

Cite this: *RSC Adv.*, 2018, 8, 19272

Received 26th March 2018

Accepted 14th May 2018

DOI: 10.1039/c8ra02610f

rsc.li/rsc-advances

Rheological phase reaction synthesis and electrochemical performance of rufigallol anode for lithium ion batteries†

Xiaoyan Han,^{ID}* Guanyu Lin, Qing Zhang and Yingkui Yang^{ID}

1,2,3,5,6,7-hexahydroxy-anthraquinone (rufigallol) and its metal–organic complex (rufigallol-Li/Ni, R-LN) were both synthesized. The electrochemical performance investigation of rufigallol and R-LN as anodes for lithium ion batteries indicates that pure rufigallol delivers high initial capacity but poor cycling stability, by contrast, the R-LN complex exhibits high initial capacity and excellent cycling stability.

Lithium ion batteries (LIBs) have proven to be clean and efficient energy-storage technologies to meet the growing demand for green and sustainable electric power storage. Considering the sustainability, low cost, abundant natural sources, structural design with tolerance for variable functional groups, fast reaction kinetics and high power density, organic electrode materials have been the most competitive alternative to traditional inorganic materials.^{1–6} Quinone organic materials with carbonyl functional group have received great concern for their high theoretical capacities and reaction reversibility. While, the dissolution of small molecule quinones in the organic electrolyte and the resulting capacity attenuation have restricted their application. Therefore, it is urgently demanded to design new organic electrode materials with high energy efficiency and good cycling stability. Among small molecule quinones, 1,2,3,5,6,7-hexahydroxy-anthraquinone (rufigallol) as a critical component has been widely used in dye industry, drug synthesis, and organic materials.^{7–10} Rufigallol was firstly synthesized in 1836.¹¹ Since then, very little efficient method for the preparation of rufigallol has been reported. Recently, Bisoyi and Kumar's research indicates that the rufigallol can be achieved by self-condensation of gallic acid in the presence of sulfuric acid under microwave-assisted, and the yield reached 86%.¹² Also, as far as we know, there is no correlative literature exists about rufigallol being used as electrode materials for lithium ion batteries.

It is well-known that the dissolution of organic molecule in the electrolyte can be effectively suppressed *via* salt formation. Therefore, constructing the metal–organic complex is an efficient way to obtain a stable and flexible framework as well as a better cycling stability.^{13–19} In our previous study,^{20,21} we have

successfully realized the improvement of cycling stability for 3,4,9,10-erythro-tetracarboxylic acid-dianhydride (NTCDA) through the introducing of Li/Ni or Co/Mn to the matrix material. The obtained metal–organic complex, namely, Li/Ni-1,4,5,8-naphthalenetetracarboxylate or Co/Mn-1,4,5,8-naphthalenetetracarboxylate, showed a high specific capacity and a good cycling stability.

Herein, we developed a simple, economical and effective rheological phase method²² to synthesize the rufigallol with high yield and investigated its electrochemical application in lithium ion batteries. In view of the easily connecting of hydroxyl groups in aromatic carbonyl compound for rufigallol with metal ions, in order to modify the cycling stability of pure rufigallol, we synthesized rufigallol-Li/Ni complex (R-LN) through the introducing of lithium and nickel by a hydrothermal method. When used as the LIBs anode, rufigallol shows initial discharge and charge capacities of 977 mA h g⁻¹ and 460 mA h g⁻¹, respectively, which is far higher than that of traditional graphite anode. The R-LN complex delivers initial a charge capacity of 560 mA h g⁻¹ and still remain at about 500 mA h g⁻¹ after 100 cycles, indicating a good electrochemical performance. Our work presents a new quinone-based organic materials with high capacity and competitive cycling stability, which enriches the organic electrode materials.

