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Supercritical-Fluid Synthesis of FeF₂ and CoF₂ Li-ion Conversion Materials

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The synthesis of the Li-ion conversion candidates, FeF_2 and CoF_2 , obtained from the single source organometallic precursors [Fe(tta)₃] (tta = C₈H₄F₃O₂S), and [Co(hfac)₂.2H₂O] (hfac = C₅H₁F₆O₂), respectively, *via* a novel supercritical fluid (SCF) method is presented. The nature of the synthesis led to

- ¹⁰ highly-crystalline FeF₂ and CoF₂ powders requiring no additional thermal treatment. The as-obtained powders were investigated for use as potential positive Li-ion conversion electrodes by means of chronopotentiometric measurements. The FeF₂ cells displayed high initial capacities following electrochemical conversion (up to ~ 1100 mAh g⁻¹ at a potential of 1.0 V *vs.* Li/Li⁺), with appreciable cyclic behaviour over 25 discharge-charge cycles. The deposition of a ~ 5 nm layer of amorphous carbon
- ¹⁵ onto the surface of the active material following SCF treatment, likely facilitated adequate electron transport through an otherwise poorly conducting FeF₂ phase. Similarly, CoF₂ cells displayed high initial capacities (up to ~ 650 mAh g⁻¹ at a potential of 1.2 V vs. Li/Li⁺), although significant capacity fading ensued in the subsequent cycles. *Ex-situ* XRD measurements confirmed a poor reversibility in the conversion sequence for CoF₂, with a complete loss of CoF₂ crystallinity and the sole presence of a

20 crystalline LiF phase following charging.

1. Introduction

Since the advent of Li-ion battery technology, considerable research effort has focused on the discovery and development of electrode materials as candidates for future Li-ion batteries. In

- ²⁵ particular, nanostructuring of conventional cathode materials, namely LiMO₂ and LiMPO₄ (M = Fe, Co, Mn, Ni *etc.*), has attracted much focus, owing to shorter Li⁺ diffusion lengths and hence, improved intercalation kinetics.^{1,2} Whilst attractive, firstrow transition metal oxides and phosphates suffer from relatively
- ³⁰ low Li⁺ ion storage capacities (e.g., ~ 170 mAh g⁻¹ in the case of LiFePO₄), and thus their specific energies are limited. The requisite properties of future Li-ion batteries such as increased capacity, enhanced cyclability, rate performance, amongst others, coupled with their ever increasing popularity as drivers for
- ³⁵ modern and demanding electronic devices and electric/hybridelectric vehicles (EV/HEVs) *etc.*, challenge researchers to seek ever more desirable materials with the properties to meet such rising demand.

In more recent times, researchers have considered the so-called 40 'conversion' electrodes as potential alternatives to the Li⁺ intercalation host. The reason for their supposition lies with the

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inherent ability of the conversion candidate to utilise the full range of oxidation states available in the corresponding metal 50 atom (Fe, Mn, Co etc.). The nature of such conversion chemistries typically results in redox transitions involving more than one electron and hence, the potential for greater specific capacities. Typical examples of conversion materials include transition metal oxides (MO_x), phosphides (MP_x), sulphides 55 (MS_x), and fluorides (MF_x), and have been the subject of recent excellent review articles.3,4 While conventional Li+ intercalation hosts operate on the physical insertion and removal of Li⁺ ions into and out of the local structure, conversion electrodes undergo their redox transitions in the absence of Li⁺ intercalation, during 60 which a complete reduction of the metal to form metallic nanoparticles of dimensions < 5 nm, embedded in a matrix of Li-X (X= O, S, O, F *etc.*), results as depicted in equation 1.4

 $\mathbf{M}_{a}\mathbf{X}_{b} + (b \cdot n)\mathbf{Li} \leftrightarrow a\mathbf{M} + b\mathbf{Li}_{n}\mathbf{X}$ (1)

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Initially, conversion materials were only considered useful as negative Li-ion electrodes owing to their relatively low operating potentials *vs*. Li/Li⁺.⁴ Metal fluorides, however, have been touted as one class of conversion materials with potential use as a 70 positive electrode since the highly ionic nature of the M-F bond results in more feasible cathodic operating potentials. However, to their detriment, such high ionicity also renders potential MF_x



candidates poorly conducting and, coupled with the formation of an insulating Li-X phase following their conversion, may lead to a reduction in the efficiency of electron transfer and hence, poor electrochemical performance.

- ⁵ Iron (II) fluoride, FeF₂, which crystallises in a rutile, tetragonal structure (space group: P_{42} /mnm), is one such MF_x conversion candidate since it offers a high theoretical capacity (~565 mAh g⁻¹), at a theoretical electrode potential of 2.66 V *vs*. Li/Li^{+.5} Such attributes could potentially lead to FeF₂ cathodes
- ¹⁰ displaying considerably higher energy densities compared to their intercalation analogues. The potentially high energy density may, however, be somewhat restrained as FeF₂ exhibits known Mottinsulator properties.⁶ Poor conductivity is an unfortunate and regular feature of many prospective Li-ion cathodes and is
- ¹⁵ typically challenged by the addition of electrically conducting agents, *e.g.*, C, Cu, Ag *etc.*, to the electrode mix, or by grafting layers of such materials to the active material surface either *insitu* or *via* post-synthetic techniques. Further to such treatments, it is accepted that a move to nanoscale dimensions may influence
- ²⁰ properties such as electrical conductivity versus their bulk counterparts. As such, a tailored synthesis of MF_x conversion materials having improved electrode properties may permit their use in Li-ion technology.
- Whilst several examples of iron (III) fluoride (FeF₃) cathodes ²⁵ have been reported in the literature, including nanowires,⁷ thinfilms,^{8,9} carbon nanocomposites,^{10–14} and the hydrated form, FeF₃·0.33H₂O,¹⁵ much less has been reported on the synthesis and electrochemical performance of the analogous FeF₂ phase alone. A recent report by Reddy et al. has shown C-FeF₂ to be a
- ³⁰ promising candidate.¹⁶ It is important to note that, although the complete reduction of FeF₃ affords higher theoretical discharge capacities (~ 712 mAh g⁻¹ vs. 565 mAh g⁻¹, respectively), at slightly higher electrode potentials (see table 1), previous work has demonstrated an increasingly ineffective reconversion
- ³⁵ process in FeF₃ following successive charge-discharge cycling.⁷ In addition, Yamakawa *et al.*¹⁷ identified that charging of FeF₃ cells following electrochemical conversion was associated with a transition from an original ReO₃-type structure to that of a defective rutile phase. FeF₃ does, however, possess the inherent
- ⁴⁰ ability to insert ~ 0.5 mol Li⁺ before the onset of electrochemical conversion, which has led to its utilisation as a more typical intercalation electrode with capacities reported approaching 200 mAh g⁻¹, at voltages which are beyond the scope of FeF₂.^{13,15,7,12} To this end, FeF₂ may be considered a somewhat more model
- ⁴⁵ conversion system, displaying reversible conversion character while beneficially involving fewer phase transitions compared to that of FeF₃. It should also be noted that the complete reduction products of FeF₂ and FeF₃ (Fe⁰ and LiF), are theoretically equivalent, indicating a predictably similar electrochemical ⁵⁰ pathway should occur during their electrochemical conversion
- (subsequent to an initial $Fe^{3+} \rightarrow Fe^{2+}$ transition in FeF₃); the mechanism of which, is still a subject of heavy scrutiny.^{18–21}

