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Analysis of the Interphase on Carbon Black Formed in High Voltage Batteries

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Carbon black (CB) additives commonly used to increase the electrical conductivity of electrodes in Li-ion batteries are generally believed to be electrochemically inert additives in cathodes. Decomposition of electrolyte in the surface region of CB in Li-ion cells at high voltages up to 4.9 V is here studied using electrochemical measurements as well as structural and surface characterizations. LiPF6 and LiClO4 dissolved in ethylene carbonate:diethylene carbonate (1:1) were used as the electrolyte to study irreversible charge capacity of CB cathodes when cycled between 4.9 V and 2.5 V. Synchronron-based soft X-ray photoelectron spectroscopy (SOXPES) results revealed spontaneous partial decomposition of the electrolytes on the CB electrode, without applying external current or voltage. Depth profile analysis of the electrolyte/cathode interphase indicated that the concentration of decomposed species is highest at the outermost surface of the CB. It is concluded that carboxylate and carbonate bonds (originating from solvent decomposition) and LiF (when LiPF6 was used) take part in the formation of the decomposed species. Electrochemical impedance spectroscopy measurements and transmission electron microscopy results, however, did not show formation of a dense surface layer on CB particles.

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The growth of earth’s population with concomitant increase in energy consumption require development of renewable energy conversion technologies coupled with advanced energy storage systems like lithium batteries.1,2 In order to increase the power density in Li-ion batteries, much research is focused on developing cathode materials that can operate at high voltages (above 4.5 V vs. Li/Li+) with a high capacity, high cycling stability, and good rate capability.1–5 However, at high voltages, all the components of positive electrodes including the Al current collector, polymer binders, conductive additives, and other possible additives have an increased risk of degradation. In addition, one of the main issues with high voltage batteries is the instability of common aprotic electrolytes at voltage above 4.5 V.6,7 The stability of the electrolyte/cathode interphase is related to the chemistry of electrolyte solvents and salts and also to the chemistry of the components of the cathode.

Carbon black (CB) additives are one of the main constituents of cathodes, added to increase the electrical percolation and thus the electrochemical conductivity.3,8–10 Though the weight percentage of CB in commercial batteries is generally very small, it composes a rather large part of the internal surface area of a cathode due to its small particle size (∼50 nm), low density, and high surface area. CBs are generally thought of being an electrochemically inert additive in cathodes, but few studies have investigated the role of CBs at high voltages and have indicated that CBs exhibit irreversible electrochemical reactions resulting in appreciable irreversible charge capacities.10–18 This charge capacity is attributed to oxidation reactions, anodic degradation of aprotic electrolytes on the surface of CBs, side reactions involving binder and salt, and intercalation of anions such as PF6− (partly reversible) and solvent molecules into graphitic layers.10–19 The oxidation voltage, decomposition products, and possible formation of a surface layer are dependent on the chemistry of electrolyte and the surface area and the surface functional groups of CBs.10–14 This is similar, but not identical, to the concept of formation of solid electrolyte interphase (SEI) on anodes. This SEI-like layer is often referred to as a solid permeable interphase (SPI)20 or cathode electrolyte interphase (CEI)21 and can ideally prevent further parasitic reactions between electrolyte and cathode materials. According to La Mantia et al. the oxidation of surface active groups occurs around 4.5–4.6 V.12 They showed that the first charge capacity is dependent on the charge rate, which could indicate the formation of a SPI layer. This surface layer was suggested to be made of Li2CO3, LiF, and polycarbonates, but was not analyzed to prove.12 In two recent studies Demeaux et al. and Syzdek et al. suggested decomposition products from the electrolyte on the surface of carbon particles at high voltages using FT-IR.11,14 A surface layer of decomposed electrolyte formed on carbon particles can decrease electronic conductivity, and thus, hinder the overall performance of the cathode.22

