Fluorescein Derivative Immobilized Optical Hydrogels: Fabrication and Its Application for Detection of H$_2$O$_2$

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Abstract: A novel fluorescein-based probe FLA-Boe was developed for detecting H$_2$O$_2$. Modified by 2-Bromomethylphenylboronic acid pinacol ester, FLA-Boe is a Fluorescein derivative with eminent photostability and remarkable H$_2$O$_2$ sensitivity and selectivity. FLA-Boe was utilized to synthesize hydrogel sensors in the manner of guest–host interaction by taking advantage of its aforementioned features. The hydrogel sensor can be used to detect H$_2$O$_2$ effectively in both flowing and static water environments with satisfactory performance. It is expected that this application may open a new page to develop a neoteric fluorescent property analysis method aiming at H$_2$O$_2$ detection.

Keywords: hydrogen peroxide; hydrogel sensor; naked-eye detection

1. Introduction

Hydrogen peroxide (H$_2$O$_2$), a typical reactive oxygen species (ROS), functions as a key throughout cell growth, proliferation, host defense, and signal transmission pathways under a physiological environment [1,2]. However, the chemical and pharmaceutical industries inevitably use hydrogen peroxide as an oxidizer for related production activities, resulting in the discharge of excessive hydrogen peroxide in wastewater, which severely impacts people’s living environment and the water treatment system [3]. Excessive H$_2$O$_2$ in the human body triggers the pathogenesis of many disorders, such as inflammation, Alzheimer’s disease, cardiovascular disease and, more seriously, cancer [4]. Consequently, it is vital to monitor H$_2$O$_2$ in the inner condition and outer environment of the human body [5]. Undoubtedly, the development of H$_2$O$_2$ fluorescent sensors has been arousing people’s interest. Thus, the design and fabrication of sensing systems with H$_2$O$_2$ responsive characteristics are key for human health and environmental protection.

Macroscopic supramolecular hydrogels are excellent candidate materials for intelligent sensing activity. In comparison with naturally existing hydrogels, such as polysaccharides and proteins, assorted synthetic hydrogels have been applied as smart materials on account of their functionalization during synthesis. Furthermore, most hydrogels are economical and environmentally friendly with transparent and stretchable features [6,7]. Nowadays, composite materials based on macroscopic supramolecular hydrogels have been investigated for optical detection. Using hydrogel sensors to detect test objects in environmental samples is a handy, easily operated, low-cost, analytical method with remarkable sensitivity [8]. Through research, it is possible to conduct naked-eye observation and direct analysis without spectroscopic devices and apparatus. Hitherto, diverse synthetic strategies using hydrogel sensors have been extensively explored and studied, accompanied by their
According to experience, device-free sensor systems aiming at colorimetric and fluorometric detection by the naked eye are of the most extensive and applicable utility ad hoc; the aforementioned hydrogel sensors are easily manipulated, and are highly adaptive to various conditions. As mentioned above, cyclodextrin moieties can be incorporated into pre-existing polymeric materials via grafting reactions. Interpenetrated networks can also be used to develop new selective and synergistic sorption capacities for specific purposes, such as the fabrication of a hydrogel sensor. Surmounting the shortcomings existing among traditional fluorescence detection methods, such as monitoring limited, complicated detection processes and man-made mistakes requires the application of hydrogel immobilized sensors with response signals that are adaptive to the living environment and that are able to simplify the detection operation and acquire more convincing and urgently needed detection results. Thus, the development of hydrogel sensors for detecting $H_2O_2$ with dual colorimetric and fluorescence signals is of profound significance. Crosslinking and grafting are applied to prepare covalently-attached cyclodextrins. The application of a supramolecular sensing system has a promising application prospect. With regard to complexity, functionality and capabilities, the evolution of cyclodextrin polymers matches the modern requirements of macromolecular materials.

