PYRIDOXAL PHOSPHATE N-OXIDE. ANALYSES OF COENZYMATIC ACTIVITIES FOR TRYPTOPHANASE AND ASPARTATE AMINOTRANSFERASE BY USE OF STOPPED FLOW METHOD

Fuminori Masugi, Toshihiko Maeda, Yutaka Sumi, Shoichi Shimizu, and Saburo Fukui

Laboratory of Industrial Biochemistry, Department of Industrial Chemistry, Faculty of Engineering, Kyoto University, Kyoto, Japan
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In order to obtain accurate information about the function of the ring-nitrogen of B6 in the enzymatic reactions dependent on the vitamin, the reaction rate leading to α-proton elimination from amino acid substrate was compared between native enzyme and holoenzyme species, reconstituted with pyridoxal phosphate N-oxide (PLP N-oxide), using a stopped flow method. In tryptophanase, the formation rate of the deprotonated Schiff's base (A_502nm complex) of the PLP N-oxide enzyme with L-alanine was about one-fifth of that of the native enzyme, being substantially consistent with the marked low catalytic activity of PL N-oxide in the corresponding nonenzymatic model reaction and of the PLP N-oxide enzyme in overall tryptophan degradation. This result strongly suggests that the electronegativity of the pyridinium-nitrogen plays a predominant role in the α-proton elimination from the substrate, one of the most important steps in overall reaction.

On the other hand, in aspartate aminotransferase (GOT) reaction, such a significant difference was not observed between the formation rate of deprotonated Schiff's base (A_402nm complex) of the native GOT with erythro-β-hydroxy-aspartate and that of the artificial GOT activated with PLP N-oxide. However, the reaction of PMP N-oxide form of GOT with α-keto acid proceeded at an extremely slow rate as compared with that of PMP form of GOT. These phenomena indicate that the lower V_max of overall GOT reaction catalyzed by the PLP N-oxide-bound holoenzyme would be accounted for by the marked smaller reactivity of PMP N-oxide form with α-keto acid. These results strongly suggest that, in GOT, the α-proton elimination from substrate is mainly mediated by a nucleophilic attack of basic amino acid residue located at an appropriate position of enzyme protein and that the introduction of an...
oxygen atom into the pyridine-nitrogen of PLP results in a conformational change of coenzyme analog-bound GOT which renders the interaction between PMP-form GOT with α-keto acid very difficult.

It is well established that vitamin B₆-catalyzed enzymatic reactions proceed via the Schiff’s base (aldimine) formation between pyridoxal phosphate (PLP) and amino acid substrate. Special properties of PLP enable the labilization of the bonds to the α-carbon of the amino acid, leading to a variety of reactions, such as transamination, decarboxylation, racemization and splitting of the Cₓ-Cᵧ bond of amino acid and so on. Model studies with analogs of pyridoxal (PL) permitted delineation of the structural features of the coenzyme required for its catalytic actions: Schiff’s base formation between the formyl group at position 4 and amino acid is greatly favored by the phenol group at position 3, and, in the resulting complex, both the pyridine-nitrogen and the azomethine-nitrogen provide electrophilic groups so placed as to weaken each of the bonds about the α-carbon atom of the amino acid. These concepts, derived largely from studies of nonezymatic model reactions, offered the basis for the general mechanism of reactions catalyzed by PLP (1, 2). In enzymatic reactions, however, it should be taken into consideration that both marked enhancement of reaction rate and strict stereospecificity are brought about by the contribution of apoprotein, as proposed by DUNATHAN (3) and other researchers.

To obtain further information on the role of pyridine-nitrogen of PLP for the exhibition of its coenzymatic activity, N-oxide of PLP (PLP N-oxide) was prepared in our laboratory (4, 5), and its catalytic activity (4, 5) as well as mode of binding to the apoproteins (6) in some enzyme systems were investigated. In general, the incorporation of oxygen atom into the pyridine-nitrogen of PLP decreased the coenzymatic activity in various B₆-enzyme systems. Moreover, a complicated phenomenon was observed in the binding step of PLP N-oxide to the apoprotein of aspartate aminotransferase (GOT) (6).

This paper describes the comparison of the catalytic activities of GOT and tryptophanase activated with PLP and with PLP N-oxide in the reaction rates up to the elimination of α-proton from the amino acids by use of a stopped flow method.