The aim of this work is to study parasitic reactions and possible formation of a surface layer on Super P, as one of the most common CB, in cathodes during the first few cycles at high voltages in Li-ion cells with two different electrolyte salts, LiPF6 and LiClO4. The surface layer on cathodes can be very thin, and thus, difficult to analyze by SEM, XRD,14 or conventional X-ray photoelectron spectroscopy (XPS). Therefore, we used synchrotron-based soft X-ray photoelectron spectroscopy (SOXPES), which is one of the most powerful techniques for studying the outermost surface of compounds, to analyze the surface of CBs. We chose to use relatively low excitation photon energies to detect both crystalline and/or amorphous chemical compounds at shallow depths from the surfaces of the cathodes. Tuning the photon energies, we could achieve a nondestructive depth profiling of the surface layer. In addition electrochemical impedance spectroscopy (EIS), transmission electron microscopy (TEM) and energy-dispersive X-ray spectroscopy (EDS) were used to characterize the change in morphology, structure, and chemistry of CB before and after charging to high voltages.

Experimental

Li-ion cells were assembled in a “Coffee-bag” (pouch cell) configuration23 using carbon black electrodes as the positive electrode, Li metal as the negative electrode, and two layers of Celgard 2500 (Monolayer Polypropylene) membrane (dried at 60 °C overnight) as the separator. Cathodes were prepared by mixing Super P carbon black (purchased from Erachem Comilog N.V.) with Kynar binder (Kynar Flex 2801, Arkema) dissolved in (N-Methyl-2-Pyrrolidone) and casting on Al foil. The ratio of carbon:binder was 70:30 by weight. The electrodes were dried at 120 °C overnight and were cut to discs of 10 mm in diameter. A total mass of 0.42 mg Super P was used for each electrode. The electrolyte mixture consisted of 1 M of either LiPF6 or LiClO4 salts dissolved in EC:DEC (vol. 1:1), where EC is ethylene carbonate and DEC is diethyl carbonate purchased from BASF. 200 µl of electrolyte were added to each cell. The electrochemical characterizations were performed using a BioLogic VMP3
multichannel analyzer. A constant current of 5 μA yielding the current density of about 6.4 μA cm⁻² and voltage limits of either 4.9–2.5 V or 4.3–2.5 V were applied for cycling cells used for SOXPES analysis. For comparison, identical cells were assembled and stored for 3 days without applying external current or voltage (hereafter referred to as "stored" electrodes). Similar cells were assembled and cycled between 5.2 V and 2.5 V for TEM and EDS analysis. Electrochemical impedance spectroscopy (EIS) measurements were performed in a three-electrode EL-CELL ECC-Combi using lithium metal as counter and reference electrodes. The cell was assembled inside the glove box using glass fiber separator (and 200 μl electrolyte solution. EIS measurements in a frequency range from 500 kHz to 10 mHz of CB electrodes were obtained before (stored) and after stepwise charge to 4.9 V. All measurements were performed at OCV after the cell had reached steady state defined by a change <5 mV/h. The impedance results were modeled using the equivalent circuit presented in Figure 1. The RₑRₑCₓA elements model the high-frequency region where R corresponds to a resistance and C corresponds to a capacitor. These values have been normalized to the geometrical surface area of the electrode (0.785 cm²). The low-frequency region has been modeled with a simplified transmission line (TL) for a porous electrode, according to De Levie model.24,25 This model involves a cylindrical pore with length L and radius r, filled with the electrolytic solution, and the electronic resistance of the electrode is assumed to be much lower than the ionic resistance of the solution (Rₑ ≤ Rₑl). In case of a non-faradaic process, the overall impedance inside the pore is equal to:

\[
Z = \frac{R_{\text{ion},L}}{j\omega C_{\text{dl}}A^{2}\pi r} \coth \left( \frac{R_{\text{ion},L} \ast j \omega C_{\text{dl}}A^{2}\pi r}{2} \right)
\]