Cobalt (II) fluoride, CoF₂, which also crystallises in tetragonal form (space group *P*₄2/mnm), is another interesting candidate not ⁵⁵ least given cobalt's extensive use in early generation Li-ion cells (as LiCoO₂). In addition, CoF₂ extends a theoretical capacity of ~ 553 mAh g⁻¹ at a slightly electrode potential of 2.74 V *vs*. Li/Li⁺, leading to theoretical energy densities as high as 1500 Wh kg⁻¹. Previous studies,²² however, have outlined an incompatibility of ⁶⁰ phase-pure thin-film CoF₂ electrodes in conventional organicbased electrolytes, which has led to their relatively low levels of evaluation as potential Li-ion conversion candidates. A partial stability of CoF₂ in organic electrolyte was achieved by depositing a layer of lithium phosphorus oxynitride (Lipon) onto

⁶⁵ the surface of CoF₂ thin-films by a pulsed laser deposition (PLD) process, where such coatings led to the limited cyclic evaluation of CoF₂ as a potential negative Li-ion electrode.²² Phase-pure CoF₂ cathodes have also been evaluated by Wall *et al.*²³ which displayed high initial capacities (~ 600 mAh g⁻¹ at 1.0 V vs.

⁷⁰ Li/Li⁺), however, such cells also displayed significant capacity decline over 25 discharge-charge cycles. CoF₂ has therefore, been somewhat disregarded as a potential conversion electrode to-date, however, a review of its electrical properties with Li⁺ is still a facet that should not be ignored.

Herein this chapter details a novel supercritical fluid (SCF) synthesis of FeF_2 and CoF_2 powders using $[Fe(tta)_3]$ (tta = 4,4,4trifluoro-1-(2-thienyl)-1,3-butanedione), as a single-source precursor for the synthesis of FeF₂ powders and (hfac 1,1,1,5,5,5-hexafluoro-2,4-[Co(hfac)₂·2H₂O] = ⁸⁰ pentanedione), similarly, for the synthesis of CoF₂ powders. Previously, [Co(hfac)2.2H2O] precursors have been used for the growth of CoO or Co₃O₄ thin films by MOCVD,²⁴ while a similar Fe-based precursor, $[Fe(acac)_3]$ (acac = pentane-2,4-dione), has been employed for the growth of Fe₂O₃ thin-films in supercritical 85 CO₂²⁵ This is the first apparent synthesis of Fe- and Cocontaining fluorides using [Fe(tta)3] and [Co(hfac)2.2H2O] as single-source precursors by direct chemical routes. The asobtained FeF₂ and CoF₂ electrodes display high initial discharge capacities (up to ~ 1100 mAh g⁻¹ at 1.0 V vs. Li/Li⁺ in the case of 90 FeF2, and ~ 600 mAh g⁻¹ at 1.2 V vs. Li/Li⁺ for CoF2).

2. Experimental Section

2.1 Supercritical Fluid Synthesis of FeF₂ Powders

FeF₂ powders were synthesised using the β -diketonate, [Fe(tta)₃] (iron (III)- thenoyltrifluoroacetone), as a single-source 95 precursor according to procedures reported in the literature.^{26,27} 0.2 g of [Fe(tta)₃] was dissolved in 15 ml of dry toluene and sonicated for 3-5 min to ensure complete dissolution. The [Fe(tta)3] precursor solution was then injected into a sealed supercritical fluid (SCF) stainless steel reaction cell (24 ml total 100 volume, High Pressure Equipment (HiP) company), and heated to 400 °C at a ramp rate of 30 °C min⁻¹, and held for 1 h. An illustrative representation of the batch SCF setup is displayed in figure 2.1. After 1 h, the SCF cell was cooled to room temperature naturally (high pressure cells should never be 105 quenched with liquids), and slowly depressurised. The FeF2 product was isolated by decantation, washed with acetone to remove trace impurities and excess solvent and air-dried at room temperature for at least 24 h, forming the final brown-coloured powder product.

110 2.2 Supercritical Fluid Synthesis of CoF₂ Powders

CoF₂ was prepared using [Co(hfac)_{2.2}H₂O] (cobalt (II)hexafluoroacetylacetonate), as a single-source precursor which was used as received from Sigma-Aldrich and treated in a similar SCF batch method as described in 2.1.1. Typically, 0.2 g of $[Co(hfac)_2.2H_2O]$ precursor was dissolved in toluene (15 ml) and sonicated for 3-5 min, before being injected into a sealed stainless steel reaction cell (24 ml total volume, High Pressure Equipment (HIP) company), and heated to 400 °C at a heating rate of 30 °C

⁵ min⁻¹ and held at this temperature for 1 h. After 1 h, the reaction cell was cooled to room temperature and slowly depressurised. The CoF₂ product was isolated by means of decantation, washed with acetone to remove trace impurity and excess solvent and airdried at room temperature for at least 24 h forming the final pink ¹⁰ coloured powder.

2.3 Material Characterisation

X-ray diffraction (XRD) measurements were performed on a Phillip's XPERT diffractometer using Cu Kα radiation (1.5406 nm) operating at 40 kV and 35 mA. Scanning electron ¹⁵ microscopy (SEM) images were collected on a FEI Quanta 650 microscope operating at 10 kV and transmission electron microscopy (TEM) images were collected on a JEOL JEM 2100 TEM operating at 200 kV. EDX spectra were collected on an Oxford Instruments (INCA) EDX system fitted to the TEM.

²⁰ Thermogravimetric analysis (TGA) was carried out at a heating rate of 40 °C min-1 in air using a Mettler Toledo TGA/DSC1 STAR system.

For the electrochemical measurements, electrode slurries were prepared by mixing the active powder, acetylene black (Alfa

- ²⁵ Aesar, corp.), and PVDF (Sigma-Aldrich) in a ratio of 75:25:5 by mass, respectively. Absolute EtOH was used to ensure adequate consistency of the slurries. The slurries were then cast on to precleaned Al-foil and dried at 120 °C for several hours before mechanically pressing and further drying at 120 °C for at least
- ³⁰ 24 h. CR2025-type coin cells were assembled in a glovebox filled with high purity Ar gas, using pure Li chips (MTI corp.), serving as the negative electrode and Celgard® 2320 PE/PP/PE membranes as separator. The electrolyte was a solution of 1M LiPF₆ in EC:DMC (50:50 v/v). Galvanostatic cycling was carried
- ³⁵ out in the potential range between 1.0 5.0 V for FeF₂ and 1.2 4.8 V vs. Li/Li⁺ for CoF₂, using a Princeton Applied Research (V3) potentio/galvanostat. Typical active loadings were in the order of 0.5 1 mg. Respective current densities and capacities were determined on the mass of active material loading.