EXPERIMENTAL PROCEDURES

Materials. Cytoplasmic and mitochondrial aspartate aminotransferases (GOTᵦ and GOTₘ) were purified from pig heart muscle by the procedure of WADA and MORINO (7). The apoenzymes were prepared according to the method of TURANO et al. (8) Apotryptophanase was obtained from E. coli B/lt-7A and converted to desired holoenzyme species according to the procedure of NEWTON et al. (9) Pyridoxal phosphate N-oxide (PLP N-oxide) was prepared by our
method (5) with a slight modification in its crystallization step. Erythro-β-hydroxy-DL-aspartate was synthesized by the method of KORNGUTH and SALLACH (10).

Protein concentration was determined by the method of LOWRY et al. (11) with crystalline bovine serum albumin as standard. Absorption spectra were recorded with a Shimadzu Multipurpose Recording Spectrophotometer 50L and their rapid time-course changes were followed using a Hitachi Rapid Scan Spectrophotometer RSP–2 equipped with a stopped flow apparatus.

THEORY

Steady state kinetics of the Schiff’s base formation

The Schiff’s base formation reaction in a nonenzymatic system should be expressed in the following general equation.

\[ P + A \xrightarrow{k_1} S + H_2O \]

where \( P \), PL or PL N-oxide; \( A \), amino acid; \( S \), Schiff’s base.

When amino acid is added to a PL or PL N-oxide solution, the change in absorbance at 420 nm (\( A_{420} \)) is related to the dissociation constant of the Schiff’s base between amino acid and PL or PL N-oxide according to the following theoretical equation. When coenzyme analog has no absorption at 420 nm:

\[ \frac{P_0}{A_{420}} = \frac{1}{\varepsilon} + \frac{K}{\varepsilon \cdot A} \]  

(1)

When coenzyme analog has absorption at 420 nm:

\[ \frac{P_0}{A_{420} - \varepsilon_0 \cdot P_0} = \frac{1}{\varepsilon - \varepsilon_0} + \frac{K}{(\varepsilon - \varepsilon_0) \cdot A} \]  

(2)

where \( P_0 \), the initial concentration of \( P \); \( K \), dissociation constant of the Schiff’s base in aqueous solution (\( K = k_2/k_1 \), \( k_2' = k_2(H_2O) \)); \( \varepsilon \) and \( \varepsilon_0 \), the apparent extinction coefficients of the Schiff’s base and the coenzyme analog at 420 nm. At high concentrations of the amino acid (\( A \)), plots of \( P_0/A_{420} \) or \( P_0/(A_{420} - \varepsilon_0 \cdot P_0) \) against \( 1/A \) are linear and have an abscissa equal to \((-1/K)\).

Kinetics of the Schiff’s base formation in nonenzymatic systems

The relation of the concentration of the Schiff’s base (\( x \)) at time \( t \) to \( t \) can be expressed by the following equation.

\[ k_2(A_0 + K)t = \ln \left( \frac{1}{1 - \frac{A_0 + K}{A_0 P_0}} \right) \]  

(3)

\((A_0 \gg P_0)\)

where \( A_0 \), the initial concentration of \( A \).
Equation 3 is the function of time (t) and the concentration of the Schiff's base (x). The Schiff’s base of PL or PL N-oxide with valine usually absorbs at 420 nm. Hence, the concentration of the Schiff’s base was determined by measuring the absorbance at 420 nm.

Kinetics of the intermediate formation in enzymatic reactions

The reaction scheme of tryptophanase is simply described as follows:

\[ E + S \xrightarrow{k_s} EX \xrightarrow{k_1} E + P \]

where E, S, EX and P represent the holoenzyme, substrate, intermediate and product, respectively.

On addition of substrate, holotryptophanase exhibits a new absorption peak between 492 and 507 nm (the exact position is different according to the kind of substrate used). These peaks disappear as the substrate is decomposed to yield reaction products. L-Alanine also gives rise to an intense absorption band at 502 nm. In contrast to the similar peak formed with the normal substrate of tryptophanase, this peak does not disappear even after long standing. The species absorbing at 502 nm represents a deprotonated enzyme-L-alanine complex (12).

