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34 Claims, 5 Drawing Sheets
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Figure 1

Figure 2
Figure 3

Figure 4
**Figure 5**

- **Unmodified**
- **Al₂O₃ modified**
- **CeO₂ modified**
- **F modified**
- **ZrO₂ modified**

**Figure 6**

- **Unmodified**
- **Al₂O₃ modified**
- **CeO₂ modified**
- **F modified**
- **ZnO modified**
- **SiO₂ modified**
Figure 7
Figure 8

Figure 9
The present invention relates in general to the field of lithium-ion cathode materials, more particularly to the use of layered lithium transition metal oxide cathode materials having the O3 type structure of LiCoO2.

BACKGROUND OF THE INVENTION

Without limiting the scope of the invention, its background is described in connection with layered lithium metal oxide electrodes and layered lithium metal ion battery electrode materials.

The miniaturization of portable electronic devices has created a necessity for smaller, lighter, more durable batteries as their power source. To address this need, a lithium ion battery has been developed having a high capacity, small size, and light weight. Generally, the lithium ion battery has electrodes that can occlude and release lithium ions without requiring electrodeposition of metal lithium. The lithium ion can migrate from the anode into the electrolyte and occlude (by intercalation) from the electrolyte at the cathode. The anode is commonly made of a carbonaceous material and the cathode is made of a lithium transition metal oxide.

Although, cathode materials of lithium metal oxides have a theoretical capacity of around 280 mAh/g, the full capacity of these materials cannot be achieved in practice, and only about 140 mAh/g can be used. Furthermore, overcharging results in lithium removal that degrades the cyclability of these materials. In addition, lithium cobalt oxide and lithium nickel oxide also may undergo a decomposition reaction on overcharge. For example, layered LiNNiO2 transforms to the spinel LiNi2O4 on heating to above 200°C; at about 245°C, the delithiated material also experiences significant oxygen generation and heat liberation due to decomposition.

SUMMARY OF THE INVENTION

The present inventors recognized that only 50% of the theoretical capacity of the currently used LiCoO2 is being used in practical lithium ion cells due to the chemical and structural instabilities at deep charge and that cobalt is relatively expensive and toxic. The inventors recognize the need for a cathode material that exhibits high capacity with lower cost and better safety features than the currently used LiCoO2 cathode.

The present invention provides layered oxide cathode compositions as well as surface modification procedures for reducing the irreversible capacity loss and increasing the discharge capacity to about the theoretical value involving the reversible extraction of close to one lithium ion per formula unit. The surface modification suppresses undesired reactions of the cathode surface with the electrolyte, alters the solid-electrolyte interface (SEI) layer, and thereby lowers the irreversible capacity loss.

Layered (1-x)Li[(Li1/3Mn2/3)xLi0.45Ni0.45Co0.10]O2 cathode materials belong to a solid solution series between Li[(Li1/3Mn2/3)0.45Li0.55]O2 and Li[(Li1/3Mn2/3)0.45Ni0.45Co0.10]O2 with the value of x between 0 and 1 and the value of y between 0 and 0.5 have been synthesized and characterized by charge-discharge measurements in lithium cells before and after modifying their bulk and surface with cationic and anionic substitutions with metals and nonmetal ions like Al3+ and F-. The surface modified cathodes show significantly lower irreversible capacity loss (ICL) and higher discharge capacity with excellent cyclability compared to the unmodified counterparts. For example, Li[(Li1/3Mn2/3)0.45Ni0.45Co0.10]O2 with x=0.4 and y=0/3 shows a remarkably high capacity of 285 mAh/g with an irreversible capacity loss of 41 mAh/g and good rate capability. This capacity value is much higher than that achieved with the previously known Li[(Li1/3+xMn2/3)xNi0.45Co0.10]O2 with x=0.5 and y=0 after similar surface modification (<255 mAh/g).

The present invention provides a method of making a (1-x)

Layered (1-x)Li[(Li1/3Mn2/3)xLi0.45Ni0.45Co0.10]O2 cathode composition by mixing two or more co-precipitated transition metal hydroxides with a lithium hydroxide and firing the two or more co-precipitated transition metal hydroxides-lithium hydroxide mixture to form a (1-x)Li[(Li1/3Mn2/3)xLi0.45Ni0.45Co0.10]O2 material having an O3 crystal structure. The two or more co-precipitated transition metal hydroxides are formed by adding one or more transition metal acetate solutions to a basic solution.

