Impact of single vs. blended functional electrolyte additives on interphase formation and overall lithium ion battery performance

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Abstract

Two functional high-voltage additives, namely 2-(2,2,3,3,3-pentafluoropropoxy)-1,3,2-dioxaphospholane (PFPOEpI) and 1-methyl-3,5-bis(trifluoromethyl)-1H-pyrazole (MBTFMP) were combined as functional additive mixture in organic carbonate-based electrolyte formulation for high-voltage lithium battery application. Their impact on the overall performance in NMC111 cathode-based cells was compared with the single-additive-containing electrolyte counterpart. The obtained results point to similar cycling performance of the additive mixture containing electrolyte formulation compared with the MBTFMP-containing cells, whereas the single PFPOEpI-containing cells displayed the best cycling performance in NMC111||graphite cells. With regard to the cathode electrolyte interphase (CEI), characterized and analyzed by means of scanning electron microscopy (SEM) and X-ray photoelectron spectroscopy (XPS), both the MBTFMP and the PFPOEpI functional additives decompose on the NMC111 surface in single-additive-containing electrolyte formulations. However, the thickness of the CEI formed in the additive mixture–containing electrolyte formulation is determined by the MBTFMP additive, whereas the PFPOEpI additive impacts a change in the composition of the CEI. Furthermore, the MBTFMP additive decomposes prior to the PFPOEpI and, therefore, dominates the cycling performance of NMC111||graphite cells containing functional additive mixture–based electrolyte. This systematic approach allows us to understand the synergistic impact of each functional additive in an electrolyte formulation containing an additive mixture and helps to identify the right additive combination for advanced electrolyte formulation as well as to elucidate whether the single-additive or the additive mixture approach is more effective for the development of advanced functional electrolytes for lithium-based cell chemistries.

Introduction

Lithium-based batteries (lithium ion batteries (LIB) and lithium metal batteries (LMB)) are still seen as the most promising electrochemical energy storage system candidates for automotive and portable applications [1–3].

However, due to the rapid increase and ever growing energy-related demands, novel, advanced electrode materials are highly demanded [4, 5], and the development of high-performing functional electrolytes is of equal crucial importance [6, 7]. Important aspects of this development, often considered as trivial and natural, comprise the study and understanding of the fundamental processes occurring at the different interfaces present in electrochemical systems, like for instance batteries by electrochemical or electroanalytical analyses [8–10]. Another aspect addresses utilizing insights on interface processes by substitution or addition of appropriate functional electrolyte components (conducting salt, solvent/co-solvent or functional/multifunctional additive) in order to tune the interface properties as needed [11–13]. In line with this strategy, the addition of at least two functional molecules with different structure to state-of-the-art (SOTA) electrolyte formulation represents an effective approach [6]. Different challenges related to the SOTA electrolyte comprising, for example, solid electrolyte interphase (SEI)
formation at the anode/electrolyte interface, cathode electrolyte interphase (CEI) formation at the cathode/electrolyte surface, safety (overcharge protection and flammability issues) can be addressed [14–19], while the bulk physical properties of the SOTA electrolyte, which consists of LiPF₆ as a conducting salt and a mixture of cyclic and linear organic carbonate–based solvents, remain mostly unaffected. The rapid increase in demand of high-energy density cells [4, 5] calls for increased cut-off cell voltage beyond 4.3 V [20, 21], which can be enabled by addition of CEI functional additives [22]. Based on general literature research and our previous work, we have identified phospholane- [22] and pyrazole-based [23] functional additives as promising candidates for high voltage application. In both cases, a beneficial impact on the overall cell performance was shown when using 0.5 wt.% of the phospholanes or 5.0 wt.% of the pyrazoles as a single additive [22, 23]. In case of the phospholane-based molecules, 2-(2,2,3,3,3-pentafluoropropoxy)-1,3,2-dioxaphospholane (PFPOEPi) (Scheme 1) was identified as the most effective functional additive among the candidates for the development of advanced high-voltage electrolytes, as it forms an effective CEI via ring opening polymerization at the cathode/electrolyte interface [22]. The pyrazole-based additive 1-methyl-3,5-bis(trifluoromethyl)-1H-pyrazole (MBTFMP) (Scheme 1) forms an effective CEI on the cathode surface as well, due to the formation of a complex with the transition metals (Ni, Co, Mn) present in the cathode [23]. To combine the advantages of the phospholane- and pyrazole-based additives with respect to high voltage application, PFPOEPi and MBTFMP were combined in an electrolyte formulation for LiNi₁/₃Mn₁/₃Co₁/₃O₂ (NMC111)||graphite cell chemistry. To identify the influence of each functional additive in the additive mixture containing electrolyte formulation, the obtained results were correlated to the single-additive electrolyte formulations. This approach allows us to determine the effect of the additive mixture vs. the single-additive approach on the overall cell performance.

