LiFePO4-Fe2P-C composite cathode: An environmentally friendly promising electrode material for lithium-ion battery

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Abstract
In this investigation, the synthesis strategy is involved the creation of LiFePO4-Fe2P-C composites with a porous conductive architecture, which includes distinct regions or clusters containing antiferromagnetic LiFePO4 in close proximity to ferromagnetic Fe2P. The microstructure is achieved by using a simple ultra-fast solvent assisted manual grinding method, combined with solid state reaction, which can replace the time-consuming high energy ball milling method. The crystalline structure, morphology, and electrochemical characterization of the synthesized product are investigated systematically. The electrochemical performance is outstanding, especially the high C rate. The composite cathode is found to display specific capacity of 167 mAh g\(^{-1}\) at 0.2 c and 146 mAh g\(^{-1}\) at 5 c after 100 cycles, respectively. At the high current density of 1700 mA g\(^{-1}\), it exhibits long-term cycling stability, retaining around 96 per cent of its original discharge capacity beyond 1000 cycles, which can meet the requirements of a lithium-ion battery for large-scale power applications. The obtained results have demonstrated that the fabrication of samples with strong and extensive antiferromagnetic and ferromagnetic interface coupling of LiFePO4/Fe2P provides a versatile strategy toward improving the electrochemical properties of LiFePO4 materials and also opens up a new window for material scientists to further study the new exchange bias phenomenon and its ability to enhance the electrochemical performance of lithium-ion battery electrode.

Keywords
material, lithium, ion, battery, c, composite, cathode, environmentally, friendly, promising, electrode, lifepo4, fe2p