The present invention also provides a cathode composition including (1-x)Li[(Li1/3Mn2/3)xLi0.45Ni0.45Co0.10]O2 having an O3 crystal structure and in some instances, when incorporated in a lithium-ion battery, the capacity is between about 250 and 300 mAh/g and an irreversible capacity loss of between about 30 and 50 mAh/g. The (1-x)Li[(Li1/3Mn2/3)xLi0.45Ni0.45Co0.10]O2 cathode composition has an x value between about 0.0 and about 1.0 and a y value between 0 and 0.5, e.g., when x=0.5 and y=0, the resulting cathode composition is Li[(Li1/3Mn2/3)0.45Ni0.45]O2 and when x=0.5 and y=0, the resulting cathode composition is Li[(Li1/3Mn2/3)0.45Ni0.45]O2. For example, the present invention includes a method of making a lithium cathode by mixing a (1-x)Li[(Li1/3Mn2/3)xLi0.45Ni0.45Co0.10]O2 cathode composition with a conductive diluent and a binder and forming the mixture into a cathode shape. The cathode shape will depend on the final application or use of the cathode, e.g., generally cylindrical or generally disk shaped.

The binder includes polytetrafluoroethylene or polyvinylidene fluoride. Suitable conductive diluent materials for this purpose include acetylene black, carbon black and graphite or a metallic powder such as powdered nickel, aluminum, titanium or stainless steel. For example, the binder may be polytetrafluoroethylene at about 1 to about
10 weight percent of the cathode mixture, the conductive diluent comprises acetylene black at about 5 to about 25 weight percent of the cathode mixture, and the Li[Li1/3Mn2/3O2] composition may be about 10 to about 95 weight percent of the cathode mixture.

The present invention provides a method of modifying a layered oxide with a metal oxide by dispersing a (1-x)Li

$\text{Li}_2\text{Mn}_2\text{O}_4$ composition in a metal alkoxide solution and then heating to obtain a surface modified layered oxide. The metal salt precursor solution includes Zr, Si, Ti or a combination thereof. A method of modifying a layered oxide cathodes by combining a layered oxide cathodes with NH$_4$HF$_2$ at a low temperature of between 300 and 500° C. is also provided.

In addition, the present invention provides a method of dispersing a (1-x)Li

$\text{Li}_2\text{Mn}_2\text{O}_4$ composition in a metal alkoxide solution. Water is added to the metal alkoxide solution, wherein the alkoxide solution is hydrolyzed and metal hydroxides are generated. The layered oxide containing the metal hydroxide is heated to form a surface modified layered oxide. The metal salt precursor solution includes Al, Ce, Zn or a combination thereof, and the metal oxide comprises Al$_2$O$_3$, CeO$_2$, ZnO or combinations thereof.

A method of modifying an anion substituted cathode composition. The mixture is fired to form a bulk modified anion substituted cathode composition. The anionic substitution compound may be lithium fluoride, lithium chloride, lithium sulfide or a combination thereof depending on the substitution desired by the skilled artisan.

The present invention also includes the composition made by this process.

BRIEF DESCRIPTION OF THE DRAWINGS

For a more complete understanding of the features and advantages of the present invention, reference is now made to the detailed description of the invention along with the accompanying figures and in which:

FIG. 1 is an XRD pattern of samples having the O3 type structure of LiCoO$_2$; FIGS. 2a and 2b are TEM images of Al$_2$O$_3$ modified Li[Li$_{0.2}$Mn$_{0.54}$Ni$_{0.13}$Co$_{0.13}$]O$_2$ at different magnifications; FIG. 3 is a graph comparing the first charge-discharge profiles of samples before and after surface modification with Al$_2$O$_3$; FIG. 4 is a graph comparing the cyclability data of samples before and after surface modification with Al$_2$O$_3$; FIG. 5 is a graph comparing the rate capabilities of Li[Li$_{0.2}$Mn$_{0.54}$Ni$_{0.13}$Co$_{0.13}$]O$_2$ before and after surface modification of Al$_2$O$_3$; FIGS. 6a, 6b and 6c are graphs comparing the first charge-discharge profiles of unmodified and surface modified samples; FIGS. 7a, 7b, and 7c are graphs comparing the cyclabilities of the unmodified and surface modified samples; FIGS. 8a and 8b are graphs comparing the first charge-discharge profiles of samples; and FIG. 9 is a graph comparing the cyclabilities of the Li[Li$_{0.17}$Al$_{0.3}$Mn$_{0.58}$Ni$_{0.25}$]O$_2$ and Li[Li$_{0.17}$Mn$_{0.58}$Ni$_{0.25}$]O$_2$ cathodes.

