Simulation and Measurement of the Current Density Distribution in Lithium-Ion Batteries by a Multi-Tab Cell Approach

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A single-layered NMC/graphite pouch cell is investigated by means of differential local potential measurements during various operation scenarios. 44 tabs in total allow for a highly resolved potential measurement along the electrodes whilst the single layer configuration guarantees the absence of superimposed thermal gradients. By applying a multi-dimensional model framework to this cell, the current density and SOC distribution are analyzed quantitatively. The study is performed for four C-rates (0.1C, 0.5C, 1C, 2C) at three temperatures (5°C, 25°C, 40°C). The maximum potential drop as well the corresponding SOC deviation are characterized. The results indicate that cell inhomogeneity is positively coupled to temperature, i.e. the lower the temperature, the more uniform the electrodes will be utilized.

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Within the past decades, demand for lithium-ion batteries in mobile applications has significantly increased. Due to their well proven performance as well as their stability in long-term usage, lithium-ion batteries became the technology of choice for electrochemical energy storage devices.1–3 Still, the specific energy density as well as cycle life are constantly being optimized by either commercializing new active materials, electrolytes and additives or by reducing the fraction of non-active parts within a battery. Often, this corresponds to thicker electrodes or larger form factors leading to capacities of up to 100 Ah per cell. In these large format cells, severe gradients in current density and temperature distribution can occur along the electrode stack,1–9 which might provoke a performance loss during operation due to inhomogeneous utilization. Also non-properly adapted thermal conditioning can have a crucial impact on the performance of larger cells.10–12

Modeling of internal distributions of potential and temperature along the electrodes is quite challenging, since even to calculate only a few cycles, a lot of computational resources are required for fully resolved models. In literature, there are many examples for spatially resolved multi-dimensional modeling approaches, which aim at representing the cell’s internal behavior in terms of potential, current density, state of charge (SOC) and temperature distribution. Unfortunately, all of these examples lack a detailed, i.e. spatially resolved experimental validation, which is capable of tracking internal variables instead of measuring the surface temperature at a few spots and considering the overall battery’s terminal voltage. Also, only a few examples of direct measurements of the internal current density distribution were published so far. Zhang et al. built a specific LFP/graphite prototype cell for this purpose. A segmented cathode was used for analyzing the current distribution during discharge at varying C-rates and temperatures. This setup allows for a precise monitoring of the current of each electrode element individually. Large deviations in SOC of up to several percent were identified during the process of discharge. However, it is not clear how the cell temperature influences these results as no data analysis is shown regarding this boundary condition. Klink et al. also built a lab-scale prototype cell, which was set up for analyzing the current density distribution through the electrode layer thickness instead along its width and height. In their work they investigated the distribution of current during the formation process of a graphite electrode. A similar approach was chosen by Ng et al., who analyzed multiple working electrodes to show that even low current densities can cause severe gradients within the electrodes. In order to gain further insights into the behavior of a commercial cylindrical cell, Osswald et al. modified a 26650 LFP/graphite cell in such way that any of the four tabs along a 1.7 m long electrode can be accessed separately, whilst still showing stable behavior for more than 200 cycles. It was shown that potential drops as large as 10 mV occur applying a discharge current of 0.1 C to the most extended tab configuration. Further, a multi-dimensional model for the modified cell was built and the local potential measurement method was proven to be applicable as a tool for current density analysis as well as model validation purposes. Minor deviations between simulation and measurement were assumed to be related to the use of approximated parameters for the cell’s heat capacity and thermal conductivity and the implementation of a simplified LFP electrode representation assuming very low diffusivity for its phase change mechanism. Waldmann et al. extended this study by experimentally investigating different tab patterns and their effect on the temperature development between the cell center and its surface. In continuation of these efforts, we studied the current density distribution along NMC/graphite electrodes in large format batteries allowing for an enhanced spatial resolution as well as an improved distinction between thermal and electrochemical processes.

For that purpose, a specific prototype cell following a form factor of 500 × 100 mm² was designed, which contains state-of-the-art electrode materials and was assembled by means of manufacturing processes at high industrial standards. By monitoring 10 tab pairs at each edge of the single-layered NMC/graphite pouch cell, the quality and resolution in measuring local potentials is maximized. Such a multi-tab cell can be used as measurement setup and validation tool for spatially resolved models in order to study effects related to electrode configuration and tab patterns, local relaxation processes26–28 local impedance effects.29 Due to the specific cell design, any thermal gradient along the electrode can be avoided and, therefore, the super-imposed effect of temperature inhomogeneity on cell performance is eliminated. A general study on the current density distribution within that cell during discharge derived from measuring local potentials and their deviation at varying C-rates and temperatures is presented here. In the second part of this work, we apply our multi-dimensional modeling framework to reproduce the measured data and investigate the

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internal current density and SOC distribution theoretically depending on temperature and applied C-rate.

