamide gel electrophoresis, or other separation techniques. DNA from the libraries or the separated DNA may be transferred to and immobilized on nitrocellulose or other suitable carrier material. In order to identify a clone or DNA that is homologous with SEQ ID NO: 1; or a subsequence thereof, the carrier material is preferably used in a Southern blot.

For purposes of the present invention, hybridization indicates that the polynucleotide hybridizes to a labelled nucleic acid probe corresponding to the mature polypeptide coding sequence of SEQ ID NO: 1; its full-length complementary strand; or a subsequence thereof; under very low to very high stringency conditions. Molecules to which the nucleic acid probe hybridizes under these conditions can be detected using, for example, X-ray film.

In one aspect, the nucleic acid probe is the mature polypeptide coding sequence of SEQ ID NO: 1. In another aspect, the nucleic acid probe is a fragment thereof. In another aspect, the nucleic acid probe is a polynucleotide that encodes the polypeptide of SEQ ID NO: 2 or SEQ ID NO: 3 or a fragment thereof. In another preferred aspect, the nucleic acid probe is SEQ ID NO: 1.
For long probes of at least 100 nucleotides in length, high to very high stringency conditions are defined as prehybridization and hybridization at 42°C in 5X SSPE, 0.3% SDS, 200 micrograms/ml sheared and denatured salmon sperm DNA, and 35% formamide, following standard Southern blotting procedures for 12 to 24 hours optimally. The carrier material is finally washed three times each for 15 minutes using 1.5X SSC (very low stringency), 0.8 SSC (low stringency), 0.4X SSC (medium low stringency), 0.2X SSC (medium-high and high stringency) or 0.1X SSC (very high stringency), 0.2% SDS at 65°C (low to medium-high stringency), and at 70°C (high and very high stringency).

For short probes of about 15 nucleotides to about 70 nucleotides in length, stringency conditions are defined as prehybridization and hybridization at about 5°C to about 10°C below the calculated Tm using the calculation according to Bolton and McCarthy (1962, Proc. Natl. Acad. Sci. USA 48:1390) in 0.9 M NaCl, 0.09 M Tris-HCl pH 7.6, 6 mM EDTA, 0.5% NP-40, 1X Denhardt's solution, 1 mM sodium pyrophosphate, 1 mM sodium monobasic phosphate, 0.1 mM ATP, and 0.2 mg of yeast RNA per ml following standard Southern blotting procedures for 12 to 24 hours optimally. The carrier material is finally washed once in 6X SCC plus 0.1% SDS for 15 minutes and twice each for 15 minutes using 6X SSC at 5°C to 10°C below the calculated Tm.

The present invention also relates to the use in animal feed or detergents of isolated polypeptides having protease activity encoded by polynucleotides having a sequence identity to the mature polypeptide coding sequence of SEQ ID NO: 1 of at least 80%, at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100%.

In particular embodiments, the parent proteases and/or the protease variants of the invention and for use according to the invention are selected from the group consisting of:

(a) Proteases belonging to the EC 3.4.21 enzyme group; and


For determining whether a given protease is a serine protease, and a family S1 protease, reference is made to the above Handbook and the principles indicated therein. Such determination can be carried out for all types of proteases, be it naturally occurring or wild-type proteases; or genetically engineered or synthetic proteases.

In a particular embodiment, the present invention also relates to a method for preparing an animal feed or feed additive, comprising preparing an animal feed or feed additive composition comprising an animal feed and a protease of selected from the group consisting of:

(i) a polypeptide of SEQ ID NO: 3;
(ii) a polypeptide having least 80%, e.g. at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or 100% sequence identity with the polypeptide of 3, and which has protease activity.

The present invention also relates to an animal feed or feed additive composition comprising an animal feed and a protease of selected from the group consisting of:

(i) a polypeptide of SEQ ID NO: 3;

(ii) a polypeptide having least 80%, e.g. at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity with the polypeptide of SEQ ID NO: 3, and which has protease activity.

In one aspect, the polypeptides differ by no more thirtytwo amino acids, e.g., by thirty amino acids, by twentyfive amino acids, by twenty amino acids, by fifteen amino acids, by ten amino acids, by eight amino acids, by seven amino acids, by six amino acids, by five amino acids, by four amino acids, by three amino acids, by two amino acids, and by one amino acid from the polypeptide of SEQ ID NO: 3.

The animal feed compositions may in particular embodiments be in the form of a pellet, a mash or liquid composition, as further described herein.

The present invention also relates to variant polypeptides having protease activity and having at least 85%, e.g., at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% sequence identity to SEQ ID NO: 3 comprising at least one substitution, deletion, and/or insertion of at least one or more (several) amino acids of SEQ ID NO: 3 or a homologous sequence thereof.

The variant polypeptide of the invention may in one embodiment have at least 86% sequence identity to SEQ ID NO: 3.

The variant polypeptide of the invention may in one embodiment have at least 87% sequence identity to SEQ ID NO: 3.

The variant polypeptide of the invention may in one embodiment have at least 88% sequence identity to SEQ ID NO: 3.

The variant polypeptide of the invention may in one embodiment have at least 89% sequence identity to SEQ ID NO: 3.

The variant polypeptide of the invention may in one embodiment have at least 90% sequence identity to SEQ ID NO: 3.
The variant polypeptide of the invention may in one embodiment have at least 91% sequence identity to SEQ ID NO: 3.

The variant polypeptide of the invention may in one embodiment have at least 92% sequence identity to SEQ ID NO: 3.

The variant polypeptide of the invention may in one embodiment have at least 93% sequence identity to SEQ ID NO: 3.

The variant polypeptide of the invention may in one embodiment have at least 94% sequence identity to SEQ ID NO: 3.

The variant polypeptide of the invention may in one embodiment have at least 95% sequence identity to SEQ ID NO: 3.

The variant polypeptide of the invention may in one embodiment have at least 96% sequence identity to SEQ ID NO: 3.

The variant polypeptide of the invention may in one embodiment have at least 97% sequence identity to SEQ ID NO: 3.

The variant polypeptide of the invention may in one embodiment have at least 98% sequence identity to SEQ ID NO: 3.

The variant polypeptide of the invention may in one embodiment have at least 99% sequence identity to SEQ ID NO: 3.

In a further embodiment, the total number of positions of the variant polypeptide of the invention (SEQ ID NO: 3) having amino acid substitutions, deletions and/or insertions is not more than 24, e.g., 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23 or 24. The amino acid changes may be of a minor nature, that is conservative amino acid substitutions or insertions that do not significantly affect the folding and/or activity of the protein; small deletions, typically of one to about 30 amino acids; small amino- or carboxyl-terminal extensions, such as an amino-terminal methionine residue; a small linker peptide of up to about 20-25 residues; or a small extension that facilitates purification by changing net charge or another function, such as a poly-histidine tract, an antigenic epitope or a binding domain.

The present invention also relates to variants for use in animal feed or detergents comprising a substitution, deletion, and/or insertion of one or more (or several) amino acids of the mature polypeptide of SEQ ID NO: 2 or a homologous sequence thereof. The total number of positions having amino acid substitutions, deletions and/or insertions in the mature polypeptide of SEQ ID NO: 2 is not more than 32, e.g., 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31 or 32. Preferably, amino acid changes are of a minor nature, that is conservative amino acid substitutions, insertions or deletions that do not significantly affect the folding and/or activity of the protein; small deletions, typically of one to about 30 amino acids; small amino- or carboxyl-terminal extensions, such as an amino-terminal methionine residue; a small linker peptide of up to about
20-25 residues; or a small extension that facilitates purification by changing net charge or another function, such as a poly-histidine tract, an antigenic epitope or a binding domain.

The present invention also relates to variants for use in animal feed or detergents comprising comprising a substitution, deletion, and/or insertion of one or more (or several) amino acids of SEQ ID NO: 3 or a homologous sequence thereof. The total number of positions having amino acid substitutions, deletions and/or insertions in SEQ ID NO: 3 is not more than 32, e.g., 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31 or 32. Preferably, amino acid changes are of a minor nature, that is conservative amino acid substitutions, insertions or deletions that do not significantly affect the folding and/or activity of the protein; small deletions, typically of one to about 30 amino acids; small amino- or carboxyl-terminal extensions, such as an amino-terminal methionine residue; a small linker peptide of up to about 20-25 residues; or a small extension that facilitates purification by changing net charge or another function, such as a poly-histidine tract, an antigenic epitope or a binding domain.

Examples of conservative substitutions are within the group of basic amino acids (arginine, lysine and histidine), acidic amino acids (glutamic acid and aspartic acid), polar amino acids (glutamine and asparagine), hydrophobic amino acids (leucine, isoleucine and valine), aromatic amino acids (phenylalanine, tryptophan and tyrosine), and small amino acids (glycine, alanine, serine, threonine and methionine). Amino acid substitutions that do not generally alter specific activity are known in the art and are described, for example, by H. Neurath and R.L. Hill, 1979, In, The Proteins, Academic Press, New York. The most commonly occurring exchanges that are expected not to alter the specific activity substantially are Ala/Ser, Val/Ile, Asp/Glu, Thr/Ser, Ala/Gly, Ala/Thr, Ser/Asn, Ala/Val, Ser/Gly, Tyr/Phe, Ala/Pro, Lys/Arg, Asp/Asn, Leu/Ile, Leu/AI, Ala/Glu, and Asp/Gly.

Alternatively, the amino acid changes are of such a nature that the physico-chemical properties of the polypeptides are altered. For example, amino acid changes may improve the thermal stability of the polypeptide, alter the substrate specificity, change the pH optimum, and the like. Essential amino acids in a parent polypeptide can be identified according to procedures known in the art, such as site-directed mutagenesis or alanine-scanning mutagenesis (Cunningham and Wells, 1989, Science 244: 1081-1085). In the latter technique, single alanine mutations are introduced at every residue in the molecule, and the resultant mutant molecules are tested for protease activity to identify amino acid residues that are critical to the activity of the molecule. See also, Hilton et al., 1996, J. Biol. Chem. 271 : 4699-4708. The active site of the enzyme or other biological interaction can also be determined by physical analysis of structure, as determined by such techniques as nuclear magnetic resonance, crystallography, electron diffraction, or photoaffinity labelling, in conjunction with mutation of putative contact site amino acids. See, for example, de Vos et al., 1992, Science 255: 306-312; Smith et al., 1992, J. Mol.
Biol. 224: 899-904; Wlodaver et al., 1992, FEBS Lett. 309: 59-64. The identities of essential amino acids can also be inferred from analysis of identities with polypeptides that are related to the parent polypeptide.

Single or multiple amino acid substitutions, deletions, and/or insertions can be made and tested using known methods of mutagenesis, recombination, and/or shuffling, followed by a relevant screening procedure, such as those disclosed by Reidhaar-Olson and Sauer, 1988, Science 241: 53-57; Bowie and Sauer, 1989, Proc. Natl. Acad. Sci. USA 86: 2152-2156; WO 95/17413; or WO 95/22625. Other methods that can be used include error-prone PCR, phage display (e.g., Lowman et al., 1991, Biochemistry 30: 10832-10837; U.S. Patent No. 5,223,409; WO 92/06204), and region-directed mutagenesis (Derbyshire et al., 1986, Gene 46: 145; Ner et al., 1988, DNA 7: 127).

Mutagenesis/shuffling methods can be combined with high-throughput, automated screening methods to detect activity of cloned, mutagenized polypeptides expressed by host cells (Ness et al., 1999, Nature Biotechnology 17: 893-896). Mutagenized DNA molecules that encode active polypeptides can be recovered from the host cells and rapidly sequenced using standard methods in the art. These methods allow the rapid determination of the importance of individual amino acid residues in a polypeptide.

The total number of amino acid substitutions, deletions and/or insertions of the mature polypeptide of SEQ ID NO: 2 is not more than 10, e.g., 1, 2, 3, 4, 5, 6, 7, 8 or 9. The total number of amino acid substitutions, deletions and/or insertions in SEQ ID NO: 3 is not more than 10, e.g., 1, 2, 3, 4, 5, 6, 7, 8 or 9. The polypeptide may be hybrid polypeptide in which a portion of one polypeptide is fused at the N-terminus or the C-terminus of a portion of another polypeptide.

The polypeptide may be a fused polypeptide or cleavable fusion polypeptide in which another polypeptide is fused at the N-terminus or the C-terminus of the polypeptide of the present invention. A fused polypeptide is produced by fusing a polynucleotide encoding another polypeptide to a polynucleotide of the present invention. Techniques for producing fusion polypeptides are known in the art, and include ligating the coding sequences encoding the polypeptides so that they are in frame and that expression of the fused polypeptide is under control of the same promoter(s) and terminator. Fusion proteins may also be constructed using intein technology in which fusions are created post-translationally (Cooper et al., 1993, EMBO J. 12: 2575-2583; Dawson et al., 1994, Science 266: 776-779).

A fusion polypeptide can further comprise a cleavage site between the two polypeptides. Upon secretion of the fusion protein, the site is cleaved releasing the two polypeptides. Examples of cleavage sites include, but are not limited to, the sites disclosed in Martin et al., 2003, J. Ind. Microbiol. Biotechnol. 3: 568-576; Svetina et al., 2000, J. Biotechnol. 76: 245-251; Rasmussen-Wilson et al., 1997, Appl. Environ. Microbiol. 63: 3488-3493; Ward et al., 1995,
Embodiments

In certain embodiments of the invention, the protease of the invention exhibits beneficial thermal properties such as thermostability, steam stability, etc. and/or pH properties, such as acid stability, pH optimum, etc.

An embodiment of the invention is isolated polypeptides having improved protease activity between pH 7 and 9, such as at pH 7.0, pH 8.0 or pH 9.0, at 25°C compared to protease 10R.

A further embodiment of the invention is isolated polypeptides having improved protease activity at e.g. 60°C or below, such as 50°C or below, 37°C or below, or between 25°C and 60°C, or between 37°C and 60°C or at 37°C, or at 50°C or at 60°C at pH 7.0 compared to protease 10R at pH 6.5.

Acidity/alkalinity properties

In certain embodiments of the invention the protease of the invention exhibits beneficial properties in respect of pH, such as acid stability, pH optimum, etc. Stability of the protease at a low pH is beneficial since the protease can have activity in the intestine after passing through the stomach. In one embodiment of the invention the protease retains >95% activity after 2 hours at pH 3 as determined using the method described in Example 3.

Temperature-activity

The temperature-activity profile of the protease may be determined as described in Example 3. Activity at high temperature (e.g. 60°C) could be beneficial to e.g. washing clothes, whereas activity at low temperatures (20-40°C) can be advantageous for low temperature washing or for the digestion of proteins in an animal.

In one embodiment, the invention comprises a protease having a temperature activity profile at pH 7.0 with relative activity of 0.15 or higher at 37°C, relative activity of 0.50 or higher at 50°C, or relative activity of 0.80 or higher at 60°C when compared to the activity of the protease at 70°C (cf. Example 3).

Thermostability

Thermostability may be determined as described in Example 10, i.e. using DSC measurements to determine the denaturation temperature, T_d, of the purified protease protein. The T_d is indicative of the thermostability of the protein: The higher the T_d, the higher the
thermostability. Accordingly, in a preferred embodiment, the protease of the invention has a \( T_d \) which is higher than the \( T_d \) of a reference protease, wherein \( T_d \) is determined on purified protease samples (preferably with a purity of at least 90% or 95%, as determined by SDS-PAGE).

In preferred embodiments, the thermal properties such as heat-stability, temperature stability, thermostability, steam stability, and/or pelleting stability as provided by the residual activity, denaturation temperature \( T_d \), or other parameter of the protease of the invention is higher than the corresponding value, such as the residual activity or \( T_d \), of the protease of SEQ ID NO: 3, more preferably at least 101% thereof, or at least 102%, 103%, 104%, 105%, 106%, 107%, 108%, 109%, or at least 110% thereof. Even more preferably, the value of the parameter, such as residual activity or \( T_d \), of the protease of the invention is at least 120%, 130%, 140%, 150%, 160%, 170%, 180%, or at least 190% of the value for the protease of SEQ ID NO: 3.

In still further particular embodiments, the thermostable protease of the invention has a melting temperature, \( T_m \) (or a denaturation temperature, \( T_d \), as determined using Differential Scanning Calorimetry (DSC) as described in example 10 (i.e. in 20 mM sodium acetate, pH 4.0), of at least 50°C. In still further particular embodiments, the \( T_m \) is at least 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99 or at least 100°C.

Steam stability

Steam stability may be determined as described in Example 11 by determining the residual activity of protease molecules after steam treatment at 85°C or 90°C for a short time.

Pelleting stability

Pelleting stability may be determined as described in Example 12 by using enzyme granulate pre-mixed with feed. From the mixer the feed is conditioned with steam to 95°C. After conditioning the feed is pressed to pellets and the residual activity determined.

Sources of Polypeptides Having Protease Activity

A polypeptide having protease activity and to be used according to the present invention may be obtained from microorganisms of any genus. For purposes of the present invention, the term "obtained from" as used herein in connection with a given source shall mean that the polypeptide encoded by a polynucleotide is produced by the source or by a strain in which the polynucleotide from the source has been inserted. In one aspect, the polypeptide obtained from a given source is secreted extracellularly.
The polypeptide may be a bacterial polypeptide. For example, the polypeptide may be a polypeptide having protease activity from a gram-positive bacterium within a phylum such as *Actinobacteria* or from a gram-negative bacterium within a phylum such as *Proteobacteria*.

In one aspect, the polypeptide is a protease from a bacterium of the class *Actinobacteria*, such as from the order *Actinomycetales*, or from the suborder *Propionibacterineae*, or from the family *Nocardioidaceae*, or from the genera *Kribbella*. In another aspect, the polypeptide is a protease from the suborder *Pseudonocardineae*, or from the family *Pseudonocardiaeae*, or from the genera *Saccharomonospora*, *Saccharopolyspora*; or *Amycolatopsis*.

Strains of these taxa are readily accessible to the public in a number of culture collections, such as the American Type Culture Collection (ATCC), Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH (DSM), Centraalbureau Voor Schimmelcultures (CBS), and Agricultural Research Service Patent Culture Collection, Northern Regional Research Center (NRRL).

The polypeptide may be identified and obtained from other sources including microorganisms isolated from nature (e.g., soil, composts, water, etc.) using the above-mentioned probes. Techniques for isolating microorganisms from natural habitats are well known in the art. The polynucleotide encoding the polypeptide may then be obtained by similarly screening a genomic or cDNA library of another microorganism or mixed DNA sample. Once a polynucleotide encoding a polypeptide has been detected with the probe(s), the polynucleotide can be isolated or cloned by utilizing techniques that are well known to those of ordinary skill in the art (see, e.g., Sambrook *et al.*, 1989, *supra*).

**Polynucleotides**

The present invention also relates to isolated polynucleotides encoding a polypeptide of the present invention and used for recombinant production of the polypeptide.

The techniques used to isolate or clone a polynucleotide encoding a polypeptide are known in the art and include isolation from genomic DNA, preparation from cDNA, or a combination thereof. The cloning of the polynucleotides from such genomic DNA can be effected, e.g., by using the well known polymerase chain reaction (PCR) or antibody screening of expression libraries to detect cloned DNA fragments with shared structural features. See, e.g., Innis *et al.*, 1990, *PCR: A Guide to Methods and Application*, Academic Press, New York. Other nucleic acid amplification procedures such as ligase chain reaction (LCR), ligation activated transcription (LAT) and polynucleotide-based amplification (NASBA) may be used. The polynucleotides may be cloned from a strain of *Saccharopolyspora*, or another related organism from the *Actinomycetales* and thus, for example, may be an allelic or species variant of the polypeptide encoding region of the polynucleotide.
The present invention also relates to isolated polynucleotides comprising or consisting of polynucleotides having a degree of sequence identity to the mature polypeptide coding sequence of SEQ ID NO: 1 of at least 80%, e.g., at least 85%, e.g., at least 87%, at least 89%, at least 90%, at least 93%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100%, with the proviso that it is not 100% identical to the mature polypeptide coding sequence of SEQ ID NO: 1, and which encode a polypeptide having protease activity.

Modification of a polynucleotide encoding a polypeptide of the present invention may be necessary for the synthesis of polypeptides substantially similar to the polypeptide. The term “substantially similar” to the polypeptide refers to non-naturally occurring forms of the polypeptide. These polypeptides may differ in some engineered way from the polypeptide isolated from its native source, e.g., variants that differ in specific activity, thermostability, pH optimum, or the like. The variant may be constructed on the basis of the polynucleotide presented as the mature polypeptide coding sequence of SEQ ID NO: 1, e.g., a subsequence thereof, and/or by introduction of nucleotide substitutions that do not result in a change in the amino acid sequence of the polypeptide, but which correspond to the codon usage of the host organism intended for production of the enzyme, or by introduction of nucleotide substitutions that may give rise to a different amino acid sequence. For a general description of nucleotide substitution, see, e.g., Ford et al., 1991, Protein Expression and Purification 2: 95-107.

The present invention also relates to isolated polynucleotides encoding polypeptides of the present invention, which hybridize under very low stringency conditions, low stringency conditions, medium stringency conditions, medium-high stringency conditions, high stringency conditions, or very high stringency conditions with (i) the mature polypeptide coding sequence of SEQ ID NO: 1, (ii) the genomic DNA sequence comprising the mature polypeptide coding sequence of SEQ ID NO: 1, or (iii) the full-length complementary strand of (i) or (ii); or allelic variants and subsequences thereof (Sambrook et al., 1989, supra), as defined herein.

In one aspect, the polynucleotide comprises or consists of SEQ ID NO: 1, the mature polypeptide coding sequence of SEQ ID NO: 1, or a subsequence of SEQ ID NO: 1 that encodes a fragment of SEQ ID NO: 2 having protease activity, such as the polynucleotide of nucleotides 595-1074 of SEQ ID NO: 1.

Nucleic Acid Constructs

The present invention also relates to nucleic acid constructs comprising a polynucleotide of the present invention operably linked to one or more (several) control sequences that direct the expression of the coding sequence in a suitable host cell under conditions compatible with the control sequences.

A polynucleotide may be manipulated in a variety of ways to provide for expression of the polypeptide. Manipulation of the polynucleotide prior to its insertion into a vector may be
desirable or necessary depending on the expression vector. The techniques for modifying polynucleotides utilizing recombinant DNA methods are well known in the art.

The control sequence may be a promoter sequence, a polynucleotide that is recognized by a host cell for expression of a polynucleotide encoding a polypeptide of the present invention. The promoter sequence contains transcriptional control sequences that mediate the expression of the polypeptide. The promoter may be any polynucleotide that shows transcriptional activity in the host cell of choice including mutant, truncated, and hybrid promoters, and may be obtained from genes encoding extracellular or intracellular polypeptides either homologous or heterologous to the host cell.

Examples of suitable promoters for directing the transcription of the nucleic acid constructs of the present invention in a bacterial host cell are the promoters obtained from the Bacillus amyloliquefaciens alpha-amylase gene (amyQ), Bacillus licheniformis alpha-amylase gene (amyL), Bacillus licheniformis penicillinase gene (penP), Bacillus stearothermophilus maltogenic amylase gene (amyM), Bacillus subtilis levansucrase gene (sacB), Bacillus subtilis xylA and xylB genes, E. coli lac operon, Streptomyces coelicolor agarase gene (dagA), and prokaryotic beta-lactamase gene (Villa-Kamaroff et al., 1978, Proc. Natl. Acad. Sci. USA 75: 3727-3731), as well as the tac promoter (DeBoer et al., 1983, Proc. Natl. Acad. Sci. USA 80: 21-25). Further promoters are described in "Useful proteins from recombinant bacteria" in Gilbert et al., 1980, Scientific American, 242: 74-94; and in Sambrook et al., 1989, supra.

