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Effect of TiDoping on the Structure and Performance of L iNi 75 Co_{0 25} O₂

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Abstract: In this study, a series of Co-doped compounds with the formula of $LNi_{(0,75-x)}Co_{0,25}Ti_xO_2$ (x = 0, 0, 1, 0, 25) was synthesized by using sol-gel method. Their structure, particle size, electrochemical properties were studied by TG, XRD, SEM and electrochemical tests. It was shown that the structure of materials changed from a hexagonal layered structure to a cubic structure as the x value increased. In particular, a mixture of hexagonal and cubic structure was presented at x = 0. 25. The electrochemical performance of the materials depended strongly on the amount of Ti doping. Possible effects on the structure and performance of LNi_(0,75-x) Co_{0,25} Ti_xO₂ due to Ti doping are briefly discussed based upon the experiment and computational results.

Key words: Lithium ion batteries, Cathode materials, Ti doping, Electrochemical performance

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1 Introduction

Lithium cobalt oxide has been widely used as a cathode (positive electrode) material for commercial secondary lithium-ion batteries due to its advantages of easy preparation, high voltage, good reversibility, and high theoretical specific capacity. A LiCoO₂ cathode with a carbon anode to make the first successful Li-DN battery [1-2], which now dominates the lithium battery market There is a limited availability of cobalt, which causes it to have a high price and also cobalt brings the environment pollution^[2]. In order to tackle the problems associated with the high cost and system instability of $LiCoO_2$, a method for the preparation of a version of the compound which has improved electrochemical characteristics and cationic substitutions on the cobalt sites has been sought extensively. Of the se sub stituted compounds, $LNi_{x}Co_{1-x}O_{2}$ has been identified as one of the most

attractive materials The layered $L Ni_x Co_{1-x}O_2$ (0 < x < 1) compounds have been studied extensively as cathode materials for lithium batteries^[6-9]. The poor thermal stability and cyclability of $L Ni_x Co_{1-x}O_2$ (0 < x < 1) demand for doping another element, such as Ti, Al, Mg, Fe, Y, Sr, etc.

Among those doping elements tried, Ti is resourceful in the earth and it has small structure changes when lithium ion intercalation and deintercalation Previous studies ^[10-13] confirmed that doped Ti can make the structure more stable, hence, improve the thermal stability and cyclability. However, previous work only investigated doped Ti with small amount (<10%). In order to further study the effect of the amount of Ti doping on the structure and performance and understand the role of Ti played in the cathode material of LN i_x Co_{1-x}O₂, we used so1-gel method to synthesize a series of LN i_h 75-x Co_{1 25} Ti_x O₂ (x = 0,

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0, 1, 0, 25). The structure, morphology, electrochemical performance and electric state of the materials at different x values both experimentally and by calculation using VASP were examined

2 Experimental

2.1 Syn thesis of Materials

Three types of L N $i_{0.75-x}$ Co_{0.25} Ti_xO₂ (x = 0, 0.1, 0.25) were prepared by sol-gel pretreatment using citric acid as a chelating agent and solid-phase formation A stoichiometric amount of lithium nitrate (LNO₃), nickel nitrate (N NO₃ · 6H₂O) and cobalt nitrate (CoNO₃ · 6H₂O) was dissolved in absolute ethyl alcohol and mixed with aqueous solution of citric acid And then a stoichiometric amount of tetrabutyl titanate (C₁₆H₃₆O₄ Ti) was added The resulting solution was stirred at 80 for more than 12 h to obtain a clear viscous gel The gelwas dried at an oven at 120

for 12 h $LNi_{0.75-x}Co_{0.25}Ti_xO_2$ (x = 0, 0.1, 0.25) were calcined at 725 for 2 h after precalcining the obtained precursor at 380 . During heating and cooling, the variation of the temperature was fixed at 1 /m in

2. 2 Characterization of Materials

The thermal analysis was carried out on a Netzsch STA 400 analyzer with 50 mL/min of flowing air and a heating rate of 10 /m in the temperature range of 323 ~ 1273 K used precursors of dry gels The structures of the materials prepared were characterized by powder X-ray diffraction (XRD), using Philips Panalytical X 'pert diffractometer with CuK radiation operated at 40 kV and 30 mA. The data were collected in the 2 range of 10 °~ 90 ° using a step size of 0. 0167 ° and a counting time of 10 s per step. The morphologies of the materials were obtained by using a scanning electron microscope (LEO 1530 field emission SEM, Oxford Instruments) which was operated at 15 kV. The samples were coated by Au before the SEM observation

Electrochemical characterization was performed with coin-type cells The cathode was prepared by mixing an 85 5 10 (w/w) ratio of active material, carbon black, and polyvinylidene fluoride binder, respectively, in *N*-methyl pyrrolidinone. Lithium metal was used as an anode and a polypropylene separator was used to separate the anode and the cathode; 1. 0 mol/L LiPF₆ dissolved in a 1–1 mixture of ethylene carbonate / diethyl carbonate was used as an electrolyte. The charge and discharge cycles were carried out at a 0–1 C-rate over a potential range between 2. 7 and 4. 4V.