### Experimental section

#### Synthesis of the phospholane and pyrazole functional additive

2-(2,2,3,3,3-Pentafluoropropoxy)-1,3,2-dioxaphospholane (PFPOEPi) and 1-methyl-3,5-bis(trifluoro-methyl)-1H-pyrazole (MBTFMP) molecules were synthesized according to our previous work [22, 23].

#### Electrolyte formulation and cell assembly

A 1-M LiPF₆ in ethylene carbonate (EC) and ethyl methyl carbonate (EMC) (1:1 by wt.) was used as a reference electrolyte (REF). The functional additive mixture containing electrolyte formulation contained 0.5 wt.% of PFPOEPi and 5 wt.% of MBTFMP in the REF. In the case of the single-additive-containing electrolyte formulations, either 0.5 wt.% PFPOEPi or 5 wt.% of MBTFMP were added to the REF. Graphite and NMC111 electrodes were provided by Customcells®. One layer of Separion® was used as a separator. Electrodes and separators were dried at 120 °C under vacuum for 12 h prior to use. Swagelok® T-cells (for cyclic voltammetry), 2032-type coin cells (for electrochemical impedance spectroscopy), and house-made pouch cells (galvanostatic cycling) were assembled in either a glove box (MBraun, H₂O and O₂ < 0.5 ppm) or in the dry room (dew point of at least −60 °C).

#### Electrochemical characterization

For the cyclic voltammetry (CV) measurements, conducted at a VSP potentiostat (BioLogic) with a scan rate of 20 μV·s⁻¹, Swagelok® T-cells were assembled in a 3-electrode setup [24]. As working electrode, either graphite or NMC111 was used, whereas Li metal served as both counter and reference electrodes. The cells were cycled for three cycles in the potential range from 3.0 to 4.6 V vs. Li|Li⁺ for NMC111 or from 0.025 to 1.5 V vs. Li|Li⁺ for graphite, starting from the open circuit potential (OCP) in both cases. The cells were charged/discharged with a C-rate of 0.1C and 0.333C in the voltage range of 3.0–4.5 V for three cycles, respectively. At the end of the charge step, a 3-h constant voltage step was included. After the formation step, the cells were cycled with 1.0C for 100 charge/discharge cycles in the voltage range from 3.0 to 4.5 V at room temperature (RT).

![Scheme 1 Chemical structures of considered high voltage functional electrolyte additives: 2-(2,2,3,3,3-pentafluoropropoxy)-1,3,2-dioxaphospholane (PFPOEPi) and 1-methyl-3,5-bis(trifluoro-methyl)-1H-pyrazole (MBTFMP)](image-url)
EIS, SEM, EDX, and XPS analyses

For postmortem studies, assembled pouch cells were galvanostatically cycled in the same voltage range (3.0 to 4.5 V) until either formation or 100 charge/discharge cycles. The cells were charged to 50% of the lithiation degree and, thereafter, disassembled. Cathode and anode were separated from each other and used in the measurements.

