Effects of cationic substitution on structural defects in layered cathode materials LiNiO₂

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The electrochemical properties of layered rock salt cathode materials are strongly influenced by defects. The three most common defects in LiNiO₂-based compounds, namely extra Ni, Li–Ni anti-site and oxygen vacancy defects have been investigated. The calculated defect formation energies are very low in LiNiO₂, consistent with the difficulty in synthesizing stoichiometric defect-free LiNiO₂. A systematic study is conducted to examine the effect of Co, Mn and Al substitution on defect formation. It is shown that the presence of Ni²⁺ in the Li layer can be rationalized using ideas of superexchange interactions. In addition, a correlation between oxygen vacancy formation energy and oxygen charge is noted. This explains the better thermal stability obtained by early transition metal or Al substitutions.

Oxygen loss is another issue in layered cathode materials. A recent study on LiNi₀.₅Mn₀.₅O₂ demonstrated that up to ~12% oxygen loss occurs depending on the synthesis conditions and that there is a strong correlation between oxygen content and electrochemical performance.³¹ Delithiated Li₁₉NiO₂ is not thermally stable. It undergoes a phase transition accompanied by oxygen evolution. It has been shown that the extent of oxygen evolution increases as x increases.³² This irreversible structural change is concomitant with oxygen loss and maybe responsible for the observed capacity fading.³³,³⁴

In order to improve the electrochemical performance of LiNiO₂, the strategy of partial substitution of Ni by other metal cations has been deployed. It is known that Co substitution gives better 2-D layered character. For LiNi₁₋ₓCoₓO₂ with x > 0.3, nickel is no longer present in the lithium layer.³⁵ As a result, the irreversibility seen at the first-cycle mentioned above disappears. By contrast, the interlayer mixing increases with Mn doping.³⁶ Nevertheless, LiNi₀.₅Mn₀.₅O₂ exhibits excellent structural stability against oxygen loss at low Li content and therefore better safety. Al doping improves the thermal stability although Ni is still found in the Li layer.³⁷,³⁸,³⁹ Cycling tests show that 10% Al suppresses all the phase transitions observed for the Li₉Ni₉O₂ system.³⁹

Although the properties produced by partial cationic substitution are well studied, the reasons why these foreign dopants produce them are not clear. In this study, first-principles calculations are performed to investigate the structural defects of Li–Ni anti-site, extra Ni and oxygen vacancy in LiNiO₂ and the effect of Ni substitution by Co (LiNi₀.₅Co₀.₅O₂), Al (LiNi₀.₅Al₀.₅O₂) and Mn (LiNi₀.₅Mn₀.₅O₂). The same structural defects in NaNiO₂ and LiCoO₂ are also calculated for comparison.

Defect formation energies

In this study, we consider the presence of extra Ni in the Li layers, the Li–Ni interlayer mixing and the oxygen loss as point

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defects in the supposedly perfect layered LiMO₂. The extra Ni defect can be considered as occurring through the following defect reaction

\[ \text{LiMO}_2^- + 2\text{Ni} \rightarrow 2\text{Ni}_\text{li} + 2\text{M}_\text{M} + \text{Li}_2\text{O} + \frac{1}{2}\text{O}_2 \]

Similarly the interlayer mixing defect occurs through the reaction

\[ \text{LiMO}_2^- \rightarrow \text{Ni}_\text{li} + \text{M}_\text{M} \]

and the oxygen vacancy defect occurs through the reaction.

\[ \text{LiMO}_2^- \rightarrow \text{Vo} + 2\text{M}_\text{M}' + \frac{1}{2}\text{O}_2 \]

In this work we define defect formation energies as the formation enthalpies of the above defect reactions at 0 K. Two assumptions are made here. First, in solid phases the volume term can be neglected and therefore the enthalpy corresponds to the internal energy. Second, defects are assumed to distribute evenly in the crystal. In the case of extra Ni defects in LiNiO₂, the defect formation energy per defect is then given as

\[
\text{DFE(extra Ni)} = \Delta G = \Delta H - T\Delta S = -E(\text{perfect}) - E(\text{NiO}) + E(\text{defective}) + \frac{1}{2}E(\text{Li}_2\text{O}) + \frac{1}{4}E(\text{O}_2)
\]

where \(E(\text{perfect})\) is the lattice energy of a perfect LiNiO₂ cell and \(E(\text{defective})\) is the lattice energy of the cell containing one extra Ni defect.

