Introduction

The lithium ion battery utilising a LiCoO$_2$ cathode, with a capacity of 160–190 mA h g$^{-1}$, was the first commercially successful chemistry released to the marketplace. LiCoO$_2$ has a layered structure in which lithium cations lie between anionic sheets containing cobalt and oxygen. Over the subsequent years of research this general structure has proved extremely versatile, with routine substitution of the Co for other transition metals to form new materials that are analogous to LiCoO$_2$. Today, the energy density of Li-ion batteries is limited by the cathode. The state-of-the-art NMC (LiNi$_{1-x}$Co$_x$Mn$_y$O$_2$) and NCA (LiNi$_{1-x}$Co$_x$Al$_y$O$_2$) materials approach 200 mA h g$^{-1}$. However, they are limited by storing electrons only on the transition metals (TM-redox). In recent years it has been shown that so called “Li-rich” cathode materials, e.g. Li$_{1.2}$Ni$_{0.2}$Mn$_{0.6}$O$_2$ and Li$_{1.2}$Ni$_{0.13}$Co$_{0.13}$Mn$_{0.54}$O$_2$, can exceed the limit of TM-redox by invoking O-redox, where electrons are stored on oxygen as well as the transition metal ions.$^{1-6}$ Capacities exceeding 250 mA h g$^{-1}$ are possible, making these materials candidates for future Li-ion batteries. This process of O-redox has been extensively characterised using X-ray absorption near edge structure (XANES), soft X-ray absorption spectroscopy (SXAS), online electrochemical mass spectrometry (OEMS), X-ray photoelectron spectroscopy (XPS), resonant inelastic X-ray scattering (RIXS) and several models exist for the nature of the oxygen redox process.$^{7-10}$ Typically, whilst the oxygen oxidation and reduction processes are reversible, a significant oxygen loss component is seen. It is known that the surface/near surface of Li transition metal oxide cathodes, not only Li-rich, are often composed of rocksalt/spinel-like structures even in the pristine materials.$^{11-17}$ For instance, Koga et al.$^8$ demonstrate a mixture of two phases formed on charging of Li$_{1.20}$Mn$_{0.54}$Co$_{0.13}$Ni$_{0.13}$O$_2$; oxygen participates in different reactions depending on whether it is located in the surface or the bulk phase. It has been proposed that O-loss from Li-rich materials on charging, inherently a surface process, results in densification forming rocksalt/spinel structures at the surface.$^{12-14,18,19}$
The consequence is a core–shell morphology to the particles. Such a morphology has implications for using techniques that are surface sensitive for probing the nature of the O-redox process.

How the electronic structure varies from the surface to the bulk in the materials at various states of charge is not well known. In this work, we use the archetypal Li-rich cathode material Li$_{1.2}$Ni$_{0.2}$Mn$_{0.6}$O$_2$ to show that the balance between O-redox and TM-redox varies significantly from the surface to the bulk and that the surface, likely owing to its greater energy and interface with the electrolyte, appears to exhibit significantly higher degrees of oxygen oxidation. The concentration gradient of oxidised oxygen observed within the particle on charge supports the already well-established model for such materials, in which a rocksalt shell encases a layered core.$^{7,9,11-15,20,21}$ We reach these conclusions through the first use of energy-tuned synchrotron X-ray photoelectron spectroscopy (XPS) with a truly wide range of excitation energies to give the most reliable synchrotron XPS with its unique capabilities to collect spectra from so tender to hard X-rays, as highlighted in Fig. 1a. As a comparison, the use of Al Kz X-ray radiation on a laboratory XPS corresponds to a depth of more than 6 nm which could not be considered surface-sensitive for the purposes of this study. Since the radial dimension of the particles (determined previously$^{19}$) in this study is only double the approximate sampling depth we can consider this technique to be relatively bulk sensitive.

We performed the energy-tuned XPS depth profiling at the 109 beamline at the Diamond Light Source taking advantage of its unique capabilities to collect spectra from soft and hard X-rays on the same sample. This allows us to map gradients in the oxygen and transition metal oxidation states for various states of charge. Additionally, excellent energy resolution can be achieved at the beamline, allowing spectral features to be resolved which may not have been possible using other sources, thus ensuring a reliable fitting of the data. The experimental results have been compared to theoretical models calculated using high level DFT methodology and this has been used to help us rationalise the difference between surface and bulk structures.

Methods

1. Materials synthesis

Li$_{1.2}$Ni$_{0.2}$Mn$_{0.6}$O$_2$. The formaldehyde-resorcinol sol gel synthetic route$^{24}$ was employed to synthesise the layered lithium nickel manganese oxide, Li$_2$Li$_{0.2}$Ni$_{0.2}$Mn$_{0.6}$O$_2$; all the reagent ratios were calculated in order to obtain 0.01 mol of the final product. Stoichiometric amounts of Li$_2$CO$_3$·2H$_2$O (99.0%, Sigma Aldrich), Mn(CH$_3$CO$_2$)$_2$·4H$_2$O (>99.0%, Sigma Aldrich) and H$_2$O (99.8%, VWR) were mixed together in a ground glass bottle to ensure a homogeneous mixture. After being heated at 120 °C for 24 h, a black solid was obtained, which was then dried at 100 °C for 24 h. The black solid was then calcined at 400 °C for 6 h under an Ar atmosphere to form the Li$_2$CO$_3$·2H$_2$O. After being ground and sieved to a size of 75–200 μm, the sample was placed in a glass crucible, which was then heated at 800 °C for 15 h under a flowing Ar atmosphere to obtain the final product. The final product was then ground and sieved to a size of 75–200 μm.

