New superconductor Li$_x$Fe$_{1+\delta}$Se ($x \leq 0.07$, $T_c$ up to 44 K) by an electrochemical route

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The superconducting transition temperature ($T_c$) of tetragonal Fe$_{1+\delta}$Se was enhanced from 8.5 K to 44 K by chemical structure modification. While insertion of large alkaline cations like K or solvated lithium and iron cations in the interlayer space, the [Fe$_2$Se$_2$] interlayer separation increases significantly from 5.5 Å in native Fe$_{1+\delta}$Se to >7 Å in K$_{Fe_{1+\delta}}$Se and to >9 Å in Li$_{Fe_{1+\delta}}$(OH)$_{Fe_{1+\delta}}$Se, we report on an electrochemical route to modify the superconducting properties of Fe$_{1+\delta}$Se. In contrast to conventional chemical (solution) techniques, the electrochemical approach allows to insert non-solvated Li$^+$ into the Fe$_{1+\delta}$Se structure which preserves the native arrangement of [Fe$_2$Se$_2$] layers and their small separation. The amount of intercalated lithium is extremely small (about 0.07 Li$^+$ per f.u.), however, its incorporation results in the enhancement of $T_c$ up to ~44 K. The quantum-mechanical calculations show that Li occupies the octahedrally coordinated position, while the [Fe$_2$Se$_2$] layers remain basically unmodified. The obtained enhancement of the electronic density of states at the Fermi level clearly exceeds the effect expected on basis of rigid band behavior.

Tetragonal iron selenide, Fe$_{1+\delta}$Se (0.01 $\leq \delta \leq 0.04$), is one of the most interesting representatives of iron-based superconductors discovered a few years ago$^{1-3}$. Although the superconducting transition temperature is rather low ($T_c = 8.5$ K for $\delta = 0.01$), Fe$_{1+\delta}$Se offers unique opportunities for structure modification which significantly enhances $T_c$ up to 40 K$^{4,5}$. The increase of $T_c$ is accompanied by changes of distances and interactions between the anti-fluorite layers [Fe$_2$Se$_2$]-: adjacent [Fe$_2$Se$_2$] layers shift by (1/4, 1/4, 0) and the interlayer space is filled by heavy alkali metals$^{4,5}$ or solvated metal cations (chemical intercalation using anoxic polar solvent$^{6,7}$). Very interesting results were obtained in hydrothermal synthesis experiments which yielded the formation of Li$_{Fe_{1+\delta}}$(OH)$_{Fe_{1+\delta}}$Fe$_{1-\delta}$Se with $T_c \approx 40$ K (refs 8,9). The Li$_{Fe_{1+\delta}}$(OH)$_{Fe_{1+\delta}}$Fe$_{1-\delta}$Se structure is a stacking arrangement of alternating [Fe$_2$Se$_2$] and [(Li,Fe)(OH)] layers, the latter acting as charge reservoirs$^9$.

The literature information on electrochemical studies of iron chalcogenides is limited to just a few papers. So, the electrochemical Li-ion intercalation into FeSe$_{1-x}$Te$_x$ results in the formation of Li$_x$FeSe$_{1-x}$Te$_x$ for 0.25 $\leq x \leq 1$ (ref. 10). However, the observed preservation of lattice parameters and $T_c$ (even at high $y$ values) are not consistent with the claimed formation of the Li-intercalated phases. Moreover, the structural characterization is insufficient. Carbon-coated FeSe-nanoparticles were investigated as an anode material for lithium-ion batteries with a sustainable reversible capacity of 340 mAh·g$^{-1}$ (ref. 11). The reversible charge-discharge cycling implicates the amorphization of the initial iron selenide and probably involves the formation of metallic Fe-nanoparticles and Li$_2$Se. Successful electrochemical intercalation of potassium and sodium into FeSe has been reported very recently$^{12}$.

In this work we have successfully modified superconducting properties of Fe$_{1+\delta}$Se for the first time by electrochemical lithium intercalation. This approach allows the preservation of the native structure and the enhancement of $T_c$ up to 44 K due to changes of the carrier concentration. The obtained results provide the opportunity to shed new light on the mechanism of superconductivity in Fe-based superconductors and opens new ways for
enhancing $T_c$. The detailed investigation of the electrochemical behavior of Fe$_{1+\delta}$Se upon lithium intercalation conditions clarifies the existing contradictions and demonstrates the potential of the electrochemical approach to modify crystal structure and properties of Fe$_{1+\delta}$Se.

