Determination of Kinetic Parameters of Enzyme-Catalyzed Reaction $A + B + C \rightarrow$ Products with the Minimum Number of Velocity Measurements

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Rapid-equilibrium rate equations are derived for the five different mechanisms for the enzymatic catalysis of $A + B + C \rightarrow$ products using a computer. These rate equations are used to determine the minimum number of velocities required to estimate the values of the kinetic parameters. The rate equation for the completely ordered mechanism involves four kinetic parameters, and the rate equation for the completely random mechanism involves eight kinetic parameters. Therefore, the four to eight kinetic parameters can be estimated by determining four to eight velocities and solving four to eight simultaneous equations. General recommendations are made as to the choices of triplets of substrate concentrations $\{[A], [B], [C]\}$ to be used to determine the velocities. The effects of 5% errors in the measured velocities, one at a time, are calculated and are summarized in tables. Calculations of effects of experimental errors are useful in choosing the triplets of substrate concentrations to be used to obtain the most accurate values of the kinetic parameters. When the kinetic parameters for $A + B + C \rightarrow$ products are to be determined for the first time, it is recommended that the program for the completely random mechanism be used because it can identify the mechanism and determine the kinetic parameters in one operation.

### Introduction

This article is concerned with the possible rapid-equilibrium rate equations for enzyme-catalyzed reactions of the type $A + B + C \rightarrow$ products. Many reactions of this type are found among the oxidoreductases, and most ligase reactions are of this type.1

Four examples of these reactions are shown in Table 1. The fact that the change in binding of hydrogen ions $\Delta N_{H}$ of the forward reaction is positive suggests that one or more hydrogen ions are bound in the rate-determining reaction.2 The effects of $\text{pH}$ in rapid-equilibrium enzyme kinetics have been discussed recently,3 and the relation between $\Delta N_{H}$ and the integer number $n$ of hydrogen ions consumed in the rate-determining reaction has been derived for $A + B \rightarrow$ products.

There are five possible rapid-equilibrium rate equations for the enzyme-catalyzed reaction $A + B + C \rightarrow$ products, with mechanisms from the completely ordered mechanism to the completely random mechanism. The completely ordered mechanism involves four kinetic parameters, and the completely random mechanism involves eight kinetic parameters. Rapid-equilibrium rate equations are derived for each of these five mechanisms using a computer. These rate equations can be used to determine the kinetic parameters with the minimum number of measured velocities.4 The concept that $N$ kinetic parameters can be calculated from $N$ velocity measurements was introduced by Duggleby5 in 1975. Before that, Frieden,6 in an article about malate dehydrogenase, showed that it was possible to determine the four kinetic parameters ($V_{AB}$, $K_{A}$, $K_{B}$, and $K_{AB}$) in the rate equation $\nu = V_{AB}(1 + K_{A}[A] + K_{B}[B] + K_{AB}[A][B])$ with four velocity measurements. This article is concerned with rapid-equilibrium rate equations because they are the simplest possible rate equations. Steady-state rate equations have more terms, but the concept of determining the kinetic parameters with the minimum number of velocity measurements also applies.

Rapid-equilibrium rate equations can be derived with a computer because Mathematica, Maple, and MatLab have the operation Solve that solves sets of simultaneous polynomial equations. General recommendations are made here as to the choice of triplets of substrate concentrations $\{[A],[B],[C]\}$ to be used to determine velocities. The objective is to use triplets of substrate concentrations that reduce the effects of experimental errors on the estimated kinetic parameters.

### Derivations of Rapid-Equilibrium Rate Equations for $A + B + C \rightarrow$ Products

When the rapid-equilibrium assumption is used, all of the reactions up to the rate-determining reaction are in equilibrium. When a system of reactions is at equilibrium, not only are the reactions used in the calculation at equilibrium, but all reactions that can be obtained by adding and subtracting these reactions are also at equilibrium. This makes it possible to calculate dissociation constants (Michaelis constants) for this larger set

### Table 1: Changes in Binding of Hydrogen Ions $\Delta N_{H}$ in Enzyme-Catalyzed Reactions as a Function of pH at 298.15 K

| Reaction                             | pH 5 | pH 6 | pH 7 | pH 8 | pH 9 |
|--------------------------------------|------|------|------|------|------|
| EC 1.4.1.1 pyruvate + NAD$_{\text{red}}$ + ammonia = alanine + NAD$_{\text{ox}}$ + H$_{2}$O | 1.00 | 1.00 | 1.01 | 1.05 | 1.36 |
| EC 1.4.1.2 + NAD$_{\text{red}}$ + ammonia = glutamate + NAD$_{\text{ox}}$ + H$_{2}$O | 1.00 | 1.00 | 1.01 | 1.05 | 1.36 |
| EC 6.3.5.4 aspartate + ATP + glutamine = asparagine + glutamate + AMP + PP$_{i}$ | 1.83 | 1.27 | 0.92 | 0.67 | 0.17 |
| EC 6.4.1.5 succinate + ATP + CoA = succinylCoA + ATP + P | 0.44 | 0.66 | 0.32 | 0.48 | 0.89 |

