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applications†

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By condensing M and TFP under solvothermal conditions, a new porous organic polymer POP_{M-TFP} was obtained. The electrode modified with triazine containing POP_{M-TFP} exhibits well-defined rapid redox processes and showed a high specific capacitance of 130.5 F g^{-1} at 2 A g^{-1} , suggesting well electrochemical performance.

Preparation of triazine containing porous organic

polymer for high performance supercapacitor

Due to the overconsumption of oil and coal over the past few decades, it is highly desirable to develop safe sources of energy, as well as the eco-friendly use of conventional energy through energy storage devices. 1,2 Supercapacitors have attracted much attention because of their high power density, long recycle life and a wide temperature range.3 Generally, supercapacitors are classified into two types: electrochemical double layer capacitor and pseudocapacitor.4-6 Electrode materials are the key ingredient for this electrochemically occurring process.7 The applied voltage depended ion adsorption on the electrode surface depending on the structure of the material used for the electrodes. To obtain a device with good performance, significant study has been paid to the exploration of redox-active materials for supercapacitors.8,9 In order to improve the electronic performance of electrode materials, the functionalized porous structure having a high surface area and hierarchical porosity is important.10

Nanoporous organic polymers with high nitrogen content have got great interest because of their applications in semi-conductors, gas storage, and electrochemical supercapacitors *etc.*¹¹⁻¹³ Recently, nitrogen-doped carbon materials exhibit good electrical properties because the pseudocapacitance effect provided by N atoms.¹⁴ Nanoporous organic polymers, such as covalent organic frameworks (COFs), covalent triazine-based frameworks (CTFs), porous organic polymers (POPs) which have controllable pore structure and heteroatom doping are promising electrode materials in supercapacitors.¹⁵⁻¹⁹ Among them, POPs which are built by the polymerization of rigid organic molecules showing good supercapacitor applications

Nowadays, nitrogen-enriched polymers attract increasing attention for application in supercapacitors.²⁵ It is a challenge to synthesize nitrogen-enriched nanoporous polymers by using cheap reagents.26 POPs or COFs are prepared by using different synthetic method such as ultrahigh vacuum surface synthesis, 27 microwave assisted solvothermal synthesis, 28 ionthermal synthesis29 and vapor-assisted conversion etc.30 Solvothermal synthesis often used to produce nanoporous materials through exploring reaction conditions appropriately.31-33 CTFs represent a class of functionalized polymers which are obtained by the trimerization of aromatic nitriles under ionthermal conditions in the presence of a catalyst. This method involves harsh experimental conditions such as very high temperature.29 Melamine is a trimer of aromatic nitriles, with a 1,3,5-triazine skeleton considered as Lewis base.34 The lone pair of electrons in N plays an important role in electrochemical applications. Introducing triazine skeleton into POP materials by using precursor bearing triazine moieties is a straightforward approach in POP synthesis.

Herein, we choose a trigonal-symmetrical nitrogencontaining melamine (M) as a monomer, which can be used to construct conjugated microporous covalent triazine-based organic polymer through the coupling with triformylphloroglucinol (TFP) monomer under solvothermal conditions without any catalysts. The doping of nitrogen atoms in POP_{M-TFP} can affect the electron distribution of the materials and further enhance the wettability and the electro-active surface area of the electrode. Otherwise, the nitrogen atom can also introduce pseudocapacitance to the supercapacitors, which are beneficial for enhancing the supercapacitor performance.

are TaPa-Py COF, HCPANIs, TpDAB and Aza-CMPs *etc.*²⁰⁻²³ POPs have micropores with high specific surface area and heteroatom doping, which should be quite conductive to the electrochemical performance when applied as electrode materials. Consequently, polytriazines with high surface area stand out due to their high nitrogen content.²⁴

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Scheme 1 Schematic representation of the structure of POP_{M-TFP} .

Scheme 1 illustrates the synthesis of POP_{M-TFP} at 160 °C. The synthesized POP_{M-TFP} was employed as an electrode material for supercapacitor application. The electrodes which are modified with POP_{M-TFP} exhibit well-defined rapid redox processes and high capacitance.

