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Structural and Electrochemical Analysis of Iron Doping in $LiNi_{0.6-x}Mn_{0.2}Co_{0.2}Fe_xO_2$ battery

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Abstract: Doping in Li-ion Battery cathode material has become an interesting subject around the world. LiNi_{0.6}Mn_{0.2}Co_{0.2}O₂ is a high energy density cathode material. According to its ability to form particles with a high degree of atomic homogeneity, the co-precipitation method was chosen for the synthesis of the material LiNi_{0.55}Mn_{0.2}Co_{0.2}Fe_{0.05}O₂ (Fe-doped NMC622). In this study, Fe was used as a doping agent because of their high-energy metal oxygen bond dissociation, Oxalic acid is used as precipitant, while ammonia is used as chelating agent. Under the air stream, the oxalate precursor obtained was sintered. The characterization of the Fe-doped NMC622 is performed. Hexagonal-layered material structure shown in x-ray diffraction patterns. XRF analysis confirmed the composition of the final product. SEM (scanning electron microscope) study displayed the material's morphology. The electrochemical performance was achieved by a charge-discharge test at 1/10 C between 2-4.8 V. The Fe doped NMC622 exhibited enhancing cycle efficiency and structural characteristic compared to the un-doped NMC622.

Keywords: NMC; cathode; doping

1. Introduction

Energy is a crucial aspect for human civilization, since almost all activities rely on the availability of energy, one of which is in the transportation sector and portable devices¹⁾. The use of fossil-fuel vehicles has begun to be replaced with more efficient and environmentally friendly electric cars. Energy storage in the form of electronic devices is needed to be efficient, lightweight, not oversized, and has high performance along with its growth. Electricity is the main catalyst for urbanization, industrialization and economic development²⁾.

A significant role in the lithium ion battery performance is cathode material. The cathode's structure is entirely responsible for the transfer process of lithium ions, both during lithium ions de-intercalation (charging) and lithium ions intercalation (discharging)³⁾. Lithium-ion batteries (LIBs) not only used for small but also medium-sized electricity storage systems, but the production of large-scale storage systems has been limited due to safety and economic concerns⁴⁾.

LiCoO₂ (lithium cobalt oxide) is the first layer

structured cathode that has been produced commercially. Since this material was deemed costly, hazardous and highly combustible when used in extreme conditions ⁵⁾, the use of this material began to reduce. With the same characteristics of operating voltage, NMC (Lithium Nickel Manganese Cobalt Oxide) has a far greater potential than LiCoO₂. Typically used LIB cathodes such as LiCoO2. are synthesized from small mineral supplies and thus simple, low-metal-free⁶⁾. Based on economic considerations, NMC has a lower price than LiCoO₂ as Ni and Mn became replacement of cobalt. Because Co is a low natural resource, a substitute cathode material to LiCoO₂. is essential⁷⁾. NMC is a lithium battery cathode material excellent features such as high capacity, better thermal and current stability, relatively high volume density and long life cycle⁸⁾.

Recently, research on lithium ion batteries has concentrated mainly on rising energy density and there's no reduction in life cycle and safety concerns. That is caused by the structural instability of anode and cathode chemical compounds. The instability of the active ingredient can cause thermal runaway cell batteries in an

uncontrolled exothermic reaction. Thermal runaway that caused by highly reactive oxygen released at the cathode, it is caused by phase transitions at high temperatures and flammable organic carbonate-based electrolyte reactions ⁹⁾. Therefore, the structural stability of active cathode with high energy densities such as LiNiMnCoO₂ (NMC) is important in order to ensure stable, high-energy cathode active materials ¹⁰⁾.

Atomic doping is a widely used process to stabilize the cathode's chemical structure. The crystal structure can be specially modified by atomic doping which implies cation substitution in the host structure. High bond dissociation energy offers a good structural stability through structural decomposition of NMC and thus inhibits oxygen release which is an unavoidable procedure ¹¹. The elements for doping that have been investigated are Al ^{12–14}, Mg ¹⁵, Ti ¹⁶, Cr ¹⁷, Mo ¹⁸, Nb ¹⁷, Sn ¹¹ and Fe ^{12,19,20}. Generally, discharge capability decreases with increased concentrations of doping elements due to the presence of doped elements which are not electrochemically active ¹¹, however, the structural stability increases ^{12,16,18,20,21}.

