Title: SUBTILASE ENZYMES OF THE I-S1 AND I-S2 SUB-GROUPS HAVING AT LEAST ONE ADDITIONAL AMINO ACID RESIDUE BETWEEN POSITIONS 97 AND 98

Abstract: Subtilases of the I-S1 and I-S2 sub-groups having an additional amino acid residue in position 97 of the active site loop (b) region from positions 95 to 103. Variant subtilases exhibit improved wash performance in a detergent in comparison to its parent enzyme.
Subtilase enzymes of the I-S1 and I-S2 sub-groups having at least one additional amino acid residue between positions 97 and 98

5 TECHNICAL FIELD

This invention relates to novel subtilase enzymes of the I-S1 and I-S2 sub-groups having at least one additional amino acid residue in position 97 of the active site loop (b) region from positions 95 to 103. These proteases are useful exhibiting excellent or improved wash performance when used in detergents; cleaning and detergent compositions. The invention further relates to genes coding for the expression of said enzymes when inserted into a suitable host cell or organism; and such host cells transformed therewith and capable of expressing said enzyme variants, and methods for producing the novel enzymes.

BACKGROUND OF THE INVENTION

In the detergent industry enzymes have for more than 30 years been implemented in washing formulations. Enzymes used in such formulations comprise proteases, lipases, amylases, cellulases, as well as other enzymes, or mixtures thereof. Commercially most important enzymes are proteases.

An increasing number of commercially used proteases are protein engineered variants of naturally occurring wild type proteases, e.g. DURAZYM® (Novo Nordisk A/S), RELASE® (Novo Nordisk A/S), MAXAPEM® (Gist-Brocades N.V.), PURAFECT® (Genencor International, Inc.).

Further a number of protease variants are described in the art, such as in EP 130756 (GENENTECH) (corresponding to US Reissue Patent No. 34,606 (GENENCOR)); EP 214435 (HENKEL); WO 87/04461 (AMGEN); WO 87/05050 (GENEX); EP 260105 (GENENCOR); Thomas, Russell, and Persht (1985) Nature 318 375-376; Thomas,
Russell, and Persht (1987) J. Mol. Biol. 193 803-813; Russel and Persht Nature 328 496-500 (1987); WO 88/08028 (Genex); WO 88/08033 (Amgen); WO 95/27049 (SOLVAY S.A.); WO 95/30011 (PROCTER & GAMBLE COMPANY); WO 95/30010 (PROCTER & GAMBLE COMPANY); US 5,543,302 (SOLVAY S.A.); EP 251 446 (GENENCOR); WO 89/06279 (NOVO NORDISK A/S); WO 91/00345 (NOVO NORDISK A/S); EP 525 610 A1 (SOLVAY); and WO 94/02618 (GIST-BROCADES N.V.).

However, even though a number of useful protease variants have been described, there is still a need for new improved proteases or protease variants for a number of industrial uses.

Therefore, an object of the present invention, is to provide improved proteases or protein engineered protease variants, especially for use in the detergent industry.

**SUMMARY OF THE INVENTION**

The present inventors have found that subtilisins wherein at least one of the active site loops are longer than those presently known, exhibit improved wash performance properties in detergent compositions. The identification thereof was done in constructing subtilisin variants, especially of the subtilisin 309 (BLSAVI or Savinase®), exhibiting improved wash performance properties in detergent compositions relative to the parent wild type enzyme. This has been described in our earlier application DK1332/97 (now published as WO 99/27082).

It has now been found that certain subtilases or variants thereof of the I-S1 (true “subtilisins”) and I-S2 (high alkaline subtilisins) sub-groups having at least one additional amino acid residue in position 97 (or rather between positions 97 and 98) of the active site loop (b) region from positions 95 to 103, exhibit surprisingly improved wash
performance in comparison to those presently known and those
described in said application. The position 97 insertion
variants described in WO 99/27082 include G97GASG, G97GAA,
G97GASG+A98S+S99G+G100A+S101A and G97GAA+A98S+S99G+S101T. These
variants are not within the scope of the present invention.

The improved proteases according to the invention may
be obtained by isolation from natural resources or by the
introduction of at least one further amino acid residue (an
insertion) in the active site loop (b) between positions 97 and
98 in a wild type subtilase (for a definition of the active
site loops and the numbering of positions see below).

Although this finding was done in subtilisin 309 it is
predicted that it will be possible to produce or isolate
similar advantageous subtilases or subtilase variants.

Furthermore it will be possible to specifically screen
natural isolates to identify novel wild type subtilases
comprising an active site loop (b) which is longer than the
corresponding active site loop in known wild type subtilases,
such as subtilisin 309, which subtilases can be considered to
have an inserted amino acid residue between positions 97 and
98, and exhibiting excellent wash performance in a detergent,
in comparison to their closest related known subtilisin, such
as subtilisin 309.

Concerning alignment and numbering reference is made to
Figs. 1, 1a, 2 and 2a below showing alignments between
subtilisin BPN’(a) (BASBPN) and subtilisin 309 (BLSAVI)(b), and
alignments between subtilisin BPN’(a) (BASBPN) and subtilisin
Carlsberg (g). These alignments are in this patent application
used as a reference for numbering the residues.

The seven active site loops (a) to (g) (including both
the end amino acid residues indicated) are here defined to
encompass the amino acid residues in the segments given below
(a) the region between amino acid residue 33 and 43;
(b) the region between amino acid residue 95 and 103;
(c) the region between amino acid residue 125 and 132;
(d) the region between amino acid residue 153 and 173;
(e) the region between amino acid residue 181 and 195;
(f) the region between amino acid residue 202 and 204;
(g) the region between amino acid residue 218 and 219.

Accordingly, in a first aspect the invention relates to an isolated (i.e. greater than 10 % pure) subtilase enzyme of the I-S1 and I-S2 sub-groups having at least one additional amino acid residue in position 97 of the active site loop (b) region from positions 95 to 103, whereby said additional amino acid residue(s) correspond to the insertion of at least one amino acid residue between positions 97 and 98, with the proviso that the subtilase enzyme does not contain the following insertions G97GASG, G97GAA, G97GAS, G97GAS+A98S+S99G, G97GGG+A98S+S99G, G97GAA+A98S+S99G, G97GASG+A98S+S99G+G100A+S101A and G97GAA+A98S+S99G+S101T.

In a second aspect the invention relates to an isolated DNA sequence encoding a subtilase variant of the invention.

In a third aspect the invention relates to an expression vector comprising an isolated DNA sequence encoding a subtilase variant of the invention.

In a fourth aspect the invention relates to a microbial host cell transformed with an expression vector according to the third aspect.

In a further aspect the invention relates to the production of the subtilisin enzymes of the invention.

The enzymes of the invention can generally be produced by either cultivation of a microbial strain from which the enzyme was isolated and recovering the enzyme in substantially pure form; or by inserting an expression vector according to
the third aspect of the invention into a suitable microbial host, cultivating the host to express the desired subtilase enzyme, and recovering the enzyme product.

Further the invention relates to a composition comprising a subtilase or subtilase variant of the invention.

Even further the invention relates to the use of the enzymes of the invention for a number of industrial relevant uses, in particular for use in cleaning compositions and cleaning compositions comprising the enzymes, especially detergent compositions comprising the subtilisin enzymes of the invention.

DEFINITIONS

Prior to discussing this invention in further detail, the following terms and conventions will first be defined.

**NOMENCLATURE OF AMINO ACIDS**

<table>
<thead>
<tr>
<th>A</th>
<th>= Ala</th>
<th>= Alanine</th>
</tr>
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<tbody>
<tr>
<td>V</td>
<td>= Val</td>
<td>= Valine</td>
</tr>
<tr>
<td>L</td>
<td>= Leu</td>
<td>= Leucine</td>
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<td>P</td>
<td>= Pro</td>
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<td>F</td>
<td>= Phe</td>
<td>= Phenylalanine</td>
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<td>W</td>
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<td>= Tryptophan</td>
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<tr>
<td>M</td>
<td>= Met</td>
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<td>G</td>
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<tr>
<td>N</td>
<td>= Asn</td>
<td>= Asparagine</td>
</tr>
<tr>
<td>Q</td>
<td>= Gln</td>
<td>= Glutamine</td>
</tr>
<tr>
<td>D</td>
<td>= Asp</td>
<td>= Aspartic Acid</td>
</tr>
</tbody>
</table>
E = Glu = Glutamic Acid
K = Lys = Lysine
R = Arg = Arginine
H = His = Histidine
5 X = Xaa = Any amino acid

NOMENCLATURE OF NUCLEIC ACIDS
A = Adenine
G = Guanine
10 C = Cytosine
T = Thymine (only in DNA)
U = Uracil (only in RNA)

NOMENCLATURE AND CONVENTIONS FOR DESIGNATION OF VARIANTS
15 In describing the various enzyme variants produced or contemplated according to the invention, the following nomenclatures and conventions have been adapted for ease of reference:

A frame of reference is first defined by aligning the isolated or parent wild type enzyme with subtilisin BPN’ (BASBPN).

The alignment can be obtained by the GAP routine of the GCG package version 9.1 to number the variants using the following parameters: gap creation penalty = 8 and gap extension penalty = 8 and all other parameters kept at their default values.

Another method is to use known recognised alignments between subtilisases, such as the alignment indicated in WO 91/00345. In most cases the differences will not be of any importance.