**Experimental**

*Design considerations.*—The cell is designed to analyze potential gradients within the current collector foils along a considerably long electrode in order to provoke a significant, i.e. detectable potential drop. As we are focusing on the cell’s length, the width is kept rather small. The dimensions of the cell’s current collectors are set to 500 × 100 mm². Even larger gradients can be provoked by longer electrodes, but in our case the size of the used climate chamber sets a restriction to that dimension. Thus, the cell length is a compromise of applicability and maximum signal amplitude. For further details of the cell configuration, see Fig. 1.

The cell comprises a LiNi0.33Mn0.33Co0.33O2 cathode and a graphite anode, as this material combination nowadays serves as a standard in many applications. The electrode loading is 2 mAh cm⁻² for the NMC and 2.2 mAh cm⁻² for the graphite electrode. That capacitive oversizing is assumed to help avoiding lithium plating during charging. Additionally, the anode is 2 mm larger in width and length than the cathode in order to suppress overvoltage peaks at the electrode edges. In total, the cell delivers about 920 mAh at a 0.1C discharge. Its nominal capacity is rated as 0.8 Ah, thus the current at 1C rate is 0.8 A.

The electrodes consist of single-sided coatings on current collector foil, which is aluminum on the cathode and copper on the anode side of the cell. Two larger tab pairs with 15 mm width are added at the smaller side of the cell and serve as reference tabs. By adding 10 tab pairs along the electrode, voltage differences can be measured during operation of the cell. At every position, the positive as well as the negative current collector potential can be measured. For improved signal quality of that measurement, large tabs (5 mm width) were connected to the current collector foil by ultrasonic welding. Additionally, we performed a study on the distribution of the contact resistances and the ohmic contributions of the current collector by analyzing a non-coated foil with exactly the same tab pattern and materials.

**Cell production.**—The NMC (111, LiNi0.33Mn0.33Co0.33O2) electrode consists of 86% active material (BASF), 6% PVdF-binder and 8% conductive additives. All percent values correspond to weight fractions. The cathode slurry was based on organic solvent and coated and dried by using a Mathis coating line. The graphite anode is made of 96% active material (Hitachi), 2% binder and 2% conductive additive. The anode slurry was based on water and coated and dried by using the same Mathis coating line. The electrolyte is 1 M LiPF6 in EC/DMC (1:1 by mass) plus 2% VC as additive. The separator is a PP monolayer (Celgard 2500). At the shorter side of the cell Al and Ni plated Cu tabs (width: 15 mm, thickness 0.2 mm) and at the longer side Al and Ni tabs (width: 5 mm, thickness 0.1 mm) were connected by ultrasonic welding. The electrodes were packed in aluminum composite foil and dried in vacuum at an elevated temperature for 48 h. The cell was filled with 10 ml of electrolyte in an argon atmosphere. The formation process of the cell was carried out by running four cycles (2 × 0.1C, 0.2C, 1C; charge: CC&CV, discharge: CC). Finally, the cell was charged to 3.7 V and degassed.

Coin cells for open-circuit voltage (OCV) measurement of NMC and graphite were produced by punching out small disks with a diameter of 14 mm from the same electrode sheets used for the multi-tab cell. The counter electrode used within the coin cells is a lithium metal foil with a thickness of 400 µm and a diameter of 15 mm. The coin cells contain the same electrode material, separator and electrolyte as the multi-tab cell and the same formation procedure was applied prior to operation. Four cells were built for each electrode. Half-cell potential measurements were performed at 25°C with a current rate of 0.02C, while applying three cycles (after initial formation cycle). The mean value between the third charge and discharge cycle corresponds to the OCV, taken as a reference for the model. Strictly speaking, averaging between a low rate discharge and charge cycle does not reflect the OCV of an electrode perfectly due to a remaining hysteresis as well as non-symmetric polarization. By application of very small currents (e.g. 0.02C), these impacts can be reduced to an acceptable level. In this case, we assume this technique to be sufficient for that type of analysis, since the main characteristics of the OCV are conserved, whereas the error is substantially small. Nevertheless, slight variations in precise positioning of graphite induced OCV changes.

