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Enhanced Fast Charging and Reduced Lithium-Plating by Laser-Structured Anodes for Lithium-Ion Batteries

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Lithium-ion batteries (LIBs) are the key-components for the success of electric vehicles and a sustainable energy economy. Their performance is highly dependent on the charge and mass transport kinetics between the porous electrodes. At low temperatures, the cell internal resistances rise due to a decreasing ionic conductivity of the liquid electrolyte, sluggish charge transfer processes on the interfacial area between the active material and the electrolyte, and slow lithium diffusion within the active material. The increasing resistances result in high overpotentials and electrode polarization during cell operation. For discharge, the consequences are a reduced deployable capacity and an amplified heat generation. However, high overpotentials during charging can cause the deposition of metallic lithium (Li-plating) onto the anode surface, specifically when the anode surface potential falls below 0 V. Li-plating has been reported to accelerate the ageing of the LIB and presents severe safety risks due to dendrite formation.

Among others, a low anode-to-cathode capacity ratio, low temperatures, high charging rates, and overcharging favor the deposition of metallic lithium, as these factors have an effect on the anode kinetics and the diffusion rate. Thus, the charging currents have to be strictly limited, especially at low temperatures. While in laboratory cells the individual electrode potentials can be measured by applying reference electrodes, the detection of Li-plating in commercial full cells is challenging. The Li-plating process is partially reversible and some of the deposited lithium re-intercalates into the graphite when a relaxation period occurs after charging. During this period, a distinctive voltage plateau can be observed, making the onset of Li-plating detectable. When measuring at very low frequencies, the ionic resistance within LIB electrodes can be assessed in situ by measuring the cell impedance using electrochemical impedance spectroscopy. The spectra represent the total impedance of the electrochemical cell. The impedance spectra consist of two partially overlapping semicircles and a straight sloping line in the complex plane, which provides a relation between the ionic conductivity $\kappa$ and the effective conductivity $\kappa_{eff}$ through the porous media. From a geometrical perspective, the tortuosity $\tau$ is a characteristic value for describing the ion transport restraint of the porous electrode microstructure, where $d_{path}$ is the actual transport path of an ion through the electrode and $d$ is the direct path between the points of de-intercalation and intercalation while charging or discharging. The tortuosity $\tau$ has been reported to have a major influence on diffusion and concentration overpotentials within the electrode structure. As the lithium transport into and out of the electrode is impeded, large concentration gradients are formed, creating a strongly inhomogeneous charge carrier distribution. This effect predominantly occurs in anodes containing flake-like graphite particles which are commonly used as anode material in LIBs due to their lower cost compared to spherical graphite, which is roughly three times more expensive due to high scrap rates in spherical graphite production. Due to the challenging process in LIB production, the preferred orientation of the graphite flakes is parallel to the current collector and perpendicular to the diffusion direction through the electrode. This generates a spatial tortuosity anisotropy and creates particularly long diffusion pathways $d_{path}$ through the electrode. Among many other parameters, the ionic resistance within LIB electrodes can be assessed in situ by measuring the cell impedance using electrochemical impedance spectroscopy (EIS), enabling a calculation of the tortuosity. By measuring the impedance of cells at different states of charge (SOC) and temperatures, conclusions concerning the kinetics of lithium ions within the electrodes can be drawn. Impedance spectra for LIB are represented in the form of Nyquist plots and typically consist of two partially overlapping semicircles and a straight sloping line in the low frequency region. As the effective impedances of cathode and anode in full cells cannot be precisely distinguished, the recorded EIS spectra represent the total impedance of the electrochemical cell. The size of the semicircles provides information about the magnitude of the resistances resulting from the charge transfer processes between the electrodes and the electrolyte as well as the passing through the solid electrolyte interface (SEI). When measuring at very low frequencies, the resulting quasi-linear slope can be related to the solid-state diffusion processes within the active material particles. At low
temperatures, low and SOCs, the charge transfer resistances within the cell rise, resulting in an increase in the overall impedance, which can be seen in the EIS spectra.

