Enhanced adsorptive removal of p-nitrophenol from water by aluminum metal–organic framework/reduced graphene oxide composite

Zhibin Wu1,2, Xingzhong Yuan1,2, Hua Zhong1,2,3, Hou Wang1,2, Guangming Zeng1,2, Xiaohong Chen4, Hui Wang1,2, Lei Zhang1,2 & Jianguang Shao1,2

In this study, the composite of aluminum metal–organic framework MIL-68(Al) and reduced graphene oxide (MA/RG) was synthesized via a one–step solvothermal method, and their performances for p–nitrophenol (PNP) adsorption from aqueous solution were systematically investigated. The introduction of reduced graphene oxide (RG) into MIL-68(Al) (MA) significantly changes the morphologies of the MA and increases the surface area. The MA/RG-15% prepared at RG-to-MA mass ratio of 15% shows a PNP uptake rate 64% and 123% higher than MIL-68(Al) and reduced graphene oxide (RG), respectively. The hydrogen bond and π–π dispersion were considered to be the major driving force for the spontaneous and endothermic adsorption process for PNP removal. The adsorption kinetics, which was controlled by film–diffusion and intra–particle diffusion, was greatly influenced by solution pH, ionic strength, temperature and initial PNP concentration. The adsorption kinetics and isotherms can be well delineated using pseudo–second–order and Langmuir equations, respectively. The presence of phenol or isomeric nitrophenols in the solution had minimal influence on PNP adsorption by reusable MA/RG composite.

Nitrophenols are widely used in petrochemical synthesis, including paints, plastics, rubber, pulp, pesticides and dyes production1. The presence of nitrophenols in the industrial wastewater has aroused great concerns in recent years due to the increase in wastewater discharge and the toxicity of nitrophenols to the receiving bodies2. In particular, the p-nitrophenol (PNP) has intensive toxic effect on methaemoglobin formation, causing liver and kidney damage, anaemia, skin and eye irritation, and systemic poisoning3,4. It has been listed as a priority pollutant by the U. S. Environmental Protection Agency (U.S. EPA)5. For years, to minimize nitrophenol pollution from wastewater, the methods of photo-degradation6, adsorption7, and chemical oxidation8, etc, have been developed. Among these methods, adsorption is considered to be a promising one due to the advantages of this method, e.g., simplicity and cost-effectiveness.

Graphene oxide (GO), a type of negatively charged colloid comprising multiple oxygenated graphene layers with one-atom thickness honeycomb lattice structure, has received great attention for pollutants removal from wastewater due to the high specific surface area and great application promise9–12. For example, Wang et al.13 used reduced graphene oxide for adsorption of phenolics and interpreted the correlation between the adsorption ability and reduction degree of graphene oxide. In our previous studies, the graphene oxide exhibited excellent efficiency for Zn2+ removal14 and a superior adsorption capacity of methylene blue was achieved by rhamnolipid functionalized graphene oxide15. In terms of the removal of PNP, Zhang et al.16 reported that the precursor for

1College of Environmental Science and Engineering, Hunan University, Changsha 410082, P. R. China. 2Key Laboratory of Environment Biology and Pollution Control, Hunan University, Ministry of Education, Changsha 410082, P. R. China. 3Department of Soil, Water and Environmental Science, the University of Arizona, Tucson, AZ85719, US. 4Hunan University of Commerce, Changsha 410205, P. R. China. Correspondence and requests for materials should be addressed to X.Y. (email: yxz@hnu.edu.cn) or H.Z (email: zhonghua@hnu.edu.cn) or G.Z. (email: zgming@hnu.edu.cn)
GO, nanographite oxide, has a maximum PNP adsorption capacity of 268.5 mg/g at 283 K and a natural pH. However, the GO is hard to separate from solution after adsorption due to the hydrophilic property. Recently, reducing the surface functional groups of graphene oxide is considered to be an effective method to decrease hydrophilicity and thus achieve better separation performance. Unfortunately, when the GO was reduced, the highest adsorption capacity of PNP was observed to be only 15.5 mg/g at 298 K and pH6. Therefore, it is a challenge to improve the adsorption performance of graphene-based materials for PNP removal.

Due to the large surface area, diverse structure, and tunable functionality, metal–organic frameworks (MOFs) have recently attracted extensive attentions in adsorption, catalysis, sensing, gas storage, and drug delivery. The MOFs of MIL–68(Al) (MA) is assembled from the infinite straight chains of corner–sharing metal–centered octahedral AlO4(OH), that is connected to each other through hydroxyl groups and terephthalate ligand. Yang et al. reported that the MIL–68(Al) has great gas adsorption due to the presence of triangular and hexagonal channels of an opening diameter (6.0 – 6.4 Å and 16 – 17 Å). Xie et al. utilized MIL–68(Al) for nitrobenzene removal from water and achieved a quite high adsorption capacity of 1130 ± 10 mg/g. The adsorption properties of MIL–68(Al) can be improved by hybridization with other materials. Han et al. chose the functionalized carbon nanotube (CNT) to composite with MIL–68(Al) and exhibited 188.7% enhanced phenol adsorption capacity from water than pristine MIL–68(Al). Although series of MOFs/graphene based composites were synthesized and used for gas adsorption, gas storage, and organic compounds adsorption, to date the adsorption of greenhouse gases or pollutants have not been reported.

In this study, the MIL–68(Al)/RG composite was synthesized using a simple solvothermal method, and its performance for adsorptive removal of PNP from water was examined. The adsorption kinetics and thermodynamics were investigated in detail. Factors that may potentially affect the adsorption process, such as pH, ionic strength, temperature, recycle number and coexistence of isomers or phenol, were also examined.

**Result and Discussion**

**Characterizations.** The morphologies of MA, RG and MA/RG observed by scanning electron microscopy (SEM) and transmission electron microscopy (TEM) are presented in Fig. 1. It can be seen that the MA (Fig. 1a) is in aggregate ball–liked particles in the form of nanosized and well–dispersed MIL–68(Al) crystallites. The results of element composition (Fig. 1i) and the maps of elements (Fig. 1j–l) obtained using EDX show that the C, O and Al are uniformly distributed on the surface of the MA/RG–15% (Fig. 1h). The TEM images (Fig. 1(m–o)) further confirm the homogeneous nucleation nods for the formation of MIL–68(Al) with precious few agglomeration, and thus lead to the formation of nanosized and well–dispersed MIL–68(Al) crystallites. The results of element composition (Fig. 1i) and the maps of elements (Fig. 1j–l) obtained using EDX show that the C, O and Al are uniformly distributed on the surface of the MA/RG–15% (Fig. 1h). The TEM images (Fig. 1(m–o)) further confirm the morphological change after the hybridization. As depicted in Fig. 1o, the MA scattered in the transparent folded RG layers of MA/RG–15% composite are plate–like particles, which seemed to be different from the parental one (Fig. 1m), attributing to the distortion force of GO for MIL–68(Al) formation by the π–π stacking interaction.

