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Construction of a Fluorescent H$_2$O$_2$ Biosensor with Chitosan 6-OH Immobilized β-Cyclodextrin Derivatives

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Abstract: In the present work, a fluorescent H$_2$O$_2$ biosensor was constructed by encapsulating fluorescent probe Rhodamine B (RhmB) in the hydrophobic cavity of the cyclodextrin (β-CD) and immobilizing catalase (CAT) on the 2-NH$_2$ of chitosan (CTS) in a chitosan 6-OH immobilized β-cyclodextrin derivative (CTS-6-CD). The inclusion complex of CTS-6-CD to RhmB (CTS-6-CD-RhmB) was prepared by a solution method. Its structure and inclusion efficiency were determined by Fourier transform infrared spectroscopy (FTIR), X-ray diffraction (XRD), and fluorescence spectroscopy (FL). CAT was immobilized on CTS-6-CD-RhmB to eventually form the functional membrane, CTS-6-CD-RhmB-CAT, via glutaraldehyde crosslinking, which was further characterized by FTIR and FL, and used as a H$_2$O$_2$ biosensor. The functional membrane was used to simultaneously oxidize and detect H$_2$O$_2$. The detection condition was optimized as pH 8, a reaction temperature of 25 °C, and an immobilized enzyme concentration of 2 × 10$^{-4}$ mol/L. The fluorescence response of the biosensor exhibited a good linear relationship with the concentration of H$_2$O$_2$ in the range of 20 mM–300 µM and the detection limit of 10$^{-8}$ mol/L.

Keywords: chitosan; catalase; β-cyclodextrin; Rhodamine B; fluorescent biosensor

1. Introduction

Reactive oxygen species (ROS) are involved in the initiation of biological effects of various factors. Their concentration can significantly affect many biological activities [1–3]. H$_2$O$_2$ is a representative of ROS. A low H$_2$O$_2$ concentration can be used as a signal transduction and expansion of the second messenger. High concentrations of H$_2$O$_2$ can cause oxidative stress and trigger a variety of diseases and physiological disorders [4,5]. Therefore, the real-time detection of H$_2$O$_2$ is highly desired to understand its relationship with human diseases, accurately diagnose diseases, and monitor disease development [6].

Various H$_2$O$_2$ detection techniques, such as spectrophotometry, high performance liquid chromatography, chemiluminescence, fluorescence spectrophotometry, electrochemical analysis, and so on, have been reported [7,8]. Among them, fluorescent biosensor has been used in a variety of fields, especially on-site H$_2$O$_2$ detection, due to its low cost, convenience, and high sensitivity. Recently, fluorescent H$_2$O$_2$ biosensors have attracted considerable attention. For example, Han [9] constructed three novel low cost and highly selective fluorescent H$_2$O$_2$ biosensors with silver nanoclusters and nucleic acid dyes. The biosensors were successfully used for the H$_2$O$_2$ detection in the presence of Fe$^{2+}$ without any complex modification required.
Chitosan (CTS) is the only natural alkaline polysaccharide which possesses a unique 2-NH\(_2\) structure. Its excellent film-forming property, degradability, and biocompatibility make it an important carrier of biosensing films [10–13]. The modified CTS can significantly improve the detection performance of various biosensors.

β-cyclodextrin (β-CD) is hydrophilic due to its external hydroxyl groups, but possesses a hydrophobic cavity with a certain size [14]. Some small molecules can be included in the cavity of β-CD or its derivatives by hydrophobic, hydrogen, and van der Waals forces to form fluorescent biosensors. The force to include the small molecules decides the selectivity of the biosensors. In addition, β-CD can improve the solubility and stability of the included compounds [15], which thus increases the fluorescence intensity of the guest molecules, and significantly improves the sensitivity of the corresponding fluorescent sensors. Thus, β-CD has been widely used in fluorescence enhancement [16,17].