Similarly the defect formation energy of interlayer mixing is

\[
E(\text{interlayer mixing}) = -E(\text{perfect}) + E(\text{defective})
\]

where \(E(\text{perfect})\) is the lattice energy of the perfect cell and \(E(\text{defective})\) is the lattice energy of the cell containing one interlayer mixing defect.

The defect formation energy of oxygen vacancy is

\[
E(\text{oxygen vacancy}) = -E(\text{perfect}) + E(\text{defective}) + \frac{1}{2}E(\text{O}_2)
\]

where \(E(\text{perfect})\) is the lattice energy of the perfect cell and \(E(\text{defective})\) is the lattice energy of the cell containing one oxygen vacancy.

For LiNiO₂, the concentrations of Ni present in the Li layers and oxygen vacancies are reported to be a few percent and are far beyond the dilute limit. However by an appropriate choice of the supercell size, the correct defect concentration can be simulated by substitution of the appropriate number of defects in the cell.

**Computational approach**

In this study, first-principles calculations are performed to investigate the structural defects of interlayer mixing, extra Ni and oxygen vacancy in layered LiNiO₂ and the effect of Ni substitution by Co (LiNi₀.₅Co₀.₅O₂), Al (LiNi₀.₅Al₀.₅O₂) and Mn (LiNi₀.₃Mn₀.₇O₂). The structural defects in layered NaNiO₂ and LiCoO₂ are also calculated for comparison.

All calculations are based on Density Functional Theory (DFT) in combination with the projector augmented wave (PAW) method. The generalized gradient approximation (GGA) is used with the Perdew–Burke–Ernzerhof functional and a Hubbard model U correction is incorporated for the d electrons to give a better description of this strongly correlated system. The U parameters used for Ni, Co and Mn are 6.5, 4.9 and 4.5 eV, respectively. These parameters are adapted from a self-consistent calculation. The plane wave energy-cutoff is set to 500 eV. For all cells, the \(k\)-point spacing is less than 0.05 Å\(^{-1}\) in the Brillouin zone. Structural optimizations were performed until the residual force acting on each ion was less than 0.01 eV Å\(^{-1}\). All calculations were carried out using the Vienna *ab initio* simulation package (VASP).

For the calculation of perfect layered LiNiO₂, a possible ground state cell with space group symmetry \(P2₁/c\) is used as the starting structure. In this \(P2₁/c\) cell the Jahn–Teller distortions of Ni\(^{3+}\) in the NiO₂ slab are in a zigzag ordering. For calculations of layered LiNi₀.₅Co₀.₅O₂, LiNi₀.₅Al₀.₅O₂ and LiNi₀.₃Mn₀.₇O₂ the two simplest in-plane cation orderings, linear and zigzag orderings, are considered as shown in Fig. 1. Supercells with 32 formula units containing 128 atoms are used in all defect calculations. The interlayer mixing defects and extra Ni defects in such a supercell correspond to a concentration of 3.125%, which is well within the experimentally reported range of defect concentration in LiNiO₂. Therefore the size of the cell is adequate for simulating the observed defects in LiNiO₂ and there is no need for extrapolation to the infinite limit.

The interlayer mixing defect in layered AMO₂ (A = Li, Na) is constructed by swapping one Ni (Co in the LiCoO₂ case) in the MO₂ slab with the nearest A ion. The extra Ni defect is constructed by replacing one A ion by Ni in the supercell. The oxygen vacancy defect is constructed by removing one oxygen atom from the supercell.

**Results and discussion**

**Properties of undoped compounds**

Before proceeding to the defect structure calculations, the crystal and electronic structures of perfect LiNi₀.₅Co₀.₅O₂, LiNi₀.₅Al₀.₅O₂ and LiNi₀.₃Mn₀.₇O₂ are first determined. In

![Fig. 1](a) Linear and (b) zigzag ordering of cations. Red denotes oxygen, blue and yellow denote the two different cations. Lithium is omitted for clarity.)
LiNi$_{0.5}$Co$_{0.5}$O$_2$ and LiNi$_{0.5}$Al$_{0.5}$O$_2$ the linear ordering of cations with space group symmetry $P2_1/m$ is found to be more energetically favorable than the zigzag ordering and is therefore used for subsequent defect calculations. In LiNi$_{0.5}$Mn$_{0.5}$O$_2$, the zigzag ordering of Ni and Mn with space group symmetry $P2_1/c$ is energetically more favorable, in agreement with a previous theoretical study.$^{25}$