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of reactions. For example, the determination of 8 velocities for the completely random mechanism makes it possible to calculate 12 dissociation constants (Michaelis constants). The number \( n' \) of reactants in the completely random mechanism is 11; these reactants are E, A, B, C, EA, EB, EC, EAB, EBC, and EABC. The total number of dissociation reactions is 12

\[
\begin{align*}
\text{E + A = EA} & \quad K_A = [E][A]/[EA] = 5 \\
\text{E + B = EB} & \quad K_B = [E][B]/[EB] = 10 \\
\text{E + C = EC} & \quad K_C = [E][C]/[EC] = 15 \\
\text{EA + B = EAB} & \quad K_{AB} = [EA][B]/[EAB] = 20 \\
\text{EA + C = EAC} & \quad K_{AC} = [EA][C]/[EAC] = 25 \\
\text{EB + A = EAB} & \quad K_{BA} = [EB][A]/[EAB] = 30 \\
\text{EB + C = EBC} & \quad K_{BC} = [EB][C]/[EBC] = 35 \\
\text{EC + A = EAC} & \quad K_{CA} = [EC][A]/[EAC] = 40 \\
\text{EC + B = EBC} & \quad K_{CB} = [EC][C]/[EBC] = 45 \\
\text{EAB + C = EABC} & \quad K_{ABC} = [EAB][C]/[EABC] = 50 \\
\text{EAC + B = EABC} & \quad K_{ACB} = [EAC][B]/[EABC] = 60 \\
\text{EBC + A = EABC} & \quad K_{BCA} = [EBC][A]/[EABC] = 70
\end{align*}
\]

These reactions are not all independent. The number of independent reactions \( R' \) can be calculated\(^2\) by using \( n' = C' + R' \), where the number of components \( C' \) is 4 (E, A, B, and C), and there are 11 reactants in eqs 1–12. The number of independent reactions is given by \( R' = n' - C' = 11 - 4 = 7 \); thus, only 7 of the dissociations in eqs 1–12 are independent. Seven is also the number of dissociation reactions in the completely random mechanism. These seven reactions are the ones for which numerical values have been arbitrarily assigned in eqs 1–5, 7, and 10 so that rapid-equilibrium velocities can be calculated. Other choices of independent variables can be made. The equilibrium constants for these 7 reactions can be used to calculate the equilibrium constants for the other 5 dissociation reactions, but we will take a different path here and discuss the estimation of all 12 equilibrium constants by determining the minimum number of velocities for the 5 mechanisms for \( A + B + C \rightarrow \text{products} \).

There are two ways that the pH can affect the rate of an enzyme-catalyzed reaction calculated using the rapid-equilibrium assumption. The more familiar way is through the pKs of the substrates, enzymatic site, and enzyme—substrate complexes. The less familiar way is through the consumption of hydrogen ions in the rate-determining reaction,\(^7,^8\) which introduces a factor of \( 10^{\text{pH}} \) into the expression for the velocity. When one hydrogen ion is consumed, \( n = -1 \) because of the convention in thermodynamics that the stoichiometric numbers of reactants are negative. The fact that there are oxidoreductase reactions for which the limiting velocity \( V_{\text{exp}} \) is proportional to \( 10^{\text{pH}} \) over a wide range of pHs is an argument for the rapid-equilibrium assumption. Only the consumption of hydrogen ions in the rate-determining reaction can give rise to the \( 10^{\text{pH}} \) factor over a range of pHs. For the calculations in this paper, \( V_{\text{exp}} \) is taken to be 1, and the effects of pH are not investigated further.

### Determination of the Kinetic Parameters \( V_{\text{exp}}, K_A, K_{AB}, \) and \( K_{ABC} \) for Mechanism I

Completely ordered mechanism I is made up of the reactions in eqs 1, 4, 10, and

\[
\begin{align*}
\text{EABC} \rightarrow \text{products} & \quad v = k_d[EABC] = \\
& \quad k_{f}[E][EABC]/[E] = V_{\text{exp}}[EABC]/[E] (13)
\end{align*}
\]

where the limiting velocity \( V_{\text{exp}} \) is equal to \( k_d[E] \), where \( k_d \) is the rate constant for the production of products and \([E]_t\) is the total concentration of enzymatic sites.