The crystalline structure of POP_{M-TFP} was first characterized by powder X-ray diffraction (XRD). The XRD pattern of POP_{M-TFP} material has been shown in Fig. 1a which reveals the crystalline nature of the material. As seen from Fig. 1a, the peaks at 5.2° , 9.5° and 27.6° are attributed to the corresponding (100), (200) and (201) planes diffractions of POP_{M-TFP} respectively. Successful condensation of M and TFP under a simple solvothermal conditions without any catalysts yields product, which has been confirmed by ^{13}C CP-MAS NMR and The Fourier transform infrared (FT-IR) spectroscopic investigations. In order to further understand the chemical environment of various types of carbon atoms in POP_{M-TFP} , ^{13}C solid state NMR spectroscopy was performed. POP_{M-TFP} exhibits a clear signal at

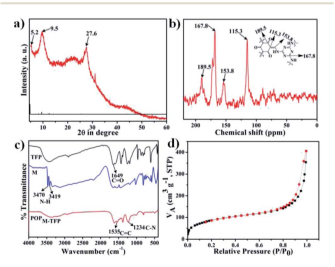


Fig. 1 (a) XRD pattern of POP_{M-TFP} (red) and simulated XRD pattern (black) of POP_{M-TFP}. (b) ^{13}C NMR spectra of POP_{M-TFP}. (c) FT-IR spectrum of M, TFP, and POP_{M-TFP}. (d) Nitrogen adsorption/desorption isotherm pattern of POP_{M-TFP} where black square indicates the adsorption and red circle represents the desorption isotherm.

167.8 ppm confirms the presence of the triazine skeletons (Fig. 1b). $^{35-37}$ The additional characteristic 13 C resonance in the C–N was observed to be located at 153.8 ppm. The characteristic peak of C=O appearing at 189.5 ppm. In summary, it can be confirmed that the M has successfully reacted with TFP to form POP_{M-TFP} material.

The FT-IR spectra of M, TFP and POP_{M-TFP} between 4000 to 400 cm⁻¹ are shown in Fig. 1c. The two separated peaks at 3470 and 3419 cm⁻¹ originate from stretching vibrations of amine (-NH₂) groups in M. The peak at 1649 cm⁻¹ belongs to stretching vibrations of C=O in TFP. The FT-IR spectra of POP_{M-TFP} shows the disappearance of the -NH₂ stretching bands of the M. The appearance of a broad -NH peak at 3379 cm⁻¹ further indicates the formation of POP_{M-TFP}. Additionally, the emergence of a new band at 1234 cm⁻¹ is characteristic of C-N stretch moieties. The peak at 1535 cm⁻¹ belongs to stretching vibrations of C=C in POP_{M-TFP}. 16,38 Further, the chemical environment of the elements present in the specimen was studied using X-ray photoelectron spectroscopy (XPS). Full scan XPS spectra of POP_{M-TFP} showed in Fig. S1[†] reveals the peaks of C, N and O. To analyze the detailed chemical states of N elements, the high-resolution XPS scan was also recorded (Fig. S2†). The N 1s spectra of POP_{M-TFP} split into two peaks located at 398.8 eV and 399.8 eV, respectively. The peak at 398.8 eV can be assigned to triazine N, while the peak at 399.8 eV can be assigned to the N in the -C=C-NH-. These peak positions are in agreement with previous reports. 20,35 The N_2 adsorption/desorption was carried out for POP_{M-TEP} in order to evaluate the permanent porosity. The N2 sorption isotherm measured at 77 K as showed in Fig. 1d. POP_{M-TEP} exhibits characteristics of type-I adsorption isotherms with sharp uptake at a low pressure region indicating the microporous nature of this material. The Brunauer-Emmett-Teller (BET) surface area of POP_{M-TFP} was found to be 317 m² g⁻¹. The pore size distribution plot has been demonstrated in Fig. S3.† The two sharp peaks at 1.5 and 2.5 nm are observed from the pore size distribution plot. Micropores of dimension 1.5 nm could be attributed to the porous network of the POP_{M-TFP}, whereas mesopores of 2.5 nm could be originated due to void spaces associated to the grain boundary.

The morphology of the synthesized POP_{M-TFP} can be well investigated by field emission scanning electron microscopy (FE-SEM) and transmission electron microscope (TEM) analysis respectively. FE-SEM images of POP_{M-TFP} at two different magnifications are shown in Fig. 2a and b. SEM images show that POP_{M-TFP} reveals irregular nanowires morphologies. The TEM characterization further confirms the nanowire structures morphology with porous architecture (Fig. 2c and d.). Fig. 2d represents the high-resolution TEM image of POP_{M-TFP} where the micropores of about 1.5 nm (white spot) are spread throughout the specimen.

