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Ganoderma lucidum bran-derived blue-emissive and green-emissive carbon dots for detection of copper ions†

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Ganoderma lucidum bran (GB) has a broad application prospect in the preparation of activated carbon, livestock feed, and biogas, but the preparation of carbon dots (CDs) from GB has never been reported. In this work, GB was applied as a carbon source and nitrogen source to prepare both blue fluorescent CDs (BCDs) and green fluorescent CDs (GCDs). The former were prepared at 160 °C for 4 h by a hydrothermal approach, while the latter were acquired at 25 °C for 24 h by chemical oxidation. Two kinds of as-synthesized CDs exhibited unique excitation-dependent fluorescence behavior and high fluorescent chemical stability. Based on the fantastic optical behavior of the CDs, they were utilized as probes for fluorescent determination of copper ions (Cu²⁺). In the range of 1–10 μ mol L⁻¹, the fluorescent intensity of BCDs and GCDs decreased linearly with the increase of Cu²⁺ concentration; the linear correlation coefficient reached 0.9951 and 0.9982, and the limit of detection (LOD) was 0.74 and 1.08 μ mol L⁻¹, respectively. In addition, these CDs remained stable in 0.001–0.1 mmol L⁻¹ salt solutions; BCDs were more stable in the neutral pH range, but GCDs were more stable in neutral to alkaline conditions. The CDs prepared from GB are not only simple and low-cost, but also can realize the comprehensive utilization of biomass.

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1. Introduction

The application of heavy metals and their compounds is more and more extensive, and the discharge of wastewater containing heavy metals is also increasing with a rapid development of chemical industry.1 Heavy metals can interact with ligands in the soil to form complexes or chelates, resulting in more excellent solubility and migration activity of heavy metals in soil. The long-term accumulation of heavy metals in the body will cause significant damage to the kidneys and bones.^{2,3} They comprise one of the most harmful pollutants to human health. Heavy metals cannot be degraded by microorganisms and are potential long-term environmental harmful substances.4 In recent years, great efforts have been made to develop efficient heavy metal detection techniques to prevent the increase of heavy metal concentrations, such as ultraviolet spectrophotometry,5 atomic absorption spectrometry,6 fluorescence spectrometry,7 inductively coupled plasma,8 atomic fluorescence spectrometry,9 and inductively coupled plasma mass spectrometry.10 Among them, fluorescence spectrometry is one of the most economical and promising strategies for detecting heavy metals.¹¹ The advantage of fluorescent spectrometry is non-destructive testing, and which can directly analyze finished products.

The increase of global population has led to a sharp growth in the demand for food, which has contributed to the accumulation of agricultural waste. 12 Agricultural waste is known to be rich in lignocellulose. Rational disposal of agricultural waste has far-reaching significance. On the one hand, it can reduce environmental pressure, on the other hand, it can create economic value. Therefore, it is necessary to develop new ways to employ agricultural waste. 13-15 A large number of fungus brans, like GB are piled up everywhere, which not only wastes resources but also pollutes the environment.16 The comprehensive utilization of agricultural wastes must be paid attention to achieving sustainable development. The GB is the residue after harvest cultivated with rice straw, wood chips and other raw materials,17 and usually composed of cellulose, hemicellulose, and lignin, each of which can be utilized as a raw material for valuable chemicals.18,19 After the edible fungi are collected, a large number of mycelium and beneficial bacteria are left on the rod, and a variety of sugars, organic acids, enzymes, and bioactive substances are produced through enzymatic hydrolysis during the growth of mycelium.20 The bacterial residue is rich in protein, cellulose, and amino acids.21 Here, the degradation of GB into high value-added products is expected.

