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Carbon dot based nucleus targeted fluorescence imaging and detection of nuclear hydrogen peroxide in living cells

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Investigation of the intracellular generation of H₂O₂, one of the most important reactive oxygen species (ROS), is crucial for preventing various diseases since it is closely linked with different physiological and complex cell signaling pathways. Despite the development of various fluorescent probes, the majority of the fluorescent probes cannot move across the nuclear membrane. However, detection of the nuclear level of H₂O₂ is very important since it can directly cause oxidative DNA damage which ultimately leads to various diseases. Therefore, in this study, *p*-phenylenediamine based carbon quantum dots (B-PPD CDs) have been synthesized and integrated with 4-formylbenzeneboronic acid as a doping agent for the detection of H₂O₂. The detection mechanism showed that, upon exposure to H₂O₂, the fluorescence of the B-PPD CDs was immediately quenched. Further investigation has been done in the *in vitro* RAW 264.7 cell line by both exogenous and endogenous exposure of H₂O₂ to demonstrate the feasibility of the method. It is shown successfully that the exogenous presence and endogenous generation of H₂O₂ in RAW 264.7 cells can be detected using B-PPD CDs. The limit of detection (LOD) was determined to be 0.242 μM. The development of such imaging probes using carbon quantum dots will lead to live-cell imaging as well as ROS detection.

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1 Introduction

The term reactive oxygen species (ROS) is used for the partially reduced intermediate products of oxygen that are generated during the mitochondrial oxidative metabolism in a living organism.^{1–3} Among reactive oxygen species (ROS), H₂O₂ is considered as one of the major ROS that participate in cellular signal transduction, differentiation, cellular proliferation, immune systems, and various physiological processes.^{4–6} The major role of H₂O₂ in the body is that it acts as an oxidative stress marker and also plays an important role in the protection against invading pathogens.⁷ However the formation of high concentrations of H₂O₂ leads to the oxidation of various biomolecules such as DNA, enzymes, many proteins, and fatty acids which results in the disruption of a balanced cell redox mechanism and ultimately causes an increased risk of many diseases including cancer, stroke, Alzheimer's disease, *etc.*^{8–11} Unlike superoxide free radicals which are extremely short-lived since they undergo rapid reduction into hydrogen peroxide *via* an enzymatic reaction of superoxide dismutase (SOD),¹² hydrogen peroxide exhibits a long life span.¹³ Due to its higher

stability, it is considered to be a well-established cellular oxidative stress biomarker.^{14,15} Therefore, accurate detection of hydrogen peroxide expedites proper diagnosis and prevention of various relative diseases.

There are various techniques for the measurement of ROS such as high-performance liquid chromatography,^{16,17} fluorescence,¹⁸ colorimetry,¹⁹ and electrochemistry,^{20,21} which have already been reported. Apart from such technologies, researchers are engaged in enzyme-based detection of H₂O₂.^{22,23} Drawbacks of such approaches involved specialized equipment and tiresome analysis as well as poor reproducibility and instability.^{24,25} Among various analytical approaches, fluorescence-based probe technology exhibits tremendous advantages in the detection of highly reactive oxygen species (hROS) because of its unique features such as excellent biocompatibility, rapid response, and high temporal and spatial resolution.^{26–29} Despite using several fluorescent probes, very few are practically applicable because of their photophysical limitations, *i.e.*, relatively short excitation/emission wavelengths which limit their application in deep tissue penetration with a minimized background.³⁰ There are some limitations of short emission wavelength probes. Some intrinsic biomolecules (such as NADH, folic acid, *etc.*), hemoglobin (Hb) and oxyhemoglobin (HbO₂) can absorb a two-photon fluorescent probe at <550 nm emission which increases the background fluorescence by decreasing the collection efficiency in deep tissue

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imaging,³¹ which causes background fluorescence in the blue to green region.³² Therefore, preparation of such probes with emission in the red-emissive region (>600 nm) is advantageous since it can avoid cellular absorption of biomolecules and facilitates deeper into tissues (higher collection efficiency) by minimizing the background fluorescence noise.³³

Carbon quantum dots (CDs) are an emerging nanomaterial and have been used as an alternative to conventional organic dyes in various applications including bioimaging, sensing, *etc.* due to their good biocompatibility,³⁴ high photostability,³⁵ high pH stability, *etc.*^{36–38}

Moreover, the application of most of the reported CDs is usually restricted because their emission usually concentrates on the blue light region.^{39–41} Red emission CDs can be prepared by the precursor PPD (*p*-phenylenediamine) which shows an emission longer than 600 nm.^{42–44} Recently Wang *et al.* reported different PPD CDs where emissions based on solvent with QY for red emission approached 34%.⁴⁵ Despite knowing this fact, the solvent-based tunable properties of CDs and red emission of PPD based CDs are still unclear.

The detection of ROS especially hydrogen peroxide has been explored by many researchers using various CDs *via* modification of their surface. Zheng *et al.*⁴⁶ successfully achieved the colorimetric detection of H₂O₂ and glucose by CQDs *via* utilizing the peroxidase-like catalytic activity. Li *et al.*⁴⁷ functionalized CQDs with boronic acid substituted bipyridinium salt to detect glucose *via* fluorescence quenching by the salt resulting in the recovery of glucose. Similarly, Qu *et al.*⁴⁸ synthesized phenylboronic acid-functionalized CQDs as a glucose-sensing system.

The nucleus of eukaryotic cells plays a crucial role in various genomic regulatory processes including the generation of reactive oxygen species such as H₂O₂, HOCl, *etc.*⁴⁹ Moreover, abnormal generation of ROS in nuclei triggers oxidative stress which ultimately leads to oxidative DNA damage, consequently linked with various disorders such as cancer, aging, Alzheimer's disease, and so on.^{8–11} Therefore, monitoring of nuclear ROS flux becomes very important in the diagnosis of such diseases. To address the issue, fluorescence tracking is a very influential tool that can detect several ROS species such as H₂O₂. Various types of fluorescence probe have been synthesized that are recognized for their high sensitivity and specific selectivity but most of them are capable of gathering only in the cytoplasm and cannot penetrate the nuclear envelope^{46–48} and therefore the limitation of diffusion from the cytoplasm to nucleoplasm sometimes prevents the status of the nuclear ROS level to be checked. However, at present, a very small number of nucleus-targeted probes have been developed.⁵⁰

In the current work, we synthesized a nucleus targeted fluorescent CD-based sensing system for hydrogen peroxide detection by taking *p*-phenylenediamine (PPD) as a precursor doped with 4-formylbenzeneboronic acid *via* the microwave-assisted method. This one-pot solvothermal boronic acid-based PPD CD preparation has been reported for the first time for hydrogen peroxide detection without requiring any further functionalization. Moreover, among various biological ROS and other biomolecules, B-PPD CDs display high specificity

and sensitivity to H₂O₂. Furthermore, B-PPD CDs have been exposed to murine macrophage cell line RAW 264.7 where the bright red fluorescence intensity of B-PPD CDs immediately decreases to a significant level after exposure to hydrogen peroxide. Such quenching effect is shown in both endogenous and exogenous H₂O₂ spikes in living RAW 264.7 cells without showing any marked cytotoxicity (Scheme 1).

