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Lithium-Rich Rock Salt Type Sulfides-Selenides (Li$_2$TiSe$_x$S$_{3-x}$): High Energy Cathode Materials for Lithium-Ion Batteries

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Abstract: Lithium-rich disordered rocksalt Li$_2$TiS$_3$ offers large discharge capacities (>350 mAh·g$^{-1}$) and can be considered a promising cathode material for high-energy lithium-ion battery applications. However, the quick fading of the specific capacity results in a poor cycle life of the system, especially when liquid electrolyte-based batteries are used. Our efforts to solve the cycling stability problem resulted in the discovery of new high-energy selenium-substituted materials (Li$_2$TiSe$_x$S$_{3-x}$), which were prepared using a wet mechanochemistry process. X-ray diffraction analysis confirmed that all compositions were obtained in cation-disordered rocksalt phase and that the lattice parameters were expanded by selenium substitution. Substituted materials delivered large reversible capacities, with smaller average potentials, and their cycling stability was superior compared to Li$_2$TiS$_3$ upon cycling at a rate of C/10 between 3.0–1.6 V vs. Li$^+$/Li.

Keywords: high-energy materials; sulfides; anionic redox; wet mechanochemistry; selenium substitution; cyclic voltammetry

1. Introduction

The rapid growth in electric vehicle market requires high performance, safe, and low-cost battery packs that should enable driving ranges exceeding 500 km. Current Lithium-ion battery positive electrode materials can realize this objective by providing specific energies that exceed 250 Wh·kg$^{-1}$ at cell level [1–4]. These materials still contain a small percentage of cobalt, and the concerns related to this critical element have increased, due to its price and availability. Recent research efforts have focused on the discovery of Co-poor or even Co-less positive electrode materials and the development of alternative high-energy materials, such as lithium sulfide and cation-disordered rocksalts [5–20]. Among them, cation-disordered rocksalts have recently received great interest, as a new generation of positive electrode materials for lithium ion batteries [16,17,21,22]. For instance, disordered rocksalt sulfides deliver high specific capacities (>400 mAh·g$^{-1}$) at relatively low operating potentials (~2.2 V vs. Li$^+$/Li), and their energy density is competitive with conventional layered materials [19]. This low operating potential may be considered an advantage of sulfide-type batteries, since they do not experience the serious electrolyte degradation problems occurring at higher voltages [23]. In addition, these materials allow the exchange of more than one Li per metal cation (multi-electrons redox reactions); thus, greater capacities could be produced [19,24,25]. Such a property was already noticed and reported in earlier studies, as TiS$_3$ was able to host three Li$^+$ ions during discharge [26,27]. However, only one lithium ion was reversibly intercalated [26].

More recently, new studies have paved the way for the discovery of novel Li-rich disordered rocksalt sulfides. Sakuda et al. highlighted that Li$_2$TiS$_3$ and Li$_3$NbS$_4$ materials with a disordered rocksalt cubic structure can provide capacities above 400 mAh·g$^{-1}$ upon cycling between 3–1.5 V vs. Li$^+$/Li [19,24]. Despite their promising reversible capacities, the retention performances of Li$_2$TiS$_3$ and Li$_3$NbS$_4$ were rather poor in conventional cells;
on the contrary, superior retention performances were achieved in solid-state cells [16]. For industrial applications, materials should have both a long shelf life and a high capacity. The retention performance of Li$_2$TiS$_3$ is a major bottleneck, and new approaches are necessary to improve this. Partial substitution mechanisms may be promising solutions for this issue. For instance, sulfur anions could be replaced with an alternative anion. Studies show that selenium substitution can improve the cycling stability of Li$_2$S cells [28,29]; therefore, such results motivated us to substitute sulfur with selenium in Li$_2$TiS$_3$. Thanks to an optimized wet milling process, two new lithium-rich rock salt type sulfides–selenides compounds (Li$_2$TiSe$_x$S$_{3-x}$) are reported in this study, for the first time. Structural investigations, as well as electrochemical tests have been performed, to compare the mechanism of lithium insertion/ extraction between these new compounds and the previously published Li$_2$TiS$_3$ cathode material [19].