Such alignments between subtilisin BPN’ (BASBPN) and subtilisin 309 (BLSAVI) and subtilisin Carlsberg (BLSCAR), respectively are indicated in Figs. 1, 1a, 2, and 2a. By this a
number of deletions and insertions will be defined in relation to BASBPN. In Fig. 1 subtilisin 309 has 6 deletions in positions 36, 58, 158, 162, 163, and 164 in comparison to BASBPN, whereas in Fig. 1a subtilisin 309 has the same deletions in positions 36, 56, 159, 164, 165, and 166 in comparison to BASBPN. In Fig. 2 subtilisin Carlsberg has one deletion in position 58 in comparison to BASBPN, whereas in Fig. 2a subtilisin Carlsberg has the one deletion in position 56 in comparison to BASBPN. These deletions are in Figs. 1, 1a, 2, and 2a indicated by asterixes (*).

The various modifications performed in a wild type enzyme is indicated in general using three elements as follows:

Original amino acid Position Substituted amino acid

The notation G195E thus means a substitution of a glycine in position 195 with a glutamic acid.

Position Substituted amino acid

In the case when the original amino acid residue may be any amino acid residue, a short hand notation may at times be used indicating only the position and substituted amino acid, such a notation is particular relevant in connection with modification(s) in homologous subtilases.

Original amino acid Position

This notation can be used when the amino acid residue(s) replacing a specific amino acid in a position is immaterial.

Position

When both the original amino acid(s) and substituted amino acid(s) may comprise any amino acid, then only the position is indicated, e.g.: 170.
Original amino acid position \{substituted amino acid\sub_{1}, \ldots, substituted amino acid\sub_{n}\}

When the original amino acid(s) and/or substituted amino acid(s) may comprise more than one, but not all amino acid(s), then the selected amino acids are indicated inside brackets \{ \},

For specific variants the specific three or one letter codes are used, including the codes Xaa and X to indicate any amino acid residue.

SUBSTITUTIONS:
The substitution of Glutamic acid for glycine in position 195 is designated as:

\begin{verbatim}
Gly195Glu  or  G195E
\end{verbatim}

or the substitution of any amino acid residue acid for glycine in position 195 is designated as:

\begin{verbatim}
Gly195Xaa  or  G195X
\end{verbatim}
or

\begin{verbatim}
Gly195  or  G195
\end{verbatim}

The substitution of serine for any amino acid residue in position 170 would thus be designated

\begin{verbatim}
Xaa170Ser  or  X170S.
\end{verbatim}
or

\begin{verbatim}
170Ser  or  170S
\end{verbatim}

Such a notation is particular relevant in connection with modification(s) in homologous subtilases (vide infra).

170Ser is thus meant to comprise e.g. both a Lys170Ser modification in BASBPN and an Arg170Ser modification in BLSAVI (cf. Fig. 1).
For a modification where the original amino acid(s) and/or substituted amino acid(s) may comprise more than one, but not all amino acid(s), the substitution of glycine, alanine, serine or threonine for arginine in position 170 would be indicated by

\[\text{Arg170}\{\text{Gly, Ala, Ser, Thr}\} \quad \text{or} \quad \text{R170}\{\text{G, A, S, T}\}\]

to indicate the variants

R170G, R170A, R170S, and R170T.

**DELETIONS:**

A deletion of glycine in position 195 will be indicated by:

Gly195* or G195*

Correspondingly the deletion of more than one amino acid residue, such as the deletion of glycine and leucine in positions 195 and 196 will be designated

Gly195*+Leu196* or G195*+L196*

**INSERTIONS:**

The insertion of an additional amino acid residue such as e.g. a lysine after G195 is:

\[\text{Gly195GlyLys} \quad \text{or} \quad \text{G195GK}; \text{ or}\]

when more than one amino acid residue is inserted, such as e.g. a Lys, Ala and Ser after G195 this is:

Gly195GlyLysAlaSer or G195GKAS

In such cases the inserted amino acid residue(s) are numbered by the addition of lower case letters to the position number of the amino acid residue preceding the inserted amino acid residue(s). In the above example the sequences 194 to 196 would thus be:
In cases where an amino acid residue identical to the existing amino acid residue is inserted it is clear that a degeneracy in the nomenclature arises. If for example a glycine is inserted after the glycine in the above example this would be indicated by G195GG. The same actual change could just as well be indicated as A194AG for the change from 194 195 196

BLSAVI  A - G - L

194 195 195a 195b 195c 196

Variant  A - G - K - A - S - L

to

194 195 195a 196

BLSAVI  A - G - L

Variant  A - G - G - L

194 194a 195 196

Such instances will be apparent to the skilled person, and the indication G195GG and corresponding indications for this type of insertions are thus meant to comprise such equivalent degenerate indications.

Some times it is desired to both perform a modification and an insertion at the same time. This situation is also covered by the present definitions. Thus S130TP indicates that the serine in position 130 has been replaced by a tyrosine and a proline has been inserted between positions 130 and 131. A more cumbersome way of describing such a variant would be: S130SP+S130T.
FILLING A GAP:

Where a deletion in an enzyme exists in the reference comparison with the subtilisin BPN' sequence used for the numbering, an insertion in such a position is indicated as:

*36Asp or *36D

for the insertion of an aspartic acid in position 36.

MULTIPLE MODIFICATIONS

Variants comprising multiple modifications are separated by pluses, e.g.:

Arg170Tyr+Gly195Glu or R170Y+G195E

representing modifications in positions 170 and 195 substituting tyrosine and glutamic acid for arginine and glycine, respectively.

or e.g. Tyr167{Gly,Ala,Ser,Thr}+Arg170{Gly,Ala,Ser,Thr} designates the variants

Tyr167Gly+Arg170Gly, Tyr167Gly+Arg170Ala,
Tyr167Gly+Arg170Ser, Tyr167Gly+Arg170Thr,
Tyr167Ala+Arg170Gly, Tyr167Ala+Arg170Ala,
Tyr167Ala+Arg170Ser, Tyr167Ala+Arg170Thr,
Tyr167Ser+Arg170Gly, Tyr167Ser+Arg170Ala,
Tyr167Ser+Arg170Ser, Tyr167Ser+Arg170Thr,
Tyr167Thr+Arg170Gly, Tyr167Thr+Arg170Ala,
Tyr167Thr+Arg170Ser, and Tyr167Thr+Arg170Thr.

This nomenclature is particular relevant relating to modifications aimed at substituting, replacing, inserting or deleting amino acid residues having specific common properties, such as residues of positive charge (K, R, H), negative charge (D, E), or conservative amino acid modification(s) of e.g. Tyr167{Gly,Ala,Ser,Thr}+Arg170{Gly,Ala,Ser,Thr}, which signifies substituting a small amino acid for another small amino acid. See section "Detailed description of the invention" for further details.
Numbering of amino acid positions/residues

If nothing else is mentioned the amino acid numbering used herein corresponds to that of the subtilase BPN' (BASBPN) sequence. For further description of the BPN' sequence see Figs. 1 and 2, or Siezen et al., Protein Engng. 4 (1991) 719-737.

Proteases

Enzymes cleaving the amide linkages in protein substrates are classified as proteases, or (interchangeably) peptidases (see Walsh, 1979, Enzymatic Reaction Mechanisms, W.H. Freeman and Company, San Francisco, Chapter 3).

Serine proteases

A serine protease is an enzyme which catalyzes the hydrolysis of peptide bonds, and in which there is an essential serine residue at the active site (White, Handler and Smith, 1973 "Principles of Biochemistry," Fifth Edition, McGraw-Hill Book Company, NY, pp. 271-272).

The bacterial serine proteases have molecular weights in the 20,000 to 45,000 Dalton range. They are inhibited by diisopropylfluorophosphatate. They hydrolyze simple terminal esters and are similar in activity to eukaryotic chymotrypsin, also a serine protease. A more narrow term, alkaline protease, covering a sub-group, reflects the high pH optimum of some of the serine proteases, from pH 9.0 to 11.0 (for review, see Priest (1977) Bacteriological Rev. 41 711-753).

Subtilases

A sub-group of the serine proteases tentatively designated subtilases has been proposed by Siezen et al., Protein Engng. 4 (1991) 719-737 and Siezen et al. Protein Science 6 (1997) 501-523. They are defined by homology analysis of more than 170
amino acid sequences of serine proteases previously referred to as subtilisin-like proteases. A subtilisin was previously often defined as a serine protease produced by Gram-positive bacteria or fungi, and according to Siezen et al. now is a subgroup of the subtilases. A wide variety of subtilases have been identified, and the amino acid sequence of a number of subtilases has been determined. For a more detailed description of such subtilases and their amino acid sequences reference is made to Siezen et al. (1997).

One subgroup of the subtilases, I-S1 or "true" subtilisins, comprises the "classical" subtilisins, such as subtilisin 168 (BSS168), subtilisin BPN', subtilisin Carlsberg (ALCALASE®, NOVO NORDISK A/S), and subtilisin DY (BSSDY).

A further subgroup of the subtilases, I-S2 or high alkaline subtilisins, is recognised by Siezen et al. Sub-group I-S2 proteases are described as highly alkaline subtilisins and comprises enzymes such as subtilisin PB92 (BAALKP) (MAXACAL®, Gist-Brocades NV), subtilisin 309 (SAVINASE®, NOVO NORDISK A/S), subtilisin 147 (BLS147) (ESPERASE®, NOVO NORDISK A/S), and alkaline elastase YaB (BSEYAB).

