Purification and some properties of carboxynorspermidine synthase participating in a novel biosynthetic pathway for norspermidine in *Vibrio alginolyticus*

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Carboxynorspermidine synthase, mediates the nicotinamide-nucleotide-linked reduction of the Schiff base H₂N(CH₂)₂N=CHCH₂CH(NH₂)COOH. This is formed from L-aspartic β-semialdehyde (ASA) and 1,3-diaminopropane (DAP) and is reduced to carboxynorspermidine [H₂N(CH₂)₂NH(CH₂)₂CH(NH₂)COOH], an intermediate in the novel pathway for norspermidine (NSPD) biosynthesis. The enzyme was purified to apparent homogeneity from *Vibrio alginolyticus* and characterized. The overall purification was about 1800-fold over the crude extract, with a yield of 33%. The enzyme displayed an apparent Mₐ of 93500 ± 1000 by gel filtration and 45100 ± 500 by SDS-PAGE, indicating that the native form is probably composed of two subunits of similar size. The specific activity of the purified enzyme was 31.0 pmol carboxynorspermidine produced min⁻¹ (mg protein)⁻¹. The enzyme was activated by dithiothreitol, and inhibited by SH-reactive compounds. The pH and temperature optima were 7.25–7.5 and 37 °C, respectively. The Kₘ value for the Schiff base was 4–68 mM, measured by varying the ASA concentration while keeping the DAP concentration constant. Putrescine was slightly active as a substrate, forming carboxyspermidine (at about 7% of the rate of DAP), but ethylenediamine, cadaverine and D-ASA were inert. The Kₘ value for NADPH was 1.51 mM. NADH was a much poorer cofactor than NADPH.

When *V. alginolyticus* was grown in the presence of 5 mM-NSPD, the specific activity of this enzyme was reduced by ~70%. NSPD also repressed two other enzymes responsible for its biosynthesis, 2,4-diaminobutyrate decarboxylase and carboxynorspermidine decarboxylase.

Introduction

The polyamine distribution patterns of members of the genera *Vibrio* and *Listonella* (formally allocated in *Vibrio*) are unique in that norspermidine (NSPD), a homologue of spermidine (SPD) with two C₃ chains, is a major triamine species, whereas putrescine (PUT) and SPD are found only in small amounts under normal growth conditions (Yamamoto et al., 1983, 1991). Experiments using radiolabelled L-aspartic acid have shown that, in some *Vibrio* species, it is converted to L-aspartic β-semialdehyde (ASA), which acts as an aminopropyl group donor in NSPD biosynthesis (Yamamoto et al., 1986a):

\[
\text{L-Aspartic } \beta\text{-semialdehyde} + \text{1,3-diaminopropane} + \text{NADPH} + \text{H}^+ \rightarrow \text{Carboxynorspermidine} + \text{NADP}^+ + \text{H}_2\text{O} \quad (1)
\]

\[
\text{Carboxynorspermidine} \rightarrow \text{Norspermidine} + \text{CO}_2 \quad (2)
\]

An analogous reaction using PUT as a diamine precursor instead of DAP, has been demonstrated for SPD biosynthesis in certain bacteria (Tait, 1976, 1985) and seedlings of a plant (Srivenugopal & Adiga, 1980). However, the enzymes responsible have been studied only in crude or partially purified preparations in experiments aimed primarily at delineating the pathway. In these studies, the Schiff base spontaneously or non-enzymically formed between ASA and PUT has been postulated as a possible substrate in an NADPH-dependent enzyme reaction.

In our previous study (Nakao et al., 1990), C-NSPD decarboxylase, the pyridoxal phosphate-dependent enzyme that catalyses reaction 2, was purified from a
slightly halophilic *Vibrio alginolyticus* to homogeneity and some of its properties were studied. This paper describes the purification and characterization of the enzyme which catalyses the reductive condensation of C-NSPD and ASA in *V. alginolyticus* and some of its properties were studied. This paper

