Date de dépôt PCT/PCT Filing Date: 1996/11/22
Date publication PCT/PCT Publication Date: 1997/05/29
Date de délivrance/issue Date: 2005/04/12
Entrée phase nationale/National Entry: 1997/07/08
N° demande PCT/PCT Application No.: JP 1996/003436
N° publication PCT/PCT Publication No.: 1997/019023
1996/01/18 (025976/1996) JP;
1996/04/12 (115428/1996) JP;
1996/05/17 (148147/1996) JP;
1996/05/21 (150127/1996) JP;
1996/06/20 (181587/1996) JP

Titre : OXYDE COMPOSITE LITHIUM-NICKEL, SON PROCEDE DE PREPARATION, ET MATERIAU ACTIF POSITIF DESTINE A UNE BATTERIE SECONDAIRE

Title: A LITHIUM NICKEL COMPLEX OXIDE, A PROCESS FOR PREPARING THE SAME AND A POSITIVE ELECTRODE ACTIVE MATERIAL FOR A SECONDARY BATTERY

Abrégé/Abstract:
A process for preparing a lithium nickel complex oxide having the formula (I) \( \text{Li}_{x+y} \text{Ni}_{x} \text{M}_{y} \text{O}_{z} \) (I) wherein \( M \) represents a metal from Al, Fe, Co, Mn or Mg, \( x=x_{1}+x_{2} \), and (i) when \( M \) is Al or Fe, then \( 0 < x \leq 2.0, x_{1} < 0 \) and \( x_{2}=x \); (ii) when \( M \) is Co or Mn, then \( 0 < x \leq 0.5, x_{1}=0 \) and \( x_{2}=x \); (iii) when \( M \) is Mg, then \( 0 < x < 0.2, 0 < x_{1} < 0.2 \) and \( 0 < x_{2} < 0.2 \); and \( y \) represents \( 0.9 \leq y < 1.3 \), comprising: (a) reacting a basic metal salt of the formula (II) \( \text{Ni}_{x} \text{M}_{y} \text{O}_{z} \text{M}^{n+} (\text{OH})_{2n-2z} (\text{A}^{m+})_{z} \text{mH}_{2} \text{O} \) (II) wherein \( M \) represents Al, Fe, Co, Mn or Mg, \( p \) is \( 2 \leq p \leq 3 \), \( \text{A}^{m+} \) is an anion having the valence \( n \), and \( x, z \) and \( m \) are positive numbers satisfying the ranges: \( 0 < x \leq 0.2; 0.03 \leq z \leq 0.3 \); and \( 0 \leq m < 2 \); with a water soluble lithium compound in an aqueous medium in a molar ratio Li/(Ni+M) of 0.9-1.3 to obtain a slurry, (b) spray drying the slurry; and (c) heating the dried material at a temperature of about 600 °C - 900 °C for at least 4 hours in an oxidative atmosphere. The crystals obtained by the process are sufficiently developed and highly purified for use as a positive electrode material for a secondary battery in which the decreased rate of discharge capacity at 100th cycle based on the initial discharge capacity is 10% or less.
**ABSTRACT**

A process for preparing a lithium nickel complex oxide having the formula (I)

\[ \text{Li}_y \text{Ni}_{1-x_1}\text{M}_{x_2}\text{O}_2 \]  (I)

wherein \( M \) represents a metal from Al, Fe, Co, Mn or Mg, \( x = x_1 + x_2 \), and

(i) when \( M \) is Al or Fe, then \( 0 < x \leq 2.0 \), \( x_1 = 0 \) and \( x_2 = x \);  
(ii) when \( M \) is Co or Mn, then \( 0 < x \leq 0.5 \), \( x_1 = 0 \) and \( x_2 = x \);  
(iii) when \( M \) is Mg, then \( 0 < x < 0.2 \), \( 0 < x_1 \leq 0.2 \) and \( 0 < x_2 \leq 0.2 \); and \( y \) represents 0.9 \( \leq y < 1.3 \), comprising:

(a) reacting a basic metal salt of the formula (II)

\[ \text{Ni}_{1-x}\text{M}^p\text{O}(\text{OH})_{2-n} (\text{A}^{n-})_{(z+1)} (\text{H}_2\text{O})_m \]  (II)

wherein \( M \) represents Al, Fe, Co, Mn or Mg, \( p \) is \( 2 \leq p \leq 3 \), \( \text{A}^{n-} \) is an anion having the valence \( n \), and \( x \), \( z \) and \( m \) are positive numbers satisfying the ranges: \( 0 < x \leq 0.2 \); \( 0.03 \leq z \leq 0.3 \); and \( 0 \leq m < 2 \); with a water soluble lithium compound in an aqueous medium in a molar ratio \( \text{Li}/(\text{Ni}+M) \) of 0.9-1.3 to obtain a slurry,

(b) spray drying the slurry; and

(c) heating the dried material at a temperature of about 600 °C - 900 °C for at least 4 hours in an oxidative atmosphere. The crystals obtained by the process are sufficiently developed and highly purified for use as a positive electrode material for a secondary battery in which the decreased rate of discharge capacity at 100th cycle based on the initial discharge capacity is 10% or less.
SPECIFICATION

A LITHIUM NICKEL COMPLEX OXIDE, A PROCESS FOR PREPARING THE SAME AND A POSITIVE ELECTRODE ACTIVE MATERIAL FOR A SECONDARY BATTERY

Technical Field

The present invention relates to a novel lithium nickel complex oxide having a high charge and discharge capacity and an excellent stability in retaining the capacity so that the degree of decrease in the capacity is little even with increasing the number of cycle, to a process for preparing the same, and to a positive electrode active material for a secondary battery.

Background Art

As small size electronic appliances have been rendered portable in recent years, there has been increased a demand for a lithium secondary battery having a small size, a light weight and a high energy density in place of nickel/cadmium battery.

As active materials of positive electrode for such a lithium secondary battery, there are known LiCoO₂ and LiNiO₂ which are layered compounds capable of being intercalated and deintercalated with lithium. Of them, LiNiO₂ is being expected owing to its higher electric capacity than LiCoO₂.
Usually, LiNiO$_2$ has been prepared by mixing a lithium component (LiOH, Li$_2$CO$_3$, LiNO$_3$ etc.) with a nickel component (hydroxide, carbonate etc.) in a powdery form and reacting the mixture by the so-called dry process, and hence the heat at an elevated temperature for a long time was required. Especially, in the case of Ni, it is hard to convert divalent one into the trivalent one and therefore the heat at an elevated temperature for a long time was essential. Consequently, the crystal growth proceeds but some of lithium is evaporated off and NiO as a by-product is formed, thereby the lowering of the purity is caused.

To the contrary, the present inventors succeeded previously in the preparation of highly purified LiNiO$_2$ having a high crystallization degree by forming a uniform precursor of Li and Ni components using a wet process and heating it for a short time as disclosed in Japanese Patent Application No. 6-80895 (Japanese Patent Kokai NO. 8-130013).

As to LiNiO$_2$, however, when much Li was evaporated off (at the time of discharging), the structure tends to become unstable owing to the two dimension structure and therefore an essential problem that the cycle property of a lithium secondary battery is bad could not be overcome completely. Although the effect of improving the cycle characteristics was recognized to be achieved to some extent by using the technique of the Japanese Patent Application No. 6-80895, it was still insufficient in a long term cycle
characteristics of more than 100 cycles.

Under such circumstance, many attempts have been made to stabilize the structure by substituting a part of nickel for other component (third component). For example, active materials of positive electrode represented by Li$_{1-x}$Ni$_x$Co$_{1-x}$O$_2$ (wherein x is 0 < x ≤ 0.75 and y is y ≤ 1) where Co was doped as solid solution into LiNiO$_2$, and Li$_{1-x}$Ni$_x$Me$_x$O$_2$ (wherein Me represents any one of Ti, V, Mn and Fe, x is 0 < x ≤ 0.6 and y is 0.2 < y ≤ 1.3) where Ti, V, Mn or Fe was doped as solid solution into LiNiO$_2$ are disclosed in Japanese Patent Kokai Nos. 63-299056 and 5-283076 respectively.

However, the process doped third component as solid solution was carried out by a process to be said the dry process described above and hence it is difficult to dope third component as solid solution homogeneously. It involved the necessities of increasing an amount of third component, heating at an elevated temperature for a long time and conducting several times of pulverization steps inevitably. And consequently, Li was evaporated off and the by-product of NiO was formed thereby the lowering of the purity was caused and sufficient improvement in cycle property could not be achieved like LiNiO$_2$ described above. Also, since the dry process required a long term heat and pulverization steps it was bad in the production efficiency and uneconomical. Furthermore, as it takes a long time for the heating inevitably in these dry processes, it was
impossible to adjust crystal size to a desired, keeping a crystallization degree and purity at high level.

Under these circumstances, some attempts have been made to prepare spherical particles for increasing a packing density. For example, a technique is disclosed in Japanese Patent Kokai No. 7-105950 for preparing spherical LiNiO$_2$ particles having 5 $\mu$m ~ 50 $\mu$m using spherical Ni(OH)$_2$. This technique is one for obtaining spherical LiNiO$_2$ using spherical Ni(OH)$_2$ as raw material by a dry process for the purpose of increasing a packing density only. But, the primary particle size and the purity of LiNiO$_2$ was not taken into consideration. The result was not satisfactory. Another technique is disclosed in Japanese Patent Kokai No. 6-333562 for preparing spherical LiNiO$_2$ having 0.1 ~ 1.1 $\mu$m with a mist dry process. In this technique, the crystal size is too fine so that the crystals are passed through a sepalator when employed as a battery and thus it is not practical for battery use. Especially in case of LiNiO$_2$, when the primary particles are too fine there is caused a problem that the storage stability is poor owing to moisture absorption so that good and stable battery characteristics cannot be obtained.

Moreover, it is known that when the battery was once placed under an elevated temperature, for example in a car in the daytime, even if it was reverted to the normal temperature, the positive electrode active material is susceptible to deterioration to a great extent and discharge
performance becomes bad so that the battery performance is decreased greatly.

As the means to prevent the deterioration of the positive electrode active material, in considering the fact that finer the primary particle size, has greater the deterioration, how large is the primary particle size of the active material to be prepared has been noticed.