One promising way for enhancing the lithium transport in LIBs is the creation of three-dimensional (3D) electrode structures, aiming at a reduction of the average electrode tortuosity. Thereby the effective diffusion of lithium in the electrolyte $D_{\text{eff}}$ can be increased according to

$$D_{\text{eff}} = \frac{\varepsilon}{\tau} D_0$$

where $\varepsilon$ is the electrode porosity and $D_0$ is the diffusion coefficient of lithium in the liquid electrolyte. The 3D-structures can be created by laser pulses which precisely remove a small fraction of the coating. It has been reported that the shortened ion transport pathways lead to a reduced cell resistance and thereby improve the discharge rate capability compared to conventional unstructured electrodes. This effect was computationally analyzed in a numerical model, precisely depicting the structure arrangement and dimensions. The introduction of additional pores by laser radiation offers a high flexibility regarding the structure type (e.g. grid, line, or hole structures) and the respective spacing between the structures. Also, the dimensions of each individual structure can be adjusted. Generally speaking, more particles are removed close to the surface of the electrode, resulting in conical structures which narrow toward deeper parts of the electrode. The size and the distribution of the structures have a strong effect on the cell internal resistance and, thus, the electrochemical performance. When evaluating structure geometries that lead to a comparable total amount of removed electrode material, finer structures have proven to be more beneficial than coarser structures. In contrast to other concepts such as co-extruded or sintered as well as 3D-printed electrodes, the laser structuring process can be applied to conventionally coated electrodes. From a production point of view, establishing the laser structuring process in the industrial fabrication of LIBs shows higher potential for in-line implementation (roll-to-roll processing) into the existing production chain and is associated with lower cost. Cost calculations with regard to laser structuring of electrodes based on scenarios by Wood et al. even predict potential savings in LIB production which may surpass the additional costs due to laser machinery, maintenance, and operation. These potential savings result from a reduction in the electrolyte wetting time and the application of electrodes with a higher areal loading and energy density.

**Objectives and Approach**

As described in the introductory section, Li-plating is one of the major obstacles in the development of LIBs with fast-charging capability. This paper presents a new approach to electrode design in order to reduce the cell impedance and, thus, the overpotentials during charging and discharging. The generation of structures in the anode by means of short laser pulses (pulse duration 150 ps) aims at improving the ion transport between anode and cathode (Fig. 1).

As structures, holes in a hexagonal pattern with a center point distance of 100 μm were created. The laser process was designed to achieve an aspect ratio of the structures of > 1.5, which is defined by the ratio of the structure depth over the structure diameter at the electrode surface. Based on previously published data, the hexagonal hole pattern represents the best geometric structure to balance the conflict between the removal of active material and the reduction of the anode tortuosity. By analyzing the electrochemical impedance spectra at temperatures of −15°C, 0°C, and 25°C, the impedances of coin cells containing laser-structured anodes (S1 and S2) were compared to their conventional counterparts (R1 and R2). Subsequently, the effects of the structuring on the discharge C-rate capability were examined. In the next step, the charging behavior was analyzed with a special focus on the detection of Li-plating. For this purpose, the voltage relaxation after charging at C-rates from 0.1C to 2C and ambient temperatures of −15°C, 0°C, and 25°C was monitored and indications of Li-plating were observed. Finally, the charging durations required to fully charge the LIBs were assessed in order to evaluate the potential for improvements in fast charging under demanding low-temperature conditions. In Fig. 1a and Fig. 1b, the internal setup of a LIB with laser-structured graphite anodes and conventional unstructured NMC cathodes is illustrated. Fig. 1c shows a top-view scanning electron microscopy (SEM) image of a laser-structured graphite anode as well as a close-up image of a single laser-induced hole structure. More detailed information on the materials and processes used can be found in the subsequent Materials and Methods section.

**Materials and Methods**

**Materials.**—All coin cell components including the coated electrode materials were purchased from commercial sources to ensure a high comparability of the base materials (type 2032 coin cell cases, springs, and spacers: MTI Corporation, USA; glass fiber separator: Type 691, VWR, USA; cathodes: Umicore NV/SA, Belgium; anodes: SEI Corporation, Japan). The electrode characteristics, as provided by the electrode manufacturers, are presented in Table I. Further details on the precise material types (e.g. type of binder, type of conductive carbon) and the respective production processes were not disclosed to the authors for reasons of confidentiality.

