Hydroxyl Radical Generation by Mammalian Catalase: A Few Experimental Evidences

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Abstract
Catalase is well known antioxidant enzyme which protects cells from the toxic effects of hydrogen peroxide. Unlike peroxidase, it catalyses the dissociation of hydrogen peroxide directly into H2O and O2. Our results show that apart from the known catalytic action, mammalian catalase can generate Hydroxyl Radicals (HRs) in experimental conditions. At low concentrations of H2O2 (<20 µM), presence of bovine liver catalase (BLC) increased the generation of HRs in reaction mixture. Differential productions of HRs due to either presence of azide or urea induced structural changes of BLC suggest that the enzyme participate in the production of HRs. The possible character is unusual in contrast to the well established fact that catalase reduce the generation of reactive oxygen species in biological system. The work is being presented with discussion on experimental limitations we faced throughout the study.

Keywords: Mammalian catalase; Free radical; Hydroxyl radical generation

Abbreviations: 3AT: 3-amino-1,2,4-triazole; BLC: Bovine Liver Catalase; Bpy: 2,2'-Bipyridyl; EDTA: Ethylene Diamine Tetra Acetic Acid; Flu: Fluorescence; HRs: Hydroxyl Radicals; NAD(P)+: Nicotinamide Adenine Dinucleotide Phosphate; NAD(P)H: reduced form of NAD(P)+; ROS: Reactive Oxygen Species; TA: Terepthalic Acid

Introduction
Catalases are generally characterized by breaking down hydrogen peroxide into H2O and O2 through the mechanism where hydrogen peroxide alternately oxidizes and reduces the haem iron at the active site (Rea n 1,2) [1,2]. Although the structure, mechanism of action and functions of catalases have been studied for many years, research continues to reveal differences in the role of mammalian catalase compared to those of prokaryotes. For example, mammalian catalase use unbound NAD(P)H to prevent substrate inactivation without compared to those of prokaryotes. For example, mammalian catalase use unbound NAD(P)H to prevent substrate inactivation without displacing catalase-bound NADP(+) [3]. Studies on catalase have gained momentum recently owing to their postulated impact in extending life-span [4-6]. Recent reviews emphasize on the need of further research to answer the unexplained findings related to mammalian catalase [3,7].

Mammalian catalase is a homotetramer with a subunit molecular mass of ~60 kDa and belongs to a group of monofunctional catalases with small subunit size. Each subunit contains a haem group at the active site, which is internally located and approachable by a passageway that becomes narrow, thereby accessible for only substrates of small size. During catalytic reaction, intermediate compound-I at low concentrations of H2O2 and in the presence of one-electron donors may give rise to the compound-II; that is an inactive form of the enzyme (Rean 3) [3]. Reaction 1, 2 and 3 shows the alterations occurs at the active site during the catalytic reaction. 'Por' denotes porphyrin ring which with iron ion forms the haem group.

- Resting state (Por+=FeIII) + H2O2 ----> Compound-I (Por+=FeIV =O) + H2O [1]
- Compound-I (Por+=FeIV =O) + H2O2 ----> Resting state (Por+=FeIII) + H2O + O2 [2]
- Compound-I (Por+=FeIV =O) + e− ----> Compound-II (Por+=FeII =O) [3]

In the present work, we have shown that mammalian catalase (bovine liver catalase) is involved in the generation of Hydroxyl Radicals (OH·). The possible character is unusual in contrast to the well-known fact that catalase reduces the generation of reactive oxygen species (ROS) in biological system.

Materials and Methods
All chemicals used for the present experiments were of analytical grade. Bovine liver catalase (BLC) and 3-amino-1,2,4-triazole (3AT) was obtained from Sigma, USA. Terepthalic acid (TA) and other chemicals were obtained from HiMedia, India.

