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Predicting the formation of mud cracks in Li-ion battery electrodes during the drying process with *in situ* X-ray computed tomography

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Drying of slurry cast electrodes is amongst the most energy intensive unit operations in battery manufacture. In spite of this, the optimisation of drying processes has been highly empirical, and there remains limited understanding of the interplay between drying dynamics and resulting microstructure. In this work, we used synchrotron X-ray computed microtomography in order to study the formation of mud cracks during the drying process, and evaluate their impact on the electrode microstructure. This was achieved by applying a reduced drying rate, which proved to be an effective means of studying the drying mechanism with a greater resolution and image contrast than otherwise possible. The rate of crack growth is measured, and the differing crack morphology resultant from changes in thickness (between 300 and 800 μm doctor blade gaps) and the presence of air bubbles in the slurry is demonstrated. Digital volume correlation is utilised to identify the specific location of crack formation before these cracks were visible, suggesting image correlation methods as an appropriate tool for process feedback in order to control or eliminate mud cracking. This new approach which enables direct quantification of the evolving microstructure during dynamic drying, in 3D, is therefore transformative in our understanding of the underlying physical processes and will guide rational optimisation of this industrially significant process.

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Broader context

Solvent-based slurry processing is the means by which the majority of battery electrodes of all chemistries are presently manufactured, typically with a slot die coater head, as part of a roll-to-roll electrode manufacturing line. These approaches are also increasingly applied in the manufacture of perovskite solar cells and other functional coatings. The slurry drying process can have a major impact on the structure and performance of coatings. One such impact is the formation of mud cracks, which result from capillary stress which builds as solvent evaporates. Here we apply *in situ* X-ray computed tomography in a time-resolved study of the electrode drying process, developing key insights necessary to better understand, control or prevent mud cracking.

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1 Introduction

Roll-to-roll fabrication of battery electrodes includes mixing, coating, drying and calendaring steps, after which the electrodes are slit and wound into cells. The drying process of Li-ion battery electrodes, including solvent recovery, has been estimated to account for 15% of total cell cost, and contributes profoundly to the mechanical and electrochemical performance of electrodes.^{1–10} Understanding, predicting and controlling this process is thus crucial to progressing knowledge of battery manufacturing. The drying process begins with a coated electrode slurry where active particles and conductive carbon are suspended in a solution of binder and solvent – typically polyvinylidene fluoride (PVDF) and *N*-methyl pyrrolidone (NMP), respectively – in conventional Li-ion positive electrodes. As solvent evaporates, the solid components consolidate such that inter-particle distances are shorter until all particles are compacted. In granular coatings, such as Li-ion battery electrodes, at the point the solvent level begins to drop below the upper layer of particles, concurrently with air intrusion into the porous structure, a capillary action of solvent towards the evaporating surface results.^{11–15} This capillary action causes capillary stress, σ_{cap} , which can be approximated by the relationship:

$$\sigma_{\text{cap}} \sim \frac{2\phi\gamma}{d} \quad (1)$$

where ϕ is a particle packing factor, γ is the drying solvent surface tension and d is the particle diameter.¹¹ For example, in aqueous silica particle coatings, capillary stress has been shown to increase sharply at the point of initial air intrusion, and then steadily reduce as solvent evaporates, with a small residual stress retained in the fully dry coating.¹⁶ Where capillary stress exceeds the fracture toughness, K , of the coating, cracks form which grow perpendicularly to the applied stress, as shown in Fig. S1.¹⁷ Fracture toughness can be related to a critical cracking thickness, h_c , above which cracking occurs such that:

$$h_c = \left(\frac{K}{1.4\sigma_{\text{cap}}} \right)^2 \quad (2)$$

where h_c is inversely proportional to solvent surface tension and proportional to particle size, and crack propagation proceeds until the applied stress no longer exceeds fracture toughness.^{11,13,15} Cracking results in a sharper decrease in the measured stress present in the coating during crack propagation.¹⁸

In Li-ion battery electrodes, the capillary action process has also been shown to cause the migration of binder and conductive carbon towards the upper region of the electrode.^{2–8,19,20} This effect, which is influenced by drying conditions such as drying temperature and time, can negatively affect adhesion to the current collector and resistivity of the electrode.^{2–10} In addition to binder migration, Jaiser *et al.* showed using cryo-SEM with EDS mapping (in work following on from studies in

silica coatings by Price *et al.*) that the formation of ‘pendular rings’ of solvent between particles in the final stage of the drying process causes the binder to precipitate largely between closely-packed active particles, whilst binder is not present in larger pore spaces.^{16,21} The conditions of the drying process are therefore a major influence on electrode performance.

As discussed above, the drying process can also result in mud cracking of Li-ion battery electrodes above a critical thickness. Mud cracking refers to the formation of cracks in the electrode during the drying process as a consequence of drying stress.²² Mud crack formation in Li-ion battery electrodes has been shown to be promoted by high drying temperatures, as well as increased electrode thickness, both of which are a hindrance to the manufacture of thick electrodes for higher energy density batteries.^{19,22–25} The inverse relationship between critical cracking thickness and solvent surface tension is also a barrier to a move towards aqueous processing and fluorine-free binders, since water has a higher surface tension than alternative organic solvents.^{14,22,23,26,27} Similarly, LFP and LMFP particles are commonly processed as primary particles hundreds of nanometres in diameter, rather than micron-scale secondary particles, so are likely to have a lower critical cracking thickness and so be even less amenable to thick slurry coating.^{28,29}

In contrast, studies by Bryntesen *et al.*, and our own work, have demonstrated a relationship between the presence of cracking and improved electrochemical performance of electrodes.^{25,30} It is believed that this improvement is due in part to enhanced ion transport in cracked electrodes, since cracks provide a directional pore channel which facilitates ion transport through the electrode thickness.^{25,31} In this respect mud cracks are similar to various pore structure engineering methodologies, such as laser ablation,^{32–36} templating,^{37–41} mechanical pore punching⁴² and acoustic patterning.⁴³ These approaches aim to provide enhanced ion transport through directional porosity, enabling improved rate performance in thick electrodes, which are otherwise hindered by poor ion transport.⁴⁴ Controlled mud cracking offers the potential to achieve this effect without additional electrode processing.⁴⁵ The motivation to understand crack formation is therefore two-fold: to prevent excessive cracking which can result in coating delamination, electronic disconnection or other electrode failure mode; and to identify and facilitate an optimal degree of cracking which enhances ion transport without harming performance.

In order to understand the mechanism of the drying process, several imaging approaches have been applied to observe structural changes. Jaiser *et al.* applied time-resolved cryo-SEM of electrode cross sections to investigate solvent binder distribution during the drying process.²¹ They also proposed heterogeneous pore emptying at the drying electrode surface with *in situ* optical fluorescence microscopy.⁵ However, the inherently three dimensional nature of the drying process limited the extent of their study. Higa *et al.* used high speed X-ray radiography to monitor the changing density of drying aqueous SiO_x slurries, and paired this *in situ* analysis with



three dimensional X-ray computed tomography (X-ray CT) to show that particles settled in a ring around a drying drop of coating.⁹ A second study on NMC-based cathodes drying showed poor image contrast of active material, and in both studies *in situ* X-ray CT was not attempted, since the drying times of several minutes were too short to carry out tomographic imaging.⁴⁶ Yang *et al.* resolved this issue using a high-speed dynamic tomography mode which achieved 0.5 s per tomograph, and applied digital volume correlation (DVC) to resulting reconstructions of a drying slurry of NMC 622 in NMP with a PVDF binder, in order to capture particle motion.⁴⁷ Whilst the rough position of active particles could be resolved, the full electrode microstructure was not visible, again due to the short drying time of under 10 minutes.