**Laser structuring of the anodes.**—A pulsed Nd:YAG fiber laser (YLP-1-150V-30, IPG Photonics Corporation, USA) with discrete tunable pulse durations from 150 ps to 5 ns was used to generate the...
Table I. Electrode specifications.

| Feature                | Cathode | Anode     |
|------------------------|---------|-----------|
| Active material        | NMC (1/1/1) | Graphite |
| Conductive carbon      | 3.0 wt% | 0.5 wt%  |
| Binder                 | 4.0 wt% | 7.0 wt%  |
| Capacity loading       | 2.748 mAh/cm² | 3.606 mAh/cm² |
| Weight loading         | 20.38 mg/cm² | 12.99 mg/cm² |
| Loading deviation      | ± 2.5% | ± 3.0%   |
| Porosity               | approx. 32% | approx. 32% |
| Electrode thickness    | 69 μm  | 70 μm    |
| Thickness deviation    | ± 2 μm | ± 2 μm   |

In Table I, we display characteristics of the electrodes used in the study. All other cell components were dried overnight at 60°C in an oven, to ensure no significant changes in the mechanical properties of the electrodes were observed. This included the binder, which is responsible for the cohesion of the electrode, was not detectably vaporized or degraded beyond the actual structure diameter. All subsequent handling processes in the assembly of the coin cells could be carried out unaltered.

Cell assembly and formation.—The coin cells (type 2032) were assembled in a dry room (dew point < -40°C). Before cell assembly, the electrodes were dried for 12 hours at 120°C in a vacuum oven. All other cell components were dried overnight at 60°C in an oven within the dry room atmosphere. Two cell types were manufactured: reference cells with conventional anodes and conventional cathodes, and cells with laser-structured anodes and conventional cathodes. The circular cathode electrodes were 14 mm in diameter while the anode coins were 15 mm in diameter to ensure a complete coverage of the cathodes by the anodes. The diameter of the glass microfiber separator was 16 mm to avoid internal short circuits. The conventional electrodes had an area specific capacity ratio between anode and cathode of approx. 1.31, while the cells comprising structured electrodes showed a capacity ratio of approx. 1.24 due to anode material removal. The electrolyte (LP572, BASF, Germany) consisted of ethylene carbonate (EC) and ethyl-methyl-carbonate (EMC) in a ratio of 3:7 with 1 M lithium-hexafluoroarsenate (LiPF6) and 2% vinylene carbonate (VC). After assembly, the cells went through formation by charging and discharging them five times at approx. 0.1 C in a laboratory battery testing system (CTS, BaSyTec, Germany). The measured capacity in the last formation cycle was used for referencing purposes, i.e. to define an individual nominal capacity for each cell. During formation, no negative influence on the initial capacity by laser structuring was observed as all the nominal cell capacities showed a negligible standard deviation of 0.048 mAh around a mean capacity of 2.93 mAh, see Table II.

Detection of Li-plating.—The tests for the detection of Li-plating were based on the identification of voltage plateaus in a relaxation phase after CCCV charging. In the CC phase, different currents were applied and also the termination criteria for the CC phase were adjusted. In case of the 0.1 C and 0.5 C charging current in the CC phase, a 0.05 C cutoff current in the CV phase was chosen, while in case of the 1 C and 2 C charging currents in the CC phase, a 0.2 C cutoff current in the CV phase was applied. The latter termination criterion for the CV phase was used in order to achieve comparable durations of the CV phases. Equal termination criteria would have resulted in very long CV phases for the faster charging processes (1 C and 2 C), which would have strongly altered the significance of the results with regard to Li-plating as lithium re-intercalation also would have happened during the CC phase.

Table II. Coulombic efficiencies (CE) of the first and second formation cycle and nominal cell capacities after the fifth formation cycle.

| Cell name | CE cycle 1 | CE cycle 2 | Capacity after formation |
|-----------|------------|------------|--------------------------|
| R1        | 87.54%     | 99.17%     | 2.96 mAh                |
| R2        | 87.59%     | 98.65%     | 2.85 mAh                |
| S1        | 86.74%     | 98.30%     | 2.92 mAh                |
| S2        | 86.86%     | 98.44%     | 2.97 mAh                |

In particular, no indications of a significantly increased SEI formation were detected as the coulombic efficiencies (CE) in the first and second formation cycles do not deviate strongly (Table II). This may be explained by the fact that, although the macroscopic and therefore exposed surface area of the electrode was enhanced, graphite particles were ultimately removed by the laser ablation process. These removed graphite particles were therefore not subject to SEI formation. A variation of the material system or the structure geometry could change the ratio of the counteracting influences regarding SEI formation described above. Further studies on SEI formation will be performed to substantiate this interpretation of the measurement data.