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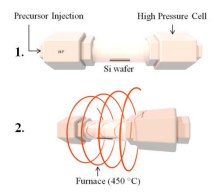
3. Results and Discussion

3.1 Synthesis and Structural Characterisation of FeF2

FeF₂ powders were prepared *via* a novel supercritical fluid synthesis using [Fe(tta)₃] as a single-source precursor, dissolved ⁴⁵ in toluene. The reaction setup is depicted in figure 1. Supercritical conditions were attained by the batch-method by sealing the as-prepared precursor sol in a high pressure reaction cell, followed by the thermal treatment at 400 °C for 1h.

- Previously, $[M(tta)_2]$ (M = Cd, Zn), single-source precursors have ⁵⁰ been used in the preparation of CdO, CdS or CdF₂ phases by a solution route,²⁴ and ZnO by a MOCVD route,²⁸ respectively, following coordination of the bis-complex precursor sphere with tmeda (N,N,N',N'-tetramethyletilendiamine), to form the singlesource precursor adduct [M(tta)₂-tmeda]. Similarly, the
- $_{55}$ precursors [Mg(hfac)_2] and [Mg(TFacac)_2], (TFacac = 1,1,1-trifluoro-2,4-pentanedione), have been used to prepare MgF_2 films by the thermally-driven disproportionation and

decomposition of the respective precursors at temperatures exceeding 500 °C.²⁹ The similarity of precursor composition in ⁶⁰ these instances outlines the role of synthetic conditions in directing the formation of either metal- oxide, sulphide or fluoride as the major product phase. Under supercritical CO₂ conditions, the non-fluorinated [Fe(acac)₃] (acac = pentane-2,4dione), single-source precursor has been used in the preparation ⁶⁵ of Fe₃O₄,²⁵ however, this is the first apparent synthesis and direct formation of FeF₂ by means of either conventional thermal treatment or *via* a supercritical route using [Fe(tta)₃] as a singlesource precursor.



70 Fig. 1 Batch SCF reaction scheme used in the preparation of FeF₂/CoF₂ powders.

To elucidate potential FeF2 formation under SCF conditions, a consideration of the thermal decomposition of [Fe(tta)₃] is necessary, where the formation of either oxide, sulphide or 75 fluoride as mixed or single phases may be expected depending on the nature of decomposition; the process of which, likely varies considerably as changes are employed to the reaction conditions (temperature, time etc.). Such phases are not easily distinguishable by means of TGA given that their expected ⁸⁰ residues are appreciably similar.²⁴ The variance in C-X (X = S, O, F), bond dissociation energies of [Fe(tta)₃] (C=O = 799 kJ mol^{-1} , C-S = 272 kJ mol^{-1} and C-F = 490 kJ mol^{-1}), could possibly indicate a preference for the formation of a dominant FeSx phase, however, XRD analysis (figure 2), of the brown powder product 85 indicated the formation of tetragonal-phase FeF2 as the singlephase reaction product (JCPDS card no.: 81-2272), with reflections at 20 values of 26.99, 33.31, 38.56, 43.17, 51.66, 55.40, 62.89, 66.10 and 69.68° corresponding to the (110), (101), (111), (210), (211), (220), (112), (301) and (202) set of planes in 90 tetragonal FeF2, respectively. No impurity peaks belonging to sulphide or oxide phases, or otherwise originating from the analogous FeF₃ or other Fe³⁺ impurities, were discernible in the XRD profile. Interestingly, Malandrino et al.²⁴ have shown that an analogous [Cd(tta)2] tmeda precursor adduct favoured the 95 formation of CdO via a conventional solution route when thermal treatment was carried out between 300 - 700 °C under flowing O2, while CdF2 was favoured under flowing N2 between 350 -450 °C and CdS favoured above a temperature of 550 °C.24 These results suggest that the formation of CdF₂ is favoured at 100 the lower temperature region (350 - 450 °C), while dissociation of the C-F bond under such conditions is likely preferential to that of C-S, which is located within the heavily stabilised 5membered ring of the respective thiophene group. Malandrino et $al.^{24}$ have also alluded to the possibility of CdS or S phase formation existing in non-crystalline form which could further react with CdF₂ to form CdS at the higher temperature regions. In the case of FeF₂, a temperature of 400 °C was used to attain

- ⁵ supercritical conditions (toluene) which likely facilitated a similar decomposition process of the [Fe(tta)₃] precursor. TGA analysis of the [Fe(tta)₃] precursor collected under flowing N₂ depicts a two-phase transition process resulting in the complete decomposition of the precursor over the temperature range
- ¹⁰ between 400 450 °C. A two-phase transition commences with a ~ 1 wt. % loss occurring at a temperature of ~ 165 °C (ascribed to the onset of precursor melting), followed by a decomposition step over the range 210 – 400 °C resulting in a further loss of ~ 93 wt. % and a residue of ~ 6 wt. %.
- ¹⁵ The formation of FeF₂ as a single phase component underlines SCF conditions were also favourable to the formation of Fe existing in the +2 oxidation state, given that no FeF₃ or other Fe³⁺ species were discernible in the XRD powder patterns; such an effect is particularly of note while the availability of F in the
- ²⁰ precursor was abundant and that the relative Gibbs energy of formation, ΔG_f , of FeF₃ is considerably lower than that of FeF₂ under standard conditions (-972 *vs.* -663 kJ mol⁻¹, respectively). It must also be suggested, however, that there lies a possibility of such Fe³⁺ phases existing in the amorphous state or critically,
- 25 existing in feature size below the X-ray coherence length used in the evaluation of the product composition.

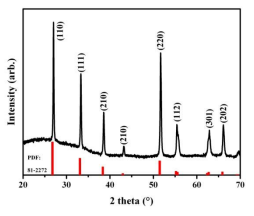


Fig 2. XRD profile of FeF₂ powders obtained from the SCF synthesis of Fe(tta)₃ at 400 °C for 1 h. The red bar makers indicate the respective positions (2θ) of FeF₂ (JCPDS: 81-2272).

The morphology of the FeF₂ powders was investigated by SEM and is displayed in figure 3. The SEM images reveal the formation of a variety of different structures consisting of smallscale nanorods which widely assembled into urchin-like features, to larger, more-elongated ribbon-like structures. The diameter of these structures varied from approximately 100 nm (short nanorod structures) to the wider, up to ~ 2 μ m sized elongated rods. TEM analysis of the FeF₂ structures (figure 3(d)), revealed a highly crystalline nature with an amorphous layer surrounding

⁴⁰ the FeF₂ domains, measuring approximately 5 nm in thickness. The clearly observable lattice fringes, which correspond to an inter-planar (d) spacing of 3.13 Å, may be indexed to the (110) set of planes in tetragonal FeF₂ (JCPDS: 81-2272), while the clearly defined lattice fringes visibly extend the length of the 45 FeF₂ structure and confirm the high crystallinity of the FeF₂ powder product. The ~5 nm amorphous layer may be attributed to the deposition of carbonaceous species following decomposition of the organometallic precursor or the organic reaction medium (toluene) used during preparation. The presence 50 of amorphous carbon is also indicated in the XRD profile (shown in figure 2), displaying a broad ($20 - 35^{\circ} 2\theta$), bending region, a feature often characteristic of non-graphitic carbon types. The presence of carbon is further confirmed by Raman analysis, displayed in figure 4, which shows the D and G bands 55 characteristic of sp³ and sp² carbon types.