"SAVINASE®"

SAVINASE® is marketed by NOVO NORDISK A/S. It is subtilisin 309 from B. Lentus and differs from BAALKP only in one position (N87S, see Fig. 1 herein). SAVINASE® has the amino acid sequence designated b) in Fig. 1.

Parent subtilase

The term "parent subtilase" describes a subtilase defined according to Siezen et al. (1991 and 1997). For further details see description of "SUBTILASES" immediately above. A parent subtilase may also be a subtilase isolated from a natural source, wherein subsequent modification have been made
while retaining the characteristic of a subtilase.
Alternatively the term "parent subtilase" may be termed "wild type subtilase".

Modification(s) of a subtilase variant

The term "modification(s)" used herein is defined to include chemical modification of a subtilase as well as genetic manipulation of the DNA encoding a subtilase. The modification(s) can be replacement(s) of the amino acid side chain(s), substitution(s), deletion(s) and/or insertions in or at the amino acid(s) of interest.

Subtilase variant

In the context of this invention, the term subtilase variant or mutated subtilase means a subtilase that has been produced by an organism which is expressing a mutant gene derived from a parent microorganism which possessed an original or parent gene and which produced a corresponding parent enzyme, the parent gene having been mutated in order to produce the mutant gene from which said mutated subtilase protease is produced when expressed in a suitable host.

Homologous subtilase sequences

Specific active site loop regions, and amino acid insertions in said loops of the subtilase SAVINASE® are identified for modification herein to obtain a subtilase variant of the invention.

However, the invention is not limited to modifications of this particular subtilase, but extends to other parent (wild-type) subtilases, which have a homologous primary structure to that of SAVINASE®. The homology between two amino acid sequences is in this context described by the parameter "identity".
In order to determine the degree of identity between two subtilases the GAP routine of the GCG package version 9.1 can be applied using the same settings as indicated above. The output from the routine is besides the amino acid alignment the calculation of the "Percent Identity" between the two sequences.

Based on this description it is routine for a person skilled in the art to identify suitable homologous subtilases and corresponding homologous active site loop regions, which can be modified according to the invention.

Wash performance

The ability of an enzyme to catalyze the degradation of various naturally occurring substrates present on the objects to be cleaned during e.g. wash or hard surface cleaning is often referred to as its washing ability, wash-ability, detergency, or wash performance. Throughout this application the term wash performance will be used to encompass this property.

Isolated DNA sequence

The term "isolated", when applied to a DNA sequence molecule, denotes that the DNA sequence has been removed from its natural genetic milieu and is thus free of other extraneous or unwanted coding sequences, and is in a form suitable for use within genetically engineered protein production systems. Such isolated molecules are those that are separated from their natural environment and include cDNA and genomic clones. Isolated DNA molecules of the present invention are free of other genes with which they are ordinarily associated, but may include naturally occurring 5' and 3' untranslated regions such as promoters and terminators. The identification of associated regions will be evident to one of ordinary skill in the art.
(see for example, Dynan and Tijan, *Nature* 316:774-78, 1985). The term "an isolated DNA sequence" may alternatively be termed "a cloned DNA sequence".

5 *Isolated protein*

When applied to a protein, the term "isolated" indicates that the protein has been removed from its native environment.

In a preferred form, the isolated protein is substantially free of other proteins, particularly other homologous proteins (i.e. "homologous impurities" (see below)).

An isolated protein is greater than 10 \% pure, preferably greater than 20 \% pure, more preferably greater than 30 \% pure, as determined by SDS-PAGE. Further it is preferred to provide the protein in a highly purified form, i.e., greater than 40\% pure, greater than 60\% pure, greater than 80\% pure, more preferably greater than 95\% pure, and even more preferably greater than 99\% pure, as determined by SDS-PAGE.

The term "isolated protein" may alternatively be termed "purified protein".

*Homologous impurities*

The term "homologous impurities" means any impurity (e.g. another polypeptide than the polypeptide of the invention) which originate from the homologous cell where the polypeptide of the invention is originally obtained from.

*Obtained from*

The term "obtained from" as used herein in connection with a specific microbial source, means that the polynucleotide and/or polypeptide is produced by the specific source, or by a cell in which a gene from the source has been inserted.
Substrate

The term "Substrate" used in connection with a substrate for a protease should be interpreted in its broadest form as comprising a compound containing at least one peptide bond susceptible to hydrolysis by a subtilisin protease.

Product

The term "product" used in connection with a product derived from a protease enzymatic reaction should in the context of this invention be interpreted to include the products of a hydrolysis reaction involving a subtilase protease. A product may be the substrate in a subsequent hydrolysis reaction.

BRIEF DESCRIPTION OF THE DRAWING

Fig. 1 shows an alignment between subtilisin BPN' (a) and Savinase® (b) using the GAP routine mentioned above.

Fig. 1a shows the alignment between subtilisin BPN' and Savinase® as taken from WO 91/00345.

Fig. 2 shows an alignment between subtilisin BPN' and subtilisin Carlsberg using the GAP routine mentioned above.

Fig. 2a shows the alignment between subtilisin BPN' and subtilisin Carlsberg as taken from WO 91/00345.

Fig. 3 shows the three dimensional structure of Savinase (Protein data bank (PDB) entry 1SVN). In the Figure the active site loop (b) is indicated.

DETAILED DESCRIPTION OF THE INVENTION

The subtilases of the invention in a first aspect relates to an isolated (i.e. greater than 10 % pure) subtilase enzyme of the I-S1 and I-S2 sub-groups having at least one additional amino acid residue in position 97 of the active site loop (b) region from positions 95 to 103, whereby said
additional amino acid residue(s) correspond to the insertion of at least one amino acid residue between positions 97 and 98.

In other words the subtilases of the invention are characterized by comprising an active site loop (b) region of more than 9 amino acid residue and wherein the additional amino acid residue is or can be considered as being inserted between positions 97 and 98 as compared to the parent or a known wild type subtilase.

A subtilase of the first aspect of the invention may be a parent or wildtype subtilase identified and isolated from nature.

Such a parent wildtype subtilase may be specifically screened for by standard techniques known in the art.

One preferred way of doing this may be by specifically PCR amplify DNA regions known to encode active site loops in subtilases from numerous different microorganism, preferably different Bacillus strains.

Subtilases are a group of conserved enzymes, in the sense that their DNA and amino acid sequences are homologous. Accordingly it is possible to construct relatively specific primers flanking active site loops.

One way of doing this is by investigating an alignment of different subtilases (see e.g. Siezen et al. Protein Science 6 (1997) 501-523). It is from this routine work for a person skilled in the art to construct PCR primers flanking the active site loop corresponding to the active site loop (b) between amino acid residue 95 to 103 in any of the groups I-S1 or I-S2, such as from BLASAVI. Using such PCR primers to amplify DNA from a number of different microorganism, preferably different Bacillus strains, followed by DNA sequencing of said amplified PCR fragments, it will be possible to identify strains which produce subtilases of these groups comprising a longer, as compared to e.g. BLASAVI, active site region corresponding to
the active site loop region from positions 95 to 103, and where an insertion can be considered to exist between positions 97 and 98. Having identified the strain and a partial DNA sequence of such a subtilase of interest, it is routine work for a person skilled in the art to complete cloning, expression and purification of such a subtilase of the invention.

However, it is envisaged that a subtilase enzyme of the invention predominantly is a variant of a parent subtilase.

Accordingly, in one embodiment the invention relates to an isolated subtilase enzyme according to the first aspect of the invention, wherein said subtilase enzyme is a constructed variant having a longer active site loop (b) than its parent enzyme by having at least one amino acid insertion between amino acid residues 97 and 98.

The subtilases of the invention exhibit excellent wash performance in a detergent, and if the enzyme is a constructed variant an improved wash performance in a detergent in comparison to its closest related subtilase, such as subtilisin 309.

Different subtilase products will exhibit a different wash performance in different types of detergent compositions. A subtilase of the invention has improved wash performance, as compared to its closest relative in a majority of such different types of detergent compositions.

Preferably a subtilase enzyme of the invention has improved wash performance, as compared to its closest relative in the detergent compositions indicated in Example 3 herein.

In order to determine if a given subtilase amino acid sequence (irrelevant whether said subtilase sequence is a parent wildtype subtilase sequence or a subtilase variant sequence produced by any other method than by site directed mutagenesis) is within the scope of the invention, the following procedure may be used:
i) align said subtilase sequence to the amino acid sequence of subtilisin BPN';

ii) based on the alignment performed in step i) identify the active site loop (b), in said subtilase sequence corresponding to the active site loop (b) region of subtilisin BPN' comprising the region (both of the end amino acids included) between amino acid residue from 95 to 103;

iii) determine if the active site loop (b) in said subtilase sequence, identified in step ii) is longer than the corresponding active site loop in subtilisin BPN' and if said prolongation corresponds to the insertion of at least one amino acid residue between positions 97 and 98.

If this is the case the subtilase investigated is a subtilase within the scope of the present invention.

The alignment performed in step i) above is performed as described above by using the GAP routine.

Based on this description it is routine for a person skilled in the art to identify the active site loop (b) in a subtilase and determine if the subtilase in question is within the scope of the invention. If a variant is constructed by site directed mutagenesis, it is of course known beforehand if the subtilase variant is within the scope of the invention.

A subtilase variant of the invention may be constructed by standard techniques known in the art such as by site-directed/random mutagenesis or by DNA shuffling of different subtilase sequences. See sections "PRODUCING A SUBTILASE VARIANT" and "Materials and methods" for further details.

