4D Analysis of the Microstructural Evolution of Si-based Electrodes during Lithiation: Time-lapse X-ray Imaging and Digital Volume Correlation

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Abstract

Silicon is a promising candidate to substitute or complement graphite as anode material in Li-ion batteries due, mainly, to its high energy density. However, the lithiation/delithiation processes of silicon particles are inherently related to drastic volume changes which, within a battery’s physically constrained case, can induce significant deformation of the fundamental components of the battery that can eventually cause it to fail.

In this work, we use non-destructive time-lapse X-ray imaging techniques to study the coupled electrochemical-mechanical phenomena in Li-ion batteries. We present X-ray computed tomography data acquired at different times during the first lithiation of custom-built silicon-lithium battery cells. Microstructural volume changes have been quantified using full 3D strain field measurements from digital volume correlation analysis. Furthermore, the extent of lithiation of silicon particles has been quantified in 3D from the grey-scale of the tomography images. Correlation of the volume expansion and grey-scale changes over the silicon-based electrode volume indicates that the process of lithiation is kinetically affected by the reaction at the Si/Li$_x$Si interface.

Keywords: Lithium-ion battery, Silicon electrode, X-ray computed tomography, Digital volume correlation

1. Introduction

Lithium-ion batteries (LIBs) are characterized by high energy density, low self-discharge, no memory effect, high efficiency and long cycle life [1]. These properties have made LIBs the preferable power source for the consumer electronics market [2], which drives forward the LIB technology together with the sectors of transportation and grid energy storage [3]. The development of LIB technology is expected to have an important effect on the popularization of environmentally friendly energy systems such as solar cells, wind power generation and electric vehicles [4]. There is, thus, a growing interest in the development of a new generation of LIBs, which should match the needs of high capacity, stable cyclability and safety of those technologies. In this context, intense research has been devoted to the investigation of materials to substitute existing electrodes, such as graphite for the negative electrode and lithium-cobalt oxide, lithium-manganese oxide and lithium iron phosphate for the positive electrode [5–7].

In the aforementioned traditional electrode materials, the lithiation process takes place by means of an intercalation mechanism. That is, ions occupy interstitial sites within a relatively rigid structures. For example, the graphene layers in the graphite electrode. In a charged battery with a graphite negative electrode, the lithium ions occupy interstitial spaces up to a stoichiometric composition of LiC$_6$, presenting a theoretical capacity of 372 mAh g$^{-1}$ [8, 9].

Lithium-alloying materials, such as silicon, tin or aluminium have been proposed as candidates to substitute or complement the graphite electrode, due to their significantly higher gravimetric capacities. Among these alloying anode materials, silicon (Si) is one of the most promising due to its exceptionally high capacity and its abundance in the earth’s crust [6]. The theoretical fully-lithiated phase, Li$_x$Si, has a capacity of approximately 4200 mAh g$^{-1}$. In practice, it has been observed that the highest lithiated phase of Si at room temperature corresponds to Li$_{3.75}$Si,

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Table 1: Comparison of anode materials for LIBs [4, 8]

<table>
<thead>
<tr>
<th>Materials</th>
<th>Li</th>
<th>C</th>
<th>Si</th>
</tr>
</thead>
<tbody>
<tr>
<td>Density (g cm(^{-3}))</td>
<td>0.53</td>
<td>2.25</td>
<td>2.33</td>
</tr>
<tr>
<td>Lithiated phase</td>
<td>Li</td>
<td>LiC(_6)</td>
<td>Li(_{1.75})Si</td>
</tr>
<tr>
<td>Gravimetric capacity (mAh g(^{-1}))</td>
<td>3862</td>
<td>372</td>
<td>3579</td>
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<tr>
<td>Volumetric capacity (mAh cm(^{-3}))</td>
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<td>840</td>
<td>2440</td>
</tr>
<tr>
<td>Volume change (%)</td>
<td>100</td>
<td>12</td>
<td>&gt;300</td>
</tr>
<tr>
<td>Potential vs. Li ((\approx) V)</td>
<td>0</td>
<td>0.05</td>
<td>0.15</td>
</tr>
</tbody>
</table>

with a specific capacity of approximately 3579 mAh g\(^{-1}\), with a number of intermediate amorphous phases with stoichiometry Li\(_x\)Si (with 0 \(\leq\) x \(\leq\) 3.75) [10–12]. Table 1 compiles some properties of the anode materials of interest in this study.