2 Experimental

2.1 Reagents and materials

p-Phenylenediamine, 4-formylbenzeneboronic acid, and H₂O₂ were purchased from Sigma-Aldrich. The RAW 264.7 macrophage cell line was purchased from NCCS Pune. Lipopolysaccharide (LPS) was purchased from Sigma-Aldrich. ProLong Gold Antifade Mountant with DAPI was obtained from Thermo Fisher Scientific, and Fluoromount aqueous mounting medium and paraformaldehyde were from Merck. The other reagents for the experiments were of analytical grade and utilized as received. Ultrapure water was produced from a Millipore-Q water purification system.

2.2 Instrumentation

The morphology of the carbon dots was examined using a TEM JEOL 2100 plus instrument. Samples were prepared by dropping ethanolic suspensions onto amorphous carbon film-coated Cu TEM grids and dried in an oven. The UV-vis spectrum was recorded using a Shimadzu UV 2600 UV-vis spectrophotometer. Fourier transform infrared spectroscopy (FTIR) analysis was carried out on a PerkinElmer instrument. Particle size and zeta potential measurements were performed on a Malvern Zetasizer Nano series Nano-ZS90. Fluorescence spectra were examined using a Jasco spectrofluorometer, FP-8300. Cell imaging was obtained using a Leica confocal laser scanning microscope (CLSM).

2.3 Synthesis and characterization of B-doped *p*-phenylenediamine based carbon quantum dots (B-PPD CDs)

p-Phenylenediamine (0.25 mM) was mixed with 4-formylbenzeneboronic acid (2.7 mg) in 50 ml of ethanol and sonicated for 15 minutes. The fine mixture was introduced into a microwave reactor at 140 °C for 15 minutes. After that, the solution was kept at room temperature for cooling down and centrifuged at 10 000 rpm for 15 minutes for exclusion of larger particles. Finally, the B-doped PPD CD solution was filtered through a 0.22 μm sterile syringe filter.

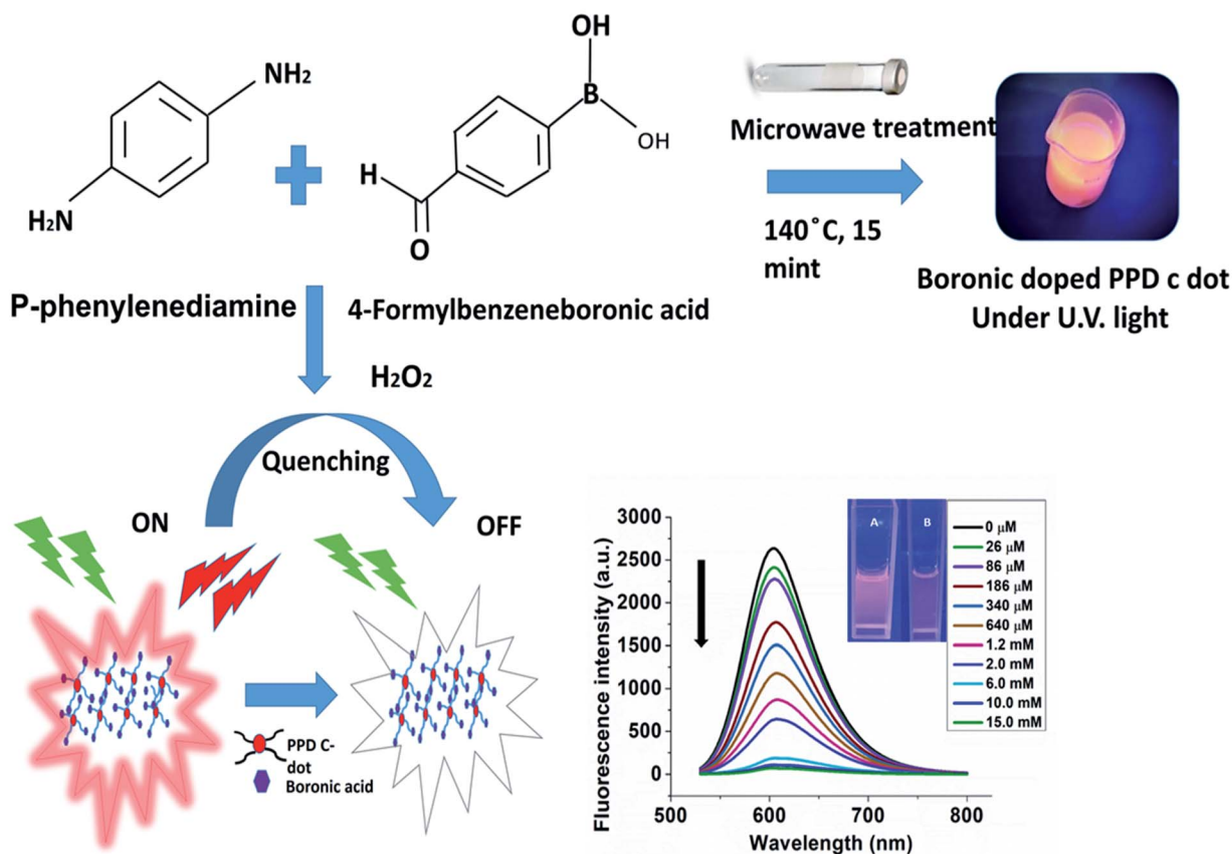
2.4 Quantum yield measurement

The quantum yield calculation was done using the following equation:

$$Q = Q_R \times (I_{IR} \times A_R/A \times \eta^2/\eta_R^2)$$

where Q denotes the quantum yield, I is the integrated photoluminescence emission intensity, A is the absorbance, and η is the refractive index. The subscript R indicates the reference or standard sample with a known quantum yield. The quantum





Scheme 1 Schematic representation of the synthesis of B-PPD CDs and a fluorescent switch on–off sensor for H_2O_2 detection.

yield measurements were performed using rhodamine B as the reference sample ($\Phi_R = 0.31$).

2.5 Fluorescence quenching of B-doped *p*-phenylenediamine based carbon quantum dots (B-PPD CDs) by H_2O_2

Different concentrations of H_2O_2 were added to the B-PPD CD solution by continuous stirring and the corresponding fluorescence spectrum was recorded.

2.6 Calculation of the limit of detection (LOD)

The fluorescence emission spectrum of B-PPD CDs was measured 25 times ($n = 25$) and the standard deviation of the blank measurement was evaluated. The fluorescence intensity was measured at 606 nm plotted against the concentration of H_2O_2 . The detection limit was calculated based on the formula $3\sigma/m$, where σ is the standard deviation of the blank measurement, m is the slope between the fluorescence intensity and H_2O_2 concentration.