2.3 Calculation

All calculations were performed in the generalized gradient approximation (GGA) to density functional theory as implemented in the Vienna Ab Initio Simulation Package (VASP). A plane-wave basis set with a kinetic-energy cutoff of 500 eV was used The reciprocal space sampling was done with a 4 \times 4 \times 4 kpoint grid for structures containing 4 grids

3 **Results and D iscussion**

Figure 1 shows the TGA-DTG curves of the drygel precursors prepared for the precalcination and calcination processes The symbols C_1 , C_2 and C_3 in the figure represent LN $i_{0.75}$ Co_{0.25} O₂, LN $i_{0.65}$ Co_{0.25} T $i_{0.10}$ and LN $i_{0.5}$ Co_{0.25} T $i_{0.25}$ O₂, respectively Below 200

, all three types of the materials had small peaks, and this weight loss of the precursors could be due to the desorption of superficial and structural water In the temperature range of $200 \sim 300$, a peak occurred due to the decomposition of nitrate and/or the dehydration of metal citrate to aconitate^[14]. Then a shap drop in weight appeared between 350 and , which is attributed to the combustion of acon-400 itate and its complex There were small changes in the temperature range of $400 \sim 600$. When the temperature was higher than 700, a small peak appeared and could be related to the decomposition of carbonate It is evident that temperature of phase transformation for C_1 was above 650 , while the temperature for the phase transformation of C_2 and C_3 were about 600 and 500 , respectively. Thereby, the thermal stability can be improved by increasing the amount of Ti doping

Based on the TGA /DTA results given in Fig 1, eight characteristic temperatures, namely, 350,



Fig 1 TG-DTG curves of the dry-gel precursor of three different amounts of Ti doped materials

380 , 500 , 550 , 400 , 450 , 600 650 and 725 , were selected and a series of XRD measurements at each temperature kept for 2 h was made ex-situ Figure 2 shows the XRD patters of C_1 , C_2 , and C_3 at eight characteristic temperatures raised consequently from 350 to 725 The precursors were first heated up to each of the selected temperatures and kept constant in the oven for 2 h, and then were analyzed by XRD at room temperature. The XRD patterns of C_1 , C_2 and C_3 kept at 725 are those of finished product It can be seen from Fig 2a



Fig 2 XRD patters of three different amount of Ti doped materials on synthesized at for 2 hours

that C_1 at 725 had the typically hexagonal structure with the space group of R-3m. For hexagonal structure, the relative intensities of XRD lines $I_{(003)} / I_{(104)}$ or $I_{(101)} / I_{(006, 102)}$ are considered to be indicators of the ordering of lithium and other transition metal cations (N i and/or Co)^[15], and the ordering of the structure can also be evaluated from the XRD spectra with the degree of either (108)/(110) or (006)/(102) peak splitting All the peaks belong to those of hexagonal structure and no other phases were presented The value of $I_{(003)} / I_{(104)}$ is 1. 0027 and (108)/(110) and (006) / (102) are splited, indicating an ordered and layered structure. As evident in Fig 2b, C_2 which calcined at 725 also had hexagonal structure with $I_{(003)} / I_{(104)}$ being 1. 027 and the degree of the peak spliting of (006) / (012) and (018) / (110)was better than that of C_1 . That is to say doped Ti can increase the ordering of lithium and other transition metal cations (Ni and/or Co) and can increase the ordering of structure. However, for C_3 at 725 (Fig 2c), two phases, cubic structure and hexagonal structure co-existed

From the whole XRD patterns of C_1 , it can be seen the courses of phase transformation from cubic structure to hexagon structure. In the temperature range of 350 ~ 600 , the XRD patters were similar, and all the peaks were characteristic of the cubic structure, the development of (003) and (104)



Fig 3 Micrographs of three types of cathode materials synthesized at 725 for 2 hours

peaks with the increase of temperature became apparent. The phase transformation took place above 650 and the hexagonal structure replaced the cubic structure. With the increase of temperature, the intensity of characteristic peaks became stronger and well-defined, the degree of the peak splitting in (006) / (012) and (018) / (110) became significant For C_2 the hexagonal structure was obtained at above 600

, both agreed well with the TGA results In the case of C_3 , two phases were presented through the whole temperature rang The results in Fig 2 suggested the better defined hexagonal and layered structure can be obtained at x = 0. 1. While two phases are presented with x = 0.25.