where \(R_{\text{ion},L}\) is the ionic resistance of the electrolyte per unit pore length (Ω cm⁻²) and \(C_{\text{dl}}\) is the electrical double layer capacitance per unit surface area (F cm⁻²). From this model we calculated the total double layer capacitance of the CB electrode \(C_{\text{dl}}\) (F). The uniform transmission line for a flooded ideally polarized porous electrode can be seen in Figure 1. This model makes a good fit of the high- and low-frequency regions of the experimental data, but shows a small deviation in the mid-frequency part. Despite this, the model has been chosen as it provides an acceptable fit with a meaningful physical interpretation to the interface between CB surface and electrolyte in the extended porous network. The double layer capacitance is related to the total surface area \(A\):

\[
C_{\text{dl}} = \varepsilon_{\text{d}} \frac{A}{\varepsilon_{0}}
\]

where \(\varepsilon_{\text{d}}\) is the dielectric constant, and \(\varepsilon\) and \(l\) is the dielectric constant and the thickness of a surface layer, respectively. After cycling, Li-ion cells were transferred to MAX-IV Laboratory where they were dismantled in an Ar-filled glove box (H₂O < 1 ppm). The carbon electrodes were washed with several drops of dimethyl carbonate (DMC). This washing is necessary to remove remaining electrolyte species in order to obtain spectra from the surface of the samples. The samples were mounted in a specially designed transfer chamber and transported to the analyzing chamber without exposure to the atmosphere. The measurements were performed using synchrotron radiation (beamline I-411) at the MAX IV Laboratory. Photons were monochromatized by a Zeiss SX-700 planar grating monochromator. Core level spectra of carbon (C 1s) spectra were measured using two different photon energies of 430 eV and 835 eV to obtain depth profile analysis of surface layer on carbon cathodes. Core level spectra of fluorine (F 1s) spectra were measured using photon energies of 835 eV. The spectra were energy calibrated using the hydrocarbon peak at 285.0 eV. The spectra were intensity normalized to 1. Curve fitting was performed using IGOR Pro, and a linear background was subtracted before the spectra were deconvoluted. TEM imaging and selected area electron diffraction (SAED) of CB was carried out by using a JEOL 3000 F equipped with a 300 kV FEG. EDS was carried out for compositional analysis by using an Oxford Instruments detector with an ultra-thin window. The analysis was performed using the Inca software. To compare the intensities in the SAED data, the patterns were circularly integrated and the full intensities were normalized to the intensity of the first diffraction ring. XRD analyses was performed using a Bruker D8 TwinTwin X-ray diffractometer operating at 40 kV and 40 mA using Cu Kα radiation (\(\lambda = 1.5406\) Å).

Results and Discussion

Figure 2a shows the first charge curves of Li-ion cells assembled using CB cathodes with LiPF₆ or LiClO₄ as the electrolyte salts. It can be seen that the LiClO₄-based cell reached the anodic limit of 4.9 V much faster than the LiPF₆-based cell. In other words, the LiClO₄ based cell provided a smaller charge capacity, about 50 mAh·g⁻¹, while the LiPF₆ cell yielded a larger charge capacity of about 800 mAh·g⁻¹. These capacities are comparable (considering differences in applied current densities) to previously reported results for Super P carbon black and noticeably smaller than capacities of similar cells with graphitic carbons or high surface area carbons like Ketjen black which provides irreversible charge capacity above 2000 mAh·g⁻¹.12,13

To investigate the reversibility of these capacities, the cells were discharged to 2.5 V and then cycled 10 times between 4.9 and 2.5 V. Figure 2b shows that the charge capacities were significantly higher than the discharge capacities indicating irreversible electrochemical processes during charge. In addition, the charge capacity decreased significantly from the first cycle to the following cycles (note that the capacities are presented on a logarithmic scale). The small discharge capacity is the same in all cycles indicating that a minor reversible reaction occurs on the CB cathodes. This reversible capacity is due to the sum of the accumulated surface capacitance and the charge transfer from any reversible electrochemical processes. The latter could be intercalation/deintercalation of PF₆⁻ or ClO₄⁻ anions to/from the positive electrode, as this can provide a small reversible capacity.5,15–19