In previous work, we have studied the three dimensional structure of mud cracks with *ex situ* X-ray CT, showing the importance of three dimensional crack structure in influencing ion transport and electrode performance, but without clear resolution of the full electrode microstructure.²⁵ In this study we extend that work, to analyse the electrode drying and mud cracking processes *in situ*, with full resolution of the electrode microstructure (Fig. 1). The relationship between mud crack formation and the local microstructural reorganisation is probed, and we demonstrate using DVC that the position of mud cracks can be predicted minutes before they are visible in tomograms.

2 Experimental

2.1 Preparation of Li-ion positive electrode slurry imaging samples

Positive electrode slurries were prepared using the active material $\text{LiNi}_{0.6}\text{Mn}_{0.2}\text{Co}_{0.2}$ (BASF), conductive additive carbon black (C-Nergy Super C65, Timcal) and polyvinylidene fluoride binder (PVDF, Solef, Solvay Specialty Polymers) dispersed in *N*-methyl pyrrolidone solvent (NMP, anhydrous 99.5%, Sigma Aldrich). The ratio of components is described in Table 1 below, where the increased C65 content and PVDF, compared with typical commercial formulations, were used in order to encourage cracking and enhance the flow behaviour of the slurry, respectively, as in our previous work.²⁵ PVDF was added to stirred NMP to prevent agglomeration and the mixture was stirred until dissolved to give an 8 wt% solution. C65 was added to a portion of this binder solution, and mixed in a planetary mixer (Thinky ARE-250, Intertronics) for 1 minute at 500 rpm and 5 minutes at 2000 rpm. NMC622 was then added alongside the additional NMP required to give the desired solids content and mixed for 10 minutes at 2000 rpm with a subsequent 3 minutes degassing at 2200 rpm in the planetary mixer. Prepared slurries were kept for a maximum of 12 hours before being replaced with a new batch to limit the effects of slurry degradation over time. To prepare samples for dynamic imaging experiments, slurries were coated with doctor blade gaps of 300, 500 and 800 μm onto a Perspex sheet with a small perforation, into which a cylindrical steel pillar of 4 mm diameter and a machined flat surface was placed. Slurry casting

used a draw down coater and a calibrated doctor blade (3600/4, Elcometer). The coated pillar was then removed from the perforated sheet by pushing it upwards from below, and transferred to the sample imaging stage (Fig. 1a). Samples were then dried at ambient temperature in the X-ray beam. Temperature measurements using a thermocouple embedded in the electrode slurry indicated that the beam induced an increase of 1–3 °C during exposure, which returned to baseline when the sample was removed from the beam. Beam induced heating affects are therefore assumed to be negligible.

2.2 Synchrotron X-ray computed tomography

Series of 3D tomograms of electrode samples of 4 mm diameter were taken at ID19 beamline for X-ray computed microtomography at ESRF, using a polychromatic beam with a peak at 70 keV. The sample was rotated through 360° with 5000 exposures of 0.015 s each, and 21 dark field and 40 flat field images. An sCMOS detector (PCOedge HS, 2560 × 2160 pixels) was used with a 5x optical magnification, giving a voxel size of 1.3 μm . The ‘half acquisition’ imaging protocol, where the sample rotation axis is offset from the detector center line and rotated over 360° instead of 180°, was used to increase the FOV to 5.15 mm × 2.6 mm (horizontal × vertical) to ensure the full 4 mm samples could be reconstructed by off-setting the axis of rotation from the centre of the field of view. Additionally, an array of 8 compound refractive X-ray lenses was used to increase X-ray flux at the sample to allow for lower exposure times and faster tomography. Tomograms were reconstructed using a standard filtered back projection algorithm using the ESRF in-house software NABU.⁴⁸

Tomographic series at Diamond Light Source (DLS) beamline i13-2 were taken using a pink beam with a peak at 30 keV and an sCMOS detector (pco.edge 5.5, 2560 × 2160 pixels) with 2x optical magnification to give an FOV of 4.5 × 3.2 mm (horizontal × vertical) and an effective pixel size of 1.625 μm , allowing for the imaging of the full sample within the field of view. Samples were therefore rotated through 180° with a step size of 0.18° and an exposure time of 0.05 s. For each tomogram 20 dark field and 20 flat field images were taken. Tomograms were reconstructed using a standard filtered back projection algorithm available in Savu.⁴⁹

2.3 Image analysis

2.3.1 Coating height and crack volume. Electrode coating heights (Fig. 2 and S2) were measured at three points across the coating and averaged, and the solvent height was taken as the highest point of the meniscus. Initial coating heights did not directly correspond to the doctor blade gap, since surface tension of the slurry in contact with the blade raised the slurry level, which reduced once the blade passed. For the 3D quantification of crack geometry (Fig. 4), tomographic images were cropped and resampled to give voxel dimensions of 5 × 5 × 5 μm in Avizo (Avizo 2023.2, ThermoFisher) in order to reduce file size and facilitate segmentation of the full volume, whilst preserving resolution of the cracks. Volumes were segmented individually in ilastik, an open source software package for the



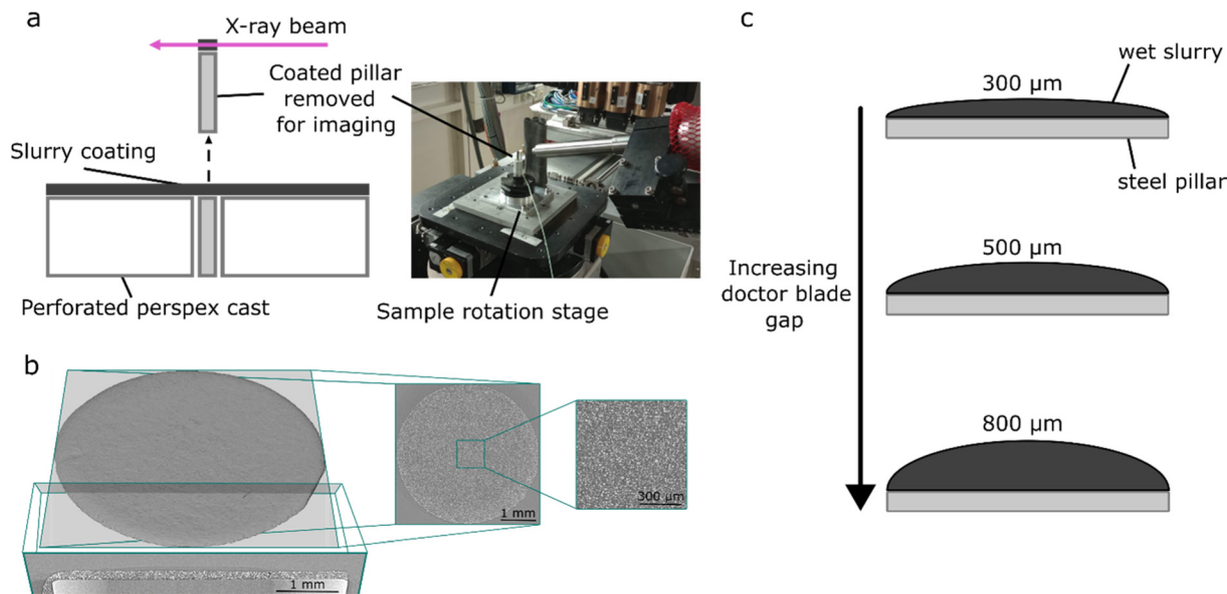


Fig. 1 (a) Schematic of the slurry coating procedure and imaging apparatus; (b) a volume rendering of the imaged coated slurry with cross-sectional cut-outs in the vertical and horizontal planes; and (c) a schematic showing the three tested doctor blade gaps coated on steel pins.

Table 1 Details of the electrode slurry formulation used in this study

Mass of solid components/%			
NMC 622	C65	PVDF	Total solid content/%
84	9	7	45

segmentation and analysis of images, using the pixel classification mode with all classifier features selected and computed in 2D.⁵⁰ Subsequent geometric analysis of segmented volumes was conducted in Avizo.