For the electrochemical impedance spectroscopy (EIS) measurements, three different coin cell setups (graphite||graphite, NMC111||graphite, NMC111||NMC111) were considered, adopting the experimental approach of Petibon et al. [25]. The measurements were conducted on a VSP potentiostat (BioLogic) in the frequency range of 1 kHz–1 MHz.

Prior to the scanning electron microscopy (SEM) measurements, the considered electrodes were washed with dimethyl carbonate (1 mL) in the glove box and, thereafter, were analyzed with an Auriga® CrossBeam workstation from Zeiss with an accelerating voltage of 3 kV and a working distance of 3 mm, using the InLens detector.

In the case of X-ray photoelectron spectroscopy (XPS), considered electrodes were transferred into the instrument (Axis Ultra DLD, Kratos, U.K.) and kept under reduced pressure (<10⁻⁸ mbar) for 12 h. Al Kα radiation with an energy of 1486.3 eV was used, and an angle of either 0° (cathode) or 45° (anode) of emission was applied. A pass energy of 160 eV at a 10-mA filament current and a 12-kV filament voltage source energy were used. The charging of the sample was compensated by the charge neutralizer. For the anodes, sputter depth profiling with a polyatomic ion gun (coronene as the ion source) was conducted. The sputter crater was 10 times larger than the selected measurement area, and the sputtering time was 60, 120, and 600 s. For each sample, two to three data points with lateral resolution of 700 μm × 300 μm were taken and arithmetically averaged. The obtained XPS spectra were fitted by using the CasaXPS software (Version 2.3.16 PR 1.6, Casa Software Ltd., U.K.) whereas the C 1 s C-H/CC peak was chosen as the internal standard for the binding energy (BE, 284.5 eV) calibration. The relative composition and thickness of the surface layer were calculated based on the work of Niehoff et al. [26, 27].

Results and discussion

For the functional additive mixture–containing electrolyte formulation with advanced performance, appropriate selection of single functional additives and their amount in the additive mixture is a priority [6]. In line with this, a concentration screening for the optimum concentration of each functional additive (phospholane and pyrazole) in the additive mixture containing electrolyte formulation was conducted (Fig. SI 1). The obtained results reveal that the combination of 0.5% PFPOEPi and 5% MBTFMP stands for the optimized additive mixture and was therefore used for further characterization and comparison to single-additive–containing analogues. To elucidate the impact of the functional additive mixture containing electrolyte formulation, galvanostatic cycling experiments were conducted and correlated to the corresponding single functional additive–containing electrolyte formulations. The obtained results are depicted in Fig. 1.

After the first charge/discharge cycle, the following order in respect to the Coulombic efficiency values was observed: REF + 0.5% PFPOEPi + 5% MBTFMP (76.3%) < REF + MBTFMP (77.3%) < REF (82.2) ≈ REF + 0.5% PFPOEPi. Here, the cell with the functional additive mixture-containing electrolyte formulation behaves similarly to the single MBTFMP-containing cell and no positive influence of the PFPOEPi functional additive is noticeable. Moreover, the cycling stability of the functional additive mixture-containing cells is similar to the single MBTFMP-containing

![Fig. 1 Coulombic efficiencies (top), specific discharge capacities (middle), and the capacity retentions (bottom) vs. cycle number of the four electrolyte formulations (1 M LiPF6 EC:EMC (1:1 by wt.) (REF), REF + 0.5% PFPOEPi, REF + 5% MBTFMP, and REF + 0.5% PFPOEPi + 5% MBTFMP) in NMC111||graphite cells at 20 °C with an upper cutoff voltage of 4.5 V](https://example.com/fig1.png)
cell, and only a slight improvement to the REF-containing cell is observed. The cycling stability of the single PFPOEPI-containing cell is much higher; however, the influence of the PFPOEPI functional additive in the additive mixture-containing electrolyte formulation seems to be negligible. To exclude the electrochemical impact of the PFPOEPI functional additive in the additive mixture–containing electrolyte formulation on the cycling performance, CV measurements of graphite||Li and NMC111||Li cells containing considered electrolyte formulations were conducted, as depicted in Fig. 2.