DETAILED DESCRIPTION OF THE INVENTION

While the making and using of various embodiments of the present invention are discussed in detail below, it should be appreciated that the present invention provides many applicable inventive concepts that can be embodied in a wide variety of specific contexts. The specific embodiments discussed herein are merely illustrative of specific ways to make and use the invention and do not delimit the scope of the invention.

To facilitate the understanding of this invention, a number of terms are defined below. Terms defined herein have meanings as commonly understood by a person of ordinary skill in the areas relevant to the present invention. Terms such as “a”, “an” and “the” are not intended to refer to only a singular entity, but include the general class of which a specific example may be used for illustration. The terminology herein is used to describe specific embodiments of the invention, but their usage does not delimit the invention, except as outlined in the claims.

As used herein the term “Ampere-hour (Ah)” refers to the units used in specifying the storage capacity of a battery. For example, a battery with 1 Ah capacity can supply a current of one ampere for one hour or 0.5 A for two hours, etc. 1 Ampere-hour (Ah) is the equivalent of 3600 coulombs of electrical charge. As used herein the term “basic” refers to an aqueous solution containing more OH" ions than H" ions. Generally, the aqueous solution has a pH greater than about 7, e.g., lithium hydroxide, sodium hydroxide, potassium hydroxide.
rubidium hydroxide, cesium hydroxide, magnesium hydroxide, calcium hydroxide, strontium hydroxide, barium hydroxide, and ammonium hydroxide.

As used herein the term “C Rate” refers to the charging or discharging rate of a cell or battery, expressed in terms of its total storage capacity in Ah or mAh. For example, a rate of 1 C means utilization of all of the stored energy in one hour; a 0.1 C means utilization of 10% of the energy in one hour and the full energy in 10 hours; and a 5 C means utilization of the full energy in 12 minutes.

As used herein the term metal oxides include precursors of the metal oxides such as nitrates, carbonates and acetates which can be converted to their corresponding metal oxides by heat treatment.

Surface modified layered Li[Li1/3Mn2/3]O2 cathodes belonging to a solid solution layer between layered Li[Li1/3,Mn2/3]O2 and Li[Ni1/3,Mn2/3]O2 exhibit a remarkably high reversible capacity of 285 mA/h with excellent cyclability and low irreversible capacity loss. This capacity value of 285 mA/h corresponds to a reversible extraction of 0.9 lithium ion per formula unit, and it is much higher than that found with any of the layered oxide compositions previously reported in the literature (e.g., <255 mA/h) at similar C rates. The surface modification with Al2O3 suppresses the reaction between the cathode surface and the electrolyte, and thereby decreases the irreversible capacity loss and increases the reversible discharge capacity. Further optimization of the layered oxide cathode compositions as well as surface modification procedures can increase the capacity to the theoretical value involving the reversible extraction of one lithium ion per formula unit.

Lithium ion batteries have become attractive for portable electronic devices such as cell phones and laptop computers due to their high energy density. However, only 50% of the theoretical capacity of the currently used LiCoO2 could be used in practical lithium ion cells (e.g., 140 mA/h) due to the chemical and structural instabilities at high charge with (1-x) < 0.5 in Li1-xCoO2. Also, cobalt is relatively expensive and toxic. These difficulties have generated enormous interest in alternative cathode hosts. In this regard, solid solutions between layered Li[Li1/3Mn2/3]O2 (commonly designated as Li2MnO3) and LiMNO3 (M=Mn, Ni, Co) have become appealing as some of them exhibit much higher capacity with lower cost and better safety features compared to the currently used LiCoO2 cathode. For example, layered oxide compositions belonging to the series (1-x)Li[Li1/3Mn2/3]O2.xLi[Mn0.5Ni0.5]O2 have been found to exhibit capacities as high as 250 mA/h on cycling them to 4.8 V.3-7

However, the discharge capacities of the (1-x)Li[Li1/3Mn2/3]O2.xLi[Mn0.5Ni0.5]O2 cathodes are often much higher than the theoretical capacity values expected based on the initial oxidation states of manganese and nickel, and the first charge profile is accompanied by an irreversible voltage plateau for oxidation involving beyond the formal oxidation states of Mn4+ and Ni4+. The voltage plateau has been attributed to an irreversible loss of oxygen from the lattice based on in-situ X-ray diffraction studies.1 The oxygen loss leads to a lowering of the oxidation states of Mn and Ni correspondingly at the end of first discharge, which facilitates a higher reversible capacity with good cyclability in the subsequent cycles as the system operates with the chemically more stable Mn3+/4+ and Ni3+/4+ couples compared to the Cu2+/3+ couple.1 More recently, an ion exchange of Li+ by H+ in addition to the loss of oxygen has been suggested from thermogravimetric analysis and mass spectrometry.12

