Effect of carboxymethyl cellulose on the flow behavior of lithium-ion battery anode slurries and the electrical as well as mechanical properties of corresponding dry layers

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ABSTRACT

We present a holistic view on the role of polymeric binders in waterborne LiB anodes, including preparation and processing of wet slurries as well as microstructure, electrical conductivity and mechanical integrity of dry electrode layers. We focus on carboxymethyl cellulose (CMC), with respect to technical application the influence of soft, nano-particulate styrene–butadiene rubber (SBR) as secondary binder is also addressed. We discuss the influence of CMC concentration, molecular weight (Mw) and degree of substitution (DS) on flow behavior of anode slurries. Rheological data are not only relevant for processing, here we use them to characterize the adsorption of CMC on active material particles and dispersion of these particles in the slurry at technically relevant concentrations. The fraction of CMC adsorbed onto graphite particles increases with increasing Mw and decreasing DS. Electrical conductivity increases with Mw, i.e. with decreasing free polymer deteriorating conductive carbon black pathways. CMC does not contribute to the adhesion of electrode layers, irrespective of Mw or DS, technically feasible adhesion is inferred by SBR. Cohesive strength of anode layers, determined here for the first time under well-defined mechanical load, increases with increasing Mw and decreasing DS, i.e. with increasing fraction of adsorbed CMC and corresponding improved particle dispersion. Strong cohesion and high electrical conductivity are correlated to an alignment of graphite particles as revealed by electron microscopy, presumably enabled by higher particle mobility in well-dispersed slurries. Accordingly, targeted choice of CMC is a valuable means to control processing, electrical conductivity and mechanical strength of LiB electrodes.
Introduction

Lithium-ion batteries (LiB) play an important role as electrochemical energy storage systems. They combine high energy and power density, making them suitable for portable electronics, hybrid/full electric vehicles and grid applications [1–5]. So far, research mainly focused on the development of new electrochemically active materials to achieve high cell performance [6–10]. In contrast, little attention has been paid to the role of binders not only for processing but also for the performance of LiB. As electrochemically inactive material, poly (vinylidene fluoride) is one of the most widely used polymeric binders for LiB electrodes. Conventionally, these binders are dissolved in organic solvents, such as N-methyl-2-pyrrolidone (NMP) [11, 12]. Since organic solvents are typically environmentally unfriendly and flammable, safety concerns and high cost are associated to their use in LiB production, particularly for the upcoming high-volume mass production of electric vehicles. Therefore, water-based electrodes have been developed as promising environmentally benign alternative [13, 14].

Carboxymethyl cellulose (CMC) and styrene butadiene rubber (SBR) are often combined as additives for aqueous anode slurries to achieve high stability and processability as well as superior electrochemical cell performance [15–20]. The present investigation focuses on CMC as binder. Additionally, nano-scale SBR particles are added during slurry preparation since this soft, rubber-like polymer is necessary to provide a technically feasible adhesion to the current collector. CMC is a linear, long-chain, water-soluble, anionic polysaccharide, consisting of two anhydroglucose units (β-linked glucopyranose residues) with three hydroxyl groups each. The hydrogen in the hydroxyl groups can be substituted by a carboxyl group (–OCH_2COO–). The average number of substituted hydroxyl groups per anhydroglucose unit is defined as the degree of substitution (DS) and hence has a maximum of 3. Rheological properties of CMC solutions strongly depend on concentration, molecular weight (M_w), DS and pH [21–26]. This also shows up in the flow behavior of battery electrode slurries. Beyond that, however, the interaction with other components in these slurries has to be considered. Lee et al. investigated the effect of CMC DS on the dispersion of graphite particles and the resulting electrochemical performance of corresponding anodes [16]. CMC with lower DS adsorbed preferentially on graphite, dispersing the particles and stabilizing the slurry against sedimentation through repulsive electrosteric interparticle forces. As a result, longer cycle life and improved electrochemical cell performance was achieved using low substituted CMC. Other studies have shown the effect of polymeric binders on particle dispersion, unveiling their influence on the electrical conductivity of electrode layers [27–29]. Proper binder to particle ratios and dispersion properties of the binder lead to the formation of a conducting network, enhancing the electron transport kinetics and yielding higher energy, power and life time performance.

