



Synergistic effect on electrochemical performance of LiFePO₄ cathodes via carbon coating and Ni²⁺ doping: a combined experimental and theoretical approach

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ABSTRACT

The development of high-power cathode materials is critical for meeting the growing demand for high-rate performance in lithium-ion batteries (LIBs) used in portable electronics, electric vehicles, and grid storage. In this work, an ultra high-rate-performance LiFePO₄ (LFP) cathode material is synthesized through a synergistic strategy combining Ni²⁺-ion doping, uniform carbon coating, and particle size reduction. Structural analysis by XRD confirms the orthorhombic phase of LFP, while FE-SEM images reveal pulverized particles in the 1–10 μm, and HR-TEM confirms homogeneous carbon coating. Rietveld refinement shows that Ni²⁺ doping reduces cell volume and lattice parameters, altering the Li-ion local environment, as further supported by density functional theory calculations. XPS analysis confirms the presence of Fe exclusively in the Fe²⁺ state. Electrochemical performance demonstrated outstanding performance with a high discharge capacity of 171 mAh g⁻¹ at C/5 rate, remarkable rate capability of 136 mAh g⁻¹ at 10C with a flat 3.3 V plateau, and over 80 % capacity retention after 200 cycles at 5C. The exceptional performance is attributed to reduced particle size, improved electrode kinetics from Ni²⁺ doping, and an enhanced Li-ion transport. These findings position Ni-LFP/C as a promising high-power cathode material for next generation LIB applications for high-power LIB applications.

1. Introduction

The olivine-type lithium iron phosphate LiFePO₄ (LFP) is one of the cathode materials being employed for Li-ion batteries (LIB) due to several advantages such as low raw material cost, environmental benignity, high structural and thermal stability and long cycle life. In addition to this, LFP displays a stable constant voltage curve at 3.45V (vs. Li/Li⁺) along with appreciable specific capacity of 170 mAh g⁻¹ [1–3]. The LFP cathode material adopts an olivine structure adopting *Pnma* space group. Each unit cell consists of one lithium (Li), iron (Fe), and phosphorus (P) atom, along with four oxygen (O) atoms. In typical LiFePO₄ structure, each iron (Fe) atom is surrounded by an oxygen octahedron. This means that there are six oxygen atoms forming an octahedral shape around each iron atom (FeO₆). The phosphate (PO₄) groups are situated between the iron-oxygen octahedra, and the phosphorus

atom (P) residing at the center bonded with four oxygen atoms. Lithium atoms are located within channels formed by the phosphate groups. These channels run along one direction [010] within the crystal lattice [4,5]. However, LFP exhibits low electronic conductivity because the structure lacks a continuous network of FeO₆ octahedra since it is blocked by the PO₄ tetrahedron [6,7]. In addition, the poor Li⁺ diffusivity arises from the fact that LFP accommodates only one-dimensional Li⁺ diffusion pathway i.e., Li⁺ moves only in the channels parallel to [010] [8,9].

To address these challenges, several key approaches are identified and put forward: one, surface coating of LFP particles with conductive carbon can improve the material electrical conductivity [10,11]; two, nano-structuring, which reduces the Li⁺ diffusion pathway thereby improving the ion diffusion [12–14] and three, substitution or doping of certain metal ions into the LFP crystal structure [15–17]. Metal ion

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insertion, either into the Li or Fe-site can enhance the diffusivity of Li^+ as well as increase the intrinsic electronic conductivity. It is believed that multi-ion doping enhances the Li^+ diffusion kinetics during charging and discharging, and also widens the Li^+ diffusion channels in the crystal structure [18,19]. Carbon-coating, Fe-site substitution, and nano-structuring have proven to be highly effective methods in addressing the limitations of LiFePO_4 cathode material. The incorporation of certain metal-cations through doping is widely accepted and often adopted for enhancing the electrochemical performance of LiFePO_4 [20,21]. Notable improvements in the electrochemical performance of doped LiFePO_4 , particularly at very low concentrations of the dopant have been confirmed in previous studies. For instance, Liu et al. [22] demonstrated that LiFePO_4/C showed an enhancement in electron conductivity by 275 times after Mg^{2+} doping. The doped material exhibited a higher discharge capacity (155 mAh g^{-1} vs. 144.6 mAh g^{-1} for pristine LFP at 0.5C) and an approximately 3.6-fold increase in the Li^+ diffusion coefficient compared to pure LFP. The authors explained this by citing shortened Fe—O and P—O bonds and lengthened Li—O bonds, which widened the pathways for Li^+ diffusion channels. Teng et al. [23] reported that Nb^{5+} doped LiFePO_4 (LiFNbPO-0.75) showed discharge capacity of $169.87 \text{ mAh g}^{-1}$ for the initial cycle at C/10, and it was able to retain its capacity for up to 99 % after 100 cycles. It was argued that doping with Nb^{5+} resulted in the widening of the crystal plane space and also led to low charge transfer impedance. Jian and Wang [24] showed that nanosized vanadium-doped LiFePO_4 exhibited a specific capacity of 142.7 mAh g^{-1} after 200 cycles, measured at 10C current rate. They observed a capacity of 162.9 mAh g^{-1} demonstrated at C/10, although the cost of dopant seems to be expensive.

Despite notable advancements, significant challenges remain in developing LiFePO_4 based cathode materials that simultaneously deliver high power, high specific capacity, and long-term cyclic stability. Key obstacles include achieving uniform and optimally thin carbon coatings, minimizing defects and impurities introduced by metal-ion doping, simplifying often complex multi-step synthesis processes, and addressing the high cost of certain dopant precursors such as V, Ti, Zr, and Nb. Additionally, developing facile, scalable, and cost-effective methods for producing nanosized LiFePO_4/C particles remains critical. Recognizing that nickel (Ni^{2+}) precursors are relatively inexpensive compared to other commonly studied dopants and that Ni^{2+} -doped LiFePO_4/C remains underexplored, we adopted a synergistic approach to synthesize a high capacity cathode material via: (i) doping a small concentration of Ni^{2+} ions at the Fe^{2+} sites using a solid-state milling approach, (ii) achieving particle size reduction, and (iii) generating a homogenous, thin carbon coating on the LiFePO_4 surface. The resulting Ni-doped LiFePO_4/C (Ni-LFP/C) material was extensively characterized by adopting both experimental and computational methods to understand the effects of Ni doping. Our analysis shows that the particle shape and particle size of Ni-LFP/C are retained even after doping. This integrated approach resulted in a high-performance electrode, delivering an appreciable capacity of 171 mAh g^{-1} at C/5, superior rate performance (136 mAh g^{-1} @10C), in addition to outstanding cyclic stability with retention of 80 %, noted at 5C after 200 cycles. These exceptional electrochemical properties underscore the potential of Ni-doped LiFePO_4/C as a superior cathode electrode material for the fabrication of high-power Li-ion battery. Moreover, the synergistic strategy demonstrated here could be extended to other cathode and anode systems, contributing to the development of next-generation high-rate energy storage technologies.

2. Experimental section

A simple ball milling route has been employed to synthesize LiFePO_4/C . It was prepared by milling Li_2CO_3 (99 %, SRL), $\text{FeC}_2\text{O}_4 \cdot 2\text{H}_2\text{O}$ (99 %, Aldrich) and $\text{NH}_4\text{H}_2\text{PO}_4$ (98–101 %, Fisher Scientific) in ethanol, using a planetary mill (pulverisette 5, FRITSCH). In addition, citric acid (99 %, SRL) was used as the precursor for carbon

coating and as a complexing agent to avoid Fe oxidation from +2 to +3 while calcining the milled material. For the doped sample Ni^{2+} -doped LiFePO_4 , $(\text{CH}_3\text{COO})_2\text{Ni} \cdot 4\text{H}_2\text{O}$ (98 %, LOBA Chemie) was used as the precursor for Ni doping. Ni-doped LFP has been synthesized by taking Li, Fe, Ni, and P in the stoichiometric molar ratio of 1:0.99:0.01:1 respectively ($\text{LiFe}_{0.99}\text{Ni}_{0.01}\text{PO}_4/\text{C}$). For planetary ball milling, stainless steel (SS) vials were used with SS balls having diameter 3 mm as grinding media with the powder to ball ratio maintained at 1:10. For the synthesis of Ni doped LFP, all the precursors including Ni dopant were dispersed into ethanol and milled at 250 rpm with the milling time and rest time of 10 and 5 min, respectively for 21 repetitions. Similarly, LiFePO_4/C has been milled without addition of Ni source. Further, the milled powders were then dried overnight at 80°C to evaporate the solvent and finally pelletized and heated to 750°C for 12 h in a tube furnace. The heating rate was maintained at 5°C min^{-1} and Ar gas was continuously purged during heating and natural cooling. As synthesized samples, LiFePO_4/C and Ni^{2+} -doped LiFePO_4 were designated as LFP/C and Ni-LFP/C, respectively.