Considering the synthesized POP_{M-TFP} with 1,3,5-triazine skeleton distributed in nanopores, this may hold promising applications in electrode materials for supercapacitors. The pore channels and triazine units in POP_{M-TFP} could render this material able to store electric energy. The electrochemical performance of POP_{M-TFP} was investigated by using a typical

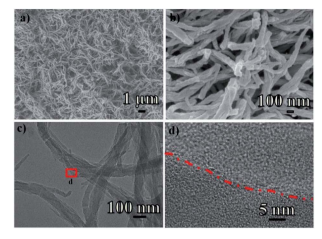


Fig. 2 (a and b) FE-SEM images of POP $_{\rm M-TFP}$ at two different magnifications. (c and d) TEM images of POP $_{\rm M-TFP}$ at two different magnifications.

three-electrode system in 2 M KOH aqueous electrolyte. The working electrodes were fabricated by mixing POP_{M-TFP}, acetylene black and binder (polyvinylidene fluoride) in a weight ratio of 80:10:10 and loaded to a piece of nickel foam. A Pt plate and Hg/HgO electrode were used as the counter electrode and reference electrode, respectively. The electrochemical charge storage activity of the POP_{M-TFP} was analysed by cyclic voltammetry (CV), galvanostatic charge discharge (GCD) and electrochemical impedance spectroscopy (EIS) in 2 M KOH electrolyte. The CV curves of POP_{M-TFP} at 5 mV s⁻¹, 10 mV s⁻¹, 20 mV s⁻¹, 50 mV s⁻¹, and 100 mV s⁻¹ are presented in Fig. 3a. The CV recorded with the as-prepared electrode over different scan rates shows redox peaks at ~0.34-0.45 V during forward and backward sweeps, suggesting its pseudo capacitive nature. The redox signature is attributed to the oxidation and reduction of the triazine skeleton in the polymer. The small separation between cathodic and anodic peaks indicated that the fast electron transfer kinetics between POP_{M-TFP} and the electrode interface. The current plateau of CV over the increasing scan

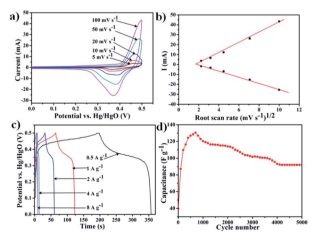


Fig. 3 (a) CV curves of POP_{M-TFP} at different scan rates in 2 M KOH electrolyte. (b) Peak current $vs.\ mV^{1/2}$ plots of POP_{M-TFP}. (c) GCD curves of POP_{M-TFP} at different current densities. (d) Cycling performance of the electrochemical capacitor based on POP_{M-TFP} at 2 A g $^{-1}$.

rate increased, suggesting fast accessible ions electrolyte through the pore channel of POP_{M-TFP} .

The specific capacitance of the electrode materials is estimated from the follow equation,³⁹

$$C_{\rm sp} = (\int I \, \mathrm{d} V) / v m V \tag{1}$$

where I represents current, V represents the potential window (V), ν indicates scan rate and m stands for the mass of active material in the electrode. The estimated specific capacitance of POP_{M-TFP} material was found to be 105.8 F g^{-1} at 5 mV s⁻¹ scan rate (Fig. S4†). With the increase of scan rate the specific capacitance was gradually decreased down. This may be due to the restriction of ion insertion into the pore channels and weak redox kinetics of POP_{M-TFP} at fast scan rates. The appearance of small humps in CV curves around 0.34-0.45 V can be attributed to the redox reactions of the heteroatom (N) functionalities of the POP_{M-TFP} and nickel foam. Additionally, the N atoms embedded in POP_{M-TFP} are desirable to improve the capacitance by enhancing the electrolyte solution wettability of the electrode materials. 40 At the slow scan rates, ions have sufficient time to arrive at the surface of POP_{M-TFP}, can be able to penetrate through channel of POP_{M-TFP}. Moreover, the peak current is proportional to the root scan rate over the range of 5 to 100 mV s⁻¹, implies that the redox reaction is diffusioncontrolled electron-transfer process presented in Fig. 3b.41 A more detailed investigations on electronic performance for its applicability as a supercapacitor electrode material is analyzed by GCD. The GCD curves of POP_{M-TFP} at various current densities (0.5 to 8 A g^{-1} , Fig. 3c) corroborate with the above CV results and display clear voltage plateaus at the same voltage position. The nonlinearity of the charge-discharge curve verifies the pseudo capacitance behavior of POP_{M-TFP} due to faradic reaction occurs at the electrode surface. Their symmetric shapes reveal the fast reaction kinetics and high electrochemical reversibility. 42 An almost negligible IR drop was observed in the GCD curve indicating a minute ohmic resistance and good capacitive performance. The specific capacitance of POP_{M-TFP} over different current densities is estimated from eqn (2),