2.7 Effect of pH

For efficient detection of H_2O_2 at physiological pH, the fluorescence intensity of B-PPD CDs was measured at different pH ranging from 4 to 9 to explore its fluorescence response at

different pH. In this study, the PL intensity was checked at an excitation wavelength of 520 nm with various H_2O_2 concentrations.

2.8 Selectivity and interference studies of interfering agents

Selectivity studies of B-PPD CDs ($20 \mu\text{g ml}^{-1}$) towards H_2O_2 were performed, and testing species including reactive oxygen species (ROS) such as H_2O_2 , *tert*-butoxy radical ($\cdot\text{OtBu}$), hypochlorite (OCl^-), *tert*butyl hydroperoxide (TBHP), hydroxyl radical ($\cdot\text{OH}$), singlet oxygen ($^1\text{O}_2$), and superoxide ($\text{O}_2^{\cdot-}$) ($200 \mu\text{M}$ for each species), amino acids (Met, Leu, and Tryp), biomolecules (ascorbic acid and glucose), other ions (Fe^{2+} , Ca^{2+} , Mg^{2+} , Cu^{2+} , Cd^{2+} , Na^+ , and K^+) and several proteolytic enzymes such as pronase, proteinase k, proteinase 3, trypsin, and cathepsin G were investigated at different concentrations.

2.9 Cell culture

The RAW 264.7 (macrophage) murine cell line was procured from the National Centre for Cell Sciences (NCCS), Pune, India. Cells were cultured and maintained in Dulbecco's Modified Eagle's Medium (DMEM) (4.5 g L^{-1}) with 10% heat-inactivated fetal bovine serum (FBS), 1% antibiotic-antimycotic, cultured in a T-75 cm^2 flask, and kept at 37°C in an incubator with 5% atmospheric CO_2 and observed daily using an inverted microscope, Leica DMi1.



2.9.1 Cell viability assay (MTT assay). RAW 264.7 cells were seeded in 96-well plates (1×10^5 per well) with a growth medium of 200 μl and incubated for 24 hours. After incubation for 24 hours, B-PPD CDs were added with varying concentrations (10, 20, 30, 40, and 50 $\mu\text{g ml}^{-1}$). The experiment was performed in triplicate. After 12 hours of drug treatment, cells were washed with phosphate buffered saline (PBS) and the old medium was replaced with fresh medium and 20 μl of 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide MTT was added to check cell viability. Cells were incubated for 4 hours followed by the addition of 100 μl DMSO, and left for 2 hours in the dark. The absorbance was measured at 570 nm using a microplate reader. Non-treated cells were considered as the control. The cell viability percentage was calculated using the following formula:

$$\text{Cell viability (\%)} = (\text{Abs sample} - \text{Abs blank} / \text{Abs control} - \text{Abs blank}) \times 100$$

2.9.2 Exogenous and endogenous detection of H_2O_2 and live-cell imaging experiments. After the cell viability test, RAW 264.7 cells were seeded in 6 well plates (at a density of 1×10^5 per well) over the sterile coverslips and allowed to adhere for 24 hours. The old cell culture medium was discarded and washed with sterile PBS. For exogenous detection of H_2O_2 , cells were pretreated with different concentrations of H_2O_2 (125 μM , 250 μM , and 500 μM) separately mixed with cell culture medium for 30 minutes. Similarly for endogenous detection, cells were incubated with LPS of varying concentrations (25 ng ml^{-1} , 50 ng ml^{-1} and 100 ng ml^{-1}) diluted with cell culture medium for 4 hours and kept at 37 $^\circ\text{C}$ in a 5% CO_2 incubator. After washing two times with PBS buffer, cells were incubated with B-PPD CDs (20 $\mu\text{g ml}^{-1}$) for another 30 min at 37 $^\circ\text{C}$. For the control experiment, RAW 264.7 cells were incubated only with B-PPD CDs (20 $\mu\text{g ml}^{-1}$) for 30 min at 37 $^\circ\text{C}$. After that cells were washed three times with PBS buffer before cell imaging experiments were performed.

3 Results and discussion

3.1 Characterization of B-doped *p*-phenylenediamine based carbon quantum dots (B-PPD CDs)

The 4-formylbenzeneboronic acid doped PPD based carbon quantum dots were characterized by UV-visible absorption spectroscopy, fluorescence spectroscopy, Fourier transform infrared spectroscopy (FTIR), dynamic light scattering (DLS) technique, zeta potential analysis, and transmission electron microscopy (TEM). The quantum yield (QY) of B-PPD CDs was found to be 5.5% with the reference rhodamine B (31%).

The optical properties of B-PPD CDs were analyzed by UV-vis absorption and fluorescence spectrum measurements. The UV-vis absorption spectra of the B-PPD CDs and undoped PPD CDs are shown in Fig. 1(A). B-PPD CDs showed two characteristic peaks at 266 and 285 nm due to the $n-\sigma^*$ transition. In contrast, the undoped PPD CDs showed peaks at 224 nm and 285 nm due to $\sigma-\sigma^*$ and $n-\sigma^*$ transitions, respectively. The photoluminescence (PL) properties of B-PPD CDs were studied by

monitoring the emission of excitation wavelengths from 400 nm to 600 nm. The stacked emission observed for B-PPD CDs for different excitations is shown in Fig. 1(B). It is evident from the stacked emission spectra that the position of the emission peak did not show a marked obvious shift, indicating that B-PPD CDs exhibited excitation-independent photoluminescence properties. However, B-PPD CDs exhibit their maximum main emission peak at 606 nm when the excitation wavelength is 520 nm. It is to be noted that under a UV lamp B-PPD CDs show red fluorescence. The inset shows a photograph of B-PPD CDs under a UV lamp. Since the emission range is above 600 nm, B-PPD CDs can be targeted for live-cell imaging without any further damage to the cell by avoiding the quenching noise from other biomolecules. So, detection using B-PPD CDs through the imaging technique is demonstrated later on.

An FTIR study was carried out on the synthesized PPD CDs and B-PPD CDs. The stacked FTIR spectra of PPD CDs and B-PPD CDs are shown in Fig. 1(C). PPD CDs show peaks at 3346 cm^{-1} due to $\text{N}-\text{H}_{\text{str}}$ and 2917 cm^{-1} due to $\text{C}-\text{H}_{\text{str}}$. The strong peak at 1061 cm^{-1} is due to $\text{C}-\text{H}_{\text{def}}$ out of plane for an aromatic ring. In addition, the characteristic peak at 1630 cm^{-1} is due to $\text{C}=\text{C}_{\text{str}}$. On the other hand, for B-PPD CDs the peak due to $\text{N}-\text{H}_{\text{str}}$ is shifted to 3408 cm^{-1} as a result of hydrogen bonding between $-\text{NH}_2$ and B. The $\text{C}-\text{H}_{\text{str}}$ peak is also shifted to 2924 cm^{-1} . All other characteristic peaks of PPD CDs, viz. those at 1381 (well resolved) and 1630 cm^{-1} due to $\text{C}-\text{N}_{\text{str}}$ and $\text{C}-\text{H}_{\text{def}}$ out of plane, respectively, are present as well in B-PPD CDs. The size of the B-PPD CDs was measured by DLS and found to be ~ 1.73 nm. The zeta potential of B-PPD CDs was evaluated and found to be +3.43 mV. The positive surface charge on B-PPD CDs is due to protonation of the $-\text{NH}_2$ group under slightly acidic conditions (Fig. 2(A) and (B)). A representative TEM image of B-PPD CDs is shown in Fig. 2(C). It is evident from the TEM image that the synthesized B-PPD CDs are of sizes below 10 nm. In addition, the high-resolution transmission electron microscopy (HRTEM) image of one quantum dot particle in the inset clearly shows the crystal plane of carbon quantum dots, hence demonstrating the crystalline nature of B-PPD CDs.