In addition to the irreversible oxygen loss during the first charge, the (1-x)Li[Li1/3Mn2/3]O2.xLi[Mn0.5, Ni0.5]O2 compositions exhibit an undesirable, irreversible capacity loss (ICL) of 40-100 mA/h in the first cycle depending on the composition upon charging to 4.8 V.3-7 The high irreversible capacity loss could originate from a reaction of the cathode surface with the electrolyte, particularly with the high cutoff charge voltages of 4.8 V. One way to overcome this is to modify the cathode surface by coating with inert oxides and minimize the cathode-electrolyte interfacial reactions. For example, the surface modification (e.g., coating) of the LiCoO2 cathode with oxides such as Al2O3, ZrO2,14 MgO,15 and SnO2 have been shown to improve the cyclability to higher cutoff charge voltages, offering an increase in reversible capacity from 140 to about 180 mA/h. Similarly, the surface modification of Li[Li1/3Ni0.2]Mn0.6O2 with Al(OH)3 improves the rate capability.18

The present invention provides a new series of cathodes (1-x)Li[Li1/3Mn2/3]O2.xLi[Mn0.5, Ni0.5]O2 that belong to a solid solution layer between layered Li[Li1/3Mn2/3]O2 and Li[Mn0.5, Ni0.5]O2, their surface modification with Al2O3, ZrO2, CeO2, SiO2, ZnO, TiO2, and SnO2 and surface and bulk modification with anions like F- and cations like Al3+, and a comparison of their electrochemical performances before and after surface or bulk modification. For a comparison, the electrochemical performance of the cobalt-free Li[Li1/3Mn0.5Ni0.2]O2 that belongs to a previously known solid solution series (1-x)Li[Li1/3Mn2/3]O2.xLi[Mn0.5,Ni0.2]O2 is also presented before and after surface modification with Al2O3. The surface modified Li[Li0.3Mn0.5Ni0.2]O2 exhibits a high capacity of 285 mA/h with excellent cyclability and a low irreversible capacity loss of 40 mA/h.

Li[Li0.3Mn0.5Ni0.2]O2 (x=0.4 and y=1/6) and Li[Li0.3Mn0.4Ni0.2]O2 (x=0.7 and y=1/6) were synthesized by a co-precipitation method. Required amounts of the transition metal acetates were dissolved in deionized water and then added drop by drop into a 0.1 M KOH solution to form the coprecipitated hydroxides of Mn, Ni, and Co. After drying overnight at 100°C in an air-oven, the coprecipitated hydroxides were mixed with required amount of lithium hydroxide, fired in air at 900°C for 24 hours, and then quenched into liquid nitrogen. The surface modification of the synthesized layered oxides was carried out by dispersing the powders in an aluminum nitrate solution, followed by adding ammonium hydroxide to precipitate aluminum hydroxide and heating the products at about 300°C in air for 4 hours so that the Al2O3 content in the final product is 3 weight percent.14 All the samples were characterized by X-ray diffraction (XRD). Lithium content was determined by atomic absorption spectroscopy. Microstructural characterizations were carried out with a JEOL 2010F high-resolution transmission electron microscope (TEM).

For example, cathodes for evaluating the electrochemical performances were prepared by mixing 75 weight percent active material with 20 weight percent acetylene black and 5 weight percent PTFE binder, rolling the mixture into thin sheets of about 0.1 mm thick, and cutting into circular electrodes of 0.64 cm² area. CR2032 coin cells were then assembled with the cathode thus fabricated, lithium anode, and 1 M LiPF6 in ethylene carbonate (EC)/dimethyl carbonate (DEC) electrolyte.
be present as an amorphous AlOOH, considering the low parameter values of the layered oxide compositions.

TABLE 2 illustrates the electrochemical cell data collected at C/20 rate and 2.0-4.8 V of the layered oxides after surface modification with Al₂O₃. All the samples have the O₃ type structure of LiCoO₂ similar to that found previously for Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃Co₀.₁₃O₂. All the XRD patterns remain unchanged after modifying the surface with Al₂O₃ as seen in Fig. 1b, indicating that the surface modification does not cause any undue bulk structural changes to the cathodes. No extra reflections corresponding to Al₂O₃ are seen, possibly due to the small quantity (e.g., about 3 weight percent) and amorphous nature.