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**Figure 1.** Schematic representation of the multi-tab cell configuration together with the dimensioning of the single-layered NMC/graphite electrode pair (top). The cell has a reference tab pair at the left and right side of the cell (Ref) and 10 pairs of positive and negative measurement tabs which are distributed equally on the top and bottom side of the cell. To assure mechanical stiffness, the cell is mounted on an acrylic back plate which is fixed to an aluminum frame. The reference tabs are connected by cable shoes which are screwed to the tabs whereas the potential measurement tabs are connected by alligator clips (bottom).
might be assigned to this recording technique. Table I summarizes the cell parameters.

**Measurement setup.**—After formation, the cell is mounted onto its fixture and electrically contacted with alligator clips which are further connected to 2 mm banana sockets. At the reference tabs, cable shoes are connected in order to assure lower resistances for the terminal current. In this study, the cell current was only applied at one of the two reference tab pairs. In order to avoid any inhomogeneity due to non-symmetric operation over many cycles, the side was switched at the end of the measurements, while the contact resistance from the cable shoes as well as the electrical performance, capacity and cell voltage were monitored. No deviations were observed, so we assume a high reproducibility of the presented measurements, independent from the reference tab configuration.

In Fig. 2, the measurement setup is shown. A Basytec CTS is used for controlling the temperature. Before any discharge operation, the cell is charged at a rate of 0.1C followed by a CV phase until the current drops below 0.01C. A relaxation period of 4 h is included between charging and discharging, ensuring enough time for compensation of any inhomogeneity.

**Model Development**

We applied our multi-dimensional modeling framework to the cell described above. Since the cell is in quasi-isothermal conditions, no thermal model is considered here. The model represents the cell domain by considering 11 equally discretized elements for the corresponding tab pair region. This implies that 11 Newman-type models are coupled to a two-dimensional current collector domain, where the negative and positive current collector potentials are calculated. The basic idea behind the model coupling is the exchange of averaged values within a discretized element Ak. First, the potential is calculated within the current collector domain and transferred to the 11 Newman models. In the same time step, the resulting current density from the Newman model is fed back to the 2D model as source term. In Fig. 3 the modeling setup is shown containing the most important equations. For more details, the reader is referred to our previous work. The relevant model parameters are summarized in Table II. The geometry corresponds to the experimental cell domain, as shown in Fig. 1 and Fig. 2. During the measurement period the cell lost about 3% of its initial capacity. To compensate for that, the initial SOC is fitted accordingly. The model is solved in commercial finite-element method software Comsol Multiphysics 5.2a, which takes about 4 min for a 1C discharge with an i5 processor and 16 GB RAM.

**Results and Discussion**

**Measured discharge profiles.**—In Fig. 4 the cell voltage at the reference terminal (a) as well as the corresponding local potentials for the negative (b) and the positive current collector (c) are plotted as a function of discharged capacity during a 0.1C discharge at 25 °C. The cell voltage shows only the signal information restricted to the terminal (reference tabs), while the local potentials illustrate the potential distribution along the electrode. In preliminary tests we confirmed that the signals on both sides of the cell are identical. Thus, the potential is shown for only one side throughout the manuscript.

It can be seen that a differing current collector thickness and conductivity lead to corresponding amplitudes of the potential drop. The positive current collector foil faces the higher loss in potential due to its lower absolute conductivity. The factor of absolute potential drop in between both foils should be about

$$\frac{\Delta U_{\text{neg}}}{\Delta U_{\text{pos}}} = \frac{20 \mu \text{m} \times 3.6 \times 10^7 \text{ S m}^{-1}}{18 \mu \text{m} \times 5.8 \times 10^7 \text{ S m}^{-1}} = 0.69$$

**Table I. Multi-tab cell parameters.**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>NMC</th>
<th>Graphite</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of single-coated layers</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Electrode thickness (µm)</td>
<td>74</td>
<td>62</td>
</tr>
<tr>
<td>Area specific capacity (mAh cm⁻²)</td>
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<td>Porosity</td>
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<td>Particle size D50 (µm)</td>
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<td>22</td>
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<td>Width (mm)</td>
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<td>Length (mm)</td>
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<td>500</td>
</tr>
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<td>Separator type</td>
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<td>Separator thickness (µm)</td>
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<td></td>
</tr>
<tr>
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<td>Copper</td>
</tr>
<tr>
<td>Current collector thickness (µm)</td>
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<td>18</td>
</tr>
<tr>
<td>Number of tab pairs</td>
<td>10</td>
<td>10</td>
</tr>
</tbody>
</table>

![Figure 2. Measurement setup applied to multi-tab cell consisting of a battery tester, multi-channel voltmeters and four PT100s on its surface. The distance in x-direction between the reference tab and the tab of the negative current collector with number 01, 05 and 10 is indicated.](image-url)
**Model Setup**

![Diagram of Model Setup](image)

**Figure 3.** Model setup applied to the Multi-Tab Cell: Eleven 1D electrochemical models (based on Newman’s work) are coupled to a 2D current collector domain by a potential-current exchange averaged within one element \( A_k \). Only the main equations are shown, for more details the reader is referred to our previous publication.

assuming common literature values for specific conductivities of copper and aluminum. Compared to the measured values in Figs. 4b and 4c, a ratio of about 0.5 can be seen in between the potential drop on the positive and negative current collector side, which confirms the assumed conductivity values. The deviation can be explained by the higher welding spot resistance at the reference tab in the positive current collector foil, yielding 1.4 mΩ in contrast to 0.2 mΩ.

