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Improved voltammetric discrimination of acetaminophen and uric acid in urine using CoO biochar nanocomposite†

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Overuse of acetaminophen (APAP) has become a severe societal burden in recent years. The rapid and reliable detection of urine APAP concentration can offer certain guidance for better management of APAP usage. This study explored the electrochemical sensing application of a novel electrocatalyst prepared from the biomass of *Elaeagnus angustifolia* gum. The biomass was first activated by ferric chloride to form a porous biomass carbon material (FBC). Then cobalt oxide (CoO) cracked nanoplate were synthesized by alkali precipitation and calcination and were then hybridized onto the biomass carbon via a simple sonication process. The electrocatalyst of CoO-FBC was characterized by scanning electron microscopy (SEM) and energy-dispersive X-ray spectroscopy (EDS), element mapping, transmission electron microscopy (TEM) and high resolution (HR-TEM), X-ray photoelectron spectroscopy (XPS), X-ray powder diffraction (XRD), thermogravimetric analysis (TGA), Raman spectroscopy, and nitrogen adsorption/desorption analysis. The CoO-FBC modified glassy carbon electrode (CoO-FBC/GCE) was characterized by various electrochemical methods including cyclic voltammetry (CV), electrochemical impedance spectroscopy (EIS) and differential pulse voltammetry (DPV). The CoO-FBC/GCE sensor was used to measure APAP in 0.1 M phosphate buffered saline (PBS) with a pH of 7.0, and with two linear sensing ranges from 1 μM to 10 μM and from 10 μM to 100 μM , with a sensitivity of 25.89 $\mu\text{A } \mu\text{M}^{-1} \text{ cm}^{-2}$ and 10.04 $\mu\text{A } \mu\text{M}^{-1} \text{ cm}^{-2}$, respectively, and a limit of detection of 0.46 μM . The unavoidable interference in measuring APAP is the inherent uric acid in urine. Uric acid and APAP exhibited adjacent and sometimes unseparable voltammetric peaks. This CoO-FBC/GCE sensor is capable of distinguishing APAP from uric acid and so APAP can be measured in human urine samples with good recoveries. This CoO-FBC/GCE sensor is a promising application for clinical diagnosis and environmental detection.

Keywords: *Elaeagnus angustifolia* gum; Ferric chloride; Polysaccharide biomass; Cobalt oxide nanoplate; Electrochemical sensing; Analgesic and antipyretic drug.

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1 Introduction

Acetaminophen (*N*-acetyl-*p*-aminophenol, APAP, also commonly known as paracetamol) is a common ingredient used for analgesic and antipyretic effects. The excessive presence of APAP in human body fluids and contamination in environmental waters can heavily influence the human body and earth ecosystem.^{1–3} The U.S. Food and Drug Administration recommends that an adult dose of APAP is

4000 milligrams per day. The most notorious side effect of the overuse of APAP is acute liver damage. Besides hepatotoxicity, other potential risks may include pregnancy interference, rare skin reactions, and so on. Currently, multiple centralized laboratory methods can quantify APAP in various samples, e.g. high-performance liquid chromatography. However, such measurement requires use of expensive instrumentation and skilled personnel. Electrochemical sensing is promising in providing rapid, specific, inexpensive, internet integration, miniaturization and fully automated means for efficient clinical diagnoses and environmental detection.^{4,5}

Biomass represents a wide range of organic components, and sustainable and recyclable natural resources in the environment. Although more or less differences might exist in such huge biomass family, the major composition of

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biomass is carbon-based. Biomass derived carbon has found a multitude of applications in energy, sensing, biomedicine, the environment and catalysis.^{6–12} *Elaeagnus angustifolia* gum is secreted in from tree trunks of *Elaeagnus angustifolia* L. and its major composition is polysaccharide. Our previous study revealed that the molecular weight of this polysaccharide is about 160 kDa. The *E. angustifolia* gum was originally used as a medicinal material. Here we explore the potential use of it as a biomass derived carbon material for electrochemical sensing applications. Biomass derived carbon has several characteristics including simple preparation, good conductance, high surface area and as a matrix for producing composites with nanomaterials, and so on.^{13,14}

In order to achieve a larger surface area, several methods have been used to activate biomass carbon (BC) material, and a porous structure is usually formed after the activation. Traditional activation adopts a strong alkali etching approach, e.g. NaOH and KOH. In recent years, new activation chemicals were selected to activate biomass derived carbon material, including chloride salts like ZnCl₂, FeCl₃, and CaCl₂, and so on. The intrinsic activating mechanism is complicated by the fact that the metal salts can form multiple materials, such as a metal hydroxide transforming into metal oxide by using a different calcination process. In addition to the chemical composition change and gas/liquid release, the activating agents may also expand or shrink and as a result, form a porous structure in the biomass material.^{15,16} For the FeCl₃ activation, FeCl₂, FeOCl, C₃N₄, and FeOOH, and so on, are formed during the calcination. The polysaccharide has a certain chemical reducibility and is able to interact with a variety of oxidative chemicals. The multiple iron species offer a complex reaction with polysaccharides which forms exquisite carbon structures.¹⁷

Transition metals have multiple orbital electrons and multiple valences which gives them a novel special catalytic function towards a mass of chemicals. Cobalt containing nanomaterials are one of the research topics for the degradation of various chemicals and sensing. Li and co-workers synthesized the Co–SnP composite, which can completely degrade APAP in Co–SnP/peroxymonosulfate system within 15 minutes. The catalyst of Co–SnP also exhibited excellent stability, reusability and universality.¹⁸ Kim and co-workers prepared a cobalt hexacyanoferrate incorporating a Fe-terephthalate metal organic framework as a robust and cost-effective acetaminophen sensing GCE modification material. The modified GCE sensor demonstrated excellent selectivity, sensitivity, stability, reproducibility.¹⁹ The Senthil Kumar group synthesized the highly uniform N-doped hollow carbon spheres with a uniform distribution of cobalt oxide materials and used them for the electrochemical sensing of APAP.²⁰ Besides a cobalt nanomaterial for APAP sensing, the Mohamed group fabricated graphite carbon nickel nanoparticles with functionalized multiwalled carbon nanotubes for the detection of an immunomodulator drug, baricitinib, which

showed a high precision in sensing baricitinib in plasma and medicinal formulations.²¹

Herein, we prepared the porous biomass carbon by ferric chloride activation of the gum derived from *E. angustifolia* using a simple pyrolysis procedure. To enhance the electrocatalytic property for use in measuring APAP concentrations, we synthesized cobalt hydroxide by alkali precipitation of Co²⁺ aqueous solution and calcined the cobalt hydroxide into a cobalt oxide cracked nanoplate. The sonication method was chosen to hybridize the cobalt oxide cracked nanoplate onto the porous biomass carbon material. This CoO-FBC/GCE sensor could sense APAP in a wide range of concentrations and was able to detect APAP in urine as well.