Table 2 compares the first charge and discharge capacities of the surface modified and unmodified Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃Co₀.₁₃O₂ before and after surface modification with Al₂O₃ at C/20 rate, and the capacity fades per cycle are given in Table 2. While both Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃Co₀.₁₃O₂, Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃Co₀.₁₃O₂ and Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃O₂ before and after surface modification with Al₂O₃. The surface modification offers the advantage of increasing the discharge capacity significantly in the newly developed cobalt-containing Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃Co₀.₁₃O₂ series compared to the cobalt-free Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃O₂ series. Moreover, the capacity of 285 mAh/g observed with Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃Co₀.₁₃O₂ is significantly higher than the values reported previously in the literature for any layered oxide composition (< about 255 mAh/g). It corresponds to a reversible extraction of 0.9 lithium ion per LiMO₂ formula unit (e.g., about 90% of theoretical capacity).

Figs. 4a, 4b, and 4c, are graphs that compare the cyclability data of all the three systems before and after surface modification with Al₂O₃ at C/20 rate, and the capacity fades per cycle are given in Table 2. While both Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃Co₀.₁₃O₂ and Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃Co₀.₁₃O₂ do not show any improvement. Additionally, there is little difference between the discharge and charge capacities after the first cycle as seen in Figs. 4a, 4b, and 4c, indicating good coulombic efficiency.

For example, the irreversible capacity loss decreases from 75 to 41 mAh/g and the discharge capacity increases from 253 to 285 mAh/g after surface modification in the case of Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃Co₀.₁₃O₂. Similarly, irreversible capacity loss decreases from 63 to 38 mAh/g and the discharge capacity increases from 227 to 250 mAh/g in the case of Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃Co₀.₁₃O₂. In contrast, while the irreversible capacity loss decreases significantly from 60 to 30 mAh/g, the capacity increases only slightly from 249 to 254 mAh/g in the case of the cobalt-free Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃O₂. The surface modification offers the advantage of increasing the discharge capacity significantly in the newly developed cobalt-containing Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃Co₀.₁₃O₂ series compared to the cobalt-free Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃O₂ series. Moreover, the capacity of 285 mAh/g observed with Li₄Li₀.₁₇Mn₀.₅₈Ni₀.₁₃Co₀.₁₃O₂ is significantly higher than the values reported previously in the literature for any layered oxide composition (< about 255 mAh/g). It corresponds to a reversible extraction of 0.9 lithium ion per LiMO₂ formula unit (e.g., about 90% of theoretical capacity).

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Al$_2$O$_3$ by dispersing the cathode powder in the corresponding metal salt precursor solution, followed by adding ammonium hydroxide to precipitate the metal hydroxides. The surface modifications with ZrO$_2$ and SiO$_2$ were carried out by dispersing the cathode powder in the corresponding metal alkoxide solution, followed by adding water to hydrolyze the alkoxide and generate the metal hydroxides. In both cases, the resulting product was heated at a temperature between 300°C and 700°C in air for 4 hours so that the metal oxide content in the final product is 3 weight percent. The surface modification with fluoride was carried out by heating the already formed layered oxide cathodes with NH$_4$HF$_2$ at a low temperature of between 300 and 500°C for about 4 hours. The first charge-discharge profiles are compared in Fig. 6.

All the surface modifications reduce the irreversible capacity loss; among them, Al$_2$O$_3$ modification is the most effective one. For example, the discharge capacity increased from 253 to 285 mAh/g and the irreversible capacity loss decreased from 75 to 41 mAh/g for the sample with x=0.4 and y=1/6. Figs. 6a, 6b, and 6c are graphs comparing the first charge-discharge profiles of unmodified and surface modified samples belonging to the series (1-x)Li$_{1/3}$Li$_{4/3}$Sn$_{0.25}$Co$_{0.13}$O$_{2}$-s at C/20 rate and 2.0-4.8V with (a) x=0.5 and y=0, (b) x=0.4 and y=1/6, and (c) x=0.4 and y=1/3.

The cyclabilities of the different modified samples are compared in the graphs of Figs. 7a, 7b, and 7c. It gives a comparison of the cyclabilities of the unmodified and surface modified samples belonging to the series (1-x)Li$_{1/3}$Li$_{4/3}$Sn$_{0.25}$Co$_{0.13}$O$_{2}$-s at C/20 rate and 2.0-4.8V for x=0.5 and y=0(Fig. 7a), x=0.5 and y=1/6(Fig. 7b), and x=0.4 and y=1/3(Fig. 7c). Samples modified with fluoride seem to have slightly better cycle performance, while the oxide modified samples have almost the same or worse cyclability than the unmodified sample. Again for the x=0.4 and y=1/6 sample modified with Al$_2$O$_3$, the discharge capacity is still as high as 268 mAh/g after 50 cycles.