Battery lifetime strongly depends on the mechanical integrity of the dry electrode. During lithium intercalation, the electrode must withstand mechanical stresses due to expansion and shrinkage of the electrochemically active material [30–32]. The volume fluctuations can induce a local delamination of the electrode layer from the current collector and cracking within the layer. Hence, considerable research work dealt with the determination of adhesive strength between electrode layer and substrate [17, 33–46]. However, the cohesive strength in the electrode layer has not received much attention yet. The scratch test has been used to characterize the mechanical strength or scratch resistance of electrodes [33, 47, 48]. Since in this test the electrode layer experiences tangential and normal forces, the measured load comprises information about the adhesive as well as the cohesive strength of the electrode. Chen et al. combined micro-scratch and digital image correlation techniques to decouple particle/particle cohesion from electrode-layer/current-collector adhesion and defined the coefficient of friction as a measure of cohesive strength [49]. Nevertheless, it remains elusive whether such local force measurements are representative for the cohesive strength of the electrode layer. Finally, indentation and tensile tests have been carried out to determine the mechanical properties of electrode layers [37, 50–52]. These studies provided some understanding about how wet processing of the slurry affects porosity and mechanical properties of the dry layer. Still, the true contribution of the binder to the cohesive strength of electrode layers is not yet fully resolved.

In this study we present a holistic view on the role of polymeric binders for LiB anodes, including...
preparation and processing of wet slurries as well as the microstructure, electrical conductivity and mechanical integrity of dry electrode layers. We first investigate the influence of CMC concentration, molecular weight and degree of substitution on flow behavior of aqueous anode slurries. Rheological data are not only relevant to judge processing behavior, here we use them to characterize the degree of adsorption of CMC on active material particles and the degree of dispersion of these particles in the slurry at technically relevant concentrations. The slurries are then cast on to glass plates and copper foils and dried to determine the electrical conductivity and the adhesive strength of the electrode layer, respectively. Furthermore, first cohesive strength data for thick anode layers comprising graphite, carbon black and CMC are presented. The adsorption behavior of CMC on the particle surface is discussed and correlated to the obtained results. Finally, we present data characterizing the cohesive strength of anode layers under well-defined mechanical load conditions for the first time and correlate these data to the electrode microstructure and its change with varying type and concentration of CMC, emphasizing its relevance for the mechanical integrity of the electrode layer beyond the mere presence of the polymeric binder. With respect to the typical composition of commercial LiB electrodes, the influence of two different SBR-binders on the mechanical, structural and electrical properties mentioned above is also discussed.

**Experimental**

**Materials**

Commercially available flake-like, synthetic graphite particles (SLP50, Imerys Graphite & Carbon, Switzerland) with a volume-based average diameter $x_{50,3} = 23 \mu m$, a specific surface area of $5.7 m^2 g^{-1}$ and density of $2.25 g cm^{-3}$ are used as active material for the preparation of aqueous anode slurries. Carbon black (CB, C-Nergy Super C65, Imerys Graphite & Carbon, Switzerland) with a primary particle size $x_{50,3} = 32 nm$, a specific surface area of $64.1 m^2 g^{-1}$ and density of $1.8 g cm^{-3}$ was added as a conductivity agent. CB particles tend to agglomerate in the slurry reaching an average agglomerate size of $5 \mu m$. Four types of sodium carboxymethyl cellulose (CMC, Sigma Aldrich, Germany) with average molecular weight ($M_w$) of 250 kDa and 700 kDa were used in this investigation. The latter had a degree of substitution (DS) of 0.9, whereas three different DS of 0.7, 0.9 and 1.2 were investigated for CMC with $M_w = 250 kDa$. Additionally, CMC with $M_w = 400 kDa$ and DS = 0.9 was purchased from Dow Wolff Cellulosics GmbH. Commercially available styrene butadiene rubber (SBR, TRD2001, JSR Micro NV., Belgium) as well as a modified version (TRD105A) were used as secondary binder. These polymers are supplied as aqueous dispersions of nanoparticles (particle volume fraction 48 vol% and 41 vol% for TRD2001 and TRD105A, respectively). According to manufacturer, TRD2001 exhibits a particle size of 170 nm and a glass transition temperature of $-2^\circ C$, whereas TRD105A has a particle size of 95 nm and a glass transition temperature of $-5^\circ C$. Small amplitude oscillatory shear experiments at fixed frequency and deformation covering a wide temperature range on dry SBR films did not show a cross-over of the storage ($G'$) and loss ($G''$) moduli at high temperatures, indicating high molecular weight of the polymers or even chemical cross-linking (see supporting information). Gravimetric measurements using toluene as solvent yielded a degree of cross-linking of 89.9% ± 0.5% and 81.5% ± 0.4% for TRD2001 and TRD105A, respectively.