In further embodiments the invention relates to 1.an isolated subtilase enzyme according to the invention, wherein said at least one inserted amino acid residue is chosen from the group comprising: T,G,A, and S;
2. an isolated subtilase enzyme according to the invention, wherein said at least one inserted amino acid residue is chosen from the group of charged amino acid residues comprising: D, E, H, K, and R, more preferably D, E, K and R;

3. an isolated subtilase enzyme according to the invention, wherein said at least one inserted amino acid residue is chosen from the group of hydrophilic amino acid residues comprising: C, N, Q, S and T, more preferably N, Q, S and T;

4. an isolated subtilase enzyme according to the invention, wherein said at least one inserted amino acid residue is chosen from the group of small hydrophobic amino acid residues comprising: A, G and V; or

5. an isolated subtilase enzyme according to the invention, wherein said at least one inserted amino acid residue is chosen from the group of large hydrophilic amino acid residues comprising: F, I, L, M, P, W and Y, more preferably F, I, L, M, and Y.

In a further embodiment, the invention relates to an isolated subtilase enzyme according to the invention, wherein said insertion between positions 97 and 98 comprises at least two amino acids, as compared to the corresponding active site loop in subtilisin BPN’.

In further embodiments the invention relates to an isolated subtilase enzyme comprising at least one insertion, chosen from the group comprising (in BASBPN numbering):

X97X{T, G, A, S}
X97X{D, E, K, R}
X97X{H, V, C, N, Q}
X97X{F, I, L, M, P, W, Y}

or more specific for subtilisin 309 and closely related subtilases, such as BAALKP, BLSUBL, and BSKSMK
Furthermore the invention relates to subtilases comprising multiple insertions in position 97, or any of the below-mentioned combinations.

It is well known in the art that a so-called conservative substitution of one amino acid residue to a similar amino acid residue is expected to produce only a minor change in the characteristic of the enzyme.

Table III below list groups of conservative amino acid substitutions.
Table III

Conservative amino acid substitutions

<table>
<thead>
<tr>
<th>Common Property</th>
<th>Amino Acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basic (positive charge)</td>
<td>K = lysine</td>
</tr>
<tr>
<td></td>
<td>H = histidine</td>
</tr>
<tr>
<td>Acidic (negative charge)</td>
<td>E = glutamic acid</td>
</tr>
<tr>
<td></td>
<td>D = aspartic acid</td>
</tr>
<tr>
<td>Polar</td>
<td>Q = glutamine</td>
</tr>
<tr>
<td></td>
<td>N = asparagine</td>
</tr>
<tr>
<td>Hydrophobic</td>
<td>L = leucine</td>
</tr>
<tr>
<td></td>
<td>I = isoleucine</td>
</tr>
<tr>
<td></td>
<td>V = valine</td>
</tr>
<tr>
<td></td>
<td>M = methionine</td>
</tr>
<tr>
<td>Aromatic</td>
<td>F = phenylalanine</td>
</tr>
<tr>
<td></td>
<td>W = tryptophan</td>
</tr>
<tr>
<td></td>
<td>Y = tyrosine</td>
</tr>
<tr>
<td>Small</td>
<td>G = glycine</td>
</tr>
<tr>
<td></td>
<td>A = alanine</td>
</tr>
<tr>
<td></td>
<td>S = serine</td>
</tr>
<tr>
<td></td>
<td>T = threonine</td>
</tr>
</tbody>
</table>

According to this principle subtilase variants comprising conservative substitutions, such as G97A+A98AS+S99G, G97S+A98AT+S99A are expected to exhibit characteristics that are not drastically different from each other.

Based on the disclosed and/or exemplified subtilase variants herein, it is routine work for a person skilled in the art to identify suitable conservative modification(s) to these variants in order to obtain other subtilase variants exhibiting similarly improved wash-performance.

According to the invention it the subtilases of the invention belong to the subgroups I-S1 and I-S2, especially subgroup I-S2, both for isolating novel enzymes of the
invention from nature or from the artificial creation of diversity, and for designing and producing variants from a parent subtilase.

In relation to variants from subgroup I-S1, it is preferred to choose a parent subtilase from the group comprising BS168 (BSSAS, BSAPRJ, BSAPRN, BMSAMP), BASBPN, BSSDY, BLSCAR (BLKERA, BLSCA1, BLSCA2, BLSCA3), BSSPRC, and BSSPRD, or functional variants thereof having retained the characteristic of sub-group I-S1.

In relation to variants from subgroup I-S2 it is preferred to choose a parent subtilase from the group comprising BSAPRQ, BLS147 (BSAPRM, BAH101), BLSAVI (BSKSMK, BAALKP, BLSUBL), BYSYAB, and BSAPRS, or functional variants thereof having retained the characteristic of sub-group I-S2.

In particular said parent subtilase is BLSAVI (SAVINASE® NOVO NORDISK A/S), and a preferred subtilase variant of the invention is accordingly a variant of SAVINASE®.

The present invention also comprises any of the above mentioned subtilases of the invention in combination with any other modification to the amino acid sequence thereof. Especially combinations with other modifications known in the art to provide improved properties to the enzyme are envisaged. The art describe a number of subtilase variants with different improved properties and a number of those are mentioned in the "Background of the invention" section herein (vide supra). Those references are disclosed here as references to identify a subtilase variant, which advantageously can be combined with a subtilase variant of the invention.

Such combinations comprise the positions: 222 (improve oxidation stability), 218 (improves thermal stability), substitutions in the Ca-binding sites stabilizing the enzyme, e.g. position 76, and many other apparent from the prior art.
In further embodiments a subtilase variant of the invention may advantageously be combined with one or more modification(s) in any of the positions: 27, 36, 57, 76, 87, 97, 101, 104, 120, 123, 167, 170, 206, 218, 222, 224, 235 and 274.


Furthermore variants comprising any of the variants S101G+V104N, S87N+S101G+V104N, K27R+V104Y+N123S+T274A, N76D+S103A+V104I or N76D+V104A or other combinations of these mutations (V104N, S101G, K27R, V104Y, N123S, T274A, N76D, V104A) in combination with any one or more of the modification(s) mentioned above exhibit improved properties.

Even further subtilase variants of the main aspect(s) of the invention are preferably combined with one or more modification(s) in any of the positions 129, 131, 133 and 194, preferably as 129K, 131H, 133P, 133D and 194P modifications, and most preferably as P129K, P131H, A133P, A133D and A194P modifications. Any of those modification(s) are expected to provide a higher expression level of a subtilase variant of the invention in the production thereof.

Accordingly, an even further embodiment of the invention relates to a variant according to the invention, wherein said modification is chosen from the group comprising: G97GA
G97GT
G97GG
G97GS
G97GD
G97GE
PRODUCING A SUBTILASE VARIANT

Many methods for cloning a subtilase of the invention and for introducing insertions into genes (e.g. subtilase genes) are well known in the art, cf. the references cited in the "BACKGROUND OF THE INVENTION" section.

In general standard procedures for cloning of genes and introducing insertions (random and/or site directed) into said genes may be used in order to obtain a subtilase variant of the invention. For further description of suitable techniques reference is made to Examples herein and Sambrook et al. (1989) Molecular cloning: A laboratory manual, Cold Spring Harbor lab., Cold Spring Harbor, NY; Ausubel, F. M. et al. (eds.) "Current protocols in Molecular Biology". John Wiley and Sons, 1995; Harwood, C. R., and Cutting, S. M. (eds.) "Molecular Biological Methods for Bacillus". John Wiley and Sons, 1990); and WO 96/34946.

Further a subtilase variant of the invention may be constructed by standard techniques for artificial creation of
diversity, such as by DNA shuffling of different subtilase genes (WO 95/22625; Stemmer WPC, Nature 370:389-91 (1994)). DNA shuffling of e.g. the gene encoding Savinase® with one or more partial subtilase sequences identified in nature to comprise an active site loop (b) regions longer than the active site loop (b) of Savinase®, will after subsequent screening for improved wash performance variants, provide subtilase variants according to the invention.

10 **EXPRESSION VECTORS**

A recombinant expression vector comprising a DNA construct encoding the enzyme of the invention may be any vector which may conveniently be subjected to recombinant DNA procedures.

The choice of vector will often depend on the host cell into which it is to be introduced. Thus, the vector may be an autonomously replicating vector, i.e. a vector which exists as an extrachromosomal entity, the replication of which is independent of chromosomal replication, e.g. a plasmid.

Alternatively, the vector may be one that on introduction into a host cell is integrated into the host cell genome in part or in its entirety and replicated together with the chromosome(s) into which it has been integrated.

The vector is preferably an expression vector in which the DNA sequence encoding the enzyme of the invention is operably linked to additional segments required for transcription of the DNA. In general, the expression vector is derived from plasmid or viral DNA, or may contain elements of both. The term, ‘operably linked’ indicates that the segments are arranged so that they function in concert for their intended purposes, e.g. transcription initiates in a promoter and proceeds through the DNA sequence coding for the enzyme.
The promoter may be any DNA sequence which shows transcriptional activity in the host cell of choice and may be derived from genes encoding proteins either homologous or heterologous to the host cell.

Examples of suitable promoters for use in bacterial host cells include the promoter of the Bacillus stearothermophilus maltogenic amylase gene, the Bacillus licheniformis alpha-amylase gene, the Bacillus amyloliquefaciens alpha-amylase gene, the Bacillus subtilis alkaline protease gene, or the Bacillus pumilus xylosidase gene, or the phage Lambda P_R or P_L promoters or the E. coli lac, trp or tac promoters.