### Table II. Model parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>NMC</th>
<th>Separator</th>
<th>Graphite</th>
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</thead>
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<tr>
<td>Open-circuit voltage</td>
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<td>measured</td>
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<td>Initial SOC</td>
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<tr>
<td>at 5°C</td>
<td>0.40(^f)</td>
<td>0.72(^f)</td>
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<tr>
<td>at 25°C</td>
<td>0.39(^f)</td>
<td>0.76(^f)</td>
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<tr>
<td>at 40°C</td>
<td>0.41(^f)</td>
<td>0.75(^f)</td>
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<tr>
<td>Theoretical max. concentration (mol m(^{-3}))</td>
<td>51500(^a)</td>
<td>31370(^a)</td>
<td></td>
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<tr>
<td>Active material volume fraction</td>
<td>0.38(^a)</td>
<td>0.45(^a)</td>
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</tr>
<tr>
<td>Solid state diffusivity (m(^2) s(^{-1}))</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>at 5°C</td>
<td>2.2e-14(^f)</td>
<td>2.2e-14(^f)</td>
<td></td>
</tr>
<tr>
<td>at 25°C</td>
<td>3e-14(^f)</td>
<td>3e-14(^f)</td>
<td></td>
</tr>
<tr>
<td>at 40°C</td>
<td>5e-14(^f)</td>
<td>5e-14(^f)</td>
<td></td>
</tr>
<tr>
<td>Reaction rate constant (m s(^{-1}))</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>at 5°C</td>
<td>0.7e-11(^f)</td>
<td>0.7e-11(^f)</td>
<td></td>
</tr>
<tr>
<td>at 25°C</td>
<td>2e-11(^f)</td>
<td>2e-11(^f)</td>
<td></td>
</tr>
<tr>
<td>at 40°C</td>
<td>4e-11(^f)</td>
<td>4e-11(^f)</td>
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<tr>
<td>Charge transfer coefficient</td>
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<td>Electrolyte salt diffusivity (m(^2) s(^{-1}))</td>
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<td></td>
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<tr>
<td>Electrolyte conductivity (S m(^{-1}))</td>
<td>Fitted function(^{31})</td>
<td>Fitted function(^{31})</td>
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<td>Electrolyte’s activity dependence</td>
<td>Fitted function(^{31})</td>
<td>Fitted function(^{31})</td>
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<tr>
<td>Electrolyte’s transference number</td>
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<td></td>
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<tr>
<td>Tortuosity</td>
<td>3.5(^c)</td>
<td>2.5(^c)</td>
<td>5(^c)</td>
</tr>
<tr>
<td>Porosity</td>
<td>0.55(^c)</td>
<td></td>
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<tr>
<td>Current collector’s conductivity (S m(^{-1}))</td>
<td>3.6e7(^{30})</td>
<td>5.8e7(^{30})</td>
<td></td>
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<tr>
<td>Temperature dependency (K(^{-1}))</td>
<td>0.004(^{30})</td>
<td>0.004(^{30})</td>
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</table>

\(^a\)assumed value.

\(^b\)estimated from Ref. 33.

\(^c\)fitted value.

**Figure 4.** Discharge profile at 0.1C and 25°C measured at reference terminal (a) and corresponding eight differential potentials for the negative (b) and the positive current collector (c) plotted versus discharge capacity. Tab numbers are highlighted on the right side.
monotonically decreasing behavior over a wide SOC range, the visible oscillations can be ascribed to the graphite’s OCV steps between the plateaus. At about 350 mAh the first characteristic voltage oscillations appear due to the change from stage 1 to stage 2 in the graphite electrode. Due to the overbalanced graphite electrode, this point is shifted to a higher SOC. It can be also observed that these oscillations are more pronounced with increasing distance to the reference tab, where the current is applied. Further, there is a slight increase in the overall slope of the differential voltage curves with increasing distance to the reference tab. This might be related to the relative polarization of the electrode section, which should be the lowest opposite the reference tab. In this region, the relative change due to an electrode OCV change is considerably large in comparison to the ohmic potential drop within the current collector, which results in a higher sensitivity to any variations in the electrode OCV.