Results and Discussion
Electrochemical intercalation of Li-ions into tetragonal Fe$_{1+\delta}$Se was performed using a single-phase polycrystalline sample and large aggregates of single crystals. The polycrystalline sample consisted of agglomerates of particles with linear sizes between 1 and 10 $\mu$m (Fig. SI1). The lattice parameters $a = 3.7720(2)$ Å and $c = 5.5231(4)$ Å obtained from powder X-ray diffraction data (PXRD, Fig. SI2) are in good agreement with the literature$^{13,14}$. According to the SEM study, crystal aggregate of tetragonal Fe$_{1+\delta}$Se with a linear size of approximately $1 \times 2$ mm contains oriented single-crystal blocks (Fig. SI3).

Cyclic voltammetry (CV A) studies of polycrystalline Fe$_{1+\delta}$Se electrodes between 1.3 – 2.0 V vs. Li/Li$^+$ (scan rate of 0.05 mV·s$^{-1}$); (b) a part of ex-situ PXRD patterns for polycrystalline Fe$_{1+\delta}$Se electrodes at different potentials during 1st cycle. Reflections of tetragonal Fe$_{1+\delta}$Se are indexed. The diffraction maximum at 2$\theta$ of approx. 23° is unidentified. Electrochemical investigations at the same conditions (potential region, scan rates) revealed the appearance of this maximum also in the patterns of electrodes based on other active materials and even of the ‘idle’ electrode. Thus, the maximum is assigned to products of the electrolyte reduction occurring at the effective electrode surface.

$\text{FeSe}^2^+ + 2e^- \rightarrow \text{Li}_2\text{Se} + \text{Fe} \quad \text{theor} = 396 \text{ mAh} \cdot \text{g}^{-1}$

Figure 1. (a) CVAs curves (first three cycles) obtained for the polycrystalline Fe$_{1+\delta}$Se electrode in the potential range 1.3 – 2.0 V vs. Li/Li$^+$ (scan rate of 0.05 mV·s$^{-1}$); (b) a part of ex-situ PXRD patterns for polycrystalline Fe$_{1+\delta}$Se electrodes at different potentials during 1st cycle. Reflections of tetragonal Fe$_{1+\delta}$Se are indexed. The diffraction maximum at 2$\theta$ of approx. 23° is unidentified. Electrochemical investigations at the same conditions (potential region, scan rates) revealed the appearance of this maximum also in the patterns of electrodes based on other active materials and even of the ‘idle’ electrode. Thus, the maximum is assigned to products of the electrolyte reduction occurring at the effective electrode surface.
The reduction process was studied in more detail by means of slow scan-rate linear voltammetry (LVS) and potentiostatic intermittent-titration technique (PITT) combined with chronoamperometry (CA). This study reveals the complex character of the 1st reduction curve including additional peaks at \( \approx 2.1, 1.8 \) and 1.6 \( \text{V} \) vs Li/Li\(^+\) (Fig. 2, Fig. S14) preceding the main conversion peak (starting at \( \approx 1.5 \text{V} \), similarly to previous observations\(^{11}\). The results of analogous investigation of carbon-free electrodes allowed attributing the rather intense peak at \( \approx 2.1 \text{V} \) to the electrolyte reduction at the carbon surface (Fig. S14). The charge passed through the cell in the potential region of 3.0–1.55 \( \text{V} \) (before the major reduction peak) corresponds to \( \approx \) 11 mAh·g\(^{-1}\) (0.055 Li\(^+\) per f.u.). Further application of CA at 1.7 \( \text{V} \) for a few days did not result in significant growth of the current. This voltage is between the peaks of the assumed Li\(^+\) ion intercalation and the main conversion process in the differential capacity plot (Fig. 2). We suppose that the amount of the intercalated Li in Li\(_x\)Fe\(_{1+\delta}\)Se at this voltage corresponds to \( x \approx 0.06 \). According to PXRD, the Fe\(_{1+\delta}\)Se phase is preserved in the PITT procedure. Even after long CA treatment at 1.7 \( \text{V} \) for a few days did not result in significant growth of the current. This voltage is between the peaks of the assumed Li\(^+\) ion intercalation and the main conversion process in the differential capacity plot (Fig. 2). We suppose that the amount of the intercalated Li in Li\(_x\)Fe\(_{1+\delta}\)Se at this voltage corresponds to \( x \approx 0.06 \). According to PXRD, the Fe\(_{1+\delta}\)Se phase is preserved in the PITT procedure. Even after long CA treatment at 1.7 \( \text{V} \) (several days up to a week) all samples exhibit the same behavior: after turning-off the electrochemical cell, the potential increases rather rapidly from 1.7 \( \text{V} \) to approx. 2–2.5 \( \text{V} \) vs Li/Li\(^+\) as in case of initial Fe\(_{1+\delta}\)Se. While this effect as the result of the spontaneous decomposition of the intercalated phase Li\(_x\)Fe\(_{1+\delta}\)Se. Indeed, magnetic measurements of the electrode treated at 1.7 \( \text{V} \) vs Li/Li\(^+\) kept inside the glove-box at room temperature for several days showed the presence of the initial Fe\(_{1+\delta}\)Se phase only.