To improve the performance of fluorescent biosensors, attempts have been made to combine the excellent properties of CTS with those of β-CD by modifying CTS with β-CD in biosensor construction. However, most studies have been focused on preparation of CTS-CD by the reaction of cyclodextrin derivatives with the highly active 2-NH\(_2\) groups on the chain of chitosan, which is unconducive to the utilization of the biologically active 2-NH\(_2\) of CTS. We previously systematically explored the preparation of various CTS-6-CD derivatives. For example, Chen et al. [18] immobilized cyclodextrin on the 6-OH of CTS by the nucleophilic substitution method, where the amino group of CTS was pre-protected with phthalic anhydride. The 2-NH\(_2\) was then deprotected with hydrazine hydrate. The resultant CTS-6-CD derivative exhibited good solubility with the biological activity of 2-NH\(_2\) unaffected. In the present work, we proposed a novel method for the construction of fluorescent H\(_2\)O\(_2\) biosensors with CTS and β-CD. A catalase was immobilized on the 2-NH\(_2\) of CTS and the fluorescent probe rhodamine B (RhmB) was encapsulated in the hydrophobic cavity of the cyclodextrin immobilized on the 6-OH of CTS to form a functional membrane CTS-6-CD-RhmB-CAT (Figure 1).

It was then used for the H\(_2\)O\(_2\) detection due to the synergistic effect of these two functional groups. Such strategy was able to simplify the detection of H\(_2\)O\(_2\) and the separation of catalase, and increase the stability of RhmB in detection. The oxidation and detection of H\(_2\)O\(_2\) were realized on the same functional membrane.

![Figure 1. Schematic diagram of experimental structure and detection process.](image-url)
2. Results and Discussion

2.1. Characterization of CTS-6-CD-RhmB

The IR, XRD, and fluorescence emission spectra of CTS-6-CD-RhmB, CTS-6-CD, and RhmB are shown in Figure 2. It is clear that the IR spectrum of CTS-6-CD-RhmB is not a superposition of those of CTS-6-CD and RhmB (Figure 2a). The benzene ring vibration peaks of RhmB at 1600 cm\(^{-1}\) and 1480 cm\(^{-1}\) disappeared and the intensity of C–C stretching vibration peak at 1250 cm\(^{-1}\) was significantly reduced after the inclusion, indicating that the hydrophobic O-benzoic acid group of RhmB was included in the cavity of \(\beta\)-CD [19]. The intensities of the absorption peaks at 1030 cm\(^{-1}\) were attributed to the pyranyl groups of \(\beta\)-CD and CTS, respectively, were sharply decreased, and became wider after the inclusion due to the “inclusion compound infrared absorption attenuates effect” [20]. The strong and broad band at 3400 cm\(^{-1}\) that was ascribed to the hydrogen bond between the hydroxyl group and amine group of CTS-6-CD was weakened after the inclusion due to the increased distance between CTS chain by the included RhmB. For the XRD spectrum, the inclusion of RhmB in CTS-6-CD led to a new extra diffraction peak at 15\(^\circ\), and two weak peaks at ~28\(^\circ\) and 35\(^\circ\). By the way, the intensity of CTS-6-CD around 2\(\theta\) = 20\(^\circ\) was decreased (Figure 2b). It can be explained that RhmB was included in CTS-6-CD via hydrogen bonding and van der Waals force, which increased the irregularity of the macromolecular chain and weakened the intensities of its diffraction signals, and the peaks assigned to the structure of RhmB stretched outside the cavity of cyclodextrin emerged. The fluorescence spectra of RhmB, CTS-6-CD, and CTS-6-CD-RhmB were compared. It could be found that RhmB exhibited a strong emission peak at 582 nm and CTS-6-CD exhibited no emission peak under the excitation at 552 nm (Figure 2c). The inclusion of RhmB in CTS-6-CD caused a blueshift of its emission peak because the inclusion limited the movement of the rhodamine molecules and thus caused certain damages to the conjugate system. These results indicate that RhmB was successfully included in CTS-6-CD.