Figure 2a shows the X–ray diffraction (XRD) patterns of MA, RG and MA/RG composites. The characteristic peaks of MA are in agreement with the previous report of MIL–68(Al), suggesting that the current material has the structure as expected. For RG, the peak at 20 = 22.2° corresponds to graphene with the interlayer distance of 0.400 nm, indicating the reduction of graphene oxide and the restoration of sp2 bonded carbon. The peak at 20 = 13.2° is produced by the residual oxygen groups such as epoxy as carbonyl groups, due to incomplete reduction. After hybridization of MA with RG, the major diffraction patterns of the composites are similar to that of the pure MA, and no diffraction peak for RG is observed in the composites, which is probably due to that the low content of RG is shielded by the attached MA particles. To testify the presence of RG, the Raman spectra (Fig. 2b) of the MA/RG composites are measured. The major characteristic peaks for RG include D peak (∼1348 cm−1) which is resulted from the breathing mode of π–point phonons of Al1g symmetry, and the G peak (∼1601 cm−1) which is from the Eg2 phonon of the sp2 hybridization. For MA, the characteristic peaks at 1616, 1475 and 1147 cm−1 correspond to the in–plane vibration of the aromatic rings in the terephthalic acid ligands, and the peaks at 869 and 631 cm−1 are associated with C–H stretching or out–of–plane vibration of the aromatic rings.

The FT–IR results for the parent materials and MA/RG composites are presented in Fig. 2c. Characteristic peaks of C–H (754 cm−1), C–O–C (991 cm−1), C–O (1512 and 1701 cm−1) and C=C (1411 and 1587 cm−1) are observed for RG. These peaks are also observed in the spectra of MA and MA/RG composites. Different from RG, the MA and MA/RG has the other peaks of C=OH for carbosyl (1274 cm−1), C=H for benzene ring (1097 cm−1) in terephthalic acid ligand and –OH (3425 cm−1). The N2 adsorption–desorption isotherms obtained at 77 K are shown in Fig. 2d. All curves are type II isotherms typically with a type H3 hysteresis, due to the presence of mesopores. It should be noted that the N2 adsorption–desorption isotherms for MA/RG were above that of the MA and RG, indicating that the specific surface area of the material is increased after MA composit with RG. The MA/RG–15% has the highest specific surface area. The porous structure parameters and the pore size distributions were obtained from the N2 adsorption data analyzed using Barrete–Joyner–Halenda (BJH) model, and the results are summarized in Table 1. The MA/RG–15% exhibits the largest volume of micro and mesoporous, which is consistent with the result of BET surface area measurement.
The surface element of MA/RG–15% is analyzed using the X–Ray photoelectron spectroscopy (XPS) and the results were presented in Fig. 3a. The atomic content of Al, C and O is 6.35%, 63.30% and 30.35%, respectively. The binding energy of the metal A12p is 74.92 eV (Fig. 3b), which is due to the formation of AlO4(OH2) in the MIL–68(Al) framework. The O1s peak (Fig. 3c) at 532.40 eV accounts for the carboxylate oxygen −COO of terephthalic acid and the residue oxygen–containing groups in RG in the composites. This is further confirmed by the C1s band (Fig. 3d), which can be divided into three peaks located at 284.59, 285.55, and 289.70 eV, corresponding to the C=CC−C, C−O and carboxylate carbon structures, respectively.

Figure 1. The SEM images of MA (a), RG (b), MA/RG–1% (c), MA/RG–5% (d), MA/RG–15% (e,h), MA/RG–25% (f), MA/RG–35% (g); The EDX spectrum (i) and elemental mapping images (j–l) of MA/RG–15%; C (j), O (k) and Al (l); The TEM images of MA (m), RG (n) and MA/RG–15% (o).

The surface element of MA/RG–15% is analyzed using the X–Ray photoelectron spectroscopy (XPS) and the results were presented in Fig. 3a. The atomic content of Al, C and O is 6.35%, 63.30% and 30.35%, respectively. The binding energy of the metal A12p is 74.92 eV (Fig. 3b), which is due to the formation of AlO4(OH2) in the MIL–68(Al) framework. The O1s peak (Fig. 3c) at 532.40 eV accounts for the carboxylate oxygen −COO of terephthalic acid and the residue oxygen–containing groups in RG in the composites. This is further confirmed by the C1s band (Fig. 3d), which can be divided into three peaks located at 284.59, 285.55, and 289.70 eV, corresponding to the C=CC−C, C−O and carboxylate carbon structures, respectively. 
The p-nitrophenol (PNP) adsorption. Adsorption kinetics and rate-control mechanism. The effects of contact time on the PNP adsorption on the RG, MA and MA/RG composites are shown in Fig. 4a. For all samples, the PNP adsorption rates decrease with time until the adsorption equilibrium is reached. After hybridization of MA with RG, the adsorption of PNP are significantly enhanced, and the MA/RG–15% exhibits the best adsorption performance of 307.38 mg/g, which is 64% and 123% higher than that of MA and RG, respectively, due to the increase of surface area. The molecular size of PNP is calculated to be 0.66 nm × 0.43 nm, which is smaller than the average diameter of the pores (Table 1) of MA/RG composites. Therefore, the PNP molecules can easily enter into the pore and access the surface, which favors the PNP adsorption.