In this study, fluorescein-based derivatives are selected as the fluorophores by structural modifications and the phenylboronic acid ester acts as the recognition site for $H_2O_2$. The collected results underpinned the development of a responsive fluorescence probe, in an aqueous medium, that has the potential for discovering hydrogen peroxide in the inner condition and outer environment of the human body. Hence a novel fluorescence probe FLA-Boe was devised and synthesized, which exhibited ideal properties for $H_2O_2$ detection. By utilizing this probe, a transparent optical hydrogel sensor with a fluorescence feature for $H_2O_2$ sensing was prepared and synthesized; the result remained effective based on this hydrogel sensor after a series of experiments were conducted. After being exposed to $H_2O_2$, the hydrogels were colored purplish red; the intensity was associated with $H_2O_2$ concentration. Thus, the hydrogel sensor features selective and integrated detection and monitoring of $H_2O_2$ within one system. This effective approach to developing a novel macroscopic hydrogel sensor provides a convenient and intelligent sensing system that is adaptive to complex aqueous environments.

2. Materials and Methods

2.1. Materials and Instrumentation

Fluorescein and 2-Bromomethylphenylboronic acid pinacol ester were purchased from Aladdin Co., Ltd. (Jinan, China). All chemical reagents reached analytical grade or obtained the highest purity availability, and were free of further purification. The preparation in terms of metal ions solutions was conducted in deionized water. $^1$H and $^{13}$C NMR spectra were recorded on a Bruker 400 MHz spectrometer ($^1$H 400 MHz; $^{13}$C 100 MHz, Jinan, China) by using DMSO-$d_6$ as solvents. FT-IR spectra were obtained on a Thermo Fisher Nicolet 6700 FT-IR spectrophotometer (Jinan, China). Q-TOF LC/MS mass spectrometry acted as the recorder for high-resolution mass spectra (HRMS). Shimadzu UV-2600 spectrophotometer was utilized to perform UV–Vis absorption spectra (Jinan, China). Fluorescence spectra were measured on Hitachi F-4600 (Jinan, China) which functions as the gauge of fluorescent intensity.

2.2. Preparation of Samples and Test Solution

Solutions of $10^{-3}$ M of the test object—NaCl, Na$_2$CO$_3$, NaNO$_3$, NaNO$_2$, Na$_2$S NaOCl, glutathione (GSH), Glutamic acid (Glu), Histidine (His), Levodopa (L-DOPA), Lactate (Lac), Fructose (Fru), Glucose (Glc), Urea, Alanine (Ala), Choline oxidase (ChOx), glucose oxidase (GOD)—were prepared in deionized water. The stock solution ($10^{-3}$ M) was prepared
by dissolving FLA-Boe in DMSO. The DMSO/H₂O mixed solution diluted the FLA-Boe solution to acquire the analytical solution (DMSO/H₂O = 1/1, v/v). The pH scope of solutions was adjusted by 4 M of hydrochloric acid aqueous solution and 4 M of sodium hydroxide aqueous solution.

2.3. Synthesis of FLA-Boe

As shown in Scheme 1, Fluorescein (3.3230 g, 1.00 mmol) and hydrazine monohydrate (50%, 3 mL) was dissolved in 100 mL of methanol, after which the mixture was heated to reflux for 4 h. Then the solvent was evaporated, and the resultant residue was subjected to column chromatography (dichloromethane/ethanol = 20/1, v/v) to obtain the intermediate N-(Fluorescein) lactam (2.9611 g, 0.86 mmol). N-(Fluorescein) lactam (3.4634 g, 1.00 mmol) and acetic acid (0.5 mL) were dissolved in 150 mL of acetone and the mixture was heated to 50 °C and stirred for 5 h. The mixture was filtered and the filter cake was washed with acetone and then purified by column chromatography (dichloromethane: ethanol = 30/1, v/v) to obtain the N-(Fluorescein) lactam-N′-methylethylidene (2.7814 g, 0.75 mmol). N-(Fluorescein) lactam-N′-methylethylidene (3.8640 g, 1.00 mmol) was added into a flask containing the mixture of 4-Bromomethylphenylboronic acid pinacol ester (2.3760 g, 0.8 mmol), K₂CO₃ (3.0 g), and 20 mL of DMF at 40 °C for 6 h, and the mixture was poured into H₂O and extracted with ethyl acetate. The organic phase was separated and dried by MgSO₄, meanwhile, the solvent was removed by vacuum distillation. FLA-Boe was obtained as the white solid after being purified by column chromatography with dichloromethane/ethanol (40/1, v/v) as eluent (2.5295 g, yield = 42%). ¹H NMR (400 MHz, DMSO-d₆, ppm): δ 9.86 (d, J = 15.1 Hz, 1H), 7.86–7.79 (m, 1H), 7.70 (d, J = 7.9 Hz, 2H), 7.58–7.52 (m, 2H), 7.46 (d, J = 7.8 Hz, 2H), 7.05 (p, J = 4.4 Hz, 1H), 6.85 (t, J = 3.7 Hz, 1H), 6.69 (dd, J = 8.7, 2.4 Hz, 1H), 6.61–6.57 (m, 1H), 6.57 (s, 1H), 6.53–6.50 (m, 1H), 6.48 (dd, J = 8.0, 5.9 Hz, 1H), 5.17 (s, 2H), 1.85 (s, 3H), 1.76 (s, 3H), 1.29 (s, 12H). ¹³C NMR (101 MHz, DMSO-d₆, ppm): δ 174.00, 160.47, 159.35, 158.81, 152.59, 152.49, 140.60, 135.09, 133.34, 130.07, 129.10, 127.35, 123.26, 112.32, 102.66, 84.17, 69.73, 65.24, 25.41, 25.15, 21.73. FT-IR (KBr, cm⁻¹): ν 2981.31 (O–H), 1696.71 (C=O), 1614.28 (C=N), 1362.12 (C=C), 1176.55 (C–O). HRMS (ESI): C₃₆H₃₅BN₂O₆ calculated for: 602.2588; Found 603.2827 for [M+H]+.