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In the new century, clean and renewable energy storage devices have become the focus of both the building industry and research development. Lithium ion batteries, as one of the most promising battery technologies, have attracted much attention due to their fast boom of market share with the most advanced electrochemical energy storage and conversion system for a wide range of green applications, including hybrid electric vehicles (HEVs), plug-in hybrid electric vehicles (PHEVs), and stationary energy storage for solar and wind electricity generation as well as smart grids. However, for EVs/HEVs applications, commercialized lithium-ion batteries do not yet meet the required combinations of high energy density, high power, and high rate capability. Apart from the search for new or improved electrode materials with higher energy densities, the enhancement of electrode capacity retention at high charge/discharge rates is one main challenges of lithium-ion nanocomposite (X mAh g⁻¹ at 10C after 1000 cycles) and by electronic conductivity and low ionic diffusion coefficient.[8,9] Applications, is its poor high-rate performance, owing to its low material, which prevents it from being used in large-scale high rate have been reported for the LiFePO₄ material through application. Recently, ultrafast charging and discharging at very high current rate have been reported for the LiFePO₄ material, in particular for high power application, because of its high theoretical capacity (170 mAh g⁻¹), acceptable operating voltage (3.4 V vs. Li⁺/Li⁻), low cost, environmental friendliness, long cycle life, cell safety, and high thermal stability.[10] Nevertheless, a major limitation of this material, which prevents it from being used in large-scale applications, is its low high-rate performance, owing to its low electronic conductivity and low ionic diffusion coefficient.[9,10] However, the long term cycling stability at high current rate is still a great challenge for this material as it is a compulsory requirement for lithium ion batteries to have long cycle life for EVs/HEVs application. Recently, ultrafast charging and discharging at very high rate have been reported for the LiFePO₄ material through creating an ion conducting lithium phosphate coating on the surface of LiFePO₄ nanoparticles but their reported cycle number is not good enough.[10] A satisfactory long term cycling stability has been achieved through the formation of mesoporous LiFePO₄/C nanocomposite (118 mAh g⁻¹ at 10C after 1000 cycles) and by creating LiFePO₄/carbon composite (~85 mAh g⁻¹ at 10C after 2400 cycles) via high-energy ball milling combined with spray-drying method.[11] Both of these reported results satisfy the long term cycling but their specific discharge capacity is not as high as we expect and thus, there is much more room for further improvement. In this investigation, our synthesis strategy has been involved to create porous conductive architecture of LiFePO₄-C composite with an antiferromagnetic (AFM) and ferromagnetic (FM) cluster interface coupling of LiFePO₄/Fe₃P through a solvent assisted modified solid state reaction method, which is quite different from the others reported in the literature. Herein, we report on “exchange bias (EB)” effect due to LiFePO₄/Fe₃P interface coupling and their enhancement in electrochemical performance of LiFePO₄/Fe₃P-C composite cathode. The electrochemical measurements demonstrated that the synthesised LiFePO₄/Fe₃P-C composite delivered a high capacity of 167 mAh g⁻¹ at 0.2C at 100th cycle and displayed long term cycling stability with a capacity retention of around 96% (131 mAh g⁻¹) even after 1000 cycles at 10C. To the best of our knowledge, this is the best high rate long-term cycling performance for LiFePO₄/C composite cathode material reported so far. The XRD patterns of the samples are shown in Figure S1 (Supporting Information). The profiles of the diffraction peaks could be indexed according to the olivine LiFePO₄ phase (JCPDS Card Number 40-1499). The illegible Fe₃P peak (2θ = 40.28°) in XRD patterns for the carbon coated samples indicate that it begins to form the iron phosphide phase (barringerite Fe₇P₃) during the annealed process and it usually exist in the form of nanosized clusters.[12] We also have collected the XRD pattern of bare-LiFePO₄ and have not found the illegible Fe₃P peaks, it means that the residue coating carbon originated from the citrate framework has acted as the reductive under Ar atmosphere during the annealed process in our synthesis system. To estimate the amount of amorphous carbon in the LiFePO₄/Fe₃P-C composites, TGA was carried out in air (Figure S2 in the Supporting Information), it was estimated that the amount of amorphous carbon in the composites were approximately 5.8 wt.% C [LiFePO₄/Fe₃P-C (1)], 10.4 wt.% C [LiFePO₄/Fe₃P-C (2)], and 19.9 wt.% C [LiFePO₄/Fe₃P-C (3)] obtained from different amount of citric acid. The specific surface areas of the synthesised products were also measured by the BET N₂ adsorption method. LiFePO₄/Fe₃P-C (1) composite shows the highest specific surface area (33.14 m² g⁻¹), while bare-LiFePO₄, LiFePO₄/Fe₃P-C (2), and LiFePO₄/Fe₃P-C (3) have specific surface areas of 1.17 m² g⁻¹, 16.74 m² g⁻¹, and 14.25 m² g⁻¹, respectively. Field emission scanning electron microscopy (FESEM) images of the bare-LiFePO₄ and LiFePO₄/Fe₃P-C composites with different carbon content are shown in Figure S1. It is obviously observed that the growth of the LiFePO₄ grains is inhibited by the formed carbon and Fe₃P during the heat treatment process. According to previous researches, the particle size and electrochemical polarization can be reduced effectively only when LiFePO₄ particle surface is coated by conductive carbon.[8,13] In Figure 1 (b-d), the FESEM observation reveals an abrupt particle growth with increasing carbon content in the sample, which maybe caused by the agglomeration of the excessive carbon in the sample where Fe₃P nanoclusters are being trapped. The porous network structure along with small particles and rough surfaces can be clearly observed in Figure 1 (b). As shown in Figure 1 (c, d), it is obvious that the porous network structure with rough surfaces disappear and agglomerated larger particles with smooth surfaces appear. FESEM high-contrast backscattered imaging (Figure 2) of sectioned powders was performed with quantitative calibration of the three most distinct phases (Fe₃P, LiFePO₄, and C) present in local regions of constant grey level (Figure S3 in the Supporting Information).
Examination of Figure 2(b-d) reveals the presence of inhomogeneous distributions of nano Fe₂P particles (white), in a highly porous architecture of LiFePO₄ (light grey) and carbon (dark grey).

Despite the inhomogeneous nature of microstructures, it was observed that the LiFePO₄-Fe₂P-C composite containing 5.8 wt.% C (Figure 2(b)) exhibited the largest fraction of local areas comprising fine distribution of Fe₂P particles in close contact with LiFePO₄ and carbon, (compare Figure 2(b) with Figure 2(c) and (d)). It was also observed that this sample had a particularly porous and highly porous conductive architecture (Figure S4 in the Supporting Information). These observations are consistent with the formation of more and strong interface coupling of LiFePO₄ (AFM) /Fe₂P(FM) in LiFePO₄-Fe₂P-C (1) sample compared with other samples. The results are also well supported by the BET surface areas were found of 33.14, 16.74, 14.25, and 1.17 m²g⁻¹ for the LiFePO₄-Fe₂P-C (1), LiFePO₄-Fe₂P-C (2), LiFePO₄-Fe₂P-C (3), and bare-LiFePO₄, respectively, and by electrochemical impedance spectroscopy (EIS) analysis (described later).