Experiment section

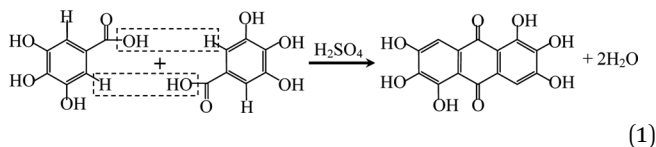
Rufigallol was synthesized from gallic acid and sulfuric acid as precursors by a simple rheological phase method. The gallic acid monohydrate was purchased from commercial sources and dehydrated before using. Proper gallic acid was added slowly in sulfuric acid at 80 °C with stirring to form a rheological mixture. Then the mixture was transferred into a Teflon container sealed in a stainless steel reactor at 120 °C for 5 h. The obtained product was separated simply by enough distilled water with vigorous stirring and subsequently filtered, washed and dried,

Key Laboratory of Catalysis and Materials Science of the State Ethnic Affairs Commission & Ministry of Education, South-Central University for Nationalities, Wuhan 430074, Hubei Province, China. E-mail: xyhan@scuec.edu.cn

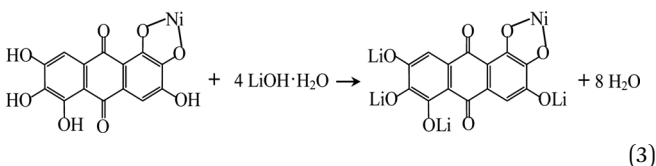
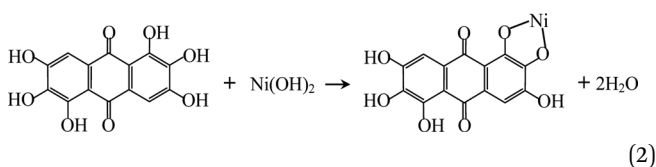
† Electronic supplementary information (ESI) available. See DOI: 10.1039/c8ra02610f



the overall yield was 95%. The reaction process is expressed as follows:



Rufigallol-Li/Ni complex (R-LN) was synthesized by a simple hydrothermal reaction by using rufigallol, Ni(OH)₂ and LiOH·H₂O as precursors. Rufigallol and Ni(OH)₂ with 1 : 1 mol ratio were mixed in ethanol solution with stirring at 70 °C for 3 h. Then, proper LiOH·H₂O (1 : 4 mol ratio) was added, and further stirred for 2 h. After that, the final R-LN product was obtained after dried at 120 °C. The complex reaction process can be expressed as follows:



The chemical structure of the product was characterized by NMR spectroscopy (¹H NMR, Bruker Avance III 400) and Fourier transform infrared spectrometer (FT-IR, Nicolet AVATAR-360). Powder X-ray diffraction (XRD) patterns were obtained by a Bruker D8-advance diffractometer (Germany) with a Ni filter and Cu K α radiation. Thermogravimetry and derivative thermogravimetry (TG/DTG) were conducted to analyze the thermal behavior of the materials in a TG thermal analyzer system (NETZSCH, TG209F3, Germany) with a nitrogen atmosphere of 20.0 ml min⁻¹, at a heating rate of 10 °C min⁻¹, up to the temperature of 800 °C. The samples were packed in an alumina crucible with a mass of about 6.0 mg. The surface morphology was observed by scanning electron microscope (SEM, Hitachi SU8010, Japan).

The battery performance was carried out using a 2016-type coin cells (CR2016), which consisted of a working electrode and a lithium foil as the counter electrode separated by a Celgard-2300 microporous membrane. The working electrode consists of active material, acetylene black and polytetrafluoroethylene (PTFE) binder with the weighting ratio of 60 : 35 : 5. The active materials loading on each electrode sheet is around 3.0 mg cm⁻². A stainless-steel mesh acted as the current collector. 1 M LiPF₆ solution dissolved in a 1 : 1 (v/v) mixture of ethylene carbonate (EC) and dimethylcarbonate (DMC) was used as the electrolyte. The cells were assembled in an argon-filled glove

box. The discharge/charge tests were conducted using a battery test system (Land CT2001A, China) with a scope of 0.01–3.0 V at a constant current density of 100 mA g⁻¹. The electrochemical impedance spectroscopy (EIS) was performed on an electrochemical working station (CHI660E) using model cells. The frequency range applied in this experiment was 0.01–10⁵ Hz and voltage amplitude was 5 mV.