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CDs is a new type of zero-dimensional carbon nanomaterials composed of ultrafine, dispersed, quasi-spherical carbon nanoparticles with a size of less than 10 nm. Compared with other nanomaterials, CDs exhibits their unique characteristics, such as low toxicity, good solubility, chemical durability and remarkable photoluminescence, thus demonstrating application prospects in many fields, such as optoelectronics medical imaging technology, environmental monitoring, chemical analysis, catalyst preparation, energy development, and so on. Many methods for preparing CDs have been proposed, which can be roughly divided into "top-down" and "bottom-up" methods. Compared with the former method, the latter is more diverse. Various biomass materials (such as watermelon peel, coconut shell, orange juice, silkworm leaves, etc.), organic molecules (such as citric acid, sugars, amino acids, ammonium citrate, etc.), and macromolecules (such as chitosan, polyethylene glycol, bovine serum albumin, etc.) can be adopted as precursors.22-25 However, synthesizing CDs from organic reagents and polymers do not conform to the concept of green chemistry. Agricultural by-products as carbon sources to prepare high-value-added carbon materials have attracted significant attention because biomass is renewable, which aligns with green chemistry and sustainable development concepts. Methods for preparing CDs include are discharge method,26 laser ablation method,27 chemical oxidation method,28 electrochemical method,29 template method,30 hydrothermal method,31 direct pyrolysis method,32 and microblog-assisted method.33 In this case, two kinds of CDs were prepared by hydrothermal and chemical oxidation, and their fluorescence quenching for different metal ions and stability were investigated. In this study, CDs with two colors were prepared by a simple method with very sensitive detection of Cu²⁺, providing a new source of carbon for the preparation of detection materials and realizing a high-value use of waste agricultural crops.

2. Experimental section

2.1 Reagents and materials

Sodium hydroxide (NaOH) and hydrogen peroxide (H_2O_2) were provided by Aladdin Industries Co., Ltd (Shanghai, China). Standard solutions of metal ions $(Zn^{2+}, Ca^{2+}, Li^+, Cd^{2+}, Co^{2+}, Cr^{3+}, Cu^{2+}, Fe^{3+}, K^+, Mg^{2+}, Na^+, Pb^{2+} and Zr^{4+})$ were prepared with ultrapure water from respective metal salts of $ZnCl_2$, $CaCl_2$, LiCl, $Cd(NO_3)_2 \cdot 4H_2O$, $CoCl_2 \cdot 6H_2O$, $Cr(NO_3)_3 \cdot 9H_2O$, $CuSO_4$, $FeCl_3 \cdot 6H_2O$, KCl, $Mg(NO_3)_2$, NaCl, $Pb(NO_3)_2$ and $ZrCl_4$, which were also from Aladdin (Shanghai). Hydrochloric acid (HCl) was provided by Sinopharm Chemical Reagent (Shanghai). GB was provided by Ningxia Academy of Agricultural and Forestry Sciences (Yinchuan, China).

2.2 Preparation of CDs

The schematic process is shown in Fig. 1. First, GB was cut into 2–3 mm and dried in an oven at 60 °C for 24 h, and then crushed by a high-energy grinder. After screened through 30 mesh, the GB powder was dispersed into 60 ml of deionized water and

placed in a reaction kettle and reacted at 160 °C for 4 h. After the reaction kettle was cooled to room temperature, the BCDs and the GB-residue were generated and separated by centrifugation. The GB-residue further reacted in a solution containing 0.2 mol L $^{-1}$ NaOH and H $_2$ O $_2$ (5:1, v/v) at 25 °C for 24 h. After the reaction, the obtained GCDs suspension was neutralized with HCl. Thus, the GCDs was fabricated by chemical oxidation. The reacted liquid is filtered through a 0.22 μm microporous membrane to remove insoluble impurities. The CDs solution was then dialyzed with a dialysis membrane (MV = 1000 Da) for 24 h. The obtained CDs were stored at 4 °C for further assays.

2.3 Characterization of material

Transmission electron microscopy (TEM) images were obtained by FEI Tecnai G2 F20 (China). Surface functional groups were determined by Fourier-transform infrared spectroscopy (FT-IR, presri-21 400–4000 cm⁻¹). X-ray photoelectron spectroscopy (XPS) was performed on ESCALAB 250XI XPS spectrometer (Thermo Scientific, USA). Fluorescence spectra were measured with a fluorescence spectrometer (RF-5301 PC, Shimadzu, Japan). X-ray diffraction was obtained by scanning with Smartlab SE (XRD, XRD-6100200 Ma, Japan). Fluorescent characteristics were analyzed by an ultraviolet-visible spectrophotometer (UV, TU-1950).