To further investigate the OCV influence, the graphite OCV (a) as well as the local potential measurement from the multi-tab cell (b) are shown in Fig. 5, plotted as a function of the specific capacity. The OCV curve in the upper part (a) corresponds to coin cell measurements at a 0.02C charge vs. Li/Li+. The lower part (b) displays a 0.1C discharge of the multi-tab cell. Both measurements were performed at 25 °C. The OCV is shifted by 14.7% to the left in order to visualize the effect of slope changes on the local potential. Since the graphite is overbalanced by 10% and an irreversible capacity loss occurs during formation, 14.7% is a fairly realistic value. Obviously, the graphite OCV stages provoke the differential voltage oscillations along the current collector foil in the multi-tab cell. As soon as an electrode section closer to the reference tab enters a higher intercalation stage, the potential difference to the next segment increases until that segment enters the same state of charge. Then, the voltage difference reduces to its initial value, which is mainly caused by an ohmic drop. Especially at lower lithiation degrees of graphite, more pronounced oscillations occur in the differential signal along the current collectors. The highlighted peak in the signal at about 0.65 mAh cm\(^{-2}\) can be clearly assigned to the phase transition from stage 1 to stage 2 in graphite. The features at a discharge capacity of about 1.3 mAh cm\(^{-2}\) reflect a more oscillating behavior due to more abrupt slope changes in the OCV and the absence of a significant plateau area. The OCV of NMC is not shown here as there are no distinct slope changes throughout the used lithiation range which would result in such an oscillating behavior.

Two major results can be concluded so far. At extended electrode length, the potential drop follows a more decreasing scheme during discharge being not only defined by the ohmic drop within the current collector foil. Slope changes within the active material’s OCV cause potential oscillations, being more pronounced at further distance to the reference terminal. When using NMC and graphite as electrodes, these oscillations can be related to the graphite’s OCV curve. A maximum potential drop of about 0.7 mV can be measured per 46 cm electrode length at a low rate of 0.1C in the positive current collector foil.

To study the rate dependency, Fig. 6 contains the signal from the farthest tab #10 for discharge current rates of 0.1C (a), 0.5C (b), 1C (c) and 2C (d) at 25 °C. In general, the ratio of potential drop within the current collector foils for differing C-rates corresponds well to Ohm’s law. Moreover, the shape of the signal depends strongly on the current rate. In the first part of the discharge process, the slope at 2C (d) seems to be shaped smoothly in comparison to that at 0.1C (a). Further, the clear peak assigned to the graphite’s phase transition is hardly visible at higher rates. We can assume that the higher potential drop along the current collector foil causes a less pronounced OCV influence and results in a flatter contour. This corresponds well to our investigation within modified commercial cells, where the same trend was observed. Especially at high DOD values of as large as 80% the potential drop characteristics change distinctly toward higher rates, which can be seen in a considerably flat profile at 2C. In general, oscillations due to the OCV influence of graphite can be only identified at substantially low C-rates such as 0.1C. Zhang et al. discussed this behavior and concluded that the OCV dependency is dominant over the potential drop along the current collector foil at lower C-rates. As soon as the rate is high enough, the ohmic drop determines the differential potentials to be more monotonically shaped. The order of magnitude of the potential loss is about 3.6 mV per 46 cm negative current collector length for 1C, resulting in a drop of approximately 0.08 mV per cm electrode length. This might be important when designing cells with electrodes of up to meters in length in cylindrical configurations while adding only a few tabs due to manufacturing costs. This critically affects the active material utilization along the electrodes. For materials with very flat OCV such as LTO or LFP, this effect is even more pronounced, since larger gradients in current density occur due to the dominating potential drop along the electrodes.

### Indication of SOC variations

Differential voltage analysis (DVA) is performed for the local potential signals along the current collector and plotted for a 0.1C discharge at 25 °C in Fig. 7. In the upper part of the diagram (a), the overview of the entire DOD range is given. Main peaks can be seen at a DOD of about 37% and at DODs higher than 70%. More detailed overviews of these DOD regions can be found in the lower part of the diagram (b and c). In contrast to (b), the curves (c) show a slight variation in the peak position of the

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Figure 5. Half-cell potential of graphite (OCV) vs. lithium measured with the aid of coin cells at 0.02C and 25 °C shifted by 14.7% due to the electrode balancing, plotted as a function of the specific capacity (a). Corresponding local potentials (\(U_{\text{neg,10}}\) and \(U_{\text{pos,10}}\)) at tab #10 of the negative and positive current collector, measured relative to reference tabs within the multi-tab cell during 0.1C discharge at 25 °C (b). The vertical lines mark the region of the phase change within the graphite OCV.