The large irreversible capacity (defined as the difference between the charge and the discharge capacity) can be explained by i) formation of a AlF₃ passivation layer on the aluminum current collector via decomposition of LiPF₆.18,27 and ii) reactions between the electrolyte and the surface functional groups on the carbon surface. In order to clarify the contribution of Al current collector corrosion to the total irreversible capacities observed for CB cathodes, identical cells were assembled using only an Al disk (as it is without any carbon black or active material on it) as cathode. Figure 2c shows that the irreversible
charge ($Q_{irr}$) is order of magnitudes higher in the CB electrode than in an Al electrode in cells with LiPF$_6$ electrolyte salt, showing that the Al current collector corrosion is not the main origin of $Q_{irr}$. However, one may argue that the higher surface area of carbon particles causes the higher degree of $Q_{irr}$ in the CB electrode. Figure 2d shows $Q_{irr}$ per surface area of the electrodes. The surface area for the Al disk is estimated to be 0.785 cm$^2$, assuming a smooth surface. In case of CB electrodes, the surface area is estimated by using a BET value of 62 m$^2$·g$^{-1}$ for CB powder, assuming that all particles are available for charge accumulation i.e. no surface blocking due to binder etc. Thus, the total surface area of CB electrode is assumed to be equal to 260 cm$^2$. Figure 2d shows that $Q_{irr}$/cm$^2$ of Al disk is significantly deceased after few cycles, which is known to be due to formation of a passivation layer of AlF$_3$. However, $Q_{irr}$/cm$^2$ of CB electrode is still about one order of magnitude higher for CB in the following cycles. This finding confirms that $Q_{irr}$ in long term cycling is originated from CB particles. Winter et al. have shown that the nature and properties of CBs such as presence of functional groups and graphitic structure influence irreversible capacities at high voltage. They have shown that thermal treatment of CB results in graphitization of CB as well as removal of functional groups, which consequently can decrease $Q_{irr}$ of CB.

To investigate a possible formation of electrolyte decomposition species in or on CB particles at high voltages, CB electrodes were analyzed using synchrotron-based SOXPES. Figure 3a shows C1s spectra of a CB cathode cycled 10 times between 4.9–2.5 V using LiPF$_6$ as the electrolyte salts. Also, C1s spectra of a CB cathode that was cycled between 4.3–2.5 V, as well as, C1s spectra of pristine and stored CB electrodes are presented in the Figure 3a. The C1s spectrum of the pristine sample contains 4 main contributions at the binding energies of 284.6 eV, 285 eV, 286.5 eV, and 290.9 eV assigned to Super P, hydrocarbons, CH$_2$ (from binder), and CF$_2$ (from binder), respectively. There are also two small contributions at binding energies of 288.7 eV and 293.5 eV representing CF and CF$_3$, respectively, in Kynar binder. Interestingly, the C1s spectrum of the stored sample shows different features compared to the spectrum of the pristine sample, as the stored sample has a higher intensity/contribution of the hydrocarbon peak (red peak at 285 eV). Also, the intensity of the peak at 286.5 eV (green peak) is increased, which indicates presence of C–O bond in the surface region of the stored CB particles. In addition, a small peak has appeared at 287.5 eV originating from O–C–O and/or C=O bonds. The changes in the spectrum of the stored sample compared to the pristine sample shows that the electrolyte solution partially decomposes in/at the surface of CB particles when the electrode is immersed in the electrolyte. It should be mentioned again that all the electrodes were washed with DMC before SOXPES measurements to make sure no electrolyte remained on the surface of electrodes. Similar, but not identical, results regarding the surface degradation on stored carbon electrodes have been observed in other studies using FTIR. Demeaux et al. proposed that polyether and polycarbonate species form on the surface of CB samples stored for longer time (4 weeks) at higher temperature (40°C). While our SOXPES confirms formation of ether-based species, we did not observe formation of carbonate-based compound on the stored samples. Also, Demeaux et al. suggested a formation of a surface layer based on scanning electron microscopy (SEM). However, Syzdek et al. did not detect any surface layer in SEM results, and this was explained by the possibly removal of any surface layer by DMC prior to imaging. The formation of decomposition species in the stored sample may take place by a redox reaction as the cells have open circuit voltage about 3 V vs. Li/Li$^+$. Compared to well-studied storage degradation of cathode materials, less attention has been paid to spontaneous reaction occurring in the surface region of stored CB. Thus, the presented results are useful for detecting the origin of surface layer on stored cathodes, because decomposition products originated from active materials and carbon black may overlap with each other.