The DNA sequence encoding the enzyme of the invention may also, if necessary, be operably connected to a suitable terminator.

The recombinant vector of the invention may further comprise a DNA sequence enabling the vector to replicate in the host cell in question.

The vector may also comprise a selectable marker, e.g. a gene the product of which complements a defect in the host cell, or a gene encoding resistance to e.g. antibiotics like kanamycin, chloramphenicol, erythromycin, tetracycline, spectinomycine, or the like, or resistance to heavy metals or herbicides.

To direct an enzyme of the present invention into the secretory pathway of the host cells, a secretory signal sequence (also known as a leader sequence, prepro sequence or pre sequence) may be provided in the recombinant vector. The secretory signal sequence is joined to the DNA sequence encoding the enzyme in the correct reading frame. Secretory signal sequences are commonly positioned 5' to the DNA sequence encoding the enzyme. The secretory signal sequence may be that
normally associated with the enzyme or may be from a gene encoding another secreted protein.

The procedures used to ligate the DNA sequences coding for the present enzyme, the promoter and optionally the terminator and/or secretory signal sequence, respectively, or to assemble these sequences by suitable PCR amplification schemes, and to insert them into suitable vectors containing the information necessary for replication or integration, are well known to persons skilled in the art (cf., for instance, Sambrook et al., op.cit.).

HOST CELL
The DNA sequence encoding the present enzyme introduced into the host cell may be either homologous or heterologous to the host in question. If homologous to the host cell, i.e. produced by the host cell in nature, it will typically be operably connected to another promoter sequence or, if applicable, another secretory signal sequence and/or terminator sequence than in its natural environment. The term "homologous" is intended to include a DNA sequence encoding an enzyme native to the host organism in question. The term "heterologous" is intended to include a DNA sequence not expressed by the host cell in nature. Thus, the DNA sequence may be from another organism, or it may be a synthetic sequence.

The host cell into which the DNA construct or the recombinant vector of the invention is introduced may be any cell which is capable of producing the present enzyme and includes bacteria, yeast, fungi and higher eukaryotic cells including plants.

Examples of bacterial host cells which, on cultivation, are capable of producing the enzyme of the invention are gram-positive bacteria such as strains of Bacillus, such as strains of B. subtilis, B. licheniformis, B. lentus, B. brevis, B.
stearothermophilus, B. alkalophilus, B. amyloliquefaciens, B. coagulans, B. circulans, B. laetus, B. megatherium or B. thuringiensis, or strains of Streptomyces, such as S. lividans or S. murinus, or gram-negative bacteria such as Echerichia coli.

The transformation of the bacteria may be effected by protoplast transformation, electroporation, conjugation, or by using competent cells in a manner known per se (cf. Sambrook et al., supra).

When expressing the enzyme in bacteria such as E. coli, the enzyme may be retained in the cytoplasm, typically as insoluble granules (known as inclusion bodies), or may be directed to the periplasmic space by a bacterial secretion sequence. In the former case, the cells are lysed and the granules are recovered and denatured after which the enzyme is refolded by diluting the denaturing agent. In the latter case, the enzyme may be recovered from the periplasmic space by disrupting the cells, e.g. by sonication or osmotic shock, to release the contents of the periplasmic space and recovering the enzyme.

When expressing the enzyme in gram-positive bacteria such as Bacillus or Streptomyces strains, the enzyme may be retained in the cytoplasm, or may be directed to the extracellular medium by a bacterial secretion sequence. In the latter case, the enzyme may be recovered from the medium as described below.

METHOD OF PRODUCING SUBTILASE

The present invention provides a method of producing an isolated enzyme according to the invention, wherein a suitable host cell, which has been transformed with a DNA sequence encoding the enzyme, is cultured under conditions permitting
the production of the enzyme, and the resulting enzyme is recovered from the culture.

When an expression vector comprising a DNA sequence encoding the enzyme is transformed into a heterologous host cell it is possible to enable heterologous recombinant production of the enzyme of the invention.

Thereby it is possible to make a highly purified subtilase composition, characterized in being free from homologous impurities.

In this context homologous impurities means any impurities (e.g. other polypeptides than the enzyme of the invention) which originate from the homologous cell where the enzyme of the invention is originally obtained from.

The medium used to culture the transformed host cells may be any conventional medium suitable for growing the host cells in question. The expressed subtilase may conveniently be secreted into the culture medium and may be recovered therefrom by well-known procedures including separating the cells from the medium by centrifugation or filtration, precipitating proteinaceous components of the medium by means of a salt such as ammonium sulfate, followed by chromatographic procedures such as ion exchange chromatography, affinity chromatography, or the like.

USE OF A SUBTILASE VARIANT OF THE INVENTION

A subtilase protease variant of the invention may be used for a number of industrial applications, in particular within the detergent industry.

Further the invention relates to an enzyme composition, which comprises a subtilase variant of the invention.

A summary of preferred industrial applications and corresponding preferred enzyme compositions is provided below.
This summary is not in any way intended to be a complete list of suitable applications of a subtilase variant of the invention. A subtilase variants of the invention may be used in other industrial applications known in the art to include the use of a protease, in particular a subtilase.

DETERGENT COMPOSITIONS COMPRISING THE MUTANT ENZYMES

The present invention comprises the use of the mutant enzymes of the invention in cleaning and detergent compositions and such compositions comprising the mutant subtilisin enzymes. Such cleaning and detergent compositions are well described in the art and reference is made to WO 96/34946; WO 97/07202; WO 95/30011 for further description of suitable cleaning and detergent compositions.

Furthermore the example(s) below demonstrate the improvements in wash performance for a number of subtilase variants of the invention.

Detergent Compositions

The enzyme of the invention may be added to and thus become a component of a detergent composition.

The detergent composition of the invention may for example be formulated as a hand or machine laundry detergent composition including a laundry additive composition suitable for pre-treatment of stained fabrics and a rinse added fabric softener composition, or be formulated as a detergent composition for use in general household hard surface cleaning operations, or be formulated for hand or machine dishwashing operations.

In a specific aspect, the invention provides a detergent additive comprising the enzyme of the invention. The detergent additive as well as the detergent composition may comprise one or more other enzymes such as a protease, a lipase, a cutinase, an amylase, a carbohydrase, a cellulase, a pectinase, a man-
nanase, an arabinase, a galactanase, a xylanase, an oxidase, e.g., a laccase, and/or a peroxidase.

In general the properties of the chosen 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.

**Proteases:** Suitable proteases include those of animal, vegetable or microbial origin. Microbial origin is preferred. Chemically modified or protein engineered mutants are included. The protease may be a serine protease or a metallo protease, preferably an alkaline microbial protease or a trypsin-like protease. Examples of alkaline proteases are subtilisins, especially those derived from Bacillus, e.g., subtilisin Novo, subtilisin Carlsberg, subtilisin 309, subtilisin 147 and subtilisin 168 (described in WO 89/06279). Examples of trypsin-like proteases are trypsin (e.g. of porcine or bovine origin) and the Fusarium protease described in WO 89/06270 and WO 94/25583.

Examples of useful proteases are the variants described in WO 92/19729, WO 98/20115, WO 98/20116, and WO 98/34946, especially the variants with substitutions in one or more of the following positions: 27, 36, 57, 76, 87, 97, 101, 104, 120, 123, 167, 170, 194, 206, 218, 222, 224, 235 and 274.

Preferred commercially available protease enzymes include Alcalase™, Savinase™, Primase™, Duralase™, Esperase™, and Kannase™ (Novo Nordisk A/S), Maxatase™, Maxacal™, Maxapem™, Properase™, Purapect™, Purapect OxP™, FN2™, and FN3™ (Genencor International Inc.).

**Lipases:** Suitable lipases include those of bacterial or fungal origin. Chemically modified or protein engineered mutants are included. Examples of useful lipases include lipases from *Humicola* (synonym *Thermomyces*), e.g. from *H. lanuginosa* (T.
lanuginosus) as described in EP 258 068 and EP 305 216 or from H. insolens as described in WO 96/13580, a Pseudomonas lipase, e.g. from P. alcaligenes or P. pseudoalcaligenes (EP 218 272), P. cepacia (EP 331 376), P. stutzeri (GB 1,372,034), P. fluorescens, Pseudomonas sp. strain SD 705 (WO 95/06720 and WO 96/27002), P. wisconsinensis (WO 96/12012), a Bacillus lipase, e.g. from B. subtilis (Dartois et al. (1993), Biochemica et Biophysica Acta, 1131, 253-360), B. stearothermophilus (JP 64/744992) or B. pumilus (WO 91/16422).


Preferred commercially available lipase enzymes include Lipolase™ and Lipolase Ultra™ (Novo Nordisk A/S).

Amylases: Suitable amylases (α and/or β) include those of bacterial or fungal origin. Chemically modified or protein engineered mutants are included. Amylases include, for example, α-amylases obtained from Bacillus, e.g. a special strain of B. licheniformis, described in more detail in GB 1,296,839.

Examples of useful amylases are the variants described in WO 94/02597, WO 94/18314, WO 96/23873, and WO 97/43424, especially the variants with substitutions in one or more of the following positions: 15, 23, 105, 106, 124, 128, 133, 154, 156, 181, 188, 190, 197, 202, 208, 209, 243, 264, 304, 305, 391, 408, and 444.