As a process for improving the storage stability or the discharge characteristics of the positive electrode active material at high temperature by making the primary particle size large, there have been reported with respect to LiCoO$_2$ series some attempts to make the primary particle size large by improving the heating condition [Japanese Patent Kokai No. 6-243897 (0.1 ~ 2.0 $\mu$m), Japanese Patent Kokai No. 6-325791 (0.01 ~ 5 $\mu$m) and Japanese Patent Kokai No. 7-14579 (0.01 ~ 5 $\mu$m)].

Also, the primary particle having an average size of higher than 2 $\mu$m is prepared by adding Bi oxide to the raw material resource as proposed in Japanese Patent Kokai No. 8-55624. Thus it is easy for LiCoO$_2$ series to make the primary particle size large.

On the other hand, there has not been found yet any example of having made the primary particle of LiNiO$_2$ such large as seen in LiCoO$_2$ series. The reason is attributed to that the synthesis of LiNiO$_2$ or third component (M) added related compound Li$_x$Ni$_{1-x}$M$_x$O$_2$ required the heating at an elevated temperature for a long time owing to
their bad reactivity as described above, thereby Li is liable to be evaporated off. And consequently the crystal growth is hard to proceed and becomes imperfect crystals having much lattice defect. For this reason, the heating has been conducted at possible high temperature within the permissible temperature range and as a result only fine primary particle of less than 1μm could be obtained.

Disclosure of The Invention

The objects of the present invention are to provide (1) a novel lithium nickel complex oxide wherein the crystals have been developed sufficiently are highly purified and whoes stability of a high charge and discharge capacity is excellent, (2) a process for preparing said lithium nickel complex oxide whereby the size and shape may be set to desired one and (3) a positive electrode active material for a secondary battery containing as an effective component this complex oxide.

The present inventors have prepared for the first time by a wet process a novel lithium nickel complex oxide represented by the following formula (I):

\[ \text{Li}_{1-x_1}\text{Ni}_{1-x_2}\text{M}_x\text{O}_2 \]  

(wherin M is Co, Al, Fe, Mg or Mn, \( x = x_1 + x_2 \), \( x \) is 0 < \( x \leq 0.5 \), \( x_1 \) is 0\( \leq x_1 < 0.2 \), \( x_2 \) is 0 < \( x_2 \leq 0.2 \) and \( y \) is 0.9 \( \leq y \leq 1.3 \)) wherein the crystals have been developed sufficiently and are highly purified, and stability of a high charge and discharge capacity is
excellent.

The process of the present invention is characterized by that the size and shape of the primary particles for the objected lithium nickel complex oxide may be set to desired one. That is, the present invention relates to a lithium nickel complex oxide which is represented by the general formula (I):

\[ \text{Li}_{y-x_1}\text{Ni}_{1-x_2}M_x\text{O}_2 \] (I)

[wherein M is one selected from the group consisting of Al, Fe, Co, Mn and Mg, \( x = x_1 + x_2 \) (herein

(i) when M is Al or Fe, \( x, x_1 \) and \( x_2 \) represent respectively \( 0 < x \leq 0.2, x_1 = 0, x_2 = x \)

(ii) when M is Co or Mn, \( x, x_1 \) and \( x_2 \) represent respectively \( 0 < x \leq 0.5, x_1 = 0, x_2 = x \)

(iii) when M is Mg, \( x, x_1 \) and \( x_2 \) represent respectively \( 0 < x \leq 0.2, 0 < x_1 < 0.2, 0 < x_2 < 0.2 \), and \( y \) represents \( 0.9 \leq y \leq 1.3 \), and which is characterized in that X-ray diffraction pattern shows a ratio in the peak intensity of the face (003) to the face (104) i.e., \( (003)/(104) \) is 1.2 or higher and a ratio in the peak intensity of the face (006) to the face (101) i.e., \( (006)/(101) \) is 0.13 or lower, said face being defined by Miller indices hkl, BET specific surface area being in the range of \( 0.1 \sim 2 \text{ m}^2/\text{g} \), the proportion of Ni\(^{3+} \) to the total Ni being 99 % by weight or higher, an average secondary particle size \( D \) being in the range of \( 5 \sim 100 \mu \text{m} \) with 10 % of the particle size distribution being 0.5D or higher and 90 % 2D
or lower, with the surface of spherical secondary particle being uneven as observed with a scanning electron microscope (SEM) and the primary particle constituting the spherical secondary particle being uniform and distributed in the range of 0.2 ~ 3.0 µm in terms of long diameter with the average diameter of 0.3 ~ 2.0 µm as observed with a SEM.

Furthermore, the present invention relates to a lithium nickel complex oxide which is represented by the general formula (I):

\[ \text{Li}_{1-x_1}\text{Ni}_{1-x_2}\text{M}_x\text{O}_2 \] (I)

[wherein M is one selected from the group consisting of Al, Fe, Co, Mn and Mg, \( x = x_1 + x_2 \) (herein

(i) ) when M is Al or Fe, \( x, x_1 \) and \( x_2 \) represent respectively \( 0 < x \leq 0.2, x_1 = 0, x_2 = x \)

(ii) ) when M is Co or Mn, \( x, x_1 \) and \( x_2 \) represent respectively \( 0 < x \leq 0.5, x_1 = 0, x_2 = x \)

( iii ) when M is Mg, \( x, x_1 \) and \( x_2 \) represent respectively \( 0 < x \leq 0.2, 0 < x_1 < 0.2, 0 < x_2 < 0.2 \), and \( y \) represents \( 0.9 \leq y \leq 1.3 \), and which is characterized in that X-ray diffraction pattern shows a ratio in the peak intensity of the face (003) to the face (104) i.e., \( (003)/(104) \) is 1.2 or higher and a ratio in the peak intensity of the face (006) to the face (101) i.e., \( (006)/(101) \) is 0.13 or lower, said face being defined by Miller indices hkl, and an average long diameter of the primary particles is in the range of 1 ~ 10 µm as observed with a SEM.

The lithium nickel complex oxide represented by the
above general formula in the present invention may be more specifically classified into ones shown by the following general formulae (I a) \sim (I c):

A lithium nickel complex oxide represented by the formula (I a):

\[ \text{Li}_{x}\text{Ni}_{1-x}\text{M}_y\text{O}_2 \]  
(I a)

(wherein M represents Al or Fe, x represents \(0 < x \leq 0.2\) and y represents \(0.9 \leq y \leq 1.3\)) is one wherein a small amount of Al or Fe has been uniformly doped as the solid solution in the trivalent form thereby the structure is stabilized and cycle characteristics is enhanced whereas any battery characteristics cannot be achieved even when Al or Fe was converted into the corresponding LiMO\(_2\).

If x is less than 0.01, it is not preferable because the amount doped of Al or Fe as the solid solution is little so that the stabilization of the structure is insufficient and the cycle property becomes bad. Also, when x exceeds 0.2, the doping is insufficient and the impurities are formed so that discharge capacity decreases rapidly and the desired high capacity is lost which is an essential function for the positive electrode active material for a lithium secondary battery in the present invention.

A lithium nickel complex oxide represented by the formula (I b):

\[ \text{Li}_x\text{Ni}_{1-x}\text{M}_y\text{O}_2 \]  
(I b)

(wherein M represents Co or Mn, x represents \(0 < x \leq 0.5\) and y represents \(0.9 \leq y \leq 1.3\)) is one wherein Co or Mn
has been doped as the solid solution in relatively large amount thereby the structure is stabilized and a high electric capacity is retained although the battery characteristics can also be achieved when Co or Mn was converted into LiMO₂.

If x is less than 0.01, it is not preferable because the amount doped of Co or Mn is little and the stabilization of the structure is insufficient. When x exceeds 0.5, in the case of Co, the amount doped is much thereby not only discharge capacity is decreased but also it is not advantageous economically because of a large amount of an expensive cobalt to be intercalated. Also, in the case of Mn, inherently discharge capacity is small. When x exceeds 0.5, the desired high capacity is lost which is an essential function for the positive electrode active material for a lithium secondary battery in the present invention.

A lithium nickel complex oxide represented by the following formula (I c) where M is Mg in the formula (I)

\[ \text{Li}_{1-x_1} \text{Ni}_{1-x_2} \text{Mg}_x \text{O}_2 \]  
(I c)

(wherein \( x = x_1 + x_2 \), x represents \( 0 < x \leq 0.2 \), \( x_1 \) represents \( 0 < x_1 < 0.2 \), \( x_2 \) represents \( 0 < x_2 < 0.2 \) and \( y \) represents \( 0.9 \leq y \leq 1.3 \) is one wherein Mg which itself has no electric characteristics has been uniformly and partially doped as the solid solution into each of Li and Ni layers thereby the structure is stabilized and the cycle characteristics is enhanced.

If x is less than 0.01, it is not preferable because
the amount doped of Mg is little and the stabilization of the structure is insufficient. When \( x \) exceeds 0.2, the doping is insufficient and the impurities are formed so that discharge capacity decreases rapidly and the desired high capacity is lost which is an essential function of the positive electrode active material for a lithium secondary battery in the present invention.

The above-mentioned lithium nickel complex oxide of the present invention is composed of spherical particle to be formed by a wet-spray dry processes as described later and large primary particle size to be formed by the subsequent press process wherein the spray dried product (spherical particle product) is employed as raw material for heating as described later.

First, the lithium nickel complex oxide of the present invention will be illustrated with respect to spherical particle one formed by spray dry process. The spherical particle has characteristic properties as shown below.

The first feature is that the purity is high so that any impurities cannot be detected by a X-ray diffractometry. And, a ratio in peak intensity of the face (003) to the face (104) of Miller indices hkl of the X-ray diffraction pattern i.e., \((003)/(104)\) is 1.2 or higher and a ratio in the peak intensity of the face (006) to the face (101) i.e., \((006)/(101)\) is 0.13 or lower, and the proportion of \( \text{Ni}^{2+} \) to the total Ni is 99 % by weight or higher, thus it is highly purified. Usually the stability of the structure is
enhanced by the substitution of a part of Ni for other component (third component) but the purity decreases relative to the substitution amount.