Alloying electrodes experience extreme volume changes during lithiation, over 300% for Si [13]. These huge volume changes produce mechanical deformations and damage to the fundamental components of a battery, causing rapid and severe capacity loss leading to battery failure. Irreversible capacity loss, particularly during the first lithiation step, has been reported to be caused mainly by: (1) the loss of active material, as the active particles crack and pulverize leading to the disconnection of the alloy particles from the current collector, binder and the conductive carbon; (2) the continuous formation of solid-electrolyte interphase (SEI) and volume changes that produce the detachment of the SEI, which reforms in every cycle causing irreversible loss [8].

Different options have been proposed to mitigate the negative effect of these volume changes. One of the most promising solutions is the use of silicon nanoparticles, nanotubes or nanowires, where the reduced particle size seems to produce more stable lithiation/delithiation cycles. However, large specific surface areas lead to high levels of SEI formation. In this context, Liu et al. [14] proposed a yolk-shell structure to mitigate volume changes; the structure consists of a carbon-based shell around each nano-sized Si particle aimed at forming a stable mechanical framework upon Si volume expansion. The well-defined void space allows the Si particles to expand freely without breaking the outer carbon shell, therefore stabilizing the solid-electrolyte interphase on the shell surface [15].

The mechanical consequences of the electrochemical processes that take place in battery electrodes play a crucial role in determining battery performance and lifetime [16]. A better understanding of these processes is essential to achieve new generations of LIBs based on alloying electrodes. In this study, we use 4D X-ray computed tomography (XCT) imaging techniques to visualize the evolution of Si-based electrodes during the first lithiation step.

XCT enables high-resolution, non-invasive, non-destructive 3D imaging for the qualitative and quantitative inspection of material structure and properties. XCT has been demonstrated for the visualization of the battery components’ micro-structural evolution at multiple length scales and as an effective tool for diagnosing battery failure mechanisms post-mortem [17, 18]. XCT has previously been used to explore the micro-structural properties of electrode materials in LIBs [19–22]. Furthermore, consecutive 3D images during the operation allows the 4D (3D plus time) analysis, including the possibility to carry out tests under in-situ or in-operando conditions (XCT during, for example, electro-chemical tests). For cells similar to those studied in this work, Gonzalez et al. [5, 23] investigated the evolution of Si electrodes during the first lithiation and Sn electrodes during the first lithiation/delithiation cycle.

In this study, we use Digital Volume Correlation (DVC) (e.g., [24, 25]) to quantify the mechanical deformation of the fundamental cell components (electrodes and separator) during the lithiation process. DVC has been demonstrated as a useful tool for the diagnosis of electrochemical devices [18, 26], but has not been used yet (to the authors knowledge) for the study of lithium-alloying electrodes.

2. Experimental and Methods

2.1. Electrode preparation and cell assembly

Lithium-silicon half cells (Fig. 1) were assembled within thin-walled X-ray transparent PFA Swagelok cells [21]. Electrode slurry was prepared by mixing micron-sized Si powder (300 mesh, 99% purity), conductive graphite and polyvinylidene fluoride (PVDF) binder in the respective weight ratios 80:10:10 with n-methyl-2-pyrrolidone (NMP) using a mortar and pestle. All chemicals used in the slurry preparation were acquired from Pi-Kem Ltd., UK. The
resulting slurry was then dip-coated onto one tip of a steel current collecting pin (ca. 3.2 mm diameter), and the coated pins were placed in a vacuum oven to dry overnight at 80 °C. The active Si loading was estimated by weighing the current collecting pins before and after dip coating. Half cells were assembled in an argon-filled glove-box (O₂ < 2 ppm, H₂O < 0.5 ppm) with metallic lithium foil counter electrode and a borosilicate flass-fibre separator (Whatmann GF-D grade, GE). The electrolyte used consists of 1M LiPF₆ dissolved in a mixture of ethylene carbonate (EC) and ethyl methyl carbonate (EMC) (in the ratio 3:7 by weight). The cells were assembled at the Electrochemical Innovation Lab of the Department of Chemical Engineering, University College London.