2. Materials and Methods

2.1. Wet Mechanochemical Synthesis of Li$_2$TiSe$_x$S$_{3-x}$

According to the synthesis procedure; 1.5 g of precursors composed of lithium sulfide (Li$_2$S, Sigma Aldrich, St. Louis, MO, USA, 98%), titanium powder (Alfa Aesar, Havrier, MA, USA, 99.98%), sulfur powder (Sigma Aldrich, 99.99%), and selenium powder (Alfa Aesar, 99.9%) precursors were placed into a 50-mL zirconia jar that contained 285 zirconia balls (5 mm in diameter). Then, an appropriate amount of anhydrous hexane (99.9%, Sigma Aldrich) was added, until covering the entire ball surface. Afterwards, the jar was closed tightly and transferred from an argon-filled glovebox to the planetary ball milling machine (Retsch PM 100). Milling was operated at 510 rpm for 20 hours. Powder was recovered in an argon-filled glovebox, to limit air contamination in the powders. Zirconia impurities due to high-energy milling were checked using inductively coupled plasma atomic emission spectroscopy measurements and were considered low (0.018 mole of ZrO$_2$ for 1 mole of Li$_2$TiS$_3$).

2.2. Structural Characterizations

X-ray diffraction (XRD) analyses were carried out on a BRUKER AXS D8 diffractometer, using Cu Kα ($\lambda = 1.5406$ Å) anticathode, where the sample was scanned in the range of 8–80° using a step size of 0.02°. As the sulfide materials are air sensitive, all of the samples were prepared in a glovebox, and the sample surface was protected with Kapton®. A signal due to Kapton® film could be detected at small 2θ angles (0–28°) in the diffraction patterns of the powders.

Scanning electron microscopy (SEM) and energy-dispersive X-ray spectroscopy (EDX) were performed with a microscope, Zeiss brand MEB LEO 1530 Gemini.

Chemical characterization by X-ray photoelectron spectroscopy (XPS) was carried out using a Versaprobe II ULVAC-PHI spectrometer. A monochromatic beam (X-ray source Al Kα 1486.6 eV) of 100 μm diameter and 25 W of power was focused on the surface of the samples. Survey spectra were measured over a spectral range of 0–1200 eV, to identify the elements present in the material using a pass energy of 117 eV, which corresponds to a resolution of 1.6 eV. High resolution spectral analyses were performed using a pass energy of 23 eV, which corresponds to a resolution of 0.5 eV. All XPS measurements were carried out in an ultrahigh vacuum chamber (7 × 10$^{-10}$ mbar). All XPS spectra binding energies were corrected using the C1s line of alkyl groups in the C-C at 285.0 eV. Curve fitting and background subtraction were accomplished using Casa XPS software. The spectra curve fitting was performed using the pseudo-Voigt function, product of Gaussian (80%), and Lorentzian (20%) distributions.

2.3. Electrochemical Characterizations

Electrode preparation: In an Ar-filled glove box, positive electrodes were prepared using 80 wt.% of the active material, 10 wt.% Carbon SuperP (Timcal, Willebroek, Belgium), 10 wt.% of PVdF (Solvay, Brussels, Belgium). An adequate amount of NMP (Sigma Aldrich)
was added, to form a uniform slurry, then it was coated on an aluminum foil (20 μm) using a 100 μm doctor blade and left to dry in the glovebox for three days. The electrodes were cut (14 mm diameter), pressed (10 t), weighed (loaded > 2 mg) in the glovebox, and dried in a Buchi® oven at 60 °C for 12 h under vacuum.

Electrochemical testing: Coin cells were prepared using 14-mm diameter positive electrodes, 16-mm diameter Li foil as a negative electrode, and 150 μL electrolyte containing 1 M LiPF₆ in EC:PC:DMC, where EC: ethylene carbonate; PC: propylene carbonate; DMC: dimethyl carbonate, in a volumetric ratio of 1:1:3, respectively. A propylene separator (Celgard® 2400, 16.5 mm diameter) and polyolefin separator (Viledon®, 16.5 mm diameter) were used in the coin cells. Three-coin cells were cycled at 22 °C, at a rate of C/10 between 3 V and 1.5 V, using ARBIN cycling instrumentation.