2.4 Fluorescent property of CDs

2.4.1 Fluorescent quantum yield CDs. Fluorescent quantum yield is an important parameter to evaluate the fluorescent ability of materials. Therefore, the fluorescent quantum yield of CDs was estimated according to the standard method. Quinine sulfate with similar excitation wavelength (366 nm) was selected as the standard reference. The fluorescent quantum yield could be calculated according to the eqn (1):

$$Y_1 = (Y_2 \times S_1 \times A_2 \times n_1^2)/(S_2 \times A_1 \times n_2^2) \tag{1}$$

where Y is the fluorescent quantum yield, S is the fluorescent peak area, A is the absorbance, and n is the refractive index of the solvent. Quinine sulfate was selected in this experiment. The literature shows that its quantum yield (Y_2) is 0.54.

2.4.2 Fluorescent selective and sensitive detection for metal ions by CDs. The fluorescent quenching degree of CDs depending on the concentration of Cu²⁺ was calculated by Stern-Volmer eqn (2):

$$F_0/F = 1 + K_{SV}C$$
 (2)

where F_0 is the fluorescent intensity of CDs without metal ions, F is CDs combined with Cu^{2+} , C is the concentration of Cu^{2+} , and K_{SV} is the quenching constant.

Results and discussion

3.1 Preparation and characterization of CDs

The synthesis of multicolor fluorescent CDs from biomass is still an exciting research field.^{34,35} The main component of biomass is lignin, which has the perfect robustness that comes

Hydrothermal

Centrifugation

BCDs

Chemical oxidation

GB powder

CDs suspension

GB-residue

GCDs

Fig. 1 Schematic diagram of preparation of BCDs and GCDs.

from its remarkable and compact morphology structure. This particular structure makes lignin difficult to pyrolysis, limiting its deep development and utilization. 36 Hydrothermal approach is one of commonly utilized methods for pyrolysis of lignocellulose under high temperature and high pressure.37 After reaction, macromolecular polymers can become fluorescent CDs.38,39 The CDs prepared by hydrothermal approach has the advantages of small particle size, uniform distribution, light particle agglomeration, and cheaper raw materials. Chemical oxidation is a method to oxidize (such as H₂O₂ and other strong oxidants) the target by losing electrons with oxidants. 40 The advantages of chemical oxidation include manageable control, convenient operation, etc. In this work, GB was selected as raw material, and its main components include cellulose, hemicellulose, and lignin. As shown in Fig. 1, BCDs was synthesized by a simple hydrothermal approach, while GBDs was prepared by chemical oxidation approach.

The fluorescent color of CDs is related to particle size, graphitization degree and functional group.⁴¹ The microscopic appearance of the two CDs was observed by TME. As shown in Fig. 2a and b, both BCDs and GCDs exhibited well dispersity, and morphology of CDs were spherical. The size distribution ranged in 1.57–2.83 nm (Fig. 2c) and 2.03–3.85 nm (Fig. 2d), respectively, indicating uniform particle size distribution of BCDs and GCDs.⁴²

The chemical structures of the BCDs and GCDs were studied by FT-IR spectra and the results are shown in Fig. 3a. A wide peak at 3413 cm⁻¹ was assigned as tensile vibration of -NH_x or -OH, while the peak at 1739 cm⁻¹ was assigned as a bending vibration of C=0.43 The absorption peak at 1646 cm⁻¹ was the tensile vibration of aromatic compound C=C. The absorption peaks at 1254 and 1044 cm⁻¹ were asymmetric stretching vibration peaks of C-O and C-H. Compared with spectrum of BCDs, the absorption peaks at 3413 cm⁻¹ in spectrum of GCDs was significantly enhanced, which was attributed to a large number of oxygen-containing or amino functional groups formed by chemical oxidation.44 The content of oxygen and nitrogen functional groups were closely related to fluorescent color of CDs. 45-47 The prepared CDs exhibited excellent compatibility with water, which was related to existence of these hydrophilic functional groups and provided favorable conditions for luminescence.