3.2 Fluorescence quenching of B-doped *p*-phenylenediamine based carbon quantum dots (B-PPD CDs) by H_2O_2

The fluorescence response or sensitivity of B-PPD CDs towards H_2O_2 was investigated by employing the titration method with H_2O_2 . The change in fluorescence intensity of B-PPD CDs was observed upon the addition of different concentrations of H_2O_2 . Fig. 3(A) shows the stacked emission spectra of B-PPD CDs (excitation at 520 nm) where the addition of H_2O_2 leads to the gradual decrease in fluorescence intensity (at 606 nm) with the increase in the concentration of H_2O_2 (3–15 mM). Furthermore, Fig. 3(B) illustrates that there is a sequential increase of fluorescence quenching efficiency of B-PPD CDs corresponding to the concentrations of H_2O_2 in the range of 3–460 μM , and upon evaluating the H_2O_2 concentration in the range of 3–110 μM , the quenching efficiency showed a linear response ($R^2 = 0.94419$)



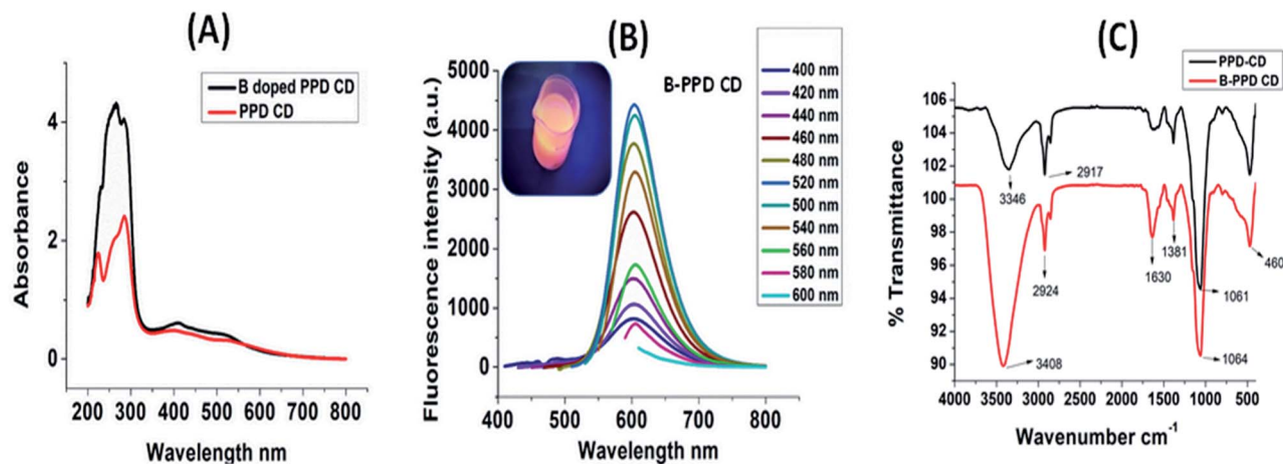


Fig. 1 (A) The UV-visible absorption spectra of B-PPD CDs and PPD CDs excited at 520 nm wavelength. (B) Fluorescence spectra of B-PPD CDs excited in the wavelength range from 400 nm to 600 nm. Inset: photograph of B-PPD CDs under a UV-visible lamp. (C) FTIR spectra of PPD CDs and B-PPD CDs.

with the fluorescence intensities of B-PPD CDs at 606 nm. The limit of detection (LOD) was calculated using the formula $LOD = 3\sigma/m$ and found to be $0.242 \mu\text{M}$. The LOD has been verified experimentally (not shown). Therefore, the minimum amount ($0.242 \mu\text{M}$) of H_2O_2 is required to trigger fluorescence quenching in the system. Thus, the result demonstrates that B-PPD CDs have high sensitivity towards H_2O_2 detection in PBS (pH 7.4).

3.3 Effect of pH

Furthermore, the applicability of B-PPD CDs was investigated by analyzing the effect of pH on the fluorescence properties of B-PPD CDs upon addition of H_2O_2 . As evident from Fig. 4(A), the decrease in the fluorescence intensity at pH 5.0 and pH 6.0 is less after the addition of H_2O_2 . At pH 4 (acidic) and pH 9