In spite of the substitution of Al, Fe, Mg, Co or Mn having been done in the present invention, highly purified complex oxide may be obtained. That is, Al or Fe which is third component takes trivalence in the structure so that the structural unstability of Ni by reversibly de-intercalation of Li can be solved. In the case of Mg, it is intercalated into each part of lithium and nickel layers thereby the structural instability is avoidable. With respect to Co or Mn, each of them has been uniformly doped whereby the structural unstability of Ni by reversibly de-intercalation of Li can be solved. The most important feature of the present invention lies in that one of these third components is doped uniformly and in a very small amount not so as to exceed the required amount.

This has a close relation with a process to be said an aqueous solution process (wet process) as described later as well as size of the primary particles as described later. The complex oxide is essentially composed of the primary particles having a specific size thereby there can be obtained composition which has a high purity and a high crystallization degree and which is stable in quality.

The second feature lies in that the complex oxide is composed of the primary particles which are uniform particles which is distributed in the range of 0.2 ~ 3.0
μm in terms of the long diameter with the average diameter of 0.3 ~ 2.0 μm as observed with a SEM.

In general, the size of the primary particle is important for a layered compound represented by LiMO₄ in the light of the reversibly de-intercalation of Li. Finer the primary particles, better ionic conductivity in the inside of the solid and Li is more reversibly de-intercalatable with the outside.

On the other hand, in regard of the crystallization degree if the primary particle is 0.2 μm or less the crystal growth does not proceeds sufficiently and the purity becomes low inevitably. Also, if it is 0.2μm or less, then the storage stability is poor owing to moisture absorbability and so good battery characteristics cannot be achieved stably. Moreover, in considering that quality is stable it is desirable that the the size of the primary particle is uniform.

As a result of having studied ardently about the primary particle size from the above viewpoints, the present inventors have found that the primary particles in the lithium nickel complex oxide of the present invention have the desirable performance when they are uniform particles which are distributed in the range of 0.2~ 3.0 μm in terms of the long diameter with the average diameter of 0.3 ~ 2.0 μm, preferably 0.3 ~ 1.0 μm, as observed with a SEM.

Third feature is that the secondary particles are made
into spherical shape by wet-spray dry processes as described later and the average size of the secondary particles D lies in the range of 5 ~ 100 μm with 10 % of the particle size distribution being 0.5D or higher and 90 % 2D or lower, and the surface is in uneven state as can be seen under observation of a SEM.

Also, the particle ratio (a ratio of the long diameter to the short diameter) of the spherical secondary particles as observed with a SEM lies in the range of a maximum of 1.5 or less and an average of 1.2 or less with 90 % or more of them being distributed in 1.3 or less, indicating that they are uniform particles even when there was included some of slightly larger particle ratio than one defined above in the complex oxide prepared during pulverization after the heating has been conducted.

It is understood from such physical properties that not only they are suitable for the closest packing density but also they have an advantage when used as battery that the contact surface with each of an electrolyte and a conductive agent becomes large so it is easy to de-intercalate Li.

The size of the spherical secondary particles can be set to the range of from 5μm to 100 μm as desired. However, an average size of about 5 ~ 30 μm is desirable for use as the battery material from the viewpoint of processibility. Also, the BET specific surface area lies in the range of 0.1 ~ 2 m²/g. When used as the battery
material, since there is no increase in the viscosity of an electrolyte the lowering of conductivity does not be caused.

The lithium nickel complex oxide represented by the general formula (I) in the present invention may be prepared by the process described below.

The lithium nickel complex oxide of the present invention may be prepared by a process which comprises reacting a basic metal salt represented by the general formula (II)

\[ \text{Ni}_{1-x}M_x(OH)_{2-n}z(A^{n-})_{\left\{\frac{z+(p_x-2)x}{n}\right\}} \cdot mH_2O \quad (II) \]

(wherein M represents one selected from the group consisting of Al, Fe, Co, Mn and Mg, p is a valence number of M and represents \(2 \leq p \leq 3\), \(A^{n-}\) is an anion having a valence of n, and x, z and m are positive numbers respectively satisfying the range of \(0 < x \leq 0.2\), \(0.03 \leq z \leq 0.3\), \(0 \leq m < 2\)), with a water soluble lithium compound in an aqueous medium in a molar ratio of Li / (Ni + M) of 0.9 \~ 1.3 to obtain a slurry, spray drying the obtained slurry, and heating the resultant dry material at a temperature of about 600°C \~ 900°C for about 4 hours or more in an oxidative atmosphere.

As the water soluble lithium compound and the basic metal salt, there may be employed one each containing an anion which is evaporated off during the heating.

As the lithium compound, there may be selected one or more from among LiOH, LiNO₃, Li₂CO₃ and hydrates thereof.

In the basic metal salt represented by the above general
formula (II) as an example of $A^-$, there may be selected from compounds shown by NO$_3^-$, Cl$^-$, Br$^-$, CH$_3$COO$^-$, CO$_3^{2-}$ and SO$_4^{2-}$.

In the above general formula (II), when M is Al or Fe, p in M$^p$ is trivalence, thereby forming hydrotalcite compounds (in the case of Fe, a part of divalent one may be contained and it causes no problem since it converts easily into trivalent one during the reaction with a lithim compound and the drying step). When M is Co or Mn, p may be divalence, trivalence or mixture thereof since it may be used equally without problem. When M is Mg, p is divalent.

Among these compounds, LiOH is preferred as the lithium compound and a basic metal nitrate where $A^-$ is NO$_3^-$ is preferred as the basic metal salt represented by the general formula (II), from the viewpoint of yield, reactivity, effective utilization of the resources and oxidation accelerating effect. This combination is particularly preferred from the viewpoint of battery characteristics.

It is recommended from the viewpoint of uniformity that the basic metal salt is in fine particles of 0.1 $\mu$m or less with respect to the crystallite of the primary particle as measured by the Scherrer's method.

Also, it is preferred from the viewpoint of surface reactivity that this fine particle has a BET specific surface area of 10 $m^2/g$ or higher, preferably 40 $m^2/g$ or higher, more preferably 100 $m^2/g$ or higher. As to the
BET specific surface area, if it is measured after an aqueous solution of a basic metal salt has been dried, then the primary particles of very fine particles aggregates during dry process, thus BET specific surface area measured is one for the aggregation. If aggregate is strong, nitrogen gas cannot enter into it so that the value of BET specific surface area becomes small. Accordingly the basic metal salt which is practically reacted with a lithium compound in aqueous solution shows a high BET specific surface area to exhibit that the surface has a high reactivity. However, BET specific surface area was set to 10\text{m}^2/g or higher from the above actual circumstances. This basic metal salt having the specific composition has a layered structure, and both the chemical composition and the crystal structure are similar to those of Ni_{1-x}M_x(OH)_2. Moreover, it is microcrystalline and the surface is highly active as described above, so that it is reacted with a lithium compound such as LiOH thereby forming an extremely desirable precursor of Li_{y-x1}Ni_{1-x2}M_xO_2.

Highly purified Li_{y-x1}Ni_{1-x2}M_xO_2 having an extremely high crystallization degree at which the present invention aims can be obtained only when such a basic metal salt having a specific composition is used. The Ni_{1-x}M_x(OH)_2 is inferior in the reactivity with the lithium compound to the basic metal salt. On the other hand, when the amount of an anion is increased, the structure of the basic metal salt is increasingly deviated from the layered structure,
and the anion inhibitive acts on the formation Li\textsubscript{1-x}Ni\textsubscript{1-x}M\textsubscript{x}O\textsubscript{2} during the heating, so that the desired compound having a high purity and an extremely high crystallization degree cannot be obtained.

The basic metal salt to be used in the present invention can be prepared by adding an alkali to an aqueous solution of Ni\textsubscript{1-x}M\textsubscript{x} salt in an amount of about 0.7 \( \sim \) 0.95 equivalent, preferably about 0.8 \( \sim \) 0.95 equivalent based on the Ni\textsubscript{1-x}M\textsubscript{x} salt and reacting at a temperature of about 80°C or lower. Examples of the alkalis to be used in the reaction include alkali metal hydroxides such as sodium hydroxide, alkaline earth metal hydroxides such as calcium hydroxide, amines and the like. In this connection, it is preferable that, after the preparation, this basic metal salt be matured at 20 \( \sim \) 70°C for 0.1 \( \sim \) 10 hours. Subsequently, any by-products are removed by washing with water and the lithium compound is added.

The drying of the slurry obtained by such a reaction is carried out by spray drying. The spray drying where drying can be instantaneously accomplished to obtain spherical particles is preferred from the viewpoint of spherical particle granulation and the uniformity of the composition (In such dry processes requiring a certain drying time as shelf type drying and band drying ones, some Li migrates into the surface of particles to result in non-uniform composition).

In case that spherical particles with uniform
composition obtained using wet and spray drying processes is subjected to the heating as it is, the heating is effected at a temperature of 600℃ ~ 800℃, preferably 700℃ ~ 750℃ for 4 hours or higher, preferably about 4 ~ 20 hours in an atmosphere of oxygen. If the heating time is more than 20 hours, not only it makes cost up but also it causes evaporation of Li thereby the proportion of trivalent Ni to the total Ni becomes rather low and so the purity becomes bad.

In the known technique by the drying process regarding the heating, both the Li and Ni components are non-uniform in any cases during the reaction where the divalent Ni is converted into trivalent Ni, and hence the heating of at least 20 hours was required for Ni which is hard to become trivalent. In the light of the fact, the process of the present invention wherein an uniform spray dried powdered product is heated as it is is very economical and advantageous.

Nextly, the complex oxide having large primary particles of the present invention, a process for preparing the same and a positive electrode active material for a secondary battery which contains as an effective ingredient said complex oxide will be illustrated more specifically.

An average long diameter of this large primary particle product is a range of 1 ~ 10 μm.

By using the spherical particle product obtained by the above-mentioned spray drying process as a positive electrode
active material for a secondary battery and setting properly the kind of a metal as third component and the amount to be doped as solid solution, remarkable improvement in cycle characteristics may be achieved sufficiently, retaining the desired battery capacity. However, the recognition regarding the safety is being requested in the public. As to the battery to be used practically there has been increased the discussion that the use of it under severe conditions in a very rare case should be taken into consideration. Under the present situation where the function for the safety as a composite battery including other battery materials (native electrode, an electrolyte, sepalator etc.) with a positive electrode active material is not matured yet, a positive electrode active material which can be used at an elevated temperature has been required strongly.

The present invention has been made to meet this requirement and provide a positive electrode active material having the following characteristic properties.

