Methods of using, storing and transporting metal protease enzyme in a stabilized form

Verfahren zur Anwendung, Lagern und Transport des Proteaseenzymes im stabilisierter Form

Méthode d’utilisation, d’approvisionnement et de transport de l’enzyme protease sous une forme stabilisée

References cited:

Note: Within nine months from the publication of the mention of the grant of the European patent, any person may give notice to the European Patent Office of opposition to the European patent granted. Notice of opposition shall be filed in a written reasoned statement. It shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).
[0001] The present invention relates to methods of using, storing and transporting a metal protease enzyme in stabili-
ized form and, more precisely, to methods of using, storing and transporting such enzyme in an aqueous solution con-
taining the enzyme. The present invention also relates to stabilized enzyme compositions.

[0002] Proteases are utilized in various fields of food industry and detergents. Where a protease is used in an aqueous
solution or a slurry, it is known that it loses its activity due to spontaneous autolysis or denaturation (pH, temperature,
etc.).

[0003] Recently, a method of synthesizing a dipeptide sweetener aspartame has been developed, using a conden-
sation reaction catalyzed by (neutral) metal proteases such as "thermolysin" (produced by Daiwa Chemical Co.) (K.
Oyama, Bioindustry, Vol. 2, No. 9, pp. 5 to 11, 1985). This aspartame synthesis process is in part based on earlier
thermolysin-catalyzed reaction of N-benzylloxy carbonyl-L-aspartic add (Z-L-Asp) with L-phenylalanine methyl ester
(L-PM). In those studies aqueous preparations of Z-L-Asp and L-PM, were incubated with a separately prepared stock
solution of thermolysin in water, and the time course of the reaction was studied. In the incubation experiments the
molar ratio of L-PM to Z-L-Asp was 2:1, and the initial molar concentration of Z-L-Asp was about 3750 times that of
thermolysin. Stabilization of thermolysin, however, was not an issue in these studies, nor could any conclusions on
thermolysin stabilization been drawn therefrom. Thus, proteases, particularly metal proteases, have become important
as synthetases for peptides in the industrial field. However, these enzymes are not stable, when the enzyme is used
or stored in an aqueous solution or slurry for a long period of time.

[0004] It is noticed that is known that the stability of thermolysin can be improved by immobilizing the enzyme, which
is then preferably being used in an organic medium (ethyl acetate). For instance, Yang et al. (Biotechnology and Bio-
engineering, Vol. 32, No. 5, 1988, pp 595-603) report some kinetic data on thermolysin-catalyzed synthesis of an as-
partame precursor by reaction of N-benzylloxy carbonyl-L-aspartic acid (Z-Asp) with excess of L-phenylalanine methyl
ester (L-PM) in ethyl acetate, which may be water-saturated. In particular, Yang et al. teach the use of immobilized
thermolysin, preferably by entrapment of the thermolysin in a hydrophobic cross-linked polyurethane. Yang et al. do
not present any evidence or suggestion for stabilisation of aqueous thermolysin-like (neutral) protease enzymes.

[0005] (Metal) Proteases are generally sold in the commercial market as a powder. A powdery enzyme easily forms
aerosols and therefore has a safety problem, since it can cause inflammation of the throat and eyes when adhered to
the mucous membranes of them. Allergic effects may also occur. Recently, therefore, (metal) protease enzymes are
handled as a liquid product or a slurry in view of the safety in handling it However, because of the stability of the enzyme
itself, such a liquid enzyme or a slurry of enzyme must be handled at a low temperature in usage, transportation and
storage, which generally results in higher costs (storage at low temperature, lower activity, etc.).

[0006] Furthermore, when a metal protease enzyme is used for the coupling reaction between N-benzylloxy-
carbonyl-L-aspartic acid (hereinafter referred to as "Z-Asp") and L- or DL-phenylalanine methyl ester (hereinafter referred to as
"PM") which is a process for production of α-L-aspartyl-L-phenylalanine methyl ester (hereinafter referred to as "As-
partame" or "APM"), the aspect of decreasing active enzyme concentration is also caused by the adsorption of enzyme
on crystals of the addition product N-benzyloxycarbonyl-Laspartyl-phenylalanine methyl ester (hereinafter referred to
"Z-APM.PM", e.g. as "Z-APM.DPM") besides the problem of stabilization of the enzyme. In case a metal protease,
such as "thermolysin" produced from Bacillus thermoproteolyticus Rokko is stored or used in the form of a solution or
slurry thereof, it is known that the enzyme rapidly deactivates if a certain concentration of calcium ions is not present.
However, in case of presence of calcium ions scaling may occur due to calcium carbonate formation in various parts
of duct lines and tanks, which will often have noticeable harmful effects in practical use of an enzyme.