Magnetic measurements of the polycrystalline Fe\(_{1+\delta}\)Se electrode treated with LVS and CA at 1.7 \( \text{V} \) vs Li/Li\(^+\) unveiled drastic changes in comparison with native Fe\(_{1+\delta}\)Se (Fig. 3). The temperature dependence of the magnetic susceptibility \( \chi(T) \) shows a strong diamagnetic signal at low temperatures with a kink at about 8 K and a broad maximum at about 70 K. The pronounced, negative slope of the magnetic susceptibility at low temperature...
indicates the presence of initial superconducting tetragonal Fe_{1+δ}Se. The broad maximum at high temperatures signals the presence of several percent of elemental iron in the system, presumably in nano-structured form. This attribution is strongly supported by magnetization curves measured at 2 and 12 K, see inset of Fig. 3. Both curves evidence hysteresis loops which are typical for iron nanoparticles below the blocking temperature. The M(H) dependence measured at 2 K is decorated with a weak diamagnetic response seen in low fields due to the presence of the superconducting phase. Thus, the redox process between 1.8 V and 1.6 V vs Li/Li⁺ is characterized by a very small capacity value and accompanied by the conservation of the Fe_{1+δ}Se structure. The observed capacity value corresponds to a small amount of Li reacted with Fe_{1+δ}Se and being intercalated as Li⁺.

The spontaneous potential increase after cell turning-off together with the magnetic measurements data can be understood taking into account the instability of the product of the electrochemical redox process if the potential is not applied.

To clarify the behavior of Fe_{1+δ}Se in the applied electrochemical potentials, in-situ PXRD studies in galvanostatic mode (GCPL) were performed. In the potential region of 2−1.5 V vs. Li/Li⁺ (corresponding to approximately 0−10 mAh·g⁻¹ in the E−C curve), the original Fe_{1+δ}Se structure exhibits no measurable alterations (Fig. 4a,b). The intensity and position of the 101 reflection remains apparently stable (taking into account experimental errors), but the 001 reflection broadens and its center of gravity shifts to slightly lower angles (Fig. 4c,d). Concordantly, the shape of the E−C curve is typical for a solid solution intercalating process involving a single phase. Further decrease of the potential (1.5−0.7 V vs. Li/Li⁺, which corresponds to 10−400 mAh·g⁻¹ on the E−C curve) results in the rapid decomposition of Fe_{1+δ}Se monitored by the drastic decrease in the intensities of both the 001 and 101 reflections (Fig. 4a). No new diffraction maxima are observed. The E−C curve reveals an almost flat plateau in this potential region, which unambiguously indicates the two-phase state of the system in accordance with reaction (1).