Scheme 1. Synthesis of FLA-Boe.

2.4. Synthesis Process of Hydrogel Sensor

The hydrogel sensors were prepared and synthesized by guest–host interaction as shown in Figure 1. At first, Acrylamide (AAm), β-cyclodextrin (β-CD) and N,N′-methylene bisacrylamide (MBA) (0.3% of monomer content) were mingled together and dissolved in DMSO solution. Then the mixture was stirred under a vacuum at 5 °C until a homogeneous solution was formed. In the next step, 100 μL of TEMED and initiator (APS,
2% of monomer content) were put into the mixture. The mixture was stirred constantly until a homogeneous solution was witnessed during the reaction, which was subsequently poured into a cylindrical mold and maintained at 50 °C for 12 h. In the following step, the hydrogel was extracted and collected from the mold and then rinsed with DMSO for 36 h. AAm-co-β-CD hydrogel was steeped in 10−2 mol/L of FLA-Boe DMSO solution for 10 h before being dialyzed in H2O for 2 h to fabricate the hydrogel sensor in accordance with the host–guest interaction.

**Figure 1.** Synthesis diagram of the hydrogel sensor.

### 3. Results

#### 3.1. Optical Properties of FLA-Boe

The optical properties of FLA-Boe were performed and analyzed with the help of UV–Vis and fluorescence spectroscopy in 1.0 × 10−3 mol/L HEPES buffer (DMSO/H2O = 1/1, v/v, pH = 7.4) through various test objects. As shown in Figure 2a, the absorption band of the solution of FLA-Boe (10−4 mol/L, DMSO/H2O, 1/1, v/v) notably reached 300–700 nm after investigation, and the addition of H2O2 performed an obvious absorption band reaching 417 nm. The result demonstrated that the FLA-Boe had a selective response to H2O2. Nevertheless, no other clear absorption band appeared after adding other test objects, thus the phenomenon proved the prominent selectivity of FLA-Boe toward H2O2.

