Green “planting” nanostructured single crystal silver

Hong Zhao1,3*, Fei Wang2, Yuesheng Ning2*, Binyuan Zhao2,3, Fujun Yin3, Yijian Lai2, Junwei Zheng1, Xiaobin Hu2, Tongxiang Fan2, Jinguo Tang4, Di Zhang2 & Keao Hu2

1College of Chemistry, Chemical Engineering and Materials Science, Soochow University, Suzhou, 215123, P. R. China, 2State Key Laboratory of Metal Matrix Composites, School of Materials Science and Engineering, Shanghai Jiao Tong University, Shanghai, 200240, P. R. China, 3Huaihai Institute of Technology, Jiangsu Marine Resources Development Research Institute, Lianyungang 222005, P. R. China, 4College of Chemistry, Chemical and Environmental Engineering, Qingdao University, Qingdao, 266071, P. R. China.

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Correspondence and requests for materials should be addressed to B.Y.Z. (byzhao@sjtu.edu.cn); J.W.Z. (jwzheng@suda.edu.cn) or T.X.F. (txfan@sju.edu.cn)

* These authors contributed equally to this work.

The past decade has witnessed the burgeoning research interest in the controlled synthesis of monodispersed noble metal nanocrystals1–4. Various nanostructures such as nanocubes5, Nanocages6, Nanoplates7, nanowires8 and nanobelts9 have been fabricated. Solution-phase methods involved the reduction of metal ions, and the growth of nanocrystals in aqueous or non-aqueous solution. They were believed to be more adaptive than gas-phase methods in preparing nanostructures with a range of morphologies in bulk quantities10. The main solution-phase methods can be categorized to (1) chemical synthesis with or without seed and light mediation, where various types of reductants and capping reagents were often utilized, and (2) hard or soft template-directed growth11. In the hard-template approach, a porous anodic aluminum oxide (AAO) membrane often served as the substrate. Since the growth of noble metals was initiated from the bottom of the channels, the synthesized product often had only one-dimensional (nanotubes, nanowires, nanorods and nanobelts etc.) morphology, and the width was confined by the size of the insulating nanochannels. The dissolution of the template with corrosive reagents also lowered the efficiency of this method. There has been very few works on the scaled production of metal nanocrystals with various morphologies using solid template12.

In our previous work13, silver nanodendrites were prepared on the outer surface of monolithic activated carbon (MAC), by simply immersing MAC into AgNO3 solution. A galvanic-cell mechanism was proposed. In this reaction mechanism (Fig. 1), there are numerous short-circuited nanobatteries composed of reductive groups on the inner surface of the microporous carbon acting as the anode, the growing silver as the cathode, and the conductive carbon skeleton as the external circuit. Considering the large BET surface of activated carbons associated with abundant functional groups, the growth capacity of this method would be quite high. However, in the previous work, only AgNO3 was used as the Ag precursor. The relatively high standard reduction potential of Ag+/Ag (0.80 V) produced a high driving force, leading to the kinetically-controlled deposition of silver in a non-equilibrium condition. This could explain why in our previous work, only dendritic silver was obtained14. In the current work, a silver complex [Ag(NH3)2]NO3 aqueous solution only. By varying the [Ag(NH3)2]NO3 concentration, silver nanoplates or nanoflowers can also be selectively obtained. The silver growth was illustrated using a galvanic-cell mechanism. The lowering of cell potential via using [Ag(NH3)2]NO3 precursor, together with the AgCl crystalline seed initiation, and the releasing of OH− in the reaction process, create a stable environment for the self-compensatory growth of silver nanocrystals. Our work revealed the great versatility of a new type of template-directed galvanic-cell reaction for the controlled growth of noble metal nanocrystals.
silver nanocrystals in a self-compensatory manner. Furthermore, the as-prepared silver nanocrystals can be easily detached from the porous carbon template since they mainly grew on the outer surface, and the MAC can be reactivated by simple post-treatment. Our method may represent a very promising template-directed method to scaling up the production of noble metal nanocrystals with controlled morphology.