[0007] The object to be attained by the present invention is to provide methods of providing a stabilized aqueous
neutral thermolysin-like metal protease enzyme, in a stabilized form without scaling, and to provide such stabilized
enzyme compositions.

[0008] Further subject matter to be attained by the present invention is to provide a method of improvement of re-
covering ratio which is achieved by the suppression of adsorption of enzyme on reaction product crystals (Z-APM.(D) 
PM).

[0009] According to the invention an N-protected amino acid (hereinafter referred to as "NPAA") is present in or
added to an aqueous solution or a slurry containing the protease in a molar concentration of said N-protected amino
acid of more than 30 times of that of the said enzyme. This is effective for preventing deactivation of metal protease
enzymes especially to noticeably improve the heat stability and storage stability of the enzymes. In particular, the effect
of N-benzylloxy carbonyl-amino acids (hereinafter referred to as "NZAA") has been found noticeable.

[0010] In addition, according to the invention a metal protease enzyme may noticeably be stabilized even in the
absence of excess calcium ions by the presence or addition of NPAA. The object of the present invention thus is
achieved in that an N-protected amino acid is present in or added to an aqueous solution or a slurry containing the
protease in at least 30 times the molar concentration thereof.
In addition, deactivation of a metal protease enzyme due to stirring in the presence of a slurry may also be noticeably inhibited by the presence or addition of NPAA, which is another characteristic feature.

Furthermore, according to the invention a metal protease enzyme may noticeably be stabilized and the recovery ratio may noticeably be improved by the presence of NPAA when such enzyme is used in the coupling reaction between Z-Asp and PM.

Specifically, the present invention advantageously can be used for storing or handling a metal protease solution or slurry in the presence of NPAA, or using this stabilizing effect during the enzymatic coupling of amino acids, especially in the preparation of aspartame. In accordance with the present invention, the storage stability of metal protease enzymes and the recovery ratio of metal protease enzymes may noticeably be improved due to a small amount of NPAA. Accordingly, decrease of the activity of these enzymes during storage is extremely small so that not only usage or storage of such enzyme for a long period of time has become possible but also the improvement of recovery ratio of metal protease enzymes can be achieved. This is especially economically advantageous for handling expensive enzymes.

Hereafter, the present invention will be explained in detail.

The metal proteases to be treated by the method of the present invention are not specifically confined, but thermolysin-like (neutral) proteases are especially effectively affected. As a metal protease, reference is made to, for example, one produced by Bacillus proteoleticus or Bacillus stearothermophilus, or one produced by expressing a gene thereof in other hosts. The enzyme solution or slurry to be used in the present invention may be a crude one containing many impurities or a purified one.

NPAA to be used as a stabilizing additive in the method of the present invention may be in the form of a salt, such as a sodium salt of NPAA or in the form of NPAA crystals. If desired, a solution or slurry containing various NPAA’s or a mixture comprising various NPAA’s may also be used. Anyway, NPAA is not specifically confined. The effects caused by NPAA are not whatsoever worsened due to presence of organic substances such as other amino acids or inorganic substances therein.

The AA’s in the NPAA’s may be all natural L-amino acids, but also D-amino acids; the acid group of the AA’s also may be in the form of an ester group, such as the benzyl ester.

As the N-protective group of NPAA, for example, benzyloxy carbonyl (Z), tertbutyloxycarbonyl (BOC), formyl (F), p-methoxybenzyloxycarbonyl (pMZ) are suitable. In particular, benzyloxy carbonyl-protected amino acids (ZAA’s) have been found to display an extremely excellent stabilizing effect. The enzyme retention percentage in the presence of NZAA can be as high as 90 % or more while that in the absence of it is about 32 % under the storage condition of 50 °C for a period of 5 hours. As examples of NZAA, mentioned are N-benzyloxy carbonyl-L-aspartic acid (also referred to as Z-Asp) where the amino acid is L-aspartic acid, N-benzyloxy carbonyl-L-glutamic acid (hereinafter referred to as Z-Glu) where the amino acid is L-glutamic acid, N-benzyloxy carbonyl-L-phenylalanine (hereinafter referred to as Z-Phe) where the amino acid is phenylalanine, and N-benzyloxy carbonylglycine (hereinafter referred to as Z-Gly) where the amino acid is glycine.

