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Hybrid poly(allylamine hydrochloride)–graphene oxide microcapsules: preparation, characterization and application in textiles with controlled release behavior†

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Permeable microcapsules are suitable forming blocks for functional materials based on controlled release. This study reports the design of a stimuli-responsive fabric coated with a type of hybrid microcapsules prepared via a layer-by-layer (LBL) approach. The resulted fabric can be used as a healthy care material. The functional microcapsules comprise polyelectrolytes poly(allylamine hydrochloride) (PAH) and graphene oxide (GO). The self-assembly of these oppositely charged surrounding components generated a hollow space to load functional molecules such as dyes, labels and drugs via a responsive trigger. The polyelectrolyte PAH ensured a robust structure and pH sensibility. GO sheets provided near-infrared (NIR) laser-induced release through photothermal effects. These hybrid microcapsules (PAH/GO)_nPAH can then be used to load *Atractylodes*, a traditional Chinese herb, and then adhere to cotton fabrics to develop functional textiles. The encapsulated *Atractylodes* can then be released from the (PAH/GO)_nPAH-coated cotton fabric. During the subsequent period of the entire process, the release rate was significantly accelerated by NIR irradiation. After washing 20–30 times, microcapsules were coated on cotton fibers. Using this technique, functional textiles for controlled release by external stimulation coupled with other synergetic effects could be obtained. Note that (PAH/GO)_nPAH microcapsules can provide sustained drug release in simulated sweat and control the release rate through NIR irradiation, which shows its potential as an external skin drug delivery system.

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1 Introduction

Microencapsulation has already been considered as an efficient technique for generating materials with a long-lasting behavior. Microcapsules usually exhibit optical or pH response due to their structural and compositional properties.^{1–4} In recent years, stimuli-responsive microcapsules have been rapidly developed. Sukhorukov *et al.* reported that UV light-responsive microcapsules can achieve the remote-controlled release of embedded materials only by external UV stimulation without direct contact or interaction. In addition to single-responsive microcapsules, multi-responsive microcapsules have drawn considerable attention.⁵ Sukhorukov *et al.* designed

a new type of inorganic/organic hybrid capsule with unique physicochemical properties for multi-conditional responsiveness to physical (ultraviolet, ultrasonic) and chemical (enzymatic treatment) stimuli. Different stimulations could be used for the controlled release of numerous substances.⁶ As reported by Willner *et al.*, a method for constructing stimuli-responsive DNA-acrylamide-based hydrogel microcapsules was produced. Preliminary studies represented that these ATP- or pH-responsive microcapsule-encapsulated doxorubicin expressed a selective cytotoxic effect on MDA-MB-231 cancer cells.⁷ In our study, we used superparamagnetic iron oxide (SPIO) and PAH to form magnetic microcapsules with low magnetization used in MRI contrast tracking, hyperthermia and drug delivery.⁸

Usually, microcapsules work as a carrier for active ingredients with releasing behavior under environmental stimulation, which can be considered as a blocking part of a stimuli-responsive material. Stimuli-responsive textile fabrics have emerged as an important aspect in functional materials, thus yielding a myriad of applications such as textile sensors, self-repairing materials and drug delivery materials.^{9–13} Different stimuli factors can be used to drive responsive textiles, including temperature, magnetic field, pH and electric field.^{14,15}

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Microcapsules sealing various substances within small vectors can be easily combined with textile fibers, which have been studied in several areas in terms of controlled release and delivery properties.^{16,17} The combination of microcapsules with fabrics is a feasible method to introduce functional chemicals into textiles.^{18–20} By changing the surrounding environment (e.g. light, pH or temperature), the functional chemicals would release the carrier that bonded with textiles. Graphene oxide (GO) is a superior carbon nanomaterial that acted as the blocking part of a carrier for controlled release due to its responsiveness to near infrared (NIR) laser. Compared with other composite structures, hollow microcapsules of GO exhibit much higher loading amount because the advanced sheet-like structure of GO stabilizes the capsule shell, thus preventing the leakage of loaded substances.^{21–23} Therefore, functional composites based on graphene coupled with other materials such as polymers have been extensively developed.^{24,25} Besides, it is necessary to study the stimuli-responsive textile which combined with GO composites. Microcapsules with pH- and NIR-responsiveness can be applied on a pure cotton fabric by coating and can become suitable materials for direct contact with human skin because a weakly acidic solution in sweat can induce the release of chemicals out of microcapsules.^{26,27} Simultaneously, NIR radiation can be considered as the external stimulation to trigger the remotely controlled release behavior of certain microcapsules coated on textiles.^{28,29}