Results

Figure 2a–c display three photographic snapshots of Ag samples growing on MAC at different time, upon immersing two pieces of small MAC (All structural parameters and elemental composition are presented in Supplementary Table S1 and S2 online) in [Ag(NH₃)₂]NO₃ aqueous solution with 5 mM concentration. (More experimental details are given in the Methods) In about 30 min, shiny Ag crystals quickly grew out of the black MAC (Fig. 2b). Soon after that, fluffy product appeared and its apparent volume became much larger than original MAC after 24 hr (Fig. 2c). SEM images of the products obtained by immersing MAC in [Ag(NH₃)₂]NO₃ solutions at (a) 100 mM, (b) 5 mM, and (c) 0.1 mM for 24 hr. The products obtained at 5 mM (Fig. 2b) consisted of highly pure nanobelts, similar to that mentioned in Fig. 2. At higher concentrations (Fig. 3a), many flower-like products were obtained, in the form of the aggregation of irregular silver nanoplates. On the other hand, at lower concentrations (Fig. 3c), well-crystallized hexagonal silver nanoplates grew on the MAC surface. The change of silver morphology with varying [Ag(NH₃)₂]NO₃ concentration was also found on other MACs (for another example, see Supplementary Fig. S1 online).

Interestingly, the MAC can be used to grow silver nanocrystals many times, with only simple post-treatment in between. Figure 4 shows that the reactivated MAC can be used to grow almost the same amount of silver samples (compare Figs. 4b and 4e) as the virgin MAC. The reactivation was performed by taking away the Ag product (Fig. 4c) by tweezers, ultrasonication the MAC in ammonium solution (5 wt%) for ca. 1 hr, washing with deionized water and drying it at 60 °C for 2 hr (Fig. 4d). It was found that the ultrasonication can also be carried out in ethanol, or even just deionized water.

To receive hints as to the underlying growth mechanism of silver nanobelts, the surface deposits on the MAC immersed in 5 mM [Ag(NH₃)₂]NO₃ solution at earlier stages were examined using SEM and XRD (Fig. 5). The black MAC quickly turned gray-yellow after immersing in the solution for ca. 1 min. SEM images (Fig. 5a) reveal that many small cubes and some right bipyramids appeared on the surface, and the corresponding XRD spectrum (Fig. 5 lower panel, curve a) indicates that they are a mixture of AgCl and Ag. AgCl is the dominant phase and Ag the minor. The cubes could be associated with fcc AgCl phase. After 10 min (Fig. 5b), the cubes almost disappeared and some hexagonal plates appeared, together with some defected pyramids and irregular structures. The corresponding XRD spectrum (curve b) indicates that the products are mainly fcc Ag crystals. The basal face of these hexagons can be ascribed to [111] plane and the sides to {100} planes of fcc Ag.

SAED pattern has a 6-fold symmetry and the d spacing of the planes nearest to the central spot is measured to be 1.44 Å, which can be assigned to the [220] reflections in a face-centered-cubic (fcc) silver whose (111) surface is perpendicular to the electron beam in TEM observation. Therefore, the photograph and EM images demonstrate that belt-like single crystal silver can be prepared in a high yield (silver product weighed ca. 9% of MAC) by simply immersing MAC in [Ag(NH₃)₂]NO₃ aqueous solution.

In addition, the morphology of single crystal silver can be controlled by varying [Ag(NH₃)₂]NO₃ concentration. Figure 3 shows SEM images of the products obtained by immersing MAC in [Ag(NH₃)₂]NO₃ solutions at (a) 100 mM, (b) 5 mM, and (c) 0.1 mM for 24 hr. The products obtained at 5 mM (Fig. 3b) consisted of highly pure nanobelts, similar to that mentioned in Fig. 2. At higher concentrations (Fig. 3a), many flower-like products were obtained, in the form of the aggregation of irregular silver nanoplates. On the other hand, at lower concentrations (Fig. 3c), well-crystallized hexagonal silver nanoplates grew on the MAC surface. The change of silver morphology with varying [Ag(NH₃)₂]NO₃ concentration was also found on other MACs (for another example, see Supplementary Fig. S1 online).

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The concomitant loss of cubes in SEM (Fig. 4b) and AgCl phase in XRD (curve b) suggested that the initially crystallized AgCl cubes underwent reduction and transformed into Ag hexagonal plates, releasing Cl\(^{-}\) ions.