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Methods

Chemicals and materials. NADP$^+$, NAD$^+$ and their reduced forms, were purchased from Oriental Yeast (Japan); allylglycine (d- and l-), bovine albumin (A-4378), dithiothreitol (DTT) and standard marker proteins for SDS-PAGE were from Sigma; DEAE-Sepharose CL-6B, Sephacyr S-200 HR and Blue Sepharose CL-6B were from Pharmacia LKB; and hydroxyapatite was from Wako (Japan). ASA and its optical isomer were prepared by ozonolysis of the corresponding forms of allylglycine, purified and stored in 4 M-HCl at 4°C (Black & Wright, 1955). The yield of ASA has been reported to be 80% and the amount stated is on the basis of this yield. Before use, a small sample of this solution was adjusted to pH 6.5-6.7 using KHCO$_3$, with stirring. C-NSPD and C-SPD were prepared as previously described (Nakao et al., 1990). Human $\gamma$-globulin ($M_\theta = 160000$), BSA (67000) and ovalbumin (45000) from Sigma, and yeast glucose-6-phosphate dehydrogenase (128000) and yeast hexokinase (104000) from Oriental Yeast were used as standard proteins for gel filtration. All other chemicals were of reagent quality.

Bacterial strain and growth conditions. *Vibrio alginolyticus* ATCC 17749 (type strain) was grown for 6-8 h as described previously (Nakao et al., 1989), and the cell pellets collected were stored at $-30$°C until enzyme purification.

Enzyme assay. Enzyme activity was determined by following the formation of C-NSPD from the Schiff base formed between ASA and DAP. The Schiff base was prepared by incubating 175 µl 250 mM-potassium phosphate buffer, pH 7.5, 50 µl 150 mM-DAP and 50 µl 150 mM-ASA at 37°C for 5 min. The assay mixture was then adjusted to a final volume of 500 µl by adding 30 µl bovine albumin solution (30 mg ml$^{-1}$), 50 µl 50 mM-NADPH, 50 µl 200 mM-DTT, distilled water and enzyme protein (0-100 µg). After incubation at 37°C for 30 min, the reaction was terminated with 0-4 ml 20% (w/v) HClO$_4$, and the appropriate amount of trypophan (25-100 nmol) was added as an internal standard. The supernatant after centrifugation at 1500g for 10 min was transferred to a reaction vial for derivatization of C-NSPD formed, followed by GC analysis with nitrogen-selective detection (Yamamoto et al., 1986a). The identity of C-NSPD was confirmed by GC-MS (Yamamoto et al., 1986a). The reproducibility of the assay method was assessed by calculating the recoveries of known amounts of C-NSPD (10-125 nmol) added to the assay mixtures denatured by the prior addition of HClO$_4$. The recovery for C-NSPD was $94 \pm 5\%$ ($n = 10$). The formation of C-NSPD was linear with respect to time (10-90 min) and the amount of enzyme protein present. The decrease in A$_{340}$ of NADPH was also confirmed with some enzyme preparations, but in these cases the cofactor was reduced to 0-5 mM. One unit of the enzyme was defined as the amount catalysing the formation of 1 µmol C-NSPD min$^{-1}$ under standard assay conditions.

Enzyme purification. Unless otherwise noted, all procedures were done at 2-4°C, and all dialyses were against 100 vols of the specified buffers for 12 h. A crude estimate of the protein content of column effluents was made by measuring A$_{280}$. For accurate measurement of the protein content of pooled fractions and final enzyme preparation, the Lowry method, or that of Bradford (1976) was used, depending upon the amount of material available.