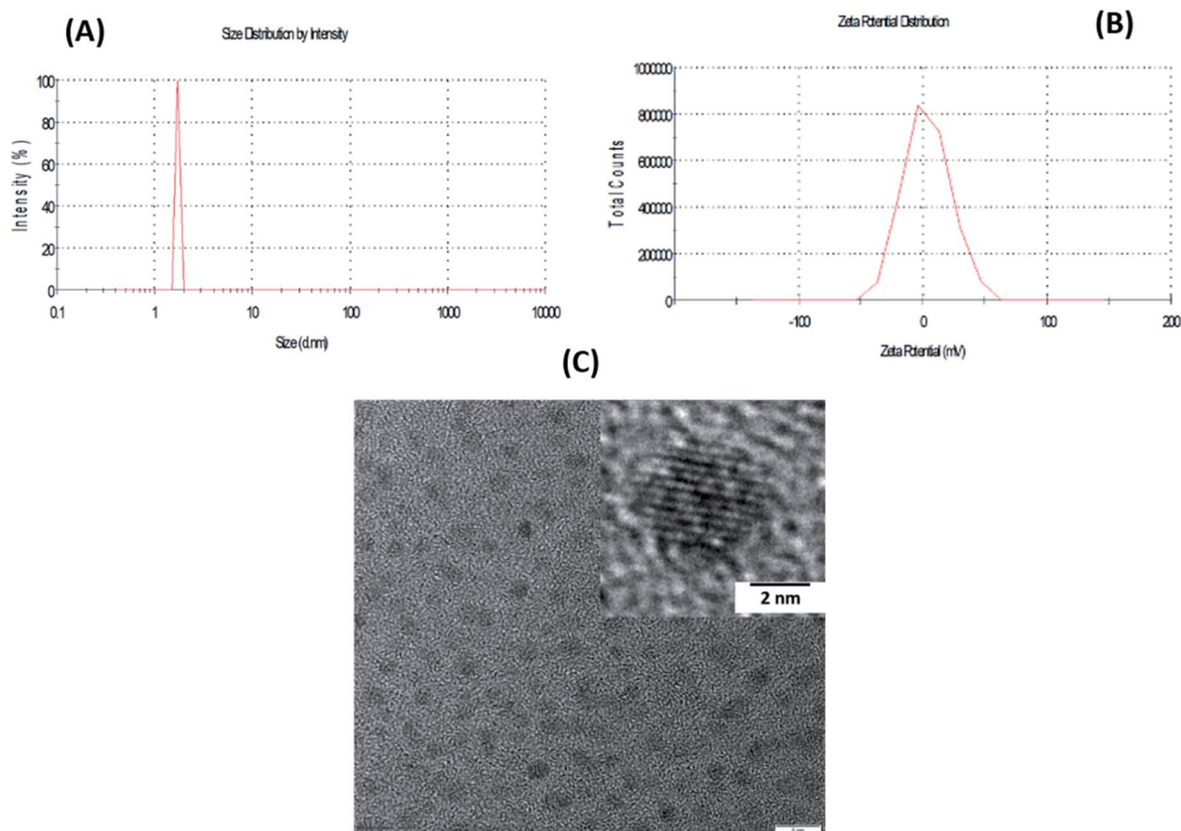


Fig. 2 (A) Particle size distribution and (B) zeta potential measured via dynamic light scattering (DLS) measurement of B-doped PPD CDs. (C) Representative TEM image of B-PPD CDs. Inset: high-resolution TEM (HRTEM) image of B-PPD CDs magnifying one quantum dot.



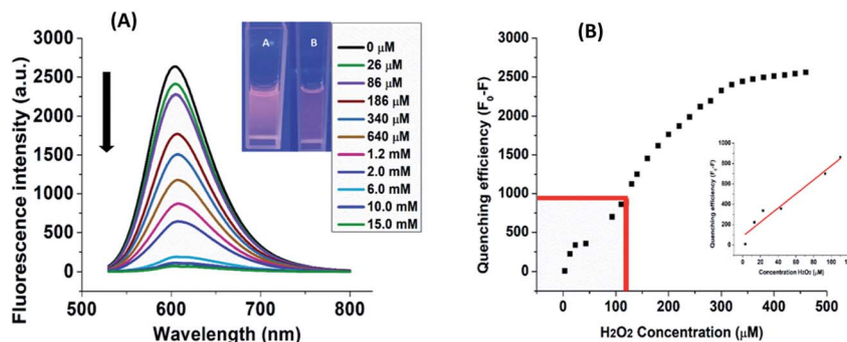


Fig. 3 (A) Quantitative analysis of fluorescence emission spectra of B-PPD CDs ($20.0 \mu\text{g ml}^{-1}$) for incubation with different concentrations of H_2O_2 at $\lambda_{\text{ex}} = 520 \text{ nm}$. (B) Fluorescence quenching efficiency ($F_0 - F$) of B-PPD CDs ($20.0 \mu\text{g ml}^{-1}$) versus various concentrations of H_2O_2 , where F_0 and F denote the fluorescence intensity without and with H_2O_2 , respectively. Inset: the linear response of quenching efficiency of B-PPD CDs against 3–110 μM H_2O_2 concentration.

(basic) the change of fluorescence intensity is more pronounced. At pH 7 and pH 8 the change of fluorescence intensity is more or less the same.

This indicates that fluorescence intensity of B-PPD CDs in the presence of H_2O_2 is more pronounced at pH 4 and at pH 9. However, fluorescent B-PPD CDs can be used in the pH range 4 to 9 for detection of H_2O_2 .

3.4 Selectivity and competitive studies of interfering agents

It is extremely necessary to demonstrate the selectivity of the PL response of B-PPD CDs towards other interfering agents. Subsequently, the B-PPD CDs ($20.0 \mu\text{g ml}^{-1}$) were exposed to

various relevant species ranging from biomolecules (glucose and ascorbic acid), amino acids (Met, Trp, and Leu), different ions (Fe^{2+} , Ca^{2+} , Mg^{2+} , Cu^{2+} , Cd^{2+} , Na^+ , and K^+), various reactive oxygen species (ROS) (*tert*-butoxy radical ($\cdot\text{OtBu}$), hypochlorite (OCl^-), *tert*butyl hydroperoxide (TBHP), hydroxyl radical ($\cdot\text{OH}$), hydrogen peroxide (H_2O_2), singlet oxygen ($^1\text{O}_2$), and superoxide ($\text{O}_2^{\cdot-}$) and various proteolytic enzymes such as pronase, proteinase k, proteinase 3, trypsin and cathepsin G to evaluate their selectivity or fluorescence response towards the above-mentioned species.^{51–53} The change of fluorescence intensity ratio in the presence of interfering species is shown in Fig. 4(B); it has been found that the B-PPD CDs demonstrated negligible