Electrochemical impedance spectroscopy.—For the EIS measurements, a potentiostat (Interface 5000E, Gamry Instruments, USA) was used. Before every measurement, a rest period of > 1 h was applied to avoid a falsification of the results by cell polarization. Every measurement started with the recording of the OCV, which was performed for 15 s. Subsequently, the impedance was recorded in the frequency range from 100 kHz to 1 Hz with 10 measurement points per decade.

Discharge rate capability tests.—To determine the discharge rate capability, the cells were cycled between 2.7 V and 4.2 V at different temperatures and increasing C-rates (Fig. 3). At a temperature of 25°C, the maximum C-rate was set to 10 C, while at 0°C and −15°C the maximum C-rate was set to 5 C in order to avoid excessive stress and cell ageing. Charging was done using a constant-current/constant-voltage (CCCV) procedure. For the corresponding discharge rates up to 1 C, an equal charging current was used in the CC phase. For discharge cycles > 1 C, the charging current was held constant at 1 C to avoid premature ageing and Li-plating. The CV phase was terminated when the charging current fell below 0.01 C. No rest period between charging and discharging was applied. Discharging was performed using a constant current (CC) procedure.

Influence of Laser Structuring on Cell Impedance

In Fig. 2, the Nyquist plots obtained from EIS measurements of LIBs with laser-structured anodes (S1, S2) and conventional anodes (R1, R2) at an OCV of 3.7 V are presented. Three ambient temperatures of 25°C, 0°C and −15°C were analyzed. Please note that the presented measurements do not aim at quantitative statements or...
models for the electrochemical processes within the cell but at a qualitative comparison of the impedances of the two different cell types at varying temperatures. As the properties of the cathodes as well as other cell parameters were kept constant in all cells, the variances in the EIS spectra can only be attributed to the additional laser-induced macroscopic pores in the anodes.

As described in the introductory section of this paper, at very low frequencies, the impedance spectra of intercalation battery electrodes represent the transport of charge carriers in the solid phases. As the graphite microstructure and, thus, the solid-state diffusion characteristics were not altered, the low frequency domain (< 1 Hz) was not considered in the presented experiments. Please also note the different scales of the individual Nyquist plots. In accordance with the literature, an increased impedance with decreasing temperature was observed. This can be attributed to slower lithium diffusion kinetics and higher charge transfer resistances within the cells. When analyzing the graphs, it becomes apparent that the impedance of cells with laser-structured anodes was considerably lower than that of cells with conventional electrodes. While no significant influence on the high frequency resistance (HFR) was noticed, the impedances in the semi-circle range of the Nyquist plots were significantly lower. This range is typically smaller than the second for both laser-structured and conventional cells. A reduction of the temperature to 0°C (Fig. 2b) led to an overall increase of the impedance, while the general shape of the graph was maintained. The impedance increased even further when the temperature was lowered to −15°C (Fig. 2a). While at 0°C both semicircles are still visible, at −15°C, the second semicircle is truncated due to the minimum applied measuring frequency of 1 Hz (the slight curvature of the plots in the direction of lower frequencies indicates that the second half circle only starts here). For all plots it becomes apparent that the impedance of cells with laser-structured anodes was considerably lower than that of cells with conventional electrodes. While no significant influence on the high frequency resistance (HFR) was noticed, the impedances in the semi-circle region of the Nyquist plots were significantly lower. This range is usually attributed to charge transfer processes, which comprise the lithium transport from the liquid into the solid phase and vice versa. Smaller semi-circles therefore mean that this charge transfer is associated with lower resistance, which may have various possible origins. On the one hand, the structuring results in a higher mean porosity, which in turn provides a better accessibility of the graphite particles for the charge transfer. Since the additional pores open toward the electrode surface due to their characteristic shape, the average porosity is further decreased in proximity of the electrode surface. This gradient in porosity additionally enhances the abovementioned effect. These findings indicate that laser structuring may have a positive influence on the charge transport behavior and, thus, on the fast charging and discharging performance of the cell. As already indicated above, a detailed modeling of the complex resistances will be carried out in future work in order to investigate the origin of the impedance reduction more closely. However, since this paper mainly aims to outline the potentials of the laser structuring of the anode with regard to fast charging, we only provide comparative observations here.

