Insertion cathode materials based on borate compounds
Florian Strauss

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Insertion cathode materials based on borate compounds

by Florian Alexander STRAUSS

PhD thesis in Chemistry

Directed by Robert Dominko and Jean-Marie Tarascon

To be presented and defended in public on November 25th, 2016

In front of the jury:

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Matériaux d’insertion de cathode d’insertion à base des borate

Par Florian Alexander STRAUSS

Thèse de doctorate de Chimie des Matériaux

Dirigée par Robert Dominko and Jean-Marie Tarascon

Présentée et soutenue publiquement le 25 November 2016

Devant un jury composée de:

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Abstract

The increased need of energy storage via the development of Li- and Na-ion batteries requires a continuous search for new positive electrode materials having higher energy density while being safe and sustainable. For this purpose we explored borate based compounds capable of reacting with Li/Na-ions in a reversible way either through intercalation/deintercalation or conversion reactions. During this survey we focused on identifying candidates possessing a polyborate anion (B$_x$O$_y$ with x > 1), that are expected to show elevated redox potentials compared to BO$_3$ based materials. Using Li$_6$CuB$_4$O$_{10}$ as a model compound we showed the possibility to achieve redox potentials of 4.2 and 3.9 V vs. Li$^+$/Li$^0$ for the α- and β-polymorphs, respectively. This redox activity was rationalized through complementary EPR spectroscopy and DFT calculations. We further reveal the structural and synthetic relation between the two polymorphs and show a surprisingly high ionic conductivity of 1.4 mS·cm$^{-1}$ at 500°C for α-Li$_6$CuB$_4$O$_{10}$, related to a structural transition. Moreover we were able to prepare two new sodium transition metal pentaborates Na$_3$MB$_5$O$_{10}$ (M = Fe, Co) possessing an open structure feasible for Na$^+$ migration. For M = Fe we observed a reversible Na intercalation at an average potential of 2.5 V vs. Na$^+$/Na$^0$, opposed to Na$_3$CoB$_5$O$_{10}$ which turned out to be electrochemical inactive. Finally deviating from classical insertion/deinsertion compounds, we studied the electrochemical driven reaction mechanism of a bismuth oxyborate Bi$_4$B$_2$O$_9$ versus Li through electrochemical measurements combined with XRD and TEM investigations. Remarkably, we found that it is possible to reversibly cycle this material between 1.7 and 3.5 V with an average redox potential of 2.3 V vs. Li$^+$/Li$^0$ with only 5wt% carbon additive and a small polarization ~300 mV. Owing to the complexity of 3d-metal borate chemistry encountered through this PhD, the chances of having a borate based positive electrode for next generation Li-ion batteries is rather slim.

Keywords: Li/Na-ion batteries, cathode materials, borates, ionic conductivity
Povzetek

Povečana potreba po shranjevanju energije in s tem razvoj Li- in Na-ionskih akumulatorjev zahteva kontinuirno iskanje novih katodnih elektrodnih materialov, ki imajo višjo energijsko gostoto in so obenem varne za uporabo ter iz široko dostopnih elementov. V ta namen smo preučevali spojine na osnovi boratov, ki lahko reverzibilno izmenjujejo Li ali Na ione s klasično vgradnjo/izgradnjo v/iz strukturo ali pa z konverzijsko elektrokemijsko reakcijo. Raziskave smo usmerili v identifikacijo materialov z poliboratnim anionom (BₓOᵧ, kjer je x > 1), od katerih se pričakuje višji redoks potencial glede na spojine, ki imajo BO₃ skupino v strukturi. Z uporabo Li₆CuB₄O₁₀ kot modelne spojine smo pokazali, da je mogoče doseči redoks potencial 4.2 in 3.9 V glede na Li⁺/Li⁰ za α- in β-polimorff. Redoks aktivnost bakra smo dodatno dokazali z EPR spektroskopijo in DFT izračuni. Nadalje smo preučevali odvisnost sinteznih pogojev in strukture dveh polimorfov, kjer smo pokazali, da ima α-Li₆CuB₄O₁₀ pri 500°C prevodnost 1.4 mS·cm⁻¹. Dodatno smo pripravili dve novi natrijivi spojini na osnovi prehodne kovine in pentaborata Na₃MB₅O₁₀ (M = Fe, Co). Za M=Fe smo izmerili reverzibilno insercijo natrija pri napetosti 2.5 V glede na Na⁺/Na⁰, medtem ko se je izkazalo, da je Na₃CoB₅O₁₀ elektrokemijsko neaktiven. Z iskanjem novih aktivnih poliboratnih spojin smo zapustili klasične materiale v katere se alkalne kovine vgrajujejo in smo se odločili za študij elektrokemijskega mehanizma konverzije Bi₄B₂O₉ glede na Li⁺/Li⁰ v povezavi z XRD in TEM meritvami. Spojino je mogoče reverzibilno galvanostatsko ciklati med 1.7 in 3.5V s povprečno napetostjo 2.3 V glede na Li⁺/Li⁰ pri čemer smo uporabili samo 5ut% ogljika, obenem pa smo dobili zelo majhno polarizacijo ~300 mV. Glede na kompleksnost boratne kemije z 3d prehodnimi kovinami s katerimi smo se ukvarjali tekom tega doktorata, so šanse, da bi imeli boratne spojine, kot katode za Li-ionske akumulatorje za naslednjo generacijo zelo majhne.

Ključne besede: Li / Na-ionske baterije, katodni materiali, boratov, ionsko prevodnost
Le besoin accru de stockage d'énergie via le développement de Li- et batteries Na-ion nécessite une recherche continue de nouveaux matériaux d'électrodes positives ayant une densité d'énergie plus élevée tout en étant sûre et durable. A cet effet, nous avons exploré les composés de base de borate capables de réagir avec Li / Na-ions de manière réversible soit par intercalation / désintercalation ou de conversion des réactions. Au cours de cette enquête, nous avons mis l'accent sur l'identification des candidats possédant un anion polyborate (BₓOᵧ avec x > 1), qui devraient montrer des potentiels redox élevés par rapport aux matériaux à base BO₃. Li₆CuB₄O₁₀ utilisant comme composé modèle, nous avons montré la possibilité d'obtenir des potentiels d'oxydo-réduction de 4.2 et 3.9 V vs. Li⁺/Li₀ les a et β polymorphes, respectivement. Cette activité redox a été rationalisée par spectroscopie EPR complémentaire et calculs DFT. Nous révélons en outre la relation structurelle et synthétique entre les deux polymorphes et montrons une conductivité ionique étonnamment élevé de 1.4 mS·cm⁻¹ à 500°C pour les α-Li₆CuB₄O₁₀, liée à une transition structurelle. De plus, nous avons pu préparer deux nouveaux pentaborates transition de sodium métalliques Na₃MB₅O₁₀ (M = Fe, Co) possédant une structure ouverte possible pour la migration Na⁺. M = Fe, nous avons observé une intercalation Na réversible à un potentiel moyen de 2.5 V par rapport à Na⁺/Na₀, opposé à Na₃CoB₅O₁₀ qui se révéla être inactif électrochimique. Enfin déviant de composés d'insertion / désinsertion de classiques, nous avons étudié le mécanisme d'un oxyborate de bismuth par rapport Bi₄B₂O₉ Li réaction électrochimique entraînée par des mesures électrochimiques combinées à XRD et TEM investigations. Fait étonnant, nous avons trouvé qu'il est possible de faire défiler ce matériau réversible entre 1.7 et 3.5 V avec un potentiel d'oxydo-réduction moyenne de 2.3 V vs. Li⁺/Li₀ avec seulement 5 wt% d'additif de carbone et une faible polarisation ~300 mV. En raison de la complexité de la chimie de borate 3d-métal rencontré par le biais de cette thèse, les chances d'avoir une électrode positive à base de borate pour les batteries Li/Na-ion de nouvelle génération est plutôt mince.

Mots clés : batteries à ions Li/Na, matériaux de cathode, borates, conductivité ionique
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1 Introduction

The energy demand of our modern society is continuously growing, however the global consumed energy is nowadays mainly produced through the combustion of fossil resources, whereas only a small proportion of 8% of the total consumed energy is supplied through renewables (Figure 1a).

It is widely accepted that geopolitical issues and global greenhouse gas emissions (in particular CO$_2$) originating from the utilization of energy production by burning fossil fuels, have a huge impact on our modern society, with future consequences which cannot yet been entirely predicted. More than 50% of the global CO$_2$ emitted, is generated by the production of electricity and heat as well as powering our global transportation (Figure 1b).\textsuperscript{1,2}

\textbf{Figure 1:} (a) Total primary global energy consumption from 1980 to 2040 by fuel (Mtoe).\textsuperscript{1} (b) World CO$_2$ emission by sector in 2013.\textsuperscript{2}

To circumvent the global dependence on fossil fuels and minimize the global warming, an increased interest has been aroused for the development of renewable energy supply for electricity generation and transportation. Over the last 20 years global renewable energy production has been strongly raised (Figure 2), not only driven by their sustainability and positive effect on the environment but also due to the continuous raise of the prices for fossil fuels (although temporary fluctuations can occur) since it is supposed that their maximum production is reached.
To change the energy supply from fossil fuels to renewables, one major issue has to be solved in order to fully implement and benefit from them. This major issue of renewable energy production is their non-continuously power supply, since the amount of energy harvested is a function of time, place, and weather conditions. In order to balance the energy production from renewables with the consumption demands, smart solutions are needed. Obviously one can think about storing excess energy in hydroelectric power stations, unfortunately their availability is limited to the local landscape, requires large areas and comes along with low roundtrip efficiencies.

Several energy storage technologies like the afore mentioned pumped hydro, compressed air, flywheels and electrochemical energy storage devices (batteries, capacitors) have been proposed in the past, however it is important to note that none of these technologies is able to manage all kinds of different demands, if comparing for instance transportation versus grid scale energy storage, but batteries may play an important role for electric powered transportation and grid support (Figure 3).
Figure 3: Comparison of discharge time and power rating for various electrochemical energy storage technologies.\(^5\)

Regarding energy supply for transportation, today’s available rechargeable batteries are still not competitive with gasoline in terms of range and costs, hence the percentage of electricity used for transportation is negligible and the prediction of the increase are poor as well (Figure 4).\(^6\) Fuel cells are considered as a serious candidate to power electric vehicles, but only in 2015 one of the first commercial fuel cell cars was unveiled by Toyota, despite 200 years of research in this field.\(^7,8\) By contrast, a broad commercialization of affordable Li-ion battery powered EVs, only about 20 years after the first commercialization of Li-ion battery itself is expected within the next 5 years.\(^9,10\)
In order to make the rechargeable battery technology serious contender for electrical energy storage (as power supply in EVs or grid support), new chemistries and battery concepts need to be explored. For example if made of earth abundant and cheap materials with high energy density, the same materials could be used for small as well as large scale applications lowering the overall production costs and coming along with a lower environmental impact.\textsuperscript{11,12}

1.1 Batteries for electrochemical energy storage

Batteries as provider of electricity have been known since the invention of the first battery, the so called ‘voltaic pile’ by the Italian physicist Alessandro Volta in 1799. This device consisted of an alternating stack of copper and zinc plates separated by the electrolyte which can be either diluted sulfuric acid or just simple a saltwater solution. This device was the first being able to supply a continuous flow of electricity, opening a variety of pathways
for new discoveries in natural science (water electrolysis, discovery of new elements by molten salt electrolysis, etc.)\textsuperscript{13}

Although this first battery had a rather short lifetime and could not be recharged, also named primary battery, it already was based on the same concept like all other batteries to date, whether primary or secondary.

In brief all batteries consist of two electrodes, a negative and a positive one (often called anode and cathode respectively, Figure 5) which are separated by an ion conductive but electrical insulating medium, the so called electrolyte. The two electrodes have a different chemical potential depending on their redox couple. When the electrodes are connected through an external circuit, the electrochemical reactions proceed at both sides, causing a flow of electrons (current) from the more negative to the more positive side. At the same time electro-neutrality is ensured by ions travelling the opposite direction through the electrolyte. If one of the redox reactions is finished, the current stops, and the battery is completely discharged. If this process can be reversed by applying an external voltage, the battery can be recharged and is called secondary battery.

\[ V_{OC} = \frac{\mu_A - \mu_C}{e} \]  

From equation 1, the open circuit potential of an electrochemical cell \( V_{OC} \) is defined by the difference between the electrochemical potentials \( \mu_A \) and \( \mu_C \) of the anode and cathode respectively, divided by the elementary charge \( e \). The voltage is limited by the energy gap between the highest occupied molecular orbital HOMO and the lowest unoccupied molecular orbital LUMO of the electrolyte (voltage window).\textsuperscript{14,15}
In general the two electrodes can be chosen from any kind of materials possessing a different chemical potential. However for practical considerations the chosen redox couple needs to fulfill several criteria like specific mass of the materials, their costs, safety and durability. The amount of electrical energy stored in batteries can be expressed in terms of unit per weight (Wh kg\(^{-1}\)) or unit per volume (Wh L\(^{-1}\)) which are linked to the battery’s voltage (V) and gravimetric capacity (Ah kg\(^{-1}\)). The last two terms are directly correlated to the battery’s chemistry and can therefore be tuned depending on requirements. Among various explored technologies, the most established and common rechargeable systems are lead-acid, nickel-cadmium, nickel-metal hydride batteries and lithium-ion batteries. A summary about their characteristics is given in Table 1.\(^{15}\)

<table>
<thead>
<tr>
<th>Battery type</th>
<th>Lead-acid</th>
<th>Nickel-cadmium</th>
<th>Nickel-metal hydride</th>
<th>Li-ion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Voltage (V)</td>
<td>2.1</td>
<td>1.3</td>
<td>1.3</td>
<td>4.1</td>
</tr>
<tr>
<td>Practical energy density</td>
<td>70 Wh/L</td>
<td>100 Wh/L</td>
<td>240 Wh/L</td>
<td>400 Wh/L</td>
</tr>
<tr>
<td>Power density (W/kg)</td>
<td>35 Wh/kg</td>
<td>35 Wh/kg</td>
<td>75 Wh/kg</td>
<td>150 Wh/kg</td>
</tr>
<tr>
<td>Cycles</td>
<td>500-800</td>
<td>2000</td>
<td>500-1000</td>
<td>400-1200</td>
</tr>
</tbody>
</table>

Table 1: Characteristic of commonly used rechargeable batteries.
Among various existing technologies (Figure. 6) Li-ion batteries have encountered the global market as the most promising one, due to their favorable energy densities and flexibility in terms of design and chemistries.\textsuperscript{14,16} A brief overview about their historical development with emphasis on different cathode materials is given in the next and following sections.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{comparison.png}
\caption{Comparison of different battery technologies in terms of their gravimetric and volumetric energy density.\textsuperscript{5}}
\end{figure}

\section*{1.2 Lithium- and sodium ion batteries}

\subsection*{1.2.1 Lithium ion batteries}

One of the main motivations employing lithium in rechargeable batteries is the fact that lithium represents the lightest metal (molar weight 6.94 g mol\textsuperscript{-1}, gravimetric density $\rho = 0.53$ g cm\textsuperscript{-3}) and owns the highest reduction potential among all alkali metals (-3.04 vs. SHE), thus potentially leading to high energy densities. Due to the incompatibility of alkali metals with aqueous electrolytes and the elevated voltage which would lead to water electrolysis, organic electrolytes have to be employed in lithium secondary batteries. In the 1970s several inorganic compounds were shown to reversibly intercalate alkali metals, especially transition metal chalcogenides.\textsuperscript{17,18,19} In 1972 Exxon demonstrated a first non-aqueous rechargeable lithium cell based on TiS\textsubscript{2} as a positive and lithium metal as the negative electrode.\textsuperscript{20} Unfortunately this concept soon encountered two main issues, first the combination of lithium
metal with a liquid electrolyte and second the dendritic lithium growth which occurred as a consequence of lithium deposition during subsequent charge-discharge cycles, leading to explosion hazards. Following this first demonstration, researchers investigated several other chalcogenides as positive electrode materials and started soon considering transition metal oxide materials showing higher capacities and redox potentials. Based on this pathway, Goodenough and coworkers proposed the family of lithium transition metal oxides Li$_x$MO$_2$ (M = Co, Ni, Mn) materials which are still used in today’s lithium- and sodium-ion batteries.\textsuperscript{21,22}

To circumvent the safety issues coming along with the use of metallic lithium as an anode, first the use of a lithium-aluminium alloy instead of pure lithium was proposed and solved the dendrite issue, but the electrodes survived only a certain number of cycles due to the large volume change during charge/discharge.\textsuperscript{23} At the early 1990s a second approach was proposed, substituting the metallic lithium by another insertion material.\textsuperscript{24,25} This concept has led to the development of the so called lithium-ion or rocking-chair technology. Because lithium does not occur in a metallic state anymore in this type of cells, the dendrite issue is solved and Li-ion cells are therefore in principle much safer than Li-metal cells. To compensate for the overall voltage loss because of the increased redox potential of the anode, high voltage cathode materials are needed. According to these new demands for the cathode, the focus of research shifted from layered- transition metal sulfides to layered-, or three-dimensional transition metal oxides.\textsuperscript{22} However it took almost ten years to commercialize the Li-ion battery. The delay was attributed to find suitable electrolytes, as well as anode materials. Only the discovery on Li-intercalation in carbonaceous materials,\textsuperscript{26} based on earlier research,\textsuperscript{27} led to the design of the first LiCoO$_2$/graphite cell (Figure 7) commercialized in 1991 by \textit{SONY}.\textsuperscript{28} This type of cell had a potential exceeding 3.6 V and gravimetric energy densities around 120-150 Wh/kg, a schematic view is drawn in Figure 7, which concept is still found in most of the today’s Li-ion cells.
1.2.2 Sodium ion batteries

Thanks to the great success of Li-ion batteries, Na-ion batteries are seriously considered for the post-lithium batteries, although the Na-ion concept is not new and has been studied along with the Li-ion chemistry. Similar chemical approaches, including synthetic strategies and characterization methods can be adopted from the field of Li-ion batteries. Despite a lower potential (~0.3 V) and energy density of Na-ion versus Li-ion batteries, the utilization of sodium may push the price per stored energy even further down, due to the natural earth abundance, which is more than 1000 times higher for sodium compared to lithium. Components and electrochemical energy storage mechanism are basically the same like for Li-ion batteries, with the main difference is that hard carbon has to be used instead of graphite as negative electrode material (Figure 9).

---

1 Hard carbon is a disordered carbon where the graphene sheets are not parallely aligned as in graphite.
1.3 Cathode materials for lithium- and sodium –ion batteries

In the following section an overview over the most studied class of cathode materials for lithium- and sodium ion-batteries is given. In particular different examples of positive electrode materials, namely transition metal oxides and important polyanionic materials are described in detail and finally conversion type materials are briefly reviewed.

Contrary to other battery technologies, lithium- and sodium-ion batteries can rely on a variety of different electrode materials, for both the positive and negative side. Obviously as shown in Figure 10, the main limiting factor for energy density is in both cases the lack of a high capacity positive electrode material. Therefore most of the work during the last decades has been focused on the search for new positive cathode materials displaying higher capacity and voltage, mainly for lithium batteries, however emerging more and more for sodium ion batteries.
Figure 9: Overview of different positive and negative electrode materials for (a) Li-ion and (b) Na-ion batteries, with respect to their redox potential and theoretical capacity. \(^{11,16,34}\)

In general an ideal positive electrode materials would possess a relative high redox potential (within the electrochemical stability of currently used electrolytes, a potential greater than 4.8 V vs. Li\(^+\)/Li\(^0\)) in combination with a high specific capacity. The specific capacity depends proportional on the number of exchanged Li-ions, and is lower, the higher the molar weight of the material. These two main requirements are by far not the only ones which need to be taken into account when searching for a new electrode material. It also should react with lithium/ sodium in a highly reversible way, preferable an insertion/ deinsertion mechanism so the crystal structure is maintained during charge discharge cycling leading to a longer lifetime of the material. Furthermore the electrode materials should be capable in conducting both, electrons and ions, since these two properties are the main limiting kinetic factors for the redox reaction and determining thus the rate capability of the material.

Besides the afore-mentioned properties, non-toxicity of the used elements, high thermal stability and lithium in the pristine material to avoid the use of lithium metal in the full cell or pre-lithiation step before the cell assembly are also important factors especially in terms of practical application.

1.3.1 Lithium- and sodium transition metal oxides

As mentioned before, following the studies of lithium intercalation in sulfides, researchers turned into the study of transition metal oxides soon, since they were expected to
present higher redox potentials versus lithium. Indeed replacing sulfur by the more electronegative oxygen enhances the ionic character of the M–X (M = transition metal, X = S, O, etc.) bond leading to an increase in the M\(^{n+}/M^{(n+1)}\) redox potential.

1.3.1.1 Layered oxides

*Goodenough* and coworkers showed in 1980 that lithium could be reversibly removed electrochemically from LiCoO\(_2\) around 3.9 V vs. Li\(^+/Li^0\), thus making it suitable cathode material,\(^{21}\) still used intensively in commercial Li-ion cells up to date. The crystal structure of layered oxides is built of a cubic close packing \(ccp\) of oxygen anions with Co\(^{3+}\) and Li\(^+\) sitting in octahedral sites in alternating layers (Figure 11). This material delivers attractive volumetric energy densities, excellent cyclability and a high rate capability, equivalent with the removal of 0.5 Li\(^+\) per Co, therefore leading to a limited capacity of \(\sim 150\) mAh/g. If more than 0.5 Li\(^+\) are electrochemically removed (going to higher voltages vs. Li), the 2D layered structure rearranges giving a \(hcp\) hexagonal close packing of the oxygen anions in CoO\(_2\).\(^{35}\) This is commonly observed for all lithium transition metal oxides Li\(_x\)MO\(_2\) for \(x<0.5\) releasing oxygen from the lattice and showing a limited thermal stability in the delithiated state. Although LiCoO\(_2\) is limited in gravimetric capacity and shows some concerns for large scale applications (thermal safety, high cost and toxicity of cobalt), *SONY* combined this material with a carbon anode to release the first commercial Li-ion battery.\(^{28}\) Recent development showed that by surface modifications or partial substitution of Co through Ni and Al (Li\(_{Ni_{0.80}Co_{0.15}Al_{0.05}}O_2\), NCA), the amount of reversibly extracted lithium could be increased up to 0.7 Li\(^+\) per formula unit, leading to reversible capacities around 185 mAh/g.\(^{36,37}\)
Figure 10: Schematic view of the crystal structures of layered LiCoO$_2$ with Li-, Co- and O atoms drawn in yellow, blue and red respectively.

Many sodium counterparts Na$_x$MO$_2$ have been investigated already since the 1970s in terms of their structure$^{38,39}$ but due to the different coordination environment of Li and Na, ion-exchange reaction between them comes along with a change in the MO$_2$ layer stacking.$^{40,41}$ All Na$_x$MO$_2$ phases reversible intercalate Na ions, but M = Mn and Co have been the most studied, showing in general a multistep voltage-composition curve with several solid-solution and biphasic domains for different x-values due to Na-vacancy ordering or gliding of the transition metal planes.$^{42,43}$ Although a variety of different compounds show a promising electrochemical behavior, the understanding and minimizing of irreversible structural changes affecting the cycling behavior represents still an issue which remains to be solved in the future along with the discovery of new compounds.$^{44}$ Along the Na-layered oxides P2-type Na$_x$[Fe$_{1/2}$Mn$_{1/2}$]O$_2$ is one of the most promising one due to the earth-abundant composition and a reversible capacity around 190 mAh/g (Figure 12).$^{45}$

Figure 11: (a) Structure of the P2-Na$_{2/3}$[Fe$_{1/2}$Mn$_{1/2}$]O$_2$ its (b) galvanostatic cycling and the corresponding capacity retention.$^{45}$

1.3.1.2 Spinel manganese oxide

Unlike cobalt and nickel, the straight forward synthesis of LiMnO$_2$ with the LiCoO$_2$ structure is not possible. It can only be obtained by ion exchange from $\alpha$-NaMnO$_2$, but
converts easily from the layered to a spinel type structure upon lithium insertion/deinsertion.\textsuperscript{46,47} By contrast for a 1:2 Mn:O ratio stable phases are formed, e.g. Li$_{0.5}$MnO$_2$ or LiMn$_2$O$_4$ with spinel type structure. This material was originally proposed as a cathode material by Thackeray et al.,\textsuperscript{22} and has been extensively studied in the Bellcore labs.\textsuperscript{48,49} Spinel LiMn$_2$O$_4$ is built up of MnO$_6$ octahedra connected through edge-sharing to forming a 3D network of conduction path for lithium ions sitting in tetrahedral sites (Figure 13a). One half of the manganese occurs in a +III oxidation state while the other half is in +IV oxidation state, thus leading to an amphoteric electrode material. This means that the material can be either oxidized leading to the formation of γ-MnO$_2$ with a potential centered around 4.1V vs. Li$^+$/Li$^0$, or reduced to Li$_2$Mn$_2$O$_4$ at an average potential of 3.1V vs. Li$^+$/Li$^0$ (Figure 13b). However for Li-ion batteries only the high-voltage plateau can be used, since there is no other source of lithium inside a full cell except the cathode material, thus limiting the practical reversible capacity to 148 mAh/g.

**Figure 12:** (a) Structure of the spinel LiMn$_2$O$_4$ where MnO$_6$ octahedra, lithium- and oxygen atoms are shown in purple, yellow and red respectively. (b) Voltage-composition curve of LixMn$_2$O$_4$ for 0<x<2.\textsuperscript{50}

Despite the slightly lower capacity compared to other lithium transition metal oxides, LiMn$_2$O$_4$ has several advantages for application in commercial cells like high voltage, non-toxicity and low-cost of manganese.\textsuperscript{51} Therefore this material has been serious considered as a material for high-power applications, prevented by unsolved issues, regarding cycling at elevated temperature (~50°C) or manganese dissolution over time.\textsuperscript{50} Several solutions regarding these issues have been proposed, including excess lithium,\textsuperscript{52} metal substitutions\textsuperscript{53} or mixing the spinel with a layered phase.\textsuperscript{54}
1.3.1.3 Li- and Na-rich layered oxides

Continuing the research for new cheap, safe and non-toxic cathode materials, two significant advances in the synthesis and electrochemistry of manganese based layered materials were made. In particular Li$_2$MnO$_3$ turned into the most interesting candidate to further improve oxide based cathode materials. It has a rocksalt structure which can be represented in the conventional layered LiMO$_2$ notation as Li[Li$_{0.33}$Mn$_{0.67}$]O$_2$ as ¼ of the Li-atoms of Li$_2$MnO$_3$ are sitting in the transition metal sheets (Figure 14).\textsuperscript{55,56} First the discovery that Li$_2$MnO$_3$, which should be electrochemically inactive for lithium insertion/extraction from a structural point of view since manganese is already in its highest oxidation state +IV, could be activated either chemically by acid leaching of Li$_2$O leading to a new compound with the stoichiometry Li$_{2-x}$MnO$_{3-x/2}$, or electrochemical by charging a Li$_2$MnO$_3$/Li cell to a high voltage (> 4.5 V vs. Li$^+/Li^0$), renewed the interest in this class of materials.\textsuperscript{57,58} Substituting Mn by other 3d transition metals like Ni and Co led to the discovery of a new class of materials xLi$_2$MnO$_3$∙(1-x)LiMO$_2$. Several groups reported capacities exceeding 250 mAh/g for this class of materials, therefore being the number one candidate as cathode materials for next generation Li-ion batteries. The unusual high capacity achieved could not be explained satisfactorily, by only taking into account the cationic redox process (M$^{n+}$ ↔ M$^{(n+1)+}$), however our group showed recently the participation of a reversible anionic redox process (O$^2-$ ↔ O$_2^{2-}$) for lithium- and sodium rich layered oxides, opening a new route in the search for high capacity cathode materials.\textsuperscript{59–61}

\begin{figure}[h]
\centering
\includegraphics[width=0.8\textwidth]{layered_structures.png}
\caption{Layered structures of (a) LiMO$_2$ (M = Co, Ni, Mn) and (b) Li$_2$MnO$_3$.\textsuperscript{62}}
\end{figure}
1.3.2 Polyanionic cathode materials and the inductive effect

Besides oxide based cathode materials, polyanionic materials consisting of a transition metal and a polyanionic group \((\text{XO}_n)^{m-}\) (\(X = \text{C, B, P, S, Si, Mo, W, etc.}\)) have gained strong interest since the discovery of electrochemically active lithium iron phosphate \(\text{LiFePO}_4\) in 1997.\(^{63}\) Although additional weight is introduced by the presence of the polyanionic group leading to a reduced gravimetric capacity, polyanionic electrode materials have several advantages as follows. First, they provide a stable anionic framework essential for long term stable cycling and safety issues (oxygen release in the case of oxides is circumvented). Second the redox potential of the \(\text{Mn}^{n+}/\text{M}^{(n-1)+}\) redox couple can be modified through the inductive effect of the polyanionic group (described in detail in the next paragraph) and leads to higher values compared to oxide compounds, and finally these class of compounds offer a very rich chemistry due to a variety of different crystal structures and possible substitutions.\(^{64,65}\)

In the following section an overview about different polyanionic materials is given, however since numerous compounds have been investigated, materials mainly based on the \(\text{Fe}^{3+}/\text{Fe}^{2+}\) redox couple are presented hereinafter.