2.1. Materials characterization

Morphological features of the developed materials were checked through Gemini 500, Carl Zeiss field emission scanning electron microscope (FE-SEM) operated at a voltage of 5 kV. For analyzing bulk chemical composition, energy dispersive X-ray spectroscopy (EDS) studies were carried out using an attached EDS unit (AMETEK, octane) to the FESEM. For TEM imaging, ethanol-dispersed samples were drop-casted onto a lacey carbon-coated Cu grid and left for solvent evaporation. The TEM images, scanning TEM (STEM), and EDS spectra and maps were obtained by employing a Jeol JEM F200 Multipurpose electron microscopy operating at 200 kV. Gatan One view camera was used to acquire TEM images. Powder X-ray diffraction experiment (XRPD) was performed using Rigaku SmartLab high-flux XRD with 45 kV working voltage and filament current of 200 mA. XRPD patterns were measured in the 2θ range of $10\text{--}100^\circ$ at room temperature. The automated multipurpose XRPD was equipped with a 9-kW rotating anode ($\text{Cu } K_\alpha$; $\lambda=1.54 \text{ \AA}$) and D/tex Ultra 250 detector having a parabolic multilayer X-ray mirror with cross beam optics. The patterns were matched using MATCH Software which employed the ICDD PDF-4+ 2022 database. For pattern refinement, general structural analysis software II (GSAS-II) was used with the Rietveld method of refinement. The patterns were refined using an orthorhombic $Pnma$ (No. 62) standard (ICDD #04-015-6173). The background correction was carried out using a Chebyshev polynomial function with up to 12 background coefficients. The reliable fits (the fits for which the Rietveld refinement R value, namely the R_{wp} and the goodness-of-fit (GOF) are close to desirable values) are only considered for analysis. N_2 physisorption isotherms were performed employing a Quantachrome surface area analyzer. Sample degassing was carried out at 140°C for 5 h before analysis. The BET isotherm equation was used for calculating the specific surface area and the DFT method was used for deriving total pore volume. Raman analysis was carried out using a Raman spectrometer from WiTec having a green laser (532 nm). Elemental oxidation states were studied through X-ray photoelectron spectroscopy using a Thermo Fischer Scientific which uses an $\text{Al } K_\alpha$ radiation. A Carbon-Sulphur analyzer (CS-444, Leco, USA) was employed for measuring the amount of carbon (wt%) in the synthesized samples. Chemical analysis of Li, Fe, P and Ni was examined through Inductively coupled plasma spectroscopy (ICP-OES, Agilent 5110). Before analysis, approximately 100 mg of the sample was digested in a 1:3 mixture of HNO_3 and HCl , and used for the measurement. Thermogravimetric analysis (TGA) of synthesized materials were tested at a temperature range of 30°C to 800°C , and under heating rate of $10^\circ\text{C}/\text{min}$ in air atmosphere.

2.2. Electrochemical characterization

Electrode sheets were coated by mixing (in wt%) 80 % cathode active material, 10 % carbon conductive agent (CNT-multiwalled, TCI C2154, 20–40 mm diam., 5–15 μm length) and 10 % polymeric binder (poly-vinylidene fluoride, pvdf, Solef® 5130/1001 Solvay) using NMP solvent (*n*-methyl 2-pyrrolidone, 98 %, LOBA Chemie) thoroughly and finally coating the wet slurry on a carbon-coated aluminum current collector (18 μm , GELON LIB group) using a doctor blade set-up (AFA-III automatic thick film coater, MTI Corp.). Before wet-slurry mixing, synthesized materials were mixed with CNT and then ball milled for a period of 2 h in a desk-top planetary ball miller (SFM-1, MTI Corp.) using ceramic vials using 3 mm ZrO_2 balls (ball:powder = 20:1). The wet thickness was set to 150 μm resulting in an mass loading of 1.5 mg cm^{-2} for working electrode, after drying at 80 °C overnight and roll pressing (hot roll press GN-MR-200H). Electrochemical validation of material was carried out using swagelok-type half-cell assembly. Li-foil was employed as a counter electrode, where GF/D glass fiber (GE healthcare) was used as the separator which was soaked with 1 M LiPF_6 (EC:DMC in 1:1 vol%, with 2 % VC) electrolyte as additive. All cells were assembled inside an Ar glovebox (Mbruan unilab pro SP), where H_2O and O_2 were maintained below 0.1 ppm.

Cyclic voltammetry (CV) experiments were demonstrated at a scan rate of 0.1 to 0.5 mV s^{-1} , between the voltage limits of 2.0–4.2 V (vs Li/

Li^+) and electrochemical impedance spectroscopy (EIS) was done under the limits of 10^{-2} Hz $\leq f \leq 10^6$ Hz with a sinusoidal amplitude of 10 mV. Here, CV and EIS experiments were done using PARSTAT MC2000 electrochemical workstation. Further, the constant-current/galvanostatic experiments (2.0–4.2 V) were performed to evaluate the rate performance and cycling stability of coated electrodes at different C-rates using Arbin instruments (BT2000). Here, C/n is the calculated current density which is needed to completely discharge or charge a cell in n h. For example, a C-rate of $C/10$ or $5C$ means that the cell is fully charged or discharged in $n = 10$ h or $n = 1/5$ h, respectively. For the long cycling experiments, the cells were cycled at $C/10$ thrice before stepping up the current rate to 1C or 5C. More importantly, active material mass was considered to calculate the specific charge/discharge capacities. All the electrochemical measurements were conducted at 28 ± 1 °C.

We performed density functional theory (DFT) [25] calculations on the LFP and Ni-doped LFP using the Vienna ab initio simulation package (VASP) [26–28]. We utilized plane-wave basis sets (with a cut-off energy of 520 eV), projected augmented wave (PAW) [29] potentials and the Hubbard U corrected [30,31] strongly constrained and appropriately normed (SCAN) [32] exchange-correlation functional. We started with the experimental olivine LFP structure, as available in the inorganic crystal structure database [33] and relaxed its atomic positions, cell

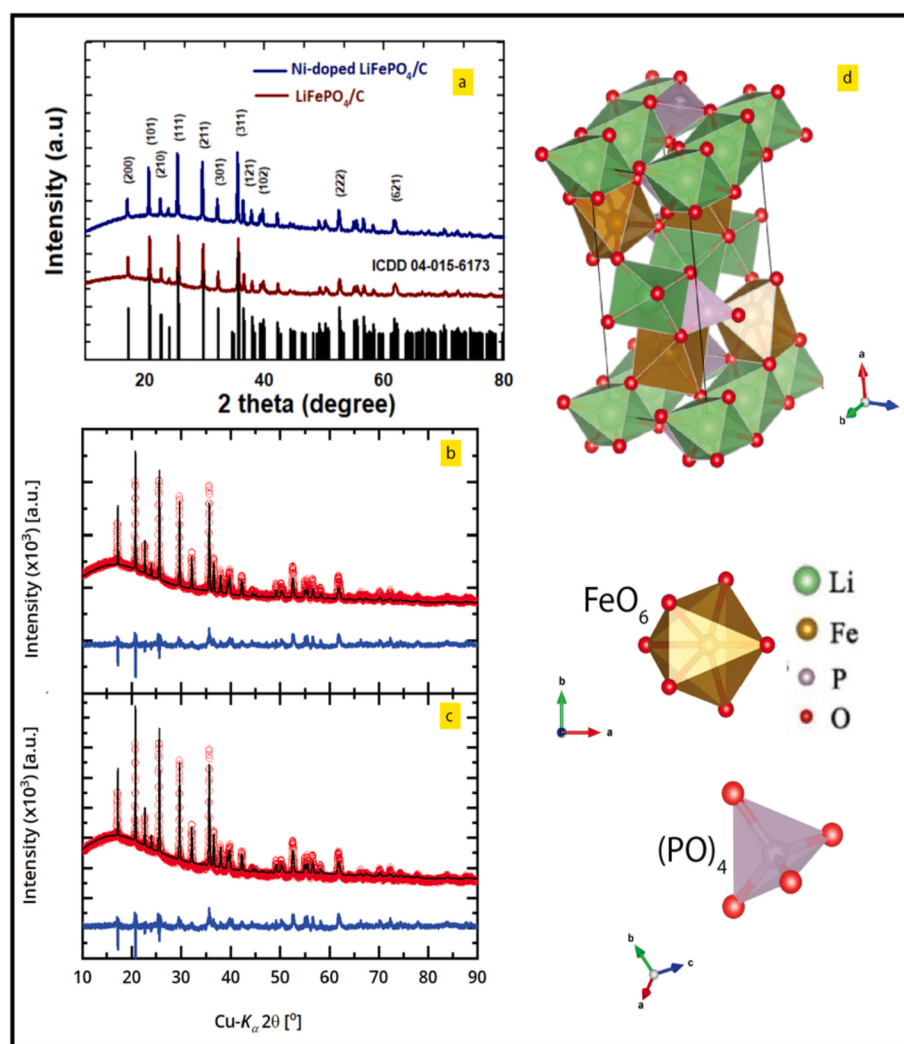


Fig. 1. XRD patterns (a) Rietveld refinement (b, c) of LiFePO_4/C and Ni-doped LiFePO_4/C ; refined crystal structure of carbon-coated Ni^{2+} doped LiFePO_4 prepared in this study (d).

shape, and cell volume without preserving any symmetry. Subsequently, we constructed a $2 \times 2 \times 1$ supercell (ensuring all lattice parameters are above 9 Å in length) of the relaxed LFP and randomly substituted one of the Fe sites with a Ni and relaxed the atomic positions only of the substituted structure. All structure relaxations were performed with an energy convergence criterion of 10^{-5} eV, and an atomic force convergence criterion of $|0.03|$ eV Å $^{-1}$. Finally we performed the density of states (DOS) calculations on the relaxed Ni-LFP and a $2 \times 2 \times 1$ supercell of relaxed pristine-LFP using an energy convergence criterion of 10^{-6} eV. All the structure relaxations and DOS calculations were performed using Γ -centred k -points with a mesh density of 48 Å $^{-1}$, and Hubbard U corrections of 3.1 for Fe and 2.5 for Ni [34,35].