Figure 6. Rate dependency of the farthest local potential (tab #10) at the negative current collector foil for 0.1C (a), 0.5C (b), 1C (c) and 2C (d) plotted versus DOD at 25 °C.
corresponding tabs. In general, the peak shifts to larger DODs and is more pronounced with increasing distance to the reference tab. If we take a closer look to the x-position of the local maximum in the DVA (refer to Fig. 7c), it turns out that the temporal DOD difference between tab 3 (yellow) and tab 10 (blue) is in the range of 0.5% to 1%. This order of magnitude corresponds well to our recent investigation of a modified commercial cell, where we proposed about 1% to 2% SOC difference within a 128 cm LFP/graphite electrode segment at a 0.1 C discharge at 25°C. However, the signal quality is too low for a reliable quantitative analysis of DOD distribution, which relates to our measurement setup. Due to the limited size of the climate chamber, the data logger had to be placed outside. As a result, the signal quality suffers from the small differences in temperature fluctuation between inside and outside of the chamber, which are small in general, but when a sub-μV resolution is required, an inevitable noise occurs. Additionally, the relatively low control rate of the climate chamber and the accompanying temperature variations caused a negative effect on the preciseness of the voltage measurement setup and prevented a reliable DVA. For that reason, the setup was optimized by attaching a copper plate which is heated and cooled by two powerful Peltier elements, which yield an absolute temperature variation of less than 0.05 K. Experimental work on this setup is ongoing. Here, we are restricted to state that at 0.1 C, deviations are visible, but can hardly be analyzed quantitatively. We can estimate the order of SOC deviation to be around 1% over a distance of approximately 26 cm (distance from tab 03 to tab 10) at a 0.1 C discharge at 25°C.

In comparison, the differential analysis at a 2C discharge and 25°C is shown in Fig. 8a. Throughout the entire DOD range nearly no characteristic peaks occur except at about 87%. The peak is highlighted in the lower part of the figure (b). It can be seen that the point of intersection varies with the position along the electrode following the same scheme as for 0.1 C, the larger the distance, the later the intersection. So far, it can be stated that a SOC variation due to a peak shift within the DVA can be identified at a discharge at 0.1 C as well as at 2 C. The intersections vary by about 0.5% to 1% DOD, comparable to the observations made at a 0.1 C discharge. As stated before, these results are rather of a qualitative nature than a quantitative one. More generally, it can be stated that the artificially prolonged LFP/graphite cell shown in Ref. 5 is capable of showing higher SOC deviations, similarly identified by peak shifts of the DVA signal. This is due to the flat LFP voltage profile as well as the total distance between the measurement points of 128 cm. To overcome this restriction related to the experiment for the multi-tab cell setup, our modeling framework was applied after validating it via the local potential measurement. Following this approach, we can give a more quantitative representation of the current density distribution within the cell in Model Analysis section.

Temperature dependency.—In Fig. 9, the temperature dependency of the local potential is presented for the farthest tab at three temperatures (5°C, 25°C and 40°C) for a rate of 0.1 C (a) and 2 C (b). Here, the DOD is defined by the capacity of the cell at the corresponding temperature and discharge rate to allow for a reliable comparison. An increase in temperature causes the local potentials to oscillate more while showing a higher absolute potential drop along the current collector foil. At 0.1 C and 5°C, almost no peak or variation in the signal can be detected, which corresponds to a very homogeneous discharge procedure without any significant disturbances along the electrode. In contrast to that, the elevated temperature at 40°C emphasizes a peak between a DOD of 30% and 40% as well as several peaks after a DOD of 60% (Fig. 9a). The peak at about 90% discharge capacity is visible for all temperatures, but is much more pronounced at 40°C. The difference between the absolute potential drops seems to have a linear correlation to the temperature difference between them. If we include an averaged linear temperature coefficient of 0.004 K⁻¹ for

![Image](https://example.com/image1.png)

**Figure 7.** Differential voltage analysis of the local potential signals at a 0.1 C discharge and 25°C (a) with two highlighted regions close to a DOD of 37% (b) and 80% (c). The line colors correspond to Fig. 4 and the maximum of tab position 03 and 10 is marked by a line.

![Image](https://example.com/image2.png)

**Figure 8.** Differential voltage analysis of the local potential signal at 2C discharge and 25°C (a) with a highlighted region close to a DOD of 87% (b).