Fig. 1 Schematic diagram representing the principles of energy-tuned X-ray photoelectron spectroscopy measurements (a) for cathodes removed from cells at various voltages along the charge–discharge curve (b). Photoelectrons of variable kinetic energy are ejected as a result of excitation by monochromatic light of different energies thus probing different depths of an active material particle. Refer to the ESI† (Note 1) for an explanation of calculated depths of origin. Red-coloured markers on the voltage profile of Li$_{1.2}$Ni$_{0.2}$Mn$_{0.6}$O$_2$ represent points at which samples were prepared for photoelectron spectroscopy measurements (pristine, charged to 4.1, 4.55, 4.65, and 4.8 V, discharged to 2 V vs. Li/Li$^{+}$).
and Ni(CH₂CO₂)₂·4H₂O (99.0%, Sigma Aldrich) were dissolved in 50 mL of water with 0.25 mmol of Li₂CO₃ (99.0%, Sigma Aldrich). At the same time 0.1 mol of resorcinol (99.0%, Sigma Aldrich) was dissolved in 0.15 mol of formaldehyde (36.5% w/w solution in water, Fluka). Once all the reagents were completely dissolved in their respective solvents, the two solutions were mixed and the mixture was vigorously stirred for one hour until the colour changed from light green to light brown. The resulting solution, containing 5% molar excess of lithium, was subsequently heated in an oil bath at 80 °C until formation of a homogeneous white gel. The gel was finally dried following a three-step process: 90 °C overnight, 500 °C for 15 hours and 800 °C for 20 hours. The obtained powder was very fine with a dark-brown colour. Composition and structure was verified as described in our previous study.³⁰

LiNi₀.₅Mn₁.₅O₄. The spinel phase material was synthesised using a resorcinol-formaldehyde sol gel method.³⁴ A 1 g batch was prepared by dissolving stoichiometric amounts of LiCH₃COO·2H₂O (99.0%, Sigma-Aldrich), Ni(CH₂CO₂)₂·4H₂O (99.0%, Sigma-Aldrich), and Mn(CH₂CO₂)₂·4H₂O (99.0%, Sigma-Aldrich) in 50 mL of distilled water containing 0.1 mol of resorcinol (99.0%, Fluka), 0.15 mol of formaldehyde (35%, containing 10% methanol as stabilizer, Sigma-Aldrich), and 0.25 mmol of Li₂CO₃ (99.9%, Sigma-Aldrich). The solution was heated at 90 °C under stirring until a gel was formed. The gel was dried and decomposed at 500 °C for 10 h, and the resulting powder was ground using a mortar and pestle. Finally, the powder was calcined at 900 °C for 15 h to obtain the final material, which had a similar particle size to the Li-rich material. All synthesis steps were carried out in air.

2. Electrode preparation

The cathode electrodes were prepared in the form of pellets composed of a mixture of active material, carbon Super P (Timcal) and polyethylene tetrafluoride (PTFE) in an 8 : 1 : 1 mass ratio, respectively. Once weighed in the correct ratio the powder mixture was homogenised with a mortar and pestle. Subsequently, 0.5 mL of isopropyl alcohol (99.5%, Sigma Aldrich) was added drop by drop to allow for a better homogeneity of the components distribution within the pellet. The wet powders were continuously kneaded in the mortar until all the IPA was evaporated and a uniform and compact pellet was attained. The pellet was then roll pressed down to a thickness of 125 μm and cut into small pieces that were dried overnight under dynamic vacuum at 80 °C before being assembled into coin cells.

3. Cell preparation

After being completely dried the electrodes were assembled into coin cells from CR2325 coin-cells parts, provided by the National Research Council of Canada, under an inert argon atmosphere inside a glove box (MBraun LABmaster sp, O₂ and H₂O < 0.1 ppm). A lithium metal disc (Ø = 18 mm, 0.1 mm thick) was used as counter electrode with one Celgard and two glass fibre discs (Ø = 22 mm, Whatman GF-D) as separator, soaked with 0.2 mL of LP30 Selectilyte electrolyte (1 M LiPF₆ in 1 : 1 w/w ethylene carbonate : dimethyl carbonate, provided by BASF).

4. Electrochemical cycling

Cells were cycled on a BioLogic VMP3 potentiostat/galvanostat at 30 °C in a thermostatically controlled oven. A constant current corresponding to 50 mA g⁻¹ was applied to the cells. The electrochemical behaviour of the materials is highly reproducible and the cells from which the samples were taken were cycled as expected.