3. Results and discussions

XRD profiles of synthesized pristine and doped carbon coated LFP (Fig. 1a) were indexed to orthorhombic structure with space group *Pnma* #62. Also, the diffraction peaks in the case of both LFP/C and Ni-LFP/C cathode materials align with the standard data (ICDD #04-015-6173) of LiFePO₄. Further, no other crystalline phases have been observed, implying that the ball-milling method combined with carbothermal reduction is apt for preparing phase pure (orthorhombic) LFP/C and Ni-LFP/C samples. Presence of controlled inert working (Ar gas flow) atmosphere during calcination and the carbon coating that creates reducing atmosphere, helps in preventing the formation of possible impurities such as Li₃Fe₂PO₄, LiFeP₂O₇, Fe₂O₃, Fe₂P etc.

Moreover, no extra diffraction peaks associated with carbon or Ni²⁺ doping are observed, suggesting that carbon in LFP/C and Ni-LFP/C is amorphous [36], and the amount of Ni²⁺ in Ni-LFP/C is be fractional. Rietveld refinements are conducted on the XRD patterns (Fig. 1(b, c)), and the corresponding results are tabulated as shown in Table 1, and the crystal structure of LFP has been shown in Fig. 1(d). The cell volumes of LFP/C and Ni-LFP/C from Rietveld refinements are found to be 292.19 (13) Å and 291.47(11) Å, respectively. Similarly, the lattice constants *a*, *b* and *c* for LFP/C and Ni-LFP/C are found to be 10.336, 6.014, 4.70 and 10.327, 6.008, 4.698 Å, respectively. It can be observed that the lattice parameters have reduced after incorporation of Ni²⁺. The doping of Ni²⁺ in the LiFePO₄ crystal structure can be ascribed to the small radius of Ni²⁺ (*r* = 69 pm), than Fe²⁺ (*r* = 74 pm) [37]. This size difference resulted in the reduction of Ni-doped LiFePO₄/C lattice parameters, which is supported by the Rietveld '*R*' values and the observed lowering in the lattice parameters (Table 1). Furthermore, the bond length averages of P—O and Fe—O reduced when Ni²⁺ is doped into LFP unit cell, whereas it increased for Li—O bonds. This adjustment in bond lengths also results in the broadening of the Li-ion diffusion path i.e., in the [010] direction of the LiFePO₄ crystal structure [38] due to Ni²⁺ doping on Fe²⁺ site of LFP. Overall, XRD, and Rietveld refinement revealed the formation of pure LFP phase without any impure phases.

From the FE-SEM micrographs captured at lower and higher

magnification as shown in Fig. 2(a-f) depicts insignificant morphological changes for both LFP/C and Ni-LFP/C samples. Similar conclusions can also be drawn regarding particle size for both pristine and doped LFP. Pristine and doped LFP samples have particle sizes around 1 μm and show pulverized and irregular spherical morphology. However, less agglomerated particles are observed for Ni-LFP/C than for pristine LFP/C. Aggregation of particles leading to tortuous travel path for e^- and Li⁺ ions results in poor kinetics for LFP/C. The reduction in agglomeration observed upon Ni doping can be attributed to the influence of Ni²⁺ ions on the crystal growth kinetics and surface energy during synthesis. Specifically, the incorporation of Ni²⁺ into the LFP lattice modifies the local bonding environment and alters the surface energy of growing crystallites [39]. This change can inhibit uncontrolled particle growth and reduce the tendency of primary particles to fuse during high-temperature calcination.

Moreover, Ni doping may introduce slight lattice strain and distortions that interfere with the oriented attachment or aggregation of nanoparticles. As a result, the final particles exhibit more uniform size distribution and reduced agglomeration. This behaviour has also been reported in other doped cathode materials [40–42], where transition metal dopants act as grain growth inhibitors by modifying diffusion pathways during synthesis.

Further, High Resolution TEM images of both the samples were presented in Fig. 3. Moreover, LFP/C reveals a primary particle size of 50 to 100 nm (Fig. 3(a1)), with a carbon coating of ~5 nm thickness (Fig. 3(b1)). Distinct lattice fringes shown in Fig. 3(c1) with *d* spacing value of 2.17 Å corresponding to (311) plane. SAED pattern shown in Fig. 3(d1-e1) confirms the polycrystalline formation. Corresponding mapping results shown in Fig. 3(f1-j1), and EDS maps (Fig. 3(k1)) confirms the presence of all the elements in the LFP/C. Core-shell structured Ni-LFP/C shown in Fig. 3(a2) depicts the uniform carbon coating around core Ni-LFP/C. Carbon coating on the particles can be observed which is having thickness of ~4–5 nm (Fig. 3(b2)). As shown in Fig. 3(c2), lattice fringes with 1.305 Å represents (211) crystal plane. Polycrystalline Ni-LFP/C formation was witnessed with SAED pattern shown in Fig. 3 (d2-e2) with presence of ordered bright spots. The EDS map (Fig. 3(f2-k2)) shows a homogeneous distribution of dopants within the Ni-LFP/C sample. The presence of thin layer homogeneous carbon coating and Ni²⁺ – ions doping in Fe²⁺ sites of LFP synergistically contributing for the increase in ionic and electronic conductivity and thereby exhibit promising electrochemical performance. The HRTEM images of Ni-LFP/C measured at different parts of the material presented in Fig.S1. The figure clearly unveils the homogeneous spreading of carbon coating and the presence of a thin layer (5 nm) of carbon all around LFP particles. The carbon coating on LFP particles can be seen as core-shell structure, core being LFP material and thin carbon layer presumed as shell on core LFP material. Despite increasing the electronic conductivity, carbon coating on LFP particles mitigate the LFP grain growth during heating, and avoid direct contact of electrodes with electrolyte. Furthermore, to assess the impact of thickness of carbon coating and its distribution, Ni-LFP/C material has been synthesized by varying the concentration (10 wt%, 15 wt%, and 20 wt%) of carbon precursor and the resulting materials were characterized by HR-TEM and C—S sulphur analysis to identify the variation in the thickness of carbon coatings and carbon content. Further, HR-TEM studies on doped LFP synthesized with different carbon concentration were shown in Fig. S2. When the lower concentration (i.e. 10 wt%) of carbon precursor used for the synthesis of Ni-LFP/C, non-uniform distribution of carbon layer on LFP particle was observed. The thickness of the carbon layer around the LFP particle is measured to be ~2 nm (Fig.S2(a1- a4)). Such kind of non-uniformity and lower carbon thickness causes poor electronic conduction and metal ion dissolution in electrolyte respectively. The HRTEM images of Ni-LFP/C with 15 wt% concentration of carbon precursor showed the presence of homogeneous carbon coating with a carbon layer of approximately 5 nm thickness on LFP particles (Fig. S2 (b1-b4)).

Table 1
Crystallographic lattice and Rietveld refinement parameters for the synthesized samples.

	LiFePO ₄ /C	Ni-LiFePO ₄ /C
Crystal structure	Orthorhombic	Orthorhombic
Space group	<i>Pnma</i> (No. 62)	<i>Pnma</i> (No. 62)
<i>a</i> (Å)	10.33(4)	10.32(4)
<i>b</i> (Å)	6.01(20)	6.00(19)
<i>c</i> (Å)	4.70(16)	4.69(18)
<i>V</i> (Å ³)	292.19(13)	291.47(11)
$\alpha = \beta = \gamma$ (°)	90	90
Average Li—O (Å)	2.07	2.16
Average Fe—O (Å)	2.22	2.15
Average P—O (Å)	1.56	1.54
<i>Z</i>	4	4
<i>R</i> _{wp} (%)	1.489	1.400
GoF (%)	2.43	2.22

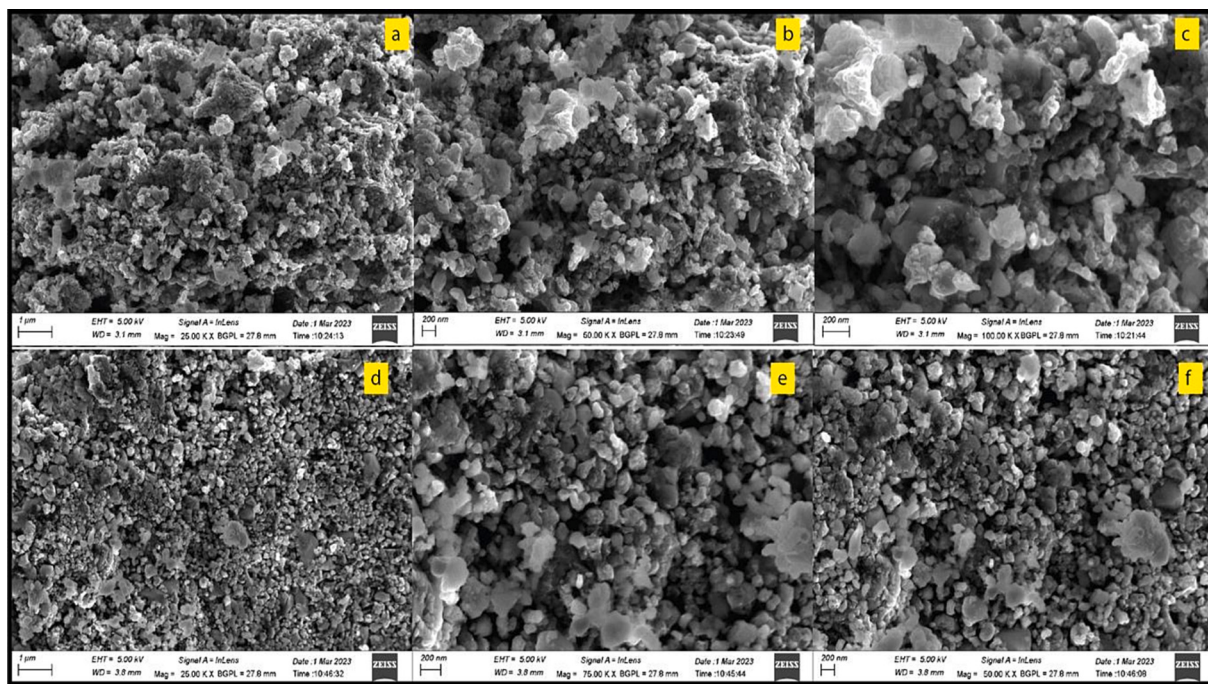


Fig. 2. FESEM images of LiFePO₄/C (a-c) and Ni-doped LiFePO₄/C (d-f).