The Pseudo–first–order16, Pseudo–second–order40 and Elovich41 equations are used to describe the adsorption kinetics (Eqs 1–3).

\[
\log(q_e - q_t) = \log q_e - \frac{k_1}{2.303} t
\]  

\[
\frac{t}{q_t} = \frac{1}{k_2 q_e} + \frac{t}{q_e}
\]  

Table 1. Parameters of the porous structure for the MA, MA/RG and RG.

| Sample       | S_{BET} (m²/g) | V (cm³/g) | V_{mic} (cm³/g) | V_{mes} (cm³/g) | V_{mac} (cm³/g) | Pore Size (nm) |
|--------------|----------------|-----------|-----------------|-----------------|-----------------|----------------|
| RG           | 18.96          | 0.055     | 0.001           | 0.022           | 0.032           | 15.18          |
| MA           | 550.03         | 0.220     | 0.002           | 0.096           | 0.122           | 21.38          |
| MA/RG–1%     | 599.26         | 0.248     | 0.014           | 0.146           | 0.088           | 9.44           |
| MA/RG–5%     | 629.52         | 0.259     | 0.013           | 0.144           | 0.102           | 9.26           |
| MA/RG–15%    | 761.97         | 0.266     | 0.016           | 0.164           | 0.086           | 9.07           |
| MA/RG–25%    | 714.77         | 0.231     | 0.015           | 0.118           | 0.098           | 9.24           |
| MA/RG–35%    | 703.33         | 0.204     | 0.011           | 0.095           | 0.098           | 11.40          |

Figure 2. The XRD spectrum (a), Raman spectrum (b), FT–IR spectrum (c), and N₂ adsorption – desorption isotherms (d) of MA/RG composites.
Figure 3. The XPS spectra and of MA/RG–15%: (a) the full XPS spectra; the core level spectra of Al2p (b), O1s (c) and C1s (d).

Figure 4. (a) The effect of time on PNP adsorption; (b) Pseudo–second–order plots for PNP adsorption; (c) Intra–particle diffusion for PNP adsorption.
Intra–particle diffusion

\[ q_i = k_d t + C_i \]  

(4)

where \( q_i \) and \( q_e \) (mg/g) are the adsorption amount of PNP at equilibrium and time \( t \) (h), respectively. \( k_d (\text{mg} / (\text{g} \cdot \text{h}^{1/2})) \) and \( k_i (\text{h}^{-1}) \) are the Pseudo–first–order and Pseudo–second–order adsorption rate constants, respectively. \( \alpha (\text{mg/g}) \) is the initial sorption rate and \( \beta (\text{g/mg}) \) is related to the extent of surface coverage and activation energy for chemisorptions. Table 2 summarizes the adsorption kinetic parameters of PNP onto the tested adsorbents. Comparing with the correlation coefficients (\( R^2 \)) of the Pseudo–first–order, Pseudo–second–order and the Elovich models, it can be concluded that the Pseudo–second–order kinetic model fits the adsorption process of all samples better than the other two. Furthermore, the deviation between calculated \( q_{\text{cal}} \) and experimental \( q_{\text{exp}} \) values of the Pseudo–second–order kinetic model are very lower, while that of the Pseudo–first–order kinetic model is very large. The fitting line of Pseudo–second–order is perfectly plotted in Fig. 4b, suggesting that the adsorption–determining factor of the PNP removal may be involve in the chemisorption.

To better understand the diffusion rate controlling procedure, the Intra–particle diffusion model\(^{42} \) is tested as Eq. (4).

\[ q_t = \frac{1}{\beta} \ln(\alpha \beta) + \frac{1}{\beta} \ln t \]  

(3)

Adsorption isotherms and thermodynamics. The adsorption isotherm investigations are carried out under different temperature with various initial PNP concentrations. As illustrated in Fig. 5(a–c), the uptake amount of PNP increases firstly with the increase of PNP concentration and then it keeps on a horizontal (the adsorption saturation stage). The reason may be that the higher initial PNP concentration, the more strength driving force provides to overcome the mass transfer resistances when the utilization of active sites do not reach adsorption

| Parameters | Adsorbents |
|------------|------------|
| \( q_{\text{exp}} \) (mg/g) | RG | MA | MA/RG –1% | MA/RG –5% | MA/RG –15% | MA/RG –25% | MA/RG –35% |
| \( k_1 (l/h) \) | 0.19 | 0.14 | 0.18 | 0.09 | 1.79 | 0.21 | 0.10 |
| \( q_{\text{cal}} \) (mg/g) | 67.92 | 77.10 | 113.27 | 134.91 | 152.10 | 108.97 | 99.62 |
| \( R^2 \) | 0.865 | 0.925 | 0.979 | 0.965 | 0.964 | 0.907 | 0.799 |
| \( k_{2} (g/(mg.h)) \) | 2.04E–02 | 1.33E–02 | 8.77E–03 | 6.97E–03 | 7.85E–03 | 1.35E–02 | 1.59E–02 |
| \( \alpha (mg/g) \) | 139.47 | 183.82 | 202.84 | 241.55 | 304.88 | 249.38 | 234.74 |
| \( \beta (g/mg) \) | 3.47E+03 | 7.08E+06 | 6.26E+04 | 3.74E+04 | 1.23E+05 | 3.19E+04 | 1.70E+05 |
| \( R^2 \) | 0.999 | 0.995 | 0.995 | 0.993 | 0.997 | 0.999 | 0.999 |
| \( k_2 (g/(mg.h)) \) | 112.89 | 78.96 | 51.12 | 47.21 | 150.90 | 115.37 | 93.72 |
| \( C_i \) | 11.44 | 94.04 | 82.86 | 110.04 | 108.00 | 76.87 | 97.01 |
| \( (R_1)^2 \) | 0.993 | 0.936 | 0.912 | 0.939 | 0.851 | 0.941 | 0.900 |
| \( k_{3d} (mg/(g.h^{1/2})) \) | 29.75 | 13.36 | 22.23 | 31.79 | 32.99 | 25.28 | 17.78 |
| \( C_2 \) | 62.87 | 122.58 | 98.79 | 113.52 | 177.10 | 161.11 | 167.76 |
| \( (R_2)^2 \) | 0.985 | 0.973 | 0.989 | 0.991 | 0.965 | 0.983 | 0.969 |
| \( k_{3d} (mg/(g.h^{1/2})) \) | 0.19 | 9.94 | 5.60 | 11.51 | 10.07 | 0.278 | 2.51 |
| \( C_3 \) | 136.75 | 139.05 | 176.42 | 188.02 | 258.61 | 245.76 | 222.52 |
| \( (R_3)^2 \) | 0.875 | 0.907 | 0.906 | 0.900 | 0.916 | 0.998 | 0.981 |

Table 2. Adsorption kinetics parameters of PNP onto adsorbents.
saturation. Besides, for MA (Fig. 5b) and MA/RG–15% (Fig. 5c), the PNP adsorption amount increases with increase of temperature, while for RG (Fig. 5a), the adsorbed PNP is slightly decrease as increase in temperature, which suggests that an endothermic process is for MA and MA/RG–15%, but an exothermic procedure is for RG in nature. Therefore, after MA incorporated with RG, rising the temperature favors the PNP adsorption44.