The results above can be summarized in a scenario for the electrochemical processes of a polycrystalline Fe_{1+δ}Se electrode. In agreement with earlier results, the major process favored by thermodynamics is the decomposition and complete reduction of Fe_{1+δ}Se to metallic Fe. Electrochemical data together with in-situ PXRD measurements unveil that at higher potentials (ca. 1.7 V vs. Li/Li⁺) a subtle redox process takes place. This process conserves the initial structure topology and leads to the formation of a ternary phase LiₓFe_{1+δ}Se with slightly larger lattice parameter c compared with the one of Fe_{1+δ}Se. The potential increase which is observed after
through the cell before the current increases corresponds to the conversion reaction (Fig. S15). The differences mainly concern a shift of the redox peaks to higher potential values. The first reduction peak, which corresponds to the suggested intercalation process, appears at \( T_1 \approx 9 \text{ K} \). The second reduction peak, which is followed by the major peak (attributed to the conversion reaction) with an onset at \( T_2 \approx 30 \text{ K} \), becomes positive in the range \( T_3 \approx 44 \text{ K} \). It becomes positive in the range \( T_3 \approx 44 \text{ K} \). The temperature dependence of the magnetic susceptibility (Fig. 5) does not show a strong diamagnetic response even at low temperatures corresponding to 100% volume fraction of the superconducting phases. At higher temperatures the magnetization curve measured at 2 K (bottom inset to Fig. 5) is typical for Fe\(_{1+\delta}\)Se single crystals. Two other anomalies can be attributed to two new superconducting phases. Some confirmation of the superconducting nature of the phases in the region \( T_1 < T < T_3 \) can be found in magnetization curves measured at 12 K and 35 K (upper inset to Fig. 5) indicating a small diamagnetic response. From the results of the chronopotentiometric experiment (Fig. S16) we assume that in the products of the electrochemical treatment of the Fe\(_{1+\delta}\)Se crystal aggregates there are single-phase domains with slightly different content of Li.

The enhancement of \( T_c \) observed for the Fe\(_{1+\delta}\)Se crystal after electrochemical treatment confirms the suggested formation of Li\(_2\)Fe\(_{1+\delta}\)Se. As additional technique to determine the amount of Li\(_{1+\delta}\) and Fe\(_{1+\delta}\)Se and the formation of metallic Fe) in the latter case can kinetically inhibit the decomposition process. Polycrystalline electrodes consist of a large amount of relatively small crystallites (1–10 \( \mu \text{m} \) in size), crystal aggregates contain several single crystals of much larger size (100–1000 \( \mu \text{m} \)). Surface effects (including the decomposition of Li\(_2\)Fe\(_{1+\delta}\)Se) and the formation of metallic Fe) in the latter case should be much less pronounced. Furthermore, the application of crystals allows to reduce the current density of carbon black and polyvinylidenefluoride in the measurement cell that makes the corresponding electrochemical side reactions negligible. The electrochemical behavior of Fe\(_{1+\delta}\)Se crystal aggregates differs only slightly from that of the polycrystalline electrodes (Fig. S15). The differences mainly concern a shift of the redox peaks to higher potential values. The first reduction peak, which corresponds to the suggested intercalation process, appears at \( ca. 1.8 \text{ V} \) vs. Li/Li\(^{+}\). It is followed by the major peak (attributed to the conversion reaction) with an onset at \( ca. 1.7 \text{ V} \). This major peak is distorted in the CVA curve because the drastic volume change causes the destruction of the crystal. \( I-t \) curves obtained by long-time potentiostatic experiment at 1.7 V are close to those of the polycrystalline material. The current (in module) increases in the course of the 30 hours experiment. The amount of charge passed through the cell before the current increases corresponds to \( ca. 0.06 \text{ Li per f.u.} \) Fe\(_{1+\delta}\)Se.