The fluorescence emission property of FLA-Boe toward different test objects (H2O2, NaCl, Na2CO3, NaNO3, NaNO2, Na2S, NaClO, glutathione (GSH), glutamic acid (Glu), histidine (His), levodopa (L-DOPA), lactate (Lac), fructose (Fru), glucose (Glc), urea, alanine (Ala), choline oxidase (ChOx), and glucose oxidase (GOD)) was analyzed and investigated in DMSO and H2O solution (1/1, v/v), as shown in Figure 2b. From the standard solution of FLA-Boe in DMSO and H2O (1/1, v/v), no distinct fluorescence emission was performed. However, as long as H2O2 was added, significant fluorescence emission was displayed, whose emission level was maximized at 526 nm, which could indirectly affect the π-electron push–pull function and resulted in the intramolecular charge transfer (ICT). Additionally, the color change of the solution from transparent to red clearly appeared (Figure 2c). For experiment comparison, other metal ions were added in the solution, and the fluorescence spectra remained unchanged. Hence the result demonstrated eminent fluorescent selectivity of FLA-Boe towards H2O2.
was distinctly boosted, which could be caused by the intramolecular charge transfer (ICT) process. Phenylboronic ester in FLA-Boe (DMSO/H2O = 1/1, v/v, λex = 417 nm) was excited at 417 nm, the fluorescence intensity evidently increased at 526 nm after the addition of H2O2. FLA-Boe rapidly responded to H2O2 and the fluorescence intensity was distinctly boosted, which could be caused by the intramolecular charge transfer (ICT) process. Phenylboronic ester in FLA-Boe was cleaved through an oxidative reaction when contacting with H2O2, which resulted in the enhancement of fluorescence intensity. As shown in Figure 3a, the fluorescence titration experiments were conducted. FLA-Boe solution was incubated by changing the amounts of H2O2 in pH 7.4 HEPES buffer solution. On the basis of the aforementioned investigation, the emission of FLA-Boe was dominated by the intense excimer emission (DMSO/H2O = 1/1, v/v, λex = 417 nm), (c) the color changes of FLA-Boe in the presence of different species in DMSO/H2O (1/1, v/v) under visible light and UV light at 365 nm.

### 3.2. Sensitivity and Selectivity

As shown in Figure 3a, the fluorescence titration experiments were conducted. FLA-Boe solution was incubated by changing the amounts of H2O2 in pH 7.4 HEPES buffer solution. On the basis of the aforementioned investigation, the emission of FLA-Boe is dominated by the intense excimer emission (DMSO/H2O = 1/1, v/v). When the probe was excited at 417 nm, the fluorescence intensity evidently increased at 526 nm after the addition of H2O2. FLA-Boe rapidly responded to H2O2 and the fluorescence intensity was distinctly boosted, which could be caused by the intramolecular charge transfer (ICT) process. Phenylboronic ester in FLA-Boe was cleaved through an oxidative reaction when contacting with H2O2, which resulted in the enhancement of fluorescence intensity. As shown in Figure 3b, the detection limit (DL) was investigated and obtained from the data, reaching 0.22 µM.

![Figure 2](image_url)

**Figure 2.** (a) UV–Vis spectra of FLA-Boe (10−4 mol•L−1) in DMSO/H2O (1/1, v/v) (λex = 417 nm), (b) fluorescence spectra of FLA-Boe in the presence of different test objects (10−5 mol•L−1) in DMSO/H2O (1/1, v/v) (λex = 417 nm), (c) the color changes of FLA-Boe in the presence of different species in DMSO/H2O (1/1, v/v) under visible light and UV light at 365 nm.

![Figure 3](image_url)

**Figure 3.** (a) Fluorescence spectra of FLA-Boe under different concentrations of H2O2 (DMSO/H2O = 1/1, v/v, λex = 417 nm), (b) calculation of detection limits of FLA-Boe for H2O2 (DMSO/H2O = 1/1, v/v).
3.3. Competition Experiments

To further check the practical applicability of receptor **FLA-Boe** as the selective fluorescent sensor for **H_2O_2**, competitive experiments were carried out with 2.0 equiv of **H_2O_2** in the presence of other test objects in an aqueous solution, as shown in Figure 4. It was interesting to note that only **H_2O_2** rendered significant fluorescence turn-on responses among all the tested objects, whereas all the coexistent test objects had no noteworthy variance with regard to fluorescence intensity. These investigated results clearly demonstrated that the sensor **FLA-Boe** for detecting **H_2O_2** had no interference and could be a good sensor for **H_2O_2** detection in aqueous media.

![Figure 4](image-url)

**Figure 4.** The fluorescence intensity contrast bars in order to investigate the interference effect of other test objects on the detection ability of **FLA-Boe** for **H_2O_2**.