$$C_{\rm sp} = \frac{(I \times \Delta t)}{\Delta V \times m} \tag{2}$$

where I is current, ΔV represents potential window (V), Δt signifies discharge time (s) and m represents the mass of active material. The highest specific capacitance of POP_{M-TFP} is calculated to be $130.5~F~g^{-1}$ at $2~A~g^{-1}$. The cycle performance of POP_{M-TFP} was evaluated by GCD test at $2~A~g^{-1}$ as showed in Fig. 4d. The highest specific capacitance $130.5~F~g^{-1}$ achieved after 130 cycling. The high specific capacitance of POP_{M-TFP} is attributed to better utilization of the porous structure, presence of triazine units, which could provide numerous active sites for ion transfer. It shows a slow loss of the specific capacitance, which reaches $120.2~F~g^{-1}$ after 1000 cycles. Long-term GCD cycling revealed a decrease of specific capacitance could be ascribed to the collapse of the electrode structures. The specific capacitances of POP_{M-TFP} were also calculated from GCDs and

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Fig. 4 Electrochemical performance of POP_{M-TFP} in an asymmetry supercapacitor cell in 2 M KOH. (a) CV curves of POP_{M-TFP} at different scan rates in the potential window of 0–1.5 V. (b) GCD profiles at various current densities. (c) Corresponding specific capacitances of the asymmetric cell. (d) Cycling performance test of POP_{M-TFP} at 0.5 A g^{-1} .

compared as showed in Fig. S5.† The highest specific capacitance of POP_{M-TFP} is calculated to be 178.0 F g^{-1} at 0.5 A g^{-1} , 137.4 F g^{-1} at 1 A g^{-1} , 130.5 F g^{-1} at 2 A g^{-1} , 86.4 F g^{-1} at 4 A g^{-1} and 76.9 F g⁻¹ at 8 A g⁻¹ respectively. With increasing the current density, the specific capacitance of POP_{M-TFP} tends to decrease. A slight decrease in the specific capacitance values with the increased current density is believed to be due to the hierarchical pore structure, which facilitates the smooth transfer of ions at higher current densities. Additionally, most of the active sites triazine moieties on the electrode surface are accessible to electrolyte. This material has the advantage as electrode material because of inherent stacking behavior between two adjacent layers, which can provide a nanoscale channel to the stacking direction. Owing to the presence of stacking phenomena, the presence of triazine moieties in the POP_{M-TFP} can expedite the ion transfer through porous channels. Extended π -conjugation and ion conduction inside the pores throughout POP_{M-TFP}, the redox processes arising from triazine units could be responsible for this high supercapacitor performance.

Furthermore, the interfacial charge transfer activity and electrocapacitive characteristics of POP_{M-TFP} are studied by EIS analysis. The impedance characteristic of the electrode is represented by a Nyquist plot (Fig. S6†). From the analysis of the Nyquist plots, POP_{M-TFP} displays a near vertical line within the low-frequency region, indicative of a well capacitive behavior. The slope of the line at low frequency region is related to the diffusive resistivity of the electrolyte ions within the nanopores. The small semicircular shapes in the EIS plots in the high frequency could be neglected indicates a low charge transfer resistance at the electrode/electrolyte interface (inset in Fig. S3†).⁴³ We also estimated the accessible electro-active sites of POP_{M-TFP} system by the method already discussed in the previous literature.^{16,20} These calculations suggest the about 1.99% of the triazine moieties are accessible.

To evaluate the energy and power density, the supercapacitor performance of POP_{M-TFP} was tested in two electrode system using 2 M KOH as electrolyte. Activated carbon and as-prepared POP_{M-TFP} were used as negative and positive electrodes in the asymmetric supercapacitor fabrication, respectively. The asymmetric supercapacitor was investigated in the potential window of 0 to 1.5 V at different scan rates (Fig. 4a). The current increases with increasing scan rate suggesting a good rate capability for energy storage. The CV curves and GCD curves (Fig. 4b) indicated their typical pseudo capacitive behaviour. The calculated specific capacitances of the supercapacitor at different current densities are summarized in Fig. 4c. The specific capacitance at current density of 0.2 A g⁻¹ was 91.7 F g⁻¹. The highest energy density and power density were calculated to be 25.8 W h kg $^{-1}$ and 727 W kg $^{-1}$, respectively (Fig. S7 \dagger). The cycle life of the asymmetric supercapacitor was further investigated at 0.5 A g-1 (Fig. 4d). The specific capacitance retained 72.5 after 4500 GCD cycles at current density of 0.5 A g⁻¹, demonstrating the outstanding stability of the device.