Figure 2a illustrates the cyclic voltammograms of the graphite||Li cells. The REF and the PFPOEPI-containing electrolytes display good compatibility with the graphite anode, whereas for the MBTFMP-containing cell, a decomposition with the peak maximum at 0.8 V vs. Li|Li⁺ is observed. These findings are in good agreement with the theoretically calculated reduction potentials, as they point out that MBTFMP decomposes prior to the other electrolyte components, whereas PFPOEPI is more reductively stable than EC and EMC [22, 23]. The CV results related to the functional additive mixture–containing electrolyte formulation, however, display no additional peak at 0.8 V vs. Li|Li⁺. This implies that PFPOEPI must somehow interfere with the decomposition reaction of MBTFMP, which leads to the suppression of the decomposition of the MBTFMP functional additive. On the oxidation side (Fig. 2b), the PFPOEPI-containing cell displays no additional decomposition peak compared with the REF-containing cell, whereas for the MBTFMP-containing and the additive mixture–containing analogues, an additional decomposition peak can be observed at the potential of 4.5 V vs. Li|Li⁺. This means that the oxidation potential of the additive mixture–containing electrolyte formulation is mainly driven by the MBTFMP molecule. When comparing these findings with the calculated oxidation potentials, a clear disagreement is observed. As the theoretically determined values indicate that PFPOEPI (oxidation potential of 5.09 V vs. Li|Li⁺ [22]) decomposes prior to MBTFMP (oxidation potential of 5.77 V vs. Li|Li⁺ [23]), the experimental results of the single-addition–containing electrolyte cells, however, reveal that the MBTFMP-containing electrolyte decomposes prior to the PFPOEPI-containing analogue. The theoretical approach of considering the oxidation potential reveals to be too simplistic as for the calculation of a single isolated molecule is used. Moreover, the calculated values suggest that PFPOEPI should have the main impact in CEI formation of the additive mixture–containing electrolyte formulation.

The obtained experimental results indicate that MBTFMP is the driving force of the decomposition reaction on the cathode, whereas PFPOEPI suppresses the decomposition of MBTFMP on the anode surface.

The electrochemical impact of PFPOEPI in the additive mixture–containing electrolyte formulation was confirmed by EIS measurements as illustrated in Fig. 3.

The pouch cells with considered electrolyte formulations were cycled for either formation state or for 100 charge/discharge cycles and thereafter, charged to 50% of the lithiation degree. To determine the influence of each electrode, symmetric graphite||graphite and NMC111||NMC111 cells as well as NMC111||graphite cells were assembled (see ref. [24]). The obtained results were interpreted according to Zhao et al. [28]. Here, the x-axis interception in the Nyquist plot represents the electrolyte resistance (Rₑₑ), the first semicircle the interface/interphase resistance (Rₑₑₑₑ, Rₑₑₑₑ, Rₑₑₑₑ) at the considered electrodes, and the second semicircle the charge transfer resistance (Rₐₑ). As the obtained cycling results reveal that the impact of the PFPOEPI additive in the additive mixture–containing electrolyte formulation is negligible; however, the obtained EIS results (Fig. 3) display an influence of the PFPOEPI additive in the additive mixture–containing electrolyte formulation. First of all, the impedance growth during prolonged cycling is driven by the cathode side for all considered electrolyte formulations as the overall impedance in the NMC111||NMC111 cells increases significantly compared with the graphite||graphite cells. When looking more closely into the NMC111||NMC111 cell results, it can be seen that the impedance of Rₑₑₑₑ is the lowest in the case of the additive mixture–containing electrolyte formulation, while the Rₑₑₑₑ for the REF- and the MBTFMP-containing cell is similar. The PFPOEPI-containing cell displays the highest Rₑₑₑₑ. In the case of the charge transfer resistance, after
formation, the REF-containing cells have the lowest resistance followed by the additive mixture counterpart, the MBTFMP- and the PFPOEPi-containing electrolyte formulation-based cells. Herein, no influence of either functional additive can be distinguished for the additive mixture–containing electrolyte formulation. However, after 100 charge/discharge cycles, the influence of each additive in the additive mixture–containing electrolyte formulation becomes observable. \( R_{\text{CEI}} \) does not increase during prolonged cycling for the additive mixture–containing electrolyte, which can be clearly assigned to the MBTFMP-containing cells, as it also displays a constant \( R_{\text{CEI}} \) during prolonged cycling, whereas \( R_{\text{CEI}} \) for the PFPOEPi-containing cells increases during prolonged cycling. In the case of \( R_{\text{ct}} \), the main influence for the increasing second semicircle for the additive mixture–containing electrolyte formulation originates from both functional additives. For