Fig. 2 shows the calculated density of states for each material. The insulating behaviour of these compounds is well reproduced with band gaps of about 0.7 eV, 0.9 eV and 1.1 eV for LiNi$_{0.5}$Co$_{0.5}$O$_2$, LiNi$_{0.5}$Al$_{0.5}$O$_2$ and LiNi$_{0.5}$Mn$_{0.5}$O$_2$, respectively. The local density of states (DOS) of Ni in LiNi$_{0.5}$Co$_{0.5}$O$_2$ shows one empty spin-up and two empty spin-down states which indicates that the electronic configuration of Ni is $t_{2g}^6e_{g}^1(S=1/2)$.
which is low-spin Ni$^{3+}$, in accordance with the calculated magnetic moment 1.12 $\mu_B$. A Jahn–Teller distortion occurs as expected for low-spin Ni$^{3+}$ as shown from the Ni–O bond lengths in Table 1. Cobalt ions are therefore anticipated to be Co$^{3+}$ for spin-down eg states seen in the local density of states of nickel in the isotropic octahedral environment with 6 identical Co$^{3+}$ of states of manganese indicate that their electronic con- temporary configuration is $t_{2g}e_g^0$ ($S = 0$) implying low-spin Co$^{3+}$, along with its calculated zero magnetic moment. Likewise, nickel ions are determined to be low spin Ni$^{2+}$ in LiNi$_{0.5}$Al$_{0.5}$O$_2$ with a Jahn–Teller distortion. Nevertheless, from the Ni$^{3+}$–O bond lengths in Table 1, it is clear that Ni$^{3+}$ displays two different modes of Jahn–Teller distortion, Q$_3$ and Q$_1$ in LiNi$_{0.5}$Co$_{0.5}$O$_2$ and LiNi$_{0.5}$Al$_{0.5}$O$_2$ respectively, as shown in Fig. 3. The Q$_3$ mode of Jahn–Teller distortion is the one observed in LiNiO$_2$. In LiNi$_{0.5}$Co$_{0.5}$O$_2$, the low-spin Co$^{3+}$ ions are very stable in the isotropic octahedral environment with 6 identical Co$^{3+}$–O$^2$ bond lengths. The structural constraint imposed by the presence of rigid Co$^{3+}$ octahedra makes the more distorted Q$_3$ mode less favourable and results in the Q$_2$ mode for distorted Ni$^{3+}$. This result is in agreement with an EXAFS study that in LiNi$_{1-x}$Co$_x$O$_2$ the Jahn–Teller distortion of NiO$_6$ octahedra is suppressed with increasing x.$^{27}$ In LiNi$_{0.5}$Mn$_{0.5}$O$_2$, two empty spin-down $e_g$ states seen in the local density of states of nickel and fully-occupied spin-down $t_{2g}$ states seen in the local density of states of manganese indicate that their electronic configurations are $t_{2g}e_g^0$ and $t_{2g}e_g^2$ corresponding to Ni$^{2+}$ and Mn$^{3+}$, in agreement with previously reported results.$^{28}$

### Influence of defect formation on cation charge state

In all LiMO$_2$ cells with extra Ni and interlayer mixing defects, the calculated magnetic moment of 1.7 $\mu_B$ ($S = 1$) for the Ni present in the Li layer along with its average Ni–O bond length 2.07 Å together imply that Nickel is present as Ni$^{2+}$. Therefore in the LiNiO$_2$, LiNi$_{0.5}$Co$_{0.5}$O$_2$ and LiNi$_{0.5}$Al$_{0.5}$O$_2$ cells with the interlayer mixing defect, in order to retain charge neutrality one Ni in the NiO$_2$ slab is oxidised from Ni$^{3+}$ to Ni$^{4+}$ with a calculated magnetic moment 0.19 $\mu_B$ ($S = 0$), as seen in the spin density contour map in Fig. 4(a), and the average Ni–O bond length of 1.89 Å. In the LiNi$_{0.5}$Mn$_{0.5}$O$_2$ cell, the interlayer mixing defect does not cause any change of charge state as nickel ions are already Ni$^{3+}$.