![Image](https://example.com/image3.png)

**Figure 9.** Temperature dependency of the local potential at the farthest tab (#10) within the negative current collector for three temperatures (5°C, 25°C and 40°C) at a 0.1 C discharge (a) and a 2 C discharge (b).
the electrical resistivity of copper,\(^3\) a temperature variation of 20 K corresponds to an overall decrease in conductivity of 8% and a difference of 15 K results in a decrease in 6%, respectively. If we take the initial voltage amplitudes of 0.33 mV, 0.36 mV and 0.38 mV from Fig. 9a as a reference for the temperatures, the difference between them corresponds quite well to the relative change in conductivity of copper:

\[
\frac{0.36 - 0.33}{0.33} \text{ mV} = 0.091
\]

\[
\frac{0.38 - 0.36}{0.36} \text{ mV} = 0.056
\]

That means that at a 0.1C discharge rate, the temperature dependent current collector resistance causes the absolute potential drop whereas other processes lead to stronger oscillations at higher temperatures. Here, we assume enhanced kinetics and mass transport to have more influence on the electrodes’ sensitivity toward small potential disturbances. Since the resistivity of aluminum has nearly the same temperature dependence of 0.004 K\(^{-1}\), these results hold for both current collector foils.

To correlate these findings with the current rate, the same set of three temperatures was applied at 2C discharge rate, which is shown Fig. 9b. As stated before, the higher the rate, the less pronounced the oscillations in the voltage signal along the electrode become. The spread in between the absolute potential drop follows the same trend as for 0.1C, namely a linear increase of potential drop depending on temperature due to the increasing current collector’s resistance. Considering the initial potential values, the difference yields 7.5% between 5°C and 20°C, and 3.3% between 20°C and 40°C (Fig. 9b). Both are slightly smaller than those at 0.1C, which might be an indicator for additional effects, mainly kinetics and mass transport. That means, the potential drop does not only follow Ohm’s law by temperature dependent resistivity, but corresponds to non-linearly increased kinetics within the active material as well as changed transport parameters of the electrolyte. Unfortunately, the cell’s current rate is limited to 2C to ensure adequate lifetime, otherwise this relation could have been stressed in terms of higher rates. Finally, the combination of low temperatures and high C-rate seems to show the lowest oscillations without any deviations. This corresponds to the findings of Zhang et al.\(^7\) and Osswald et al.,\(^19\) where the current density distribution is more uniform at lower temperatures.

Model Analysis.—Current Rate Dependency.—In Fig. 10 the comparison of simulated and measured data for 0.5C, 1C and 2C at 25°C is shown. In general, the model is capable of representing the cell voltage at the reference tabs precisely, which is shown in the upper part (a) of Fig. 10. At a discharge rate of 0.5C, the modeled cell potential drops to the cutoff voltage slightly earlier than in the measurement. Additionally, the simulation is not able to fit the first part of the discharge at 2C. This combination suggests a superimposition of non-ideally suited mass transport parameters and the electrodes’ OCV
or its initial concentrations, respectively. Still, we face a limitation in terms of using measured transport parameters from the electrolyte applied to the cell. In this study, a comparable set of fitted parameters from literature is transferred to our cell setup, assuming sufficient representation of any concentration dependent electrolyte transport limitation. Further, although the OCV of both electrodes was measured with coin cells versus lithium, the initial stoichiometry remains a fitting parameter. Additionally, some contribution from the edge area of the electrodes as well as side reactions within the cell might influence this initial stoichiometry parameter. In sum, without having precise transport parameters, kinetic parameters as well as morphologic information of the electrode, an ambiguity inevitably remains. But in this case, we focus rather on the local distribution of variables such as the OCV characteristics of the active material, in contrast to that, the measured local potentials as illustrated in Fig. 9 show a larger amplitude in the oscillating behavior at 40°C, which is more distinct compared to 25°C. This cannot be represented by the model in full spread. Though, a general statement in terms of temperature dependency of the current density distribution can be deduced. For higher cell temperatures, the inhomogeneity of the current density distribution increases, ranging from 5% at 5°C up to nearly 11% at 40°C, referred to a DOD of 90% (see Fig. 11c).

Preliminary sensitivity studies revealed that the reaction rate constant and its influence on the exchange current density in the Butler-Volmer kinetics affect the amplitude of the visible oscillations in the local potential measurement (refer to Fig. 11b) as well as in the current density distribution (refer to Fig. 11c). During the measurement, the cell was kept at the listed temperatures for several weeks. Unfortunately, that led to a capacity fade superimposed onto these results. At the end of the measurements, we recognized a distinct rise of the absolute potential drop correlated to the cathode, whereas the spread in between the local potentials remained the same. Hence, we assume that additional contact resistances between the aluminum foil and the cathode coating increase and interfere our interpretation of the reaction rate constant as well as its temperature dependency. That might be the reason for the model’s underestimation of the potential oscillation and the small difference of the current density variations between 25°C and 40°C (refer to Fig. 11c). Since the cell is still under operation, we can clarify that point of view as soon as we open up the cell.