Examples of suitable promoters for directing the transcription of the nucleic acid constructs of the present invention in a filamentous fungal host cell are promoters obtained from the genes for Aspergillus nidulans acetamidase, Aspergillus niger neutral alpha-amylase, Aspergillus niger acid stable alpha-amylase, Aspergillus niger or Aspergillus awamori glucoamylase (glaA), Aspergillus oryzae TAKA amylase, Aspergillus oryzae alkaline protease, Aspergillus oryzae triose phosphate isomerase, Fusarium oxysporum trypsin-like protease (WO 96/00787), Fusarium venenatum amyloglucosidase (WO 00/56900), Fusarium venenatum Daria (WO 00/56900), Fusarium venenatum Quinn (WO 00/56900), Rhizomucor miehei lipase, Rhizomucor miehei aspartic proteinase, Trichoderma reesei beta-glucosidase, Trichoderma reesei cellobiohydrolase I, Trichoderma reesei cellobiohydrolase II, Trichoderma reesei endoglucanase I, Trichoderma reesei endoglucanase II, Trichoderma reesei endoglucanase III, Trichoderma reesei endoglucanase IV, Trichoderma reesei endoglucanase V, Trichoderma reesei xylanase I, Trichoderma reesei xylanase II, Trichoderma reesei beta-xylodidase, as well as the NA2-tpi promoter (a modified promoter including a gene encoding a neutral alpha-amylase in Aspergillus in which the untranslated leader has been replaced by an untranslated leader from a gene encoding triose phosphate isomerase in Aspergillus; non-limiting examples include modified promoters including the gene encoding neutral alpha-amylase in Aspergillus niger in which the untranslated leader has been replaced by an untranslated leader from the
gene encoding triose phosphate isomerase in *Aspergillus nidulans* or *Aspergillus oryzae*); and mutant, truncated, and hybrid promoters thereof.

In a yeast host, useful promoters are obtained from the genes for *Saccharomyces cerevisiae* enolase (ENO-1), *Saccharomyces cerevisiae* galactokinase (GAL1), *Saccharomyces cerevisiae* alcohol dehydrogenase/glyceraldehyde-3-phosphate dehydrogenase (ADH1, ADH2/GAP), *Saccharomyces cerevisiae* triose phosphate isomerase (TPI), *Saccharomyces cerevisiae* metallothionein (CUP1), and *Saccharomyces cerevisiae* 3-phosphoglycerate kinase. Other useful promoters for yeast host cells are described by Romanos *et al.*, 1992, *Yeast* 8: 423-488.

The control sequence may also be a suitable transcription terminator sequence, which is recognized by a host cell to terminate transcription. The terminator sequence is operably linked to the 3'-terminus of the polynucleotide encoding the polypeptide. Any terminator that is functional in the host cell of choice may be used in the present invention.

Preferred terminators for filamentous fungal host cells are obtained from the genes for *Aspergillus nidulans* anthranilate synthase, *Aspergillus niger* glucoamylase, *Aspergillus niger* alpha-glucosidase, *Aspergillus oryzae* TAKA amylase, and *Fusarium oxysporum* trypsin-like protease.

Preferred terminators for yeast host cells are obtained from the genes for *Saccharomyces cerevisiae* enolase, *Saccharomyces cerevisiae* cytochrome C (CYC1), and *Saccharomyces cerevisiae* glyceraldehyde-3-phosphate dehydrogenase. Other useful terminators for yeast host cells are described by Romanos *et al.*, 1992, *supra*.

The control sequence may also be a suitable leader sequence, when transcribed is a nontranslated region of an mRNA that is important for translation by the host cell. The leader sequence is operably linked to the 5'-terminus of the polynucleotide encoding the polypeptide. Any leader sequence that is functional in the host cell of choice may be used.

Preferred leaders for filamentous fungal host cells are obtained from the genes for *Aspergillus oryzae* TAKA amylase and *Aspergillus nidulans* triose phosphate isomerase.

Suitable leaders for yeast host cells are obtained from the genes for *Saccharomyces cerevisiae* enolase (ENO-1), *Saccharomyces cerevisiae* 3-phosphoglycerate kinase, *Saccharomyces cerevisiae* alpha-factor, and *Saccharomyces cerevisiae* alcohol dehydrogenase/glyceraldehyde-3-phosphate dehydrogenase (ADH2/GAP).

The control sequence may also be a polyadenylation sequence, a sequence operably linked to the 3'-terminus of the polynucleotide and, when transcribed, is recognized by the host cell as a signal to add polyadenosine residues to transcribed mRNA. Any polyadenylation sequence that is functional in the host cell of choice may be used.

Preferred polyadenylation sequences for filamentous fungal host cells are obtained from the genes for *Aspergillus oryzae* TAKA amylase, *Aspergillus niger* glucoamylase, *Aspergillus*
nidulans anthranilate synthase, Fusarium oxysporum trypsin-like protease, and Aspergillus niger alpha-glucosidase.

Useful polyadenylation sequences for yeast host cells are described by Guo and Sherman, 1995, Mol. Cellular Biol. 15: 5983-5990.

The control sequence may also be a signal peptide coding region that encodes a signal peptide linked to the N-terminus of a polypeptide and directs the polypeptide into the cell's secretory pathway. The 5'-end of the coding sequence of the polynucleotide may inherently contain a signal peptide coding sequence naturally linked in translation reading frame with the segment of the coding sequence that encodes the polypeptide. Alternatively, the 5'-end of the coding sequence may contain a signal peptide coding sequence that is foreign to the coding sequence. The foreign signal peptide coding sequence may be required where the coding sequence does not naturally contain a signal peptide coding sequence. Alternatively, the foreign signal peptide coding sequence may simply replace the natural signal peptide coding sequence in order to enhance secretion of the polypeptide. However, any signal peptide coding sequence that directs the expressed polypeptide into the secretory pathway of a host cell of choice may be used.

Effective signal peptide coding sequences for bacterial host cells are the signal peptide coding sequences obtained from the genes for Bacillus NCIB 11837 maltogenic amylase, Bacillus licheniformis subtilisin, Bacillus licheniformis beta-lactamase, Bacillus stearothermophilus alpha-amylase, Bacillus stearothermophilus neutral proteases {nprT, nprS, nprM}, Bacillus Clausii subtilisin, and Bacillus subtilis prsA. Further signal peptides are described by Simonen and Palva, 1993, Microbiological Reviews 57: 109-137.

Effective signal peptide coding sequences for filamentous fungal host cells are the signal peptide coding sequences obtained from the genes for Aspergillus niger neutral amylase, Aspergillus niger glucoamylase, Aspergillus oryzae TAKA amylase, Humicola insolens cellulase, Humicola insolens endoglucanase V, Humicola lanuginosa lipase, and Rhizomucor miehei aspartic proteinase.

Useful signal peptides for yeast host cells are obtained from the genes for Saccharomyces cerevisiae alpha-factor and Saccharomyces cerevisiae invertase. Other useful signal peptide coding sequences are described by Romanos et al., 1992, supra.

The control sequence may also be a propeptide coding sequence that encodes a propeptide positioned at the N-terminus of a polypeptide. The resultant polypeptide is known as a proenzyme or propolypeptide (or a zymogen in some cases). A propolypeptide is generally inactive and can be converted to an active polypeptide by catalytic or autocatalytic cleavage of the propeptide from the propolypeptide. The propeptide coding sequence may be obtained from the genes for Bacillus subtilis alkaline protease {aprE}, Bacillus subtilis neutral protease {nprT},
Myceliophthora thermophila laccase (WO 95/33836), Rhizomucor miehei aspartic proteinase, and Saccharomyces cerevisiae alpha-factor.

Where both signal peptide and propeptide sequences are present at the N-terminus of a polypeptide, the propeptide sequence is positioned next to the N-terminus of a polypeptide and the signal peptide sequence is positioned next to the N-terminus of the propeptide sequence.

It may also be desirable to add regulatory sequences that allow the regulation of the expression of the polypeptide relative to the growth of the host cell. Examples of regulatory systems are those that cause the expression of the gene to be turned on or off in response to a chemical or physical stimulus, including the presence of a regulatory compound. Regulatory systems in prokaryotic systems include the lac, tac, and trp operator systems. In yeast, the ADH2 system or GAL1 system may be used. In filamentous fungi, the Aspergillus niger glucoamylase promoter, Aspergillus oryzae TAKA alpha-amylase promoter, and Aspergillus oryzae glucoamylase promoter may be used. Other examples of regulatory sequences are those that allow for gene amplification. In eukaryotic systems, these regulatory sequences include the dihydrofolate reductase gene that is amplified in the presence of methotrexate, and the metallothionein genes that are amplified with heavy metals. In these cases, the polynucleotide encoding the polypeptide would be operably linked with the regulatory sequence.

Expression Vectors

The present invention also relates to recombinant expression vectors comprising a polynucleotide of the present invention, a promoter, and transcriptional and translational stop signals. The various nucleotide and control sequences may be joined together to produce a recombinant expression vector that may include one or more (several) convenient restriction sites to allow for insertion or substitution of the polynucleotide encoding the polypeptide at such sites. Alternatively, the polynucleotide may be expressed by inserting the polynucleotide or a nucleic acid construct comprising the sequence into an appropriate vector for expression. In creating the expression vector, the coding sequence is located in the vector so that the coding sequence is operably linked with the appropriate control sequences for expression.

The recombinant expression vector may be any vector (e.g., a plasmid or virus) that can be conveniently subjected to recombinant DNA procedures and can bring about expression of the polynucleotide. The choice of the vector will typically depend on the compatibility of the vector with the host cell into which the vector is to be introduced. The vector may be a linear or closed circular plasmid.

The vector may be an autonomously replicating vector, i.e., a vector that exists as an extrachromosomal entity, the replication of which is independent of chromosomal replication, e.g., a plasmid, an extrachromosomal element, a minichromosome, or an artificial chromosome. The vector may contain any means for assuring self-replication. Alternatively, the vector may be
one that, when introduced into the host cell, is integrated into the genome and replicated together with the chromosome(s) into which it has been integrated. Furthermore, a single vector or plasmid or two or more vectors or plasmids that together contain the total DNA to be introduced into the genome of the host cell, or a transposon, may be used.

The vector preferably contains one or more (several) selectable markers that permit easy selection of transformed, transfected, transduced, or the like cells. A selectable marker is a gene the product of which provides for biocide or viral resistance, resistance to heavy metals, prototrophy to auxotrophs, and the like.

Examples of bacterial selectable markers are the dal genes from *Bacillus subtilis* or *Bacillus licheniformis*, or markers that confer antibiotic resistance such as ampicillin, chloramphenicol, kanamycin, or tetracycline resistance. Suitable markers for yeast host cells are ADE2, HIS3, LEU2, LYS2, MET3, TRP1, and URA3. Selectable markers for use in a filamentous fungal host cell include, but are not limited to, *amdS* (acetamidase), *argB* (ornithine carbamoyltransferase), *bar* (phosphinothricin acetyltransferase), *hph* (hygromycin phosphotransferase), *niaD* (nitrate reductase), *pyrG* (orotidine-5'-phosphate decarboxylase), *sC* (sulfate adenyltransferase), and *trpC* (anthranilate synthase), as well as equivalents thereof.

Preferred for use in an *Aspergillus* cell are the *amdS* and *pyrG* genes of *Aspergillus nidulans* or *Aspergillus oryzae* and the *bar* gene of *Streptomyces hygroscopicus*.

The vector preferably contains an element(s) that permits integration of the vector into the host cell's genome or autonomous replication of the vector in the cell independent of the genome.

For integration into the host cell genome, the vector may rely on the polynucleotide's sequence encoding the polypeptide or any other element of the vector for integration into the genome by homologous or non-homologous recombination. Alternatively, the vector may contain additional polynucleotides for directing integration by homologous recombination into the genome of the host cell at a precise location(s) in the chromosome(s). To increase the likelihood of integration at a precise location, the integrational elements should contain a sufficient number of nucleic acids, such as 100 to 10,000 base pairs, 400 to 10,000 base pairs, and 800 to 10,000 base pairs, which have a high degree of sequence identity to the corresponding target sequence to enhance the probability of homologous recombination. The integrational elements may be any sequence that is homologous with the target sequence in the genome of the host cell. Furthermore, the integrational elements may be non-encoding or encoding polynucleotides. On the other hand, the vector may be integrated into the genome of the host cell by non-homologous recombination.

For autonomous replication, the vector may further comprise an origin of replication enabling the vector to replicate autonomously in the host cell in question. The origin of replication may be any plasmid replicator mediating autonomous replication that functions in a
cell. The term "origin of replication" or "plasmid replicator" means a polynucleotide that enables a plasmid or vector to replicate in vivo.

Examples of bacterial origins of replication are the origins of replication of plasmids pBR322, pUC19, pACYC177, and pACYC184 permitting replication in E. coli, and pUB110, pE194, pTA1060, and pAMβI permitting replication in Bacillus.

Examples of origins of replication for use in a yeast host cell are the 2 micron origin of replication, ARS1, ARS4, the combination of ARS1 and CEN3, and the combination of ARS4 and CEN6.

Examples of origins of replication useful in a filamentous fungal cell are AMA1 and ANSI (Gems et al., 1991, Gene 98: 61-67; Cullen et al., 1987, Nucleic Acids Res. 15: 9163-9175; WO 00/24883). Isolation of the AMA1 gene and construction of plasmids or vectors comprising the gene can be accomplished according to the methods disclosed in WO 00/24883.

More than one copy of a polynucleotide of the present invention may be inserted into a host cell to increase production of a polypeptide. An increase in the copy number of the polynucleotide can be obtained by integrating at least one additional copy of the sequence into the host cell genome or by including an amplifiable selectable marker gene with the polynucleotide where cells containing amplified copies of the selectable marker gene, and thereby additional copies of the polynucleotide, can be selected for by cultivating the cells in the presence of the appropriate selectable agent.

The procedures used to ligate the elements described above to construct the recombinant expression vectors of the present invention are well known to one skilled in the art (see, e.g., Sambrook et al., 1989, supra).

**Host Cells**

The present invention also relates to recombinant host cells, comprising a polynucleotide of the present invention operably linked to one or more (several) control sequences that direct the production of a polypeptide of the present invention. A construct or vector comprising a polynucleotide is introduced into a host cell so that the construct or vector is maintained as a chromosomal integrant or as a self-replicating extra-chromosomal vector as described earlier. The term "host cell" encompasses any progeny of a parent cell that is not identical to the parent cell due to mutations that occur during replication. The choice of a host cell will to a large extent depend upon the gene encoding the polypeptide and its source.

The host cell may be any cell useful in the recombinant production of a polypeptide of the present invention, e.g., a prokaryote or a eukaryote.

The prokaryotic host cell may be any gram-positive or gram-negative bacterium. Gram-positive bacteria include, but are not limited to, *Bacillus, Brevibacillus, Clostridium, Geobacillus*, 35
Lactobacillus, Lactococcus, Paenibacillus, and Streptomyces. Gram-negative bacteria include, but are not limited to E. coli, and Pseudomonas.

The bacterial host cell may be any Bacillus cell including, but not limited to, Bacillus alkalophilus, Bacillus amyloliquefaciens, Bacillus brevis, Bacillus circulans, Bacillus clausii, Bacillus coagulans, Bacillus firmus, Bacillus licheniformis, Bacillus megaterium, Bacillus pumilus, Bacillus stearothermophilus, Bacillus subtilis, and Bacillus thuringiensis cells. Specifically preferred host cells are Bacillus subtilis and Bacillus licheniformis cells.

The bacterial host cell may also be any Streptococcus cell including, but not limited to, Streptococcus equisimilis, Streptococcus pyogenes, Streptococcus uberis, and Streptococcus equi subsp. Zooepidemicus cells.

The bacterial host cell may also be any Streptomyces cell including, but not limited to, Streptomyces achrromogenes, Streptomyces avermitilis, Streptomyces coelicolor, Streptomyces griseus, and Streptomyces lividans cells.

The introduction of DNA into a Bacillus cell may, for instance, be effected by protoplast transformation (see, e.g., Chang and Cohen, 1979, Mol. Gen. Genet. 168: 111-15), by using competent cells (see, e.g., Young and Spizizen, 1961, J. Bacteriol. 81: 823-829, or Dubnau and Davidoff-Abelson, 1971, J. Mol. Biol. 56: 209-221), by electroporation (see, e.g., Shigekawa and Dower, 1988, Biotechniques 6: 742-751), or by conjugation (see, e.g., Koehler and Thorne, 1987, J. Bacteriol. 169: 5271-5278). The introduction of DNA into an E. coli cell may, for instance, be effected by protoplast transformation (see, e.g., Hanahan, 1983, J. Mol. Biol. 166: 557-580) or electroporation (see, e.g., Dower et al., 1988, Nucleic Acids Res. 16: 6127-6145).

The introduction of DNA into a Streptomyces cell may, for instance, be effected by protoplast transformation and electroporation (see, e.g., Gong et al., 2004, Folia Microbiol. (Praha) 49: 399-405), by conjugation (see, e.g., Mazodier et al., 1989, J. Bacteriol. 171: 3583-3585), or by transduction (see, e.g., Burke et al., 2001, Proc. Natl. Acad. Sci. USA 98: 6289-6294). The introduction of DNA into a Pseudomonas cell may, for instance, be effected by electroporation (see, e.g., Choi et al., 2006, J. Microbiol. Methods 64: 391-397) or by conjugation (see, e.g., Pinedo and Smets, 2005, Appl. Environ. Microbiol. 71: 51-57). The introduction of DNA into a Streptococcus cell may, for instance, be effected by natural competence (see, e.g., Perry and Kuramitsu, 1981, Infect. Immun. 32: 1295-1297), by protoplast transformation (see, e.g., Catt and Jollick, 1991, Microbiol 68: 189-207), by electroporation (see, e.g., Buckley et al., 1999, Appl. Environ. Microbiol. 65: 3800-3804) or by conjugation (see, e.g., Clewell, 1981, Microbiol. Rev. 45: 409-436). However, any method known in the art for introducing DNA into a host cell can be used.

The host cell may also be a eukaryote, such as a mammalian, insect, plant, or fungal cell.
The host cell may be a fungal cell. "Fungi" as used herein includes the phyla Ascomycota, Basidiomycota, Chytridiomycota, and Zygomycota (as defined by Hawksworth et al., in Ainsworth and Bisby's Dictionary of The Fungi, 8th edition, 1995, CAB International, University Press, Cambridge, UK) as well as the Oomycota (as cited in Hawksworth et al., 1995, supra, page 171) and all mitosporic fungi (Hawksworth et al., 1995, supra).

The fungal host cell may be a yeast cell. "Yeast" as used herein includes ascosporogenous yeast (Endomycetales), basidiosporogenous yeast, and yeast belonging to the Fungi Imperfecti (Blastomycetes). Since the classification of yeast may change in the future, for the purposes of this invention, yeast shall be defined as described in Biology and Activities of Yeast (Skinner, F.A., Passmore, S.M., and Davenport, R.R., eds, Soc. App. Bacteriol. Symposium Series No. 9, 1980).

The yeast host cell may be a Candida, Hansenula, Kluyveromyces, Pichia, Saccharomyces, Schizosaccharomyces, or Yarrowia cell such as a Kluyveromyces lactis, Saccharomyces carlsbergensis, Saccharomyces cerevisiae, Saccharomyces diastaticus, Saccharomyces douglasii, Saccharomyces kluveri, Saccharomyces norbensis, Saccharomyces oviformis, or Yarrowia lipolytica cell.

The fungal host cell may be a filamentous fungal cell. "Filamentous fungi" include all filamentous forms of the subdivision Eumycota and Oomycota (as defined by Hawksworth et al., 1995, supra). The filamentous fungi are generally characterized by a mycelial wall composed of chitin, cellulose, glucan, chitosan, mannan, and other complex polysaccharides. Vegetative growth is by hyphal elongation and carbon catabolism is obligately aerobic. In contrast, vegetative growth by yeasts such as Saccharomyces cerevisiae is by budding of a unicellular thallus and carbon catabolism may be fermentative.

The filamentous fungal host cell may be an Acremonium, Aspergillus, Aureobasidium, Bjerkandera, Ceriporiopsis, Chrysosporium, Coprinus, Coriolus, Cryptococcus, Filibasidium, Fusarium, Humincola, Magnaporthe, Mucor, Myceliophthora, Neocallimastix, Neurospora, Paecilomyces, Penicillium, Phanerochaete, Phlebia, Piromyces, Pleurotus, Schizophyllum, Talaromyces, Thermoascus, Thielavia, Tolypocladium, Trametes, or Trichoderma cell.

For example, the filamentous fungal host cell may be an Aspergillus awamori, Aspergillus foetidus, Aspergillus fumigatus, Aspergillus japonicus, Aspergillus nidulans, Aspergillus niger, Aspergillus oryzae, Bjerkandera adusta, Cerioporiopsis aneirina, Ceriporiopsis caregea, Ceriporiopsis gilvescens, Ceriporiopsis pannocinta, Ceriporiopsis rivulosa, Ceriporiopsis sub Rufa, Ceriporiopsis subvermispora, Chrysosporium inops, Chrysosporium keratinophilum, Chrysosporium lucknowense, Chrysosporium merdarium, Chrysosporium pannicola, Chrysosporium queenslandicum, Chrysosporium tropicum, Chrysosporium zonatum, Coprinus cinereus, Coriolus hirsutus, Fusarium bactridioides, Fusarium cerealis, Fusarium crookwellense, Fusarium culmorum, Fusarium graminearum, Fusarium graminum, Fusarium
heterosporum, Fusarium negundi, Fusarium oxysporum, Fusarium reticulatum, Fusarium roseum, Fusarium sambucinum, Fusarium sarcochroum, Fusarium sporotrichioides, Fusarium sulphureum, Fusarium torulosum, Fusarium trichothecioides, Fusarium venenatum, Humicola insolens, Humicola lanuginosa, Mucor miehei, Myceliophthora thermophila, Neurospora crassa, Penicillium purpurogenum, Phanerochaete chrysosporium, Phlebia radiata, Pleurotus eryngii, Thielavia terrestris, Trametes villosa, Trametes versicolor, Trichoderma harzianum, Trichoderma koningii, Trichoderma longibrachiatum, Trichoderma reesei, or Trichoderma viride cell.


Methods of Production

The present invention also relates to methods of producing a polypeptide of the present invention, comprising: (a) cultivating a cell, which in its wild-type form produces the polypeptide, under conditions conducive for production of the polypeptide; and (b) recovering the polypeptide. In one aspect, the cell is of the genus Saccharomonospora. In a more preferred aspect, the cell is a Saccharomonospora viridis cell.

The present invention also relates to methods of producing a polypeptide of the present invention, comprising: (a) cultivating a recombinant host cell of the present invention under conditions conducive for production of the polypeptide; and (b) recovering the polypeptide.

The host cells are cultivated in a nutrient medium suitable for production of the polypeptide using methods well known in the art. For example, the cell may be cultivated by shake flask cultivation, and small-scale or large-scale fermentation (including continuous, batch, fed-batch, or solid state fermentations) in laboratory or industrial fermentors performed in a suitable medium and under conditions allowing the polypeptide to be expressed and/or isolated. The cultivation takes place in a suitable nutrient medium comprising carbon and nitrogen sources and inorganic salts, using procedures known in the art. Suitable media are available from commercial suppliers or may be prepared according to published compositions (e.g., in catalogues of the American Type Culture Collection). If the polypeptide is secreted into the
nutrient medium, the polypeptide can be recovered directly from the medium. If the polypeptide is not secreted, it can be recovered from cell lysates.

More details are provided in the Sections above and in the Section on "Nucleic Acid Constructs, Expression Vectors, Recombinant Host Cells, and Methods for Production of Proteases" below.

The polypeptide may be detected using methods known in the art that are specific for the polypeptides. These detection methods may include use of specific antibodies, formation of an enzyme product, or disappearance of an enzyme substrate. For example, an enzyme assay may be used to determine the activity of the polypeptide.