It should be mentioned that the Cl\(^{-}\) ions for the formation of AgCl in the first step (Fig. 5a) were inherent in our MAC samples. The content of the Cl\(^{-}\) in MAC should be kept at a relatively low level (preferably less than 0.2 at% as revealed by our XPS results in Supplementary Table S2). It is deduced that Ag\(^{+}\) released from [Ag(NH\(_3\)_2]\(^{+}\) first complexed with Cl\(^{-}\) ions on MAC, forming AgCl cubes on the surface. Then, the AgCl crystallites were gradually reduced, generating numerous Ag hexagonal plate seeds while releasing Cl\(^{-}\) ions. The formation of Ag hexagonal plates could be explained by preferential adsorption of Cl\(^{-}\) on the [111] basal surfaces of fcc Ag\(^{2+}\), and the stacking faults on the side surfaces parallel to [111] surfaces\(^{19,20}\). Subsequently, Ag atoms may directly deposit on the Ag hexagonal sides due to the edge defects and associated higher surface energy. For comparison, we have also immersed Cl\(^{-}\) ions on MAC, forming AgCl cubes on the surface. Therefore, it seems that crystalline seeds are indispensable for the growth of well-crystallized nanoparticles in our reaction system.

With the growing process continued, many silver nanobelts and large nanosheets appeared in ca. 30 min (Fig. 5c). At the same time, it became quite difficult to find silver hexagonal plates on the MAC surface. Therefore, it is suggested that the hexagonal plates obtained at earlier stages still grow along the [100] hexagonal plate sides, either on all six sides to form nanosheets, or on only two neighboring sides and grow into nanobelts. Notably, some irregular Ag nanosheets with short nanobelt tapes and/or indentations were frequently found in this sample (Supplementary Fig. S3 online). The reason for the formation of such irregular products remains unclear. It is reminiscent of a study where similar silver structures were also found when using NH\(_3\)·H\(_2\)O to tune the pH value\(^{46}\), inferring that NH\(_3\)·H\(_2\)O may play a role in etching the silver nanosheets. Such nanosheets mostly disappeared in the later growing process, and silver nanobelts became the only dominant products after 12 hr (Fig. 5d).

**Discussion**

It is of paramount importance to know what caused the reduction of Ag\(^{+}\). In the preparation of silver nanostructures, various reducing agents have been employed, such as polyols\(^{21}\), citrate\(^{22}\), ascorbic acid\(^{23}\) etc. Other reducing/transformation protocols include more reactive metals (termed as galvanic replacement)\(^{24}\), colloidal reflux\(^{10}\), light mediation\(^{25}\) and electrochemical reduction\(^{26}\). However, in our work, all these reducing forces have been avoided. The silver growth simply proceeded on the surface of an MAC immersed in a freshly-prepared [Ag(NH\(_3\)_2]NO\(_3\) solution. The MAC had been vigorously ultrasonicated in DI water to remove physically-adsorbed species. And the experimental setup was shunned off common laboratory light. Therefore, the reduction could only be exerted by chemically (probably covalently) bound functional groups on the porous MAC surface\(^{27,28}\).

To identify the reductive groups in our reaction system, FTIR was performed on virgin, used, and post-treated (by ammonium) MAC samples. These samples were ultrasonicated in fresh water, dried, ground into powders, mixed with KBr and pressed into small discs prior to FTIR characterization (more experimental details are given in the Methods). Unfortunately, it was not always successful in catching the spectral difference between the virgin and used MAC samples (possible reasons will be discussed below). Often all spectra look quite similar. Figure 6 displays a set of FTIR spectra through which the chemical reaction on the activated carbon may be inferred. In virgin MAC (curve a), the most intense peak at 3429 cm\(^{-1}\) can be assigned to alcohol or phenol –OH stretching. The two peaks at 2858, 2960 cm\(^{-1}\) may be attributed to symmetric, asymmetric –CH\(_2\) stretching respectively. The peaks at 1633 cm\(^{-1}\) could be ascribed to C=C or C=O stretching. After the growth of silver (curve c), there is significant alleviation of –OH stretching (3429 cm\(^{-1}\)) intensity, especially when compared to –CH\(_2\) stretching (2858 and 2960 cm\(^{-1}\)). The peak intensity at 1633 cm\(^{-1}\) is also fairly reduced. This suggested that OH groups were involved in the chemical reaction. At the same time, two new peaks appeared in the used activated carbon (curve c). According to the literatures on the characterization of chemically modified porous carbon\(^{29}\), the peak at 1732 cm\(^{-1}\) can be ascribed to carboxylic O=C=O asymmetric stretching, and the broad shoulder peak at 3174 cm\(^{-1}\) is attributable to –OH stretching in COOH groups as a result of H-bonding. Therefore, the FTIR results suggested that there was an oxidative transformation from hydroxyl and/or carboxyl groups to carboxylic groups in the process of silver growth. Given the existence of O\(_2\) in the atmosphere and in the pores of the microporous carbon, the hydroxyl groups may be
Figure 5 | SEM images and XRD spectra showing the growth process of Ag nanobelts on monolithic activated carbon (MAC). The samples were obtained after immersing monolithic activated carbon (MAC) in \([\text{Ag(NH}_3\text{)}_2\text{NO}_3\text{]}\) solution (5 mM, 30 ml) for (a) 1 min; (b) 10 min; (c) 30 min; (d) 12 hr. SEM images for (a), (b) and (c) were directly taken on MAC surface, while SEM image (d) was obtained from detached Ag sample. XRD samples (a) and (b) were prepared by grinding MAC into powder, while XRD sample (d) was detached Ag product.