Fig 4 Cyclic discharge curves of the first two cycles of three different amounts of Ti doped materials

		Material		
		C_1	C_2	C_3
First	Charge capacity /mAh \cdot g ⁻¹	193	201	170
	Discharge capacity /mAh \cdot g ⁻¹	172	190	140
	Loss(%)	12 0	6	18
Second	Charge capacity /mAh \cdot g ⁻¹	178	181	123
	Discharge capacity /mAh · g ⁻¹	166	179	115

Tab 1 Summary of cycling performance from Fig 4





Fig 5 DOS of three different amount of Ti doped materials a)

a) \sim c) lithiation, d) \sim f) delithialization

-20 -15 -10

0 -5 0 Energy/eV

Figure 3 is the SEM micrographs of C_1 , C_2 , and C_3 which calcined at 725 for 2 h after precalcining

the obtained precursor at 380 . It can be seen that the shape of the particles for C_1 and C_2 was regular

10 15

5

with clear edges (Fig 3a and 3b), indicating good crystal structure. The particle sizes for C_2 are smaller and more uniform than those of C_1 . The morphology revealed C_3 indistinct edges of particles and rough surfaces, hence poor structure. The SEM observation is consistent with the XRD results discussed before.

Figure 4 compares the first charge and discharge curves of these three materials For C_1 and C_2 , they both had plateaus at about 3. 6 V, but for C_3 , there was no obvious plateaus The charge/discharge capalities and there losses are summarized in Table 1. It can be seen that for C_1 the first charge capacity was 193 mAh/g and the first cycle loss was 12%. The first charge capacity for C_2 reached 201 mAh/g while which is higher than that of C_1 , the first cycle loss maintained at 6%, suggesting that the cyclability of C_2 is better than that of C_1 . The first charge capacity for C_3 was 23mAh/g lower and the first cycle loss was 6% higher than those of C_1 , suggesting poor electrochem ical performance

The density of states for L i_vN i_{1,75-x}Co_{0,25} Ti_xO₂ (x =0, 0.1, 0.25) when y = 1 and y = 0 were calculated and the results are provided in Fig 5. When the lithium ion deintercalation occerred, the energy of Fermi $(E_{\rm F})$ was lower and the Ni-e_g orbit moved to a higher lever of $E_{\rm F}$, indicating the degree of polarization on NiO and Co-O increased, hence, the valence of N i ion changed from +Ni() to +Ni(). The doping with small amount of Ti (x = 0, 1) increased the DOS of Ni- e_g (Fig 5b), implying that the valences of Ni are + Ni() and + Ni() with less N_{1}^{3+} and more N_{1}^{2+} . The presence of N_{1}^{2+} made the structure more stable. However, when x = 0.25, the DOS of Ni- e_g decreased, indicating that the high doping with Ti is detrimental to the structure of material The calculated results agreed well with the experimental results

4 Conclusion

Three types of materials L N $i_{0.75-x}$ Co_{0.25} T i_x O₂ (x

=0, 0. 1, 0. 25) were prepared by sol-gel method and analyzed by TG, XRD and SEM for the structure and morphology information. The electrochemical performance electron structure were also examined. The results showed that the thermal stability can be improved by doping Ti Better defined hexagonal structure can be obtained at x = 0. 1 with good electrochemical performance. When x = 0. 25, two phase were presented, which resulted in poor electrochemical performance. The calculation in the density of states revealed the change of Ni ion valence during delithializating

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钛掺杂对 L Ni 75 Co 25 O2结构与性能的影响

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摘要: 应用溶胶 凝胶法合成 LNi_(0.75-x) Co_{0.25} Ti_xO₂ (x = 0, 0.1, 0.25)系列正极材料,其结构、形貌、粒度、电 化学性能由 TG、XRD、SEM和电池充放电测试表征研究表明,材料的电化学性能与钛掺杂量密切相关.在钴 含量不变的情况下,随着 Ti含量 (x)的增加,材料由六方层状结构逐渐向立方结构转变,x = 0.25时,出现了 立方相与六方相共存.根据实验和理论计算结果简要讨论了钛掺杂对正极材料 LNi_{0.75} Co_{0.25}O₂结构和电化学 性能的影响.

关键词: 锂离子电池;正极材料;钛掺杂;电化学性能