Magnetic measurements of the Fe\(_{1+\delta}\)Se crystal aggregates after treatment at 1.7 V indicate the presence of inhomogeneous superconducting phases. The temperature dependence of the magnetic susceptibility (Fig. 5) demonstrates a strong diamagnetic response (Meissner effect) at low temperatures corresponding to 100% volume fraction of the superconducting phases. At higher temperatures the \( \chi(T) \) curve shows three kinks at \( T_1 \approx 9 \text{ K} \), \( T_2 \approx 30 \text{ K} \) and \( T_3 \approx 44 \text{ K} \). The first reduction peak, which corresponds to the suggested intercalation process, appears at \( ca. 1.8 \text{ V} \) vs. Li/Li\(^{+}\). It is followed by the major peak (attributed to the conversion reaction) with an onset at \( ca. 1.7 \text{ V} \). This major peak is distorted in the CVA curve because the drastic volume change causes the destruction of the crystal. \( I-t \) curves obtained by long-time potentiostatic experiment at 1.7 V are close to those of the polycrystalline material. The current (in module) increases in the course of the 30 hours experiment. The amount of charge passed through the cell before the current increases corresponds to \( ca. 0.06 \text{ Li per f.u.} \) Fe\(_{1+\delta}\)Se.

In-situ PXRD data together with the magnetic measurements indicate bulk (i.e. not surface) formation of the superconducting phase which is structurally similar to tetragonal Fe\(_{1+\delta}\)Se. In case of tetragonal Fe\(_{1+\delta}\)Se, such a \( T_c \) increase is mainly achieved by electron doping\(^{4,7}\). The electrochemical insertion of Li into the Fe\(_{1+\delta}\)Se structure can be considered as one possible way of electron doping. Due to the close-packed features of the Fe\(_{1+\delta}\)Se structure (defect anti-fluorite blocks), there are tetrahedral and distorted octahedral voids (Fig. 6). For tetrahedral coordination, the calculated Li–Se distances (\( ca. 2.28 \text{ Å} \)) are significantly shorter than those in the Li\(_2\)Se structure.

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**Figure 5.** Temperature dependence of the magnetic susceptibility of initial Fe\(_{1+\delta}\)Se crystal aggregates (open blue triangles) and that after treatment at 1.7 V vs. Li/Li\(^{+}\) (solid black circles) measured in the ZFC regime at 0.01 T. The insets show the field dependence of magnetization at temperatures below the superconducting transition measured at 12 K and 35 K (top) and 2 K (bottom).
(2.6 Å)\textsuperscript{18}. For octahedral voids, the estimated Li$-$Se distances (ca. 2.57–2.95 Å) are comparable to the ones in Li-Se, while the Li$-$Fe distances (ca. 2.39 Å) are too short. However, shift of the Li in [001] direction results in a square-pyramidal coordination of Li with longer Li$-$Fe contacts.

A similar effect is observed in LiFeAs (PbClF type)\textsuperscript{19}, where the anti-PbO-like topology is combined with a square-pyramidal Li\textsuperscript{+} coordination. Here, Li cations stabilize the whole structure because of Fe\textsuperscript{3+}/Fe\textsuperscript{2+} reduction.

It should be noted that the size of the anti-fluorite [Fe\textsubscript{2}As\textsubscript{2}] layer in LiFeAs is very close to that of the [Fe\textsubscript{1+δ}Se\textsubscript{2}] layer. Obviously, complete Li insertion into Fe\textsubscript{1+δ}Se would correspond to the complete reduction Fe\textsuperscript{3+}/Fe\textsuperscript{2+}, thereby limiting the amount of inserted Li.

The total energy for different model structures and Li concentration \(x\) was calculated to investigate the stability of Li\textsubscript{x}FeSe with Li atoms occupying different sites. From chemical experience, it is expected that Li occupies tetrahedral or octahedral voids between the Se atoms (Fig. 6). Without relaxation of Se coordinates, we find a binding energy of slightly more than 1 eV per Li atom occupying the octahedral void, essentially irrespective of the Li concentration (Table SI2). For the tetrahedral position, the binding energy is reduced considerably to less than 0.5 eV per Li. Due to this large energy difference and the large Li mobility, lithium occupation of the tetrahedral void can be safely discarded near the equilibrium. Relaxing all atoms without symmetry constraint (space group P\textsubscript{1}) in a Li\textsubscript{1/18}FeSe super cell (Fig. 6) and with a Li concentration close to the experimental value, the binding energy increases (Table SI2) by about 0.2 eV per Li. The predominant structural feature is an off-center Li position (Fig. 6), and a smoothing of the Fe-Se layers. The fourfold symmetry of the Li position and its environment however, is essentially preserved.