![EIS spectra of the four electrolyte formulations](image-url)

**Fig. 3** EIS spectra of the four electrolyte formulations (1 M LiPF₆, EC:EMC (1:1 by wt.) (REF), REF + 0.5% PFPOEPi, REF + 5% MBTFMP, and REF + 0.5% PFPOEPi + 5% MBTFMP) in graphite||graphite (top), NMC111||graphite (middle), and NMC111||NMC111 (bottom) cells after formation (left column) as well as after 100 charge/discharge cycles (right column) at 20 °C
PFPOE–Pi-containing cells, $R_{\text{ct}}$ does not increase, whereas a drastic increase for the MBTFMP-containing cells is noticeable during prolonged cycling. The synergistic effect of the phospholane and pyrazole functional additives in the additive mixture–containing electrolyte formulation leads to an $R_{\text{ct}}$ which lies between the values of the single PFPOE–Pi- and MBTFMP-containing cells. With this, we can conclude that the PFPOE–Pi additive has a positive influence on the resistance of the additive mixture–containing electrolyte formulation; however, as the cycling results reveal, an impact on the decreased $R_{\text{ct}}$ is not observable. Moreover, the influence of the MBTFMP additive on the constant $R_{\text{CEI}}$ seems to be more important. In the case of the graphite|graphite cells, the impact of the PFPOE–Pi additive in the additive mixture–containing electrolyte formulation is more pronounced. After formation, the following order of the overall impedance can be observed: $\text{REF} + 5\% \text{ MBTFMP} < \text{REF} + 0.5\% \text{ PFPOE–Pi} < \text{REF} + 5\% \text{ PFPOE–Pi}$. After 100 charge/discharge cycles, the overall impedance increases and is the same as for the PFPOE–Pi-containing cells after 100 charge/discharge cycles. This impact of PFPOE–Pi additive in the additive mixture–containing electrolyte formulation displays only a minor impact, as the significant increase in the NMC111|graphite cells is driven by the cathode surface and is mainly responsible for the cycling performances displayed in Fig. 1.

To elucidate the impact of MBTFMP and PFPOE–Pi on the CEI formation, postmortem analysis was performed by means of SEM and XPS analyses of the NMC111 electrodes. The obtained SEM results are depicted in Fig. 4.

The obtained SEM images of the NMC111 electrodes with REF after formation show presence of decomposition products on the electrode surface, whereas after 100 charge/discharge cycles, the electrode particles are completely covered by a layer of decomposition products. In the case of the phospholane-containing cells after formation, additional spherically shaped decomposition products can be observed. For the PFPOE–Pi-containing cell, the electrode surface after 100 charge/discharge cycles appears to be unaffected compared with the cathode surface after formation. For the MBTFMP-containing cells, after formation, small decomposition spots can be observed, and their amount does not increase during prolonged cycling. In the case of the additive mixture–containing electrolyte formulation–based cells, the surface of the NMC111 electrode is covered by decomposition products after formation. After 100 charge/discharge cycles, decomposition products are still observable for the additive mixture–containing electrolyte formulation and remain unaffected during prolonged cycling. As the same behavior could be observed for the single-additive–containing cells, an influence of individual additives on CEI formation could not be distinguished.