The polypeptide may be recovered using methods known in the art. For example, the polypeptide may be recovered from the nutrient medium by conventional procedures including, but not limited to, centrifugation, filtration, extraction, spray-drying, evaporation, or precipitation.

The polypeptide may be purified by a variety of procedures known in the art including, but not limited to, chromatography (e.g., ion exchange, affinity, hydrophobic, chromatofocusing, and size exclusion), electrophoretic procedures (e.g., preparative isoelectric focusing), differential solubility (e.g., ammonium sulfate precipitation), SDS-PAGE, or extraction (see, e.g., Protein Purification, J.-C. Janson and Lars Ryden, editors, VCH Publishers, New York, 1989) to obtain substantially pure polypeptides.

In an alternative aspect, the polypeptide is not recovered, but rather a host cell of the present invention expressing a polypeptide is used as a source of the polypeptide.

Plants

The present invention also relates to plants, e.g., a transgenic plant, plant part, or plant cell, comprising an isolated polynucleotide of the present invention so as to express and produce the polypeptide in recoverable quantities. The polypeptide may be recovered from the plant or plant part. Alternatively, the plant or plant part containing the polypeptide may be used as such for improving the quality of a food or feed, e.g., improving nutritional value, palatability, and rheological properties, or to destroy an antinutritive factor.

The transgenic plant can be dicotyledonous (a dicot) or monocotyledonous (a monocot). Examples of monocot plants are grasses, such as meadow grass (blue grass, Poa), forage grass such as Festuca, Lolium, temperate grass, such as Agrostis, and cereals, e.g., wheat, oats, rye, barley, rice, sorghum, and maize (corn).

Examples of dicot plants are tobacco, legumes, such as lupins, potato, sugar beet, pea, bean and soybean, and cruciferous plants (family Brassicaceae), such as cauliflower, rape seed, and the closely related model organism Arabidopsis thaliana.

Examples of plant parts are stem, callus, leaves, root, fruits, seeds, and tubers as well as the individual tissues comprising these parts, e.g., epidermis, mesophyll, parenchyme,
vascular tissues, meristems. Specific plant cell compartments, such as chloroplasts, apoplasts, mitochondria, vacuoles, peroxisomes and cytoplasm are also considered to be a plant part. Furthermore, any plant cell, whatever the tissue origin, is considered to be a plant part. Likewise, plant parts such as specific tissues and cells isolated to facilitate the utilization of the invention are also considered plant parts, e.g., embryos, endosperms, aleurone and seeds coats.

Also included within the scope of the present invention are the progeny of such plants, plant parts, and plant cells.

The transgenic plant or plant cell expressing a polypeptide may be constructed in accordance with methods known in the art. In short, the plant or plant cell is constructed by incorporating one or more (several) expression constructs encoding a polypeptide into the plant host genome or chloroplast genome and propagating the resulting modified plant or plant cell into a transgenic plant or plant cell.

The expression construct is conveniently a nucleic acid construct that comprises a polynucleotide encoding a polypeptide operably linked with appropriate regulatory sequences required for expression of the polynucleotide in the plant or plant part of choice. Furthermore, the expression construct may comprise a selectable marker useful for identifying host cells into which the expression construct has been integrated and DNA sequences necessary for introduction of the construct into the plant in question (the latter depends on the DNA introduction method to be used).

The choice of regulatory sequences, such as promoter and terminator sequences and optionally signal or transit sequences, is determined, for example, on the basis of when, where, and how the polypeptide is desired to be expressed. For instance, the expression of the gene encoding a polypeptide may be constitutive or inducible, or may be developmental, stage or tissue specific, and the gene product may be targeted to a specific tissue or plant part such as seeds or leaves. Regulatory sequences are, for example, described by Tague et al., 1988, Plant Physiology 86: 506.

For constitutive expression, the 35S-CaMV, the maize ubiquitin 1, and the rice actin 1 promoter may be used (Franck et al., 1980, Cell 21: 285-294; Christensen et al., 1992, Plant Mol. Biol. 18: 675-689; Zhang et al., 1991, Plant Cell 3: 1155-1165). Organ-specific promoters may be, for example, a promoter from storage sink tissues such as seeds, potato tubers, and fruits (Edwards and Coruzzi, 1990, Ann. Rev. Genet. 24: 275-303), or from metabolic sink tissues such as meristems (Ito et al., 1994, Plant Mol. Biol. 24: 863-878), a seed specific promoter such as the glutelin, prolamin, globulin, or albumin promoter from rice (Wu et al., 1998, Plant Cell Physiol. 39: 885-889), a Vicia faba promoter from the legumin B4 and the unknown seed protein gene from Vicia faba (Conrad et al., 1998, J. Plant Physiol. 152: 708-711), a promoter from a seed oil body protein (Chen et al., 1998, Plant Cell Physiol. 39: 935-941), the
storage protein napA promoter from Brassica napus, or any other seed specific promoter known in the art, e.g., as described in WO 91/14772. Furthermore, the promoter may be a leaf specific promoter such as the rbcS promoter from rice or tomato (Kyozuka et al., 1993, Plant Physiol. 102: 991-1000), the chlorella virus adenine methyltransferase gene promoter (Mitra and Higgins, 1994, Plant Mol. Biol. 26: 85-93), the aldP gene promoter from rice (Kagaya et al., 1995, Mol. Gen. Genet. 248: 668-674), or a wound inducible promoter such as the potato pin2 promoter (Xu et al., 1993, Plant Mol. Biol. 22: 573-588). Likewise, the promoter may be inducible by abiotic treatments such as temperature, drought, or alterations in salinity or induced by exogenously applied substances that activate the promoter, e.g., ethanol, oestrogens, plant hormones such as ethylene, abscisic acid, and gibberellic acid, and heavy metals.

A promoter enhancer element may also be used to achieve higher expression of a polypeptide in the plant. For instance, the promoter enhancer element may be an intron that is placed between the promoter and the polynucleotide encoding a polypeptide. For instance, Xu et al., 1993, supra, disclose the use of the first intron of the rice actin 1 gene to enhance expression.

The selectable marker gene and any other parts of the expression construct may be chosen from those available in the art.

The nucleic acid construct is incorporated into the plant genome according to conventional techniques known in the art, including Agrobacterium-mediated transformation, virus-mediated transformation, microinjection, particle bombardment, biolistic transformation, and electroporation (Gasser et al., 1990, Science 244: 1293; Potrykus, 1990, Bio/Technology 8: 535; Shimamoto et al., 1989, Nature 338: 274).

Presently, Agrobacterium tumefaciens-mediated gene transfer is the method of choice for generating transgenic dicots (for a review, see Hooykaas and Schilperoort, 1992, Plant Mol. Biol. 19: 15-38) and can also be used for transforming monocots, although other transformation methods are often used for these plants. Presently, the method of choice for generating transgenic monocots is particle bombardment (microscopic gold or tungsten particles coated with the transforming DNA) of embryonic calli or developing embryos (Christou, 1992, Plant J. 2: 275-281; Shimamoto, 1994, Curr. Opin. Biotechnol. 5: 158-162; Vasil et al., 1992, Bio/Technology 10: 667-674). An alternative method for transformation of monocots is based on protoplast transformation as described by Omirulleh et al., 1993, Plant Mol. Biol. 21: 415-428. Additional transformation methods for use in accordance with the present disclosure include those described in U.S. Patent Nos. 6,395,966 and 7,151,204 (both of which are herein incorporated by reference in their entirety).

Following transformation, the transformants having incorporated the expression construct are selected and regenerated into whole plants according to methods well known in the art. Often the transformation procedure is designed for the selective elimination of selection.
genes either during regeneration or in the following generations by using, for example, co-
transformation with two separate T-DNA constructs or site specific excision of the selection
gene by a specific recombinase.

In addition to direct transformation of a particular plant genotype with a construct
prepared according to the present invention, transgenic plants may be made by crossing a plant
having the construct to a second plant lacking the construct. For example, a construct encoding
a polypeptide can be introduced into a particular plant variety by crossing, without the need for
ever directly transforming a plant of that given variety. Therefore, the present invention
encompasses not only a plant directly regenerated from cells which have been transformed in
accordance with the present invention, but also the progeny of such plants. As used herein,
progeny may refer to the offspring of any generation of a parent plant prepared in accordance
with the present invention. Such progeny may include a DNA construct prepared in accordance
with the present invention, or a portion of a DNA construct prepared in accordance with the
present invention. Crossing results in the introduction of a transgene into a plant line by cross
pollinating a starting line with a donor plant line. Non-limiting examples of such steps are further
articulated in U.S. Patent NO: 7,151,204.

Plants may be generated through a process of backcross conversion. For example, plants include plants referred to as a backcross converted genotype, line, inbred, or hybrid.

Genetic markers may be used to assist in the introgression of one or more transgenes of
the invention from one genetic background into another. Marker assisted selection offers
advantages relative to conventional breeding in that it can be used to avoid errors caused by
phenotypic variations. Further, genetic markers may provide data regarding the relative degree
of elite germplasm in the individual progeny of a particular cross. For example, when a plant
with a desired trait which otherwise has a non-agronomically desirable genetic background is
crossed to an elite parent, genetic markers may be used to select progeny which not only
possess the trait of interest, but also have a relatively large proportion of the desired
germplasm. In this way, the number of generations required to introgress one or more traits into
a particular genetic background is minimized.

The present invention also relates to methods of producing a polypeptide of the present
invention comprising: (a) cultivating a transgenic plant or a plant cell comprising a
polynucleotide encoding the polypeptide under conditions conducive for production of the
polypeptide; and (b) recovering the polypeptide.

**Compositions**

The present invention also relates to compositions comprising a protease of the present
invention. Preferably, the compositions are enriched in such a protease. The term "enriched"
indicates that the protease activity of the composition has been increased, e.g., with an enrichment factor of at least 1.1.

In one aspect, the composition comprises an isolated polypeptide having protease activity, selected from the group consisting of:

(a) a polypeptide having at least 80%, e.g., at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or 100% sequence identity to SEQ ID NO: 3;

(b) a polypeptide encoded by a polynucleotide that hybridizes under medium stringency conditions, medium-high stringency conditions, high stringency conditions or very-high stringency conditions with:

(i) the mature polypeptide coding sequence of SEQ ID NO: 1; and/or

(iii) the full-length complementary strand of (i);

(c) a polypeptide encoded by a polynucleotide having at least 80%, e.g., at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or 100% sequence identity to the mature polypeptide coding sequence of SEQ ID NO: 3;

(d) a variant comprising a substitution, deletion, and/or insertion of one or more (several) amino acids of SEQ ID NO: 3; and

(e) a fragment of a polypeptide of (a), (b), (c) or (d), that has protease activity.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 85% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 86% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 87% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 88% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 89% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 90% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 91% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 92% sequence identity to the polypeptide of SEQ ID NO: 3.
An embodiment of the invention is a composition comprising an isolated polypeptide having at least 93% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 94% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 95% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 96% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 97% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 98% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having at least 99% sequence identity to the polypeptide of SEQ ID NO: 3.

An embodiment of the invention is a composition comprising an isolated polypeptide having 100% sequence identity to the polypeptide of SEQ ID NO: 3.

In one aspect, the composition comprises or consists of the amino acid sequence of SEQ ID NO: 3 or an allelic variant thereof; or is a fragment thereof having protease activity. In another aspect, the composition comprises or consists of the mature polypeptide of SEQ ID NO: 2. In a further aspect, the composition comprises or consists of the polypeptide of SEQ ID NO: 3. In another aspect, the composition comprises or consists of amino acids 1 to 160 of SEQ ID NO: 2, amino acids 5 to 154 of SEQ ID NO: 2, or amino acids 10 to 149 of SEQ ID NO: 2. In another aspect, the polypeptide comprises or consists of amino acids 1 to 160 of SEQ ID NO: 3, amino acids 5 to 154 of SEQ ID NO: 3, or amino acids 10 to 149 of SEQ ID NO: 3.

In an embodiment, the variant comprising a substitution, deletion, and/or insertion of one or more (several) amino acids of SEQ ID NO: 3 has at least 80%, e.g., at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, but less than 100% sequence identity to SEQ ID NO: 3.

The present invention also relates to compositions comprising isolated polypeptides having protease activity that are encoded by polynucleotides that hybridize under medium stringency conditions, medium-high stringency conditions, high stringency conditions or very-high stringency conditions with (i) the mature polypeptide coding sequence of SEQ ID NO: 1, and/or (ii) the full-length complementary strand of (i) (J. Sambrook, E.F. Fritsch, and T. Maniatis, 1989, Molecular Cloning, A Laboratory Manual, 2d edition, Cold Spring Harbor, New York).
The present invention further relates compositions comprising isolated polypeptides that
differ by no more than thirtytwo amino acids, e.g., by thirty amino acids, by twentyfive amino
acids, by twenty amino acids, by fifteen amino acids, by ten amino acids, by eight amino acids,
by seven amino acids, by six amino acids, by five amino acids, by four amino acids, by three
amino acids, by two amino acids, and by one amino acid from the polypeptide of SEQ ID NO: 3.

The present invention also relates to compositions comprising variants
comprising a substitution, deletion, and/or insertion of one or more (or several) amino acids of
SEQ ID NO: 3 or a homologous sequence thereof. The total number of positions having amino
acid substitutions, deletions and/or insertions in SEQ ID NO: 3 is not more than 32, e.g., 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31 or 32. Preferably, amino acid changes are of a minor nature, that is conservative amino acid
substitutions, insertions or deletions that do not significantly affect the folding and/or activity of
the protein; small deletions, typically of one to about 30 amino acids; small amino- or carboxyl-
terminal extensions, such as an amino-terminal methionine residue; a small linker peptide of up
to about 20-25 residues; or a small extension that facilitates purification by changing net charge
or another function, such as a poly-histidine tract, an antigenic epitope or a binding domain.

In a preferred embodiment, the composition is an animal feed composition or additive,
comprising at least one fat soluble vitamin. In another preferred embodiment, the composition is
an animal feed composition or additive, comprising at least one water soluble vitamin. In a
further preferred embodiment, the composition is an animal feed composition or additive,
comprising at least one trace mineral. In another embodiment, the animal feed composition
comprises one or more further enzymes, wherein the further enzymes are selected from the
group comprising of amylases; phytases; xylanases; galactanases; alpha-galactosidases;
proteases, phospholipases; and beta-glucanases, or any mixture thereof. In another
embodiment, the animal feed additive comprises one or more further enzymes, wherein the
further enzymes are selected from the group comprising of amylases; phytases; xylanases;
galactanases; alpha-galactosidases; proteases, phospholipases; and beta-glucanases, or any
mixture thereof.

In another preferred embodiment, the composition is a detergent composition which may
be used in laundry, laundering, hard surface cleaning and/or dishwash. In an embodiment, the
detergent compositon comprises one or more detergent components as defined herein. In another embodiment, the detergent composition comprises one or more additional enzymes
selected from the group comprising of proteases, amylases, lipases, cutinases, cellulases,
endoglucanases, xyloglucanases, pectinases, pectin lyases, xanthanases, peroxidaes,
haloperoxxygenases, catalases and mannanases, or any mixture thereof. In a further
embodiment, the detergent compositon comprises one or more detergent components as
defined herein and one or more additional enzymes selected from the group comprising of

46
The composition may comprise a protease of the present invention as the major enzymatic component, e.g., a mono-component composition. Alternatively, the composition may comprise multiple enzymatic activities, such as an aminopeptidase, amylase, carboxydrase, carboxypeptidase, catalase, cellulase, chitinase, cutinase, cyclodextrin glycosyltransferase, deoxyribonuclease, esterase, alpha-galactosidase, beta-galactosidase, glucoamylase, alpha-glucosidase, beta-glucosidase, haloperoxidase, invertase, laccase, lipase, mannosidase, oxidase, pectinolytic enzyme, peptidoglaminase, peroxidase, phytase, polyphenoloxidase, proteolytic enzyme, ribonuclease, transglaminase, or xylanase. The additional enzyme(s) may be produced, for example, by a microorganism such as bacteria or fungi or by plants or by animals. The compositions may be prepared in accordance with methods known in the art and may be in the form of a liquid or a dry composition. For instance, the composition may be in the form of a granulate or a microgranulate. The protease may be stabilized in accordance with methods known in the art.

**Detergent Compositions**

In one embodiment, the invention is directed to detergent compositions comprising an enzyme of the present invention in combination with one or more detergent components. The choice of detergent components is within the skill of the artisan and includes conventional ingredients, including the exemplary non-limiting components set forth below.

The choice of components may include, for textile care, the consideration of the type of textile to be cleaned, the type and/or degree of soiling, the temperature at which cleaning is to take place, and the formulation of the detergent product. Although components mentioned below are categorized by general header according to a particular functionality, this is not to be construed as a limitation, as a component may comprise additional functionalities as will be appreciated by the skilled artisan.

The cleaning process or the textile care process may for example be a laundry process, a dishwashing process or cleaning of hard surfaces such as bathroom tiles, floors, table tops, drains, sinks and washbasins. Laundry processes can for example be household laundering, but it may also be industrial laundering. Furthermore, the invention relates to a process for laundering of fabrics and/or garments where the process comprises treating fabrics with a washing solution containing a detergent composition, and at least one protease of the invention. The cleaning process or a textile care process can for example be carried out in a machine washing process or in a manual washing process. The washing solution can for example be an aqueous washing solution containing a detergent composition.
The fabrics and/or garments subjected to a washing, cleaning or textile care process of the present invention may be conventional washable laundry, for example household laundry. Preferably, the major part of the laundry is garments and fabrics, including knits, woven, denims, non-woven, felts, yarns, and towelling. The fabrics may be cellulose based such as natural cellulosics, including cotton, flax, linen, jute, ramie, sisal or coir or manmade cellulosics (e.g., originating from wood pulp) including viscose/rayon, ramie, cellulose acetate fibers (tricell), lyocell or blends thereof. The fabrics may also be non-cellulose based such as natural polyamides including wool, camel, cashmere, mohair, rabbit and silk or synthetic polymer such as nylon, aramid, polyester, acrylic, polypropylene and spandex/elastane, or blends thereof as well as blend of cellulose based and non-cellulose based fibers. Examples of blends are blends of cotton and/or rayon/viscose with one or more companion material such as wool, synthetic fibers (e.g., polyamide fibers, acrylic fibers, polyester fibers, polyvinyl alcohol fibers, polyvinyl chloride fibers, polyurethane fibers, polyurea fibers, aramid fibers), and cellulose-containing fibers (e.g., rayon/viscose, ramie, flax, linen, jute, cellulose acetate fibers, lyocell).

The last few years there has been an increasing interest in replacing components in detergents, which is derived from petrochemicals with renewable biological components such as enzymes and polypeptides without compromising the wash performance. When the components of detergent compositions change new enzyme activities or new enzymes having alternative and/or improved properties compared to the common used detergent enzymes such as proteases, lipases and amylases is needed to achieve a similar or improved wash performance when compared to the traditional detergent compositions.

The invention further concerns the use of proteases of the invention in a proteinaceous stain removing processes. The proteinaceous stains may be stains such as food stains, e.g., baby food, sebum, cocoa, egg, blood, milk, ink, grass, or a combination hereof.

Typical detergent compositions includes various components in addition to the enzymes, these components have different effects, some components like the surfactants lower the surface tension in the detergent, which allows the stain being cleaned to be lifted and dispersed and then washed away, other components like bleach systems removes discoulour often by oxidation and many bleaches also have strong bactericidal properties, and are used for disinfecting and sterilizing. Yet other components like builder and chelator softens, e.g., the wash water by removing the metal ions form the liquid.

In a particular embodiment, the invention concerns the use of a composition comprising a protease of the invention, wherein said enzyme composition further comprises at least one or more of the following a surfactant, a builder, a chelator or chelating agent, bleach system or bleach component in laundry or dish wash.

In one embodiment of the present invention, the a polypeptide of the present invention may be added to a detergent composition in an amount corresponding to 0.001-200 mg of

48
protein, such as 0.005-100 mg of protein, preferably 0.01-50 mg of protein, more preferably 0.05-20 mg of protein, even more preferably 0.1-10 mg of protein per litre of wash liquor.

A composition for use in automatic dishwash (ADW), for example, may include 0.0001%-50%, such as 0.001-%-20%, such as 0.01%-10%, such as 0.05-5% of enzyme protein by weight of the composition.

A composition for use in laundry granulation, for example, may include 0.0001 %-50%, such as 0.001 %-20%, such as 0.01%-10%, such as 0.05%-5% of enzyme protein by weight of the composition.

A composition for use in laundry liquid, for example, may include 0.0001 %-10%, such as 0.001-7%, such as 0.1%-5% of enzyme protein by weight of the composition.

The enzyme(s) of the detergent composition of the invention may be stabilized using conventional stabilizing agents, e.g., a polyol such as propylene glycol or glycerol, a sugar or sugar alcohol, lactic acid, boric acid, or a boric acid derivative, e.g., an aromatic borate ester, or a phenyl boronic acid derivative such as 4-formylphenyl boronic acid, and the composition may be formulated as described in, for example, W092/1 9709 and W092/1 9708.

In certain markets different wash conditions and, as such, different types of detergents are used. This is disclosed in e.g. EP 1 025 240. For example, in Asia (Japan) a low detergent concentration system is used, while the United States uses a medium detergent concentration system, and Europe uses a high detergent concentration system.

A low detergent concentration system includes detergents where less than about 800 ppm of detergent components are present in the wash water. Japanese detergents are typically considered low detergent concentration system as they have approximately 667 ppm of detergent components present in the wash water.

A medium detergent concentration includes detergents where between about 800 ppm and about 2000ppm of detergent components are present in the wash water. North American detergents are generally considered to be medium detergent concentration systems as they have approximately 975 ppm of detergent components present in the wash water.

A high detergent concentration system includes detergents where greater than about 2000 ppm of detergent components are present in the wash water. European detergents are generally considered to be high detergent concentration systems as they have approximately 4500-5000 ppm of detergent components in the wash water.

Latin American detergents are generally high suds phosphate builder detergents and the range of detergents used in Latin America can fall in both the medium and high detergent concentrations as they range from 1500 ppm to 6000 ppm of detergent components in the wash water. Such detergent compositions are all embodiments of the invention.

A polypeptide of the present invention may also be incorporated in the detergent formulations disclosed in WO97/07202, which is hereby incorporated by reference.
Surfactants

The detergent composition may comprise one or more surfactants, which may be anionic and/or cationic and/or non-ionic and/or semi-polar and/or zwitterionic, or a mixture thereof. In a particular embodiment, the detergent composition includes a mixture of one or more nonionic surfactants and one or more anionic surfactants. The surfactant(s) is typically present at a level of from about 0.1% to 60% by weight, such as about 1% to about 40%, or about 3% to about 20%, or about 3% to about 10%. The surfactant(s) is chosen based on the desired cleaning application, and includes any conventional surfactant(s) known in the art. Any surfactant known in the art for use in detergents may be utilized.

When included therein the detergent will usually contain from about 1% to about 40% by weight, such as from about 5% to about 30%, including from about 5% to about 15%, or from about 20% to about 25% of an anionic surfactant. Non-limiting examples of anionic surfactants include sulfates and sulfonates, in particular, linear alkylbenzenesulfonates (LAS), isomers of LAS, branched alkylbenzenesulfonates (BABS), phenylalkanesulfonates, alpha-olefinsulfonates (AOS), olefin sulfonates, alkene sulfonates, alkane-2,3-diylibis(sulfates), hydroxyalkanesulfonates and disulfonates, alkyl sulfates (AS) such as sodium dodecyl sulfate (SDS), fatty alcohol sulfates (FAS), primary alcohol sulfates (PAS), alcohol ethersulfates (AES or AEOS or FES, also known as alcohol ethoxysulfates or fatty alcohol ether sulfates), secondary alkanesulfonates (SAS), paraffin sulfonates (PS), ester sulfonates, sulfonated fatty acid glycerol esters, alpha-sulfo fatty acid methyl esters (alpha-SFMe or SES) including methyl ester sulfonate (MES), alkyl- or alkenylsuccinic acid, dodecenyl/tetradecenyl succinic acid (DTSA), fatty acid derivatives of amino acids, diesters and monoesters of sulfo-succinic acid or soap, and combinations thereof.