1.3.2.1 NASICON-type compounds \(\text{A}_x\text{M}_2(\text{XO}_4)_3\) (\(\text{A} = \text{Li, Na}\))

In the 1980s it was demonstrated for the first time possible electrochemical reversible intercalation of \(\text{Li}\) or \(\text{Na}\) into 3D framework \(\text{NASICON (NA-SuperIonic CONductors)}\) phases \(\text{Fe}_2(\text{MoO}_4)_3\) (Figure 14) and \((\text{Li}, \text{Na})\text{Ti}_2(\text{PO}_4)_3\).\(^{66-68}\) In general they can be expressed by formula \(\text{A}_x\text{MM'}(\text{XO}_4)_3\) and are built up on a framework of \(\text{MO}_6\) and \(\text{M'O}_6\) octahedra sharing all their corners with \(\text{XO}_4\) tetrahedras, with alkali metal ions sitting interstitial spaces, enabling fast ion conduction.
Later Goodenough showed by substitution of the polyanionic group of Li$_3$Fe$_2$(XO$_4$)$_3$ (X = As, P, Mo, S), that the redox potential of the Fe$^{3+}$/Fe$^{2+}$ couple could be shifted up 3.6 V vs. Li$^+/Li^0$ (Figure 16a).$^{69,70}$ Given that all three compounds possess more or less the same metal to oxygen distance, they concluded that this increase in redox potential was due to the large electronegativity of the SO$_4^{2-}$ group, thus increasing the ionic character of the metal oxygen bond. The more electronegative X is, the stronger the covalent X─O bond, the weaker the M─O bond leading to a higher potential of the Fe$^{3+}$/Fe$^{2+}$ redox couple (Figure 16b), giving birth to the concept of inductive effect for polyanionic type compounds.

**Figure 14:** Crystal structure of the NASICON type compound Fe$_2$(MoO$_4$)$_3$ along the (a) a- and (b) b-axis. FeO$_6$ octahedra and MoO$_4$ tetrahedra are shown in brown and purple respectively.

**Figure 15:** Schematic representation of the inductive effect. (a) MO$_6$ octahedra connected over one bridging oxygen to the XO$_4$ tetrahedra. (b) Depending on the electronegativity of the polyanionic group, the redox potential of the Fe$^{3+}$/Fe$^{2+}$ can be shifted up from 2.8 to 3.6 V vs. Li$^+/Li^0$. $^{71}$

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**Figure 14:** Crystal structure of the NASICON type compound Fe$_2$(MoO$_4$)$_3$ along the (a) a- and (b) b-axis. FeO$_6$ octahedra and MoO$_4$ tetrahedra are shown in brown and purple respectively.

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Using this concept, the different redox potentials for different polyanionic environment could be sufficiently explained and new potential cathode materials with attractive voltage could be identified, however suffering from the high molecular weight thus theoretical capacities around 120 mAh/g. Subsequently the family was extended to phosphate based NASICON-type phases having the general formula Li$_x$MM’(XO$_4$)$_3$, summarized in Figure 17.

![Figure 16: Different NASICON-type compounds with a Li$_x$MM’(XO$_4$)$_3$ composition and different $M^{(n+1)+}/M^{n+}$ redox couples.](image)

Consequently this new concept called for investigating all different kind of polyanionic structures in combination with different transition metals, the most popular being iron due to its cost and safety (non-toxic element) advantages. In the course of this study Goodenough and coworkers identified a new compound lithium iron phosphate LiFePO$_4$ as a potential cathode material.

1.3.2.2 Phosphate based materials AMPO$_4$ and A$_2$MP$_2$O$_7$ (A = Li, Na; M = Mn, Fe, Co)

Lithium iron phosphate LiFePO$_4$ (LFP), the most famous compound among the family of phosphor-olivines, also known as the mineral *triphylite*, crystallizes in an orthorhombic space group and consists of edge sharing FeO$_6$ octahedra which are ordered in parallel layers. These layers are bridged by PO$_4$-groups forming tunnels along the [010] axis which is also the
preferred path for the 1D Li-ion conduction (Figure 18). The theoretical capacity of LFP is 170 mAh/g, comparable to that of layered oxides, but with lower redox potential of 3.45 V vs. Li⁺/Li⁰.

Figure 17: Structure of LiFePO₄ viewed along the (a) c- and the (b) b-axis. FeO₆ octahedras, PO₄ groups, oxygen and lithium atoms are represented in brown, purple, red and yellow.

Initially Goodenough et al. were only able to achieve ~80% of the theoretical capacity at slow charge/discharge rates. Then it was shown that the full capacity could be achieved by the nanosizing approach and nanocoating with carbon, even at fast rates for several hundred of cycles (Figure 19), hence commercialized by A123 in 2005. ⁷³–⁷⁶

Figure 18: Voltage-composition curve for galvanostatic cycling of carbon coated LiFePO₄. ⁷⁴

Based on the promising results for LFP, other phosphor-olivines LiMPO₄ (M = Mn, Co, Ni) were also investigated as potential electrode materials, while the latter two exhibit a
redox potential outside of the stability window of the common used liquid organic electrolytes\(^2\) (Table 2).\(^7^7\)

**Table 2: Redox potentials versus lithium and theoretical capacities for different transition metal phopho-olivines.**

<table>
<thead>
<tr>
<th>Phospho-olivine</th>
<th>Redox potential (V vs. Li(^+/Li^-))</th>
<th>Theoretical capacity (mAh/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LiFePO(_4)</td>
<td>3.45</td>
<td>170</td>
</tr>
<tr>
<td>LiMnPPO(_4)</td>
<td>4.1</td>
<td>171</td>
</tr>
<tr>
<td>LiCoPO(_4)</td>
<td>4.8</td>
<td>167</td>
</tr>
<tr>
<td>LiNiPO(_4)</td>
<td>5.1</td>
<td>167</td>
</tr>
</tbody>
</table>

Following that success for LFP in Li-ion batteries, the search for a Na equivalent was obvious, however the thermodynamically stable phase NaFePO\(_4\) does not crystallize in the *triphylite*- but rather in the *maricite* structure (Figure 19). The fact that Na\(^+\) ions are trapped in cavities inside the *maricite* structure renders this material not suitable as electrode material since it does not show any reversible redox behavior.\(^7^8\) However the *triphylite* NaFePO\(_4\) could only be prepared starting from LFP through chemical/ electrochemical ion exchange.\(^7^9^-^8^1\)

**Figure 19: Structure of maricite NaFePO\(_4\) along the (a) c- and (b) b-axis. FeO\(_6\) octahedra, PO\(_4\) tetrahedra, oxygen- and sodium-atoms are represented in brown, purple, red and blue respectively.**

\(^2\) Commonly used electrolytes for Li-ion batteries (a) 1M LiPF\(_6\) in ethylene carbonate EC / dimethylcarbonate DMC 1/1 w/w or (b) 1M LiPF\(_6\) in ethylene carbonate EC/ dimethylcarbonate DMC/ propylene carbonate PC 1/1/3 w/w/w.
Finally to explore other phosphate based cathode materials, which display a higher voltage in contrast to the PO$_4^{3-}$ compounds, so called pyrophosphate or diphosphate group [P$_2$O$_7$]$^{4-}$ have drawn attention to battery researchers.

Preliminary studies have focused on the electrochemical activity of pyrophosphate based compounds LiMP$_2$O$_7$ where M represents a trivalent metal e.g. Fe$^{3+}$ or V$^{3+}$. Starting on Li insertion for LiFeP$_2$O$_7$, this compounds showed a redox process for around 3.0 V vs. Li$^+/Li^0$ and a limited capacity of around 80 mAh/g in the case of M = Fe. Following these first investigations, the interest turned into lithium pyrophosphates containing a divalent metal and two lithium in the pristine material A$_2$MnP$_2$O$_7$ (A = Li, Na), where it should be theoretically possible to extract two Li$^+$ per transition metal.

A variety of different compounds, their solid solutions and polymorphs with the general formula A$_2$MP$_2$O$_7$ (M = Mn, Fe, Co, Ni; A = Li, Na) have been reported in literature. Figure 20a shows the structure of Li$_2$FeP$_2$O$_7$, crystallizing in a monoclinic space group P2$_1$\(/a$. Fe occupies three different crystallographic sites, one of them FeO$_6$ octahedra, the other two distorted FeO$_5$ trigonal bipyramids. Li is coordinates by oxygen forming LiO$_4$ tetrahedra and LiO$_5$ trigonal bipyramids leading to a 2D network along the b-c-plane (Figure 20a). Yamada’s group showed that it is possible to extract approximately 1Li$^+$ reversibly, giving around 110 mAh/g at an average redox potential of 3.5 V vs. Li$^+/Li^0$ playing with the Fe$^{3+}$/Fe$^{2+}$ redox couple, the highest redox potential observed for Fe$^{3+}$/Fe$^{2+}$ in phosphate based materials (500 mV higher than for LFP).

**Figure 20:** (a) Crystal structure of Li$_2$FeP$_2$O$_7$ (Li in green, FeO$_5$, FeO$_6$ in brown and P$_2$O$_7$ in purple) and (b) galvanostatic cycling vs. Li at a C/20 rate.
Following the work done on Li$_2$FeP$_2$O$_7$, the Na counterpart Na$_2$FeP$_2$O$_7$ was studied as a potential cathode material for Na-ion batteries. It adopts a triclinic space group P-1 and is built from corner sharing FeO$_6$-FeO$_6$ [Fe$_2$O$_{11}$] dimers, which are interconnected by [P$_2$O$_7$] units to create three-dimensional zigzag channels for Na$^+$-ion migration (Figure 21a). This enables to reach a reversible capacity around 80 mAh/g at an redox potential of 3.0V vs. Na$^+/Na^0$ (Figure 26b).\textsuperscript{89–91}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure21.png}
\caption{(a) Crystal structure of Na$_2$FeP$_2$O$_7$ (Na in yellow, FeO$_6$ in green and P$_2$O$_7$ in purple. (b) The voltage capacity curve.\textsuperscript{92}}
\end{figure}

Despite the theoretical possibility of two-electron reaction per transition metal (theoretical capacity 220 mAh/g), more than a one-electron process per transition metal was never observed and other transition metals like Co or Mn show limited electrochemical activities versus Na or Li (<80 mAh/g).\textsuperscript{87,88,93,94}

All the polyanionic based cathode materials presented before come along with moderate capacities and lower redox potentials if compared to state-of-the-art oxide based materials. To circumvent this disadvantage different solutions have been proposed through the past years. First, increase the specific capacity by lowering the molecular weight of the polyanionic group, or achieving more than 1 Li$^+$ transfer per transition metal. Second, increase the redox potential and hence the energy density of the material, applying strongly electronegative polyanionic groups. Both of the mentioned directions were further explored with borates and silicates for the former, and fluorosulfates and fluorophosphates for the latter one, discussed in the following sections.
1.3.2.3 Silicates Li$_2$MSiO$_4$

Following the path to achieve more than one Li$^+$ exchange per transition metal, the family of lithium transition metal silicates Li$_2$MSiO$_4$ (M = Fe, Mn, Co) was early identified as a candidate to master this challenge and intensive research efforts have been put into afterwards, revealing limited electrochemical performance.$^{95,96}$

The crystal structures of Li$_2$MSiO$_4$ consist of a distorted hcp of the oxygen anions with ½ of the tetrahedral sites occupied by Li, Si an M, where a variety of different polymorphs is possible depending on the occupancy of the tetrahedral sites. All of them are related to the structure of Li$_3$PO$_4$, either the low temperature- or the high-temperature polymorph, denoted as $\beta$- and $\gamma$-Li$_3$PO$_4$ respectively. The only difference in between all these polymorphs is basically the ordering/distribution of the cations within the tetrahedral sites, thus leading to a large variety (for sake of clarity only the two existing polymorphs of Li$_2$FeSiO$_4$ are shown in Figure 22).$^{97-99}$

![Figure 22: Two known polymorphs of Li$_2$FeSiO$_4$ crystallizing in (a) P2mn21 and (b) P21/n space group. FeO$_4$ tetrahedra, SiO$_4$ tetradedra and Li are drawn in brown, blue and yellow respectively.](image)

Regarding the application of silicates as cathode material, one of the drawbacks of this family is the low intrinsic conductivity at room temperature ($5\cdot10^{-16}$ S/cm for Li$_2$MnSiO$_4$ and $6\cdot10^{-14}$ S/cm for Li$_2$FeSiO$_4$).$^{100}$ This limitation could be overcome through an optimized synthesis including nanoparticles and conductive carbon coating.$^{101-103}$ Although optimized, it is not possible to extract more than 1Li$^+$ per formula unit affiliated to the Fe$^{3+}$/Fe$^{2+}$ redox couple at 3.1 and 2.8 V vs. Li$^+$/Li$^0$ for the initial and subsequent charge/discharge cycles (Figure 23) leading to a reversible capacity 120-140 mAh/g. The change in potential from the
first to the subsequent oxidations is due to a structural rearrangement after the first oxidation, minimizing the repulsion between Fe$^{3+}$ and Si$^{4+}$ in the charged state. Theoretical the second redox process involving the Fe$^{4+}$/Fe$^{3+}$ couple should show up at 4.7 V vs. Li$^+$/Li$^0$ (limit of the stability window of common electrolytes) but was never observed.\textsuperscript{99}

\textbf{Figure 23:} Voltage-composition curve of Li$_2$FeSiO$_4$ versus lithium at a C/20 rate.\textsuperscript{100}

In contrast to Li$_2$FeSiO$_4$, the other members of the silicate family Li$_2$MnSiO$_4$ and Li$_2$CoSiO$_4$ have not shown promising electrochemical activity. Additionally to electronic and ionic limitations of Li$_2$MnSiO$_4$, Mn$^{3+}$(d4) cations are not stable in a tetrahedral environment, leading to a serious distortion of the crystallinity during the first oxidation and a poor reversibility on following cycles. Even for the solid-solution member Li$_2$Mn$_{0.2}$Fe$_{0.8}$SiO$_4$ it was show that only Fe is taking part in the reversible redox process, hence Mn being non active.\textsuperscript{104}

\textbf{1.3.2.4 Sulfate based materials}

As already observed by Padhi et al. in the late 1990’s, the redox potential of iron could be shifted up ~0.8 V by replacing the phosphate in NASICON type compounds by a sulfate group (Figure 16). Following this work a variety of new materials out of the sulfate family were proposed as cathode materials for Li- and Na-ion batteries. To achieve even higher voltage compared to bare sulfate polyanionic compounds, the introduction of fluorine into the structure should lead to elevated redox potentials as F is even more electron withdrawing.
The first fluorosulfate reported in literature was LiMgSO\textsubscript{4}F with \textit{tavorite} structure, however ceramic methods to synthesize LiFeSO\textsubscript{4}F failed.\textsuperscript{105} Only eight years later \textit{tavorite} LiFeSO\textsubscript{4}F could be stabilized using an ionothermal route at low temperatures.\textsuperscript{106} As it turned out this material decomposes around 375°C and is water soluble, explaining why it was not possible to synthesize LiFeSO\textsubscript{4}F by conventional ceramic synthesis or solution chemistry in aqueous media. In addition, the use of monohydrate iron(II) sulfate precursor FeSO\textsubscript{4}·H\textsubscript{2}O was essential to obtain LiFeSO\textsubscript{4}F, since the structure of FeSO\textsubscript{4}·H\textsubscript{2}O and \textit{tavorite} LiFeSO\textsubscript{4}F a closely related. During the reaction of LiF with FeSO\textsubscript{4}·H\textsubscript{2}O in ionic liquid media \textit{1-Ethyl-3-methylimidazolium bis(trifluoromethylsulfonyl) imide} (EMI-TFSI) the water molecules linked to iron are replaced by fluorine and lithium is inserted into the structure for charge balance (Figure 24).\textsuperscript{107}

\textit{Tavorite} LiFeSO\textsubscript{4}F operates at a redox potential of 3.6 V vs. Li\textsuperscript{+}/Li\textsuperscript{0} and delivers a reversible capacity of 135 mAh/g (exchange of 0.8 Li\textsuperscript{+}, Figure 25b). However, contrary to LFP no carbon coating is needed to achieve this electrochemical performance, making LiFeSO\textsubscript{4}F a serious contender if it would not undergo fast degradation in ambient atmosphere.\textsuperscript{108}

Following the synthetic strategy developed for \textit{tavorite} LiFeSO\textsubscript{4}F, a great variety of other different compounds with the same stoichiometry AMSO\textsubscript{4}F (A = Li, Na, K; M = Fe, Co, Ni, Mn, Zn) crystallizing in different polymorphs could be prepared.\textsuperscript{109,110} During this in-depth examination another polymorph for LiFeSO\textsubscript{4}F was revealed, crystallizing in a \textit{triplite} structure (Figure 25c) and displaying with 3.9 V vs. Li\textsuperscript{+}/Li\textsuperscript{0} (Figure 25d) the highest redox potential for Fe-based polyanionic compounds ever reported.\textsuperscript{111,112}
Since the presence of fluorine in a cathode material could be problematic from the toxicity point of view, the replacement of F\(^{-}\) with OH\(^{-}\) was investigated. Starting from an investigation of the mixed F/OH compound FeSO\(_4\)F\(_{1-y}\)OH\(_{y}\),\(^{113}\) led to the discovery of a layered materials LiFeSO\(_4\)OH which showed good electrochemical performance versus lithium around 3.6 V with an average capacity of ~110 mAh/g.\(^{114}\)

As described in previous sections, it is possible to achieve high redox potentials for insertion type positive electrode materials, close to the boarder of the stability window of standard electrolytes (max. 4.8 V vs. Li\(^+\)/Li\(^0\)), however the specific capacity and energy density still remains low compared to the negative side of a lithium ion battery.
1.3.3 Conversion type cathode materials

All the before discussed cathode materials store electrical energy through an insertion/deinsertion (topotactic) reaction of Li/Na ions in a host structure (Figure 26). To further push the development of Li-ion batteries, reversible conversion reactions have been intensively investigated since the 2000’s. The following section will give a short introduction and overview about conversion cathode materials for Li- and Na-ion batteries.

Figure 26: Schematic illustration of insertion and conversion type reaction investigated as electrode materials for Li- and Na-ion batteries.\(^\text{15}\)

In order to use all the possible oxidation states of the TM, thus increasing the specific capacity drastically, conversion type reactions have been investigated since decades for primary batteries exclusively,\(^\text{115}\) because it was believed that such reactions are irreversible. Besenhard et al. however showed in 1978 a reversible conversion reaction for a non-transition metal sulfide, namely Bi\(_2\)S\(_3\). During discharge elemental bismuth Bi\(^0\) and LiS were formed, nevertheless during the subsequent charge only about 75% could be converted back into Bi\(_2\)S\(_3\). Furthermore the reversibility of the discharge-charge could be demonstrated only for one cycle.\(^\text{116}\)

The first real reversible conversion reaction could be demonstrated for TM oxides in 2000, drawing renewed attention on the investigation of conversion type electrode materials for rechargeable batteries. The conversion process of these compounds can be described as
follows: the metal M is reduced to elemental M\(^0\) and simultaneously the corresponding lithium compound LiX is formed (equation 1, X = O, S, P, etc.).

\[ M_aX_b + ne^- + nLi^+ \xrightarrow{\text{charge}} aM^0 + bLi_nX \]  
(2)

Although the conversion reaction of the TM-oxides delivers much higher gravimetric capacities compared to insertion type electrode materials (Figure 27), the redox potential versus lithium of 1 to 1.5 V vs. Li\(^+/Li^0\), is too low for practical applications.\(^{117}\)

![Figure 27: (a) Voltage composition curves for different TM oxides and (b) the corresponding capacity retention. Co\(_3\)O\(_4\) is shown to proof that not only divalent TM oxides can be applied. The inset show the rate capability of CoO.\(^{117}\)](image)

In order to increase the redox potential, more ionic TM compounds like the corresponding sulfides,\(^{118}\) -nitrides\(^{119}\) and especially fluorides\(^{120,121}\) were investigated for conversion reaction versus lithium. Among them iron fluoride FeF\(_3\) is one of the most promising one, but due to its insulating nature it has to be “activated” to be used as electrode material. This activation is usually achieved by forming an iron fluoride carbon composite FeF\(_3\)/C (ca. 20 wt% carbon) during a prolonged high energy milling. Cycling this composite versus lithium between 1.5 and 4.5 V, around 600 mAh/g capacity can be achieved (Figure 28).
Figure 28: (a) Voltage versus time curve of a FeF$_3$-activated carbon nanocomposite ball milled for 6h and (b) the corresponding capacity retention for different kind of carbon.\textsuperscript{120}

The energy density of this FeF$_3$/C composite is rather low due to the high carbon content in the composite without considering the low gravimetric density of FeF$_3$ itself compared to other positive electrode materials (Table 3).\textsuperscript{120,122} However one main drawback of FeF$_3$ cathode composites is the large polarization, which has been explained by the different reaction pathways during discharge/charge.\textsuperscript{123}

<table>
<thead>
<tr>
<th>Compound</th>
<th>Redox potential vs. Li$^+$/Li$^0$ (V)</th>
<th>Gravimetric capacity (mAh/g) theo/exp Mn$^+\rightarrow$M$^0$</th>
<th>Gravimetric energy density (Wh/kg) theo</th>
<th>Volumetric density (g/cm$^3$)</th>
<th>Volumetric energy density (Wh/L)</th>
<th>Voltage hysteresis (V)</th>
<th>Carbon percentage (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LiCoO$_2$</td>
<td>3.9</td>
<td>274/148</td>
<td>1068</td>
<td>5.1</td>
<td>5447</td>
<td>0.01</td>
<td>5</td>
</tr>
<tr>
<td>LiFePO$_4$</td>
<td>3.4</td>
<td>170/165</td>
<td>589</td>
<td>3.6</td>
<td>2120</td>
<td>0.02</td>
<td>5</td>
</tr>
<tr>
<td>FeF$_3$</td>
<td>2.74</td>
<td>713/600</td>
<td>1954</td>
<td>3.87</td>
<td>7562</td>
<td>0.8-1.6</td>
<td>20</td>
</tr>
<tr>
<td>BiF$_3$</td>
<td>3.18</td>
<td>303/230</td>
<td>964</td>
<td>5.32</td>
<td>5128</td>
<td>0.5-0.7</td>
<td>20</td>
</tr>
<tr>
<td>CuF$_2$</td>
<td>3.55</td>
<td>528/250</td>
<td>1874</td>
<td>4.23</td>
<td>7927</td>
<td>0.8</td>
<td>20</td>
</tr>
</tbody>
</table>

In order to achieve higher energy densities and maintain the benefit of high specific capacity, bismuth fluoride\textsuperscript{121,125} and bismuth oxyfluoride\textsuperscript{126} turned out as interesting candidates. In 2005 it was shown that a BiF$_3$/C composite could deliver a gravimetric capacity...
of around 230 mAh/g at an average redox potential of 3 V vs. Li⁺/Li⁰, however fast capacity fading (Figure 29).  

**Figure 29:** (a) Galvanostatic curve of BiF₃ versus lithium for different current rates and (b) the capacity retention for ab BiF₃/C ball milled for different times composite with 20wt% carbon. ¹²¹,¹²⁵

### 1.4 Borate based cathode materials for lithium- and sodium-ion batteries

Boron, the fifth element of the periodic table, doesn’t occur in its elemental form in nature, only as its oxide form. More than 200 borate minerals are found in earth’s crust, so called borates, but only a few of them are of commercial interest: *borax, kernite, ulexite* and *colemanite* (Table 4). ¹²⁷

**Table 4:** The four main borate based minerals found in earth’s crust with their chemical formula and the main deposit location. ¹²⁷

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Formula</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>Borax</td>
<td>Na₂B₄O₇10H₂O</td>
<td>USA, Turkey, Argentinia</td>
</tr>
<tr>
<td>Kernite</td>
<td>Na₂B₄O₇·4H₂O</td>
<td>USA</td>
</tr>
<tr>
<td>Ulexite</td>
<td>NaCaB₄O₇·8H₂O</td>
<td>Turkey, South America</td>
</tr>
<tr>
<td>colemanite</td>
<td>Ca₃B₆O₁₁·5H₂O</td>
<td>Turkey</td>
</tr>
</tbody>
</table>

Starting in the 19th century, borates based glasses have been studied for their optical properties and later on for alkali ion conduction. ¹²⁸,¹²⁹ Nevertheless crystalline borate based
materials were mainly investigated because of its non-linear optical properties and as host structures for light emitting phosphors. Since the discovery of $\beta$-BaB$_2$O$_4$ and LiB$_3$O$_5$ in the mid 1980’s (which are up to now still the most frequently used borate based non-linear optical properties (NLO) materials) an increased interest in the scientific community has led to the discovery of a large variety of borate based compounds.$^{132}$ It is believed that these optical properties derive from the unique crystal- and electronic resulting from the small boron atoms inside an oxide matrix.$^{133}$

**1.4.1 Why borate based materials?**

As mentioned above, borate based minerals can form a variety of different structures, therefore an uncountable number of borate based materials has been reported in literature. In terms of application of borates as electrode materials for Li-ion batteries, they present two main advantages. First of all, in the context of the polyanionic concept for cathode materials, the borate anion represent the lightest anionic group among all others (Table 5), hence having the possibility to increase the gravimetric capacity significant in contrast to phosphates or silicates.