When, further increasing the carbon content (i.e. 20 wt%), the formation of thicker carbon coating of ~10 nm observed on LFP particle as shown in Fig. S2(c1 and c4). In general, the thickness and distribution of carbon greatly influence the materials physico-chemical characteristics and its electrochemical performance as reported previously for carbon coated LFP [12]. Homogeneous carbon coating on the particles aids in improving the electronic conductivity by reducing the interparticle resistance for electron flow and thus improve the electrochemical performance. Further, ICP-OES analysis of Ni-LFP resulted in weight percent (wt%) of Fe, Li, Ni, and P elements as follows, i.e., 30.46, 4.16, 0.32, and 16.16 respectively. The carbon content measured by C—S analysis, for Ni-LFP/C with 10 wt%, 15 wt%, and 20 wt% carbon precursor is 2.8 %, 3.7 % and 4.3 %. The high carbon content resulting in thicker carbon coating on LFP particles leads to slower Li-ion diffusion and thus low energy density, as it possesses a high amount of porous carbon. Hence, among all Ni-LFP/C materials, Ni-LFP/C material with 15 wt% carbon concentration possessing uniform carbon coating with optimized thickness is expected to result in enhanced performance. Accordingly, in view of the benefits and shortcomings of carbon, Ni-LFP/C with 15 wt% carbon concentration is considered as material of interest in the current study. The ICP-OES analysis shows Li, Fe, and P in the amount of 4.2, 34.1 and 19.6 % for respectively. The ICP-OES analysis as well as the EDS mapping images shows the existence of Ni in very dilute concentration.

The XPS analysis is performed to identify the nature of elements and to determine the oxidation of both samples Fig. 4(a, b). For the sample LFP/C, concerned peaks for Fe 2p_{3/2} and Fe 2p_{1/2} are located at 711.39, and 724.45 eV sequentially. The deconvoluted core level Fe 2p spectra (Fig. S4) with presence of Fe 2p_{3/2}: ~709–711 eV, and Fe 2p_{1/2}: ~722–725 eV with difference in the binding energy of 13 eV clearly shows the presence of Fe in the Fe²⁺ oxidation state. Additionally, the XPS survey spectrum has been added to the Supporting Information (Fig. S5) to confirm the elemental composition and to demonstrate the absence of any unexpected elements. Similarly, for Ni-LFP/C the peaks are followed at 711.22 and 724.28 eV for Fe 2p_{3/2} and Fe 2p_{1/2}, respectively. This shows that Fe in both the synthesized samples (with and without doping) are present in divalent state [43]. Regarding the Ni²⁺ signal, we note that Ni was not clearly detected in the XPS spectra,

likely due to its low doping concentration (1 mol%) and the surface-sensitive nature of XPS. Since Ni is present in a very small amount and primarily incorporated within the bulk crystal structure, its surface concentration may fall below the XPS detection limit. However, the presence of Ni in the material has been independently confirmed through ICP-OES analysis, which provides more accurate bulk elemental quantification. Fig. 5(a, b)) shows the isotherms derived from N₂ adsorption-desorption experiments. Both the cathode materials show a type IV isotherm with H3 hysteresis loop [44]. The LFP/C cathode material shows a BET surface area of 61.6 m² g⁻¹ with mean pore radius of 81.3 Å, while the Ni- LFP/C cathode material shows a SSA of 70.57 m² g⁻¹ with an average pore radius of 68.9 Å. The increased specific surface area in Ni-doped LiFePO₄/C can be explained by two synergistic effects. First, nickel species act as a catalyst during calcination, promoting partial graphitization and modifying the carbonization pathway of the carbon precursor; catalytic graphitization produces thinner, more ordered carbon and alters surface morphology of the particles. Second, Ni incorporation into the LFP lattice induces local strain and modifies surface energies during crystallization, which suppresses grain growth and particle coalescence, yielding smaller primary particles and less agglomeration. Together these effects produce a higher surface area and a carbon coating with improved electronic connectivity, both of which contribute to the observed enhanced rate performance [39,45]. In addition, the pore volume (DFT method) of LFP/C and Ni-LFP/C is 0.205 and 0.213 cc g⁻¹, respectively.

The mesopores likely resulted from the expulsion of gas (CO₂) during the carbothermal reduction with citric acid, leaving behind a high number of small voids. The TGA thermogram for both pristine and doped samples are displayed in Fig. 5 (c, d). We can observe some phase transition regions in both the thermograms. During the initial phase, from RT-300 °C. The material undergoes dehydration and loses all water molecules. Thereafter, the material is oxidized to form Li₃Fe₂(PO₄)₃ and Fe₂O₃, which can be observed from the weight gain around 400–600 °C. Lastly, the material experiences weight loss after complete combustion of carbon around 600–800 °C [46]. Corresponding particle size histograms (Fig. 5(e,f)) measured from DLS for LFP/C and Ni-LFP/C representing the μm (D₅₀) sized particles formation as discussed earlier. Raman spectroscopy was employed to gain insights into uniformity of

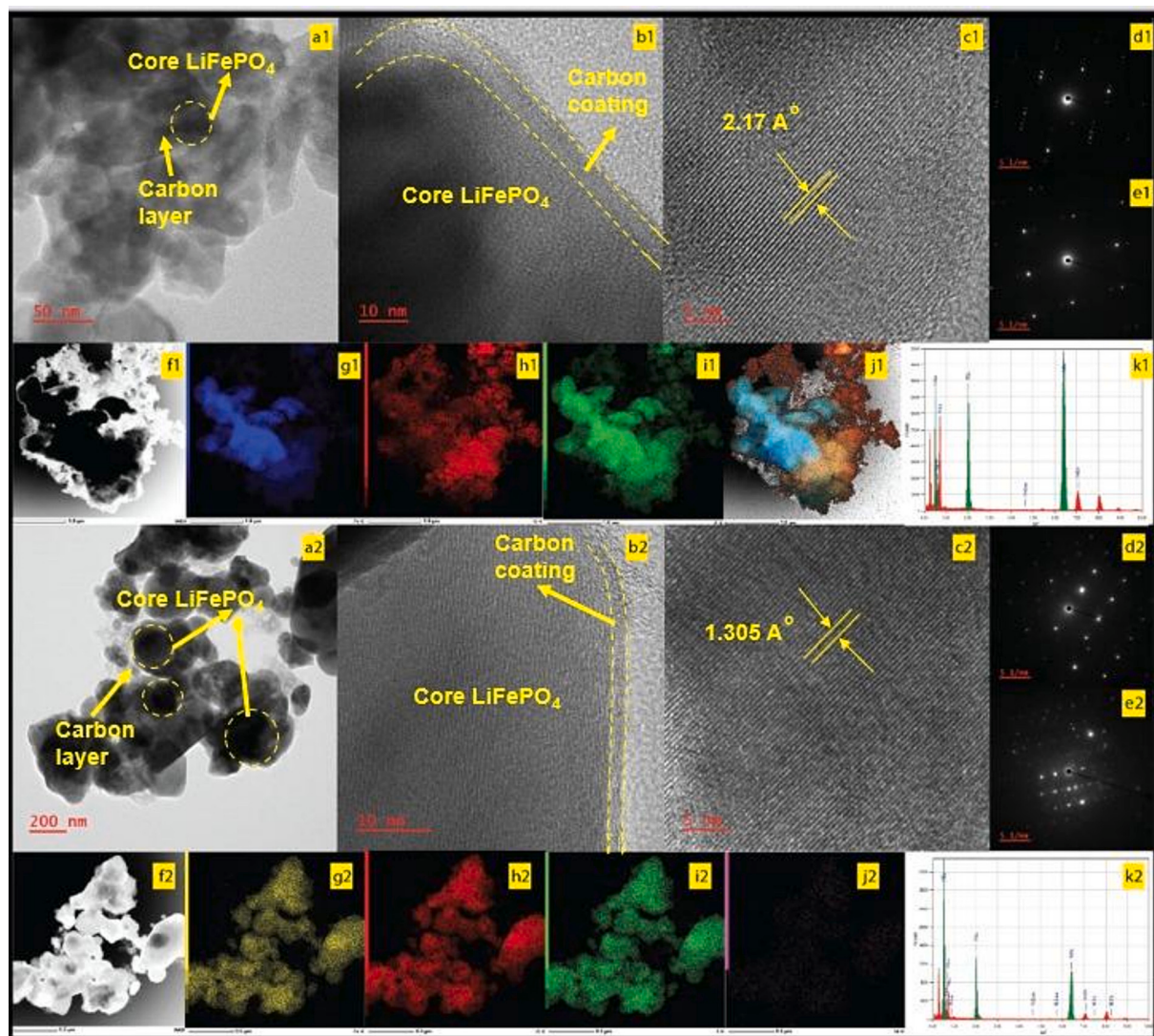


Fig. 3. HR-TEM analysis of LiFePO₄/C (a1-k1) and Ni-doped LiFePO₄/C (a2-k2).