In cells with the extra Ni defect, since one Li$^+$ is replaced by Ni$^{2+}$, one metal ion in the MO$_2$ layer must be reduced to maintain charge neutrality. In the LiNiO$_2$, NaNiO$_2$, LiNi$_{0.5}$Co$_{0.5}$O$_2$ and LiNi$_{0.5}$Al$_{0.5}$O$_2$ cells, it is the Ni in the MO$_2$ layer that gets reduced from Ni$^{3+}$ to Ni$^{2+}$ with a calculated magnetic moment of 1.7 $\mu_B$ ($S = 0$) and an average Ni–O bond length of 2.07 Å. The change of preferred charge state on Ni rather than Co in LiNi$_{0.5}$Co$_{0.5}$O$_2$ is probably due to the relatively stable electronic configuration of Co$^{3+}$ ($t_{2g}e_g^0$). In LiNi$_{0.5}$Mn$_{0.5}$O$_2$, the charge state of Ni$^{2+}$ cannot be reduced further and therefore the charge compensation accompanied by the extra Ni defect takes place on manganese with Mn$^{4+}$ → Mn$^{3+}$. Fig. 4(b) shows the case of the extra-Ni defect in LiNi$_{0.5}$Mn$_{0.5}$O$_2$. The $e_g$ orbital character on Ni$^{2+}$ ($t_{2g}^0e_g$) can be seen from the shape of spin density pointing towards oxygen ions. Similarly, the spin density on Mn$^{4+}$ ($t_{2g}^1e_g$) pointing away from the oxygen represents the $t_{2g}$ orbital character. The Mn ion showing the different shape of spin density is the one that is reduced from Mn$^{4+}$ to Mn$^{3+}$.

For cells with the oxygen vacancy defect, two metal ions in the MO$_2$ layer next to the oxygen vacancy site are reduced to keep the charge neutrality. Fig. 4(c) clearly shows that two Co$^{3+}$ ions are reduced to Co$^{2+}$ upon the removal of one oxygen ion.

### Stability of defects and the effect of cation substitution

The calculated defect formation energies in LiNiO$_2$ are shown in Fig. 5. The calculated formation energies of the three defects in LiNiO$_2$ are all small, ranging from approximately 0.3 to 1.0 eV. This is consistent with the difficulty in synthesizing stoichiometric defect-free LiNiO$_2$. It is possible to rationalize these results using the idea of superexchange interactions.$^{29-31}$ Both Ni$^{2+}$ ($t_{2g}e_g$) and Ni$^{3+}$ ($e_g^0e_g$) have fully filled $t_{2g}$ states but partially filled $e_g$ states. Consequently the 180° Ni–O–Ni superexchange is much stronger than the 90° Ni–O–Ni superexchange plus direct exchange. This means there is a larger energy gain through orbital interactions when Ni–O–Ni is in the 180° configuration than the 90° configuration. In fact it has been shown that there is a tendency for Ni ions to locate as second-nearest neighbors (180° Ni–O–Ni configuration) in the cation sublattice of the rocksalt structure due to the energy gain from the 180° superexchange interaction.$^{32}$ Therefore the presence of Ni ions in the Li layers of LiNiO$_2$ can be viewed as being stabilized by the 180° Ni–O–Ni superexchange interaction, giving rise to low formation energies for both interlayer mixing and the extra Ni defects. However, the extra Ni defect is the most
favorable and therefore is the predominant defect species in LiNiO₂.

In LiNi₀.₅Co₀.₅O₂ and LiNi₀.₅Al₀.₅O₂, due to the linear cation ordering in the transition metal plane, there are two inequivalent Li sites on which to place the Ni in the interlayer mixing and the extra Ni defects, as shown in Fig. 6. These are referred to as configurations A and B. In LiNi₀.₅Mn₀.₅O₂, the zigzag ordering of Ni and Mn also results in two inequivalent Li sites referred to as A and B. Similarly in the cells of LiNi₀.₅Co₀.₅O₂, LiNi₀.₅Al₀.₅O₂ and LiNi₀.₅Mn₀.₅O₂, there are two inequivalent oxygen ions, one bonding with two Ni and one bonding with one Ni, in the cell, which can be removed to create the oxygen vacancy. We refer to the removal of the oxygen bonded to two Ni as configuration A and the removal of the oxygen bonded to only one Ni as configuration B.