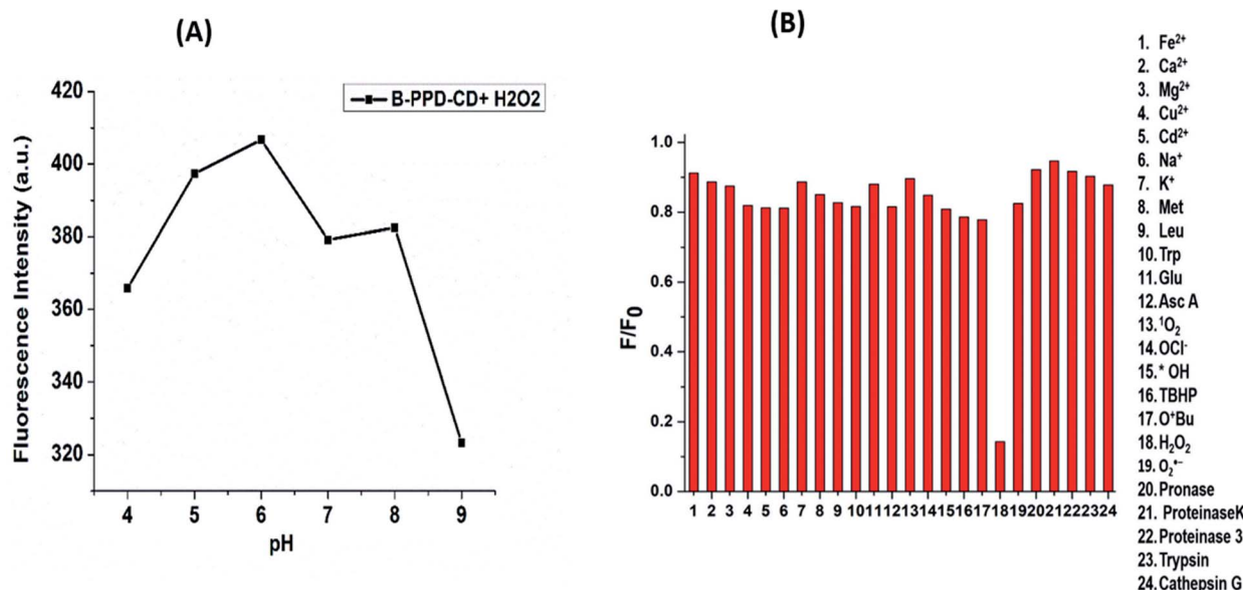


Fig. 4 (A) Effect of pH on the fluorescence intensity of B-PPD CDs ($20.0 \mu\text{g ml}^{-1}$) at an excitation wavelength of 520 nm in the presence of H_2O_2 . (B) Selectivity and competitive studies of B-PPD CDs investigating the fluorescence intensity ratio (F/F_0) of B-PPD CDs, where F and F_0 are the fluorescence intensity of B-PPD CDs ($20.0 \mu\text{g ml}^{-1}$) in the presence and absence of various relevant interfering species, each at 200 μM , including biomolecules (glucose and ascorbic acid), amino acids (Met, Trp, and Leu), different ions (Fe^{2+} , Ca^{2+} , Mg^{2+} , Cu^{2+} , Cd^{2+} , Na^+ , and K^+), various reactive oxygen species (ROS) (*tert*-butoxy radical ($\cdot\text{OtBu}$), hypochlorite (OCl^-), *tert*butyl hydroperoxide (TBHP), hydroxyl radical ($\cdot\text{OH}$), hydrogen peroxide (H_2O_2), singlet oxygen ($^1\text{O}_2$), and superoxide ($\text{O}_2^{\cdot-}$) and various proteolytic enzymes such as pronase, proteinase k, proteinase 3, trypsin and cathepsin G.



fluorescence changes upon the addition of the above-mentioned interfering agents. On the other hand, incorporation of H_2O_2 triggered a significant decrease in fluorescence emission of B-PPD CDs, from which it could be concluded that B-PPD CDs were successfully achieved to detect H_2O_2 with high selectivity without any effects of interfering agents.

3.4.1 Cell viability assay. Depending upon the observations, based on the photoluminescence spectra of B-PPD CDs towards the selective and quick sensing of H_2O_2 , we further investigated the detection of H_2O_2 by B-PPD CDs *via* both exogenous and endogenous induction in RAW 264.7 cells. At first, we proceeded to check the effect of B-PPD CDs on cell viability. Cell viability was analyzed by performing an MTT assay in RAW 264.7 cells. RAW 264.7 cells were treated with different concentrations of B-PPD CDs ($10 \mu\text{g ml}^{-1}$, $20 \mu\text{g ml}^{-1}$, $30 \mu\text{g ml}^{-1}$, $40 \mu\text{g ml}^{-1}$, and $50 \mu\text{g ml}^{-1}$) as shown in Fig. 5 where they demonstrated that the cells exhibited 94% cell viability in $10 \mu\text{g ml}^{-1}$ and viability of the cells is maintained up to 85% even at $40 \mu\text{g ml}^{-1}$ concentrations; however, the viability is decreased to 75% at $50 \mu\text{g ml}^{-1}$ concentration. Nevertheless, the RAW 264.7 cells exhibited good viability (above 80%) even up to $40 \mu\text{g ml}^{-1}$ concentration after the 12 h incubation period. The MTT result showed that B-PPD CDs can be utilized for targeted H_2O_2 levels in nuclear imaging.

3.5 Nuclear colocalization of B-PPD CDs with the commercially available nuclear dye

To examine the nuclear localization of B-PPD CDs, RAW 264.7 cells were co-incubated with B-PPD CDs for 30 minutes and a commercial prominent nuclear staining dye, DAPI, for fluorescence co-localization imaging. Fig. 6 shows that the co-stained RAW 264.7 cells exhibited blue fluorescence with DAPI and emitted red fluorescence with B-PPD CDs when excited at 405 nm and 514 nm, respectively. Furthermore, the merged images proved that incubation with B-PPD CDs for 30 minutes leads to strong fluorescence due to successful penetration of the nucleus and staining of the nucleus specifically,

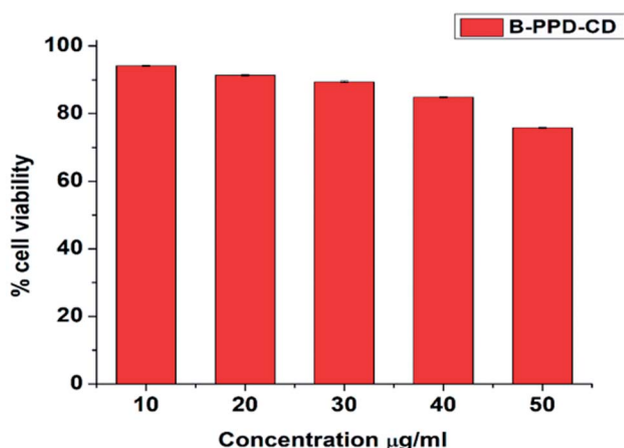


Fig. 5 Cell viability percentage in RAW 264.7 cells after 12 h incubation of B-PPD CDs at different concentrations from 10 to $50 \mu\text{g ml}^{-1}$.

which indicated an excellent ability for targeted nucleus staining without showing any unspecific binding to the cytoplasm. The result hence suggested that by maintaining good biocompatibility and low cost, B-PPD CDs prepared by simple one-pot synthesis possessed excellent nuclear staining properties comparable to those of commercial nuclear stain DAPI.

3.6 Exogenous and endogenous detection of the nuclear H_2O_2 level in the RAW 264.7 cell

To confirm the sensing capacity of B-PPD CDs towards H_2O_2 in living cells, RAW 264.7 cells have been exposed to different concentrations ($125 \mu\text{M}$, $250 \mu\text{M}$, and $500 \mu\text{M}$) of exogenous H_2O_2 by separate incubation while maintaining $20 \mu\text{g ml}^{-1}$ concentration of B-PPD CDs (Fig. 7) and the result clearly showed that there is a concentration-wise gradual decrease in fluorescence intensity as compared to the control where the cells have been incubated with the B-PPD CDs alone ($20 \mu\text{g ml}^{-1}$). Furthermore, inspired by the above results, we tried to perform endogenous generation of H_2O_2 by RAW 264.7 cells.

Since a previous report has suggested that LPS upregulated the production of H_2O_2 ,⁵⁴ cells have been allowed to incubate with LPS of varying (25 ng ml^{-1} , 50 ng ml^{-1} and 100 ng ml^{-1}) concentrations and the fluorescence becomes quenched gradually with an increase in concentration in the LPS treated cells when compared with the control group (B-PPD CDs alone). The outcome of the study (Fig. 8) showed that B-PPD CDs not only crossed the cell membrane but could also invade and stain specifically the nucleus, thereby sensing the H_2O_2 level inside the nucleus by quenching the fluorescence of B-PPD CDs. There are some earlier reports which investigated the activity and levels of H_2O_2 in the nucleus that cause severe oxidative damage to DNA.^{8–11} Therefore, the result suggests that superior biocompatibility, targeted nucleus staining, and specific response to H_2O_2 makes B-PPD CDs a robust system for the detection of H_2O_2 in living cells.