2.3.2 Electrode porosity. For the analysis of the change in porosity of the through the thickness of the full electrode (Fig. 2c and d), a series of reconstructed volumes from the first 15 scans of 300 μm electrodes, those prior to crack initiation, were subdivided into 16 equal volumes using a Matlab script to allow for segmentation whilst preserving the original voxel size. Full electrode volumes were segmented using ilastik, with two of the sixteen subvolumes were used as the training datasets, one taken from the interior and one from the edge of the sample, and the trained models were applied to the remaining fourteen subvolumes. Segmented subvolumes were recombined and image analysis was conducted in Avizo. A further subvolume, of dimensions $543 \times 511 \times 83 \mu\text{m}^3$, of the 300 μm electrode was cropped adjacent to a crack (Fig. 6) to analyse the impact of crack growth on porosity. The original voxel size of 1.3 μm was retained for this analysis, allowing for accurate estimation of particle and pore volume, given they are typically of the order of 10–20 μm in diameter.

2.3.3 Particle packing density. For the analysis of the particle packing density in Fig. S5 in cracked and uncracked

regions of the 300 μm electrode, the full electrode, immediately prior to crack initiation, was segmented in two-phases using ilastik as described above for porosity calculation. Watershed segmentation was then performed on the binary segmentation using open-source Python libraries including scikit-image and SciPy. A Euclidean distance transform was first applied to the segmented regions, followed by Gaussian smoothing to suppress noise. Local maxima within the distance map were used as markers to seed the watershed algorithm, yielding a 16-bit labelled segmentation of individual particles. A custom 3D filter was then applied to the segmented and labelled particle images. The filter convolves a cubic kernel ($18 \times 18 \times 18$ voxels) across the image, and for each voxel computes a local statistic within the kernel region. Two types of output were generated: (i) the number of distinct particles present within the local neighborhood, and (ii) the cumulative particle volume within the neighborhood. The resulting images therefore encode, at each voxel, either the local particle count or the local particle volume, depending on the chosen mode of the filter.

The fine cracks present in the electrode after 49 minutes of drying were segmented in ilastik, preserving voxel size such that the segmented image could be used as a mask to separate cracked and uncracked regions from localised particle number and volume maps. The greyscale histogram, plotted as a distribution, was used to compare particle packing density in cracked and uncracked regions.

2.3.4 Digital volume correlation (DVC). To create the displacement maps the Iterative Digital Volume Correlation algorithm in Avizo 2023 was used which estimates 3-D displacement and by correlating a reference image with a ‘deformed’ image (after a time step where movement has occurred), and mini-



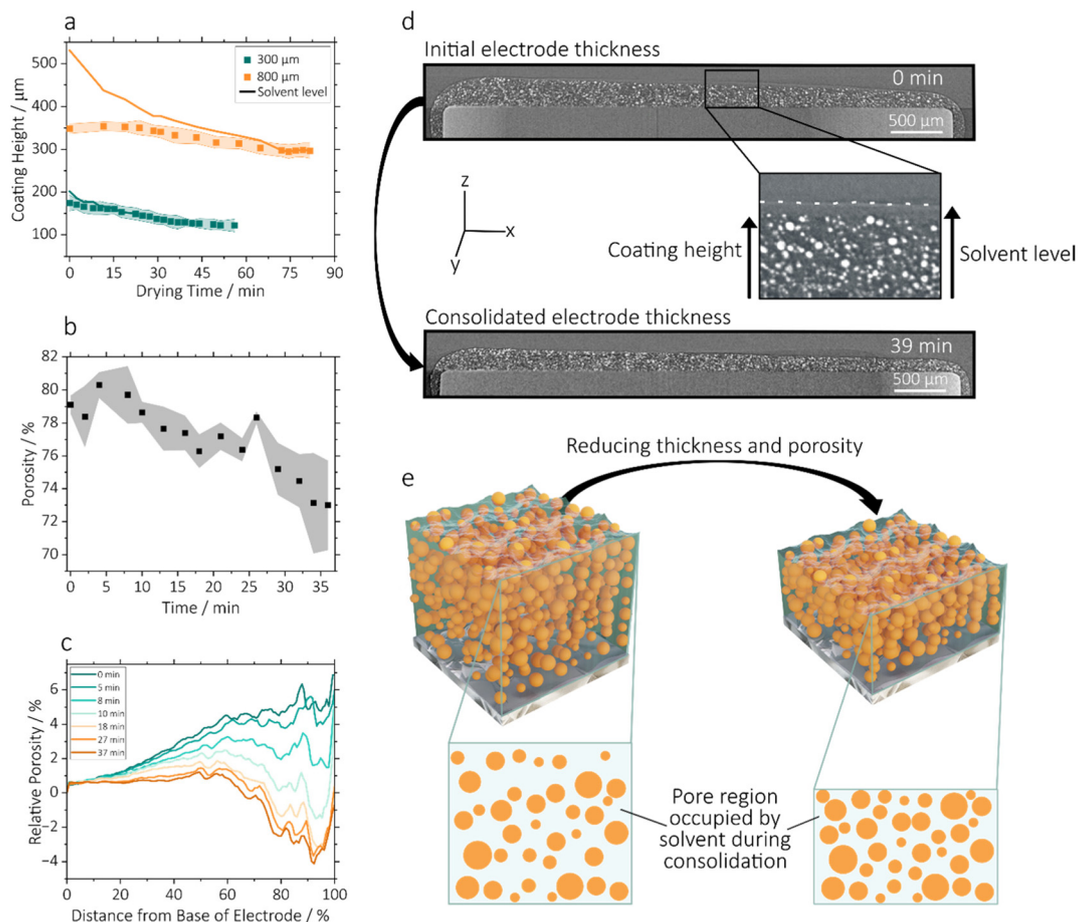


Fig. 2 Analysis of electrode consolidation and densification prior to crack formation. (a) Plot showing the change in coating height over time for 2 coating thicknesses, where solid lines indicate the solvent level above the coating and the shaded area indicates standard deviation. (b) the average porosity over time, during electrode consolidation, for the 300 μm coating, where the shaded area represents the standard deviation from 3 measurements; (c) the relative porosity as a function of distance from the base of the electrode coating, with a range of drying times shown; (d) 2D slices in the x - z plane from the centre of the reconstructed X-ray CT volume for the 300 μm coating showing initial and consolidated electrode thickness, with an additional cut out showing the solvent above the solid electrode material at the beginning of drying, indicated by a white dashed line; (e) indicative volumes showing the reduction in coating thickness and densification of solid material during this consolidation stage of the drying process, before cracking, where the solvent still occupies the space between active particles.

mining a correlation cost *via* an iterative solver (Gauss-Newton), updating nodal parameters until convergence, with a process analogous to that in Tonge *et al.*⁵¹ Regularisation and continuity constraints were used to improve robustness in high displacement regions. The iterative algorithm was chosen due to large discrepancies in displacement between the cracked region and the far-field displacements. The initial reference image was at 34 minutes, the last tomogram before cracking initiated, and subsequent iterations were used with images at 37, 39 and 42 minutes. The subset size was 1.3 μm , the same as the voxel size in the tomography image and this was constant throughout the experiment.