To gain the insight into adsorption thermodynamic behavior, the Langmuir45 and Freundlich18 isotherm models are used to analyze the equilibrium data according to the Eqs (5) and (6), respectively.

\[
\frac{C_e}{q_e} = \frac{C_e}{q_{\text{max}}} + \frac{1}{q_{\text{max}} K_L}
\]

\[
\ln q_e = \ln K_F + \frac{1}{n} \ln C_e
\]

where, \(C_e\) is the equilibrium concentration (mg/L) of the PNP, \(q_e\) is the amount (mg/g) of the PNP adsorbed at equilibrium and \(q_{\text{max}}\) is the maximum adsorption capacity (mg/g), \(K_L\) (L/mg) is the Langmuir constants related to energy of the adsorption. \(K_F\) (L/mg) and \(1/n\) are Freundlich constants giving an indicator of the adsorption capacity and the adsorption intensity, respectively. Table 3 lists isotherm parameters of Langmuir and Freundlich

![Figure 5](image_url). The PNP adsorption by RG (a), MA (b) and MA/RG–15% (c) at different temperature; The Langmuir isotherm model for PNP adsorption by RG (d), MA (e) and MA/RG–15% (f).
isotherms for the PNP adsorption. It can be found that, for MA, RG or MA/RG, the regression coefficients $R^2$ obtained from Langmuir model are much higher than that from Freundlich isotherm model, which suggests that the adsorption of PNP is best fitted with the Langmuir isotherm and the adsorption behavior is governed by monolayer adsorption on a homogenous surface. The linear relation between $Ce/qe$ of Langmuir model is well ploted in Fig. 5(d–f). The maximum uptake capacity ($qm,cal$) calculated from Langmuir model of the MA/RG composite at different temperature are much higher than that of the parent MA and RG. The comparisons of PNP adsorption maximum capacity with various adsorbents previously reported are listed in Table 4. It can be seen that the MA/RG composite exhibits superior PNP uptake capacity than NiAl-layered double hydroxide, alumina hollow microspheres, NH2-MIL-101(Al), carbon nanotube, nanographite oxide and graphene, indicating that the MA/RG has great potentials for PNP removal from contaminated water.

To gain insight into the essential feature of Langmuir isotherm, a dimensionless constant separation factor ($RL$) is tested as the following equation 1:

$$RL = \frac{1}{1 + KcC_0}$$

where, $Kc$ (L/mg) is the Langmuir constant and $C_0$ (mg/L) is the initial concentration of PNP in the liquid phase. The value of $R_L$ indicates whether the type of the Langmuir isotherm is unfavorable ($R_L > 1$), linear ($R_L = 1$), favorable ($0 < R_L < 1$) or irreversible ($R_L = 0$). From Table 3, the $R_L$ values for MA, RG and MA/RG are between 0 and 1, especially for MA/RG, the $R_L$ values are smaller than that for the monomers, indicating that a more favorable adsorption process is for the PNP adsorption onto the MA/RG composites.

Table 3. Isotherm parameters for the adsorption of PNP onto adsorbents.

| Adsorbents | $T$ (K) | $q_{max}$ (mg/g) | $Kc$ (L/mg) | $R_L$ | $R^2$ | $1/n$ | $Kc$ (L/mg) | $R^2$ |
|------------|--------|----------------|-----------|-------|-------|-------|-----------|-------|
| RG         | 303    | 175.44        | 0.04      | 0.998 | 0.18  | 58.55 | 0.997     |       |
|            | 313    | 173.31        | 0.04      | 0.999 | 0.20  | 54.95 | 0.990     |       |
|            | 323    | 175.75        | 0.03      | 1.0   | 0.996 | 0.25  | 40.65     | 0.986  |
| MA         | 303    | 271.00        | 0.05      | 0.997 | 0.23  | 76.56 | 0.896     |       |
|            | 313    | 305.81        | 0.05      | 0.993 | 0.27  | 69.46 | 0.908     |       |
|            | 323    | 335.57        | 0.04      | 0.991 | 0.26  | 76.59 | 0.940     |       |
| MA/RG      | 303    | 332.23        | 0.14      | 0.998 | 0.19  | 122.03| 0.724     |       |
|            | 313    | 336.70        | 0.16      | 0.999 | 0.19  | 129.51| 0.681     |       |
|            | 323    | 353.36        | 0.14      | 0.998 | 0.20  | 128.05| 0.715     |       |

Table 4. Maximum adsorption capacities for PNP ontovarious adsorbents.

| Sorbents                                      | $T$ (K) | $q_{max}$ (mg/g) | Ref.  |
|-----------------------------------------------|--------|----------------|-------|
| NiAl-layered double hydroxide                 | 303    | 77.70          | 1     |
| Alumina hollow microspheres                   | 303    | 217.40         | 46    |
| NH2-MIL-101(Al)                               | 303    | 195.52         | 47    |
| Copper-based MOFs (HKUST-1)                   | 293    | 372.00         |       |
| Carbon nanotube                               | 293    | 206.00         | 3     |
| Nanographite oxide                            | 303    | 264.90         | 16    |
| Graphene                                      | 298    | 15.50          | 18    |
| Reduced graphene oxide                        | 303    | 175.44         | In this study |
| MIL-68(Al)                                    | 303    | 271.00         | In this study |
| MIL-68(Al)/Reduced graphene oxide             | 303    | 332.23         | In this study |

Table 4. Maximum adsorption capacities for PNP on various adsorbents.
As presented in Table 5, the value of $\Delta H^0$ is positive for MA and MA/RG composite, and negative for RG, which indicates that the PNP adsorption on MA and MA/RG is an endothermic nature, but on the RG is an exothermic process. This coincides with the result of temperature effects. Furthermore, the positive (or negative) value of $\Delta S^0$ reflects an increase (or decrease) in randomness at the solid/solution interface for MA and MA/RG (or RG) during the PNP adsorption process.40

### Effects of pH and ionic strength

The effect of pH ranging from 3.0 to 11.0 on the adsorption capacity ($q_e$) of MA/RG–15% is shown in Fig. 6a. The amount of PNP adsorbed on MA/RG–15% increases with increasing initial pH of up to 5, and then decreases with further increase in pH. The effect of pH on PNP adsorption by MA/RG is mainly resulted from the distribution of two PNP species, the molecular and the anionic. The PNP is a water-soluble solid that is moderately acidic in water ($pK_a = 7.15$). The molecular PNP can be effectively adsorbed onto MA/RG at a lower pH by hydrophobic interaction and $\pi-\pi$ interactions with the aromatic moieties of the PNP and the aromatic matrix of the MA/RG. However, when the pH value is below or equal to 4, a damage of MOFs starts to appear, leading to the weak binding force for PNP adsorption.32 The PNP exists as a phenolate anion when the pH is higher than the $pK_a$ (7.15) and is hydrophilic in nature, which cannot be effectively loaded onto MA/RG composite due to the weak electron accepting ability of benzene ring in PNP for donor–acceptor interaction with MA/RG. Furthermore, the zeta potentials measurement (Fig. 6a) shows that the surface charge of MA/RG is negative when pH is greater than or equal to 9, which implies that the electrostatic repulsion between PNP anions and the MA/RG surface increases with increasing pH value, resulting in the decrease of PNP uptake amount to MA/RG adsorbent.