To get a deeper understanding of the impact of each functional additive on CEI formation and composition, XPS analysis was performed. The corresponding XPS results of the composition and the thickness of the CEI of the NMC111 electrodes determined by the mathematical approach developed by Niehoff et al. [26, 27] are depicted in Fig. 5.

After formation, a significant difference between the formed CEI on the NMC111 surface, driven by the different electrolyte formulations, can be observed (REF + 0.5% PFPOE–Pi < REF + 5% MBTFMP = REF + 0.5% PFPOE–Pi + 5% MBTFMP). During 100 charge/discharge cycles, the CEI thickness of the MBTFMP-containing cell remains unaffected, whereas the thickness of CEI of the PFPOE–Pi-containing cells significantly increases. In the case of the additive mixture–containing electrolyte formulation, only a slight increase in the CEI thickness from formation to 100 charge/discharge cycles is observed. This results in CEI thicknesses of the following order: $\text{REF} + 0.5\% \text{ PFPOE–Pi} < \text{REF} + 5\% \text{ MBTFMP} < \text{REF} + 0.5\% \text{ PFPOE–Pi} + 5\% \text{ MBTFMP}$, indicating that the thickness of the CEI in the additive mixture–containing electrolyte formulation is mainly determined by the MBTFMP additive. When looking into the composition of the CEI, a significant influence of PFPOE–Pi additive in the additive mixture–containing cells can be observed. Firstly, a smaller content of LiF for the additive mixture–containing electrolyte formulation after formation as well as after 100 charge/discharge cycles compared with the MBTFMP-containing cells is noticeable. The decreased LiF content is probably due to the presence of the PFPOE–Pi additive, as the cells containing only PFPOE–Pi display a very low content of LiF in the CEI. Secondly, in the case of the MBTFMP and the additive mixture electrolyte–containing cells a nitrogen peak can be observed. In both cases, this means that the MBTFMP additive decomposes on the NMC111 electrode surface. Due to the fact that the nitrogen concentration does not change during prolonged cycling in both cases it can be assumed that the decomposition of MBTFMP functional additive occurs during formation. In addition, the thickness is similar in both cases and indicates that the same amount of MBTFMP additive decomposes on the NMC111 surface. The CEI of the MBTFMP-containing cells displays a CF$_3$ thickness of 0.04 nm after 100 charge/discharge cycles, whereas the additive mixture–containing composition products
electrolyte formulation has a thickness of 0.08 nm after 100 charge/discharge cycles. In addition, an increase of the CF₃ content in the CEI layer from formation to 100 charge/discharge cycles is observable while the nitrogen content is constant. This means that the increase in the CF₃ content does not originate from the MBTFMP molecule. Due to the fact that the PFPOEPi molecule possesses a CF₃ group in the structure, the increase of the CF₃ content in the CEI layer in the additive mixture–containing electrolyte formulation can be clearly assigned to the decomposition of the PFPOEPi additive on the NMC111 surface. Furthermore, an increase in the C–O content in the CEI layer from 0.69 nm for the MBTFMP-containing cell to 0.88 nm for the additive mixture–containing electrolyte formulation after formation and 100 charge/discharge cycles.

