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Investigating lithium-ion battery materials during overcharge-induced thermal runaway: An *operando* and multi-scale X-ray CT study

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Catastrophic failure of lithium-ion batteries occurs across multiple length scales and over very short time periods. A combination of high-speed *operando* tomography, thermal imaging and electrochemical measurements is used to probe the degradation mechanisms leading up to overcharge-induced thermal runaway of a LiCoO$_2$ pouch cell, through its interrelated dynamic structural, thermal and electrical responses. Failure mechanisms across multiple length scales are explored using a *post-mortem* multi-scale tomography approach, revealing significant morphological and phase changes in the LiCoO$_2$ electrode microstructure and location dependent degradation. This combined *operando* and multi-scale X-ray computed tomography (CT) technique is demonstrated as a comprehensive approach to understanding battery degradation and failure.

Introduction

Lithium-ion batteries are ubiquitous energy storage devices in portable electronics owing to their high energy and power densities. The synergy between lithium-ion batteries and clean, renewable energy sources to achieve global CO$_2$ emission targets, also makes them the technology of choice for the rapidly expanding market for electric and hybrid electric vehicles. However, for many advanced applications, lithium-ion batteries are required to operate safely over a range of temperatures and during high charge and discharge rates. Although failure is rare the consequences are severe and several well publicised incidents involving catastrophic battery failures have highlighted the need to further reduce the risk associated with battery packs and modules.

Overcharge of lithium-ion batteries poses a significant safety risk as chemical and electrochemical reactions can occur between cell materials, and is one of the primary causes of thermal runaway, as seen in the recent spate of failures associated with charging self-balancing scooters; consequently, overcharge tests have been integrated into safety test standards for lithium-ion batteries. The magnitude of the energy released during failure increases with increasing state-of-charge (SOC), as the chemical energy stored in the electrode materials is released, hence thermal runaway resulting from overcharge is particularly catastrophic.

At a critical temperature, thermal runaway takes place when a chain of exothermic reactions between the electrodes and electrolyte occurs. For example, the LiCoO$_2$ positive electrode undergoes exothermic decomposition at elevated temperatures through a series of reduction steps in the presence of electrolyte. There have been numerous studies focusing on the reaction pathways of positive electrode materials during failure, but a spatial and...

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temporal understanding of the reaction processes and their effects on particle microstructure, rate of reaction and critical temperatures reached during failure, is yet to be achieved.

It is well known that the microstructure of electrode materials plays an important role in determining the performance of cells, but the role of particle morphology in the safety of cells is not yet well understood. Jiang and Dahn used accelerated rate calorimetry to show the effect of particle size on the onset temperature of thermal runaway, demonstrating that a larger particle size is more thermally stable. More recently, Gedar et al. demonstrated through thermogravimetric analyses that the mass loss associated with the decomposition of LiCoO$_2$ increases linearly with surface area at 400 °C. A higher mass loss corresponds to increased heat generation during failure, and increased gas (including O$_2$) evolution, further fuelling decomposition of the electrolyte; smaller particles were shown to release more oxygen and have an oxygen deficiency at their surface.

The fast reaction kinetics between the electrolyte and the positive electrode lead to generation of heat at a very high rate, which can result in internal temperatures of commercial cells reaching in excess of 1000 °C, damaging and by-passing integrated safety devices, which poses a great risk to modular systems where cell-to-cell propagation of failure is a major concern. Thus, the reliability and safety of lithium-ion batteries is considered a function of the entirety of the cell or modular system. Understanding the link between behaviour on the microscale and the catastrophic effects observed for full cells and modular systems is imperative to progress towards safer battery designs. With the advancement of operando X-ray imaging techniques, tracking the propagation of failure at high frame rates and across multiple length scales is now possible. X-ray computed tomography (CT) has previously been used to perform multi-length scale analyses of battery materials, and has also been demonstrated as an effective diagnostic tool for lithium-ion batteries during, and after failure.