*Table 5: Comparison of the most common polyanions in terms of the molecular weight.*

<table>
<thead>
<tr>
<th>Polyanion</th>
<th>Molecular weight [g/mol]</th>
</tr>
</thead>
<tbody>
<tr>
<td>BO$_3$$^-$$^-$</td>
<td>59</td>
</tr>
<tr>
<td>CO$_3$$^-$$^-$</td>
<td>60</td>
</tr>
<tr>
<td>SiO$_4$$^+$</td>
<td>92</td>
</tr>
<tr>
<td>PO$_4$$^+$</td>
<td>95</td>
</tr>
<tr>
<td>SO$_4$$^-$</td>
<td>96</td>
</tr>
</tbody>
</table>

A second reason, to explore borate based cathode materials is the very rich chemistry since boron can adopt trigonal planar- or tetrahedral oxygen coordination (Figure 30), leading to unique condensed polyanions.$^{132,134}$
Driven by these two points, the investigation of borates as cathode materials took its course starting around 2000, however involving exclusively compounds based on the trigonal planar BO$_3$ group, the so called ortho borate group.

### 1.4.2 Lithium transition metal borates

The first transition metal borate compounds LiMBO$_3$ were investigated since the late 1990’s for their optical and structural properties (M = Mg, Co, Cd, Zn). It was first in 2001 when Legagneur et al. introduced the family of lithium 3d-metal borates LiMBO$_3$ (M = Fe, Mn, Co) as potential cathode materials for Li-ion batteries, with theoretical capacities exceeding 200 mAh/g. The three compounds were prepared by classical ceramic synthesis at high temperatures. The crystal structures for M = Mn, Fe, Co crystallizing in a monoclinic space group are shown in Figure 31, and are built up of edge sharing [MO$_5$] trigonal-bipyramids running along the c-axis. These columns are connected through planar BO$_3$-groups and split lithium sites in tetrahedral coordination. In general the three compounds differ only in the size of the trigonal-bipyramid depending on the transition metal ion. Only for M = Mn a hexagonal polymorph is reported (Figure 31d) which can be obtained from the monoclinic form if heated above 550°C.
**Figure 31:** Crystal structures of LiMBO$_3$ for (a) $M = \text{Mn}$ (monoclinic), (b) Fe, (c) Co and (d) Mn (hexagonal). Oxygen, lithium and boron atoms are shown in red, yellow and green respectively.

Initially Legagneur et al. reported an electrochemical negligible activity versus lithium even if very slow current rates were applied (Figure 32).

**Figure 32:** Voltage composition curve of LiMBO$_3$ cycled with different C-rates for (a) hexagonal-Mn, C/60, (b) Fe, C/250 and (c) Co, C/100.\textsuperscript{138}

It took around one decade to optimize synthesis conditions,\textsuperscript{139–142} so that by preparing nano-sized particles\textsuperscript{143–146} or controlling antisite-disorder\textsuperscript{147,148} the experimental capacities could be significantly improved. However only for LiFeBO$_3$ the theoretical capacity of 220
mAh/g could be achieved. In contrast the theoretical capacities could not be reached for LiMBO₃ (M = Mn, Co) (Figure 33).

![Figure 33: Galvanostatic charge-discharge curves for (a) LiCoBO₃, (b) LiFeBO₃, (c) m-LiMnBO₃ and (d) h-LiMnBO₃ cycled at a C/20 rate versus lithium.]

After the initial report of Yamada at al. where he almost achieved the theoretical capacity for LiFeBO₃, this material turned into a serious earth abundant cathode material for lithium ion batteries, confirmed by following publications. As mentioned in the initial paper, LiFeBO₃ displays a redox potential versus lithium of around 2.8 V vs. Li⁺/Li₀, rendering this compound moisture sensitive. Therefore this material has to be handled carefully in inert atmosphere during the whole synthesis and cell preparation process. If not, it undergoes a fast degradation process as studied deeply in literature leading to a considerably reduced capacity.

Besides the above describe class of lithium transition metal borates LiMBO₃ (M = Mn, Fe, Co) only one more compound has been reported in literature as a candidate for cathode materials in Li-ion batteries, a lithium manganese borate LiₓMn(BO₃)₂. This material crystallizes in a triclinic space group having Mn²⁺ in a tetrahedral environment surrounded by planar BO₃ groups and lithium in tetrahedral sites. The authors claim to be able to reach a
reversible capacity of around 120 mAh/g with an average redox potential centered around 3 V vs. Li$^+$/Li$^0$ (Figure 34), but again this material has to be handled thorough under inert atmosphere since it is decomposed by moisture.$^{153}$ However no following publication reporting on the synthesis and electrochemical activity on this material was able to confirm their results so far.

Figure 34: (a) Structure of Li$_7$Mn(BO$_3$)$_3$ viewed along the a-axis. Oxygen, lithium boron and manganese are represented in red, yellow, green and purple respectively. (b) Galvanostatic charge-discharge curve for Li$_7$Mn(BO$_3$)$_3$ between 4.7 and 1.7 V vs. Li$^+$/Li$^0$ with 10 mA/g.$^{153}$

1.4.3 Borophosphates

The low potential $\sim$2.8 V vs. Li$^+$/Li$^0$ of LiFeBO$_3$ prevents it from being a commercial challenger to state-of-the-art cathode materials. Following the polyanionic concept, borophosphates, a combination of borate- and phosphate groups could offer elevated redox potentials compared to pure borate based compounds. Generally speaking borophosphates are compounds out of the system M$_x$O$_y$-B$_2$O$_3$-P$_2$O$_5$-(H$_2$O) with complex anionic structures built up of BO$_3$, BO$_4$ and PO$_4$ anions and their partially protonated species, thus offering an incredible huge amount of possible structures and compounds.$^{154}$ Known already for one century, only about one decade ago a systematic investigation of this class of inorganic compounds has been started and pushed forward mainly by Kniep and coworkers.$^{155,156}$

Borophosphates are usually prepared using hydrothermal synthesis methods leading often to open framework and microporous structures, investigated in the past for their optical and catalytic properties.$^{157-161}$ These open structures providing possible Li- and Na-ion migration pathways made borophosphate materials attractive candidates as possible cathode
materials for Li- and Na-ion batteries. Recently two examples reported in literature show indeed possible extraction/insertion of Li- or Na-ions at redox potentials slightly higher than \( \text{LiFeBO}_3 \), nevertheless the capacity (even theoretical) is relatively low due to the high molecular weight of this class of materials (Figure 35).\textsuperscript{162,163}

![Figure 35](image)

**Figure 35:** (a) Galvanostatic charge/discharge curve of \( \text{Li}_{0.8}\text{Fe}((\text{H}_2\text{O})_2[\text{BP}_2\text{O}_8] \cdot \text{H}_2\text{O} \) versus lithium and (b) voltage composition curve of \( \text{NH}_4_{0.75}\text{Fe}((\text{H}_2\text{O})_2[\text{BP}_2\text{O}_8] \cdot 0.25\text{H}_2\text{O} \) versus sodium.\textsuperscript{162,163}

### 1.5 Motivation and aim of the thesis

As mentioned, lithium transition metal borates \( \text{LiMBO}_3 \) can be attractive candidates as polyanionic cathode materials for Li-ion batteries. It has been shown in the case of \( \text{LiFeBO}_3 \) that high gravimetric capacities (~200 mAh/g) are accessible if \( \text{BO}_3 \) is used instead of \( \text{PO}_4 \) or \( \text{SO}_4 \) as the polyanionic group.\textsuperscript{139} Despite the weight gain coming along with borates, in any case the redox potential will be lower compared to phosphate \( \text{PO}_4 \) or sulfate \( \text{SO}_4 \) based materials, which it not surprising if one takes a look at the Pauling electronegativity of the following atoms:\textsuperscript{164}

\[
\text{B}(2.09) < \text{C}(2.55) < \text{P}(2.19) < \text{S}(2.58) < \text{O}(3.44) < \text{F}(3.98).
\]

Recently it was shown in the case of \( \text{LiFePO}_4 \), that the redox potential of 3.45 V vs. \( \text{Li}^+/\text{Li}^0 \) could be shifted up around 500 mV by using a pyrophosphate \( [\text{P}_2\text{O}_7]^– \) based material \( \text{Li}_2\text{FeP}_2\text{O}_7 \).\textsuperscript{86} Although the voltage increase is not outstanding and its origin is not fully understood yet,\textsuperscript{165} it was a first hint to apply this concept to the exploration of borate based cathode materials beyond \( \text{BO}_3 \) based materials.
Introduction

Even though pyroborate (or diborate)-$[\text{B}_2\text{O}_3]^{4-}$ and other various polyborate ($[\text{B}_5\text{O}_{10}]^{3-}$, $[\text{B}_3\text{O}_8]^{3-}$, etc.) based compounds are well known in literature, there is certainly a lack of compounds suitable as potential cathode materials.\textsuperscript{166–169} Only two alkali-3d-metal pyroborate materials, NaScB$_2$O$_5$ and Li$_6$CuB$_4$O$_{10}$ and no other alkali-3d-metal polyborate based material was reported so far.\textsuperscript{170–173} This was an inspiration to get involved in the search for new pyroborates as described in the following sections.

Thus, the work presented in this thesis was aimed to design new borate based cathode materials for lithium and sodium ion batteries. To achieve this goal a synthetic exploration combined with fundamental examination of structure-properties relationship has been conducted. The following manuscript presents the main results obtained during this study, and is divided into five chapters.

First an introduction into electrochemical energy storage in batteries (emphasis on Lithium- and Sodium ion batteries) with a brief overview about the development and state-of-the-art cathode materials is given. Furthermore the motivation to explore explicit borate based materials will be pointed out and the approaches applied during this work will be described in detail.

Secondly the experimental techniques and approaches employed within this thesis will be described in details, to inspire further studies or experimental work in this field.

The third chapter shows as a proof of concept the investigation of the structural electrochemical properties of a lithium copper borate Li$_6$CuB$_4$O$_{10}$, displaying a high redox potential of 4.25 V vs. Li/Li$^+$ and a surprisingly high ionic conductivity around 1.4 mS/cm at 500°C.

The fourth chapter focuses on new compounds synthesized, namely sodium transition metal pentaborates Na$_3$MB$_5$O$_{10}$ (M = Fe, Co). Their crystal structures are solved and electrochemical properties versus sodium are revealed, showing that it is possible to reversible desodiate Na$_3$FeB$_5$O$_{10}$ at elevated temperatures (55°C) at a redox potential of 2.5 V vs Na$^+/\text{Na}$ (≈30 mAh/g) whereas Na$_3$CoB$_5$O$_{10}$ is electrochemically inactive, due to kinetic limitations within these materials, revealed by AC and DC conductivity measurements.

To overcome the limitations of insertion/ deinsertion reactions for borate based compounds, a bismuth oxyborate Bi$_4$B$_2$O$_9$ is tested as a conversion type cathode material versus lithium. The reaction mechanism is revealed and it is shown that this material undergoes a conversion reaction around 2.3 V vs. Li$^+/\text{Li}$ with surprisingly little carbon additive and no need for nano-sized particles as usual for conversion type cathode materials.
Finally all the results presented within this thesis work are summarized and compared to materials studied up to now for the same purpose. At the end a broader context and outlook regarding borate based materials for energy storage applications is given.
2 Search for new pyroborate $\text{B}_2\text{O}_5$ based compounds

It is nowadays established that by modifying the polyanion of a cathode material, elevated redox potentials versus Li/Na can be achieved. Following this direction, we tried in this thesis to increase the potential of borate based materials, since for instance LiFeBO$_3$ shows a voltage of 2.8 V which is rather low compared to other polyanionic cathode materials. To increase the redox potential condensed polyanions can be introduced, as previously recalled in switching from LiFePO$_4$ to Li$_2$FeP$_2$O$_7$.

Thus, for Fe-based materials, the expected potential should be well above 2.8 V vs. Li$^+$/Li$^0$ if going from BO$_3$ to B$_2$O$_5$ or even B$_2$O$_5$F.

We have followed this strategy and our results are reported in this chapter which first gives a general overview about the reactivity of borates and possible synthetic routes for crystalline borate materials and then focuses on the experimental description of our first synthetic approaches, theoretically leading to pyroborate cathode materials.

2.1 A few synthesis considerations

Given by the nature of boric acid H$_3$BO$_3$ (mainly used precursor for borates) which contains one boron atom connected to three hydroxyl groups, borate based compounds can be synthesized through different synthetic methods. It is known that H$_3$BO$_3$ can easily undergo dehydration/ polymerization reactions leading to unpredictable crystal structures. As schematically shown in Figure 36, a complete- or a partial dehydration of H$_3$BO$_3$ can be realized, leading to new polyanions that can be viewed as a polymerization of the ortho borate group BO$_3$. These different synthetic routes can involve more or less all techniques used in solid state/ inorganic chemistry including high temperature ceramic-, or low temperature mechanochemical or solvothermal methods. Some of these approaches will exemplary be described in the next sections, whether successful or not, however with the aim to possibly provide ideas for future synthetic work.
Figure 36: Schematic illustration of the synthetic concept applied within this thesis. Starting from boric acid different ways can be chosen, involving different synthetic approaches and temperatures. For reasons of clarity only ortho- and pyroborate based groups are shown. However one has to keep in mind that borate based species can not only polymerize in two-, but also in three-dimensions involving trigonal BO\textsubscript{3} and tetrahedral BO\textsubscript{4} coordinated species.

Dealing with cathode materials, compounds like A\textsubscript{2}MB\textsubscript{2}O\textsubscript{5} (A = Li, Na; M = Mn, Fe, Co) are of interest. However as known from preliminary work,\textsuperscript{174} a lot of difficulties were encountered on the way to stabilize these phases. Even if different type of metathesis reactions starting from pyroborate based precursors (M\textsubscript{2}B\textsubscript{2}O\textsubscript{5} or Li\textsubscript{4}B\textsubscript{2}O\textsubscript{5})\textsuperscript{168,175,176} were tried, no progress was made. Since these preliminary results were not promising, other model compounds were needed, and along that line Li\textsubscript{0.8}M\textsubscript{2.1}B\textsubscript{2}O\textsubscript{5}F, a lithium containing fluoro pyroborate was recently published.\textsuperscript{177} If Mg\textsuperscript{2+} could be replaced by a redox active divalent 3d metal, it should be theoretically possible to achieve a capacity of 121 mAh/g by reversible extracting 0.8 Li\textsuperscript{+}. Needless to say that the capacity value is not outstanding, however it could act as a proof of concept that pyroborate based materials offer elevated redox potentials versus Li compared to LiMBO\textsubscript{3}. In the following sections different synthetic approaches are described as attempts to synthesize Li\textsubscript{0.8}M\textsubscript{2.1}B\textsubscript{2}O\textsubscript{5}F (M = Mn, Fe, Co).
2.2 Ceramic synthesis

A straightforward solid state synthesis was originally reported by Wang et al. consisting of mixing stoichiometric amount of lithium carbonate Li$_2$CO$_3$, magnesium fluoride MgF$_2$, magnesium nitrate Mg(NO$_3$)$_2$·6H$_2$O and boric acid H$_3$BO$_3$ with a subsequent annealing at 750°C for 60 h in a platinum crucible with intermediate regrinding. So our first attempts consisted in replacing MgF$_2$ by a 3d metal fluoride MF$_2$ (M = Co, Mn, Fe) and Mg(NO$_3$)$_2$·6H$_2$O through the corresponding oxalate.$^3$ However this first approach failed entirely, instead more thermodynamically stable M$_2$B$_2$O$_5$ and lithium fluoride LiF were formed in all cases, guiding us to directly start from the pyroborate precursor M$_2$B$_2$O$_5$ and react it with LiF (equation 3). Therefore the approach was again unsuccessful for M = Mn; Fe, Co as more stable LiMBO$_3$ and Li$_2$B$_4$O$_7$ with other unidentified impurity phases were formed. Note that in the case of M = Mg the desired phase was formed.

\[
\text{LiF} + M_2B_2O_5 \rightarrow LiM_2B_2O_5F \quad (3)
\]

This unfruitful experiments let us to explore unconventional synthetic approaches, involving more reactive precursors and low temperature solution based approaches, with the aim to stabilize metastable phases which are not accessible through high temperature ceramic methods.

2.3 Low temperature synthesis

Given the difficulties to obtain alkali-3d metal pyroborates we decided to start from more reactive borate based precursors to form in situ a pyroborate. One way is to use a precursor containing a borate hydroxide group which can undergo a in situ dehydration reaction upon thermal treatment as shown in Figure 36.

Magnesium borate hydroxide MgBO$_2$OH is well known as the mineral szaibelyite and the preparation of MBO$_2$OH (M = Mg, Mn, Fe, Co) was reported.$^{178}$ These compounds which contain a [B$_2$O$_4$OH] structural unit closely related to a pyroborate B$_2$O$_5$ group, are synthesized

$^3$ 3d metal nitrates were not considered since the nitrous gases evolved during heating easily oxidize the divalent metal.
hydrothermally at temperatures ranging from 180 to 200°C with reaction times between 5 and 10 days.

Using a stoichiometric mixture of MgBO₂OH and LiF (5-10 mol% excess), Li₀.₈Mg₂.₁B₂O₅F could be synthesized at temperatures around 600°C contrary to the reported synthesis (equation 4). Switching to MBO₂OH (M = Mn, Fe, Co) didn’t bring the desired success, driving us to consider different reaction media. As it is known ionic liquids can promote reaction between two insoluble solids. We used a solvothermal approach in 1-Ethyl-3-methyl imidazolium bis (trifluoro methyl sulfonyl) imide (EMI-TFSI) where MBO₂OH was dispersed together with AF (A = Li, Na, K) and exposed to temperatures around 250°C for 24 h. Unfortunately for all these reactions MF₂ and AMF₃ were formed.

\[
\text{LiF} + 2 \text{MBO}_2\text{OH} \rightarrow \text{LiM}_2\text{B}_2\text{O}_5\text{F} + \text{H}_2\text{O} \quad (4)
\]

Facing this failure, we next considered Tris (pentafluorophenyl) borane TPFPB as a “catalyst” to drive the reaction at temperatures below 100°C. TPFPB well known in organometallic catalysis as strong Lewis acid and is able to “dissolve” LiF in organic solvents by forming a [TPFPB─F]⁻ anion releasing Li⁺ as a counter ion (Figure 37).

Our idea was to dissolve stoichiometric amounts of LiF and TPFPB in organic solvents like propylene carbonate PC, to achieve a reaction with MBO₂OH (equation 5), however no reactivity at all was observed in this system.

\[
\text{Li}^+ + \text{TPFPB} \cdot \text{F} + 2 \text{MBO}_2\text{OH} \rightarrow \text{LiM}_2\text{B}_2\text{O}_5\text{F} + \text{TPFPB} \cdot \text{H}_2\text{O} \quad (5)
\]

At this stage bearing in mind the difficulties encountered in preparing fluoroborates we decided to switch our efforts towards pure pyroborates.
There are only two alkali-3d-metal pyroborate compounds reported so far: a) sodium scandium pyroborate NaScB$_2$O$_5$ and b) lithium copper pyroborate Li$_6$CuB$_4$O$_{10}$ for which two polymorphs $\alpha$ and $\beta$ were reported. As Sc is not redox active, particular attention has been given to Li$_6$CuB$_4$O$_{10}$ which contains two pyroborate units and Cu$^{2+}$ as the redox active transition metal. Combining both features, theoretical high redox potentials are expected for the reversible extraction/insertion of Li, hence this material was subjected to a thoroughly investigation of its electrochemical and structural properties, although its theoretical capacity associated to the Cu$^{3+}$/Cu$^{2+}$ redox couple is only 87 mAh/g.
3 Structural and electrochemical properties of a lithium copper pyroborate Li$_6$CuB$_4$O$_{10}$

The synthesis of Li$_6$CuB$_4$O$_{10}$ was first reported in 2006 by Pan et al.\textsuperscript{171} but already mentioned in Sparta’s PhD thesis in 2003. The synthesis of this phase was done by the authors through mixing stoichiometric amounts of Li$_2$CO$_3$, CuO and H$_3$BO$_3$ and annealing the mixture at 590°C for 2-3 days, denoted as $\alpha$-Li$_6$CuB$_4$O$_{10}$ afterwards.\textsuperscript{173} To obtain a better crystallized product our synthesis consisted in mixing stoichiometric amounts of the same precursors with mortar and pestle, followed by a decomposition of H$_3$BO$_3$ at 450°C for 4 h and annealing the mixture for 48 h at 650°C. A royal blue powder of pure $\alpha$-Li$_6$CuB$_4$O$_{10}$ was obtained, showing a particle size from 20 to 50 μm with no signs of impurities (Figure 38a).

Later in 2013, Kuratieva et al. isolated a crystal of another Li$_6$CuB$_4$O$_{10}$ polymorph denoted as $\beta$-Li$_6$CuB$_4$O$_{10}$, in exploring the Li$_2$O-CuO-B$_2$O$_3$ ternary system.\textsuperscript{172} Through the course of this thesis we found that bulk samples of $\beta$-Li$_6$CuB$_4$O$_{10}$ can be obtained from $\alpha$-Li$_6$CuB$_4$O$_{10}$ if the latter is annealed for at least 9 days at 500°C in air with a particle size distribution from 10 to 50 μm,.

![Figure 38: Representative SEM images of (a) $\alpha$-Li$_6$CuB$_4$O$_{10}$ and (b) $\beta$-Li$_6$CuB$_4$O$_{10}$. Insets show a magnified view on the crystallites and a colored picture of the as-prepared powder.](image)

The different color of both polymorphs ($\alpha$-Li$_6$CuB$_4$O$_{10}$: royal blue, $\beta$-Li$_6$CuB$_4$O$_{10}$: pale blue) let us consider a slightly different oxidation state of Cu as well a slightly different stoichiometry. To check both of these issues electron paramagnetic resonance (EPR)
spectroscopy was conducted (experimental details see annexes chapter 7.3.2). The room temperature EPR spectrum for both polymorphs are identical and show a signal for Cu$^{2+}$ (3d$^9$) of the same amplitude (Figure 39). Moreover the two α- and β-polymerm have the same stoichiometry as determined from inductive coupled plasma optical emission spectroscopy (ICP-OES) (Table 6).

**Table 6:** Theoretical and experimental values obtained from the ICP-OES elemental analysis of Li, Cu and B for α- and β- Li$_6$CuB$_4$O$_{10}$.

<table>
<thead>
<tr>
<th>Atom</th>
<th>theoretical stoichiometry</th>
<th>Wt% theoretical</th>
<th>Wt% exp. α-LCB</th>
<th>Stoichiometry exp. α-LCB</th>
<th>Wt% exp. β-LCB</th>
<th>Stoichiometry exp. β-LCB</th>
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</thead>
<tbody>
<tr>
<td>Li</td>
<td>6</td>
<td>13.5</td>
<td>12.1</td>
<td>5.4</td>
<td>12.1</td>
<td>5.4</td>
</tr>
<tr>
<td>Cu</td>
<td>1</td>
<td>20.6</td>
<td>20.4</td>
<td>1</td>
<td>20.6</td>
<td>1</td>
</tr>
<tr>
<td>B</td>
<td>4</td>
<td>14.02</td>
<td>13.5</td>
<td>3.9</td>
<td>13.7</td>
<td>3.9</td>
</tr>
<tr>
<td>O</td>
<td>10</td>
<td>51.87</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

*Figure 39:* EPR spectra recorded at room temperature for both polymorphs of Li$_6$CuB$_4$O$_{10}$.

The differences in terms of the crystal structure, relationship and electrochemical properties of the both polymorphs will be described consecutively.
3.1 Structure and polymorphism

Two structural models were reported for Li$_6$CuB$_4$O$_{10}$, the first one by Pan et al. (α-polymorph) and the second one by Kuratieva et al. (β-polymorph) both crystallizing in a triclinic unit cell. Their structural models were originally derived from single crystal diffraction and reported without any structural or synthetic relationship between them. For α-Li$_6$CuB$_4$O$_{10}$ the XRD pattern can be indexed using Pan’s triclinic cell, with the exception of weak peaks which could not be indexed suggesting a superstructure. This superstructural peaks (Figure 40, inset) were already mentioned in Sparta’s PhD thesis and explained in a structural model including a tripled unit cell. This structural model fits perfectly the recorded XRD pattern (Figure 40a) with lattice parameters listed in Table 7. Regarding β-Li$_6$CuB$_4$O$_{10}$ the XRD pattern could be proper fitted using the triclinic model described by Kuratieva et al. (Figure 40b) with lattice parameters shown in Table 8.

![Figure 40: Rietveld refinement of the synchrotron XRD pattern of (a) α- and (b) β- Li$_6$CuB$_4$O$_{10}$. The red circles, black line and the bottom grey line represent the observed, calculated and difference pattern respectively. Bragg positions are marked with blue vertical thick bars.](image)
**Lithium copper pyroborate Li\(_6\)CuB\(_4\)O\(_{10}\)**

**Table 7: Structural parameters and atomic positions of \(\alpha\)-Li\(_6\)CuB\(_4\)O\(_{10}\).**

<table>
<thead>
<tr>
<th>Atom</th>
<th>Wyckoff site</th>
<th>(x)</th>
<th>(y)</th>
<th>(z)</th>
<th>B(Å(^2))</th>
</tr>
</thead>
<tbody>
<tr>
<td>B1</td>
<td>2i</td>
<td>0.0852(19)</td>
<td>0.4463(9)</td>
<td>0.3336(7)</td>
<td>0.39(14)</td>
</tr>
<tr>
<td>B2</td>
<td>2i</td>
<td>0.1779(20)</td>
<td>0.6796(10)</td>
<td>0.4822(7)</td>
<td>0.65(16)</td>
</tr>
<tr>
<td>B3</td>
<td>2i</td>
<td>0.190(2)</td>
<td>0.6607(10)</td>
<td>0.1849(7)</td>
<td>1.31(19)</td>
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<tr>
<td>B4</td>
<td>2i</td>
<td>0.276(2)</td>
<td>0.8972(10)</td>
<td>0.3284(7)</td>
<td>1.15(19)</td>
</tr>
<tr>
<td>B5</td>
<td>2i</td>
<td>0.4190(20)</td>
<td>0.7715(9)</td>
<td>0.9908(7)</td>
<td>0.66(17)</td>
</tr>
<tr>
<td>B6</td>
<td>2i</td>
<td>0.4720(19)</td>
<td>0.0004(9)</td>
<td>0.1452(6)</td>
<td>0.20(15)</td>
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<tr>
<td>Cu1</td>
<td>1a</td>
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<td>0</td>
<td>0</td>
<td>0.55(3)</td>
</tr>
<tr>
<td>Cu2</td>
<td>2i</td>
<td>0.3000(2)</td>
<td>0.31614(10)</td>
<td>0.67137(8)</td>
<td>0.43(2)</td>
</tr>
<tr>
<td>Li1</td>
<td>2i</td>
<td>0.038(2)</td>
<td>0.8018(12)</td>
<td>0.8149(9)</td>
<td>-0.08(20)</td>
</tr>
<tr>
<td>Li2</td>
<td>2i</td>
<td>0.068(4)</td>
<td>0.3786(17)</td>
<td>0.0345(12)</td>
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<tr>
<td>Li3</td>
<td>2i</td>
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<td>0.0713(14)</td>
<td>0.8176(10)</td>
<td>2.5(3)</td>
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<tr>
<td>Li4</td>
<td>2i</td>
<td>0.221(3)</td>
<td>0.9290(15)</td>
<td>0.6293(10)</td>
<td>1.7(3)</td>
</tr>
<tr>
<td>Li5</td>
<td>2i</td>
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<td>0.1367(12)</td>
<td>0.4883(9)</td>
<td>0.3(2)</td>
</tr>
<tr>
<td>Li6</td>
<td>2i</td>
<td>0.368(3)</td>
<td>0.4314(12)</td>
<td>0.9148(9)</td>
<td>1.0(2)</td>
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<tr>
<td>Li7</td>
<td>2i</td>
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<td>0.1719(12)</td>
<td>0.3115(9)</td>
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<tr>
<td>Li8</td>
<td>2i</td>
<td>0.414(3)</td>
<td>0.5709(13)</td>
<td>0.6101(9)</td>
<td>1.8(3)</td>
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<tr>
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<td>2i</td>
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<td>0.3997(14)</td>
<td>0.1973(9)</td>
<td>1.9(3)</td>
</tr>
<tr>
<td>O1</td>
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<td>0.5885(5)</td>
<td>0.0998(4)</td>
<td>0.55(10)</td>
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<tr>
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<td>0.2575(5)</td>
<td>0.5419(3)</td>
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<td>O3</td>
<td>2i</td>
<td>0.1071(10)</td>
<td>0.7874(5)</td>
<td>0.2506(3)</td>
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<tr>
<td>O4</td>
<td>2i</td>
<td>0.1604(11)</td>
<td>0.8056(5)</td>
<td>0.9530(4)</td>
<td>0.74(10)</td>
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<tr>
<td>O5</td>
<td>2i</td>
<td>0.1579(10)</td>
<td>0.0242(5)</td>
<td>0.3609(3)</td>
<td>0.26(9)</td>
</tr>
<tr>
<td>O6</td>
<td>2i</td>
<td>0.1917(10)</td>
<td>0.5242(5)</td>
<td>0.6976(4)</td>
<td>0.47(9)</td>
</tr>
<tr>
<td>O7</td>
<td>2i</td>
<td>0.1846(9)</td>
<td>0.0295(5)</td>
<td>0.1315(3)</td>
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<tr>
<td>O8</td>
<td>2i</td>
<td>0.2232(10)</td>
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<td>0.40(9)</td>
</tr>
<tr>
<td>O9</td>
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<td>0.2392(11)</td>
<td>0.5498(5)</td>
<td>0.4203(4)</td>
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<tr>
<td>O10</td>
<td>2i</td>
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<td>0.5606(4)</td>
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<tr>
<td>O11</td>
<td>2i</td>
<td>0.3435(11)</td>
<td>0.9223(5)</td>
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<tr>
<td>O12</td>
<td>2i</td>
<td>0.4361(10)</td>
<td>0.1228(5)</td>
<td>0.9209(3)</td>
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<tr>
<td>O13</td>
<td>2i</td>
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<td>0.6251(4)</td>
<td>0.48(9)</td>
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<tr>
<td>O14</td>
<td>2i</td>
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<td>0.0435(4)</td>
<td>0.97(12)</td>
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<tr>
<td>O15</td>
<td>2i</td>
<td>0.4624(10)</td>
<td>0.6166(5)</td>
<td>0.2042(3)</td>
<td>0.24(8)</td>
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Table 8: Structural parameters and atomic positions of β-Li₆CuB₄O₁₀.