Commercially available amylases are Duramyl™, Termamyl™, Fungamyl™ and BAN™ (Novo Nordisk A/S), Rapidase™ and Purastar™ (from Genencor International Inc.).

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,


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

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™ (Novo Nordisk 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 e.g. as a granulate, a liquid, a 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.

The detergent composition of the invention may be in any convenient form, e.g., a bar, a tablet, a powder, a granule, a paste or a liquid. A liquid detergent may be aqueous, typically containing up to 70 % water and 0-30 % organic solvent, or non-aqueous.

The detergent composition comprises one or more surfactants, which may be non-ionic including semi-polar and/or anionic and/or cationic and/or zwitterionic. The surfactants are typically present at a level of from 0.1% to 60% by weight.

When included therein the detergent will usually contain from about 1% to about 40% of an anionic surfactant such as linear alkylbenzenesulfonate, alpha-olefinsulfonate, alkyl sulfate (fatty alcohol sulfate), alcohol ethoxysulfate, secondary alkanesulfonate, alpha-sulfo fatty acid methyl ester, alkyl- or alkenylsuccinic acid or soap.

When included therein the detergent will usually contain from about 0.2% to about 40% of a non-ionic surfactant such as
alcohol ethoxylate, nonylphenol ethoxylate, alkylpolyglycoside,
aluminium, ethoxylated fatty acid monoethanolamide, fatty acid monoethanolamide, polyhydroxy alkyl fatty acid amide, or N-acyl N-alkyl derivatives of glucosamine
5 ("glucamides").

The detergent may contain 0-65 % of a detergent builder or complexing agent such as zeolite, diphosphate, triphosphate, phosphonate, carbonate, citrate, nitrilotriacetic acid, ethylenediaminetetraacetic acid, diethylenetriaminepentaacetic acid, alkyl- or alkenylsuccinic acid, soluble silicates or layered silicates (e.g. SKS-6 from Hoechst).

The detergent may comprise one or more polymers. Examples are carboxymethylcellulose, poly(vinylpyrrolidone), poly (ethylene glycol), poly(vinyl alcohol), poly(vinylpyridine-N-oxide), poly(vinylimidazole), polycarboxylates such as polycrylic acid copolymers and lauryl methacrylate/acrylic acid copolymers.

The detergent may contain a bleaching system which may comprise a H₂O₂ source such as perborate or percarbonate which may be combined with a peracid-forming bleach activator such as tetraacetylene diamine or nonoxoxybenzenesulfonate. Alternatively, the bleaching system may comprise peroxyacids of e.g. the amide, imide, or sulfone type.

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 e.g. WO 92/19709 and WO 92/19708.

The detergent may also contain other conventional detergent ingredients such as e.g. fabric conditioners including
clays, foam boosters, suds suppressors, anti-corrosion agents, soil-suspending agents, anti-soil redeposition agents, dyes, bactericides, optical brighteners, hydrotropes, tarnish inhibitors, or perfumes.

It is at present contemplated that in the detergent compositions any enzyme, in particular the enzyme of the invention, may be added in an amount corresponding to 0.01-100 mg of enzyme protein per liter of wash liquor, preferably 0.05-5 mg of enzyme protein per liter of wash liquor, in particular 0.1-1 mg of enzyme protein per liter of wash liquor.

The enzyme of the invention may additionally be incorporated in the detergent formulations disclosed in WO 97/07202 which is hereby incorporated as reference.

LEATHER INDUSTRY APPLICATIONS

A subtilase of the invention may be used in the leather industry, in particular for use in depilation of skins.

In said application a subtilase variant of the invention is preferably used in an enzyme composition which further comprises another protease.

For a more detailed description of suitable other proteases see section relating to suitable enzymes for use in a detergent composition (vide supra).

WOOL INDUSTRY APPLICATIONS

A subtilase of the invention may be used in the wool industry, in particular for use in cleaning of clothes comprising wool.

In said application a subtilase variant of the invention is preferably used in an enzyme composition which further comprises another protease.
For a more detailed description of suitable other proteases see section relating to suitable enzymes for use in a detergent composition (*vide supra*).

The invention is described in further detail in the following examples which are not in any way intended to limit the scope of the invention as claimed.

**MATERIALS AND METHODS**

10 **STRAINS:**


*B. lentus* 309 and 147 are specific strains of *Bacillus lentus*, deposited with the NCIB and accorded the accession numbers NCIB 10309 and 10147, and described in US Patent No. 3,723,250 incorporated by reference herein.

*E. coli* MC 1000 (*M. J. Casadaban and S. N. Cohen* (1980); *J. Mol. Biol.* **138** 179-207), was made *r<sup>-</sup>,m<sup>+</sup>* by conventional methods and is also described in US Patent Application Serial No. 039,298.

20 **PLASMIDS:**

pJS3: *E. coli* - *B. subtilis* shuttle vector containing a synthetic gene encoding for subtilase 309. (Described by *Jacob Schiødt et al.* in Protein and Peptide letters **3**:39-44 (1996)).

pSX222: *B. subtilis* expression vector (Described in WO 96/34946).

**GENERAL MOLECULAR BIOLOGY METHODS:**

Unless otherwise mentioned the DNA manipulations and transformations are performed using standard methods of molecular biology (*Sambrook et al.* (1989) Molecular cloning: A laboratory manual, Cold Spring Harbor lab., Cold Spring Harbor,

Enzymes for DNA manipulations are used according to the specifications of the suppliers.

ENZYMES FOR DNA MANIPULATIONS

Unless otherwise mentioned all enzymes for DNA manipulations, such as e.g. restriction endonucleases, ligases etc., are obtained from New England Biolabs, Inc.

PROTEOLYTIC ACTIVITY

In the context of this invention proteolytic activity is expressed in Kilo NOVO Protease Units (KNPU). The activity is determined relatively to an enzyme standard (SAVINASE®), and the determination is based on the digestion of a dimethyl casein (DMC) solution by the proteolytic enzyme at standard conditions, i.e. 50°C, pH 8.3, 9 min. reaction time, 3 min. measuring time. A folder AF 220/1 is available upon request to Novo Nordisk A/S, Denmark, which folder is hereby included by reference.

A GU is a Glycine Unit, defined as the proteolytic enzyme activity which, under standard conditions, during a 15 minutes' incubation at 40°C, with N-acetyl casein as substrate, produces an amount of NH₃-group equivalent to 1 mmole of glycine.

FERMENTATION:

Fermentations for the production of subtilase enzymes are performed at 30°C on a rotary shaking table (300 r.p.m.) in 500 ml baffled Erlenmeyer flasks containing 100 ml BPX medium for 5 days.

Consequently in order to make an e.g. 2 liter broth 20 Erlenmeyer flasks are fermented simultaneously.

MEDIA:

BPX Medium Composition (per liter)

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potato starch</td>
<td>100 g</td>
</tr>
<tr>
<td>Ground barley</td>
<td>50 g</td>
</tr>
<tr>
<td>Soybean flour</td>
<td>20 g</td>
</tr>
<tr>
<td>Na$_2$HPO$_4$ x 12 H$_2$O</td>
<td>9 g</td>
</tr>
<tr>
<td>Pluronic</td>
<td>0.1 g</td>
</tr>
<tr>
<td>Sodium caseinate</td>
<td>10 g</td>
</tr>
</tbody>
</table>

The starch in the medium is liquefied with α-amylase and the medium is sterilized by heating at 120°C for 45 minutes. After sterilization the pH of the medium is adjusted to 9 by addition of NaHCO$_3$ to 0.1 M.
EXAMPLE 1

CONSTRUCTION AND EXPRESSION OF ENZYME VARIANTS:

SITE-DIRECTED MUTAGENESIS:

Subtilase 309 site-directed variants of the invention comprising specific insertions in the active site loop (b) between positions 97 and 98 are made by traditional cloning of DNA fragments (Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd Ed., Cold Spring Harbor, 1989) produced by PCR of oligos containing the desired insertions (see below).

The template plasmid DNA is pJS3, or an analogue of this containing a variant of Subtilase 309.

Insertions are introduced by oligo directed mutagenesis to the construction of G97GX insertion variants (X = any amino acid residue inserted between positions 97 and 98) resulting in G97GX Subtilase 309 variants.

The Subtilase 309 variants are transformed into E. coli. DNA purified from an over night culture of these transformants were transformed into B. subtilis by restriction endonuclease digestion, purification of DNA fragments, ligation, transformation of B. subtilis. Transformation of B. subtilis is performed as described by Dubnau et al., 1971, J. Mol. Biol. 56, pp. 209-221.

LOCALIZED RANDOM MUTAGENESIS IN ORDER TO INSERT RANDOM INSERTIONS IN A LOCALIZED REGION:

LOCALIZED RANDOM MUTAGENESIS IN ORDER TO INSERT RANDOM INSERTIONS IN A LOCALIZED REGION:

The overall strategy used to perform localized random mutagenesis is:

A mutagenic primer (oligonucleotide) is synthesized corresponding to the DNA sequence flanking the sites of insertion, separated by the DNA base pairs defining the insertion.
Subsequently, the resulting mutagenic primer is used in a PCR reaction with a suitable opposite primer. The resulting PCR fragment was purified and digested by endonucleases and cloned into the E. coli - B. subtilis shuttle vector (see below).

Alternatively, and if necessary, the resulting PCR fragment is used in a second PCR reaction as a primer with a second suitable opposite primer to allow digestion and cloning of the mutagenized region into the shuttle vector. The PCR reactions are performed under normal conditions.