The effects of partial substitution of Al$^{3+}$ or Li$^+$ in the transition metal layer and partial substitution of F$^-$ for O$^2-$ in Li$_{1/3}$Li$_{4/3}$Sn$_{0.25}$Co$_{0.13}$O$_{2}$-s (x=0.5 and y=0 sample) have also been investigated in detail. The fluorine substitution was accomplished by firing the coprecipitated transition metal hydroxides with a mixture of required amounts lithium fluoride and lithium hydroxide at 900°C for 24 hours and then quenching into liquid nitrogen. The aluminum substitution was accomplished by incorporating a required amount of Al$^{3+}$ ions into the mixed transition metal ion solution before forming the coprecipitated hydroxides, followed by firing the coprecipitated hydroxides with lithium hydroxide at 900°C for 24 hours and then quenching into liquid nitrogen. It was found that cationic and anionic substitutions influence the performance of the cathodes dramatically.

Figs. 7a and 7b are graphs comparing the first charge-discharge profiles of Li$_{1/3}$Li$_{4/3}$Al$_{0.17}$Ni$_{0.58}$Co$_{0.13}$O$_{2}$-s in Fig. 8a and Li$_{1/3}$Li$_{4/3}$Al$_{0.17}$Ni$_{0.58}$Co$_{0.13}$O$_{2}$-s in Fig. 8b. As shown in Fig. 8a, a small amount of substitution decreases the capacity. According to the cyclic voltammetry (CV) studies, Al and F substitutions strongly alter the chemical environment of Li$_{1/3}$Li$_{4/3}$Al$_{0.17}$Ni$_{0.58}$Co$_{0.13}$O$_{2}$ and make it more difficult for the oxygen ion to be removed from the lattice.

Fig. 9 is a graph that compares the cyclabilities of the aluminum- and fluorine-substituted samples, Li$_{1/3}$Li$_{4/3}$Al$_{0.17}$Ni$_{0.58}$Co$_{0.13}$O$_{2}$ and Li$_{1/3}$Li$_{4/3}$Al$_{0.17}$Ni$_{0.58}$Co$_{0.13}$O$_{2}$-s. Although the substitutions lead to a decrease in capacity values, the fluorine substitution leads to a better cyclability. The decrease in capacity is due to a decrease in the ability of the samples to lose oxygen from the lattice during the first charge. It is interesting to note that an incorporation of a very small amount (<0.05%) of foreign ions like Al$^{3+}$ or F$^-$ results in a significant change in the ability to lose oxygen from the lattice on charging. Thus, cationic and anionic substitutions could be used effectively to control the irreversible loss of oxygen from the lattice and the electrochemical performance factors such as reversible capacity, cyclability, and irreversible capacity loss. It could also help to tune the chemical stability and safety. The surface modification to reduce the irreversible capacity loss could also be pursued with other materials like carbon, metal, and other oxides like SnO$_2$, CoO, NiO, and CuO. Combining the ion substitution and the surface modification may produce materials that have less oxygen loss, lower IRC, and high capacity.
variation of error for the device, the method being employed to
to determine the value, or the variation that exists among the
study subjects.
As used in this specification and claim(s), the words “comprising” (and any form of comprising, such as “comprise” and “comprises”), “having” (and any form of having, such as “have” and “has”), “including” (and any form of including, such as “includes” and “include”) or “containing” (and any form of containing, such as “contains” and “contain”) are inclusive or open-ended and do not exclude additional, unre-
cited elements or method steps.
The term “or combinations thereof” as used herein refers to all permutations and combinations of the listed items preceding the term. For example, “A, B, C, or combinations thereof” is intended to include at least one of: A, B, C, AB, AC, BC, or ABC, and if order is important in a particular context, also BA, CA, CB, CBA, BCA, ACB, BAC, or CAB. Continuing with this example, expressly included are combinations that contain repeats of one or more item or term, such as BB, AAA, MB, BABC, AABCCC, CBBAAA, CABAABB, and so forth. The skilled artisan will understand that typically there is no limit on the number of items or terms in any combination, unless otherwise apparent from the context.

All of the compositions and/or methods disclosed and claimed herein can be made and executed without undue experimentation in light of the present disclosure. While the compositions and methods of this invention have been described in terms of preferred embodiments, it will be apparent to those of skill in the art that variations may be applied to the compositions and/or methods and in the steps or in the sequence of steps of the method described herein without departing from the concept, spirit and scope of the invention. All such similar substitutes and modifications apparent to those skilled in the art are deemed to be within the spirit, scope and concept of the invention as defined by the appended claims.