Surprisingly, the C1s spectrum of the CB cathode cycled 10 times up to 4.3 V present similar features to the C1s spectrum of pristine sample indicating that no major decomposition product is present in the CB cathode when cycled up to 4.3 V. The C1s spectrum consists of extra minor contribution from O–C–O and/or C=O bonds compared to the pristine sample. The results could suggest that the decomposition species in the surface of the stored CB diminishes when the CB cathode is cycled up to 4.3 V. This can, for example, occur by oxidation and/or dissolution/desorption.

Cycling the CB cathode between 4.9–2.5 V using LiPF$_6$ electrolyte, we could detect an increase in the relative intensity of peaks at 287.5 eV and 290.9 eV. The former originates from O–C–O and/or C=O bonds while the latter from carbonate species (CO$_3$).
increase in the relative intensity of these peaks implies that the CB cathode decompose EC:DEC solvent partially when the cell was cycled to 4.9 V.

To obtain a depth profiling of the surface layer, the CB cathodes were analyzed using two different photon energies of 430 eV and 835 eV, as presented by the black and red spectra, respectively, in Figure 3b. For the pristine CB electrode, the C1s spectra look similar using these two photon energies, as expected. A minor difference is that the C1s spectrum measured with lower photon energy shows slightly more contribution from binder (peaks at 286.5 eV, and 290.9 eV), indicating that concentration of binder is slightly higher on top surface. The stored and cycled samples display that the C1s spectra are more similar to the pristine sample for the higher photon energy of 835 eV. The spectra obtained with the higher photon energy are originated from increased depth of about 5–7 nm. Therefore, the highest concentration of electrolyte decomposition species are found in a thin, about 1–3 nm, surface region of the CB.

Figure 4a shows C1s spectra of CB electrodes stored and cycled to 4.9 V using LiClO4 in EC:DEC electrolyte. Both spectra demonstrate presence of decomposed electrolyte species on the surface of the sample. However, compared to the LiPF6-based samples, the spectra of LiClO4-based sample show a less degree of difference from the spectrum the pristine sample. Also, the spectra of LiClO4-based samples measured by two photon energies of 430 eV and 835 eV (black and red spectra respectively in Figure 4b) are more similar than those for LiPF6-based samples (Figure 3b). This means that a smaller amount of degradation products are formed in the surface of CB cathodes in LiClO4-based cells compared to the LiPF6-based cells. This is in agreement with the capacity results presented in Figure 2, which indicated that charge capacity of LiClO4-based cells is smaller than that of LiPF6-based cells.

The F1s spectra of all the samples look more or less similar since the peak in the spectra are substantially originated from binder of electrodes (Figure 5). The peak at 688 eV mainly represent bond between F and C in the Kynar binder. However, a closer look at the spectra reveals that a small shoulder is present at 685 eV in the spectra of stored and cycled samples when LiPF6 was used as the electrolyte salt. This small contribution represents LiF formed due to partial decomposition of LiPF6, suggesting that decomposed LiPF6 salt is found in the surface region of the CB.

As described in detail above decomposed electrolyte species are found in the surface region of the CB particle according to the SOX-PES results. The structure of these decomposed electrolyte species is, however, not clear from the SOXPES results. It could for example be speculated that the species are present as layers partly covering the CB surfaces, or alternatively that the species are integrated into the CB top surface layer. To address this question, high-resolution TEM (HRTEM) was applied and HRTEM images of the pristine and charged CB electrodes are shown in Figure 6. From these images no surface layers on the CB particles can be observed in any of the samples. However, the internal structure of the pristine CB particles indicates the presence of graphitic structures approximately in the direction along the surface of the primary CB particles, while this structure is less pronounced for the charged sample (HRTEM image of the stored sample is presented in Figure S1 displaying slightly changes between pristine and stored CB particles). These results could indicate a loss in crystallinity after storing and charging. Also, for the charged sample the surface of particle became less sharp and in many areas the damaged graphitic was observed. EDS analysis showed that phosphorus and fluorine signals increase from stored to charged samples, suggesting that more PF6− originated compounds are present in the charged sample (see Figure 7). These results together with the...
Figure 4. (a) Deconvoluted C1s spectra of stored and cycled CB cathodes using LiClO4 in EC:DEC electrolyte. The spectra were measured by photon energy of 430 eV. (b) C1s spectra of the same samples measured using two photon energies of: 430 eV (black) and 835 eV (red).