5. X-ray photoelectron spectroscopy

Cells were disassembled under an inert argon atmosphere inside a glovebox (O₂ and H₂O < 0.1 ppm) and the cathodes washed in dry dimethyl carbonate (<10 ppm H₂O) three times before being dried under vacuum for 5 minutes. Samples were mounted on copper plates using copper tape and loaded into a UHV transfer suitcase to be transported from the glovebox (at the home laboratory in Oxford) to the synchrotron. Hard X-ray photoelectron spectroscopy (HAXPES) and soft X-ray photoelectron spectroscopy (SOXPES) were performed at the Surface and Interface Structural Analysis beamline (I09) at Diamond Light Source (Oxfordshire, UK). In the hard X-ray regime, excitation energies of 2350 eV and the 3rd order light 7050 eV were used on all samples. To minimize radiation damage of the samples and to probe a large area of the sample to reduce the possibility of heterogeneity between particles, the beam was spread out using defocused settings; the spot at the sample is estimated to be approximately 300 μm (H) and up to 1 mm (W). Photons were monochromatized by a Si(111) double-crystal monochromator. In the soft X-ray regime, 1090 eV was used for more surface sensitive measurements. This branch of the beam line uses a plane grating monochromator. No charge neutralizer was used during the measurements. The spectra were recorded using a hemispherical VG Scienta EW4000 analyser set to a pass energy of 50 eV for soft X-rays and 200 eV for hard X-rays. Binding energies are calibrated to the C=C peak, from the carbon black in the electrodes, at 284.0 eV in the C 1s spectra (see ESI, Note 3). The software package CasaXPS was used for analysis of photoelectron spectroscopy data and the Gaussian/Lorentzian peak shape GL(30) was used throughout the fitting. Replicate measurements on another separate set of samples demonstrated to us that the results and trends observed are reproducible.

6. Ab initio simulations

All calculations were performed using spin-polarised density functional theory (DFT) implemented in the Vienna ab initio simulation package (VASP).³² The Heyd–Scuseria–Ernzerhof (HSE06) hybrid functional³³ was used to describe the electron exchange and correlation, with a hybrid mixing parameter of 0.2. It was shown that the application of this functional is necessary for the accurate description of the oxygen hole states³⁴–³⁶ and redox activity.³⁷–³⁹ The projector augmented plane-wave method (PAW)⁴⁰ was used for the interactions between
valence and core electrons, while the valence electrons are described as plane waves with a cut-off energy of 520 eV. The k-space was sampled with an automatically generated k-point mesh using a spacing smaller than 0.05 Å⁻¹. All atomic coordinates and lattice parameters were allowed to fully relax until the residual forces were smaller than 0.05 eV Å⁻¹. Similar computational techniques have been applied in successful studies of other electrode materials.29–45

The structure of the Li-excess material was simulated with the composition Li₁.₁₇⁻Ni₀.₃₅Mn₀.₃₈O₂. The 2 × 1 × 2 supercell of Li₂MnO₃ was modified to contain 24 formula units of layered Li₁.₁₇⁻Ni₀.₃₅Mn₀.₃₈O₂, while the cation disordered rocksalt was created based on an Fm̅m̅m LiMnO₂ structure containing 12 formula units. The cation ordering within the pristine materials and the Li⁺ arrangement within the delithiated structures of the layered oxide were generated following the methodology described by Seo et al.4 and using the Python Materials Genomics tool (pymatgen).46 Core level binding energies (E_{cl}) of the O 1s electrons were evaluated with the initial state approximation and shifted with respect to a reference system (Li₂MnO₃ [ref. 51]) into the experimentally observed energy region.46–49 Full description of the applied methodology is in the ESI† (Note 10). It is important to note that the calculated core level energies are not suited for predictions of absolute values. Nevertheless, since the simulated core level shifts are accurate, these calculations assist in the interpretation of the experimental data by differentiating between types of oxygen species and environments at an atomistic level.

Results

7. Photoelectron spectra of Li-rich Li₁.₅Ni₀.₂Mn₀.₆O₂

Fig. 1b shows the voltage profiles for the Li₁.₂Ni₀.₃Mn₀.₆O₂ cathode material studied. The red points indicate the voltages of samples analysed by photoelectron spectroscopy. The samples include the pristine electrode, charged to 4.1 V (during Ni oxidation), charged to 4.55 V (beyond the onset voltage typically associated with oxygen oxidation, and after Ni oxidation),25,51 charged to 4.65 V (further along the plateau), the top charge of 4.8 V, and discharged to 2 V.

O 1s. Photoelectron spectra were collected at three different photon energies from surface sensitive soft X-ray up to hard X-ray energies. Fig. 2 shows the fitted O 1s spectra for pristine, charged and discharged electrodes for the three analysis depths. In the pristine material with the lowest excitation energy, several peaks have been used in the fitting of the spectrum. The complete details of the fitting procedure are given in the ESI† (Note 2). The lowest energy peak (≈529 eV, green in figure) is a typical response for oxygen in a transition metal oxide. At higher binding energies two additional peaks are used (≈531.4 and ≈532.7 eV, grey); these peaks represent surface-bound species such as oxygen in carbonates and hydroxides. As the excitation energy is increased from 1.09 to 2.35 and then to 7.05 keV, the signal for the surface-bound species reduces significantly, confirming their presence only at the very surface of the particle as terminating species or a sparse coverage. The position of these peaks is constant throughout the series of spectra, as there is no indication to show that these species undergo any chemical change; for example, if there was significant electrolyte oxidation at the surface during charge then we would also expect to see changes in the C 1s spectra, which we do not (see ESI†, Note 3). At the highest photon energy of 7.05 keV, it is clear that the majority of the ejected electrons are from the lattice oxygens in the bulk material, with little response observed from the surface-bound species.

A positive shift (~0.15–0.3 eV) of the transition metal oxide peak (O₂⁺) is observed in the surface spectra, compared with the bulk, at all voltages. This shift has been determined to be a true chemical effect rather than a result of charging or an instrumental artefact (see ESI†, Note 4). Such a small shift represents the slightly more oxidised environment of oxygens (O₁⁻/₂⁻) existing in the near-surface region or under-coordinated oxygens at the very surface making up the various facets of the particle. It is also possible for phases other than the layered structure to exist at some surface facets, as has been reported previously, even for pristine materials.11–17 For example, in the disordered rocksalt structure, cation mixing occurs between the transition metal and lithium layers, forming a greater number of Li–O–Li bonds than in the layered structure. This greater coordination of oxygen by lithium ions results in a larger degree of ionic bonding and hence less electron density on oxygen. This leads to the positive shift of the O²⁻ peak observed at the surface.