In systems having no excess of calcium ions, the enzyme retention percentage was about 44 % in the absence of NPAA under the storage condition of 40 °C for a period of 5 hours. In the same system, however, deactivation of a metal protease enzyme was noticeable inhibited by addition of NPAA. In particular, when NZAA was added to the system, almost no protease inactivation occurred. The results show that a metal protease enzyme solution is extremely stable during storage or can be used at room temperature or higher for a long period of time due to the presence of NPAA. Thus, the addition of NPAA is economically highly advantageous in treating and handling an expensive enzyme.

In accordance with the present invention, a metal protease enzyme is quite stable during handling even in the absence of excess calcium ions. Therefore, the present invention is also advantageous for preventing troubles to be caused by calcium ions, such as scaling.

It may be noticed here that JP-A-62269689 mentions stabilization of alkaliproteases useful for detergents by adding a specific amount of a reversible inhibitor such as chymostatin or Z-Phe. The present inventors, however, have observed that addition of NPAA’s to other protease enzymes such as papain, chymotrypsin or substilisin (all of them not being metal proteases), which according to the state of the art can be used in enzymatic APM-synthesis, does not result in any significant stabilization of the enzyme.

Furthermore, when the metal protease enzyme is used for the coupling reaction between Z-Asp and PM, the lowering of the enzyme recovery may noticeable be improved very much by the presence of NPAA in an aqueous solution or slurry of an enzyme.

The molar concentration of NPAA to be present in the aqueous solution or slurry of a metal protease enzyme for storing and transporting the enzyme in accordance with the present invention should be more than 30 times of that of the said enzyme, preferably more than 50 times of that of the enzyme.

On the other hand, when a metal protease enzyme is used for the coupling reaction between Z-Asp and PM and Z-Asp is used as NPAA, the molar concentration of Z-Asp that should be present in the aqueous solution or slurry of the enzyme after the coupling reaction should be more than 500 times of that of said enzyme, and preferably more
than 1000 times of that of said enzyme. In general this means that the concentration of Z-Asp should be more than 15 mmol/l, preferable more than 30 mmol/liter.

[0024] In the coupling reaction between Z-Asp and PM, the recovery ratio of enzyme was about 90% when the concentration of Z-Asp which was present in the aqueous solution or slurry of enzyme after the coupling reaction was 50 mmol/l (molar concentration 1724 times of the enzyme concentration).

[0025] On the other hand, the recovery ratio of enzyme was only about 50% when the concentration of Z-Asp which was present in the aqueous solution or slurry of enzyme after the coupling reaction was 10 mmol/l (molar concentration 345 times of the enzyme concentration).

[0026] As mentioned above, the metal protease enzyme may noticeably be stabilized by the presence of NPAA in an aqueous solution or a slurry of such enzyme, and the recovery ratio of such enzyme is also noticeably improved by the presence of NPAA in an aqueous solution or a slurry of enzyme when said enzyme is used for the coupling reaction between Z-Asp and PM.

[0027] Using of a metal protease enzyme in the coupling reaction for the production of aspartame is therefore very much improved if the molar concentration of an N-protected amino acid such as Z-Asp, is maintained at above 500 times that of the enzyme by adding an N-protected amino acid to the coupling reaction system during and/or in the final stage of the coupling reaction.

[0028] Next, the present invention will be explained in more detail by way of the following examples, which, however, are not intended to restrict the scope of the present invention.

Example 1:

[0029] 5 g of powdery Thermoase (crude thermolysin, trade name by Daiwa Chemical Co.; purity about 20 %) was suspended in one liter of water containing CaCl₂2H₂O (6.8 mmol/l) (the Thermoase concentration being 0.029 mmol/liter), and Z-Asp was added thereto until a determined concentration (2.62 mmol/liter or 24.7 mmol/liter, or in other words having a molar concentration of 90 respectively 850 times the concentration of enzyme) was reached with the pH of the resulting solution being adjusted to be 5.0 by adding 1 N NaOH aqueous solution. For testing the enzyme retention percentage (i.e. the active enzyme concentration), the enzyme solution was put in a separable flask having a volume of 2 liters and controlled to have a temperature of 70°C with stirring at 240 rpm. This was sampled at regular intervals and the amount of the remaining enzyme in each sample was measured by HPLC (High Pressure Liquid Chromatography). The results obtained are shown in Table 1.