The standard curve of the cyclodextrin clathrate to RhmB is shown in the main curve of Figure 2d. 1/(F – F\(_0\)) exhibited a linear relationship with 1/C\(_0\), indicating the inclusion ratio was 1:1. The inclusion constant K was calculated to be 3.74 × 10\(^3\), indicating that the inclusion reaction occurred spontaneously at room temperature.

The inclusion efficiency was calculated to be 44.6% based on the load capacity of CTS-6-CD of 212.16 \(\mu\)mol/g and the inclusion amount of 94.57 \(\mu\)mol/g. Since cyclodextrin was immobilized on the CTS molecular chain, the degree of freedom of its molecular movement was limited. In addition, it tends to curl in solvents, and is thus partially shielded. Therefore, the inclusion efficiency of CTS-6-CD was lower than that of free \(\beta\)-CD, but still reached ~50%. It was calculated from the standard curve of inclusion effect in the inserting figure of Figure 2d according to the formula in Section 3.5 that the maximum inclusion amount of RhmB in free \(\beta\)-CD is ~150 \(\mu\)mol/g. The inclusion amount of RhmB in the immobilized \(\beta\)-CD reached 63% of the maximum inclusion amount of RhmB in free \(\beta\)-CD, indicating that the inclusion complex was able to maintain the sensitivity and accuracy of the \(\beta\)-CD based sensor.

CTS-6-CD has excellent film-forming properties due to its CTS macromolecule chains, which can increase the stability of its modified electrode. The 2-NH\(_2\) group on the CTS backbone in CTS-6-CD provides a binding site for multiple enzymes. The hydrophobic cavity of the cyclodextrin in CTS-6-CD provides a suitable environment for the inclusion of many small molecules. The immobilization promotes chemical stability. Therefore, CTS-6-CD is an excellent material for formation of single or composite functional membranes.
2.2. Characterization of CTS-6-CD-RhmB-CAT

The immobilization of CAT on CTS-CD-RhmB weaken the IR absorption peak of the rocking vibration of -NH2 at 1580 cm\(^{-1}\), but significantly enhanced the stretching vibration absorption peak of C-N at 1640 cm\(^{-1}\) (Figure 3a). It can be explained that the 2-NH2 in CTS was occupied by glutaraldehyde to form a Schiff base [21,22]. The absorption peak at 1720 cm\(^{-1}\) was attributed to the suspended aldehyde group that was not linked with the enzyme [23]. A new peak appeared in the amide I band at 1660 cm\(^{-1}\) due to the stretching vibration of carbonyl in the skeleton peptide chain [24]. As shown in Figure 3b, the addition of H\(_2\)O\(_2\) significantly reduced the fluorescence intensity of CTS-6-CD-RhmB-CAT due to the static fluorescence quenching where H\(_2\)O\(_2\) was decomposed by CAT into hydroxyl radicals, and the included RhmB reacted with the hydroxyl radicals to form a non-fluorescent substance. These results indicate that CTS-6-CD-RhmB-CAT was successfully prepared.

Figure 2. Characterization of CTS-6-CD-RhmB (a) Infrared spectrum of CTS-6-CD-RhmB, CTS-6-CD and RhmB; (b) XRD spectra of CTS-CD-RhmB and CTS-6-CD; (c) Fluorescence spectroscopy of CTS-6-CD-RhmB; (d) \(1/\Delta F\) vs. \(1/C\) curve and standard curve of inclusion effect of cyclodextrin on RhmB (the inserting figure).

Figure 3. Characterization of CTS-6-CD-RhmB-CAT (a) Infrared spectrum of CTS-6-CD-RhmB-CAT and CTS-6-CD-RhmB; (b) The influence of the addition of H\(_2\)O\(_2\) on fluorescence spectroscopy of CTS-6-CD-RhmB-CAT.
2.3. Fluorescence Detection of H$_2$O$_2$ by CTS-6-CD-RhmB-CAT

The condition for the fluorescence detection of H$_2$O$_2$ by CTS-6-CD-RhmB-CAT was optimized to evaluate the detection performance.