2.3.5 Addressing limitations of the experimental setup. In order to carry out X-ray CT imaging of the drying process with resolution of the electrode microstructure, it was necessary to dry small, circular samples on a rigid steel pin, at a lower temperature than is typical (ambient temperature rather than

over 80 $^{\circ}\text{C}$). Despite this low temperature and resultant slow drying rate, bulk particle consolidation occurred gradually, in line with observations in previous studies at higher drying rates.^{9,46,47} The 4 mm circular sample geometry may have caused the appearance of a layer of solvent above the solid electrode material, and also likely results in a greater degree of lateral solvent flow than would be observed in a bulk electrode sample. However, neither of these differences appeared to affect the formation of cracks, which were qualitatively consistent with our previous report, where samples were dried at 80 $^{\circ}\text{C}$, and showed similar quantitative trends.²⁵ Fig. S7 shows the relationship between doctor blade wet gap and dry coating thickness for the formulation used in this work with two different formulations tested in bulk electrodes. It is clear that the electrodes in the present study fall within the range expected based on this previous work.²⁵ Delamination did however appear more severe in the measured samples than in



previous work on typical aluminium current collectors, because the steel pin was inflexible and unable to deform in response to the drying stress, as well as having a greater surface roughness than a typical current collector foil.²⁵ This effect is also likely to have increased cracking severity. The behaviours observed in this system are therefore considered to be generalisable to more rapidly dried electrodes, with some minor divergences.

3 Results and discussion

For X-ray CT imaging of the electrode drying process, electrode slurries were coated using a typical draw down coater and doctor blade with gaps of 300 μm , 500 μm , and 800 μm onto steel pins of 4 mm diameter as shown in Fig. 1a and c. Samples were dried at ambient temperature over the course of 1–3 hours whilst imaging was carried out continuously, with interruptions only for dark and flat field images. In general, where cracks form they are clearly interconnected with the pore network, but are distinct from pores in size and morphology.^{25,31} In this work, cracks were defined as roughly vertical channels (perpendicular to the plane of the coating) which opened during drying, without solid material impeding their connection to the upper electrode surface. Film thickness and crack formation were studied in all three electrodes, whilst further detailed analysis of the electrode microstructure during drying and cracking, as well as DVC analysis, was carried out for the 300 μm electrode.

3.1 Initial electrode drying process

Before cracks initiate, the electrode undergoes an initial phase of drying where the coating reduces in thickness and active particles move closer together, until the active particles have fully consolidated but are still immersed in solvent. This process is represented in Fig. 2e.²⁰ Fig. 2a and Fig. S2 in the SI show the reduction in height of the three drying coating thicknesses during this period. In images of the drying 300 μm electrode, two levels of material were visible – a solid layer of electrode active material, and a solvent meniscus above the electrode, which are represented as solid lines in Fig. 2a. In the 300 μm electrode, this excess solvent layer was 30 μm thick and began reducing thickness immediately after coating. On meeting the upper solid material in the electrode, the solvent level could no longer be resolved by X-ray CT due to the highly X-ray attenuating nature of the NMC622 active material. Concurrently, the solid phase thickness reduced linearly to a final thickness of 129 μm after 37 minutes.

The solvent level was also clearly visible above the solid electrode material in the 800 μm coatings, which is shown in X-ray CT cross sections in Fig. 2d. The much larger gap between solvent and solid layers (of 180 μm) in the 800 μm coating likely resulted from a combination of the tendency of the solvent to form a curved meniscus on 4 mm pins, causing separation between solvent and solid material, a behaviour more marked at higher thickness, and possibly a degree of

slurry degradation over time which caused poorer slurry phase suspension (electrode slurries were retained for 12 hours during the experiment before being replaced). In the 800 μm electrode, the solid layer did not change in thickness until 24 minutes, at which point the excess solvent had reduced from 180 μm to 50 μm above the solid layer. The solid layer then reduced in a linear fashion until the coating thickness stopped reducing (in other words, the electrode was consolidated) after 72 minutes and at 298 μm thick. The 300 μm electrode, which began with an excess solvent layer only 30 μm thick, began reducing thickness linearly in the solid phase immediately after coating, to a final thickness of 129 μm after 37 minutes. Imaging of the 500 μm failed during the first 35 minutes of the drying process, due to an error in the centre of rotation, and the coating had largely consolidated once imaging was successful. The coating height for this sample is shown in Fig. S2b. The experiments reveal a complex interplay between slurry age, coated thickness and solvent height.

In addition to evaluating solvent height and macro-crack behaviour, the experimental approach employed in this study enables the analysis of the porosity within the electrode microstructure. Fig. 2b and c represent the porous phase of the composite electrode (where porosity is defined here for simplicity as the non-active particle volume of the coating, shown in Fig. 2e) of the 300 μm electrode during the initial 37 minutes of the drying process. Fig. 2b shows overall porosity and Fig. 2c shows the porosity as a function of the percentage distance from the base of the electrode coating, relative to the porosity at the base of the electrode. Porosity data was extracted from segmented tomograms at each time step. In Fig. 2b the porosities given are an average of those determined from three machine learning-based segmentations of each volume, in order to adapt to the challenge of segmenting a large series of images where the materials are in motion. The total porosity of the coating reduced from 79% to 73% over the course of this initial drying period. This relatively high porosity reflects the large proportion (9%) of low-density porous conductive carbon binder domain (CBD) in the slurry formulation (Table 1), which cannot be resolved directly by X-ray CT, and is therefore included within the pore phase. Porosity calculations continued up to 37 minutes, since beyond this time cracking occurred, so porosity changes no longer related to vertical electrode consolidation. Further porosity changes are analysed in Fig. 6.

In order to achieve comparison despite changing electrode thickness and variation in image characteristics between time steps, the x and y axes in Fig. 2c are normalised. The bottom of the electrode was chosen as the normalised 0 point, since theoretically its porosity shouldn't change over the drying time, as active particles were resting on the bottom surface at the time of coating. This principle can be seen in the cut-outs in Fig. 2e. At the start of the drying process the porosity was greater with increased distance from the base of the electrode, from which observation it can be inferred that the active particles had a greater degree of compaction at the bottom of the coating, and were suspended more sparsely towards the



upper surface. As drying progressed this trend reduced rapidly towards a roughly even distribution of porosity through the thickness, as can be seen between 8 and 10 minutes. Between 10 and 37 minutes the decrease in porosity in upper portions of the electrode continued, such that up to 50% of the full distance from the base of the electrode, the porosity remained roughly constant, but closer to the upper surface porosity reduced sharply, reversing the original trend. This difference was stark, with a minimum porosity at the top of electrode 4% lower than that at the base of the electrode after 37 minutes of drying. A reduced porosity at the upper surface would clearly have negative impacts on ion transport, and this effect may have been a consequence of the meniscus of the coating having lower initial density of active material causing greater compaction at the highest point as it dried. This further highlights the importance of consistent coating thickness. The steep increase in porosity in the top 5–10% of the electrode coating was due to the tendency of porosity towards 100% above the upper surface.

3.2 Growth of mud cracks

After particle consolidation, which occurs in the initial drying stage described above, mud-cracks may initiate in the electrode, particularly in thick electrode coatings. Experimental results, shown in Fig. 3, showed that crack initiation occurred after the drying electrode thickness ceased reducing (39 and 74 minutes for the 300 μm and 800 μm electrodes respectively). In the 500 μm electrode, crack initiation occurred at 45 minutes, which event is shown in Fig. S3h. This is in agreement with studies by Moorhead and Price *et al.*, who studied ceramic and silica particulate coatings respectively, showing that crack initiation is expected to occur at the point where solvent drops below the upper layer of active particles, when there is a sharp increase in stress acting on the coating.^{16,18}