Preparation of the reagents
5 mM stock solution of TA was prepared in 50 mM sodium phosphate buffer (NaH2PO4/Na2HPO4) at pH 7.4. As per requirement it was diluted in the same buffer to prepare 0.5 mM TA working solution. 100 µM FeSO4.7H2O, 100 mM 2,2'-bipyridyl (Bpy), 10 mM 3AT and 10 mM sodium azide (NaN3) solutions were prepared in distilled water. After measuring initial concentration spectrophotometrically with help of molar extinction coefficient (43.6 M-1cm-1 at λ315, H2O2 was diluted to get 10,100 and 1000 µM concentrations. 40 mg BLC (2,950 units/mg of protein) was dissolved in 10 ml of 50 mM sodium phosphate buffer (pH 7.4). After that it was centrifuged at 3500 g for 10 minutes to get clear solution.

Detection of hydroxyl radical generation
TA was used as a specific fluorescent probe for the detection of HRs. Nonfluorescent compound TA reacts with hydroxyl radical to form a stable and highly fluorometric product 2-OH-TA (λex 315 nm, λem 395 nm).

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nm, λ 426 nm) [8,9]. In reaction mixture, HRs were generated by the well known Fenton’s reaction (Fe²⁺ + H₂O₂ → Fe³⁺ + OH⁻ + OH•). Both substrates of the Fenton’s reaction (H₂O₂ and FeSO₄·7H₂O; 100 µM each) were added consecutively in 2.5 ml of 0.5 mM TA working solution. To check the non-linearity of the curve 50, 100 and 200 µl quantity of the substrates were used. The reaction mixture was incubated at 37°C for 15 min. At the end of reaction, fluorescence intensity was measured with a Jasco FP-6300 spectrophuorometer in a 4 cm² quartz cuvette. Readings were taken at λ 315 nm, λ max 426 nm, band width 5 nm both for excitation and emission; and with high sensitivity. The fluorescence intensity was the measure of hydroxyl radicals (HRs) generated by the Fenton’s reaction.

**Effect of chelator on generation of HRs by free iron**

Bipyridyl, citric acid, fructose, EDTA·Na₂, glycine, L-arginine and L-lysine were used to analyse the relative strength of iron chelation. Free amino acids have been found effective for chelating iron in various studies [10,11]. 2 ml of 0.5 mM TA and 50 µl of 100 µM FeSO₄·7H₂O were added in seven test-tubes. 50 µl of each chelator (10 mM) was added in separate test tube followed by addition of 250 µl of 0.6% H₂O₂. It was incubated for 15 minutes and HR generation was measured as described in previous subheading.

**HR generation by catalase**

To show the HR generation by catalase following experiment was designed. As shown in the Table 1, 100 mM Bpy, BLC, 10 mM 3AT, 10 mM NaN₃ and 1000 µM H₂O₂ were added in 0.5 mM TA solution to prepare series of reaction mixtures (Rₓ). Final volumes were made equal by water and incubated for 10 minutes at 37°C before adding up 1000 µM H₂O₂. In addition, the R -4, 5 and 6 were prepared simultaneously with 10 and 100 µM H₂O₂. The three different concentrations (10, 100 and 1000 µM) provided 0.2, 2.0 and 20 µM H₂O₂ in final volume of Rₓ. After all additions reaction mixtures were incubated overnight at 37°C. Fluorescence was measured at λ max 315 nm, λ max 426 nm, band width 5 nm both for excitation and emission; and with high sensitivity. The detection of Hydroxyl Radicals (HRs) by TA is a highly sensitive and specific method and it has been used to detect HR generation in nonenzymatic systems [8,9]. In the present study we have used it for the first time for detection of HRs generated by catalase. TA is a perfect qualitative reagent and can also quantify HRs however in present study it was used to detect relative HR generation in different reaction mixtures.

**Determining the fluorescence spectrum of catalase in presence of urea**

To confirm the structural changes of BLC fluorescent spectra were recorded in each set of reaction mixtures prepared to see the effect of various concentrations of urea on HR generation by catalase. At λ max 280 nm fluorescence emission spectra was recorded (λ ex 270-400 nm) at medium sensitivity.