In summary, by a simple solvothermal synthesis method, we have successfully synthesized a crystalline triazine containing porous organic polymer POP_{M-TFP} through simple condensation reaction between M and TFP using dimethyl sulfoxide solvent without any catalysts. Further, we have used POP_{M-TFP} as an electrode material for supercapacitor application. The electrochemical measurement of POP_{M-TFP} material showed well electro-chemical performance in supercapacitors. The wonderful electrochemical properties of the POP_{M-TFP} can be ascribed to the synergistic effect of the π -conjugated triazine contained POP_{M-TFP} and nanopores structure. Active sites triazine moieties which put up redox behavior distributed in the pore channels. These findings will meet the requirements like energy storage ability and help more energy efficient electrode materials in the near future.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- 1 B. Dunn, H. Kamath and J.-M. Tarascon, *Science*, 2011, 334, 928.
- 2 X. Liu, N. Wen, X. Wang and Y. Zheng, ACS Sustainable Chem. Eng., 2015, 3, 475.
- 3 M. Beidaghi and Y. Gogotsi, Energy Environ. Sci., 2014, 7, 867.
- 4 A. M. Khattak, H. Yin, Z. A. Ghazi, B. Liang, A. Iqbal, N. A. Khan, Y. Gao, L. Li and Z. Tang, RSC Adv., 2016, 6, 58994
- 5 J. P. Paraknowitsch and A. Thomas, *Energy Environ. Sci.*, 2013, **6**, 2839.

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306.

- 7 Y. Kou, Y. Xu, Z. Guo and D. Jiang, Angew. Chem., Int. Ed., 2011, **50**, 8753.
- 8 C. Wang, Y. Xi, M. Wang, C. Zhang, X. Wang, Q. Yang, W. Li, C. Hu and D. Zhang, *Nano Energy*, 2016, 28, 115.
- 9 C. R. Mulzer, L. Shen, R. P. Bisbey, J. R. McKone, N. Zhang, H. D. Abruna and W. R. Dichtel, *ACS Cent. Sci.*, 2016, 2, 667.
- 10 S.-Y. Ding and W. Wang, Chem. Soc. Rev., 2013, 42, 548.
- 11 K. Sakaushi, G. Nickerl, H. C. Kandpal, L. Cano-Cortes, T. Gemming, J. Eckert, S. Kaskel and J. van den Brink, *J. Phys. Chem. Lett.*, 2013, 4, 2977.
- 12 S. Hug, L. Stegbauer, H. Oh, M. Hirscher and B. V. Lotsch, *Chem. Mater.*, 2015, 27, 8001.
- 13 L. Hao, B. Luo, X. Li, M. Jin, Y. Fang, Z. Tang, Y. Jia, M. Liang, A. Thomas, J. Yang and L. Zhi, *Energy Environ. Sci.*, 2012, 5, 9747.
- 14 T. Lin, I. W. Chen, F. Liu, C. Yang, H. Bi, F. Xu and F. Huang, Science, 2015, 350, 1508.
- 15 L. Xu, Y. Yu, J. Lin, X. Zhou, W. Q. Tian, D. Nieckarz, P. Szabelski and S. Lei, *Nanoscale*, 2016, 8, 8568.
- 16 C. R. DeBlase, K. E. Silberstein, T. Thanh-Tam, H. D. Abruna and W. R. Dichtel, *J. Am. Chem. Soc.*, 2013, 135, 16821.
- 17 C. R. DeBlase, K. Hernandez-Burgos, K. E. Silberstein, G. G. Rodriguez-Calero, R. P. Bisbey, H. D. Abruna and W. R. Dichtel, ACS Nano, 2015, 9, 3178.
- 18 L. Hao, J. Ning, B. Luo, B. Wang, Y. Zhang, Z. Tang, J. Yang, A. Thomas and L. Zhi, J. Am. Chem. Soc., 2015, 137, 219.
- 19 Y. Su, Y. Liu, P. Liu, D. Wu, X. Zhuang, F. Zhang and X. Feng, *Angew. Chem.*, *Int. Ed.*, 2015, **54**, 1812.
- 20 A. M. Khattak, Z. A. Ghazi, B. Liang, N. A. Khan, A. Iqbal, L. Li and Z. Tang, *J. Mater. Chem. A*, 2016, 4, 16312.
- 21 V. Sharma, S. Khilari, D. Pradhan and P. Mohanty, *RSC Adv.*, 2016, **6**, 56421.
- 22 B. C. Patra, S. Khilari, L. Satyanarayana, D. Pradhan and A. Bhaumik, *Chem. Commun.*, 2016, 52, 7592.
- 23 X. Zhan, Z. Chen and Q. Zhang, *J. Mater. Chem. A*, 2017, 5, 14463.
- 24 J. Zhang, Z. Wang, L. Li, J. Zhao, J. Zheng, H. Cui and Z. Zhu, *J. Mater. Chem. A*, 2014, **2**, 8179.