On charge, a shift to a lower binding energy (~528.8 eV) of the oxide (O²⁻) peak for all excitation energies can be attributed to the removal of Li⁺ ions from around O²⁻ in the structure. It is expected that fewer Li⁺ ions coordinated to the O²⁻ would result in more charge density on the oxide, leading to a negative shift on charging the cathode. The reverse is observed on discharge, when it is expected that lithium is inserted into the structure.

The oxidation of O²⁻ to some kind of oxidised condition O³⁻ is typically reported23–25 as a relatively large positive shift which can be represented by a new peak (purple in figure, 530 eV) fitted in the spectra to higher binding energy of the O²⁻ peak. This new peak can be resolved in the spectra for the electrodes at all potentials and excitation energies as the material is charged up to 4.8 V. The relative intensity for the O³⁻ peak is seen to be much smaller at 4.1 V than for higher voltages, where the oxygen redox processes are expected to a larger extent; this will be discussed further later. Apart from consistency with previous studies, the O³⁻ peak assignment is clear when compared with spectra for a material which does not exhibit oxygen redox activity and where such a peak would not be expected. The spinel-phase LiNi₀.₃Mn₁.₅O₄ (Fig. 3), which offers robust high-voltage cycling and is not known to exhibit oxygen redox activity, does not show the formation of the O³⁻ species on charge. This provides further confirmation that the new peak fitted for the Li-rich material is attributable to the species O³⁻ and is a strong indication for the material undergoing oxygen oxidation on charge.

On lithiation of the layered oxide, the area for the O³⁻ peak reduces almost to nothing and an increase in the signal from surface-bound species is evident. This suggests a highly (but not completely) reversible process for the oxygen redox reaction.
While extensive decomposition of the electrolyte is unlikely at 2 V vs. Li/Li⁺, it may be possible for a minor surface layer to form, detectable using the most surface-sensitive photon energies. The binding energy of the main surface species present on discharge is identical to that of the surface species already fitted in the spectra of the other samples, suggesting that lithium carbonate or similar is present. A peak at 55.6 eV corresponding to Li₂CO₃ is also present in the Li 1s spectrum of the discharged sample (ESI, † Note 5).

Further analysis in the ESI † (Note 6) explores the necessity to include the O²⁻ peak in the fitting model. Moreover, an alternative approach to the analysis (ESI, † Note 7), with the presumption of a uniform O⁰⁻ distribution from surface to bulk, results in a poorly fitted model to the experimental data and inconsistencies with previously reported observations.⁹

Fig. 4 shows the percentage O²⁻ out of the total oxide contribution (O²⁻/ [O²⁻ + O⁰⁻]%) at each depth, based on peak area analysis, as a function of voltage. The plot indicates that the proportion of O²⁻ is greater near to the surface for each of the charged samples. This demonstrates a strong depth dependency for the oxidation state of oxygen in this material during charge, with a greater concentration of O²⁻ near to the particle surface. While we might expect the ratio to increase across the oxygen redox plateau, in fact it drops slightly at the top of charge (4.8 V). This is consistent with the previously observed loss of oxygen from the structure at high voltages after the oxygen redox plateau.⁵⁵,⁵⁶

Ni 2p. The state of nickel in the structure is also observed (Fig. 5) to be dependent on depth. For the pristine material, the binding energy for the Ni 2p₃/₂ peak is higher at the surface than in the bulk (854.9 eV at 1.09 keV vs. 854.3 eV at 7.05 keV). Typically a positive BE shift can be associated with oxidation, but assignment of nickel oxidation states from the Ni 2p core level is not so straightforward due to the complex multiplet splitting observed for this transition. Therefore, changes in peak shape and relative BE shifts are considered much more important than absolute peak positions when analysing this core level.⁵⁵,⁵⁶ Many of the spectra have peak shapes that resemble those for previously reported Ni²⁺-containing materials, suggesting that, for the pristine material and for surface measurements of charged samples, nickel is likely in the 2⁺ oxidation state.

The most significant observation that we can make of the peak shape is in the series of spectra at 7.05 keV photon energy. On charge, particularly at 4.1 V but also at 4.55, 4.65, and 4.8 V, a shoulder (blue region) is observed at the higher BE side of the Ni 2p₃/₂ peak, which is also present in the spectra measured at 2.35 keV. Upon discharge the shape returns to one identical to that of the pristine sample. For surface measurements, we observe an opposite shift, to lower binding energies, for the charged samples, and can be indicative of a reduction of nickel at the near-surface. These features will be analysed further later in the discussion.
Mn 2p and Mn 3s. The oxidation state of the manganese in the Li-rich compound is expected to remain in the +4 state during charge. For this reason, any charge compensation from Mn during lithium extraction-insertion is not anticipated.

Fig. 3 O 1s photoelectron spectra for pristine and charged (4.75 V and 5 V) high voltage spinel LiNi_{0.5}Mn_{1.5}O_{4} electrodes, measured using photon energies of 1.09, 2.35, and 7.05 keV. Only one peak can be fitted for structural oxygen (O^{2-}, green), while surface-bound species (grey) show greatest intensity in near-surface measurements. Increasing intensity of the surface-bound species on charge is consistent with electrolyte oxidation/decomposition at the surface of the active material particles. Refer to the ESIt (Note B) for further details. Raw data (red points) and the fitted model (black solid line), both without background subtracted, are superimposed and vertically offset from the component peaks.

