Thermal Dissociation and Conformational Lock of Superoxide Dismutase

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The kinetics of thermal dissociation of superoxide dismutase (SOD) was studied in 0.05 M Tris-HCl buffer at pH 7.4 containing 10^{-4} M EDTA. The number of conformational locks and contact areas and amino acid residues of dimers of SOD were obtained by kinetic analysis and biochemical calculation. The cleavage bonds between dimers of SOD during thermal dissociation and type of interactions between specific amino acid residues were also simulated. Two identical contact areas between two subunits were identified. Cleavage of these contact areas resulted in dissociation of the subunits, with destruction of the active centers, and thus, lost of activity. It is suggested that the contact areas interact with active centers by conformational changes involving secondary structural elements.

Keywords: Biochemical calculation, Conformational lock, Contact area, Kinetics, Superoxide dismutase (SOD), Thermal dissociation

Introduction

Superoxide dismutases (Cu, Zn-SODs) have been found in the cytoplasm of all the eukaryotic cells and in the periplasm of several bacterial species (Bannister et al., 1987; Kroll et al., 1995). Eukaryotic Cu, Zn-SODs are homodimers that contain one atom of zinc and one atom of copper per subunit and catalyze the dismutation of the superoxide anion at a diffusion-limited rate enhanced by electrostatic guidance of the substrate to the active site (Desideri et al., 1992). Cu, Zn-SODs possess a very compact structure that is highly resistant to denaturation by urea, SDS and proteolytic degradation. Several factors are thought to contribute to the enzyme stability, including the prosthetic metal ions (Forman and Fridovich, 1973), the intra-subunit disulfide bond (Abemethy et al., 1974), and the close packing of the hydrophobic interface between the subunits and the two halves of the β-barrel core (Getzoff et al., 1989). Amino acid sequence comparisons and the analysis of the three dimensional structure of the dimeric enzyme from Pleiogonathi have shown that the prokaryotic and eukaryotic Cu, Zn-SODs share a conserved ligand stereochemistry and a very similar monomer fold, based on a flattened Greek-key eight-stranded β-barrel (Bordo et al., 1994; Bourne et al., 1996; Imlay and Imlay, 1996; Battistoni et al., 1996).

Since the discovery of Cu, Zn-SOD, there has been serious in understanding how the dimeric structure contributes to the high catalytic efficiency and the remarkable stability of this class of enzymes. All attempts to obtain monomeric Cu, Zn-SODs by treatments with detergents (Rigo et al., 1978) or site-directed substitutions of hydrophobic residues at the dimer interface (Bertiini et al., 1994) provided also enzymes that display very low catalytic activity and gross alterations of spectroscopic properties. These dramatic changes probably reflect changes in the tertiary structure consequent to rearrangements of the solvent-exposed hydrophobic dimer interface.

The properties of interfaces in oligomeric enzymes and their influence on catalytic activity can be studied and explained by two independent methods, involving the use of the structural and kinetics data. Poltorak et al. (1999a; 1999b) reported that, in alkaline phosphatase from different sources, the same results that in both methods are in reasonable agreement. In our previous paper (Moosavi-Nejad et al., 2003), we had put forward a mechanism of Lentil seedling amine oxidase denaturation based on structural data. In
Materials and Methods

Materials Superoxide dismutase (SOD) from bovine erythrocyte [E.C: 1.15.1.1] was a product of Sigma (S2515). Pyrogallol, Tris, EDTA and HCl were the products of Merck.

Enzyme assay The activity of superoxide dismutase from bovine erythrocyte was determined by using an improved pyrogallol autoxidation inhibition assay (Marklund and Marklund, 1974; Zou et al., 1986; Luo et al., 1987; Yuan and Gao, 1997). Superoxide (O_2^-) is generated indirectly at alkaline pH by the action of oxygen on pyrogallol. SOD reacts with the superoxide and this slows down the rate of formation of the o-hydroxy-o-benzoquinone and other polymer products (Liang et al., 2000). One unit of SOD is defined as the amount of enzyme that reduces the rate of autoxidation of pyrogallol by 50%. (Marklund and Marklund, 1974; Zou et al., 1986; Luo et al., 1987; Yuan and Gao, 1997; Liang et al., 2000).