We also calculated the stability of Li\textsubscript{x}FeSe (\(x = 1/18 = 0.056\), near the experimental Li content) with respect to its decomposition into Li\textsubscript{2}Se, Fe and FeSe. Li\textsubscript{1/18}FeSe is not stable and thus should, under equilibrium conditions, decompose towards FeSe, Li\textsubscript{2}Se and Fe with an energy gain between 0.20 eV (for non-magnetic Fe clusters) and 0.42 eV (for magnetic Fe clusters) per Li (Table SI2). This is in agreement with the available experimental data. The calculated decomposition energy is only slightly dependent on the magnetic state of Fe in the clusters formed during the decomposition. For the experimentally observed superparamagnetic iron clusters the energy should be in-between the two limiting cases.

For a better understanding of the superconducting properties of Li\textsubscript{x}Fe\textsubscript{1+δ}Se, the change of the electronic density of states near the Fermi level \(E_F\) upon Li insertion is a crucial parameter. Comparison of the calculated DOS (Fig. 7 and Table SI2) for Li\textsubscript{1/18}FeSe and the parent compound FeSe yields three main results: (i) Surprisingly, the insertion of Li leads to a sizeable re-structuring of the DOS which goes beyond the changes expected on basis of a rigid band picture and results in an increase of the DOS at \(E_F\). (ii) The total band width is basically unchanged. (iii) The main increase of the DOS at \(E_F\) originates from more distant Se and Fe neighbors and not from the closer Li

Figure 6. The optimized 3 × 3 supercell of the tetragonal structure of Li\textsubscript{1/18}FeSe (left top; the 4-fold symmetry around Li (red) is essentially preserved, relaxation of Fe (purple) and Se (green) is taking place basically for nearest neighbors, only) with Li filling tetrahedral voids (right top) and octahedral ones (ideal–left bottom, optimized–right bottom). Relaxing all atoms with respect to the total energy shifts Li off-center and reduces the undulation of the Fe-Se layers (right bottom).
environment. This demonstrates that the additional electron donated by the Li ion is rather delocalized. However, a possible formation of polarons (beyond the DFT calculations) might modify this picture.

The observed changes in the electronic structure evidence the formation of a ternary compound which is chemically different from Fe\(_{1+\delta}\)Se. The appearance of a DOS spike at the Fermi level is in line with the observed higher critical temperature of the superconductivity.

In summary, our results are in line with earlier findings that the Fe\(_{1+\delta}\)Se structure pattern has a strong tendency to preserve the electron count of the [Fe\(_2\)Se\(_2\)] layers. Thus, the homogeneity range of these structure units with respect to heterovalent substitution or the insertion of electropositive metals is rather limited. Even in case of structure alterations which result in substantial increases of the \(T_c\) values, the charge of the anti-fluorite layer changes only marginally\(^7,8\). Our finding suggests that the significant \(T_c\)-enhancement of Fe\(_{1+\delta}\)Se can be achieved by a merely minute reduction of the Fe cations without increasing the [Fe\(_2\)Se\(_2\)]-interlayer separation. Therefore, it appears promising to investigate if the carrier concentration (or the iron valence) influences the \(T_c\)-value of layered iron-based selenides to a much larger extent than the interlayer separation.

Methods
Preparation. Polycrystalline sample of Fe\(_{1+\delta}\)Se was prepared under argon atmosphere using iron pieces (99.995\%) and selenium shots (99.999\%). Glassy carbon crucibles with a lid were filled with Fe/Se mixture with molar ratio close to 1 : 1 (Fe : Se = \(z\) : 1 and 0.98 \(\leq z \leq\) 1.02), placed into vacuumed quartz ampoule and subjected to high-temperature annealing (for details see ref. 13). Large crystals of Fe\(_{1+\delta}\)Se were prepared using the KCl/AlCl\(_3\) flux at 427 °C in evacuated quartz ampoules in constant temperature gradient\(^20\).