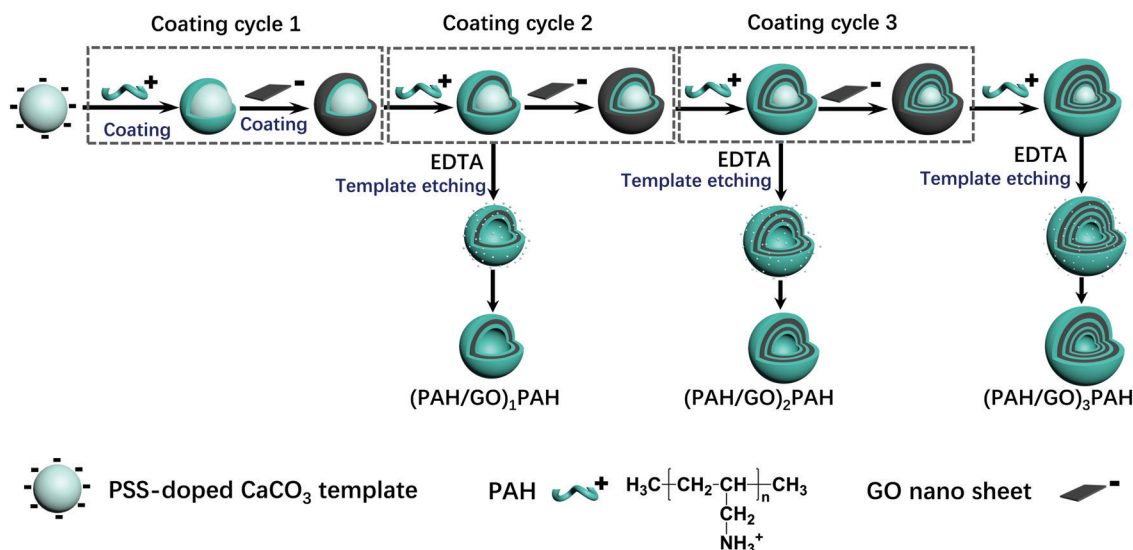
Atractylodes, beneficial for invigorating the spleen and eliminating dampness, is extensively distributed in China. As a type of Chinese traditional medicine, *Atractylodes* has been extensively used for treating rheumatic diseases, digestive disorders, hepatic protection and influenza.³⁰ The major ingredient effectively works for releasing pain from arthritis.³¹ Besides, it is reported that *Atractylodes* has pharmaceutical activities such as anticancer activities, activities on nervous, activities on gastrointestinal systems and activities on cardiovascular system.³²

Furthermore, *Atractylodes* is a suitable ingredient which can be loaded into microcapsules for controlled release. This study presents a type of self-assembled hybrid microcapsules comprising poly(allylamine hydrochloride) (PAH) and GO nanosheets using a layer-by-layer technique. Under NIR irradiation, these hybrid microcapsules can be easily coated on cotton fabrics to manufacture functional textiles with a controlled releasing behavior. The GO nanosheet is considered as a functional layer between polyelectrolytes layers due to its responsiveness to NIR. This microcapsule-coated fabric can be envisioned as a functional textile for healthy care treatment. Importantly, these microcapsule-coated cotton fabrics can be used as a type of responsive NIR-triggered textile with the controlled release of *Atractylodes* for treating both rheumatism and arthritis. Combining photothermal effect and remotely controlled release by an external stimuli factor, i.e., NIR laser, this *Atractylodes*-microcapsule-coated fabric generated an innovative functional textile for health care treatment.

2 Experimental

2.1 Materials

Poly(allylamine hydrochloride) (PAH) and poly(styrene sulfonic acid) sodium salt (PSS $M_w \approx 70$ kDa) were purchased from Alfa Aesar (Tianjin) Co., Ltd. Graphene oxide (GO) was purchased from Nanjing Pioneer Nanomaterials Co. Ltd. *Atractylodes* was purchased as a herbal medicine from Tianjin Yongchun Pharmacy and ground into a powder before use. An epoxyethane quaternary ammonium salt was used as a crosslinker. The other reagents were of analytical grade. Standard 100% cotton fabric (weight = 106.6 g m⁻²; warp = 133 yarns per inch; weft = 72 yarns per inch; thickness = 0.21 mm) was purchased from Tianyi printing and dyeing company in Tianjin, China. A hemocytometer was purchased from Changde BKMAM Biotechnology Co., Ltd.



Scheme 1 Schematic of (PAH/GO)_nPAH microcapsules prepared via a layer-by-layer assembly.



2.2 Preparation of graphene oxide nanosheets

Graphene oxide (GO) was dispersed in distilled water (1.0 mg mL^{-1}) under ultrasonication for 10–15 min for three times, followed by the addition of 500 mg NaOH. The mixture was maintained at 45°C for 4 h under stirring. Then, the solution was sonicated using an ultrasonic homogenizer system (JY92-IIN, Ningbo Scientz Biotechnology Co. Ltd) for 2 h. The parameters were set as 70% of output, 2 s of “operation on” and 4 s of “operation off”. Subsequently, the suspension was neutralized by dialysis (3500 Da) against deionized water for 48 h with frequent water-related changes. Finally, the neutralized GO suspension was sonicated for 2 h, followed by centrifugation, and the resultant nano GO solution was stored at 4°C for further use.

2.3 Preparation of (PAH/GO)_nPAH microcapsules

$\text{Ca}(\text{NO}_3)_2 \cdot 4\text{H}_2\text{O}$ solution (100 mL, 0.025 M) comprising 200 mg of PSS ($M_w \approx 70 \text{ kDa}$) was quickly poured into Na_2CO_3 (100 mL, 0.025 M) under stirring. After 15–30 min, the precipitated CaCO_3 particles were washed and collected. CaCO_3 templates were then dispersed into the PAH solution (10 mL, 1.0 mg mL^{-1} , and 0.5 M NaCl) for 15 min under continuous stirring (120 rpm). The resultant particles were washed three times and collected using centrifugation (8000 rpm for 1 min). Then, the coated particles were incubated in GO solution (10 mL, 0.1 mg mL^{-1}) using the same procedure. The multilayer structure was formed by the alternative assembly of corresponding materials, and the hollow hybrid microcapsules were obtained by etching CaCO_3 templates using EDTA (30 mL, 0.1 M, pH 7.0). The process is shown in Scheme 1.