The assay is performed in 1ml of 0.05 M Tris-HCl buffer at pH 8.2 containing 10^{-7} M EDTA at 27°C in a thermostated Shimadzu UV-Vis-3100 spectrophotometer. The standard reaction mixture usually contained a certain concentration pyrogallol, which produces a rate of autoxidation of 0.07 absorbance unit per min measured at 325 nm. Under this condition, the amount of superoxide dismutase required to reduce the rate to 0.035 absorbance unit per min is defined as one unit of activity (Marklund and Marklund, 1974; Zou et al., 1986; Luo et al., 1987; Yuan and Gao, 1997; Liang et al., 2000).

Determination of optimum temperature Optimum temperature (T_{opt}) is defined as the maximum temperature at which the activity of the enzyme does not change during the incubation time (Segel, 1999). SOD samples at a concentration of 4.9 \times 10^{-5} M were incubated in multiItem III thermostat (Pharmacia Biotech, Uppsala, Sweden) at different temperatures (from 27°C to 77°C). A standard mixture of SOD, EDTA and HCl buffers at pH 7.4 containing 10^{-7} M EDTA for 150 minutes, cooled rapidly to 27°C, and assayed for residual activity.

Thermal dissociation of SOD SOD samples at a concentration of 4.9 \times 10^{-5} M, in 30 mM Tris-HCl buffer at pH 7.4 containing 0.1 mM EDTA, were incubated separately at 75 and 77°C. At suitable intervals, usually 5 minutes, samples were removed and cooled rapidly to 27°C and assayed for activity.

Biochemical computation by using liggplot and spdbv LIGPLOT (v.4.4.2) is a program for automatically plotting protein-ligand interactions, which automatically generates schematic diagrams of protein-ligand interactions from a given PDB file (Wallace et al., 1999). The interactions were taken into consideration as hydrogen bonds and hydrophobic contacts. Hydrogen bonds are indicated by dashed lines between the atoms involved, while hydrophobic contacts are represented by an arc with spokes radiating toward the ligand atoms they make contact with. The contact atoms are shown with spokes radiating back. The operating manual is located at the site: http://www.biochem.ucl.ac.uk/bsm/ligplot/manual/. Swiss-PdbViewer was used to deduce structural alignments and compare their active sites or any other relevant parts. Swiss-models was used to generate models, as it is possible to thread a protein primary sequence onto a 3D template and get an immediate feedback of how well the threaded protein will be accepted by the reference structure before submitting a request to build missing loops and refine sidechain packing. Swiss-PdbViewer can also read electron density maps, and provides various tools to build into the density. In addition, various modelling tools are integrated and command files for popular energy minimization packages can be generated. (Guex and Peitsch, 1997).

Results and Discussion

The properties of interfaces in oligomeric enzymes and their influence on catalytic activity can be studied and explained by two independent methods, involving the use of the structural data and kinetics data. Poltorak et al. (1999a, 1999b) reported that, in alkaline phosphatase from different sources, the same results in both methods are in reasonable agreement. They proposed (Poltorak et al., 1999b) that the catalytic reaction rate is a measure of active form concentration under given conditions, e.g. saturation with substrate. The simple thermal dissociation mechanism is:

\[ E_2 \leftrightarrow 2E_1 \leftrightarrow 2E_{\text{dim}} \]  \hspace{1cm} (scheme 1)

\( E_2 \) is a monomer capable of reversible transformation of the initial structure, \( E_{\text{dim}} \), the product of irreversible transformation (under given conditions) of \( E_1 \). The protein \( E_{\text{dim}} \) is unable to reassociate and form a functionally active protein complex \( E_2 \) (dimer form). Usually, there are several intermediate active ‘m’ forms of dimeric protein.

\[ E_{\text{dim}} \leftrightarrow E_2^{(1)} \leftrightarrow E_2^{(2)} \leftrightarrow E_2^{(3)} \ldots \leftrightarrow E_2^{(m)} \leftrightarrow K_{\text{dim}} \]  \hspace{1cm} (scheme 2)

Poltorak et al. (1999b) proposed that in the event of variations of the rate constants for the reciprocal transitions of various forms of catalytically active protein, m, the real number of forms is always greater than a certain quantity n that describes the irreversible transitions of n enzyme dimeric forms with similar rate constants. To determine this value from the kinetic curves it is convenient to introduce the dimensional ratio of period of induction \( \tau_1 \) and period of thermal dissociation \( \tau_2 \) shown in Fig. 1.

\[ \delta = \frac{\tau_1}{\tau_2 - \tau_1} - R - 1 \]  \hspace{1cm} (1)

where \( R \) is shown in the ordinate of Fig. 1.