Here, we combine high-speed operando synchrotron X-ray CT with thermal and electrochemical measurements to link, and assess, the observed morphological, thermal and electrical response of a commercial LiCoO$_2$ battery during overcharge-induced failure. A multi-scale comparative analysis of the cell in its fresh and failed state is also performed to elucidate failure mechanisms across multiple length scales, from individual electrode particles to the full cell architecture. Image-based quantification is used to determine the change in electrode particle morphology via a particle size distribution analysis, and the effects of particle morphology on the failure mechanism of lithium-ion batteries are discussed.

**Experimental**

**Electrical abuse testing and operando synchrotron CT**

Fast tomographic imaging was performed at beamline ID15A at The European Synchrotron (ESRF). An in-situ containment system was designed to allow simultaneous high speed X-ray CT and thermal imaging (Figure 1a) and is discussed in more detail in a previous work. The overcharge abuse test was performed on a Turnigy nanotech 160 mAh lithium-ion pouch cell (dimensions 39mm × 12 × 8mm) which consists of a LiCoO$_2$ positive electrode, polymer separator and graphite negative electrode. The Turnigy cell is designed to cater for high charge (10 C) and discharge (30 C) rates (where a C-rate is the rate at which a battery is discharged relative to its maximum capacity). In this study the battery was overcharged from 100 % SOC (4.2 V) at a constant current of 3 A (18.75 C) until failure, which is well outside the safe operating limits recommended by manufacturers.

Use of an electrical slip ring (P4+ Compact Slip Ring, Moog, UK) built into the ID15A rotation stage (ABR1000, Aerotech, USA) allowed simultaneous rotation of the sample (for X-ray CT) and electrical charging. The cell voltage plot during overcharge is shown alongside the surface temperature profile in Figure 1b; the temperature presented in the plot was taken as an average over an area ca. 9 mm$^2$ on the surface of the cell (shown in Figure 1a).

The surface temperature of the battery was recorded using a thermal camera (FLIR SC5000MB, FLIR Systems, France). The thermal camera was set to the calibrated range of 15 °C to 200 °C during these experiments. A uniform layer of high emissivity black paint (with a calibrated emissivity of 0.96 over the range 40 °C – 180 °C) was applied to the surface of the battery before imaging. The FLIR SC5000MB thermal camera has an extended wavelength detector to capture infra-red wavelengths between 2.5 μm and 7 μm. In the calibrated range, the camera has a noise equivalent
temperature difference < 20 mK. The manufacturer specifies a measurement accuracy of ± 1 °C or ± 1 % of the temperature in degrees Celsius. Images were recorded at 25 Hz and the real time movie is provided as Supplementary Movie 1.

The Turnigy cell was imaged using a 76 keV monochromatic synchrotron X-ray beam. The X-ray absorption signal was converted into a visible light signal by a LuAG:Ce scintillator screen and then collected by a 1X macro objective. A high-speed PCO Dimax CMOS camera (PCO AG, Germany) was used to record the images. The field of view (FOV) was 10.5 mm × 7.6 mm, which consisted of 960 × 700 pixels (horizontal × vertical) and gave a pixel resolution of 10.9 µm. The sample rotation axis was at the edge of the FOV, such that through a 360 ° rotation it was possible to double the size of the 3D image in the horizontal direction30. The resulting 3D reconstruction was then 1785 × 1785 × 700 pixels (19.2 mm × 19.2 mm × 7.6 mm). Each tomogram consisted of 2 × 2000 half projections using an exposure time of 0.7 ms. The acquisition time for each tomogram was 2.8 s and one tomogram was captured every 40 s.