It is important to check the number of independent equilibrium expressions used in the derivation of the reaction velocity \( v \). The number \( n' \) of reactants in mechanism I is seven (E, A, B, C, EAB, and EABC). The number of components \( C' \) is four (E, A, B, and C). The number of independent equilibrium equations \( R' \) is three, and therefore, \( n' = C' + R' = 7 + 4 = 11 \). To obtain the expression for the equilibrium concentration of EABC, three equilibrium equations and one conservation equation are used in Solve. This operation is shown in the Appendix. The rapid-equilibrium velocity of the reaction \( A + B + C \rightarrow \text{products} \) is given by \( v = k_d[EABC] \). Therefore, the \([E]_t\) in the expression for \([EABC] \) obtained using Solve is replaced by \( V_{\text{exp}} \), and the symbol for the limiting velocity in the forward direction is \( V_{\text{exp}} = k_d[E] \). In Mathematica, \( V_{\text{exp}} \) is represented by \( v_{\text{exp}} \) because capital letters are used for operations. The rapid-equilibrium velocity for mechanism I obtained in this way is

\[
\begin{align*}
&\quad v = \\
&\quad [A][B][C]V_{\text{exp}} \\
&\quad [A][B][C] + [A][B]K_{ABC} + [A]K_{AB}K_{ABC} + K_AK_{AB}K_{ABC} (14)
\end{align*}
\]

An advantage in deriving the rapid-equilibrium rate equation with a computer is that the equation is in computer-readable form. If \( v \) is measured at four triplets of substrate concentrations \([A],[B],[C]\), the four versions of rate equation 14 can be solved for the four kinetic parameters. Since there are experimental errors in the measured velocities, the triplets of substrate concentrations used should be chosen so as to reduce the effects of these experimental errors on the calculated kinetic parameters. There are also experimental errors in the concentrations of the substrates, but these errors are independent of the choices of substrate concentrations. The effects of 5% errors in substrate concentrations are 5% errors in Michaelis constants. There are other possible errors in velocity measurements that have to be discussed for specific enzyme kinetic studies.

The effects of choices of substrate concentrations have been discussed for \( A + B \rightarrow \text{products} \).\(^4\) Therefore, the calculations here on mechanism I have been made with the following triplets of substrate concentrations \([A],[B],[C]\): \(\{200,200,200\}, \{0.5,200,200\}, \{200,0.5,200\}, \text{ and } \{200,200,0.5\}\). The concentration 200 is hopefully significantly higher than the largest
Michaelis constant, and the 0.5 is hopefully significantly lower than the smallest Michaelis constant. The velocity for the first triplet has the predominate effect in determining $V_{\text{fexp}}$. The velocity for the second triplet has the predominate effect in determining $K_A$. The velocity for the third triplet has the predominate effect in determining $K_{AB}$. The velocity for the fourth triplet has the predominate effect in determining $K_{ABC}$. The velocities for the four triplets of substrate concentrations calculated using eq 14 with the specified values of $V_{\text{fexp}}$ and the Michaelis constants (see eqs 1, 4, and 10) are $v_1 = 0.8383$, $v_2 = 0.7313$, $v_3 = 0.1198$, and $v_4 = 0.01279$. Since $v_4$ is very low, it may be necessary to raise $[E]_t$.

The Mathematica program calckinparsABC1 for calculating the four kinetic parameters in mechanism I from four measured velocities is given in the Appendix. This program is based on the Solve operation. The effects of 5% errors in the four measured velocities, one at a time, calculated using this program, are given in Table 2.

Increasing the high substrate concentrations and reducing the low substrate concentrations at which velocities are calculated will reduce these errors somewhat. Calculations with 5% errors are arbitrary, and the errors can be increased without changing the general conclusions.