3. Results

To prepare selenium substituted Li₂TiS₃ compositions (Li₂TiSeₓS₃₋ₓ), we developed a flexible and easy process utilizing ball-milling, in which precursors are used in their neutral form, as neither Li₂Se nor TiSe₂ precursors were commercially available [30]. Our synthesis process does not require applying multiple steps to prepare the ceramic powders [28]. The powder diffraction patterns of the three resulting powders (Li₂TiSeₓS₃₋ₓ) are presented in Figure 1a.

With our original synthesis process (wet mechanochemistry), Li₂TiS₃ was successfully prepared as a disordered rocksalt phase. Selenium substituted samples showed a similar peak profile as Li₂TiS₃, indicating that Li₂TiSeₓS₃₋ₓ compositions were also prepared in disordered rocksalt phase.

At first observation, the diffraction peaks of Li₂TiSeS₂ and Li₂TiSe₂S patterns shifted to small 2θ angles, indicating that the lattice parameters expanded with selenium substitution. This behavior was expected, as Se²⁻ is bigger than S²⁻. The lattice parameters of Li₂TiSeₓS₃₋ₓ powders were refined with Rietveld refinement in the Fullprof program [31]. All of the peak profiles are illustrated in Figure 1d–f, and the refined parameters and the resulting structure is shown in Figure 2. The lattice parameters of Li₂TiS₃ (5.0831 (8) Å) were found to be slightly bigger than those previously reported (5.05 Å) [16]. Second, lattice parameters of Li₂TiSeS₂ and Li₂TiSe₂S were found (5.1729 (8) Å) and (5.2459 (1) Å), respectively.

In addition, the peak intensity ratios of Li₂TiSeS₂ and Li₂TiSe₂S differed from the peak intensity ratio of Li₂TiS₃ powder, as can be clearly seen on the intensity of the (311) and (222) reflections around 60°. We, thus, simulated the diffraction patterns of Li₂TiSeS₂ and Li₂TiSe₂S using the refined cell parameters and a similar structure as for Li₂TiS₃ (Figure 1b). The same evolution in peak intensity ratios was observed with selenium substitution, confirming the Se substitution. For the Rietveld refinements, owing to the poor crystallinity of the material and the presence of a broad signal coming from the Kapton® protective film, it was decided to not refine the atomic occupancies and keep the model structures. However, without further refining, the agreement between the observed and calculated pattern was already quite good and confirmed that Se substitution had occurred in these samples.

The Figure 3 shows SEM images of Li₂TiSeₓS₃₋ₓ powders.

All powders were composed of agglomerates of primary nanoparticles, and, curiously, the Li₂TiSeₓS₃₋ₓ agglomerates tended to have a plate-like morphology.

The electrochemical performances of Li₂TiS₃ and selenium-substituted samples (Li₂TiSe₂S and Li₂TiSe₂S) were investigated and compared in half-cells (Figure 4).

The open circuit potential of Li₂TiS₃ was measured as 2.31 V at the beginning of the cycling. When the Li₂TiS₃ was charged with a C-rate of C/10 until 3 V, the 1.77 Li⁺ ions were extracted from Li₂TiS₃ and a capacity of 300 mAh·g⁻¹ was delivered. During discharge, 2.0 Li⁺ ions were inserted into the structure, the composition changed into Li₂.75TiS₃, and a capacity of 339 mAh·g⁻¹ was delivered. The average charge and discharge potentials were measured at 2.46 V and 2.23 V, respectively, and a summary of the results is shown in Table 1.
Table 1. Summary of electrochemical performances of Li$_2$TiSe$_{3-x}$S$_x$ (0 ≤ x ≤ 2) materials.

<table>
<thead>
<tr>
<th>Composition</th>
<th>Lattice Parameter a (Å)</th>
<th>1st Charge Capacity (mAh·g$^{-1}$)</th>
<th>1st Discharge Capacity (mAh·g$^{-1}$)</th>
<th>Average Charge Potential (V vs. Li$^+$/Li)</th>
<th>Average Discharge Potential (V vs. Li$^+$/Li)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li$_2$TiS$_3$</td>
<td>5.0831</td>
<td>300</td>
<td>339</td>
<td>2.46</td>
<td>2.23</td>
</tr>
<tr>
<td>Li$_2$TiSeS$_2$</td>
<td>5.1729</td>
<td>179</td>
<td>310</td>
<td>2.34</td>
<td>2.06</td>
</tr>
<tr>
<td>Li$_2$TiSe$_2$S</td>
<td>5.2459</td>
<td>149</td>
<td>379</td>
<td>2.24</td>
<td>1.98</td>
</tr>
</tbody>
</table>