Figure 5. F1s spectra of pristine, stored, and cycled CB cathodes using LiPF6 and LiClO4 as the electrolyte salt.

SOXPES shows that the decomposed electrolyte species are present, not as layers partly covering the CB particles, but integrated into the surface of the CB. To evaluate the possible loss of crystallinity indicated from the HRTEM images, SAED data for all three electrodes are presented in Figure 8 along with normalized circular integration of the SAED patterns. Three main peaks can be distinguished in the pristine electrode corresponding to the (002), (100) and (110) reflections indicating that a partially graphitic structure exists in SP, which is in agreement with XRD results presented in Figure S2. The broad peaks are indicative of low-graphitized CB with short-range crystalline domains.17 Figure 8 shows that storing and charging CB leads to broadening of the diffraction peaks which indicates a loss in crystallinity, consistent with the HRTEM images. The distance between the sheets in the graphitic domains, which is obtained by the position of the d_{002} reflection, is determined to 3.3 ± 0.1 Å for the pristine electrode. This value is slightly increased to 3.5 ± 0.1 Å for the charged electrodes. This increase is much smaller than the increase expected for complete intercalation of PF6− anions into the graphitic domain (i.e. 4.5 Å).19 Finally, TEM images were recorded at lower magnification to evaluate the overall structure of the three samples. This analysis showed that the mean particle sizes slightly changed (see Figure 9). In the pristine electrode, the mean particle size is 33.2 ± 1.2 nm, in the stored electrode it is 35.3 ± 1.0 nm and after cycling (at the charged stat) the value has increased to 40.2 ± 1.6 nm. The loss of crystallinity and particle swelling of the stored and charged CB samples could possibly be explained partly by absorption of electrolyte solution followed by structural rearrangement of the internal CB structure. Also, the integration of decomposed electrolyte species to CB particles could influence CB particles.

The impedance result of a stored CB electrode is presented in Figures 10a and 10b. The Nyquist plot consists of a large semicircle in the high frequency range (between 10 kHz and 10 Hz) and an almost vertical tail in the low frequency range. The high frequency side of the semicircle intersects with the x-axis at 5 Ωcm² (Rq). This value originates mainly from the ionic resistance of the electrolyte between
Figure 6. HRTEM images of CB samples: (a-c) pristine and (d-f) charged.

Figure 7. Normalized EDS spectra of stored (solid) and charged (dash) electrodes. A larger content of F and P was found in the charged electrode. Al originates from the preparation step when CB was scratched from the Al current collector. Cr and Au originate from the TEM grid.

Figure 8. SAED patterns of (a) pristine, (b) stored, and (c) charged CB samples. Corresponding intensity profiles from the SAED patterns are shown in frame (d). The intensities are normalized to the intensity of the first diffraction ring (002). Inset of frame (d) shows a zoom-in of the first diffraction peak.