Frozen cells (200 g wet wt) of *V. alginolyticus* were suspended in 800 ml 20 mM-Tris/HCl, pH 7.5, containing 1 mM-DTT (buffer A), followed by disruption with an ultrasonic homogenizer in a salt/ice bath. The cell debris was removed by centrifugation at 4000g for 30 min, and a small portion of the supernatant (crude extract) was saved for determining the activity after dialysis against two changes of buffer A. The supernatant was fractionated with ammonium sulphate (50-70%), and the precipitate was dissolved in a minimum volume of buffer A. After dialysis of this solution against two changes of the same buffer, the precipitate that appeared during dialysis was removed by centrifugation. The clear supernatant (60 ml) was divided into two equal portions, each of which was applied to a DEAE-Sepharose CL-6B column (2.2 x 25 cm) equilibrated with buffer A. After washing the column with buffer A containing 0.2 M-NaCl (300 ml), elution was done with a 500 ml linear gradient of 0-2-0.4 M-NaCl in buffer A at a flow rate of 15 ml h$^{-1}$. Active fractions eluting at 0.31-0.33 M-NaCl were combined (40 ml) and then dialysed against 10 mM-potassium phosphate, pH 7.5, containing 1 mM-DTT (buffer B). The dialysed solution was placed on a column (2.65 x 204 cm) of hydroxyapatite equilibrated with buffer B. After washing the column with the same buffer (330 ml), the enzyme was eluted with a 500 ml linear gradient of 10-50 mM-potassium phosphate, pH 7.5, at a flow rate of 10 ml h$^{-1}$. Active fractions eluting at 31-38 mM-potassium phosphate were pooled (100 ml) and then dialysed against buffer B. The enzyme solution was applied to a Blue Sepharose CL-6B column (2 x 15 cm) equilibrated with buffer B containing 0.1 M-NaCl, and the column was washed with two volumes of the same buffer. The enzyme was eluted with a 300 ml linear gradient of 0-1 M-NaCl at a flow rate of 10 ml h$^{-1}$. Active fractions eluting at 0.31-0.49 M-NaCl were pooled (70 ml), and dialysed against buffer A containing 0.02% NaN$_3$. The dialysed solution was concentrated to about 0.5 ml in an Amicon 8050 ultrafiltration cell with a YM-10 membrane, and subsequently in an Amicon Centricron 30, and was then stored at 4°C.

Analytical methods for the purified enzyme. Electrophoresis on a 7.5% (w/v) polyacrylamide slab gel under non-denaturing conditions was done as described by Williams & Reisfeld (1964) except that DTT was added to buffers for gel preparation to give a final concentration of 1 mM. For monitoring the enzyme activity after electrophoresis, the gel was sliced and macerated in buffer A, and left overnight at 4°C to elute the enzyme. SDS-PAGE was done in a 10% (w/v) polyacrylamide gel according to Laemmli (1970). Before loading, the enzyme sample was heated for 3 min with 2% (w/v) SDS and 5% (w/v) 2-mercaptoethanol in a boiling-water bath. Proteins were visualized by staining with Coomassie Brilliant Blue R-250.

Analytical gel filtration for $M_\theta$. Determination of the native enzyme was done by the use of a Sephacryl S-200 HR column (2.1 x 88 cm) calibrated with the standard proteins, as described by Andrews (1965). The column was equilibrated and eluted with 20 mM-Tris/HCl, pH 7.5, containing 0.4 M-NaCl, 1 mM-DTT and 0.02% NaN$_3$.

Determination of $pI$. A complete loss of enzyme activity was observed upon both isoelectric focusing and chromatofocusing. Therefore, the pI of the purified enzyme was determined by a batchwise method using DEAE-Sepharose CL-6B according to the supplier's instructions. Briefly, 1 ml resin was placed in each of ten 5 ml test tubes, equilibrated with 0.01 M-sodium acetate buffer adjusted to pH 3.8-4.7 with 0.1 pH unit increase per tube, and then suspended in 1 ml of the same buffer containing 1 mM-DTT. Into each suspension was stirred 2.5 µg (10 µl) of the purified enzyme. After stirring for 10 min at 4°C, the enzyme activity in the supernatants was measured.
Table 1. Purification of C-NSPD synthase from V. alginolyticus

A typical purification from 200 g wet wt of cells is shown.

<table>
<thead>
<tr>
<th>Step</th>
<th>Total protein (mg)</th>
<th>Total activity (U)</th>
<th>Specific activity [U (mg protein)-¹]</th>
<th>Purification (-fold)</th>
<th>Recovery (%)</th>
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<tr>
<td>Crude extract</td>
<td>4445</td>
<td>74·1</td>
<td>0·0167</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>50-70% Ammonium sulphate</td>
<td>2649</td>
<td>71·9</td>
<td>0·0271</td>
<td>1·6</td>
<td>97</td>
</tr>
<tr>
<td>DEAE-Sepharose CL-6B</td>
<td>163</td>
<td>101·4</td>
<td>0·662</td>
<td>37·2</td>
<td>137</td>
</tr>
<tr>
<td>Hydroxyapatite</td>
<td>12·8</td>
<td>70·4</td>
<td>5·5</td>
<td>329·3</td>
<td>95</td>
</tr>
<tr>
<td>Blue Sepharose CL-6B</td>
<td>0·8</td>
<td>24·8</td>
<td>31·0</td>
<td>1856</td>
<td>33</td>
</tr>
</tbody>
</table>