When included therein the detergent will usually contain from about 0% to about 10% by weight of a cationic surfactant. Non-limiting examples of cationic surfactants include alkyldimethylethanolamine quat (ADMEAQ), cetyltrimethylammonium bromide (CTAB), dimethyldistearylammonium chloride (DSDMAC), and alkylbenzyl(dimethylammonium, alkyl quaternary ammonium compounds, alkoxylated quaternary ammonium (AQA) compounds, and combinations thereof.

When included therein the detergent will usually contain from about 0.2% to about 40% by weight of a non-ionic surfactant, for example from about 0.5% to about 30%, in particular from about 1% to about 20%, from about 3% to about 10%, such as from about 3% to about 5%, or from about 8% to about 12%. Non-limiting examples of non-ionic surfactants include alcohol ethoxylates (AE or AEO), alcohol propoxylates, propoxylated fatty alcohols (PFA), alkoxylated fatty acid alkyl esters, such as ethoxylated and/or propoxylated fatty acid alkyl esters, alkylyphenol ethoxylates (APE), nonylphenol ethoxylates (NPE), alkylpolyglycosides (APG), alkoxylated amines, fatty acid monoethanolamides (FAM), fatty acid diethanolamides (FADA), ethoxylated fatty acid monoethanolamides (EFAM), propoxylated fatty acid monoethanolamides (PFAM), polyhydroxy alkyl fatty acid amides, or /V-acyl /V-alkyl derivatives
of glucosamine (glucamides, GA, or fatty acid glucamide, FAGA), as well as products available under the trade names SPAN and TWEEN, and combinations thereof.

When included therein the detergent will usually contain from about 0% to about 10% by weight of a semipolar surfactant. Non-limiting examples of semipolar surfactants include amine oxides (AO) such as alkylidimethylamineoxide, /V-(coco alkyl)/V./V.-dimethylamine oxide and /V-(tallow-alkyl)/V./V.-bis(2-hydroxyethyl)amine oxide, fatty acid alkanolamides and ethoxylated fatty acid alkanolamides, and combinations thereof.

When included therein the detergent will usually contain from about 0% to about 10% by weight of a zwitterionic surfactant. Non-limiting examples of zwitterionic surfactants include betaine, alkylidimethylbetaine, sulfobetaine, and combinations thereof.

Hydrotropes

A hydrotrope is a compound that solubilises hydrophobic compounds in aqueous solutions (or oppositely, polar substances in a non-polar environment). Typically, hydrotropes have both hydrophilic and a hydrophobic character (so-called amphiphilic properties as known from surfactants); however the molecular structure of hydrotropes generally do not favor spontaneous self-aggregation, see e.g. review by Hodgdon and Kaler (2007), Current Opinion in Colloid & Interface Science 12: 121-128. Hydrotropes do not display a critical concentration above which self-aggregation occurs as found for surfactants and lipids forming miceller, lamellar or other well defined meso-phases. Instead, many hydrotropes show a continuous-type aggregation process where the sizes of aggregates grow as concentration increases. However, many hydrotropes alter the phase behavior, stability, and colloidal properties of systems containing substances of polar and non-polar character, including mixtures of water, oil, surfactants, and polymers. Hydrotropes are classically used across industries from pharma, personal care, food, to technical applications. Use of hydrotropes in detergent compositions allow for example more concentrated formulations of surfactants (as in the process of compacting liquid detergents by removing water) without inducing undesired phenomena such as phase separation or high viscosity.

The detergent may contain 0-5% by weight, such as about 0.5 to about 5%, or about 3% to about 5%, of a hydrotrope. Any hydrotrope known in the art for use in detergents may be utilized. Non-limiting examples of hydrotropes include sodium benzene sulfonate, sodium p-toluene sulfonate (STS), sodium xylene sulfonate (SXS), sodium cumene sulfonate (SCS), sodium cymene sulfonate, amine oxides, alcohols and polyglycolethers, sodium hydroxynaphthoate, sodium hydroxynaphthalene sulfonate, sodium ethylhexyl sulfate, and combinations thereof.

Builders and Co-Builders

The detergent composition may contain about 0-65% by weight, such as about 5% to about 45% of a detergent builder or co-builder, or a mixture thereof. In a dish wash detergent,
the level of builder is typically 40-65%, particularly 50-65%. The builder and/or co-builder may particularly be a chelating agent that forms water-soluble complexes with Ca and Mg. Any builder and/or co-builder known in the art for use in laundry detergents may be utilized. Non-limiting examples of builders include zeolites, diophosphates (pyrophosphates), triphosphates such as sodium tripolyphosphate (STP or STPP), carbonates such as sodium carbonate, soluble silicates such as sodium metasilicate, layered silicates (e.g., SKS-6 from Hoechst), ethanolamines such as 2-aminoethan-1-ol (MEA), diethanolamine (DEA, also known as iminodiethanol), triethanolamine (TEA, also known as 2,2',2''-nitrilotriethanol), and carboxymethyl inulin (CMI), and combinations thereof.

The detergent composition may also contain 0-20% by weight, such as about 5% to 10%, of a detergent co-builder, or a mixture thereof. The detergent composition may include include a co-builder alone, or in combination with a builder, for example a zeolite builder. Non-limiting examples of co-builders include homopolymers of polyacrylates or copolymers thereof, such as poly(acrylic acid) (PAA) or copoly(acrylic acid/maleic acid) (PAA/PMA). Further non-limiting examples include citrate, chelators such as aminocarboxylates, aminopolycarboxylates and phosphonates, and alkyl- or alkenylsuccinic acid. Additional specific examples include 2,2',2''-nitrilotriacetic acid (NTA), ethylenediaminetetraacetic acid (EDTA), diethylenetriaminepentaacetic acid (DTPA), iminodisuccinic acid (IDS), ethylenediamine-N,N'-diacetic acid (EDDA), methylglycinediacetic acid (MGDA), glutamic acid-N,N-diacetic acid (GLDA), 1-hydroxyethane-1,1-diphosphonic acid (HEDP), ethylenediaminetetra-(methyleneephosphonic acid) (EDTMPA), diethylenetriaminepentakis(methyleneephosphonic acid) (DTPMPA or DTMPA), N-(2-hydroxyethyl)iminodiacetic acid (EDG), aspartic acid-N,N'-diacetic acid (ASDA), aspartic acid-N,N'-diacetic acid (ASMA), aspartic acid-N,N'-diacetic acid (ASDP), aspartic acid-N,N'-diacetic acid (ASMP), iminodisuccinic acid (IDA), N-(2-sulfomethyl)-aspartic acid (SMAS), N-(2-sulfomethyl)-aspartic acid (SEAS), N-(2-sulfomethyl)-N,N'-glutamic acid (SMGL), N-(2-sulfomethyl)-N,N'-glutamic acid (SEG), N,N'-methyleneiminodiacetic acid (MIDA), a-alanine-N,N'-diacetic acid (a-ALDA), serine-N,N'-diacetic acid (SEDA), isoserine-N,N'-diacetic acid (ISDA), phenylalanine-N,N'-diacetic acid (PHDA), anthranilic acid-N,N'-diacetic acid (ANDA), sulfanilic acid-N,N'-diacetic acid (SLDA), taurine-N,N'-diacetic acid (TUDA) and sulfomethyl-N,N'-diacetic acid (SMDA), N-(2-hydroxyethyl)-ethyldenediamine-N,N'-diacetic acid (HEDTA), diethanolglycin (DEG), diethylenetriamine penta(methyleneephosphonic acid) (DTPMP), aminotris(methyleneephosphonic acid) (ATMP), and combinations and salts thereof. Further exemplary builders and/or co-builders are described in, e.g., WO 09/102854, US 5977053

Bleaching Systems

The detergent may contain 0-50% by weight, such as about 0.1% to about 25%, of a bleaching system. Any bleaching system known in the art for use in laundry detergents may be utilized. Suitable bleaching system components include bleaching catalysts, photobleaches, bleach activators, sources of hydrogen peroxide such as sodium percarbonate and sodium
perborates, preformed peracids and mixtures thereof. Suitable preformed peracids include, but
are not limited to, peroxycarboxylic acids and salts, percarbonic acids and salts, perimidic acids
and salts, peroxymonosulfuric acids and salts, for example, Oxone (R), and mixtures thereof.
Non-limiting examples of bleaching systems include peroxide-based bleaching systems, which
may comprise, for example, an inorganic salt, including alkali metal salts such as sodium salts
of perborate (usually mono- or tetra-hydrate), percarbonate, persulfate, perphosphate,
persilicate salts, in combination with a peracid-forming bleach activator. The term bleach
activator is meant herein as a compound which reacts with peroxygen bleach like hydrogen
peroxide to form a peracid. The peracid thus formed constitutes the activated bleach. Suitable
bleach activators to be used herein include those belonging to the class of esters amides,
imides or anhydrides. Suitable examples are tetracetylene diamine (TAED), sodium 4-
[(3,5,5-trimethylhexanoyl)oxy]benzene sulfonate (ISONOBS), diperoxy dodecanoic acid, 4-
(dodecanoyloxy)benzenesulfonate (LOBS), 4-(decanoyloxy)benzenesulfonate, 4-
(decanoyloxy)benzoate (DOBS), 4-(nonanoyloxy)-benzenesulfonate (NOBS), and/or those
disclosed in W098/17767. A particular family of bleach activators of interest was disclosed in
EP624154 and particularly preferred in that family is acetyl triethyl citrate (ATC). ATC or a short
chain triglyceride like triacetin has the advantage that it is environmental friendly as it eventually
degradates into citric acid and alcohol. Furthermore acetyl triethyl citrate and triacetin has a good
hydrolytical stability in the product upon storage and it is an efficient bleach activator. Finally
ATC provides a good building capacity to the laundry additive. Alternatively, the bleaching
system may comprise peroxyacids of, for example, the amide, imide, or sulfone type. The
bleaching system may also comprise peracids such as 6-(phthalimido)peroxyhexanoic acid
(PAP). The bleaching system may also include a bleach catalyst. In some embodiments the
bleach component may be an organic catalyst selected from the group consisting of organic
catalysts having the following formulae:

(i)

(ii)

(iii) and mixtures thereof; wherein each R^1 is independently a branched alkyl group
containing from 9 to 24 carbons or linear alkyl group containing from 11 to 24 carbons,
preferably each R^1 is independently a branched alkyl group containing from 9 to 18 carbons or
linear alkyl group containing from 11 to 18 carbons, more preferably each R^1 is independently
selected from the group consisting of 2-propylethyl, 2-butyloctyl, 2-pentynonyl, 2-hexyldecyl,
n-dodecyl, n-tetradecyl, n-hexadecyl, n-octadecyl, iso-nonyl, iso-decyl, iso-tridecyl and iso-
pentadecyl. Other exemplary bleaching systems are described, e.g. in WO2007/087258,
example be sulfonated zinc phthalocyanine.
Polymers

The detergent may contain 0-10% by weight, such as 0.5-5%, 2-5%, 0.5-2% or 0.2-1% of a polymer. Any polymer known in the art for use in detergents may be utilized. The polymer may function as a co-builder as mentioned above, or may provide antiredeposition, fiber protection, soil release, dye transfer inhibition, grease cleaning and/or anti-foaming properties. Some polymers may have more than one of the above-mentioned properties and/or more than one of the below-mentioned motifs. Exemplary polymers include (carboxymethyl)cellulose (CMC), polyvinyl alcohol (PVA), poly(vinylpyrrolidone) (PVP), poly(ethyleneglycol) or poly(ethylene oxide) (PEG), ethoxylated poly(ethyleneimine), carboxymethyl inulin (CMI), and polycarboxylates such as PAA, PAA/PMA, poly-aspartic acid, and lauryl methacrylate/acrylic acid copolymers, hydrophobically modified CMC (HM-CMC) and silicones, copolymers of terephthalic acid and oligomeric glycols, copolymers of poly(ethylene terephthalate) and poly(oxyethene terephthalate) (PET-POET), PVP, poly(vinylimidazole) (PVI), poly(vinylpyridine-/V-oxide) (PVPO or PVPNV) and polyvinylpyrrolidone-vinylimidazole (PVPVI). Further exemplary polymers include sulfonated polycarboxylates, polyethylene oxide and polypropylene oxide (PEO-PPO) and diquaternium ethoxy sulfate. Other exemplary polymers are disclosed in, e.g., WO 2006/130575. Salts of the above-mentioned polymers are also contemplated.

Fabric hueing agents

The detergent compositions of the present invention may also include fabric hueing agents such as dyes or pigments, which when formulated in detergent compositions can deposit onto a fabric when said fabric is contacted with a wash liquor comprising said detergent compositions and thus altering the tint of said fabric through absorption/reflection of visible light. Fluorescent whitening agents emit at least some visible light. In contrast, fabric hueing agents alter the tint of a surface as they absorb at least a portion of the visible light spectrum. Suitable fabric hueing agents include dyes and dye-clay conjugates, and may also include pigments. Suitable dyes include small molecule dyes and polymeric dyes. Suitable small molecule dyes include small molecule dyes selected from the group consisting of dyes falling into the Colour Index (C.I.) classifications of Direct Blue, Direct Red, Direct Violet, Acid Blue, Acid Red, Acid Violet, Basic Blue, Basic Violet and Basic Red, or mixtures thereof, for example as described in WO2005/03274, WO2005/03275, WO2005/03276 and EP1876226 (hereby incorporated by reference). The detergent composition preferably comprises from about 0.00003 wt% to about 0.2 wt%, from about 0.00008 wt% to about 0.05 wt%, or even from about 0.0001 wt% to about 0.04 wt% fabric hueing agent. The composition may comprise from 0.0001 wt% to 0.2 wt% fabric hueing agent, this may be especially preferred when the composition is in the form of a unit dose pouch. Suitable hueing agents are also disclosed in, e.g. WO 2007/087257 and WO2007/087243.
Additional Enzymes

The detergent additive as well as the detergent composition may comprise one or more [additional] enzymes such as a protease, lipase, cutinase, an amylase, carbohydrase, cellulase, pectinase, mannanase, arabinase, galactanase, xylanase, oxidase, e.g., a laccase, and/or peroxidase.

In general the properties of the selected enzyme(s) should be compatible with the selected detergent, (i.e., pH-optimum, compatibility with other enzymatic and non-enzymatic ingredients, etc.), and the enzyme(s) should be present in effective amounts.

Cellulases: Suitable cellulases include those of bacterial or fungal origin. Chemically modified or protein engineered mutants are included. Suitable cellulases include cellulases from the genera Bacillus, Pseudomonas, Humicola, Fusarium, Thielavia, Acremonium, e.g., the fungal cellulases produced from Humicola insolens, Myceliophthora thermophila and Fusarium oxysporum disclosed in US 4,435,307, US 5,648,263, US 5,691,178, US 5,776,757 and WO 89/09259.

Especially suitable cellulases are the alkaline or neutral cellulases having color care benefits. Examples of such cellulases are cellulases described in EP 0 495 257, EP 0 531 372, WO 96/1 1262, WO 96/29397, WO 98/08940. Other examples are cellulase variants such as those described in WO 94/07998, EP 0 531 315, US 5,457,046, US 5,686,593, US 5,763,254, WO 95/24471, WO 98/12307 and PCT/DK98/00299.

Example of cellulases exhibiting endo-beta-1,4-glucanase activity (EC 3.2.1.4) are those having described in WO02/099091.

Other examples of cellulases include the family 45 cellulases described in W096/29397, and especially variants thereof having substitution, insertion and/or deletion at one or more of the positions corresponding to the following positions in SEQ ID NO: 8 of WO 02/099091 : 2, 4, 7, 8, 10, 13, 15, 19, 20, 21, 25, 26, 29, 32, 33, 34, 35, 37, 40, 42, 42a, 43, 44, 48, 53, 54, 55, 58, 59, 63, 64, 65, 66, 67, 70, 72, 76, 79, 80, 82, 84, 86, 88, 90, 91, 93, 95, 95d, 95h, 95j, 97, 100, 101, 102, 103, 113, 114, 117, 119, 121, 133, 136, 137, 138, 139, 140a, 141, 143a, 145, 146, 147, 150e, 150j, 151, 152, 153, 154, 155, 156, 157, 158, 159, 160c, 160e, 160k, 161, 162, 164, 165, 168, 170, 171, 172, 173, 175, 176, 178, 181, 183, 184, 185, 186, 188, 191, 192, 195, 196, 200, and/or 20, preferably selected among P19A, G20K, Q44K, N48E, Q119H or Q146 R.

Commercially available cellulases include Celluzyme™, and Carezyme™ (Novozymes A/S), Clazinase™, and Puradax HA™ (Genencor International Inc.), and KAC-500(B)™ (Kao Corporation).

Proteases: Suitable proteases include those of bacterial, fungal, plant, viral or animal origin e.g. vegetable or microbial origin. Microbial origin is preferred. Chemically modified or protein engineered mutants are included. It may be an alkaline protease, such as a serine protease or a metalloprotease. A serine protease may for example be of the S1 family, such as
trypsin, or the S8 family such as subtilisin. A metalloproteases protease may for example be a thermolysin from e.g. family M4 or other metalloprotease such as those from M5, M7 or M8 families.

The term "subtilases" refers to a sub-group of serine protease according to Siezen et al., Protein Engng. 4 (1991) 719-737 and Siezen et al. Protein Science 6 (1997) 501-523. Serine proteases are a subgroup of proteases characterized by having a serine in the active site, which forms a covalent adduct with the substrate. The subtilases may be divided into 6 sub-divisions, i.e. the Subtilisin family, the Thermitase family, the Proteinase K family, the Lantibiotic peptidase family, the Kexin family and the Pyrolysin family.

Examples of subtilases are those derived from Bacillus such as Bacillus lentus, B. alkalophilus, B. subtilis, B. amyloliquefaciens, Bacillus pumilus and Bacillus gibsonii described in; US7262042 and WO09/021867, and subtilisin lentus, subtilisin Novo, subtilisin Carlsberg, Bacillus licheniformis, subtilisin BPN', subtilisin 309, subtilisin 147 and subtilisin 168 described in WO89/06279 and protease PD138 described in (WO93/18140). Other useful proteases may be those described in WO92/175177, WO01/016285, WO02/026024 and WO02/016547. Examples of trypsin-like proteases are trypsin (e.g. of porcine or bovine origin) and the Fusarium protease described in WO98/06207, WO94/25583 and WO05/040372, and the chymotrypsin proteases derived from Cellumonas described in WO05/052161 and WO05/052146.

A further preferred protease is the alkaline protease from Bacillus lentus DSM 5483, as described for example in WO95/23221, and variants thereof which are described in WO92/21760, WO95/23221, EP1921147 and EP1921148.

Examples of metalloproteases are the neutral metalloprotease as described in WO07/044993 (Genencor Int.) such as those derived from Bacillus amyloliquefaciens.


Suitable commercially available protease enzymes include those sold under the trade names Alcalase®, Duralase™, Durazym™, Relase®, Relase® Ultra, Savinase®, Savinase®
Ultra, Primase®, Polarzyme®, Kannase®, Liquanase®, Liquanase® Ultra, Ovozyme®, Coronase®, Coronase® Ultra, Neutrase®, Everlase® and Esperase® (Novozymes A/S), those sold under the tradename Maxatase®, Maxacal®, Maxapem®, Purafect®, Purafect Prime®, Preferenz™, Purafect MA®, Purafect Ox®, Purafect OxP®, Puramax®, Properase®, Effectenz™, FN2®, FN3®, FN4®, Excellase®, Opticlean® and Optimase® (Danisco/DuPont), Axapem™ (Gist-Brocades N.V.), BLAP (sequence shown in Figure 29 of US5352604) and variants hereof (Henkel AG) and KAP (Bacillus alkalophilus subtilisin) from Kao.

Lipases and Cutinases: Suitable lipases and cutinases include those of bacterial or fungal origin. Chemically modified or protein engineered mutant enzymes are included.

Examples include lipase from Thermomyces, e.g. from T. lanuginosus (previously named Humicola lanuginosa) as described in EP258068 and EP305216, cutinase from Humicola, e.g. H. insolens (WO96/13580), lipase from strains of Pseudomonas (some of these now renamed to Burkholderia), e.g. P. alcaligenes or P. pseudoalcaligenes (EP218272), P. cepacia (EP331376), P. sp. strain SD705 (WO95/06720 & WO96/27002), P. wisconsinensis (WO96/12012), GDSL-type Streptomyces lipases (WO 10/065455), cutinase from Magnaporthe grisea (WO 10/1 07560), cutinase from Pseudomonas mendocina (US5,389,536), lipase from Thermobifida fusca (WO 11/084412), Geobacillus stearothermophilus lipase (WO 11/084417), lipase from Bacillus subtilis (WO 11/084599), and lipase from Streptomyces griseus (WO1 1/150157) and S. pristinaespiralis (WO12/1 37147).

Other examples are lipase variants such as those described in EP407225, WO92/05249, WO94/01541, WO94/25578, WO95/14783, WO95/30744, WO95/35381, WO95/22615, WO96/00292, WO97/04079, WO97/07202, WO00/34450, WO00/60063, WO01/92502, WO07/87508 and WO09/1 09500.

Preferred commercial lipase products include include Lipolase™, Lipex™; Lipolex™ and Lipoclean™ (Novozymes A/S), Lumafast (originally from Genencor) and Lipomax (originally from Gist-Brocades).

Still other examples are lipases sometimes referred to as acyltransferases or perhydrolases, e.g. acyltransferases with homology to Candida antarctica lipase A (WO10/1 11143), acyltransferase from Mycobacterium smegmatis (WO05/56782), perhydrolases from the CE 7 family (WO09/67279), and variants of the M. smegmatis perhydrolase in particular the S54V variant used in the commercial product Gentle Power Bleach from Huntsman Textile Effects Pte Ltd (WO1 0/1 00028).

Amylases: Suitable amylases which can be used together with XX of the invention may be an alpha-amylase or a glucoamylase and may be of bacterial or fungal origin. Chemically modified or protein engineered mutants are included. Amylases include, for example, alpha-amylases obtained from Bacillus, e.g., a special strain of Bacillus licheniformis, described in more detail in GB 1,296,839.
Suitable amylases include amylases having SEQ ID NO: 3 in WO 95/10603 or variants having 90% sequence identity to SEQ ID NO: 3 thereof. Preferred variants are described in WO 94/02597, WO 94/18314, WO 97/43424 and SEQ ID NO: 4 of WO 99/019467, such as variants with substitutions in one or more of the following positions: 15, 23, 105, 106, 124, 128, 133, 154, 156, 178, 179, 181, 188, 190, 197, 201, 202, 207, 208, 209, 211, 243, 264, 304, 305, 391, 408, and 444.

Different suitable amylases include amylases having SEQ ID NO: 6 in WO 02/010355 or variants thereof having 90% sequence identity to SEQ ID NO: 6. Preferred variants of SEQ ID NO: 6 are those having a deletion in positions 181 and 182 and a substitution in position 193.