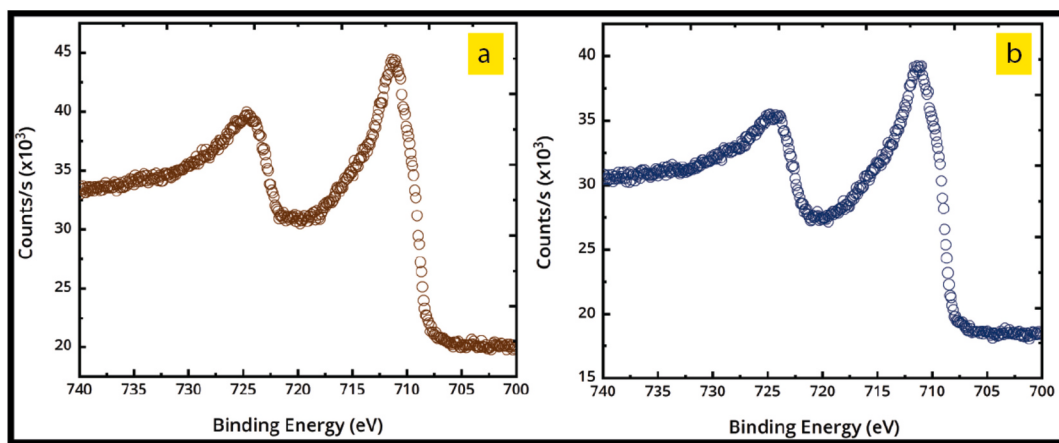


Fig. 4. XPS core-level Fe 2p scan of LiFePO₄/C (a) and Ni-doped LiFePO₄/C (b).

carbon coating for both the samples. As depicted in Fig. 5(g,h), Raman spectra resulting in peaks in the wave number ranging from ~ 100 – 700 cm^{-1} indicates Fe—O vibrational peaks, whereas Raman shift at 900 – 1100 cm^{-1} is due to PO₄ vibrations respectively for LiFePO₄/C and

Ni-doped LiFePO₄/C. Both samples exhibited D and G bands of carbon at following wavenumber i.e., at 1300 , and 1600cm^{-1} consecutively [47,48]. Additionally, the I_D/I_G ratio, which is considered as an indicator of degree of carbon graphitization measured to be 0.99 and 0.96

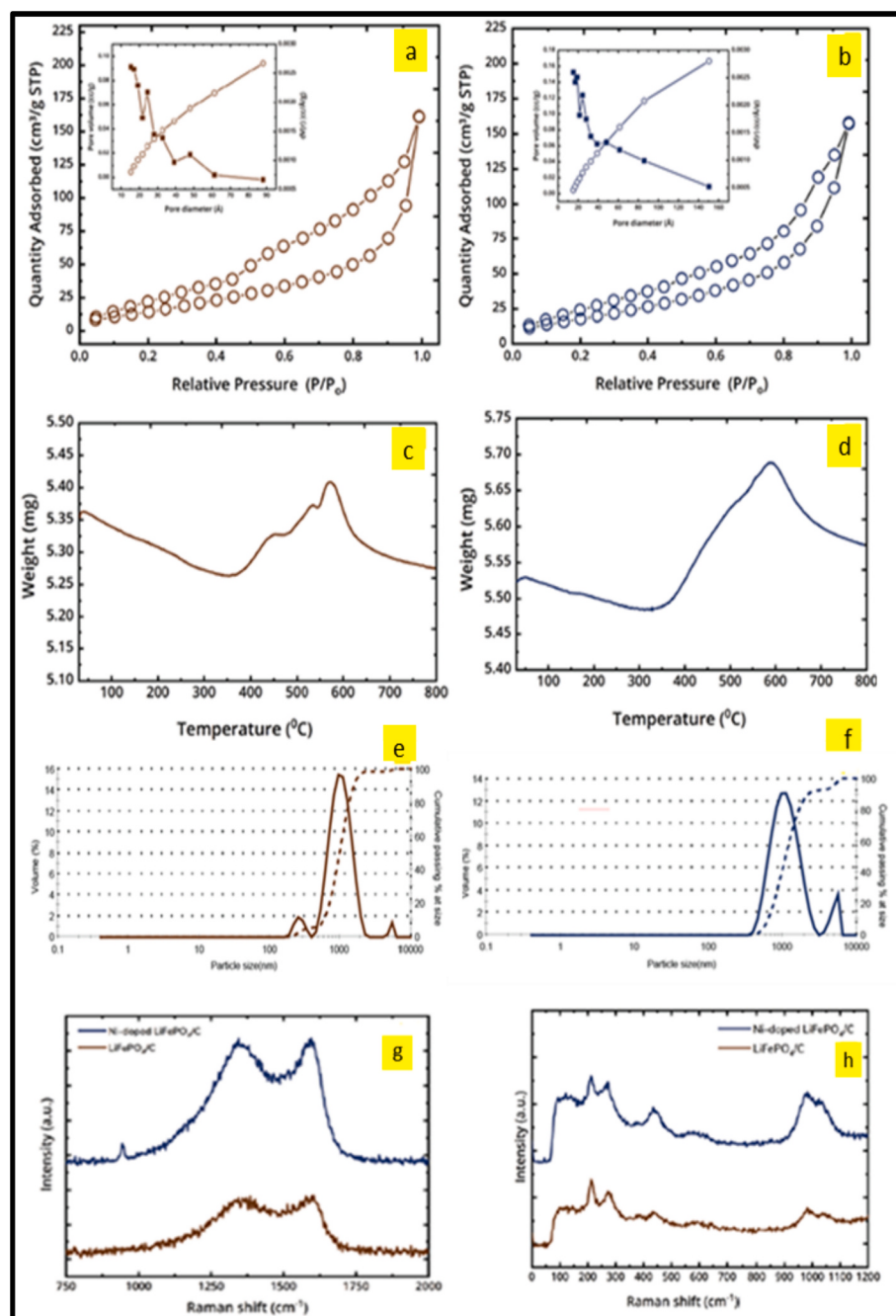


Fig. 5. BET isotherms (a-LFP/C, b-Ni-LFP/C), TGA (c-LFP/C, d-Ni-LFP/C), particle size distribution (e-LFP/C, f-Ni-LFP/C) and Raman spectra (g, h) of the prepared samples.

for LFP/C and Ni-LFP/C respectively. Following studies indicate that Ni-LFP/C exhibits a higher degree of graphitization, contributing to enhanced electronic conductivity and superior electrochemical performance [49,50]. The reason for higher degree of graphitization in the Ni doped material could be related to catalytic graphitization occurring during calcination, facilitated by the presence of Ni metal-ions, that used as a dopant source in the present study [51]. Though an amorphous layer of carbon forms during the initial calcination process, continuous calcination allows amorphous layers to dissolve into nickel catalyst and form as graphitic layer carbon. Further, these layers will be moved towards the particle surface and act as bridges for electron transfer. Nevertheless, the quantity and quality of coating carbon coating performs a significant role in the electrochemical properties of material. If the carbon coating is thin and non-homogenous, then the transport of

electrons is affected in the uncoated region as it slows down Li⁺ diffusion. The thin layer carbon coating also results in metal dissolution in electrolyte and will reduce the cycle life. If thicker carbon coating forms, then the diffusion of Lithium ion through it will be slow. In addition to doping and carbon coating, the particle size of materials presents a vital role in enhancing the performance of material. With an optimal particle size, Li-ion diffusion channel length is reduced. Thus, the particle dimensions are playing an active role in achieving high-rate performance [52] in LIB during the charge/discharge process.

Cyclic voltammograms of pristine and doped LFP are displayed in Fig. 6 (a1, b1). Two samples with and without doping display the typical oxidation-reduction curves of Fe^{2+/3+} occurring around 3.4 V. The Ni-LFP/C electrode exhibits an oxidation reaction peak of Fe (Fe^{2+/3+}) at 3.5 V, while reduction reaction peak (Fe^{3+/2+}) at 3.4 V. For the LFP/C