Figure 6 | FTIR spectra of monolithic activated carbon (MAC) for (a) virgin MAC; (b) post-treated MAC; (c) used MAC. For (b), the treatment was performed by ultrasonicating MAC in ammonium solution (wt 5%) for ca. 1 hour, flushed with plenty of deionized water, and dried.
first oxidized to aldehyde by O2, and then to carboxylic acid by galvanic reaction, in agreement with the mechanism proposed by Xia et al. for their polyol approach15. The rich functional groups contained in our activated carbon can be associated with its biological source. After the detachment of Ag, it can be seen that OH groups were regenerated by ammonium ultrasonication and drying (curve b), which may explain why we obtained silver crystals with similar amount after the treatment (Fig. 4).

In our previous work13, we have proposed a galvanic-cell mechanism for the preparation of silver nanodendrites on MAC (Fig. 1). Considering the large BET surface of MAC, silver nanocrystals can be produced in quite high yield, which was not clearly expressed in that work. For example, for the MAC used in this work, it is estimated that 0.35 g silver nanocrystal can be produced on 1 g MAC, provided that all the reductive groups on the inner pores can be oxidized through galvanic-cell mechanism (see Supplementary Information for estimation method). If only the reductive groups on the outer surface can be oxidized for the preparation of silver, then its yield would not exceed 1 × 10−6 g. In this work, we have obtained ca. 0.11 g silver on 1 g MAC in one batch, implying ca. 31.5% available reductive groups in the inner pores have been electrochemically oxidized. However, the reacted functional groups only constitute 9.7% of all oxygen-containing functional groups, which is difficult to tell by FTIR in each experimental trial.

To understand the electrochemical behavior of the MAC, cyclic voltammetry (Fig. 7) was conducted in a three-electrode configuration in a 50-ml beaker where 0.1 M NaNO3 was used as the electrolyte. MAC was ground into fine powders, mixed with ethanol, and loaded onto the surface of a glassy carbon working electrode. (The details of the experimental setup was given in the Methods) Two types of MAC used in this work were characterized. Both had one anodic (at ~0.22 V) and two cathodic peaks (at ~0.17 V and ~0.08 V). The intensity of these three peaks attenuated simultaneously with the experiments going on, indicating that the peak at 0.22 V is the anodic counterpart to either of the two cathodic peaks. Consider the IR results discussed above, the anodic peak at 0.22 V was tentatively assigned to R-CHO (H2O R-COOH + 2H+ + 2e → R-COOH + 2H+ + 2e → R-CHO + H2O respectively. However, more investigation is still required for the verification of this assignment. Anyway, our CV results did confirm that functional groups on MAC can be electrochemically converted in reducing Ag+ for the growth of Ag nanostructures.