<table>
<thead>
<tr>
<th>Time (hr)</th>
<th>Retention Percentage (%) of Enzyme</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No Addition</td>
</tr>
<tr>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>1</td>
<td>68</td>
</tr>
<tr>
<td>2</td>
<td>53</td>
</tr>
<tr>
<td>3</td>
<td>47</td>
</tr>
<tr>
<td>4</td>
<td>39</td>
</tr>
<tr>
<td>5</td>
<td>35</td>
</tr>
<tr>
<td>7</td>
<td>31</td>
</tr>
</tbody>
</table>

[0030] As is noted from Table 1 above, the enzyme was noticeably stabilized due to addition of Z-Asp.

Example 2:

[0031] To 100 ml of water containing 0.029 mmol/liter of powdery Thermoase (crude thermolysin, trade name by Daiwa Chemical Co.; purity about 20 %) and 10 mmol/liter of CaCl₂, were gradually added an amount of NPAA till a determined concentration (10 mmol/liter) was reached and 0.1 N NaOH aqueous solution were gradually added, the pH being adjusted to be 6.0. The thus prepared enzyme solution was put in a water bath controlled to have a temperature of 70°C. The enzyme solution was then sampled at regular intervals, and the amount of the remaining enzyme in each sample was measured by HPLC. The results obtained are shown in Table 2 below.

As is noted from Table 1 above, the enzyme was noticeably stabilized due to addition of Z-Asp.
As is noted from Table 2 above, the enzyme was noticeably stabilized due to addition of NPAA, especially NZAA (Z-Asp, Z-Asp-OBzl, Z-Glu, Z-Phe and Z-Gly).

Example 3:

5 g of powdery Thermoase was dissolved in one liter of water containing CaCl$_2$·2H$_2$O (CaCl$_2$ concentration = 0.6 mmol/liter) or of water containing Z-Asp (Z-Asp concentration = 30 mmol/liter), each with a Thermoase concentration of being 0.029 mmol/liter, and the pH of the resulting solutions was adjusted to be 6.0 with 1 N NaOH aqueous solution. The enzyme solution was put in a separable flask having a volume of 2.0 liters and controlled to have a temperature of 40°C with stirring at 200 rpm. This was sampled at regular intervals and the amount of the remaining enzyme in each sample was measured by HPLC. The results obtained are shown in Table 3.

The experiments of table 3 show that addition of Z-Asp is as effective as the use of 0.6 mmol/liter CaCl$_2$ solution; without such excess of Ca$^{2+}$ less scaling will occur.

Example 4:

An amount of NPAA was added to 100 ml of an aqueous solution containing 0.029 mmol/liter of powdery Thermoase.
Thermoase (not containing excess Ca) until a determined concentration (10 mmol/liter) was reached, in the same manner as in Example 2, the pH of the system being adjusted to be 6. This was put in a water bath controlled to have a temperature at 40°C.

[0038] The enzyme solutions were sampled at regular intervals, and the amount of the remaining enzyme was measured by HPLC. The results obtained are shown in Table 4 below.

<table>
<thead>
<tr>
<th>N-Protected Amino Acid (NPAA)</th>
<th>Retention Percentage(%) of Enzyme</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 hr</td>
</tr>
<tr>
<td>No Addition</td>
<td>100</td>
</tr>
<tr>
<td>Z-Asp</td>
<td>100</td>
</tr>
<tr>
<td>Z-Asp-OBzl(*)</td>
<td>100</td>
</tr>
<tr>
<td>Z-Glu</td>
<td>100</td>
</tr>
<tr>
<td>Z-Phe</td>
<td>100</td>
</tr>
<tr>
<td>Z-Gly</td>
<td>100</td>
</tr>
<tr>
<td>BOC-Asp</td>
<td>100</td>
</tr>
<tr>
<td>BOC-Glu</td>
<td>100</td>
</tr>
<tr>
<td>BOC-Phe</td>
<td>100</td>
</tr>
<tr>
<td>BOC-Gly</td>
<td>100</td>
</tr>
<tr>
<td>BOC-Cys-SBzl(*)</td>
<td>100</td>
</tr>
<tr>
<td>pMZ-Asp</td>
<td>100</td>
</tr>
<tr>
<td>pMZ-Glu</td>
<td>100</td>
</tr>
<tr>
<td>F-Asp-OBzl(*)</td>
<td>100</td>
</tr>
</tbody>
</table>