**SOC dependency.** In addition to the analysis of the current density distribution, the SOC distribution within the electrode can be calculated by the model. Fig. 12 shows the simulated absolute SOC deviation in percent points along the electrode for all C-rates. Generally, the deviation increases during discharge, whereby small variations occur due to the OCV characteristics of the active material, in this case mainly due to graphite. Interestingly, the inhomogeneity declines after reaching its maximum at about 70% of the discharge time. This is in correspondence with the findings of Zhang et al. They also observed a homogenization period at the end of the discharge, where regions far from the reference tabs deliver higher current densities. Especially at high rates, whether for discharge or charge conditions, this gives implications about the effect of a constant current protocol on electrode utilization within substantially prolonged electrodes. The maximum deviation is in the range of about 1.2% (Fig. 12c) and occurs at a DOD value of about 70%. An effect of varying temperature on the SOC distribution is visible, but is rather small at lower C-rates. In contrast to that, a clear dependency can be seen at 2°C. At 40°C a deviation of about 1.3% occurs, whereas at 5°C a substantially smaller value of 0.7% appears. Further, the deviation remains at a value of about 0.5% for all temperatures at the end of the discharge. Contrarily, at 5°C the cell gets nearly homogenized at the end of discharge. That means, in combination with higher current rates, the temperature has a major effect on the cell inhomogeneity and a uniform active material utilization. Especially for interrupted discharge profiles at a DOD of 70%, the effect is most pronounced. If normalized to its electrode length, a maximum deviation of about 0.6% can be identified for the distance of 46 cm at 1C and 25°C, which corresponds to 0.013 percent per cm. In our ongoing study, we are investigating the influence of a thermal gradient on this result. It can be assumed that a higher temperature at the reference tabs will cause an even more inhomogeneous SOC distribution along the electrodes.
Figure 11. Simulation (solid lines) and measurement (symbols) of cell voltage (a), measured and simulated potentials at local measurement points #01, #05 and #10 (b), and simulated current densities at the corresponding points (c) for 1C at temperatures of 5°C, 25°C and 40°C.

It is worth stating here that the current collector foils used in our cell are relatively thick and are usually applied to double coated electrode sheets. Thinner current collector foils were not available at the time. This implies that the potential drop along the electrodes will more or less double throughout the discharge procedure which involves a more inhomogeneous utilization of the electrodes. It can be therefore assumed that the SOC variation will also increase when a double coated electrode or half the current collector thickness is utilized.

Conclusions

In this study, the method of using local potentials as internal state variables for gathering insights into the current density and SOC distribution within lithium-ion batteries was applied to a prototype NMC/graphite pouch cell. With a size of 500 × 100 mm² and 44 tabs, the cell allows for a detailed differential voltage measurement along the electrodes during common operation scenarios. Due to its single-layered cell design, the heat capacity is low in contrast to the heat exchange via the cell’s surface and tabs, so that a constant temperature is maintained even at high discharge rates of up to 2C. We performed an analysis for four discharge rates and three temperatures, applying a standard tab configuration to the pouch cell. Additionally, our multi-dimensional modeling framework was used for simulating the internal current density and SOC distribution, after being validated by means of the local potential measurements along the electrode. For parameterization, most of the geometric cell values are measured, whereas the electrolyte properties and electrode kinetic parameters are taken from literature and fitted depending on C-rates and temperature. The study illustrates that the current density peaks are following the electrodes’ characteristics of the OCV, clearly visible at lower C-rates. As soon as the C-rate is large enough, the peaks vanish and the local potential is predominantly shaped by the ohmic drop along the current collector foil. Extended by the model results, our study further reveals that temperature has a positively coupled effect on the cell inhomogeneity during discharge. This can be related to enhanced kinetics and mass transport within the electrode and separator domain as well as an increased current collector foil resistance. The deviation in SOC was about 0.013 percent per cm at a 1C discharge at 25°C and 70% DOD, taken from the maximum difference between the first tab (#01) and the last tab (#10). Generally, the SOC distribution is most inhomogeneous at a DOD of 70% and equalizes at the end of the discharge, where only a small deviation remains. Our results were compared to preliminary studies which used modified LFP/graphite cells. Further
studies will focus on more precise electrochemical parameter determination in order to clarify their sensitivity regarding current density and SOC distribution. Additionally, subjecting the cell to defined thermal gradients to influence the SOC inhomogeneity should provide further insights.

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References