Other amylases which are suitable are hybrid alpha-amylase comprising residues 1-33 of the alpha-amylase derived from B. amyloliquifaciens shown in SEQ ID NO: 6 of WO 2006/066594 and residues 36-483 of the B. licheniformis alpha-amylase shown in SEQ ID NO: 4 of WO 2006/066594 or variants having 90% sequence identity thereof. Preferred variants of this hybrid alpha-amylase are those having a substitution, a deletion or an insertion in one of more of the following positions: G48, T49, G107, H156, A181, N190, M197, I201, A209 and Q264. Most preferred variants of the hybrid alpha-amylase comprising residues 1-33 of the alpha-amylase derived from B. amyloliquifaciens shown in SEQ ID NO: 6 of WO 2006/066594 and residues 36-483 of SEQ ID NO: 4 are those having the substitutions:

M197T;

H156Y+A181T+N190F+A209V+Q264S; or

G48A+T49I+G107A+H156Y+A181T+N190F+I201 F+A209V+Q264S.

Further amylases which are suitable are amylases having SEQ ID NO: 6 in WO 99/019467 or variants thereof having 90% sequence identity to SEQ ID NO: 6. Preferred variants of SEQ ID NO: 6 are those having a substitution, a deletion or an insertion in one or more of the following positions: R181, G182, H183, G184, N195, I206, E212, E216 and K269. Particularly preferred amylases are those having deletion in positions R181 and G182, or positions H183 and G184.

Additional amylases which can be used are those having SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 2 or SEQ ID NO: 7 of WO 96/023873 or variants thereof having 90% sequence identity to SEQ ID NO: 1, SEQ ID NO: 2, SEQ ID NO: 3 or SEQ ID NO: 7. Preferred variants of SEQ ID NO: 1, SEQ ID NO: 2, SEQ ID NO: 3 or SEQ ID NO: 7 are those having a substitution, a deletion or an insertion in one or more of the following positions: 140, 181, 182, 183, 184, 195, 206, 212, 243, 260, 269, 304 and 476. More preferred variants are those having a deletion in positions 181 and 182 or positions 183 and 184. Most preferred amylase variants of SEQ ID NO: 1, SEQ ID NO: 2 or SEQ ID NO: 7 are those having a deletion in positions 183 and 184 and a substitution in one or more of positions 140, 195, 206, 243, 260, 304 and 476.
Other amylases which can be used are amylases having SEQ ID NO: 2 of WO 08/153815, SEQ ID NO: 10 in WO 01/66712 or variants thereof having 90% sequence identity to SEQ ID NO: 2 of WO 08/153815 or 90% sequence identity to SEQ ID NO: 10 in WO 01/66712. Preferred variants of SEQ ID NO: 10 in WO 01/66712 are those having a substitution, a deletion or an insertion in one of more of the following positions: 176, 177, 178, 179, 190, 201, 207, 211 and 264.

Further suitable amylases are amylases having SEQ ID NO: 2 of WO 09/061380 or variants having 90% sequence identity to SEQ ID NO: 2 thereof. Preferred variants of SEQ ID NO: 2 are those having a truncation of the C-terminus and/or a substitution, a deletion or an insertion in one of more of the following positions: Q87, Q98, S125, N128, T131, T165, K178, R180, S181, T182, G183, M201, F202, N225, S243, N272, N282, Y305, R309, D319, Q320, Q359, K444 and G475. More preferred variants of SEQ ID NO: 2 are those having the substitution in one of more of the following positions: Q87E,R, Q98R, S125A, N128C, T131I, T165I, K178L, T182G, M201L, F202Y, N225E,R, N272E,R, S243Q,A,E,D, Y305R, R309A, Q320R, Q359E, K444E and G475K and/or deletion in position R180 and/or S181 or of T182 and/or G183. Most preferred amylase variants of SEQ ID NO: 2 are those having the substitutions:

N128C+K178L+T182G+Y305R+G475K;
N128C+K178L+T182G+F202Y+Y305R+D31 9T+G475K;
S125A+N128C+K178L+T182G+Y305R+G475K; or
S125A+N128C+T131I+T165I+K178L+T182G+Y305R+G475K wherein the variants are C-terminally truncated and optionally further comprises a substitution at position 243 and/or a deletion at position 180 and/or position 181.

Other suitable amylases are the alpha-amylase having SEQ ID NO: 12 in WO01/66712 or a variant having at least 90% sequence identity to SEQ ID NO: 12. Preferred amylase variants are those having a substitution, a deletion or an insertion in one of more of the following positions of SEQ ID NO: 12 in WO01/66712: R28, R118, N174; R181, G182, D183, G184, G186, W189, N195, M202, Y298, N299, K302, S303, N306, R310, N314; R320, H324, E345, Y396, R400, W439, R444, N445, K446, Q449, R458, N471, N484. Particular preferred amylases include variants having a deletion of D183 and G184 and having the substitutions R118K, N195F, R320K and R458K, and a variant additionally having substitutions in one or more position selected from the group: M9, G149, G182, G186, M202, T257, Y295, N299, M323, E345 and A339, most preferred a variant that additionally has substitutions in all these positions.

Other examples are amylase variants such as those described in WO20 11/098531, WO201 3/001 078 and WO201 3/001 087.
Commercially available amylases are Duramyl™, Termamyl™, Fungamyl™, Stainzyme™, Stainzyme Plus™, Natalase™, Liquozyme X and BAN™ (from Novozymes A/S), and Rapidase™, Purastar™/Effectenz™, Powerase and Preferenz S100 (from Genencor International Inc./DuPont).

Peroxidases/Oxidases: Suitable peroxidases/oxidases include those of plant, bacterial or fungal origin. Chemically modified or protein engineered mutants are included. Examples of useful peroxidases include peroxidases from Coprinus, e.g., from C. cinereus, and variants thereof as those described in WO 93/24618, WO 95/10602, and WO 98/15257.

Commercially available peroxidases include Guardzyme™ (Novozymes A/S).

The detergent enzyme(s) may be included in a detergent composition by adding separate additives containing one or more enzymes, or by adding a combined additive comprising all of these enzymes. A detergent additive of the invention, i.e., a separate additive or a combined additive, can be formulated, for example, as a granulate, liquid, slurry, etc. Preferred detergent additive formulations are granulates, in particular non-dusting granulates, liquids, in particular stabilized liquids, or slurries.

Non-dusting granulates may be produced, e.g., as disclosed in US 4,106,991 and 4,661,452 and may optionally be coated by methods known in the art. Examples of waxy coating materials are poly(ethylene oxide) products (polyethyleneglycol, PEG) with mean molar weights of 1000 to 20000; ethoxylated nonylphenols having from 16 to 50 ethylene oxide units; ethoxylated fatty alcohols in which the alcohol contains from 12 to 20 carbon atoms and in which there are 15 to 80 ethylene oxide units; fatty alcohols; fatty acids; and mono- and di- and triglycerides of fatty acids. Examples of film-forming coating materials suitable for application by fluid bed techniques are given in GB 1483591. Liquid enzyme preparations may, for instance, be stabilized by adding a polyol such as propylene glycol, a sugar or sugar alcohol, lactic acid or boric acid according to established methods. Protected enzymes may be prepared according to the method disclosed in EP 238,216.

Adjunct materials

Any detergent components known in the art for use in laundry detergents may also be utilized. Other optional detergent components include anti-corrosion agents, anti-shrink agents, anti-soil redeposition agents, anti-wrinkling agents, bactericides, binders, corrosion inhibitors, disintegrants/disintegration agents, dyes, enzyme stabilizers (including boric acid, borates, CMC, and/or polyols such as propylene glycol), fabric conditioners including clays, fillers/processing aids, fluorescent whitening agents/optical brighteners, foam boosters, foam (suds) regulators, perfumes, soil-suspending agents, softeners, suds suppressors, tarnish inhibitors, and wicking agents, either alone or in combination. Any ingredient known in the art for
use in laundry detergents may be utilized. The choice of such ingredients is well within the skill of the artisan.

**Dispersants:** The detergent compositions of the present invention can also contain dispersants. In particular powdered detergents may comprise dispersants. Suitable water-soluble organic materials include the homo- or co-polymeric acids or their salts, in which the polycarboxylic acid comprises at least two carboxyl radicals separated from each other by not more than two carbon atoms. Suitable dispersants are for example described in Powdered Detergents, Surfactant science series volume 71, Marcel Dekker, Inc.

**Dye Transfer Inhibiting Agents:** The detergent compositions of the present invention may also include one or more dye transfer inhibiting agents. Suitable polymeric dye transfer inhibiting agents include, but are not limited to, polyvinylpyrrolidone polymers, polyamine N-oxide polymers, copolymers of N-vinylpyrrolidone and N-vinylimidazole, polyvinyloxazolidones and polyvinylimidazoles or mixtures thereof. When present in a subject composition, the dye transfer inhibiting agents may be present at levels from about 0.0001% to about 10%, from about 0.01% to about 5% or even from about 0.1% to about 3% by weight of the composition.

**Fluorescent whitening agent:** The detergent compositions of the present invention will preferably also contain additional components that may tint articles being cleaned, such as fluorescent whitening agent or optical brighteners. Where present the brightener is preferably at a level of about 0.01% to about 0.5%. Any fluorescent whitening agent suitable for use in a laundry detergent composition may be used in the composition of the present invention. The most commonly used fluorescent whitening agents are those belonging to the classes of diaminostilbene-sulphonic acid derivatives, diarylpyrazoline derivatives and bisphenyl-distyryl derivatives. Examples of the diaminostilbene-sulphonic acid derivative type of fluorescent whitening agents include the sodium salts of: 4,4'-bis-(2-diethanolamino-4-anilino-s-triazin-6-ylamino) stilbene-2,2'-disulphonate; 4,4'-bis-(2,4-dianilino-s-triazin-6-ylamino) stilbene-2,2'-disulphonate; 4,4'-bis-(2-anilino-4(N-methyl-N-2-hydroxy-ethylamino)-s-triazin-6-ylamino) stilbene-2,2'-disulphonate, 4,4'-bis-(4-phenyl-2,1,3-triazol-2-yl)stilbene-2,2'-disulphonate; 4,4'-bis-(2-anilino-4(1-methyl-2-hydroxy-ethylamino)-s-triazin-6-ylamino) stilbene-2,2'-disulphonate and 2-(stilbyl-4"-naptho-1 ,2,4,5)-1 ,2,3-trizole-2"-sulphonate. Preferred fluorescent whitening agents are Tinopal DMS and Tinopal CBS available from Ciba-Geigy AG, Basel, Switzerland. Tinopal DMS is the disodium salt of 4,4'-bis-(2-morpholino-4 anilino-s-triazin-6-ylamino) stilbene disulphonate. Tinopal CBS is the disodium salt of 2,2'-bis-(phenyl-styryl) disulphonate. Also preferred are fluorescent whitening agents is the commercially available Parawhite KX, supplied by Paramount Minerals and Chemicals, Mumbai, India. Other fluorescers suitable for use in the invention include the 1-3-diaryl pyrazolines and the 7-alkylaminocoumarins. Suitable fluorescent brightener levels include lower levels of from about 0.01%, from 0.05, from about 0.1 or even from about 0.2 wt % to upper levels of 0.5 or even 0.75 wt%.

**Soil release polymers:** The detergent compositions of the present invention may also include one or more soil release polymers which aid the removal of soils from fabrics such as
cotton and polyester based fabrics, in particular the removal of hydrophobic soils from polyester based fabrics. The soil release polymers may for example be nonionic or anionic terephthalate based polymers, polyvinyl caprolactam and related copolymers, vinyl graft copolymers, polyester polyamides see for example Chapter 7 in Powdered Detergents, Surfactant science series volume 71, Marcel Dekker, Inc. Another type of soil release polymers are amphiphilic alkoxylated grease cleaning polymers comprising a core structure and a plurality of alkoxylate groups attached to that core structure. The core structure may comprise a polyalkylenimine structure or a polyalkanolamine structure as described in detail in WO 2009/087523 (hereby incorporated by reference). Furthermore random graft co-polymers are suitable soil release polymers Suitable graft co-polymers are described in more detail in WO 2007/138054, WO 2006/108856 and WO 2006/1 13314 (hereby incorporated by reference). Other soil release polymers are substituted polysaccharide structures especially substituted cellulose structures such as modified cellulose derivatives such as those described in EP 1867808 or WO 2003/040279 (both are hereby incorporated by reference). Suitable cellulosic polymers include cellulose, cellulose ethers, cellulose esters, cellulose amides and mixtures thereof. Suitable cellulosic polymers include anionically modified cellulose, nonionically modified cellulose, cationically modified cellulose, zwitterionically modified cellulose, and mixtures thereof. Suitable cellulosic polymers include methyl cellulose, carboxy methyl cellulose, ethyl cellulose, hydroxyl ethyl cellulose, hydroxyl propyl methyl cellulose, ester carboxy methyl cellulose, and mixtures thereof.

Anti-redeposition agents: The detergent compositions of the present invention may also include one or more anti-redeposition agents such as carboxymethylcellulose (CMC), polyvinyl alcohol (PVA), polyvinylpyrrolidone (PVP), polyoxyethylene and/or polyethylene glycol (PEG), homopolymers of acrylic acid, copolymers of acrylic acid and maleic acid, and ethoxylated polyethyleneimines. The cellulose based polymers described under soil release polymers above may also function as anti-redeposition agents.

Other suitable adjunct materials include, but are not limited to, anti-shrink agents, anti-wrinkling agents, bactericides, binders, carriers, dyes, enzyme stabilizers, fabric softeners, fillers, foam regulators, hydrotrpops, perfumes, pigments, sod suppressors, solvents, and structurants for liquid detergents and/or structure elasticizing agents.

Formulation of detergent products

The detergent composition of the invention may be in any convenient form, e.g., a bar, a homogenous tablet, a tablet having two or more layers, a pouch having one or more compartments, a regular or compact powder, a granule, a paste, a gel, or a regular, compact or concentrated liquid. There are a number of detergent formulation forms such as layers (same or different phases), pouches, as well as forms for machine dosing unit.

Pouches can be configured as single or multicompartment. It can be of any form, shape and material which is suitable for hold the composition, e.g. without allowing the release of the
composition from the pouch prior to water contact. The pouch is made from water soluble film which encloses an inner volume. Said inner volume can be divided into compartments of the pouch. Preferred films are polymeric materials preferably polymers which are formed into a film or sheet. Preferred polymers, copolymers or derivatives thereof are selected polyacrylates, and water soluble acrylate copolymers, methyl cellulose, carboxy methyl cellulose, sodium dextrin, ethyl cellulose, hydroxyethyl cellulose, hydroxypropyl methyl cellulose, malto dextrin, poly methacrylates, most preferably polyvinyl alcohol copolymers and, hydroxypropyl methyl cellulose (HPMC). Preferably the level of polymer in the film for example PVA is at least about 60%. Preferred average molecular weight will typically be about 20,000 to about 150,000. Films can also be of blend compositions comprising hydrolytically degradable and water soluble polymer blends such as polyactide and polyvinyl alcohol (known under the Trade reference M8630 as sold by Chris Craft In. Prod. Of Gary, Ind., US) plus plasticisers like glycerol, ethylene glycerol, Propylene glycol, sorbitol and mixtures thereof. The pouches can comprise a solid laundry cleaning composition or part components thereof or a liquid cleaning composition or part components separated by the water soluble film. The compartment for liquid components can be different in composition than compartments containing solids. Ref: (US2009/001 1970 A1).

Detergent ingredients can be separated physically from each other by compartments in water dissolvable pouches or in different layers of tablets. Thereby negative storage interaction between components can be avoided. Different dissolution profiles of each of the compartments can also give rise to delayed dissolution of selected components in the wash solution.

A liquid or gel detergent, which is not unit dosed, may be aqueous, typically containing at least 20% by weight and up to 95% water, such as up to about 70% water, up to about 65% water, up to about 55% water, up to about 45% water, up to about 35% water. Other types of liquids, including without limitation, alkanols, amines, diols, ethers and polyols may be included in an aqueous liquid or gel. An aqueous liquid or gel detergent may contain from 0-30% organic solvent. A liquid or gel detergent may be non-aqueous.

Laundry Soap Bars

The enzymes of the invention may be added to laundry soap bars and used for hand washing laundry, fabrics and/or textiles. The term laundry soap bar includes laundry bars, soap bars, combo bars, syndet bars and detergent bars. The types of bar usually differ in the type of surfactant they contain, and the term laundry soap bar includes those containing soaps from fatty acids and/or synthetic soaps. The laundry soap bar has a physical form which is solid and not a liquid, gel or a powder at room temperature. The term solid is defined as a physical form which does not significantly change over time, i.e. if a solid object (e.g. laundry soap bar) is placed inside a container, the solid object does not change to fill the container it is placed in. The bar is a solid typically in bar form but can be in other solid shapes such as round or oval.
The laundry soap bar may contain one or more additional enzymes, protease inhibitors such as peptide aldehydes (or hydrosulfite adduct or hemiacetal adduct), boric acid, borate, borax and/or phenylboronic acid derivatives such as 4-formylphenylboronic acid, one or more soaps or synthetic surfactants, polyols such as glycerine, pH controlling compounds such as fatty acids, citric acid, acetic acid and/or formic acid, and/or a salt of a monovalent cation and an organic anion wherein the monovalent cation may be for example Na\(^+\), K\(^+\) or NH\(_4^+\) and the organic anion may be for example formate, acetate, citrate or lactate such that the salt of a monovalent cation and an organic anion may be, for example, sodium formate.

The laundry soap bar may also contain complexing agents like EDTA and HEDP, perfumes and/or different type of fillers, surfactants e.g. anionic synthetic surfactants, builders, polymeric soil release agents, detergent chelators, stabilizing agents, fillers, dyes, colorants, dye transfer inhibitors, alkoxylated polycarbonates, suds suppressers, structurants, binders, leaching agents, bleaching activators, clay soil removal agents, anti-redeposition agents, polymeric dispersing agents, brighteners, fabric softeners, perfumes and/or other compounds known in the art.

The laundry soap bar may be processed in conventional laundry soap bar making equipment such as but not limited to: mixers, plodders, e.g. a two stage vacuum plodder, extruders, cutters, logo-stampers, cooling tunnels and wrappers. The invention is not limited to preparing the laundry soap bars by any single method. The premix of the invention may be added to the soap at different stages of the process. For example, the premix containing a soap, an enzyme, optionally one or more additional enzymes, a protease inhibitor, and a salt of a monovalent cation and an organic anion may be prepared and and the mixture is then plodded. The enzyme and optional additional enzymes may be added at the same time as the protease inhibitor for example in liquid form. Besides the mixing step and the plodding step, the process may further comprise the steps of milling, extruding, cutting, stamping, cooling and/or wrapping.

Granular detergent formulations


WO2013/189972
WO2010105905, WO2010105910, WO2010105813, WO2010112528, WO2010120863,
WO2010108002, WO2010111365, WO2010108000, WO2010107635, WO2010099091,
WO2010033976, WO2010033746, WO2010033747, WO2010033897, WO2010033979,
WO2010030540, WO2010030541, WO2010024467, WO2010024469,
WO2010024470, WO201010025161, WO2010014395, WO2010044905, WO2010145887,
WO2010142503, WO2010122051, WO2010102861, WO2010099997, WO2010084039,
WO2010076292, WO2010069742, WO2010069718, WO2010069957, WO2010057784,
WO2010054986, WO20100018043, WO2010003783, WO2010003792, WO2010102316,
WO2010142539, WO2010118959, WO2010115813, WO2010105942, WO2010105961,
WO20100105962, WO20100094356, WO20100084203, WO20100078979, WO20100072456,
WO20100069905, WO20100076165, WO20100072603, WO20100066486, WO20100066631,
WO20100066632, WO20100063689, WO20100060821, WO2010049187, WO20100031607,
WO2010000636.

Washing Method

The detergent compositions of the present invention are ideally suited for use in laundry applications. Accordingly, the present invention includes a method for laundering a fabric. The method comprises the steps of contacting a fabric to be laundered with a cleaning laundry solution comprising the detergent composition according to the invention. The fabric may comprise any fabric capable of being laundered in normal consumer use conditions. The solution preferably has a pH of from about 5.5 to about 8. The compositions may be employed at concentrations of from about 100 ppm, preferably 500 ppm to about 15,000 ppm in solution. The water temperatures typically range from about 5°C to about 90°C, including about 10°C, about 15°C, about 20°C, about 25°C, about 30°C, about 35°C, about 40°C, about 45°C, about 50°C, about 55°C, about 60°C, about 65°C, about 70°C, about 75°C, about 80°C, about 85°C and about 90°C. The water to fabric ratio is typically from about 1:1 to about 30:1.

In particular embodiments, the washing method is conducted at a pH of from about 5.0 to about 11.5, or in alternative embodiments, even from about 6 to about 10.5, such as about 5 to about 11, about 5 to about 10, about 5 to about 9, about 5 to about 8, about 5 to about 7, about 5.5 to about 11, about 5.5 to about 10, about 5.5 to about 9, about 5.5 to about 8, about 5.5 to about 7, about 6 to about 11, about 6 to about 10, about 6 to about 9, about 6 to about 8, about 6 to about 7, about 6.5 to about 11, about 6.5 to about 10, about 6.5 to about 9, about 6.5 to about 8, about 6.5 to about 7, about 7 to about 11, about 7 to about 10, about 7 to about 9, or about 7 to about 8, preferably about 5.5 to about 9, and more preferably about 6 to about 8.

In particular embodiments, the washing method is conducted at a degree of hardness of from about 0°dH to about 30°dH, such as about 1°dH, about 2°dH, about 3°dH, about 4°dH, about 5°dH, about 6°dH, about 7°dH, about 8°dH, about 9°dH, about 10°dH, about 11°dH,
about 12°dH, about 13°dH, about 14°dH, about 15°dH, about 16°dH, about 17°dH, about 18°dH, about 19°dH, about 20°dH, about 21°dH, about 22°dH, about 23°dH, about 24°dH, about 25°dH, about 26°dH, about 27°dH, about 28°dH, about 29°dH, about 30°dH. Under typical European wash conditions, the degree of hardness is about 15°dH, under typical US wash conditions about 6°dH, and under typical Asian wash conditions, about 3°dH.

The present invention relates to a method of cleaning a fabric, a dishware or hard surface with a detergent composition comprising a protease of the invention.

A preferred embodiment concerns a method of cleaning, said method comprising the steps of: contacting an object with a cleaning composition comprising a protease of the invention under conditions suitable for cleaning said object. In a preferred embodiment the cleaning composition is a detergent composition and the process is a laundry or a dish wash process.

Still another embodiment relates to a method for removing stains from fabric which comprises contacting said a fabric with a composition comprising a protease of the invention under conditions suitable for cleaning said object.

In a preferred embodiment the compositions for use in the methods above further comprises at least one additional enzyme as set forth in the “other enzymes” section above, such as an enzyme selected from the group consisting of carbohydrases, peptidases, proteases, lipases, cellulase, xylanases or cutinases or a combination hereof. In yet another preferred embodiment the compositions comprises a reduced amount of at least one or more of the following components a surfactant, a builder, a chelator or chelating agent, bleach system or bleach component or a polymer.

Also contemplated are compositions and methods of treating fabrics (e.g., to desize a textile) using one or more of the protease of the invention. The protease can be used in any fabric-treating method which is well known in the art (see, e.g., U.S. Patent No. 6,077,316). For example, in one aspect, the feel and appearance of a fabric is improved by a method comprising contacting the fabric with a protease in a solution. In one aspect, the fabric is treated with the solution under pressure.

In one embodiment, the protease is applied during or after the weaving of textiles, or during the desizing stage, or one or more additional fabric processing steps. During the weaving of textiles, the threads are exposed to considerable mechanical strain. Prior to weaving on mechanical looms, warp yarns are often coated with sizing starch or starch derivatives in order to increase their tensile strength and to prevent breaking. The protease can be applied to remove these sizing protein or protein derivatives. After the textiles have been woven, a fabric can proceed to a desizing stage. This can be followed by one or more additional fabric processing steps. Desizing is the act of removing size from textiles. After weaving, the size coating should be removed before further processing the fabric in order to ensure a
homogeneous and wash-proof result. Also provided is a method of desizing comprising enzymatic hydrolysis of the size by the action of an enzyme.