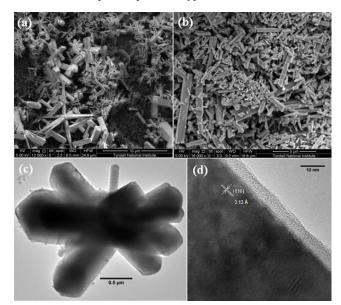


Fig 3. (a,b) SEM images of FeF₂ obtained from [Fe(tta)₃] at 400 °C in supercritical toluene (c,d) TEM images of FeF₂ obtained under the same conditions. The lattice spacing indicated in figure 5.2(d) may be indexed to the (110) set of planes in FeF₂ (JCPDS: 81-2272).

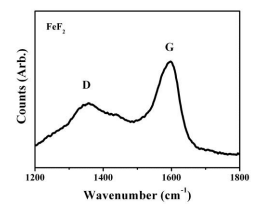


Fig 4. Raman spectra of a FeF_2 powder sample collected over the range 1200 – 1800 cm⁻¹, showing D (sp³) and G (sp²) bands characteristic of carbon types.

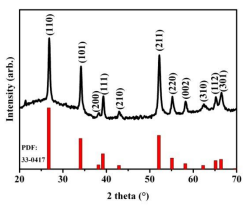
65 2.2 Synthesis and Characterisation of CoF₂ Powders

 CoF_2 powders were synthesised under a similar SCF method following the thermal decomposition of the [Co(hfac)₂·2H₂O] in supercritical toluene at 400 °C. The XRD profile of the asobtained, pale-pink coloured CoF₂ powder is shown in figure 5.

30

The crystal reflections at 20 values of 26.83, 34.14, 39.26, 42.90, 52.19, 55.17, 58.21, 62.22, 65.26 and 66.50° correspond to (110), (101), (111), (210), (211), (220), (002), (310), (112) and (301) set of planes in tetragonal CoF₂ (JCPDS: 33-0417), respectively, and

- $_{\rm 5}$ confirms the presence of CoF₂ existing as the major product phase. A similar bending region in the XRD profile (20 – 35° 2 θ) was also observed, and is ascribed to the existence of amorphous carbon. The thermal decomposition of [Co(hfac)₂·2H₂O] is predictably less complex than that of [Fe(tta)₃] given the likely
- ¹⁰ possibility of forming just two separate phases as the major product (Co_xO_y and CoF_x), as opposed to a third MS_x phase. In this instance, the formation of CoF_2 was pronounced under supercritical conditions at 400 °C with such conditions favouring the formation of Co in the +2 oxidation state while the
- ¹⁵ availability of F in the [Co(hfac)₂·2H₂O] precursor (12 F⁻ per formula unit), was markedly abundant. The possibility of forming Co_xO_y phases was also considered, however, the apparent lack of oxide formation as a result of supercritical processing (in what is a closed reaction system), may be explained by the lack
- ²⁰ of available oxygen existing in the reaction medium (toluene), coupled with the strong dissociation energy of the C=O bond of the organometallic precursor (C=O = 799 kJ mol⁻¹). Such an effect would also describe the lack of Fe_xO_{x+1} phases resulting from the supercritical processing of [Fe(tta)₃] under similar
- ²⁵ conditions. A direct comparison of the relative Gibbs' energy of formation for the respective phases (Co_xO_y, CoF₂), is difficult in this instance, given the large deviation from standard conditions under SCF processing.



 $_{30}$ Fig. 5 XRD profile of CoF $_2$ powders obtained from the SCF synthesis of Fe(hfac) $_2$ at 400 °C for 1 h. The red bar makers indicate the respective positions (20) of CoF $_2$ from PDF pattern no. 33-1074

SEM analysis of the CoF₂ powder samples indicated the formation of near-uniform spherical morphologies, ranging in ³⁵ diameter from 0.7 – 1.8 µm as shown in figures 6(a) and (b). Closer inspection of the CoF₂ spheres revealed a distinctive surface roughness, indicating that the spheres are comprised of a series of smaller, nanoscale features of an acicular-like crystal type. Higher resolution TEM images displayed in figures 6(c) ⁴⁰ and 6(d), confirmed that the larger CoF₂ spheres formed from a series of dendritic-like structures, whose feature sizes ranged between 20 – 30 nm in diameter. The nanoscale featuring of the CoF₂ powders in this instance may enhance the size of the ⁴⁵ counterpart; such properties may possibly favour Li⁺ transport by facilitating improved electrolyte access. It must also be considered, however, that such increases in the available surface area compared to that of the bulk phase may augment the formation of unwanted side interactions with the electrolyte, ⁵⁰ resulting in increases in respective decomposition products.

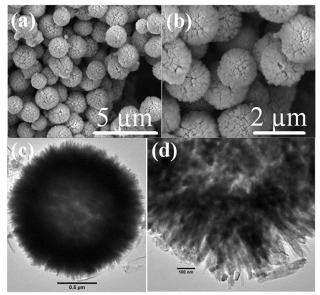


Fig. 6 (a,b) SEM images of CoF₂ spheres obtained at 400 °C from [Co(hfac)₂.2H₂O] using supercritical toluene and (c,d) TEM images of CoF₂ spheres obtained under the same conditions.

55 2.3 Li⁺ Induced Conversion of FeF₂ Powders

Table 1 summarises the theoretical electrode potentials and capacities of Fe- and Co- fluorides, outlining their practical use as potential positive electrodes in Li-ion technology; theoretical electrode potentials, E(V), are given in the literature.⁵ Our study ⁶⁰ has focused on the synthesis of FeF₂ and CoF₂ powders and their use as prospective Li-ion cathodes since the equilibrium potentials of FeF₂ and CoF₂ fall within feasible limits for such applications. Although the corresponding tri-fluoride phases afford higher theoretical energy densities given their increased ⁶⁵ working potentials and capacities (table 1), much less has been reported on the model MF₂ conversion systems alone.

Table 1. The theoretical capacity and electrode potentials, E(V), of Fe- and Co- fluorides.

Compound	Oxidation state	Theoretical capacity (mAh g ⁻¹)	E(V)
FeF ₂	+2	571	2.66
FeF ₃	+3	712	2.74
CoF ₂	+2	553	2.85
CoF ₃	+3	694	3.61

The equilibrium potential of FeF₂ is predicated from thermodynamic considerations at 2.66 V *vs.* Li/Li⁺ (table 1). Such potentials are considerably higher than those afforded by the ferric oxides, sulfides and phosphides, outlining the practical ⁷⁵ use of FeF₂ as a positive Li-ion electrode.^{30,4} The premise of the FeF₂ conversion mechanism assumes the reversible reaction as depicted in equation 2. Although the complete mechanism is

electrode-electrolyte interface versus the respective bulk

somewhat bypassed in this form, the formation of LiF and Fe⁰ accounts for the complete reduction of Fe²⁺ in the original species via a 2 e⁻ process, resulting in a theoretical capacity of ~ 571 mAh g⁻¹. Such capacities are considerably higher than those s afforded by conventional intercalation hosts such as LiFePO₄ (~ 170 mAh g⁻¹), and LiCoO₂ (~ 120 mAh g⁻¹) and lead to vast specific energies exceeding 1500 Wh kg⁻¹.