3.7 Mechanistic insight

In general, there is a lot of debate about the exact origin of PL in CDs. One PL model has been proposed for CDs based on the bandgap transitions in conjugated π -domains. Another model proposes that it could be related to surface defects on the CDs.⁵⁵ The PL observed in PPD is believed to be due to surface states. The PL is due to the presence of different emissive traps due to various functionalities, excitons of carbon, *etc.* as well as due to electron-hole recombination pathways. Boron is known as a notable electron-deficient compound and it takes part in charge transfer especially in an excited state. When doped with the boron compound (4-formylbenzeneboronic acid), the higher energy electron of CDs can move to the electron-deficient part of the B thus enhancing the PL intensity. The observed increase in PL intensity can also be attributed to passivation with oxygenated functional groups, *e.g.*, $-\text{C}=\text{O}$ and OH .^{56,57} B-PPD CDs have a positive zeta potential value, indicating the electron-deficient nature of the B-PPD CDs. An earlier report has suggested that the active oxygen species (AOS) commonly known as the reactive oxygen species, generated from H_2O_2 when it comes into contact



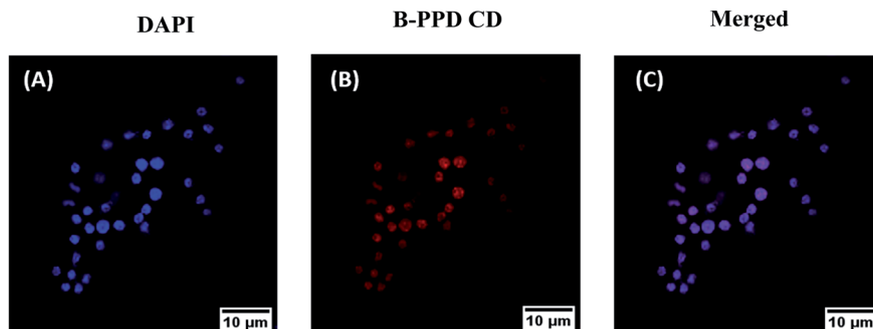


Fig. 6 Confocal laser scanning microscopy (CLSM) images of RAW 264.7 cells co-stained with DAPI and B-PPD CDs. The RAW 264.7 cells were incubated with B-PPD CDs ($20 \mu\text{g ml}^{-1}$) for 30 min and DAPI for 15 min at 37°C . (A) DAPI, (B) B-PPD CDs and (C) merged image of (A) DAPI and (B) B-PPD CDs. The excitation and emission ranges for DAPI and B-PPD CDs were around 405–460 nm and 514–603 nm, respectively. Cells were observed under $63\times$ magnification.

with the quantum dots (QDs), readily captures electrons at the conduction bands of QDs, which results in inhibition of radiative recombination of the photoinduced electrons and holes and causes a decrease in fluorescence intensity.⁵⁸

However, the quenching effect of different active oxygen species on B-PPD CDs is negligible in comparison with H_2O_2 as shown in Fig. 4(B). The selectivity to H_2O_2 can be explained by

putting forward the charge transfer mechanism. It has been assumed that in the presence of H_2O_2 (analyte), there is a charge transfer from H_2O_2 to the electron-deficient boron atom of B-PPD CDs to form stable B–O coordination bonds leading to quenching of PL intensity.⁵⁹ This implies that the doped boron played an important role in the charge transfer mechanism by functioning as a good electron acceptor.

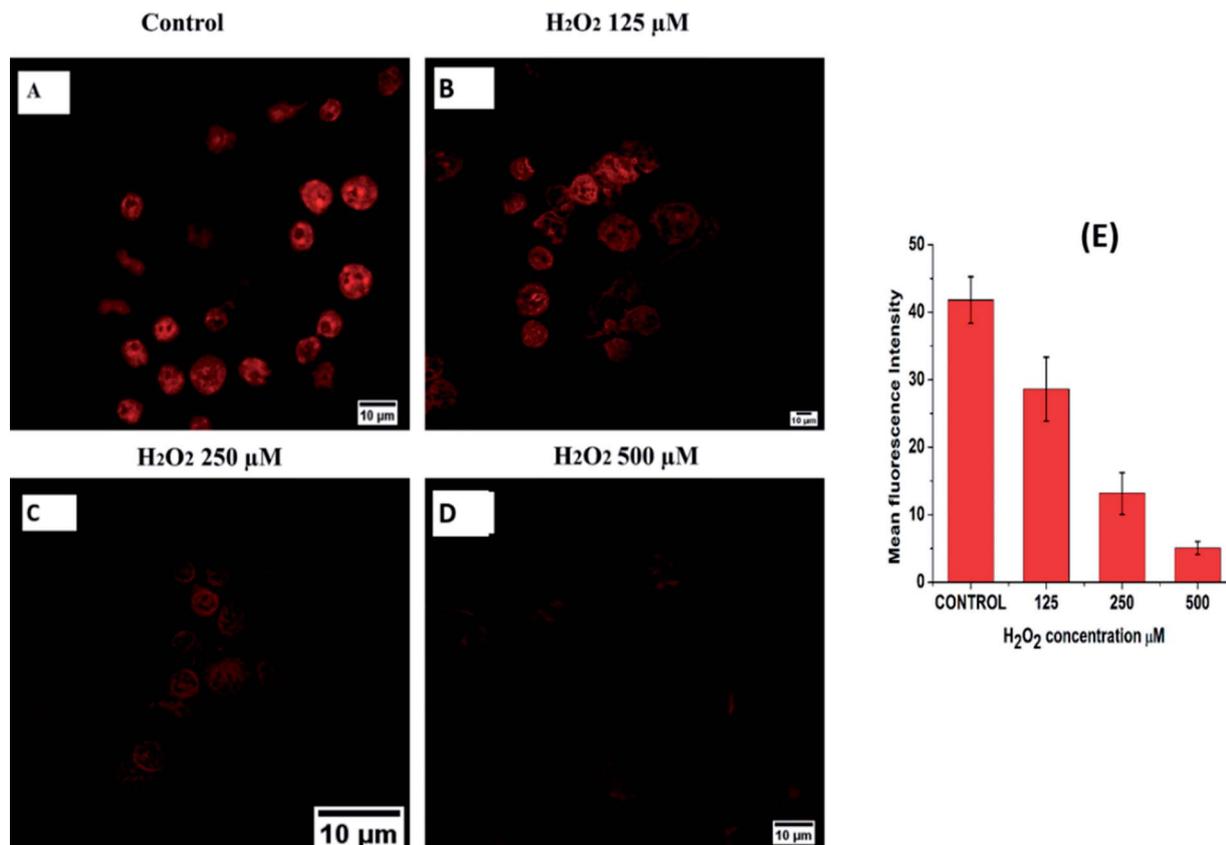


Fig. 7 Confocal fluorescence microscopy images of RAW 264.7 cells incubated with $20 \mu\text{g ml}^{-1}$ B-PPD CDs for 30 min. (A) Without H_2O_2 and (B) treated with $125 \mu\text{M}$, (C) $250 \mu\text{M}$ and (D) $500 \mu\text{M}$ H_2O_2 . Figure (E) presents the mean fluorescence intensity (MFI) of (A–D). Fluorescence intensities were measured using Image J software. The excitation and emission range for B-PPD CDs was around 514–603 nm. Cells were observed under $63\times$ magnification.