The phase structure analysis of BCDs and GCDs was carried out by XRD technique, as shown in Fig. 3b. A hump-shaped diffraction peak was located at 22.33°, which was assigned as

characteristic peak of crystal planes of carbon material (002), indicating that the diffraction peaks of BCDs and GCDs were still mainly graphite-like structures.⁴⁸ The stacking between layers in CDs was similar to graphite, which means that highly conjugated sp² carbon atoms were closely stacked, demonstrating the rich graphene structure of CDs. However, compared with the spectrum of BCDs, the diffraction peaks at 22.33° was non-obvious changed in the spectrum of GCDs, indicating that the graphitization degree were similar.⁴⁹

XPS was performed to investigate the surface chemical composition, and the high-resolution full spectra of BCDs and GCDs are shown in Fig. 4. In the N 1s spectrum of BCDs (Fig. 4a), the peak at 398.0 eV belonged to N-H group, while in the N 1s spectrum of GCDs (Fig. 4d), the increase in the nitrogen content of GCDs due to the new chemical bond C-N at 400.2 eV after oxidation of GCDs leads to the color change of GCDs. The peaks at 398.4 and 400.2 eV belonged to C-N and N-H groups. 50 In Fig. 4b, the peaks at 284.8, 287.4 and 289.2 eV were assigned as C=C/C-C, C-O and C=O group in BCDs, respectively. In C1s XPS spectrum of GCDs (Fig. 4e), the peaks at 284.2, 286.6 and 288.8 eV belonged to C=C/C-C, C-O and C=O group, respectively. The peaks at 532.7 and 533.0 eV belonged to C=O and C-O groups, respectively, in O 1s spectrum of BCDs (Fig. 4c). The peaks at 532.1 and 533.5 eV belonged to C=O and C-O groups in BCDs (Fig. 4f).51,52 These results indicated that BCDs and GCDs possessed almost same functional groups, the existence of which also proved that CDs was not a complete sp² structure, but doped with nitrogen and oxygen atoms.

3.2 Fluorescent property of CDs

The formation of natural substances is complex, mainly connected by hydrogen bonds and various chemical bonds.⁵³ Deionized water under high pressure and high temperature provides the impetus for the breaking of chemical bonds, thus making GB degradation and destruction under high temperature and pressure to form oligomers and micro-molecules, which contain carboxyl groups and other tubular energy groups in these micro-molecules, making the solution acidic, under continuous high temperature and pressure carbohydrates are further degraded to form more under continuous high temperature and pressure, carbohydrates are further degraded to form more water-soluble small molecules, which are aggregated to form polymers and become macromolecular compounds, and finally these macromolecular compounds

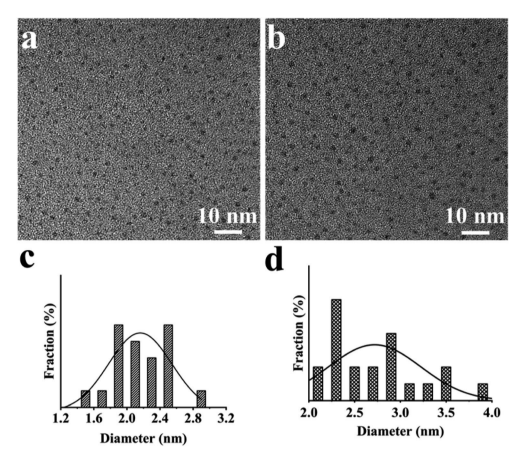


Fig. 2 TEM image of (a) BCDs and (b) GCDs, particle size distribution diagram of (c) BCDs and (d) GCDs.

form CDs in solution, in which the aqueous solution not only acts as an oxidant and catalyst, but also helps to form CDs. 54,55 Then tiny molecular substances were successively converted into BCDs through polymerization and condensation. The GCDs was prepared by adding a certain proportion of NaOH and $\rm H_2O_2$ to the GB-residue, thus fluorescent color was changed by a simple chemical oxidation.