**Determination of the Kinetic Parameters** $V_{\text{fexp}}, K_A, K_B, K_{AB},$ and $K_{ABC}$ for Mechanism II

For mechanism II, reaction 2 is added to the reactions in mechanism I. The number $N'$ of reactants is eight. The number of components $C'$ is four (E, A, B, and C). The number of independent equilibrium equations $R'$ is four, and therefore, $N' = C' + R' = 8 = 4 + 4$. To calculate the equilibrium concentration of EABC using Solve, four equilibrium equations and one conservation equation are used. This yields the following expression for the rapid-equilibrium velocity.

$$V = \frac{[A][B][C]V_{\text{fexp}}K_B}{[B]K_AK_{AB}K_{ABC} + [A][B][C]K_B + [A][B]K_{ABC}K_B + [A]K_{AB}K_{ABC}K_B + K_AK_{AB}K_{ABC}K_B}$$

When $v$ is measured at five triplets of substrate concentrations $[[A],[B],[C]]$, the five versions of rate eq 15 can be solved for the five kinetic parameters. The calculations here for mechanism II have been made with the following triplets of concentrations $[[A],[B],[C]]$: \{[200,200,200],[0.5,200,200],[200,0.5,200],[200,200,0.5], and [0.5,0.5,200] \}. The velocities for the five triplets calculated using eq 15 and the Michaelis constants involved are $v_1 = 0.8322$, $v_2 = 0.2054$, $v_3 = 0.1196$, $v_4 = 0.01224$, and $v_5 = 0.01224$. Since $v_4$ and $v_5$ are very low, it may be necessary to raise $[E]_t$, for these two measurements.

The program calckinparsABC2 (not in the Appendix) was written to calculate the five kinetic parameters from five measured velocities. This program has the same structure as calckinparsABC1 given in the Appendix, but there are five lines, which are different versions of eq 15. The effects of 5% errors in velocities, one at a time, were calculated using this program, and they are summarized in Table 3.

Three of the reactions in mechanism II almost form a cycle; they are reactions 1, 2, and 4.

$$K_A$$

$E + A = EA$

$$+ + +$$

$B B$

$$||K_B ||K_{AB}$$

$EB + A = EAB$

$$K_{BA}$$

This cycle is completed with $EB + A = EAB$. Therefore, the value of $K_{BA}$ can be calculated from values of $K_A, K_B,$ and $K_{AB}$ determined using the five measured velocities.

$$K_{BA} = K_AK_{AB}/K_B$$

Equations of this type are well-known in enzyme kinetics. Thus, when the enzyme-catalyzed reaction follows mechanism II and there are no experimental errors, $K_{BA} = 5 \times 20/10 = 10$. The values of $K_{BA}$ when there are 5% errors in the velocities, one at a time, are given later in Table 7. This is the first illustration here of the fact that when a system of reactions is at equilibrium, not only the reactions used to calculate the equilibrium composition are at equilibrium, but also any reaction that can be obtained by adding and subtracting the reactions used to calculate the equilibrium concentrations is also at equilibrium.

**Determination of the Kinetic Parameters** $V_{\text{fexp}}, K_A, K_B, K_C, K_{ABC},$ and $K_{AB}$ for Mechanism III

For mechanism III, reaction 3 is added to the reactions in mechanism II. The number $N'$ of reactants is nine. The number of components $C'$ is four (E, A, B, and C). The number of independent equilibrium equations $R'$ is five, and therefore, $N' = C' + R' = 9 = 4 + 5$. To calculate the equilibrium concentration of EABC using Solve, five equilibrium equations and one conservation equation are used. This yields the following expression for the rapid-equilibrium velocity
\[ v = \frac{[A][B][C]V_{\text{exp}}K_BK_C}{D_1 + D_2} \]

\[ D_1 = [C]K_AK_{AB}K_{ABC}K_B + [B]K_AK_{AB}K_{ABC}K_C + [A][B]K_ABK_{ABC}K_C \]

\[ D_2 = [A][B]K_ABK_{ABC}K_C + [A]K_{AB}K_{ABC}K_BK_C + K_AK_{ABC}K_{ABC}K_BK_C \]

When \( v \) is measured at six triplets of substrate concentrations \([A],[B],[C]\), the seven versions of the rate equation can be solved for the seven kinetic parameters. The calculations for mechanism III have been made with the following triplets of concentrations: \(200,200,200\), \(0.5,200,200\), \(200,0.5,200\), \(200,200,0.5\), \(5,5,5\), and \(5,5,5\). The reason for the particular sixth triplet is that the velocity is most sensitive to the change in concentration of a reactant is low (in comparison with its Michaelis constant) when the concentration of a reactant is low (in comparison with its Michaelis constant) that the velocity is most sensitive to the change in concentration of that reactant. However, \(0.5,0.5,0.5\) yields an extremely low velocity. Therefore, all three concentrations are increased, and \(5,5,5\) is used for the sixth triplet. The velocities for the six triplets calculated using eq 18 and the specified values of \(v_1 = 0.8281\), \(v_2 = 0.1389\), \(v_3 = 0.9353\), \(v_4 = 0.01224\), \(v_5 = 0.05545\), and \(v_6 = 0.01145\). Mathematica program calckinparsABC3 was written to calculate the six kinetic parameters from six measured velocities. This program has the same structure as calckinparsABC1 given in the Appendix, but there are seven lines, which are different versions of eq 19. The effects of 5\% errors in velocities, one at a time, were calculated using this program, and they are summarized in Table 4.