Enhanced Discharge Rate Capability

In the previous section, significantly reduced impedances for the laser-structured electrodes were determined by EIS. The amount of charge provided by lithium-ion cells during discharge depends on the discharge current. This interrelation has its origin in the cell-internal resistances. A reduction in these resistances therefore leads to an increase in the amount of charge provided. To quantify the influence of structuring, discharge rate tests were performed on cells with conventional anodes and the results were compared to cells with laser-structured anodes at different temperatures (25°C, 0°C, −15°C) (Fig. 3). To ease the comparison, the provided discharge capacities were normalized to the value measured at a low discharge rate of 0.1C in the first discharge cycle after cell formation. Since both the conventional and the laser-structured cells showed only a very small deviation in capacity (cp. Table II), we do not consider the validity of the results as falsified by the normalization. Low temperatures generally limit the rate capability as charge and mass transport processes decelerate. The measurements presented here are in agreement with this relationship. At all temperatures, enhancements in rate capability were observed due to laser structuring. The level of improvement through laser structuring varied in strong dependence of the temperature. At 25°C, a maximum increase of approx. 9% in the delivered charge could be achieved by anode structuring. At 0°C the maximum increase was approx. 21% compared to unstructured cells, while at −15°C the maximum improvement was approx. 27%. Depending on the temperature, these maxima were measured at different C rates. While the best results at −15°C were already observed at 2C, the maxima at 0°C and at 25°C were shifted toward 4C and 8C respectively. This trend is shown in Fig. 3d. Moreover, an explicit maximum of improvement in C-rate capability was observed at all temperatures and the achievable improvements decreased again at very high C-rates. This indicates that the positive impact of anode structuring is visible at temperature and material specific discharge rates. At even higher C-rates, other limitations predominate and the associated homogenization of the charge carrier distribution in the electrolyte and in the electrode decreases.

It should be noted that the removal of a small fraction of the graphite coating (reduction of the ratio between anode and cathode capacity from 1.31 to 1.24, see materials and methods section) results in a lower mass loading of the anode, which should generally lead to a slightly improved C-rate capability, even without the structures. Investigations to distinguish between the influence of mass loading reduction and the benefit of the structures by precise adjustment of the electrode parameters are currently underway and will be published shortly. However, previously published work on the electrochemical simulation of pore morphology modifications by laser structuring indicates that the
electrode structure itself has a decisive influence on the cell performance. Also note, that anode overbalancing is typically used to prevent the occurrence of Li-plating. However, in the following section, we show that despite lower anode overbalancing, the structuring results in reduced Li-plating during fast charging. Laser structuring may therefore be a solution for eliminating the need for excessive overbalancing in future cell design.

Reduction of Lithium-Plating

As described in the introduction to this paper, there are several ways to electrochemically detect the occurrence of Li-plating in full cells. The measurement method described below to detect Li-plating was adapted from Uhlmann et al. and is illustrated in Fig. 4. The voltage relaxation of the full cell during a rest period after CCCV charging is considered. Depending on whether or not metallic lithium is plated at the anode during the charging process, there are clearly distinguishable relaxation voltage curves.

If no Li-plating occurs, the voltage converges quasi-exponentially against a threshold value. This relaxation has its origin in the homogenization of the cell polarization in the electrodes and in the electrolyte. However, if Li-plating occurs during charging, these homogenization processes are superimposed by lithium re-intercalation in the relaxation phase (de-plating). The metallic lithium adhering to the anode surface decomposes and intercalates into the anode within a specific timespan that correlates with the amount of plated lithium. Since this process takes place at a different potential level, a characteristic transient voltage plateau emerges, compare Fig. 4. The voltage curves presented in Fig. 5 show the relaxation after previous charging (CCCV) at a temperature of 0°C for the two structured and the two conventional cells. These environmental conditions already pose a challenge in the field of electric mobility. At a temperature of 25°C, either no Li-plating could be provoked or detected during these tests (charging up to 2C). Within preliminary tests, it could be shown that with a further increase of the charging current the upper cutoff voltage of 4.2 V was reached very quickly. Since the cells still had a very low SOC at that time, only a small amount of lithium had been transferred to the anode at all. However, since Li-plating occurs particularly at high SOCs (due to the approximation of the anode potential toward 0 V vs. lithium at high degrees of lithiation), no Li-plating could be detected. Therefore, the fast charge measurements at 25°C will not be discussed further.