(*) Bzl : Benzyl Ester

[0039] As is noted from Table 4 above, the enzyme was noticeably stabilized due to addition of NPAA, especially NZAA (Z-Asp, Z-Asp-OBzl, Z-Glu, Z-phe and Z-Gly).

Example 5:

[0040] 5 g of powdery Thermoase (purity = about 20 %) was suspended in 750 g of water containing 0.1 % CaCl₂·2H₂O in separable flask having a volume of 2 liters, and 250 g of Celite (produced by Johns Manvill Corp., Grade: Standard Super-Cel) was added thereto to form a uniform slurry. This was controlled to have a temperature of 40°C, the Thermoase concentration being 0.029 mmol/liter.

[0041] Z-Asp was added to this until a determined concentration (30 mmol/liter) was reached, which was then adjusted to have pH of 7.3 with 1 N aqueous NaOH solution. This was stirred with a stirrer equipped with motor at a rate of 240 rpm and was sampled at regular intervals. The retention percentage of enzyme in each sample was measured by HPLC. The results obtained are shown in Table 5 below.

<table>
<thead>
<tr>
<th>Time</th>
<th>Retention Percentage (%) of Enzyme</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No Addition of Z-Asp</td>
</tr>
<tr>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>1</td>
<td>85</td>
</tr>
<tr>
<td>3</td>
<td>70</td>
</tr>
<tr>
<td>5</td>
<td>60</td>
</tr>
<tr>
<td>7</td>
<td>55</td>
</tr>
</tbody>
</table>

[0042] As is noted from Table 5 above, the enzyme which is influenced and deactivated in the slurry was noticeably stabilized by addition of Z-Asp.
Example 6:

[0043] 5 g of powdery Protease TD (product by Amano Pharmaceutical Co.; metal protease produced by Bacillus stearothermophilus; purity = about 20 %) was suspended in one liter of water containing 0.1 % CaCl₂-H₂O, with the concentration of Protease TD being 0.029 mmol/liter. Z-Asp was added to this until a determined concentration (10 mmol/liter) was reached, which was then adjusted to have pH of 5.0 with 1 N aqueous NaOH solution. The resulting solution was put in a separable flask having a volume of 2 liters and controlled to have a temperature of 70°C with stirring at 240 rpm. This was sampled at determined intervals, and the amount of the remaining enzyme in each sample was measured by HPLC. The results obtained are shown in Table 6 below.

<table>
<thead>
<tr>
<th>Time (hr)</th>
<th>Retention Percentage (% of Enzyme)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No Addition of Z-Asp</td>
</tr>
<tr>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>1</td>
<td>84.5</td>
</tr>
<tr>
<td>2</td>
<td>71.9</td>
</tr>
<tr>
<td>3</td>
<td>62.9</td>
</tr>
<tr>
<td>4</td>
<td>53.4</td>
</tr>
<tr>
<td>5</td>
<td>45.5</td>
</tr>
<tr>
<td>7</td>
<td>36.0</td>
</tr>
</tbody>
</table>

Example 7:

[0045] 943 g of Z-Asp aqueous solution (the amount of Z-Asp was 1.0 mol) and 1258 g of DL-PM aqueous solution (the amount of PM was 2.0 mol) was mixed, warmed to 40°C, and the aqueous solution was prepared by the adjustment of pH of this solution using 25% NaOH solution, and this solution was used as the substrate solution.

[0046] Separately 1.28 g of CaCl₂·2H₂O and 127 g of NaCl was dissolved in 1242 g of purified water, and 45 g of thermoase was dissolved in this aqueous solution, and this aqueous solution was used as the aqueous solution of an enzyme.

[0047] The substrate solution and the aqueous solution were mixed in the separable-flask of 5 liters which was settled in a water-bath maintained at 40°C, and pH of this mixed solution was adjusted to be 6.0 and the coupling reaction between Z-Asp and PM was started.