That is, the composition of the positive electrode active material is similar to that of the above spherical particle product and the characteristic properties is characterized by that a ratio in peak intensity of the face (003) to the face (104) of Miller indices hkl of the X-ray diffraction pattern i.e., (003)/(104) is 1.2 or higher and a ratio in the peak intensity of the face (006) to the face (101) i.e., (006)/(101) is 0.13 or lower, and an average
long diameter of the primary particle product is a range of
$1 \sim 10 \ \mu m$, more preferably $2 \sim 10 \ \mu m$, as measures with
a SEM.

Said lithium nickel complex oxide is a novel one wherein
an average long diameter of the primary particles in the
sufficiently developed crystals is large and which are
excellent in stability.

Also, it is preferable that the positive electrode
active material having large primary particles is highly
purified to such extent as the proportion of $\text{Ni}^{2+}$ to the
total $\text{Ni}$ is 99% by weight or higher like the spray dried
particle product described above. It should have a BET
specific surface area of $0.01 \sim 1 \ m^2/g$, thus it has
adversely small BET specific surface area to that extent
that particles became larger. When used as the battery
material, it does not cause any raise in the viscosity of
the electrolyte and the reactivity with the electrolyte is
small.

The lithium nickel complex oxide having a large primary
particle in the present invention can be obtained by
reacting a basic metal salt represented by the above-
mentioned general formula (II) as the starting material
with a water soluble lithium compound in an aqueous medium
under condition such that a molar ratio of $\text{Li} / (\text{Ni} + \text{M})$
becomes $0.9 \sim 1.3$, spray drying the obtained slurry, and
subjecting the spray dried product to press molding.

As the process for preparing the positive electrode
active material, there may be applied one wherein the spray
dried product is subjected to press molding and then the
heating is conducted and an alternative one wherein after
the spray dried product has been once heated as it is,
resultant heat-treated product is subjected to press
molding and then re-heating is conducted.

First process is one which comprises subjecting the
spray dried product to press molding and then firing the
resultant molded material at a temperature of about 600°C
∼ 900°C for about 4 hours or more in an oxidative
atmosphere.

The spherical particle product obtained by the above-
mentioned spray dried process is excellent in flowability,
molding and filling properties, and may be pressed into a
shape as it is according to the conventional manner, for
example under a static pressure of usually 500 ∼ 3,000 kg
/㎝², preferably 800 ∼ 1,500 kg/㎝² using a Brinell
hardness tester. The press molding is of a great
significance in that moving distance among molecules becomes
short and thereby the crystal growth during the firing is
accelerated.

The press molded product obtained thusly can be heated
as it is.

The heating is effected at a temperature of usually 600
∼ 900°C, preferably 750°C ∼ 800°C for a period of usually
4 hours or higher, preferably 10 ∼ 72 hours under an
atmosphere of oxygen.
The crystal growth of primary particles during the heating is accelerated by a press molding and thereby obtaining the complex oxide having larger primary particles.

The process wherein this spherical particle product is hired through the press molding makes it possible to prepare a complex oxide having a high purity and a high crystallization degree even when the heating time of more than 20 hours was taken unlike the process wherein the spherical particle product is heated as it is for the reason that the contact area between particles becomes small because of compressed material so that evaporation of Li and formation of by-product NiO do not occur.

Second process for preparing the desired complex oxide is one wherein the spray dried product is heated at a temperature of 600°C ~ 900°C for 0.5 hour or more under an atmosphere of oxygen, the obtained heat-treated product after pulverized if necessary is pressed into a shape in a similar manner as in the first process, and then re-heated at a temperature of 600°C ~ 900°C for 1 hour or more under an atmosphere of oxygen. This process has an advantage that the total time required for the heated process may be shortened in comparison with the first process. In this connection, even when the spherical particle product which may be obtained by the above-mentioned spray drying process was heated for a long time as it was, the primary particles become large no longer. The heat-treatment for
20 hours gives the average particle size of 2μm or less while the heat-treatment for 72 hours gives the average particle size of about 2 ~ 3 μm or less.

Also, even when the press molding process was applied to the powdered complex oxide obtained by the prior art drying process, the average particle size of the 20 hours heated product is 1 μm or less, and there is almost no appreciable change in the particle size even when the heating time was merely extended further as shown in Comparative Examples described later.

Therefore, the combined process of the wet, spray drying and press molding ones involved in the present invention is extremely advantageous in making the primary particle larger.

In addition, although a distinct cause why it occurs has not been clear yet, the moving distance between powder-powder and that between Li and Ni molecules are considered to be decreased by the pressing procedure according to the present process, so that the evaporation of Li and formation of the by-product NiO may be suppressed and the desired complex oxide having a high purity and a high crystallization can be obtained even when the heat process was conducted for a long time.

As is apparent from the Examples described later, the complex oxide having larger primary particle is poor in the reactivity and has a small battery capacity owing to the larger primary particle but the stability (cycle property
and high temperature stabilities) is enhanced to that extent.

The heat-treated product composed of lithium nickel complex oxide obtained thusly may be properly pulverized into the desired size depending on use.

As is apparent from the Examples and Comparative Examples described later, when the lithium nickel complex oxide of the present invention is used as a positive electrode active material for a secondary battery, high discharge capacity of 130 ~ 200 mAh/g may be achieved and the decrease rate in discharge capacity is less than 10 % even after 100 cycles, indicating that it can be effectively utilized because of stability.

Brief Description of the Drawings

Fig. 1 is X-ray diffraction pattern (XRD) of the complex oxide obtained in Example 1.

Fig. 2 is SEM photograph (150 magnifications) of the complex oxide obtained in Example 1.

Fig. 3 is SEM photograph (30,000 magnifications) of the complex oxide obtained in Example 1.

Fig. 4 is particle size distribution of the complex oxide obtained in Example 1.

Fig. 5 is XRD of the complex oxide obtained in Example 5.

Fig. 6 is SEM photograph (10,000 magnifications) of the complex oxide obtained in Example 5.
Fig. 7 is XRD of the complex oxide obtained in Example 6.

Fig. 8 is SEM photograph (150 magnifications) of the complex oxide obtained in Example 6.

Fig. 9 is SEM photograph (30,000 magnifications) of the complex oxide obtained in Example 6.

Fig. 10 is particle size distribution of the complex oxide obtained in Example 6.

Fig. 11 is XRD of the complex oxide obtained in Example 9.

Fig. 12 is SEM photograph (10,000 magnifications) of the complex oxide obtained in Example 9.

Fig. 13 is XRD of the complex oxide obtained in Example 10.

Fig. 14 is SEM photograph (30,000 magnifications) of the complex oxide obtained in Example 10.

Fig. 15 is particle size distribution of the complex oxide obtained in Example 10.

Fig. 16 is XRD of the complex oxide obtained in Example 13.

Fig. 17 is SEM photograph (10,000 magnifications) of the complex oxide obtained in Example 13.

Fig. 18 is XRD of the complex oxide obtained in Example 14.

Fig. 19 is SEM photograph (150 magnifications) of the complex oxide obtained in Example 14.

Fig. 20 is SEM photograph (10,000 magnifications) of
the complex oxide obtained in Example 14.

Fig. 21 is particle size distribution of the complex oxide obtained in Example 14.

Fig. 22 is XRD of the complex oxide obtained in Example 16.

Fig. 23 is SEM photograph (1,000 magnifications) of the complex oxide obtained in Example 16.

Fig. 24 is SEM photograph (10,000 magnifications) of the complex oxide obtained in Example 16.

Fig. 25 is particle size distribution of the complex oxide obtained in Example 16.

Fig. 26 is XRD of the complex oxide obtained in Example 19.

Fig. 27 is SEM photograph (10,000 magnifications) of the complex oxide obtained in Example 19.

Fig. 28 is XRD of the complex oxide obtained in Example 21.

Fig. 29 is SEM photograph (50 magnifications) of the complex oxide obtained in Example 21.

Fig. 30 is SEM photograph (30,000 magnifications) of the complex oxide obtained in Example 21.

Fig. 31 is particle size distribution of the complex oxide obtained in Example 21.

Fig. 32 is XRD of the complex oxide obtained in Comparative Example 1.

Fig. 33 is SEM photograph (1,000 magnifications) of the complex oxide obtained in Comparative Example 1.
Fig. 34 is SEM photograph (10,000 magnifications) of the complex oxide obtained in Comparative Example 1.

Fig. 35 is particle size distribution of the complex oxide obtained in Comparative Example 1.

Fig. 36 is XRD of the complex oxide obtained in Comparative Example 5.

Fig. 37 is SEM photograph (3,500 magnifications) of the complex oxide obtained in Comparative Example 5.

Fig. 38 is SEM photograph (10,000 magnifications) of the complex oxide obtained in Comparative Example 5.

Fig. 39 is XRD of the complex oxide obtained in Comparative Example 15.

Fig. 40 is SEM photograph (20,000 magnifications) of the complex oxide obtained in Comparative Example 15.

Best Modes for Carrying Out the Invention

The present invention is more specifically illustrated by the following examples.

The respective methods for measuring the BET specific surface area, for measuring primary particle size by Scherrer’s method and for measuring trivalent nickel carried out with respect to the heat-treated product are as shown below.

[A method for measuring BET specific surface area]

A given amount of the heat-treated product was weighed in a cell as the sample to be employed and degassed by heating under the flow of a mixed gas of 30 % nitrogen and
70% helium, and thereafter specific surface area was measured according to the BET one point continuous flowing method. As BET specific surface area measuring apparatus, "MONOSORB" a product of Yuasa Ionics Co., Ltd. was employed.

[Scherrer's method]

It is a method wherein the size of crystallite is calculated by the following equation (1) on the assumption that the width of diffraction pattern is dependent on the size of crystallite alone owing to uniformity in the size of crystallite and no strain in the crystals.

\[ D_{hkl} = \frac{(k \lambda)}{(\beta \cos \theta)} \]  (Equation 1)

wherein \( D_{hkl} \) (angstrom) represents size of crystallite in vertical direction to the face (hkl), \( \lambda \) (angstrom) the wavelength of X-ray, \( \beta \) (rad) the width of diffraction pattern, \( \theta \) (°) angle of diffraction, and \( k \) a constant.