Following this strategy a localized random library was constructed in SAVINASE wherein insertions were introduced in the active site loop region between positions 97 and 98.

The mutations were introduced by mutagenic primers (see below), so that all 20 amino acids are represented (N = 25% of A, T, C, and G; whereas S = 50% C and G. The produced PCR fragment were extended towards the N-terminal of Savinase by another round of PCR by combination of a overlapping sequence with a PCR-fragment produced by PCR-amplification with primers; 5' CTA AAT ATT CGT GGT GGC GC 3' (sense) and 5' GAC TTT AAC AGC GTA TAG CTC AGC 3' (antisense). The extended DNA-fragments were cloned into the Hind III- and Mlu I-sites of the modified plasmid pJS3 (see above), and ten randomly chosen E. coli colonies were sequenced to confirm the mutations designed.

The mutagenic primer (5' GCT GAG CTA TAC GCT GTT AAA GTC CTA GGG NNS GCG AGC GGT TCA GGT TC 3' (sense)) were used in a PCR reaction with a suitable anti-sense opposite primer, situated downstream of the Mlu I site in pJS3 (e.g. 5' - CCC TTT AAC CGC ACA GCG TTT -3' (anti-sense)) and the plasmid pJS3 as template. This resulting PCR product was cloned into the pJS3 shuttle vector by using the restriction enzymes Hind III and Mlu I.
The random library was transformed into E. coli by well known techniques.

The library prepared contained approximately 100,000 individual clones/library.

Ten randomly chosen colonies were sequenced to confirm the mutations designed.

In order to purify a subtilase variant of the invention, the *B. subtilis* pJS3 expression plasmid comprising a variant of the invention was transformed into a competent *B. subtilis* strain and was fermented as described above in a medium containing 10 μg/ml Chloramphenicol (CAM).

**EXAMPLE 2**

PURIFICATION OF ENZYME VARIANTS:

This procedure relates to purification of a 2 liter scale fermentation for the production of the subtilases of the invention in a Bacillus host cell.

Approximately 1.6 liters of fermentation broth are centrifuged at 5000 rpm for 35 minutes in 1 liter beakers. The supernatants are adjusted to pH 6.5 using 10% acetic acid and filtered on Seitz Supra S100 filter plates.

The filtrates are concentrated to approximately 400 ml using an Amicon CH2A UF unit equipped with an Amicon S1Y10 UF cartridge. The UF concentrate is centrifuged and filtered prior to absorption at room temperature on a Bacitracin affinity column at pH 7. The protease is eluted from the Bacitracin column at room temperature using 25% 2-propanol and 1 M sodium chloride in a buffer solution with 0.01 dimethylglutaric acid, 0.1 M boric acid and 0.002 M calcium chloride adjusted to pH 7.

The fractions with protease activity from the Bacitra-cin purification step are combined and applied to a 750 ml Sephadex G25 column (5 cm dia.) equilibrated with a buffer containing 0.01 dimethylglutaric acid, 0.2 M boric acid and 0.002 M calcium chloride adjusted to pH 6.5.
Fractions with proteolytic activity from the Sephadex G25 column are combined and applied to a 150 ml CM Sepharose CL 6B cation exchange column (5 cm dia.) equilibrated with a buffer containing 0.01 M dimethylglutaric acid, 0.2 M boric acid, and 0.002 M calcium chloride adjusted to pH 6.5.

The protease is eluted using a linear gradient of 0-0.1 M sodium chloride in 2 litres of the same buffer (0-0.2 M sodium chloride in case of Subtilisin 147).

In a final purification step protease containing fractions from the CM Sepharose column are combined and concentrated in an Amicon ultrafiltration cell equipped with a GR81PP membrane (from the Danish Sugar Factories Inc.).

By using the techniques of Example 1 for the construction and fermentation, and the above isolation procedure the following subtilisin 309 variants were produced and isolated:

G97GT
G97GS
G97GD
G97GE
G97GP
G97GG
G97GH
G97GI
G97GA
G97GT+Y167A
G97GP+A98T

These variants were found to exhibit better wash performance than Savinase in a preliminary assay.
EXAMPLE 3

WASH PERFORMANCE OF DETERGENT COMPOSITIONS COMPRISING ENZYME VARIANTS

The following examples provide results from a number of washing tests that were conducted under the conditions indicated.

MINI WASH

WASH CONDITIONS:

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<tr>
<td>Detergent Dosage</td>
<td>4g/l</td>
<td>1g/l</td>
</tr>
<tr>
<td>Wash Temp</td>
<td>30°C</td>
<td>25°C</td>
</tr>
<tr>
<td>Wash Time</td>
<td>30min</td>
<td>10min</td>
</tr>
<tr>
<td>Water hardness</td>
<td>18°dH (Ca²⁺/Mg²⁺ = 5:1)</td>
<td>6°dH (Ca²⁺/Mg²⁺ = 2:1)</td>
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<tr>
<td>PH</td>
<td>Not adjusted</td>
<td>Not adjusted</td>
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<tr>
<td>Enzyme conc.</td>
<td>1, 2, 5, 10, 30 nM</td>
<td>1, 2, 5, 10, 30 nM</td>
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<tr>
<td>Test system</td>
<td>150 ml glass beakers with a stirring rod</td>
<td>150 ml glass beakers with a stirring rod</td>
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<tr>
<td>Textile/volume</td>
<td>5 textile pieces (Ø 2.5 cm) in 50 ml detergent</td>
<td>5 textile pieces (Ø 2.5 cm) in 50 ml detergent</td>
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<tr>
<td>Test Material</td>
<td>EMPA116</td>
<td>EMPA117</td>
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DETERGENTS:

The detergents used are obtained from supermarkets in Denmark (OMO, datasheet ED-9745105) and the USA (Wisk, datasheet ED-9711893), respectively. Prior to use all enzymatic activity in the detergents is inactivated by micro wave treatment.
SWATCHES;

The swatches used are EMPA116 and EMPA117, obtained from EMPA Testmaterialen, Movenstrasse 12, CH-9015 St. Gall, Switzerland.

REFLECTANCE

Measurement of reflectance (R) on the test material is done at 460 nm using a Macbeth ColorEye 7000 photometer. The measurements are done according to the manufacturers protocol.

EVALUATION

The evaluation of the wash performance of a subtilase is determined by either the improvement factor or the performance factor for the subtilase investigated.

The improvement factor, IF_{Dose/response}, is defined as the ratio between the slopes of the wash performance curves for a detergent containing the subtilase investigated and the same detergent containing a reference subtilase at the asymptotic concentration of the subtilase goes to zero.

\[ \text{IF}_{\text{Dose/response}} = \frac{a}{a_{\text{ref}}} \]

The wash performance is calculated according to the formula I:

\[ R = R_0 + \frac{a \cdot \Delta R_{\text{max}} \cdot c}{\Delta R_{\text{max}} + a \cdot c} \quad (I); \]

where

R is the wash performance in reflectance units; \( R_0 \) is the intercept of the fitted curve with y-axis (blind); a is the slope of the fitted curve as c \( \to 0 \); c is the enzyme
concentration; and $\Delta R_{\text{max}}$ is the theoretical maximal wash effect as $c \to \infty$.

The performance factor, $P$, is calculated according to the formula II

$$P = \frac{R_{\text{Variant}} - R_{\text{Blank}}}{R_{\text{Savinase}} - R_{\text{Blank}}} \quad (\text{II})$$

where $R_{\text{variant}}$ is the reflectance of test material washed with 10nM variant; $R_{\text{savinase}}$ is the reflectance of test material washed with 10nM Savinase; $R_{\text{blank}}$ is the reflectance of test material washed with no enzyme.

US (detergent: OMO, Swatch: EMPA116)

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<th>Variant</th>
<th>IFDose/response</th>
<th>$P$</th>
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<td>G97GA</td>
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US (detergent: US Wisk, Swatch: EMPA117)

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<th>Variant</th>
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<td>G97GA</td>
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<tr>
<td>G97GP+A98T</td>
<td>-</td>
<td>1.51</td>
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<td>G97GAA+A98S+S99G+</td>
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<td>1.28</td>
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<td>S101T*</td>
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</table>

*Disclosed in WO 99/27082

The subtilases of the inventions are thus seen to exhibit improved wash performance in comparison to Savinase*. 
CLAIMS

1. A Subtilase enzyme of the I-S1 or I-S2 sub-group having at least one additional amino acid residue in position 97 of the active site loop (b) region from position 95 to 103, whereby said additional amino acid residue(s) correspond to the insertion of at least one amino acid residue between positions 97 and 98, with the proviso that the subtilase enzyme does not contain the following insertions G97GASG, G97GAA, G97GAS,
G97GASG+A98S+S99G+G100A+S101A or G97GAA+A98S+S99G+S101T.

2. The isolated subtilase enzyme of claim 1, wherein said subtilase enzyme is a constructed variant having at least one inserted amino acid residue between positions 97 and 98 of a precursor subtilase.

3. The isolated subtilase enzyme of claim 1 or 2 selected from the group consisting of

\[ X97\{A,T,G,S\}, \]
\[ X97\{D,E,K,R\}, \]
\[ X97\{H,V,C,N,Q\}, \text{ and} \]
\[ X97\{F,I,L,M,P,W,Y\}. \]

4. The isolated subtilase enzyme of claim 3, wherein said at least one additional or inserted amino acid residue is selected from the group consisting of T, G, A, and S.