Fig. 4 Analysis of O 1s spectra from Fig. 2 for Li_{x}Ni_{0.2}Mn_{0.8}O_{2} electrodes at various voltages. The top panel shows a plot of the voltage profile for the first cycle as a function of the molar fraction of lithium, xLi (based on charge passed). The bottom panel shows the percentage O^{2-} out of the total oxide contribution (O^{2-}/[O^{2-} + O^{n-}]*100), based on peak areas, for each excitation energy (1.09, 2.35, and 7.05 keV) as a function of xLi.

Fig. 5 Ni 2p photoelectron spectra for pristine, charged (4.1, 4.55, 4.65, 4.8 V vs. Li/Li+) and discharged (2 V) Li_{1.2}Ni_{0.2}Mn_{0.6}O_{2} samples measured using excitation energies of 1.09, 2.35, and 7.05 keV. The Ni 2p_{3/2} and associated satellite region is shown as the most informative for determining oxidation state changes. Vertical lines indicate binding energy shifts of the peaks at each excitation energy. The orange coloured region in the bulk measurement represents the state of nickel detected in the pristine material, whilst the blue coloured regions indicate the formation of a shoulder at higher binding energy for some spectra.

Fig. 6a presents the Mn 2p spectra for the pristine materials, samples charged up to 4.8 V and then discharged. For the bulk measurement, very little change in the Mn 2p_{3/2} peak shape or
position is observed, a clear indication that the manganese is not undergoing any redox process on cycling and is stable as Mn$^{4+}$ in the bulk. At the surface, there is little variation during cycling, apart from a shoulder appearing on discharge at the lower binding energy side of the Mn 2p$_{3/2}$ peak. This is suggestive of a partial reduction of Mn$^{4+}$, likely to Mn$^{3+}$, at the surface. An indication of the Mn oxidation state can also be made by analysis of the Mn 3s spectra. A splitting of peaks in the Mn 3s spectra results from a 3s electron coupling with the 3d valence electrons, once the first 3s electron has been ejected.$^{55,57}$ This coupling can be in either a parallel (like spin) or antiparallel (opposite spin) manner and, therefore, results in two different final states giving rise to two different electron energies. The peak splitting can be used as an indication of the manganese oxidation state, as presented in Fig. 6b. Typically, a peak splitting of 4.5 eV indicates manganese in an oxidation state of +4, while a splitting of 5.4 eV is indicative of a +3 oxidation state; values in between these can suggest some mixture of oxidation states. In the bulk, the splitting ranges between 4.41 and 4.62 eV, confirming the presence of Mn$^{4+}$ over the whole charge/discharge cycle, while a slightly more reduced state is apparent at the surface. On discharge, a mixture of Mn$^{4+}$ and Mn$^{3+}$ is indicated by this analysis. This is supported by Mn 3p spectra (ESI,† Note 5), which shows a negative shift on charge and reversal on discharge, leaving a component at lower binding energy.

**Multiple cycles.** After 10 cycles (discharged state, 2 V), a small peak can be fitted in O 1s spectra (Fig. 7) for the O$^{2-}$ species.
Oxygen redox activity measured using excitation energies of 1.09, 2.35, and 7.05 keV. Peaks are fitted for $O^2-$ (green), $O^{4-}$ (purple), and surface-bound species (grey), according to the procedure in the ESI† (Note 2). Raw data (red points) and the fitted model (black solid line), both without background subtracted, are superimposed and vertically offset from the component peaks.

Fig. 7  O 1s photoelectron spectra for the Li$_{1.2}$Ni$_{0.2}$Mn$_{0.6}$O$_2$ samples at 10 cycles (discharged state, 2 V) and 10.5 cycles (charged state, 4.8 V) measured using excitation energies of 1.09, 2.35, and 7.05 keV. Peaks are fitted for $O^2-$ (green), $O^{4-}$ (purple), and surface-bound species (grey), according to the procedure in the ESI† (Note 2). Raw data (red points) and the fitted model (black solid line), both without background subtracted, are superimposed and vertically offset from the component peaks.

near to the surface, the contribution of which increases slightly at greater depths. This indicates that the process is not completely reversible after many cycles and that some of the oxidation product remains in the bulk of material, consistent the formation of a core–shell structure. After a further charge (10.5 cycles, 4.80 V), the peak area for $O^{4-}$ increases as observed in the first cycle but not to the same extent. The $O^{4-}$ proportion ($O^{4-}$/[O$^2-$ + $O^{4-}$%]), determined by peak area, for the surface measurement is calculated at 22.8% after 11 charges versus 43.2% after one charge, indicating a marked decrease in the oxygen redox activity after 11 charges. This reduced activity is expected from the lower capacity and operating voltage exhibited after 10 cycles (see ESI†, Note 9).