| No errors | 1.00 | 5.0 | 20.0 | 10.0 | 15.0 | 35.0 |
|-----------|------|-----|------|------|------|------|
| 1.05 × \(v_1\) | 1.06 | 4.9 | 20.1 | 9.7  | 14.7 | 37.1 |
| 1.05 × \(v_2\) | 1.00 | 5.3 | 19.9 | 11.7 | 15.8 | 35.2 |
| 1.05 × \(v_3\) | 1.00 | 5.8 | 18.4 | 10.8 | 16.1 | 35.3 |
| 1.05 × \(v_4\) | 0.99 | 5.1 | 21.2 | 10.1 | 15.2 | 32.7 |
| 1.05 × \(v_5\) | 1.00 | 4.7 | 20.8 | 9.0  | 16.1 | 34.7 |
| 1.05 × \(v_6\) | 1.00 | 4.4 | 19.7 | 8.8  | 12.3 | 35.1 |

Thus, when the enzyme-catalyzed reaction follows mechanism IV and there are no experimental errors, \(K_{CA} = 5 \times 25/15 = 8.333\). The values of \(K_{CA}\) when there are 5\% errors in the velocities, one at a time, are given later in Table 7.

Another three reactions in this mechanism almost form a cycle; they are reactions 1, 3, and 5.

When \( v \) is measured at seven triplets of substrate concentrations \([A],[B],[C]\), the seven versions of the rate equation can be solved for the seven kinetic parameters. The calculations for mechanism IV have been made with the following triplets of concentrations: \(200,200,200\), \(0.5,200,200\), \(200,0.5,200\), \(200,200,0.5\), \(5,5,5\), and \(5,5,5\). The velocities calculated with these seven triplets are \(v_1 = 0.7421\), \(v_2 = 1.362\), \(v_3 = 0.1500\), \(v_4 = 0.01222\), \(v_5 = 0.04231\), \(v_6 = 0.01076\), and \(v_7 = 0.02332\). Mathematica program calckinparsABC4 was written to calculate the seven kinetic parameters from seven measured velocities. This program has the same structure as calckinparsABC1 given in the Appendix, but there are seven lines, which are different versions of eq 19. The effects of 5\% errors in velocities, one at a time, were calculated using this program, and they are summarized in Table 5.

Three of the reactions in this mechanism almost form a cycle; they are reactions 1, 3, and 5.

\[
K_A
\]
\[
E + A = EA
\]
\[
+ +
\]
\[
C \quad C
\]
\[
||K_C \quad ||K_{AC}
\]
\[
EC + A \rightarrow EAC
\]

\[
K_{CA}
\]

Therefore, values of \(K_{CA}\) can be calculated from experimental values of \(K_A\), \(K_{AC}\), and \(K_C\).

\[
K_{CA} = K_AK_{AC}/K_C
\]

Thus, when the enzyme-catalyzed reaction follows mechanism IV and there are no experimental errors, \(K_{AC} = 20 \times 35/25 = 28\). The values of \(K_{AC}\) when there are 5\% errors in the velocities, one at a time, are given later in Table 7.

\[
K_{AB}
\]
\[
EA + B = EAB
\]
\[
+ +
\]
\[
C \quad C
\]
\[
||K_{AC} \quad ||K_{ABC}
\]
\[
EAC + B \rightarrow EABC
\]

\[
K_{ABC}
\]

Therefore, values of \(K_{ABC}\) can be calculated from experimental values of \(K_{AB}\), \(K_{AC}\), and \(K_{ABC}\).