The Si electrodes were tested against lithium metal. Accordingly, in these half cells, the lithium metal acts as the negative electrode and the silicon as the positive electrode [27].

2.2. Electrochemical and XCT tests

An assembled cell with theoretical capacity of ca. 7.45 mAh (calculated based on the Si content and assuming lithiation to Li₁₃₋₇₅Si), was galvanostatically discharged using a potentiostat (Ivium Compactstat, Ivium Technologies). XCT was performed using Xradia Versa MicroXCT-520 laboratory tomograph at the 4D-Imaging lab (Division of Solid Mechanics, Lund University). The cell was partially discharged at successive intervals in 10 consecutive steps at 25 μA. The first step lasted 10 h and the electrode was lithiated 3.36 %. Consecutive steps had 20 h duration each, and the electrode was lithiated ≈ 6.72 % in each step. Further lithiation of the cell after the 10th step was not possible, probably due to breakdown of the separator, as discussed in the results sections. Figure 2 shows the cell potential readings for the 10 steps of partial lithiation of the Si electrode.

A total of 11 XCT scans were carried out (one for the initial state and after each of the 10 partial discharge steps). For the scans, the source and the detector were set at 15 mm distance (in front of and behind to, respectively) the center of the sample, achieving a pixel size of 1.7 μm using a 4× objective. Scans were carried out with a source tube...
voltage of 45 kV with exposure time of 30 s per projection, acquiring 2001 projections per scan. The reconstructed
tomograms consisted of 16 bit grey-scale image stacks of 2000 × 2000 pixels per image.

2.3. DVC analysis

DVC was used to quantify the micro-structural evolution of the cells between the successive discharge steps. The
DVC technique consists of the comparison of two volumes (3D image acquired via XCT) to identify the displacement
field that maps the deformed volume on to the original volume. The DVC provides full 3D tensor displacement fields,
which are then interpreted following continuum mechanics to provide 3D tensor-stain maps.

The DVC analysis was carried out using the DVC software Tomowarp2 [28], Python-based code developed be-
tween the Universities of Grenoble and Lund. This DVC algorithm includes sub-pixel resolution of the displacements
and consists of the following steps. First, a 3D grid of nodes is defined in the reference volume. Around each node, a
correlation window (subvolume centred on the node in the original volume) is defined. The corresponding correlation
window is sought within the deformed volume, over a 3D region denoted as search window (see Fig. 2.3.a). During
the search, the subvolumes are compared by a correlation coefficient calculated at each position.

One of the challenges addressed in this research is to investigate whether it is possible to use DVC to follow the
volume changes in the Si electrode during the lithiation process, which is undergoing chemical transformation, with
associated attenuation changes, in addition to displacements. In this context, it is assumed that the lithiation steps
applied to the cells are small enough to allow a proper correlation between two consecutive images in the sequence
of acquisitions. Consequently, it should be possible to carry out a study of the incremental deformation of the battery
during the lithiation process.

3. Results and Discussion

3.1. XCT results

Figures 4 and 5 show, respectively, 6 vertical and 3 horizontal cross-section images from the 11 acquired image
volumes. In both figures the grey-scale range has been adjusted to the threshold limit between 13750 and 18250 (16
bit). This threshold leaves the lithium metal near the lower saturation limit. Materials with higher density, such as Si
particles and glass fibre separator, are marked by high grey-scale values. A small number of particles are observed
with X-ray attenuation much higher than Si (bright saturated in the images). These particles were assumed to be
impurities in the original Si powder.