Results and discussion

Fig. 1 shows the ¹H NMR (400 MHz, DMSO-d₆) of the synthesized rufigallol and R-LN. As can be seen from Fig. 1a, the relevant analysis data of the synthesized rufigallol are given below. δ : 7.24 (s, 2H, C-H), 9.92 (s, 2H, 2,6-OH), 10.73 (s, 2H, 3,7-OH), 12.93 (s, 2H, 1,5-OH). The results indicate that the 1,2,3,5,6,7-hexahydroxy-anthraquinone (rufigallol) is successfully synthesized by the rheological phase method. In Fig. 1b, for R-LN, the peaks of hydroxyl hydrogen (1,2,3,5,6,7-OH) sharply decrease or even disappear, and the peaks of aromatic hydrogen at C-2 and C-8 (anthraquinone ring) obviously weaken, which indicate the loss of hydrogen and the introduction of the metal ions.

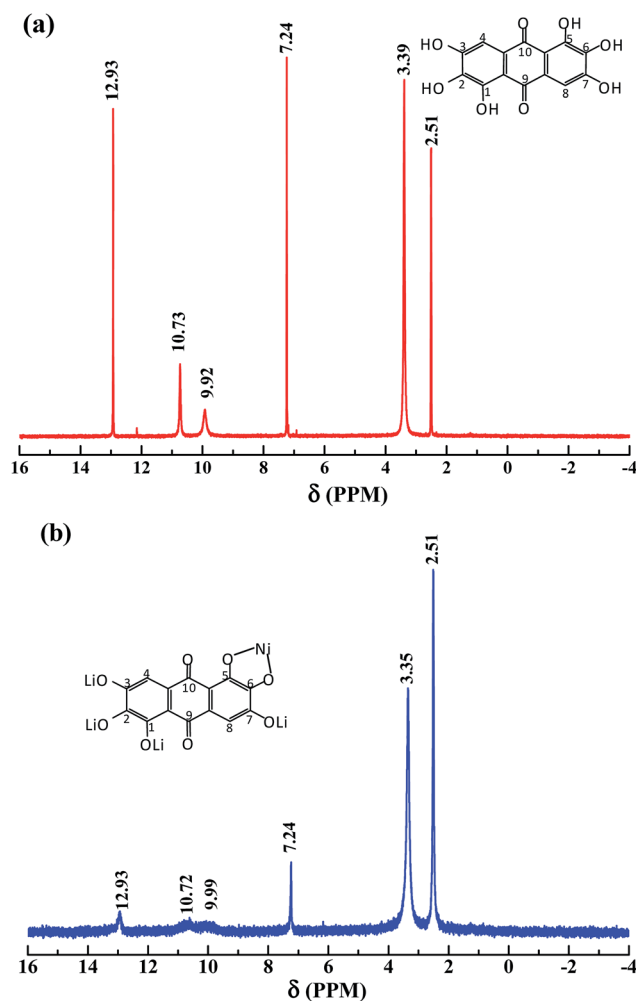


Fig. 1 ¹H NMR for the synthesized rufigallol (a) and R-LN (b).



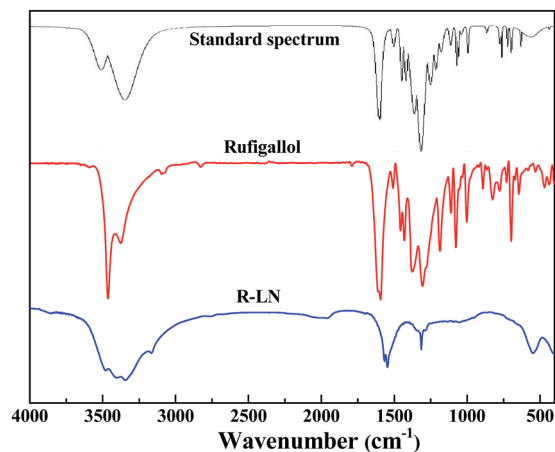


Fig. 2 Infrared spectra of rufigallol and R-LN. The standard IR absorption spectrum of rufigallol is obtained from Wiley Subscription Services, Inc. (US).