For scheme 2, the exact analytical solution gives that value \( \delta \) is depending only of the number of steps before loss of activity m.
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\[ \delta = e^{-n} \left[ 1 + \frac{(n-1)^{n-2}}{(n-2)!} + \sum_{m=1}^{n-2} \left( \frac{(n-1)^m}{m!} \right) - 1 \right] \]  

(2)

where \( m \) is the total transition steps before dissociation of enzyme and \( n \) is the minimum transition steps which may be happened.

However, the exact calculation of \( n \) is practically impossible from this expression. For this reason, it was approximated by the empirical equation (Poltorak et al., 1999b)

\[ \delta = \frac{0.13(n-1)}{1 - 0.05n} \]  

(3)

or

\[ n = \frac{0.13 + \delta}{0.13 - 0.05\delta} \]  

(4)

Such calculation gives only limiting minimal value of \( n \) for scheme 2 (Poltorak et al., 1999b). These make it possible to estimate the minimal number of steps for a dimeric protein conversion by dissociation into catalytically inactive monomers.

To explain experimental data, Poltorak et al. (1998; 1999a; 1999b) proposed the phenomenon of “conformational lock” as an interprotein contact with possible partial breaks (loosening of binding between globules). Each successive state of the interprotein binding in an oligomeric enzyme cannot be achieved without the break of the preceding one. It has been shown that in some oligomeric enzymes, such as alkaline phosphatase, several contact areas are separated, and conformational lock can open stepwise through these areas.

The plot of residual activity of SOD versus temperature shows that the activity remained constant at incubation temperatures below 70°C. However, above 73°C, the activity rapidly reduces. The \( T_{\text{opt}} \) of SOD is calculated as 73°C (see Fig. 2).

The \( \delta \) and R values and calculated minimum thermal dissociative steps in the thermal dissociation of SOD based on scheme 3 was calculated by using the method of Poltorak et al. (1998; 1999a; 1999b) which is shown in Fig. 3. The quantitative values of these parameters show that minimum steps during the dimeric SOD thermal dissociation is two, which is consistent with the results obtained by crystal structure using methods of X-ray (PDB 1CBJ). The theoretical kinetics and processes of thermal dissociation and denaturation of SOD can be described as Scheme 3.

From the PDB information, SOD from bovine erythrocyte (PDB 1CBJ) consists of two subunits. Each subunit consists of 151 amino acid residues. There are two contact areas between the two subunits. There is a Greek key eight-stranded \( \beta \)-barrel in each subunit of SOD. This structure is very important for the biological function of SOD.
al., 1996; Battistoni et al., 1998). The Greek key eight-stranded β-barrel consists of β1(T2-L8), β2(Q15-K23), β3(T26-G35), β4(G39-Q47), β5(G91-D99), β6(G80-A87), β7(R113-E119), β8(S140-A150). The contact areas of SOD subunits are shown in the Table 1 and Fig. 4. The amino acid residues at contact areas and their interactions were presented at Figures 4 and 5.

The two subunits of SOD molecule make contact with each other through two identical contact areas by H-bonds and hydrophobic interactions: First, the residue of Ile149 on the subunit A interacts with the residues Gly112 and Gly49 of chain B through H-bonds and Lys151 on subunit A connect with Asp50 on subunit B through a H₂O molecule by H-bonds (see Figs 4 and 5). Second, the residues Val15, Val17, Thr17, Val146, Gly148, Ile149, Ala150 and Lys151 of subunit A interact with residues Phe48, Gly49, Asp50, Asn51, Thr52, Ile111, Gly112 and Arg113 of chain B through hydrophobic interactions (see Figs 4 and 5). When the thermal dissociation occurs, one of two contact areas may be cleaved (first intermediate) at step one, then the second contact area probably cleaved (second intermediate). Finally, the two subunits of SOD can be separated during thermal dissociation or conformational lock, and the dimeric SOD may be dissociated into monomeric structure (see scheme 3). Meanwhile, the spatial structure along the contact area may change. These spatial structural changes, for example, β1, β4, β7 and β8, might induce the reduction or lose of activity of SOD.