2.4 Characterization of (PAH/GO)_nPAH microcapsules

The TEM images of (PAH/GO)_nPAH microcapsules samples were recorded using a transmission electron microscope (TEM) (Hitachi H7650, HITACHI). The samples for the TEM analysis were prepared by dropping a diluted microcapsule solution onto the standard 400-mesh TEM carbon-coated Cu grid, and then the solvent was evaporated at room temperature before imaging. The morphology of (PAH/GO)_nPAH microcapsules was characterized by a scanning electron microscope (SEM) (Hitachi S4800, HITACHI) under an

accelerating voltage of 20 kV. Moreover, the particle size of PSS-doped CaCO_3 particles was measured using a laser diffraction analyzer (LA-300, HORIBA). Raman spectra were recorded using a Raman microscope with a Renishaw CCD detector (British Renishaw Company). Fourier transform infrared (FTIR) spectroscopy was performed on a Nicolet iS 50 (ThermoFisher Technology) spectrometer to obtain the vibration modes of functional groups within $400\text{--}4000 \text{ cm}^{-1}$.

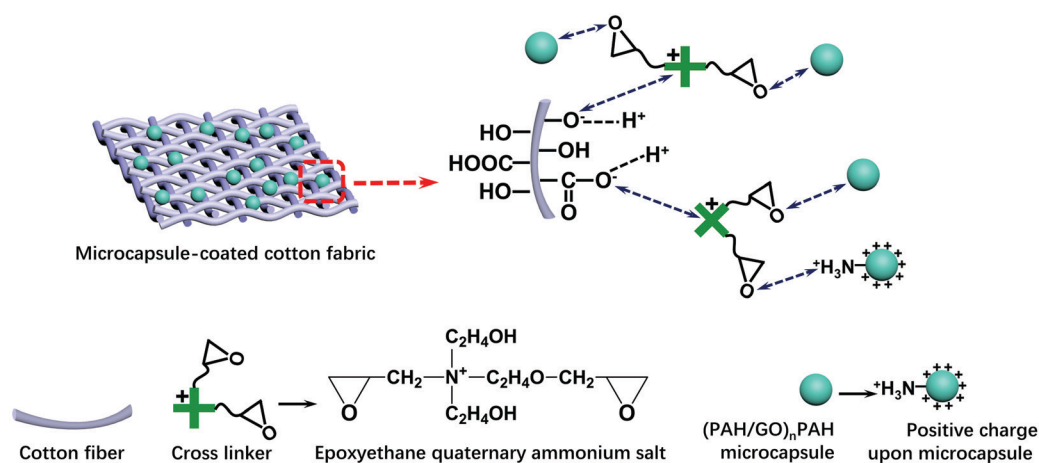
2.5 NIR-triggered RhB release from (PAH/GO)_nPAH microcapsules

(PAH/GO)_nPAH microcapsules (2 mg mL^{-1} , 1.0×10^8) were then washed and centrifuged three times ($7000 \text{ rpm min}^{-1}$, 1 min). Then, the washed microcapsules were incubated in a Rhodamine B (RhB) solution (2 mL, 1 mg mL^{-1}) at 25°C for 24 h. The resultant microcapsules were centrifuged and washed with PBS (pH 7.4) to calculate the loading efficiency (eqn (1-1)). The release experiment of (PAH/GO)_nPAH microcapsules was performed in two buffers (pH 5.8 and pH 7.4), and the release with the intermittent treatment by NIR laser was tested. The NIR laser was purchased from Changchun New Industries Optoelectronics Technology Co., Ltd (PAH/GO)_nPAH microcapsules, which were irradiated with a NIR laser of 808 nm at 0.5 W cm^{-2} for 1 min. The release behavior was as follows: RhB-loaded microcapsules ($\sim 1.0 \times 10^8$) were mixed with PBS (5 mL, pH 5.8 and 7.4) and 3 mL of supernatant was obtained at different time intervals for absorbance measurement ($\lambda = 554 \text{ nm}$, UV-1200 spectrophotometer, Mapada Instrument Co., Ltd), and then the supernatant was poured back into the initial solution. Finally, the cumulative release was obtained (eqn (1-2)).

$$\text{Loading efficiency (\%)} = \frac{C_0 V_0 - C_1 V_1}{C_0 V_0} \times 100\% \quad (1-1)$$

$$\text{Release rate (\%)} = \frac{C_t V_t}{C_0 V_0 - C_1 V_1} \times 100\% \quad (1-2)$$

where C_0 (mg mL^{-1}) and V_0 (mL) are the initial concentration and volume of the RhB solution; C_1 (mg mL^{-1}) and V_1 (mL) are the concentration and volume of the RhB solution after the incubation of microcapsules; and C_t (mg mL^{-1}) and V_t (mL) are the concentration and volume of the RhB solution at different time intervals