Fig. 2 presents the FT-IR spectra of the synthesized rufigallol and R-LN. The synthesized rufigallol can be best characterized by FTIR spectra, which is well consistent with the standard spectrum. Rufigallol shows distinct peaks near 1305 cm^{-1} and 1595 cm^{-1} . The obvious decrease and red shift of these two peaks in R-LN complex is caused by the interaction of C–O with Li or Ni atom, suggesting that the Li or Ni atom exists near the C–O of rufigallol and thus affects the molecular vibration. The O–H stretching and bending modes of R-LN (around 3400 cm^{-1} and 700 cm^{-1}) sharply decrease indicates that the Li or Ni atom substitutes the H in O–H band to form O–Li band (or O–Ni band).

Fig. 3 shows the XRD patterns of the synthesized rufigallol and R-LN. All diffraction peaks of rufigallol are very sharp, which indicates a good crystallinity. The characteristic diffraction peaks are well indexed to the triclinic crystal system and $P\bar{1}$ (no. 2) space group, indicating a pure rufigallol phase of the product. However, R-LN shows poor diffraction peaks which are different from those of rufigallol. No peaks for element Li or Ni can be observed, which indicates that the Li or Ni atom should interconnect with rufigallol molecule in R-LN complex. The XRD results accord well with those from ^1H NMR and FT-IR.

Fig. 4 presents the TG-DTG curves of rufigallol and R-LN. The specific temperature and weight losses are labeled on TG-DTG curves. Generally, the little weight loss below $200\text{ }^\circ\text{C}$ is probably due to volatilization of adsorbed water. The comparison of the TG data for the synthesized rufigallol and R-LN reveals that the materials show different thermal behaviors. As shown in Fig. 4a, the thermal decomposition of rufigallol proceeds in two steps: dehydration and decomposition of the rufigallol organics. The decomposition of rufigallol occurs from $300\text{ }^\circ\text{C}$ to $670\text{ }^\circ\text{C}$ with a weight loss of about 93%, and above $670\text{ }^\circ\text{C}$ the decomposition reaction is complete. R-LN presents three distinct peaks on the DTG curve at $395\text{ }^\circ\text{C}$, $595\text{ }^\circ\text{C}$ and $650\text{ }^\circ\text{C}$, respectively (see Fig. 4b). It can be seen that the thermal decomposition of R-LN proceeds in three main steps. The weight loss occurs from $180\text{ }^\circ\text{C}$ to $410\text{ }^\circ\text{C}$ (31.5%) can be attributed to the decomposition of the organic composition in R-LN. When the temperature increases to $800\text{ }^\circ\text{C}$,

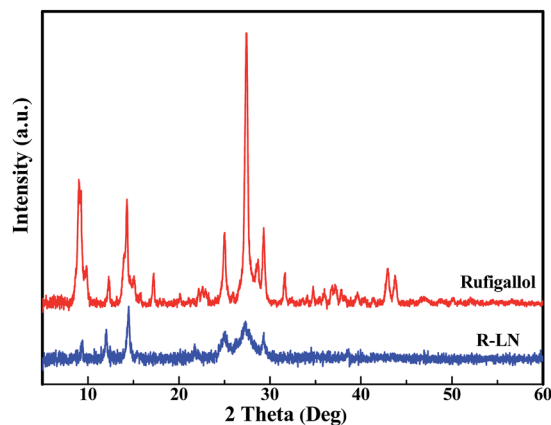


Fig. 3 XRD patterns of rufigallol and R-LN.

the weight loss is about 30.5%, which can be ascribed to the formation of lithium and nickel oxides.

Fig. 5 shows the morphology of rufigallol and R-LN. Typical SEM image shows that the rufigallol is spindle-shaped bulk with a length of $30\text{--}50\text{ }\mu\text{m}$ (Fig. 5a), while its higher magnification SEM image indicates that the spindle-shaped rufigallol is composed of dendritic agglomerates (Fig. 5b). As shown in Fig. 5c and d, the R-LN complex exhibits a porous structure, which consists of dispersed nano-particles with a diameter of about 50 nm . This nano-porous structure material is expected to exhibit good electrochemical behavior for lithium ion batteries.