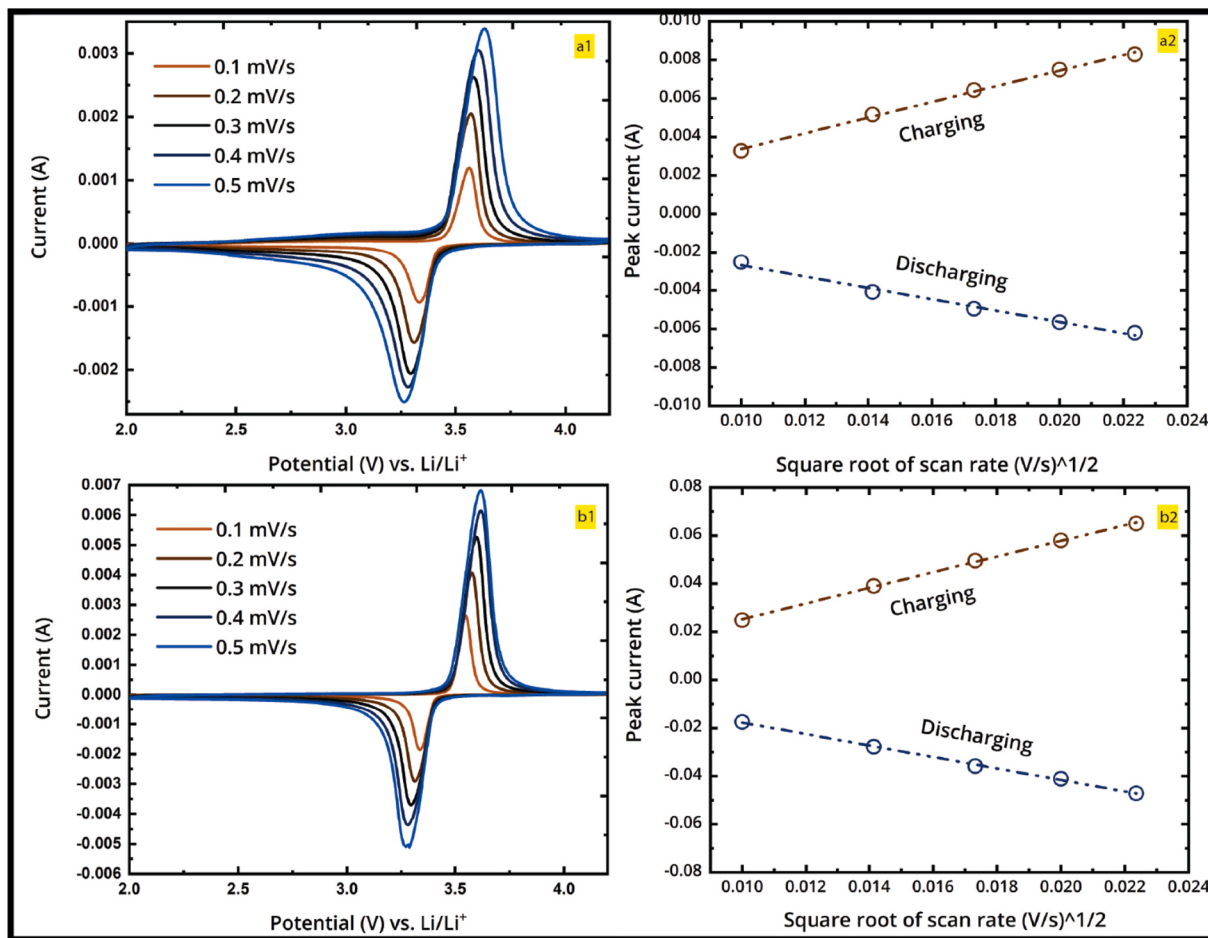


Fig. 6. Cyclic voltammetry scans of LiFePO₄/C (a1) and Ni-doped LiFePO₄/C (b1) electrodes measured at sweep rates; Linear curve plotted for peak current vs. square root of sweep rate for LiFePO₄/C (a2) and Ni-doped LiFePO₄/C (b2).

electrode, oxidation and reduction peaks of Fe were observed at 3.6 V and 3.4 V (for 0.1 mV s⁻¹ scan rate) [20,53]. The doping of Ni²⁺ has resulted in weaker polarization and enhanced reversibility of the Li-ions. This is also expected to increase the Li⁺ diffusion kinetics. Cyclic voltammetry showing no peaks regarding Ni²⁺ implies that the dopant is not entirely participating in the redox reactions [52]. Furthermore, diffusion coefficient (D_{Li}) measured for Li⁺ in LFP/C and Ni-LFP/C samples are estimated from the CV curves using Randles-Sevcik equation:

$$I_p = 2.687 \times 10^5 n^3 D^{1/2} A C \gamma^{1/2} \quad (4.1)$$

As shown in Eq. 4.1, I_p represents the peak current (A), n denotes the no. of electrons transferred ($n=1$) for $\text{LiFePO}_4 \leftrightarrow \text{FePO}_4 + e^- + \text{Li}^+$, A represents the area of the electrode (cm²), C denotes Li-ion density (mol cm⁻³) and γ represents the scan rate (V s⁻¹). Fig. 6 (a2, b2) show linear change in the current (I_p) vs square root of sweep rate ($\gamma^{1/2}$) for LFP/C and Ni-LFP/C electrodes. This implies the strong structural stability of active material during continuous lithiation and de-lithiation [54,55]. The D_{Li} for LFP/C and Ni-LFP/C samples are 9.1×10^{-11} , and 5.6×10^{-9} cm² s⁻¹ correspondingly. Notably, the Ni-LFP/C sample exhibits two orders of higher Li⁺ diffusion kinetics, indicating superior stability and reversibility. The increase in lithium-ion diffusion coefficient can be attributed to widening of the diffusion channel along [010] direction, with increase in Li—O bond lengths, and decrease in the bond lengths of P—O and Fe—O.

The charge and discharge profiles of LFP/C reveal that it delivers specific discharge capacities of 163, 160, 155, 133, and 120 mAh g⁻¹ at C/5, 1C, 2C, 5C, and 10C, sequentially (Fig. 7(a1)). On the other hand,

the Ni-LFP/C (15 wt%) electrode shows discharge specific capacities of 171, 166, 161, 150, and 136 mAh g⁻¹ at the same C rates (Fig. 7(a2)).

The theoretical capacity of LiFePO₄ is approximately 170 mAh g⁻¹, corresponding to the full extraction of one Li⁺ ion per formula unit. In our study, the Ni-doped LiFePO₄/C sample delivered a slightly higher specific capacity of 171 mAh g⁻¹ at C/5.

This marginal increase can be attributed to the synergistic effects of Ni doping, uniform carbon coating, and the incorporation of MWCNTs, which together enhance the electronic conductivity and improve the utilization of the active material. Ni doping may facilitate more efficient lithium-ion transport and reduce polarization, allowing the electrode to access near-complete lithium extraction more effectively. Additionally, surface capacitive contributions from the carbon matrix and MWCNTs may provide a minor boost to the overall capacity, especially at lower current rates. Further, half-cell results of Ni-LFP/C materials with different concentration of carbon precursors are shown Fig.S3. Ni-LFP/C with lower carbon content (10 wt%) delivers discharge capacities of 142, 137, 127, 119, and 107 mAh g⁻¹ at 0.1C, 0.2C, 1C, 2C, and 5C accordingly. While with high carbon content (20 wt%) exhibits inferior performance, i.e. 117, 127, 117, 111, and 100 mAh g⁻¹ measured at similar C rates, compared to other two (10 and 15 wt%) Ni-LFP/C materials. The lower capacities of Ni-LFP/C (10 wt% and 20 wt%) could be attributed to non-homogeneous and thicker coating on LFP particles, which results in low electronic and ionic conductivity and thus results in poor electrochemical performance. Ni-LFP/C (15 %) exhibiting good rate capability, excellent cyclic stability and high specific capacity, owing to homogeneous and thin layer carbon coating is found to be consistent with the literature reported previously [12]. Impressively, the

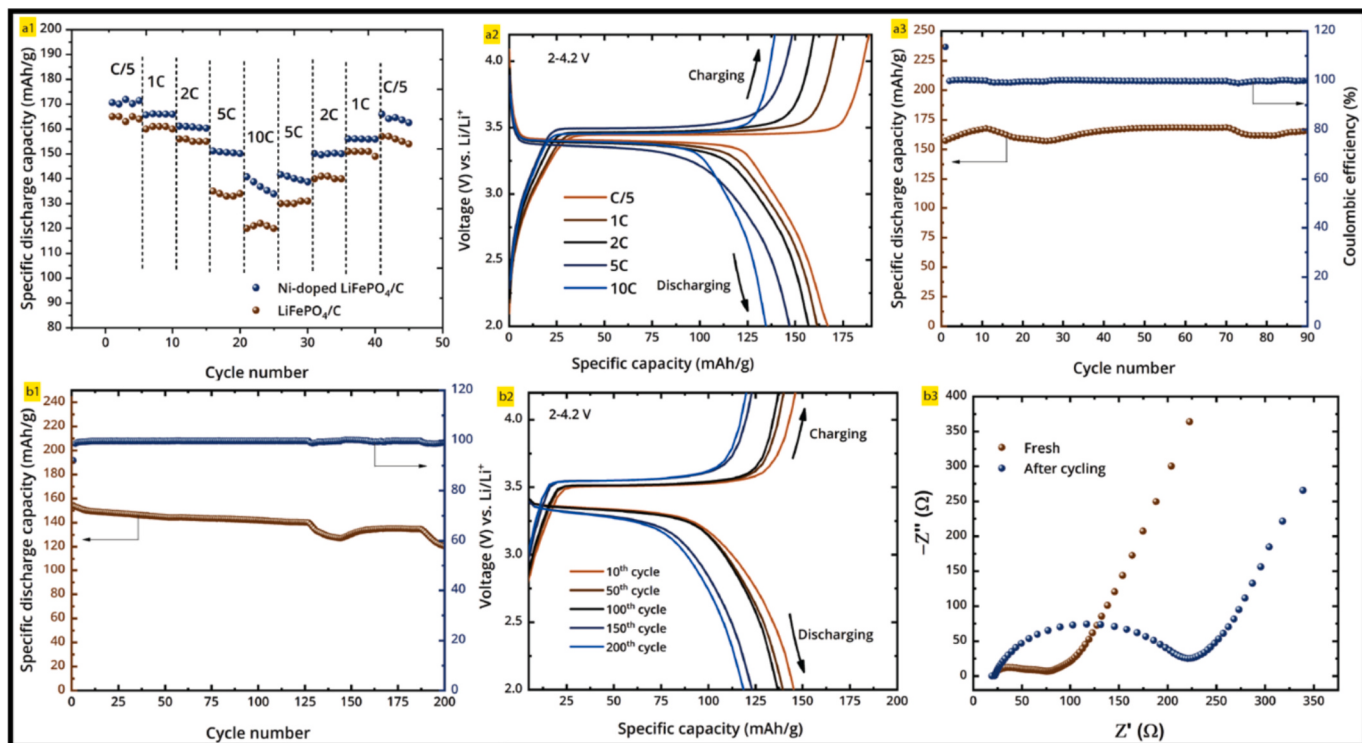


Fig. 7. Rate capability of LiFePO₄/C and Ni-LFP/C electrode (a1), Charge-Discharge studies (a2), Cyclic stability of Ni-LFP/C electrodes at 1C (a3), and 5C (b1); Charge-Discharge studies at 5C (b2) EIS analysis of fresh and cycled Ni-doped LiFePO₄/C electrodes (b3).