<table>
<thead>
<tr>
<th>Atom</th>
<th>Wyckoff site</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>B(Å²)</th>
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<tr>
<td>Cu1</td>
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<td>0</td>
<td>0</td>
<td>0.94(3)</td>
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<tr>
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<td>2i</td>
<td>0.6452(9)</td>
<td>0.2087(4)</td>
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<tr>
<td>O2</td>
<td>2i</td>
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<td>0.7959(4)</td>
<td>0.1849(3)</td>
<td>0.29(4)</td>
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<tr>
<td>O3</td>
<td>2i</td>
<td>0.6794(9)</td>
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<td>0.29(4)</td>
</tr>
<tr>
<td>O4</td>
<td>2i</td>
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<td>0.2653(3)</td>
<td>0.29(4)</td>
</tr>
<tr>
<td>O5</td>
<td>2i</td>
<td>0.9405(9)</td>
<td>0.7180(4)</td>
<td>0.4458(3)</td>
<td>0.29(4)</td>
</tr>
<tr>
<td>B1</td>
<td>2i</td>
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<td>0.2213(8)</td>
<td>0.2137(6)</td>
<td>0.47(9)</td>
</tr>
<tr>
<td>B2</td>
<td>2i</td>
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<td>0.8476(9)</td>
<td>0.3145(6)</td>
<td>0.47(9)</td>
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<tr>
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<td>2i</td>
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<td>0.63(11)</td>
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<tr>
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<td>2i</td>
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<td>0.3551(8)</td>
<td>0.63(11)</td>
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<tr>
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<td>2i</td>
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<td>0.6482(12)</td>
<td>0.1214(8)</td>
<td>0.63(11)</td>
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</tbody>
</table>

The basic building unit of both polymorphs can be described as a [Cu(B₂O₅)₂]⁶⁻ structural unit, which consists of a Cu²⁺ square planar coordinated by two pyroborate B₂O₅ groups (Figure 41). In the α-polymorph 2 independent Cu sites are present, contrary to the β-polymorph which contains only one crystallographic site for Cu.

Figure 41: View of the [Cu(B₂O₅)₂]⁶⁻ structural unit in (a) α-Li₆CuB₄O₁₀(Cu1 and Cu2) and (b) β-Li₆CuB₄O₁₀(Cu1). Copper, boron and oxygen atoms are shown in blue, green and red respectively.
These units are stacked independently in a parallel way on top of each other along crystallographic directions, with Li sitting in tetrahedral sites in between these units (Figure 42). The $\alpha$-polymorph presents 9 different Li-sites whereas in the $\beta$-polymorph Li occupies 3 different sites. The main difference between the two polymorphs is located in the way Li connects the $[\text{Cu}(\text{B}_2\text{O}_5)_2]^6^-$ units, which forms layers along the ac-plane in $\beta$-$\text{Li}_6\text{CuB}_4\text{O}_{10}$ compared to $\alpha$-$\text{Li}_6\text{CuB}_4\text{O}_{10}$ (Figure 42b, d).

![Figure 42: Structure of (a, b) $\alpha$- and (c, d) $\beta$- $\text{Li}_6\text{CuB}_4\text{O}_{10}$ along different crystallographic directions. Copper, boron and oxygen are drawn in blue, green and red. Li, Cu, B and O atoms are drawn in yellow, blue, green and red respectively.](image)

Although in the initial publication from Pan et al. no phase transition was observed with differential scanning calorimetry (DSC) for $\alpha$-$\text{Li}_6\text{CuB}_4\text{O}_{10}$, Sparta mentioned a reversible transition around 350°C. To check this point in detail DSC measurements coupled with thermogravimetric analysis (TGA) were measured with a STA 449C Netzsch (experimental details see annexes chapter 7.3.1). A heating rate of 2°C/min was applied under
flowing air using around 15-20 mg of the pristine material between room temperature and 500 or 650°C for the α- and β-polymorph respectively.

For α-Li₆CuB₄O₁₀ one large endothermic phase transition is visible during heating followed by a very tiny one at 346 and 372°C respectively. These transitions are perfectly reversible upon cooling since equivalent exothermic peaks are visible, suggesting a structural transition between a low temperature (LT) and a high temperature (HT) form of α-Li₆CuB₄O₁₀ (Figure 43a). Note that this reversible structural transition between an LT-α and a HT-α form was further supported by the complementary TGA measurement where only a negligible mass change was observed, most likely due to the removal of absorbed water during heating of the sample (Figure 43c).

Regarding β-Li₆CuB₄O₁₀ one broad endothermic phase transition is present with a maximum at 570°C upon heating (Figure 43b), whereas during cooling a smaller but well-defined exothermic peak shows up at 330°C. The huge temperature difference of the peaks in the DSC measurement during heating and cooling indicated two different structural transitions. Based on the observed color change of the sample from pale- to royal blue, we could assume a change from the β- to the α-polymorph on cooling. The structural nature of this transition was further confirmed through the TGA measurement whereas in the case of α-Li₆CuB₄O₁₀ again no mass change is observed (Figure 43d).
Figure 43: DSC and TGA curve for (a,c) α- and (b,d) β-Li₆CuB₄O₁₀ respectively. The black and the orange line are recorded during heating and cooling respectively. LT- and HT- refers to the low temperature- and high temperature form of α-Li₆CuB₄O₁₀.

To get more insights into the structural changes taking place during the phase transitions observed in DSC, in situ synchrotron XRD pattern were recorded (experimental details see annexes chapter 7.2.2) for α- and β-Li₆CuB₄O₁₀, during heating, and after the sample was cooled down to room temperature (Figure 44, 45). As seen in Figure 44, upon heating a gradual shift of the reflections around 3.5 and 5.1° 2θ is visible, till a striking jump at 350°C, where a sudden strong shift is observed. All the patterns can be indexed with the same triclinic unit cell as for the room temperature one, but with different lattice parameters $a=4.90122$ Å, $b=9.18486$ Å, $c=14.49619$ Å, $\alpha=105.5898^\circ$, $\beta=98.0462(11)^\circ$ and $\gamma=94.1337^\circ$ for the pattern recorded at T=500 °C. Note that after the 350°C phase transition the superstructure reflections between 5.8 and 6.8° disappear. This gives the possibility to index the patterns with the smaller unit cell reported by Pan. This corresponds to an increase of
Lithium copper pyroborate Li$_6$CuB$_4$O$_{10}$

the $a$ and $c$ lattice parameters and a decrease of $b$ as compared to the room temperature phase (Figure 45). Overall, the volume is enlarged by 4\% to reach 618.248 Å$^3$ at 500 °C (Figure 45a). All the changes in the XRD pattern are perfectly reversible if the sample is cooled down to room temperature, in agreement with the DSC measurements (Figure 43a). Note that we were not able to stabilize the HT α-Li$_6$CuB$_4$O$_{10}$ at RT by quenching from temperatures above 500°C even if liquid nitrogen was used.

Figure 44: In situ synchrotron XRD pattern for α-Li$_6$CuB$_4$O$_{10}$ during heating up to 500°C and after cooled down to 100°C. A global view is given in (a) while (b) and (c) highlight the grey marked areas.
Figure 45: Change in lattice parameters as a function of temperature for α-Li$_6$CuB$_4$O$_{10}$ during heating as derived from in situ high temperature XRD. The grey shaded bar marks the temperature region where the structural transition from the LT- to the HT α-Li$_6$CuB$_4$O$_{10}$ occurs. (a) Volume of the unit cell and angles shown in the inset. (b) Variation of unit cell parameters.

If β-Li$_6$CuB$_4$O$_{10}$ is gradually heated up to 600°C (Figure 46), first a slight shift of the reflections is observed up to 550°C coming along with the linear increase of the lattice parameters $a$, $b$, $c$ (Figure 47a) and a decrease of all angles (Figure 47c) followed by a transformation from the β- to the α-polymorph starting around 550°C. The XRD pattern at this temperature can be indexed with the same triclinic unit cell used at room temperature for β-Li$_6$CuB$_4$O$_{10}$, with increased lattice parameters $a$= 3.37791 Å, $b$= 6.73946 Å, $c$= 9.33539 Å, $\alpha$= 78.66432°, $\beta$= 89.15504° and $\gamma$= 85.41164° leading to a unit cell volume of 207.709 Å$^3$ which is ~3% enlarged compared to the one at 30°C. Already visible at 550°C, reflections of the HT α-polymorph start to show up and are constantly growing, becoming the major phase at 600°C. After the sample was cooled down to room temperature an almost pure LT α-Li$_6$CuB$_4$O$_{10}$ is obtained, confirming the phase transition from β to HT α around 550°C as seen in the corresponding DSC measurement (Figure 43b).
Figure 46: (a) In situ synchrotron XRD pattern for $\beta$-Li$_6$CuB$_4$O$_{10}$ during heating up to 600°C and after cooled down to room temperature with (b) a magnification of the grey shaded region marked in (a). The brown asterisks mark the appearance of reflections from the HT $\alpha$-Li$_6$CuB$_4$O$_{10}$ phase.

Figure 47: Change of the lattice parameters of $\beta$-Li$_6$CuB$_4$O$_{10}$ with temperature. (a) Shows the evolution of the unit cell volume, (b) unit cell edges and (c) the unit cell angles.

From the complementary XRD and DSC measurements, the relationship between the two known polymorphs of Li$_6$CuB$_4$O$_{10}$ could be established and a schematic overview is shown in Figure 48. From our experimental observations it seems like the direct synthesis of $\beta$-Li$_6$CuB$_4$O$_{10}$ at 500°C starting from the ternary Li$_2$O-CuO-B$_2$O$_3$ system is not possible or requires infinite long annealing times. However as described above, the $\beta$-polymorph could be obtained if $\alpha$-Li$_6$CuB$_4$O$_{10}$ is annealed for at least 9 days at temperatures not above 500°C. If $\beta$-Li$_6$CuB$_4$O$_{10}$ is exposed to temperatures higher than 550°C, a rapid biphasic phase transformation back to $\alpha$-Li$_6$CuB$_4$O$_{10}$ occurs. Regarding the $\alpha$-polymorph, whenever exposed...
to temperatures above 350°C a HT form is observed, which turns back into LT \( \alpha\)-Li₆CuB₄O₁₀ if the temperature drops below 350°C.

![Schematic representation of the synthetic relationship different polymorphs of Li₆CuB₄O₁₀.](image)

**Figure 48:** Schematic representation of the synthetic relationship different polymorphs of Li₆CuB₄O₁₀.

### 3.2 Electrochemical characterization

#### 3.2.1 Activity versus lithium

Both Li₆CuB₄O₁₀ polymorphs were tested versus Li in Swagelok type cells (previously mixed with 20 wt% carbon SP in a SPEX high energy miller in argon). The positive electrode was separated from the negative electrode through two sheets of glass fiber separator soaked with electrolyte (detailed description of the setup see annexes chapter 7.1).

Initially we used a 1M solution of LiPF₆ in a mixture of ethylene carbonate (EC), dimethyl carbonate (DMC) and propylene carbonate (PC) in a weight ratio of 1/1/3 respectively (commercial name LP100) as electrolyte. However, as we observed a fast degradation of the electrolyte during galvanostatic cycling, leading to dry cells after only a few cycles, we screened different electrolytes towards their oxidative stability. Two carbonate based electrolytes and one composed of a mixture of a 1M LiTFSI solution in the ionic liquid N-Propyl-N-methyl pyrrolidinium bis(trifluoro methane sulfonyl) imide (PYR₃TFSI) were tested. This was done by means of cyclic voltammetry of \( \alpha\)-Li₆CuB₄O₁₀/Li half cells with a scan speed of 0.1 mV·s⁻¹ in a potential window from 4.5 to 1.0 V starting on oxidation (Figure 49). One can clearly see a pronounced degradation for the carbonate based electrolytes, as the anodic currents is higher compared to the ionic liquid based one (Figure 49 inset). Such degradation is even more pronounced for the cathodic current. As a result of this electrolyte screening, the ionic liquid based electrolyte described above was selected for all electrochemical experiments further described in this section.
Lithium copper pyroborate Li₆CuB₄O₁₀

Figure 49: Cyclic voltammetry of α-Li₆CuB₄O₁₀/Li half cells in a potential window from 4.5 to 1.0 V during the first cycle.

A typical voltage composition curve of a α-Li₆CuB₄O₁₀/Li half-cell is shown in Figure 50. The voltage rapidly increases up to 3.8 V and then smoothly slopes up to the cutoff voltage of 4.5 V. The derivative plot of the first charge shows the presence of two oxidation peaks located at ~3.8 and ~4.25 V vs Li⁺/Li⁰ with the first one shifted by ~0.1 V during the following cycles. As only one single redox center is present in α-Li₆CuB₄O₁₀, the two peaks observed in the derivative curve can be most likely correlated to the Cu1 and Cu2 crystallographic site. Around 0.9 Li⁺ ions are removed on charge, part of which come from the structure and the rest being associated to electrolyte decomposition, however upon subsequent discharge only 0.3 Li⁺ are reinserted. During reduction the derivative curve shows two peaks at ~4.2 V and ~4.0 V vs Li⁺/Li⁰ which are remarkably more close together compared to the corresponding peaks on oxidation pointing towards an irreversible structural transformation during the first charge, however we don’t have a full explanation for this phenomenon yet. Furthermore the large amount of irreversible loss during the first and following cycles can be partially attributed to catalytic electrolyte decomposition as well as irreversible Li loss, since on subsequent cycles between 4.5 and 2.5 V (Figures 50) only 0.3 Li⁺ can be reversibly inserted leading to an reversible capacity of 20 mAh/g (Figure 50a inset) fading down about 50% during 10 cycles.
To confirm the redox activity versus Li being correlated to the Cu$^{3+}$/Cu$^{2+}$ redox couple, we decided to use EPR spectroscopy to monitor the Cu$^{2+}$ evolution since Cu$^{2+}$ (3d$^9$) is EPR active, whereas Cu$^{3+}$ (3d$^8$) is EPR silent. Measured spectra of the pristine- and cycled samples are shown in Figure 51 and show similar EPR spectra with axial symmetry ($g_\parallel=2.29$, $g_\perp=2.07$), signature of Cu$^{2+}$ (3d$^9$), and small sharp peaks ($g=2.00$) arising from the conductive carbon SP (delocalized electrons) (experimental details see annexes chapter 7.3.2).\textsuperscript{179} After oxidation to 4.5 V a decrease of the Cu$^{2+}$ signal is observed (~50%), indicating the formation of Cu$^{3+}$. On subsequent reduction to 2.5 V, a part of the initial intensity (~80%) is recovered suggesting an incomplete reversibility of the Cu$^{3+}$/Cu$^{2+}$ redox process as suggested by galvanostatic measurements.
Figure 51: Room temperature EPR spectra of \( \alpha\)-Li\(_6\)CuB\(_4\)O\(_{10} \) ball-milled with carbon at pristine (black), charged (4.5 V, red) and discharged (2.5 V, purple) states.

To rationalize such a Cu\(^{3+}/\)Cu\(^{2+} \) redox couple, density functional calculations (DFT+\( U \)) were performed (detailed description see annexes chapter 7.3.6).\(^{180} \) First of all, using a \( U \) parameter at 4 eV for copper (correction factor taken from literature for Cu d-electrons),\(^{180} \) the structure of \( \alpha\)-Li\(_6\)CuB\(_4\)O\(_{10} \) could be well reproduced with less than 2% difference. By considering different delithiation paths from the pristine material, a potential at 3.6 V was obtained, which is about 0.3 to 0.5 V lower than the experimental one, however a pronounced distortion of the CuO\(_4\) square plane is observed for the delithiated phase after the structure was relaxed (Figure 52). This distortion was only accessible from DFT calculations, since we were not able to refine the atomic positions from our \textit{in situ} XRD experiment (Figure 53). To confirm the possibility of having Cu\(^{3+} \) at 4.5 V, the projected density of states (DOS) and crystal orbital overlap population (COOP) of \( \alpha\)-Li\(_6\)CuB\(_4\)O\(_{10} \) were calculated (Figure 52a). Clearly, the band below the Fermi level, involved in the oxidation process, has an important oxygen contribution and a Cu-O anti-bonding character as seen in the COOP. To probe the redox process the hole Fukui function of \( \alpha\)-Li\(_6\)CuB\(_4\)O\(_{10} \) was computed (difference in the charge density induced by the removal of electrons in Li\(_6\)CuB\(_4\)O\(_{10} \), Figure 52b) and the electron Fukui function of the Li\(_3\)CuB\(_4\)O\(_{10} \) phase as found by DFT calculation (difference in the charge density induced by the addition of electrons in Li\(_3\)CuB\(_4\)O\(_{10} \), Figure 52c). The comparison of both Fukui functions together with Highest Occupied Molecular Orbital (HOMO) of square planar CuO\(_4\) (Figure 52d) indicates that the distortion of the Cu square plane upon delithiation allows an orbital mixing between Cu d orbitals and the non-bonding O
p orbitals (which is forbidden in a regular square planar due to symmetry). This distortion re-equilibrates the weight of Cu/O in the redox band in order to involve primarily Cu in the redox process and stabilizes oxygen atoms in the structure.\textsuperscript{181}

\textbf{Figure 52:} (a) Cu- and O-projected density of states (DOS) and Crystal orbital overlap population (COOP) for the $\alpha$-Li$_6$CuB$_4$O$_{10}$ phase. The bands lying just below the Fermi level (at 0 eV) show an important O(2p) contribution and a strong Cu-O covalency as highlighted by the computed hole-Fukui function (b) shown in the CuO$_4$ square plane. Blue and red spheres represent copper and oxygen atoms respectively; the yellow volume corresponds to electron density depletion (holes) while the small cyan regions correspond to electron density accumulation (electrons) due to orbital polarization. (c) Electron-Fukui function computed in the Li$_5$CuB$_4$O$_{10}$ phases where the Cu square plane is distorted. The yellow volume corresponds to electron density accumulation (electrons) while the small cyan regions correspond to electron density depletion (hole) due to orbital polarization. It shows a more pronounced implication of the metal thanks to the distortion. (d) Illustration of the local electronic levels involved in the redox band of the Li$_6$CuB$_4$O$_{10}$ system, the Highest Occupied Molecular Orbital (HOMO).
To follow the structural changes during Li uptake/removal in $\alpha$-Li$_6$CuB$_4$O$_{10}$, \textit{in situ} XRD was performed using a homemade electrochemical cell equipped with a beryllium window transparent to X-rays (experimental details see annexes chapter 7.1).\textsuperscript{182} XRD patterns were collected for every 0.05 Li removed/inserted (Figure 53). During charge no major changes with the exception of i) two reflections centered around 19.50° which merge gradually together (Figure 53b), and of ii) a reflection at 27.75° is shifted towards 28.0° (Figure 53b).

On the subsequent discharge down to 2.5 V no reverse changes can be observed (Figure 53a, green XRD pattern). Only if continuing the discharge from 2.5 to 1.0 V, a plateau around 1.5 V appears, leading to the insertion of additional Li-ions and the complete reverse of the \textit{in situ} XRD patterns. More specifically the XRD pattern of the phase charged to 4.5 V can be indexed with the same triclinic cell as pristine Li$_6$CuB$_4$O$_{10}$ but with different lattice parameters (Table 9), since there is a contraction of the $a$ axis and an extension of the $b$ and $c$ axis. During Li reinsertion, the unit cell volume is slightly enlarged if compared to the pristine value (599 Å$^3$ versus 595 Å$^3$, see Table 9), which is consistent with the nominal composition of material at the end of discharge to 1 V of “Li$_{6.5}$CuB$_{4}$O$_{10}$” (see Figure 53a). In general the Li-extraction/insertion in $\alpha$-Li$_6$CuB$_4$O$_{10}$ represents a reversible biphasic processes from a structural point of view.
Figure 53: (a) In situ XRD patterns during charge and subsequent discharge in relation to the voltage versus time curve. (b) The reversible changes of the reflections indicated centered around 19.5 and 25.0°.

Table 9: Lattice parameters and the corresponding voltage of α-Li₆CuB₄O₁₀ during galvanostatic charge and following discharge.

<table>
<thead>
<tr>
<th></th>
<th>a (Å)</th>
<th>b (Å)</th>
<th>c (Å)</th>
<th>α (°)</th>
<th>β (°)</th>
<th>γ (°)</th>
<th>V (Å³)</th>
</tr>
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<tbody>
<tr>
<td>Discharged 1.0V</td>
<td>4.829(5)</td>
<td>9.251(5)</td>
<td>14.021(5)</td>
<td>104.36(5)</td>
<td>96.538(5)</td>
<td>94.472(5)</td>
<td>599.244(5)</td>
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</tbody>
</table>
To further explore the electrochemical behavior, Li/Li$_6$CuB$_4$O$_{10}$ cells were either started on oxidation or reduction using a C/50 rate. As shown in Figure 5a,c, similar curves are observed for both independently of what we started on charge or discharge. Equally both cells show similar capacity fading upon subsequent cycling (Figure 5b,d inset). Between 4.5 and 2.5 V. Turning to the lower voltage part of the curve note that there is the appearance of a plateau around 1.5 V which smoothly decreased down to 1 V with the concomitant uptake of additional Li. Nearly around 1 Li is additionally inserted around 1.5 V if the cell is initially started in discharge (Figure 5c), however the corresponding derivative plots represent similar voltages for the oxidation/ reduction process (Figure 5b,d). The inserted Li can be fully removed when the cell is charged back to ~4.0 V with additional 0.7 Li being removed when the cell voltage is further increased to 4.5 V (Figure 5a,c). Although not fully understood yet, we believe that this low voltage process is more related to a Li-driven conversion than insertion/ deinsertion mechanism in agreement with the absence of additional Bragg peaks in the in situ XRD when the cell is discharged.
Figure 54: Voltage composition curve (a) and (c) and the corresponding derivative plot (b) and (d) for galvanostatic cycling of $\alpha$-Li$_6$Cu$_4$B$_4$O$_{10}$ in a potential window from 4.5 to 1.0 V started either upon oxidation or reduction. The specific charge and discharge capacities versus cycle number are given in the insets of (b) and (d).

At this stage it is worth to mention the observation of a reddish color in the inside of the cells disassembled cycled down to 1 V for a few cycles. We hypothesize the formation of elemental copper. To check this issue, cells which have been cycled for 5 times were disassembled within the dry box in their discharge state and the recovered powders were washed with DMC, dried in vacuum and examined by TEM (experimental details see annexes chapter 7.3.5). Figure 55 clearly indicates the appearance of nanograins (size between 1 and 5 nm, too small to be detected by XRD), distributed over the surface of Li$_6$Cu$_4$B$_4$O$_{10}$ carbon composite particles. The chemical nature of these nanograins was identified as pure Cu by both, single area electron diffraction (SAED) and energy dispersive X-ray spectroscopy (EDS) measurements. This extrusion is most likely responsible for the fast capacity fading (90% fading during 15 cycles, Figure 54 b,d inset) leading to a partial amorphization of the material as deduced by XRD.
Figure 55: TEM image of α-Li₆CuB₄O₁₀ carbon composite discharged to 1.0 V. (a) shows an overview of agglomerated particles whereas (b) highlights one particle where copper nanograins can be seen on the surface, as confirmed by the corresponding EDS mapping shown below for copper (pale blue) and oxygen (red).

Turning to β-Li₆CuB₄O₁₀, a typical voltage composition curve is shown in Figure 56. After a quick rise to 3.8 V, the curve smoothly slopes up to 4.5 V, showing one well defined oxidation domain centered around 4.2 V as seen in the derivative plot (Figure 56b) most likely associated to the single Cu crystallographic site. Initially ~0.8 Li⁺ can be extracted and only 0.2 Li⁺ can be reinserted around 3.8 V suggesting irreversible Li extraction owing to copious electrolyte oxidation. During the following cycle only 0.4 Li⁺ can be extracted and again 0.2 Li⁺ can be reinserted, however the onset of the oxidation potential is shifted up around 0.5 V and the reduction potential is shifted down to 3.6 V, indicating irreversible structural modifications during the first cycle.
3.2.2 Ionic conductivity

The poor electrochemical activity of this phase is somewhat surprising because it consists of an open structure and possesses a large amount of Li with additionally absence of thermodynamic limitations as deduced by DFT. This led us to hypothesize that the phase poor electrochemical activity was mainly due to poor ionic conductivity of Li$^+$ in Li$_6$CuB$_4$O$_{10}$. To check the capability of the structure towards Li conduction, bond valence energy landscapes (BVEL) were generated for the α- and β-polymorph, based on the methodology developed by S. Adams (for details see annexes chapter 7.3.7). Figure 57 shows the calculated BVEL maps representing the Li$^+$ conduction pathways, taking into account the polarizability of the Li$^+$ species and the influence of the counterions of the structure considering anionic neighbors up to 8 Å at 1.0 eV above the minimum energy (Figure 57). One can first note a huge difference between the LT-α and β-polymorph, whereas the latter one shows only limited 2D pathways in contrast to LT-α where a 3D Li conduction network is visible. Regarding the HT form of α-Li$_6$CuB$_4$O$_{10}$, clearly a 3D network is visible slightly enhanced compared to the LT form. The results from BVEL calculations suggest an ionic conductivity which is increased in the order from β << LT-α < HT-α.
Figure 57: Bond valence energy landscape (BVEL) for the different polymorphs of Li$_6$CuB$_4$O$_{10}$, LT- and HT refers to the Low- and high-temperature form of α-Li$_6$CuB$_4$O$_{10}$. The energy values chosen for the plots are 1.0 eV above the minimum energy. Iso-energy yellow domains indicate paths for Li conduction. Slight differences in the Li conduction pathways between the LT and HT form of α-Li$_6$CuB$_4$O$_{10}$ are marked with brown arrow and circles.