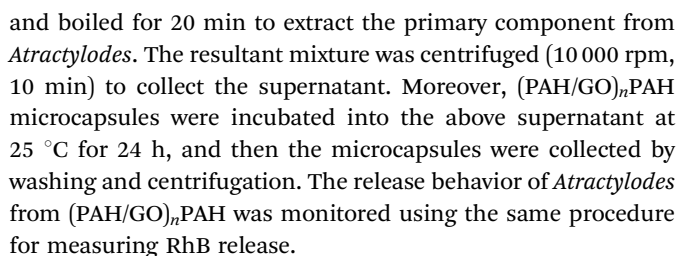


Scheme 2 Combination of (PAH/GO)_nPAH microcapsules with cotton fabric through an epoxyethane quaternary ammonium salt as cross linkers.



bind microcapsules with cotton fabrics. Scheme 2 shows the combination of (PAH/GO)_nPAH microcapsules with cotton fabrics using a cross-linker. The morphologies of microcapsule-coated cotton fabrics were characterized, and the coating percentage of the microcapsule-coated fabrics was calculated as follows:

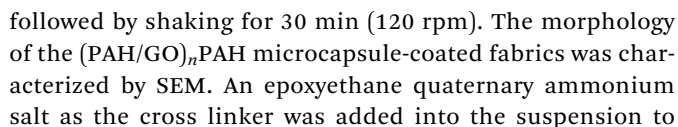
For the release behavior of *Atractylodes*, the powder of *Atractylodes* was immersed in deionized water (250 mL, 20 mg mL⁻¹)


$$\text{Coating percentage (\%)} = \frac{N_0 - N_1}{N_0} \times 100\% \quad (1-3)$$

where N_0 is the initial microcapsules number added into the system, and N_i is the microcapsules number in the system after coating on the cotton fabric. The number of microcapsules was calculated using hemocytometers.

The positively charged part NR_4^+ of the cross linker was bonded with $-\text{COO}^-$ ($-\text{COOH}$) or $-\text{O}^-$ ($-\text{OH}$) groups of cotton fiber, and the reaction equations are as follows:

The ethylene oxide groups interacted with the -NH_3^+ group of the outermost layer PAH of the $(\text{PAH/GO})_2\text{PAH}$



microcapsules. The following reaction (eqn (1-5)) is the formation of the secondary amine and hydroxyl group between epoxyethane and primary amine.³³ The reaction equation is as follows:



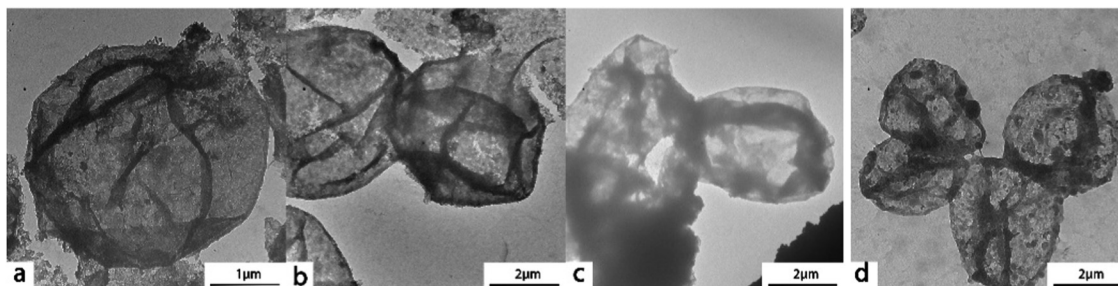


Fig. 1 TEM images of (PAH/GO)₁PAH (a and b), (PAH/GO)₂PAH (c), (PAH/GO)₃PAH (d).

Furthermore, the resultant secondary amine can react with ethylene oxide *via* a similar route, and the reaction is shown as follows: (PAH/GO)_nPAH-coated cotton fabrics were washed with 100 mL deionized water (30 °C, 10 min) for 5, 10, and 20 times to investigate the washing fastness of the coated fabrics. A piece of the coated fabric was fixed on the Al post using a conductive carbon tape, and Au and Pd were sputtered to make a conductive coating for SEM testing under an accelerating voltage of

20 kV. The coating effect of microcapsules on cotton fabrics was measured *via* FTIR spectroscopy.

3 Results and discussion

3.1 Morphology of (PAH/GO)_nPAH microcapsules

The size of (PAH/GO)_nPAH microcapsules depends on the size of PSS-doped CaCO₃ templates. The size distribution histograms

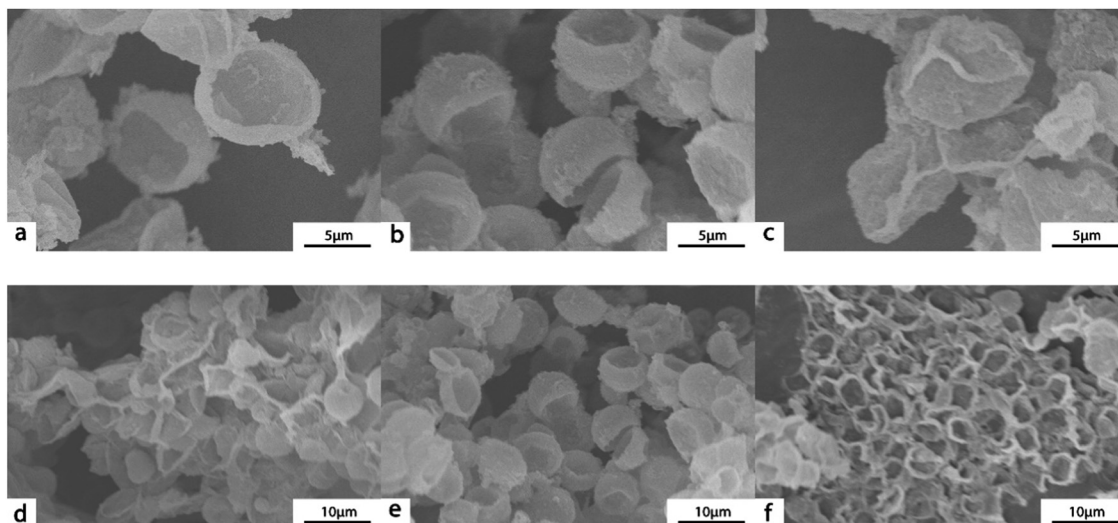


Fig. 2 SEM images of (PAH/GO)₁PAH (a and d), (PAH/GO)₂PAH (b and e), (PAH/GO)₃PAH (c and f).