Preliminary magnetic measurement investigations (Figure 3) revealed additional information concerning structural evolution of the samples. However, so far there has been no experimental determination of exchange bias (EB) effect of LiFePO₄/Fe₂P interface coupling in LiFePO₄ materials, even though the magnetic structure and properties of LiFePO₄ have been re-examined theoretically and experimentally. The exchange interaction at the interface between a ferromagnetic (FM) and antiferromagnetic (AFM) component often results in an interesting phenomenon called “exchange bias” (EB), which is manifested by a shift in the hysteresis loop along the field axis when the system is cooled down in an external magnetic field.[16,17] Comparing to other samples, the huge shift in the magnetic hysteresis loop is observed for the LiFePO₄-Fe₂P-C (1) sample, in which the magnitude of the shift in the field axis is defined as the EB (exchange bias) field, \( -H_E = H_1 + H_2 / 2 \), where \( H_1 \) and \( H_2 \) are the left and right coercive fields, respectively.[18] is shown in Figure 3 (a-d). The maximum value of \( H_E \) is 634 Oe at 500 Oe cooling field for the LiFePO₄-Fe₂P-C (1) sample, which is larger than the value for other samples at 5K. The comparison of the EB effect among the samples indicates that the EB effect is stronger for the LiFePO₄-Fe₂P-C (1) sample and is in ascending order of LiFePO₄-Fe₂P-C (1)>LiFePO₄-Fe₂P-C (2)> LiFePO₄-Fe₂P-C (3)> Bare-LiFePO₄. The reason could be attributed to the largest fraction of local areas comprising fine distribution of Fe₂P particles in close contact with LiFePO₄ which leads to the formation of more and strong interface coupling of LiFePO₄ (AFM) /Fe₂P(FM) in LiFePO₄-Fe₂P-C (1) sample compared with other samples. The electrochemical performances of the prepared samples were evaluated systematically using CR 2032 coin cells. The short term cycle life performances for the bare-LiFePO₄ and LiFePO₄-Fe₂P-C composite electrodes at 10 C charge/discharge rates are shown in Figure 4 (a). The initial discharge capacities were measured to be 43, 59, 89, and 137 mAh g⁻¹ with a capacity retention of 40, 56, 84, and 136 mAh g⁻¹ at the 120⁰ cycle at 10 C rate for the bare-LiFePO₄, LiFePO₄-Fe₂P-C (3), LiFePO₄-Fe₂P-C (2), and LiFePO₄-Fe₂P-C (1) electrodes, respectively. The electrochemical performance among the carbon coated samples is in ascending order of LiFePO₄-Fe₂P-C (1) > LiFePO₄-Fe₂P-C (2) > LiFePO₄-Fe₂P-C (3). Electrode, composed of LiFePO₄-Fe₂P-C (1) (5.8 wt.%C) shows the best electrochemical performances even at high current density.
of 10 C. In order to fully estimate the electrochemical performance of the LiFePO$_4$-Fe$_2$P-C (1) (5.8 wt.%C) composite electrode, the cycling behaviours at different current densities of 0.2, 2, 5 and 10 C were measured at the 100$^{th}$ cycle and their corresponding charge-discharge voltage profile is shown in Figure 4 (c). The LiFePO$_4$-Fe$_2$P-C (1) (5.8 wt.%C) composite electrode shows long and flat voltage plateaus in the 3.4-3.5 V range and the small voltage difference between the charge-discharge plateaus is representing of its good kinetics. This observation is also supported by the cyclic voltammogram (CV curve) shown in Figure 4 (d). The well defined sharp redox peaks in the range of 3.26-3.70 V should be attributed to the Fe$^{3+}$/Fe$^{2+}$ redox couple reaction, corresponding to lithium extraction and insertion in LiFePO$_4$ crystal structure.$^{[2]}$ The 100$^{th}$ cycle discharge capacities were measured to be 167 mAh g$^{-1}$ at 0.2 C, 159 mAh g$^{-1}$ at 2 C, 146 mAh g$^{-1}$ at 5 C, and 136 mAh g$^{-1}$ at 10 C for the LiFePO$_4$-Fe$_2$P-C (1) (5.8 wt.%C) electrode, respectively. At the low current density of 0.2 C (5 hours charge and 5 hours discharge), the obtained discharge capacity (167 mAh g$^{-1}$) is very close to the theoretical capacity of LiFePO$_4$ (170 mAh g$^{-1}$). Even at the high current rate of 10 C (6 minutes for charging and 6 minutes for discharging), a capacity of 136 mAh g$^{-1}$ is still obtained, demonstrating that the LiFePO$_4$-Fe$_2$P-C (1) (5.8 wt.%C) composite can tolerate high rate charge and discharge. The capacity fading is observed only ~18 % with the increasing of charge-discharge rate from 0.2 to 10 C.