The effect of Co substitution can be seen in Fig. 5(a). It is first noted that the defect formation energies of the interlayer mixing and the extra Ni defects in LiNi₀.₅Co₀.₅O₂ with configuration A are lower than in LiNiO₂. This is unexpected since it is known experimentally that Co substitution in LiNiO₂ suppresses the presence of Ni in the Li layer. Nevertheless the formation energies of the interlayer mixing and extra Ni defects are higher in configuration B than configuration A by about 300 meV and 360 meV respectively. The result can also be rationalized by considering superexchange interactions. As seen in Fig. 6(a), in configuration A, the Ni in the Li layer forms six 180° Ni–O–Ni chains. In configuration B, the six 2nd-nearest-neighbours in the cation sublattice, of the Ni in the Li layer are Co³⁺, forming 180° Ni–O–Co chains which do not give rise to the 180° superexchange interaction due to the empty eg orbitals of Co³⁺ (t^6_2g^0_e^0_g).
cobalt substitution would be to screen the superexchange interaction. This is different from a previous proposed size effect, and destabilizes the presence of Ni in the Li layer.

This again suggests that the presence of Ni\textsuperscript{2+} in the Li layer is stabilized by the 180° Ni–O–Ni superexchange interaction. The higher number of the 180° Ni–O–Ni chains gives rise to the lower energy. In the real LiNi\textsubscript{0.5}CoO\textsubscript{2} compound, the Co\textsuperscript{3+} ions distribute randomly in the MO\textsubscript{2} slab and the main effect of (a) Co and (b) Al substitution on the calculated defect formation energies.

Since there are no d electrons in the Al\textsuperscript{3+} ion, there can be no superexchange interaction between Al\textsuperscript{3+} and Ni\textsuperscript{2+}. The effect of Al substitution on defect formation energies is therefore expected to be similar to that of Co substitution since Al substitution should also effectively screen the Ni–O–Ni superexchange interaction. Indeed by adopting the linear cation ordering in the LiNi\textsubscript{0.5}Al\textsubscript{0.5}O\textsubscript{2} cell (Fig. 1(a)), as shown in Fig. 5(b) the calculated formation energies of the interlayer mixing and the extra Ni defects are very similar to those in LiNi\textsubscript{0.5}CoO\textsubscript{2}. Defects of configuration A are also more favourable than configuration B due to the stabilisation by the exchange interaction. However, unlike in LiNi\textsubscript{1−x}Co\textsubscript{x}O\textsubscript{2} with x > 0.3, neither the interlayer mixing defect nor extra Ni defects are observed. Experimentally 5% of extra-nickel ions are still found in the lithium layer in LiNi\textsubscript{1−x}Al\textsubscript{x}O\textsubscript{2} with 0.1 < x < 0.5. This is because Al tends to segregate to interfaces\textsuperscript{34} and hence a core–shell structure can be formed\textsuperscript{34} in LiNi\textsubscript{0.5}Al\textsubscript{0.5}O\textsubscript{2}. Consequently, the extra-Ni and Li–Ni anti-site defects can still occur in Ni-rich domains in LiNi\textsubscript{0.5}Al\textsubscript{0.5}O\textsubscript{2} as in LiNiO\textsubscript{2}, where the presence of Ni in the Li layer can be stabilized by the 180° Ni–O–Ni exchange interaction.