To confirm the predictions given by the BVEL calculations, combined AC and DC impedance measurements were conducted on a pellet with ionically blocking platinum electrodes in a temperature range of 150 up to 500°C in air. A pellet suitable for the conductivity measurement was prepared according to the following procedure. Stoichiometric amounts of Li$_2$CO$_3$, CuO and H$_3$BO$_3$ were ground with mortar and pestle and heated up to 450°C for 6 h in air. After regrinding, the powder was pressed into a pellet with 13 mm diameter and a thickness of ca. 1 mm by applying approximately 3 t pressure. The pellet was then sintered for 48 h at 650°C in air, and placed in the apparatus (detailed description of the setup please see annexes chapter 7.1.3). The obtained density was around 65% of the theoretical density.
First of all, the AC impedance spectra were recorded between 140 and 400°C with a voltage amplitude of 10 mV in a frequency range from 30 MHz to 0.1 Hz. Both Li₆CuB₄O₁₀ polymorph spectra present a nearly perfect semi-circle with a linear Warburg diffusion resistance tail at low temperatures (~180°C) (Figure 58a, b). With increased temperature the shape of the spectra is preserved for the β-polymorph as opposed to the α-polymorph where for temperatures higher than 350°C, an almost straight line with a slope ≈45° is visible (Figure 58c), indicating an unexpected high ionic conductivity after the structural transition.

The depressed semi-circles derived from the complex AC impedance spectra were fitted by an R₀ - R₁/CPE₁ (Figure 58 a,b inset) equivalent circuit, where R₀ represents an initial shift of impedance arc from zero, R₁ is the sum of the grain boundary and bulk resistance and CPE₁ is a constant phase element.

To clearly separate the electronic from the ionic contribution to the total conductivity of this material, DC polarization measurements on Li₆CuB₄O₁₀ samples were done (experimental details see chapter 7.1.4). After applying a potential across the pellet and following the current response, a sharp increase followed by a fast decay to a steady state value was observed (Figure 59), representing the pure electronic conduction of the material. The electronic resistance was then calculated according to Ohm’s law (equation 6).

\[ R = \frac{U}{I} \] (6)
Lithium copper pyroborate Li₆CuB₄O₁₀

From the experimental values for the AC and DC resistance, the corresponding conductivity σ was calculated (details see annexes chapter 7.1.3) and summarized in the Arrhenius plot shown in Figure 60. For α-Li₆CuB₄O₁₀ the values for conductivity increase from 3·10⁻⁸ to 1.4·10⁻² S·cm⁻¹ as the temperature raises from 140 and 500°C. This increase is not continuous but enlists two different regimes; a low temperature and a high temperature one, corresponding to activation energies of 1.07 and 0.52 eV respectively. Both regimes referring to the LT-α and HT-α polymorph, are separated by a strikingly jump in conductivity between 320 and 370°C, the same temperature range over which a structural transition was previously shown to occur by both, DSC and synchrotron XRD experiments. In contrast, the conductivity for β-Li₆CuB₄O₁₀ increases linearly from 5.5·10⁻¹⁰ to 7.9·10⁻⁷ S·cm⁻¹ for 180 and 390°C respectively and is associated to an activation energy of 0.92 eV. A linear extrapolation of log (σ) = f(1000/T) at 25°C gives room temperature ionic conductivities in the range of 10⁻¹³ S·cm⁻¹ for the both polymorphs, explaining their similar poor electrochemical activity versus lithium.

Regarding electronic conductivity, the values for α-Li₆CuB₄O₁₀ versus temperature slightly increase linearly from 2.3·10⁻⁷ to 1.1·10⁻⁶ S·cm⁻¹ for 172 and 492°C, respectively, related to an activation energy of 0.17 eV (Figure 60). For β-Li₆CuB₄O₁₀ the electronic conductivity rises more significant from 5.3·10⁻¹⁰ to 1.1·10⁻⁸ S·cm⁻¹ as the temperature raises from 180 to 390°C respectively with an activation energy of 0.57 eV.

The difference between the ionic- and electronic part of the conductivity for both Li₆CuB₄O₁₀ polymorphs indicates that the strong increase in conductivity we measured (especially for α- Li₆CuB₄O₁₀) reflects the material’s ionic conductivity. Overall, the low

Figure 59: (a) DC polarization of a pellet of α-Li₆CuB₄O₁₀ with 1.0 V at 172°C using Pt-electrodes.

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room temperature ionic conductivity for both polymorphs imply a low Li-ion mobility which partially explains the encountered difficulties to reversibly remove Li from this material. The ionic conductivity of the HT-α, which is $10^{-2}$ S·cm$^{-1}$ could be quite attractive for applications if we could achieve such values at RT.

Figure 60: Arrhenius plot of the AC- and DC conductivity of α- and β-\(\text{Li}_6\text{CuB}_4\text{O}_{10}\) and a schematic view of the sample pellet between two ionically blocking Pt-electrodes used for AC- as well as for DC measurements.

This calls for further effort in lowering the transition temperature. Work on this direction was realized via chemical substitution of Cu (through Al, Mg, etc.), but without consequent success.
3.3 Conclusion

In this section Li₆CuB₄O₁₀ was initially chosen as a model compound to show that elevated redox potentials can be achieved within the Li 3d metal borate family. Initially TGA-DSC and in situ synchrotron XRD established the relationship between the α- and β-polymorph of Li₆CuB₄O₁₀. In terms of synthesis and structure we show that both polymorphs react towards Li around 4.25 and 3.9 V for α- and β-Li₆CuB₄O₁₀ respectively. Such redox activity associated to the Cu³⁺/Cu²⁺ redox couple, lying well above lithium copper oxides (Li₂CuO₂ ≈3.1 V, LiCuO₂ ≈3.0 V),¹⁸⁴ but below the one of lithium copper oxysulfate Li₂Cu₂O(SO₄)₂ (4.7 V) whose potential is governed by anionic redox processes (Figure 61a).¹⁸⁵,¹⁸⁶ The participation of the Cu³⁺/Cu²⁺ redox couple in the insertion/ deinsertion reaction of Li between 4.5 and 2.5 V was confirmed by EPR spectroscopy and rationalized through DFT calculations. As the voltage window is enlarged down to 1.0 V, a conversion type reaction takes place, leading to the extrusion of copper nanograins as observed with TEM. Moreover the limitation of the electrochemical performance of these materials, were addressed using AC impedance spectroscopy and DC polarization with complementary BVEL calculations, revealing strong kinetic limitations as seen from the low ionic/ electronic conductivity at RT in combination with high activation energies ~1.0 eV. Surprisingly due to a structural transition around 350°C, α-Li₆CuB₄O₁₀ shows a high ionic conductivity of 1.4 mS·cm⁻¹ at 500°C with a low activation energy of 0.52 eV, comparable to other solid state Li-ion conductors (Garnet-type Li₇La₃Zr₂O₁₂, lithium borohydride LiBH₄ and lithium yttrium borate Li₆Y(BO₃)₃, Figure 61b).¹⁸⁷–¹⁹⁰ This unexpected finding may renew the interest in borate based ionic conductors, prompting to shift the phase transition to lower temperatures applying suitable doping strategies etc.
Figure 61: (a) Schematic illustration of the Cu$^{3+}$/Cu$^{2+}$ redox potentials versus Li for different compounds including α- and β-\( \text{Li}_6\text{CuB}_4\text{O}_{10} \). Note that the potential in quotation marks was derived from DFT calculations but never experimentally observed. (b) Comparison of the ionic conductivity of α-\( \text{Li}_6\text{CuB}_4\text{O}_{10} \) with different borate based materials and a Garnet type solid electrolyte.
4 Synthesis, structural and electrochemical properties of sodium transition metal pentaborates \( \text{Na}_3\text{MB}_5\text{O}_{10} \) (\( M = \text{Fe}, \text{Co} \))

4.1 Synthesis and structure

In the previous chapter we have demonstrated through the synthesis of \( \text{Li}_6\text{CuB}_4\text{O}_{10} \) the feasibility to obtain high potentials in borate compounds. Therefore at the same time we show the difficulty in making new pyroborates. This was an impetus to explore other borate anions, hence our interest for the family of sodium metal pentaborates \( \text{Na}_3\text{MB}_5\text{O}_{10} \). Solid state synthesis and structures have been reported to prepare these phases with \( M = \text{Mg}, \text{Ca}, \text{Zn} \). Their preparation consists briefly in mixing stoichiometric amounts of \( \text{Na}_2\text{CO}_3 \), \( \text{M(II)carbonate/nitrate} \) and \( \text{H}_3\text{BO}_3 \) followed by annealing of the mixture at temperatures between 650-700°C in air for several days.\(^{191,192}\) Our first attempts by replacing the \( \text{M(II)}- \)carbonate/nitrate through \( \text{Co/Fe(II)-oxalate} \) and heating the mixture under argon to avoid oxidation of the transition metal were not successful. Using \( \text{NaOH} \) instead of \( \text{Na}_2\text{CO}_3 \), led to the formation of \( \text{Na}_3\text{FeB}_5\text{O}_{10} \), but was again not successful in the case of \( \text{Co} \). \( \text{Na}_3\text{CoB}_5\text{O}_{10} \) could solely be obtained if \( \text{Co(OH)}_2 \) was used as precursor. If the oxalate was used, we always end up with \( \text{Co metal} \) due to the easy reduction of \( \text{Co}^{2+} \) in Co-oxalate if heated under argon. Since the precursors are prone to react with ambient atmosphere, all steps of the synthesis were carried out in an argon filled glove box or under argon flow in a tube furnace. All chemicals were stored in argon, sodium hydroxide \( \text{NaOH} \) and boric acid \( \text{H}_3\text{BO}_3 \) were dried prior to be used at 200°C for 4 h in vacuum, and at 55°C for 24 h in air respectively.

In general, stoichiometric amounts of \( \text{NaOH} \), \( \text{Fe(C}_2\text{O}_4)\cdot2\text{H}_2\text{O} \) or \( \text{Co(OH)}_2 \) and \( \text{H}_3\text{BO}_3 \) were mixed together for 15 min in argon using a SPEX milling apparatus. We observed that the mechanochemical process causes most likely a reaction between the precursors which solidified within the milling jar, yielding in an amorphous intermediate product. The resulting solid was then reground with a mortar and pestle, and heated with a rate of 10°C/min up to 700°C for 1 h under argon flow. Once the synthesis is finished, the formed product \( \text{Na}_3\text{FeB}_5\text{O}_{10} \) was immediately transferred to the glove box for further use, since we observed degradation in ambient atmosphere. At the opposite, \( \text{Na}_3\text{CoB}_5\text{O}_{10} \) is stable in air. Both
compounds possess particle sizes ranging from 1 to 50 μm as shown in Figure 62 with grey and blue color for M=Fe and Co respectively.

![Figure 62: SEM images of (a) Na₃FeB₅O₁₀ and (b) Na₃CoB₅O₁₀.](image)

The structure of Na₃FeB₅O₁₀ was examined by means of synchrotron XRD (Figure 63a). Besides a few weak peaks attributed to an Fe₃BO₅ impurity (~4%), all peaks could be perfectly indexed in an orthorhombic unit cell having a Pbcα space group with lattice parameters a= 7.95493(1) Å, b= 12.30669(1) Å and c= 18.03978(1) Å and a unit cell volume V=1766.073 Å³ (Table 10), based on a structural model reported for Na₃MgB₅O₁₀. All atoms are placed in general positions 8c with one crystallographic site for iron, three for sodium, 5 for boron and 10 for oxygen. The final atomic positions are given in Table 10 and their atomic positions within a unit cell are illustrated in Figure 64a,b. Iron atoms are located in the center of slightly distorted tetrahedra (Fe-O distances ranging from 1.970 to 2.116 Å) with an average bond length of 2.0243 Å, in well agreement with reported Fe-O in tetrahedral oxygen coordination.
Figure 63: Synchrotron XRD pattern and the corresponding Rietveld refinement of (a) Na₃FeB₅O₁₀ and (b) Na₃CoB₅O₁₀. The red crosses, black-, and green line represent the observed, calculated and difference patterns, respectively. The positions of the Bragg reflections are shown as vertical blue bars. For Na₃FeB₅O₁₀, the lower set of tick marks is Fe₃BO₅ (~4 w%).

All four oxygen ligands of the FeO₄ tetrahedra belong to terminal oxygen atoms of distinct pentaborate anions [B₅O₁₀]⁵⁻, which consists of one BO₄ tetrahedra (average bond length 1.4799 Å) surrounded by 4 trigonal planar BO₃ groups forming pyroborate units [B₂O₅]⁺ (Figure 64). The FeO₄ tetrahedra are connected through the pentaborate anions to form a 2D layer along the ab-plane (Figure 64c). These layers are stacked along the c-axis with sodium atoms sitting between the layers as well as in channels along the a-axis.
Table 10: Structural parameters and atomic positions of Na₃FeB₅O₁₀.

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<tr>
<th>Atom</th>
<th>Wyckoff site</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>B (Å²)</th>
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<td>8c</td>
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<td>0.0800(3)</td>
<td>1.97(7)</td>
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<td>8c</td>
<td>0.5902(11)</td>
<td>0.1222(5)</td>
<td>0.0472(4)</td>
<td>1.53(6)</td>
</tr>
<tr>
<td>O4</td>
<td>8c</td>
<td>0.7243(9)</td>
<td>0.2498(6)</td>
<td>0.1277(3)</td>
<td>1.53(6)</td>
</tr>
<tr>
<td>O5</td>
<td>8c</td>
<td>0.8771(9)</td>
<td>0.1416(5)</td>
<td>0.0435(3)</td>
<td>1.53(6)</td>
</tr>
<tr>
<td>O6</td>
<td>8c</td>
<td>0.5657(8)</td>
<td>0.2707(5)</td>
<td>0.2427(4)</td>
<td>1.53(6)</td>
</tr>
<tr>
<td>O7</td>
<td>8c</td>
<td>0.5371(8)</td>
<td>0.3980(5)</td>
<td>0.1466(3)</td>
<td>1.53(6)</td>
</tr>
<tr>
<td>O8</td>
<td>8c</td>
<td>0.4969(8)</td>
<td>0.4514(5)</td>
<td>0.2724(4)</td>
<td>1.53(6)</td>
</tr>
<tr>
<td>O9</td>
<td>8c</td>
<td>0.5762(10)</td>
<td>0.3344(5)</td>
<td>0.3713(3)</td>
<td>1.53(6)</td>
</tr>
<tr>
<td>O10</td>
<td>8c</td>
<td>0.4102(9)</td>
<td>0.5690(5)</td>
<td>0.1844(3)</td>
<td>1.53(6)</td>
</tr>
</tbody>
</table>
The Fe-Mössbauer spectrum for as synthesized Na$_3$FeB$_5$O$_{10}$ is shown in Figure 65 (details see annexes chapter 7.3.3). The majority of the Fe signal in Na$_3$FeB$_5$O$_{10}$ can be fitted with a single doublet, confirming the occurrence of one single crystallographic site for Fe in Na$_3$FeB$_5$O$_{10}$. The isomer shift IS = 1.02 mm/s and a quadrupole splitting QS = 2.10 mm/s are typical of Fe$^{2+}$ in tetrahedral coordination.$^{19,20}$ Note that the Mössbauer spectrum contains about 4% of Fe$_3$BO$_5$ impurity, in agreement with the values obtained from the XRD pattern (Figure 63a).

Regarding Na$_3$CoB$_5$O$_{10}$, the corresponding synchrotron XRD pattern and the Rietveld refinement are shown in Figure 63b. All peaks could be perfectly indexed using a structural model reported for KCdB$_5$O$_{10}$ by replacing K and Cd through Na and Co respectively.$^{194}$ The
Na$_3$CoB$_5$O$_{10}$ compound crystallizes in a monoclinic space group $P2_1/n$ with lattice parameters $a = 6.65143(1)$ Å, $b = 18.20986(1)$ Å, $c = 7.80513(1)$ Å and $\beta = 114.79220(1)^\circ$ resulting in a unit cell volume $V = 858.240$ Å$^3$ (Table 11). The atomic positions within the unit cell are shown in Figure 66. All atoms are placed in general position 4$e$ with one crystallographic site for cobalt, three for sodium, five for boron and ten for oxygen (refined atomic positions are given in Table 11). Cobalt is in a slightly distorted tetrahedral oxygen environment with average Co-O bond lengths of 1.98 Å in agreement with values reported in the case of other compounds possessing Co$^{2+}$ in tetrahedral environment.$^{195}$ All four oxygen ligands belong to terminal oxygen atoms of distinct pentaborate anions $[\text{B}_5\text{O}_{10}]^{5-}$, connecting the CoO$_4$ tetrahedra to form a 2D layer along the $ac$-plane. The layers are stacked along the $b$-axis with sodium atoms placed in between the layers as well as in channels.

Figure 66: Structure of Na$_3$CoB$_5$O$_{10}$ viewed along different crystallographic directions. Oxygen, boron, sodium and cobalt atoms indicated in red, green, yellow and blue respectively.
### Table 11: Structural parameters and atomic positions of Na₃CoB₅O₁₀

<table>
<thead>
<tr>
<th>Atom</th>
<th>Wyckoff site</th>
<th>x</th>
<th>y</th>
<th>Z</th>
<th>B (Å²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co1</td>
<td>4e</td>
<td>0.4938(2)</td>
<td>0.09529(6)</td>
<td>0.12100(16)</td>
<td>0.97(3)</td>
</tr>
<tr>
<td>Na1</td>
<td>4e</td>
<td>1.0215(5)</td>
<td>0.06927(15)</td>
<td>0.1537(4)</td>
<td>1.31(7)</td>
</tr>
<tr>
<td>Na2</td>
<td>4e</td>
<td>0.5356(5)</td>
<td>0.07253(15)</td>
<td>-0.2967(4)</td>
<td>2.25(9)</td>
</tr>
<tr>
<td>Na3</td>
<td>4e</td>
<td>0.2007(5)</td>
<td>0.24423(14)</td>
<td>-0.0962(4)</td>
<td>0.85(3)</td>
</tr>
<tr>
<td>B1</td>
<td>4e</td>
<td>0.7050(15)</td>
<td>0.2127(5)</td>
<td>-0.0092(14)</td>
<td>0.7(2)</td>
</tr>
<tr>
<td>B2</td>
<td>4e</td>
<td>0.1192(16)</td>
<td>0.0766(4)</td>
<td>-0.2204(12)</td>
<td>0.5(2)</td>
</tr>
<tr>
<td>B3</td>
<td>4e</td>
<td>0.1200(19)</td>
<td>0.1643(6)</td>
<td>-0.4532(14)</td>
<td>1.9(3)</td>
</tr>
<tr>
<td>B4</td>
<td>4e</td>
<td>0.2734(16)</td>
<td>0.1881(6)</td>
<td>0.3233(13)</td>
<td>1.7(3)</td>
</tr>
<tr>
<td>B5</td>
<td>4e</td>
<td>0.8089(15)</td>
<td>0.0738(5)</td>
<td>0.4833(12)</td>
<td>0.52(20)</td>
</tr>
<tr>
<td>O1</td>
<td>4e</td>
<td>0.3823(8)</td>
<td>0.3659(3)</td>
<td>-0.0583(6)</td>
<td>0.85(3)</td>
</tr>
<tr>
<td>O2</td>
<td>4e</td>
<td>0.6864(9)</td>
<td>0.1334(2)</td>
<td>0.0034(6)</td>
<td>0.85(3)</td>
</tr>
<tr>
<td>O3</td>
<td>4e</td>
<td>0.6234(8)</td>
<td>0.0399(2)</td>
<td>0.3567(6)</td>
<td>0.85(3)</td>
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<tr>
<td>O4</td>
<td>4e</td>
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<td>0.1274(3)</td>
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<td>0.85(3)</td>
</tr>
<tr>
<td>O5</td>
<td>4e</td>
<td>0.2347(8)</td>
<td>0.1406(2)</td>
<td>-0.5576(7)</td>
<td>0.85(3)</td>
</tr>
<tr>
<td>O6</td>
<td>4e</td>
<td>0.2214(8)</td>
<td>0.0375(2)</td>
<td>-0.0479(6)</td>
<td>0.85(3)</td>
</tr>
<tr>
<td>O7</td>
<td>4e</td>
<td>0.1189(7)</td>
<td>0.2431(3)</td>
<td>-0.4250(6)</td>
<td>0.85(3)</td>
</tr>
<tr>
<td>O8</td>
<td>4e</td>
<td>1.0892(9)</td>
<td>-0.0428(2)</td>
<td>0.3357(7)</td>
<td>0.85(3)</td>
</tr>
<tr>
<td>O9</td>
<td>4e</td>
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<td>0.1712(2)</td>
<td>0.1747(6)</td>
<td>0.85(3)</td>
</tr>
<tr>
<td>O10</td>
<td>4e</td>
<td>-0.2188(8)</td>
<td>0.2304(2)</td>
<td>-0.1428(7)</td>
<td>0.85(3)</td>
</tr>
</tbody>
</table>

Regarding the structures of Na₃MB₅O₁₀ (M = Fe, Co) described above, clearly significant similarities despite their different space groups are obvious. Note that the lattice basis vectors of both structures are related through the following matrix:

$$
\begin{pmatrix}
a_{Fe} \\
b_{Fe} \\
c_{Fe}
\end{pmatrix} = 
\begin{pmatrix}
1 & 0 & 1 \\
0 & 1 & 0 \\
-1 & 0 & 1
\end{pmatrix}
\begin{pmatrix}
a_{Co} \\
b_{Co} \\
c_{Co}
\end{pmatrix}
$$
The thermal stability of Na$_3$FeB$_5$O$_{10}$ and Na$_3$CoB$_5$O$_{10}$ was investigated using TGA-DSC analysis (details see annexes chapter 7.3.1). Measurements were recorded during heating a sample of around 20-30 mg under flowing air from RT to 850°C with a rate of 5°C/min (Figure 67a,b). For Na$_3$FeB$_5$O$_{10}$ a broad exothermic peak is observed around 640°C in the DSC curve, which is most likely associated with the decomposition of Na$_3$FeB$_5$O$_{10}$ into iron oxide Fe$_2$O$_3$ and Na$_2$B$_4$O$_7$. This is confirmed by the XRD pattern of the Na$_3$FeB$_5$O$_{10}$ sample recovered after 12h at 650°C in air (Figure 67c) which shows reflections corresponding to Fe$_2$O$_3$ and Na$_2$B$_4$O$_7$. The formation of Fe$_2$O$_3$ is consistent with the 2% weight gain that is visible in the TGA curve when the temperature increases from 500°C to 650°C. If the temperature is further raised the DSC plot shows a sharp endothermic peak ~710°C which can be correlated to the melting of the Na$_2$B$_4$O$_7$ formed during the decomposition of Na$_3$FeB$_5$O$_{10}$. In the case of Na$_3$CoB$_5$O$_{10}$ only one large endothermic peak is observed in the DSC curve ~770°C, referring to the melting point of the compound. Note that the initial mass loss ~1% for Na$_3$CoB$_5$O$_{10}$ can most likely be correlated to the removal of absorbed moisture.
Figure 67: TGA-DSC curves of (a) Na₃FeB₅O₁₀ and (c) Na₃CoB₅O₁₀ recorded in air flow with a heating rate of 5°C/min. (c) XRD patterns of Na₃FeB₅O₁₀ taken before and after annealing for 12h at 650°C in air.

4.2 Electrochemical characterization

4.2.1 Activity versus sodium

Given the possibility of feasible Na⁺ mobility due to the open structure of Na₃MB₅O₁₀, we decided to check the electrochemical activity of this compound towards Na. The experiments were conducted in Swagelok type half cells versus metallic sodium using a 1M solution of NaClO₄ in a 1/1 (w/w) mixture of EC and DMC with 1 wt.% fluoro ethylene carbonate (FEC) (details see annexes chapter 7.1). The positive electrode powder was
prepared by mixing the active material with 20 wt.% carbon SP with mortar and pestle under inert atmosphere for 10 min.

A typical voltage composition curve of Na$_3$FeB$_5$O$_{10}$ versus Na with a C/40 rate at 55°C is shown in Figure 68a. Initially the potential rapidly increases up to ~4 V, and then continuously increases but at a lower slope till 4.5 V with a corresponding removal of ~0.6 Na$^+$. During following discharge, a steep potential drop down to ~2 V is observed, and it is followed by a smooth voltage decrease till 1.5 V with in total the uptake of ~0.25 Na$^+$. Note that the amount of reversible Na$^+$ inserted in the initial cycle compared to the second one is much lower, indicating an irreversible Na loss during the first oxidation. However during conventional galvanostatic cycling no defined potential for the Fe$^{3+}$/Fe$^{2+}$ redox couple could be unambiguously determined. Thus we decided to conduct galvanostatic intermittent titration technique (GITT) measurements by applying a positive or negative current pulse for 4 h (corresponding to the extraction/insertion of 0.1 Na$^+$) alternating with open circuit voltage (OCV) periods of 12 h between 4.5 and 1.5 V at 55°C. As seen in Figure 68b, after each current pulse, the potential initially drops and increases strongly afterwards revealing an equilibrium potential of 2.5 V vs. Na$^+$/Na$^0$ pointing towards a biphasic process for Na extraction/insertion. As in the GITT curve large overpotentials in the range from 0.5 to almost 2 V during charge/discharge are observed, suggesting a strong kinetic limitation for Na$^+$ diffusion. At this point we suppose that these limitations are routed either in the poor ionic/electronic transport properties of one of the phases present during the biphasic process (Na$_3$FeB$_5$O$_{10}$ vs. Na$_x$FeB$_5$O$_{10}$).
Figure 68: (a) Voltage-composition curve of Na$_3$FeB$_5$O$_{10}$ versus Na cycled at a C/40 rate between 4.5 and 1.5 V vs. Na$^+$/Na$^0$ at 55°C. (b) GITT measurement with 4 h current pulse and 12 h open circuit voltage (OCV) period during the first charge and discharge.