NMC622 cathode materials have been known as next generation cathode materials due to their higher capacity and lower raw material costs than commercialized cathode materials such as NMC532. Many manufacturing companies have also made every attempt to commercialize and adapt the NMC622 to the lithium ion battery technology, such as the energy storage system (ESS).

Transition metals that have high dissociation energy and are quite easy to find in the interesting nature to study is Fe. Previous studies using Fe as doping element has performed. Wilcox synthesized Li(Ni_{0.4}Co_{0.15}Fe_{0.05}Mn_{0.4})O₂ using combustion method in the process with glycine nitrate. Cells containing Li(Ni_{0.4}Co_{0.15}Fe_{0.05}Mn_{0.4})O₂ cycled to 4.3 V showed reduced efficiency relative to Li(Ni_{0.4}Co_{0.15}Fe_{0.05}Mn_{0.4})O₂ while iron should be electroactive in the range of operating voltages and there is no rate improvement impact 12). Park also used Fe as doping element to synthesize $LiNi_1/_3Co_1/_3Mn_1/_3Fe_x(OH)_2$ from LIB that have been used and using hydroxide co-precipitation. Fe 's positive effects on NMC 's electrochemical and thermal properties make it simpler to extract Fe from spent LIBs in leach liquors. The previous study in NMC442 and NMC111, so it can be a novelty to use Fe as a doping in NMC622 that has higher capacity and lower cost, since Indonesia is the first rank in nickel source so if we use more nickel in battery can be good advantages. Manganese and Cobalt also have some contribution in NMC, so it maintained that way.

Thus there is no research that used Fe as a doping on NMC622 batteries using co-precipitation process. Direct doping during co-precipitation is kind of method that more efficient compared to the conventional sequential doping technique.

In this study, we used batch co-precipitation method to

NMC synthesis with perform iron doping (LiNi_{0.55}Mn_{0.2}Co_{0.2}Fe_{0.05}O₂ or Fe-doped NMC622), in an attempt to reduce the complexity of the previously mentioned current processes. Technical grade materials like Li, Ni, Mn, Co and Fe are used for co-precipitation; ammonia used as a chelating agent; and oxalic acid for a precipitation agent. As far as we consider, this approach has never been reported in previous studies. The product's electrochemical performance was tested with graphite as a commercial counter anode in a full cell cylindrical battery. The implementation on NMC and commercial NMC samples of the output values is comparable.

2. Materials and Method

2.1 Tool

The synthesize of Fe-doped NMC622 cathode have used several laboratory tools such as vacuum oven, 200 filter mesh, beaker glass, filter paper, bottle flask, measuring glass, pipette dropper, mortar and pestle, magnetic stirrer, spoon, vacuum mixing, automatic coating machine, hot rolling machine, glove box, furnace. X-Ray Diffractometer (XRD), XRF analyzer, FTIR spectrometer, and Scanning Electron Microscope (SEM) and Battery Analyzer are used for product characterization instruments.

2.2 Material

The primary materials used in the production of Fedoped NMC622 cathode batteries are Ni₂SO₄ from Zenith (High Purity Grade), CoSO₄ technical grade (Rubamin, India), FeSO₄ (Pudak, China). Additional ingredients are Ammonia as chelating agents, oxalic acid as precipitant, AB, PVDF, KS6, LiOH S.A. 98% (Leverton, US), LiPF₆/EC:DEC(3:7) electrolyte, graphite anodes, PP separators.

2.3 Experimental Method

2.3.1 Synthesis $Li[Ni_{0.55}Mn_{0.2}Co_{0.2}Fe_{0.05}]O_2$ (Fe-doped NMC622)

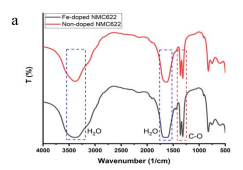
Synthesis of LiNi_{0.55}Mn_{0.2}Co_{0.2}Fe_{0.05}O₂ or Fe-doped NMC622 cathode material by preparing raw materials and additives. The composition we planned is Ni 0.55, Mn 0.2, Co 0.2 and Fe 0.05, it is the modification of NMC622. The metal sulfate (Ni, Co, Mn, Fe) were diluted in distilled water to get 2 M concentration. Using a magnetic stirrer, the solution was stirred to form homogenous solution. Then ammonia and oxalic acid were added simultaneously (equimolar amount) to pH = 2 until a pale cloudy slurry is formed. The slurry was vigorously mixed while the pH and temperature of 60°C was maintained for 2 hours. After the co-precipitation process, the precipitate is taken by passing it into the paper filter and washed until the filtrate is neutral. The precursor is dried and mixed with LiOH using mortar and pestle. LiOH is in 3% excess. Presintering is carried out at 480 °C for 6 hours, and 12 hours at 800 ° C sintering.