**Low Temperature Uses**

It was surprisingly found that the proteases of the present invention - were actually performing relatively better at low temperature, e.g., temperatures of about 40°C or below than at higher temperatures, e.g., of about 60°C or above when tested in AMSA as described in the below Examples.

Moreover, in a particularly preferred embodiment the proteases of the invention perform relatively better than a well known subtilisin protease such as Savinase at a wash temperature of about 40°C or below when tested in AMSA as described herein.

Thus, in one embodiment of the invention concerns a method of doing laundry, dish wash or industrial cleaning comprising contacting a surface to be cleaned with a protease of the invention, and wherein said laundry, dish wash, industrial or institutional cleaning is performed at a temperature of about 40°C or below. One embodiment of the invention relates to the use of a protease of the invention in laundry, dish wash or a cleaning process wherein the temperature in laundry, dish wash, industrial cleaning is about 40°C or below.

In another embodiment, the invention concerns the use of a protease of the invention in a protein removing process, wherein the temperature in the protein removing process is about 40°C or below.

The present invention also relates to the use in laundry, dish wash or industrial cleaning process of a protease of the invention having at least one improved property compared to Savinase and wherein the temperature in laundry, dish wash or cleaning process is performed at a temperature of about 40°C or below.

In each of the above-identified methods and uses, the wash temperature is about 40°C or below, such as about 39°C or below, such as about 38°C or below, such as about 37°C or below, such as about 36°C or below, such as about 35°C or below, such as about 34°C or below, such as about 33°C or below, such as about 32°C or below, such as about 31°C or below, such as about 30°C or below, such as about 29°C or below, such as about 28°C or below, such as about 27°C or below, such as about 26°C or below, such as about 25°C or below, such as about 24°C or below, such as about 23°C or below, such as about 22°C or below, such as about 21°C or below, such as about 20°C or below, such as about 19°C or below, such as about 18°C or below, such as about 17°C or below, such as about 16°C or below, such as about 15°C or below, such as about 14°C or below, such as about 13°C or below, such as about 12°C or below, such as about 11°C or below, such as about 10°C or below, such as about 9°C or below, such as about 8°C or below, such as about 7°C or below,
such as about 6°C or below, such as about 5°C or below, such as about 4°C or below, such as about 3°C or below, such as about 2°C or below, such as about 1°C or below.

In another preferred embodiment, the wash temperature is in the range of about 5-40°C, such as about 5-30°C, about 5-20°C, about 5-10°C, about 10-40°C, about 10-30°C, about 10-20°C, about 15-40°C, about 15-30°C, about 15-20°C, about 20-40°C, about 20-30°C, about 25-40°C, about 25-30°C, or about 30-40°C. In a particular preferred embodiment the wash temperature is about 30°C.

In particular embodiments, the low temperature washing method is conducted at a pH of from about 5.0 to about 11.5, or in alternative embodiments, even from about 6 to about 10.5, such as about 5 to about 11, about 5 to about 10, about 5 to about 9, about 5 to about 8, about 5 to about 7, about 5.5 to about 11, about 5.5 to about 10, about 5.5 to about 9, about 5.5 to about 8, about 5.5, to about 7, about 6 to about 11, about 6 to about 10, about 6 to about 9, about 6 to about 8, about 6 to about 7, about 6.5 to about 11, about 6.5 to about 10, about 6.5 to about 9, about 6.5 to about 8, about 6.5 to about 7, about 7 to about 11, about 7 to about 10, about 7 to about 9, or about 7 to about 8, preferably about 5 to about 9, and more preferably about 6 to about 8.

In particular embodiments, the low temperature washing method is conducted at a degree of hardness of from about 0°dH to about 30°dH, such as about 1°dH, about 2°dH, about 3°dH, about 4°dH, about 5°dH, about 6°dH, about 7°dH, about 8°dH, about 9°dH, about 10°dH, about 11°dH, about 12°dH, about 13°dH, about 14°dH, about 15°dH, about 16°dH, about 17°dH, about 18°dH, about 19°dH, about 20°dH, about 21°dH, about 22°dH, about 23°dH, about 24°dH, about 25°dH, about 26°dH, about 27°dH, about 28°dH, about 29°dH, about 30°dH. Under typical European wash conditions, the degree of hardness is about 15°dH, under typical US wash conditions about 6°dH, and under typical Asian wash conditions, about 3°dH.

**Uses**

The present invention is directed to methods for using the polypeptides having protease activity, or compositions thereof. The invention may be used in compositions thereof in the laundering of textiles and fabrics, such as house hold laundry washing and industrial laundry washing. The invention is directed to methods for using the compositions thereof in hard surface cleaning such as automated dish washing (ADW), car wash and cleaning of industrial surfaces.

The present invention is also directed to methods for using the proteases having protease activity in animal feed, as well as to feed compositions and feed additives comprising the proteases of the invention.
Use of Proteases of the Invention in Animal Feed

The term animal includes all animals. Examples of animals are non-ruminants, and ruminants. Ruminant animals include, for example, animals such as sheep, goats, and cattle, e.g. beef cattle, cows, and young calves. 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, ducks and chicken (including but not limited to broiler chicks, layers); horses (including but not limited to hotbloods, coldbloods and warm bloods), young calves; and fish (including but not limited to salmon, trout, tilapia, catfish and carp); and crustaceans (including but not limited to shrimps and prawns).

The term feed or feed composition means any compound, preparation, mixture, or composition suitable for, or intended for intake by an animal. In the use according to the invention the protease can be fed to the animal before, after, or simultaneously with the diet. The latter is preferred.

In a particular embodiment, the protease, in the form in which it is added to the feed, or when being included in a feed additive, is well-defined. Well-defined means that the protease preparation is at least 50% pure as determined by Size-exclusion chromatography (see Example 12 of WO 01/58275). In other particular embodiments the protease preparation is at least 60, 70, 80, 85, 88, 90, 92, 94, or at least 95% pure as determined by this method.

A well-defined protease preparation is advantageous. For instance, it is much easier to dose correctly to the feed a protease that is essentially free from interfering or contaminating other proteases or other proteins in general. The term dose correctly refers in particular to the objective of obtaining consistent and constant results, and the capability of optimising dosage based upon the desired effect.

For the use in animal feed, however, the protease need not be that pure; it may e.g. include other enzymes, in which case it could be termed a protease preparation.

The protease preparation can be (a) added directly to the feed (or used directly in a protein treatment process), or (b) it can be used in the production of one or more intermediate compositions such as feed additives or premixes that is subsequently added to the feed (or used in a treatment process). The degree of purity described above refers to the purity of the original protease preparation, whether used according to (a) or (b) above.

Protease preparations with purities of this order of magnitude are in particular obtainable using recombinant methods of production, whereas they are not so easily obtained and also subject to a much higher batch-to-batch variation when the protease is produced by traditional fermentation methods. Such protease preparation may of course be mixed with other enzymes to obtain a preparation with two or more purified enzymes with different or similar activities.
The substrate protein may be an animal protein, such as meat and bone meal, feather meal, and/or fish meal; or it may be a vegetable protein.

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, sunflower seed, cotton seed, 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, triticale, sorghum, dried distillers grains with solubles (DDGS) and microalgae.

In a particular embodiment of a treatment process the protease(s) in question is affecting (or acting on, or exerting its hydrolyzing or degrading influence on) the proteins, such as vegetable proteins or protein sources. To achieve this, the protein or protein source is typically suspended in a solvent, e.g. an aqueous solvent such as water, and the pH and temperature values are adjusted paying due regard to the characteristics of the enzyme in question. For example, the treatment may take place at a pH-value at which the activity of the actual protease is at least 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, or at least 90%. Likewise, for example, the treatment may take place at a temperature at which the activity of the actual protease is at least 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, or at least 90%. The above percentage activity indications are relative to the maximum activities. The enzymatic reaction is continued until the desired result is achieved, following which it may or may not be stopped by inactivating the enzyme, e.g. by a heat-treatment step.

In another particular embodiment of a treatment process of the invention, the protease action is sustained, meaning e.g. that the protease is added to the proteins, but its hydrolyzing influence is so to speak not switched on until later when desired, once suitable hydrolysing conditions are established, or once any enzyme inhibitors are inactivated, or whatever other means could have been applied to postpone the action of the enzyme.

In one embodiment the treatment is a pre-treatment of animal feed or proteins for use in animal feed, i.e. the proteins are hydrolysed before intake.
The term improving the nutritional value of an animal feed means improving the availability of nutrients in the feed. In this invention, improving the nutritional values refers in particular to improving the availability of the protein fraction of the feed, thereby leading to increased protein extraction, higher protein yields, and/or improved protein utilization. When the nutritional value of the feed is increased, the protein and/or amino acid digestibility is increased and the growth rate and/or weight gain and/or feed conversion (i.e., the weight of ingested feed relative to weight gain) of the animal might be improved.

The protease can be added to the feed in any form, be it as a relatively pure protease, or in admixture with other components intended for addition to animal feed, i.e., in the form of animal feed additives, such as the so-called pre-mixes for animal feed.

In a further aspect, the present invention relates to compositions for use in animal feed, such as animal feed, and animal feed additives, e.g., premixes.

Apart from the protease of the invention, the animal feed additives of the invention contain at least one fat-soluble vitamin, and/or at least one water-soluble vitamin, and/or at least one trace mineral, and/or at least one macro mineral.

Further, optional, feed-additive ingredients are colouring agents, e.g., carotenoids such as beta-carotene, astaxanthin, and lutein; stabilisers; growth improving additives and aroma compounds/flavourings, e.g., creosol, anethol, deca-, undeca- and/or dodeca-lactones, ionones, irone, gingerol, piperidine, propylidine phthalate, butyldiene phthalate, capsaicin and/or tannin; antimicrobial peptides; polyunsaturated fatty acids (PUFAs); reactive oxygen generating species; also, a support may be used that may contain, for example, 40-50% by weight of wood fibres, 8-10% by weight of stearine, 4-5% by weight of curcuma powder, 4-58% by weight of rosemary powder, 22-28% by weight of limestone, 1-3% by weight of a gum, such as gum arabic, 5-50% by weight of sugar and/or starch and 5-15% by weight of water.

A feed or a feed additive of the invention may also comprise at least one other enzyme selected from amongst phytase (EC 3.1.3.8 or 3.1.3.26); xylanase (EC 3.2.1.8); galactanase (EC 3.2.1.89); alpha-galactosidase (EC 3.2.1.22); further protease (EC 3.4), phospholipase A1 (EC 3.1.1.32); phospholipase A2 (EC 3.1.1.4); lysophospholipase (EC 3.1.1.5); phospholipase C (3.1.4.3); phospholipase D (EC 3.1.4.4); amylase such as, for example, alpha-amylase (EC 3.2.1.1); and/or beta-glucanase (EC 3.2.1.4 or EC 3.2.1.6).

In a particular embodiment, the feed or a feed additive of the invention also comprises a phytase (EC 3.1.3.8 or 3.1.3.26).

In a particular embodiment, the feed or a feed additive of the invention also comprises a xylanase (EC 3.2.1.8).

A feed or a feed additive of the invention may also comprise at least one probiotic or direct-fed microbial (DFM) optionally together with one or more other enzymes selected from amongst phytase (EC 3.1.3.8 or 3.1.3.26); xylanase (EC 3.2.1.8); galactanase (EC 3.2.1.89);
alpha-galactosidase (EC 3.2.1.22); further protease (EC 3.4), phospholipase A1 (EC 3.1.1.32); phospholipase A2 (EC 3.1.1.4); lysophospholipase (EC 3.1.1.5); phospholipase C (3.1.4.3); phospholipase D (EC 3.1.4.4); amylase such as, for example, alpha-amylase (EC 3.2.1.1); and/or beta-glucanase (EC 3.2.1.4 or EC 3.2.1.6).

The direct feed microbial may be a bacterium from one or more of the following genera: Lactobacillus, Lactococcus, Streptococcus, Bacillus, Pediococcus, Enterococcus, Leuconostoc, Carnobacterium, Propionibacterium, Bifidobacterium, Clostridium and Megasphaera or any combination thereof, preferably from Bacillus subtilis, Bacillus licheniformis, Bacillus amylobiiquefaciens, Enterococcus faecium, Enterococcus spp, and Pediococcus spp, Lactobacillus spp, Bifidobacterium spp, Lactobacillus acidophilus, Pediococcus acidilactici, Lactococcus lactis, Bifidobacterium bifidum, Propionibacterium thoenii, Lactobacillus farcininus, lactobacillus rhamnosus, Clostridium butyricum, Bifidobacterium animalis spp. animalis, Lactobacillus reuteri, Bacillus cereus, Lactobacillus salivarius spp. salivarius, Megasphaera elsdenii, Propionibacteria sp and more preferably from Bacillus subtilis strains 3A-P4 (PTA-6506); 15A-P4 (PTA-6507); 22C-P1 (PTA-6508); 2084 (NRRL B-500130); LSSA01 (NRRL-B-50104); BS27 (NRRL B-501 05); BS 18 (NRRL B-50633); and BS 278 (NRRL B-50634).

In a particular embodiment these other enzymes are well-defined (as defined above for protease preparations).

Examples of antimicrobial peptides (AMP's) are CAP18, Leucocin A, Tritrpticin, Protegrin-1, Thanatin, Defensin, Lactoferrin, Lactoferricin, and Ovispirin such as Novispirin (Robert Lehrer, 2000), Plectasins, and Statins, including the compounds and polypeptides disclosed in WO 03/044049 and WO 03/048148, as well as variants or fragments of the above that retain antimicrobial activity.

Examples of antifungal 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 WO 02/090384.

Examples of polyunsaturated fatty acids are C18, C20 and C22 polyunsaturated fatty acids, such as arachidonic acid, docosohexaenoic acid, eicosapentaenoic acid and gamma-linoleic acid.

Examples of reactive oxygen generating species are chemicals such as perborate, persulphate, or percarbonate; and enzymes such as an oxidase, an oxygenase or a syntethase.

Usually fat- and water-soluble vitamins, as well as trace minerals form part of a so-called premix intended for addition to the feed, whereas macro minerals are usually separately added to the feed. Either of these composition types, when enriched with a protease of the invention, is an animal feed additive of the invention.

In a particular embodiment, the animal feed additive of the invention is intended for being included (or prescribed as having to be included) in animal diets or feed at levels of 0.01
to 10.0%; more particularly 0.05 to 5.0%; or 0.2 to 1.0% (% meaning g additive per 100 g feed).
This is so in particular for premixes.

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 pantothenate, e.g. Ca-D-pantothenate.
Examples of trace minerals are manganese, zinc, iron, copper, iodine, selenium, and cobalt.
Examples of macro minerals are calcium, phosphorus and sodium.

The nutritional requirements of these components (exemplified with poultry and piglets/pigs) are listed in Table A of WO 01/58275. Nutritional requirement means that these components should be provided in the diet in the concentrations indicated.

In the alternative, the animal feed additive of the invention comprises at least one of the individual components specified in Table A of WO 01/58275. At least one means either of, one or more of, one, or two, or three, or four and so forth up to all thirteen, or up to all fifteen individual components. More specifically, this at least one individual component is included in the additive of the invention in such an amount as to provide an in-feed-concentration within the range indicated in column four, or column five, or column six of Table A.

In a still further embodiment, the animal feed additive of the invention comprises at least one of the below vitamins, preferably to provide an in-feed-concentration within the ranges specified in the below Table 1 (for piglet diets, and broiler diets, respectively).

Table 1: Typical vitamin recommendations

<table>
<thead>
<tr>
<th>Vitamin</th>
<th>Piglet diet</th>
<th>Broiler diet</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vitamin A</td>
<td>10,000-15,000 IU/kg feed</td>
<td>8-12,500 IU/kg feed</td>
</tr>
<tr>
<td>Vitamin D3</td>
<td>1800-2000 IU/kg feed</td>
<td>3000-5000 IU/kg feed</td>
</tr>
<tr>
<td>Vitamin E</td>
<td>60-100 mg/kg feed</td>
<td>150-240 mg/kg feed</td>
</tr>
<tr>
<td>Vitamin K3</td>
<td>2-4 mg/kg feed</td>
<td>2-4 mg/kg feed</td>
</tr>
<tr>
<td>Vitamin B1</td>
<td>2-4 mg/kg feed</td>
<td>2-3 mg/kg feed</td>
</tr>
<tr>
<td>Vitamin B2</td>
<td>6-10 mg/kg feed</td>
<td>7-9 mg/kg feed</td>
</tr>
<tr>
<td>Vitamin B6</td>
<td>4-8 mg/kg feed</td>
<td>3-6 mg/kg feed</td>
</tr>
<tr>
<td>Vitamin B12</td>
<td>0.03-0.05 mg/kg feed</td>
<td>0.015-0.04 mg/kg feed</td>
</tr>
<tr>
<td>Niacin (Vitamin B3)</td>
<td>30-50 mg/kg feed</td>
<td>50-80 mg/kg feed</td>
</tr>
<tr>
<td></td>
<td>Feed 1 (mg/kg)</td>
<td>Feed 2 (mg/kg)</td>
</tr>
<tr>
<td>----------------------</td>
<td>---------------</td>
<td>---------------</td>
</tr>
<tr>
<td>Pantothenic acid</td>
<td>20-40</td>
<td>10-18</td>
</tr>
<tr>
<td>Folic acid</td>
<td>1-2</td>
<td>1-2</td>
</tr>
<tr>
<td>Biotin</td>
<td>0.15-0.4</td>
<td>0.15-0.3</td>
</tr>
<tr>
<td>Choline chloride</td>
<td>200-400</td>
<td>300-600</td>
</tr>
</tbody>
</table>

The present invention also relates to animal feed compositions. Animal feed compositions or diets have a relatively high content of protein. Poultry and pig diets can be characterised as indicated in Table B of WO 01/58275, columns 2-3. Fish diets can be characterised as indicated in column 4 of this Table B. Furthermore such fish diets usually have a crude fat content of 200-310 g/kg. WO 01/58275 corresponds to US 09/779334 which is hereby incorporated by reference.

An animal feed composition according to the invention has a crude protein content of 50-800 g/kg, and furthermore comprises at least one protease as claimed herein.

Furthermore, or in the alternative (to the crude protein content indicated above), the animal feed composition of the invention has a content of metabolisable energy of 10-30 MJ/kg; and/or a content of calcium of 0.1-200 g/kg; and/or a content of available phosphorus of 0.1-200 g/kg; and/or a content of methionine of 0.1-100 g/kg; and/or a content of methionine plus cysteine of 0.1-150 g/kg; and/or a content of lysine of 0.5-50 g/kg.

In particular embodiments, the content of metabolisable energy, crude protein, calcium, phosphorus, methionine, methionine plus cysteine, and/or lysine is within any one of ranges 2, 3, 4 or 5 in Table B of WO 01/58275 (R. 2-5).

Crude protein is calculated as nitrogen (N) multiplied by a factor 6.25, i.e. Crude protein (g/kg) = N (g/kg) × 6.25. The nitrogen content is determined by the Kjeldahl method (A.O.A.C., 1984, Official Methods of Analysis 14th ed., Association of Official Analytical Chemists, Washington DC).


The dietary content of calcium, available phosphorus and amino acids in complete animal diets is calculated on the basis of feed tables such as Veevoedertabel 1997, gegevens over chemische samenstelling, verteerbaarheid en voederwaarde van voedermiddelen, Central Veevoederbureau, Runderweg 6, 8219 pk Lelystad. ISBN 90-72839-13-7.
In a particular embodiment, the animal feed composition of the invention contains at least one vegetable protein as defined above.

The animal feed composition of the invention may also contain animal protein, such as Meat and Bone Meal, Feather meal, and/or Fish Meal, typically in an amount of 0-25%. The animal feed composition of the invention may also comprise Dried Distillers Grains with Solubles (DDGS), typically in amounts of 0-30%.

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 0-25% fish meal; and/or 0-25% meat and bone meal; and/or 0-20% whey.

Animal diets can e.g. be manufactured as mash feed (non-pelleted) or pelleted feed. Typically, the milled feed-stuffs are mixed and sufficient amounts of essential vitamins and minerals are added according to the specifications for the species in question. Enzymes can be added as solid or liquid enzyme formulations. For example, for mash feed a solid or liquid enzyme formulation may be added before or during the ingredient mixing step. For pelleted feed the (liquid or solid) protease/enzyme preparation may also be added before or during the feed ingredient step. Typically a liquid protease/enzyme preparation is added after the pelleting step. The enzyme may also be incorporated in a feed additive or premix.

The final enzyme concentration in the diet is within the range of 0.01-200 mg enzyme protein per kg diet, for example in the range of 0.5-25 mg enzyme protein per kg animal diet.

The protease should of course be applied in an effective amount, i.e. in an amount adequate for improving protein hydrolysis, protein and amino acid digestibility, and/or improving nutritional value of feed. It is at present contemplated that the enzyme is administered in one or more of the following amounts (dosage ranges): 0.01-200; 0.01-100; 0.5-100; 1-50; 5-100; 10-100; 0.05-50; or 0.10-10 - all these ranges being in mg protease protein per kg feed (ppm).

For determining mg protease protein per kg feed, the protease is purified from the feed composition, and the specific activity of the purified protease is determined using a relevant assay (see under protease activity, substrates, and assays). The protease activity of the feed composition as such is also determined using the same assay, and on the basis of these two determinations, the dosage in mg protease protein per kg feed is calculated.

The same principles apply for determining mg protease protein in feed additives. Of course, if a sample is available of the protease used for preparing the feed additive or the feed, the specific activity is determined from this sample (no need to purify the protease from the feed composition or the additive).
Nucleic Acid Constructs, Expression Vectors, Recombinant Host Cells, and Methods for Production of Proteases

The present invention also relates to nucleic acid constructs, expression vectors and recombinant host cells comprising such polynucleotides encoding the proteases of the invention.

The present invention also relates to methods of producing a protease, comprising: (a) cultivating a recombinant host cell comprising such polynucleotide; and (b) recovering the protein.

The protein may be native or heterologous to a host cell. The term "protein" is not meant herein to refer to a specific length of the encoded product and, therefore, encompasses peptides, oligopeptides, and proteins. The term "protein" also encompasses two or more polypeptides combined to form the encoded product. The proteins also include hybrid polypeptides and fused polypeptides.

Preferably, the protein is a protease. For example, the protein may be a hydrolase, such as a proteolytic enzyme or protease.

The gene may be obtained from any prokaryotic, eukaryotic, or other source.

The present invention is further described by the following examples that should not be construed as limiting the scope of the invention.

EXAMPLES

Materials and Methods

Wash assays

Automatic Mechanical Stress Assay (AMSA) for laundry

In order to assess the wash performance in laundry washing experiments are performed, using the Automatic Mechanical Stress Assay (AMSA). With the AMSA, the wash performance of a large quantity of small volume enzyme-detergent solutions can be examined. The AMSA plate has a number of slots for test solutions and a lid firmly squeezing the laundry sample, the textile to be washed against all the slot openings. During the washing time, the plate, test solutions, textile and lid are vigorously shaken to bring the test solution in contact with the textile and apply mechanical stress in a regular, periodic oscillating manner. For further description see WO02/42740 especially the paragraph "Special method embodiments" at page 23-24.

The wash performance is measured as the brightness of the colour of the textile washed. Brightness can also be expressed as the intensity of the light reflected from the sample
when illuminated with white light. When the sample is stained the intensity of the reflected light is lower than that of a clean sample. Therefore the intensity of the reflected light can be used to measure wash performance.

Colour measurements are made with a professional flatbed scanner (Kodak iQsmart, Kodak, Midtager 29, DK-2605 Brandby, Denmark), which is used to capture an image of the washed textile.