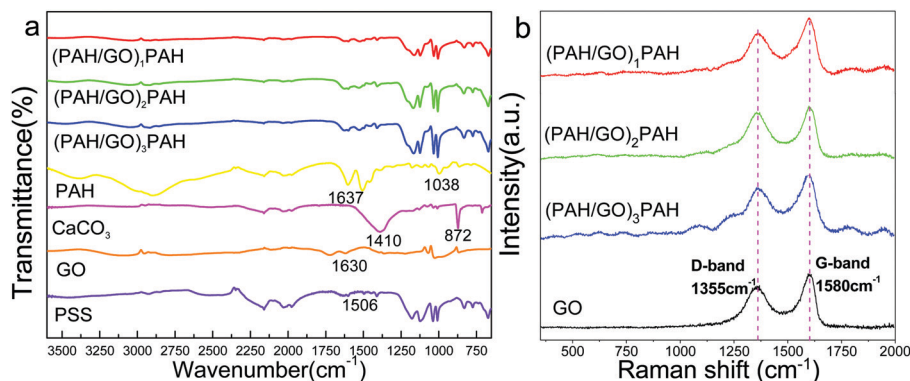


Fig. 3 FTIR spectra (a) and Raman spectra (b) of (PAH/GO)_nPAH microcapsules.



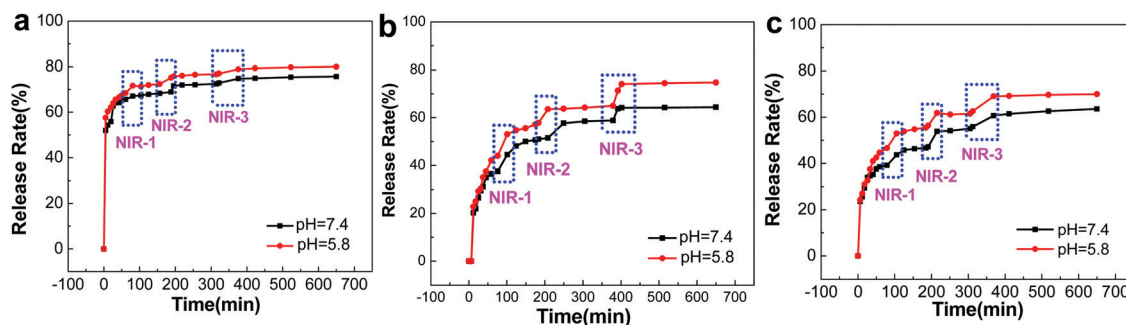


Fig. 4 NIR-triggered RhB release from (PAH/GO)₁PAH (a), (PAH/GO)₂PAH (b) and (PAH/GO)₃PAH (c) microcapsules. Each irradiation (blue zones) is with 808 nm NIR laser at 0.5 W cm^{-2} for 1 min.

(Fig. S1, ESI[†]) demonstrated that the diameter of the CaCO_3 template ranged from $5 \mu\text{m}$ to $7 \mu\text{m}$. These monodispersed spherical PSS-doped CaCO_3 particles with a negative charge were suitable as sacrificial templates for forming microcapsules *via* the LBL assembly. The morphology of (PAH/GO)_nPAH has been shown in TEM images (Fig. 1). Microcapsules with different layers were prepared with the diameter of about $\sim 5 \mu\text{m}$. The central CaCO_3 templates were efficiently etched, generating a hollow structure as vectors for loading either drugs or dyes. Note that the SEM images confirmed the evidence of hollow structure through the visible collapse in the shell of the microcapsules during the drying process (Fig. 2). Furthermore, the diameter of microcapsules indicated by SEM images was $\sim 5 \mu\text{m}$, in agreement with the TEM results.

3.2 Structure of (PAH/GO)_nPAH microcapsules

The FTIR spectra of (PAH/GO)₁PAH, (PAH/GO)₂PAH, (PAH/GO)₃PAH, PAH, CaCO_3 , GO and PSS are shown in Fig. 3a. The peaks located at $3540\text{--}3125 \text{ cm}^{-1}$ were assigned to the N-H group. Similarly, the peaks at $1690\text{--}1620 \text{ cm}^{-1}$ were attributed to the stretching vibration of C=O ; they suggested supramolecular interaction between the COO^- of GO and NH_3^+ of PAH, leading to the formation of “salt-bridge”. A strong absorption at 872 cm^{-1} corresponded to CO_3^{2-} , while the peak observed at 1410 cm^{-1} was attributed to the stretching vibration of C-O.³⁴ The characteristic bands of PSS at 1038 cm^{-1} and 1123 cm^{-1}