In this case, the reaction scheme can be described as follows:

\[ E + S' \xrightarrow{k_1} EX' \]

where E, S' and EX' represent the holoenzyme, L-alanine and the deprotonated enzyme-L-alanine complex, respectively. When L-alanine (S') is added to an enzyme solution (e0), the change in absorbance at 502 nm is related to the dissociation constant of EX’ (K = k2/k1) as follows:

\[ \frac{e_0}{A_{502}} = \frac{1}{\epsilon} + \frac{K}{\epsilon \cdot S'} \quad (S' \gg e_0) \]  

(4)

where \( \epsilon \) is the apparent extinction coefficient of the complex absorbing at 502 nm.

The relation of the concentration of EX’ (x) at time t to t is expressed by the following equation:

\[ k_1(e_0 + K) t = \ln \left( \frac{1}{S_0 + K} \right) \quad (S' \gg e_0) \]  

(5)

When erythro-ß-hydroxyaspartate is added to PLP-form of GOT, a new absorption peak appears at 492 nm. This species represents a deprotonated enzyme-erythro-ß-hydroxyaspartate complex (12). Equations 4 and 5 are also used for the 492-nm complex formation in GOT.
RESULTS

Steady state of the Schiff’s base formation of pyridoxal N-oxide with valine

The Schiff’s base formation of PL and of PL N-oxide with valine of various concentrations was studied spectrophotometrically at different pHs. Figure 1 shows the results obtained according to Equations 1 or 2. From the data depicted in Fig. 1, the dissociation constants and apparent extinction coefficients of the Schiff’s bases are given in Table 1. In general, the Schiff’s base of PL N-oxide exhibited smaller dissociation constants than that of PL in the pH region tested. The former showed the smallest value at pH 7 (9.52×10⁻³), while the latter was most stable at pH 9 (1.41×10⁻²).

Table 1. Dissociation constants and extinction coefficients of the Schiff’s base formed between valine and PL or PL N-oxide at various pHs.

|          | pH 6   | pH 7   | pH 8   | pH 9   | λ_max |
|----------|--------|--------|--------|--------|-------|
| PL       | K_diss | 2.00   | 2.22×10⁻¹| 3.58×10⁻²| 1.73×10⁻²| 415   |
|          | ε×10⁻³ | 6.66   | 6.66   | 6.66   | 6.66   |
| PL N-oxide| K_diss | 4.76×10⁻²| 9.52×10⁻⁴| 1.22×10⁻²| 1.41×10⁻²| 409   |
|          | ε×10⁻³ | 3.27   | 5.86   | 5.67   | 3.71   |

Fig. 1. Schiff’s base formation between PL or PL N-oxide and DL-valine under various pH conditions. PL (1.5×10⁻⁴M) (left) and PL N-oxide (1.5×10⁻⁴M) (right) were incubated with various concentrations of DL-valine at different pHs, respectively. The absorbance at 420 nm of the Schiff’s bases formed was plotted against the concentrations of DL-valine according to Eq. 1. The buffer solution used was 0.05 M K-phosphate.
The rate constant of the Schiff's base formation

In order to determine the rate constant for the Schiff's base formation, log $\frac{1}{[1 - (A + K) \cdot x/A \cdot P]}$ was plotted against time ($t$) according to Eq. 3. The results obtained are illustrated in Fig. 2. The forward and reverse rate constants ($k_1$, $k_2$) of the Schiff's base formation were calculated from Fig. 2 by using the dissociation constants in Table 1. The forward reaction constant ($k_1$) is higher at pH 8 than at pH 7 in the cases of both PL and PL N-oxide. At both pH 7 and 8, the $k_1$ values for PL N-oxide were markedly large as compared with those for PL, whereas the $k_2$ values for PL N-oxide-valine aldimine were smaller than those for PL-valine aldimine (Table 2). Accordingly, it is concluded that PL N-oxide can form the Schiff's base with an appropriate amino acid such as valine more
Easily than PL and the resulting complex of PL N-oxide is less dissociable than that of PL at the neutral and weak alkaline pH regions where the GOT and tryptophanase reactions are carried out.