REFERENCES


What is claimed is:
1. A method, comprising: adding two or more transition metal acetate solutions to a basic solution to form two or more co-precipitated transition metal hydroxides comprising two or more hydroxides of Mn, Ni or Co;
mixing the two or more co-precipitated transition metal hydroxides with lithium hydroxide; and
heating the two or more co-precipitated transition metal hydroxides-lithium hydroxide mixture to form a (1-x)Li [Li1/3Mn2/3]O2.xLi[Mn05-yNi05-yCo2y]O2 material having an O3 crystal structure, where x is between 0.0 and 1.0 and y is from 0.0 to less than 0.5.
2. The method of claim 1, wherein the basic solution is selected from the group consisting of lithium hydroxide, sodium hydroxide, potassium hydroxide, rubidium hydroxide, cesium hydroxide, magnesium hydroxide, calcium hydroxide, strontium hydroxide, barium hydroxide, ammonium hydroxide and combinations thereof.
3. The method of claim 1, further comprising drying the two or more co-precipitated transition metal hydroxides at 90-120°C. for between 12 and 36 hours.
4. The method of claim 1, wherein the heating occurs at between 800-1000°C. for 12 to 36 hours.
5. The method of claim 2, wherein the (1-x)Li [Li1/3Mn2/3]O2.xLi[Mn05-yNi05-yCo2y]O2 material is selected from the group consisting of Li1/3Mn2/3O2, Li1/3Mn2/3O2.4Li0.5Mn0.5Ni0.5O2, Li1/3Mn2/3O2.4Li0.5Mn0.5Ni0.5O2 and mixtures thereof.
6. The method of claim 2, further comprising incorporating the (1-x)Li [Li1/3Mn2/3]O2.xLi[Mn05-yNi05-yCo2y]O2 material into a lithium-ion battery to produce a capacity of between about 250 and 300 mAh/g and an irreversible capacity loss of between about 20 and 50 mAh/g.
7. The method of claim 1, wherein the (1-x)Li [Li1/3Mn2/3]O2.xLi[Mn05-yNi05-yCo2y]O2 material is selected from the group consisting of Li1/3Mn2/3O2, Li1/3Mn2/3O2.4Li0.5Mn0.5Ni0.5O2, Li1/3Mn2/3O2.4Li0.5Mn0.5Ni0.5O2 and mixtures thereof.
8. A cathode composition, comprising: (1-x)Li [Li1/3Mn2/3]O2.xLi[Mn05-yNi05-yCo2y]O2 having an O3 crystal structure, wherein x is between 0.0 and 1.0, y is between 0.0 and 0.5, and the (1-x)Li [Li1/3Mn2/3]O2.xLi[Mn05-yNi05-yCo2y]O2 has a capacity between about 250 and 300 mAh/g and an irreversible capacity loss of between about 20 and 50 mAh/g when incorporated in a lithium ion battery.
9. The cathode composition of claim 8, wherein the (1-x)Li [Li1/3Mn2/3]O2.xLi[Mn05-yNi05-yCo2y]O2 is incorporated into a lithium-ion battery.
10. The cathode composition of claim 8, wherein the (1-x)Li [Li1/3Mn2/3]O2.xLi[Mn05-yNi05-yCo2y]O2 is selected from the group consisting of Li1/3Mn2/3O2, Li1/3Mn2/3O2.4Li0.5Mn0.5Ni0.5O2 and Li1/3Mn2/3O2.4Li0.5Mn0.5Ni0.5O2 and mixtures thereof.
11. The method of claim 1, further comprising: 

mixing the Li(1-x)Li[Li1/3Mn2/3O2]xLi[Co1-xNi1-xMn1-yMno.5-yO2] material with a conductive diluent and a binder to provide a mixture; and 

forming the mixture into a general cathode shape to provide a cathode.

12. The method of claim 11, wherein the binder comprises powdered polytetrafluoroethylene or polyvinylidene fluoride, and the conductive diluent is selected from the group consisting of acetylene black, carbon black, graphite, nickel powder, aluminum powder, titanium powder, stainless steel powder and combinations thereof.

13. The method of claim 11, wherein the binder comprises powdered polytetrafluoroethylene at about 1 to about 10 weight percent of the mixture, the conductive diluent comprises acetylene black at about 5 to about 25 weight percent of the mixture, and the (1-x)Li[Li1/3Mn2/3O2]xLi[Co1-xNi1-xMn1-yMno.5-yNio.5-yCo1-y]O2 material comprises about 70 to about 95 weight percent of the mixture.