[A method for measuring \( \text{Ni}^{3+} \)]

\( \text{Ni}^{3+} \) is expressed in percentage based on the total Ni. It was measured by oxidation-reduction titration. 0.2 gram of a sample was dissolved in 0.25 M FeSO₄·3.6 N H₂SO₄ solution, and 2 ml of a concentrated phosphoric acid was added to the solution. The mixture was titrated with a 0.1 N KMnO₄ solution. Blank test was carried out in the same manner as in the above. The \( \text{Ni}^{3+} \) % was calculated by the following equation 2. In the equation 2, \( f \) represents the factor of the 0.1 N KMnO₄ solution, \( X_0 \) the titer of the blank test (ml), \( X \) the titer of the sample (ml), \( m \) the
amount of the sample (g), and B the content of Ni (%) and A 5.871.
\[ fX (X_0 - X) \times A \times 10 ] / (m \times B) \quad \text{(Equation 2)} \]

Example 1

There was prepared a mixed aqueous solution of 2.0 mol/l aluminum nitrate and nickel nitrate wherein the molar ratio of Al / (Ni + Al) was 0.03. This mixed aqueous solution and 2.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 50°C for about 60 minutes with vigorous stirring while pH was kept to 9.0.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni$_{0.97}$Al$_{0.03}$(OH)$_2$(NO$_3$)$_{0.03}$ slurry. A 3.5 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Al) was 1.05 with respect to the (Ni + Al) content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the firing product was Li$_{0.03}$Ni$_{0.97}$Al$_{0.03}$O$_2$.

Example 2

There was prepared a mixed aqueous solution of 2.0 mol/l aluminum nitrate and nickel nitrate wherein the molar ratio
of Al / (Ni + Al) was 0.02. This mixed aqueous solution and 2.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 50°C over the period of about 60 minutes with vigorous stirring while pH was kept to 9.5.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni_{0.88}Al_{0.12}(OH)_2(NO_3)_0.02 slurry. A 3.5 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Al) was 1.03 with respect to the (Ni + Al) content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was Li_{1.02}Ni_{0.88}Al_{0.12}O_2.

Example 3

There was prepared a mixed aqueous solution of 2.0 mol/l aluminum nitrate and nickel nitrate wherein the molar ratio of Al / (Ni + Al) was 0.1. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C over the period of about 60 minutes with vigorous stirring while pH was kept to 8.5.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni_{0.88}Al_{0.12}(OH)_2(NO_3)_0.1 slurry. A 3.5 mol/l
aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Al) was 1.05 with respect to the (Ni + Al) content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was LiNi$_{0.8}$Al$_{0.2}$O$_2$.

Example 4

There was prepared a mixed aqueous solution of 2.0 mol/l aluminum nitrate and nickel nitrate wherein the molar ratio of Al / (Ni + Al) was 0.2. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C over the period of about 60 minutes with vigorous stirring while pH was kept to 8.5.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni$_{0.8}$Al$_{0.2}$(OH)$_2$(NO$_3$)$_{0.2}$ slurry. A 3.5 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Al) was 1.05 with respect to the (Ni + Al) content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750 °C for
5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was Li$_{1.8}$Ni$_{0.8}$Al$_{0.2}$O$_2$.

Example 5

The spray dried product obtained in Example 3 was pressed into a shape under a static pressure of 1,000 kg/cm$^2$ using Brinell hardness tester. The shaped form was put in an alumina boat and fired at 750°C for 72 hours in an atmosphere of oxygen in a tube furnace. After cooling on standing, the heat-treated product was pulverized to obtain powdery LiNi$_{0.8}$Al$_{0.1}$O$_2$.

Example 6

There was prepared a mixed aqueous solution of 2.0 mol/l ferric nitrate and nickel nitrate wherein the molar ratio of Fe / (Ni + Fe) was 0.03. This mixed aqueous solution and 2.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 50°C over the period of about 60 minutes with vigorous stirring while pH was kept to 9.5.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni$_{0.8}$Fe$_{0.8}$(OH)$_2$(NO$_3$)$_{0.03}$ slurry. A 3.5 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Fe) was 1.03 with respect to the (Ni + Fe) content in the above slurry to effect reaction. Thereafter,
the reaction mixture was spray dried. The obtained dry material was put in a alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was \( \text{Li}_{1.02}\text{Ni}_{0.97}\text{Fe}_{0.03}\text{O}_2 \).

Example 7

There was prepared a mixed aqueous solution of 2.0 mol/l ferric nitrate and nickel nitrate wherein the molar ratio of Fe / (Ni + Fe) was 0.1. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C with vigorous stirring while pH was kept to 8.5.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l \( \text{Ni}_{0.9}\text{Fe}_{0.1}\text{(OH)}_2\text{(NO}_3\text{)}_{0.1} \) slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Fe) was 1.05 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in a alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was \( \text{Li}_{x}\text{Ni}_{0.9}\text{Fe}_{0.1}\text{O}_2 \).

Example 8
There was prepared a mixed aqueous solution of 2.0 mol/l ferric nitrate and nickel nitrate wherein the molar ratio of Fe / (Ni + Fe) was 0.2. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C with vigorous stirring while pH was kept to 8.5.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni$_{0.8}$Fe$_{0.2}$$(OH)_2(\text{NO}_3)_2$ slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Fe) was 1.05 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in a alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was LiNi$_{0.8}$Fe$_{0.2}$O$_2$.

Example 9

The spray dried product obtained in Example 7 was pressed into a shape under a static pressure of 1,000 kg/cm$^2$ using Brinell hardness tester. The shaped form was put in an alumina boat and heated at 750 °C for 20 hours in an atmosphere of oxygen in a tube furnace. After cooling on standing, the heat-treated product was pulverized to obtain
powdery LiNi_{0.5}Fe_{0.5}O_{2}.

Example 10

There was prepared a mixed aqueous solution of 1.0 mol/l manganese nitrate and nickel nitrate wherein the molar ratio of Mn / (Ni + Mn) was 0.03. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C with vigorous stirring while pH was kept to 8.5.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni_{0.97}Mn_{0.03}(OH)_{1.87}(NO_{3})_{0.03} slurry. (This basic nitrate salt was dried and the resultant dry one has a BET specific surface area of 147.18 m²/g and a primary particle size (crystal lattice) of 44.1 angstroms as the value calculated by Scherrer's method from the half value width of the peak appeared at near 2θ = 60° of XRD ). A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Mn) was 1.05 with respect to the Ni content in the above suspension to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750 °C for 5 hours in an atmosphere of oxygen in a tube furnace and pulverized in a mortarto obtain powdery LiNi_{0.97}Mn_{0.03}O_{2}.
Example 11

There was prepared a mixed aqueous solution of 1.0 mol/l manganese nitrate and nickel nitrate wherein the molar ratio of Mn / (Ni + Mn) was 0.1. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25 °C with vigorous stirring while pH was kept to 8.5.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni$_{0.8}$Mn$_{0.2}$(OH)$_{1.8}$(NO$_3$)$_{0.1}$ slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Mn) was 1.05 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was LiNi$_{0.8}$Mn$_{0.2}$O$_2$.

Example 12

There was prepared a mixed aqueous solution of 1.0 mol/l manganese nitrate and nickel nitrate wherein the molar ratio of Mn / (Ni + Mn) was 0.4. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25 °C with vigorous stirring while pH was kept to 8.5.
The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni$_{0.8}$Mn$_{0.4}$(OH)$_{1.7}$(NO$_3$)$_{0.3}$ slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Mn) was 1.05 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product (powder) was LiNi$_{0.8}$Mn$_{0.4}$O$_2$.

Example 13

The spray dried product obtained in Example 11 was pressed into a shape under a static pressure of 1,000 kg/cm$^2$ using Brinell hardness tester. The shaped form was put in an alumina boat and heated at 750 °C for 20 hours in an atmosphere of oxygen in a tube furnace. After cooling on standing, the heat-treated product was pulverized to obtain powdery LiNi$_{0.8}$Mn$_{0.4}$O$_2$.

Example 14

There was prepared a mixed aqueous solution of 2.0 mol/l cobalt nitrate and nickel nitrate wherein the molar ratio of Co / (Ni + Co) was 0.03. This mixed aqueous solution and 2.0 mol/l aqueous sodium hydroxide solution were
simultaneously added at 40 °C over the period of about 60 minutes with vigorous stirring while pH was kept to 9.0.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni\(_{0.87}\)Co\(_{0.13}\)(OH)\(_2\)(NO\(_3\))\(_{0.83}\) slurry. A 3.5 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Co) was 1.05 with respect to the (Ni + Co) content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was Li\(_{1.03}\)Ni\(_{0.87}\)Co\(_{0.13}\)O\(_2\).

Example 15

There was prepared a mixed aqueous solution of 1.0 mol/l cobalt nitrate and nickel nitrate wherein the molar ratio of Co / (Ni + Co) was 0.1. This mixed aqueous solution and 2.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C with vigorous stirring while pH was kept to 8.5.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni\(_{0.87}\)Co\(_{0.13}\)(OH)\(_2\)(NO\(_3\))\(_{0.83}\) slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio

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of Li / (Ni + Co) was 1.05 with respect to the Ni content in the above suspension to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750 °C for 5 hours in an atmosphere of oxygen in a tube furnace to obtain powdery LiNi_{0.8}Co_{0.2}O_{2}.

Example 16

There was prepared a mixed aqueous solution of 1.0 mol/l cobalt nitrate and nickel nitrate wherein the molar ratio of Co / (Ni + Co) was 0.2. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C with vigorous stirring while pH was kept to 8.5.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni_{0.8}Co_{0.2}(OH)_{1.9}NO_{3} slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Co) was 1.05 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace to obtain powdery LiNi_{0.8}Co_{0.2}O_{2}.

Example 17
There was prepared a mixed aqueous solution of 1.0 mol/l cobalt nitrate and nickel nitrate wherein the molar ratio of Co / (Ni + Co) was 0.3. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C with vigorous stirring while pH was kept to 8.5.

The obtained precipitate was filtered out, washed with water and suspended in water thereby obtaining a 1 mol/l Ni$_{0.7}$Co$_{0.3}$(OH)$_{1.7}$(NO$_3$)$_{0.3}$ slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwise to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Co) was 1.05 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750 °C for 5 hours in an atmosphere of oxygen in a tube furnace to obtain powdery LiNi$_{0.7}$Co$_{0.3}$O$_2$.