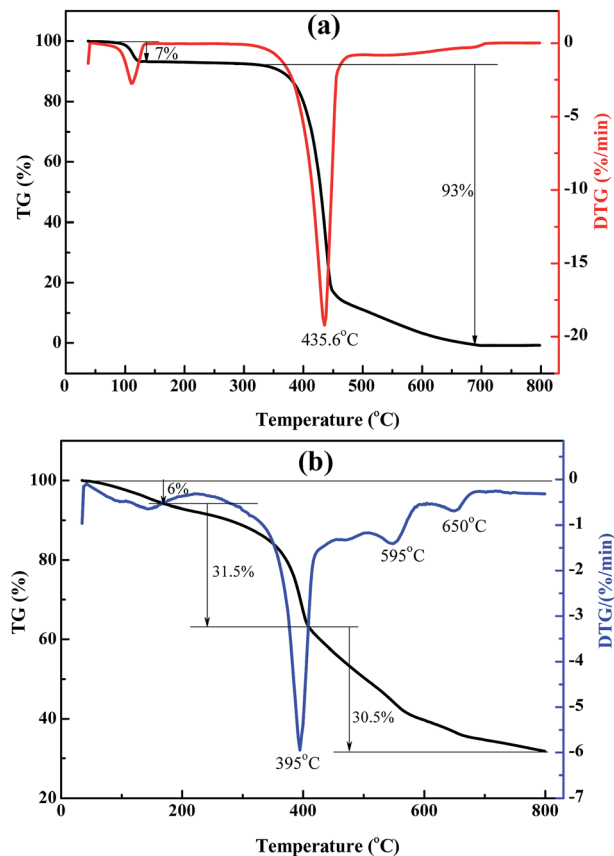


Fig. 4 TG-DTG curves of rufigallol (a) and R-LN (b).



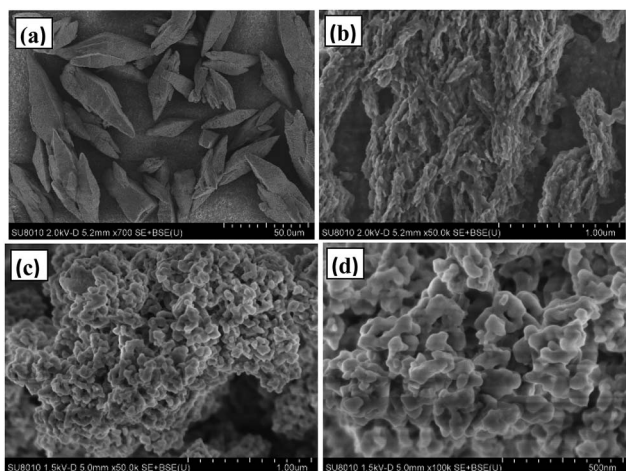


Fig. 5 SEM images of rufigallol (a and b) and R-LN (c and d).

The synthesized rufigallol and R-LN are tested as anode materials for lithium ion batteries by using galvanostatic discharge/charge cycling. Fig. 6a–e shows the electrochemical performance of rufigallol and R-LN electrode at different current densities between 0.01 V and 3.0 V. In Fig. 6a and b, rufigallol and R-LN show similar typical discharge/charge curves, indicating similar electrochemical behaviors. Fig. 6c compares the cycling performance of rufigallol and R-LN at 100 mA g⁻¹. There is a little capacity reduction in the first few cycles of both rufigallol and R-LN, while the discharge and charge capacities increase gradually in the following cycles.