8. Electronic structure simulations of layered and rocksalt structures

To complement and help interpret the spectroscopic measurements, we performed high level ab initio simulations including complex calculations of binding energies. The pristine material was investigated both as a layered oxide and a cation disordered rocksalt structure with the composition of Li$_{1.17}$Ni$_{0.25}$Mn$_{0.58}$O$_2$, which is a generally accepted simulation model. The cation disordered rocksalt structure serves as a representative surface phase, which is particularly consistent with previous reports of rocksalt formation on charge, but which may also be present to an extent on the pristine material. We note that the surface structure may not only be limited to the rocksalt phase, but that there may also be other phases present at the surface of the particles. The intention for these simulations is to demonstrate shifts in binding energies dependent on atomic arrangement in different structures of the same composition. The final simulated layered structure was fully consistent with previous studies, in which the excess Li ions in the transition metal layers are arranged in a honeycomb structure surrounded with either six Mn ions, or five Mn ions and one Ni ion (details in ESI†, Note 10). The rocksalt structure ($Fd\overline{3}m$) was considered fully disordered with the same transition metal ratio as in the layered structure. We find that the disordered rocksalt form, as a bulk phase, is calculated to have a higher total energy than the layered structure by approximately 0.5 eV f.u. $^{-1}$, which falls in the same range as previous reports on ordered/disordered LiMnO$_2$ materials.

Fig. 8 shows the simulated XPS binding energies for the O 1s core level of the two structure types of the pristine material, broken down into different transition metal oxide environments. For the rocksalt structure, many of the environments have higher binding energy than those in the layered structure. Therefore, we find a net positive shift in the calculated mean binding energy upon cation disordering: the mean core level binding energies are 530.5 eV and 530.8 eV for layered and cation disordered structures, respectively. The simulated energy difference agrees with the experimentally measured value (0.3 eV) between the surface and bulk, although we note that the calculated XPS energies are not sufficiently accurate for a quantitative comparison. In addition, the calculated normal Gaussian distribution of binding energies is broader for the disordered structure, which is also consistent with our experimental XPS data.

By plotting a cumulative Gaussian distribution for each type of oxygen coordination and for every oxygen within the structure we observe that the broadening is the consequence of a greater number of possible oxygen environments within the cation disordered rocksalt phase. Oxygen coordinated to four Li ions and two Mn ions are the most common in both structures of Li$_{1.17}$Ni$_{0.25}$Mn$_{0.58}$O$_2$. However, there are a total of four different coordination environments around oxygen within the layered-oxide model, while the rocksalt phase contains seven possible octahedral configurations. In addition, half of the oxygen environments of the layered structure appear to have linear Li–O–Li bonds, whereas every oxygen has such bonding within the rocksalt structure. According to Seo et al., the linear Li–O–Li coordination is a structural requirement for oxygen oxidation upon delithiation. The presence of rocksalt phases at the surface may therefore, additionally, promote the formation of oxidised oxygens.

Four delithiated phases of layered Li$_{1.17-x}$Ni$_{0.25}$Mn$_{0.58}$O$_2$ ($x = 0.0, 0.5, 0.67, 0.83$ and $1.0$) were modelled and their O 1s core level binding energies calculated. Several different Li$^+$ arrangements were considered to generate structures for the delithiated phases. According to previous studies, defect spinel-like structures...
features are possible on Li extraction, leading to Li⁺ in tetrahedral coordination. To allow for this possibility our delithiated structures contain Li⁺ ions in both octahedral and tetrahedral coordination. The simulated charging profile (shown in Fig. 9a) has the characteristic voltage plateau around 4.4–4.5 V in the 0.5 ≤ x ≤ 1.0 region.

The oxidation state of ions can best be related to their calculated magnetic moments as this reflects the spin density. Fig. 9b–d show the average magnetic moments of every ion type during the charging process. Our results indicate the oxidation of a small number of anions even at the beginning of the voltage plateau (x = 0.5), accompanied by the partial oxidation of Ni²⁺.

With further extraction of lithium, the number of oxidised oxygen gradually increases and all Ni²⁺ ions oxidise. In contrast, the oxidation state of the Mn⁴⁺ ions is unchanged. While the Mn⁴⁺ has a constant value (2.9 µB), the magnetic moment for Ni varies between ~1.7 and 0.0 µB depending on the lithium content. The change in average magnetic moment for oxygen is small in magnitude, but detectable, and corresponds to one quarter of the oxygens being oxidised. This is in good agreement with the experimental bulk O¹⁻ signal. In general, the experimental data and models on oxygen redox activity are supported by these simulation results.

The oxygen redox activity within the delithiated structures is further investigated through analysis of the partial density of states and spin density (Fig. 10). Such analysis helps to support the modelling of the delithiated states where localised hole states are observed, and demonstrates consistency with other studies. It was previously shown that localised hole states on the oxygen have 2p character and usually appear in the lowest unoccupied bands. This aspect is illustrated more clearly by plotting the partial density of states for only the oxygens that are identified as O¹⁻ based on their magnetic moments (Fig. 10a). Fig. 10b shows the charge density associated with the lowest unoccupied states, indicating that these oxygens are indeed associated with localised 2p character and, interestingly, point in the direction of an unoccupied (or vacant) Li site along an Li–O–Li axis.

Ab initio simulations were also used to evaluate the core level binding energies of every oxygen for Li₁.₁₇₋ₓNi₀.₂₅Mn₀.₅₈O₂ structures with varying Li content (x = 0.0, 0.5, 0.67, 0.83 and 1.0), and are summarised in Table 1; the methodology used is described in the ESI† (Note 10). The results reveal a distinct energy difference between the core level energies of the two different oxygen species: the lattice O²⁻ peak is found to fall...
around 530.2–531.0 eV, whereas the oxidised oxygen always appear at higher binding energies of \( \approx 532.0 \) eV. We conclude that the difference between calculated mean values (i.e. the peak separation) is around 1.1–1.5 eV, which is in agreement with the experimental peak difference; these results again provide quantitative support for our experimental analysis of the oxygen redox activity.