Results

Establishment of enzyme assay conditions

Preliminary experiments demonstrated that enzyme activity exhibited an upward curvature with respect to the amount of protein, whenever the components required were added consecutively in an ordinary manner, even if the order of addition was changed. Further, activity decreased in proportion to the concentration of ASA, even when DAP was present in a large excess. These results suggested that the enzyme might be directly inactivated by free ASA. It was also suspected that ASA could react with the NH₂ group of the cofactor. To overcome these problems, prior formation of the Schiff base was introduced. However, since it was difficult to measure the absolute yield of the Schiff base formation, we assumed that under the conditions used, the reaction between ASA and DAP runs rapidly to completion, and that there is little chance that both DAP NH₂ groups react with ASA. In fact, Schiff base formation, as judged by the amount of C-NSPD formed, reached a maximum within a short period of time (3 min) and did not change over an incubation of at least 20 min. This may also indicate that the reversibility of the Schiff base reaction was slight. Furthermore, the enzyme appeared to be stabilized by bovine albumin. Enzyme preparations thus stabilized displayed a linear response passing through the origin with respect to the amount of protein and incubation time.

Purification of C-NSPD synthase

The synthase was a soluble intracellular enzyme. When purified, the final yield of enzyme activity was 33% with a specific activity of 31·0 U (mg protein)-¹ (Table 1), which represented an approximately 1800-fold purification over the crude extract. DEAE-Sepharose chromatography led to an increase (36%) in the total enzyme activity of the crude extract, with a remarkable increase in purification. The increase in the total activity suggests that an enzyme inhibitor(s) may have been removed in the ammonium sulphate fraction. Hydroxyapatite chromatography also resulted in a high improvement in purification with a good yield, but there were still several minor contaminating proteins. The enzyme was further purified with affinity chromatography through a Blue Sepharose column. Unlike C-SPD synthase from Lathyrus sativus seedlings (Srivenugopal & Adiga, 1980), the V. alginolyticus enzyme was so tightly bound to the column that it was not eluted with 5 mM-NAD⁺ or 1-5 mM-NADP⁺. A gradient elution with NaCl from 0·1 to 1 M dislodged the enzyme and appeared to be highly effective in obtaining the pure enzyme.

Properties of the purified enzyme

Stability. The final preparation at a protein concentration of about 1·6 mg ml⁻¹ in 20 mM-Tris/HCl, pH 7·5, containing 1 mM-DTT and 0·02% NaN₃, lost about 15% of its activity when stored at 4°C for a week. Addition of NADPH to this solution gave no significant stabilization of the enzyme. In contrast, DTT, at a concentration (1 mM) much lower than that needed for enzyme activation, had a protective effect on the enzyme during purification and storage. Freezing and thawing of the enzyme resulted in almost complete loss of activity. The enzyme was unstable in buffers of pH < 6·5; at pH 6·5, 50% loss of activity was observed within 12 h at 4°C.

Purity and Mₜ determination. On native PAGE, the purified enzyme migrated as a single species, which coincided with the enzyme activity, and it also gave only one band on SDS-PAGE. On the calibrated column of Sephacryl S-200 HR, the Mₜ was estimated to be 93500 ± 1000. The Mₜ corresponding to the SDS-PAGE single band was estimated to be 45100 ± 500. Thus, the enzyme appears to consist of two subunits of identical or very similar Mₜ.

Cofactor requirements. The enzyme was completely
inactive unless NADPH or NADH was added. Plots of enzyme activity as a function of NADPH or NADH concentration displayed typical Michaelis-Menten kinetics and double-reciprocal plots were used to estimate $K_m$ and $V$ values. NADPH gave a $K_m$ of 1.51 mM, with a $V$ of 31.0 pmol C-NSPD formed min$^{-1}$ (mg protein)$^{-1}$, whereas NADH gave a $K_m$ of 2.97 mM, with a $V$ of 13.5 pmol C-NSPD formed min$^{-1}$ (mg protein)$^{-1}$. These data show that the former is the preferred cofactor under the conditions used. Use of DTT became increasingly essential as the protein became purer. Without addition of DTT to the assay mixture, only 7% of the maximum activity was detected in the purified enzyme. The addition of DTT to the assay mixture, however, greatly activated the enzyme, maximum activation (about 14-fold) being observed at 20 mM.