2 Results and discussion

2.1 Material characterization

Synthesized biomass derived carbon electrocatalyst materials of BC, FBC and CoO-FBC were checked using SEM, elemental mapping, TEM and HR-TEM techniques. As shown in Fig. 1a–c, the unactivated BC displays some mesopores on the surface, but a lot of solid parts remained. In comparison, the ferric chloride activated FBC shows pipe like etched micropores when compared to BC. A high degree of porous structure could be seen, and the inner parts are also highly porous. In the CoO-FBC SEM image, many cobalt oxide cracked nanoplates are attached onto the FBC's surface in random directions, demonstrating the successful incorporation of the CoO cracked nanoplate onto the FBC. In the magnified SEM image (Fig. S1†), irregular cracked holes are present in the hexagonal plate. These holes are assumed to come from the cobalt hydroxide shrinkage cracking during the calcination. These holes expose more cobalt atoms on the nanoplate surface. The hexagonal nanoplates have a side length of about 200 nm and a thickness of about 20 nm. In Fig. 1d–g a larger view of the electrocatalyst is shown for the elemental mapping. The CoO-FBC of (d) shows the presence of (e) the cobalt element, (f) the oxygen element and (g) the carbon element. The cobalt and oxygen elements are mainly from the cobalt oxide cracked nanoplate. The inverse deep red carbon signal area reflects the biomass carbon material. The elemental mapping of BC and FBC, and the EDS spectra of all three biomass derived carbon materials can be found in Fig. S2.† Fig. 1h–j are the TEM images of BC, FBC and CoO-FBC, respectively. The BC TEM image shows the solid structure of the carbon material. Whereas the exterior part of the FBC is etched in an unorganized way as shown in Fig. 1i, the rich internal pipe-like hollow pore structure can still be clearly recognized. The CoO-FBC shows the successful hybridization of the CoO nanoplate on the FBC surface. The internal pipe-like hollow pores remained after the CoO decoration shown in Fig. 1j and the cracked nanoplate structure is clearly seen. The HR-TEM image of the CoO in the CoO-FBC is shown in Fig. 1k, and the interplanar crystal spacing was measured and found to be 0.66 nm, which is in



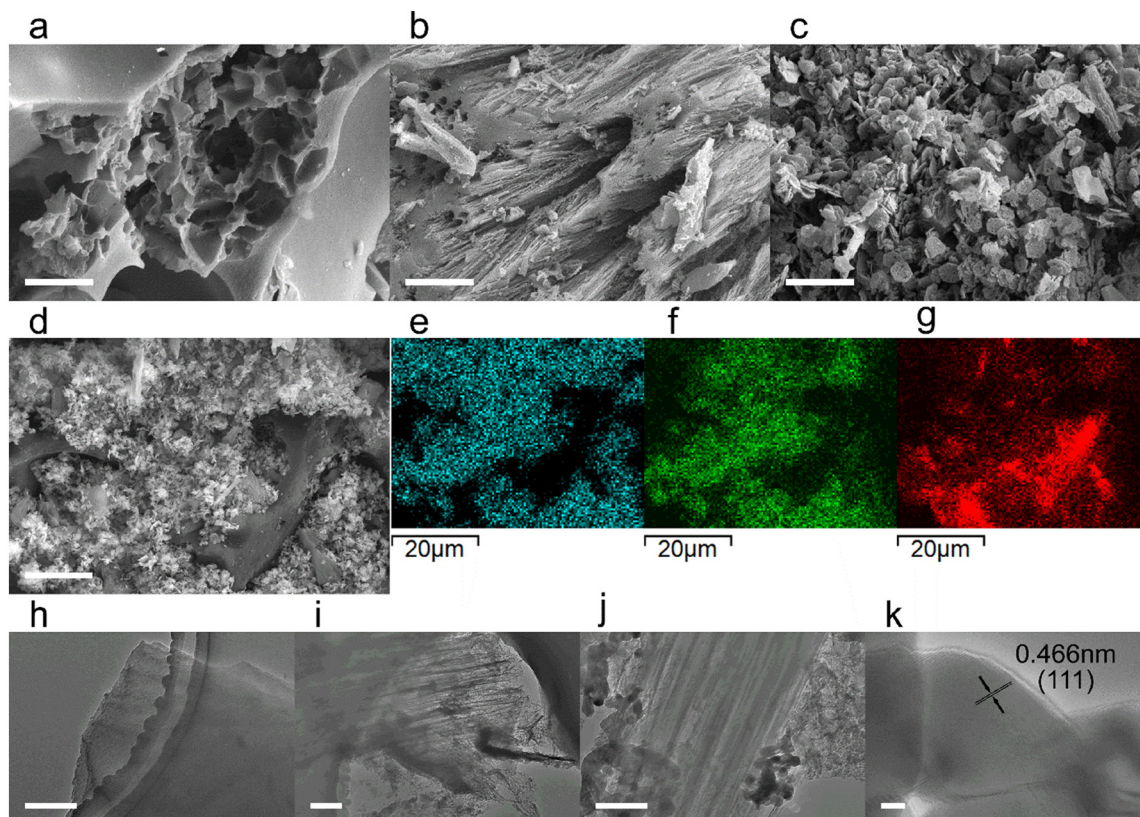


Fig. 1 SEM images of (a) BC; (b) FBC; (c) CoO-FBC; SEM image of (d) CoO-FBC and corresponding elemental mapping of (e) cobalt, (f) oxygen, (g) carbon; TEM images of (h) BC; (i) FBC; (j) CoO-FBC; (k) HR-TEM image of the enlarged CoO in CoO-FBC. Scale bar in (a–c) is 2 μm , in (d) is 10 μm , in (h–j) is 200 nm, and in (k) is 5 nm.

agreement with the (111) plane of the Co_3O_4 (PDF-#43-1003). The results of the XRD analysis are shown in Fig. 2c.

Fig. 2a shows the nitrogen adsorption/desorption curves of the prepared biomass derived materials. Obviously, all three carbon materials display characteristic features of type I and IV isotherms (according to IUPAC). The hysteresis loop is in the range of 0.45 to 1.0 (P/P_0). The porous structure and the successful activation of the biomass carbon by the ferric chloride method are confirmed. The unactivated BC has the smallest calculated Brunauer–Emmett–Teller (BET) surface area of 342 $\text{m}^2 \text{g}^{-1}$, and the FBC has the largest calculated BET surface area of 894 $\text{m}^2 \text{g}^{-1}$, which is 2.6 times larger than the BC's surface area. After the use of sonication to incorporate the cobalt oxide onto the FBC, the CoO-FBC has a decreased calculated BET surface area of 429 $\text{m}^2 \text{g}^{-1}$. The decrease is assumed to come from the blocking by the cobalt oxide cracked nanoplate. From the measured pore size distribution (Fig. S3†), the mesopores with a size close to 17 nm are the major pores in the FBC and CoO-FBC, whereas the BC does not show any mesopores. After the nanoplate hybridization, the pore size near 17 nm is lowered.