were attributed to the S-O stretching vibration deformation and symmetric stretching of S=O groups, respectively.³⁵ The peaks at 2923 cm^{-1} were assigned to the asymmetric vibration of CH_2 groups in PAH-Cl. The presence of protonated NH_3^+ groups in PAH-Cl was confirmed by bands due to the asymmetric vibration of these species, which can be observed at 1612 cm^{-1} .³⁶ The bands associated with oxygen-containing functional groups in GO at 1720 , 1401 , and 1070 cm^{-1} were particularly assigned to the vibration of C=O in carboxylic acid/or the carbonyl, C-O-H and C-O groups.³⁷ GO is hydrophilic because of the presence of polar oxygen-containing groups. The band observed at 1621 cm^{-1} was the stretching vibration of C=C . These peaks indicated efficient interaction between PAH and GO, generating a hollow structure of PAH/GO microcapsules. Furthermore, the Raman shift peaks are observed at 1355 cm^{-1} and 1594 cm^{-1} , which represented the D-band and G-band of graphene, confirming the presence of GO in PAH/GO microcapsules (Fig. 3b). Here, the D-band was attributed to the structural disorder and defects induced on the sp^2 hybridization, while the G-band represented the planar configuration of the sp^2 bonded carbon that constitutes GO.^{38,39} The peaks at $1000\text{--}1100 \text{ cm}^{-1}$ can be assigned to the S=O stretching on PSS.⁴⁰

3.3 NIR-triggered RhB release from (PAH/GO)_nPAH microcapsules

A significant advantage of GO-based microcapsules is that due to the unique optical properties of GO, the release of chemicals

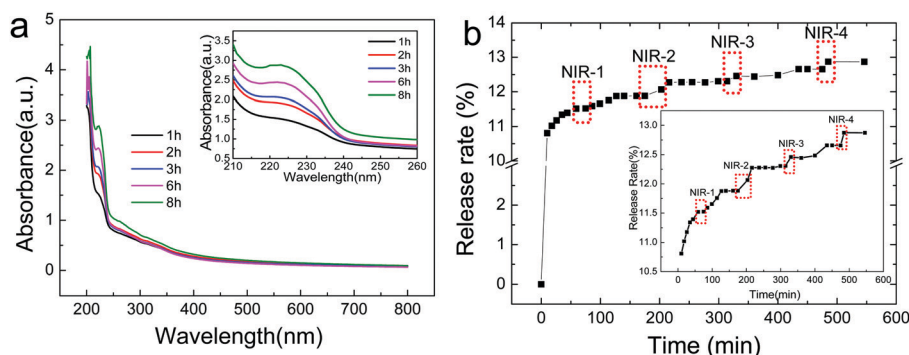


Fig. 5 Release of *Atractylodes* from (PAH/GO)₂PAH microcapsules at pH = 7.4 buffer. (a) UV-visible spectra of *Atractylodes* releasing from microcapsules with an inset of magnified image for peaks at 222 nm . (b) NIR-triggered *Atractylodes* release from (PAH/GO)₂PAH microcapsules with a calibration inlet. Each irradiation (red zones) is with the 808 nm NIR laser at 0.5 W cm^{-2} for 1 min.



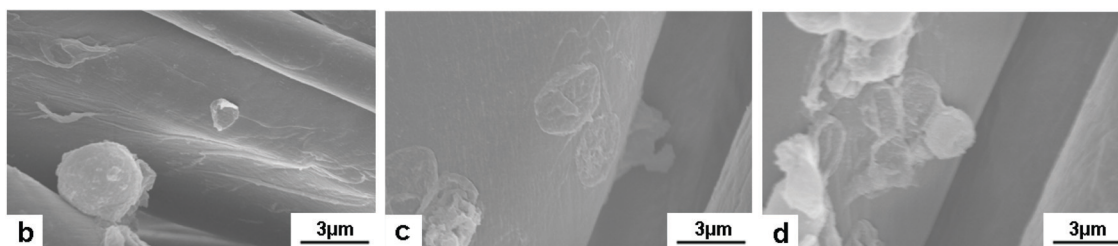
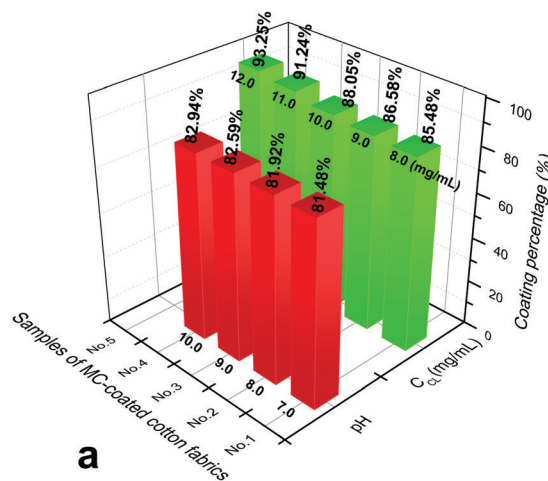


Fig. 6 Coating of (PAH/GO)₂PAH microcapsules on cotton fibers. (a) Effect of pH and crosslinker (CL) on coating percentage; SEM images of the microcapsule-coated cotton fiber in different coating media, (b) pH = 7.0, (c) pH = 10.0 and (d) CL concentration of 12.0 mg mL⁻¹ at pH = 7.0. The initial number of microcapsules was 1.0×10^8 .