In the UV-vis spectrum of BCDs (Fig. S1†), the peaks at 225 and 240 nm belonged to $\pi \to \pi^*$ electronic transition caused by C=C, while in the UV-vis spectrum of GCDs, the peaks at 212 and 237 nm also belonged to $\pi \to \pi^*$ electronic transition caused by C=O.^{22,56} Moreover, the fluorescent properties of CDs were also related to particle size, functional groups and graphitization degree. As mentioned above, the size distribution of BCDs ranged in 1.57–2.83 nm, while that of GCDs ranged in

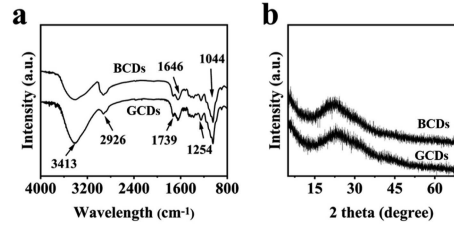


Fig. 3 (a) FT-IR spectra and (b) XRD diffraction of CDs.

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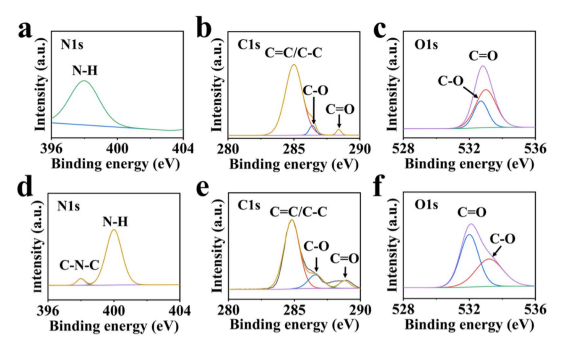


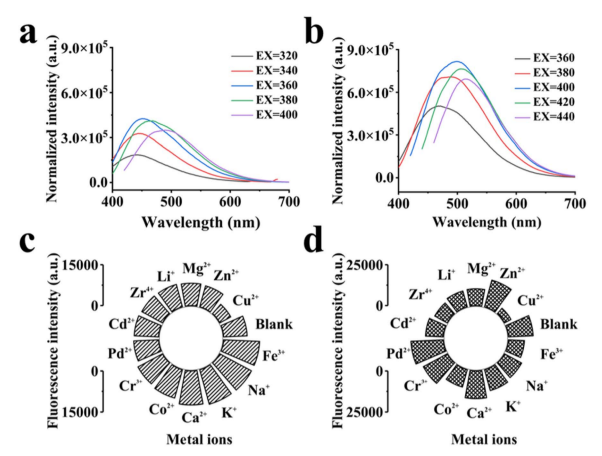
Fig. 4 High resolution XPS spectra of (a and d) N 1s, (b and e) C 1s, (c and f) O 1s of (a, b and c) BCD and (d, e and f) GCD.

2.03–3.85 nm. It was speculated that different fluorescent color was related to the size of particles, and a red shift of luminescence color of CDs occurred with the increase of particle size. ⁴² The variation of emission wavelengths of BCDs and GCDs is related to the size variation of both CDs and the content of graphitic nitrogen atoms in the carbon core. The graphitic nitrogen content can increase the domain stiffness and thus reduce the nonradiative relaxation through the molecular vibrations on the surface of CDs. Thus, particle size and graphitic nitrogen content synergistically control the emission wavelength of CDs and play an important role in the color variation of CDs. ^{50,57,58}

3.2.1 Fluorescent quantum yield of CDs. As presented in Fig. 5a and b, the excitation wavelength of BCDs was 320, 340, 360, 380, 400 nm, its emission wavelength was 440, 445, 450, 465, 490 nm, respectively. The excitation wavelength of GCDs were 360, 380, 400, 420, 440 nm, its emission wavelength were 470, 490, 500, 510, 520 nm, respectively. The fluorescent emission spectra of CDs under different excitation wavelength show that CDs has excitation dependence, which was entirely consistent with the excitation dependence reported by Ganjkhanlou et al.59 Fluorescent quantum yield is an essential parameter for evaluating the fluorescent ability of materials. Therefore, quinine sulfate with a similar excitation wavelength was selected as the standard sample to calculate fluorescent quantum yield of CDs according to eqn (1). The fluorescent quantum yield of BCDs and GCDs were 4.6% and 2.6%, respectively.