Fig. 2b displays the Raman spectra of the biomass derived materials. The typical D band and G band of most carbon materials at around 1335 cm^{-1} and 1587 cm^{-1} are clearly seen. The D band signal represents the disordered carbon atom, originating from the carbon distortions and edges in

the material structure. The G band signal represents the ordered graphitic carbon atom, originating from the vibration in the 2D hexagonal lattice of the sp^2 carbon atom. Therefore, the band intensity ratio of I_D/I_G which reflects the disorderedness of the carbon atom in the carbon material. The I_D/I_G values of BC, FBC and CoO-FBC were 1.031, 1.036 and 1.026, respectively. After the activation, the I_D/I_G value of FBC increased slightly, and this came from the added etched porous inner pipeline holes and the outer surface. After the cobalt oxide nanoplate incorporation, the I_D/I_G value of CoO-FBC decreased to 1.026, which was smaller than that of BC. The diminution was speculated to come from the carbon edges forming physical and chemical bonds with the nanoplate, which reduce the carbon material's edges.²²

Fig. 2c displays the XRD spectrum of the biomass derived materials. The BC shows the characteristic hard carbon diffraction peaks of (002), (g002) and (101) at 2θ of 22.7, 27.0 and 44.7, respectively. The (g002) is attributed to the crystallographic plane (002) reflection of graphitic carbon.^{23,24} As the BC has been calcined at 800 $^\circ\text{C}$ for 1 h, the polysaccharide may form certain sugar crystals and after calcination, transforms into graphitic carbon. The peak near the 2θ theta of 12.7 has rarely been seen reported in literature, and we only found one study about a cellulose biomass membrane that also exhibited this peak.²⁵ We assume that this peak might be related to the sugar crystals



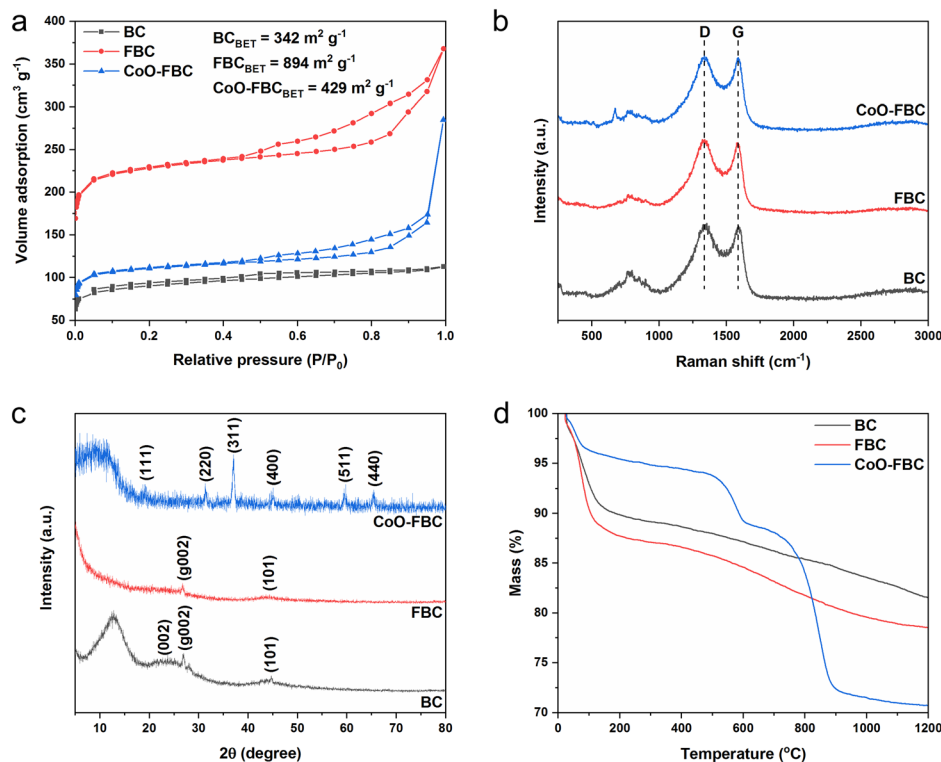
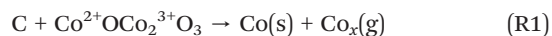


Fig. 2 BC, FBC and CoO-FBC characterizations of (a) nitrogen adsorption/desorption curve and the calculated BET surface area; (b) Raman spectrum, with a laser wavelength of 532 nm; (c) XRD spectrum, Cu-K α radiation, $\lambda = 1.5406 \text{ \AA}$; (d) TGA curves, $10 ^{\circ}\text{C min}^{-1}$, air flow rate of 30 mL min^{-1} combined with a nitrogen flow rate of 20 mL min^{-1} .

formed during the fabrication procedure. The FBC still has the diffraction peaks of (g002) and (101), but the peak of (002) become much less apparent when compared to that of BC. The mesopores formed may cause this diminished peak. The CoO-FBC was discovered to agree well with the pattern of Co_3O_4 (PDF-#43-1003), showing the characteristic diffraction peaks of (111), (220), (311), (400), (511) and (440). The thermal stability of the biomass carbon material was checked using TGA, as shown in Fig. 2d. In order to resolve better of the material degradation process *versus* temperature increment, a total flow of 60% air and 40% nitrogen (protective gas) at the flow rate of 50 mL min^{-1} was used in the experiment. During the initial heating stage of $50 ^{\circ}\text{C}$ to $100 ^{\circ}\text{C}$, the mass loss (about 10% loss) mainly came from the desorption of the physisorbed water. Starting from $200 ^{\circ}\text{C}$, the BC and FBC decomposed slowly and stably until $1200 ^{\circ}\text{C}$ was reached, and about 80% of the mass still remained. This mass loss came from the burning of the carbon material, and the 40% protective gas slows down the burning speed. Rather than a quick and fast mass loss by using pure air, the adoption of partial protective gas resolved the thermal degradation curve better.

As can be seen from the TGA curve of the CoO-FBC, the mass loss explicitly requires detailed inspection. In the preparation of the CoO-FBC, an equal weight of FBC and CoO were added for the preparation. The CoO cracked nanoplate was calcined at $500 ^{\circ}\text{C}$. Therefore, before $500 ^{\circ}\text{C}$, the mass loss of FBC was about 2.5

times higher than the mass loss of CoO-FBC. For example, the mass loss of FBC at $200 ^{\circ}\text{C}$, $300 ^{\circ}\text{C}$ and $400 ^{\circ}\text{C}$ was 12.28%, 12.91% and 13.41%, respectively, and the mass losses of CoO-FBC at $200 ^{\circ}\text{C}$, $300 ^{\circ}\text{C}$ and $400 ^{\circ}\text{C}$ were 4.59%, 5.19% and 5.56%, respectively. At this range, the major mass loss came from the FBC part of the CoO-FBC. At the range from $500 ^{\circ}\text{C}$ to $600 ^{\circ}\text{C}$, the CoO-FBC experienced a quick mass loss, and at the range from $700 ^{\circ}\text{C}$ to $900 ^{\circ}\text{C}$, the CoO-FBC experienced another quick mass loss. The cobalt oxide is regarded as Co_3O_4 from the XRD spectrum, the composition of which is considered to be $\text{Co}^{2+}\text{OCo}_2^{3+}\text{O}_3$. According to the mass loss calculation (Table S1†), the first mass loss is assumed to be from the reduction of the Co^{2+}O component by the carbon, and the second mass loss is assumed from the reduction of the $\text{Co}_2^{3+}\text{O}_3$ component by the carbon. The reaction equation is described as:²⁶



The reaction is achieved at higher temperature (over $600 ^{\circ}\text{C}$), but the nanomaterial condition of CoO may lower the reaction temperature to $600 ^{\circ}\text{C}$. The separate reduction reaction may also be caused by the different reaction activation energy. The partial protective gas and flow rate may also lead to such a separate reduction procedure. At $1200 ^{\circ}\text{C}$, the CoO-FBC has a mass percentage of about 70%.