In order to confirm the Fe$^{3+}$/Fe$^{2+}$ redox process, ex situ Mössbauer spectra were recorded for charged and discharged samples of Na$_3$FeB$_5$O$_{10}$ (details see annexes chapter 7.3.3). To do so, cells that have been fully charged (4.5 V) or discharged (1.5 V) were disassembled inside the glove box, the electrode powder was recovered, washed twice with DMC and dried under vacuum prior to examination. As shown in Figure 69 at the end of charge, the Fe$^{2+}$ doublet is reduced (49 at%) to the benefit of an additional contribution of Fe$^{3+}$ (51 at%) which is diminished upon subsequent discharge, although not fully reversible as 27 at% of Fe$^{3+}$ are remaining (together with 73 at% Fe$^{2+}$). The percentage of Fe$^{3+}$ and Fe$^{2+}$ given in Table 12 are in agreement with the amount of extracted/inserted Na compared to the voltage composition curve (Figure 68a).
Figure 69: Room temperature ex situ Mössbauer spectra recorded for Na$_3$FeB$_5$O$_{10}$ at 4.5 V (charged) and at 1.5 V (discharged).

Table 12: Fitted Mössbauer parameters of the pristine sample shown in Figure 69 and the spectra for the cycled samples given in Figure 73. IS, QS, LW and area are the isomer shift, the quadrupole splitting, the linewidth and the relative areas respectively.

<table>
<thead>
<tr>
<th>Sample</th>
<th>IS (mm/s)</th>
<th>SQ (mm/s)</th>
<th>LW (mm/s)</th>
<th>at%</th>
<th>Attribution</th>
</tr>
</thead>
<tbody>
<tr>
<td>pristine</td>
<td>1.02</td>
<td>2.10</td>
<td>0.26</td>
<td>100</td>
<td>Fe$^{2+}$</td>
</tr>
<tr>
<td>4.5 V</td>
<td>1.01</td>
<td>2.13</td>
<td>0.36</td>
<td>49</td>
<td>Fe$^{2+}$</td>
</tr>
<tr>
<td></td>
<td>0.33</td>
<td>0.71</td>
<td>0.61</td>
<td>51</td>
<td>Fe$^{3+}$</td>
</tr>
<tr>
<td>1.5 V</td>
<td>1.00</td>
<td>2.07</td>
<td>0.35</td>
<td>73</td>
<td>Fe$^{2+}$</td>
</tr>
<tr>
<td></td>
<td>0.31</td>
<td>0.73</td>
<td>0.42</td>
<td>27</td>
<td>Fe$^{3+}$</td>
</tr>
</tbody>
</table>

Na$_3$CoB$_5$O$_{10}$ was equally tested versus Na using a similar protocol as Na$_3$FeB$_5$O$_{10}$, however no electrochemical activity was observed. This does not come as a surprise as it is well known from DFT calculations that the redox potential of a 3d metal redox couple M$^{3+}$/M$^{2+}$ in a family of polyanionic compounds increases in the order from Fe < Mn < Co < Ni with differences of ~1.5 V if going from Fe$^{3+}$/Fe$^{2+}$ to Co$^{3+}$/Co$^{2+}$ in olivine-type materials
Sodium transition metal pentaborate $\text{Na}_3\text{MB}_5\text{O}_{10}$ for instance.\(^{196}\) Thus we believe that even such a compound will present an activity we won’t be able to reach it with the present electrolytes.

### 4.2.2 Ionic conductivity

At this stage an intriguing question regards the poor electrochemical performance of $\text{Na}_3\text{MB}_5\text{O}_{10}$ in light of its open structure. We hypothesize that such poor activity were routed in the poor electronic and ionic transport of the material. To check this hypothesis, BVEL calculations were generated according to the method developed by S. Adams to reveal possible ion diffusion pathways for $\text{Na}^+$ (details see annexes chapter 7.3.7).\(^{183}\) To determine the possible transport ways, the theoretical oxidation state of the mobile species ($\text{Na}^+$) is calculated as a function of the position inside the structure. Positions with only a small deviation (typically 10%) from the oxidation state are regarded as a part of the $\text{Na}^+$ diffusion pathway and are visualized in Figure 70 for $\text{Na}_3\text{MB}_5\text{O}_{10}$ (M =Fe (a), Co (b)). For $\text{Na}_3\text{FeB}_5\text{O}_{10}$, 1D $\text{Na}^+$ conduction paths, involving Na1 and Na3 sites appear along the a-axis at 1.00 eV above the minimum energy\(^4\) however Na2 is sterically hindered. Note that to involve also Na2 into the $\text{Na}^+$ migration network, an energy of 1.7 eV above the minimum is required, hence Na2 is most likely not contributing to the materials ionic conductivity. For $\text{Na}_3\text{CoB}_5\text{O}_{10}$ a percolation energy of 1.09 eV above the minimum is required to get fully connected pathways for Na migration involving all Na sites. Moreover the $\text{Na}^+$ conduction pathways are lying in between the layers consisting of the MO$_4$ tetrahedra connected through the pentaborate B$_5$O$_{10}$ units, implying an anisotropic $\text{Na}^+$ conduction.

\(^4\) Minimum energy refers to the calculated ground energy of the imaginary mobile $\text{Na}^+$ ion in the structure.
Figure 70: Bond valence energy landscape (BVEL) of (a) $\text{Na}_3\text{FeB}_5\text{O}_{10}$ and (b) $\text{Na}_3\text{CoB}_5\text{O}_{10}$. The percolation energies are given in the figure. Iso-energy blue domains indicate paths for Na (yellow spheres) conduction.

To experimentally validate the results derived from the BVEL calculations, complementary AC/DC measurements on sintered pellets of $\text{Na}_3\text{MB}_5\text{O}_{10}$ between ionically blocking Pt electrodes were done. Pellet suitable for the measurement were prepared by pressing around 300 mg of powdered sample in a 10 mm die. The pellet was annealed for 124 hours at 650°C under argon, or for 124 hours at 600°C in a vacuum sealed tube for M = Co and M = Fe, respectively. The obtained density was around 65 to 75 % of the theoretical one deduced from structural data. After placing the pellet in the apparatus, AC impedance spectra were recorded for temperatures ranging from 250 up to 400°C while applying a voltage amplitude of 200 mV in a frequency range between 30 MHz and 0.01 Hz during heating under argon flow (detailed description of the setup see annexes chapter 7.1.3).

First of all, the AC complex impedance spectra for $\text{Na}_3\text{MB}_5\text{O}_{10}$ (M = Fe, Co) present a depressed semicircle with a Warburg diffusion resistance tail over the whole measured temperature range, suggesting the AC conductivity to be mostly ionic (Figure 71). The spectra can be expressed with an equivalent circuit $R_0 - R_1/CPE_1 - R_2/CPE_2$ (Figure 71 inset) where each parallel R/CPE circuit gives raise to one semicircle (schematically plotted as grey lines in Figure 71). For each semicircle the resistance $R_1$ and $R_2$ (intercept with the x-axis, indicated in Figure 71) plus the corresponding capacitance $C_1$ and $C_2$ can be obtained from the fitting parameters (details see annexes chapter 7.1.3), whereas the latter one usually differs
from $10^{-12}$ to $10^{-9}$ F for the bulk and grain boundary contribution respectively. This allows us to separate between both contributions, corresponding to $R_1$ and $R_2$ for the bulk- and grain boundary resistance.

![Complex Impedance Spectra](image)

**Figure 71**: Representative complex impedance spectra of (a) Na$_3$FeB$_5$O$_{10}$ and (b) Na$_3$CoB$_5$O$_{10}$ recorded at 345 and 340°C respectively. The equivalent circuits are shown as inset, and the corresponding bulk ($R_1$) - and grain boundary ($R_2$) resistance are shown schematically as grey semicircles.

To clearly separate between ionic and electronic conductivity, DC polarization measurements were conducted (Figure 72) on sintered Na$_3$MB$_5$O$_{10}$ (M = Fe, Co) pellets sandwiched between two Pt-electrodes. After applying a constant potential of 0.5 V across the pellet, the current response was recorded. First a sharp increase followed a slow decay to a steady state value was observed. Initially, both ions and electrons account for the measured current, whereas the ionic current decays to zero after waiting a sufficiently long time, as Na$^+$ gets blocked at the Pt-electrodes. Electrons however are conducted through Pt, thus the steady-state current can be related to the pure electronic conduction of the material. As seen in Figure 72, the time to reach a constant current value took 4 times longer for Na$_3$CoB$_5$O$_{10}$ compared to Na$_3$FeB$_5$O$_{10}$ implying faster Na$^+$ migration for the latter one. The pure electronic resistance was calculated according to Ohm’s law from the steady state current value and the applied potential.
From the experimentally determined values for the AC and DC resistance, the corresponding conductivity was derived and the activation energy was obtained using the Arrhenius equation (details see annexes chapter 7.1.3). All results obtained from AC and DC measurements are summarized in the Arrhenius plot for Na$_3$FeB$_5$O$_{10}$ and Na$_3$CoB$_5$O$_{10}$ as shown in Figure 73a and b, respectively.

For Na$_3$FeB$_5$O$_{10}$ one can first note a linear increase of the bulk ionic conductivity between 250 and 380°C from $1.5\cdot 10^{-8}$ and $1.9\cdot 10^{-7}$ S·cm$^{-1}$ with an activation energy of 0.65 eV (Figure 73a). Conductivities associated to the grain boundaries also increase in the same temperature range from $1.7\cdot 10^{-7}$ and $4.7\cdot 10^{-7}$ S·cm$^{-1}$ correlated to an activation energy of 0.24 eV. Obviously the grain boundary conductivity is higher than the bulk one. One possibility to account for this observation is most likely rooted in the existence of a minor Fe$_3$BO$_5$ impurity (4%) which sits at the grain boundaries within our sample. The interaction between such an ionically insulating impurity and Na$_3$FeB$_5$O$_{10}$ could then lead to a redistribution of ions in the narrow interface region introducing structural defects/ disorder which lead as a consequence to a non-uniform charge distribution. This so called space-charge region can strongly improve the ionic conductivity through interfacial transport, hence our experimental values for the grain boundary conductivity may be overestimated.$^{199-203}$ The electronic conductivity increases between 250 to 350°C from $6.3\cdot 10^{-10}$ to $1.0\cdot 10^{-8}$ S·cm$^{-1}$ with a corresponding activation energy of 0.72 eV. By extrapolating the high temperature part of the curves to RT,
conductivities of $1.1 \cdot 10^{-10}$, $7.6 \cdot 10^{-9}$ and $1.2 \cdot 10^{-14}$ S·cm$^{-1}$ for the bulk-, grain boundary- and electronic conductivity are obtained respectively.

Regarding Na$_3$CoB$_5$O$_{10}$, the bulk ionic conductivity increases linearly from $6.9 \cdot 10^{-8}$ to $9.8 \cdot 10^{-7}$ S·cm$^{-1}$ between 280 and 390°C with a corresponding activation energy of 0.73 eV. The absolute values are about one order of magnitude lower than the values obtained for the bulk conductivity ($5.9 \cdot 10^{-9}$ to $8.6 \cdot 10^{-8}$ S/cm), corresponding to an activation energy of 0.83 eV. As the materials appears to be free of impurities, we hypothesize these lower values for the grain boundary conductivity arise from structural disorder/defects at the grain interfaces, impeding the ionic conduction, although an insulating amorphous layer at the grain surface cannot be excluded either.\textsuperscript{204-206} The electronic conductivity increases between 290 to 390°C from $1.1 \cdot 10^{-9}$ to $2.2 \cdot 10^{-8}$ S·cm$^{-1}$ respectively, correlated to an activation energy of 0.97 eV. Linear extrapolation of the experimental values leads to RT conductivities of $4.0 \cdot 10^{-13}$, $9.1 \cdot 10^{-17}$ and $5.4 \cdot 10^{-17}$ S·cm$^{-1}$ for the bulk-, grain boundary- and electronic conductivity, respectively.

For comparison, it should be recalled that the RT bulk ionic conductivity values for Na$_3$FeB$_5$O$_{10}$ are one only order of magnitude lower than the one of LiFePO$_4$ ($\approx 2 \cdot 10^{-9}$ S·cm$^{-1}$)\textsuperscript{106} and two order of magnitudes higher than Li$_2$FeSiO$_4$ ($\approx 6 \cdot 10^{-14}$ S·cm$^{-1}$),\textsuperscript{100} both of which can reversibly insert one Li per unit formula. Having this in mind, the poor electrochemical performance of Na$_3$FeB$_5$O$_{10}$ comes still as a surprise since it should not be limited by the ionic transport. Furthermore, it is worth mentioning that the electronic conductivity of LiFePO$_4$ is higher than that of Na$_3$FeB$_5$O$_{10}$. This may explain the poor electrochemical activity as we need ionic an electronic conductivity to electrochemically trigger insertion/deinsertion reactions. Such a limitation can also be structural related with the under-stoichiometric Na$_3$-$x$FeB$_5$O$_{10}$ phase being thermodynamically unstable, the reason why for instance the de-insertion of Li from Li$_2$FeSiO$_4$ could not be extended below one, as confirmed by DFT calculations. Turning back to the GITT measurement (Figure 68b), one can note that the equilibrium potential reaches a plateau at 2.5 V vs. Na$^+/Na^0$ which is indicative of a two-phase Na insertion/deinsertion reaction, hence one can assume the formation of a second phase Na$_x$FeB$_5$O$_{10}$ which is most likely the limiting factor for electrochemical activity, similar to what was observed in the case of LiFeBO$_3$.\textsuperscript{151,207}
**Figure 73:** Arrhenius plot of the AC and DC conductivity of (a) $\text{Na}_3\text{FeB}_5\text{O}_{10}$ and (b) $\text{Na}_3\text{CoB}_5\text{O}_{10}$ derived in the temperature range from around 250 to 400°C during heating in argon flow.

### 4.3 Conclusion

In this section were report about the synthesis, structure and electrochemical properties of two new sodium transition metal pentaborates $\text{Na}_3\text{MB}_5\text{O}_{10}$ (M = Fe, Co) where M = Fe presents the first ever reported compound out of the ternary Na$_2$O-FeO-B$_2$O$_3$ system. The structures were solved by means of synchrotron XRD. They consist of layers built up by MO$_4$ tetrahedra connected through pentaborate units B$_5$O$_{10}$ with sodium sitting in voids inside these layers as well as in between them. Complementary galvanostatic- and *ex situ* Mössbauer measurements have confirmed a Fe$^{3+}$/Fe$^{2+}$ redox activity centered around 2.5 V vs. Na$^+$/Na$^0$ for $\text{Na}_3\text{FeB}_5\text{O}_{10}$ in contrast to $\text{Na}_3\text{CoB}_5\text{O}_{10}$ which is electrochemical inactive. Combined BVEL calculations and AC/DC conductivity measurements reveal strong kinetic limitations for $\text{Na}_3\text{CoB}_5\text{O}_{10}$, reflected in the low RT ionic- and electronic conductivities of $4.0 \times 10^{-13}$ and $5.4 \times 10^{-17}$ S·cm$^{-1}$ respectively. $\text{Na}_3\text{FeB}_5\text{O}_{10}$ in contrast shows RT ionic conductivity of $1.1 \times 10^{-10}$ S·cm$^{-1}$ comparable to LiFePO$_4$, however one limiting factor is nested in the much lower RT electronic conductivity ($1.2 \times 10^{-14}$ S·cm$^{-1}$). An additional limitation for the electrochemical performance could equally be due to the structural /thermodynamic instability of the intermediate phase $\text{Na}_x\text{FeB}_5\text{O}_{10}$ formed during the two-phase (de)sodiation reaction.
Although a reversible electrochemical activity could be demonstrated for Na$_3$FeB$_5$O$_{10}$, this material cannot be considered for practical applications. Note that our attempts to circumvent the poor electronic conductivity by employing carbon coated particles failed. Nevertheless, these results call for further exploration of borate based materials in terms of their ionic mobility, despite difficulties encountered during the design of new compounds out of this family.

In general we have shown possible insertion/deinsertion reaction for Li and Na in borate based materials beyond LiMBO$_3$ in the last two chapters, however the limited electrochemical activity led us to consider other reaction mechanism, namely conversion type reactions as explored in the next chapter.
5 Study of the electrochemical driven reaction mechanism of bismuth oxyborate Bi$_4$B$_2$O$_9$ versus lithium

5.1 Synthesis and structure

In the previous chapters, we have shown through the synthesis of Li$_6$CuB$_4$O$_{10}$ the feasibility to obtain high redox potentials for borate materials, however our further exploration for new pyroborate based compounds has encountered strong synthetic difficulties. Nevertheless, this had led us to the synthesis of two new sodium 3d metal pentaborates Na$_3$MB$_5$O$_{10}$ (M = Fe, Co) with poor electrochemical performances. Therefore we considered another reaction mechanism, namely conversion type reactions. They differ from classical insertion/ deinsertion reactions in which Li$^+$/Na$^+$ is incorporated into a host structure while the metal M changes its oxidation state to account for charge balance. In a reversible conversion reaction, M is fully reduced upon lithiation and forms nanoparticles embedded in an amorphous matrix composed of Li$_n$X (X = O$^{2-}$, S$^{2-}$, F$^-$, P$_3$$^-$$^3$, etc.) (equation 7). Upon charge this process is reversible to a certain extend.

$$M_{a}X_{b} + n e^- + n Li^+ \leftrightarrow a M^0 + b Li_nX \quad (7)$$

An interesting tool to tune the redox potential of electrode materials relies on the inductive effect. In short the ionicity of the M─X bond influences the average redox potential of the conversion reaction with compounds having the highest ionicity showing the highest redox potentials. As fluorine is the most electronegative element, fluorides were widely studied in the past, however one has to note that highly ionic compounds are usually poor electronic conductors.\cite{123,208-212}. Therefore in most cases these materials have to be “activated” through a prolonged high energy milling process with a large amount of carbon to achieve good electronic contact. Worthwhile mentioning that this addition of carbon diminishes drastically the volumetric energy density which is one problem regarding applications. Thus the quest for denser materials, with the most promising among them being bismuth fluoride BiF$_3$ (5.23 g·cm$^{-3}$). But again due to the insulating nature of this compound, a relatively large polarization was observed (~1 V), hence we decided to switch to Bi-borates. Owing to the weaker inductive effect of the borate BO$_3$ group compared to fluorine, they should show a lower polarization while keeping the benefit of the high volumetric density (around 5 to 8
Bismuth oxyborate Bi$_4$B$_2$O$_9$ g·cm$^{-3}$), comparable to BiF$_3$. Among all reported Bi-borates, Bi$_4$B$_2$O$_9$ presents the highest theoretical capacity correlated to the reduction of Bi$^{3+}$ to Bi$^0$ (~320 mAh/g) therefore, we chose this material to investigate its reaction mechanism versus Li. So far this materials was solely investigated for its structural and photocatalytical properties.

Although the theoretical capacity is rather low for conversion type electrode materials (usually > 600 mAh/g), we strongly emphasize the fact that this material presents a very high volumetric density ~8.2 g·cm$^{-3}$, hence being favorable from the viewpoint of volumetric energy density.

Bi$_4$B$_2$O$_9$ was obtained by mixing stoichiometric amounts of bismuth oxide Bi$_2$O$_3$ and boric acid H$_3$BO$_3$ with a mortar and pestle and annealing the mixture at 600°C for 36 h in air with one intermediate regrinding. After the synthesis a pale white powder was recovered, consisting of particles with sizes ranging from 5 to 30 μm (Figure 74, inset).

![Figure 74: XRD pattern and Rietveld refinement of Bi$_4$B$_2$O$_9$. Observed, calculated pattern and their difference are shown as red crosses, black- and green line respectively. Bragg reflections are indicated as vertical blue bars. The inset show a representative SEM image of the as synthesized compound with two different magnifications.](image)

The corresponding XRD pattern and the Rietveld refinement are shown in Figure 74 and the refined atomic positions are given in Table 13. All peaks could be indexed in a monoclinic space group P2$_1$/c with the unit cell parameters a = 11.11652(4) Å, b = 6.63300(3)
Å, c = 11.04575(4) Å, β = 91.04795(16)° and V = 814.331(5) Å³, similar to those reported in literature.\textsuperscript{213,214} The atomic arrangement within the unit cell is shown in Figure 75, built up of 4 different crystallographic sites for Bi, 2 for B and 9 for O. Among them Bi4 and Bi2 form BiO\textsubscript{4}-, and Bi1 and Bi3 form BiO\textsubscript{5} asymmetric truncated polyhedra respectively, with Bi-O distances ranging from 2.150 to 2.492 Å. Boron is trigonal planar coordinated by oxygen forming regular BO\textsubscript{3}\textsuperscript{3-} triangles (average B-O bond distance 1.3396 Å) which are stacked on top of each other perpendicular to \textit{b}-axis (Figure 75a). The structure also contains three O atoms (O1, O2, O4) which are non-coordinated to boron, only bond to Bi atoms with bond length ranging from 2.177 to 2.381 Å, referring Bi\textsubscript{4}B\textsubscript{2}O\textsubscript{9} to oxyborate which can be more precisely expressed in the formula Bi\textsubscript{4}O\textsubscript{3}(BO\textsubscript{3})\textsubscript{2}.

\textbf{Figure 75:} Crystal structure of Bi\textsubscript{4}B\textsubscript{2}O\textsubscript{9} shown along the (a) \textit{b}- and (b) \textit{c}-axis. Bismuth atoms/polyhedral, boron and oxygen are shown in purple, green and red respectively.
Table 13: Structural parameters and atomic positions of Bi$_4$B$_2$O$_9$.

<table>
<thead>
<tr>
<th>Atom</th>
<th>Wyckoff site</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>B (Å$^2$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bi1</td>
<td>2a</td>
<td>0.9920(1)</td>
<td>0.4822(2)</td>
<td>0.3460(1)</td>
<td>0.19(3)</td>
</tr>
<tr>
<td>Bi2</td>
<td>2a</td>
<td>0.8083(1)</td>
<td>0.0480(2)</td>
<td>0.4920(1)</td>
<td>0.36(3)</td>
</tr>
<tr>
<td>Bi3</td>
<td>2a</td>
<td>0.5014(1)</td>
<td>0.4236(2)</td>
<td>0.1597(1)</td>
<td>0.18(3)</td>
</tr>
<tr>
<td>Bi4</td>
<td>2a</td>
<td>0.3402(1)</td>
<td>0.4366(2)</td>
<td>0.4825(1)</td>
<td>0.31(4)</td>
</tr>
<tr>
<td>O1</td>
<td>2a</td>
<td>0.5003</td>
<td>0.1076</td>
<td>0.1050</td>
<td>1.89</td>
</tr>
<tr>
<td>O2</td>
<td>2a</td>
<td>0.3833</td>
<td>0.0994</td>
<td>0.4944</td>
<td>1.99</td>
</tr>
<tr>
<td>O3</td>
<td>2a</td>
<td>0.8652</td>
<td>0.1597</td>
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</tr>
<tr>
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<td>0.1577</td>
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<td>1.96</td>
</tr>
<tr>
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<td>2a</td>
<td>0.1879</td>
<td>0.2030</td>
<td>0.1427</td>
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<td>0.3676</td>
<td>2.81</td>
</tr>
<tr>
<td>O7</td>
<td>2a</td>
<td>0.1472</td>
<td>0.2420</td>
<td>0.3527</td>
<td>2.14</td>
</tr>
<tr>
<td>O8</td>
<td>2a</td>
<td>0.3433</td>
<td>0.3144</td>
<td>0.2792</td>
<td>1.30</td>
</tr>
<tr>
<td>O9</td>
<td>2a</td>
<td>0.7152</td>
<td>0.3195</td>
<td>0.1549</td>
<td>2.00</td>
</tr>
<tr>
<td>B1</td>
<td>2a</td>
<td>0.7534</td>
<td>0.2356</td>
<td>0.2633</td>
<td>2.00</td>
</tr>
<tr>
<td>B2</td>
<td>2a</td>
<td>0.2244</td>
<td>0.2520</td>
<td>0.2533</td>
<td>2.15</td>
</tr>
</tbody>
</table>

The thermal stability of Bi$_4$B$_2$O$_9$ was investigated using TGA-DSC analysis (for details see annexes chapter 7.3.1). Measurements were recorded during heating a sample of around 20-30 mg under flowing air from RT to 500°C with a rate of 5°C/min (Figure 76). In the whole temperature range the mass of the sample stays constant and no peak in the DSC curve indicative of a possible structural transition is visible, as already suggested through XRD measurements at 200 and 450°C by Filatov et al.. In general Bi$_4$B$_2$O$_9$ is stable up to 500°C in air.
Figure 76: TGA-DSC curve of Bi₄B₂O₉ recorded in air from RT to 500°C during heating under air flow.

5.2 Electrochemical characterization

Given the possibility of the reduction from Bi³⁺ to metallic Bi⁰ and expected higher redox potentials than pure oxides owing to the inductive effect of the borate group, we measured the electrochemical activity of Bi₄B₂O₉ versus lithium. Electrochemical testing was done in Swagelok half cells using a 1M solution of LiPF₆ in EC/DMC/PC 1/1/3 (w/w/w) as electrolyte soaked into two sheets of glass fiber which separate the positive electrode from metallic lithium. The positive electrode powder was prepared by mixing Bi₄B₂O₉ with 5 wt.% carbon for 10 min using a SPEX high energy miller.

First Bi₄B₂O₉/Li half cells were cycled between 0.0 and 3.5 V at a C/10 rate in order to ensure a complete reduction of the material. A typical voltage composition curve, together with the corresponding derivative plot and the capacity retention are shown in Figure 77. First of all one can note that the initial discharge corresponds to the reaction of 24 mol of Li⁺ with 1 mol of Bi₄B₂O₉. Such a Li uptake is associated with three different voltage regimes. At the beginning, a flat plateau at 1.8 V is visible concomitant with the uptake of ~8 Li⁺, followed by an S-shaped curve centered at 1.4 V and corresponding to the uptake of 4 Li⁺. Finally the potential reaches a low voltage flat plateau around 0.8 V, accompanied with the uptake of ~12 Li⁺. On subsequent charge, only ~9 Li⁺ can be extracted, solely correlated to the redox activity ~0.8 V (Figure 77b, inset). The irreversible loss in capacity between the first discharge and first charge is quite important as it is about 60%. Moreover the reversible
capacity decays initially very fast upon cycling while becoming almost negligible after 5 cycles, indicative of a large irreversible Li-reactivity.

Figure 77: (a) Voltage-composition curve of Bi$_4$B$_2$O$_9$/Li half-cell and (b) the capacity retention and the corresponding derivative plot of voltage-composition curve is show as inset.

The Li$^+$ uptake during the first discharge can be explained by first considering a reduction of Bi$_4$B$_2$O$_9$ to metallic Bi involving the reaction with 12 Li$^+$ (equation 8), and second an alloying of the as-formed Bi with lithium to yield Li$_3$Bi (equation 9). By summing these two reactions, one exactly reaches the value of 24 Li$^+$ incorporated during the initial reduction (equation 10), consistent with the electrochemical data (Figure 77a).

$$Bi_4B_2O_9 + 12 Li^+ + 12 e^- \leftrightarrow 3 Li_2O + 2 Li_3BO_3 + 4 Bi^0 \quad (8)$$

$$4 Bi^0 + 12 Li^+ + 12 e^- \leftrightarrow 4 Li_3Bi \quad (9)$$

$$Bi_4B_2O_9 + 24 Li^+ + 24 e^- \leftrightarrow 3 Li_2O + 2 Li_3BO_3 + 4 Li_3Bi \quad (10)$$

To get insights from a structural point of view, *ex situ* XRD pattern were recorded for Bi$_4$B$_2$O$_9$ powder recovered at different stages of discharge and charge. To do so Bi$_4$B$_2$O$_9$ /Li half cells discharged to 1.0, 0.0 and charged at 3.5 V were disassembled inside the glove box, the recovered powder was washed twice with DMC and dried under vacuum prior to
examination. As seen in Figure 78 at 1.0 V one can note a strong decrease in intensity and a highly pronounced broadening of the main reflections around a 2θ value of 28.6°, suggesting a strong amorphisation/decomposition of the pristine phase. Based on the electrochemical data this phase should have fully decomposed leading to a featureless XRD. This is not the case and the main reason is most like rooted in the presence of secondary Li consuming reactions leading to the formation of a Li containing solid electrolyte interface (SEI) or a Li containing organic layer around the active material.