3.4. Detection Mechanism of FLA-Boe

The presupposition of the **FLA-Boe** probe design aims to block the possible ICT process by protecting the hydroxyl group with boronate ester. Hence, Fluorescein was selected as the fluorophore and phenylboronic acid pinacol ester acted as the receptor unit. The inferred detection mechanism of the probe **FLA-Boe** was recommended in Scheme 2. Due to the protection mechanism of the boronate ester on the hydroxyl group, the ICT process of **FLA-Boe** was hindered, which demonstrates no fluorescence emission of **FLA-Boe** was triggered. While **FLA-Boe** was exposed to **H_2O_2**, the decomposition of the boronate ester group was taken and the hydroxyl group was released; then the ICT process of Fluorescein was recovered, through which the strong fluorescence emission from Fluorescein was inevitably triggered. The FT-IR and HRMS tests were investigated to deduce the rational mechanism. In order to verify the response mechanism in detail, the generation of Compound 2—the product of **FLA-Boe** and **H_2O_2** reaction—was monitored by means of HRMS. As shown in Figure 5a, FT-IR spectrum results confirmed the reasonableness of the aforementioned response mechanism hypothesis; after the addition of the **H_2O_2**, a notable hydroxyl peak appeared at around 3250 cm\(^{-1}\). Moreover, the HRMS result proved that Compound 2 was the product of the **FLA-Boe** and **H_2O_2** reaction (Figure 5b). The inference concerning the fluorescence response mechanism of **FLA-Boe** to **H_2O_2** was preliminarily confirmed from these results of the experiments.
3.5. The pH Effect

The influence and phenomenon of the probe **FLA-Boe** to H$_2$O$_2$ were analyzed in different pH conditions. As shown in Figure 6, within the scale of pH 3.0–4.0, the probe failed to respond notably to H$_2$O$_2$. While **FLA-Boe** was exposed to H$_2$O$_2$ within the scope of pH 5.0–8.0, the decomposition of the boronate ester group was taken, then the ICT process of Fluorescein was recovered, through which the strong fluorescence emission from Fluorescein was triggered. As a result, the **FLA-Boe** could respond efficiently to H$_2$O$_2$ in the pH range of 5.0–8.0. The fluorescence intensity reached 581 nm and the response of the probe to H$_2$O$_2$ gradually receded after the pH value was improved to 8.0–9.0. Thus,
the phenomenon illustrated that the probe FLA-Boe had eminent fluorescent responses to H2O2 from weakly acidic to weakly alkaline conditions.

![Figure 6. Fluorescence response of FLA-Boe as a function of pH (3.0–9.0) in the absence and presence of H2O2 (DMSO/H2O = 1/1, v/v, λex = 417 nm).](image)

3.6. The Fabrication of Hydrogel Sensor

The work noted that the formation of the β-CD-FLA-Boe inclusion complex was distinctly performed by 1H NMR; meanwhile, the inclusion procedure was demonstrated preliminarily. The 1H NMR spectrum of β-CD during the appearance and disappearance of FLA-Boe was shown in Figure 7. The integral signal strengths in the 1H NMR spectrum were investigated. The inclusion of β-CD with FLA-Boe exhibited small changes, and a conspicuous influence was shown on the chemical shifts of protons of β-CD-FLA-Boe, which varied notably (Table 1), revealing that these protons primarily contributed to the inclusion with FLA-Boe. More supporting evidence regarding the inclusion of FLA-Boe in the central cavity of β-CD was acquired from 1H NMR studies. In comparisons of the free β-CD and FLA-Boe, the chemical shifts of the inclusion complexes were investigated, demonstrating that FLA-Boe had mutually interacted with β-CD.

![Figure 7. 1H NMR spectra of pure β-CD and β-CD-FLA-Boe.](image)
Table 1. $^1$H NMR chemical shifts (δ, ppm) of pure β-CD and β-CD-FLA-Boe.

| Proton         | 1    | 2    | 3    | 4    | 5    | 6    |
|----------------|------|------|------|------|------|------|
| β-CD           | 5.74 | 5.72 | 5.68 | 4.83 | 4.47 | 3.63 |
| β-CD-FLA-Boe   | 5.76 | 5.74 | 5.70 | 4.84 | 4.49 | 3.65 |

3.7. Detection Performance of the Hydrogel Sensor

The hydrogel sensor responded conspicuously to H$_2$O$_2$ with prominent selectivity and sensitivity due to the role of ICT. To ascertain the capability and effectiveness of the hydrogel sensor, hydrogel detection limits were adjusted and confirmed among varied concentrations of H$_2$O$_2$ solutions, from $10^{-4}$ to $10^{-7}$ mol/L. By observation, the color of the hydrogel sensor could still be notably changed when H$_2$O$_2$ concentration reached $10^{-6}$ mol/L. The color of the hydrogel sensor changed when H$_2$O$_2$ concentration was $10^{-4}$, $10^{-5}$, and $10^{-6}$ mol/L for 50 s, 4 min, and 20 min, respectively. In Figure 8, the hydrogel sensor performed visible red light in the aqueous solution environment, which was the same as FLA-Boe's; under 365 nm UV light, the color of the hydrogel sensor in the aqueous solution also conspicuously changed.