The lithiation process is clearly accompanied by a significant volume expansion of the Si-based electrode. The
volume of the electrode at 64.5 % lithiation is nearly three times that of the initial state. The separators mechanical
stability contributes immensely to the safety and performance of LIB cells. The huge volume changes induced the
rupture of the separator in the middle part with through-going cracks. It was impossible to further lithiate the electrode
over the value of 64.5 % due, presumably, to a short-circuit between the two electrodes.

In the cell studied here, the Si electrode was tested against a lithium metal electrode, whose volume decreases as
the lithium oxidises into lithium ions. During the lithiation, the separator suffered a vertical displacement towards the
lithium electrode, pushed by the growing Si-based electrode. The separator also suffered compression, indicating than
the expansion of the the Si electrode exceeds the volume released at the lithium metal electrode.
Figure 4: (a-f) Vertical cross sections of the cell at different lithiation degrees. 6 images (16 bits images and grey-scale of bulk with threshold limits 13750 - 18250) are shown within a total of 11 XCT acquisitions. Inset brackets indicate the acquisition number and the degree of lithiation of the electrode. Yellow rectangle in (d) indicates the limits of the subvolume analysed in Fig. 7 (g) Colour depth histogram of a subvolume in the Si-based electrode, capturing the change in attenuation at different states of electrode lithiation. Coloured boxes, to the right, indicate the subvolumes over the expanding electrode where the colour depth histograms were computed.
Moreover, in the tested cell, the separator exhibited uneven initial thickness indicating certain level of compression in the central part of the cell. This initial compression, resulting from the manufacturing process, is related to a decrease of the separator’s porosity in that central region. As a consequence of the the local decrease of the lithium diffusivity in that central region, lithium ions seems to go across the separator preferentially through the outer region. The lithiation front (i.e., the front of decrease of the attenuation in the images) is observed to progress in radial direction from the upper-outer towards the inner-central part of the electrode.

Despite the narrow grey-scale range needed to appreciate contrast between the phases, the change of the attenuation in the Si electrode during the lithiation process is clearly observed. As the lithiation takes place, the Si particles react with the lithium ions to form lithiated-Si phases (Li\textsubscript{x}Si) which have lower X-ray attenuation and, therefore, appear darker. The impurities did not experience either volume nor attenuation changes during the process. Figure 4.g shows the grey-scale histogram for subvolumes corresponding to the same region in the Si electrode (i.e. excluding the separator and the lithium foil region) taking into account the dilation of the electrode. The fresh cell presents a histogram with two overlapped peaks centred on 15800 and 17000. The lower grey-scale peak is related to the electrode matrix (mainly consisting of conductive carbon, binder and pores filled with electrolyte) and the outer boundary of the silicon particles. The peak in the histogram centred on a grey-scale of at 17000 is related to the core of the silicon particles (non-lithiated Si). The histogram evolution during the lithiation process shows a clear decrease of the attenuation, towards a single peak centred on approximately 14900, which is inferred to correspond to the lithiated-Si phase.

As the lithiated-Si phase has an X-ray attenuation factor very similar to the electrode’s conductive matrix, it is easier to distinguish the shrinking Si core of the particles than the expansion of the outer lithiated-Si region. Figure 6 shows a selected particle at 3 different states of electrode lithiation. The chosen particle, which is also indicated in Fig. 5, does not experience significant attenuation and volume changes until the fifth acquisition. It must be noted that the degree of lithiation of the particle is not the same as the lithiation degree of the bulk electrode, since the lithiation does not take place homogeneously within the cell.