**Fluorescence (Flu. of one urea solution with BLC – Flu. of same urea solution without BLC) at each urea concentration was calculated.**

**Results**

The detection of Hydroxyl Radicals (HRs) by TA is a highly sensitive and specific method and it has been used to detect HR generation in nonenzymatic systems [8,9]. In the present study we have used it for the first time for detection of HRs generated by catalase. TA is a perfect qualitative reagent and can also quantify HRs however in present study it was used to detect relative HR generation in different reaction mixtures.

Figure 1 shows the units of fluorescence in the reaction mixture which was directly proportional to the amount of HRs produced by the Fenton’s reaction. At a given concentration of Fe²⁺/H₂O₂ system, iron chelating agents like glycine, fructose, L-arginine and L-lysine did not change HR generation significantly whereas presence of EDTA and citric acid increased the production of HRs at least by two folds (data not shown). Presence of Bpy reduced HR generation maximum when compared with other chelating agents. The concentration of Bpy was 200 µl.

**Effects of various concentrations of urea on generation of HRs by catalase**

To confirm the generation of HRs by catalase it (HR) was measured in presence of different concentrations of urea. 0.1, 0.2, 0.3 and 4 M urea solutions were prepared in 50 mM phosphate buffer at pH 7.4. 1.8 ml of each urea solution was taken in separate test-tube with 50 µl BLC. In the same way another set of test-tubes was prepared by adding 50 µl phosphate buffer instead of BLC. Thus at each given concentration of urea two test-tubes were prepared with and without BLC. After 10 minutes incubation at room temperature, 200 µl TA (5 mM), 200 µl Bpy, 50 µl H₂O₂ (100 µM) were added in each test-tube. Fluorescence was measured at room temperature with λ 326 nm, λ max 426 nm, band width 5 nm both for excitation and emission; at medium sensitivity. Δ

<table>
<thead>
<tr>
<th>No.</th>
<th>Reaction mixtures (Rₓ)</th>
<th>Fluorescence units</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>2.0 ml TA + 200 µl Bpy + 50µl H₂O₂</td>
<td>50µl H₂O₂</td>
</tr>
<tr>
<td>2.</td>
<td>2.0 ml TA + 200 µl Bpy + 10µl 3AT + 50µl H₂O₂</td>
<td>50µl H₂O₂</td>
</tr>
<tr>
<td>3.</td>
<td>2.0 ml TA + 200 µl Bpy + 10µl NaNO₃ + 50µl H₂O₂</td>
<td>50µl H₂O₂</td>
</tr>
<tr>
<td>4.</td>
<td>2.0 ml TA + 200 µl Bpy + 50µl BLC</td>
<td>50µl H₂O₂</td>
</tr>
<tr>
<td>5.</td>
<td>2.0 ml TA + 200 µl Bpy + 250µl BLC + 10µl 3AT</td>
<td>50µl H₂O₂</td>
</tr>
<tr>
<td>6.</td>
<td>2.0 ml TA + 200 µl Bpy + 250µl BLC + 10µl NaNO₃ + 50µl H₂O₂</td>
<td>50µl H₂O₂</td>
</tr>
</tbody>
</table>

Table 1: Fluorescence probe-TA, iron chelating agent-Bpy, enzyme-BLC and enzyme inhibitor-NaNO₃/3AT were added in six different test tubes according to following table. Reaction mixtures were incubated for 15 minutes at 37°C and afterwards H₂O₂ was added in each tube.
100 times more compared to Fe2+ in the reaction mixture (Figure 1) where it reduced the HR generation up to 55-60%.