**Sample preparation**

Anode slurries with a constant solid volume fraction $\phi_S = 20$ vol% and a constant graphite to CB volume ratio of 46.5 were studied. The CMC concentration was varied in a range $\phi_{CMC} = 2.5$–4.0 vol% for different molecular weight and degree of substitution. Here, $\phi_{CMC}$ refers to the amount of polymer in the dry electrode layer. First, CMC was dissolved in distilled water and homogenized with a 55 mm propeller mixer at 1200 rpm for 30 min. Carbon black, followed by graphite particles, were dispersed in the aqueous CMC solution using a 50 mm dissolver disk at 1600 rpm for 5 min each. Only part of the total amount of water needed for the slurry was initially used to prepare the CMC solution in order to ensure particle deagglomeration and ideal slurry homogeneity. After addition of the solid particles, the remaining amount of water was added stepwise in four intervals of 5 min each until reaching the desired solids content [53]. For anode slurries including SBR,
the secondary polymer was added after dispersion of the solid particles and finally the desired solids content was adjusted by adding the appropriate amount of water. Slurries including both polymer binders were studied at fixed CMC concentration $\phi_{\text{CMC}} = 2.5 \text{ vol\%}$ and constant $M_w = 700 \text{ kDa}$ and DS = 0.9 but varying SBR concentration in a range $\phi_{\text{SBR}} = 1.0-5.0 \text{ vol\%}$.

Slurries were coated on glass plates and 10 $\mu$m thick copper foil (SE-Cu, Schlenk Metallfolien GmbH & Co. KG, Germany) for electrical conductivity and adhesive strength measurements, respectively. Therefore, a doctor blade (ZUA 2000, Zehntner GmbH, Switzerland) with a coating width of 60 mm and a coating gap of 300 $\mu$m was applied. Subsequently, the wet film was dried at 60 $^\circ$C for 30 min. Additionally, thick electrode layers were prepared for the determination of the cohesive strength in the dry electrode layer. Slurries were poured into a $26 \times 49 \times 11$ mm$^3$ ($W \times L \times H$) silicone mold and dried at 60 $^\circ$C for 24 h, during the first 3 h of drying an aluminum foil covered one third of the area to suppress the Marangoni effect. The dry layers were then cut and grinded into smaller samples with defined shape. First, the samples were brought into shape using sandpaper with rough texture and then smoothed applying fine sandpaper. Different sample shapes were used to determine the cohesive strength of the electrode layer depending on the applied load type as summarized in Table 1.

**Sample characterization**

**Rheological measurements**

The flow behavior of the anode slurries was characterized applying rotational steady shear measurements. A stress controlled rheometer (Physica, MCR 501, Anton Paar GmbH, Germany) was used to determine the viscosity $\eta$ in the shear stress range $0.1 \text{ Pa} < \tau < 1000 \text{ Pa}$. All measurements were performed with a 5 min waiting period before starting the measurement to ensure structure recovery using a 25 mm diameter plate-plate geometry and 1 mm gap width at 20 $^\circ$C.

**Electrical conductivity tests**

The four-point resistivity test was conducted on a self-manufactured set-up equipped with four measuring probes (S45G, Uwe Electronic GmbH, Germany). Measurements were carried out at five different positions of the 60 mm $\times$ 120 mm anode layer applying five different currents and recording the corresponding voltage. An average value of the electrical conductivity was calculated for a probe spacing of 30 mm according to Smits [54].

**Mechanical properties**

The adhesive strength of the electrode layer to the copper foil was characterized using a 90$^\circ$-peel-test based on the standard DIN 28510-1. A universal testing machine (Texture Analyzer TA.XT plus, Stable Micro Systems, UK) was used to peel off the copper foil. Specimen with a width of 25 mm and a length of 60 mm were fixed to the measuring plate with double-sided adhesive tape (Universal, Tesa SE, Germany). A 500 g plate was set on top of the electrode layer as a pre-compression step to ensure homogeneous contact between adhesive tape and electrode layer. All measurements were performed at a constant peel velocity of 5 mm s$^{-1}$ using a 5 kg load cell force sensor (max. force: 5 kg, force sensitivity: 0.1 g). The measured tensile force was normalized to the line width. This line load was used as a measure for adhesive strength.

The cohesive strength of the electrode layer was determined employing standard test methods for the characterization of mechanical properties. Compressive $\sigma_{c,\text{max}}$ and flexural strength $\sigma_{b,\text{max}}$ (4-point bending) measurements were conducted using a universal testing machine (Texture Analyzer TA.XT plus, Stable Micro Systems, UK) based on the standard DIN 51104 and DIN 843-1, respectively. Additionally, tensile strength $\sigma_{t,\text{max}}$ tests were carried out using custom-made measuring fixtures and DIN ISO
EN 527-4 based sample geometries. Further, the torsional strength \( \sigma_{\text{tor,max}} \) was obtained from dynamic mechanical analysis using a stress-controlled rheometer (Physica, MCR 301, Anton Paar GmbH, Germany) equipped with a 12 mm solid rectangular fixture (SRF 12). Therefore, oscillatory stress-sweep measurements were carried out at a constant frequency of 0.1 Hz until exceeding the destruction-free linear viscoelastic region, indicating the maximum torsional strength of the sample.

Microstructure

The porosity \( \varepsilon \) of the electrode layers was determined from the Archimedes density according to DIN EN 993-1 and DIN EN 993-18. Samples of dry electrodes were vacuum infused with epoxy resin, ground with SiC paper and polished with a diamond suspension to investigate the microstructure of the electrode layers by means of scanning electron microscopy (SEM) using a LEO1530 microscope (Carl Zeiss AG, Germany).