Figure 4a shows the effects of pH on the fluorescent response of CTS-6-CD-RhmB-CAT to H$_2$O$_2$. It is clear that CTS-6-CD-RhmB-CAT exhibited the strongest response to H$_2$O$_2$ at pH ~8. It can be explained that H$_2$O$_2$ was decomposed into hydroxyl radicals under the catalysis of catalase. The hydroxyl radicals oxidized RhmB and quench its fluorescence. Catalases have the highest activity under the biological condition. The catalase produced the highest amount of hydroxyl radical at pH ~8. Therefore, the pH for the H$_2$O$_2$ detection by CTS-6-CD-RhmB-CAT was optimized as 8.

The effects of reaction time on the fluorescence quenching of CTS-6-CD-RhmB-CAT by H$_2$O$_2$ were also investigated. It can be seen from Figure 4b that the fluorescence intensity remained stable within 10 min, significantly reduced at 15 min, and remained stable after 25 min reaction. The CTS-6-CD-RhmB-CAT exhibited certain fluorescence even after the reaction, indicating that the reaction between H$_2$O$_2$ and the enzyme was complete. Therefore, the time for the reaction between CTS-6-CD-RhmB-CAT and H$_2$O$_2$ was optimized to be 25 min. Su et al. [25] successfully synthesized a fluorescent gold nanoclusters and stabilized it with lysozyme under alkaline conditions for the diction of alkaline protease. They found that the optimal reaction time for the reaction was 3 h. In contrast, the biosensors we constructed in the present work significantly shortened the detection time and realized rapid detection for H$_2$O$_2$.

Figure 4c shows the effects of temperature on the H$_2$O$_2$ detection. The fluorescence quenching effect dramatically increased with the increase of temperature from 0 °C to 25 °C. It can be explained
that the collision frequency between the catalase immobilized on the chitosan derivative and H₂O₂ was low at low temperatures, e.g., the collision between fluorescent substance and target molecules was low, resulting in weak fluorescence quenching effect. The collision frequency between the immobilized catalase and H₂O₂ increased with the increase of temperature, resulting in stronger fluorescence quenching effects. As the temperature further increased to 60 °C, no significant change was found in the fluorescence intensity. In the temperature range of 25–60 °C, the fluorescence quenching effect increases with the increase of temperature. Meanwhile, the fluorescence emission efficiency and fluorescence intensity of CTS-6-CD-RhmB-CAT in the solution decrease with the increase of temperature due to the decomposition of RhmB at high temperatures. In addition, optimum active temperature of the catalase is 25 °C. High temperatures can deactivate the enzyme, leading to a decreased capacity to produce hydroxyl radicals. Therefore, the detection temperature was optimized at 25 °C.

CTS-6-CD-RhmB-CAT concentration can directly affect the sensitivity and resolution of the biosensor. A low concentration leads to low sensitivity towards the target substances. Extremely high concentrations reduce the efficiency of the sensor and utilization of the derivative. Therefore, the optimization of CTS-6-CD-RhmB-CAT concentration is of paramount importance. Figure 4d shows the effects of CTS-6-CD-RhmB-CAT concentration on H₂O₂ detection. The fluorescence quenching effect linearly increased with the increase of CTS-6-CD-RhmB-CAT concentration from 0.1 to 200 μmol/L. Further increasing CTS-6-CD-RhmB-CAT concentration showed no significant effect on the fluorescence quenching effect, indicating increasing the concentration of CTS-6-CD-RhmB-CAT to over 200 μmol/L had no significant influence on the detection sensitivity. Therefore, the optimal CTS-6-CD-RhmB-CAT concentration was determined to be 200 μmol/L.

Based on these results, the optimal condition for the detection of H₂O₂ by the CTS-6-CD-RhmB-CAT biosensor can be summarized as follows: pH 8.0, the reaction temperature of 25 °C, and the CTS-6-CD-RhmB-CAT concentration of 200 μmol/L.

2.4. Quantitative Detection of H₂O₂

The fluorescence responses of CTS-6-CD-RhmB-CAT biosensor to different concentrations of H₂O₂ under the optimal condition are shown in Figure 5.