Following crack initiation, shown in Fig. 3, in the case of the 300 μm and 500 μm electrodes, crack propagation was immediate, with cracks progressing from near the electrode surface downwards towards the substrate. Crack volume increased linearly in both cases until 75 minutes, as shown in Fig. 4g, in which the 3D cracking intensity factor (3D CIF; defined in previous work as the crack volume fraction²⁵) is plotted against drying time.²⁵ In both electrodes, the crack volume increased at approximately the same rate, peaking at a 3D CIF of 18.7% and 19.0% for 300 μm and 500 μm electrodes respectively. Though similar in volume, the crack morphology showed clear structural differences, with a greater degree of crack branching and inter-connectedness in the 300 μm electrode. In the 500 μm electrodes cracks developed which were larger with a lesser degree of cross-linking, as has been shown previously in Li-ion battery electrodes and a number of other particulate coating systems, where thicker coatings have larger crack spacings.^{12,25,52,53} The 300 μm electrode also formed a number of fine radial cracks around the outer edge of the electrode, not present in thicker coatings. These radial cracks are considered to be stochastic, rather than a consequence of

lateral solvent transport towards the edges during drying, as has been noted previously in drying droplets, since this mechanism would result in crack initiation at the droplet edge, which is evidently not the case in this study.^{53,54}

The air bubble present in the 800 μm electrode (Fig. 3b) which occurred due to incomplete degassing of the sample, resulted in a different pattern of crack initiation. In the first scan at $t = 0$ (Fig. S3o and q) the bubble had a maximum height and width of 84 μm and 229 μm respectively, which increased to 125 μm and 405 μm after 19 minutes. A second smaller bubble with a maximum height and width of 63 μm and 109 μm respectively was visible from the beginning of drying, and did not change in size during the initial phase of the drying process described. The larger bubble retained consistent geometry between 19 and 55 minutes, at which point air was seen to escape from the bubble towards the upper surface, which caused permanent deformation of the highly viscous, partially-dried coating. This feature underwent no further growth until 74 minutes, during which the electrode coating continued to consolidate and contained an excess of solvent (Fig. 2a). As in the other electrodes, once consolidation had completed, crack propagation occurred, forming a connection between the upper surface and the bubble, and extending across the electrode.

As a consequence, cracking in the 800 μm electrode was starkly morphologically different, and much greater in volume (the final 3D CIF was 41.5%), than the two thinner coatings, though the rate of increase of the 3D CIF was similar. The air bubble acted as the nucleation point of a crack when the drying capillary stress increased, concentrating stress in a single region. A single large crack subsequently formed through the centre of the electrode, with branching at either end. Two further examples of crack initiation at bubbles are shown in Fig. S4, where both electrodes were coated with a 500 μm doctor blade gap. The first (Fig. S4a) had a single small bubble and the other (Fig. S4b) contained seven bubbles of varying sizes. In both cases cracks nucleated exclusively from bubbles, and the largest four bubbles in Fig. S4b had similar protrusions connecting them to the electrode surface, seven minutes before crack initiation. The final crack morphology differed from the electrode of the same blade gap without a bubble shown in Fig. 4d, with fewer, less connected cracks, concentrated around the position of the bubbles. This suggests that the morphology of the cracking in the 800 μm electrode (Fig. 4f and Fig. S3o–v), where only a single crack was observed, likely resulted due to the bubble, rather than as a consequence of the trend towards fewer, larger cracks with increasing coating thickness. Although bubbles were not the target of this study, they are known to arise in manufacturing and their presence in this study has provided valuable mechanistic insight.

As shown in Fig. 3a, cracks propagate downwards from the upper electrode surface, and it was believed that they may initiate and propagate where there were less closely-packed particles since, as noted above, the binder forms between particles which are close together at the end of the drying



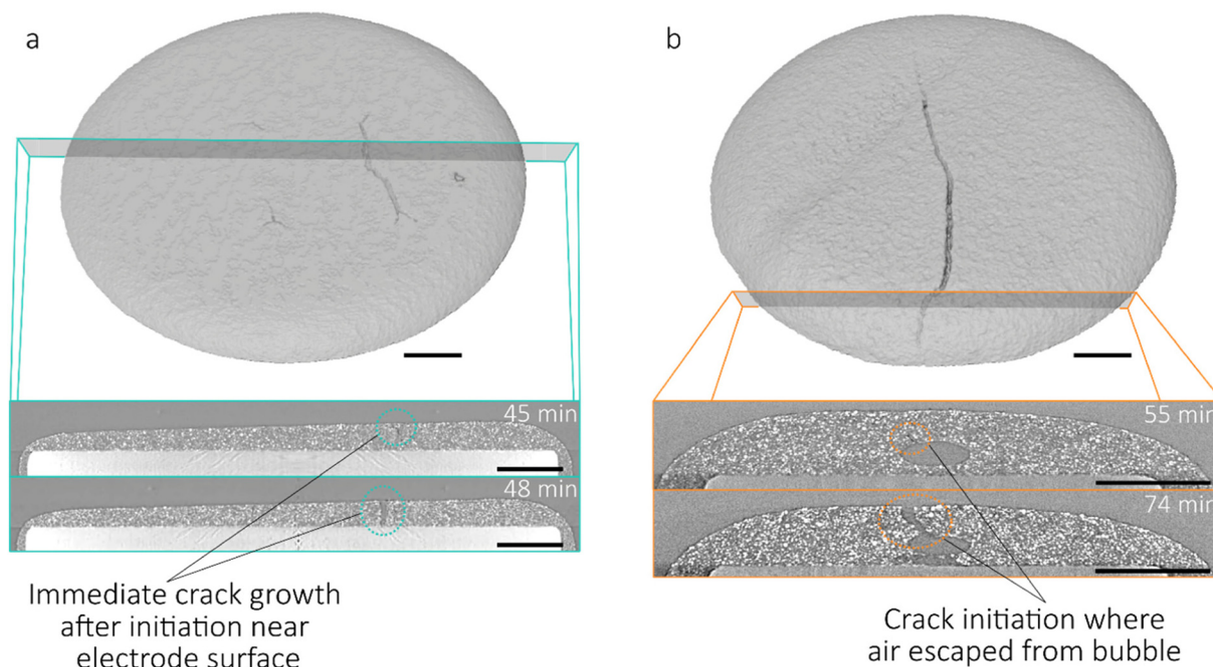


Fig. 3 3D volume renderings of (a) the 300 μm electrode and (b) the 800 μm electrode, each with two 2D slices from the reconstructed volumes showing initiation and growth of cracks. The scale bars represent 0.5 mm.

process.²¹ The temporal resolution (two minutes per image) of the tomographic imaging was insufficient to capture the precise position of crack initiation, so in order to investigate the principle that local binder formation may determine crack nucleation position, an analysis of the volume and number of neighbouring particles for each voxel in cracked and uncracked regions, as a proxy for inter-particle distance, was carried out. This analysis, which is shown in Fig. S5, found no clear correlation between crack position and the density of particles in a region. Whilst cracks may initiate at such structural features, they may be too small in comparison to the full volume to resolve by this method.

3.3 Delamination from the current collector

Alongside cracking, delamination from the substrate, in this case a steel pin, was observed. As highlighted in Fig. 5, this delamination began where cracks had grown downwards to reach the substrate, and the junctions of cracks with the substrate acted as nucleation points for delamination, which in essence is an interfacial cracking between the coating and current collector. Delamination is therefore a related but distinct process from the formation of vertical mud cracks, and the 3D CIF does not capture their different contributions to overall crack volume. Further examples of delamination are shown in Fig. S3(e, l and u).

Fig. 4h–j show the 2D CIF, which represents the cracking area fraction for each horizontal slice through the coating thickness, enabling the disentanglement of the magnitudes of mud cracking and delamination. Since the coating thickness did not change measurably in the period during which crack-

ing occurred, distance from the electrode base can be given in absolute terms. In the 300 μm electrode, between crack initiation at 39 minutes and 56 minutes, the 2D CIF remained fairly even through the thickness, with a slight increase towards the upper surface, in line with *ex situ* observations in previous work, indicating an absence of delamination.²⁵ Between 63 and 78 minutes a small increase in the 2D CIF at the current collector was seen, indicating a delamination distinct from the reduction of contact resultant between electrode coating and current collector resulting from vertical cracking. Delamination events are seen clearly in the 2D image slices from 51 and 82 minutes (Fig. S3e and g). Concurrent with this delamination, far larger increases in the 2D CIF close to the electrode surface were observed, as 2D CIF increased at the surface increased from 11% to 20.5% at the upper surface between 56 and 63 minutes. This trend developed through 70 and 78 minutes of drying, with the appearance of a region of near constant 2D CIF between 15 μm and 60 μm from the electrode base where no substantial increase was seen over time, and large increases in 2D CIF at the electrode surface, suggesting the vertical mud cracking had ceased, but drying stresses acting in parallel to vertical cracks continued to cause delamination.