Results and discussion

Rheological characterization

The viscosity \( \eta \) of anode slurries with and without CB as a function of the shear stress \( \tau \) was compared to that of corresponding pure CMC solutions to characterize their flow behavior and the influence of solid particles on the rheological properties of the slurries. Figure 1 shows such flow curves for slurries including CMC with \( M_w = 700 \text{ kDa} \) and \( \text{DS} = 0.9 \) at \( \phi_{\text{CMC}} = 4 \text{ vol\%} \). The CMC concentration refers to the amount of CMC in the dry electrode layer. The slurries show higher viscosity than the pure CMC solution. The relative viscosity \( \eta_{\text{rel}} = \eta_{\text{slurry}} / \eta_{\text{solution}} \) for the slurry comprising graphite and CB is \( \eta_{\text{rel}} \approx 2–3 \) throughout the investigated shear stress range as expected for well dispersed non-Brownian hard sphere suspensions with \( \phi_{\text{p}} = 20 \text{ vol\%} \) [55]. For the slurry containing only graphite, the low shear viscosity is substantially higher (\( \eta_{\text{rel}} \approx 10 \)), indicating incomplete dispersion of particles, i.e. agglomerates immobilizing part of the solvent still exist. At high shear stresses these agglomerates are broken up due to sufficient energy input and the viscosity of both slurries matches.

The influence of DS and \( M_w \) of CMC on the low shear (\( \tau = 1 \text{ Pa} \)) and high shear (\( \tau = 500 \text{ Pa} \)) viscosity of anode slurries was investigated at different CMC concentrations. The low shear viscosity provides information about the particle dispersion in the slurry, whereas the high shear viscosity at constant \( \phi_{\text{p}} \) reflects the amount of CMC dissolved in the fluid phase. The low shear viscosity as well as the high shear viscosity values of anode slurries slightly increase with increasing DS at constant \( M_w \), suggesting that a lower DS results in slightly better particle dispersion and lower amount of free CMC in the fluid phase (see supporting information) as expected due to the higher degree of CMC adsorption on particle surfaces at low DS [16].

The low and high shear viscosity of slurries at different \( \phi_{\text{CMC}} \) and constant \( \text{DS} = 0.9 \) is displayed as a function of \( M_w \) in Fig. 2a and b, respectively. Instead of a monotonic increase of viscosity with \( M_w \) according to a superior thickening efficiency of longer polymer chains we find, that both low and high shear viscosity increases with increasing \( M_w \) up to 400 kDa and then levels off for all investigated \( \phi_{\text{CMC}} \) [56]. This result can be related to the adsorption behavior of CMC on graphite and CB particles. To gain better insight into the \( M_w \) dependence of CMC adsorption on the suspended particles, the relative viscosity at high shear stress (\( \tau = 500 \text{ Pa} \)) \( \eta_{\text{rel,x}} = \eta_{x,\text{suspension}} / \eta_{x,\text{solvent}} \) was plotted against \( M_w \) as shown in Fig. 2c. The characterization of the CMC adsorption behavior using this rheological
approach renders a significant advantage over common techniques like optical methods or adsorption isotherms, which require highly dilute systems. In contrast, rheological measurements enable the determination of CMC adsorption on the solid particles surface at technically relevant particle loadings. The relative high shear viscosity $\eta_{rel,x}$ monotonically decreases with increasing $M_w$, indicating a reduced fraction of free polymer in the solvent when using high $M_w$ CMC. Consequently, the high shear viscosity of the slurry does not increase monotonically with $M_w$. Obviously, the effect of reduced free polymer in solution outweighs the slight increase in effective $\phi_p$ due to the CMC adsorption layer and the effect is independent of $\phi_{CMC}$ in the concentration range investigated here. It must be noted that the addition of SBR does not affect the flow behavior of the slurry. The contribution of the SBR nanoparticles to the overall particle volume fraction is negligible in relation to the active material content.