 $FeF_2 + 2Li^+ + 2e^- \rightarrow Fe^0 + 2LiF$ (2)

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The first discharge curves of the FeF₂ cells cycled at a lower cut-off potential of 1.0 *vs.* Li/Li⁺ at the current densities of 10 and 20 mA g⁻¹ (~ 0.017 and 0.035 C rate, respectively), are displayed in figure 7(a). Both discharge profiles show a sloping voltage ¹⁵ region commencing at ~ 2.6 V *vs.* Li/Li⁺ (open circuit voltage = 2.95 V), which accounts for some ~ 50 mAh g⁻¹ of the total capacity obtained in the first cycle. A portion of this sloping region may be attributed to the formation of SEI layers on the active material surface: Initially, such passivation layers were ²⁰ only considered for lithium or carbon anodes, however, research has confirmed the existence of the SEI layer on cathodic surfaces,

- including that of FeF_x.¹⁷ Such occurrences may arise as a result of spontaneous contact with electrolyte or predictably, during normal electrochemical operation, however, no structural ²⁵ confirmation of this layer is presented. Additionally, a number of
- other possible mechanisms have been predicted which could account for the initial sloping region, preceding the onset of electrochemical conversion. Although the existence of impurity Fe³⁺ phases, including that of FeF₃, were not detected in the XRD
- ³⁰ profile (figure 5), a reduction of trace amounts of amorphous Fe³⁺ phases must be considered; indeed, the reduction of Fe³⁺ phases is known to occur in the 3.0 2.0 V region and is thus a distinct possibility.¹⁷ Conversely, a number of conversion-competing processes have also been proposed,^{17,18} although there remains
- ³⁵ some degree of uncertainty as to their proposition. Yamakawa *et* $al.^{17}$ studied the local structures formed as a result of cycling FeF_x cathodes by Li⁶ and F¹⁷ solid-state NMR coupled with *ex-situ* XRD measurements, and concluded that the early stages of discharge in FeF₂ involved both intercalation and conversion
- ⁴⁰ processes. It is generally accepted that reduction of Fe^{2+} to Fe^{1+} is unlikely *via* simple 1×1 tunnelling, thus any possible intercalation is likely to proceed following reduction of Fe^{3+} impurity, or *via* the formation of intermediate species formed as a result of a disproportionation reaction of FeF_2 to form Fe^0 and ⁴⁵ Fe^{3+} species, outlined in equations 3 - 6.¹⁷

$$FeF_2 \rightarrow Fe_{1-x}F_2 + Fe \tag{3}$$

$$\operatorname{Fel}_{x\Gamma^{2}} + {}_{x}\operatorname{El} \xrightarrow{} \operatorname{El}_{x}\operatorname{Fel}_{x\Gamma^{2}}$$

$$\operatorname{Fel}_{x\Gamma^{2}} + {}_{x}\operatorname{El}_{x}\operatorname{Fel}_{x\Gamma^{2}}$$

$$\operatorname{Fel}_{x\Gamma^{2}} + {}_{x}\operatorname{El}_{x}\operatorname{Fel}_{x\Gamma^{2}} + {}_{x}\operatorname{El}_{x}\operatorname{Fe$$

$$50 \quad \text{Li}_{2x}\text{Fe}_{1-x}\text{F}_{2} \rightarrow 2_{y}\text{Li}\text{F} + \text{Li}_{2x-2y}\text{Fe}_{1-x}\text{F}_{2-2y} \tag{6}$$

The first step (equation 3) proposed by Yamakawa *et al.*,¹⁷ involves the lithium-driven displacement of Fe from the FeF₂ lattice, forming Fe⁰ and Fe³⁺, whilst also maintaining a rutile ⁵⁵ Fe_{1-x}F₂ structure. Such structures are thought to accommodate the insertion of Li⁺ ions in a more typical intercalative fashion. The possibility of further Li⁺ insertion to form Li_x[Li_xFe_{1-x}]F₂ is also considered. The reactions such as those proposed in

equations (3) to (6), are believed to fall within the 3.0 - 2.0 V 60 voltage window and could potentially compete with an onset of the conversion reaction outlined earlier in equation (2). In support of this fact, molecular dynamic simulations have also predicted early intercalation in FeF₂ cathodes by means of analysing Li⁺ exposure to the low energy (001) and (110) 65 surfaces of FeF2, where it was identified that a competitive process existing between Li⁺ intercalation along [001] channels and that of electrochemical conversion results.¹⁸ Such mechanisms are suggested to be widely governed by the rate of discharge and crystal orientation of the FeF2 local structure 70 during exposure to Li⁺.¹⁸ Such a competitive mechanism is, however, challenged somewhat by experimental observation that FeF₂ undergoes a rapid structural conversion mechanism in the absence of intercalation to the corresponding Fe⁰ and LiF phases with no Fe⁺ intermediaries.^{20,19} A recent communication by 75 Wang et al.²¹ has also depicted the conversion process of FeF2 for the first time by in-situ TEM observation. The authors found a surprisingly fast conversion process occurs (several minutes for FeF₂ nanoparticles), forming small diameter (1 - 3 nm) metallic Fe⁰ nanoparticles, surrounded by an amorphous LiF phase which ⁸⁰ ensued as a result of a propagation of a reaction front through the active FeF₂ domains. The absence of any identifiable Fe³⁺ phases by EELS measurements suggested that the conversion mechanism followed an intermediary pathway although, notably, the authors also suggest the possibility of Li⁺ intercalation 85 resulting in supersaturation of the FeF2 host resulting in a transformation from a crystalline rutile structure to that of an amorphous phase which could be responsible for the lack of detection of a lithiated (intercalated), phase by means of electron diffraction measurements.²¹ Following on from the early sloping 90 region in our discharge profile (figure 7), a long plateauing region develops which is attributed to the relevant phase transition(s) associated with the electrochemical conversion reaction of FeF₂. At the lower current density of 10 mA g^{-1} (~ 0.035 C rate, 1C = 571 mA g^{-1}), the onset of plateauing ensues from ~ 1.65 V before 95 showing very gradual decline over the remainder of the discharge process. After reaching a potential limit of 1.0 V, the capacity obtained following the first discharge cycle was ~ 1100 mAh g⁻¹, representing approximately twice that of the theoretical capacity assumed for a 2 e⁻ conversion process in FeF₂. On increasing the 100 current rate to 20 mA g⁻¹, the discharge profile displayed a gradual decline over the first ~ 400 mAh g⁻¹, followed by an inflection from ~ 1.5 V before a slight plateau region ensued nearer the low-voltage limit. After reaching a potential limit of 1.0 V, the capacity returned was some 590 mAh g⁻¹, representing 105 only a slight over-capacity with respect to theoretical maxima (571 mAh g⁻¹); in both cases, the high capacities returned outlines the highly promising potential of FeF2 during early stages of electrochemical cycling. The disparity in capacity and the nature of discharge between the comparative current densities does, 110 however, indicate a large degree of sensitivity of the conversion process to the applied discharge rate. Such a result is not surprising given the current understanding of the relatively sluggish kinetics of the conversion sequence, coupled by a predictably high incidence of overpotential which arises from 115 poor electronic conductivity and polarisation effects.