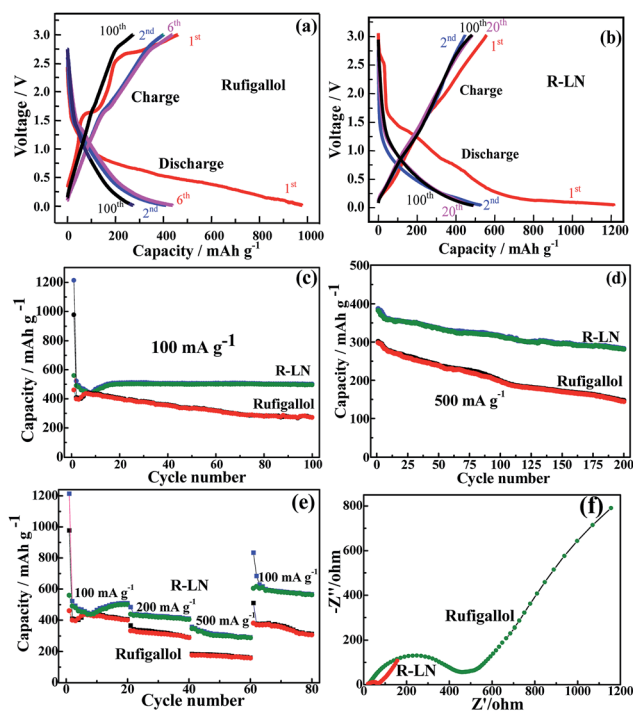


Fig. 6 Electrochemical characterization: (a) and (b) discharge/charge profiles of rufigallol and R-LN at 100 mA g⁻¹; (c) and (d) cycling performance at different current densities; (e) rate capability; (f) EIS of rufigallol and R-LN.

Rufigallol shows initial discharge and charge capacities of 977 mA h g⁻¹ and 460 mA h g⁻¹, respectively. However, the charge capacity rapidly decreases to 270 mA h g⁻¹ in 100 cycles, suggesting a poor cycling stability. In comparison, the R-LN complex shows a better cycling property. It delivers a initial charge capacity of 560 mA h g⁻¹, and still remains about 500 mA h g⁻¹ after 100 cycles, indicating a better cycling stability. Even at a high current density of 500 mA g⁻¹, the R-LN electrode also exhibits high reversible capacity than pure rufigallol, a high charge capacity of 281 mA h g⁻¹ can be still maintained after 200 cycles (Fig. 6d), which should be ascribed to the reduced dissolution of small organic molecular materials and enhanced metal–organic stable structure.

In addition to its improved high capacity, the R-LN complex also exhibits significant improvement in the rate capability. In Fig. 6e, the rate capability of the synthesized rufigallol and R-LN is studied. R-LN achieved the higher capacity under the same current density, showing the better rate performance than pure rufigallol. After 20 cycles, the related charge capacities of R-LN at current densities of 100, 200 and 500 mA g⁻¹ are settled on about 500, 404 and 288 mA h g⁻¹, respectively. A reversible initial charge capacity of 604 mA h g⁻¹ is recovered as the current density returns to 100 mA g⁻¹ and a reversible charge capacity of 563 mA h g⁻¹ can be recovered after 20 additional cycles, demonstrating a good reversibility of R-LN. Nevertheless, with increasing current density to 500 mA g⁻¹, the rufigallol anode deliver poor initial capacity (177 mA h g⁻¹), and after 80 cycles at different current densities, only 66% of the initial capacity is recovered as the current density returns to 100 mA g⁻¹. The good rate performance of R-LN anode is apparently resulted from its greatly reduced electrochemical impedance. As shown in Fig. 6f, the electrochemical impedance of R-LN anode is smaller than that of pure rufigallol anode. The greatly reduced impedance of the R-LN anode is obviously due to its metal–organic stable structure, which can effectively improve the ionic conductivity. Moreover, the R-LN complex with nano-porous structure can provide abundant reaction areas and ionic transport channels. Meanwhile, compared with other metal–organic electrode materials recently reported (Table S1†), the R-LN complex displayed much better cycling stability and rate capability.