Figure 1. (a)/XRD patterns of Li$_2$TiSe$_{3-x}$S$_x$ (0 ≤ x ≤ 2) materials—(b)/Simulated XRD patterns of Li$_2$TiSe$_{3-x}$S$_x$ powders and (c)/experimental and simulated peak intensity ratio of (311) and (222) planes, I$_{311}$/I$_{222}$ for Li$_2$TiS$_3$, Li$_2$TiSeS$_2$, and Li$_2$TiSe$_2$S powders—(d–f)/XRD pattern of Li$_2$TiSe$_{3-x}$S$_x$ (0 ≤ x ≤ 2) Rietveld refinement in the Fullprof program. Brackets with asterisk indicate the zone where the protective Kapton® foil signal prevented the refinement from being carried out.
with 2·(0.885−0.123) Refined parameters of Li2TiSexS3

<table>
<thead>
<tr>
<th>Formula</th>
<th>Space group</th>
<th>cell parameters</th>
<th>atom</th>
<th>Wyckoff position</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>occupancy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li2TS3</td>
<td>Fm-3m</td>
<td>a=b=c=5.0813(8) Å</td>
<td>Li1</td>
<td>4a</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.66</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Ti1</td>
<td>4a</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.33</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Se1</td>
<td>4b</td>
<td>1/2</td>
<td>1/2</td>
<td>1/2</td>
<td>1.00</td>
</tr>
<tr>
<td>Li2TiSeS2</td>
<td>Fm-3m</td>
<td>a=b=c=5.1729(8) Å</td>
<td>Li1</td>
<td>4a</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.66</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Ti1</td>
<td>4a</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.33</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Se1</td>
<td>4b</td>
<td>1/2</td>
<td>1/2</td>
<td>1/2</td>
<td>0.33</td>
</tr>
<tr>
<td>Li2TiSe2S</td>
<td>Fm-3m</td>
<td>a=b=c=5.2459(1) Å</td>
<td>Li1</td>
<td>4a</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.66</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Ti1</td>
<td>4a</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.33</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Se1</td>
<td>4b</td>
<td>1/2</td>
<td>1/2</td>
<td>1/2</td>
<td>0.33</td>
</tr>
</tbody>
</table>

Figure 2. (a) Refined parameters of Li2TiSexS3−x (0 ≤ x ≤ 2) materials and (b) Li2TiSeS2 structure.

Figure 3. SEM images of Li2TiSexS3−x (0 ≤ x ≤ 2) materials.

Figure 4. Galvanostatic curves of (a) Li1.23TiS3, (b) Li1.23TiSeS2, and (c) Li1.23TiSe2S cathode materials in half-cell at a rate of C/10 between 3 and 1.5 V vs. Li+/Li, the first three cycles were represented in the figures.

The Li2TiS3 material prepared using our original synthesis process delivered greater charge capacities, but slightly smaller discharge capacities, than the Li2TiS3 material previously reported [16]. Moreover, we observed that Li2TiS3 provided a greater discharge capacity at the initial cycle, but the charge–discharge curve was fully reversible at the end of the second and third cycles.

\[
\text{Charge: Li}_2\text{Ti}^{4+}\text{S}_3^{2−} \rightarrow \text{Li}_0.23\text{Ti}^{4+}(\text{S}^{2−})_{1.23}(\text{S}_2^{2−})_{0.885} + 1.77\text{Li}^{+} + 1.77\text{e}^{−} \tag{1}
\]

\[
\text{Discharge: Li}_{0.23}\text{Ti}^{4+}(\text{S}^{2−})_{1.23}(\text{S}_2^{2−})_{0.885} + 2\text{Li}^{+} + 2\text{e}^{−} \rightarrow \text{Li}_{2.23}(\text{Ti}^{3+})_{x}(\text{Ti}^{3+})_{1−x}(\text{S}^{2−})_{3−y}(\text{S}_2^{2−})_{y} \tag{2}
\]