3.2.2 Fluorescent selective and sensitive detection for metal ions by CDs. The selectivity of BCDs and GCDs for 13 kinds of metal ions was investigated, as shown in Fig. 5c and d. Both BCDs and GCDs generated fluorescence in the blank solution with strong intensity. When BCD was added into

aqueous solutions of Fe³⁺, Na⁺, K⁺ and Ca²⁺, respectively, the fluorescence signals were remarkably enhanced by 53%, 53%,52%, 44%, while adding Co²⁺, Cr³⁺, Pd²⁺ and Cd²⁺ led to slightly decrease of fluorescent intensity of 24%, 18%, 5.9%, 3.6%. When GCDs was added into aqueous solutions of Ca²⁺, Cr3+, and Pd2+, respectively, the fluorescence signals were remarkably enhanced by 9.6%, 31%, 11%, while adding Fe³⁺, Na⁺, K⁺, Co²⁺, Cd²⁺, Zr⁴⁺, Li⁺, Mg²⁺ and Zn²⁺ led to slightly decrease of fluorescent intensity of 32%, 34%, 43%, 28%, 28%, 13%, 22%, 36%. The CDs with $\pi \to \pi^*$ electronic transition were easier to enhance fluorescence, and fluorescence intensity will generally increase as conjugation degree of the system increases. This was mainly because increasing the molar absorption coefficient of fluorescent substances was conducive to generating more excited molecules, which was conducive to the occurrence of fluorescence. Weakening fluorescence response was because CDs and metal ions encounter and collide with less energy loss, or CDs and metal ions react to form a non-luminous coordination compound. The strong response of BCDs (47%) and GCDs (71%) to Cu²⁺ was mainly attributed to oxygen-containing group and nitrogen-containing groups on the surface CDs.59 The electrons were attributed to Cu2+ as an electron donor. The lone pair electrons of the nitrogen atom shown a good affinity for Cu²⁺, which could be attributed to the behavior of the characteristic pair of nitrogen atom and Cu²⁺.60 The quenching constant is an important means to measure fluorescent sensitivity. The quenching constant was an important means of determining fluorescent sensitivity, which were calculated according to Stern-Volmer equation (eqn (2)). The quenching constant of BCDs and GCDs were 0.7×10^7 and 2.3 \times 10⁷, respectively, demonstrating perfect fluorescent response of CDs to Cu²⁺.



Fluorescence emission spectra of (a) BCDs and (b) GCDs, fluorescence intensity of (c) BCDs and (d) GCDs with 13 kinds of metal ions.

3.2.3 Quantitative measurement and quenching mechanism of CDs. The fluorescent response of BCDs and GCDs to different concentrations of Cu²⁺ were tested, as shown in Fig. 6a and b. The fluorescent intensity of BCDs and GCDs decreased with an increase of Cu²⁺ concentration. The linear relationship between fluorescent intensity of BCDs with Cu²⁺ concentration and that of GCDs are shown in Fig. 6c and d, and the linear correlation coefficients R were 0.9951 and 0.9982, respectively. The limit of detection (LOD) of BCDs and GCDs were 0.74 \times 10^{-3} and $1.08\times 10^{-3}\,\mu\text{mol}\,\text{L}^{-1}\text{,}$ when the concentration of Cu^{2^+} ranged in 1-5 µmoL L⁻¹, which provided a new idea for the preparation of carbon-based dual-functional materials. Table 1 lists several CDs materials with fluorescent properties for selective detection of Cu²⁺. Yan et al.⁶¹ prepared CDs by grafting 7-diethylaminocoumarin-3-carbohydrazide onto the surface of glyoxylic acid-modified. The CDs through amidation reaction, within the range of 0-10 μ mol L⁻¹, LOD was 0.21 μ mol L⁻¹. Ma et al. 62 prepared CDs as a potential probe for metal ion sensing by a simple stirring treatment of AC with sulfuric acid, ranging from 0 to 500 μ mol L⁻¹, the LOD was 2.40 μ mol L⁻¹. Zeng et al. 63 utilized 2,4-difluorobenzoic acid and 1-lysine as precursors to synthesize nitrogen fluoride co-doped CDs by hydrothermal method. Within the concentration range of 0-200 μ mol L⁻¹, LOD was 0.22 μ mol L⁻¹. When concentration of Cu²⁺ within the range of 1-10 μmol L⁻¹, LOD of BCDs and GCDs were lower than other materials. The fluorescent detection method based