Morphologies of the fresh electrode and the retrieved electrode after electrochemical cycles of rufigallol and R-LN were observed through SEM. Fig. S1† shows that the R-LN complex can maintain its nano-porous structure before and after electrochemical cycles (Fig. S1c and d†). From this viewpoint, the stable electrode structural of R-LN should greatly contribute to the excellent cycling performance of the electrode. The fresh rufigallol electrode consists dispersed particles (Fig. S1a†), but these particles aggregate after electrochemical cycles (Fig. S1b†), which may be the main reason for their rapid capacity fading and poor rate performance. To study the structural changes of electrodes, the XRD patterns of the fresh electrode and the retrieved electrode after electrochemical cycles were depicted in Fig. S2.† After the electrochemical cycles, no new diffraction peak appears, and the existing peaks do not change or shift, reflecting the very stable crystallographic



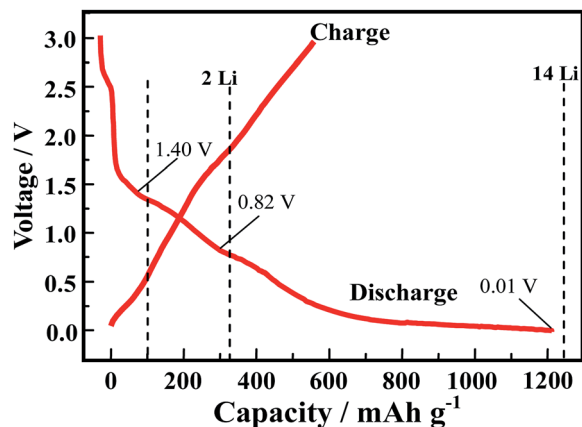
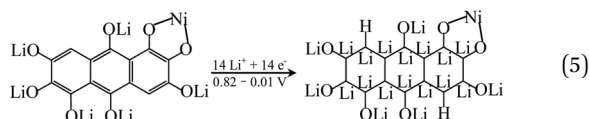
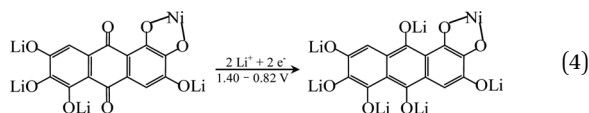


Fig. 7 Initial discharge/charge curves of R-LN at 100 mA g⁻¹.

structure of R-LN during electrochemical processes (Fig. S2a†). However, in Fig. S2b,† for the fresh rufigallol electrode, the distinct peaks at 9.15° and 27.6° disappeared in the retrieved electrode, suggesting the structural change of rufigallol after electrochemical cycles.

In order to explore the lithium storage mechanism, the first discharge/charge curves of R-LN is investigated as shown in Fig. 7. The first discharge curve of R-LN possesses four plateaus. The two small plateaus above 1.4 V lead to the irreversible capacity loss (about 100 mA h g⁻¹), which disappears in the following cycles and result in a capacity reduction. Considering that for one molecule of R-LN, one Li ion insertion contributes a theoretical capacity of 69.7 mA h g⁻¹, and the discharge capacity of R-LN is 1214 mA h g⁻¹, the above data show that each R-LN molecule is capable of accepting 16 Li ions. According to our previous research work,⁶ the actual Li insertion reaction in R-LN includes two major steps: the initial two Li ions inserted at a potential of above 0.82 V can be attributed to a lithium enolization reaction at the two carbonyl oxygens of R-LN, and then the electrochemical lithium addition reaction to the 14 unsaturated carbons occurs at below 0.82 V. The Li insertion reaction of R-LN can be expressed as follows:



Conclusions

In this work, rufigallol has been successfully synthesized by the rheological phase method. Its metal-organic complex, namely, rufigallol-Li/Ni (R-LN), was also synthesized by hydrothermal method. The rufigallol and R-LN were both characterized as

anodes for lithium ion batteries. The electrochemical investigation shows that, as anode material for lithium ion batteries, pure rufigallol shows high initial capacity but poor cycling stability, while the R-LN complex based on rufigallol exhibits high initial capacity and excellent cycling stability. Such optimized electrochemical properties of R-LN complex can be attributed to the metal-organic framework structure, which enhances the structure stability and promotes the cycling stability of small organic molecular materials. The stable framework can slow down the volume collapse and mitigate dissolution in electrolyte. Moreover, the R-LN complex with nano-porous structure, which may be contribute to the insertion/extraction of Li ions. Importantly, the strategy in this work can be extended to fabricate other quinone materials towards high lithium storage performance. Ultimately, the conveniently-synthesized rufigallol with high yield and the observation of improved electrochemical performance by introducing of metal demonstrates that quinone-based materials with stable metal-organic structure is a prospective class of organic electrode materials for lithium ion batteries. Thus, with sufficient effort and improved experimental technique these materials may possess broader practical application prospects than other organic electrode materials.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by the National Natural Science Foundation of China (Grant No. 21503282 and 51673061).