Figure 9. TEM images of (a) pristine, (b) stored, and (c) charged CB electrodes.
the cathode and the Li reference electrode. The semicircle (R_{Al,CAl}) can be assigned to the interface between the aluminum current collector and the porous carbon network. The low-frequency part (T_{LS}) of the Nyquist plot shows a long capacitive tail, which refers to the double layer capacitance of the ion-blocking CB surface in the porous electrodes. Since the measurements are obtained at OCV no charge transfer reaction is expected. The Bode plot in Figure 10b shows the frequency domain of the impedance response. The impedance response from a CB electrode before, during, and after stepwise charge to 4.9 V is shown in Figure 10c. All measurements were performed at OCV after the cell had reached steady state. An increase in the semicircle and a small change in the angle of the tail are observed. Fitting of each plot was performed and the values for R_{Al}, C_{Al}, R_{ion,L}, and C_{dl} are extracted and shown as a function of the charge potential in Figure 11. The Al/CB interface resistance (R_{Al}) remains constant up to 4.3 V after which it increases slowly until 4.85 V and suddenly increases significantly at 4.9 V. The capacitance of the Al/CB interface (C_{Al}) is almost constant until 4.8 V after which it suddenly decreases. The increase in R_{Al} (Figure 11a) could be explained by the growing of resistive surface layers between aluminum and carbon particles. From 4.0 V and above an AlF_{3} surface layer will form on aluminum which fits with the increase in the slope above 4.3 V. A decrease in C_{Al}
(Figure 11c) after 4.8 V supports that an interphase layer is formed. A formation of an AlF$_3$ layer will lead to an increase of the thickness of double layer and to a reduction of dielectric constant $\varepsilon$ (EC and DEC have $\varepsilon$ equal to 89.6 and 3.12 respectively, while AlF$_3$ has a $\varepsilon$ equal to 2.2). According to equation 2 these factors will both lead to a reduction of $C_{dl}$. The ionic resistance of the electrolyte in the pores ($R_{ion}$) is almost constant until 4.85 V after which it suddenly increases at 4.9 V. This increase is possibly linked to a change in the pore structure induced by the particle growth. Finally, the carbon surface double layer capacitance ($C_{dl}$) increases steadily from 0.95 to 1.25 mF at 4.85 V after which it keeps a constant value (Figure 11d). Normalization of the initial carbon double layer capacitance to the total surface area of the electrode (260 cm$^2$), gives a $C_{dl}$,equal to 3.6 $\mu$F cm$^{-2}$. This value is close to the values found in literature i.e. 5 to 10 $\mu$F cm$^{-2}$. The actual surface area is slightly smaller due to blocking with PVDF binder and isolated particles not connected to the conductive network, and this can explain the smaller value. According to equation 2, the capacitance is directly correlated to a change in the surface area. As indicated from TEM analysis the CB particles swell from 35 to 40 nm, increasing their surface area with 30%. This is in agreement with the observed increase of $C_{dl}$ by assuming that the conductive network is intact. Alternatively, increased wetting of the electrode network could also explain such an increase in $C_{dl}$. Therefore, since no decrease in double layer capacitance was observed, the EIS result cannot prove the formation of a surface layer on top of the CB particles.

Conclusions

Synchrotron-based SOXPEX results revealed that spontaneous decomposition of electrolyte solution (LiPF$_6$ dissolved in EC:DEC) occurs in the surface region of carbon black particles stored in the electrolyte in the absence of external potential and current. Consequently, mainly hydrocarbons and ether species (C–O) originated from EC:DEC, and also, LiF originated from LiPF$_6$ decomposition are formed in the surface of the stored electrode. The decomposition species diminish when the electrode is cycled between 2.5–4.3 V, which suggests that they were oxidized/desorbed when electrochemical processes started. Increased anodic potential to 4.9 V leads to irreversible charges, and thus, to an increase in the impedance of the carbon cathode and to formation of a quite different composition of degraded electrolyte species compared to that formed in the stored electrode. The electrolyte/cathode interphase formed at high voltages consists of relatively higher contribution from O–C–O and/or C=O bonds as well as carbonates compounds (CO$_3$) originating from the solvent, in addition to hydrocarbons, ethers and LiF. Depth profile analysis of the interphase indicated that the concentration of the decomposed species is highest at the outermost CB surface (1–3 nm). TEM and EIS showed no distinctive surface layer indicating that the electrolyte degradation products are likely integrated into the surface region of the CB. The findings suggest that cathode/electrolyte interphases commonly observed on high voltage cathodes may originate from reactions between carbon black and electrolyte and not necessarily between active material and electrolyte.

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