Example 18

There was prepared a mixed aqueous solution of 1.0 mol/l cobalt nitrate and nickel nitrate wherein the molar ratio of Co / (Ni + Co) was 0.4. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C with vigorous stirring while pH was kept to 8.5.

The obtained precipitate was filtered out, washed with water and suspended in water thereby obtaining a 1 mol/l
Ni\textsubscript{0.8}Co\textsubscript{0.2}(OH)\textsubscript{1.7}(NO\textsubscript{3})\textsubscript{0.3} slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Co) was 1.05 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace to obtain powdery LiNi0\textsubscript{0.8}Co0\textsubscript{0.2}O\textsubscript{2}.

Example 19

There was prepared a mixed aqueous solution of 1.0 mol/l cobalt nitrate and nickel nitrate wherein the molar ratio of Co / (Ni + Co) was 0.2. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C with vigorous stirring while pH was kept to 11.5.

The obtained precipitate was filtered out, washed with water and suspended in water thereby obtaining a 1 mol/l Ni\textsubscript{0.8}Co\textsubscript{0.2}(OH)\textsubscript{1.7}(NO\textsubscript{3})\textsubscript{0.3} slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Co) was 1.05 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained spray-dried product was pressed into a shape under a static pressure of 1,300kg/cm\textsuperscript{2} Brinell hardness tester. The shaped form was
put in an alumina boat and heated at 750 °C for 72 hours in an atmosphere of oxygen in a tube furnace. After cooling on standing, the heat-treated product was pulverized to obtain powdery LiNi₀.₈Co₀.₂O₂.

Example 20

The spray dried product obtained in Example 15 was pressed into a shape under a static pressure of 1,000 kg/cm² using Brinell hardness tester. The shaped form was put in an alumina boat and heated at 750 °C for 20 hours in an atmosphere of oxygen in a tube furnace. After cooling on standing, the heat-treated product was pulverized to obtain powdery LiNi₀.₈Co₀.₂O₂.

Example 21

There was prepared a mixed aqueous solution of 1.0 mol/l magnesium nitrate and nickel nitrate wherein the molar ratio of Mg / (Ni + Mg) was 0.049. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25 °C with vigorous stirring while pH was kept to 11.0.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni₀.₈₈Mg₀.₁₂(OH)₁.₁(NO₃)₂.₃ slurry. (The BET specific surface area of the dried basic nitrate salt was 169.4 m²/g and its primary particle size (crystal lattice) was 32.3 angstroms as the value calculated by the
Scherrer's method from the half value width of the peak appeared at near $2\theta = 60^\circ$ of XRD). A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Mg) was 1.0 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The spray dried material was put in an alumina boat, heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace and pulverized in a mortar to obtain powdery LiNi$_{0.87}$Mg$_{0.13}$O$_2$.

Example 22

The spray dried product obtained in Example 15 was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen. After cooling on standing, the firing product was pulverized and pressed into a shape under a static pressure of 1,300 kg/cm$^2$ using Brinell hardness tester. The press shaped form was once more put in an alumina boat and heated at 800 °C for 1 hour in an atmosphere of oxygen in a tube furnace. After cooling on standing, the heat-treated product was pulverized to obtain powdery LiNi$_{0.8}$Co$_{0.2}$O$_2$.

Comparative Example 1

1,900 ml of 1.0 mol/l aqueous sodium hydroxide solution was added under stirring to 500 ml of 2.0 mol/l aqueous nickel nitrate solution, thus the molar ratio of Na/Ni
being 1.9. The resultant reaction mixture was filtered, washed with water and suspended in water, thereby obtaining 1 mol/l Ni(OH)$_{1.97}$ (NO$_3$)$_{0.03}$ slurry. A 3.5 mol/l aqueous lithium hydroxide solution was added dropwisely in an amount such that the atomic ratio of Li/Ni was 1.05 with respect to the Ni content in the slurry to effect a reaction. Thereafter, the reaction mixture was spray dried. The obtained dried gel was put in an alumina boat, and heated at 750 °C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was Li$_{1.02}$NiO$_2$.

Comparative Example 2

1.05 mol sodium hydroxide, 0.97 mol nickel hydroxide and 0.03 mol aluminum hydroxide were sufficiently mixed in dry state and pulverized in a mortar, and then pelletized into size of 14 mm of a diameter × 2 mm of a thickness. The resultant pellets were heated at 750 °C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was Li$_{1.04}$Ni$_{0.97}$Al$_{0.03}$O$_2$.

Comparative Example 3

1.05 mol sodium hydroxide, 0.97 mol nickel hydroxide and 0.03 mol ferric oxide were sufficiently mixed in dry state and pulverized in a mortar, and then pelletized into
size of 14 mm of a diameter × 2 mm of a thickness. The resultant pellets were heated at 750 °C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was Li$_{1.04}$Ni$_{0.97}$Fe$_{0.03}$O$_2$.

Comparative Example 4

1.05 mol sodium hydroxide, 0.97 mol nickel hydroxide and 0.03 mol manganese dioxide were sufficiently mixed in dry state and pulverized in a mortar, and then pelletized into size of 14 mm of a diameter × 2 mm of a thickness. The resultant pellets were heated at 750 °C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was Li$_{1.04}$Ni$_{0.97}$Mn$_{0.03}$O$_2$.

Comparative Example 5

1.05 mol sodium hydroxide, 0.97 mol nickel hydroxide and 0.03 mol cobalt hydroxide were sufficiently mixed in dry state and pulverized in a mortar, and then pelletized into size of 14 mm of a diameter × 2 mm of a thickness. The resultant pellets were heated at 750 °C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was Li$_{1.04}$Ni$_{0.97}$Co$_{0.03}$O$_2$.

Comparative Example 6
1.05 mol sodium hydroxide, 0.97 mol nickel hydroxide and 0.03 mol magnesium oxide were sufficiently mixed in dry state and pulverized in a mortar, and then pelletized into size of 14 mm of a diameter × 2 mm of a thickness. The resultant pellets were heated at 750 °C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was Li_{1.04}Ni_{0.97}Mg_{0.03}O_2.

Comparative Example 7

There was prepared a mixed aqueous solution of 2.0 mol/l aluminum nitrate and nickel nitrate wherein the molar ratio of Al / (Ni + Al) was 0.3. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C with vigorous stirring while pH was kept to 8.5.

The obtained precipitate was filtered out, washed with water and suspended in water thereby obtaining a 1 mol/l Ni_{0.7}Al_{0.3}(OH)_2(NO_3)_2.3 slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Al) was 1.05 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was
LiNi\textsubscript{0.7}Al\textsubscript{0.3}O\textsubscript{2}.

Comparative Example 8

There was prepared a mixed aqueous solution of 2.0 mol/l ferric nitrate and nickel nitrate wherein the molar ratio of Fe / (Ni + Fe) was 0.3. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C with vigorous stirring while pH was kept to 8.5.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni\textsubscript{0.7}Fe\textsubscript{0.3}(OH)\textsubscript{2}(NO\textsubscript{3})\textsubscript{0.3} slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Fe) was 1.05 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in a alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace.

The chemical composition of the heat-treated product was LiNi\textsubscript{0.7}Fe\textsubscript{0.3}O\textsubscript{2}.

Comparative Example 9

There was prepared a mixed aqueous solution of 1.0 mol/l magnesium nitrate and nickel nitrate wherein the molar ratio of Mg / (Ni + Mg) was 0.3. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution.
were simultaneously added at 25°C with vigorous stirring while pH was kept to 11.0.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni$_{0.7}$Mg$_{0.3}$(OH)$_{1.7}$(NO$_3$)$_{0.3}$ slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Mg) was 1.0 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace to obtain powdery LiNi$_{0.7}$Mg$_{0.3}$O$_2$.

Comparative Example 10

There was prepared a mixed aqueous solution of 1.0 mol/l manganese nitrate and nickel nitrate wherein the molar ratio of Mn / (Ni + Mn) was 0.6. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C with vigorous stirring while pH was kept to 8.5.

The obtained reaction mixture was filtered, washed with water and suspended in water thereby obtaining a 1 mol/l Ni$_{0.4}$Mn$_{0.6}$(OH)$_{1.7}$(NO$_3$)$_{0.3}$ slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Mn) was 1.05 with respect to the Ni content.
in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained dry material was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace to obtain powdery LiNi_{0.4}Mn_{0.6}O_{2}.

Comparative Example 11

There was prepared a mixed aqueous solution of 1.0 mol/l cobalt nitrate and nickel nitrate wherein the molar ratio of Co / (Ni + Co) was 0.6. This mixed aqueous solution and 1.0 mol/l aqueous sodium hydroxide solution were simultaneously added at 25°C with vigorous stirring while pH was kept to 8.5.

The obtained precipitate was filtered out, washed with water and suspended in water thereby obtaining a 1 mol/l Ni_{0.4}Co_{0.6}(OH)_{1.7}(NO_3)_{0.3} slurry. A 3.0 mol/l aqueous lithium hydroxide solution was added dropwisely to the obtained slurry in an amount such that the atomic ratio of Li / (Ni + Co) was 1.05 with respect to the Ni content in the above slurry to effect reaction. Thereafter, the reaction mixture was spray dried. The obtained spray-dried product was put in an alumina boat and heated at 750°C for 5 hours in an atmosphere of oxygen in a tube furnace, and then pulverized in a mortar to obtain powdery LiNi_{0.4}Co_{0.6}O_{2}.

Comparative Example 12
1.05 mol sodium hydroxide, 0.9 mol nickel hydroxide and 0.1 mol aluminum hydroxide were sufficiently mixed in dry state and pulverized in a mortar, and then pressed into a shape under a static pressure of 1,000 kg/cm² using Brinell hardness tester. The shaped form was put in an alumina boat and heated at 750 °C for 20 hours in an atmosphere of oxygen in a tube furnace. After cooling on standing, the heat-treated product was pulverized to obtain powdery LiNi_{0.5}Al_{0.5}O_2.

Comparative Example 13

1.05 mol sodium hydroxide, 0.9 mol nickel hydroxide and 0.1 mol ferric oxide were sufficiently mixed in dry state and pulverized in a mortar, and then pressed into a shape under a static pressure of 1,000 kg/cm² using Brinell hardness tester. The shaped form was put in an alumina boat and heated at 750 °C for 20 hours in an atmosphere of oxygen in a tube furnace. After cooling on standing, the heat-treated product was pulverized to obtain powdery LiNi_{0.5}Fe_{0.5}O_2.