As shown by Fig. 7(a) there is no significant difference in configurations A and B for the formation energy of the interlayer mixing and the extra Ni defects in LiNi\textsubscript{0.5}Mn\textsubscript{0.5}O\textsubscript{2}. The formation energy of the interlayer mixing is markedly lower than that of the extra-Ni defect. This is consistent with the experimentally observed high concentration of interlayered mixing defects in LiNi\textsubscript{0.5}Mn\textsubscript{0.5}O\textsubscript{2}. Also the formation energy of the interlayer mixing defect is lower by about 0.3 eV than that of LiNiO\textsubscript{2}. Unlike in LiNi\textsubscript{0.5}Co\textsubscript{0.5}O\textsubscript{2} and LiNi\textsubscript{0.5}Al\textsubscript{0.5}O\textsubscript{2} where the 180° Ni–O–Co and 180° Ni–O–Al interactions are absent, in LiNi\textsubscript{0.5}Mn\textsubscript{0.5}O\textsubscript{2} the electronic configuration of Mn\textsuperscript{4+} (t\textsubscript{2g}\textsuperscript{6}) could give rise to moderate 180° Ni\textsuperscript{2+}–O\textsuperscript{2−}–Mn\textsuperscript{4+} interactions.\textsuperscript{35} Consequently, although the number of 180° Ni–O–Ni interactions is reduced due to Mn substitution, the presence of Ni\textsuperscript{2+} can be stabilized not only by the 180° Ni–O–Ni interaction but also by the 180° Ni\textsuperscript{2+}–O\textsuperscript{2−}–Mn\textsuperscript{4+} interaction. Moreover, since the ionic radius of Ni\textsuperscript{2+} is similar to Li\textsuperscript{+}, these ions can exchange sites readily without significant rearrangement of the surrounding atomic positions. No charge compensation is necessary to create the interlayer mixing defect in LiNi\textsubscript{0.5}Mn\textsubscript{0.5}O\textsubscript{2}. In contrast to the interlayer mixing defect, the defect formation energy for the extra Ni defect is much higher in LiNi\textsubscript{0.5}Mn\textsubscript{0.5}O\textsubscript{2} compared to LiNiO\textsubscript{2}. The probable reason for this is the reduction of Mn\textsuperscript{4+} to Mn\textsuperscript{3+}, which is the charge compensation accompanying the extra-Ni defect, is considerably less favorable than the reduction of Ni\textsuperscript{3+} to Ni\textsuperscript{2+} due to the stable electronic configuration of Mn\textsuperscript{4+} (t\textsubscript{2g}\textsuperscript{6}).

In NaNiO\textsubscript{2}, there is a structural constraint arising from the large ionic size of Na\textsuperscript{+}. It is shown in Table 2 that the LiO\textsubscript{6} octahedron must undergo significant distortion for the zigzag ordering of the Ni\textsuperscript{3+} Jahn–Teller distortions or charge
disproportionation \( \text{Ni}^{3+} \rightarrow \text{Ni}^{2+} + \text{Ni}^{4+} \) in the NiO\(_2\) layer to happen. However, the larger Na\(^+\) ion fills up the interlayer space and so forbids the zigzag ordering of the Ni\(^{3+}\) Jahn–Teller distortions or charge disproportionation \( \text{Ni}^{3+} \rightarrow \text{Ni}^{2+} + \text{Ni}^{4+} \) in the NiO\(_2\) layer. Hence the Ni\(^{3+}\) Jahn–Teller distortions in NaNiO\(_2\) are forced to align collinearly as observed experimentally, which results in undistorted NaO\(_6\) octahedra. This gives a good 2-D layered character and is less susceptible to defects as shown by the high defect formation energies for NaNiO\(_2\) compared to LiNiO\(_2\) in Fig. 7(b). Because of the dramatic difference in ionic radii between Na and first-row transition metal ions, the size effect dominates the interactions between cations and consequently all NaMO\(_2\) form perfect layered structures.

**Oxygen vacancy**

Fig. 8 shows the calculated defect formation energies of the oxygen vacancy plotted against the oxygen charge calculated using the Bader analysis.\(^{35-36}\) The formal charge on oxygen is \(-2\) in highly ionic compounds. However, in transition metal oxides, there is a considerable overlap between the oxygen 2p and metal 3d orbitals, particularly for late transition metals or metals with high charge states. This is reflected in the calculated oxygen charge as shown in Fig. 8, from left to right (LiAlO\(_2\) → LiNiO\(_2\) and LiCoO\(_2\) → Li\(_{0.5}\)CoO\(_2\)) the decrease of calculated oxygen charge is a consequence of the increase in overlap between oxygen 2p and metal ion 3d orbitals or equivalently greater metal–oxygen covalency. A correlation can be clearly seen between the formation energy of the oxygen vacancy defect and the calculated oxygen charge. Also, as shown in Fig. 5 and 7, in LiNi\(_{0.5}\)Co\(_{0.5}\)O\(_2\), LiNi\(_{0.5}\)Al\(_{0.5}\)O\(_2\) and LiNi\(_{0.5}\)Mn\(_{0.5}\)O\(_2\), the defect formation energy for removing the oxygen bonded to two Ni (configuration A) is lower than the oxygen bonded to one Ni (configuration B). The oxygen bonded to two Ni has a lower charge. The smaller the oxygen charge is, the easier it seems to be to remove the oxygen. It has previously been suggested that the strength of the metal–oxygen bond depends on the effective charge on oxygen.\(^{37}\) In addition when the charge on oxygen ions is low, there would be a tendency for them to form peroxide at the surface as suggested by Goodenough et al.\(^{38}\) and then dissociate through the following reaction:

\[
2(O_2)^{2-} = 2O_2^{2-} + O_2
\]