The formation of $A_{502}$ complex in PLP N-oxide-bound tryptophanase

The appearance of the absorption spectra of the deprotonated alanine-tryptophanase complex was studied using the native holoenzyme and the holoenzyme activated with PLP N-oxide with different concentrations of L-alanine (Fig. 3). According to Eq. 4, the apparent dissociation constants were determined by plotting $1/A_{502}$ against $1/(\text{L-alanine})$ (Fig. 4, Table 3). The apparent dissociation constant of PLP N-oxide enzyme-L-alanine complex was about five times

Fig. 3. Absorption spectra of tryptophanase-L-alanine complexes. Upper; holotryptophanase (1.42 mg/ml) was incubated with various concentrations of L-alanine at pH 8.0 (0.1 M phosphate buffer). Lower; holotryptophanase reconstituted with PLP N-oxide (11 mg/ml) was incubated with L-alanine in the same way.
Fig. 4. Time course of $A_{502}$ complex formation. Upper; holotryptophanase (2.96 mg/ml) and L-alanine (0.1 M) were mixed instantaneously by using a stopped flow apparatus equipped with flow generator. The change in absorbance at 502 nm with time was recorded by using a memoryscope. Lower; holotryptophanase reconstituted with PLP N-oxide (6.7 mg/ml) and L-alanine (0.5 M) were treated in the same way.

Table 3. Dissociation constant and rate constants of ECS' complex between tryptophanase and L-alanine.

$$EC + S' \quad \frac{k_1}{k_{-1}} \quad ECS'$$

$$K_{diss.} = \frac{k_{-1}}{k_1}$$

|                  | $K_{diss.}$ | $k_1 (1/\text{m}\cdot\text{sec})$ | $k_{-1} (1/\text{sec})$ |
|------------------|------------|----------------------------------|------------------------|
| PLP N-oxide enzyme | 0.22       | 1.85                             | 0.407                  |
| PLP enzyme       | 0.042      | 9.53                             | 0.396                  |

as large as that of PLP enzyme-L-alanine complex. In order to determine the rate constant of 502 nm complex formation, the absorption at 502 nm at time $(t)$ was followed by using a stopped flow apparatus (Fig. 4). According to Eq. 5, the rate constant of 502 nm complex formation was calculated from the above-mentioned apparent dissociation constant (Table 3). The forward rate constant for the formation of PLP N-oxide enzyme-L-alanine complex was about one-fifth that of PLP enzyme-L-alanine complex. On the other hand, the reverse rate constant was almost equal in both PLP enzyme and PLP N-oxide enzyme.

**The formation of $A_{492}$ complex in PLP N-oxide-bound GOT**

Figure 5 shows the absorption spectra of $A_{492}$ complexes of PLP-bound GOT.
Fig. 5. Absorption spectra of GOT-erythro-β-hydroxyaspartate complexes. Upper; holo-GOT (5 mg/ml) was incubated with various concentrations of erythro-β-hydroxyaspartate at pH 7.5 (0.05 M phosphate buffer). Lower; holoGOT reconstituted with PLP N-oxide (5 mg/ml) was incubated with erythro-β-hydroxyaspartate in the same way.

(native GOT) and of PLP N-oxide-bound GOT using various concentrations of erythro-β-hydroxyaspartate. These spectra were obtained after standing the mixture of the enzyme and the substrate analog for 3 min. According to Eq. 4, the reciprocal of absorption at 492 nm was plotted against the reciprocal of erythro-β-hydroxyaspartate concentration (Fig. 6). From Fig. 6, the apparent dissociation constants of PLP-bound GOT-pseudosubstrate complex and of PLP N-oxide-
Fig. 6. Equilibrium of holoGOT-erythro-β-hydroxyaspartate complex formation at pH 7.5. The absorbance at 492 nm was plotted against the concentration of erythro-β-hydroxy according to Eq. 4. Closed circle, PLP N-oxide-enzyme; open circle, PLP-enzyme.

|          | PLP  | PLP N-oxide |
|----------|------|-------------|
| $K_{\text{eq}} \times 10^4$ | 2.50 | 1.22        |
| $K_{\text{eq}} \times 10^{-4}$ | 1.93 | 0.50        |

bound GOT-pseudosubstrate complex were estimated graphically as $2.50 \times 10^{-4}$ M and $1.93 \times 10^{-4}$ M, respectively. Next, the kinetics of the $A_{492}$ complex formation was studied. The time-course change of the absorbance at 492 nm was recorded using the stopped flow apparatus equipped with a memoryscope immediately after rapid mixing of PLP enzyme or PLP N-oxide enzyme with the substrate analog. As shown in Fig. 7, there was no marked difference between the PLP enzyme reaction and PLP N-oxide enzyme reaction. In the case of PLP enzyme, $A_{492}$ complex was formed instantaneously after mixing the enzyme and the pseudosubstrate. After several milliseconds, the absorbance increased further, suggesting the formation of an unidentified state having a higher extinction coefficient at 492 nm. After 60 milliseconds, the absorbance at 492 nm gradually decreased and reached a constant level within 2.5 seconds. Thereafter, no change in the absorbance was observed at least during the subsequent 20 min.