can be remotely controlled by external stimulation, *e.g.* NIR radiation.^{41,42} Rhodamine B (RhB) was employed as a model molecule for investigating the permeability of (PAH/GO)_nPAH microcapsules; the standard curve of Rhodamine B is shown in the ESI† (Fig. S2). RhB was mixed with (PAH/GO)_nPAH microcapsules in PBS (pH 5.8) at room temperature for 24 h. Then, microcapsules were washed by PBS (pH 7.4) to remove excess RhB. The cumulative release of RhB from microcapsules with or without the NIR laser (0.5 W cm⁻², 808 nm, irradiation for 1 min) treatment was consequently tested *via* UV-vis spectroscopy for RhB-loaded microcapsules in solutions (pH 5.8 and 7.4). For microcapsules with three different layers, it was obvious that the release of all microcapsules at pH 5.8 was higher than that at pH 7.4, confirming pH-responsiveness of PAH/GO microcapsules. The intermolecular interaction between -COOH (of GO) and -NH₂ (of PAH) would be weakened in acidic solutions, and is beneficial for the permeable behavior. Interestingly, GO microcapsules showed a physically triggered release where NIR is considered to induce local heating through the photothermal conversion of GO, thus generating damage within the GO/PAH shell of microcapsules.^{43,44} Under the radiation of NIR laser, GO in the microcapsule layers heated its surroundings, leading to the relaxation of polyelectrolyte molecule chains and the disintegration of intermolecular interaction. This NIR irradiation contributed considerably to the accelerated release from GO microcapsules. The increase in release areas are highlighted in blue. And for microcapsules with different numbers of layers, this trend is similar (Fig. 4). For microcapsules (PAH/GO)_nPAH with

n (layers) of 1, 2, and 3, the release percentage was 75.73%, 65.02%, and 63.56% at pH 7.4 and 80.14%, 75.35%, and 70.01% at pH 5.8. Note that the release rate decreases by increasing the layer number of microcapsules. The release rate is closely related to the thickness of the shell of microcapsules. Additional layers represent an extended route distance for diffusion. Moreover, there is additional blocking effect for releasing agents getting out of the microcapsules with additional layers.^{45,46}

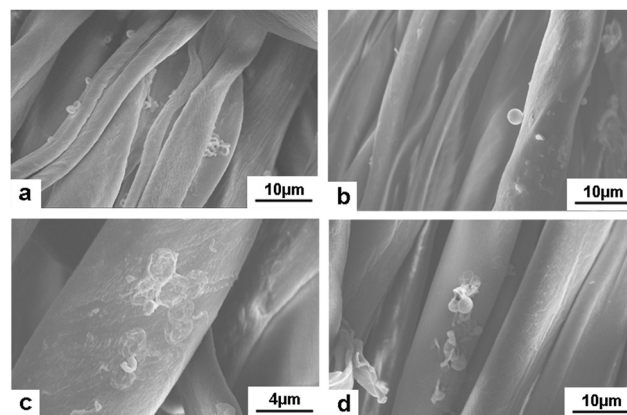


Fig. 7 SEM images of the microcapsule-coated cotton fibers after washing. (a and b) coating at pH = 10.0, washing 20 and 30 times; (c and d) coating at presence of 12.0 mg mL⁻¹ CL, washing 20 and 30 times. The initial number of microcapsules was 1.0×10^8 .



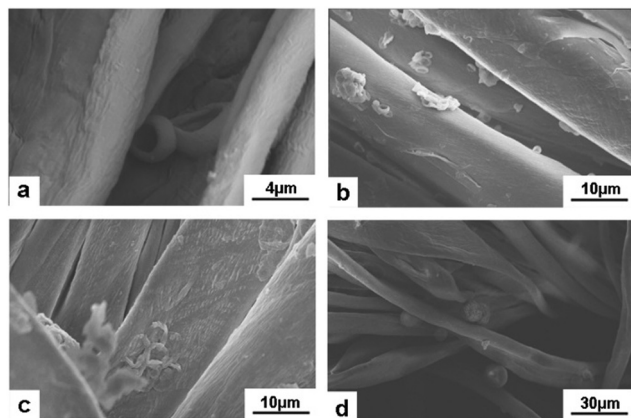


Fig. 8 *Atractylodes*-loaded microcapsules coated on cotton fibers at pH = 7.0 (a) and after washing 5 (b), 10 (c) and 20 (d) times.

3.4 NIR-triggered *Atractylodes* release from (PAH/GO)_nPAH microcapsules

Atractylodes is a natural herb with complex components. Here, a UV spectrophotometer was used to monitor the releasing behavior of *Atractylodes* from (PAH/GO)_nPAH microcapsules, and the variation in *Atractylodes* releasing at different time intervals can be monitored by the maximum adsorption peak at 220 nm (Fig. 5a). The standard curve of the *Atractylodes* solution (Fig. S3, ESI†) was determined by measuring the absorbance of the *Atractylodes* solution with different concentrations at 220 nm. Then, the release of the *Atractylodes* solution in (PAH/GO)₂PAH microcapsules with the intermittent treatment by the NIR laser (0.5 W cm⁻², 808 nm, irradiating for 1 min) was monitored. As shown in Fig. 4, because NIR has already been known to cause local heating in response to the GO photo-thermal conversion, the diffusion of *Atractylodes* out of (PAH/GO)₂PAH microcapsules significantly accelerated (the red zones in Fig. 5b). During the entire release, it could be found that this enhancement of the release rate triggered by NIR in the plain stage (later period) was more obvious than that in the beginning period with a high permeability rate. This indicated that NIR was an efficient external stimulus promoting the availability of *Atractylodes* and demonstrating the potential application of *Atractylodes*-encapsulated textiles.