To shed more light on the adsorption behavior of CMC on graphite particles a series of graphite suspensions with $\phi_p$ ranging from $10^{-2}$ to 20 vol% was investigated with respect to their zero-shear viscosity $\eta_{0,suspension}$ normalized to the zero-shear viscosity $\eta_{0,solvent}$ of the used solvent, an aqueous 1 vol% CMC ($M_w = 700$ kDa, $DS = 0.9$) solution. The selected CMC concentration allows for a conclusive investigation of the adsorption behavior since higher concentrations result in a weaker change in relative viscosity, since a high amount of CMC is dissolved in the continuous phase, whereas lower concentrations would aggravate assessment of reliable relative viscosity data. The zero-shear viscosity of the CMC-solution as well as the graphite suspensions was obtained as the limiting constant viscosity value obtained at vanishing shear stresses. The relative viscosity $\eta_{r,0} = \eta_{0,suspension}/\eta_{0,solvent}$ of these graphite suspensions as a function of particle concentration is shown in Fig. 2d. This quantity exhibits a clear minimum around $\phi_p = 0.2$ vol%, i.e. at a CMC to graphite volume ratio of 5 with a minimum value $\eta_{r,0} \approx 0.6$. In very dilute suspensions ($\phi_p \approx 10^{-2}$ vol%), however, $\eta_{r,0}$ approaches one, as expected. Interestingly, $\eta_{r,0}$ values are below 1 for $\phi_p < 5$ vol%. As confirmed by numerous theoretical and...
experimental studies the relative viscosity \( \eta_r \) of suspensions increases with increasing particle loading and \( \eta_r > 1 \) always holds [57–60]. Therefore, another physical mechanism must control the low shear viscosity of the dilute graphite slurries investigated here. Li et al. showed a decrease of zeta potential for low concentrated graphite and CB suspensions upon addition of CMC, suggesting polymer adsorption on the particle surface [61]. Accordingly, the addition of graphite particles to the CMC solution decreases the polymer concentration in the fluid phase. The corresponding drop in \( \eta_{0,\text{suspension}} \) is more pronounced than the weak increase in viscosity expected due to the added graphite, leading to relative viscosities below 1. At higher graphite concentration (\( \phi_p > 5 \text{ vol\%} \)), however, \( \eta_{r,0} \) strongly increases because of the strong contribution of the particles to the suspensions viscosity. For \( \phi_p = 20 \text{ vol\%} \), \( \eta_{r,0} \) is even somewhat higher than 3, the value expected for non-Brownian hard sphere suspensions [55], and this is attributed to an increase in effective particle volume fraction due to the adsorbed CMC layer. In this concentration range the increase in \( \eta_{0,\text{suspension}} \) with increasing particle concentration dominates over the drop in solvent viscosity due to the increasing fraction of CMC adsorbed to the particles when \( \phi_p \) increases.

**Electrical conductivity**

Anode slurries were coated on glass plates and dried to investigate their electrical conductivity. Figure 3a shows corresponding data for layers including different CMC concentration and constant DS = 0.9 as a function of \( M_w \). Note, the conductivity is due to the formation of a conductive CB particle network [27, 28]. Anode layers including only graphite exhibit overall lower values and an analog dependence on CMC concentration, \( M_w \) and DS as the electrical conductivity of anodes comprising graphite and CB. Irrespective of CMC molecular weight, conductivity of the dry anode layers clearly decreases with increasing polymer concentration as expected since the polymer can deteriorate and interrupt conductive pathways. However, the polymer molecular weight obviously has a significant impact on conductivity too. At given \( \phi_{\text{CMC}} \), the absolute values obtained for the layers including the CMC with \( M_w = 700 \text{ kDa} \) are about 50% higher than that for the layers prepared using CMC with \( M_w = 250 \text{ kDa} \). It is well known from literature and it is also obvious from our rheological investigations discussed above that the mass of polymer adsorbed to the particle surface increases with increasing molecular weight [62–64]. Consequently, the fraction of CMC randomly distributed in the dried samples decreases with increasing \( M_w \) and it seems this fraction deteriorates the electrical conductivity of the anode layers investigated here. This new insight into the adsorption behavior of CMC with different polymer chain-length and its relationship to particle dispersion and electrical conductivity offers a path to increase conductivity through targeted choice of CMC molecular architecture at constant \( \phi_{\text{CMC}} \). Hence, power and energy density of corresponding cells might be improved. Finally, it should be noted that the degree of substitution has no significant effect on dry anode layer conductivity irrespective of CMC concentration (data not shown). This is in good agreement with rheological data indicating a weak effect of DS on low and high shear viscosity of the wet anode slurry.

**Figure 3** Electrical conductivity of dry electrode layers as a function of \( M_w \) including CMC at different concentrations and constant DS = 0.9 (a) as well as of \( \phi_{\text{SBR}} \) including CMC with constant \( \phi_{\text{CMC}} = 2.5 \text{ vol\%} \), \( M_w = 700 \text{ kDa} \) and DS = 0.9 but varying SBR-type.
Considering the high conductivity values of anodes including CMC at low concentration and high molecular weight, the influence of SBR type and concentration on the electrical conductivity of anode layers was investigated at constant $\phi_{\text{CMC}} = 2.5 \text{ vol\%}$, $M_w = 700 \text{ kDa}$ and $DS = 0.9$ (Fig. 3b). Increasing $\phi_{\text{SBR}}$ leads to a monotonic decrease of electrical conductivity. Rheological data of corresponding wet slurries showed no significant effect of SBR on the flow behavior, i.e. variation of $\phi_{\text{SBR}}$ hardly affects particle dispersion and polymer adsorption. Hence, the decay in electrical conductivity is assumed to be a consequence of randomly distributed isolating polymer particles, blocking conductive pathways. Interestingly, anodes including TRD105A yield overall higher values than those of anodes based on TRD2001. This might be due to the smaller particle size or due to proprietary differences in polymer architecture or specific electrical properties. Elucidating this is, however, beyond the scope of this work focusing on the role of CMC as binder for LiB electrodes.