Multi-scale laboratory X-ray micro-CT

Using material extracted from the same Turnigy cell imaged in the synchrotron, tomographic reconstructions of varying sample size and resolution were produced using lab-based X-ray CT systems (Zeiss Xradia Versa 520 and Zeiss Xradia Ultra 810, Carl Zeiss XRM, Pleasanton, CA, USA). The specific imaging properties for each scan are provided as Supplementary Information. Materials were imaged with a pixel resolution of 63.1 nm (Zeiss Xradia Ultra 810), 0.36 µm (Zeiss Xradia Versa 520) and 7.92 µm (Zeiss Xradia Versa 520). The X-ray CT system (Zeiss Xradia Ultra 810) which achieved a resolution of 63.1 nm uses a chromium target with an accelerating voltage of 35 kV and tube current of 25 mA. The characteristic spectrum from the Cr target is quasi-monochromatic around 5.4 keV. The CT system used for the remaining scans (Zeiss Xradia Versa 520) had a characteristic spectrum from a tungsten target; the accelerating voltage and tube current are user defined and determine the peak intensity of the bremsstrahlung and photon flux. The accelerating voltage and tube current were chosen based on the X-ray absorption coefficients of the samples. The transmission images from all scans were reconstructed using...
a commercial software package (Zeiss XMReconstructor), which uses an algorithm based on standard filtered back-projection.

Data processing

The reconstructed tomograms were processed using Avizo Fire 9 software (FEI VSG, France). An edge preserving non-local means filter was applied to the images to reduce noise while maintaining phase boundaries for segmentation. Phases were separated based on grey scale values, where the highly attenuating materials are displayed as white and weakly attenuating materials appear as dark grey. Measurements of sample porosity and particle size distribution were performed using Avizo Fire 9’s label analysis tool. The temperature data and thermal imaging videos were extracted using FLIR’s Altair software.

Results and discussion

Voltage, temperature and operando tomography

During overcharge of lithium-ion batteries, a sequence of events related to the evolution of voltage, temperature and chemistry of the cell occurs leading up to thermal runaway and failure. In Figure 1b, distinctive features can be identified in the temperature and voltage curves. For the initial 110 s (Region 1), the external temperature of the cell increases from 20 °C to ca. 40 °C. This initial heating is expected to derive primarily from irreversible heat generation mechanisms, such as Ohmic losses, which are most prevalent at high C rates. A plateau in cell voltage is observed after 110 s (Region 2 in Figure 1b), followed by a significant increase in the rate of temperature rise after 120 s. The simultaneous voltage plateau and temperature rise indicates the initiation and progression of the decomposition / formation of the solid electrolyte interphase (SEI). At high temperatures (> 80 °C), the SEI becomes unstable and self-discharge initiates. Lee et al. suggested that the lithium becomes more active within the carbon lattice at temperatures above 80 °C, causing a large reduction in charge transfer resistance. The combination of the accelerated loss of lithium and the instability of the SEI layer results in an increasing rate of the exothermic SEI decomposition / formation reactions and consumption of lithium.

After ca. 80 s, corresponding to an external temperature of 40 °C, the pouch of the cell has begun to swell (Figure 2) as gas is generated from reduction of the electrolyte and decomposition of the SEI. After 120 s, severe delamination is observed around the separator regions in the outer layers. Within Region 3 in Figure 1b, a distinctive peak in cell voltage occurs over the space of ca. 40 s, consistent with the behaviour observed by Belov and Yang. In Supplementary Movie 1, the pouch structure is seen to diverge from its original packaged structure, which corresponds with this voltage peak, as observed in Figure 1b. It is expected that the rise in voltage is caused by the resistance increase associated with the gas pockets forming between the active layers, and the subsequent fall in voltage by the decrease in resistance when the pouch bursts and the gases are channelled away. The tightly wound active layers channel gases out into the exterior pouch when the pouch begins to lose its original package structure and starts to swell.

Finally, within Region 4 in Figure 1b, the cell voltage rises sharply to the maximum voltage of the potentiostat, the most likely cause of which is the shutdown of the separator and damage to the internal circuit caused by the rupture of
the cell at that time. The tomogram taken at 160 s in Figure 2 shows delamination of the LiCoO$_2$ layers from the aluminium current collector in various places, which may be evidence of the electrolyte vaporising or initial stages of decomposition of the LiCoO$_2$ resulting in gas pockets forming between the aluminium current collector and the LiCoO$_2$ layer. This may also be due to the current collector being oxidised by the electrolyte, or the products of electrolyte decomposition$^{40, 41}$.