A similar pattern could be seen in the 500 μm and 800 μm electrodes (Fig. 4i and j respectively), with three distinct regions within the 2D CIF profile: delamination at the base; a plateau in the middle; and greater separation at the electrode surface. Delamination occurred sooner and with greater severity in thicker electrodes; whilst for the 300 μm electrode delamination in excess of the 2D CIF plateau was notable after



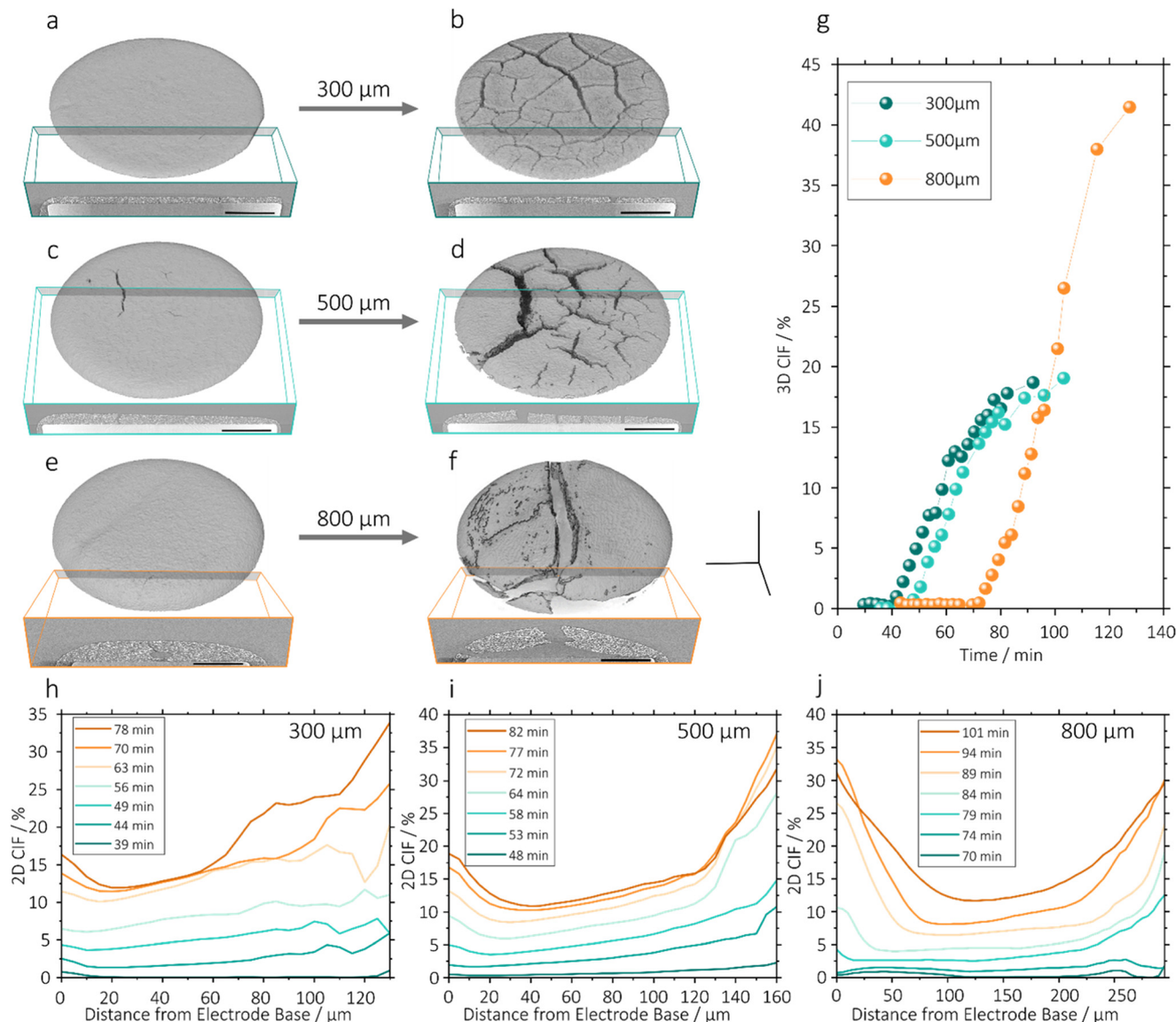


Fig. 4 Volume renderings of crack growth for electrode coatings of 300 μm , 500 μm and 800 μm doctor blade gap, where (a, c and e) show crack initiation and (b, d and f) show final cracking extent, respectively. For each volume, a vertical cut out showing a single slice of the reconstructed X-ray CT image. The scale bars and axes represent 1 mm. Plot (g) shows the increase of 3D CIF over time for each thickness, whilst (h–j) show the 2DCIF as a function of distance from the electrode base for coatings of 300 μm , 500 μm and 800 μm doctor blade gap respectively.

20 minutes of cracking, it occurred within 10 and 9 minutes for the 500 μm and 800 μm electrodes respectively. The maximum 2D CIF values at the electrode base were 16%, 19% and 33% for electrodes in ascending order of thickness. Peak 2D CIF values at the electrode surface were 34%, 37% and 30% for 300 μm , 500 μm and 800 μm electrodes respectively. The trend, showing higher 2D CIF at the surface than the base, was reversed at the surface of the 800 μm electrode after 89 minutes of drying, with delamination making an increased contribution to the overall 3D CIF. At some points the curves overlap, showing decreasing 2D CIF over time, which is likely the result of the degree of uncertainty in segmentation. These uncertainties are greater at the electrode surface and where

cracks are wider, owing to challenges differentiating between cracks and the empty space above the electrode. In Fig. 4j the two small peaks seen from 70–74 minutes, centred at 40 μm and 255 μm from the current collector, correspond to the two air bubbles present in the slurry (Fig. S3o and p). As cracking progressed at 79 minutes these signals were no longer apparent in the data, instead forming part of the baseline 2D CIF.

The higher degree of delamination in thicker electrodes indicates a greater stress resulting from the drying process, which even where cracking does not occur is likely to cause crimping around the edge of coatings on a conventional metal foil current collector. The increase in 2D CIF close to the upper electrode surface towards the end of the drying process



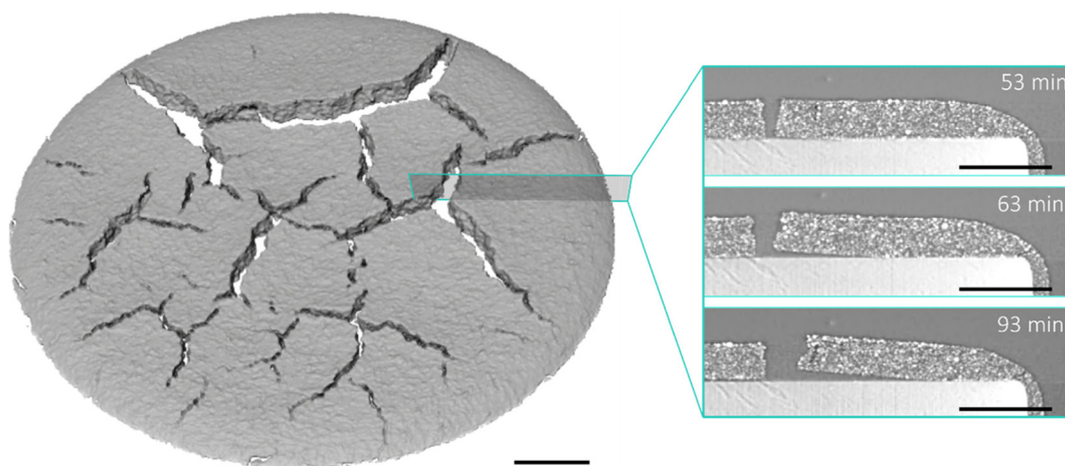


Fig. 5 3D volume rendering of the 500 μm electrode 93 minutes after drying (left) and three segments from 2D radiographs of a region of interest showing the progression of delamination of the electrode coating from the pin at three time steps. The scale bar for the 3D volume represents 1 mm, whilst those for the 2D cut-outs show 0.25 mm.