In the case of PLP N-oxide-bound GOT, the time-course change in the absorbance at 492 nm exhibited a substantially similar pattern, except that the state of having a high extinction coefficient at 492 nm observed in the case of PLP enzyme was not detected. The time-course study indicated that at least three steps would be involved in the reaction sequence leading to the deprotonated enzyme-substrate analog complex in both the native enzyme and the coenzyme analog-bound holoenzyme reactions, and that there exists no significant difference in the reaction rate up to the formation of the deprotonated complex between two enzymes.
Fig. 7. Time-course of GOT-erythro-β-hydroxyaspartate complex formation. Upper; holoGOT (5 mg/ml) and erythro-β-hydroxyaspartate (1 mM) were mixed instantaneously by using a stopped flow apparatus equipped with flow generator. The absorbance at 492 nm was recorded by using a memoryscope. Lower; holoGOT reconstituted with PLP N-oxide (5 mg/ml) and erythro-β-hydroxyaspartate (2 mM) were treated in the same manner.

As reported previously (4, 5), the activity of GOT reconstituted with PLP N-oxide was below one-half that of the native GOT in the overall reaction rate. This discrepancy could be accounted for by a significantly low reactivity of the GOT reconstituted with pyridoxamine phosphate N-oxide (PMP N-oxide) with α-ketoglutarate, the reverse reaction from PMP-form GOT to PLP-form GOT. As shown in Fig. 8, the addition of α-ketoglutarate to the PMP N-oxide-bound GOT resulted in a very slow change in the spectrum, indicating that the reaction
Fig. 8. Effect of α-ketoglutarate on the PMP N-oxide form of GOT. PMP N-oxide form of GOT (2.5 × 10⁻⁴ M) was incubated with α-ketoglutarate (10 mM) in 0.1 M phosphate buffer (pH 7.0) at 37°C. Curves 1, 2, and 3 are the spectra measured at 5 min, 1.5 hr and 6 hr after mixing.

rate is markedly low as compared with that of the native PMP-form enzyme. Namely, it would be concluded that this step is rate-limiting in the overall reaction mediated by the coenzyme analog-bound GOT.

DISCUSSION

In model experiments of B₄ enzymecatalyzed reactions, it is clearly established that the pyridine-nitrogen, together with 4-formyl group and 3-phenolic hydroxyl group, is essential for the catalytic activity of PL(P). Initial activation of an

![Figure 9: Two proposed mechanisms for the α-proton elimination from amino acid substrate.](image-url)
amino acid occurs through the formation of an aldimine (Schiff's base) between the amino acid and PL(P). The conjugated system of double bonds extending from the \(\alpha\)-carbon atom of the amino acid to the pyridinium-nitrogen of PL(P) results in a reduced electron density about the \(\alpha\)-carbon atom, thus weakening each of the bonds (Fig. 9a). One might assume that the same structural features that permit PL(P) to catalyze a given model reaction would be prerequisite to catalysis of the corresponding enzymatic reaction. In the case of enzymatic reaction, however, contribution of enzyme protein, especially that of a nucleophilic attack of the basic amino acid located at an appropriate position of enzyme, seems to take an important role for the labilization of proton attached to the \(\alpha\)-carbon atom of the amino acid substrate (Fig. 9b).