For the galvanic-cell mechanism (Fig. 1), some factors can influence the silver growth process, such as the cell potential, the density of reductive groups, the mobility of charge carriers (ions and electrons), the surface and pore structure of MAC, some accelerating/retarding species adsorbed on MAC or contained in the solution, and the solution temperature etc. Among them, the cell potential is perhaps the key influential factor, which in turn is influenced by the substrate type (the anode), the identity and concentration of the silver precursor (the cathode). For example, the galvanic-displacement growth of Ag on Al or Zn substrates tends to produce dendritic or flower-like products, since elemental Al and Zn are much more reactive than Ag, which lead to high cell potentials31–33. On Cu micro-particles14, the particle size, the identity and concentration of silver precursor can all influence the product morphology. Large Cu particle size (10 μm) and high concentration of AgNO3 mainly resulted in dendritic products. On smaller Cu particles (1 μm), silver nanobelts, nanodiscs were obtained in AgNO3 solution at higher (10 mM, ΔE = 0.35 V), lower (1 mM, ΔE = 0.29 V) concentrations, respectively. Replacing AgNO3 by [Ag(S2O3)2]− complex (5 mM, ΔE = 0.18 V) led to uniform deposition of Ag nanoshells on Cu. It is reasoned that lower cell potential resulted in lower driving force and more thermodynamically-controlled reaction, and more uniform growth of the crystals. In the current work, instead of AgNO3 used in our previous work13, a silver complex [Ag(NH3)2]NO3 was used as the silver precursor. In addition, the inherent Cl− in MAC produced AgCl cubes in the initial reaction step (Fig. 4a). Compare the standard reduction potential of Ag+/Ag (0.80 V), [Ag(NH3)2]−/Ag (0.38 V) and AgCl/Ag (0.22 V), we could see there is significant decrease in the reduction potential. Moreover, the high solubility of [Ag(NH3)2]− and the complexation structure may provide a good buffer solution of Ag+ near the Ag deposition region. Our adjustment made it possible for the quasi-equilibrium growth of well-crystallized silver nanostructures, avoiding the growth of dendritic crystals, which are normally obtained in an unstable reaction condition where the diffusion of the solute ions is the limiting factor14.

At the quasi-equilibrium growth condition, by varying the precursor concentration, the cell potential can be finely tuned. As derived from Nernst Equation, the decrease of the precursor concentration by one order will result in the decrease of cell potential by ca. 0.06 V, which had been large enough for producing silver crystals with different morphology, as discussed above14. In this work, by setting the [Ag(NH3)2]NO3 concentration at 100 mM, 5 mM, 0.1 mM, silver crystals in the form of nanoflower, nanobelt,
be considered. According to the anode reaction R-CHO + H2O → R-COOH + 2H+ + 2e, the acidity of the solution would be improved if there were no other influential factors. On the other hand (Fig. 8), the hydrolysis of [Ag(NH3)2]+ precursor, [Ag(NH3)2]+ + 2H2O → Ag+ + 2NH4+ + 2OH−, can produce OH− ions, which may somewhat neutralize the acidity with the progress of galvanic deposition. Furthermore, it has been reported that alkaline treatment of activated carbon can regenerate –OH functional groups on the activated carbon surface. In this way, the declining of reductive groups can be hindered, keeping the cell potential at a relatively stable value during silver growth. Hence, a self-compensatory growth of Ag nanocrystals can be realized at a quasi-equilibrium condition. We have also found that pure Ag2O is a very good precursor for the growth of silver nanobelts on MAC in a high yield, and the proposed galvanic-cell mechanism is further studied by analyzing the surface functional groups and the electrochemical behavior of the activated carbon electrode. It is argued that the lowering of the cell potential by using [Ag(NH3)2]+ complex precursor, the AgCl crystalline seed-initiation, and the regeneration of the reductive groups facilitate the self-compensatory growth of Ag nanoparticles. As the silver nanocrystals grew out of environmental-friendly MAC template in aqueous solution without any organic reductants or capping agents, this process is quite like planting seeds and producing harvests abundantly in a field. It is anticipated that this work will pave the way for customizing the production of various kinds of noble metal nanocrystals in a green mode and at an industrial scale.

Methods

Chemical and materials. Monolithic Activated Carbon (MAC) samples, derived from corn straw, were provided by Qufu Tembton Technology Co. Ltd. Shandong, China. Their structural and compositional information is presented in Supplementary Table S1 and Table S2 online. AgNO3, ammonia water (25–28 wt%), anhydrous ethanol were purchased from Shanghai Chemical Reagent Co. Ltd. All chemicals were of analytical grade and used without further purification. AgNO3 solution was prepared by continuously dropping ammonia water into 1 M AgNO3 until the dissolution of the brown Ag2O precipitate. [Ag(NH3)2]NO3 solutions at lower concentrations were obtained by dilution with deionized (DI) water accordingly. DI water was generated from TTL-30C Ultrapure Water Generator, with electrical resistivity of 18.2 MΩ·cm.