![Visible light](image1)

![UV light at 365 nm](image2)

Figure 8. The color of the hydrogel sensor in the presence of different concentrations of H$_2$O$_2$ under visible light and UV light at 365 nm.

A comparison of the results indicates that the Fluorescein derivative immobilized system in this work exhibits better ability for H$_2$O$_2$ detection compared to the other previously reported systems (Table 2). In addition, many of the reported sensors involve poorer reuse performance. On the other hand, the present sensing system is cost-effective, selective, exhibits simple naked-eye sensing, and is suggested to be utilized to detect H$_2$O$_2$ in a real water environment.

Table 2. Comparison table for hydrogel sensor systems.

| No. | Sensing Probe         | Detection Limit | Fabrication Method                                      | Ref   |
|-----|-----------------------|-----------------|---------------------------------------------------------|-------|
| 1   | Thiazolidinone        | 40 mM           | Self-assembly                                           | [23]  |
| 2   | Phenothiazine         | 0.1 mM          | Covalent coupling                                       | [24]  |
| 3   | Pyrene                | 0.18 M          | Reversible addition-fragmentation chain-transfer        | [25]  |
| 4   | 2-Thiohydantoin moieties | 25 μM         | Radical polymerization                                  | [26]  |
| 5   | Fluorescein           | 0.22 μM         | Host–guest interaction                                  | This work |
3.8. Universality of the Hydrogel Sensor

Inspired by the remarkable function of the hydrogel sensor in solution for H$_2$O$_2$, the response of the hydrogel sensor for H$_2$O$_2$ in flowing and static water environments was further investigated. The hydrogel sensor was colored from transparent to light aubergine (about 6 min) after the sensor was exposed to the flowing H$_2$O$_2$ solution ($10^{-4}$ mol/L). The color changed to sharp aubergine after 10 min. In a large container insulated from the atmosphere, the hydrogel sensor was exposed to a static H$_2$O$_2$ solution. An interesting fact was that when the gel was steeped in $10^{-7}$ mol/L H$_2$O$_2$ solution for 3 h, the hydrogel sensor changed color. During the experiment for detection performance, the hydrogel sensor had no response when contacting a small quantity of H$_2$O$_2$ solution whose concentration was no more than $10^{-7}$ mol/L. In accordance with the experimental phenomenon, it was summarized that a cumulative effect may appear in the hydrogel sensor that could result in the reduction of the detection limit to some extent. Thus, analyzing the sensing performance and process of the hydrogel sensor can open a new avenue for developing significant, long continuous, and fast-responding test strips for precisely detecting H$_2$O$_2$ by convenient naked-eye observation, which is of particular importance in the immediate monitoring of H$_2$O$_2$ in both natural aqueous environments and liquid environments around chemical industrial regions. Meanwhile, observing color changes by the naked eye simplifies and speeds up the detection of H$_2$O$_2$. Therefore, this technological proposal will be a reliable method and a suitable selection for the detection of H$_2$O$_2$ in aqueous environments.

4. Conclusions

In summary, based on the H$_2$O$_2$ probe FLA-Boe, we developed an easily operated and commonly applicable strategy to synthesize a H$_2$O$_2$ responsive hydrogel sensor with on–off switching and color-changing fluorescence features. Hydrogel sensors were utilized smoothly for the determination of trace amounts of H$_2$O$_2$ in water samples, which indicates that hydrogel sensors are available for use by common people and scientists depending on practical requirements. Additionally, this method has potential applications in establishing novel monitoring and detection mechanisms and systems that cover detection systems for flowing and static water environments, etc. It is expected that this method may provide diverse and novel strategies to exploit smarter systems that include functions, such as synergistic visual detection and efficient sensing.

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