As the CV results of the graphite||Li cells (Fig. 2a) and the EIS results (Fig. 3) of the graphite||graphite cells reveal, the PFPOEPi additive displays a more significant impact on the anode side than on the cathode side in the additive mixture–containing electrolyte formulation. For this reason, the SEI on the graphite anode was analyzed by means of SEM, and the obtained results are depicted in Fig. 6. The SEM images of the graphite electrodes reveal decomposition products after formation cycles in the case of REF, whereas after 100 charge/discharge cycles, the surface is completely covered with decomposition products. When comparing the PFPOEPi-containing cells with the REF-containing cells after formation, more decomposition products for PFPOEPi-containing cells are observed, whereas after 100 charge/discharge cycles, the surface of the graphite electrode cycled with the REF displays more decomposition products compared with the PFPOEPi-containing cells. In the case of the MBTFMP-containing cells, electrolyte decomposition products are observed; however, their amount does not increase during prolonged cycling and, therefore, less decomposition products are present on the graphite surface compared with the cells containing REF after 100 charge/discharge cycles. In addition, the amount of the decomposition products is similar to the PFPOEPi-containing cells. For the additive mixture electrolyte formulation–containing cells, decomposition products are observable in a low amount after formation. Their amount however, increases during prolonged cycling. The decomposition amount is less than with the REF-containing cells, but more than in the PFPOEPi- and MBTFMP-containing cells.

To evaluate the impact of each additive in the additive mixture containing electrolyte formulation on the thickness performance of the cell with additive mixture–containing electrolyte formulation is the same as for the MBTFMP-containing cell, it can be concluded that the impact of the PFPOEPi additive, which changes the composition of the CEI, is negligible. Furthermore, the CEI thickness, induced by the MBTFMP additive, is the main driving force for the cycling performance of the additive mixture containing cells. In line with this conclusion, it can be deduced that the MBTFMP additive has the main impact on the CEI formation and dynamics. The positive effects of the decomposition of the PFPOEPi additive on the NMC111 electrode surface are suppressed, which is the reason why the addition of the PFPOEPi additive to the MBTFMP additive–containing electrolyte formulation has no positive impact on cycling performance.

As the CV results of the graphite||Li cells (Fig. 2a) and the EIS results (Fig. 3) of the graphite||graphite cells reveal, the PFPOEPi additive displays a more significant impact on the anode side than on the cathode side in the additive mixture–containing electrolyte formulation. For this reason, the SEI on the graphite anode was analyzed by means of SEM, and the obtained results are depicted in Fig. 6. The SEM images of the graphite electrodes reveal decomposition products after formation cycles in the case of REF, whereas after 100 charge/discharge cycles, the surface is completely covered with decomposition products. When comparing the PFPOEPi-containing cells with the REF-containing cells after formation, more decomposition products for PFPOEPi-containing cells are observed, whereas after 100 charge/discharge cycles, the surface of the graphite electrode cycled with the REF displays more decomposition products compared with the PFPOEPi-containing cells. In the case of the MBTFMP-containing cells, electrolyte decomposition products are observed; however, their amount does not increase during prolonged cycling and, therefore, less decomposition products are present on the graphite surface compared with the cells containing REF after 100 charge/discharge cycles. In addition, the amount of the decomposition products is similar to the PFPOEPi-containing cells. For the additive mixture electrolyte formulation–containing cells, decomposition products are observable in a low amount after formation. Their amount however, increases during prolonged cycling. The decomposition amount is less than with the REF-containing cells, but more than in the PFPOEPi- and MBTFMP-containing cells.

To evaluate the impact of each additive in the additive mixture containing electrolyte formulation on the thickness
and composition of the formed SEI, XPS analysis was carried out and the obtained results are illustrated in Fig. 7.