To extract a value for the light intensity from the scanned images, 24-bit pixel values from the image are converted into values for red, green and blue (RGB). The intensity value (Int) is calculated by adding the RGB values together as vectors and then taking the length of the resulting vector:

\[
\text{Int} = \sqrt{r^2 + g^2 + b^2}.
\]

Table 2: Composition of model detergents and test materials

| Laundry powder model detergent A          | Sodium citrate dihydrate 32.3%  
|                                         | Sodium-LAS 24.2%  
|                                         | Sodium lauryl sulfate 32.2%  
|                                         | Neodol 25-7 (alcohol ethoxylate) 6.4%  
|                                         | Sodium sulfate 4.9%  
| Laundry liquid model detergent B         | Water 30.63%  
|                                         | Sodium hydroxide 2.95%  
|                                         | Dodecylbenzensulfonic acid 11.52%  
|                                         | Fatty acids (Soya) 5.50%  
|                                         | Propane-1,2-diol (MPG) 5.05%  
|                                         | Water 17.38%  
|                                         | C13-alcohol ethoxylate, 10.50%  
|                                         | Diethylenetriaminepentakis (methylenephosphonic acid) (DTMPA) 3.08%  
|                                         | Triethanolamine (TEA) 2.22%  
|                                         | Fatty acids (Coco) 4.50%  
|                                         | Sodium citrate monohydrate 1.00%  
|                                         | Ethanol 4.63%  
|                                         | Syntran 5909 (opacifier) 0.30%  
|                                         | Perfume 0.35%  
| Test material                           | PC-03 (Chocolate-milk/ink on cotton/polyester)  
|                                         | C-10 (Oil/milk/pigment on cotton)  
|                                         | PC-05 (Blood/milk/ink on cotton/polyester)  
|                                         | EMPA117EH (Blood/milk/ink on cotton/polyester)  

77
Test materials are obtained from Center For Testmaterials BV, P.O. Box 120, 3133 KT Vlaardingen, the Netherlands and EMPA Testmaterials AG, Movenstrasse 12, CH-9015 St. Gallen, Switzerland.

Protease assays

5 Suc-AAPF-pNA assay:
   pNA substrate : Suc-AAPF-pNA (Bachem L-1400).
   Temperature : Room temperature (25°C)
   Assay buffers : 100mM succinic acid, 100mM HEPES, 100mM CHES, 100mM CABS,
                   1mM CaCl$_2$, 150mM KCl, 0.01% Triton X-100 adjusted to pH-values 2.0,
                   3.0, 4.0, 5.0, 6.0, 7.0, 8.0, 9.0, 10.0, and 11.0 with HCl or NaOH.

20 µl protease (diluted in 0.01% Triton X-100) was mixed with 100 µl assay buffer. The assay was started by adding 100 µl pNA substrate (50mg dissolved in 1.0ml DMSO and further diluted 45x with 0.01 % Triton X-100). The increase in OD$_{405}$ was monitored as a measure of the protease activity.

10 Protazyme AK assay:
   Substrate : Protazyme AK tablet (cross-linked and dyed casein; from Megazyme)
   Temperature : controlled (assay temperature).
   Assay buffer : 100mM succinic acid, 100mM HEPES, 100mM CHES, 100mM CABS,
                   1mM CaCl$_2$, 150mM KCl, 0.01% Triton X-100, pH 6.5 or pH 7.0.

A Protazyme AK tablet was suspended in 2.0ml 0.01% Triton X-100 by gentle stirring. 500 µl of this suspension and 500 µl assay buffer were dispensed in an Eppendorf tube and placed on ice. 20 µl protease sample (diluted in 0.01% Triton X-100) was added. The assay was initiated by transferring the Eppendorf tube to an Eppendorf thermomixer, which was set to the assay temperature. The tube was incubated for 15 minutes on the Eppendorf thermomixer at its highest shaking rate (1400 rpm.). The incubation was stopped by transferring the tube back to the ice bath. Then the tube was centrifuged in an ice cold centrifuge for a few minutes and 200 µl supernatant was transferred to a microtiter plate. OD$_{650}$ was read as a measure of protease activity. A buffer blind was included in the assay (instead of enzyme).

Suc-AAPX-pNA assay:
   pNA substrates : Suc-AAPA-pNA (Bachem L-1775)
                  Suc-AAPR-pNA (Bachem L-1720)
                  Suc-AAPD-pNA (Bachem L-1835)
Suc-AAPI-pNA (Bachem L-1790)
Suc-AAPM-pNA (Bachem L-1395)
Suc-AAPV-pNA (Bachem L-1770)
Suc-AAPL-pNA (Bachem L-1390)
Suc-AAPE-pNA (Bachem L-1710)
Suc-AAPK-pNA (Bachem L-1725)
Suc-AAPF-pNA (Bachem L-1400)

Temperature : Room temperature (25°C)
Assay buffer : 100mM succinic acid, 100mM HEPES, 100mM CHES, 100mM CABS,
10 mM CaCl$_2$, 150mM KCl, 0.01% Triton X-100, pH 9.0.

20µl protease (diluted in 0.01% Triton X-100) was mixed with 100µl assay buffer. The assay was started by adding 100µl pNA substrate (50mg dissolved in 1.0ml DMSO and further diluted 45x with 0.01 % Triton X-100). The increase in OD$_{405}$ was monitored as a measure of the protease activity.

**o-Phthaldialdehyde (OPA) assay:**

This assay detects primary amines and hence cleavage of peptide bonds by a protease can be measured as the difference in absorbance between a protease treated sample and a control sample. The assay is conducted essentially according to Nielsen et al. (Nielsen, PM, Petersen, D, Dampmann, C. Improved method for determining food protein degree of hydrolysis. J Food Sci, 2001, 66: 642-646).

500 µl of sample is filtered through a 100 kDa Microcon centrifugal filter (60 min, 11,000 rpm, 5°C). The samples are diluted appropriately (e.g. 10, 50 or 100 times) in deionizer water and 25 µl of each sample is loaded into a 96 well microtiter plate (5 replicates). 200 µl OPA reagent (100 mM di-sodium tetraborate decahydrate, 3.5 mM sodium dodecyl sulphate (SDS), 5.7 mM di-thiothreitol (DDT), 6 mM o-Phthaldialdehyde) is dispensed into all wells, the plate is shaken (10 sec, 750 rpm) and absorbance measured at 340 nm.

**Strain**

The strain *Saccharomonospora viridis* DSM 43017 was obtained from DSMZ (Deutsche Sammlung von Mikroorganismen und Zellkulturen, Braunschweig - Germany). According to Pati *et al.*, 2009, *Stand. Genomic Sci.* 1:141-149, the strain was collected prior to 1963 from Irish peat.

**Example 1: Expression of S1 protease 1 from *Saccharomonospora viridis* (SEQ ID NO: 1)**

A linear integration vector-system was used for the expression cloning of the S1 protease 1 from *Saccharomonospora viridis* (SEQ ID NO: 1). The linear integration construct
was a PCR fusion product made by fusion of the gene between two *Bacillus subtilis* homologous chromosomal regions along with a strong promoter and a chloramphenicol resistance marker. The fusion was made by SOE PCR (Horton, R.M., Hunt, H.D., Ho, S.N., Pullen, J.K. and Pease, L.R. (1989) Engineering hybrid genes without the use of restriction enzymes, gene splicing by overlap extension Gene 77: 61-68). The SOE PCR method is also described in patent application WO 2003/095658. The gene was expressed under the control of a triple promoter system (as described in WO 1999/43835), consisting of the promoters from *Bacillus licheniformis* alpha-amylase gene (amyL), *Bacillus amyloliquefaciens* alpha-amylase gene (amyQ), and the *Bacillus thuringiensis* cryIIA promoter including stabilizing sequence. The gene coding for chloramphenicol acetyl-transferase was used as marker (described in e.g. Diderichsen, B.; Poulsen,G.B.; Joergensen,S.T. 1993, *Plasmid*, "A useful cloning vector for *Bacillus subtilis*" 30:312). The final gene constructs were integrated on the *Bacillus* chromosome by homologous recombination into the pectate lyase locus.

The gene encoding the S1 protease 1 from *Saccharomonospora viridis* was amplified from chromosomal DNA of the strain *Saccharomonospora viridis* DSM 43017 with gene specific primers containing overhang to the two flanking fragments. The upstream and downstream flanking fragments were amplified from genomic DNA of the strain iMB1 361 (described in patent application WO 2003/095658). The S1 protease 1 was expressed with a *Bacillus clausii* secretion signal (with the following amino acid sequence: MKKPLGKIVASTALLISVAFSSSIASA) replacing the native secretion signal.

The 2 vector fragments and the gene fragment was subjected to a Splicing by Overlap Extension (SOE) PCR reaction to assemble the 3 fragments into one linear vector construct. An aliquot of the PCR product was transformed into *Bacillus subtilis*. Transformants were selected on LB plates supplemented with 6 µg of chloramphenicol per ml. A recombinant *Bacillus subtilis* clone containing the integrated expression construct was grown in liquid culture. The enzyme containing supernatant was harvested and the enzyme purified as described in Example 2.

**Example 2: Purification of the S1 protease 1 from *Saccharomonospora viridis***

The culture broth was centrifuged (20000 x g, 20 min) and the supernatant was carefully decanted from the precipitate. The supernatant was filtered through a Nalgene 0.2µm filtration unit in order to remove the rest of the *Bacillus* host cells. The pH of the 0.2µm filtrate was adjusted to pH 4.5 with 20% CH3COOH and the conductivity was adjusted to the same conductivity as the conductivity of 20mM CH3COOH/NaOH, 50mM H3B03, 1mM CaCl2, pH 4.5 by dilution with deionised water. The adjusted filtrate was applied to a SP-sepharose FF column (from GE Healthcare) equilibrated in 20mM CH3COOH/NaOH, 50mM H3B03, 1mM CaCl2, pH 4.5. After washing the column extensively with the equilibration buffer, the protease was eluted with a linear NaCl gradient (0 → 0.5M) in the same buffer over five column volumes. Fractions
from the column were analysed for protease activity (using the Suc-AAPF-pNA assay at pH 9) and peak-fractions were pooled. The protease pool was diluted 10x with deionised water and pH was adjusted to pH 9 with 3M Tris-base. The adjusted pool was applied to a MEP Hypercel column (from Pall Corporation) equilibrated in 50mM Tris/HCl, 2mM CaCl₂, pH 9.0. After washing the column extensively with the equilibration buffer, the protease was step-eluted with 50mM CH₃COOH/NaOH, 2mM CaCl₂, pH 4.5. Fractions from the column were analysed for protease activity (using the Suc-AAPF-pNA assay at pH 9) and peak-fractions were further analysed by SDS-PAGE. The fractions where only one band was seen on the coomassie stained SDS-PAGE gel were pooled and were used for further characterization.

Example 3: Characterization of the S1 protease 1 from *Saccharomonospora viridis*

The Suc-AAPF-pNA assay was used for obtaining the pH-activity profile and the pH-stability profile (residual activity after 2 hours at indicated pH-values). For the pH-stability profile the protease was diluted 10x in the different assay buffers to reach the pH-values of these buffers and then incubated for 2 hours at 37°C. After incubation, the pH of the protease incubations was adjusted to the same pH-value by dilution in the pH 9.0 assay buffer. Residual activities were measured at pH 9.0 relative to a sample, which was kept at stable conditions (5°C, pH 9.0). The Protazyme AK assay was used for obtaining the temperature-activity profile at pH 7.0. The Suc-AAPX-pNA assay and ten different Suc-AAPX-pNA substrates were used for obtaining the P₁-specificity of the enzymes at pH 9.0. The results are shown in tables 3-6 and figures 1-4.

Table 3: pH-activity profile at 25°C

<table>
<thead>
<tr>
<th>pH</th>
<th>S1 protease 1 from <em>Saccharomonospora viridis</em></th>
<th>Protease 10R</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>0.00</td>
<td>-</td>
</tr>
<tr>
<td>3</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>4</td>
<td>0.00</td>
<td>0.02</td>
</tr>
<tr>
<td>5</td>
<td>0.03</td>
<td>0.07</td>
</tr>
<tr>
<td>6</td>
<td>0.15</td>
<td>0.21</td>
</tr>
<tr>
<td>7</td>
<td>0.56</td>
<td>0.44</td>
</tr>
<tr>
<td>8</td>
<td>1.00</td>
<td>0.67</td>
</tr>
<tr>
<td>9</td>
<td>1.00</td>
<td>0.88</td>
</tr>
</tbody>
</table>
Table 4: pH-stability profile (residual activity after 2 hours at 37°C)

<table>
<thead>
<tr>
<th>pH</th>
<th>S1 protease 1 from Saccharomonospora viridis</th>
<th>Protease 10R</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>0.46</td>
<td>0.78</td>
</tr>
<tr>
<td>3</td>
<td>1.02</td>
<td>1.03</td>
</tr>
<tr>
<td>4</td>
<td>1.06</td>
<td>0.99</td>
</tr>
<tr>
<td>5</td>
<td>1.05</td>
<td>1.00</td>
</tr>
<tr>
<td>6</td>
<td>1.02</td>
<td>1.03</td>
</tr>
<tr>
<td>7</td>
<td>1.05</td>
<td>1.01</td>
</tr>
<tr>
<td>8</td>
<td>1.06</td>
<td>0.98</td>
</tr>
<tr>
<td>9</td>
<td>1.03</td>
<td>0.99</td>
</tr>
<tr>
<td>10</td>
<td>1.00</td>
<td>0.99</td>
</tr>
<tr>
<td>11</td>
<td>1.06</td>
<td>0.86</td>
</tr>
</tbody>
</table>

Note: activities are residual activities relative to a sample, which was kept at stable conditions (5°C, pH 9.0).

Table 5: Temperature activity profile at pH 7.0 or pH 6.5

<table>
<thead>
<tr>
<th>Temp (°C)</th>
<th>S1 protease 1 from Saccharomonospora viridis (pH 7)</th>
<th>Protease 10R (pH 6.5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>0.00</td>
<td>0.01</td>
</tr>
<tr>
<td>25</td>
<td>0.06</td>
<td>0.02</td>
</tr>
<tr>
<td>37</td>
<td>0.20</td>
<td>0.06</td>
</tr>
<tr>
<td>50</td>
<td>0.57</td>
<td>0.13</td>
</tr>
<tr>
<td>60</td>
<td>0.93</td>
<td>0.35</td>
</tr>
</tbody>
</table>

Note: activities are relative to the optimal pH for the enzyme.
Table 6: \( P_1 \)-specificity on 10 Suc-AAPX-pNA substrates at pH 9.0 (25°C)

<table>
<thead>
<tr>
<th>Suc-AAPX-pNA</th>
<th>S1 protease 1 from Saccharomonospora viridis</th>
<th>Protease 10R</th>
</tr>
</thead>
<tbody>
<tr>
<td>Suc-AAPA-pNA</td>
<td>0.03</td>
<td>0.13</td>
</tr>
<tr>
<td>Suc-AAPR-pNA</td>
<td>0.04</td>
<td>0.09</td>
</tr>
<tr>
<td>Suc-AAPD-pNA</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Suc-AAPI-pNA</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Suc-AAPM-pNA</td>
<td>0.45</td>
<td>0.78</td>
</tr>
<tr>
<td>Suc-AAPV-pNA</td>
<td>0.00</td>
<td>0.01</td>
</tr>
<tr>
<td>Suc-AAPL-pNA</td>
<td>0.20</td>
<td>0.18</td>
</tr>
<tr>
<td>Suc-AAPE-pNA</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Suc-AAPK-pNA</td>
<td>0.02</td>
<td>0.08</td>
</tr>
<tr>
<td>Suc-AAPF-pNA</td>
<td>1.00</td>
<td>1.00</td>
</tr>
</tbody>
</table>

Note: activities are relative to the best substrate (Suc-AAPF-pNA) for the enzyme.

Other characteristics for the S1A protease 1 from *Saccharomonospora viridis*

Inhibitors: CI-2A and SSI.

Determination of the N-terminal sequence by EDMAN degradation was: MDVIGGN.

The relative molecular weight as determined by SDS-PAGE was approx. \( M_r \approx 20\text{kDa} \).

The molecular weight determined by intact molecular weight analysis was 16027.3Da.

The mature sequence (from mass spectrometry data and EDMAN degradation data and DNA sequence):

\[
\text{MDVIGGNAYMGNGGRCSVGVFTQGGFVTAGHCGTTGTSTSSPSGTFAAGSSFPGDYAFVR}
\text{TGSGDTRLPWNMYNGSARWSGSSVAPVGSSICRSSTTGGWHCGQVQAFNQTVRAYEGTV}
\text{TGLTRTNCAEPGDSGFSISGNQAGMTSGSGNCTF} \quad (\text{SEQ ID NO: 3})
\]

The calculated molecular weight from this mature sequence was 16027.4Da.
Example 4: Soybean-maize meal activity assay

An end-point assay using soybean-maize meal as substrate was used for obtaining the pH activity profile of the proteases at pH 3-7.

Substrate: Soybean meal-maize meal mixed in a 30:70 ratio.

Assay buffers: 9 buffers containing 100 mM succinic acid, 100 mM HEPES, 100 mM CHES, 100 mM CAPS, 1 mM CaCl2, 150 mM KCl, 0.01% Triton X-100 were prepared and adjusted using HCl or NaOH to a pH value such that after soybean-maize meal substrate (1 g) had been mixed with assay buffer (10 mL) to give a slurry, the final pH of the slurry was one of the following pH's: 3.0, 4.0, 5.0, 6.0, 7.0, 8.0, 9.0, 10.0 and 11.0.

Substrate slurry (2 mL) was mixed for 30 min before protease addition and incubation for 3 hours at 40°C (500 rpm). Protease (200 mg enzyme protein/kg dry matter) was dissolved in 100 μL 100 mM sodium acetate buffer (9.565 g/L NaOAc, 1.75 g/L acetic acid, 5 mM CaCl2, 0.01% BSA, 0.01% Tween20, pH 6.0) and added. Samples were centrifuged (10 min, 4000 rpm, 0°C) and the supernatants collected for analysis using the o-Phthalaldehyde (OPA) assay.

The results are shown in Table 7 below and figure 5. The proteolytic activity of the S1 protease 1 from *Saccharomonospora viridis* on soybean-maize meal increases with increasing pH from pH 3 to pH 7. The activity at pH 6-7 is somewhat higher than for protease 10R indicating that the S1 protease 1 from *Saccharomonospora viridis* might have the potential to be more efficient at protein hydrolysis in the small intestine of pigs and poultry where pH is around 7, in the crop of poultry where pH is in the range 4-6 and in the stomach of pigs where pH shortly after feeding can be as high as pH 6-7.

Table 7: Protease activity (OD₅₆₅ X dilution factor) on soybean-maize meal at pH 3.0, 4.0, 5.0, 6.0 and 7.0 (40°C).

<table>
<thead>
<tr>
<th>pH</th>
<th>S1 protease 1 from <em>Saccharomonospora viridis</em></th>
<th>Protease 10R</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Average</td>
<td>Standard deviation</td>
</tr>
<tr>
<td>3.0</td>
<td>0.13</td>
<td>0.01</td>
</tr>
<tr>
<td>4.0</td>
<td>0.37</td>
<td>0.03</td>
</tr>
<tr>
<td>5.0</td>
<td>0.79</td>
<td>0.08</td>
</tr>
<tr>
<td>6.0</td>
<td>2.00</td>
<td>0.04</td>
</tr>
<tr>
<td>7.0</td>
<td>3.32</td>
<td>0.23</td>
</tr>
</tbody>
</table>
Example 5: *In vitro* digestion assay

An *in vitro* digestion assay was used to evaluate the effect of the S1 protease 1 from *Saccharomonuspora viridis* on a feed substrate (soybean meal-maize meal mixed in a 30:70 ratio) in a setup designed to simulate digestion in monogastric animals.

The incubation process consisted of a gastric digestion phase with porcine pepsin (SP7000, Sigma-Aldrich, St. Louis, MO, USA) at pH 3 followed by a short duodenal incubation at pH 3.8 and a small intestinal incubation with pancreatin (8xUSB, P-7545, Sigma-Aldrich, St. Louis, MO, USA) at pH 7.0.

The *in vitro* digestion was performed using an automated system based on a Gilson liquid handler (Biolab, Denmark). For each sample 0.8 g feed was weighed into a tube and all tubes were placed in the liquid handler (40°C, 500 rpm). Additions of solutions as well as pH measurements were performed automatically. At time 0 min, 4.1 mL HCl (24 mM CaCl₂) was added to reach pH 3.0 in the solution. At time 30 min 0.5 mL HCl (24 mM CaCl₂, 3000 U pepsin/g feed) and 100 µL of a 100 mM sodium acetate buffer (258.6 g NaOAc per litre, 0.57% acetic acid, pH 6.0) was added. At time 90 min 900 µL NaOH was added to reach pH -3.8 and at time 120 min 400 µL of a 1 M NaHCO₃ solution containing 6.5 mg pancreatin/g feed was added leading to pH 6.8 in the solution. The pH was measured at time 30, 60, 90, 115, 120 and 180 min. The test proteases (100 mg enzyme protein/kg feed) were added via the 100 µL NaOAc buffer at time 30 min.

The level of soluble crude protein (N x 6.25) measured using a LECO FP-528 protein/nitrogen analyzer, was used as an indication of protease efficacy in the assay. Primary amines were analyzed using the o-Phthalaldehyde (OPA) assay and the absorbance values were used to calculate the degree of protein hydrolysis (DH) according to:

\[
DH (%) = 100 \times \frac{h}{htot},
\]

where \(h_{tot}\) is the total number of peptide bonds per protein equivalent, here the value for soy was used (7.8 g equivalents per kg protein) according to Adler-Nissen (J. Enzymic Hydrolysis of Food Proteins. Elsevier Applied Science Publishers. 1986). \(h\) is the number of hydrolyzed bonds expressed as:

\[
h = \frac{\text{serine-NH}_2 - \beta}{a \text{ meqv/g protein}},
\]

where \(a = 0.970\) and \(\beta = 0.342\) according to Adler-Nissen (Determination of the degree of hydrolysis of food protein hydrolysates by trinitrobenzenesulfonic acid. Journal of Agricultural and Food Chemistry, 27: 1256-1262. 1979). Serine-NH₂ is calculated as:
Serine - \(NH_2\) = \(\frac{(OD_{blank} - OD_{sample})}{(OD_{standard} - OD_{blank})}\) x 0.9516 meqv/L x 0.1 x 100 / X 

where serine-NH\(_2\) = meqv serine-NH\(_2\)/g protein; X = g sample; P = protein % in sample and 0.1 is the sample volume in litres (L).

The results are shown in tables 8 and 9 below. The S1 protease 1 from *Saccharomonomospora viridis* numerically increased the amount of soluble protein as well as the degree of protein hydrolysis in the samples indicating proteolytic activity of the S1 protease 1 from *Saccharomonomospora viridis* on top of the endogenous proteases in the assay. It was not possible to statistically differentiate between the two proteases and hence they must be considered to act equally good in the assay indicating also an equal potential to hydrolyze protein *in vivo*.

Table 8: The level of soluble protein as percent of total protein *in vitro* digestion samples after treatment with S1 protease 1 from *Saccharomonomospora viridis* or protease 10R

<table>
<thead>
<tr>
<th>Enzyme (mg enzyme protein/kg feed)</th>
<th>Soluble protein of total (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Average</td>
</tr>
<tr>
<td>No enzyme</td>
<td>91.30(^b)</td>
</tr>
<tr>
<td>Saccharomonomospora viridis (100)</td>
<td>94.41(^{ab})</td>
</tr>
<tr>
<td>Protease 10R (100)</td>
<td>96.18(^a)</td>
</tr>
</tbody>
</table>

\(^a\)\(^b\) Values not connected by the same superscript letters are statistically different (P<0.05) as determined by the Tukey Kramer test (a = 0.05) provided by the ANOVA procedure (SAS Institute Inc.).