on BCDs and GCDs has good sensitivity and could be utilized for quantitative analysis and selective detection of Cu²⁺.

Fluorescence quenching effect is related to the yoke structure of Cu²⁺. CDs are formed by the polymerization of a series of aromatic compounds. When Cu²⁺ aqueous solution was added, an electron transfer occurred and the light generated by the CDs was absorbed by Cu2+. To further verify this deduction, the fluorescence lifetime was detected (Fig. S2†). The fluorescence lifetime of the material with the addition of Cu²⁺ was increased. The experimental results show that, the fluorescence quenching of BCDs and GCDs is caused by dynamic quenching. The oxygen-containing groups (OH, C=O, COOH) present on the surface of CDs can interact with metal ions (especially Cu²⁺), resulting in CDs behaving as chelators. Usually, bursts are observed due to electron or energy transfer between the excited CDs and the metal ions. In this case, the burst phenomenon is due to electron transfer rather than through energy transfer. In addition, the decay of the fluorescence lifetime of CDs in the absence and presence of Cu²⁺ was calculated. The fluorescence lifetimes of BCDs and GCDs were 1.26 and 1.21 ns, while the lifetimes after the addition of Cu²⁺ were 9.15 and 4.57 ns. The fluorescence lifetime of CDs increased after the addition of Cu²⁺. This result suggests that the electron transfer from the excited CDs to the metal ion vacancy orbitals leads to a strong nonradiative electron/hole complex, aggregation-induced emission burst in the CDs. 64,68,69

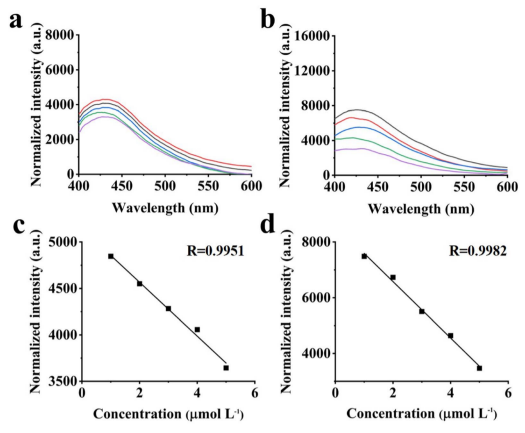


Fig. 6 Fluorescence emission spectra of (a) BCDs and (b) GCDs with different concentrations of Cu^{2+} , linear response curve of (c) BCDs and (d) GCDs for detection of Cu^{2+} .

3.2.4 Effect of salt solution on fluorescent response of CDs. The salt resistance of BCDs and GCDs were tested. The NaCl of different concentrations was utilized as the interferent to detect fluorescent intensity of BCDs and GCDs. As shown in Fig. 7a and b, the concentration of NaCl increased from 0.001 to 0.1 mol $\rm L^{-1}$ in turn. In the range of 0.001–0.1 mmol $\rm L^{-1}$ of NaCl, the fluorescent intensity of BCDs and GCDs maintains a stable state. In conclusion, BCDs and GCDs show excellent stability, which makes BCDs and GCDs have good application prospects in detection. 70