The thermal decomposition of Li$_{0.5}$CoO$_2$ can occur at temperatures above 200 °C$^{13}$ and the reaction between Li$_{0.5}$CoO$_2$ and solvent (ethylene carbonate), to produce CO$_2$ and H$_2$O, can occur at temperatures as low as 130 °C$^{10, 13}$, according to equation (1).

\[
\text{Li}_{0.5}\text{CoO}_2 + 0.1\text{C}_2\text{H}_4\text{O}_3 \rightarrow 0.5\text{LiCoO}_2 + 0.5\text{CoO} + 0.3\text{CO}_2 + 0.2\text{H}_2\text{O}
\]

Although the surface temperature of the cell appears to be too low for the decomposition of the LiCoO$_2$ to occur, as Lin et al. suggested$^{42}$, the cathode side with the aluminium current collector is most susceptible to high temperatures during overcharge abuse. It is also recognised that the temperature measured at the surface of the pouch cell is not deemed representative of the temperature of the internal active materials, due to the thermal insulating effect of the gas between the active materials and the surface of the pouch.

Fluctuations in the surface temperature of the cell occur after around 160 s as the cell ruptures (Supplementary Movie 1), at which point the unstable electrodes are exposed to air and the process of thermal runaway is accelerated. The heat generated from the reaction between the LiCoO$_2$ and electrolyte, along with the decomposition of the SEI layer provide heat for the occurrence of thermal runaway which soon ensues, following which the surface temperature of the cell exceeds the limit of the thermal camera’s calibrated range (Region 5 in Figure 1b).

**Post-mortem CT and battery architecture**

For comparison, the full cell was imaged in its fresh state (Figure 3a) and also post-mortem (Figure 3b), with a pixel size of 7.92 µm. The structure of the complete cell before and after thermal runaway is compared and the extent of structural damage and material degradation investigated. The aluminium (Al) phase was segmented (blue phase in Figure 3b), and is shown to have agglomerated at the top and bottom of the wound electrode layers. The rapid gas generation from the decomposition of the LiCoO$_2$ and electrolyte is likely to have carried the molten Al to the ends of the roll as the gas was channeled from the inner layers to the outer pouch. The porous structure of the Al globules at either end of the cell supports the hypothesis that gas played a part in carrying the molten metal. The uneven gas generation may also have had a peristaltic effect between the tightly wound layers, forcing the molten Al to either end.
current collecting tab appears to have reached the melting point of copper (1085 °C). The temperature range between 660 °C and 1085 °C presents the possibility of further decomposition reactions occurring, evidence of which is found upon further investigation of the individual LiCoO₂ particles post-mortem.

Micro-CT and particle structure degradation

The size and shape of particles have been shown to play a significant role in the thermal stability of electrode materials and influence the magnitude of catastrophic failure upon thermal runaway.¹⁶,¹⁷ The decomposition reactions of LiCoO₂ to Co₃O₄, CoO and further to Co metal occur at elevated temperatures and in the presence of organic electrolyte at the surface of the particles. For example, at > 150 °C, the following reduction reactions can occur where solv. indicates the reaction step occurs in the presence of electrolyte solvent.¹⁰

\[
\text{Li}_{0.5}\text{CoO}_2 \xrightarrow{>150^\circ C} 0.5\text{LiCoO}_2 + \frac{1}{6}\text{Co}_3\text{O}_4 + \frac{1}{6}\text{O}_2 \tag{2}
\]

\[
\text{Co}_3\text{O}_4 \xrightarrow{>150^\circ C, \text{solv.}} 3\text{CoO} + \frac{1}{2}\text{O}_2 \tag{3}
\]

\[
\text{CoO} \xrightarrow{>150^\circ C, \text{solv.}} \text{Co} + \frac{1}{2}\text{O}_2 \tag{4}
\]

Additionally, CO₂ evolved from degradation reactions (1) and combustion reactions of the solvent and separator, could react with any remaining LiCoO₂ particles according to (5), resulting in additional formation of Co metal.