[0048] The concentration of thermoase was 0.029 mmol/l as in the previous examples.

[0049] The stirring of the mixed solution was carried out with stirring speed of 130 rpm till 3.5 hours after the start of the coupling reaction, and was carried out with stirring speed of 30 rpm during 3.5 hours to 7 hours.

[0050] The yield of Z-APM.(D)PM based on the starting amount of Z-Asp was 80.5% after 7 hours, and the concentration of Z-Asp which remained after the coupling reaction was 55.9 mmol/l.

[0051] The thermoase which were recovered after the coupling reaction was 41.4 g and the recovery ratio was about 92%.

Comparative Example 1

[0052] 943 g of Z-Asp aqueous solution (the amount of Z-Asp was 1.0 mol) and 1572 g of DL-PM aqueous solution (the amount of PM was 2.5 mol) was mixed, warmed to 40°C, and the aqueous solution was prepared by the adjustment of pH of this solution using 25% NaOH solution, and this solution was used as the substrate solution.

[0053] Separately 1.28 g of CaCl₂·2H₂O and 127 g of NaCl was dissolved in 1242 g of purified water, and 45 g of thermoase was dissolved in this aqueous solution, and this aqueous solution was used as the aqueous solution of an enzyme.

[0054] In this Comparative Example the higher amount of PM as compared with Example 7 results in a lower amount of Z-Asp remaining after the coupling. It can be seen that this results in a significantly decreased thermoase recovery ratio.

[0055] The substrate solution and the aqueous solution were mixed in the separable-flask of 5 liters which was set in the water-bath maintained at 40°C, and pH of this mixed solution was adjusted to be 6.0 and the coupling reaction between Z-Asp and PM was started.
The stirring of the mixed solution was carried out with stirring speed of 130 rpm till 3.5 hours after the start of the coupling reaction, and was carried out with stirring speed of 30 rpm during 3.5 hours to 7 hours.

The yield of Z-APM/(D)PM based on the starting amount of Z-Asp was 95.2% after 7 hours, and the concentration of Z-Asp which remained after the coupling reaction was 10.5 mmol/l.

The thermoase which were recovered after the coupling reaction was 19.5 g, and the recovery ratio was about 44%.

Claims

1. Method of providing a stabilized aqueous neutral thermolysin-like metal protease enzyme, characterized in that an N-protected amino acid is present in or added to an aqueous solution or a slurry containing the protease in a molar concentration of said N-protected amino acid of more than 30 times of that of the said enzyme.

2. Method of providing a stabilized aqueous protease enzyme according to claim 1, characterized in that the N-protected amino acid is an N-benzyloxycarbonylamino acid.

3. Method of providing a stabilized aqueous protease enzyme according to any of claims 1 or 2, characterized in that the protease is thermolysin.

4. Method of providing a stabilized aqueous protease enzyme according to any of claims 1 to 3, characterized in that the molar concentration of said N-protected amino acid is more than 500 times of that of the said enzyme.

5. Method of enzymatic peptide coupling of an N-protected amino acid and an amino acid ester using an aqueous neutral thermolysin-like metal protease enzyme, characterized in that the molar concentration of the N-protected amino acid being coupled is more than 500 times of that of the said enzyme.

6. Method of enzymatic peptide coupling according to claim 5, characterized in that the molar concentration of the N-protected amino acid being coupled is maintained at above 500 times that of the said enzyme by adding said N-protected amino acid to the coupling reaction system during and/or in the final stage of the coupling reaction.

7. Method of enzymatic peptide coupling according to any of claims 5 or 6, characterized in that N-benzyloxycarbonyl-aspartic acid and phenylalanine methyl ester are being coupled.

8. Aqueous neutral thermolysin-like metal protease enzyme composition only containing water, a neutral thermolysin-like protease enzyme and an N-protected amino acid, the molar concentration of said N-protected amino acid being more than 30 times of that of the said enzyme, optionally calcium chloride and some pH-adjusting chemical.

9. Enzyme composition according to claim 8, characterized in that the enzyme is thermolysin.

Patentansprüche


2. Verfahren zum Bereitstellen eines stabilisierten wässrigen Proteaseenzymes nach Anspruch 1, dadurch gekennzeichnet, daß die N-geschützte Aminosäure eine N-Benzylxycarbonylaminosäure ist.