As reported in our previous paper (4), PL N-oxide showed markedly lower catalytic activities in various nonenzymatic model systems such as \(\alpha,\beta\)-elimination reactions of tryptophan and serine and transamination reaction between glutamate. The lower activity of PL N-oxide was attributed to the decreased electronegativity of pyridine-nitrogen, as discussed in our previous paper (4).

| Table 4. Coenzymatic parameters of PLP N-oxide for several PLP-dependent enzymes\(^a\) |
|---------------------------------|-----------------|-----------------|
|                                | \(K_{\text{on}}\) (\(\mu\)M) | \(V_{\text{max}}\) (relative) |
|                                | PLP enzyme | PLP N-oxide enzyme | PLP N-oxide enzyme |
| GOT\(_m\) (pig heart)\(^{1\dagger}\) | 1.19 | 1.19 | 0.45 |
| GOT\(_s\) (pig heart)\(^{1\dagger}\) | 1.14 | 1.14 | 0.43\(^b\) |
| \(d\)-Serine dehydratase (E. coli)\(^{2\dagger}\) | ++ + | ++ | 0.9 |
| Tryptophanase (E. coli)\(^{1\dagger}\) | 4.5 | 5.0 | 0.65 |
| Tyrosine phenol-lyase (E. intermedia)\(^{3\dagger}\) | 2.2 | 6.3 | 0.82 |
| Aspartate \(\beta\)-decarboxylase (P. dacunhae)\(^{4\dagger}\) | Decarboxylation | 0.16 |
|                                | Desulfination | 0.26 |
|                                | Transamination | 0.31 |

\(^a\) In the case of muscle phosphorylase, PLP N-oxide was converted into PLP (Pfeuffer, T., Ehrlich, J., and Helmreich, E., Biochemistry, 11, 2125 (1972)).

\(^b\) In the case of GOT\(_s\), the incorporation of one more mole of PLP N-oxide resulted in enhancement of the activity of the PLP N-oxide enzyme. The activated enzyme showed an almost similar \(V_{\text{max}}\) to that of the native enzyme (5).

In various enzyme system, PLP N-oxide served as inferior coenzymes as listed in Table 4. For example, the activity of GOT\(_m\) and GOT\(_s\) reconstituted with PLP N-oxide was below one-half that of the PLP-enzymes, although binding of one mole of PLP N-oxide to a certain lysine residue at non-catalytic site rendered
the activity of PLP N-oxide-bound GOT, almost equal to that of the native enzyme (6).

In this paper, a stopped flow method was employed to compare PLP-enzyme and PLP N-oxide-enzyme in the reaction rate to form the deprotonated aldimine. In the case of tryptophan degradation, the relative activity of PLP N-oxide enzyme in overall enzymatic reaction was ca. 65%, which was consistent with the relative activity of PL N-oxide to that of PL in the corresponding nonenzymatic reaction. The stopped flow study employed in this paper showed that the rate constant of α-hydrogen elimination from L-alanine in PLP N-oxide-bound tryptophanase was remarkably small as compared with that in PLP-bound tryptophanase. These results strongly suggest that the participation of the electronegativity of pyridine-nitrogen is most important for the α-proton elimination from substrate in this system. In the case of GOT reaction, however, such a significant difference was not observed between the apparent rate of α-proton elimination in PLP-enzyme and in PLP N-oxide-enzyme, in contrast to the marked differences in both nonenzymatic and overall enzymatic reactions. This fact suggests that the electron-attracting force of the pyridinium-nitrogen of PLP does not play the important part in the α-proton elimination step of GOT as expected hitherto. The low reaction rate of the overall GOT reaction mediated by the PLP N-oxide enzyme could be explained by the fact that the formation of PLP N-oxide-GOT was very difficult when PMP N-oxide-GOT was incubated with α-ketoglutarate. As mentioned in our preceding paper (6), the $K_{\text{coenzyme}}$ value of PMP N-oxide for GOT was remarkably higher than that of PMP and the resulting PMP N-oxide-bound holoGOT showed an extremely low $V_{\text{max}}$ value as compared with the native enzyme. From these phenomena, it would be concluded that, in GOT, the function of the pyridine-nitrogen of PLP in the α-proton elimination of amino acid substrate is significantly weak compared with that of the nucleophilic attack of the basic amino acid residue located at an appropriate position of the enzyme protein. The introduction of oxygen atom into the pyridine-nitrogen not only has some influence on the binding of the coenzyme analog with apoGOT but also leads to variations in the conformation of the resulting holoenzyme species. This conformational distortion would render the interaction between the amino group of PMP N-oxide form and the keto acid substrate more difficult than the case of the PMP enzyme, thus resulting in the decrease of the overall reaction rate.

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