Fig. 6 shows the evolution of the particle through the lithiation and, in particular, shows detail in the reducing size of the unlithiated core. This figure is constructed as follows: first, the selected particle was identified and isolated in the chosen acquisitions (7, 9 and 11) and subvolumes around the particle were extracted. Subsequently, the subvolume was thresholded to make voxels with a grey-scale value less than 16250 transparent in the final volume renderings shown in the figure. The resultant visible volume of the particle (in Fig. 6.a-c) which corresponds to the unlithiated Si in the particle was approximately 18000, 15000 and 9000 voxels respectively (the latter being the combined volume of the two pieces of the divided particle in Fig. 5.c).

Figure 6.d shows the same particle shown in Fig. 6.c, but constructed using a dilated mask, to cover not only the segmented particle but also the surrounding of that particle. The surrounding region is faded (presented with certain level of transparency) in order to distinguish the inner and the outer part. The fade function is included in the inset.
histogram. Fig. 6.d shows that there is a clear difference in the attenuation level of the particle between the inner unlithiated core and the outer lithiated region, and therefore, it is possible to make a segmentation between the Si core and the lithiated-Si phase.

3.2. DVC results

In this section, results from the DVC analysis of the sequence of acquisitions are presented. The results were obtained using a 3D grid with 25 voxels node spacing and a DVC window of ±25 voxels in X, Y and Z. Correlation windows were, therefore, cubic subvolumes of 51 voxels side (≈ 86 µm side). The DVC computation was carried out over a subvolume located in the central part of cell with size 1000 × 1000 × 500 voxels (≈ 0.845 mm³), including the Si-based electrode, the separator and a fraction of the lithium foil (see Fig. 7.d).

Figure 7: DVC results between the acquisitions 6 (≈ 30.9 %) and 7 (≈ 37.6 %). (a) 3D volume strain field. (b) local grey-scale changes at the same positions as the local DVC subvolumes. Axis in mm. X and Y axis referred to the centre of the cylindrical cell. Z axis referred to the current collector below the Si-based electrode. (c) Local relation between the volumetric strain and the grey-scale changes for the electrode region. d) 3D rendering of the subvolume used for the DVC analysis. The limits of the subvolume are indicated in the figures 4.d and 5.b

Figure 7.a shows the results obtained using the DVC algorithm between two consecutive acquisitions of the cell, namely acquisitions 6 and 7, corresponding to global lithiation degree of ≈ 30.9 % and ≈ 37.6 %, respectively. In Fig.
methods. The grey-scale change follows a linear trend with respect to the global lithiation degree. Figure 7.a-b correspond to the Si-based electrode. The DVC (Fig. 7.a) indicates positive volumetric strain (expansion) during the lithiation in this region. Meanwhile, Fig. 7.b shows that there is a clear decrease in the grey-level (decrease in X-ray attenuation) in the electrode. This is consistent with the global results presented in Fig. 4.g and in correspondence with the Si lithiation producing a decrease in the particle density.

It is possible to study the relation between the volumetric strain and the grey-scale changes in a local scale, i.e. at the different positions of the electrode where the degree of lithiation may be different due to the in-homogeneous lithiation. Fig. 7.c shows an example of the local relation between the volumetric-strain and the grey-scale changes in the region of interest selected within the Si electrode, for the same values presented in Fig. 7.a. At a given step, the local grey-scale change and volumetric strain are linearly related.

Based on the different mechanical behaviour of the Si-based electrode and the separator, it is possible to do a segmentation between the two in the volumetric strain map. In particular, we consider that the region of the subvolumes exhibiting positive volumetric strain (expansion) belongs to the electrode. Figure 8 shows the results of the DVC for each pair of consecutive acquisitions, showing only the section experiencing volume expansion. The figure shows that the local volumetric strain in the Si electrode increases as the lithiation progresses. The global incremental increase in lithiation over each of these steps is the same, but it can be seen that the degree of volumetric strain calculated over each increment increases as the total degree of lithiation increases.