with 2·(0.885 − y) + x = 2.
Such greater discharge capacity at the initial cycle was previously reported for both \( \text{Li}_2\text{TiS}_3 \) and other compositions (\( \text{Li}_3\text{NbS}_4 \) and \( \text{Li}_3\text{SnS}_4 \)) \[16,24,25\]. We have to keep in mind that this phenomenon will require the addition of “lithium sacrificial salt” inside Li-ion cells if this material is used in front of a graphite anode instead of metal lithium. Another way to overcome this phenomenon could be to use the material in all solid-state batteries, with metal lithium as anode. In all cases, the origin of the extra capacity remains unknown, and further structural analyses are needed to explain it.

We explain the possible redox mechanism in \( \text{Li}_2\text{TiS}_3 \) with a hypothesis whereby all the atoms are stabilized at their valence states (Ti\(^{4+} \) and S\(^{2-} \)) in the pristine electrode, and there is no loss of sulfur atoms during the charge–discharge process. Based on this hypothesis, only sulfur redox should be active during charging, since Ti is already at its maximum valence of Ti\(^{4+} \). Therefore, charge capacity may have been produced by anionic (sulfur) redox, with some part of the S\(^{2-} \) atoms oxidized into S\(_2\)^2 (Equation (1)). Moreover, the discharge capacity could have been provided by either anionic redox (S\(^{2-} / \text{S}_2\)^2) or both anionic and cationic (Ti\(^{4+} / \text{Ti}^{3+} \)) redox processes. A reversible sulfur redox process was previously mentioned in the literature \[32\]. Reversible formation and dissociation of covalent S-S bonds in an \( \text{Li}_2\text{TiS}_3 \) electrode were detected in both pair distribution function analyses and ab initio molecular dynamics calculations, and this was later attributed to reversible sulfur redox processes.

\( \text{Li}_2\text{TiSeS}_2 \) and \( \text{Li}_2\text{TiSe}_2\text{S} \) cells were also tested with the same cycling schedule applied to \( \text{Li}_2\text{TiS}_3 \) cells, and 179 mAh g\(^{-1} \) charge and 310 mAh g\(^{-1} \) discharge capacities were delivered between 3 and 1.5 V vs. \text{Li}^+/\text{Li}\. The average charge and discharge potentials were measured to be 2.34 V and 2.06 V, respectively, indicating that the average potentials are reduced by selenium substitution. At the end of the first cycle, we again detected a large discharge capacity. More than 2.0 Li\(^+ \) ions were inserted into the structure, and the composition changed into \( \text{Li}_{3.08}\text{TiSeS}_2 \), which can be regarded as the average (theoretical) composition. Such capacity was even greater than the theoretical capacity of \( \text{Li}_2\text{TiSeS}_2 \) cells (261 mAh g\(^{-1} \) based on two electron exchange processes). At the subsequent cycles, a reversible cycling curve was observed. If we keep the same hypothesis that we previously used to explain the redox process of \( \text{Li}_2\text{TiS}_3 \) cells, the charge capacity should be produced by either sulfur (S\(^{2-} / \text{S}_2\)^2) or selenium (Se\(^{2-} / \text{Se}_2\)^2) redox processes, which is active between 3 and 1.5 V \[29,30\]. Here, again, the discharge capacity should be provided by either anionic (S\(^{2-} / \text{S}_2\)^2 or Se\(^{2-} / \text{Se}_2\)^2) or both anionic and cationic (Ti\(^{4+} / \text{Ti}^{3+} \)) redox processes.

\( \text{Li}_2\text{TiSeS}_2 \) cells delivered charge and discharge capacities of 149 mAh g\(^{-1} \) and 379 mAh g\(^{-1} \), respectively. Much lower average charge and discharge potentials (2.24 V and 1.98 V) were detected. At the end of the initial discharge, 3.44 Li\(^+ \) ions were inserted into the cubic rocksalt structure of \( \text{Li}_2\text{TiSe}_2\text{S} \) and the theoretical composition became equivalent to \( \text{Li}_{4.17}\text{TiSe}_2\text{S} \). We again detected extra discharge capacity, and this was even greater than the theoretical capacity of \( \text{Li}_2\text{TiSe}_2\text{S} \) (213 mAh g\(^{-1} \) based on two electron exchange processes). Now, we cannot explain more than a three Li\(^+ \) uptake with the same hypothesis; a combination of anionic and Ti\(^{3+} / \text{Ti}^{4+} \) redox processes. In the cycling curve of \( \text{Li}_2\text{TiSe}_2\text{S} \) cells, we observed that the second and third cycles were reversible; however, smaller charge and discharge capacities, as well as a rapid capacity fading, were detected.