- 25 Y. Li, S. Zheng, X. Liu, P. Li, L. Sun, R. Yang, S. Wang, Z.-S. Wu, X. Bao and W.-Q. Deng, *Angew. Chem.*, *Int. Ed.*, 2018, 57, 7992.
- 26 W. Wang, Y. Yuan, F.-X. Sun and G.-S. Zhu, *Chin. Chem. Lett.*, 2014, 25, 1407.
- 27 B. Yang, J. Bjork, H. Lin, X. Zhang, H. Zhang, Y. Li, J. Fan, Q. Li and L. Chi, J. Am. Chem. Soc., 2015, 137, 4904.
- 28 H. Wei, S. Chai, N. Hu, Z. Yang, L. Wei and L. Wang, *Chem. Commun.*, 2015, **51**, 12178.
- 29 P. Kuhn, M. Antonietti and A. Thomas, *Angew. Chem., Int. Ed.*, 2008, 47, 3450.
- 30 D. D. Medina, J. M. Rotter, Y. Hu, M. Dogru, V. Werner, F. Auras, J. T. Markiewicz, P. Knochel and T. Bein, J. Am. Chem. Soc., 2015, 137, 1016.
- 31 C. S. Diercks and O. M. Yaghi, Science, 2017, 355, 923.
- 32 F. J. Uribe-Romo, J. R. Hunt, H. Furukawa, C. Klock, M. O'Keeffe and O. M. Yaghi, *J. Am. Chem. Soc.*, 2009, **131**, 4570.
- 33 M. G. Schwab, B. Fassbender, H. W. Spiess, A. Thomas, X. Feng and K. Muellen, *J. Am. Chem. Soc.*, 2009, **131**, 7216.
- 34 N. Sahiner, S. Demirci and K. Sel, J. Porous Mater., 2016, 23, 1025.
- 35 X. Huang, Z. Wu, H. Zheng, W. Dong and G. Wang, *Green Chem.*, 2018, **20**, 664.
- 36 Y. Cui, Z. Ding, X. Fu and X. Wang, Angew. Chem., Int. Ed., 2012, 51, 11814.
- 37 B. Jurgens, E. Irran, J. Senker, P. Kroll, H. Muller and W. Schnick, J. Am. Chem. Soc., 2003, 125, 10288.
- 38 S. Chandra, T. Kundu, K. Dey, M. Addicoat, T. Heine and R. Banerjee, *Chem. Mater.*, 2016, **28**, 1489.
- 39 Y. Ma, W. Chen, P. Zhang, F. Teng, J. Zhou, X. Pan and E. Xie, RSC Adv., 2014, 4, 47609.
- 40 J.-W. Jeon, R. Sharma, P. Meduri, B. W. Arey, H. T. Schaef, J. L. Lutkenhaus, J. P. Lemmon, P. K. Thallapally, M. I. Nandasiri, B. P. McGrail and S. K. Nune, ACS Appl. Mater. Interfaces, 2014, 6, 7214.
- 41 W. Lv, M. Guo, M.-H. Liang, F.-M. Jin, L. Cui, L. Zhi and Q.-H. Yang, *J. Mater. Chem.*, 2010, **20**, 6668.
- 42 F. Zhao, Y. Wang, X. Xu, Y. Liu, R. Song, G. Lu and Y. Li, *ACS Appl. Mater. Interfaces*, 2014, **6**, 11007.
- 43 J. Romero, D. Rodriguez-San-Miguel, A. Ribera, R. Mas-Balleste, T. F. Otero, I. Manet, F. Liscio, G. Abellan, F. Zamora and E. Coronado, *J. Mater. Chem. A*, 2017, 5, 4343.