However, one might expect that during initial lithiation Li2O and Li3BO3 are formed (equation 8). This decomposition seems to be reasonable as similar reactions were observed in the case of other TM borates.215 Now the question arises, why these conversion products are not visible in the XRD patterns. Either because their amorphous/ nanocrystalline nature, or the large difference of X-ray scattering between Li2O/ Li3BO3 (light elements) and heavy atoms such as Bi. To check the morphology of the active material discharged to 1.0 V, TEM images were recorded. As shown in Figure 78a one can clearly see nanoparticles with the size ranging from 2 to 10 nm embedded in an amorphous matrix. To analyze the crystal structure of the nanoparticles, selected area electron diffraction (SAED) was performed showing diffraction rings which could be attributed to a trigonal R-3m symmetry indicative of the presence of metallic α-Bi (Figure 78b). To analyze the elemental distribution of the composite we used EDX analysis. Note that the analysis of Li and B is not possible due to the low energy of the emitted X-rays which cannot be detected with the instrument. However the nanoparticles were confirmed to consist of metallic Bi and tend to form agglomerates (Figure 78c). Furthermore in contrast to the locally defined appearance of Bi, the contribution of oxygen seems to be homogenous over the whole area, supporting the existence of an evenly distributed amorphous Li2O/ Li3BO3 matrix (Figure 78c).
Figure 78: Bi$_4$B$_2$O$_9$/C sample discharged to 1.0 V. (a) Representative TEM image, (b) SAED pattern of the nanoparticles with the simulated diffraction profile of α-Bi shown for comparison and (c) another TEM image and the corresponding EDX map of bismuth (red) and oxygen (blue).

For samples fully reduced to 0.0 V, sharp reflections of Li$_3$Bi are observed from XRD. On subsequent charge to 3.5 V peaks of metallic Bi$^0$ show up in the XRD pattern (Figure 79). This is consistent with our assumptions of a reversible alloying reaction between Li and Bi as stated in equation 9 associated to the redox process around 0.8 V seen in the derivative plot (Figure 79b). The reversible electrochemical driven alloying/ dealloying process of Bi with Li has been reported. Nevertheless the reversibility here is much lower compared to literature which originates most likely from the progressive separation/ isolation of the active material (Bi/ Li$_3$Bi) through the surrounding components formed during the initial conversion process (Li$_2$O, Li$_3$BO$_3$, and organic layers) impeding ionic/ electronic transport.

A still pending question deals with inability to trigger the conversion reaction (equation 8) once Li$_3$Bi has been formed (equation 9). We believe that once Li$_3$Bi is formed, the de-alloying reaction leads to large Bi$^0$ crystallites (evidenced by the sharp reflections from
XRD) separated from the remaining Li$_2$O and Li$_3$BO$_3$ matrix. Thus in light of this lack of ionic/electronic percolation through the composite, the simultaneous delithiation of the Li-containing matrix (Li$_2$O/ Li$_3$BO$_3$) and oxidation of the Bi crystallites is not possible any longer.

Figure 79: XRD pattern of Bi$_4$B$_2$O$_9$ discharged to 1.0 and 0.0 V and after the subsequent charge to 3.5V.

As the reversibility of Bi$_4$B$_2$O$_9$/Li half-cells cycled between 0.0 and 3.5 V is very poor, we investigated the effect of higher discharge cutoff potentials in order to exclude the alloying/dealloying part and trigger only the conversion process.

Bi$_4$B$_2$O$_9$/Li half-cells were assembled and cycled between 1.0 and 3.5 V. A typical voltage-composition curve together with the corresponding derivative plot and the capacity retention are shown in Figure 80. During the initial reduction 8 Li$^+$ are incorporated via a flat plateau at 1.8 V with an additional uptake of 4 Li$^+$ during a S-shaped voltage profile centered around 1.4 V. Upon following charge, around 8 Li$^+$ can be reversible extracted associated to two different oxidation processes at average redox potential of 1.5 and 2.5 V (Figure 80b). During the first discharge the theoretical capacity for Bi$^{3+}$ to Bi$^{0}$ of 320 mAh/g is reached, however upon following charge, ~33% irreversible capacity remains, corresponding to a resulting charge capacity of ~214 mAh/g. Moreover the capacity is fading very fast upon cycling, decreasing down to 50% of the initial value after 3 cycles.
In short, if the discharge is limited to 1.0 V, the initial reduction processes around 1.8 and 1.5 V become reversible with the former one shifted up to 2.2 V through the subsequent discharges (Figure 80b). This potential difference between the initial and following discharge is typically observed for conversion type materials since different reaction pathways occur during the initial and subsequent reductions. During the first discharge, the potential drop is larger as an higher activation energy is necessary to trigger the conversion reaction. After this formation step, the second and subsequent discharges will be kinetically less limited owing to the nano character of the formed composite hence the discharge potential is increased.²¹⁸

The two different redox processes occurring at average voltages of 1.5 and 2.5 V suggest the presence of two different reactions. We assume that during the initial reduction at 1.8 V metallic Bi and Bi₂O₃ nanoparticles are formed and are accompanied with the formation of amorphous matrix of Li₃BO₃ according to equation 11. Upon further discharge the remaining Bi₂O₃ is reduced to Bi⁰ and amorphous Li₂O around 1.5 V (equation 12). If both reactions are summed up, the total amount of reacted Li is consistent with the value deduced from the electrochemical curve during the initial reduction (Figure 80a).
Although we do not have evidence for the existence of Bi$_2$O$_3$ from electron microscopy at present, our hypothesis of starting the formation of Li$_3$BO$_3$ and then Li$_2$O (equation 11, 12) seems to be reasonable, and similar observations have been made for the bismuth oxyfluoride BiO$_{0.5}$F$_2$/BiOF system. Moreover this system was shown to react with Li via a conversion reaction, but the conversion of the fluoride occurs first, followed by the conversion of the as-formed bismuth oxide. Further literature reports assume the conversion reaction of Bi$_2$O$_3$ versus Li around 1.2/1.4 V and 1.8 V for α-Bi$_2$O$_3$ and δ-Bi$_2$O$_3$ respectively. So in light of these results we believe that the reduction of Bi$_2$O$_3$ (equation 12) starts most likely before all Bi$_4$B$_2$O$_9$ is converted (equation 11).

On subsequent galvanostatic cycling we assume the reaction defined in equation 13 to be partially reversible as Bi$_4$B$_2$O$_9$ is formed back on oxidation, although the nano sized morphology of the active material composite prevents us to fully support this statement (Figure 81).

\[
\text{Bi}_4\text{B}_2\text{O}_9 + 12 \text{Li}^+ + 12 e^- \leftrightarrow 3 \text{Li}_2\text{O} + 2 \text{Li}_3\text{BO}_3 + 4 \text{Bi}^0 \quad (13)
\]

![TEM image of a Bi$_4$B$_2$O$_9$/C sample discharged to 1.0 V and recharged to 3.5 V.](image)

**Figure 81:** TEM image of a Bi$_4$B$_2$O$_9$/C sample discharged to 1.0 V and recharged to 3.5 V.

Although it is well known δ-Bi$_2$O$_3$ is only stable above 730°C, Bervas *et al.* claim on the basis of ex *situ* XRD that this particular phase exists as an intermediate product during the reduction of BiO$_{0.5}$F$_2$.  

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5 Although it is well known δ-Bi$_2$O$_3$ is only stable above 730°C, Bervas *et al.* claim on the basis of ex *situ* XRD that this particular phase exists as an intermediate product during the reduction of BiO$_{0.5}$F$_2$.  

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101
So far we have investigated the electrochemical behavior of Bi$_4$B$_2$O$_9$ versus Li in two different potential regimes (0.0-3.5 V and 1.0-3.5 V). It reveals an electrochemically driven reaction with a large amount of Li that cumulates conversion and alloying processes, but both of them being poorly reversible. Hence our desire to further increase the lower cutoff potential to 1.7 V in order to exclude the 1.5 V redox activity Figure 82. After the initial discharge plateau at 1.8 V coming along with the uptake of ~8 Li$^+$, there is a shift of the electrochemical curve to an S-shape on subsequent charge to 3.5 V. Approximately 5 Li$^+$ can be reversible extracted, hence an irreversible capacity of 35% still remains. Upon on following discharge, the potential drops vertically reaching a pseudo-plateau at 2.2 V as visualized by a sharp reduction peak in the derivate plot (Figure 82b). Once the first cycle is achieved, the average voltage of the cell is about 2.3 V with a reversible capacity of 150 mAh/g that stays at least stable over 10 charge/discharge cycles.

![Figure 82: (a) Voltage-composition curves and (b) the corresponding derivative plots and capacity retention of a Bi$_4$B$_2$O$_9$/Li half-cell cycled between 1.7 and 3.5 V.](image)

The ex situ XRD pattern taken on a sample discharged to 1.7 V still showed remaining reflections of pristine Bi$_4$B$_2$O$_9$ (Figure 83). To check if these reflections remain upon further cycling, additional ex situ XRD patterns were recorded on the active material after the second and fifth discharge for cells cycled between 1.7 and 3.5 V. As shown in Figure 83 a complete amorphization was not achieved until the 5$^{th}$ discharge/charge cycles. This suggests a gradual consumption of the material during galvanostatic cycling, as already described by cycling LiCoO$_2$ to lower potentials.
Figure 83: XRD pattern of Bi$_4$B$_2$O$_9$ discharged to 1.7 V after the first, second and fifth cycle.

To get more insights into the morphology of the active materials composite, samples initially discharged to 1.7 V were examined by means of TEM and EDX spectroscopy. As shown in Figure 84a nanoparticles (2 to 10 nm) embedded in an amorphous matrix are visible in parallel with remaining large crystallites of Bi$_4$B$_2$O$_9$ (Figure 84a, indicated with orange arrow). The nanoparticles were analyzed through SAED and are shown to crystallize in an R-3m space group referring most likely to α-Bi. Using EDX mapping, we determined the composition of the nanoparticles and confirmed them to be metallic Bi. However, we could not probe the composition of the amorphous part of the sample, most likely composed of Li$_2$O/ Li$_3$BO$_3$, since EDX analysis cannot spot light elements. Although we have claimed that during the reduction of Bi$_4$B$_2$O$_9$ there is the partial formation of Bi$_2$O$_3$ (equation 11), we could not reach evidence from electron microscopy yet. Thus we assume Bi$_2$O$_3$ to be amorphous,\textsuperscript{222,223} but further characterization like pair distribution function (PDF) analysis or X-ray absorption (XAS) measurements are necessary for more information regarding the local structure of Bi.
We have presently tested the electrochemical reaction of Bi$_4$B$_2$O$_9$ versus Li in three different potential windows between 0.0 and 3.5 V. In the following we mainly address the influence of the lower cutoff potential (0.0, 1.0 and 1.7 V). If the discharge cutoff voltage is raised from 0.0 to 1.7 V the capacity retention seems to be enhanced (Figure 85). Several parameters can account for such difference. First of all if the galvanostatic cycling is conducted in a more narrow potential range, there is less electrolyte decomposition associated to the formation of organic layers (parasitic reactions) with the end result being a minimization of the particle surface. Secondly, the faster capacity decay for lower cutoff potentials can be partially correlated to a stronger phase separation. As shown in Figure 86a and b the bright regions denoted to the amorphous Li$_3$BO$_3$/ Li$_2$O matrix is increased from the sample recovered at 1.7 V compared to 1.0 V. These stronger separation/ textural changes

**Figure 84:** Bi$_4$B$_2$O$_9$/C sample discharged to 1.7 V. (a) Representative TEM image where the orange arrow indicates large particle of not reacted Bi$_4$B$_2$O$_9$. (b) SAED pattern of the nanoparticles with the simulated diffraction profile of α-Bi shown for comparison and (c) another TEM image and the corresponding EDX map of bismuth (red) and oxygen (cyan).
lower the ionic/electronic transport, hence a large fraction of the active material is not accessible for the conversion reaction anymore.

**Figure 85**: (Capacity retention of Bi$_4$B$_2$O$_9$ mixed for 10 min with 5 wt% carbon and cycled with a C/10 rate in different potential window indicated in the plot. To have a better overview, the first discharge/charge capacities are excluded.

**Figure 86**: Typical HRTEM images of the active material discharged at (a) 1.7 V and (b) 1.0 V.

Now, at this stage that there are still two important remaining questions regarding the electrochemical performance of Bi$_4$B$_2$O$_9$ versus Li remains. They are 

1) the surprisingly small polarization (~300 mV) and 2) the tiny amount of carbon additive necessary for cycling. At
this point the alloying/ dealloying process was not anymore of interest due to the poor reversibility at low voltage. So from now on, all electrochemical measurements were conducted above 1.0 V. To address these questions, potentiostatic intermittent titration technique (PITT) measurements were implemented to Bi$_4$B$_2$O$_9$/Li half-cells, with 10 mV steps after a current decay corresponding to a C/10 rate in a voltage window from 1.0 to 3.5 V. The shape of the current response can be used to distinguish between two phases and single phase electrochemical processes. For instance a single phase process (solid solution) is usually characterized by a fast current relaxation after each voltage step. This is in contrast to a biphasic process (phase transformation) where a local maximum in the current curve appears in the chronoamperometry plot. For a single phase reaction, the rate limiting step is the chemical diffusion, whereas in the biphasic case, the phase transformation is the limiting factor.

Turning to the PITT data (Figure 87), we initially decrease the potential stepwise from 1.95 to 1.85 V with a fast current relaxation for each voltage step (Figure 87a). In this window ~0.5 Li$^+$ are incorporated, with the current response pointing towards a single phase process. Afterwards a large plateau located around 1.8 V is observed (Figure 87b) correlated to a multiphasic process supported through the successive bell-shape current responses (Figure 87b). This confirms our assumption regarding the decomposition of Bi$_4$B$_2$O$_9$ into amorphous Li$_3$BO$_3$, Bi$^0$ and Bi$_2$O$_3$. Between x ≈ 8 and 11, again a nice S-shape voltage profile is visible centered around 1.4 V (Figure 87c). It is representative of a monophasic process where we hypothesize a reduction of the remaining Bi$_2$O$_3$ to Bi$^0$. If we consider that this reaction is being governed by the Bi$^{3+}$/Bi$^0$ redox couple accompanied with a multiphase process, a plateau-like response should be visible, in contrast to what we found. Bearing in mind the formation of Bi/ Bi$_2$O$_3$ nanocomposite at 1.7 V (Figure 87a), this discrepancy does not come as a surprise, since it is well known that the particle size can strongly affect the electrochemical signatur. and this could be at the origin of the sloping electrochemical trace.
Lastly, turning back to the sloping anomaly at the beginning of the voltage-composition curve, time has come to check if Li-insertion into Bi$_4$B$_2$O$_9$ is taking place here. *In situ* XRD measurements up to a Li uptake of $x = 1$ was made. For that, a homemade cell equipped with a Be window transparent to X-rays was started with a C/10 rate on discharge and XRD pattern were collected for every change in Li stoichiometry of 0.1. As shown in Figure 88 no shift of the peaks is observed, which could have being indicative of a Li insertion into Bi$_4$B$_2$O$_9$. Furthermore neither an amorphization nor a secondary phase is detected. As we could not extract any information at this point, we followed the first discharge by means of *in situ* XRD down to 1.7 V.
Figure 88: In situ XRD of the initial discharge of a Bi₄B₂O₉/Li half-cell cycled with a C/10 rate till a Li uptake of \(x = 1.0\). XRD pattern were recorded each 0.1 Li\(^+\). A magnification of the main reflections is presented in (a) and an overview over the whole XRD pattern and the corresponding voltage-composition curve is shown in (b).

As shown in Figure 89 the intensity of the Bragg reflections of pristine Bi₄B₂O₉ is reduced during discharge and charge indicating a gradual consumption of the pristine material which undergoes a partial conversion during the initial reduction to 1.7 V. The loss in intensity during the first cycle is exemplarily shown on the main reflection (Figure 89a), with its intensity reduced by 60% and then another 10% upon successive charge (Figure 89b). The presence of pristine reflections after the initial lithiation does not come as a surprise as we can expect an incomplete consumption of the pristine Bi₄B₂O₉ when the reduction is limited to 1.7 V.
Figure 89: In situ XRD of the initial discharge/charge cycle of a Bi$_4$B$_2$O$_9$/Li half-cell cycled with a C/10 rate in a potential window between 1.7 and 3.5 V. XRD pattern were recorded each 0.1 Li$^+$, however for reasons of clarity only each 5$^{th}$ pattern is shown in (c). A magnification of the main reflections is presented in (a) and their integrated intensity of the main reflection is shown in (b).

To grasp further fundamentals insights into the process occurring at the beginning of the discharge, Bi$_4$B$_2$O$_9$/Li cells were discharged with different C-rates, ranging from 1C to C/100 (Figure 90a). When the C-rate decreases we note a more defined pseudo-plateau around 1.95 V which disappears for rates higher than C/5 indicating kinetic limitations of this process. Since in situ XRD ruled out a Li$^+$ insertion process, we assigned the initial pseudo-plateau region to electrochemical oxygen removal leading to the formation of Li$_2$O on the particle surface.
Figure 90: Initial discharge of Bi$_4$B$_2$O$_9$/carbon composite. In (a) the influence of different C-rates is shown for a milling time of 10 min, whereas in (b) the impact of the milling time is presented.

To check this hypothesis we next investigated the effect of the material surface area. Therefore Bi$_4$B$_2$O$_9$ was ball milled with 5 wt.% carbon for 10, 30 and 60 min, leading to smaller particles and hence a higher surface area, as witnessed by a progressive broadening of the XRD reflections (Figure 91). We found that the anomaly at initial state of discharge is extended from $x \approx 0.4$ up to $x \approx 2.5$ as the particles are downsized (Figure 90b). This result strongly points towards an electrochemical driven oxygen removal from Bi$_4$B$_2$O$_9$ from near surface regions.

Figure 91: XRD pattern of Bi$_4$B$_2$O$_9$ ball milled for different times with 5 wt% carbon.
To corroborate our hypothesis of oxygen removal from the particle surface let us recall that the voltage-composition profile of Bi$_4$B$_2$O$_9$ versus Li presents a similar shape to that of BIMEVOX phases (Bi$_4$V$_2$O$_{11}$, Bi$_{3.6}$Pb$_{0.4}$V$_2$O$_{11-y}$, Bi$_4$V$_{1.8}$Cu$_{0.2}$O$_{11-y}$), hence suggesting similar electrochemical reactivity. Early work dealing with the reduction of Bi$_4$V$_2$O$_{11}$ has revealed an initial decomposition reaction that was shown to result from the oxygen removal process of the pristine material yielding in Bi$_4$V$_{10.66}$ and amorphous Li$_2$O formed around the particles.\textsuperscript{226} According to equation 14 we assume similar reaction happens in Bi$_4$B$_2$O$_9$ leading to an oxygen deficient material Bi$_4$B$_2$O$_9$-$\frac{x}{2}$ in which the less negative charge is compensated by a decrease in the Bi oxidation state from +3 to +1, however spectroscopic techniques are needed to determine it exactly.

\[ \text{Bi}_4\text{B}_2\text{O}_9 + x\text{Li}^+ + xe^- \rightarrow [\text{Bi}_{4-x}^{3+}\text{Bi}_x^{1+}]\text{B}_2\text{O}_{9-x} + \left(\frac{x}{2}\right)\text{Li}_2\text{O} \quad (14) \]

Based on our assumption of oxygen removal from Bi$_4$B$_2$O$_9$, we are now finally able to rationalize the two remaining questions, dealing with the surprisingly small polarization of ~300 mV, which is among the smallest reported for conversion type materials,\textsuperscript{228} and the low amount of carbon additive necessary for cycling. Compounds with less electronegative anions were experimentally shown to have lower polarization; the reason why hydrides display lower polarization (0.2 V)\textsuperscript{228,229} than phosphides (0.4 V),\textsuperscript{230,231} oxides (0.9 V)\textsuperscript{232,233} and fluorides (1.1 V).\textsuperscript{15,212} Along this line of thinking, Bi$_4$B$_2$O$_9$ should show a polarization comparable to oxides. However it is well known that oxygen vacancies can promote high ionic- and/or electronic conductivity in metal oxides and namely Bi-oxides.\textsuperscript{219,234,235} Thus we hypothesize that the creation of oxygen vacancies in Bi$_4$B$_2$O$_9$ during cycling is responsible for a great enhanced ionic diffusion and electronic conduction. This hypothesis is able to account for both experimental observations, the small polarization as well as the ability to reversible cycle Bi$_4$B$_2$O$_9$ with only 5 wt% conductive carbon additive.

To check the importance of ionic and electronic conductivity in Bi$_4$B$_2$O$_9$, we carried out AC/DC impedance measurements. For that a sintered pellet of Bi$_4$B$_2$O$_9$ was prepared by pressing ~300 mg of powder in a 10 mm die at a pressure of around 2.5 tons followed by an annealing for 24h at 600°C in air (around 65 to 75 % of theoretical density was obtained). After placing the pellet in the apparatus between two Pt-electrodes, AC impedance spectra were recorded for temperatures ranging from 340 up to 430°C while applying voltage amplitude of 500 mV in a frequency range between 30 MHz and 0.01 Hz during heating in air (detailed description of the setup see annexes chapter 7.1.3).
First of all, the AC complex impedance spectra for Bi$_4$B$_2$O$_9$ presents a slightly depressed semicircle over the whole measured temperature range with the absence of a Warburg diffusion resistance tail implying poor ionic conductivity. The spectra can be expressed with an equivalent circuit $R_0-R_1$/CPE$_1$--$R_2$/CPE$_2$ (Figure 92 inset) where each parallel $R$/CPE circuit gives raise to one semicircle (schematically plotted as grey lines in Figure 92). For each semicircle the resistance $R_1$ and $R_2$ (intercept with the x-axis, indicated in Figure 92) plus the corresponding capacitance $C_1$ and $C_2$ can be obtained from the fitting parameters (details see annexes chapter 7.1.3). Typically capacitive values around $10^{-12}$ and $10^{-9}$ F refer to the bulk- and grain boundary contribution respectively, allowing us to differ between them.\textsuperscript{197,198}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{impedance_spectra.png}
\caption{Representative AC complex impedance spectra of Bi$_4$B$_2$O$_9$ measured at 384°C in air. The semi circles corresponding to the bulk- and grain boundary resistance are schematically drawn as grey lines, and the real resistance values are indicated with green arrows.}
\end{figure}
To clearly separate the ionic and electronic contributions to the overall conductivity, galvanostatic DC polarization measurements were conducted (Figure 93) on sintered Bi$_4$B$_2$O$_9$ pellets sandwiched between two ionically blocking Pt-electrodes. After applying a constant current, both ions and electrons are moving, however the blocking of ions gives rise to a gradient of the chemical potential. At sufficiently long waiting time, a stationary state is reached, and the chemical potential gradient is compensated by an electrical potential gradient denoted to the polarization voltage. This polarization voltage yields to the resistance of the non-blocked charge carriers which are in our case are electrons, hence refering to the pure electronic conduction of the material. Next we applied a constant current of 5 nA across the pellet, and monitored the potential response. The DC polarization plot shown in Figure 93 indicates that a true steady state potential value is instantaneously reached implying that the charge transport in Bi$_4$B$_2$O$_9$ is carried out by electrons/ holes.

From the experimentally determined values for the AC and DC resistance, the corresponding conductivity was derived and the activation energy was obtained using the Arrhenius equation (details see annexes chapter 7.1.3). All results obtained from AC and DC measurements are summarized in the Arrhenius plot as shown in Figure 94.

The bulk ionic conductivity increases linearly from $3.77 \cdot 10^{-8}$ up to $5.31 \cdot 10^{-7}$ S·cm$^{-1}$ for 344 and 426°C (Table 14), respectively and it is associated to an activation energy of 1.28 eV. The grain boundary conductivity increases also linearly in the same temperature range
from $5.5 \cdot 10^{-10}$ to $2.43 \cdot 10^{-8}$ S·cm\(^{-1}\) (Table 14) correlated to an activation energy of 1.75 eV, however around one magnitude of order lower than the bulk conductivity. We hypothesize that the lower values for grain boundary- compared to the bulk contribution can be correlated to the formation of disorder/ defects at the grain interfaces, impeding ionic conduction.\textsuperscript{206,237,238} Regarding the electronic conductivity, it also increases linearly from $2.74 \cdot 10^{-9}$ to $1.46 \cdot 10^{-8}$ S·cm\(^{-1}\) (Table 14) in the temperature range from 364 to 406°C with an activation energy of 1.46 eV. Extrapolation of the experimental values leads to RT conductivities of $9.1 \cdot 10^{-19}$, $6.1 \cdot 10^{-22}$ and $1.1 \cdot 10^{-24}$ S·cm\(^{-1}\) for the bulk-, grain boundary- and electronic conductivity respectively. The poor extrapolated RT electronic conductivity is in agreement with the large optical bandgap of 3.02 eV calculated for Bi\(_4\)B\(_2\)O\(_9\), implying strong insulating properties.\textsuperscript{214}

\textit{Table 14:} Values for bulk-, grain boundary and electronic conductivity of Bi\(_4\)B\(_2\)O\(_9\) extracted from the AC and DC impedance measurements, recorded at certain temperatures.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>(\sigma_{\text{AC}}) (bulk) (S·cm(^{-1}))</th>
<th>(\sigma_{\text{AC}}) (grain boundaries) (S·cm(^{-1}))</th>
<th>(\sigma_{\text{DC}}) (electronic) (S·cm(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>344</td>
<td>$3.77 \cdot 10^{-8}$</td>
<td>$5.5 \cdot 10^{-10}$</td>
<td>-</td>
</tr>
<tr>
<td>364</td>
<td>$8.87 \cdot 10^{-8}$</td>
<td>$2.01 \cdot 10^{-9}$</td>
<td>$2.74 \cdot 10^{-9}$</td>
</tr>
<tr>
<td>384</td>
<td>$1.70 \cdot 10^{-7}$</td>
<td>$6.25 \cdot 10^{-9}$</td>
<td>$5.15 \cdot 10^{-9}$</td>
</tr>
<tr>
<td>406</td>
<td>$3.95 \cdot 10^{-7}$</td>
<td>$1.28 \cdot 10^{-8}$</td>
<td>$1.46 \cdot 10^{-8}$</td>
</tr>
<tr>
<td>426</td>
<td>$5.31 \cdot 10^{-7}$</td>
<td>$2.43 \cdot 10^{-8}$</td>
<td>-</td>
</tr>
</tbody>
</table>
Figure 94: Arrhenius plot of the AC and DC conductivity of Bi$_4$B$_2$O$_9$ derived in the temperature range from around 340 to 430°C during heating in air.