\[
K_{ABC} = K_ABK_{AC}/K_{ABC}
\]
TABLE 5: Effects of 5% Errors in Seven Measured Velocities for Mechanism IV

| \( V_{\text{exp}} \) | \( K_A \) | \( K_{AB} \) | \( K_B \) | \( K_C \) | \( K_{ABC} \) | \( K_{AC} \) |
|----------------|--------|--------|--------|--------|--------|--------|
| No errors     | 1.00   | 5.0    | 20.0   | 10.0   | 15.0   | 35.0   | 25.0   |
| 1.05 × \( v_1 \) | 1.07   | 4.9    | 20.0   | 9.8    | 14.8   | 37.4   | 25.0   |
| 1.05 × \( v_2 \) | 1.00   | 5.3    | 19.9   | 11.7   | 15.8   | 35.2   | 25.0   |
| 1.05 × \( v_3 \) | 0.99   | 4.9    | 20.2   | 10.2   | 14.5   | 34.7   | 26.9   |
| 1.05 × \( v_4 \) | 0.99   | 4.9    | 21.5   | 10.0   | 15.0   | 32.7   | 25.4   |
| 1.05 × \( v_5 \) | 1.00   | 5.0    | 20.1   | 9.2    | 17.4   | 34.8   | 24.9   |
| 1.05 × \( v_6 \) | 1.00   | 4.2    | 20.1   | 8.6    | 11.9   | 35.1   | 25.2   |
| 1.05 × \( v_7 \) | 1.00   | 5.8    | 18.3   | 10.8   | 16.2   | 35.3   | 22.8   |

TABLE 6: Effects of 5% Errors in Eight Measured Velocities for Mechanism V

| \( V_{\text{exp}} \) | \( K_A \) | \( K_{AB} \) | \( K_B \) | \( K_C \) | \( K_{ABC} \) | \( K_{AC} \) |
|----------------|--------|--------|--------|--------|--------|--------|
| No errors     | 1.00   | 5.0    | 20.0   | 10.0   | 15.0   | 35.0   | 25.0   |
| 1.05 × \( v_1 \) | 1.07   | 5.0    | 20.0   | 9.9    | 14.8   | 37.4   | 25.0   |
| 1.05 × \( v_2 \) | 1.00   | 5.0    | 20.0   | 9.8    | 14.7   | 34.8   | 25.0   |
| 1.05 × \( v_3 \) | 0.99   | 4.9    | 20.3   | 9.9    | 14.3   | 34.7   | 26.9   |
| 1.05 × \( v_4 \) | 0.99   | 4.9    | 21.5   | 9.6    | 14.7   | 32.6   | 31.0   |
| 1.05 × \( v_5 \) | 1.00   | 5.3    | 20.0   | 10.5   | 18.6   | 35.0   | 29.5   |
| 1.05 × \( v_6 \) | 1.00   | 4.0    | 20.1   | 8.0    | 11.5   | 34.9   | 30.5   |
| 1.05 × \( v_7 \) | 1.00   | 5.9    | 18.3   | 10.8   | 16.3   | 35.3   | 29.9   |
| 1.05 × \( v_8 \) | 1.00   | 5.2    | 19.9   | 11.6   | 15.8   | 35.2   | 26.7   |

Determination of the Kinetic Parameters \( V_{\text{exp}}, K_A, K_B, K_C, K_{AB}, K_{AC}, K_{BC}, \) and \( K_{ABC} \) for Mechanism V

For the completely random mechanism V, reaction 7 is added to the mechanism. The number \( N' \) of reactants is 11. The number of components \( C' \) is four (E, A, B, and C). The number of independent equilibrium relations \( R' \) is seven, and therefore, \( N' = C' + R' = 11 = 4 + 7 \). To calculate the equilibrium concentration of EABC using Solve, seven equilibrium equations and one conservation equation are required. This yields the following expression for the velocity

\[
v = \frac{[A][B][C]V_{\text{exp}}K_{AB}K_{BC}K_{AC}}{D_1 + D_2 + D_3}
\]

\[
D_1 = [C]K_AK_{AB}K_{ABC}K_BK_{BC}K_{AC} + [B][C]K_AK_{AB}K_{ABC}K_{AC}K_C + [B]K_BK_{ABC}K_{AC}K_{BC}K_C
\]

\[
D_2 = [A][C]K_AK_{ABC}K_BK_{BC}K_{AC} + [A][B][C]K_{ABC}K_{BC}K_{AC} + [A][B]K_{ABC}K_{AC}K_{BC}K_C
\]

\[
D_3 = [A]K_{ABC}K_{AC}K_{BC}K_{AC} + K_AK_{ABC}K_{BC}K_{AC} + K_AK_{ABC}K_{AC}K_{BC}K_C
\]

When \( v \) is measured at eight triplets of substrate concentrations \([A][B][C]\), the eight versions of rate equation 24 can be solved for the eight kinetic parameters.