The surface valence states of the characteristic elements of the biomass derived carbon materials were analysed using the



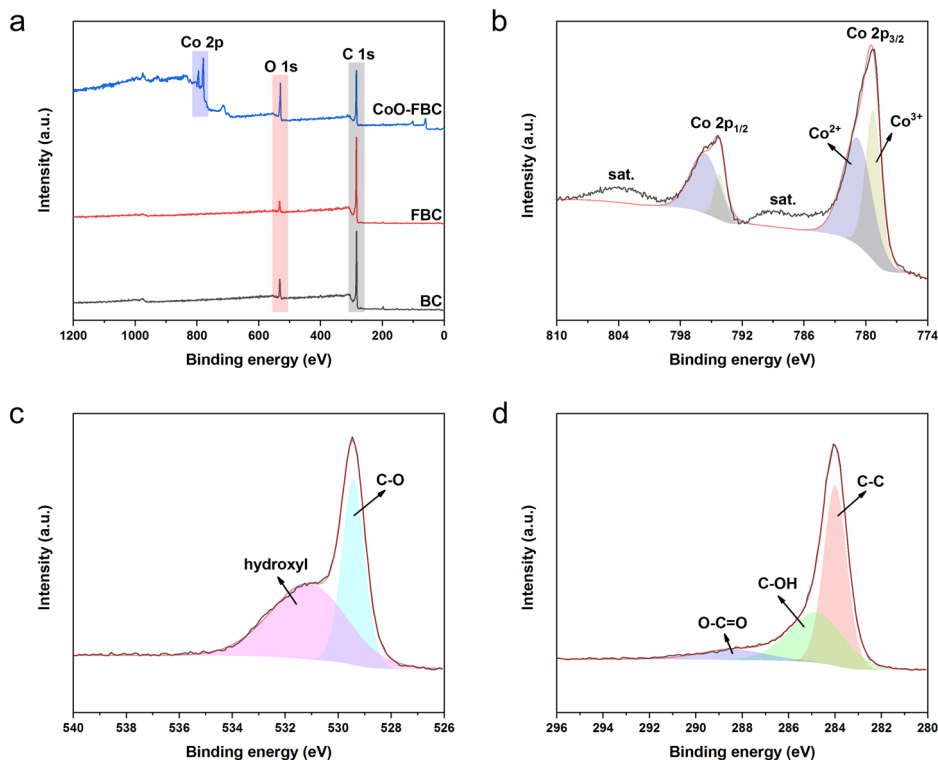


Fig. 3 XPS (a) survey scan spectra of BC, FBC and CoO-FBC; (b) Co 2p scan of CoO-FBC; (c) O 1s scan of CoO-FBC; (d) C 1s scan of CoO-FBC.

XPS measurements. Fig. 3a shows the XPS survey scans of the biomass carbon materials, and all of them share the peaks of C 1s and O 1s at 284 eV and 530 eV, respectively. The CoO-FBC additionally displays the Co 2p peaks at 780 eV and 795 eV. The elemental contents measured from XPS are listed in Table S2,[†] and the minute N element is assigned to the inherent element in the *E. angustifolia* gum. Whereas the minute chloride element is supposed to be a residual from the hydrochloride washing. Fig. 3b–d show the XPS spectrum of CoO-FBC. In Fig. 3b, the two Co 2p peaks could be deconvoluted into two peaks. The two fitted peaks for Co 2p_{3/2} were assigned to Co³⁺ (779 eV) and Co²⁺ (781 eV). The peak of Co 2p_{1/2} was assigned to Co³⁺ (794 eV) and Co²⁺ (796 eV). In the hydrothermal synthesis of cobalt hydroxide, the cobalt salt used is CoCl₂, then after the calcination at 500 °C, the cobalt hydroxide was dehydrated and became partially oxidized into Co₃O₄. Therefore, the Co²⁺ and Co³⁺ valence states were detected. This result is also in accordance with the XRD spectrum and TGA analysis. There were two satellite peaks centred at approximately 788 eV and 804 eV. These satellite peaks belong to the Co²⁺ oxidation state, and thus confirm the chemical nature of the CoO.²⁷

Fig. 3c displays the O 1s scan of CoO-FBC, where the peak is divided into two peaks at 529 eV and 531 eV. The 529 eV peak is related to the Co–O bond, whereas the 531 eV is related to the hydroxyl species. Fig. 3d shows the C 1s scan of the CoO-FBC sample, and the three deconvoluted peaks are at 284 eV, representing the C–C bond, at 284.9 eV, representing the C–OH bond, and at 288.6 eV, representing the O–C=O bond.^{28,29} It is assumed that after the sonication assisted incorporation of CoO

onto FBC, some of the ethanol molecule may interact with the cobalt oxide cracked nanoplate and cause such a signal. Also, the calcination at 500 °C may also leave some cobalt hydroxide in the nanoplate.

2.2 Electrochemical characterization

The biomass derived carbon materials are fabricated onto a GCE for electrochemical analysis. Fig. 4a shows the CV spectra of bare GCE and the fabricated GCE in [Fe(CN)₆]^{3–/4–} solution. The *I*_{pa} for bare GCE, BC/GCE, FBC/GCE and CoO-FBC/GCE were 78.73 μA, 89.25 μA, 97.68 μA and 95.74 μA, respectively. The unactivated BC/GCE CV curve resembles the bare GCE CV curve, but with a slightly increasing *I*_{pa}. The FBC/GCE shows an apparently larger CV area and an increasing *I*_{pa}. The enlarged CV area originates from the porous structure and the high surface area, whereas the CoO-FBC/GCE has a slightly smaller *I*_{pa} than that of FBC/GCE. The reason for this may be due to the coverage of the FBC surface by the cobalt oxide cracked nanoplate and decreasing of the material surface area.

The corresponding GCEs were also tested using EIS analysis as shown in Fig. 4b. The high-frequency region is zoomed in and shown in Fig. S4.[†] The charge transfer resistance *R*_{ct} of bare GCE, BC/GCE, FBC/GCE and CoO-FBC/GCE were simulated and calculated to be 710.6 ohm, 303.2 ohm, 8.1 ohm and 4.9 ohm, respectively. Although the BC is unactivated, the modification of the biomass carbon is still able to lower the interface resistance. The FBC/GCE and CoO-FBC/GCE have equivalent *R*_{ct} and these