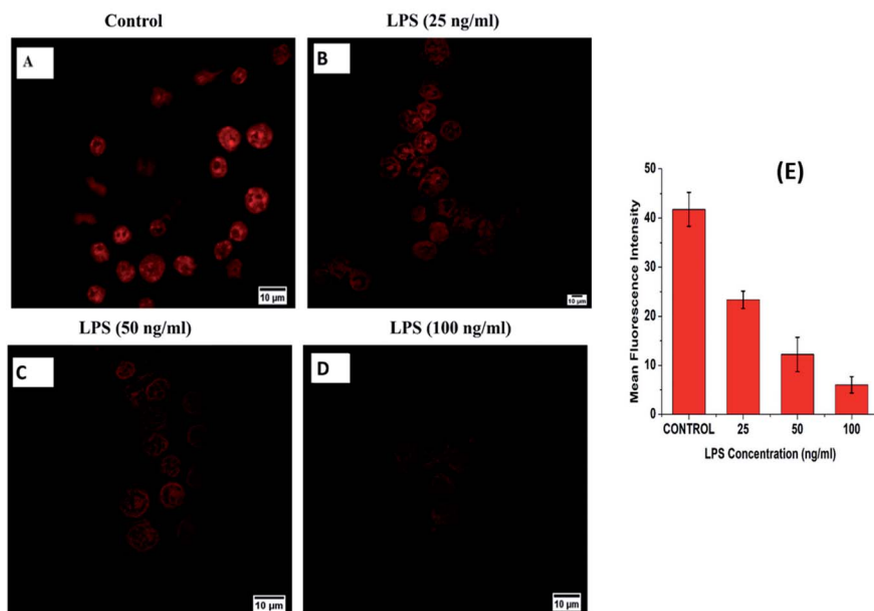
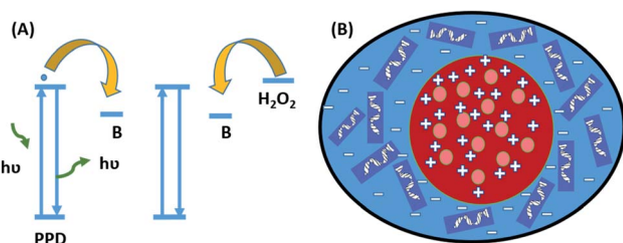


Fig. 8 Confocal fluorescence microscopy images of RAW 264.7 cells incubated with $20 \mu\text{g ml}^{-1}$ B-PPD CDs for 30 min. (A) Without H_2O_2 and (B) treated with 25 ng ml^{-1} , (C) 50 ng ml^{-1} and (D) 100 ng ml^{-1} LPS. Figure (E) shows the mean fluorescence intensity (MFI) of (A–D). Fluorescence intensities were measured using Image J software. The excitation and emission range for B-PPD CDs was around 514–603 nm. Cells were observed under $63\times$ magnification.



Scheme 2 (A) Fluorescence enhancement and quenching of B-PPD CDs. (B) Penetration of B-PPD CDs (Red) inside the nucleus (Blue).

What makes B-PPD CDs an effective nucleus-targeted staining reagent? Some well-known commercially available nuclear staining dyes such as DAPI, Hoechst 33258, and other DAPI analogs share a similar structure (that contains a phenyl ring) with B-PPD CDs. The structural resemblance of B-PPD CDs due to the phenyl ring and small size of the particles ($\sim 1.73 \text{ nm}$) may assist them to enter into the nucleus through the nuclear pores which are about 10 nm in size as demonstrated by confocal microscopy. It has been reported that these nuclear staining dyes exhibited a strong affinity for binding into the minor groove side of the B-DNA molecule specifically in the A–T base region.⁶⁰ This is due to the narrower A–T regions compared with G–C resulting in close-fitting of the flat aromatic ring within the groove; the positive charge of DAPI ($+2$)⁶¹ causes an electrostatic interaction with more electronegative potential in the minor groove of A–T regions allowing the dye to settle deeper into the groove.⁶² Although the actual mechanism of the nucleus-targeting ability of B-PPD CDs needs to be further investigated, it can be assumed that the positive zeta potential

($\zeta = +3.43 \text{ mV}$) of B-PPD CDs imparts a high affinity towards the electronegative region of DNA by electrostatic interactions.⁶³ It may be presumed that the binding of B-PPD CDs with DNA causes the formation of hydrogen bonds between the amine group of B-PPD CDs and the oxygen atoms of thymine, while bifurcated hydrogen bond formation is found between the indole nitrogen of DAPI, netropsin, Hoechst 33258, and thymine oxygen atoms of the two central base pairs.⁶⁴ Thus B-PPD CDs can be used as a nucleus-targeted staining reagent and as a good sensor for monitoring the level of H_2O_2 inside living cells (Scheme 2).

4 Conclusion

In summary, we have synthesized successfully boron-doped *p*-phenylenediamine based carbon quantum dots (B-PPD CDs) as a fluorescent probe in a single-step process. The B-PPD CDs exhibited good biocompatibility and high selectivity against H_2O_2 . *In vitro* studies demonstrated that B-PPD CDs can detect the presence of H_2O_2 . The limit of detection (LOD) was calculated to be $0.242 \mu\text{M}$. The feasibility of B-PPD CDs as a fluorescent probe in *in vivo* studies on the detection of H_2O_2 in cells was also demonstrated successfully. Confocal laser scanning microscopy (CLSM) imaging of B-PPD CDs in the RAW 264.7 cell line showed their targeted nucleus staining and efficient detection of the H_2O_2 level in the nucleus by quenching the fluorescence signal. In addition, they can detect LPS upregulated production of H_2O_2 in cells showing the versatility of the method. Hence, the current study could impart an additional clue for further investigation of DNA damage by oxidative stress.



Abbreviations

ROS	Reactive oxygen species
B-PPD	B-doped <i>p</i> -phenylenediamine based carbon
CDs	quantum dots
CLSM	Confocal laser scanning microscope
LPS	Lipopolysaccharide

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Author contributions

KP: conceptualization, formal analysis, investigation, methodology, and writing – original draft, review and editing.

RRS: formal analysis, investigation, and methodology.

SD: investigation and methodology.

RD: supervision and writing – original draft, review and editing.

DC: conceptualization, formal analysis, supervision, and writing – original draft, review and editing.

Conflicts of interest

The authors declare no competing financial interest.

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