To describe the possible redox processes taking place in \( \text{Li}_2\text{TiSe}_x\text{S}_{3-x} \) cells, we conducted cyclic voltammetry measurements (Figure 5).

\( \text{Li}_2\text{TiS}_3 \) cells showed one oxidative and one reductive peak that resulted from sulfur redox reaction (S\(^{2-} / \text{S}_2\)^2 and S\(_2\)^2 / S\(^{2-} \)) at 2.69 V and 2.28 V. Moreover, 2.51 V charge and 2.20 V discharge potentials were detected in \( \text{Li}_2\text{TiSe}_2\text{S} \) cells, and 2.43 V charge and 2.12 V discharge potentials were detected in \( \text{Li}_2\text{TiSe}_2\text{S} \) cells, in accordance with previous results showing a reduced working potential for Se-substituted materials. During discharge, the apparition of a second reduction peak could be observed as a shoulder of the main peak, only for the Se-substituted materials. This peak is reversible and can be observed in the second cycle of \( \text{Li}_2\text{TiSe}_x\text{S}_{3-x} \) cells. Therefore, we can assume that the substitution of S by
Se in the Li$_2$TiSe$_x$S$_{3-x}$ materials leads to the activation of a new redox process, which is the origin of the extra capacity.

![Figure 5](image_url)  
**Figure 5.** Redox potentials comparison of Li$_2$TiS$_3$, Li$_2$TiSeS$_2$, and Li$_2$TiSe$_2$S electrodes at the first (5 $\mu$V·s$^{-1}$) (a), and second (10 $\mu$V·s$^{-1}$) (b) cycles between 3 and 1.5 V vs. Li$^+$/Li.

To investigate the effect of this supplementary process on the cycling stability of the materials, we carried out galvanostatic cyclings with two different low voltage cut-offs: one with a cut-off at 1.5 V, allowing the cycling on the supplementary process; and one at 2 V, to avoid the major part of this process. The results presented in Figure 6 show that the two substituted samples clearly benefited from the reduced voltage window. The improvement is more important for Li$_2$TiSe$_2$S, for which the low potential process represents a bigger part of the redox processes, with the capacity retention at the 14th cycle improving from 12% to 76%. It is, thus, reasonable to assume that the supplementary process is a major factor in the performance degradation of these materials. The stability of the non-substituted material was not significantly modified; this was expected, as no major redox process occurs in the 1.5–2 V region for this material.

![Figure 6](image_url)  
**Figure 6.** Capacity retention (%) of Li$_2$TiS$_3$, Li$_2$TiSeS$_2$, and Li$_2$TiSe$_2$S cells cycled at a rate of C/10 in different cycling windows: (a) 3–1.5 V and (b) 3–2.0 V vs. Li$^+$/Li.

To investigate why Se-containing material cells failed upon cycling, and to get some hint of the nature of the low-potential process, we conducted ex situ structural analyses of the electrodes at different states of charge. The results of Li$_2$TiS$_3$, Li$_2$TiSeS$_2$, and Li$_2$TiSe$_2$S are shown in Figure 7. For both samples, we observed a reversible structural change in the diffraction patterns of the electrodes: the cubic structure became amorphous at the end of the charge, then it recrystallized to a disordered rocksalt form at the end of the discharge. Such flexible structural change in electrodes was previously reported for Li$_2$TiS$_3$ [19]. Differently from Li$_2$TiS$_3$, metallic selenium was detected in selenium substituted electrodes at the end of the charge, and disappeared at the end of the discharge (Figure 7).
Moreover, the morphologies of electrodes at the different states of charge were observed with scanning electron microscopy (SEM). In Figure 8, we can see similar electrode morphologies after charge and discharge of the Li$_2$TiS$_3$ electrodes. In both Li$_2$TiSeS$_2$ and Li$_2$TiSe$_2$S electrodes, a columnar phase was observed at charged electrodes, whereas only one phase was observed in the discharged electrodes. Such a result is in accordance with the ex situ XRD result; metallic selenium (Se$_0$) was formed at the end of the charge and no longer detectable at the discharged state. This disappearing could be due to the dissolution of the Se in the form of lithium polyselenides, which would explain the low potential process. To confirm the nature of this extra phase, a Li$_2$TiSe$_2$S electrode was maintained for 24 h at 3.0 V in a potentiostatic mode, to favor the hypothetical formation of Se, and then observed by microscopy. Large agglomerates were observed, which had different habitus from the starting phase (Figure 9). Their analysis by EDX confirmed that these particles were solely composed of Se.