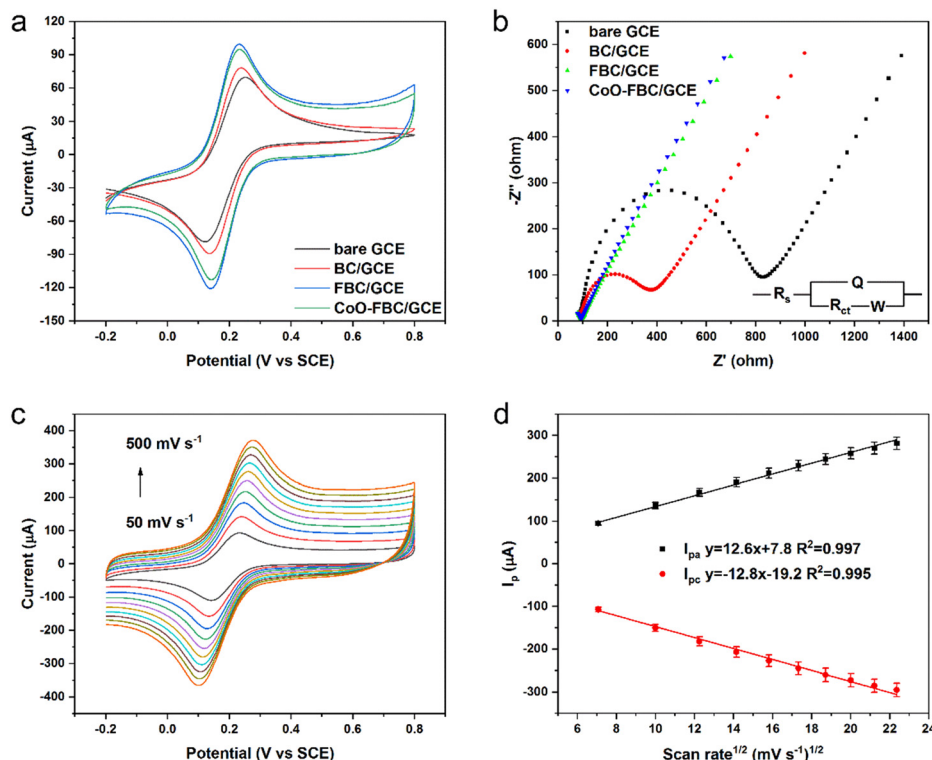


Fig. 4 (a) CV spectra of the bare GCE and biomass derived carbon electrocatalyst modified GCE in 5 mM $[\text{Fe}(\text{CN})_6]^{3-/4-}$, 0.1 M KCl, and a scan rate of 50 mV s^{-1} ; (b) EIS spectra of the bare GCE and biomass derived carbon electrocatalyst modified GCE in 5 mM $[\text{Fe}(\text{CN})_6]^{3-/4-}$, 0.1 M KCl (frequency range from 100 kHz to 0.1 Hz); (c) the CoO-FBC/GCE CV spectrum in 5 mM $[\text{Fe}(\text{CN})_6]^{3-/4-}$, 0.1 M KCl with a changing scan rate; (d) the linear fit of the I_{pa} and I_{pc} current versus the square root of the scan rate in (c).

are far smaller than those of the other two GCEs. The incorporation of the cobalt oxide cracked nanoplate did not affect the FBC's functionality too much. Although the nanoplate sits on the FBC surface, the hollow holes may still provide sufficient contact for the reactants in the solution when assessing the biomass carbon. As cobalt oxide is a semiconductor material, it showed a limited influence on the conductance of the CoO-FBC biomass carbon.

In Fig. 4c, the CV spectra of the CoO-FBC/GCE was obtained with varying scan rates (100 mV s^{-1} , 150 mV s^{-1} , 200 mV s^{-1} , 250 mV s^{-1} , 300 mV s^{-1} , 350 mV s^{-1} , 400 mV s^{-1} , 450 mV s^{-1} and 500 mV s^{-1}). The redox pair $[\text{Fe}(\text{CN})_6]^{3-/4-}$ showed the redox peaks' potential positions to be at about 0.23–0.28 V and at 0.10–0.14 V. As the scan rate became faster, the CV spectrum showed a larger redox potential difference and a larger response current. Fig. 4d is the linear fit relationship between the I_{pa} and I_{pc} currents and the square root of the scan rate in Fig. 4c, indicated that the redox reaction of $[\text{Fe}(\text{CN})_6]^{3-/4-}$ electrocatalyzed by CoO-FBC was a diffusion controlled process. The linear fit R^2 of both I_{pa} and I_{pc} were 0.997 and 0.994. The electrochemically active surface area of CoO-FBC was calculated by the Randles–Sevcik equation:³⁰

$$I_p = 2.69 \times 10^5 A \cdot D^{1/2} \cdot n^{3/2} \cdot \nu^{1/2} C \quad (1)$$

where I_p is the peak current (mA), A is the electrochemical active surface area (cm^2), D is the diffusion coefficient of the

molecule in the bulk solution ($6.67 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$ for potassium ferricyanide), n is the number of electrons that participated in the reaction, ν is the scan rate (V s^{-1}), C is the concentration of the bulk solution (M).

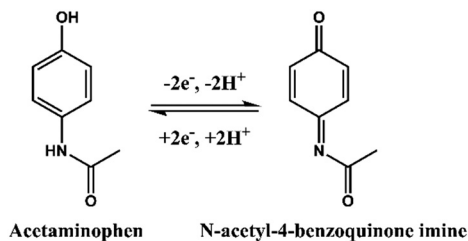
The electrochemically active surface area of the bare GCE, BC/GCE, FBC/GCE and CoO-FBC/GCE sensor towards $[\text{Fe}(\text{CN})_6]^{3-/4-}$ were calculated by an average value using both I_{pa} and I_{pc} to give 0.104 cm^2 , 0.117 cm^2 , 0.135 cm^2 , 0.130 cm^2 , respectively. Compared to the bare GCE, the modified GCEs all had a larger electrochemically active surface area. In particular, the CoO-FBC/GCE had a slightly decreased value than that of FBC/GCE, which was caused by the coverage of the cobalt oxide cracked nanoplate on FBC.

2.3 Electrochemical analysis of APAP

As shown in Scheme 1, the chemical formula of APAP is $\text{C}_8\text{H}_9\text{NO}_2$, and it can be electrochemically catalysed into *N*-acetyl-4-benzoquinone imine ($\text{C}_8\text{H}_7\text{NO}_2$) at the oxidation potential near to the 0.3 V to 0.4 V range, together with 2 hydrogen atoms and 2 electrons were lost. This process is reversible with *N*-acetyl-4-benzoquinone imine reduced back to APAP in the potential near to the 0.2 V to 0.3 V region.

To study the electrochemical response of APAP, the CV spectra of bare GCE and the fabricated GCE in APAP solution were measured as shown in Fig. 5a. Fig. S5† shows the CV spectra of the bare GCE and biomass derived carbon





Scheme 1 Electrochemical redox reaction of acetaminophen and N-acetyl-4-benzoquinone imine.

electrocatalyst modified GCE in 0.1 M PBS solution, at a scan rate of 50 mV s^{-1} . The CV areas slowly increased in the sequence of bare GCE, BC/GCE, FBC/GCE and CoO-FBC/GCE.