**Mechanical properties and electrode microstructure**

**Adhesion**

The influence of CMC on the adhesive strength between electrode layer and copper foil was investigated using the 90°-peel test. In all cases complete detachment of the anode layer from the current collector was observed (see supporting information). Figure 4a shows the line load of the measured samples over DS at constant $M_w = 250 \text{ kDa}$, whereas Fig. 4b displays line load data as function of $M_w$ at constant DS = 0.9, in both cases $\phi_{\text{CMC}}$ was varied between 2.5 and 4 vol%. Considering the experimental uncertainty of the measurements there is no significant dependence of adhesive strength on $M_w$ or DS. In all investigated cases, the adhesive strength between electrode layer and current collector lies between 0.6 and 1.8 N m$^{-1}$, and it must be noticed that the obtained values are more than one order of magnitude lower than that for water-based anodes containing an additional latex or rubber as binder system in combination with CMC [35, 43, 65]. This clearly shows that CMC hardly contributes to adhesion and that the addition of a second polymer is essential for the bond strength between electrode layer and current collector. Peel tests performed for SBR films on copper foils showed an average line load of $948 \pm 47 \text{ N m}^{-1}$ and $832 \pm 53 \text{ N m}^{-1}$ for TRD2001 and TRD105A, respectively, whereas CMC films, irrespective of $M_w$ and DS, yielded only an average of $2.3 \pm 0.7 \text{ N m}^{-1}$. Accordingly, the intrinsic contribution of SBR to the adhesion of a LiB electrode is dramatically higher than that of CMC. The addition of SBR significantly enhances the adhesion between anode layer and current collector, increasing linearly with incrementing $\phi_{\text{SBR}}$ (see supporting information). SBR concentrations above 3.0 vol% yield adhesive strength values typical for aqueous anode layers reported in the literature [35, 65] (Fig. 4b).

**Cohesion**

In this section the cohesive strength of thick electrode layers is discussed focusing on the effect of added CMC. Different well-defined mechanical load types were employed to determine the critical stress at which the structure collapses. Results for layers including different concentrations of CMC with $M_w = 250 \text{ kDa}$ and DS = 0.9 are summarized in Fig. 5. Irrespective of load type the respective maximum stress increases almost linearly with increasing $\phi_{\text{CMC}}$. The absolute value of $\sigma_{\text{max}}$ depends on the nature of the load, i.e. it depends on whether the material is being pushed together, pulled apart, bent or twisted. The relative change in failure stress, however, is almost independent of load type. All test methods yield reliable and reproducible results and are therefore, suitable to characterize the cohesive strength of thick electrode layers. For the first time, cohesive and adhesive strength of electrode layers is determined independently. This should help to understand the delamination and cracking behavior due to cyclic lithium intercalation. For sake of convenience we will discuss the compressive strength in the subsequent part of this section. The other methods yielded similar trends upon variation of sample composition (see supporting information).

The porosity of thick electrode layers, used to investigate the cohesive strength, was measured and compared to that of thin layers used for electrical conductivity and adhesion measurements. The overall porosity of all samples was between 55 and 58%, showing no particular influence of CMC concentration, $M_w$, or DS. Similar results were reported in the literature for other water-borne LiB anodes. Thus, a
significant influence of the sample porosity on the measured cohesive strength of thick electrode layers can be disregarded. Additionally, SEM images of thin and thick electrode layers exhibit no recognizable difference in their microstructure (see supporting information). Accordingly, the measured cohesive strength of thick electrode layers, investigated in this work, is assumed to be representative for conventional electrode layers.

Figure 6a displays the compressive strength of thick electrode layers including CMC with $M_w = 250$ kDa as a function of DS and Fig. 6b shows the compressive strength as a function of $M_w$ at constant DS = 0.9. In both cases data were obtained at various $\phi_{\text{CMC}}$ between 2.5 and 4 vol%. First, it should be noted that the cohesive strength of pure CMC films neither depends on $M_w$ nor on DS as directly confirmed by tensile test results obtained for the CMC grades investigated here (see supporting information).

The cohesive strength of electrode layers, however, strongly depends on these molecular parameters. On one hand, it significantly decreases with increasing DS. This decay is more pronounced at higher $\phi_{\text{CMC}}$, and for DS = 1.2 the compressive strength hardly varies with $\phi_{\text{CMC}}$. Adsorption isotherms for CMC on graphite particles show higher adsorption values for CMC with lower DS [16] corresponding to a lower solubility in water. Our results clearly show that this also has an effect on mechanical strength of the electrode layer and we assume that this is due to different particle dispersion and dry electrode layer microstructure depending on DS.