**Reversibility of the reaction.** Incubation of the purified enzyme in the pH range 6–8, with cofactor (NADP$^+$ or NAD$^+$) concentrations up to 5 mM and substrate (C-NSPD) concentrations up to 10 mM produced no observable increase in $A_{340}$ (indicative of C-NSPD oxidation).

**Effects of pH and temperature.** Activity was maximal in the range pH 7.25–7.5, assuming that there was little change in Schiff base formation in the pH range tested (6.25–8.0). The alkaline side of the pH optimum diminished gradually, while the acid side dropped off relatively sharply; 85% and 18% of the maximum activity being observed at pH 8.0 and 6.25, respectively. This is mainly due to the instability of the enzyme in acid, as described above. The optimum temperature for enzyme activity in the standard assay was 37 °C; 78% and 59% of the maximum activity remained at 30 °C and 45 °C, respectively, but no activity was observed above 50 °C.

**Saturation kinetics and substrate specificity.** To determine the kinetics of the enzyme for the ASA–DAP Schiff base, the concentration of ASA was varied from 0.25–15 mM, while the concentration of DAP was kept at 15 mM. The $K_m$ and $V$ values (determined from Lineweaver–Burk plots) were 4.68 mM and 35.0 pmol C-NSPD min$^{-1}$ (mg protein)$^{-1}$, respectively. Substitution of D-ASA for L-ASA resulted in no appearance of the peak corresponding to C-NSPD, while the concentration of DAP was kept at 15 mM. PUT (10 mM) as substrate yielded C-SPD at a rate only 7% of that of DAP. Ethylenediamine, 1,2-diaminopropane and cadaverine were totally inactive as judged by $A_{340}$ measurements. When added to the standard assay mixture at a high concentration (10 mM), these diamines caused 10–20% inhibition of C-NSPD formation.

**Inhibitors and other properties.** Even in the presence of 20 mM-DTT, 5 mM-N-ethylmaleimide and 5 mM-iodoacetamide inhibited enzyme activity by 83% and 24%, respectively. This, along with stimulation of the enzyme by DTT, indicates that C-NSPD synthase is a sulphhydryl enzyme. Enzyme activity was inhibited 51% and 10% by 5 mM-NADP$^+$ and 10 mM-NADP, respectively, but NAD$^+$, ATP and SPD were not inhibitory at these concentrations. Na$^+$ and K$^+$ at 1 mM concentration had no effect on enzyme activity. There was no evidence for activity being dependent on metal cations such as Ca$^{2+}$, Mg$^{2+}$, Fe$^{3+}$, Fe$^{2+}$, Cu$^+$, Cu$^{2+}$, Mn$^{2+}$ and Zn$^{2+}$, and these cations were not inhibitory. EDTA (1 mM) showed no significant stimulatory or inhibitory effect on enzyme activity. The pI of the enzyme was judged to be 4.2–4.3 since it released from Sepharose CL-6B at this pH range.

**Effect of addition of NSPD to growth medium on the activities of the enzymes involved in NSPD biosynthesis.**

To determine whether exogenous NSPD would affect the expression of NSPD biosynthetic enzymes, the organism was grown in the absence (control) or presence of 5 mM-NSPD, and the specific activities of DABA decarboxylase, C-NSPD synthase and C-NSPD decarboxylase were measured in the crude extracts. For comparison, SPD was also examined. The organism was grown for 6 h in 200 ml synthetic medium containing 0.5% NaCl (Yamamoto et al., 1983), and all crude extracts, including the control, were prepared in 20 mM-Tris/HCl, pH 7.5, containing 1 mM-DTT and 0.1 mM-EDTA, then dialyzed for 12 h against two changes of the same buffer containing 0.5 M-NaCl. The inclusion of NaCl in the dialysis buffer was effective in almost completely removing exogenous NSPD or SPD. The activities of the two decarboxylases were assayed as previously described.
(Nakao et al., 1989, 1990). As shown in Table 2, addition of NSPD significantly decreased the specific activity in all enzymes, compared with the control. Interestingly, SPD also showed a repressive effect on these enzymes, but to a lesser extent than NSPD. The above findings were not the result of an inhibition of stimulation of the bacterial growth, since the cultures grown in the presence of NSPD or SPD showed no significant difference in total cellular protein compared with the control.