Experimental findings presented in Figure 2 indicate that catalase generates Hydroxyl Radicals (HRs) when react with low concentrations (<20 µM) of H2O2. The initial fluorescence of the reagent blank may be due to contamination of iron in the reaction mixture participating through Fenton’s reaction (Rm -1). Enzyme inhibitors did not show any change in fluorescence activity of reagent (Rm -2 and 3). Presence of BLC (Rm -4) increased the HR generation; thus the fluorescence was enhanced by ~90 units. In Rm -6, NaN3 amplified the generation of HRs. 3AT did not alter the HR generation significantly (Rm-5).

HR generation by catalase (Δfluorescence units) was measured at different concentrations of urea (Figure 3). The generation of fluorescence due to catalase was 22 units in absence of urea. It was increased up to 30.3 units at 2 M urea and afterwards decreased up to 15.1 units at 4 M urea. In the same solutions protein fluorescence spectra were recorded (Figure 4). A protein fluorescence spectrum is widely used as a tool to monitor changes in proteins and to make inferences regarding local structure and dynamics [12]. Peak of fluorescence spectra of BLC shifted towards 350nm with increasing concentration of urea. It confirms the structural changes of BLC occurred in the presence of urea.

**Discussion**

During catalytic reaction, Hydroxyl radicals (HRs) are always thought to be generated due to impurities (mainly free iron ions) in reaction mixture [13]. As such, catalases are so well known for their antioxidant nature that perhaps no one looked for their oxidant role. Catalase activity is always measured in term of rate of either H2O2 degradation or O2 formation. The half life of HR is also very less (10-9 sec) [9]. Perhaps these are the reasons that the oxidant character of mammalian catalase was missed.

The linear increase in fluorescence with increasing amount of HRs generated by Fenton’s reaction confirmed the validity of the TA method (Figure 1). Both substrates of Fenton’s reaction (FeSO4 and H2O2) when added in equimolar concentrations produced HRs. Increase in concentration of these substrates in reaction mixture proportionally generates more HRs and accordingly they (HRs) produced more fluorescence by reacting with TA.

In our experiments, generation of HRs in the reaction mixture can occur by two possible sources; one is the free iron ion (Fe2+) and the other is by catalase. The bidentate chelator Bpy predominantly binds Fe2+ and avoids its participation in Fenton’s reaction. In reaction mixture, the HR generation by free iron was reduced by addition of Bpy so that small variations in fluorescence due to possible HR generation by the enzyme can be detected efficiently. The presence of Bpy will not affect generation of HR by catalase as the haem iron is present in deep buried active site which is not accessible to Bpy. Other well established chelators like desferrioxamine preferentially binds Fe3+ (product of Fenton’s reaction) hence they were purposefully avoided [14].

The presence of BLC increased the HRs generation in reaction mixture (Rm -4). It was required to confirm that HRs is being generated during catalytic reaction.
at the active site of the enzyme. Therefore further experiments were performed. Inhibitors were used to block the active site and structural changes were induced to modify near vicinity of active site.

His75 is present near the active site and it is involved in the mechanism of catalytic reaction. 3AT (3-amino-1,2,4-triazole) reacts with His75 and compound-I (intermediate of catalytic reaction) to inhibit catalytic reaction1. Sodium azide (NaN3) also acts as an inhibitor by converting compound-I into inactive compound-II [15]. The active site of catalase is accessible to both, 3AT and NaN3, so the production of HRs should be altered by the presence of these inhibitors of catalytic reaction.

Mechanism of azide inhibition (Rea\(^a\) 4 and 5) that showed that NaN3 promote the formation of inactive compound-II [15].

\[
\text{Compound I} + N_3^- + H^+ \longrightarrow \text{Compound II} + N_3^2 \quad [4]
\]

\[
\text{Compound II} + N_3^- + H^+ \longrightarrow \text{Catalase} \quad Fe^{3+} + N_3^2 + H_2O \quad [5]
\]

The formation of azidyl radical was found to be carbon monoxide-insensitive, indicated that resting state enzyme was not involved in the reaction. Detection of free radical formation provided evidence for one-electron oxidation/reduction mechanisms (Rea\(^a\) 3). It has been therefore proposed that the oxidation of azide by catalase proceeds by a cycle similar to peroxidases, involving Compound-I and Compound-II [15,16]. Most peroxidases, including lactoperoxidase, contain ferriprotoporphyrin IX as a prosthetic group. The formation of hydroxyl radical has been detected during the irreversible inactivation of lactoperoxidase in the presence of excess H\(_2\)O\(_2\) [17,18].