Ni-LFP/C electrode demonstrates higher performance than the LFP/C electrode, including specific capacity and rate capability (136 mAh g⁻¹ at 10C). This enhanced rate performance for Ni-LFP/C is supported from improved intrinsic electronic conduction from the uniform dopant distribution and the widened [010] Li⁺ diffusion channel achieved through controlled doping. For assessing the cycling stability of Ni-LFP/C electrode, the cells underwent charging and discharging at C/10 for the first four cycles, followed by 1C cycling for 90 cycles (Fig. 7(a3)). The Ni-LFP/C electrode exhibits excellent cyclic stability without any capacity fading as shown in Fig. 7(b1). Ni-LFP/C electrode demonstrates appreciable cycling performance at 5C current rate with cycling as depicted in Fig. 7(b2). Notably, after 200 charge-discharge cycles, the Ni-LFP/C electrode retains a specific discharge capacity of ~120 mAh g⁻¹ at 5C, representing more than 80 % capacity retention. We have compared the electrochemical performance of Ni-LFP/C reported in the present study with metal ion doped LFP available in the state of art literature as shown in the Table S1. Among the doped LFP cathode materials listed in Table S1, Ni-doped LFP exhibited prominent rate performance than others. Further, it also shown exceptional rate performance of 136 mAh g⁻¹ at 10C (6 min for charging/discharging) with a stable potential curve at 3.3 V, owing to formation of pure LFP with smaller particle size of ~1 μm. Also, the presence of homogeneous and thin layer (~4 nm) carbon coating around LFP particles enhances the electronic conductivity of LFP and regulates the LFP grain growth during heating by maintaining the reducing atmosphere. Further, the uniform distribution of Ni²⁺ in LFP enhances the electronic and Li-ion conductivity of Ni-LFP/C owing to its high electronegativity over Fe and widening of Li-ion diffusion channels along {010} direction by reducing the bonds lengths of Fe—O and P—O, increasing the bond length of Li—O in crystal structure of LFP upon Ni²⁺ ion doping into Fe²⁺ ions of LFP. Thus, the above-mentioned physico-chemical characteristics of Ni doped LFP evident for exhibiting superior electrochemical performance compared with other doped LFP materials demonstrated in the literature. The enhanced cycling performance of the Ni-LFP/C electrode is because of its phase purity, uniform dopant distribution, and increased

electronic and ionic conductivity facilitated by the thin layer of carbon-coating and Ni²⁺ doping. The analysis on EIS data represented in Fig. 7(b3) can be viewed in the further discussions.

When Ni²⁺-ions doped into Fe²⁺-ions site of LFP, the average bond distances of the phosphorous-oxygen (P—O) reduced from 1.56 to 1.54 Å, and iron-oxygen (Fe—O) bond lengths reduced to 2.22 from 2.15 Å, whereas the average bond length of lithium-oxygen (Li—O) increased from 2.07 to 2.16 Å. The shortened P—O bond results in increased stability of the phosphate (PO₄)³⁻ framework that improves the cyclic performance of LFP [22,56]. Also, the reduction of Fe—O and P—O bonds, and expansion of Li—O bonds pave the way for widening of Li⁺ diffusion channel along [010] direction results in fast diffusion of Li⁺ [57].

Our DFT calculations reveal a contraction in the Ni—O octahedron compared to the Fe—O octahedra (~5.3 % decrease in the NiO₆ octahedral volume compared to FeO₆) in Ni-LFP, in qualitative agreement with our Rietveld refinement (Table 1). This contraction causes a slight change in the (about 0.01 Å) bond distances between some Fe—O, Li—O, and P—O bonds for octahedra and tetrahedra that surround the Ni, although these changes are too small to explain the enhancement of Li diffusion. However, Ni-doping facilitates an expansion of ~1.4 % in the volume of the intermediate tetrahedral site (4.22 Å³ in Ni-LFP versus 4.16 Å³ in LFP), that crucially influences the migration barrier that Li-ions have to overcome while diffusing within the LFP structure [58]. This tetrahedral site is illustrated by yellow edges in panels a and b of Fig. 8. Given that an expansion of the intermediate site during Li⁺ migration should reduce electrostatic repulsion from nearby transition metal and P⁵⁺ cations [58], we expect Ni-addition to cause an associated decrease in the Li-migration barrier, resulting in an improvement in Li-diffusivity and power performance that is consistent with our experimental observation. Another factor that can contribute to electrode performance with doping is changes to electronic structure and/or conductivity. Thus, we calculated the electronic DOS in both pristine LFP and Ni-doped LFP, which are plotted in panels c and d of Fig. 8, correspondingly. Blue, orange, red, and green curves indicate Fe d, O p, P

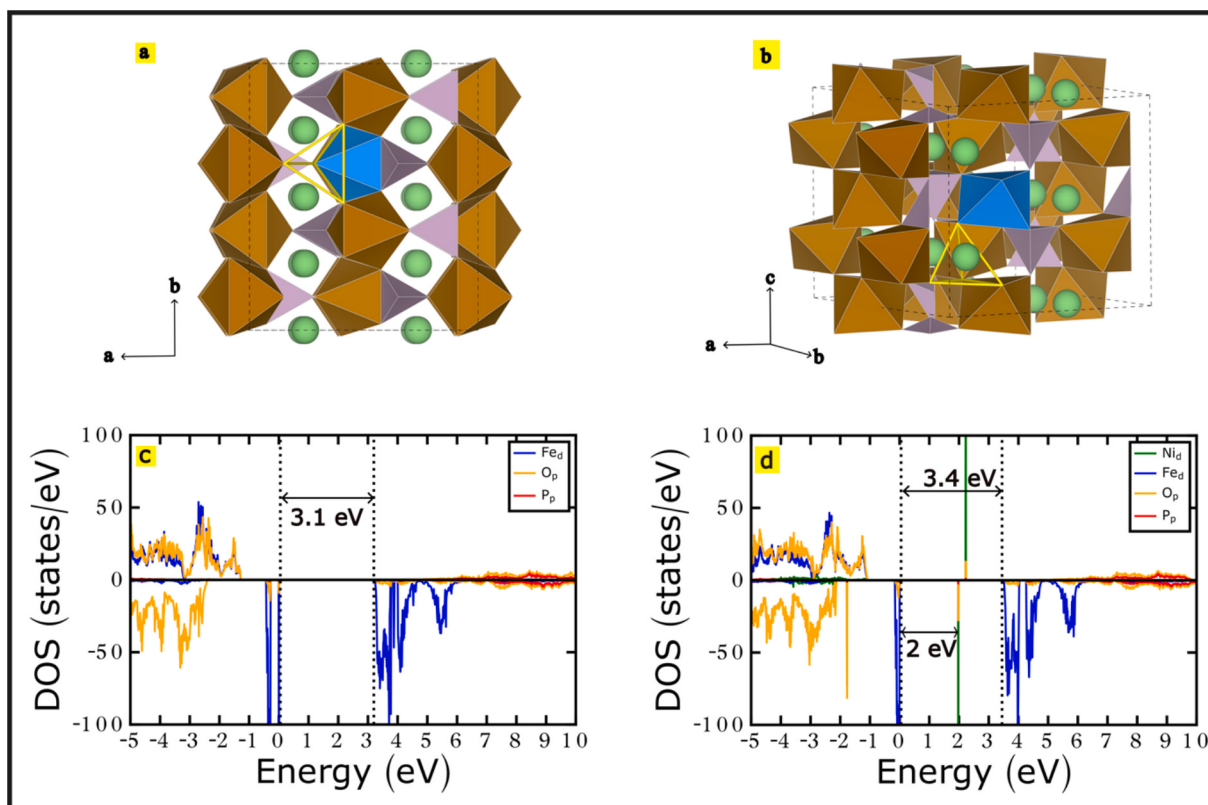


Fig. 8. Unit cell of the Ni-doped LFP marked by dashed lines with (a) c-axis going into the figure and (b) perspective view where Li ions are green spheres, Fe—O octahedra (Fe in the centre and O at the vertices) are brown, Ni—O octahedra (Ni in the centre and O at the vertices) are blue and the tetrahedral void through which Li migrates has yellow edges. Density of states (DOS) of (c) LFP (d) Ni-doped LFP. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

p , and Ni d states, respectively, while the vertical dotted lines signify valence and conduction band edges. The zero on the energy scale is arbitrarily set to the valence band maximum. Importantly, our DOS calculations predict a band gap of 3.1 eV in pristine-LFP, which underestimates the experimental gap of ~ 3.8 eV [59], but is expected given that DFT is a ground-state theory known to underestimate band gaps. Importantly, Ni-addition introduces states within the gap that are ~ 2 eV from the valence band edge and ~ 1 eV from the conduction band edge. Thus, Ni-doping promotes the availability of free electrons in the structure and acts as a n -type dopant. As a result, Ni-doping enhances the electronic conductivity for LFP, which may contribute to both better cycle life and power performance.