Preparation of silver nanocrystals. MAC was cut into small pieces and ultrasonicated in DI water to remove contaminants. The silver nanocrystals were prepared by immersing MAC into freshly-prepared [Ag(NH3)2]NO3 solution at different concentrations in the dark to avoid silver photoreduction. All experiments were carried out at room temperature (ca. 25°C). The Ag products were detached from MAC with tweezers. For the reactivation of MAC, the used MAC was ultrasonicated in ammonium solution (5 wt%) for ca. 1 hr, washed with DI water and dried at 60°C for 2 hr.

Characterization. Photograph. All photographs in this work were taken with a digital camera (OLYMPUS SZ-10).

SEM. Scanning electron microscope (SEM) experiments were performed on JEOL JSM-6390LV instrument. The images were taken in secondary electron mode, with an accelerating voltage of 15 kV, working distance of ca. 7 mm. Samples were mounted on Al stages with conductive carbon tapes.

TEM. Transmission electron microscope (TEM) and selected-area electron diffraction (SAED) images were collected in a Philips CM10 instrument at an accelerating voltage of 160 kV. TEM samples were prepared by sonicating Ag product in
anhydrous ethanol for 5 min, taking some suspension droplets and drying in a carbon-film coated copper grid.

XRD. The powder X-ray diffraction (XRD) pattern was recorded by using a Bruker AXS D8 X-ray diffractometer with Cu Kα radiation (λ = 1.5418 Å). The measurement parameters were set at 2θ = 20°–200° range, 0.02° sampling frequency, 2/min scanning speed, 30 kV tube voltage and 40 mA tube current, respectively.

FTIR. Fourier Transform Infrared (FTIR) spectra were collected by using a Bruker Tensor 27 instrument within the range of 400–4000 cm⁻¹. 16 scans and 4 cm⁻¹ resolutions were applied. For specimen preparation, MAC samples were dried, ground in an agate mortar, thoroughly mixed with KBr at an approximate ratio of 1:100 and pressed into small discs at 10 tons for 5 min.

CV. Cyclic voltammetric (CV) measurements were performed by using a CHI 660 d electrochemical analyzer (CH Instruments, Shanghai Chenghua Co.) in a conventional three-electrode configuration. A Pt plate auxiliary electrode, a saturated calomel reference electrode (SCE), and 0.1 M NaNO₃ electrolyte solution were used. For the preparation of working electrode, MAC was ground into powder, and 4 mg of the powder and 1.6 mL anhydrous ethyl alcohol were thoroughly mixed by sonication for 1 hr to generate an evenly distributed suspension. Then 4 μL of the resulting suspension was laid on the surface of the pre-cleared glassy carbon electrode (3 mm in diameter, 0.07 cm²). After drying at room temperature, 3 μL Naflon (5.0 wt%) solution was covered on the surface of the working electrode and allowed to dry again.

The as-prepared working electrode had a specific MAC loading of ca. 0.14 mg/cm². All the electrochemical measurements were carried out at 30 ± 1°C.

BET. To determine the surface areas of MAC, the N₂ adsorption-desorption isotherms were measured with an accelerated surface area and porosimetry system (ASAP Tristar, Micromeritics). A small piece of ca. 0.25 g MAC was used as the sample. Prior to the measurements, the samples were dried at 100°C for 2 hours, and then outgassed at 623 K under < 6.67 × 10⁻³ Pa pressure, for at least 4 h. The nitrogen adsorption-desorption data were recorded at a liquid nitrogen temperature of 77 K. The nitrogen adsorption isotherm was measured over a relative pressure (p/p₀) range, from approximately 0.05 to 1. The BET surface area was calculated using the BET equation from the selected N₂ adsorption data, within a range of relative pressure, p/p₀ from 0.1 to 0.25. Barrett-Joyner-Halenda (BJH) method was applied to obtain the pore size distribution in the mesopore range, and t-plot method to that in micropore range (< 17 Å).

XPS. X-ray photoelectron spectroscope (Thermo Scientific ESCALAB 250) was applied to quantify elemental composition (except H). The XPS spectra were obtained using a monochromated Al Kα radiation (1486.6 eV), which was operated at 15 kV and 14 mA. The pass energy for wide scans was set at 160 eV, while the narrow scans at 40 eV. The C(1s) electron binding energy peak was set at 284.6 eV for calibration purposes.

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