3. Verfahren zum Bereitstellen eines stabilisierten wässrigen Proteaseenzymes nach einem der Ansprüche 1 oder 2, dadurch gekennzeichnet, daß die Protease Thermolysin ist.

4. Verfahren zum Bereitstellen eines stabilisierten wässrigen Proteaseenzymes nach einem der Ansprüche 1 bis 3, dadurch gekennzeichnet, daß die molare Konzentration der N-geschützten Aminosäure mehr als 500 mal höher als die des Enzyms ist.
5. Verfahren zur enzymatischen Peptidkuppung einer N-geschützten Aminosäure und eines Aminosäureesters mit einem wässrigen neutralen Thermolysin-ähnlichen Metallproteaseenzym, **dadurch gekennzeichnet**, daß die molare Konzentration der zu kuppelnden N-geschützten Aminosäure mehr als 500 mal höher als die des Enzyms ist.

6. Verfahren zur enzymatischen Peptidkuppung nach Anspruch 5, **dadurch gekennzeichnet**, daß die molare Konzentration der zu kuppelnden N-geschützten Aminosäure bei etwa 500 mal höher gehalten wird als die des Enzyms, wobei die N-geschützte Aminosäure zu dem Kuppelungsreaktionssystem während und/oder im Endzustand der Kuppungsreaktion zugegeben wird.

7. Verfahren zur enzymatischen Peptidkuppung nach einem der Ansprüche 5 oder 6, **dadurch gekennzeichnet**, daß N-Benzylxocarbonylasparaginsäure und Phenylalaninemethylster gekuppelt werden.


9. Enzymzusammensetzung nach Anspruch 8, **dadurch gekennzeichnet**, daß das Enzym Thermolysin ist.

**Revendications**

1. Méthode de préparation d'un enzyme protéase métallique de type thermolysine neutre aqueuse stabilisée, **caractérisée en ce que** un acide aminé N-protégé est présent dans ou ajouté à une solution ou une bouillie aqueuse avec une concentration molaire dudit acide aminé N-protégé plus de 30 fois celle dudit enzyme.

2. Méthode de préparation d'un enzyme protéase aqueuse stabilisée selon la revendication 1, **caractérisée en ce que** l'acide aminé N-protégé est acide N-benzyloxycarbonylaminé.

3. Méthode de préparation d'un enzyme protéase aqueuse stabilisée selon l'une quelconque des revendications 1 ou 2, **caractérisée en ce que** la protéase est la thermolysine.

4. Méthode de préparation d'un enzyme protéase aqueuse stabilisée selon l'une quelconque des revendications 1 à 3, **caractérisée en ce que** la concentration molaire dudit acide aminé N-protégé est plus de 500 fois supérieure à celle dudit enzyme.

5. Méthode de couplage peptidique enzymatique d'un acide aminé N-protégé et d'un ester d'acide aminé utilisant un enzyme protéase métallique de type thermolysine neutre aqueuse, **caractérisée en ce que** la concentration molaire de l'acide aminé N-protégé en cours de couplage est plus de 500 fois supérieure à celle dudit enzyme.

6. Méthode de couplage peptidique enzymatique selon la revendication 5, **caractérisée en ce que** la concentration molaire de l'acide aminé N-protégé en cours de couplage est maintenue plus de 500 fois supérieure à celle dudit enzyme en ajoutant ledit acide aminé N-protégé au système de réaction de couplage pendant et/ou dans le stade final de la réaction de couplage.

7. Méthode de couplage peptidique enzymatique selon l'une quelconque des revendications 5 ou 6, **caractérisée en ce que** l'acide N-benzyloxycarbonylaspartique et l'ester méthylé de la phénylalanine sont couplés.

8. Composition d'enzyme protéase métallique de type thermolysine neutre aqueuse ne contenant que de l'eau, un enzyme protéase métallique de type thermolysine neutre et un acide aminé N-protégé, la concentration molaire dudit acide aminé N-protégé étant supérieure à plus de 30 fois celle dudit enzyme, et facultativement du chlorure de calcium et un produit chimique d'ajustement du pH.

9. Composition d'enzyme selon la revendication 8, **caractérisée en ce que** l'enzyme est la thermolysine.