The calculations here for mechanism V have been made with the following triplets of substrate concentrations: \{200,200,200\}, \{0.5,200,200\}, \{200,0.5,200\}, \{200,200,0.5\}, \{5,5,50\}, \{5,5,5\}, \{200,5,5\}, and \{5,200,5\}. The velocities calculated with these eight triplets are \( v_1 = 0.7113, v_2 = 0.03260, v_3 = 0.1498, v_4 = 0.02221, v_5 = 0.03851, v_6 = 0.01050, v_7 = 0.02329, \) and \( v_8 = 0.03830 \). A program calcinparaABC5 was written to calculate the eight kinetic parameters from eight measured velocities. This program has the same structure as calcinparaABC1 given in the Appendix, but there are eight lines, which are different versions of eq 24. The effects of 5% errors in velocities, one at a time, were calculated using this program, and they are summarized in Table 6.
TABLE 8: Effects of 5% errors in 8 velocities when calckinparsABC5 is applied to velocities for the completely-ordered mechanism

| Vexp | KA | KAB | KB | KC | KABC | KBC | KAC |
|------|----|-----|----|----|------|------|------|
| 1.00 | 5.0 | 20.0 | -8 $\times 10^3$ | 2 $\times 10^4$ | 5 $\times 10^3$ | 35.0 | 19.7 |
| 1.05 $\times v_1$ | 1.06 | 5.0 | 20.0 | 3 $\times 10^3$ | 2 $\times 10^4$ | 35.0 | -2 $\times 10^3$ |
| 1.05 $\times v_2$ | 1.00 | 5.0 | 20.0 | -2 $\times 10^3$ | 2 $\times 10^4$ | 35.0 | 2.4 |
| 1.05 $\times v_3$ | 0.99 | 4.9 | 21.5 | 4 $\times 10^2$ | 5 $\times 10^3$ | 32.7 | -1 $\times 10^4$ |
| 1.05 $\times v_4$ | 1.00 | 5.1 | 20.0 | -1 $\times 10^4$ | -4 $\times 10^2$ | 35.0 | -6.1 |
| 1.05 $\times v_5$ | 1.00 | 4.3 | 20.0 | 1 $\times 10^3$ | 3 $\times 10^2$ | 35.0 | -46.7 |
| 1.05 $\times v_6$ | 1.00 | 5.7 | 18.5 | -3 $\times 10^3$ | -6 $\times 10^3$ | 35.3 | -334 |
| 1.05 $\times v_7$ | 1.00 | 5.1 | 20.0 | -3 $\times 10^3$ | 2 $\times 10^4$ | 35.1 | -208 |

The determinations of equilibrium constants from the rate equations for the five mechanisms can be summarized as follows:

I: $K_A, K_{AB}, K_{ABC}$
II: $K_A, K_{B}, K_{AB}, K_{ABC}$, and indirectly $K_{BA}$
III: $K_A, K_B, K_C, K_{AB}, K_{ABC}$, and indirectly $K_{BA}, K_{CA}, K_{ACB}$
IV: $K_A, K_B, K_C, K_{AB}, K_{AC}, K_{ABC}$, and indirectly $K_{BA}, K_{CA}, K_{ACB}$
V: $K_A, K_B, K_C, K_{AB}, K_{AC}, K_{BC}, K_{ABC}$, and indirectly $K_{BA}, K_{CA}, K_{ACB}$, and $K_{CB}$

Use of calckinparsABC5 to determine the Mechanism When A + B + C $\rightarrow$ Products Is Studied for the First Time

An earlier article\(^4\) has shown that more general programs for calculating kinetic parameters can be used to determine the kinetic parameters for mechanisms with fewer kinetic parameters. This is demonstrated here by applying calckinparsABC5 to velocities calculated using the completely ordered mechanism. To do this, four more velocities have to be measured on A + B + C $\rightarrow$ products with the completely ordered mechanism in order to use calckinparsABC5. These additional velocities are $v(5,5,50) = 0.1370, v(5,5,5) = 0.01563, v(5,5,50) = 0.2725$, and $v(5,5,50) = 0.1064$. The first four velocities are given in the description of Table 2. The effects of 5% errors in velocities, one at a time, were calculated using calckinparsABC5, and they are summarized in Table 8.

This table shows that when the program for the completely random mechanism (calckinparsABC5) is applied to experimental data for the completely ordered mechanism, the expected values are obtained for $V_{exp}$, $K_A$, $K_{AB}$, and $K_{ABC}$, but unreasonable values are obtained for $K_B$, $K_C$, $K_{BC}$, and $K_{AC}$. This is the way the program communicates that the velocity data is for the completely ordered mechanism. The fact that $K_{AB}, K_{C}, K_{BC}$, and $K_{AC}$ are unreasonable is also shown by the sensitivity of these four kinetic parameters to 5% errors in measured velocities. Other more general programs can be used in this way. For example, calckinparsABC4 can be applied to experimental data for mechanism I. Application of less general programs to velocity data on more complicated mechanisms involves dropping measured velocities and does not provide useful information.