Fig. 5a shows the voltage curve after a 0.1C charge, Fig. 5b after a 0.5C charge, Fig. 5c after a 1C charge, and Fig. 5d after a 2C charge. For the two slower charging processes, the termination criterion of the CV phase was when the charge current fell below 0.05C, while for the two faster charging processes, the termination criterion of the CV phase was set to 0.2C. As the re-intercalation of lithium also occurs during the CV phase prior to the rest period, the latter termination criterion was chosen so that all CV phases lasted for a similar amount of time. Thus, even while fast charging, a significant re-intercalation of lithium during the CV phase could be avoided, which ensures the comparability of the results. After a 0.1C charge (Fig. 5a), both the structured and the
Figure 4. Schematic illustration of the Li-plating process with the corresponding voltage profile; in the rest period subsequent to charging, a voltage plateau is visible if Li-plating occurred during the previous charging process; this plateau forms due to the re-intercalation of the plated lithium into graphite, which takes place on a different potential level.

Figure 5. Voltage relaxation of conventional and laser-structured cells after CCCV charging at 0°C; in the CC phase different currents were applied: a) 0.1C current in CC phase, 0.05C cutoff current in CV phase; b) 0.5C current in CC phase, 0.05C cutoff current in CV phase; c) 1C current in CC phase, 0.2C cutoff current in CV phase; d) 2C current in CC phase, 0.2C cutoff current in CV phase; the onset of Li-plating was observed at significantly higher charging C-rates for laser-structured cells than for conventional cells taken from the voltage plateau caused by lithium re-intercalation; at very high charging C-rates, Li-plating is strongly reduced; voltage relaxation is generally lower for laser-structured cells, indicating a better fast charging capability.
unstructured cells showed a voltage curve indicating that no Li-plating occurred during charging. However, a small difference of the final voltage after relaxation could be measured. While the OCV of the structured cells relaxed toward a voltage of approx. 4.18 V, this value was 4.17 V for the unstructured cells. This indicates that already at low charge currents of only 0.1C a stronger polarization occurred in the conventional cells and that this polarization could be reduced by laser structuring. The course of the voltage during relaxation after a charge with a current of 0.5C (Fig. 5b) was very similar. No indications of Li-plating were observed. For charging with a current of 1C (Fig. 5c), significant differences in the voltage curves were visible. While the cells with laser-structured anodes maintained a quasi-exponential voltage relaxation, the conventional cells showed a distinct voltage plateau. This plateau lasted for about 30 minutes before the voltage relaxed against a final OCV in the range between 4.08 V and 4.10 V indicating the occurrence of Li-plating during charging. Furthermore, the duration of the persistence of the plateau gives an indication about the severity of Li-plating. The longer the period of re-intercalation, the more lithium was metallically deposited during the charging process. A charging current of 1C at 0°C which already led to strong Li-plating for the conventional cells was still uncritical for laser-structured cells. This demonstrates the high potential of laser structuring, as safety risks and ageing originating from Li-plating are reduced, which suggests that increased charging currents may be used in fast-charging applications. Li-plating was observed for both cell types at a charging current of 2C (Fig. 5d), whereby the amount of deposited lithium varied significantly. The re-intercalation plateau was present for more than one hour for the conventional cells, whereas it was considerably shortened (approx. 30 minutes) for the structured cells.

The effects described above become even more pronounced when the time derivative of the voltage curve is depicted. The corresponding differential voltage curves are shown in Fig. 6. The end of the voltage plateau, which is expressed as a change in the slope of the voltage curve (cp. Fig 5), then appears as a local minimum and thus allows quantification of the completion of the lithium re-intercalation. In accordance to the slopes in Fig. 5a and Fig. 5b no pronounced minima are visible after charging with 0.1C and 0.5C. On the contrary, Fig. 6c clearly shows that a small amount of metallic lithium was plated, leading to a local minimum in the case of the structured cells after a 1C charge. However, the degree of the Li-plating of the unstructured cells is distinctly higher. Similar characteristics can be observed after a 2C charge.