The as obtained sample was ground and sieved using 300 mesh SS-filter. Filtered powder were tested using XRF, XRD, FTIR and SEM. Then, Before the coating process NMCF sample, PVDF and AB were dried in an oven. NMC: AB: PVDF with a ratio of 92:3:5 were quickly mixed and dispersed in NMP solvent. The mixing slurry was coated on both al foil and dried for one night in the vacuum oven.

2.3.2 Fe-doped NMC622 Characterization

An XRD (X-Ray Diffractometer) from D2 Phaser Bruker, Germany, was used for the analysis of the crystal structure of the product using CuK α radiation $\lambda = 1.54$ Å with 2θ range of $10-80^{\circ}$ and a scanning rate of 0.02° per second. A scanning electron microscope (SEM) (JEOL JSM-6510LA, Japan) was used in the morphs of the sample. The substances Ni, Mn, Co and Fe were analyzed using X-Ray Fluorescence (Bruker XRF Spectrometer, Germany) in powder form. Fourier Transformed Infrared Spectroscopy (Shimadzu FTIR spectrometer, Japan) was used for analyzed the functional groups and surface composition of the material. Depending on the characterization result, the best possible synthesis material continued battery performance testing in cylindrical cell battery using a NEWARE Battery Analyzer and BTS program with a voltage window of 2.5– 4.3 V at 0.1 C or 20 mA/g (1 $C_{theoretical} = 200 \text{ mAh/g}$).

3. Result and Discussion



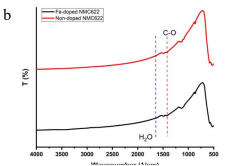


Fig. 1: FTIR spectra Fe-doped NMC622 Precursor (a) and Fe-doped NMC622 Product (b)

The synthesis of Fe-doped NMC622 materials is achieved by a process of co-precipitation. At precipitation, oxalic acid is used as precipitant and as chelating agents

was using ammonia to control the coagulation process on the NMC cathode.

Fig. 1a shows the absorbance spectrum of the Fe-doped NMC622 and Non-doped NMC622 precursors The same pattern has shown The bending vibration of the water molecule can be seen from the big peak around 3340cm⁻¹ and 1600cm⁻¹. The sharp peaks located at around 1300cm⁻¹ correlate to the Carbon-Oxygen bond. Overall, the sample's infrared absorbance spectrum is same as FeC₂O₄.2H₂O, which indicates that the precipitation of NMC-oxalate occurs with the reaction referred to in equation 1-2. After adding excess lithium, the precursors are prepared for sintering. In sintering, the Hydrogen-Oxygen-Hydrogen and the Carbon-Oxygen bonds are lost. This could be used to answer that by sintering shows the loss of bonds at the precursors as shown in Fig. 1b ²²).

$$xNi^{2+}_{(aq)} + yMn^{2+}_{(aq)} + zCo^{2+}_{(aq)} + NH_4OH_{(aq)} \rightleftharpoons Ni_xMn_yCo_z(NH_3)^{2+}_{(aq)} + H_2O_{(l)} \tag{1} \label{eq:1}$$

$$Ni_x Mn_y Co_z (NH_3)^{2+}{}_{(aq)} + H_2 C_2 O_{4(aq)} \\ \longrightarrow Ni_x Mn_y Co_z C_2 O_4 \downarrow {}_{(s)} + NH_4 + {}_{(aq)} \\ \tag{2}$$