**Discussion**

The high resolution O 1s photoelectron spectra demonstrate that there are two discrete species of oxygen, \( \text{O}^2^- \) and an oxidised oxide ion \( \text{O}^{n-} \). This is consistent with the model for O-redox described in our previous paper, in which we showed experimentally that the holes generated on oxide ions upon oxidation are localised at specific oxygens in the lattice, those coordinated by ionic bonds with Li and Mn. However, the specific degree of oxidation cannot be confirmed from the XPS
results, as is also the case in previous studies where the oxidised oxygen is described as either O$^{2-}$, O$^-$, or peroxo-like.$^{1,2,23}$ The observation of the two species at 4.1 V suggests that the onset of oxygen oxidation can occur in the near-surface region at lower voltages than previously thought, while the major onset at 4.55 V is consistent with previous reports for bulk measurements. An advantage to using the surface-sensitive soft X-ray XPS is that this initial onset at the surface can be detected while exclusively bulk-sensitive techniques such as transmission-mode absorption spectroscopies may not have detected such a signal. The O$^{2-}$ ion signal exhibits a slightly higher binding energy at the surface than in the bulk, consistent with a variation in the electronic structure at the near-surface compared with the bulk. This observation is in good accord with the established core–shell model for these materials, particularly for delithiated phases, but additionally for pristine materials which have been shown to exhibit alternative phases (e.g. rocksalt) to the bulk.$^{11,17}$ This has been particularly reported from STEM experiments for the material in the present study and similar Li-rich materials, in the pristine state and after cycling of the electrode.$^{7,9,11-15,20,21}$ Our calculations of mean core level binding energies for rocksalt and layered structures, identified by the DFT studies in this work, support the shifts in binding energies observed experimentally, which are attributed to a core–shell particle structure. A broadening of the O$^{2-}$ peak (increase in FWHM) at the surface in the pristine sample (1.13 eV for the 1.09 keV; 1.03 eV for the 7.05 eV) represents the greater disorder and more numerous oxygen environments at the near-surface, consistent with a rocksalt layer. The slightly lower binding energy for O$^{2-}$ at high voltage can be interpreted as due to the loss of Li$^+$ ions from the coordination environment of the O$^{2-}$ ions, resulting in an upward shift in the O 2p states. A depth-dependant shift in the Ni 2p spectra for the pristine material is also consistent with a core–shell structure, while shifting in surface spectra on charge points towards further structural rearrangements. Nickel is observed to be most active in the bulk layered phase with a clear change of peak shape occurring at 4.1 V on charge, indicative of an oxidation process likely from Ni$^{2+}$ to Ni$^{3+}$/Ni$^{4+}$. This feature is still evident, but to a lesser extent, upon continued charging up to 4.8 V, after which the peak shape returns to that of the pristine material on discharge suggesting a reduction of the nickel. The calculated average magnetic moment of nickel in the disordered rocksalt structure (1.4 $\mu_B$) is slightly lower than in the layered structure (1.7 $\mu_B$), suggesting a different structural phase or partially oxidised Ni ions close to the surface, which accounts for slightly different binding energies at the surface to the bulk for Ni in the pristine material.

However, for the charged electrodes we observe what is likely attributed to a reduction of nickel at the surface. This is consistent with previous reports of nickel reduction being associated with greater oxygen oxidation activity at the surface of particles.$^{24}$ The calculated average magnetic moment of manganese in the rocksalt is 3.2 $\mu_B$, while for the layered structure it is 2.9 $\mu_B$, indicative of manganese in a partially reduced state close to the surface of the pristine material. This is supported experimentally, primarily by a mixture of Mn$^{3+}$ and Mn$^{2+}$ states at the surface determined by the Mn 3s peak splitting, but also by Mn 2p BE shifting and peak shape changes from surface to bulk for the pristine material. Mn is stable during charging remaining in the Mn$^{3+}$ state, as predicted by our calculations of magnetic moments. However, on discharge we observe from both the Mn 2p and Mn 3s analysis a reduction of Mn close to the surface, further supporting the core–shell model. In addition, the reduction of Mn near the surface upon discharge could be consistent with activation of Mn due to oxygen loss, formation of a rocksalt layer, or dissolution.