Comparative Example 14

1.05 mol sodium hydroxide, 0.9 mol nickel hydroxide and 0.1 mol manganese dioxide were sufficiently mixed in dry state and pulverized in a mortar, and then pressed into a shape under a static pressure of 1,000 kg/cm² using Brinell hardness tester. The shaped form was put in an alumina
boat and heated at 750 °C for 20 hours in an atmosphere of oxygen in a tube furnace. After cooling on standing, the heat-treated product was pulverized to obtain powdery \( \text{LiNi}_{0.8}\text{Mn}_{0.2}\text{O}_2 \).

Comparative Example 15

1.05 mol sodium hydroxide, 0.9 mol nickel hydroxide and 0.1 mol cobalt hydroxide were sufficiently mixed in dry state and pulverized in a mortar, and then pressed into a shape under a static pressure of 1,000 \( \text{kg/f} \) using Brinell hardness tester. The shaped form was put in an alumina boat and heated at 750 °C for 20 hours in an atmosphere of oxygen in a tube furnace. After cooling on standing, the heat-treated product was pulverized to obtain powdered \( \text{LiNi}_{0.8}\text{Co}_{0.2}\text{O}_2 \).

Tables 1 and 2 show the ratios in X-ray peak intensity (003)/(104) and (006)/(101) calculated from the X-ray diffraction patterns, BET specific surface area, \( \text{Ni}^{3+} \) ratio (%), an average secondary particle size as measured by laser microtrack and long diameter of primary particle obtained from the SEM photograph, measured with respect to each of the compound oxides obtained in Examples 1 ~ 22 and Comparative Examples 1 ~ 15.
Table 1
The physical properties of the complex oxides obtained in Examples 1 ~ 22

<table>
<thead>
<tr>
<th>Example No.</th>
<th>Ni(^{2+}) (%)</th>
<th>BET m(^2)/g</th>
<th>Ratio of I(003)/I(104)</th>
<th>Ratio of I(006)/I(101)</th>
<th>Average Particle Size (µ)</th>
<th>Average Long Diameter of Primary Particle as Observed by SEM (µ)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>99.4</td>
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<td>1.43</td>
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<tr>
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<tr>
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<td>6.30</td>
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<td>0.39</td>
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<td>1.304</td>
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</tr>
<tr>
<td>22</td>
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<td>1.4</td>
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Table 2
The physical properties of the complex oxides obtained in Comparative Examples 1 ~ 15

<table>
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<th>Comparative Example No.</th>
<th>Ni^{3+} (%)</th>
<th>BET m^2/g</th>
<th>Ratio of I(003)/I(104)</th>
<th>Ratio of I(006)/I(101)</th>
<th>Average Particle Size (μ)</th>
<th>Average Long Diameter of Primary Particle as Observed by SEM (μ)</th>
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<tbody>
<tr>
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</table>

Also, the primary particle size of the basic metal salt (II) employed in the wet process is shown in Table 3.
Table 3

The physical properties of the basic metal salt employed in the reaction

<table>
<thead>
<tr>
<th>Example No.</th>
<th>Primary Particle Size/Scherrer's method (Angstrom)</th>
</tr>
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<tbody>
<tr>
<td>Example 1</td>
<td>34.1</td>
</tr>
<tr>
<td>Example 2</td>
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<td>32.3</td>
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<tr>
<td>Example 22</td>
<td>38.8</td>
</tr>
</tbody>
</table>
The long diameter of the primary particle size for all of the spray dried and heat-treated complex oxides were in the range of 0.2 \( \sim \) 3.0 \( \mu \)m.

Experiment 1

The following battery test (charge and discharge test) was carried out using the complex oxides obtained in Examples 1, 2, 6 and 14 and Comparative Examples 1, 2, 3 and 5.

For a positive electrode material, each of the lithium nickel complex oxides obtained in said Examples and said Comparative Examples was mixed with a conductive binder (polytetrafluoroethylene - acetylene black) in a weight ratio of 2:1. Thereafter, the mixture was formed into pellets each having a thickness of 0.5 mm and a diameter of 18 mm. The resultant pellets were press-bonded at a pressure of 1 t/cm\(^2\) to an expanded stainless steel mesh by means of a pressing machine, thereby obtaining a positive electrode molding.

As a negative electrode material, a disk of 18 mm in diameter blanked from a lithium metal sheet was used. The positive electrode molding was put in a coin type cell composed of stainless steel, and an electrolyte in which 1 mol/l LiPF\(_6\) was dissolved in a 1:4 mixture of propylene carbonate and ethylene carbonate was poured into the cell in an appropriate amount. A separator, the above negative electrode and its case were piled in this order on the
electrolyte, thereby obtaining lithium secondary battery for test. All of the above procedures were conducted in an atmosphere of argon. The performance of the positive electrode active material was evaluated by repeating charging and discharging the thus obtained lithium secondary battery and measuring the initial discharge capacity and the discharge capacity decrease attributed to the repetition of charging and discharging. The above charging and discharging were performed under a voltage control between 3 V and 4.3 V at a constant current of 1 mA.

The results of the battery test [initial discharge capacity (mAh/g), discharge capacity at 100th cycle (mAh/g) and decrease rate of discharge capacity at 100th cycle (%)] are shown in Tables 4 and 5.
### Table 4

<table>
<thead>
<tr>
<th>Example No.</th>
<th>Initial Discharge Capacity (mAh/g)</th>
<th>Discharge Capacity at 100th Cycle (mAh/g)</th>
<th>Decrease Rate of Discharge Capacity at 100th Cycle (%)</th>
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</thead>
<tbody>
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<td>14</td>
<td>160.2</td>
<td>146.2</td>
<td>8.7</td>
</tr>
</tbody>
</table>

### Table 5

<table>
<thead>
<tr>
<th>Comparative Example No.</th>
<th>Initial Discharge Capacity (mAh/g)</th>
<th>Discharge Capacity at 100th Cycle (mAh/g)</th>
<th>Decrease Rate of Discharge Capacity at 100th Cycle (%)</th>
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<tbody>
<tr>
<td>1</td>
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<td>5</td>
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</table>
Experiment 2

The following battery test (charge and discharge test) was carried out using the complex oxides obtained in Examples 1 ~ 22 and Comparative Examples 1 ~ 15.

For a positive electrode material, each of the lithium nickel compound oxides obtained in the above Examples and Comparative Examples, etylene black as a conductive agent and polytetrafluoroethylene as a binder were mixed in a ratio of 88 : 6 : 6 % by weight. Thereafter, the mixture was subjected to compression molding on stainless steel mesh, thereby obtaining pellets each having a thickness of 5 mm and a diameter of 18 mm. The resultant pellets were dried at 200°C for 2 hours, and used as a positive electrode material.

As a negative electrode material was used a rolled lithium metal sheet pressed-bonded to a stainless substrate. As a diaphragm, porous membrane made of polypropylene (trade name “Cell Guard”, a product of Hoechst Japan Co., Ltd.) and a glass filter-paper filter were used. There was used an electrolyte in which 1 M LiClO₄ was dissolved in an ethylene carbonate/dimethylmethoxyethane mixture (1:1 weight ratio). The procedures up to completion from setup of a test cell (semi-open type cell) were conducted in an argon replaced dry box. The charging and discharging for this lithium battery were performed under a voltage control between 3 V and 4.3 V at a constant current density of 0.4 mA/cm².
The results of the battery test [initial discharge capacity (mAh/g), discharge capacity at 100th cycle (mAh/g) and decrease rate of discharge capacity at 100th cycle (%)] are shown in Tables 6 and 7.
<table>
<thead>
<tr>
<th>Example No.</th>
<th>Initial Discharge Capacity (mAh/g)</th>
<th>Discharge Capacity at 100th Cycle (mAh/g)</th>
<th>Decrease Rate of Discharge Capacity at 100th Cycle (%)</th>
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<tr>
<td>Example No.</td>
<td>Initial Discharge Capacity (mAh/g)</td>
<td>Discharge Capacity at 100th Cycle (mAh/g)</td>
<td>Decrease Rate of Discharge Capacity at 100th Cycle (%)</td>
</tr>
<tr>
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<tr>
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</tr>
</tbody>
</table>

Experiment 3

As an index indicating that the complex oxides obtained in the present invention are stable at an elevated
temperature, an exothermic reaction temperature of the positive electrode material after charging was measured by the following method.

Using the test cell prepared in Experiment 2, the positive electrode material after charging was subjected to thermal analysis by DSC ("THERMOFLEX TAS200", a product of Rigaku Co., Ltd.) under an inert gas to measure the exothermic reaction temperature.

As a result, the peak of exothermic heat was recognized to appear at 208.6°C for Comparative Example 1, 224.3°C for Example 15 and at 237.6°C for Example 20.

The temperature rise in the exothermic peak by introduction of Co etc. into LiNiO₂ was recognized. Furthermore, it is understood that the positive electrode material composed of larger primary particle size even among those having the same composition gives higher the exothermic peak temperature, thereby the stability of the positive electrode material at an elevated temperature is enhanced.