An XRF study has confirmed the composition of active cathode synthesized materials. The sample formed for Fedoped NMC622 is 59.94 % Ni, 16.53 % Mn, 18.64 % Co, and 4.89 % Fe, as shown in Table 1. The sample formed for non-doped NMC622 is 59.09 % Ni, 19.89 % Mn, and 21 % Co. The compositions are significantly lower than originally planned values and lithium composition has been set as a mole ratio of 1.03 for transition metals in active cathode materials. It is clearly seen that the composition of Ni is 59.94% among quarternary metals, which is close to the target value of 55%, while the composition of Mn and Co is 16.53% and 18.64% respectively, which is less related to the target value of 20%, and the composition of Fe atoms is 4.89%, which is close to the target value of 5%.. Based on Patnaik²³, K_{sp} value of Ni, Co, Mn and Fe hydroxide is 2.0 x 10⁻¹⁵, 1.9 x 10^{-13} , 1.6 x 10^{-15} , and 8.0 x 10^{-16} respectively. When compared to Ni, Co and Fe, Mn faced the most concentration depreciation. Since Mn has the highest Ksp value (solubility product) relative to the Ksp value of Ni and Co, so that the deposition yield from Mn is lower²⁴). Fe concentration is the closest to target value because it has the lowest solubility, so the precipitation process show more effective.

Table 1. Molar fraction of ternary metal of Fe-doped NMC622 and non-doped NMC622

Cample	Element					
Sample						
	Ni (%)	Mn(%)	Co(%)	Fe(%)		
Fe-doped NMC622	59.94	16.53	18.64	4.89		
Non- doped NMC622	59.90	19.89	21.10	-		

Fig. 2 shown X-ray diffraction patterns of Fe-doped NMC622 and non-doped NMC622 powder. With group space R3 m (No. 166) $^{25)}$, the pattern reflexes indexed to the α -NaFeO2-like hexagonal structure. The reflexes obtained at 38° and 65° 20 in (006)/(102) and (108)/(110) show well crystallized materials $^{26)}$.

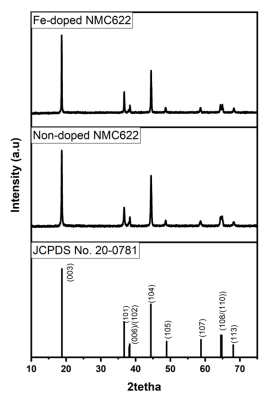


Fig. 2 XRD analysis of Fe-doped NMC622

Table 2 lattice parameters of samples with Fe doping and pure

Sample	a	c	c/a	I003/I104
Fe-doped				
NMC622	2.868	14.285	4.979	1.622599
Non-				
doped				
NMC622	2.870	14.142	4.927	1.370756

Ni²⁺ may have a bad effect on the performance of Fedoped NMC622 cathode material during phase formation, mostly because of the close radii, Li atomic radii replacing Li with Ni²⁺. A stable Ni²⁺ ion blocking the Li-ion pathway is very difficult to oxidize, inhibits its intercalation and deintercalation process which results in low electrochemical efficiency. It is called by cation mixing, and the intensity ratio of I(003)/I(104) evaluation may indicate its degree or level ²⁷⁾. I(003)/I(104) intensity ratio is normally used to specify the level of cation mixing in the Li-layers ²⁸⁾. If the value of I(003)/I(104) is higher than 1.2, cation mixing

is considered poor and normally the layered structure is perfect. I(003)/I(104) intensity ratio value in the Fe-doped material is 1.622 and non-doped material is 1.37, indicating a lower cation mixing. Lattice parameters of Fe-doped NMC622 and non-doped NMC622 shown in Table 2. The lattice expansion was slightly increased in the presence of Fe doping, which indicates that the Fe²⁺ ion had been doped into the Li (Ni_{0.6}Co_{0.2}Mn_{0.2}) O2 lattice during the calcination process. Because metal ions, for example, Ni²⁺ (0.069 nm), Co³⁺ (0.0545 nm), and Mn⁴⁺ (0.054 nm), have smaller ionic radii than Fe2 +, when replaced by Fe²⁺ ions (0.076 nm), Fe²⁺ will enlarge the lattice parameter. This is also the same as that done by previous studies with Mg and Zn doping²⁹).