Further, the high electronegativity of Ni compared to Fe (Electronegativity order: Mn < Fe < Co < Ni) [60] helps Ni^{2+} to behave like a strong pillar in the crystal structure. Ni^{2+} possessing high electron affinity helps to form a strong bond with electrons and stabilize the LFP structure without any deformation during the charging and discharging process. Thus, the low atomic radii (i.e. 0.69 Å) of Ni^{2+} compared to higher atomic radii (0.74 Å) of Fe^{2+} and high electronegativity of Ni^{2+} ions LFP results in the reduction in the unit cell dimensions and improved electronic conductivity, which attributed to the enhanced electrochemical performance of Ni-LFP/C. EIS analysis was carried out to investigate the charge-transfer resistance of the Ni-doped LiFePO₄/C electrode before and after long cycle performance (5C for 200 cycles). EIS data of the Ni-LFP/C electrode (Fig. 7(b3)) exhibit a half-circle in the medium frequency, and an inclined line (at 45°) in the low-frequency range. The EIS data shows that the charge-transfer resistance increased from 73 to 222 Ω after long-term cycling at high current rate. This might be because of certain side reactions with Ni-LFP/C and the electrolyte, which results in higher impedance inside the cell.

The overall post-mortem analysis shows that the Ni-LFP/C electrode

is quite stable even after fast charging for several hundreds of cycles. From the FE-SEM it is observed that, compared to pristine electrodes (Fig. 9(a1-a2), PC-washed cycled electrodes (Fig. 9(b1-b2)) display very minor structural transformations. The FESEM images of the cycled electrode also show small cracks and separator material intact with the active material. The cracks shown here might have occurred due to long-term cycling or while disassembling/washing the electrode. The microcracks observed in the FESEM image of the PC-washed, cycled electrodes in Fig. 9(b2) are primarily the result of repeated volume changes during high-rate cycling. These cracks are generally considered detrimental to long-term performance, as they can disrupt the structural integrity of the electrode, degrade electrical contact between active material and the conductive network, and hinder efficient Li^{+} diffusion. Over extended cycling, this can contribute to capacity fading and reduced Coulombic efficiency. The Raman (Fig. 9(c1)) and XRD (Fig. 9(c2)) spectra show no change or addition in the peaks, depicting no structural transformations or significant chemical reactions. Further, benchmarking studies were carried out using commercial LFP as shown in Fig. S6. Electrochemical tests show that (Fig. S6) the Ni-doped LFP synthesized in this work delivers a specific capacity of 168 mAh g⁻¹ at 1C, whereas commercial LFP exhibits a lower capacity of 132 mAh g⁻¹ at 1C. This enhanced capacity reflects a significant improvement in energy density, which is crucial for developing high-capacity lithium-ion cells. In addition to improved energy density, the Ni-doped LFP also demonstrates superior rate performance. At a high current rate of 10C, it maintains a specific capacity of 136 mAh g⁻¹, indicating excellent high-rate capability and fast charge-discharge performance. These results suggest that Ni-doping not only boosts the intrinsic energy storage capability of LFP but also enables rapid power delivery and reduced charging time, which are highly desirable for modern high-performance lithium-ion batteries.

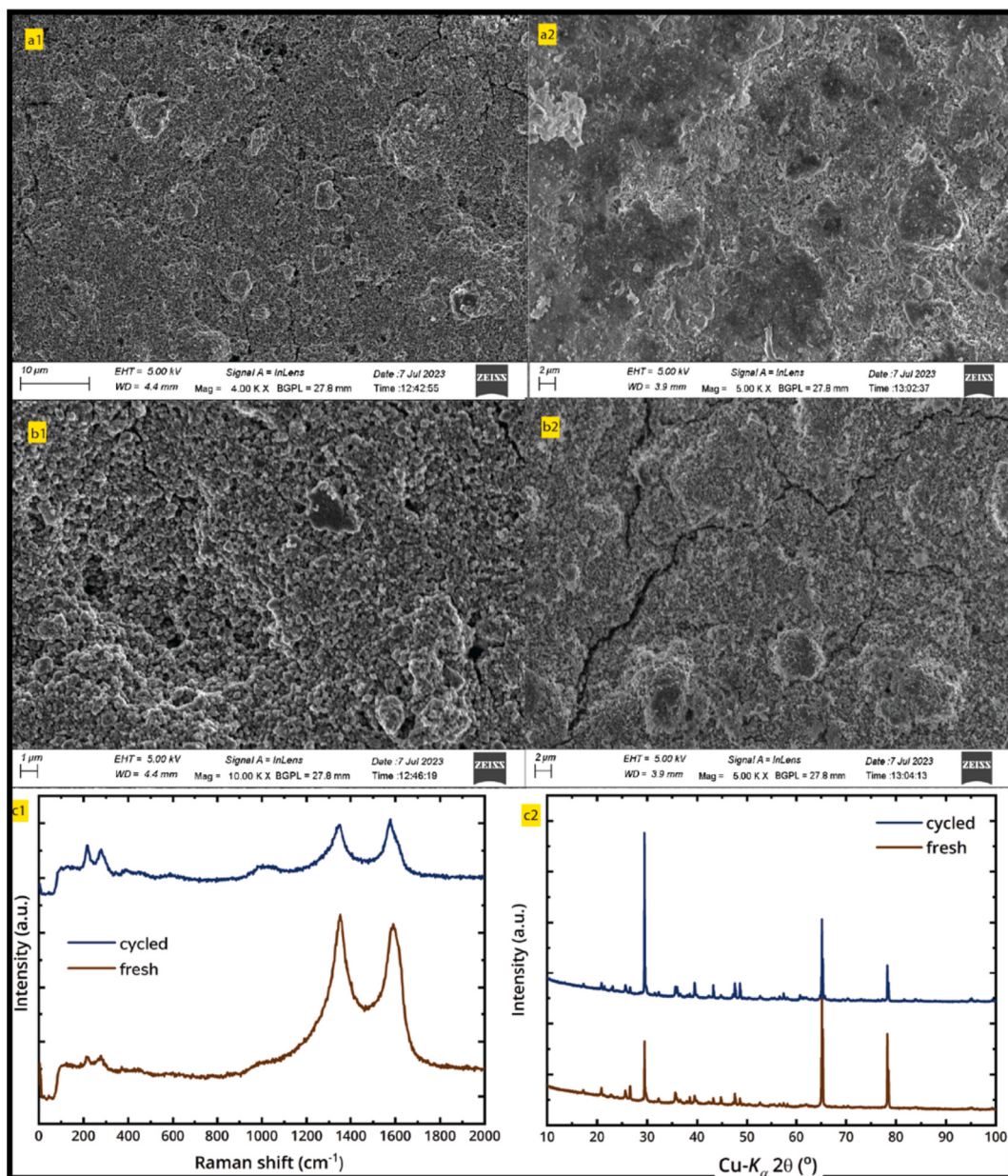


Fig. 9. FESEM images (a1,a2 and b1, b2), Raman spectra (c1) and XRD patterns (c2) of Ni-LFP/C electrode after 200 cycles at 5C.

4. Conclusions

In the present study, we successfully synthesized carbon-coated Ni^{2+} -doped LiFePO_4 (Ni-LFP/C) using a simple, cost-effective, and scalable solid-state ball milling process, achieving a pure olivine-type LFP phase. Morphological analysis revealed a pulverized structure characteristic of the ball milling route, with particle sizes estimated to be around 1 μm . High-resolution TEM-EDX mapping confirmed uniform dopant distribution, while XPS analysis indicated that iron remained in the divalent oxidation state. Cyclic voltammetry results showed reduced electrode polarization and lower charge transfer resistance due to Ni^{2+} doping. Galvanostatic charge-discharge (GCD) tests demonstrated high discharge capacities of 171, 166, 161, 150, and 136 mAh g^{-1} at C/5, 1C, 2C, 5C, and 10C, consecutively. Notably, at a high current rate of 5C, Ni-LFP/C retained over 80 % of its capacity after 200 cycles. In conclusion, Ni^{2+} -doped LiFePO_4/C exhibited markedly enhanced electrochemical performance, including excellent rate capability, high specific capacity, and exceptional cycle stability. This improved performance is owed to

the substitution doping of Fe^{2+} by Ni^{2+} , which led to a reduced unit cell volume, shortened P—O and Fe—O bonds, and elongated Li—O bonds. These structural changes expanded the Li^+ diffusion channels in the [010] direction, resulting in a two-order magnitude increase in lithium-ion diffusion kinetics ($5.6 \times 10^{-9} \text{ cm}^2 \text{ s}^{-1}$) compared with pure LFP ($9.1 \times 10^{-11} \text{ cm}^2 \text{ s}^{-1}$). The exceptional rate performance could be ascribed to the combined effects of Ni^{2+} doping, smaller particle size, homogeneous thin carbon coating, and reduced charge transfer resistance. Overall, this study presents a synergistic approach to synthesize high-performance Ni-LFP/C cathodes, well-suited for the fabrication of high-rate lithium-ion batteries used in electric vehicles, portable electronics, and grid energy storage applications. Moreover, the synergistic approach demonstrated here can be extended to other cathode and anode materials for the development of next-generation high-power LIBs.

CRediT authorship contribution statement

Shreyas J. Kashyap: Writing – review & editing, Writing – original

draft, Validation, Methodology, Investigation, Data curation. **Ch. Gowthami:** Writing – review & editing, Validation. **Sougat Purohit:** Validation, Software. **Gopalakrishnan Sai Gautam:** Validation, Software. **Vadali V.S.S. Srikanth:** Supervision, Investigation, Conceptualization. **R. Vijay:** Project administration, Funding acquisition. **Tata N. Rao:** Project administration, Funding acquisition. **Srinivasan Anandan:** Supervision, Resources, Conceptualization.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.jelechem.2025.119423>.

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