At a temperature of −15°C, all charge and mass transport processes are significantly slower. Preliminary tests showed that charging with...
Figure 7. Voltage relaxation (a, b) and corresponding differential voltage relaxation (c, d) of conventional and laser-structured cells after charging with different currents at −15°C; the beneficial effects of laser structuring seen at 0°C are far less pronounced at −15°C indicating that other shares of the cell resistance become dominant.

1C and 2C lead to a fast increase in cell voltage and to a premature reach of the upper cutoff voltage. Thereby, no SOC higher than 50% could be reached in the CC phase. In this SOC range, no evidence of Li-plating was detectable. The authors attribute this to the low SOC of the cells at which Li-plating is unlikely to occur. Therefore, only the relaxation processes after 0.1C and 0.5C charging are described below. The results are presented together with the time derivative of the voltage curves in Fig. 7. It becomes evident that at a charging current of 0.1C, Li-plating did not occur in either the conventional cells or the cells with structured anodes. This conclusion can be drawn from the voltage relaxation (Fig. 7a) as well as the time derivative (Fig. 7c), where no local minimum is apparent. When charged with 0.5C, both cell types showed indications of Li-plating, expressed by the distinctive voltage plateaus (Fig. 7b) and the respective local minima in the differential voltage curves (Fig. 7d). Again, the trend toward reduced Li-plating of the structured cells can be observed, but less pronounced than at an ambient temperature of 0°C. This may indicate that other cell internal processes, e.g., lithium diffusion within the graphite particles or at the phase boundaries between the electrolyte and the active materials, have a limiting effect. Thus, the ion transport in the electrolyte within the tortuous electrode structure no longer appears to be the limiting process. It should be noted that the re-intercalation process generally takes much longer at −15°C.

After charging with 0.5C the re-intercalation takes approx. 1.5 hours for the structured cells and approx. 2 hours for the conventional cells. More detailed measurements with a higher resolution in the examined charging rates are required to determine the C-rate at which Li-plating first occurs at −15°C for both cell types.

Improvement of Fast Charging

As described in the previous sections, laser structuring reduces the mean tortuosity of the anode, resulting in lower overpotentials and lithium concentration gradients during charging and discharging. The lower overpotentials imply that the upper cutoff voltage during CC charging is reached at a higher SOC for cells with laser-structured anodes for a given charging current. As a result, the duration of the subsequent CV phase is reduced since a smaller amount of charge is necessary to reach an SOC of 100%. The respective charging times in the CC and CV phase were measured for the four cells S1, S2, R1, and R2. The results are shown in Fig. 8 for an ambient temperature of 0°C.

The laser structuring of the anodes and the associated change in the CC/CV ratio led to a significantly reduced time to achieve an SOC of 100%. While the conventional cells needed about 2.4 hours to be fully charged at a charge current of 0.5C, the structured cells required only 2.0 hours (Fig. 8a). To further investigate this, the SOC at which the
CV phase begins was examined (Fig. 8b). The results show that the CV phase does not start before an SOC of approx. 90% compared to approx. 88% for the conventional cells. Since the charge current drops rapidly at the beginning of the CV phase, substantially more time is required to reach the final SOC of 100%. This comparison proves that the laser structuring of anodes can have a significant positive influence on the charging time of lithium-ion cells, even in operating conditions that do not cause Li-plating.

Conclusions and Future Work

In this publication, the influence of laser structuring of the graphite anodes on the operating characteristics of LIBs was assessed. By using electrochemical impedance spectroscopy it was shown that the cell impedance and specifically the charge transfer resistance was significantly reduced by laser structuring. This was attributed to a reduced mean tortuosity that facilitated lithium transport as well as an increased mean porosity which provided a higher surface area for charge transfer. The laser structuring had a positive effect on both the discharge and charge process of the cell. For discharging, it was demonstrated that the amount of capacity and therefore energy provided was distinctly increased. This effect was particularly pronounced at low temperatures. The maximum improvements in deployable capacity were observed at varying discharge C-rates depending on the ambient temperature. At this point we would like to point out once again that some of these improvements were due to the slight reduction in the active material amount of capacity and therefore energy provided was distinctly increased. This effect was particularly pronounced at low temperatures. The maximum improvements in deployable capacity were observed at varying discharge C-rates depending on the ambient temperature. For charging, it was shown that the maximum improvements in deployable capacity were observed at various charging C-rates depending on the ambient temperature.

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