Discussion

Under normal growth conditions, DAP was first detected in cells 2-5 h (the period of rapid growth) after inoculation. This appearance of DAP coincided with enhanced production of NSPD, the end product in this pathway (Yamamoto et al., 1986b). The enzyme which decarboxylates L-DABA to DAP has previously been purified (Nakao et al., 1989) and the biosynthetic pathway for L-DABA in V. alginolyticus is currently under investigation. On the other hand, appreciable incorporation of the radiolabel from L-[14C]aspartic acid into NSPD demonstrates that this bacterium can produce ASA from L-aspartic acid (Yumoto et al., 1986b). Thus, the present study was initiated to purify and characterize the enzyme which catalyses the reductive condensation of ASA and DAP to give C-NSPD, as an intermediate of NSPD biosynthesis. However, the enzyme was inhibited whenever it was incubated with free ASA, as has been reported for Escherichia coli aspartase (EC 4.3.1.1) (Yumoto et al., 1982). In this study, prior formation of the Schiff base was therefore adopted to facilitate purification of the enzyme. This inhibition by free ASA was probably due to its modification of the ε-NH₂ group of lysine and/or the SH group of cysteine near to, or in, the active site of the enzyme. Therefore, it is conceivable that, when ASA is being formed from L-aspartic acid in cells, the aldehyde is kept in a masked state to prevent interaction with the synthase. Once the Schiff base is formed with DAP, it will be accepted by C-NSPD synthase.

The highly effective activation by DTT is a distinctive feature of this enzyme. The DTT requirement and the optimal pH are similar to those of C-SPD synthase, an analogous enzyme which has been partially purified (50-fold) from L. sativus (Srivengopal & Adiga, 1980). While C-SPD synthases from this plant and from Micrococcus denitrificans and Rhodopseudomonas spheroides (Tait, 1976) can utilize only NADPH as a cofactor, the purified C-NSPD synthase from V. alginolyticus can utilize NADH in addition to NADPH, although it shows a marked preference for the latter. Furthermore, the L. sativus C-SPD synthase utilizes DAP with only 30% efficiency of PUT, at an equivalent concentration, indicating that the enzymes clearly differ in substrate specificity. The purified C-NSPD synthase did not catalyse the oxidative cleavage of C-NSPD to ASA and DAP (reverse reaction) under the conditions examined, whether NADPH or NADH was used as a cofactor.

Previously, C-NSPD decarboxylase has been shown to act on C-SPD at a rate comparable with that of C-NSPD to produce SPD (Nakao et al., 1990). Yet the cellular content of SPD, when the bacterium is grown in a defined medium free from SPD, is less than one-tenth that of NSPD at any growth phase (Yamamoto et al., 1986b). However, the present study indicated that C-NSPD synthase was far more active with the Schiff base formed between DAP and ASA than with that between PUT and ASA. Moreover, formation of SPD in this bacterium can reasonably be attributed to two enzymes responsible for NSPD biosynthesis, as the activity of the classical aminopropyltransferase (EC 2.5.1.16), which utilizes decarboxylated S-adenosylmethionine as an aminopropyl donor to give SPD, could not be detected in cell extracts.

Microbial enzymes involved in the conventional route for polyamine biosynthesis are under regulatory control by end products such as SPD (Tabor & Tabor, 1985). This may also be true of the V. alginolyticus enzymes. The reduction in specific activity of all three enzymes involved in NSPD biosynthesis when NSPD is added to the growth medium, indicates its physiological importance in this bacterium. The repression of NSPD biosynthetic enzymes by SPD is suggested by the observation that its presence in the growth medium causes a significant decrease in the cellular concentration of NSPD in Vibrio species (Yamamoto et al., 1991). However, whether there is an actual decrease in the amount of enzyme protein (i.e., repression of enzyme synthesis) or whether there is an induction of modulator(s) such as antizymes (Panagiotidis & Canelaklis, 1984), which can inactive these enzymes, is unknown.

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References