The I_{pa} for bare GCE, BC/GCE, FBC/GCE and CoO-FBC/GCE were $9.99 \mu\text{A}$, $10.62 \mu\text{A}$, $25.66 \mu\text{A}$ and $57.87 \mu\text{A}$, respectively. The cobalt atom is considered to possess a specially enhanced electrocatalytic interaction towards the APAP molecule, which explains the intensified I_{pa} current response.³¹

The cracks in the nanoplate increase the surface cobalt atom exposure to the APAP molecule. The cobalt oxide is a semiconductor and is able to transfer the electrons efficiently to the biomass carbon matrix. All of these contribute to the almost doubled I_{pa} current response compared to the I_{pa} current response of FBC/GCE. Fig. 5b and c show the changing scan rate of the CoO-FBC/GCE CV spectra and the linear fit relationship between the I_{pa} and I_{pc} current and the

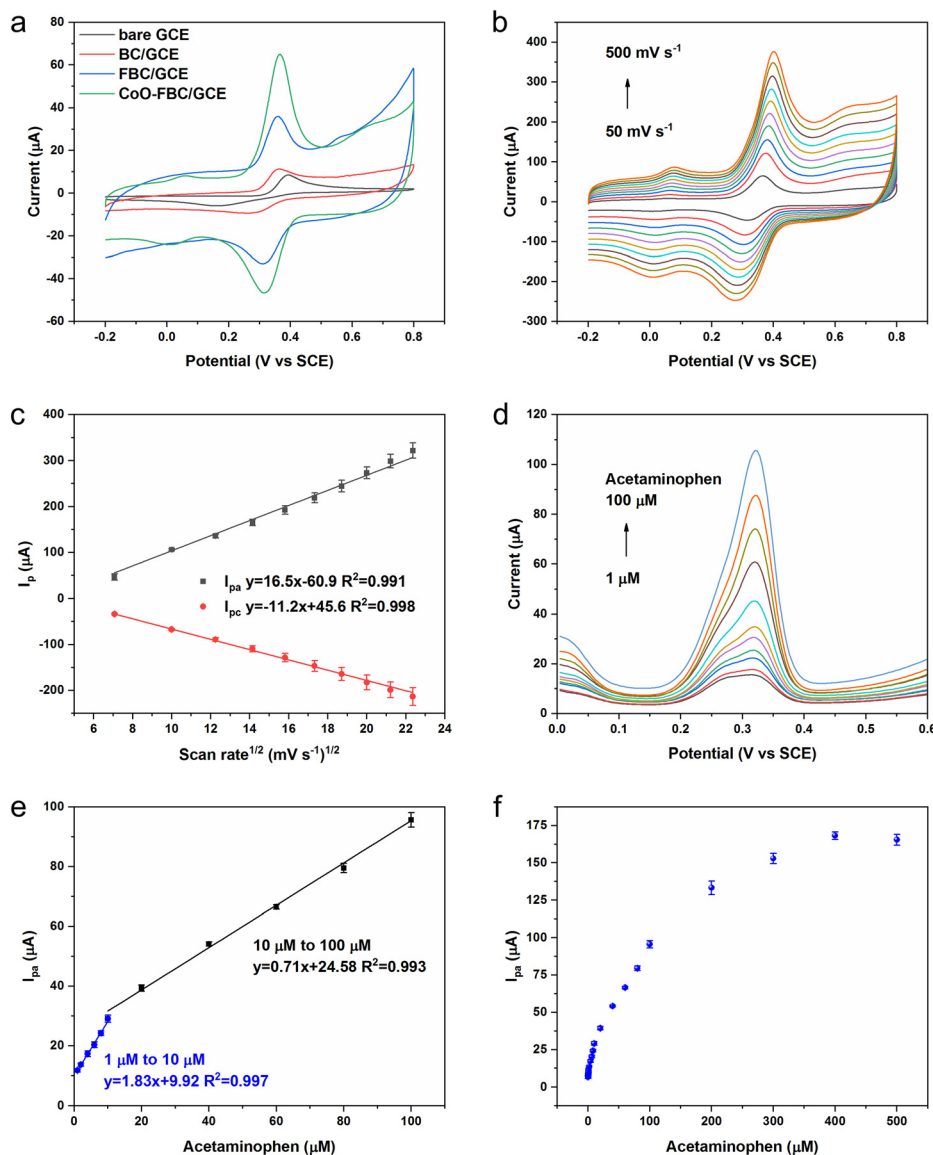


Fig. 5 (a) CV spectra of the bare GCE and biomass derived carbon electrocatalyst modified GCE in APAP solution ($500 \mu\text{M}$ in 0.1 M PBS , scan rate 50 mV s^{-1}); (b) CoO-FBC/GCE CV spectrum in $500 \mu\text{M}$ APAP in 0.1 M PBS with a changing scan rate; (c) the linear fit of the I_{pa} and I_{pc} current versus the square root of the scan rate in (b); (d) DPV measurement of APAP in 0.1 M PBS with varying concentrations of APAP from $1 \mu\text{M}$ to $100 \mu\text{M}$; (e) linear fit of DPV I_{pa} current versus the APAP concentrations in (d); (f) DPV I_{pa} current of APAP solution from $1 \mu\text{M}$ to $500 \mu\text{M}$.



square root of the scan rate in the APAP solution. Similar to the CoO-FBC/GCE CV scan in $[\text{Fe}(\text{CN})_6]^{3-/4-}$ solution, the CV scans of CoO-FBC/GCE in the APAP solution also show that the redox reaction of APAP electrocatalyzed by CoO-FBC is also a diffusion controlled process. Based on eqn (1), the electrochemically active surface area of the bare GCE, BC/GCE, FBC/GCE and CoO-FBC/GCE sensor towards APAP were calculated using the I_{pa} current response and found to be 0.013 cm^2 , 0.014 cm^2 , 0.033 cm^2 , 0.075 cm^2 , respectively. In Fig. 5b, there is a pair of redox peaks at about 0.01 V and 0.08 V, which are obvious at high scan rates. As discussed in the materials characterization section, the CoO is assumed to be Co_3O_4 (PDF-#43-1003). This redox peak is assumed to come from the oxidation and reduction from the Co^{2+} and Co^{3+} ions on the CoO nanoplate surface. There is also another oxidation peak of 0.65 V at high scan rates, which may be related to the $\text{Co}^{2+}/\text{Co}^{3+}$ redox process. The interaction of $\text{Co}^{2+}/\text{Co}^{3+}$ redox with the biochar may also form such peaks.^{32,33}

Fig. 5d and e display the DPV measurement of CoO-FBC/GCE towards various concentrations of APAP solutions. Two linear ranges from $1 \text{ }\mu\text{M}$ to $10 \text{ }\mu\text{M}$ and $10 \text{ }\mu\text{M}$ to $100 \text{ }\mu\text{M}$ were found, with a sensitivity of $25.89 \text{ }\mu\text{A } \mu\text{M}^{-1} \text{ cm}^{-2}$ and $10.04 \text{ }\mu\text{A } \mu\text{M}^{-1} \text{ cm}^{-2}$, respectively, with a limit of detection of $0.46 \text{ }\mu\text{M}$. In the low APAP concentrations, the APAP oxidation peak of 0.32 V is deconvoluted into a mixture of peaks near 0.28 V to 0.32 V. We assumed that this might also come from the electrocatalyst multicomponent nanostructures. There were CoO nanoplates and biomass carbon present in the electrocatalyst surface. The APAP molecule might be oxidized at the nanoplate surface and the biomass carbon surface, thus resulting in the mixture of peaks around the range of 0.28 V to 0.32 V. Also different catalytic capabilities of the multicomponents existed and would also contribute to such a phenomenon. Fig. 5f shows the I_{pa} current of the DPV measurement. As the APAP concentration increases from $100 \text{ }\mu\text{M}$ to $500 \text{ }\mu\text{M}$, the current response slowly became saturated and the $500 \text{ }\mu\text{M}$ APAP signal shows a slightly decreasing value. This slight saturation trend may be caused by the overwhelming number of APAP molecules at the

electrocatalyst surface, leading to the lowering of the electrocatalytic reaction.