It is clear that the fluorescence intensity of CTS-6-CD-RhmB-CAT was gradually decreased with the increase of H₂O₂ concentration (Figure 5). CTS-6-CD-RhmB-CAT exhibited a strong fluorescence emission at 552 nm that varied with the change of H₂O₂ concentration. To more intuitively determine the relationship between fluorescence intensity (FI) and H₂O₂ concentration (C), the fluorescence
intensity at 552 nm was plotted vs. H$_2$O$_2$ concentration (Figure 6). The result indicates that the fluorescence intensity of the CTS-6-CD-RhmB-CAT biosensor has an excellent linear relationship with H$_2$O$_2$ concentration in the range of 20 mM–300 µM with the fitting equation

$$FI = -0.22C + 575.80 \ (R = 0.9919)$$

Based on the linear regression curve, the detection limit of the H$_2$O$_2$ biosensor was determined to be $10^{-8}$ mol/L. Jiang et al. prepared an electrochemical H$_2$O$_2$ sensor with chitosan, carbon nanotubes, and immobilized catalase. The sensor exhibited a detection limit of 2.5 µmol and a linear detection range of 5–50 µM. Sedigheh et al. fabricated an electrochemical H$_2$O$_2$ sensor with a detection limit of 8.7 µmol and a linear detection range of 10–100 µmol by immobilizing catalase on a multi-walled nanotube and thionine membrane [26]. Compared with these H$_2$O$_2$ sensors, our CTS-6-CD-RhmB-CAT functional membrane biosensor has a simple structure, but with a lower detection limit and much wider detection range.

![Figure 6. Linear relationship between fluorescence intensity and H$_2$O$_2$ concentration.](image)

### 3. Materials and Methods

#### 3.1. Materials

Chemical grade rhodamine B (RhmB) was purchased from Tianjin Shibo Chemical Co., Ltd. (Tianjin, China). β-Cyclodextrin (A.R.) was supplied by Tianjin Kwangfu Fine Chemical Industry Research Institute (Tianjin, China). Chitosan (CTS) with a deacetylation degree of 95% and molecular weight of $1.1 \times 10^6$ was provided by Zhejiang Aoxing Biochemical Co., Ltd. (Yuhuan, Zhejiang, China). Catalase (3500 units/mg) was purchased from Aladdin Industrial Co., Ltd. (Shanghai, China). The chitosan 6-OH-loaded cyclodextrin derivative (CTS-6-CD) was prepared as described previously [18]. The loading of cyclodextrin was determined to be 212.16 µmol/g. Glutaraldehyde was purchased from Tianjin Fuchen Chemical Co., Ltd. (Tianjin, China). Other reagents were all analytical grade and used as received.

#### 3.2. Synthesis of CTS-6-CD-RhmB

CTS-6-CD was dissolved in ethylenediamine-ethanol (v:v = 1:1) solution. Excessive RhmB was added to the CTS-6-CD solution and refluxed in an oil bath at 60 °C for 6 h. The produced precipitate was washed with secondary deionized water and ethylene glycol until the wash solution became neutral, and low-temperature vacuum dried to afford CTS-6-CD-RhmB inclusion compound.
3.3. Synthesis of CTS-6-CD-RhmB-CAT

The obtained CTS-6-CD-RhmB was dissolved in 1% acetic acid aqueous solution at \( \frac{m_{\text{CTS-6-CD-RhmB}}}{m_{\text{Acetic acid}}} = 3\% \), allowed to swell for 3 h, and neutralized with a sodium hydroxide solution. Glutaraldehyde was then added to the neutralized CTS-6-CD-RhmB solution with a mass fraction of 2.5%, allowed to react for 5 min, mixed with a 2.5% (w/v) catalase (CAT) solution, and allowed to react at 4 °C for 120 min. The product was vacuum filtered and washed with distilled water and phosphate buffer to afford CTS-6-CD-RhmB-CAT.

3.4. Characterization of CTS-6-CD-RhmB and CTS-6-CD-RhmB-CAT

The IR spectra of CTS-6-CD-RhmB and CTS-6-CD-RhmB-CAT was recorded on a NicoLet NEXUS-470 Fourier transform infrared spectroscopy using KBr pellets.