Basic characterization. Microstructure investigation of polycrystalline samples and crystals was performed with JEOL JSM 5510 (LaB\(_6\) cathode, 30 kV) scanning microscope. Phase identification and lattice parameters determination (for polycrystalline sample and crystals) were performed using room-temperature X-ray powder diffraction data obtained by image plate Guinier camera Huber G670, (CoK\(_\alpha_1\) radiation, \(\lambda = 1.78892\) Å). Crystalline Ge (\(a = 5.6576\) Å, ref. 21) was used as internal standard. Ex-situ X-ray diffraction data for polycrystalline electrodes were collected in air using Bruker D8-Advance diffractometer (CuK\(_\alpha_1\) radiation, \(\lambda = 1.540598\) Å, LynxEye PSD) in reflection mode. The electrodes previously were covered by air-protective one side sticky tape.

Electrochemical processing. Electrochemical treatments were performed in two-electrode setup with metallic Li as counter and reference electrodes. The commercial electrolyte (1 mol L\(^{-1}\) LiPF\(_6\) in a mixture of dimethyl carbonate and ethylene carbonate (1: 1 by volume, Merck) was used. The glass fiber was applied as separator. Working electrode (in case of polycrystalline samples) was prepared by screen printing the slurry containing the active material, carbon black and PVDF (mass ratio 85 : 5 : 10) in N-methylpyrrolidone onto Al-substrate. Load of active material was approx. 1 mg·cm\(^{-2}\). In a case of Fe\(_{1+\delta}\)Se crystals, one crystal was placed by its flat face between current collector and separator directly. Electrochemical cells were assembled and disassembled in Ar atmosphere. Potentiostat/galvanostat Biologic VMP-3 was used for data collecting. Several electrochemical techniques were applied: linear and cyclic voltammetry (LVs and CVA) between 1.0 V and 2.5 V vs. Li/Li\(^+\) with scan rate of 0.02 mV·s\(^{-1}\) and 0.05 mV·s\(^{-1}\); potentiostatic intermittent titration technique (PITT) with 0.01 V step.
in combination with chronoamperometry (CA) at selected potentials; and galvanostatic charge with potential limitation (GCL) at current density of approx. 20 mA·g⁻¹.

**In-situ powder X-ray diffraction.** The experiment was carried out on polycrystalline electrodes. The two-electrode electrochemical cell was analogous to the mentioned above but with Be window at the anode side and porous polypropylene separator (“UFIM”, Russia) instead of glass fiber. The diffraction data were collected using Bruker D8-Advance diffractometer (CuKα radiation, λ = 1.540598 Å, LynxEye PSD) in reflection mode. During the experiment, a constant current of ~15 μA (approx. 20 mA·g⁻¹) was applied at the working electrode; diffraction patterns were collected every 30 min until the potential decreased to 0.7 V vs. Li/Li⁺ and the electrochemical cell was switched off.

**Magnetic measurements.** Magnetisation was measured by vibration magnetometer (PPMS-9T, Quantum Design) applying fields of 10–1000 Oe. The specimens (polycrystalline samples, polycrystalline electrodes, crystals aggregates) were put inside plastic holders and then placed into the magnetometer in Ar atmosphere.

**Chemical analysis.** The lithium/iron ratio in electrochemically treated electrodes was analyzed by ICP-MS by means of ICP-MS spectrometer ELAN DRC II.

**Band structure calculations.** Relativistic density functional (DFT) electronic structure calculations were performed using the full-potential local-orbital FPLO code²²,²³ (version fplo 14.00–47). For the exchange-correlation potential, within the local density approximation (LDA) and the general gradient approximation (GGA), the parametrization of Perdew–Wang²⁴ and Perdew–Burke–Ernzerhof²⁵ were chosen, respectively. To obtain precise total energy and band structure information, the calculations were carried out on well converged meshes of up to 3888 k points depending on the cell volume. For all calculations, the respective experimental lattice parameters have been used.

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**Author Contributions**

A.M.A., E.V.A. and Y.G. designed the research, O.A.D., K.A.D. and A.M.A. performed electrochemical studies and characterizations, C.K. and U.S. prepared polycrystalline sample of Fe₁₋ₓSe, D.A.C. prepared crystal
aggregates of Fe$_{1+\delta}$Se, K.V.Z., O.S.V. and A.N.V. performed physical measurements, H.R. made electronic structure calculations, all authors contributed to the text of the manuscript.

**Additional Information**

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