This is consistent with experimental results that the temperature for oxygen evolution on heating (i.e. the thermal stability) decreases as \(x\) decreases in layered Li\(_x\)MO\(_2\).\(^{39-41}\) It seems that low oxygen charge/high metal–oxygen covalency causes the chemical instability of an oxide compound against oxygen loss. A recent study has also proposed that a greater metal–oxygen covalency promotes the surface oxygen evolution reaction which involves the creation of surface oxygen vacancies.\(^{42}\)

On comparing LiNiO\(_2\) with LiCoO\(_2\) there is no noticeable difference in the oxygen charge, but the defect formation energy

| Structure                        | Li–O bond lengths (Å) |
|----------------------------------|------------------------|
| \(C2/m\) (collinear ordering of the Ni\(^{3+}\)) | 2.11 × 4               |
| Jahn–Teller distortions          | 2.13 × 2               |
| \(P2_1/c\) (zigzag ordering of the Ni\(^{3+}\)) | 2.04 × 2               |
| Jahn–Teller distortions          | 2.10 × 2               |
| \(P2/c\) (charge disproportionation) | 2.03 × 2               |
| Ni\(^{3+}\) → Ni\(^{2+}\) + Ni\(^{4+}\) | 2.08 × 2               |
| Ni\(^{3+}\) → Ni\(^{2+}\) + Ni\(^{4+}\) | 2.19 × 2               |

Fig. 7 Calculated defect formation energies in (a) LiNi\(_{0.5}\)Mn\(_{0.5}\) and (b) NaNiO\(_2\) compared to LiNiO\(_2\).

Fig. 8 The correlation between oxygen charge and defect formation energy for an oxygen vacancy for a series of structures.
of the oxygen vacancy in LiCoO₂ is significantly higher than in LiNiO₂ (by ~1.2 eV). This is probably due to the relatively stable electronic configuration of low-spin Co²⁺. Therefore by creating an oxygen vacancy, it costs more energy to reduce Co³⁺ to Co²⁺ than to reduce Ni³⁺ to Ni²⁺ in LiNiO₂. Although the defect formation energy of an oxygen vacancy in LiCoO₂ is markedly higher than in LiNiO₂, it drops drastically by ~1.5 eV in Li₅ₓCoO₄ upon the removal of half the lithium ions. This can again be explained by the decrease of oxygen charge that is associated with the creation of Co²⁺ ions.

Given this correlation between oxygen charge and the defect formation energy of the oxygen vacancy, doping with a more electron-positive cation should mitigate the oxygen loss in layered LiₓMO₂ compounds and result in better thermal stability. Indeed doping with Mn⁴⁺ decreases the oxygen loss in Li[Ni₀.₃₃Mn₀.₆₇₋ₓTiₓ]O₂.

Conclusions

All the calculated formation energies for the various LiₓMO₂ compounds are consistent with experimental results. It is demonstrated that the defect formation energies in LiNiO₂ are low, in agreement with the experimental difficulty of synthesizing stoichiometric defect-free LiNiO₂. The presence of Ni in the Li layer can be rationalized in terms of the 180° Ni–O–Ni superexchange interaction. Substituting Ni with Co in the MO₂ layer screens the 180° Ni–O–Ni configurations and thus effectively reduces the concentration of Ni in Li layers. A correlation between the defect formation energy of the oxygen vacancy and oxygen charge (as measured from a Bader analysis) is reported. It appears that the smaller the oxygen charge/higher metal–oxygen covalency, the lower the oxygen vacancy formation energy. This can explain the thermal instability of LiₓCoO₂ and LiₓNiO₂ at low x, as well as the improved electrochemical behavior in Al, Mg or early transition metal doped LiMO₂. In the quest for designing better cathode materials, the use of high electropositive cations is highly desirable.

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