![Figure 4](image.png)

**Figure 4.** Short term cycle life performance (a), long term cycle life performance beyond 1000 cycles at 10 C for the LiFePO$_4$-Fe$_2$P-C (1) electrode (b), the 100$^{th}$ cycle galvanostatic charge-discharge profiles at the different current densities from 0.2 to 10 C between 4.3 and 2.5 V for LiFePO$_4$-Fe$_2$P-C (1) electrode (c), cyclic voltammogram of LiFePO$_4$-Fe$_2$P-C (1) electrode at a scan rate of 0.1 mVs$^{-1}$ (d), EIS spectra of the bare-LiFePO$_4$ and LiFePO$_4$-Fe$_2$P-C electrodes and the equivalent circuit (inset) used to fit the impedance data (e).

Furthermore, our composite electrode was life tested at a high current density of 1700 mA g$^{-1}$ (10 C rate) for a long term cycling as batteries are required to operate at high current density and to have a cycle life of more than 2000 cycles for EVs/HEVs applications.$^{[14]}$ Therefore, we cycled LiFePO$_4$-Fe$_2$P-C (1) (5.8 wt.%C) electrode at 10 C rate (6 minutes for charging and 6 minutes for discharging) for 1000 cycles (Figure 4 (b)). Surprisingly, the LiFePO$_4$-Fe$_2$P-C (1) (5.8 wt.%C) electrode exhibited superior electrochemical performance with a capacity retention of around 96% (131 mAh g$^{-1}$) of its original discharge capacity after 1000 cycles at high current rate of 10 C. Such outstanding electrochemical performance certainly can meet the demands of many high power applications. However, to understand the effect of LiFePO$_4$/Fe$_2$P interface coupling along with carbon coating on the charge transfer resistance of electrodes, ac impedance measurements were carried out at room temperature (Figure 4 (e)). The impedance curves show one compressed semicircle in the medium-frequency region, which could be assigned to the charge-transfer resistance ($R_t$). The spike at the low frequency end indicates the Warburg impedance ($W$) of long-range lithium-ion diffusion.$^{[19-21]}$ The charge transfer resistance ($R_t$) was calculated to be 148 $\Omega$ cm$^{-2}$ for the bare-LiFePO$_4$, 28 $\Omega$ cm$^{-2}$ for the LiFePO$_4$-Fe$_2$P-C (1), 37 $\Omega$ cm$^{-2}$ for the LiFePO$_4$-Fe$_2$P-C (2), and 60 $\Omega$ cm$^{-2}$ for the LiFePO$_4$-Fe$_2$P-C (3) electrodes, respectively. According to the previous research,$^{[22]}$ higher carbon content sample shows lower charge transfer resistance ($R_t$) and generally this trend is also logical. Under this consideration, the $R_t$ should be in order of LiFePO$_4$-Fe$_2$P-C (3) (19.9 wt.% C) < LiFePO$_4$-Fe$_2$P-C (2) (10.4 wt.% C) < LiFePO$_4$-Fe$_2$P-C (1) (5.8 wt.% C), but the reality is inverse LiFePO$_4$-Fe$_2$P-C (1) < LiFePO$_4$-Fe$_2$P-C (2) < LiFePO$_4$-Fe$_2$P-C (3). At this point, we strongly believe that this $R_t$ is not only influenced by the carbon content but also strongly influenced by the interface coupling of LiFePO$_4$/Fe$_2$P clusters. Obviously, LiFePO$_4$-Fe$_2$P-C (1) sample exhibits more and strong interface coupling of antiferromagnetic (AFM) and ferromagnetic (FM) clusters than that of other samples which increase the effective interface areas, facilitate charge transfer more faster, and reduce the charge transfer resistance, leading to the huge shift in the magnetic hysteresis loop.$^{[23,24]}$ So, the excellent electrochemical performances of the LiFePO$_4$-Fe$_2$P-C (1) (5.8 wt.%C) composite could be attributed to the porous conductive architecture with huge and strong interface coupling of LiFePO$_4$/Fe$_2$P that increase the contact area among the carbon, Fe$_2$P clusters, and LiFePO$_4$ particles, providing multi-dimension channels for charge transfer and reduce the resistances for lithium ion migration. Moreover, the composite with porous architecture can suck up electrolyte to shorten enormously the diffusive distance of lithium ion.