Effect of presence of NaCl, KCl and BaCl$_2$ on the PNP adsorption is depicted in Fig. 6b. It can be seen that the amount of PNP adsorbed on MA/RG is firstly decrease (or increase), and then increase (or decrease) with the increase in the concentration of NaCl, KCl and BaCl$_2$ (or KCl). This depends on the strength of two different interaction mechanisms. On the one hand, the Na$^+$, K$^+$ and Ba$^{2+}$ are trended to compete with PNP for the active site of the MA via donor–acceptor interaction, making an adverse condition for PNP adsorption. On the other hand, a favorable factor is also provided by the salt–out effect between PNP molecule and salt. This effect illustrates that solubility of non electrolyte organic compounds in solution generally displays an inverse dependency on ionic strength. Namely, in the presence of dissolved ions, water is less ordered and compressible, and the cavity volume available to accommodate PNP molecules is reduced. Thus, the solubility of PNP neutral molecule is decreased in solution, and the more decrease in solubility, the more increase is in the sorption of PNP by enhanced diffusion.41

### Recyclability of MA/RG and PNP adsorption mechanism

The regenerability of MA/RG–15% was examined using methanol and ethanol. As seen in Fig. 6c, the PNP adsorption performance of MA/RG–15% regenerated by methanol is superior to that renewed by ethanol. After five adsorption/desorption cycles using methanol, the PNP uptake amount still remains at 271.82 mg/g, indicating that the methanol is very suitable for the regeneration of MA/RG, and the MA/RG could be a cost-effective and promising adsorbents for PNP removal. To verify the adsorption mechanisms, the FT–IR spectra (Fig. 6d) of PNP, MA/RG, mixture of MA/RG and PNP solid, and the MA/RG loaded with PNP are conducted. For PNP, the triplet bands (689, 753 and 853 cm$^{-1}$) are the ring C–H vibrations and the bands at 1101 and 1161 cm$^{-1}$ are aromatic ring vibrations; the O–H bending vibration locates at 1211 cm$^{-1}$ (and 3080 ~ 3500 cm$^{-1}$) and the phenolic C–O stretch is presented at 1337 cm$^{-1}$; the bands at 1278 and 1442 cm$^{-1}$ are due to the vibration of NO$_2$ group; the bands at 1501 and 1594 cm$^{-1}$ belong to the $\pi-\pi$ stretching of the PNP and the aromatic matrix of the MA/RG. However, when the pH value is below or equal to 4, a damage of MOFs starts to appear, leading to the weak binding force for PNP adsorption.32 The PNP exists as a phenolate anion when the pH is higher than the $pK_a$ (7.15) and is hydrophilic in nature, which cannot be effectively loaded onto MA/RG composite due to the weak electron accepting ability of benzene ring in PNP for donor–acceptor interaction with MA/RG. Furthermore, the zeta potentials measurement (Fig. 6a) shows that the surface charge of MA/RG is negative when pH is greater than or equal to 9, which implies that the electrostatic repulsion between PNP anions and the MA/RG surface increases with increasing pH value, resulting in the decrease of PNP uptake amount to MA/RG adsorbent.

Effect of presence of NaCl, KCl and BaCl$_2$ on the PNP adsorption is depicted in Fig. 6b. It can be seen that the amount of PNP adsorbed on MA/RG is firstly decrease (or increase), and then increase (or decrease) with the increase in the concentration of NaCl, KCl and BaCl$_2$ (or KCl). This depends on the strength of two different interaction mechanisms. On the one hand, the Na$^+$, K$^+$ and Ba$^{2+}$ are trended to compete with PNP for the active site of the MA via donor–acceptor interaction, making an adverse condition for PNP adsorption. On the other hand, a favorable factor is also provided by the salt–out effect between PNP molecule and salt. This effect illustrates that solubility of non electrolyte organic compounds in solution generally displays an inverse dependency on ionic strength. Namely, in the presence of dissolved ions, water is less ordered and compressible, and the cavity volume available to accommodate PNP molecules is reduced. Thus, the solubility of PNP neutral molecule is decreased in solution, and the more decrease in solubility, the more increase is in the sorption of PNP by enhanced diffusion.

### Table 5. Thermodynamic parameters for PNP adsorption.

| Adsorbents | $T$(K) | $K_c$ | $\Delta G^0$ (kJ/mol) | $\Delta S^0$ (J/(K mol)) | $\Delta H^0$ (kJ/mol) |
|------------|-------|------|----------------------|------------------------|---------------------|
| RG         | 303   | 0.56 | −3.38                | 23.51                  | −10.57              |
|            | 313   | 0.55 | −3.34                |                        |                     |
|            | 323   | 0.48 | −2.91                |                        |                     |
| MA         | 303   | 1.19 | −4.90                | 53.41                  | 11.36               |
|            | 313   | 1.20 | −5.21                |                        |                     |
|            | 323   | 1.26 | −5.97                |                        |                     |
| MA/RG      | 303   | 1.93 | −6.40                | 35.38                  | 4.31                |
|            | 313   | 2.05 | −6.78                |                        |                     |
|            | 323   | 2.10 | −7.11                |                        |                     |

As presented in Table 5, the value of $\Delta H^0$ is positive for MA and MA/RG composite, and negative for RG, which indicates that the PNP adsorption on MA and MA/RG is an endothermic nature, but on the RG is an exothermic process. This coincides with the result of temperature effects. Furthermore, the positive (or negative) value of $\Delta S^0$ reflects an increase (or decrease) in randomness at the solid/solution interface for MA and MA/RG (or RG) during the PNP adsorption process.40