3.5 Microcapsules for fabric coating

Microcapsules were considered as good materials for textile modification. Certain functional ingredients can be loaded into the hollow space of microcapsules and then be connected with textile fibers through adsorption, electrostatic interaction and chemical bonding. Due to the carboxyl and hydroxyl groups on the surface, cellulose fiber was negatively charged, which can bond with the amino group of the outermost PAH layer of GO/PAH microcapsules with a positive charge. (PAH/GO)₂PAH was then used to study the microcapsule-coated cotton fibers. In alkaline conditions, COOH⁻ and OH⁻ of cotton fiber was more ionized, improving the bonding with -NH₃⁺ of (PAH/GO)₂PAH. Moreover, in order to improve the interaction between (PAH/GO)₂PAH and cotton fiber, an epoxyethane quaternary ammonium salt was used as the cross linker (CL) efficiently binding (PAH/GO)₂PAH microcapsules with cotton fibers. The positively charged group NR₄⁺ of the cross-linker interacted with -COO⁻ (-COOH) or -O⁻ (-OH) groups of the cotton fiber, while the ethylene oxide groups bound with the -NH₃⁺ groups of the outermost layer PAH of the (PAH/GO)₂PAH microcapsules. As shown in Fig. 6, the bonding percentage of microcapsule with cotton fibers increased with the increase in pH or CL concentration. Without any CL, the number of coated microcapsules reached a maximum at pH 10. Moreover, compared with pH, the enhancement in CL on connection between microcapsules and cotton fiber was much more remarkable, indicating strong interaction by chemical bonding. At a CL of 12.0 mg mL⁻¹, the number of coated microcapsules was maximum. The morphology of cotton fiber modified with microcapsules was shown in the SEM image. The number of microcapsules connected with cotton fibers increased in the presence of alkaline media or CL. After washing 30 times, there were microcapsules on cotton fibers, which were more obvious when 12.0 mg mL⁻¹ CL was used (Fig. 7). The number and percentage of microcapsules on the fabric are determined using hemocytometers.

Moreover, *Atractylodes*-loaded microcapsules were coated on cotton fibers. As shown in Fig. 8, (PAH/GO)₂PAH was already coated on fibers. After washing by 5, 10, and 20 times, there were microcapsules coated on the textile. Fig. 9 showed the FTIR spectra of cotton fibers coated with *Atractylodes*-loaded microcapsules. The peaks at 1033 cm⁻¹ and 670 cm⁻¹ represented the Ar-H in-plane vibration and Ar-H out-of-plane bending, indicating

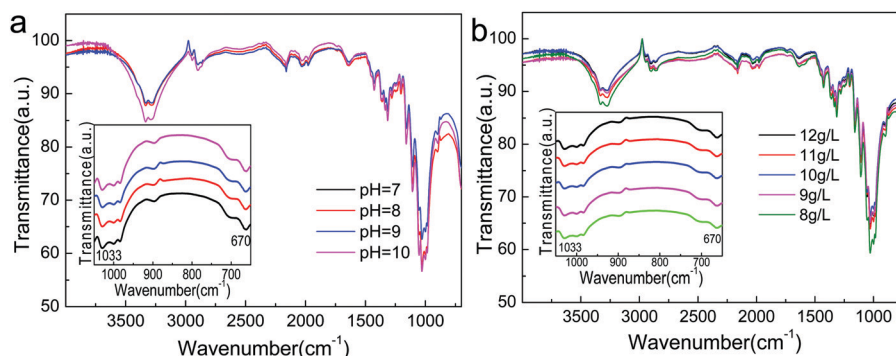


Fig. 9 FTIR spectra of *Atractylodes*-loaded microcapsules coated on cotton fibers at different pH (a) and CL concentrations (b).



(PAH/GO)₂PAH microcapsules were successfully connected with a cotton fiber. Moreover, the intensity of these peaks was enhanced with the increase in pH and CL concentration, confirming the results of SEM images.

4 Conclusion

In summary, (PAH/GO)_nPAH microcapsules were prepared *via* an LBL self-assembly. The electrostatic interaction of oppositely charged PAH and GO nanosheets provided a robust shell with a hollow space centre to load drugs or dyes for controlled release through pH stimuli and NIR irradiation. NIR-triggered GO layer provided local heating and generated on-demand drug/dye release from (PAH/GO)_nPAH microcapsules. Then, the hybrid PAH/GO microcapsules can be coated on the cotton fabric to manufacture a medical care textile with both pH and NIR responsiveness. This study suggested a method to prepare stable GO composite microcapsules, which can be used as media to introduce chemicals to textiles, thus generating functional materials with controlled release behaviour under remote external stimulation. The microcapsule-coated textile can provide a new material beneficial to transdermal drug therapy and photothermal physiotherapy as well as a shield against the discomfort from the local overheating aroused by NIR irradiation. Moreover, the synergistic effect of photothermal treatment and herb care can lead to a potential healthcare textile with promising applications.

Conflicts of interest

There are no conflicts of interest to declare.

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