Notes and references

- M. Armand, S. Grugeon, H. Vezin, S. Laruelle, P. Ribière, P. Poizot and J.-M. Tarascon, *Nat. Mater.*, 2009, **8**, 120.
- Y. Liang, Z. Tao and J. Chen, *Adv. Energy Mater.*, 2012, **2**, 742.
- Z. Song and H. Zhou, *Energy Environ. Sci.*, 2013, **6**, 2280.
- J. Wang, X. M. Wang, H. F. Li, X. W. Yang and Y. G. Zhang, *J. Electroanal. Chem.*, 2016, **773**, 22.
- B. Häupler, A. Wild and U. S. Schubert, *Adv. Energy Mater.*, 2015, **5**, 1402034.
- X. Y. Han, G. Y. Qing, J. T. Sun and T. L. Sun, *Angew. Chem., Int. Ed.*, 2012, **51**, 5147.
- E. S. B. Ferriara, A. N. Hulme, H. McNab and A. Quye, *Chem. Soc. Rev.*, 2004, **33**, 329.
- E. A. Hillard, F. C. De Abreu, D. C. M. Ferreira, G. Jaouen, M. O. F. Goulart and C. Amatore, *Chem. Commun.*, 2008, **23**, 2612.
- S. Setia, A. Soni, M. Gupta, S. Sidiq and S. K. Pal, *Liq. Cryst.*, 2013, **40**, 1364.
- J. A. Paquette, R. E. Yardley, J. W. Y. Yu, S. H. Eichhorn and K. Maly, *New J. Chem.*, 2016, **40**, 5985.
- V. Robiquet and U. die Gallussäure, *Liebigs Ann.*, 1836, **19**, 204.
- H. K. Bisoyi and S. Kumar, *Tetrahedron Lett.*, 2007, **48**, 4399.



- 13 J. J. Ma, H. J. Wang, X. R. Liu, L. D. Lu, L. Y. Nie, X. Yang, Y. Q. Chai and R. Yuan, *Chem. Eng. J.*, 2017, **309**, 545.
- 14 J. Xue, C. Fan, Q. J. Deng, M. J. Zhao, L. P. Wang, A. J. Zhou and J. Z. Li, *Electrochim. Acta*, 2016, **219**, 418.
- 15 L. Q. Mu, Y. X. Lu, X. Y. Wu, Y. J. Ding, Y. S. Hu, H. Li, L. Q. Chen and X. J. Huang, *Green Energy & Environment*, 2018, **3**, 63.
- 16 Y. Wang, Q. T. Qu, G. Liu, V. S. Battaglia and H. H. Zheng, *Nano Energy*, 2017, **39**, 200.
- 17 L. S. Shen, H. W. Song and C. X. Wang, *Electrochim. Acta*, 2017, **235**, 595.
- 18 L. P. Wang, J. Zou, S. L. Chen, J. Y. Yang, F. Z. Qing, P. Gao and J. Z. Li, *Electrochim. Acta*, 2017, **235**, 304.
- 19 L. P. Wang, C. X. Mou, B. Wu, J. Xue and J. Z. Li, *Electrochim. Acta*, 2016, **196**, 118.
- 20 X. Y. Han, H. K. Mao and H. W. Liu, *J. Electroanal. Chem.*, 2017, **802**, 89.
- 21 X. Y. Han, F. Yi, T. L. Sun and J. T. Sun, *Electrochem. Commun.*, 2012, **25**, 136.
- 22 J. T. Sun, W. Xie, L. J. Yuan, K. L. Zhang and Q. Y. Wang, *J. Mater. Sci. Eng. B*, 1999, **64**, 157.