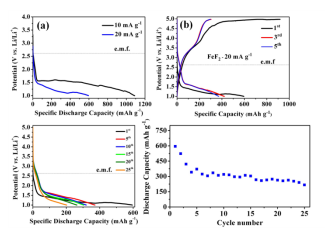


Fig. 7 (a) 1st discharge curves of FeF₂powders to 1.0 V vs. Li/Li⁺ at the current densities of 10 (black line) and 20 (blue line) mA g⁻¹, (b) 1st, 3rd and 5th discharge-charge curves obtained at a constant current of 20 mA g⁻¹, (c) 25-cycle discharge curves at the same current rate and (d) discharge capacity as a function of cycle number

The first, third and fifth discharge-charge (voltage) curves for the FeF₂ cells taken at a current density of 20 mA g⁻¹ over the same ¹⁰ potential range are displayed in figure 7(b), while the 25-cycle discharge curves are shown in figure 7(c), depicting the evolution of the discharge process. The average discharge potential displayed increases slightly with respect to the initial conversion reaction (black line, figure 7(b)), which is likely due to the

- ¹⁵ dramatically reduced active material size, resulting as a consequence of electrochemical conversion, thus leading to enhanced efficiency in the subsequent electrochemical conversion cycles. Notably, the first charge curve to 5.0 V (black line), displays a somewhat significant overcharge region, the nature of
- 20 which may be ascribed to increases in SEI formation at elevated potential. Such overcharge, however, remains relatively insignificant following the first cycle which would also corroborate its attribution to instances of SEI formation during the first charge cycle. While potentially detrimental, the high
- ²⁵ charge voltage was necessary in order to attain appreciable cycling for our FeF₂ powders (see later). Also apparent, is the distinct observation of disparity between the relevant shapes of the discharge and charge curves, a feature of hysteresis which is both a common and undesirable feature of the conversion ³⁰ electrodes.⁴

Previously, it was shown that charging of FeF₃ cells at potentials up to 4.0 V results in the recovery of a lithiated-Li_{0.5}FeF₃ phase while further charging of FeF₃ cells to 4.5 V results in an initial recovery of a defect tri-rutile FeF₃ phase.⁷ To

- ³⁵ this end, capacity fading has been expressed as a consequence of a poor reversibility in conversion sequence and has been observed by *ex-situ* TEM/electron diffraction measurements, which depicts the co-existence of α -Fe and LiF phases existing with FeF₃ after successive (50) charge cycles.⁷ Such incidences
- ⁴⁰ may suggest higher charging potentials may be required over time. In our case, charging the FeF₂ cells below 4.5 V, even at the relatively low current density of 20 mA g⁻¹, resulted in poor Coloumbic efficiency (Q_{charge}/Q_{discharge}), leading to poor capacity retention over relatively few discharge-charge cycles (not 45 shown).

Figure 7(d) displays the cycling performance of our FeF₂ powders at a current density of 20 mA g⁻¹. The specific capacity of the first discharge at a potential limit of 1.0 V was ~ 590 mAh g⁻¹, representing a conversion of approximately 2.1 Li⁺ per 50 formula unit FeF₂. In the second cycle, a reduction in capacity is observed after formation of the SEI layer, however, the discharge capacity at this stage is still largely agreeable with the theoretical approximations based on a total reduction of Fe^{2+} in FeF_2 (~ 571 mAh g⁻¹). The midpoint voltage displayed over the discharge 55 curve was ~ 1.38 V, therefore representing a second-cycle energy density of 715 Wh kg⁻¹; energy densities of this magnitude are considerably higher than those which may be afforded by conventional Li-ion intercalation cathodes such as LiFePO4 (~ 610 Wh kg⁻¹). A degree of capacity fading was evident over the 60 first four cycles, however, some stabilisation of capacity was attained notably over the first 15 cycles at ~ 300 mAh g⁻¹, while a discharge capacity of ~ 220 mAh g⁻¹ was returned following 25 charge-discharge cycles, representing a final conversion of approximately 1.1 Li⁺ per formula unit FeF₂. Such capacities are 65 largely similar to C-FeF2 nanocomposites discharged to 1.3 V after the same number of cycles,16 and still superior to theoretical capacities exhibited by some of the more conventional intercalation hosts, e.g., LiCoO2 (~ 120 mAh g-1) and LiFePO4 (~ 170 mAh g⁻¹); the lower average working voltage and low current 70 rate of FeF2 remains, however, potentially significant.

Although the discharge capacity of FeF2 remained appreciable over the cycling period, the large degree of overpotential leads to large round-trip energy inefficiencies which may currently preclude their use in commercial environments. In addition, the 75 high charging potentials required may lead to dramatic increases in electrolyte oxidation products, e.g., FeF₆, LiF₆, LiP_xO_yF_z, resulting in abject increases in SEI formation which could potentially lead to increased levels of capacity fading. In addition, the rate capability of FeFx-based electrodes, owing to 80 their intrinsically poor conductivity and the relatively limited kinetics of the conversion sequence, is also an issue facing the utilisation of FeF_x and other similar conversion candidates. The data reported in this work so far was collected at the relatively low specific currents of 10 and 20 mA g⁻¹, while cycling at an 85 elevated current density of 200 mA g⁻¹ (~ 0.35 C rate), resulted in a discharge capacity of just ~ 72 mAh g⁻¹ after 25 cycles (not shown). FeF₂, therefore, may find limited application in future high-rate Li-ion batteries due to its sluggish and complex conversion chemistries. Such examples depend on rapid Li+ 90 diffusion kinetics which are ill-afforded in this case, and may depend on suitable open host structuring. Although the conversion kinetics of FeF2 are demonstrated to be cumbersome at room temperature, both Reddy¹⁶ and Badway et al.¹⁰ have shown a vast improvement of an equivalent carbon-composite 95 FeF₂ cathodes operating at 40 and 70 °C, repectively. Interestingly, the operation of FeF2 cathodes at 70 °C was largely similar to those of C/FeF3 cells cycled at 25 °C, indicating the rate limiting steps during conversion of FeF_x lie at potentials < 2.0 V.

2.4 Li⁺ Induced conversion of CoF₂ Powders

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Theoretically, CoF_2 extends the possibility of delivering a high capacity of 553 mAh g⁻¹ at a slightly increased electrode potential

compared to the FeF₂ (2.85 V vs. Li/Li⁺; see table 1), resulting in potentially high energy densities exceeding 1500 Wh kg⁻¹. The acicular-like CoF₂ powders prepared by our SCF route (displayed in the SEM images, figure 4), are a potentially useful morphology

- ⁵ in electrode technology, and beyond, owing to an increase in the tapped density provided by their spherical structuring, while in addition, the nanoscale structuring which comprises the overall CoF₂ form, allows for an increased size of the electrode/electrolyte interface, and offering shorter Li⁺ diffusion
- ¹⁰ paths. While such characteristics have important implications for conventional intercalation hosts, such as favourable Li⁺ transport, much less is known about such influences on conversion-type electrodes which undergo significant morphological changes as a result of their conversion. To this end, the influence of starting
- ¹⁵ morphology on the first and subsequent discharge cycles remains unclear; however, it is reasonable to stipulate that an overall reduction in the size of the active material prior to conversion may facilitate an enhancement in the respective conversion kinetics. While these features are implicated before the initial
- ²⁰ discharge state, much less consideration is given to these effects after successive electrochemical cycling in which the active metallic species are already cycled between significantly smaller nano-sized features. Recently, Li *et al.*⁷ reported the ability of FeF₃ nanowire architectures to largely maintain their overall
- $_{25}$ structuring on cycling by forming a network of interconnected α -Fe domains, surrounded by a matrix of LiF. Equivalent data has not been presented for CoF₂ architectures, although a similar pathway is predicted.