His44, His46 and His118 are located at active site of SOD from bovine erythrocyte, and are very important for the catalytic function of SOD (Hough et al., 2000; Miller, 2004). From the PDB information of SOD (PDB no:1CBJ), His44 and His46 are located on the β4(G39-Q47), His118 is located on the β7(R113-E119), the spatial structure changes of β4, β7 might cause spatial change of the active site and lead to the reduction or lose of activity of SOD. On the other hand, conformational change of β8(S140-A150) may affect the mobility of loops II (residues Glu119-Leu142), and the mobility in the Glu119-Leu142 loop region is very important for catalytic function of Cu, Zn-SOD (Hough and Hasnain, 1999; Hough et al., 2000; Hough and Hasnain, 2003). Moreover, there is a disulfide bridge between Cys144 and Cys55 in each subunit of SOD, and Cys144 is located on β8.

Fig. 3. Kinetics of thermal dissociation of SOD at different initial temperatures: 75°C (▲); 77°C (■). Conditions of thermal dissociation: [SOD]₀ = 4.9 × 10⁻⁷ M in 0.05 M pH 7.4 Tris-HCl buffer containing 10⁻⁴ M EDTA. R = 1 + δ is the ordinate intercept produced by a tangent to the inflection point of the kinetic curve; δ is a dimensionless parameter used to calculate a minimal number of active intermediate enzyme forms. And δ₇₅°C = 0.08, n₇₅°C = 1.67 ≈ 2; δ₇₇°C = 0.09, n₇₇°C = 1.75 ≈ 2.

Scheme 3. Hypothetical kinetics and processes of thermal dissociation and denaturation of SOD. Where E₂(3) is the native dimer with two intact contacts, E₂(2) and E₂(1) are the labile dimer (two intermediates), and E₁ is the inactive or weak active monomer. Eₐ is the denatured monomer.

Table 1 Contact areas and residues involved in conformational lock

<table>
<thead>
<tr>
<th>Contact area</th>
<th>Residues in chain A</th>
<th>Residues in chain B</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Val5,Val7, Thr17</td>
<td>Phe48-Gly49-Asp50-Asn51-Thr52, Ile111-Gly112-Arg113</td>
</tr>
<tr>
<td></td>
<td>Val146, Gly148-Ile149-Ala150-Lys151</td>
<td>Ile111-Gly112-Arg113, Val5,Val7, Thr17</td>
</tr>
<tr>
<td>II</td>
<td>Phe48-Gly49-Asp50-Asn51-Thr52, Ile111-Gly112-Arg113</td>
<td>Val146, Gly148-Ile149-Ala150-Lys151</td>
</tr>
</tbody>
</table>

*Conformational lock in SOD consists of two identical contact areas.
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It is reported that the intra-subunit disulfide bond is very important for maintaining the spatial structure and compactness of SOD (Abernethy et al., 1974). The spatial structure change of $\beta_8$ (S114-A150) may also affect the disulfide bond, which lead to the reduction of compactness and stability of SOD (Abernethy et al., 1974). It can also be seen that residues Val5 and Val7 are located on $\beta_1$, while residues Lys151, Ala150, Ile149, Gly148, Val146 are located at $\beta_8$. These residues are very close to the C-terminal and N-terminal (see Fig. 4). The disruption of conformational lock and dissociation of contact areas might induce the unfolding and lose of catalytic activity for SOD.

Structural analysis of the dimeric SOD showed that the two identical contact areas between the two subunits of SOD interact with elements of secondary structure of active centers and Greek-key eight-stranded $\beta$-barrel. This may be helpful to maintain the catalytic activity, spatial structure, compactness and stability of SOD, that is in agreement with the works of Rigo et al. (1978) and Bertini et al. (1994) that reported the monomeric SOD displays very low catalytic activity. Dissociation of these contacts might destroy the spatial structure of active center and the Greek-key eight-stranded $\beta$-barrel. The total number of contacts in the conformational lock is two and this is in good agreement with kinetic data, giving the same minimal number of intermediates during the thermal dissociation of SOD.

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References