likely has two components. In part, a separation between opposite crack walls is driven by the delamination at the base of the electrode, and this in turn increases 2D CIF at the upper surface. However increased 2D CIFs near the upper electrode surface were also observed in previous work in the absence of delamination, so it can be concluded that additional expansion occurs towards the upper region of the electrode.²⁵ This through thickness analysis also suggests that crack growth does not continue below the surface when it has stopped at the surface. Surface optical imaging can therefore be used reliably to determine the period during which cracks grow, which may enable in-line surface characterisation to predict 3-dimensional features.

3.4 Impact of cracking on local microstructure

Fig. 6 shows an analysis of the impact of cracking on the porosity of the remainder of the electrode. The porosity of a $300 \times 300 \mu\text{m}$ square of the 300 μm electrode between a number of cracks, cropped between 30–110 μm of the electrode thickness to exclude the top and bottom of the electrode (Fig. 5a), was analysed during crack propagation. As was expected, the total porosity of the cropped region reduced as cracking progressed, at a decreasing rate between 39 and 78 minutes (Fig. 6b), which corresponds well with the plateauing of the overall increase of 3D CIF shown in Fig. 4g. Peaks and troughs, which can be seen in Fig. 6c, which occur consistently at each time step, appear to shift towards the current collector as crack growth progresses. This indicates a compaction of the electrode which could result from additional contraction due to drying, but is more probably caused by greater crack opening at the upper surface (Fig. 4h), which resulted in a downward compression of solid material. The anomalous characteristics of data points at 61 and 66 minutes resulted from more rapid movements of the electrode which appeared to occur during this period, resulting in less clear imaging of active particles,

reducing segmentation accuracy and increasing the calculated overall porosity. The compaction of the pore phase adjacent to cracks identified here is likely to limit ion transport in the bulk electrode which, despite the enhanced performance in cracked electrodes noted previously may cause increased inhomogeneity in ion transport and active a particle utilisation.

3.5 Predicting cracking with DVC

This compaction of particles suggests that volume correlation may yield insights into the movement of particles during mud cracking. Fig. 7 shows the combined displacement maps generated by DVC analysis of the 300 μm electrode during the scan prior to measured crack opening (37 minutes) and the subsequent two scans (39–42 minutes), and maps of displacement separated into its x -, y - and z -direction components (x and y are in plane of the coating direction, and z is the electrode height) are shown in Fig. S6. DVC displacement maps show the local displacement, given in μm , within the electrode by comparing a reference image (an initial X-ray CT scan) with a deformed image (the subsequent scan, where movement has occurred). In this case, for instance, the displacement map for 37 minutes was generated by comparison of local voxel intensities at 37 minutes (the deformed image) with those of the preceding scan at 34 minutes (the reference image). Three cracks are identified within white ellipses in Fig. 7, which show that displacement was observable by DVC prior to cracks becoming visible within the raw image. In displacement maps at 34 (Fig. S6) and 37 minutes, there was a y -direction displacement of approx. 5 μm across the full slice, due to slight fluctuations in image intensity causing the artefact during DVC computation. At 37 minutes, therefore, the combined displacement from the first crack was 1–2 μm above the background level, with a largely x - y character and very little z -component. This far smaller displacement than is likely to be observed by visual



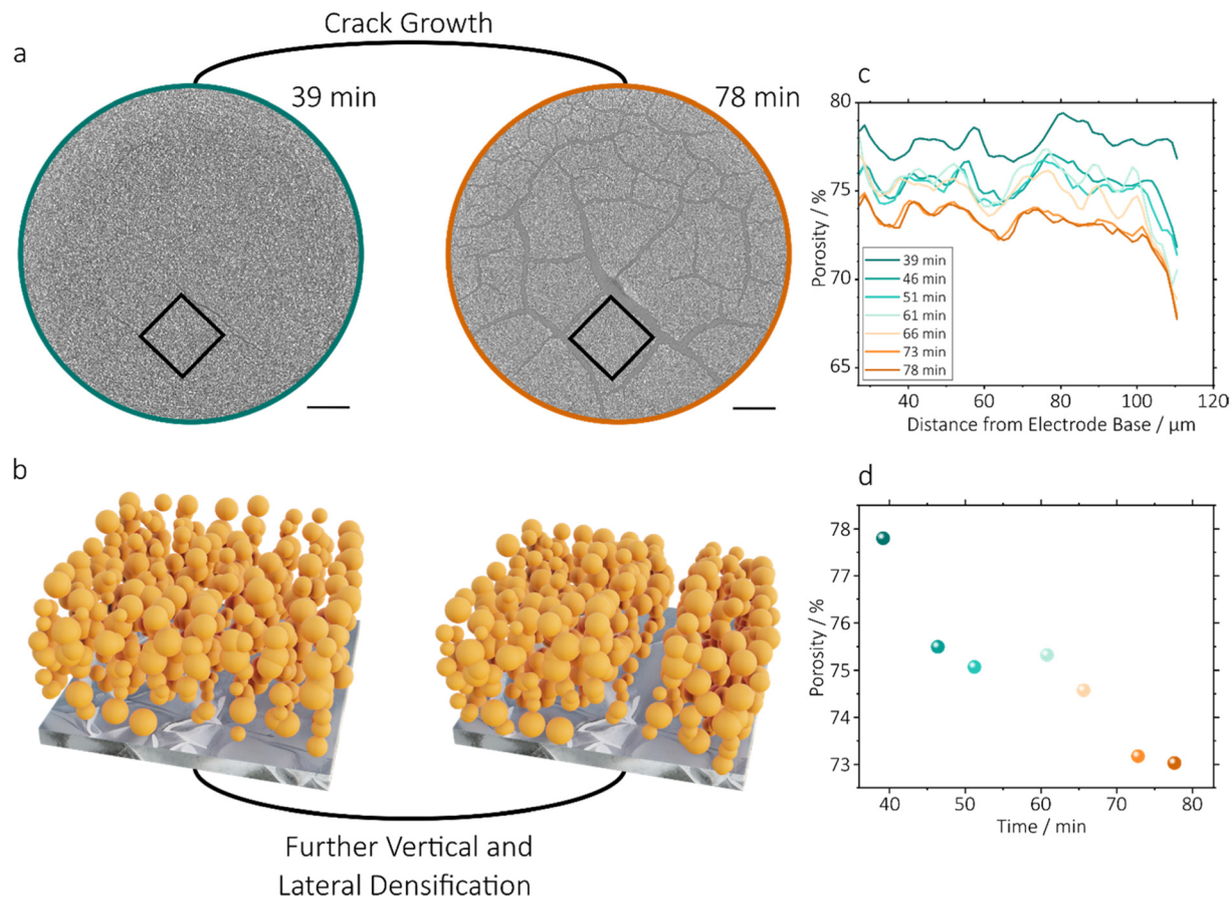


Fig. 6 Plots describing the change in porosity over time for a drying electrode coating of 300 μm thickness, where (a) the region analysed at 39 and 78 minutes, with scale bars indicating 0.5 mm; (b) is an indicative diagrams demonstrating a single crack opening and compressing adjacent particles closer together; (c) shows porosity as a function of distance from the base of the electrode over time; and (d) shows total porosity as a function of time.

comparison highlights the value of DVC for early identification of crack opening.