The overall electrochemical conversion reaction of CoF₂ is ³⁰ outlined in equation (7), where the expected sequence affords LiF and Co⁰ as the reduction species in a 2 e⁻ process. Previous reports of CoF₂ architectures operating as conversion candidates have thus far been limited due to an instability of CoF₂ in conventional liquid organic electrolytes. Fu *et al.*²² prepared

- ³⁵ LiPON-coated CoF₂ thin-films by a PLD method, which delivered a capacity in the first discharge of ~ 600 mAh g⁻¹ at the anodic potential limit of 0.01 V vs. Li/Li⁺, while the introduction of their pure CoF₂ thin-films to the electrolyte solution resulted in severe instability. In this case, the introduction of LiPON
- ⁴⁰ coatings to the CoF₂ thin-film surface was thought to facilitate a reduction in the unfavourable interaction with liquid electrolyte, serving as a physical barrier layer in preventing its dissolution. The reduction species were confirmed by *ex-situ* XRD measurements, signified by LiF (220) and Co (111) reflections in
- ⁴⁵ the XRD profile.²² Recently, Wall *et.* al^{23} prepared both CoF₂/C and Co/LiF/C composite cathodes without such protective features; although the CoF₂/C composite displayed higher initial capacity than that of the Co/LiF/C composite (~600 and ~150 mAh g⁻¹ in the first discharge cycle, respectively), the Co/LiF/C
- ⁵⁰ composites displayed good capacity retention while the CoF₂/C composites attained just 15 % capacity retention within the cycling period (50 cycles). The application of Co/LiF in composite form has also been displayed with meritable results.³¹
- In the case of our SCF-prepared CoF₂ powders, little visible ⁵⁵ interaction with the electrolyte solution (LiPF₆ dissolved in EC:DMC 50:50 vol. %) was evident. The apparent stability of the CoF₂ powders in electrolyte was ascribed to the potential formation of carbon surfaces following the decomposition of both

organometallic precursor and residual toluene during synthetic ⁶⁰ processing. To this end, evaluation by galvanostatic cycling was possible (figure 8), although consideration must be given to the long term influence of stability of CoF₂ with progressive electrochemical cycling in liquid organic electrolytes.

Limited data is available thus far with regards to detailed 65 conversion mechanisms in CoF₂, although it has been proposed that CoF₂ extends an incomplete conversion mechanism with a further possibility of reaction between the newly formed nanograins of metallic Co⁰ with LiF (equation 7), the process of which is outlined in equations 8 & 9. ²²

- $CoF_2 + 2Li + 2e^- \leftrightarrow Co + 2LiF$ (7)
- xCo + LiF \leftrightarrow (1 x)LiF + xFCo + xLi⁺ (8)
- $(1-x)\text{LiF} + x\text{FCo} + x\text{Li}^+ + xe^- \leftrightarrow x\text{Co} + \text{LiF}$ (9)

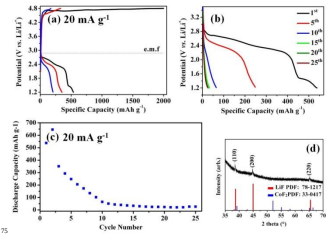


Fig. 8 CoF₂: (a) 1st, 3rd and 5th voltage curves at a current density of 20 mA g⁻¹ (b) 25 cycle discharge curves at the same rate (c) discharge capacity as a function of cycle number and (d) ex-situ XRD pattern of CoF₂ cells collected at the 25th charge-cycle

Figure 8(a) shows the initial discharge-charge curves of the CoF₂ powders prepared by SCF synthesis. At a discharge potential limit of 1.2 V in the first cycle, a capacity of ~ 535 mAh g^{-1} was obtained, representing ~ 97 % of the theoretical discharge capacity with respect to a full 2 e⁻ conversion reaction of CoF₂ (~ 85 553 mAh g⁻¹). Thus far, first discharge capacities of this magnitude for CoF₂ conversion systems have only recently been reported.²³ A similar capacity (~ 600 mAh g⁻¹), was also obtained at the anodic potential limit of 0.01 V,²² although the use of such voltages preclude their direct comparison. The first 90 discharge profile extends a dual plateauing system with such regions extending from ~ 2.7 and 1.5 V vs. Li/Li+. The first plateauing region (~ 2.7 - 2.3 V), accounts for some ~ 400 mAh g⁻¹ of the total discharge capacity obtained, while the lower plateauing region (~ 1.5 - 1.2 V), is significantly shorter, 95 accounting for ~ 130 mAh g⁻¹. Oxidation (fluorination) steps are discernable in the voltage curves, appearing as a sloping region from 4.3 - 4.6 V followed by a long, flat charge profile occurring from 4.6 - 4.8 V. The total charge capacity obtained at the upper cut-off potential was ~ 2000 mAh g-1, representing significant 100 overcharge compared to that of the first discharge capacity (535 mAh g⁻¹). Such an event indicates that complete charging of the CoF_2 cells overlaps the potential regions where the formation of electrolyte oxidation products is known to occur, through which may culminate in concomitant increases in unwanted electrolytic side reactions. Interestingly, the discharge capacity increased in