As discussed in our previous study, the average male urine APAP concentration is recognized as $200 \text{ }\mu\text{M}$. The average male urine uric acid (UA) concentration is recognized to be $2000 \text{ }\mu\text{M}$. Uric acid is a frustrating interference when carrying out the electrochemical sensing of urine APAP concentrations.³⁴ Because urine also contains a number of other biomolecules and inorganic components, the use of diluted urine should alleviate the potential sensing noise. We would use male urine diluted 10 times for the real sample measurement and based on the 10 times dilution, we would choose certain chemical concentrations. In the selectivity experiments, we chose the interfering chemicals to be uric acid, ascorbic acid, urea and glucose using equivalent concentrations of $200 \text{ }\mu\text{M}$. The average uric acid in the 10 times diluted urine was set as $20 \text{ }\mu\text{M}$. The average uric acid in the 10 times diluted urine was set as $200 \text{ }\mu\text{M}$. The selectivity experiment used the simulated 10 times diluted urine that was a much cleaner solution, and the APAP concentration was set as $10 \text{ }\mu\text{M}$.

Fig. 6a shows the DPV curves for the selected interferences. The ascorbic acid, urea and glucose show no response in the potential range. The APAP molecule is oxidized at the potential near 0.31 V, whereas the uric acid molecule is oxidized at the potential near 0.25 V. The uric acid peak and the APAP peak could be separated distinctly in the all the interference mixtures (APAP, uric acid, ascorbic acid, urea and glucose) and the mixture of APAP and uric acid solution. The current response may have been affected a little, but the deviation can be corrected by reconstruction under the designed conditions and applications. Fig. 6b displays the DPV curve of the diluted urine samples spiked by APAP. The diluted urine provides a current response near 0.24 V, which is probably coming from the uric acid. The spiked APAP ($100 \text{ }\mu\text{M}$) urine sample exhibited two current responses, of which one is from the uric acid near 0.24 V, and the other one is from the APAP near 0.32 V. If the original urine APAP is in the range from

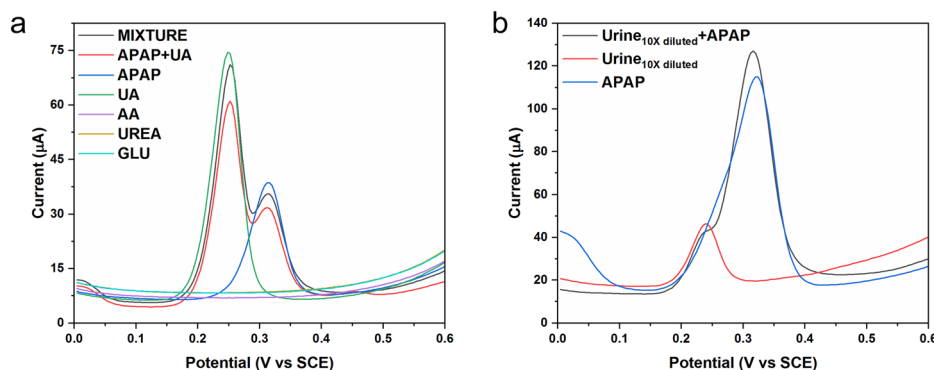


Fig. 6 (a) The selectivity DPV measurement of the CoO-FBC/GCE against interferences of uric acid (UA), ascorbic acid (AA), urea (UREA) and glucose (GLU), all the interference solutions concentration are $200 \text{ }\mu\text{M}$ in 0.1 M PBS, and the APAP concentration is $10 \text{ }\mu\text{M}$ in 0.1 M PBS; (b) the DPV measurement of the urine sample, where the urine is diluted 10 times in 0.1 M PBS, and the APAP concentration is $100 \text{ }\mu\text{M}$ in 0.1 M PBS.



100 μM to 1 mM, the designed CoO-FBC/GCE would be capable of detecting the 10 times diluted urine APAP. Considering the situation of abuse of APAP intake, and considering the uric acid interference, the CoO-FBC/GCE sensor could fulfil the needs of clinical diagnosis. Further improvements can be done to enhance the CoO-FBC/GCE performance. For example, the biomass activation procedure, the cobalt oxide cracked nanoplate size and morphology, the sonication step and the biomass carbon to cobalt oxide ratio, co-doping with other transitional elements by alkali co-precipitation and so on could be all be considered for an in-depth study to achieve sensors with superior capability.

The cycling experiment result is shown in Fig. S6.† After 10 continuous CV cycles, the CoO-FBC/GCE still obtained about 82.1% of its first sensing signal. Also, the electrocatalyst was stored in a general room environment for over 3 months, there was no negative effect on the use of the sensor, and because the major composition of the electrocatalyst is cobalt oxide and carbonized biochar, the sensor remains very stable. The modified CoO-FBC/GCE could recover a similar current signal when compared with the results of experiments carried out 3 months previously. We think the CoO-FBC/GCE sensor has good stability, reproducibility, repeatability.

2.4 Comparison of other works

A survey of related research on nanomaterials' electrochemical sensing of APAP is summarized in Table 1. From the recent relevant studies, we can see various nanomaterials have been used, including a series of carbonaceous materials such as carbon nanotubes, graphene oxide, cellulose, porous carbon nanorods and so on. In addition, carbon containing materials from the pyrolysis of metal organic frameworks, and a series of nanomaterials using gold nanoparticles, zinc oxide nanoparticles, transition element co-doped nanomaterial, bi-metallic transition metal alloy, cobalt-zinc metal organic framework derived material, quantum dots and so on, have been hybridized and utilized for electrochemical catalytic sensing of APAP in a number of applications. Compared to the cited

references, the biomass of *E. angustifolia* gum is a natural accessible abundant resource, and the hollowness of the cobalt oxide cracked nanoplate can be modulated by modifying the fabrication procedures. No complicated preparation steps are involved. The sensitivity of the CoO-FBC/GCE sensor is good. It is also worth mentioning that in this work, the uric acid response peak and the APAP response peak could be separated well and recognized in the rational physiological concentration range.

3 Conclusions

We have developed a novel cobalt oxide cracked nanoplate decorated on biomass derived carbon electrocatalyst material for electrochemical sensing of APAP. The biomass derived carbon is prepared by the ferric chloride activation of *E. angustifolia* gum, a major composition of which is polysaccharides. The cobalt oxide cracked nanoplate is synthesized by alkali precipitation of cobalt chloride in aqueous solution and then by calcination. Simple sonication assisted hybridization of biomass carbon and cobalt oxide cracked nanoplate is adopted. The prepared CoO-FBC/GCE exhibits two linear sensing range from 1 μM to 10 μM and 10 μM to 100 μM , with sensitivities of 25.89 $\mu\text{A } \mu\text{M}^{-1} \text{ cm}^{-2}$ and 10.04 $\mu\text{A } \mu\text{M}^{-1} \text{ cm}^{-2}$, and a limit of detection of 0.46 μM . This CoO-FBC electrocatalyst is sufficient for sensing 10 times diluted APAP spiked urine samples and shows promise for integration into a disposable sensor for clinical diagnosis.