At 39 and 42 minutes, the background displacement observed outside cracking regions was 0 μm , so displacement magnitude could be determined directly from the map values. In these maps, larger cracks became visible, again showing that displacement could be identified before each crack itself was clearly visible in the raw image. Longer, wider cracks showed larger displacement magnitudes, and a more rapid growth rate. Whilst the first, smallest crack had a maximum displacement of 7 μm , the largest crack (approximately 3 mm in total length) had a maximum displacement of 10 μm at 39 minutes. Additionally, hot spot regions of high displacement were identified, with a maximum combined value of 13 μm at 42 minutes. The largest displacement values tended to be focused at the centre of the cracks' length, rather than at the opening tip of the cracks, where the perpendicular driving force of cracking acts (Fig. S1). At all time-steps the z-direction displacement was lesser in magnitude, with a maximum of 5 μm , and was entirely negative, indicating the downwards movement of material towards the electrode substrate.

This corroborates the observation in Fig. 6c, which suggested that cracking resulted in the forcing of some electrode material downwards. It is not believed that this effect is a continuation of the settling of particles during drying, since the coating thickness reached a plateau prior to crack initiation.

The displacement fields around cracks were observed are over a much longer length scale and encompass regions that had not yet cracked. DVC displacement could therefore be observed earlier and more clearly than cracks themselves. Given previous work correlating 3D crack structure with surface cracking, this work suggests digital image correlation, the 2D equivalent of DVC, may be a viable tool to rapidly identify crack formation before crack growth has caused critical damage to the electrode.²⁵ Coupled with an understanding of processing parameters which influence cracking, such as a coating thickness and drying temperature, this approach may enable the rapid feedback and control to halt cracking, particularly during the initial period of process optimisation during battery manufacturing. An *in situ* analysis of crack distribution and rate of opening may also enable the responsive



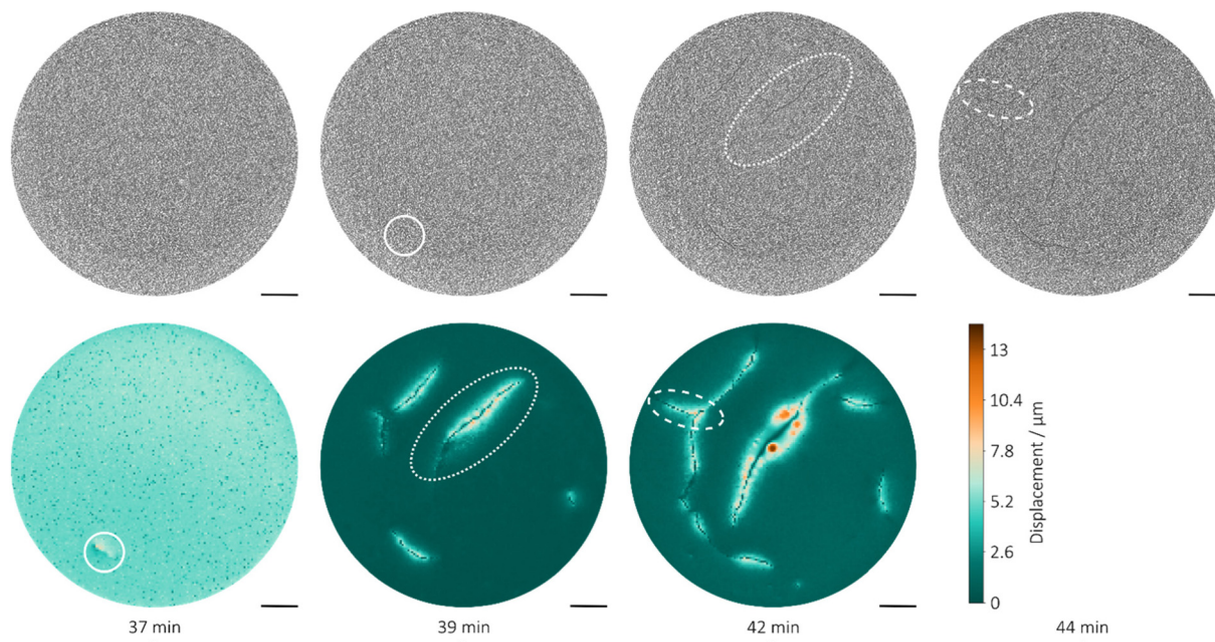


Fig. 7 DVC heat maps of total displacement for the 300 μm electrode between 37–42 minutes from a representative central slice of the volume, compared with the unprocessed slice from the reconstructed X-ray volume. Three regions indicated within white ellipses show that displacement was visible by DVC prior to the crack in that region being evident in the raw CT data.

tailoring of drying conditions to control, rather than totally eliminate, cracking.

4 Conclusions

In this work the first three dimensional time-resolved imaging of the microstructure and the mud cracking process of a Li-ion electrode during the drying process was reported using synchrotron X-ray computed tomography. This was achieved by drying at ambient temperature over 1–3 hours, coupled with the high X-ray flux of synchrotron sources, which was shown to be a highly effective approach to image changes to the electrode microstructure in detail. As in previous reports, cracking was shown to initiate at the time that the solid material had consolidated and the solvent level dropped below the upper layer of active particles. Cracking was confirmed to increase in intensity with increased coating thickness, and the cracks formed in thicker coatings were fewer in number and greater in volume. The presence of air bubbles was shown to act as a nucleating point for cracks, and concentrated the crack volume in a smaller region, which highlights the importance of thorough degassing of electrode slurries prior to drying. Furthermore, cracks which reached the electrode substrate were shown to cause the nucleation of delamination processes, which resulted in additional damage to the electrode structure. The growth of cracking was shown to compress the surrounding active material, reducing the porosity of these regions, and the combination of open pore channels (cracks) and otherwise constricted porosity may result in increased inhomogeneity of

active material utilisation in cracked electrodes. Finally, digital volume correlation was demonstrated as an effective analytical tool for the identification of crack position prior to major crack growth, and the potential of image correlation approaches as tools for process feedback and control was discussed.

These experiments provide new mechanistic insight into the drying process, which remains one of the most stubbornly energy intensive operations in battery manufacture. Whilst the experimental conditions studied here demand the use of model samples, we have corroborated our findings with studies of more conventional drying setups to provide generalisable insight. Moreover, we propose future directions for integration of advanced metrology within the manufacturing processes, which can be validated against these laboratory measurements. Collectively, this provides a pathway to optimise the manufacturing process, and the resultant electrode microstructure towards high performance and sustainable battery manufacture.

Author contributions

WJD and ARTM conceptualised and led the experiments, in which they were supported by ICA, VK, KW, PPP, BL, RSY, ZZ, JP, IK and GG. Writing and data analysis was carried out primarily WJD, with additional analysis and interpretation by ST, MPJ, ARTM, LB, and KC aided in data visualisation. AJER, JBR, DC, TSM and PRS supervised research, acquired funding and



supported in project administration. The work was reviewed and edited by ARTM, KW, JBR, RJ, TSM, PPP, BL and PRS.

Conflicts of interest

There are no conflicts of interest to declare.

Data availability

The authors affirm that the data supporting the discussion and conclusions are fully available within the article and in the supplementary information (SI). Raw X-ray CT data is available from the Diamond Light Source data gateway under experiment number MG33315-1, and from the ESRF data repository through <https://doi.esrf.fr/10.15151/ESRF-ES-1308732998>. Supplementary information is available. See DOI: <https://doi.org/10.1039/d5eb00201j>.

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