### Recyclability of MA/RG and PNP adsorption mechanism

The regenerability of MA/RG–15% was examined using methanol and ethanol. As seen in Fig. 6c, the PNP adsorption performance of MA/RG–15% regenerated by methanol is superior to that renewed by ethanol. After five adsorption/desorption cycles using methanol, the PNP uptake amount still remains at 271.82 mg/g, indicating that the methanol is very suitable for the regeneration of MA/RG, and the MA/RG could be a cost-effective and promising adsorbents for PNP removal. To verify the adsorption mechanisms, the FT–IR spectra (Fig. 6d) of PNP, MA/RG, mixture of MA/RG and PNP solid, and the MA/RG loaded with PNP are conducted. For PNP, the triplet bands (689, 753 and 853 cm$^{-1}$) are the ring C–H vibrations and the bands at 1101 and 1161 cm$^{-1}$ are aromatic ring vibrations; the O–H bending vibration locates at 1211 cm$^{-1}$ (and 3080 ~ 3500 cm$^{-1}$) and the phenolic C–O stretch is presented at 1337 cm$^{-1}$; the bands at 1278 and 1442 cm$^{-1}$ are due to the vibration of NO$_2$ group; the bands at 1501 and 1594 cm$^{-1}$ belong to the $\pi-\pi$ stretching of the PNP and the aromatic matrix of the MA/RG. However, when the pH value is below or equal to 4, a damage of MOFs starts to appear, leading to the weak binding force for PNP adsorption.32 The PNP exists as a phenolate anion when the pH is higher than the $pK_a$ (7.15) and is hydrophilic in nature, which cannot be effectively loaded onto MA/RG composite due to the weak electron accepting ability of benzene ring in PNP for donor–acceptor interaction with MA/RG. Furthermore, the zeta potentials measurement (Fig. 6a) shows that the surface charge of MA/RG is negative when pH is greater than or equal to 9, which implies that the electrostatic repulsion between PNP anions and the MA/RG surface increases with increasing pH value, resulting in the decrease of PNP uptake amount to MA/RG adsorbent.

Effect of presence of NaCl, KCl and BaCl$_2$ on the PNP adsorption is depicted in Fig. 6b. It can be seen that the amount of PNP adsorbed on MA/RG is firstly decrease (or increase), and then increase (or decrease) with the increase in the concentration of NaCl, KCl and BaCl$_2$ (or KCl). This depends on the strength of two different interaction mechanisms. On the one hand, the Na$^+$, K$^+$ and Ba$^{2+}$ are trended to compete with PNP for the active site of the MA via donor–acceptor interaction, making an adverse condition for PNP adsorption. On the other hand, a favorable factor is also provided by the salt–out effect between PNP molecule and salt. This effect illustrates that solubility of non electrolyte organic compounds in solution generally displays an inverse dependency on ionic strength. Namely, in the presence of dissolved ions, water is less ordered and compressible, and the cavity volume available to accommodate PNP molecules is reduced. Thus, the solubility of PNP neutral molecule is decreased in solution, and the more decrease in solubility, the more increase is in the sorption of PNP by enhanced diffusion.41
well known that such relationship can be affected by the availability of electron density in the donor and the electron affinity of the acceptor. The nitro–substituted benzene ring can act as electron acceptor due to the low electron density caused by strong electron withdrawing ability of NO$_2$ group reducing the overall electron density in the \( \pi \)-system of the aromatic ring\(^3\). As shown in Fig. 6e, the adsorption yield of nitrophenols and phenol onto MA/RG composite in case of single component is in the order of o–nitrophenol (ONP) \( > \) PNP \( > \) m–nitrophenol (MNP) \( \gg \) phenol, indicating that the nitro substituent causes the increment in electronic acceptance of aromatic ring on nitrophenol. In fact, the \( \pi \)-electron–rich regions in graphene layers also can interact with electron acceptor substance by \( \pi-\pi \) dispersion interaction\(^4\). For the multi aperture adsorbent, it is often assumed that the \( \pi-\pi \) interaction is stronger in the small pores. This agrees with the pore size distribution (Table 1) that the larger volume of micropore, the higher adsorption capacity is for PNP onto MA/RG samples. Besides, the C=C vibrations (Fig. 6d) in aromatic ring at 1587 and 1594 cm\(^{-1}\) for MA/RG and PNP are located in the uniform at the band of 1600 cm\(^{-1}\), indicating that the electron–rich regions in graphene layers interact with the \( \pi \) electron of the aromatic ring of PNP via stacking the center of the aromatic ring of the molecule on top of a graphene carbon atom and the benzene ring of the PNP on top of the graphene hexagon\(^1\). Owe to such force existence (Fig. 6f), it is possible to simultaneously remove isomeric nitrophenols from water. As presented in Fig. 6e, in the binary system (ONP + PNP, MNP + PNP), the MA/RG composite exhibits excellent affinity to each component with negligible change compared to that in single solution.
Conclusions
The MIL–68(Al)/reduced graphene oxide (MA/RG) composite is successfully synthesized via a simple solvo-
thermal method. The presence of reduced graphene oxide (RG) changes the morphology and surface area of
composites, but not the crystalline structure. The surface area of composite is firstly increase and then decrease
with increasing RG, and the PNP adsorption capacity exhibits the same trend with the maximum uptake calcu-
lated from Langmuir model of 332.23 mg/g for MA/RG–15% at 303 K, which is much super than the MA and RG
individual. This good performance is linked to improvement of porosity, the hydrogen bond and π–π dispersion
interaction between PNP and the composite. The solution pH, ionic strength, temperature and initial PNP con-
centration extremely affect the PNP adsorption, but the presence of phenol and isomerism nitrophenols has a
slightly influence on PNP removal. The adsorption process involved in film–diffusion and intra–particle diffusion
bays well with the Pseudo–second–order model and the Langmuir model. The coupling of MOFs with reduced
graphene oxide provides a favorable pathway to synthesis high reusability and effective adsorbent forsimulta-
aneous removal of nitrophenols from wastewater.