On the other hand, the cohesive strength of the thick dry layers increases with increasing $M_w$, particularly at high $\phi_{\text{CMC}}$. The high cohesive strength obtained at high $M_w$ again seems to be related to the higher fraction of CMC adsorbed on the surface of the graphite particles (s. Figure 2c). We hypothesize that the CMC molecular weight and the related variation in adsorption to the graphite and CB particles has a strong impact on particle dispersion as already indicated by its impact on slurry viscosity and in turn on microstructure and mechanical strength of the dry layer. This will be discussed in the next section. These findings indicate that electrode cohesion can be significantly increased without
increasing CMC concentration but through proper choice of molecular structure.

Finally, it should be noted that SBR does not contribute to the cohesive strength of thick anode layers irrespective of amount or type of added SBR (see supporting information). This is partly due to its lower intrinsic strength. Tensile tests performed on pure SBR films based on the standard DIN ISO EN 527-4 yielded intrinsic mechanical strength values of $4.3 \pm 0.7$ Nm m$^{-2}$ and $9.2 \pm 1.4$ Nm m$^{-2}$ for TRD2001 and TRD105A, respectively, whereas for CMC values $\geq 15$ N mm$^{-2}$ were obtained, irrespective of $M_w$ and DS. In addition, previous work [66] indicated that the addition of nanoparticulate secondary polymers hardly affects the electrode microstructure, which in turn controls the cohesive strength.

**Microstructure**

As already mentioned above, the porosity of all investigated samples was in the range between 55 and 58% irrespective of CMC concentration, $M_w$ or DS. In more detail, SEM images of thick electrode layers were taken to investigate their microstructure. As shown by compression tests, a significant influence of $M_w$ on the cohesive strength is obtained for $\phi_{\text{CMC}} = 4.0$ vol% and DS = 0.9. Hence, SEM images of thick layers including 4 vol% CMC with $M_w = 250$ kDa and $M_w = 700$ kDa (DS = 0.9 in both cases) were compared (Fig. 7a, b). Electrodes including the low molecular-weight CMC show a random orientation of the flake-shaped graphite particles, whereas corresponding electrodes containing high molecular-weight CMC exhibit a clear particle alignment. Image processing was carried out for both images to quantify this observation. Individual particles and particle collectives were detected and their angle of orientation in relation to a centered coordinate system was noted, as marked in red (Fig. 7a, b). The particle frequency as a function of the orientation angle for samples including CMC with $M_w = 250$ kDa, $M_w = 400$ kDa and $M_w = 700$ kDa at constant DS = 0.9 is shown in Fig. 7c, d and e, respectively. For the $M_w = 250$ kDa sample, image analysis yields a broad distribution of particle orientation i.e. a random isotropic layer structure. In contrast, high molecular-weight CMC ($M_w = 700$ kDa) leads to particle alignment as confirmed by a narrow distribution of particle orientation angle. This kind of organized microstructure apparently correlates to the high cohesion values obtained for thick electrode layers based on $M_w = 700$ kDa CMC. Anodes including CMC with $M_w = 400$ kDa show an intermediate degree of particle orientation relative to those of anodes based on low and high $M_w$. This systematic change in particle alignment correlates to the linear increase of cohesive strength with increasing polymer chain-length, i.e. particle orientation.

Furthermore, SEM images of thick electrode layers including CMC at $\phi_{\text{CMC}} = 4$ vol% and constant $M_w = 250$ kDa, but varying DS (DS = 0.7, DS = 0.9, and DS = 1.2) were compared to investigate the influence of DS on the microstructure of these layers. The corresponding histograms are shown in Fig. 7c, f and g. Electrode layers based on CMC with DS = 0.7 result in a narrow distribution of the particle orientation angles, indicating clear particle alignment. Intermediate DS leads to a weak particle alignment towards 100°, whereas the highest DS results in randomly oriented particles, i.e. a broad uniform distribution of orientation angles. Analog with the results
obtained for electrode layers comprising CMC with different $M_w$, the particle orientation in the electrode layer correlates to the cohesive strength when using different DS. These results demonstrate that the cohesive strength of electrode layers is determined by the microstructure formed due to the added CMC. This structure and hence the mechanical strength of the layers strongly depends on $M_w$ and DS of the added CMC, although the intrinsic mechanical strength of CMC itself does not depend on these parameters. The high degree of particle alignment at high $M_w$ and low DS is presumably promoted by the lower viscosity of the continuous phase of the corresponding slurry as a result of the higher fraction of CMC adsorbed on the particles.
Conclusions

Here we investigated the effect of CMC, which is widely used as a binder and rheology control agent in waterborne LiB electrode slurries, on the flow behavior of anode slurries as well as on the electrical and mechanical properties and finally also the microstructure of the corresponding dry layers. CMC concentration, \( M_w \) and DS were systematically varied in a wide technically relevant range. In addition, the influence of SBR type and concentration on the features mentioned above was discussed. This holistic approach establishes new insights into the polymer adsorption behavior on active particles at industrially relevant slurry concentrations as well as clear correlations between particle dispersion, layer microstructure as well as electrical conductivity and mechanical integrity of the electrode.