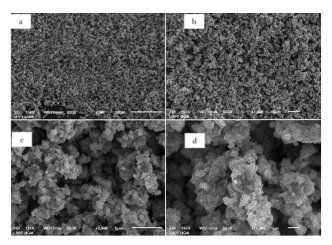


Fig. 3 SEM images of Fe-doped NMC622 (a) at 500 x magnification, (b) at 1000 x magnification, (c) at 5000 x magnification, (d) at 10,000 magnification

Fig. 3 Presents the synthesized Fe-doped NMC622 particles at various magnification. Based on the SEM images, the product has an average primary particle size of 0.509 micrometers with the range of 0.203-1.015 micrometers. Based on Fig. 3d, size of the secondary polyhedral shaped particles is approximately 1.35-6.09 micrometers and the average size of the secondary particles is 3.47 micrometers. Secondary particle is the agglomerate of the primary particle. Using Fe as doped metal in NMC622 have made smaller primary particle size.

The NMC622 sample with Fe doping has an average size that is relatively smaller than that of previous studies using the co-precipitation method on non-doped material³⁰⁾, which, as previously reported, could be caused by Fe doped which may reduce particle size ³¹⁾. Smaller particles however have advantages, as well as larger surface area and faster lithium ion transfer rather than larger particles ²⁷⁾.

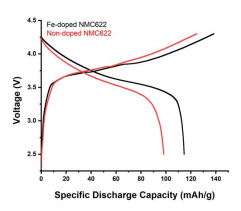


Fig. 5 Charge-discharge analysis of Fe-doped NMC622

Fig. 4 demonstrates the charge and discharge of the asprepared Fe-doped NMC622 and non-doped NMC622 under ambient conditions in a range of voltage 2.5 V-4.3 V using 0.1 C current. The first charge of the as-prepared Fe-doped NMC622 and non-doped NMC622 was resulted a charge capacity of 140 mAh/g and 125mAh/g followed by its first 114.59 mAh/g and 98.33 mAh/g discharge capacity, respectively. Theoretical battery capacity for NMC is 200 mAh/g. Previous study by Park, NMC333 with added 5% Fe have discharge capacity 162.0 mAh/g ²⁰⁾. In the case of fe-doping, the primary explanation for lower discharge capacity in Ni-rich materials is due to increased Li-diffusion hindrance and increased oxidation capability of the Fe ions 32,33). However, because the structure of the cathode, depending on the re-conversion reaction, varies during each charging and discharge process, the efficiency of the cathode depends significantly on the different treatments, such as the mixing of electrode materials, the current rate and the temperature³⁴⁾.

The cycle performance of the active cathode material in range of voltage 2.5 and 4.3 V at a constant current density of 90 mA/g (0.2C) shown in Fig. 5. a shows the performance of the Fe-doped NMC622 and Non-doped NMC622 cycles. The normalized capacity in b is the actual capacity, the initial capacity, of each sample. Fedoped NMC622 and non-doped NMC622 have different cycling behaviour for 50 cycles at a rate of 0.2 C and have a retention rate of 85.76 % and 62.84 %. Capacity decline occurred during the initial cycles; however, after three to four cycles, capacity is considered stable after 50 cycles for Fe-doped NMC622. But NMC without doping has lower capacity retention and poor stability. In addition, Fe doping obtained the structural stability required for improved cycling 35). shows that the Fe doped NMC622 has a stable coulombic efficiency. The performance capacity of the battery cell at various charging / discharge rates has been calculated in order to check the efficiency of the battery³⁶).

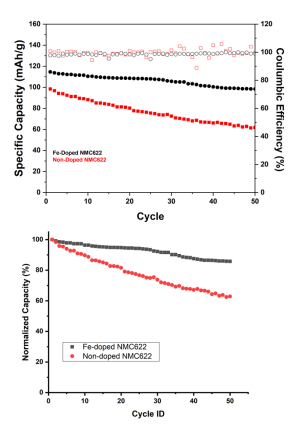


Fig. 4 Cycling performance (discharge capacity and coulombic efficiency) and (b) the normalized cycle performance of Fedoped NMC622 and Non-doped NMC622 at 0.2C

4. Conclusion

The NMC622 cathode which was doped with Fe has successfully synthesized using direct coprecipitation process and studied about electrochemical performance and structural effects of doping. Structural studies show dopants have an effect on particle size. The initial discharge capacity of Fe-doped NMC622 and non-doped NMC622 is 114.59 and 98.33 mAh/g, respectively, while the retention capacity at 0.2 C' for 50 cycles is 85.76 % and 62.84 % respectively. The results show for certain if Fe doping can improve the stability of NMC622. However, future developments in the optimization of process variables are needed in order to obtain better material. Further work is required in the future to obtain a better and more reliable discharge capacity.

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