As shown in Figure 3, increase in fluorescence in the presence of azide (R\(_m\)=6) confirms that similar to lactoperoxidase, catalase was inhibited by azide and HRs was produced in the reaction mixture. If the HR generation was due to impurities in reaction mixture the same effect would be generated in R\(_m\)=3. 3AT require H\(_2\)O\(_2\) generation in reaction mixture at constant rate and a sensitive as well as specific fluorescent probe for free radicals in HRs. If HRs in reaction mixture were generated by catalase, it should be affected by structural changes of the enzyme. Different concentrations of urea were used to induce structural changes in the enzyme. At each concentration of urea, reaction mixture was prepared with and without addition of BLC. It was followed by the measurement of fluorescence intensity and calculation of ΔFluorescence (ΔFlu) as described in materials and methods. If the HR generation was only due to impurities, there should be equal production of HRs (ΔFlu) in reaction mixture (Figure 3). The HR generation was monitored only up to 4 M urea concentration because, as shown in Figure 4, at 5 M and above urea concentration there was a sharp increase in protein fluorescence. It indicates the major changes in protein structure like complete unfolding.

Our preliminary data like a) in presence of BLC the generation of HRs in reaction mixture was increased and b) differential productions of HRs due to either presence of azide or urea induced structural changes of BLC; indicate that the BLC participate in the generation of HRs at low concentrations (<20 µM) of H\(_2\)O\(_2\). However, the present study had certain limitations. It is required to have a system free from impurities to study the generation of HRs by catalase. The effect of free iron (participation in Fenton’s reaction) increased by many folds at high concentrations (mM) of H\(_2\)O\(_2\). The measurement of catalase activity at low concentrations (µM) of H\(_2\)O\(_2\) requires very sensitive instruments. Maintaining low concentrations of H\(_2\)O\(_2\) in reaction mixture is another difficulty because it is unstable at 37°C, optimum temperature for enzyme activity. Due to these limitations we could measure neither catalase activity under the conditions used for HR formation nor the generation of HRs at high concentrations of H\(_2\)O\(_2\). The fluorescence product 2-OH-TA is not available commercially therefore we measured relative HR generation in different reaction mixtures. We sincerely admit that we were unable to quantify the concentrations of HRs produced in different experiments and due to that, could not use statistical calculations. For each value, same experiment was repeated three times simultaneously and average of results was used for data interpretation. All experiments were repeated once again after a gap of one month and the pattern of findings were same. The quantification of HRs produced by catalase was also required for the study of reaction kinetics.

The mammalian catalase has been described “a venerable enzyme with new mysteries”. There are several findings related to mammalian catalase which are still unexplained [3,4,7]. For example, in specific conditions enzyme consume H\(_2\)O\(_2\) as its only substrate without generating O\(_2\) [19]. Recent studies provide evidence of interlink between intracellular catalase, amyloid aggregation and increased ROS generation; and role of these interactions in Alzheimer disease [20]. HR generation can be a new feature of mammalian catalase which could provide explanation of unexplained findings. It requires further studies coupled with advanced techniques to establish our findings completely.

**Conclusion**

HR generation by BLC catalase is a new possible product. A system free from impurities, generation of low concentrations of H\(_2\)O\(_2\) at constant rate and a sensitive as well as specific fluorescent probe for detection and quantification of HRs are prerequisites to confirm the findings. If characterized completely, the new character will provide the opportunity to explore the role of catalase again as an ‘HR producer’ in cell signaling, aging and apoptosis making it more relevant in human health and diseases.

**References**