The XRD patterns of CTS-6-CD and CTS-6-CD-RhmB were collected on an ALC-100.4 polycrystalline X-ray diffractometer (Beijing Sartorius Instrument Systems Co., Ltd., Beijing, China) in the \( \theta \) range of 0°–60°.

The fluorescent emission spectra of CTS-6-CD-RhmB-CAT and CTS-6-CD-RhmB were measured under the excitation at 552 nm with both excitation and the emission slit widths set to 2 nm using a 970CRT type fluorescence spectrophotometer (Shanghai Precision Science Instrument Co., Ltd., Shanghai, China).

3.5. Determination of Inclusion Constant, Inclusion Amount, and Inclusion Efficiency

The reaction between \( \beta \)-CD and RhmB can be expressed as

\[
\text{CD} + n\text{RhmB} = [\text{RhmB}]_n - \text{CD}
\]

The equilibrium constant can be calculated as

\[
K = [C_{\beta-\text{CD-RhmB}}] \cdot [\text{CD}]^{-1} \cdot [\text{RhmB}]^{-n}
\]

If the inclusion ratio is 1:1, then

\[
1/(F - F_0) = 1/(\alpha C_{\text{RhmB}0} C_{\beta0} K) + 1/(\alpha C_{\text{RhmB}0})
\]

where \( C_{\text{RhmB}0} \) is the initial concentration of RhmB (10^{-5} \text{ mol/L}), \( C_{\beta0} \) is the initial concentration of \( \beta \)-CD (10^{-4} \text{ mol/L}), and \( F - F_0 \) is the difference between the absorptions before and after \( \beta \)-CD added.

To calculate the inclusion constant, \( K \), \( \beta \)-CD/RhmB dimethyl sulfoxide solutions with different mole ratios were prepared and their absorptions were measured in the range of 300–700 nm with a TU-1810 UV–Vis spectrophotometer (General Analytical Instrument Co., Ltd., Beijing, China) to establish a fitting curve of \( 1/(F - F_0) \) vs. \( \beta \)-CD concentration. The \( \beta \)-CD concentration of 0 mol/L was used as the initial condition.

To determine the inclusion amount, RhmB solutions of different concentrations in diluted sulfuric acid were prepared, and measured for their fluorescent emission to establish a standard curve equation of

\[
F = 419.93 C_{\text{RhmB}} + 52.87 \quad (R^2 = 0.9985)
\]

The concentration of RhmB in the inclusion solution was obtained by fitting the standard curve equation. The inclusion amount was then calculated using

\[
I = CV/Mm \quad (1)
\]

where \( I \) is the amount of guest molecules per gram of the inclusion product (\( \mu \text{mol/g} \)), \( C \) is the concentration of RhmB obtained from the standard curve equation (mg/mL), \( V \) is the volume of the
inclusion compound solution, M is the molecular weight of the guest molecule, and m is the mass of clathrate compound.

4. Conclusions

Based on the excellent properties of chitosan, the high bioactivity of 2-NH₂ on the skeleton of chitosan, and the hydrophobic cavity of cyclodextrin, a functional membrane CTS-CD-RhmB-CAT was prepared and its application in rapid H₂O₂ detection was explored. RhmB was included in CTS-6-CD derivative by a solution method. The clathrate product with glutaraldehyde was further linked with CAT to form a fluorescent biosensor for quantitative detection of H₂O₂. The detection condition was optimized as pH 8, the reaction temperature of 25 °C, and the CTS-6-CD-RhmB-CAT concentration of 200 µmol/L. The fluorescence response of the fluorescent biosensor exhibited a good linear relationship with H₂O₂ concentration in the range of 20 µM–300 µM and the detection limit is 10⁻⁸ mol/L, indicating the high sensitivity of the biosensor. Our work provides a new approach to the application of the important chitosan derivative, CTS-6-CD.

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Conflicts of Interest: The authors declare no conflict of interest.

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