Discussion

The determination of kinetic parameters using the minimum number of velocity measurements makes it possible to study the kinetics of more complicated enzyme-catalyzed reactions. This method has already been illustrated with A + B $\rightarrow$ products\(^4\) when the experimental data follows rapid-equilibrium rate equations. Another advantage of deriving rate equations with a computer is that complicated rate equations are obtained in computer-readable form so that they do not need to be typed into the computer. The effects of experimental errors in velocities on the kinetic parameters are smaller when the substrate concentrations are chosen according to the general recommendations given here. The objective in this article is not to find the best set of triplets of substrate concentrations but to apply some general principles that indicate what can be done.

When the kinetic parameters for A + B + C $\rightarrow$ products are to be determined for the first time, it is recommended that the program for the completely random mechanism be used because it can identify the mechanism and determine the kinetic parameters in one operation. When the enzyme catalysis follows the completely random mechanism, all of the reactions 1–12 reactions are at equilibrium, not just those in the mechanism used to calculate the equilibrium concentration of EABC.

When the values of kinetic parameters have been calculated with one minimal set of measured velocities, it is recommended that the calculation be repeated with a second set of substrate concentrations to be sure the same values (within the expected experimental errors) are obtained. The precision of kinetic parameters can be improved by making replicate measurements of the velocities.

The steady-state rate equation for ordered A + B + C $\rightarrow$ products has seven terms in the denominator,\(^9\) but the rapid-equilibrium rate equation for the completely random mechanism has eight terms in the denominator. The number of terms in the denominator is useful in distinguishing between the five mechanisms discussed here. Usually, rapid-equilibrium rate equations have fewer denominator terms than steady-state rate equations. Cornish-Bowden\(^10\) gives a good discussion of reactions with three or more substrates and calls attention to the fact that Frieden\(^11\) was the first to call attention to the importance of the missing terms in the denominator of a rate equation.

The calculations in this paper support the following conclusions:

1. Computer programs using the minimum number of velocities can be used to calculate kinetic parameters for five rapid-equilibrium mechanisms of the reaction A + B + C $\rightarrow$ products.
2. The effects on the kinetic parameters of experimental errors in the velocities and substrate concentrations can be calculated, one at a time.
3. Equilibrium constants can be calculated for reactions not included in the mechanism used to derive the rapid-equilibrium rate equation.
4. More general computer programs can be used to identify a simpler mechanism and determine the kinetic parameters for the simpler mechanism.

Appendix

For mechanism I, the expression for the equilibrium concentration of EABC is derived by applying Solve in Mathematica...
Kinetic Parameters of Enzyme-Catalyzed A + B + C → Products

The following program uses Solve to calculate the kinetic parameters from four velocity measurements for mechanism I:

\[ \text{In: } \text{Solve}\left\{ \begin{align*}
{k_a} & = e^a e_a, \quad k_{AB} = e^a b e_{ab}, \\
{k_{ABC}} & = e^a b c e_{abc}, \\
{et} & = e + e_a + e_b + e_{abc}, \\
{e_{abc}, e, e_a, e_b} & \end{align*} \right\} \]

Out: \[[\{e_{abc} -> (a b c et) / (a b c + a b k_{ABC} + a k_A k_{ABC} + k_A k_{ABC} + k_A k_{ABC})\}]\]

Lower case type is used for equilibrium constants and concentrations of reactants because in Mathematica the name of a command like Solve begins with a capitol letter. In order to obtain the expression for the velocity \( v_1 \), it is necessary to replace \( et \) with \( vfexp \).

\[ v_1 = (a b c vfexp) / (a b c + a b k_{ABC} + a k_{ABC} + k_{ABC}) \]

The velocity of \( A + B + C \rightarrow \) products with specified kinetic parameters is given by

\[ \text{In: } \text{v1funconc} = v_1 / vfexp \rightarrow 1.00008, k_{A} \rightarrow 19.9884, k_{ABC} \rightarrow 35.0096 \]

Out: \[[\{vfexp, k_{A}, k_{AB}, k_{ABC}\}]\]

The full set of calculations for this article is freely available at MathSource (http://library.wolfram.com/infocenter/MathSource).

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