Table 9: Degree of protein hydrolysis (DH) *in vitro* digestion samples after treatment with S1 protease 1 from *Saccharomonomospora viridis* or protease 10R

<table>
<thead>
<tr>
<th>Enzyme (mg enzyme protein/kg feed)</th>
<th>DH (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Average</td>
</tr>
<tr>
<td>No enzyme</td>
<td>29.90(^{bc})</td>
</tr>
<tr>
<td>Saccharomonomospora viridis (100)</td>
<td>30.38(^{ab})</td>
</tr>
<tr>
<td>Protease 10R (100)</td>
<td>31.90(^a)</td>
</tr>
</tbody>
</table>

\(^a\)\(^b\)\(^c\) Values not connected by the same superscript letters are statistically different (P<0.05) as determined by the Tukey Kramer test (a = 0.05) provided by the ANOVA procedure (SAS Institute Inc.).
Example 6: Proteolytic activity on crop, gizzard and ileum digesta

Crop, gizzard and ileum digesta material from 21 day old broiler chickens fed a corn-soy diet was collected; freeze dried and ground using a small coffee mill. The ground samples were suspended (47% w/v) in the following buffers and left to hydrate at 4°C over night (no stirring):

- **Crop buffer:** 100 mM HEPES, 1 mM CaCl₂·2 H₂O, 150 mM KCl, 0.01% Triton X-100, adjusted to pH 5 using HCl
- **Gizzard buffer:** 100 mM succinic acid, 1 mM CaCl₂·2 H₂O, 150 mM KCl, 0.01% Triton X-100, adjusted to pH 1.67 using HCl
- **Ileum buffer:** 100 mM HEPES, 1 mM CaCl₂·2 H₂O, 150 mM KCl, 0.01% Triton X-100, adjusted to pH 7.2 using HCl

The resulting pH after hydration over night was: pH 5 in crop samples; pH 3 in gizzard samples; and pH 7 in ileum samples. The suspensions were brought to 40°C and dispensed into test tubes. Three tubes representing blank (T₀) were immediately centrifuged (3000 x g, 0°C, 10 min) and supernatants frozen. Either enzyme (200 mg enzyme protein/kg substrate) in 50 µL 100 mM sodium acetate buffer (9.565 g/l NaOAc, 1.75 g/l acetic acid, 5 mM CaCl₂, 0.01% BSA, 0.01% Tween20, pH 6.0) or just sodium acetate buffer (50 µL) for the Blank samples was added to the tubes and the crop and ileum samples were incubated for 3 hours (T₃) while the gizzard samples were incubated for 1 hour (T₁) at 40°C while shaking (500 rpm). The samples were centrifuged (3000 x g, 0°C, 10 min) and supernatants recovered and frozen. The proteolytic activity was determined by analyzing primary amines using the o-phthaldialdehyde (OPA) assay.

The results are shown in Table 10. For each of the digesta types (crop, gizzard and ileum) there was a significant difference between the level of primary amines in the blank T₀ sample and the blank samples incubated for 1 or 3 hours (Table 9). This difference can be ascribed to activity of proteases present in the substrate and originating from either the diet raw materials or the animal. During incubation of the crop digesta the S₁ protease 1 from *Saccharomonospora viridis* further increased the level of primary amines compared to the blank sample incubated 3 hours, demonstrating that the protease had a proteolytic activity on this substrate under the given conditions. It was not possible to distinguish between the activity of the S₁ protease 1 from *Saccharomonospora viridis* and Protease 10R indicating that both proteases have an equal potential to degrade protein in the crop of broilers. No proteolytic effect could be shown on gizzard digesta, which was also not expected due to the pH activity properties of the proteases. Using ileum digesta as substrate a numerical effect of both the S₁ protease 1 from *Saccharomonospora viridis* and Protease 10R was shown indicating that both
proteases might be able to degrade protein which has not been digested and utilized by the broiler chicken.

Table 10: Proteolytic activity of S1 protease 1 from Saccharomonospora viridis compared to Protease 10R when incubated with broiler digesta and expressed as level of primary amines measured by the OPA assay (OD\textsubscript{340} x dilution factor).

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Crop (3 hours)</th>
<th>Gizzard (1 hour)</th>
<th>Ileum (3 hours)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blank T\textsubscript{0}</td>
<td>2.21 ± 0.02\textsuperscript{c}</td>
<td>2.95 ± 0.02\textsuperscript{b}</td>
<td>9.37 ± 0.08\textsuperscript{b}</td>
</tr>
<tr>
<td>Blank</td>
<td>3.54 ± 0.02\textsuperscript{b}</td>
<td>3.85 ± 0.08\textsuperscript{a}</td>
<td>14.40 ± 1.03\textsuperscript{a}</td>
</tr>
<tr>
<td>Saccharomonospora viridis</td>
<td>3.77 ± 0.02\textsuperscript{a}</td>
<td>3.78 ± 0.06\textsuperscript{a}</td>
<td>14.83 ± 0.45\textsuperscript{a}</td>
</tr>
<tr>
<td>Protease 10R</td>
<td>3.85 ± 0.09\textsuperscript{a}</td>
<td>3.87 ± 0.21\textsuperscript{a}</td>
<td>14.74 ± 0.12\textsuperscript{a}</td>
</tr>
</tbody>
</table>

\textsuperscript{a,b,c} Values within a column that are not connected by the same superscript letters are statistically different as determined by the Tukey Kramer test (a=0.05) provided by the ANOVA procedure (SAS Institute Inc.).

Example 7: AMSA wash performance of S1 protease 1 from Saccharomonospora viridis using a liquid and power detergent

The wash performance of S1 protease 1 from Saccharomonospora viridis was tested using a liquid detergent and a powder detergent at 2 different wash temperatures on 3 different technical stains using the Automatic Mechanical Stress Assay.

The experiments were conducted as described in the AMSA for laundry method using a single cycle wash procedure, with the detergent composition and swatches described in table 2 and the experimental conditions as specified in table 11 below.

Table 11: Experimental conditions for AMSA for tables 12 and 13

<table>
<thead>
<tr>
<th>Test solution</th>
<th>2.5 g/L powder model detergent A or 8 g/L liquid model detergent B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test solution volume</td>
<td>160 micro L</td>
</tr>
<tr>
<td>pH</td>
<td>As is</td>
</tr>
<tr>
<td>Wash time</td>
<td>20 minutes</td>
</tr>
<tr>
<td>Temperature</td>
<td>20°C or 40°C</td>
</tr>
<tr>
<td>Water hardness</td>
<td>15°dH</td>
</tr>
<tr>
<td>Protease concentration</td>
<td>0 (blank) or 30 nM</td>
</tr>
</tbody>
</table>
Water hardness was adjusted to 15°dH by addition of \( \text{CaCl}_2 \), \( \text{MgCl}_2 \), and \( \text{NaHCO}_3 \) (\( \text{Ca}^{2+} : \text{Mg}^{2+} : \text{CO}_3^{2-} = 4:1:7.5 \)) to the test system. After washing the textiles were flushed in tap water and dried.

<table>
<thead>
<tr>
<th>Swatch</th>
<th>Detergent A (2.5 g/L at 20°C)</th>
<th>Detergent B (8 g/L at 20°C)</th>
<th>Detergent A (2.5 g/L at 40°C)</th>
<th>Detergent B (8 g/L at 40°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PC-03</td>
<td>9</td>
<td>8</td>
<td>36</td>
<td>30</td>
</tr>
<tr>
<td>C-10</td>
<td>10</td>
<td>7</td>
<td>28</td>
<td>22</td>
</tr>
<tr>
<td>PC-05</td>
<td>43</td>
<td>37</td>
<td>40</td>
<td>70</td>
</tr>
</tbody>
</table>

The results show that detergent containing S1 protease 1 from *Saccharomonospora viridis* is more effective at removing stains compared to detergent without any protease. S1 protease 1 from *Saccharomonospora viridis* is also very effective at removing blood/milk/ink stains even at 20°C.

<table>
<thead>
<tr>
<th>Swatch</th>
<th>Detergent A (2.5 g/L at 20°C)</th>
<th>Detergent B (8 g/L at 20°C)</th>
<th>Detergent A (2.5 g/L at 40°C)</th>
<th>Detergent B (8 g/L at 40°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PC-03</td>
<td>108%</td>
<td>120%</td>
<td>118%</td>
<td>112%</td>
</tr>
<tr>
<td>C-10</td>
<td>128%</td>
<td>154%</td>
<td>105%</td>
<td>98%</td>
</tr>
<tr>
<td>PC-05</td>
<td>92%</td>
<td>103%</td>
<td>107%</td>
<td>104%</td>
</tr>
</tbody>
</table>

The results show that detergent containing S1 protease 1 from *Saccharomonospora viridis* is generally more effective at removing stains compared to detergent containing protease 10R, and is especially more effective at removing oil/milk/pigment stains at 20°C.

**Example 8: AMSA wash performance of S1 protease 1 from *Saccharomonospora viridis* in different water hardness's and protease concentrations using a liquid detergent**

The wash performance of S1 protease 1 from *Saccharomonospora viridis* was tested using a liquid detergent in 3 different water hardnesses and 2 different enzyme concentrations on 3 different technical stains using the Automatic Mechanical Stress Assay.
The experiments were conducted as described in the AMSA for laundry method using a single cycle wash procedure, with the detergent composition and swatches described in table 2 and the experimental conditions as specified in table 14 below.

**Table 14: Experimental conditions for AMSA for tables 15, 16 and 17**

<table>
<thead>
<tr>
<th>Test solution</th>
<th>2 g/L liquid model detergent B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test solution volume</td>
<td>160 micro L</td>
</tr>
<tr>
<td>pH</td>
<td>As is</td>
</tr>
<tr>
<td>Wash time</td>
<td>20 minutes</td>
</tr>
<tr>
<td>Temperature</td>
<td>40°C</td>
</tr>
<tr>
<td>Protease concentration</td>
<td>0 (blank), 5 nM or 30 nM</td>
</tr>
<tr>
<td>Swatch</td>
<td>EMPA117EH, PC-03, C-10</td>
</tr>
</tbody>
</table>

Water hardness was adjusted to 6, 16 or 24°dH by addition of CaCl$_2$ and MgCl$_2$ (Ca$^{2+}$:Mg$^{2+}$ = 5:1) to the test system. After washing the textiles were flushed in tap water and dried.

**Table 15: Delta intensity enzyme value of detergent containing S1 protease 1 from *Saccharomonospora viridis* or Savinase compared to detergent without protease on EMPA117EH swatches at 40°C**

<table>
<thead>
<tr>
<th>Enzyme conc.</th>
<th>5nM</th>
<th>5nM</th>
<th>5nM</th>
<th>30nM</th>
<th>30nM</th>
<th>30nM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water hardness</td>
<td>6°dH</td>
<td>16°dH</td>
<td>24°dH</td>
<td>6°dH</td>
<td>16°dH</td>
<td>24°dH</td>
</tr>
<tr>
<td>Savinase</td>
<td>-1</td>
<td>2</td>
<td>14</td>
<td>21</td>
<td>21</td>
<td>51</td>
</tr>
<tr>
<td><em>Saccharomonospora viridis</em></td>
<td>28</td>
<td>21</td>
<td>17</td>
<td>67</td>
<td>58</td>
<td>52</td>
</tr>
</tbody>
</table>

The results show that detergent containing S1 protease 1 from *Saccharomonospora viridis* is especially effective at removing blood/milk/ink on cotton/polyester stains in low to medium water hardnesses both compared to detergent without protease and to detergent containing Savinase.
The results show that detergent containing \textit{S1} protease 1 from \textit{Saccharomonospora viridis} is especially effective at removing chocolate-milk/ink on cotton/polyester stains in low to medium water hardnesses both compared to detergent without protease and to detergent containing Savinase.

Example 9: Evaluation of the Stability of \textit{S1} protease 1 from \textit{Saccharomonospora viridis} in liquid detergent using AMSA

The stability of the \textit{S1} protease 1 from \textit{Saccharomonospora viridis} in detergent was tested by examining the wash performance of the detergent with protease using an Automatic Mechanical Stress Assay at 2 different wash temperatures. 3 different stability conditions were tested, which are:

- the protease was added to the detergent composition immediately before wash;
- the protease was pre-incubated with the detergent for 48 hours at 25°C; and
The wash liquor was pre-incubated for 30 minutes at 40°C before starting the wash.

The experiments were conducted as described in the Automatic Mechanical Stress Assay (AMSA) for laundry method using a single cycle wash procedure, with the detergent composition and swatches described in table 2 and the experimental conditions as specified in table 18 below.

Table 18: Experimental conditions for AMSA for table 19

<table>
<thead>
<tr>
<th>Test solution</th>
<th>8 g/L liquid model detergent B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test solution volume</td>
<td>160 micro L</td>
</tr>
<tr>
<td>pH</td>
<td>As is</td>
</tr>
<tr>
<td>Wash time</td>
<td>20 minutes</td>
</tr>
<tr>
<td>Temperature</td>
<td>20°C or 40°C</td>
</tr>
<tr>
<td>Water hardness</td>
<td>15°dH</td>
</tr>
<tr>
<td>Protease concentration</td>
<td>0 (blank) or 30 nM</td>
</tr>
<tr>
<td>Swatch</td>
<td>PC-05</td>
</tr>
</tbody>
</table>

Water hardness was adjusted to 15°dH by addition of CaCl₂, MgCl₂, and NaHCO₃ (Ca²⁺:Mg²⁺:C0₃²⁻ = 4:1:7.5) to the test system. After washing the textiles were flushed in tap water and dried.

Table 19: Delta intensity value of detergent containing S₁ protease 1 from *Saccharomonospora viridis* compared to detergent without protease on a PC-05 swatch

<table>
<thead>
<tr>
<th></th>
<th>Δ Wash performance at 20°C</th>
<th>Δ Wash performance at 40°C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fresh enzyme</td>
<td>½-hr pre incubation at 40°C</td>
</tr>
<tr>
<td><em>Saccharomonospora viridis</em></td>
<td>33 ± 9</td>
<td>41 ± 1</td>
</tr>
</tbody>
</table>

The results show that detergent containing S₁ protease 1 from *Saccharomonospora viridis* has the same wash performance after 48 hours storage at 20°C in liquid detergent as the fresh enzyme which is added to the detergent immediately prior to the wash. This shows that
under these conditions the S1 protease 1 from *Saccharomonospora viridis* shows detergent stability.

Moreover, the results show that detergent containing S1 protease 1 from *Saccharomonospora viridis* has the same wash performance after a 30 minutes pre-incubation of the wash liquor at 40°C as wash liquor prepared with fresh enzyme added to the detergent immediately prior to the wash. This shows that under these conditions the S1 protease 1 from *Saccharomonospora viridis* shows in-wash stability.

**Example 10: Thermostability**

An aliquot of the protein sample of protease (purified as described in Example 2) is either desalted or buffer-changed into 20 mM Na-acetate, pH 4.0 using a prepacked PD-10 column or dialysed against 2 x 500 ml 20 mM Na-acetate, pH 4.0 at 4°C in a 2-3h step followed by an overnight step. The sample is 0.45 μm filtered and diluted with buffer to approx. 2 A280 units. The dialysis buffer is used as reference in Differential Scanning Calorimetry (DSC). The samples are degassed using vacuum suction and stirring for approx. 10 minutes.

A DSC scan is performed on a MicroCal VP-DSC at a constant scan rate of 1.5 °C/min from 20-90 °C. Data-handling is performed using the MicroCal Origin software (version 4.10), and the denaturation temperature, T_d (also called the melting temperature, T_m) is defined as the temperature at the apex of the peak in the thermogram.

**Example 11: Steam stability**

Residual activity of the protease after steam treatment may be evaluated using the following assay.

In these experiments a modified set-up is used whereby the steam is provided from a steam generator and led into the box. The samples placed on a plate are inserted into the box through a drawer when the temperature has reached ca. 93-94°C. Upon the insertion of the samples the temperature drops 4 °C. Incubation is performed for 30 seconds while the temperature remains approximately constant at 90°C. Thereafter the plate is quickly removed from the box, the samples placed on ice, re-suspended and evaluated with respect to protease activity using the Suc-AAPF-pNA or o-Phthaldialdehyde (OPA) assay. Each enzyme sample is compared to a similar sample that had not been steam treated in order to calculate residual activity.

**Example 12: Pelleting stability tests**

The enzyme granulation is performed in a manner as described in US No. patent 4,106,991, Example 1. The obtained granulate is 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
is coated with palm oil and calcium carbonate in a manner as described in US patent No. 4,106,991, Example 22.

Approximately 50 g enzyme granulate is pre-mixed with 10 kg feed for 10 minutes in a small horizontal mixer. This premix is mixed with 90 kg feed for 10 minutes in a larger horizontal mixer. From the mixer the feed is led to the conditioner (a cascade mixer with steam injection) at a rate of approximately 300 kg/hour. The conditioner heats up the feed to 95°C (measured at the outlet) by injecting steam. The residence time in the conditioner is 30 seconds. From the conditioner the feed is 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 are placed in an air cooler and cooled for 15 minutes.

The protease activity is measured using the Suc-AAPF-pNA assay prior to pelleting and in the feed pellets after pelleting. Pelleting stability is determined by comparing the protease activity in pelleted feed relative to the activity in non-pelleted feed.
What is claimed is:

1. Use of an isolated polypeptide having protease activity, selected from the group consisting of:
   
   (a) a polypeptide having at least 80%, e.g. at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% sequence identity to the polypeptide of SEQ ID NO: 3;
   
   (b) a polypeptide encoded by a polynucleotide that hybridizes under medium stringency conditions, medium-high stringency conditions, high stringency conditions or very-high stringency conditions with:
      
      (i) the mature polypeptide coding sequence of SEQ ID NO: 1, and/or
      
      (ii) the full-length complementary strand of (i);
   
   (c) a polypeptide encoded by a polynucleotide having at least 80%, e.g. at least 85%, at least 86%, at least 87%, at least 88%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% sequence identity to the mature polypeptide coding sequence of SEQ ID NO: 1;
   
   (d) a variant of the polypeptide of SEQ ID NO: 3 comprising a substitution, deletion, and/or insertion of one or more (e.g. several) positions; and
   
   (e) a fragment of the polypeptide of (a), (b), (c), or (d) that has protease activity in animal feed and detergent compositions.

2. The use according to claim 1, wherein the polypeptide comprises or consists of SEQ ID NO: 2.

3. The use according to claim 1, wherein the polypeptide comprises or consists of SEQ ID NO: 3.

4. The use according to any of claims 1-3, which is a variant of SEQ ID NO: 2 or SEQ ID NO: 3 comprising a substitution, deletion, and/or insertion of one or more (e.g. several) amino acids.
5. A variant polypeptide having protease activity and having at least 85%, e.g. at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% sequence identity to SEQ ID NO: 3 comprising at least one substitution, deletion, and/or insertion of at least one or more (several) amino acids of SEQ ID NO: 3.

6. A composition comprising an isolated polypeptide having protease activity, selected from the group consisting of:
   (a) a polypeptide having at least 80%, e.g., at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 3;
   (b) a polypeptide encoded by a polynucleotide that hybridizes under medium stringency conditions, medium-high stringency conditions, high stringency conditions or very-high stringency conditions with:
       (i) the mature polypeptide coding sequence of SEQ ID NO: 1; and/or
       (ii) the full-length complementary strand of (i);
   (c) a polypeptide encoded by a polynucleotide having at least 80%, e.g., at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to the mature polypeptide coding sequence of SEQ ID NO: 3;
   (d) a variant comprising a substitution, deletion, and/or insertion of one or more (several) amino acids of SEQ ID NO: 3; and
   (e) a fragment of a polypeptide of (a), (b), (c) or (d), that has protease activity.

7. The composition of claim 6, wherein the polypeptide comprises or consists of SEQ ID NO: 2.

8. The composition of claim 6, wherein the polypeptide comprises or consists of SEQ ID NO: 3.

9. The composition of any of claims 6-8, which is a variant of SEQ ID NO: 2 or SEQ ID NO: 3 comprising a substitution, deletion, and/or insertion of one or more (e.g. several) amino acids.
10. An isolated polynucleotide encoding the polypeptide indicated in any of claims 1-9 with the proviso that it is not 100% identical to SEQ ID NO: 1 or the mature polypeptide coding part thereof.

11. A nucleic acid construct or expression vector comprising the polynucleotide of claim 10 operably linked to one or more (several) control sequences that direct the production of the polypeptide in an expression host cell.

12. A recombinant expression host cell comprising a polynucleotide of claim 11 operably linked to one or more control sequences that direct the production of the polypeptide.

13. The host cell of claim 12, wherein the host is a bacterium, such as a Bacillus or Streptomyces; a fungus, such as Aspergillus, or a yeast, such as Candida, Kluyveromyces, Pichia, Saccharomyces, Schizosaccharomyces or Yarrowia.

14. The host cell of claim 13, wherein the host is a Bacillus, such as Bacillus subtilis, B. licheniformis, B. clausii, B. megaterium, B. pumilus, B. stearothermophilus, or B. Thuringiensis.

15. The host cell of claim 13, wherein the host is a Streptomyces, such as Streptomyces lividans, Streptomyces coelicolor, Streptomyces avermitilis, or Streptomyces griseus.

16. A polypeptide having protease activity and encoded by a polynucleotide according to claim 10 or produced by a nucleic acid construct or host cell of any of claims 11 to 15 with the proviso that the polypeptide is not 100% identical to SEQ ID NO: 3.

17. A method of producing the polypeptide indicated in any of claims 1 to 9, comprising:
   (a) cultivating a cell, which in its wild-type form produces the polypeptide, under conditions conducive for production of the polypeptide; and
   (b) recovering the polypeptide.

18. A method of producing the polypeptide indicated in any of claims 1 to 9, comprising:
   (a) cultivating a host cell of any of claims 9 to 12 under conditions conducive for production of the polypeptide; and
   (b) recovering the polypeptide.

19. Use of at least one polypeptide indicated in any one of claims 1 to 9;
in animal feed;
in animal feed additives;
in the preparation of a composition for use in animal feed;
for improving the nutritional value of an animal feed;
for increasing digestible and/or soluble protein in animal feed;
for increasing the degree of hydrolysis of proteins in animal diets; and/or
for the treatment of proteins.

20. A method for improving the nutritional value of an animal feed, wherein at least one polypeptide indicated in any one of claims 1 to 9 is added to the feed.

21. An animal feed additive comprising
   at least one polypeptide indicated in any one of claims 1 to 9; and
   at least one fat-soluble vitamin, and/or
   at least one water-soluble vitamin, and/or
   at least one trace mineral.

22. The animal feed additive of claim 21, wherein the animal feed comprises one or more further enzymes, wherein the further enzymes are selected from the group comprising of amylases; phytases; xylanases; galactanases; alpha-galactosidases; proteases, phospholipases; and beta-glucanases, or any mixture thereof.

23. An animal feed having a crude protein content of 50 to 800 g/kg and comprising at least one polypeptide indicated in any one of claims 1 to 9.

24. A method for the treatment of proteins, comprising the step of adding at least one polypeptide indicated in any one of claims 1 to 9 to at least one protein or protein source.

25. The method of claim 24, wherein soybean or soybean meal is included amongst the at least one protein source.

26. A detergent composition comprising at least one polypeptide indicated in claims 1 to 9 and one or more detergent components.

27. The detergent composition of claim 26 for use in laundry, laundering, hard surface cleaning and/or dish wash.
28. The detergent composition of any of claims 26 to 27, wherein the composition comprises one or more further enzymes selected from the group comprising of proteases, amylases, lipases, cutinases, cellulases, endoglucanases, xyloglucanases, pectinases, pectin lyases, xanthanases, peroxidaes, haloperoxygenases, catalases and mannanases, or any mixture thereof.
Figure 3

Figure 4
Figure 5

Protease 10R

S1 protease 1 from Saccharomonospora viridis