\[
2\text{LiCoO}_2 + \text{CO}_2 \xrightarrow{>150^\circ C, \text{solv.}} \text{Li}_2\text{CO}_3 + 2\text{Co} + \frac{3}{2}\text{O}_2 \tag{5}
\]

Most of these reactions occur at the interface between the transition metal oxide and electrolyte. Larger particle surface area increases the probability of surface degradation reactions; hence, with reduced particle size the specific surface area exposed to the electrolyte increases and the rate at which exothermic degradation reactions occur would...
be greater. Post-mortem micro-CT of the LiCoO$_2$ particles shows severe microstructural degradation (Figure 5). Evidence of a surface layer of Co metal on the electrode particles is seen in Figure 5a. Figure 5b shows a greyscale 2D slice with the highly attenuating Co layer (white) at the surface of the particle. A semi-transparent 3D rendering of the particle (Figure 5c) shows the presence of Co (teal) on the surface and Co channels within the particle. As a result of changes in density of materials during decomposition, a reduction in particle volume and delamination of phases is expected to occur. The density of Co metal is almost twice that of LiCoO$_2$ and accordingly, there is evidence of delamination of the Co layer from the bulk of the particle, shown in Figure 5(d-e). During thermal runaway, this delamination would have exposed further surface material to exothermic reduction.

From attenuation coefficients of expected decomposition products (see Supplementary Information) the composition of the particle surface layer and interior can be inferred; the difference in greyscale between the Co surface layer and the particle interior suggests that most of the particle consists of CoO, which is consistent with previous XRD experiments$^{10}$. The volume fraction of Co metal in the remaining cathode material is calculated to be 9% (Figure 5f). SEM images and EDS (Figure 6a,b) show that the surface of the particle consists of Co globules rather than the smooth surface presented in Figure 5; this is due to the limited resolution of the X-ray CT technique to resolve fine surface features$^{26}$. The surface of the LiCoO$_2$ particles after thermal runaway appears relatively rough when compared to the fresh sample presented in Figure 6c, which is ascribed to shrinkage of the Co-oxide as it is reduced, and delamination and agglomeration of Co metal. Additionally, subsurface Co metal seams are present in the tomograms which could also be explained by the infiltration of electrolyte through micro-cracks and fractures, which could not be resolved in the CT images.

Without excess electrolyte and high temperature (> 350 °C) it is expected that the air-stable Co$_3$O$_4$ and CoO would not be reduced to Co$^{33}$. Three samples were taken from subsurface layers (Region A in Figure 3b), each of which showed the presence of Co on the surface when analysed under CT; however, a sample taken from the surface of the cell (region B in Figure 3b) did not show any presence of Co on the surface of the particles (Figure 7). This may have been due to the outer regions having a lower local temperature (due to enhanced heat rejection) and insufficient quantities of electrolyte for the final reduction step to occur. As shown in a previous study$^{15}$, temperature gradients of > 700 °C can occur over as little as 7 mm during thermal runaway, where the outer regions of the cell remain relatively cool and the inner regions can reach > 1000 °C. Hence, at the outer regions the conditions (temperature and retention of electrolyte) are less favourable to sustaining the propagation of thermal runaway, making the extent of exothermic degradation location dependent.

In contrast to the subsurface samples, the surface sample presented in Figure 7 contains numerous fractured particles. As Geder et al.$^{17}$ demonstrated, a reduced particle size (and increased specific surface area) results in unfavourable consequences, such as a higher rate of heat generation and lower onset temperatures for thermal runaway.
Particle size distribution

Mass loss occurs at each reduction step according to equations (2) – (4), and the electrode particles are expected to reduce in mass and volume, and increase in density. The particle size distribution (PSD) of a fresh sample is compared with that of both post-mortem samples; the inner sample (showing presence of a Co surface layer) and the outer sample (showing significant cracking and the absence of Co).