14. A method, comprising: 

dispersing a (1-x)Li[Li1/3Mn2/3O2]xLi[Co1-xNi1-xMn1-yMno.5-yNio.5-yCo1-y]O2 composition in a metal salt precursor solution to form a first combination, wherein x is between 0.0 and 1.0 and y is between 0.0 and 0.5; 

adding ammonium hydroxide to the first combination to precipitate a metal hydroxide, wherein a layered oxide containing the metal hydroxide is formed; and 

heating the layered oxide containing the metal hydroxide, wherein a surface modified layered oxide is formed having a capacity between about 250 and 300 mAh/g and an irreversible capacity loss of between about 20 and 50 mAh/g when incorporated in a lithium ion battery.

15. The method of claim 14, wherein the metal salt precursor solution comprises Al, Ce, Zn, Sn or combinations thereof.

16. The method of claim 14, wherein the layered oxide containing the metal oxide is heated at a temperature between about 300° C. and about 700° C. for between about 1 and 6 hours.

17. A method, comprising: 

co-precipitating a metal hydroxide mixture comprising hydroxides of two or more transition metal ions and one or more metal ions; 

forming a mixture comprising the co-precipitated metal hydroxide mixture and lithium hydroxide; and 

heating the mixture at a temperature between about 800° C. and about 1000° C. to form a cationic substituted bulk modified cathode composition of (1-x)Li[Li1/3Mn2/3O2]xLi[Co1-xNi1-xMn1-yMno.5-yNio.5-yCo1-y]O2, wherein x is between 0.0 and 1.0, and y is from 0.0 to less than 0.5.

18. The method of claim 17, wherein the one or more transition metal ions are selected from the group consisting of Mg, Ti, V, Cr, Fe, Cu, Zn, Zr, Nb, Mo, W, Ga, Ca, and Cd and combinations thereof.

19. The method of claim 17, further comprising: 

disposing the cationic substituted bulk modified cathode composition in a metal salt precursor solution to form a first combination; 

adding ammonium hydroxide to the first combination to precipitate a metal hydroxide, wherein a layered oxide containing the metal hydroxide is formed; and 

heating the layered oxide containing the metal hydroxide, wherein a surface modified layered oxide is formed.

20. A cathode composition formed by the method of claim 17.

21. The method of claim 17, further comprising an anionic substitution compound.

22. The method of claim 21, wherein the anionic substitution compound comprises lithium fluoride, lithium chloride, lithium sulfide or a combination thereof.

23. The method of claim 17, further comprising forming the cathode composition into a cathode.

24. The method of claim 19, wherein the metal salt precursor solution comprises a member selected from the group consisting of selected from Al, Ce, Zn, Zr, Si, Ti, Sn and combinations thereof.

25. The method of claim 24, wherein the surface modified layered oxide has a capacity between about 250 and 300 mAh/g and an irreversible capacity loss of between about 20 and 50 mAh/g when incorporated in a lithium ion battery.

26. The method of claim 24, wherein the layered oxide containing the metal oxide is heated at a temperature between about 300° C. and about 700° C. for between about 1 and 6 hours.

27. The method of claim 24, wherein the metal hydroxide is selected from the group consisting of aluminum hydroxide, cerium hydroxide, zirconium hydroxide, silicon hydroxide, titanium hydroxide and zinc hydroxide.

28. The method of claim 24, wherein the metal hydroxide is aluminum hydroxide.

29. The method of claim 1, further comprising: 

co-precipitating a metal hydroxide mixture comprising hydroxides of two or more transition metal ions and one or more metal ions; 

forming a mixture comprising the co-precipitated metal hydroxide mixture and lithium hydroxide; and 

heating the mixture at a temperature between about 300° C. and about 700° C. to form a cationic substituted bulk modified cathode composition of (1-x)Li[Li1/3Mn2/3O2]xLi[Co1-xNi1-xMn1-yMno.5-yNio.5-yCo1-y]O2, wherein x is between 0.0 and 1.0, and y is from 0.0 to less than 0.5.

30. The method of claim 29, wherein the surface modified layered oxide has a capacity between about 250 and 300 mAh/g and an irreversible capacity loss of between about 20 and 50 mAh/g when incorporated in a lithium ion battery.

31. The method of claim 29, wherein the layered oxide containing the metal oxide is heated at a temperature between about 300° C. and about 700° C. for between about 1 and 6 hours.

32. The method of claim 29, wherein the metal salt precursor solution comprises a member selected from the group consisting of selected from Al, Ce, Zn, Zr, Si, Ti, Sn and combinations thereof.

33. The method of claim 29, wherein the metal hydroxide is selected from the group consisting of aluminum hydroxide, cerium hydroxide, zirconium hydroxide, silicon hydroxide, titanium hydroxide and zinc hydroxide.

34. The method of claim 29, wherein the metal hydroxide is aluminum hydroxide.

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