In conclusion, our results demonstrated that the fabrication of composite microstructure provides versatile strategies toward improving the electrochemical properties of LiFePO$_4$ materials at higher C rate which can meet the requirements of a lithium-ion battery for large-scale power applications.

**Experimental Section**

A simple ultrafast solvent assisted manual grinding method, combined with solid state reaction has been developed to synthesize LiFePO$_4$-Fe$_2$P-C composite with a porous conductive architecture. The grinding method involves here replaces time-consuming high energy ball milling method. Li$_2$CO$_3$, FeCO$_3$:2H$_2$O, and NH$_4$PO$_4$ in a stoichiometric molar ratio of 1:1:3 were used as starting materials and citric acid (C$_6$H$_8$O$_7$) used as a reducing agent and carbon source as well. The reactants for preparing the precursor were ground thoroughly, and a slurry was made by mortar and pestle in acetone solvent to ensure intimate and homogeneous mixing at the atomic level. The slurry was then dried in oven at 60$^\circ$C to remove acetone from the slurry. To decompose the carbonate, oxalate, and phosphate, the dried mixture was placed in a tube furnace and heat-treated at 350$^\circ$C for 10 h in argon flow. The resultant powders were cooled to room temperature and thoroughly regrind. The powders were again calcined at 600$^\circ$C for 10 h under argon flow. Thus the bare-LiFePO$_4$ and LiFePO$_4$-Fe$_2$P-C composite containing 5.8 wt.% C [LiFePO$_4$-Fe$_2$P-C (1)], 10.4 wt.% C [LiFePO$_4$-Fe$_2$P-C (2)], and 19.9 wt.% C [LiFePO$_4$-Fe$_2$P-C (3)] were obtained using different amount of citric acid.

X-ray diffraction (XRD) data were collected on a GBC MMA generator. BET surface area of the synthesized materials was measured by a NOVA 1000 high speed gas sorption analyzer (Quantachrome Corporation, USA). Magnetic measurements were performed using a physical properties measurement system (PPMS) 14T magnetometer. The amounts of amorphous carbon in the composite samples were estimated using a
onto aluminium foil substrates with an area of 1 cm$^2$. The coated electrodes were uniformly spread with N,N-dimethylformamide (NMP, Sigma-Aldrich, anhydrous, 99.5 %). The slurry was dried in a vacuum oven at 100 °C for 24 h and then pressed under a pressure of 8 MPa. CR 2032 coin-type cells were assembled in an Ar-filled glove box (Mbraun, Unilab, Germany). The electrochemical coin cells contained a working electrode, lithium foil as the counter electrode and reference electrode, porous polypropylene as the separator, and 1 M LiPF$_6$ in a 50:50 (v/v) mixture of ethylene carbonate and dimethyl carbonate (MERCK KgaA, Germany) as the electrolyte. The cells were galvanostatically charged and discharged in the range of 4.3-2.5 V at different rates of 0.2 -10C using a computer-controlled charger system manufactured by Neware Battery Testers. Cyclic voltammetry (with a scan rate of 0.1 mV s$^{-1}$ between 4.3 and 2.5 V (versus Li/Li$^+$)) and electrochemical impedance spectroscopy (EIS) were performed on the electrodes using a CHI 660C electrochemistry workstation. The AC amplitude was 5 mV, and the frequency range applied was 100 kHz - 0.01 Hz.

Keywords: manual grinding · LiFePO$_4$-Fe$_2$P-C composite · Fe$_2$P phase · LiFePO$_4$/Fe$_2$P interface coupling · Li-ion batteries

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