The particle volumes used for the PSD of the three samples were 24,897 µm$^3$ (fresh), 34,907 µm$^3$ (inner), and 33,785 µm$^3$ (outer). The PSD for each sample is presented in Figure 8. In its fresh state the PSD centres around a mean diameter of 3.87 µm with a relatively low standard deviation. There is a significant decrease in the mean diameter of the remaining particles for both post-mortem samples to approximately half their original size (to 1.99 µm and 1.97 µm for the inner and outer samples respectively), but the spread of data in the PSD greatly differs (Figure 8). The PSD for the inner sample shows a single peak which has shifted below 2 µm, which is due to a combination of particle shrinkage during phase transitions and the presence of significant debris consisting of fractured particles and delaminated Co. In contrast, a bimodal PSD is observed for the outer sample (cracked particles). When compared to the PSD of the fresh sample, the twin peaks correspond to the occurrence of particle cracking, where the majority of the distribution is shifted to a smaller equivalent particle diameter. However, a second peak which consists of particles < 1 µm in diameter contains the highest frequency of particles. This second peak may reflect smaller particle fragments from the fractured larger particles. Due to the limited resolution of the X-ray CT technique, the specific surface area of the particles could not be accurately determined; however, the dramatic reduction in particle size observed between the fresh and outer samples is expected to significantly affect the rate of decomposition of the electrode at elevated temperatures. Additional surface area would be exposed to reaction with the electrolyte and the smaller fragmented particles would exhibit a lower onset temperature$^{16, 17}$. For example, as demonstrated by Jiang and Dahn$^{16}$, a reduction in particle size of LiCoO$_2$ from 5 µm to 0.8 µm results in decreased thermal stability, with lower onset temperatures.

Although it is well known that the particle shape and size significantly affect the onset temperature and rate of heat generation during thermal runaway, it is not well understood how the evolving microstructure and likelihood of particle fracture influence these properties leading up to and during the runaway process. Further work is required to quantify the change in particle structure based on the conditions to which the particles are subjected, and the effect of initial particle morphology on the rate and quantity of heat generation during thermal runaway are required. As X-ray CT techniques continue to achieve higher spatial and temporal resolution, the dependence of morphological evolution on particle size, and its consequent effect on the magnitude of failure, could also be explored via operando micro-CT. A large particle size with low surface area may provide a more thermally stable electrode, but can also result in a reduction of the cell’s power density; hence a balance between battery performance and safety may exist. Understanding the mechanisms through which particle architecture and properties change during thermal runaway is also essential to progress towards an optimised particle size distribution for both performance and safety.

Figure 8. Particle size distributions for LiCoO$_2$ particles from a fresh sample (top), a sample taken from the inner regions of the failed overcharged cell showing evidence of complete reduction to Co metal (middle), and a sample taken from the outer region of the failed overcharged cell showing evidence of particle cracking with no formation of Co (bottom).
Conclusions

This comprehensive approach of combined high speed operando X-ray CT and thermal imaging, followed by post-mortem multi-scale X-ray CT and SEM analyses, has provided new insights into the failure mechanisms of lithium-ion batteries. The effects of heating and gas generation on the architecture of the cell have been explored during failure via high-speed operando tomography, showing electrode layer delamination, gas pocket formation and consequent swelling of the pouch. Multi-scale comparative analyses of the complete cell, as well as separate components of the cell, before and after thermal runaway, have elucidated key phenomena that contribute to the propagation and severity of thermal runaway; the presence of agglomerated aluminium at both ends of the cell demonstrates the effective heat transfer throughout the cell via molten Al during failure.

X-ray CT of the electrode particle microstructures in samples taken from different locations within the cell presents valuable information on the local kinetics and progress of exothermic reduction reactions at the positive LiCoO2 electrode. The volume fraction of Co, and its presence on the surface of the electrode particles, indicates that the rate of exothermic reduction steps, and therefore heat generation, is dependent on the surface area and PSD of the positive electrode. The extreme thermal and electrical conditions during overcharge-induced thermal runaway cause the particle morphology, and hence the PSD and specific surface area, to change.

The results of this work show that particle microstructure can significantly influence the extent and rate of exothermic degradation reactions during thermal runaway, and demonstrate that X-ray CT could be used to elucidate the relationship between electrode microstructure and overall safety of commercial cells.

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