Shear rheometry revealed that both, low and high shear viscosity strongly increase with \( M_w \) up to 400 kDa and then level off irrespective of the amount of added CMC but, as expected, the absolute viscosity values monotonically increase with \( \phi_{\text{CMC}} \). Moreover, the relative high shear viscosity \( \eta_{r,\infty} \) is essentially independent of \( \phi_{\text{CMC}} \) but monotonically decreases with increasing \( M_w \). We attribute this effect to the partial adsorption of CMC on the solid particles and the corresponding loss of polymer in the solvent. Obviously, the loss of dissolved polymer is more pronounced for higher \( M_w \), as expected [62–64]. This phenomenon was further supported considering the relative viscosity of graphite suspension in a CMC solution covering a broad range of particle loading. The higher affinity of CMC to the graphite surface at lower DS also shows up in the rheological data. Finally, the addition of SBR does not affect the flow behavior of anode slurries.

The electrical conductivity decreased with increasing \( \phi_{\text{CMC}} \) as expected, but strikingly conductivity substantially increased with increasing \( M_w \) at fixed \( \phi_{\text{CMC}} \). Since the fraction of adsorbed CMC increases with increasing \( M_w \), we hypothesize that the remaining free polymer deteriorates the formation of a percolating CB network and hence the conductivity of the dry anode layers. Accordingly, adding CMC with high \( M_w \) at constant \( \phi_{\text{CMC}} \) results in an increase of electrical conductivity, which is associated to cell power and energy density. The weak changes of CMC adsorption with varying DS did not show up in dry layer conductivity. As expected, increasing SBR concentration also resulted in a monotonic decay of electrical conductivity.

The adhesive strength of dry layers including CMC as only binder component to copper foils is about two orders of magnitude lower than that of conventional, water-based anode layers including a secondary binder and hardly varies with \( \phi_{\text{CMC}} \), \( M_w \) or DS. Obviously, CMC does not contribute to the adhesive strength of LiB anode layers. Addition of SBR leads to a linear increase of adhesion at the interface, highlighting the role of rubber-like polymer binders.

The cohesive strength of dry electrode layers has to balance the mechanical stress during cyclic intercalation of Li-ions in the layers, and is supposed to be related to cracking and mechanical failure of battery electrodes. We investigated this phenomenon employing different load types to characterize the critical stress \( \sigma_{\text{max}} \) at which thick electrode layers break. As expected, \( \sigma_{\text{max}} \) significantly increases with \( \phi_{\text{CMC}} \) and all employed test methods yielded reliable results. Preliminary porosity measurements and SEM-images confirmed, that the microstructure of the thick layers was similar to that of the thin layers investigated regarding electrical conductivity and adhesive strength. Therefore, we consider the results of these cohesion tests relevant also for thin LiB electrode layers, rendering first cohesion values for water-based anodes determined under well-defined mechanical load conditions independent of electrode adhesive strength.

Although the mechanical strength of pure CMC films neither depends on \( M_w \) nor on DS, we found a strong dependence of compressive strength of electrode layers on both parameters: \( \sigma_{\text{max}} \) decreases with increasing DS but strongly increases with increasing \( M_w \), i.e. with increasing fraction of adsorbed CMC. Apparently, the degree of CMC adsorption on graphite particles also must have a strong impact on the microstructure determining the mechanical strength of the dry layers. This change in microstructure was directly visualized using scanning electron microscopy. Image analysis of these micrographs revealed a strong orientation of graphite particles for layers including CMC with high \( M_w \) or low DS, i.e. when a larger fraction of CMC is adsorbed on the particle surface. This effect of CMC adsorption also seems to be decisive for the electrical conductivity of these layers. Contrary to its effect on adhesive strength, the addition of SBR has no influence on the cohesive strength.
In summary, the variation of CMC architecture (Mw, DS) and the corresponding change in adsorption on graphite particles not only affects the flow behavior and hence processing properties of the anode slurries but also the microstructure of the dry layers and hence their electrical conductivity as well as cohesive strength. The holistic study presented in this work renders a comprehensive and systematic understanding of the role of CMC in water-based anodes for LiB, unveiling clear correlations between polymer/particle interactions and resulting electrode properties. Accordingly, the targeted choice of CMC is supposed to be decisive not only for fabrication but also for electrical performance and long-term mechanical stability of LiB electrodes.

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