Figure 8.k presents the computed accumulated values for the volume expansion within the region of electrode studied as a function of the average degree of lithiation. A polynomial fitting curve is included, indicating that the computed volumetric strain relates to the bulk electrode lithiation degree in a quadratic manner. The volume expansion increases with lithiation, meaning that the Si electrode exhibits more volume expansion at higher degrees of lithiation. At ≈ 64.5% electrode lithiation, the volume increment is calculated to be ca. 155%. Extrapolating the second degree curve to 100 lithiation gives a predicted volume expansion of 350, which is consistent with the volume expansion reported in the literature [5, 6] for the full lithiation of Si particles.

Figure 8.k also shows the global grey-scale changes in the electrode, referred to the original average grey-scale value. Local grey-scale changes were computed in two different ways: using each of the corresponding subvolume from the 3D strain analysis, and then averaged for the entire electrode at each lithiation step, and averaging from the histogram values presented in Fig. 4.g. Local grey-scale results show strong agreement between the two methods. The grey-scale change follows a linear trend with respect to the global lithiation degree.

The lithiation process of silicon particles can be considered as the combination of different processes in series; namely: (1) the redox reaction at the electrolyte/silicon interface, (2) the diffusion of lithium through the lithiated phase, and (3) the chemical reaction at the Si/Li, Si interface [11]. As the lithiation process is carried out galvanostatically, the overall rate of the reduction reaction of lithium on the surface of the particles along the electrode is constant. However, due to the observed in-homogeneous transport of lithium ions through the separator, the rate of lithiation of the particles depends on the position in the Si-based electrode. The particles in the outer region lithiate faster than the particles in the centre of the cell.

XCT results showed that it is possible to distinguish between the non-lithiated core and the lithiated shell with diffusing lithium atoms. DVC results, in turn, show that the volume changes of the Si electrode increase with the lithiation degree in a quadratic manner, while the grey scale decreases linearly. These observations suggest that: (i) the change in local grey-scale value can be used as a proxy for the local lithiation degree; (ii) the lithiation process is affected by the available surface area at the Si/Li, Si interface where the lithiation reaction (i.e., the breaking of Si-Si bonds and formation of Li-Si bonds) takes place, since, if the lithiation rate is constant, the change in surface area will be quadratic.
4. Conclusions

This work presents results for XCT observations of the 4D microstructural evolution of Si-based porous electrodes during the first lithiation process. The presented results demonstrate that it is possible to carry out the quantification of the micro-structural evolution of lithium battery components using XCT imaging in combination with DVC.

In Si-based porous electrodes, as the lithiation takes place, the X-ray attenuation decreases and the volume of the active particles expands. Both effects are efficiently tracked by XCT and DVC. In the cell tested here, the coupled electrochemical and mechanical phenomena causes the breakdown of the separator, and the eventual short-circuit of the cell. Furthermore, the non-homogeneous compression of the separator is related to preferential paths for the transport of lithium ions during the cell operation.

The quantifications of the volume and grey-scale changes from the images show that the volume changes of the Si electrode increase with the lithiation degree in a quadratic manner, while the grey scale decreases linearly. These observations suggest that: (i) the change in local grey-scale value can be used as a proxy for the local lithiation degree; (ii) the lithiation process is affected by the available surface area at the Si/Li$_x$Si interface where the lithiation reaction takes place. Furthermore, extrapolation of the quadratic trend in volume change to 100% lithiation indicates a maximum volume expansion of $\approx 350\%$, which suggests that operating the cell below certain threshold limit of the lithiation degree may increase the durability and increase the specific capacity of the cells.

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References


Figure 8: Volumetric strain computed via DVC, corresponding to each pair of consecutive acquisitions of the tested cell (indicated in brackets). Only the region exhibiting volume expansion (the Si electrode) is shown here. The color bar is common for all images, units in percentage of volume change. Axis in mm. X and Y axis referred to the centre of the cylindrical cell. Z axis referred to the current collector below the Si-based electrode. (k) Volume increment of the electrode (blue-circles, left axis) and grey-scale changes (right axis) with referred to the original non-lithiated electrode. Grey-scale changes computed through data from the histogram in Fig. 4.g (green-squares) and averaging the local grey-scale changes computed via the DVC code (red-triangles)