Methods
Materials. Graphite powder (particle size < 30 µm) was purchased from Tianjin Kermer Chemical Regent
Ltd (Tianjin, China). Terephthalic acid (99%), N, N–Dimethylformamide (DMF) and phenol (99.5%) was from
Synpharm Chemical Regent Co, Ltd (Shanghai, China). The AlCl₃·6H₂O was from Tianjin Hengxing Chemical
Reagent Co, Ltd (Tianjin, China). The p–nitrophenol (PNP, 99.8%) was from Shanghai Shan Pu Chemical Co,
Ltd (Shanghai, China). The o–nitrophenol (ONP, 98%) and m–nitrophenol (MNP, 99%) were from Xiya chemical
industry Co, Ltd (Shandong, China). All other chemicals (analytical grade) were used without any other purifica-

Synthesis of MIL–68 (Al)/reduced graphene oxide composites. The MIL–68(Al), or MA, was
synthesized based on the method described by Yang et al. The graphene oxide was prepared by a modified
Hummers method. The MIL–68(Al)/reduced graphene oxide (MA/RG) composite was synthesized by a simple
one–pot solvothermal method. Specifically, a certain amount of graphene oxide was added into 120 mL DMF and
the mixture was sonicated for 30 min to form the brownish yellow suspension. Then 2 g of terephthalic acid and
1.95 g of AlCl₃·6H₂O were added. After the regents dissolved, the mixture was transferred to a 500 mL of single
neck ground–in round flask with an Allihn condenser on top. The flask was set in an oil bath for reaction at 403 K
18.5 h. After that, the suspension was centrifuged, and the precipitation was washed with DMF and methanol four
times. Finally, the product was dried at 423 K under vacuum for 3 h. The final products were labeled as
MA/RG–1%, MA/RG–5%, MA/RG–15%, MA/RG–25% and MA/RG–35% with the ratio of RG to MIL–68(Al)
at 0.01, 0.05, 0.15, 0.25 and 0.35, respectively. The reduced graphene oxide (RG) was synthesized via solvothermal
method without terephthalic acid and AlCl₃·6H₂O addition.

Characterization methods. The surface morphologies of the as–prepared materials were observed using the
Environmental scanning electron microscope (SEM, FEI QuANTA 200, USA) equipped with an energy
dispersive X–ray (EDX) spectroscopy, and Transmission electron microscopy (TEM, Tecnai G2F20 S–TWIN,
USA). The X–ray diffraction (XRD) patterns were obtained using Bruker AXS D8 Advance diffractometer with
Cu–Ka beam source (λ = 1.541 Å). Raman spectra were obtained using JobinYvon Micro–Raman Spectroscopy
(RamLab–010), equipped with a holographic grating of 1800 lines/mm and a He–Ne laser (633 nm, spot size ~
1 µm) as excitation source. Fourier transform infrared spectrum (FT–IR) measurements were conducted by using
Nicolet 5700 Spectrometer in KBr pellet at room temperature. The N₂ adsorption–desorption for Brunauer–
Emmette–Teller (BET) specific surface area and pore size measurement were conducted by using automatic sur-
face analyzer (Quantachrome, USA). The surface elemental composition analyses were conducted based on the
XPS spectra (Thermo Fisher Scientific–K–Alpha 1063, UK) with a resolution of 0.5 eV. The zeta potentials of MA/
RG particles in solutions at pH from 3.0 to 11.0 (adjust by NaOH or HCl) was determined with a zeta potential
meter (Zetasizer Nano–ZS90, Malvern).

Adsorption of p–nitrophenol. Batch experiments were conducted to determine the kinetics and isotherms
for p–nitrophenol (PNP) adsorption by MA/RG, and the factors affecting the adsorption as well. All the tests were
performed in triplicates. For adsorption kinetics study, the 100 mg of adsorbents was added into 250 mL PNP solu-
tion with PNP concentration of 200 mg/L. The mixture was shaken on a gyratory shaker at 303 K and 160 rpm.
At predetermined time intervals (from 1 min to 24 h), 4 mL of mixture was withdrawn using a 5 mL of pipette. The
experiments of adsorption isotherms were conducted at different temperatures (298, 308 and 318 K), for which
5 mg of adsorbents (RG, MA or MA/RG–15%) adding to 25 mL of solution with PNP concentration between
50 mg/L and 300 mg/L. The effect of pH on the PNP adsorption was examined by agitating the mixture of 5 mg
of adsorbents (RG, MA or MA/RG–15%) and 25 mL of PNP solution (100 mg/L) at 303 K under 160 rpm. The influence of phenol, ONP or MNP on PNP adsorption at 303 K were investigated using 30 mL of the solution that contains 50 mg/L phenol (or ONP, or MNP)
and 50 mg/L PNP and 10 mg of MA/RG–15%. For adsorption mechanism analysis, the solution containing only
phenol, ONP, MNP or PNP were used. The regeneration of MA/RG was conducted by agitating PNP-loaded MA/
RG in absolute methanol (or ethanol) with solid concentration of 500 mg/L at 303 K for 3 h. Adsorption tests on
renewed MA/RG were performed at 303 K in the solution containing 200 mg/L of PNP and 500 mg/L of MA/RG.
After the solid–liquid separation at the end of adsorption, the concentrations of phenol, ONP, MNP and PNP in
the supernatant was measured using UV spectrophotometer (UV–2550, SHIMADZU, Japan) at 270, 278, 273 and 317 nm, respectively. The adsorptive quantity was calculated as follows:

\[ q_e = \frac{C_0 - C_t}{m} \cdot V \]  

(10)

where \( q_e \) (mg/g) is the adsorption quantity; \( C_0 \) and \( C_t \) are the pollution concentration of initial and interval time (mg/L), respectively; \( V \) (L) is the volume of solution; and \( m \) (g) is the weight of adsorbent.