Experimentally, we observe a higher proportion of O$^{2-}$ at the surface than in the bulk in the charged states, characterised by the higher intensity of the O$^{2-}$ peaks at the surface, relative to the O$^{2-}$ peaks. Energy-tuned XPS is used as an inherently surface-sensitive (nm scale, up to 29 nm in this case) technique as opposed to a bulk–whole electrode-sensitive technique (>100 $\mu$m) in order to probe a large series of individual particles within the electrode. This gives us confidence that the concentration gradient observed on the scale of the particles rather than a result of the solid-state diffusion effects across the electrode thickness. Our results suggest that oxygen oxidation occurs preferentially in the particle surface layers, with an O$^{2-}$ concentration gradient existing through to the particle bulk. The signal for O$^{2-}$ in the surface measurement upon charge (43% at 4.8 V) cannot account for the whole O$^{2-}$ signal observed in the bulk measurement (27% at 4.8 V), despite the exponential decay of electron escape depth exhibited in XPS measurements, since the range of probing depth is so large (4 nm vs. 29 nm). A thorough calculation demonstrating that part of the O$^{2-}$ signal is originating from the bulk is provided in ESI, Note 1.† Therefore, whilst at each voltage a greater proportion of O$^{2-}$ is observed close to the surface, the results indicate that a significant amount of the oxidised oxygen is formed in the bulk. This would be expected from the substantial charge compensation attributed to O-redox. The concentration gradient observed for the oxidised oxygen is highly consistent with the well-established core–shell model for the material, which is characterised by the formation of a substantial rocksalt phase surface layer 2–3 nm in thickness as seen in STEM images (ESI,† Note 11). However, with part of the oxidised oxygen signal originating from the bulk, we can say that it is not only the rocksalt layer which contains the O$^{2-}$ species. This points to the well-established core–shell structure of such materials not being directly coupled to the oxidised oxygen concentration gradient observed in this study. This is despite the rocksalt surface layer likely exhibiting a higher probability of oxygen redox, due to the greater proportion of Li–O–Li bonding compared with the layered structure. More likely, the concentration gradient observed is as a result of the oxygen at the near-surface being oxidised first as lithium continues to be inserted from the electrolyte. As diffusion of lithium from surface to bulk proceeds, so then the bulk oxygen is partially oxidised as well. This mechanism is presented schematically in Fig. 11. It was shown that after 10 cycles there is a greater retention of O$^{2-}$ in the bulk than at the surface. We see this as particularly significant and suggests that diffusion limitations of the structure prevent oxygen loss from the bulk. This further supports the
greater activity at the near-surface region and agrees well with the core–shell model. We expect that, close to the end of charge after extensive delithiation of the structure, the possible locations for oxygen oxidation (e.g. Li–O–Li configurations) will be fewer leading to the reduced possibility for oxygen oxidation. This is likely to contribute to the limitation of the oxygen redox reaction or oxygen loss from the structure, and limit the charge capacity of the electrode. It is probable that the particle structure is more complicated than what has been determined in this study, for example with certain facets of a particle surface demonstrating a high affinity for oxygen oxidation than others. However, by using a large spot size for our experiments we have measured an average state over many particles within each sample and therefore present an overall picture but with a clear surface vs. bulk trend.

These results additionally highlight the importance of making such measurements using multiple excitation energies over a wide range to probe the structure from the very surface through to the bulk. The lowest excitation energy of 1.09 keV used in this study is able to probe almost exclusively the 2–3 nm rocksalt layer present at the surface of the particles, whilst the use of only one ‘mid-range’ energy on a laboratory XPS, corresponding to a depth of more than 6 nm does not represent the near-surface region. Similarly, employing only hard X-ray energies, as in previous studies, disregards surface processes, creating ambiguity and not revealing the true nature of the materials and processes being investigated.

Conclusions

To conclude, a depth dependency for the oxygen oxidation reaction has been demonstrated by using energy-tuned photoelectron spectroscopy to probe the Li-rich layered Li₁₋₂Ni₀.₂₋₃Mn₀.₆O₂ cathode material. As expected and reported elsewhere, this material exhibits a high degree of redox activity by oxygen in the lattice, being oxidised from O²⁻ to a species we designate O⁴⁻ on charge; both are detected as discrete species in our photoelectron spectroscopy measurements. This is in contrast to the oxygen spectra obtained for the oxygen redox inactive high voltage spinel LiNi₀.₃Mn₁.₃O₄, for which no O⁴⁻ species is detected. A concentration gradient of the O⁴⁻ species is observed in the layered material with a higher concentration at the surface of particles compared with the bulk in the charged electrodes. Our findings support an already well-established core–shell particle structure but appear to show that the concentration gradient is not directly coupled to the 2–3 nm rocksalt shell often observed. Measurements indicate a variation between surface and bulk in the electronic structure of the oxide or the presence of alternative surface phases already on the pristine material, but as would be particularly expected of the charged material. High level ab initio calculations support this observation, where the O²⁻ of the rocksalt structure, representing a possible surface layer, has a higher mean binding energy than that for the layered structure. Calculated core level binding energy differences for O³⁻ and O⁴⁻ species are in very close agreement with experimental analysis. The simulations on delithiated phases show oxidation of nickel, with negligible change to the manganese oxidation state, in good agreement with bulk photoelectron spectroscopy measurements and previous studies using alternative bulk-sensitive spectroscopies. By employing a spectroscopic technique sensitive to both the surface and the bulk, we have been able to reveal the depth-dependent oxygen redox activity of this material, which is consistent with the existing core–shell model. This methodology could be applied to similar such materials in order to further understand their charge compensation mechanisms.

Supporting data

Supporting research data has been deposited in the Oxford Research Archive and is available under this DOI: 10.3287/bodleian:8grVPv41r

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

P. G. B. is indebted to the Engineering and Physical Sciences Research Council (EPSRC), including the SUPERGEN Energy Storage Hub [EP/L019469/1], Enabling Next Generation Lithium Batteries [EP/M009521/1], Henry Royce Institute for Advanced Materials [EP/R00661X/1, EP/S019367/1, EP/R010145/1] and the Faraday Institution All-Solid-State Batteries with Li and Na Anodes [FIRG007, FIRG008] for financial support. The authors gratefully acknowledge the MCC/Archer consortium (EP/L000202/1) and the Swedish National Graduate School in Neutron Scattering (SwedNess), part of the Swedish Foundation for Strategic Research (SSF). We thank Diamond Light Source for access to beamline I09 (proposal numbers NT14733-1 and SI16629-1) that contributed to the results presented here. We are grateful to Dr Tien-Lin Lee, Dr Christoph Schlueter and Dr Pardeep Thakur Kumar for their assistance.

References