4 Experimental

4.1 Chemicals

All the chemicals used were of analytical grade and used as received. Acetaminophen and phosphate buffered saline tablets are from Aladdin. The PBS solution used throughout the experiments is prepared using the PBS tablets. Ferric chloride hexahydrate ($\text{FeCl}_3 \cdot 6\text{H}_2\text{O}$) and potassium ferricyanide ($\text{K}_3\text{Fe}(\text{CN})_6$) are from the Tianjin Baishi Chemical Co., Ltd. The cobalt chloride (CoCl_2) is from Alfa Aesar. Sodium hydroxide (NaOH) and hydrogen chloride (HCl) are from Tianjin Xinbote Chemical Co., Ltd. Urea is from the

Table 1 Comparison with other research on using nanomaterials for electrochemical sensing of APAP

Electrocatalyst	Sensing range (μM)	Limit of detection (nM)	Sensitivity ($\mu\text{A } \mu\text{M}^{-1} \text{ cm}^{-2}$)	Applications	Ref.
COF@AuNPs@MWCNTs	0.1 to 500, 500 to 1200	80	0.72, 0.30	Human serum	35
Gd ₂ ZnMnO ₆ /ZnO	0.090 to 900	25	2.75, 0.38	Human urine, plasma and pharmaceuticals	36
FeM@porous carbon nanorod	0.001 to 0.8	0.59	540.70	Pharmaceutics	37
ZnONPs/SBA-15	4 to 32	110	0.69	Pharmaceutics	38
CoNC@rGO	0.5 to 50	67	8.46	Environmental water	39
GO/MnO ₂ : h-MoO ₃	0.06 to 10, 20 to 80	13.3	28.11, 18.27	Pharmaceutics	40
Laser-induced rGO	0.099 to 1987	5.2	2.73	River water, pharmaceuticals and human urine	41
MWCNT and poly(neutral red)	2 to 70	15	56.40	Pharmaceutics	42
Ag-Fe ₃ O ₄ /nanocellulose	0.05 to 15	8	152.31	Human blood and urine	43
Ti ₃ C ₂ QDs/Fe-NC	0.50 to 210	30	6.53	Pharmaceutics and lake water	44
CoO-FBC	1 to 10, 10 to 100	460	10.04, 25.89	Human urine	This work



Tianjin Tianda Fine Chemical Plant. Potassium chloride is from Tianjin Bodi Chemical Co., Ltd. Glucose and potassium ferrocyanide trihydrate ($K_4Fe(CN)_6 \cdot 3H_2O$) are from the Tianjin Hedong District Hongyan Reagent Factory. Uric acid and ascorbic acid are from Shanghai Macklin Biochemical Technology Co., Ltd. The GCE, platinum wire electrode, platinum plate electrode, saturated calomel electrode (SCE) and aluminium slurry are from Wuhan Gaoss Union Technology Co., Ltd. All the aqueous solutions were prepared with water (18.2 M Ω cm) from a Millipore system. Dried *E. angustifolia* gum was provided by the Key Laboratory of Chemistry of Plant Resources in Arid Regions, and the State Key Laboratory Basis of Xinjiang Indigenous Medicinal Plants Resource Utilization. The human urine sample was provided by a healthy male volunteer in the laboratory, who did not consume any medicine containing APAP for at least a month before providing the urine.

4.2 Preparation procedures

Cobalt oxide preparation. Cobalt chloride (15 mmol) was dissolved in 30 mL of water and mixed well, and then sodium hydroxide (30 mmol) dissolved in 30 mL of water was then dropped into the cobalt chloride solution with vigorous stirring. The mixture was stirred for 2 h and then placed in the oven and heated at 80 °C for 5 h. The precipitate was purified and collected by multiple washes using water and centrifugation. The precipitate was dried in an oven and heated at 50 °C for 12 h. The product was further calcined in a muffle furnace (500 °C, 3 h, 5 °C min⁻¹) to obtain the cobalt oxide.

BC preparation. *E. angustifolia* gum (1 g) was dissolved in 10 mL of water and stirred for 2 h. Then the mixture was dried in an oven at 60 °C until fully dried. The dried biomass was carbonized at 800 °C for 3 h with a heating rate of 5 °C min⁻¹ under N₂ flow. The carbonized product was washed with HCl solution (15 wt%), purified using water, and dried in a vacuum oven at 60 °C for 24 h to obtain the BC.

FBC preparation. Ferric chloride hexahydrate (2 g) was dissolved in 10 mL of water. Then *E. angustifolia* gum (1 g) was added into the solution and mixed well. Then the mixture was dried in an oven at 60 °C until fully dried. The dried mixture was carbonized at 800 °C for 3 h with a heating rate of 5 °C min⁻¹ under N₂ flow. The carbonized product was washed with HCl solution (15 wt%), purified using water, and dried in a vacuum oven at 60 °C for 24 h to obtain the FBC.

CoO-FBC preparation. FBC (200 mg) was added to 20 mL ethanol and mixed by sonication for 1 h. Then cobalt oxide (200 mg) was added into the mixture and mixed by sonication for another 5 h. The product was dried in an oven (80 °C) to remove ethanol before calcination in muffle furnace (120 °C, 4 h, 5 °C min⁻¹).

4.3 Fabrication of GCE

Before modification, the GCE (3 mm in diameter) was polished with 0.5 μ m and 0.05 μ m alumina slurries to obtain a mirror-like surface. The electrode was then cleaned in ethanol and

water by sonication and dried in an ambient environment. The corresponding electrocatalyst aqueous solution (2 mg mL⁻¹) was sonicated for 30 min to form a homogeneous solution, and then 5 μ L of the solution was dropped onto the GCE surface and dried in an ambient environment.

4.4 Instruments

A field-emission scanning electron microscope (Zeiss MERLIN VP Compact, Germany) equipped with an energy-dispersive X-ray spectrometer (Oxford Instruments X-Max, UK) was used to characterize the morphology, elemental composition and elemental mapping of the electrocatalyst. X-ray photoelectron spectroscopy (Thermo Scientific K-Alpha, USA) was used to perform the surface analysis of the electrocatalyst. Raman spectra of the samples were obtained using a spectrometer (Horiba LabRAM HR800, France). The thermogravimetric analysis was performed using a Netzsch STA 449 F5 Jupiter, Germany and a Thermo Fisher Scientific Nicolet iS50 FTIR spectrometer, USA. The X-ray powder diffraction measurements were recorded on a diffractometer (Bruker D8 Advance, Germany, Cu-K α radiation). The nitrogen adsorption/desorption analysis was carried out using a Quantachrome Autosorb IQ, USA.

An electrochemical workstation (CHI-760E, Shanghai, Chenhua, China) and a workstation (RST 5000C, Zhengzhou Shiruisi Instrument Technology Co., Ltd, Zhengzhou, China) were used in the electrochemical experiments. A three-electrode system was used for the CV and DPV scans, with an SCE, a platinum wire electrode, and GCE. For the EIS measurements, a three-electrode system of an SCE electrode, platinum plate electrode (10 mm $\times$ 10 mm $\times$ 0.1 mm), and a GCE were used.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts to declare.

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