5. The isolated subtilase enzyme of claim 3, wherein said at least one additional or inserted amino acid residue is selected from the group of charged amino acid residues consisting of D,E,H,K, and R, more preferably D,E,K and R.
6. The isolated subtilase enzyme of claim 3, wherein said at least one additional or inserted amino acid residue is selected from the group of hydrophilic amino acid residues consisting of C,N,Q,S and T, more preferably N,Q,S and T.

7. The isolated subtilase enzyme of claim 3, wherein said at least one additional or inserted amino acid residue is selected from the group of small hydrophobic amino acid residues consisting of A,G and V.

8. The isolated subtilase enzyme of claim 3, wherein said at least one additional or inserted amino acid residue is selected from the group of large hydrophobic amino acid residues consisting of F,I,L,M,P,W and Y, more preferably F,I,L,M, and Y.

9. The isolated subtilase enzyme according to any of the preceding claims, wherein said at least one additional or inserted amino acid residue, comprises more than one additional or inserted amino acid residue in the active site loop (b).

10. The isolated subtilase enzyme according to any of claims 1-8, which comprises one additional or inserted amino acid residue in the active site loop (b).

11. The subtilase variant of any of the preceding claims, wherein said insertion(s) between positions 97 and 98 are combined with one or more further modification(s) in any other position(s).

12. The subtilase variant of claim 11, wherein said further modification(s) are in one or more of the positions 27, 36, 57,
76, 87, 97, 101, 104, 120, 123, 167, 170, 206, 218, 222, 224, 235 and 274.

13. The subtilase variant of any of the preceding claims, wherein said modification(s) is/are combined with modification(s) in one or more of the positions 95, 96, 97, 98, 99, 100, 101, 102, and 103.

14. The subtilase of any of the preceding claims, wherein the subtilase, or if the subtilase is a variant, the parent subtilase belongs to the sub-group I-S1.

15. The subtilase of claim 14, wherein the parent subtilase is selected from the group consisting of ABSS168, BASBPIN, BSSDY, and BLSCAR, or functional variants thereof having retained the characteristic of sub-group I-S1.

16. The subtilase according to any of claims 1-15, wherein the subtilase, or if the subtilase is a variant, the parent subtilase belongs to the sub-group I-S2.

17. The subtilase of claim 16, wherein the parent subtilase is selected from the group consisting of BLS147, BLS309, BAPB92, TVTHER and BSYAB, or functional variants thereof having retained the characteristic of sub-group I-S2.

18. The isolated subtilase enzyme of claim 3, 16 or 17 selected from the group consisting of

   G97GA,
   G97GT,
   G97GG,
   G97GS,
   G97GD,
G97GE,
G97GK,
G97GR,
G97GH,
G97GV,
G97GC,
G97GN,
G97GQ,
G97GF,
G97GI,
G97GL,
G97GM,
G97GP,
G97GW, and
G97GY.


20. The subtilase variant of any of the claims 16 to 18, wherein said further modification(s) are selected from the group consisting of S101G+V104N, S87N+S101G+V104N, K27R+V104Y+N123S+T274A, N76D+S103A+V104I or N76D+V104A, or other combinations of these mutations (V104N, S101G, K27R, V104Y, N123S, T274A, N76D, V104A), in combination with any one or more of the substitutions, deletions and/or insertions mentioned in any of claims 1 to 14.

21. The subtilase variant of any of the claims 16 to 18, wherein said further modification(s) are selected from the
group further consisting of P129K, P131H, A133P, A133D and A194P.

22. The variant according to any of the preceding claims comprising the modification selected from the group consisting of G97GT+Y167A, G97GP+A98T and G97GA.

23. A subtilase belonging to the I-S1 sub-group having the amino acid sequence:

```
10  1  10  20  30
40  50  60
70  80  90
97a 100 110 120
130 140 150
160 170 180
190 200 210
220 230 240
250 260 270
275
```

or a homologous subtilase having an amino acid sequence comprising a position 97a amino acid residue and exhibiting an
identity of more than 70%, 75%, 80%, 85%, 90%, or 95% therewith.

24. A subtilase belonging to the I-S2 sub-group having the
5 amino acid sequence:

1 10 20 30
40 50 60
L-D-T-G-I-*S-T-H-P-D-L-N-I-R-G-G-A-S-P-V-P-G-E-P-*S-T-Q-D-
70 80 90
97a 100 110 120
130 140 150
160 170 180
190 200 210
220 230 240
250 260 270
275

or a homologous subtilase having an amino acid sequence comprising a position 97a amino acid residue and exhibiting an identity of more than 70%, 75%, 80%, 85%, 90%, or 95% therewith.

25. The subtilase variant of claims 23 or 24, wherein X in position 97a is selected from the group consisting of T, A, G, S, and P.
26. An isolated DNA sequence encoding a subtilase or a subtilase variant of any of the claims 1 to 25.

27. An expression vector comprising an isolated DNA sequence of claim 26.

28. A microbial host cell transformed with an expression vector of claim 27.

29. The microbial host of claim 28, which is a bacterium, preferably a Bacillus, especially B. lentus.

30. The microbial host of claim 28, which is a fungus or yeast, preferably a filamentous fungus, especially an Aspergillus.

31. A method for producing a subtilase or a subtilase variant of any of claims 1 to 25, wherein a host of any of claims 28 to 30 is cultured under conditions conducive to the expression and secretion of said variant, and the variant is recovered.

32. A composition comprising a subtilase or a subtilase variant according to any of claims 1 to 25.

33. The composition according to claim 32, which additionally comprises a cellulase, lipase, cutinase, oxidoreductase, another protease, or an amylase.

34. The composition according to claim 32 or 33, wherein the composition is a detergent composition.

35. Use of a subtilase or a subtilase variant according to any of claims 1 to 25 or an enzyme composition according to claims 32 or 33 in a laundry and/or a dishwash detergent.
Fig. 1
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Fig. 2
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Fig. 2a

SUBSTITUTE SHEET (RULE 26)
SEQUENCE LISTING

Novo Nordisk A/S

Subtilase enzymes of the I-S1 and I-S2 sub-groups having at least one additional amino acid residue between positions 97 and 98

5926.204

PatentIn Ver. 2.1

1 Bacillus licheniformis

Ala Gln Thr Val Pro Tyr Gly Ile Pro Leu Ile Lys Ala Asp Lys Val 1 5 10 15

Gln Ala Gln Gly Phe Lys Gly Ala Asn Val Lys Val Ala Val Leu Asp 20 25 30

Thr Gly Ile Gin Ala Ser His Pro Asp Leu Asn Val Val Gly Gly Ala 35 40 45

Ser Phe Val Ala Gly Glu Ala Tyr Asn Thr Asp Gly Asn Gly His Gly 50 55 60

Thr His Val Ala Gly Thr Val Ala Ala Leu Asp Asn Thr Thr Gly Val 65 70 75 80

Leu Gly Val Ala Pro Ser Val Ser Leu Tyr Ala Val Lys Val Leu Asn 85 90 95

Xaa Ser Ser Gly Ser Gly Thr Tyr Ser Gly Ile Val Ser Gly Ile Glu 100 105 110

Trp Ala Thr Thr Asn Gly Met Asp Val Ile Asn Met Ser Leu Gly Gly 115 120 125

Pro Ser Gly Ser Thr Ala Met Lys Gln Ala Val Asp Asn Ala Tyr Ala 130 135 140

Arg Gly Val Val Val Ala Ala Ala Gly Asn Ser Gly Ser Ser Gly 145 150 155 160

Asn Thr Asn Thr Ile Gly Tyr Pro Ala Lys Tyr Asp Ser Val Ile Ala 165 170 175

Val Gly Ala Val Asp Ser Asn Ser Asn Arg Ala Ser Phe Ser Ser Val 180 185 190

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Ala Ala Gln 275

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200

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213> Bacillus lentus

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Thr Gly Ile Ser Thr His Pro Asp Leu Asn Ile Arg Gly Gly Ala Ser 35 40 45
Phe Val Pro Gly Glu Pro Ser Thr Gln Asp Gly Asn Gly His Gly Thr 50 55 60
His Val Ala Gly Thr Ile Ala Ala Leu Asn Asn Ser Ile Gly Val Leu 65 70 75 80
Gly Val Ala Pro Ser Ala Glu Leu Tyr Ala Val Lys Val Leu Gly Xaa 85 90 95
Ala Ser Gly Ser Gly Ser Val Ser Ser Ser Ile Ala Gln Gly Leu Glu Trp 100 105 110
Ala Gly Asn Asn Gly Met His Val Ala Leu Ser Leu Gly Ser Pro 115 120 125
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Ser Tyr Pro Ala Arg Tyr Ala Asn Ala Met Ala Val Gly Ala Thr Asp 165 170 175
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SUBSTITUTE SHEET (RULE 26)
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Leu Tyr Gly Ser Gly Leu Val Asn Ala Glu Ala Ala Thr Arg
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# INTERNATIONAL SEARCH REPORT

**A. CLASSIFICATION OF SUBJECT MATTER**

**IPC7:** C12N 9/54, C12N 9/58 // C11D 003/386

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

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

**IPC7:** C12N

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

SE, DK, FI, NO classes as above

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

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

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Further documents are listed in the continuation of Box C. See patent family annex.

Date of the actual completion of the international search: 12 Sept. 2000

Date of mailing of the international search report: 19-09-2000

Name and mailing address of the ISA:
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Authorized officer

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