Figure 7. Ex situ X-ray diffractograms of (a) Li$_2$TiS$_3$, (b) Li$_2$TiSeS$_2$, and (c) Li$_2$TiSe$_2$S electrodes at different states of charge: pristine, at the end of charge (3 V), and at the end of discharge (1.6 V).

Figure 8. SEM images of Li$_2$TiS$_3$, Li$_2$TiSeS$_2$, and Li$_2$TiSe$_2$S electrodes at different states of charge: pristine, at the end of charge (3 V) and at the end of discharge (1.6 V).
First, it is possible to notice that the Ti3s signals in the Se-substituted samples are very weak. We suggest the hypothesis that this phenomenon is due to the Se3d cross-section, which is higher than the Ti3S one (respectively, 0.934 vs. 0.473). As seen in the figure, the binding energy of Se3d (Se^{2-}) was rather lower than Se^{0}, and, unfortunately, Li1s and Se3d binding energies were quite close to each other (detected between 52 eV and 55 eV, respectively). Even if the Li1s peak overlaps with that of Se3d, the cross section (sensitivity factor) of Se3d (0.821) is much higher than Li1s (0.028). Therefore, even if it is difficult to attribute the Li1S/Se3d peaks, due to possible polarization when S atoms are substituted by Se atoms (environment modification), the peak is mainly related to Se3d.

The core peak of Se^{0} was detected at 56 eV. A similar binding energy of Se^{0} was reported previously [34]. Overall, no signature of selenium precursors was detected in the pristine powders. To rule out the possible presence of Se in the pristine material, which would have stayed unnoticed due to the nano-size or to an amorphous nature, and would then follow an Ostwald ripening process or an agglomeration of particles in the electrolyte to form bigger particles [33], we performed high-resolution X-ray photoelectron spectroscopy (XPS) on the pristine powders.

Figure 10 shows the Se3d core peak of Li_{2}TiS_{3}, Li_{2}TiSeS_{2}, and Li_{2}TiSe_{2}S powders and selenium precursor. The four spectra were calibrated based on C 1s core peak at 284.8 eV.
XPS spectra, and, consequently, we conclude that there was no metallic selenium remaining in the powders after the synthesis. The origin of metallic selenium seen at the end of the charge could result from the selenium redox activity; therefore, further XPS analyses on selenium-sulfide electrodes at different states of charge are required to investigate this fact.

4. Conclusions
In summary, selenium-substituted samples (Li$_2$TiSeS$_2$ and Li$_2$TiSe$_2$S) were synthesized in disordered rocksalt phase, with a flexible and versatile synthesis process. Unlike Li$_2$TiS$_3$ cells, Li$_2$TiSeS$_2$ and Li$_2$TiSe$_2$S provided larger discharge capacities than the theoretical capacities. In cyclic voltammetry tests, Li$_2$TiSeS$_2$ and Li$_2$TiSe$_2$S showed different oxidative and reductive potentials from Li$_2$TiS$_3$, indicating a different redox activity. We also detected a second phenomenon that leads to extra discharge capacities. XPS and SEM and XRD ex situ studies showed that this extra capacity was coming from the activity of metallic Se that was formed during the first charge in these substituted samples and led to the formation of soluble polyselenides during the next discharge. The possible shuttle mechanism known for these species can be the origin of the low cycle life of these samples when cycled at low potential. Further structural studies are needed to elucidate the redox activities of Li$_2$TiSe$_x$S$_{3-x}$.

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