The obtained SEI thickness values considering the additive mixture–containing electrolyte formulation reveal an impact of both functional additives on SEI formation and dynamics. The SEI thickness of anodes from the additive mixture–containing electrolyte formulation is between those of anodes of the single-additive–containing cells after formation. After 100 charge/discharge cycles, the SEI thickness of the all three additive–containing cells is similar after 100 charge/discharge cycles. Moreover, the composition of the additive mixture–containing electrolyte formulation in respect to the organic and inorganic parts is similar to the MBTFMP-containing cell after 100 charge/discharge cycles. Herein, the increase in the SEI thickness is not as pronounced as for the PFPOEPi-containing cells, which is influenced by the PFPOEPi additive. Furthermore, the presence of PFPOEPi additive impacts the increase of the organic content of the SEI after formation compared with the MBTFMP cells. However, the organic layer is thinner than for the PFPOEPi-containing cells. The influence of additives is indirect as the N 1s and C 1 s core spectra (Fig. 7b, c) reveal no decomposition of the PFPOEPi or the MBTFMP additive on the electrode surface. With this in mind, it can be concluded that the impact of the functional

Fig. 7 a Determined thicknesses of the organic and inorganic parts of the SEI, b N 1 s core spectra, and c C 1 s core spectra of the considered electrolyte formulations (1 M LiPF 6 EC:EMC (1:1 by wt.) (REF), REF + 0.5% PFPOEPi, REF + 5% MBTFMP, and REF + 0.5% PFPOEPi + 5% MBTFMP) after formation and 100 charge/discharge cycles.
additives on improved cycling performance is determined by their decomposition and formation of the CEI at the cathode surface. Furthermore, when looking into the combination of the additives PFPOEPi and MBTFMP, an influence on both electrodes is observable. The impact of the additive PFPOEPi is low on both interphases, whereas the decomposition of the MBTFMP additive dominates at the cathode surfaces. As the EIS results reveal, the cathode side in the battery cell is mainly responsible for the cycling performance and, therefore, the additive MBTFMP mainly determines the cycling performance of the additive mixture containing electrolyte formulation.

**Conclusion**

A functional additive mixture–containing electrolyte formulation consisting of a phospholane- and a pyrazole-based additive was systematically investigated for high voltage application in LIBs and correlated to the single-additive–containing electrolyte formulations to determine the influence of each additive in the additive mixture–containing electrolyte and to evaluate whether the single-additive or the additive mixture approach is more effective for the development of advanced LIBs. The obtained results revealed similar cycling performance of the additive mixture–containing electrolyte formulation to the MBTFMP-containing cells. The reason behind this observation is related to formation of the respective interphases at anode and cathode. In the case of SEI formation, the synergistic effect of the two functional additives was well observed after the formation step and after 100 charge/discharge cycles. However, the considered additives do not directly decompose at the graphite surface, and SEI formation is only indirectly influenced. EIS analysis points out that the MBTFMP additive impacts the formation of an effective interphase (also observable in the XPS results), whereas the PFPOEPi additive is responsible for a decreased \( R_{\text{ct}} \) in the additive mixture–containing cells. However, as the cycling performance of the additive mixture–containing cells is similar to the MBTFMP-containing cells, the resistance originating from the CEI is apparently more relevant for the cycling performance than the reduced \( R_{\text{ct}} \). Moreover, the obtained XPS data reveal that both functional additives in the additive mixture–containing electrolyte formulation decompose at the NMC111 surface. The MBTFMP additive mainly determines the thickness of the CEI, but still the PFPOEPi additive influences the CEI composition. By correlating these findings with the galvanostatic cycling results, it can be concluded that changes in CEI composition induced by the PFPOEPi additive have no impact on the overall cycling performance.

In addition, this work shows that the polymerization reaction of the phospholane molecule via ring opening is more effective for the formation of an effective CEI as the complex formation of the pyrazole-based molecule with the transition metals present in the NMC111 surface. This systematical approach reveals that the selection of the right additive mixture, the optimum amount of each functional additive, and the understanding of their effectiveness and performance are essential for the development of advanced electrolyte formulations for lithium-based batteries. This work is also a most illustrative example for the power of electrochemical and analytical techniques [9, 10] to identify the “best” material or material combination for a specific electrochemical application.

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