References
1. Sun, Y., Zhou, J., Cai, W., Zhao, R. & Yuter, J. Hierarchically porous NiAl-LDH nanoparticles as highly efficient adsorbent for p-nitrophenol from water. Appl Surf Sci 349, 897–903 (2015).
2. Xue, G., Gao, M., Gu, Z., Luo, Z. & Hu, Z. The removal of p-nitrophenol from aqueous solutions by adsorption using gemini surfactants modified montmorillonites. Chem Eng J 218, 223–231 (2013).
3. Wang, J.-J. et al. Removal and Adsorption of p-Nitrophenol from Aqueous Solutions Using Carbon Nanotubes and Their Composites. J Nanomater 2014, 1–9 (2014).
4. Hamidouche, S. et al. Simultaneous sorption of 4-nitrophenol and 2-nitrophenol on a hybrid geocomposite based on surfactant-modified pillared-clay and activated carbon. Chem Eng J 279, 964–972 (2015).
5. Eckenfelder, W. W. Industrial Water Pollution Control Vol. 1 (ed W. W. Eckenfelder) Ch. 1, 1–4 (McGraw-Hill Science, 1999).
6. Tuo, Y. et al. Microbial synthesis of Pd/Fe3O4, Au/Fe3O4, and PdAu/Fe3O4 nanocomposites for catalytic reduction of nitroaromatic compounds. Sci Rep 5, 15315 (2015).
7. Jing, Q., Yu, Z., Lin, D., Zhu, L. & Yang, K. Enhanced sorption of naphthalene and p-nitrophenol by nano-SiO2 modified with a cationic surfactant. Water Res 47, 4006–4012 (2013).
8. Ribeiro, R. S., Silva, A. M. T., Figueiredo, J. L., Faria, J. L. & Gomes, H. T. Removal of 2-nitrophenol by catalytic wet peroxide oxidation using carbon materials with different morphological and chemical properties. Appl Catal B: Environ 140–141, 356–362 (2013).
9. Rybas, G. Z., Deliyanni, E. A. & Matis, K. A. Graphene oxide and its application as an adsorbent for wastewater treatment. J Chem Technol Biotechnol 89, 196–205 (2014).
10. Wang, F. et al. A graphene oxide/amidoxime hydrogel for enhanced uranium capture. Sci Rep 6, 19367 (2016).
11. Wang, H. et al. Three dimensional graphene based materials: Synthesis and applications from energy storage and conversion to electrochemical sensor and environmental remediation. Adv Colloid Interface Sci 221, 41–59 (2015).
12. Wang, H. et al. Graphene-based materials: fabrication, characterization and application for the decontamination of wastewater and wastegases and hydrogen storage/generation. Adv Colloid Interface Sci 195–196, 19–40 (2013).
13. Wang, X. et al. Correlation between the adsorption ability and reduction degree of graphene oxide and tuning of adsorption of phenolic compounds. Carbon 69, 101–112 (2014).
14. Wang, H. et al. Adsorption characteristics and behaviors of graphene oxide for Zn(II) removal from aqueous solution. Appl Surf Sci 279, 432–440 (2013).
15. Wu, Z. et al. Adsorptive removal of methylene blue by rhamnolipid-functionalized graphene oxide from wastewater. Water Res 67, 330–344 (2014).
16. Zhang, B., Li, F., Wu, T., Sun, D. & Li, Y. Adsorption of p-nitrophenol from aqueous solutions using nanographite oxide. Colloids Surf A: Physicochem Eng Asp 464, 78–88 (2015).
17. Gao, W., Alemany, L. B., Ci, L. & Ajayan, P. M. New Insights into the Structure and Reduction of Graphite oxide. Nat Chem 1, 403–408 (2009).
18. Ismail, A. I. Thermodynamic and kinetic properties of the adsorption of 4-nitrophenol on graphite from aqueous solution. Can J Chem 109, 1139 (2015).
19. Qiu, X., Wang, X. & Li, W. Y. Controlled growth of dense and ordered metal–organic framework nanoparticles on graphene oxide. Chem Commun 60, 1–3 (2012).
20. Bandosz, T. J. & Petit, C. MOF/graphite oxide hybrid materials: exploring the new concept of adsorbents and catalysts. Adsorption 17, 5–16 (2010).
21. Wang, H. et al. Facile synthesis of amino-functionalized titanium metal-organic frameworks and their superior visible-light photocatalytic activity for Cr(VI) reduction. J Hazard Mater 286, 187–194 (2015).
22. Seoane, B., Sebastian, V., Tejeda, C. & Coronas, J. Crystallization in THF: the possibility of one-pot synthesis of mixed matrix membranes containing MOF MIL-68(Al). CrystEngComm 15, 9483 (2013).
23. Yang, Q. et al. Probing the adsorption performance of the hybrid porous MIL-68(Al): a synergetic combination of experimental and modelling tools. J Mater Chem 22, 10210 (2012).
24. Applied Catalysis B: Xie, L., Liu, D., Huang, H., Yang, Q. & Zhong, C. Efficient capture of nitrobenzene from waste water using metal–organic frameworks. Chem Eng J 246, 142–149 (2014).
25. Han, T. et al. Synthesis of CNT@MIL-68(Al) composites with improved adsorption capacity for phenol in aqueous solution. Chem Eng J 275, 134–141 (2015).
26. Ahmed, I. & Jiang, S. H. Composites of metal–organic frameworks: Preparation and application in adsorption. Mater Today 17, 136–146 (2014).
27. Huang, W. et al. Preparation and Adsorption Performance of GrO@Cu-BTC for Separation of CO2/CH4. Ind Eng Chem Res 53, 11176–11184 (2014).
28. Ahmed, I., Khan, N. A. & Jiang, S. H. Graphite oxide/metal-organic framework (MIL-101): remarkable performance in the adsorptive denitrogenation of model fuels. Ind Eng Chem 52, 1415–14161 (2013).
29. Zhou, X. et al. A novel MOF/graphene oxide composite GrO@MIL-101 with high adsorption capacity for acetone. J Mater Chem A 2, 4722 (2014).
30. Liu, S. et al. Nanosized Cu-MOFs induced by graphene oxide and enhanced gas storage capacity. Energy Environ Sci 6, 818 (2013).
31. Wang, H. et al. Synthesis and applications of novel graphitic carbon nitride/metal-organic frameworks mesoporous photocatalyst for dyes removal. Appl Catal B: Environ 174–175, 445–454 (2015).
32. Zu, D.-D., Lu, L., Liu, X.-Q., Zhang, D.-Y. & Sun, L.-B. Improving Hydrothermal Stability and Catalytic Activity of Metal–Organic Frameworks by Graphite Oxide Incorporation. J Phys Chem C 118, 19910–19917 (2014).
33. Banerjee, P. C. et al. Electrochemical capacitance of Ni-doped metal organic framework and reduced graphene oxide composites: more than the sum of its parts. ACS Appl Mater Interfaces 7, 3655–3664 (2015).
34. Chen, W. & Yan, L. Preparation of graphene by a low-temperature thermal reduction at atmosphere pressure. Nanoscale 2, 559–563 (2010).
35. Petit, C., Mendoza, B. & Bandosz, T. J. Hydrogen sulfide adsorption on MOFs and MOF/graphite oxide composites. Chemphyschem 11, 3678–3684 (2010).
Enhanced adsorptive removal of p-nitrophenol from water by aluminum oxide–organic framework composites: Wu, Z.

Author Contributions
The authors Z.W. and X.Y. designed experiments and wrote the manuscript; Z.W., X.Y. and H.Z. carried out experiments; Z.W. and H.W. analyzed adsorption experimental results. G.Z. and X.C. analyzed characterization data. H.W., L.Z. and J.S. assisted with adsorption experiments. All authors discussed the results and commented on the manuscript.

Additional Information
Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Wu, Z. et al. Enhanced adsorptive removal of p-nitrophenol from water by aluminum oxide–organic framework composites. Sci. Rep. 6, 25638; doi: 10.1038/srep25638 (2016).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third-party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/