3.2.5 Effect of pH on fluorescent response of CDs. In order to further explored stability of CDs in an aqueous solution, the effects of different pH conditions on fluorescent intensity were investigated. As shown in Fig. 7c and d, when the pH value of

the solution changes from 2 to 12, fluorescent signal of BCDs was strongest under neutral conditions. The fluorescent response of BCDs was significantly weakened at pH 2–4 and 10–12, which was attributed to the protonation of hydroxyl groups under acidic conditions or the precipitation of hydroxyl groups with other substances under alkaline conditions. The fluorescent signal of GCDs significantly weakened with the increase of acidic solution, which was also due to protonation of hydroxyl groups. The fluorescent intensity hardly fluctuates wildly at a pH of 5–9. Under acidic conditions, the fluorescent intensity of GCDs gradually decreased with the increase of acidic concentration. However, the fluorescent intensity of GCDs was slightly enhanced under alkaline conditions, which

Table 1 Comparison of the reported materials for fluorescent detection of Cu²⁺

Material	Linear range $(\mu \mathrm{mol~L}^{-1})$	LOD ($\mu mol~L^{-1}$)	Ref.
Coumarin-functionalized CDs	0-10	0.21	61
Prepared from activated carbon CDs	0-500	2.40	62
Prepared from 2,4-difluorobenzoic acid CDs	0-200	0.22	55
Tea-CDs	0-170	0.051	65
Banana-CDs	0-12.5	0.004	66
Lignin-CDs	0-30	0.085	67
BCDs	1-5	0.74×10^{-3}	This work
GCDs	1–5	1.08×10^{-3}	This work

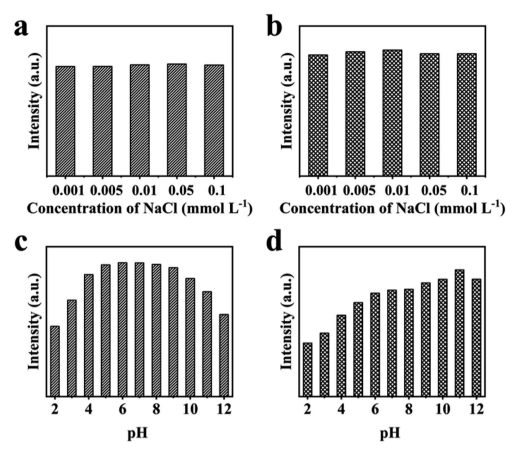


Fig. 7 Effects of (a and b) concentration of NaCl and (c and d) pH of solution on fluorescence response of (a and c) BCDs and (b and d) GCDs.

might be due to the different preparation methods from BCDs.⁷⁰ The GCDs was prepared by chemical oxidation under alkaline conditions, and GCDs suspension was also alkaline. The fluorescent intensity of BCDs for fluorescent detection of Cu²⁺ in a neutral solution was enhanced. The stability of BCDs was excellent in the pH range of 5–9, and the fluorescent intensity of GCDs was not affected by pH in the range of 7–12.

Conclusion

Two kinds of CDs were obtained from GB by hydrothermal and chemical oxidation. This method of using biomass two-step method to form two color CDs has not been reported before. It is found that the luminescent properties of CDs were related to surface functional groups or graphitization, and shown good luminescence stability. Surface state modification opens up a new research direction for polychromatic luminescence. The blue fluorescence was successfully converted to green fluorescence by adjusting the hydrogen bond on the CDs surface. The CDs of two colors was sensitive to Cu²⁺, and LOD were 0.74 and 1.08 μmol L⁻¹, respectively, within the concentration range of 1–10 $\mu mol \ L^{-1},$ and correlation coefficient was as high as 0.9951 and 0.9982. In addition, these CDs were still stable in different 0.001-0.1 mmol L⁻¹ salt solutions, BCDs was more stable in the neutral pH range, and GCDs was more stable in neutral to alkaline conditions. This work proposes a feasible method for

the high value utilization of waste crops and a new raw material for the preparation of detection materials. In our future research work, we will focus on finding greener preparation methods and more sensitive detection materials.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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