5.3 Conclusion

Within this chapter, bismuth oxyborate Bi$_4$B$_2$O$_9$ was studied towards its reaction mechanism versus lithium between 0.0 and 3.5 V. This material was initially chosen due to its very high density (~8.2 g·cm$^{-3}$) in order to maximize the volumetric energy density, which is presently very low for other conversion type cathode materials as they are usually mixed with ~20 wt% carbon. In the course of this study we find that it is possible to reversible cycle Bi$_4$B$_2$O$_9$ versus Li, via combined conversion/ alloying processes between 0.0 and 3.5 V (Figure 95a). We show that depending on the lower cutoff voltage, the alloying/ dealloying of Bi with Li can be circumvented and only a conversion reaction of Bi$_4$B$_2$O$_9$ can be triggered. For this reaction the reversibility between 1.0 and 3.5 V can be enhanced if the discharge is limited to 1.7 V (Figure 95b), where the material can be cycled at an average potential of 2.3 V, with therefore a fast capacity decay. Most surprisingly a small amount of carbon additive as little as 5 wt% was necessary to achieve electrochemical activity. Interestingly our
combined AC/DC impedance measurement reveal RT conductivities of $1.1 \times 10^{-24}$, $9.1 \times 10^{-19}$ and $6.1 \times 10^{-22}$ S·cm$^{-1}$ for the grain boundary-, bulk- and electronic conductivity respectively, extrapolated from high temperature measurements. On the basis of this measurement we suppose Bi$_4$B$_2$O$_9$ to be a very poor conductor for ionic and electronic transport.

However, these insulating properties are in sharp contrast to our experimental observation as Bi$_4$B$_2$O$_9$ needs only 5 wt% carbon additive for achieving attractive electrode performance polarization-wise. We believe this effect to be rooted in the Li-driven formation of Bi$_4$B$_2$O$_{9-x/2}$ at the surface of the particles showing better electronic/ ionic conductivity than the pristine material. It should be recalled that minor addition of Li to Li$_4$Ti$_5$O$_{12}$ (LTO) trigger drastically the transport properties as this compound nearly evolves from a semiconductor to a metal. We believe a similar phenomenon to be present in Bi$_4$B$_2$O$_9$ but related to oxygen content contrary to LTO.

Furthermore we observed a surprisingly small voltage polarization of ~300 mV where we hypothesize that the small value and reversibility of the conversion reaction versus Li, is most likely correlated to the appearance of Bi$_2$O$_3$ and oxygen deficient phases (in Bi$_2$O$_3$ and Bi$_4$B$_2$O$_9$) during the conversion, lowering the kinetic barriers for ionic/ electronic transport.

![Figure 95](image-url): (a) Schematic illustration of the different voltage-composition curve of Bi$_4$B$_2$O$_9$ versus Li depending on the lower cutoff potential referring to the conversion- and alloying part of the electrochemical reaction versus Li. (b) Capacity retention depending on the voltage window applied for galvanostatic cycling.

The presence of carbon within our electrodes prevents the determination of the transport properties for the oxygen deficient phase Bi$_4$B$_2$O$_{9-x/2}$. To circumvent this difficulty we have tried to prepare this non-stoichiometry by heating Bi$_4$B$_2$O$_9$ in an Ar/H$_2$ atmosphere at
moderate temperatures ~350°C. When doing so we observed the formation of metallic Bi\(^0\) as deduced from XRD. Milder reducing agents are needed. One possibility hence most probably more soft synthesis approaches are necessary. One possibility might be the reaction of Bi\(_4\)B\(_2\)O\(_9\) with CaH\(_2\) in sealed quartz tubes which allows to further decrease the reaction temperature.
6 General conclusion

This thesis was aimed to explore new borate based positive electrode materials for Li- and Na-ion batteries. Previous studies have shown very attractive electrochemical properties for LiFeBO$_3$ but also revealed its major limitations related to i) a low redox potential versus Li (~2.8 V), ii) fast degradation in ambient atmosphere and iii) poor kinetics. Moreover, it was also reported, that the redox potential of the same 3d metal can be shifted up by moving for instance from PO$_4$ to P$_2$O$_7$. This was an impetus for us to deviate from BO$_3$ based materials, and embark into an exploration of the polyborates (e.g. pyroborate B$_2$O$_5$), with the aim to design new high voltage cathode materials.

We initially focused in the synthetic exploration of pyroborate B$_2$O$_5$ and pyroborate fluoride B$_2$O$_5$F compounds, however during this survey, we encountered several synthetic difficulties in stabilizing the desired phases, even though a large variety of high- and low temperature synthetic approaches were tested. After this unsuccessful exploration, we selected Li$_6$CuB$_4$O$_{10}$ as a model compound to show the feasibility to achieve elevated redox potential towards Li in a pyroborate-based compound; such a high potential being routed in the presence of both Cu as a 3d-metal and of B$_2$O$_5$ groups as polyanions. Through this study, we first established the structural and synthetic relationship between the two reported structural polymorphs ($\alpha$- and $\beta$- Li$_6$CuB$_4$O$_{10}$) by means of thermogravimetric analysis coupled with differential scanning calorimetry (TGA-DSC) and high temperature (HT) synchrotron X-ray diffraction (XRD) measurements. Moreover we have shown that both polymorphs react with Li in a reversible way through a classical insertion/ deinsertion reaction with average redox potentials of 4.25 and 3.9 V vs. Li$^+$/Li$^0$ for $\alpha$- and $\beta$- Li$_6$CuB$_4$O$_{10}$ respectively. This activity was correlated to the Cu$^{3+}$/Cu$^{2+}$ redox couple through combined electron paramagnetic resonance (EPR) spectroscopy and density functional (DFT) calculations. Aside from this high voltage activity we reveal a conversion type reaction if the voltage is lowered down to 1.0 V, leading to the extrusion of elemental Cu nanograins as deduced by transmission electron microscopy (TEM) imaging. Finally to get an explanation for the poor electrochemical performance, bond valence energy landscape (BVEL) calculations in combination with complementary AC/DC conductivity measurements were conducted. Both polymorphs show very poor ionic/ electronic conductivities at room temperature (RT) and high activation energies for Li$^+$ migration (1.07 and 0.92 eV for the $\alpha$- and $\beta$-phase respectively). In particular the RT ionic conductivity is in the order of $10^{-13}$ S·cm$^{-1}$. 

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for the α- and β-polymorph, and electronic conductivities are around $10^{-8}$ and $10^{-13}$ S·cm$^{-1}$ respectively. Despite these very low values, we surprisingly reveal the existence of a structural transition of α-Li$_6$CuB$_4$O$_{10}$ at 350°C into a HT polymorph, with this transition be accompanied with a drastic increase in the ionic conductivity, reaching values of 1.4 mS·cm$^{-1}$ at 500°C associated to an activation energy for Li$^+$ mobility of 0.52 eV.

As we have shown the feasibility to obtain high redox potentials versus Li with borates, but also encountered strong difficulties in obtaining new phases, we exploited other borate anions, hence our interest for the family of sodium metal pentaborates Na$_3$MB$_5$O$_{10}$. We were able to stabilize two new compounds out of this family for M = Fe and Co and solve their crystal structure with the help of synchrotron XRD. These materials are basically built up of layers consisting of MO$_4$ tetrahedra connected through B$_5$O$_{10}$ units with sodium sitting in voids inside these layers as well as in between them. In light of such structural features favorable to Na$^+$ mobility, we examined their electrochemical activity versus Na and show a redox potential of 2.5 V vs. Na$^+$/Na$^0$ correlated to the Fe$^{3+}$/Fe$^{2+}$ redox couple as confirmed by ex situ Mössbauer spectroscopy. In light of a bulk ionic conductivity of 7.6·10$^{-10}$ S·cm$^{-1}$ similar to what was measured for LiFePO$_4$, we deduce that the poor electrochemical performance of Na$_3$FeB$_5$O$_{10}$ is most likely nested in its much lower RT electronic conductivity (1.2·10$^{-14}$ S·cm$^{-1}$). In contrast Na$_3$CoB$_5$O$_{10}$ is electrochemical inactive as expected from its very low RT ionic- and electronic conductivity of 4.0·10$^{-13}$ S·cm$^{-1}$ and 5.4·10$^{-17}$ S·cm$^{-1}$ respectively. Although Na$_3$FeB$_5$O$_{10}$ cannot be considered for practical applications, it stands as the first compound out of the Na$_2$O-FeO-B$_2$O$_3$ ternary system which shows a reversible reactivity towards sodium.

Owing to difficulties in designing new insertion type borate based cathode materials, we then considered other type of reaction mechanisms, namely conversion reactions. In this context we studied the conversion reaction of a bismuth oxyborate Bi$_4$B$_2$O$_9$ versus Li. This particular material was chosen as it combines elevated redox potentials versus Li due to the inductive effect of the borate group. Moreover, its high density (~8.2 g·cm$^{-1}$) offers an alternative to combat the practical low volumetric energy density of conversion type electrode materials. The material was studied by means of electrochemical methods combined with TEM and XRD measurements. Throughout the study we found that it shows an electrochemical activity centered at 2.3 V, vs. Li$^+$/Li$^0$, that is associated to the reversible reduction of Bi$^{3+}$ to Bi$^{0}$ and the formation of amorphous Li$_3$BO$_3$ and Li$_2$O. Surprisingly, such an activity comes out with a polarization of solely ~300 mV that is among the smallest reported for conversion materials, while we have only used 5wt% carbon and without
General conclusion

nanosizing. We rationalize this finding with first an electrochemical driven oxygen removal from the particle surface of Bi$_4$B$_2$O$_9$ enabling strong enhanced charge/ mass transport properties through the grain boundaries so the actual conversion reaction can proceed. After the initial formation step, a nanocomposite is achieved, in which we hypothesize the appearance of oxygen vacancies in Bi$_4$B$_2$O$_9$/ Bi$_2$O$_3$ plays a crucial role to promote ionic conductivity, hence responsible for the low polarization. If the potential window is enlarged to 0.0 - 3.5 V an additional alloying/ dealloying reactions between Li and Bi are observed at around 0.8 V. Therefore, there are poorly reversible due to very strong textural changes.

Overall within this thesis we have shown the feasibility of achieving high redox potentials versus Li in the borate family as exemplified with Li$_6$CuB$_4$O$_{10}$. As we have discussed in chapter one, the potential of a given M$^{(n+1)+}$/M$^{n+}$ redox couple is highly related to the ionicity of the M─O bond, described by the inductive effect. It was argued in literature that using condensed anionic species (BO$_3$ vs. B$_2$O$_5$ or PO$_4$ vs. P$_2$O$_7$, etc.) should further strengthen the inductive effect hence resulting in higher potentials. This explanation is consistent with our experimental finding for Li$_6$CuB$_4$O$_{10}$, however several other parameters influencing the redox potential have to be taken into account as well, namely the TM polyhedral connectivity and coordination number, and the nature of the TM itself. The situation becomes even more complicated in the particular case of borates, as it is well known that boron can adopt trigonal planar BO$_3$ as well as tetrahedral oxygen BO$_4$ coordination which can additionally be connected in an unpredictable way forming complex polyanions.

Although we are far away from designing tailor-made borates for cathode application, the multifaceted borate chemistry is prone to form open crystal structures with possible pathways for alkali-in migration. In the course of our exploration we succeeded in synthesizing two new pentaborates Na$_3$MB$_5$O$_{10}$ (M = Fe, Co) possessing an open structure with the possibility of 1D- and 2D Na$^+$ migration. However their poor electrochemical performance/ inactivity reminds us that searching for new electrode materials does not systematically led to outstanding compounds. Nevertheless during the rationalization of our findings in terms of electrochemical activity, other interesting properties popped up such as an unexpected high Li$^+$ ionic conductivity in HT-α-Li$_6$CuB$_4$O$_{10}$ described above. However, it out of the scope of this thesis due to times constraints to fully correlate this finding to the crystal structure, for which exact solution HT neutron measurements are required.

Finally through the exploration and study of the structure-property relationships in borates, we have encountered synthetic difficulties somewhat compensated by other interesting physical properties which could render these materials attractive for a broader
audience, beside the battery community. Moreover we hope to encourage researcher to still hunt for new borate based compounds, as their crystal structure can be hardly predicted so unexpected properties remain to be discovered.
7 Annexes

7.1 Electrochemical characterization

To characterize the electrochemical properties of possible electrode materials, all tests were done in 2 electrode Swagelok® half cells. A schematic view of the setup is given in Figure 96 where the positive electrode material is separated by two sheets of glass fiber discs (Whatman GF/D borosilicate) soaked with electrolyte from the anode (lithium or sodium metal). The entire assembly of the cells was conducted in an air- and water free atmosphere inside a glove box (MBraun, Germany, O₂ and H₂O < 0.1 ppm), the positive electrode material was prepared prior to use by mixing an appropriate amount of carbon (carbon SP) with the as synthesized compound by means of ball milling or hand grinding.

Figure 96: Schematic view of a Swagelok type 2 electrode cell.

For in situ XRD experiments a special homemade cell was used, adapted to the sample holder of the diffractometer (Figure 97).²⁴⁰ The assembly of parts is basically the same as for normal Swagelok type cells shown above with the main difference that the current collector for the positive electrode materials is made of Beryllium, almost transparent to X-rays. If voltages above ~3.5 V vs. Li⁺/Li⁰ are applied, the Be window is protected through a 3 μm thick aluminium foil (Goodfellow) to prevent Be from oxidation. As this type of cell setup is capable of a higher loading of the active material compared to the normal Swagelok setup,
this kind of cells were also used to prepare large quantities of \textit{ex situ} samples of positive electrode materials for further characterization. In that case the Be window was replaced by stainless steel covered with aluminium foil.

![Diagram of a modified Swagelok type 2 electrode cell](image)

1) Current collector  
2) PTFE seal  
3) Rubber seal  
4) Beryllium window  
5) electrode material  
6) Separator  
7) Li/Na metal disc  
8) Stainless steel disc  
9) Spring  
10) Mylar foil

\textbf{Figure 97:} Schematic view of a modified Swagelok type 2 electrode cell used for \textit{in situ} XRD experiments.

If not other stated, all electrochemical tests were carried out with a \textit{VMP3} potentiostat/galvanostat (\textit{BioLogic S.A., Claix, France}) at room temperature.

### 7.1.1 Galvanostatic techniques

Typically all materials were first tested in galvanostatic mode, enabling to trace the voltage-composition curve $V = f(x)$ of a certain compound. First information can be extracted from the shape of the curve (S-shape for a solid-solution process vs. plateau for a single-phase process). Furthermore, by calculated the derivative of the voltage-composition curve $-\frac{\delta x}{\delta V} = f(V)$, the redox potentials and its change upon subsequent cycling of an electrode material can be determined more precisely.
However if a cell is cycled in classical galvanostatic mode, the system in general not in equilibrium and the redox potentials are shifted compared to the theoretical one. Thus, to determine the thermodynamic equilibrium potential more precisely, galvanostatic cycling with relaxation was done. The principle of this measurement consists of alternating times of normal galvanostatic charge/ discharge steps followed by an open circuit voltage (OCV) period, where the system is allowed to reach the equilibrium potential.

### 7.1.2 Potentiostatic techniques

A potentiostatic intermittent titration technique (PITT) experiment consists in increasing the potential applied to the cell by small steps of a few mV (~10 mV) and following the evolution of the current response until it reaches a limit value (corresponding to a certain C-rate). Such an experiment provides information about the charge/discharge mechanism (solid solution vs. biphasic process) depending on the shape of the current decay.\textsuperscript{224}

### 7.1.3 Electrochemical impedance spectroscopy

AC impedance measurements were carried out using a BioLogic MTZ-35 setup with ionically blocking platinum electrodes equipped with a HTF-1100 furnace. Impedance measurements were recorded from 30 MHz to 0.01 Hz in a certain temperature range in argon or air applying a voltage amplitude from 10 to 200 mV.

Electrochemical impedance spectroscopy (EIS) is a widely used method, able to determine a variety of different phenomena ranging from electrochemical reactions, ionic diffusion etc. However, within the context of this thesis this technique is exclusively used for the determination of transport properties of solids (ionic/ electronic). In general a sinusoidal voltage is applied over a wide frequency range, and the current response is followed. The measured impedance usually contains resistive (R) and capacitive (C)/ inductive components and the data is drawn in the form of imaginary $Z''$ (capacitive) versus real $Z'$ (resistive) impedance, the so called Nyquist plot. Each parallel R/C element results in a perfect semicircle from which both values (R and C) can be extracted (Figure 98), where R values are obtained from the intercept of the semicircle with the $Z'$-axis.\textsuperscript{197}
Figure 98: Representative Nyquist plot with the equivalent circuit to referring to bulk- and grain boundaries respectively. (Adapted from Ref.\textsuperscript{[197]}).

In practice, deviations from ideal semicircles are often observed in the Nyquist plot of real materials, which can be explained by the overlap of different phenomena in the sample. This effect can be taken into account in the model of the equivalent circuit by using a constant phase element (CPE). From the fitting parameters Q, R and n of the CPE, the capacitance of the relaxation process can be estimated according to the following equation:

$$C = (R^{1-n} \cdot Q)^{\frac{1}{n}}$$  \hspace{1cm} (15)

In this study conductivity measurements were carried out using a BioLogic MTZ-35 impedance analyzer with ionically blocking electrodes, equipped with an HTF-1100 furnace upon heating and cooling under a flow of argon. For AC measurements, a voltage amplitude ranging usually from 10 to 200 mV over a frequency range from 35 MHz to 0.1 Hz was applied and the current response was followed.

The depressed semi-circles derived from the complex AC impedance spectra were fitted with a corresponding equivalent circuit, where $R_0$ represents an initial shift of impedance arc from zero, $R_1$ is the bulk- and $R_2$ grain boundary resistance (in case it is
possible to distinguish between them), CPE stands for a constant phase element. The conductivity \( \sigma \) was calculated from the resistance \( R \), the pellet’s thickness \( l \), and area \( A \) according to the following equation:

\[
\sigma = \frac{l}{A \cdot R} \tag{16}
\]

The activation energy \( E_a \) (eV) for \( \text{Li}^+ \) migration and electron conduction was calculated from fitting the experimentally derived values using the Arrhenius equation,

\[
\sigma_T = \sigma_0 \cdot e^{-\left(\frac{E_a}{k_B T}\right)} \tag{17}
\]

where \( \sigma_T \) (S·cm\(^{-1}\)·K) is the temperature dependent conductivity, \( \sigma_0 \) (S·cm\(^{-1}\)) is a pre-exponential factor, and \( k_B \) is the Boltzmann constant. Note that the conductivity values are not corrected by the pellet porosities, thus the values are underestimated.

### 7.1.4 Direct current polarization measurements

The current/voltage of a mixed conductor if a DC potential \( (U) \) or current \( (I) \) is applied corresponds to the dc resistance \( (R_{DC}) \), where \( R_{el} \) and \( R_{ion} \) represent the partial electronic and ionic resistances respectively:

\[
\frac{I}{U} = \frac{1}{R_{DC}} = \frac{1}{R_{el}} + \frac{1}{R_{ion}} \tag{18}
\]

The partial resistances can be extracted when selective reversible or blocking electrodes are used (Pt as reversible electrode for electrons and blocking electrode for ions). The blocking of one charge carrier leads to a polarization and a gradient of potential of the blocked component. At sufficiently long waiting times, a steady state is reached which accounts only for the electronic part of the charge transport, if ionically blocking electrode are used.\(^{236}\)
Out of the steady state current value the corresponding resistance can be calculated according to Ohm’s law:

\[ R = \frac{U}{I} \]  \hspace{1cm} (19)

The corresponding conductivity \( \sigma \) was calculated according to the formula given in equation 19. DC polarization measurements were done on similar pellets used for AC impedance spectroscopy with platinum blocking electrodes, by applying a constant voltage or current till the current- or potential response reached a constant value, respectively. The same setup like for the AC impedance measurements was used, with the difference that the applied potential/ current was controlled/ recorded with a VMP3 potentiostat (BioLogic S.A., Claix, France).
7.2 Structural characterization

X-ray diffraction (XRD) was used within this thesis as a regular characterization method to verify the purity of the samples or to perform Rietveld refinement of structural models proposed in literature. All powder patterns were refined using the Rietveld method\textsuperscript{241} as implemented in the FullProf program.\textsuperscript{242}

The crystal structures show in this thesis were drawn with the \textit{VESTA} visualization program.\textsuperscript{243} Data that were not obtained from our refinements was received as CIF files from the \textit{ICSD} database.\textsuperscript{244}

7.2.1 Laboratory XRD

Laboratory XRD patterns were recorded in Bragg-Brentano geometry with a \textit{Bruker D8 Advance} diffractometer equipped with a copper source ($\lambda_{\text{Cu-Ka1}} = 1.54056$ Å, $\lambda_{\text{Cu-Ka2}} = 1.54439$ Å) and a LynxEye detector. If samples were air/moisture sensitive, a homemade sample holder was used which, covered with a Kapton window, which could be sealed air tight inside the glove box.

Temperature-controlled XRD experiments were carried out at the same diffractometer described above, equipped with an \textit{Anton Paar HTK1200} furnace chamber.

7.2.2 Synchrotron XRD

In order to get more precise structural information of the samples, selected samples were measured either at the 11-BM beamline at the Argonne National Laboratory through the mail-in service of the 11-BM beamline of the Advanced Photon Source (APS) at Argonne National Laboratory (ANL, Argonne, USA) with a wavelength of $\lambda = 0.4139$ Å.

HT synchrotron XRD was conducted at the ID 22 – High resolution powder diffraction beamline at the European Synchrotron Radiation Facility (ESRF) using a wavelength of $\lambda = 0.4001$ Å.
7.3 Other physical characterization

7.3.1 Thermal analysis

Thermal stability and phase transition were investigated using thermogravimetric analysis coupled with differential scanning calorimetry (TGA-DSC). The data was recorded with a STA 449C Netzsch apparatus. Around 20-30 mg of sample was utilized which was heated/ cooled in flowing air or argon with heating rates from 1 to 10°C/min.

7.3.2 EPR spectroscopy

EPR spectroscopy is a technique to study materials with unpaired electrons. As unpaired electrons possess a magnetic moment, they adopt discrete energy levels, if an external magnetic field is applied (also called Zeeman effect). To induce a transition between these two energy levels, the probe is subjected to microwave radiation. The absorption signal presents the energy required for the transition, and is described as its first derivative (Figure 100).

![Figure 100: Energy levels for an electron with spin (S= ±1/2) in an applied magnetic field B, where it can adopt two different states (α and β), and the corresponding absorption spectrum and its first derivative.](image)
One important parameter which can be extracted from EPR measurements, is the so-called g-factor. It can give information about the electronic structure of the paramagnetic center.

The spectra were recorded at room temperature on a Bruker ELEXSYS E580 spectrometer using X-band continuous wave (CW) mode. Microwave power and modulation amplitude were respectively set to 2 mW and 1 G. Converse time and time constant were set to be 40.96 ms and 20.48 ms, respectively. The weight of samples and measuring Q values were normalized for the spectra to analyze the intensity of all paramagnetic species.

### 7.3.3 Mössbauer spectroscopy

Mössbauer spectroscopy is a versatile spectroscopic technique which can provide information about the chemical, structural magnetic properties of a material, mainly used to probe $^{57}$Fe. It is based on the “Mössbauer effect”, a recoilless gamma ray emission and absorption, which implies a transition between the ground- and a first excited state of a nucleus. Typically three types of nuclear interactions are observed, which are also called hyperfine interactions: i) isomeric shift, ii) quadrupole splitting and iii) magnetic splitting (Figure 101). If source and absorber nucleus are identical, a simple spectrum is observed, with a single absorption peak at 0 mm/s velocity. Depending on the electronic configuration or the ligand environment, the absorption peak or the center of the spectrum is shifted from zero respectively, referring to the isomeric shift and quadrupole splitting. Typically the Mössbauer spectrum of Fe$^{2+}$ presents an isomer shift around 1.0 mm/s and a quadrupole splitting ranging from 0.2 to 3.5 mm/s, while the doublet of Fe$^{3+}$ has an isomer shift close to 0.3-0.4 mm/s and a quadrupole splitting smaller than 2 mm/s.\textsuperscript{246,247}
Figure 101: The energy changes are different in the source (S) and absorber (A) as a result of different electron densities at the source and absorber nuclei, the result is manifested as isomer shift in the Mössbauer spectrum.247

$^{57}$Fe Mössbauer spectra were collected using 20-30 mg/cm² powder. The spectrometer is operated in the constant acceleration mode and in transmission geometry. A $^{57}$Co(Rh) source with a nominal activity of 370 MBq was used. The source and the absorber were always kept at 295 K. All the isomer shifts are given relative to $\alpha$-Fe standard at 295 K.

7.3.4 Scanning- and transmission electron microscopy

High Resolution transmission electron microscopy (TEM) and scanning transmission electron microscopy (STEM) was done in collaboration with two different persons. In both cases samples for TEM was prepared by grinding the material in an agate mortar and dipping a Cu holey carbon TEM grid into the powder. The microscope was equipped with a high brightness source X-FEG Schottky and a Cs probe corrector (DCOR). The chemical mapping EDS is acquired with Super-X detector system based on 4 windowless Silicon Drift Detector (SDD), which enhanced acquisition efficiency and spatial resolution. All observations were performed at 200 kV on nickel TEM grids with carbon films (300 mesh).

Further TEM data (SAED patterns, STEM-EDX elemental maps and the HRTEM images) were acquired on an FEI Osiris microscope operated at 200 kV and equipped with a Super-X EDX detector. In this case a special Gatan vacuum transfer holder was used for the analysis.
7.3.5 Density functional theory calculations

Spin-polarized DFT+U calculations were performed using the VASP code (Vienna Ab initio Simulation Package)\textsuperscript{248,249} and using the rotationally invariant Dudarev method\textsuperscript{250} and the generalized gradient approximation with PBE functional to describe electron exchange and correlation.\textsuperscript{251} The effective Hubbard corrections were taken from the literature for the copper d-electrons ($U_{\text{eff}}=4.0$ eV).\textsuperscript{252} A plane-wave cut-off of 600 eV was used to define the basis set, with well-converged k-point sampling for each compound. The COOPs were computed using the Lobster program developed by Dronskowski and coworkers.\textsuperscript{253–255}

7.3.6 Bond valence energy landscape calculations

The Bond Valence Energy Landscape (BVEL) method allows the calculation and visualization of ion diffusion pathways of a mobile species in a 3D framework and is often applied for identifying potentially interesting ionic conductors. The BVEL method is based on the Bond Valence Sum (BVS) approach, which calculates the oxidation state of an atom as a function of the distances to its neighboring atoms and their oxidation state, and is an extension of the Bond Valence Sum Maps (BVSM). To determine the possible ion transport pathways in a structure, the theoretical oxidation state of the mobile ion is calculated as a function of a position in the structure, where positions with a low valence mismatch (low deviation from the ideal oxidation state $\sim$10\%) are regarded as part of the diffusion pathway. In contrast to the BVS approach, the BVEL calculations transform the valence units (v.u.) given for the BVS into energy units (eV). Furthermore, soft bond valence (softBV) parameters are applied, which allow to take into account the polarizability of the mobile species as well as the influence of the counterions on the ion mobility. Further included in the BVEL calculations is the cut-off length to the surrounding coordination spheres, which in our case is set to 8 Å.\textsuperscript{183,256}

BVEL calculations are performed by the BondStr software as implemented in the Fullprof suite.\textsuperscript{242} The program calculates the isosurfaces in the structure, which are accessible for the mobile ion and which must be infinitely connected in at least one direction to permit diffusion through the structure. From the difference of the calculated minimum energy isosurface and the minimum energy value inside this isosurface, we obtain the “activation energy” necessary for the ion diffusion in this structure. The isosurface can be visualized with the VESTA program.
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