Industrial Applicability

As explained above, according to the present invention there can be provided a novel lithium nickel complex oxide represented by the general formula:

\[ \text{Li}_{7-x_1}\text{Ni}_{1-x_2}\text{M}_x\text{O}_2 \]

(wherein M represents one selected from the group consisting of Al, Fe, Co, Mn and Mg, \( x = x_1 + x_2, 0 < x_1 \)).
\[ 0.2 \leq 0.2, \quad 0 < x_2 \leq 0.5, \quad 0 < x \leq 0.5, \quad \text{and} \quad 0.9 \leq y \leq 1.3 \), wherein the crystals have been sufficiently developed and are highly purified, and a positive electrode active material for a secondary battery whose stability of high discharge capacity is excellent.
CLAIMS

1. A process for preparing a lithium nickel complex oxide represented by the general formula (I)

\[ \text{Li}_{y-x_1}\text{Ni}_{1-x_2}M_x\text{O}_2 \] (I)

wherein M represents a metal selected from the group consisting of Al, Fe, Co, Mn and Mg, \( x = x_1 + x_2 \) and

(i) when M is Al or Fe, then \( x, x_1 \) and \( x_2 \) represent respectively \( 0 < x \leq 0.2, x_1 = 0, \) and \( x_2 = x; \)

(ii) when M is Co or Mn, then \( x, x_1 \) and \( x_2 \) represent respectively \( 0 < x \leq 0.5, x_1 = 0, \) and \( x_2 = x; \)

(iii) when M is Mg, then \( x, x_1 \) and \( x_2 \) represent respectively \( 0 < x \leq 0.2, 0 < x_1 \leq 0.2 \) and \( 0 < x_2 \leq 0.2; \) and \( y \) represents \( 0.9 \leq y \leq 1.3, \)

said process comprising the steps of:

(a) reacting a basic metal salt represented by the general formula (II)

\[ \text{Ni}_{1-x}M^p_x(\text{OH})_{2-n_z}(\text{A}^{n^-})_{z+\lfloor p x-2 x_1/n \rfloor} \cdot m\text{H}_2\text{O} \] (II)

wherein M represents a metal selected from the group consisting of Al, Fe, Co, Mn and Mg, \( p \) is a valence number of M with \( 2 \leq p \leq 3, \) \( \text{A}^{n^-} \) is an anion having a valence of \( n, \) and \( x, z \) and \( m \) are positive numbers respectively satisfying the ranges of \( 0 < x \leq 0.2, 0.03 \leq z \leq 0.3, 0 \leq m < 2; \) with a water soluble lithium compound in an aqueous medium in a molar ratio of \( \text{Li}/(\text{Ni}+\text{M}) \) of \( 0.9 \sim 1.3 \) to obtain a slurry,
(b) spray drying the obtained slurry; and

(c) heating the resultant dry material at a temperature of about 600°C ~ 900°C for about 4 hours or more in an oxidative atmosphere.

2. A process for preparing a lithium nickel complex oxide represented by the general formula (I)

\[
\text{Li}_{y-x_1}\text{Ni}_{1-x_2}M_x\text{O}_2
\]  

(I)

wherein M represents a metal selected from the group consisting of Al, Fe, Co, Mn and Mg, \(x = x_1 + x_2\), and

(i) when M is Al or Fe, then \(x, x_1\) and \(x_2\) represent respectively \(0 < x \leq 0.2, x_1 = 0, \) and \(x_2 = x\);

(ii) when M is Co or Mn, then \(x, x_1\) and \(x_2\) represent respectively \(0 < x \leq 0.5, x_1 = 0, \) and \(x_2 = x\);

(iii) when M is Mg, then \(x, x_1\) and \(x_2\) represent respectively \(0 < x \leq 0.2, 0 < x_1 \leq 0.2\) and \(0 < x_2 \leq 0.2\); and \(y\) represents \(0.9 \leq y \leq 1.3\),

said process comprising the steps of:

(a) reacting a basic metal salt represented by the general formula (II)

\[
\text{Ni}_{1-xM^P}x(\text{OH})_{2-nz}(\text{A}^n^-)^{[2+(p_x-2z)/n]} \cdot m\text{H}_2\text{O}
\]  

(II)

wherein M represents a metal selected from the group consisting of Al, Fe, Co, Mn and Mg, \(p\) is a valence number of M with \(2 \leq p \leq 3\), \(A^n^-\) is an anion having a valence of \(n\), and \(x, z\) and \(m\) are positive numbers respectively satisfying the
ranges of $0 < x \leq 0.2$, $0.03 \leq z \leq 0.3$, $0 \leq m < 2$; with a water
soluble lithium compound in an aqueous medium in a molar ratio
of $Li/(Ni+M)$ of $0.9 \sim 1.3$ to obtain a slurry,

(b) spray drying the obtained slurry;

(c) subjecting the spray dried product to press molding;
and

(d) heating the molded product at a temperature of about
$600^\circ C \sim 900^\circ C$ for about 4 hours or more in an oxidative
atmosphere.

3. A process for preparing a lithium nickel complex oxide
represented by the general formula (I)

$$Li_{y-x_1}Ni_{1-x_2}M_xO_2$$  (I)

wherein $M$ represents a metal selected from the group consisting
of Al, Fe, Co, Mn and Mg, $x = x_1 + x_2$, and

(i) when $M$ is Al or Fe, then $x$, $x_1$ and $x_2$ represent
respectively $0 < x \leq 0.2$, $x_1 = 0$, and $x_2 = x$;

(ii) when $M$ is Co or Mn, then $x$, $x_1$ and $x_2$ represent
respectively $0 < x < 0.5$, $x_1 = 0$, and $x_2 = x$;

(iii) when $M$ is Mg, then $x$, $x_1$ and $x_2$ represent respectively
$0 < x \leq 0.2$, $0 < x_1 \leq 0.2$, and $0 < x_2 \leq 0.2$; and $y$ represents
$0.9 \leq y \leq 1.3$,

said process comprising the steps of:

(a) reacting a basic metal salt represented by the general
formula (II)

67
\[ \text{Ni}_{1-x}\text{M}^p \ x(\text{OH})_{2-nz}(\text{A}^{n-})_{[z+(p-px)/n]} \cdot \text{mH}_2\text{O} \quad \text{(II)} \]

wherein M represents a metal selected from the group consisting of Al, Fe, Co, Mn and Mg, p is a valence number of M with \( 2 \leq p \leq 3 \), \( \text{A}^{n-} \) is an anion having a valence of \( n \), and \( x \), \( z \) and \( m \) are positive numbers respectively satisfying the ranges of \( 0 < x \leq 0.2, \ 0.03 \leq z \leq 0.3, \ 0 \leq m < 2 \); with a water soluble lithium compound in an aqueous medium in a molar ratio of \( \text{Li}/(\text{Ni}+\text{M}) \) of \( 0.9 \sim 1.3 \) to obtain a slurry,

(b) spray drying the obtained slurry;

(c) heating the spray dried product at a temperature of about \( 600^\circ\text{C} \sim 900^\circ\text{C} \) for about 0.5 hour or more in an oxidative atmosphere;

(d) pulverizing the resultant heat-treated product;

(e) subjecting the pulverized heat-treated product to press molding; and

(f) re-heating the molded product at a temperature of about \( 600^\circ\text{C} \sim 900^\circ\text{C} \) for about 1 hour or more in an oxidative atmosphere.

4. The process of claim 1 wherein M of the general formulae (I) and (II) is Al or Mg.

5. The process of claim 2 wherein M of the general formulae (I) and (II) is Al or Mg.

6. The process of claim 3 wherein M of the general formulae (I) and (II) is Al or Mg.
7. A positive electrode active material for a secondary battery which contains as an active ingredient the lithium nickel complex oxide prepared by the process as claimed in any one of claims 1 to 6.

8. The positive electrode active material for a secondary battery as claimed in claim 7 in which the decrease rate of discharge capacity at 100th cycle based on the initial discharge capacity is 10% or less.
Fig. 1
XRD of the complex oxide obtained in Example 1
Fig. 2
SEM photograph (150 magnifications) of the complex oxide obtained in Example 1

Fig. 3
SEM photograph (30,000 magnifications) of the complex oxide obtained in Example 1
Fig. 4

Particle size distribution of the complex oxide obtained in Example 1
Fig. 5

XRD of the complex oxide obtained in Example 5
Fig. 6
SEM photograph (10,000 magnifications) of the complex oxide obtained in Example 5
Fig. 7

XRD of the complex oxide obtained in Example 6
Fig. 8
SEM photograph (150 magnifications) of the complex oxide obtained in Example 6

Fig. 9
SEM photograph (30,000 magnifications) of the complex oxide obtained in Example 6
Fig. 10

Particle size distribution of the complex oxide obtained in Example 6
Fig. 11

XRD of the complex oxide obtained in Example 9
Fig. 12

SEM photograph (10,000 magnifications) of the complex oxide obtained in Example 9
Fig. 13

XRD of the complex oxide obtained in Example 10
Fig. 14

SEM photograph (30,000 magnifications) of the complex oxide obtained in Example 10
Fig. 15

Particle size distribution of the complex oxide obtained in Example 10
Fig. 16

XRD of the complex oxide obtained in Example 13
Fig. 17

SEM photograph (10,000 magnifications) of the complex oxide obtained in Example 13
Fig. 18

XRD of the complex oxide obtained in Example 14
Fig. 19
SEM photograph (150 magnifications) of the complex oxide obtained in Example 14

Fig. 20
SEM photograph (10,000 magnifications) of the complex oxide obtained in Example 14
Fig. 21

Particle size distribution of the complex oxide obtained in Example 14

Amount of particle

Cumulative particle amount
Fig. 22

XRD of the complex oxide obtained in Example 16
Fig. 23
SEM photograph (1,000 magnications) of the complex oxide obtained in Example 16

Fig. 24
SEM photograph (10,000 magnications) of the complex oxide obtained in Example 16
Fig. 25

Particle size distribution of the complex oxide obtained in Example 16
Fig. 26

XRD of the complex oxide obtained in Example 19.
Fig. 27

SEM photograph (10,000 magnifications) of the complex oxide obtained in Example 19
Fig. 28

XRD of the complex oxide obtained in Example 21
Fig. 29
SEM photograph (50 magnifications) of the complex oxide obtained in Example 21

Fig. 30
SEM photograph (30,000 magnifications) of the complex oxide obtained in Example 21
Fig. 31

Particle size distribution of the complex oxide
obtained in Example 21
Fig. 32

XRD of the complex oxide obtained in Comparative Example 1
Fig. 33

SEM photograph (1,000 magnifications) of the complex oxide obtained in Comparative Example 1

Fig. 34

SEM photograph (10,000 magnifications) of the complex oxide obtained in Comparative Example 1
Fig. 35

Particle size distribution of the complex oxide obtained in Comparative Example 1

[Diagram showing particle size distribution with cumulative particle amount on the x-axis and amount of particle on the y-axis.]
Fig. 36

XRD of the complex oxide obtained in Comparative Example 5
Fig. 37
SEM photograph (3,500 magnifications) of the complex oxide obtained in Comparative Example 5

Fig. 38
SEM photograph (10,000 magnifications) of the complex oxide obtained in Comparative Example 5
Fig. 39
XRD of the complex oxide obtained in Comparative Example 15
Fig. 40

SEM photograph (20,000 magnifications) of the complex oxide obtained in Comparative Example 15