- ⁵ the second cycle to ~ 650 mAh g⁻¹, which is significantly higher than predicted; the possibility of an insertion/interaction of metallic Co with LiF to form a CoLiF intermediary has been proposed,²² although there is no evidence for this in our study. The third and subsequent discharge cycles offered declining
- ¹⁰ capacity performance, however, the dual-plateauing regions which extend throughout the discharge profile offer somewhat similarity to those of the first cycle and suggests a similar yet increasingly inefficient sequence of electrochemical reductions occurred. Fu *et al.*²² have shown that the first discharge reaction
- ¹⁵ (equation (7)) is largely irreversible, while the charge reaction of Co^0 and LiF affords CoF_x (here, x < 1), with only ~ 50% of the Co^0 and LiF re-converted to CoF_2 in the re-conversion process. The poor reversibility in the conversion sequence would account for the > 70% loss of capacity experienced following the first few
- ²⁰ cycles, which results in a return of ~ 100 mAh g⁻¹ after 10 cycles, while further cycling afforded yet further capacity decline, resulting in just ~ 30 mAh g⁻¹ following the cycling period (25 cycles). In support of a poorly reversible sequence, the final phases in the charged CoF₂ cells were identified by *ex-situ* XRD
- ²⁵ measurements (figure 8 (d)). The collected XRD patterns indicated a loss of CoF₂ crystallinity following cycling, with no discernible peaks originating from the starting CoF₂ phase. Broad peaks identifiable at 20 values of 38.5, 44.8 and 65.3° may, however, be indexed to the crystal reflections of the (110), (200)
- ³⁰ and (220) set of planes in LiF (PDF 78-1217). The apparent loss of CoF₂ crystallinity as detected by XRD measurements would indicate that all of the original CoF₂ phase participated in the earlier conversion (discharge) reaction, while the charging process was either incomplete, or that the resultant CoF_x phase
- ³⁵ that reformed was amorphous or sufficiently small so as to exceed the X-ray coherence length. Further, no reflections originating from Co phases could be identified to indicate the reversibility of the system. The indication of crystalline LiF existing in the charged state, however, supports the notion of a
- ⁴⁰ poorly reversible conversion mechanism as proposed by Fu *et* $al.^{22}$ An *in-situ* analysis of CoF₂ conversion with Li⁺ is therefore preferential in order to best understand the mechanistic and structural evolution in the conversion sequence. While the specific capacity and operating voltage of CoF₂ remains
- $_{45}$ agreeable with magnitudes that theoretical considerations would predict, improved electrochemical performance of CoF₂, particularly pertaining to extended cycle life, may result from a complete understanding of the conversion sequence. The optimisation of CoF₂ electrodes, such as the grafting of
- ⁵⁰ electrically conducting layers to the CoF₂ surface, could result in enhanced conversion kinetics and a more favourable reconversion process, while the development and utilisation of alternative electrolyte compositions more suited to the operating potentials could allow further study of the respective cycling
- ⁵⁵ processes. A thorough understanding of the conversion mechanisms and the respective kinetics involved may allow for complete utilisation of CoF₂ with stable performance as a future Li-ion electrode.

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60 3. Conclusions

In conclusion, a method of synthesising FeF₂ powders using a novel SCF route using [Fe(tta)₃] as a single-source precursor in toluene at 400 °C was presented. CoF₂ powders were also prepared under a similar approach, using [Co(hfac)₂.2H₂O] ⁶⁵ serving as single-source precursor. This represents the first display of utilising such precursors in the formation of highly crystalline MF₂ powders under supercritical conditions. TEM analysis revealed an amorphous layer (~ 5 nm), surrounding the FeF₂ domains, which was ascribed to the deposition of ⁷⁰ carbonaceous species following the synthetic treatment under SCF conditions. Due to the intrinsically insulating nature of FeF₂, the conducting carbon layers surrounding the FeF₂ domains

high initial energy capacities (up to 1100 mAh g⁻¹ and 595 mAh ⁷⁵ g⁻¹ at 10 and 20 mA mA g⁻¹, respectively), and are relatively stable over 25 discharge-charge cycles. A second discharge energy density of 810 Wh kg⁻¹, which is significantly higher than those afforded by the conventional intercalation electrodes such as LiFePO₄ and LiCoO₂, outlines the high-potential of FeF₂

likely enabled adequate electron transfer. The FeF₂ cells display

- ⁸⁰ cathodes. The kinetics of the FeF₂ conversion sequence were appreciably poor at room temperature, however, resulting in diminished electrochemical performance at increased discharge rates. In addition, the relatively low average working potential of FeF₂ (a deviation of > 1.0 V per the calculated e.m.f), coupled ⁸⁵ with a large overpotential remains pertinent. Further work is required in order to assess the potential of FeF₂ cathodes with
- improved rate performance, which may be achieved by further reducing the active particle size or through the introduction of dopant species.
- ⁹⁰ CoF₂ spheres, which were comprised of nanorods in the order of 20 nm diameter, also displayed appreciable capacity during early stages of electrochemical cycling, however, an unfavorable re-conversion mechanism and a possible ill-reaction with the organic electrolyte rendered the SCF prepared phase-pure CoF₂ a
- ⁹⁵ poor Li-ion electrode after relatively few electrochemical cycles. However, through the advancement of electrolyte compositions which are stable at higher voltages, or *via* composite formation, CoF₂ cannot be disregarded as a potential positive electrode at this stage.

100 4. References

- 1. P. Poizot, S. Laruelle, S. Grugeon, L. Dupont, and J. M. Tarascon, *Nature*, 2000, **407**, 496–9.
- 2. J. M. Tarascon and M. Armand, *Nature*, 2001, **414**, 359–67.
- 3. R. Malini, U. Uma, T. Sheela, M. Ganesan, and N. G. ¹⁰⁵ Renganathan, *Ionics*, 2008, **15**, 301–307.
 - 4. J. Cabana, L. Monconduit, D. Larcher, and M. R. Palacín, *Adv. Mat.*, 2010, **22**, E170–92.
 - 5. G. G. Amatucci and N. Pereira, *J. Fluorine Chem.*, 2007, **128**, 243–262.

6.	G. Valerio, M. Catti, and R. Dovesi, <i>Physical Review B</i> , 1995, 52 .
7.	L. Li, F. Meng, and S. Jin, Nano Lett., 2012, 12, 6030-7.
8. 5	Y. Makimura, a. Rougier, L. Laffont, M. Womes, JC. Jumas, JB. Leriche, and JM. Tarascon, <i>Electrochem. Commun.</i> , 2006, 8 , 1769–1774.

- Y. Makimura, A. Rougier, and J.-M. Tarascon, *Appl. Surf. Sci.*, 2006, **252**, 4587–4592.
- 10. F. Badway, F. Cosandey, N. Pereira, and G. G. Amatucci, *J. Electrochem. Soc.*, 2003, **150**, A1318.
- 11. I. Plitz, F. Badway, J. Al-Sharab, a. DuPasquier, F. Cosandey, and G. G. Amatucci, *J. Electrochem. Soc.*, 2005, **152**, A307.
- N. Yabuuchi, M. Sugano, Y. Yamakawa, I. Nakai, K. Sakamoto, H. Muramatsu, and S. Komaba, *J. Mater. Chem.*, 2011, **21**, 10035.
- 13. S.-W. Kim, D.-H. Seo, H. Gwon, J. Kim, and K. Kang, *Adv. Mat.*, 2010, **22**, 5260–4.
- S.-T. Myung, S. Sakurada, H. Yashiro, and Y.-K. Sun, J. Power Sources, 2013, 223, 1–8.
- 20 15. C. Li, L. Gu, S. Tsukimoto, P. a van Aken, and J. Maier, Adv. Mat., 2010, 22, 3650–4.
- M. A. Reddy, B. Breitung, V. S. K. Chakravadhanula, C. Wall, M. Engel, C. Kübel, A. K. Powell, H. Hahn, and M. Fichtner, *Adv. Energy Mater.*, 2013, 3, 308–313.
- 25 17. N. Yamakawa, M. Jiang, B. Key, and C. P. Grey, J. Am. Chem. Soc., 2009, 131, 10525–36.
- 18. Y. Ma and S. H. Garofalini, J. Am. Chem. Soc., 2012, **134**, 8205–11.
- 19. S. Rangan and R. Thorpe, J. Phys. Chem. C, 2012.
- F. Wang, R. Robert, N. a Chernova, N. Pereira, F. Omenya, F. Badway, X. Hua, M. Ruotolo, R. Zhang, L. Wu, V. Volkov, D. Su, B. Key, M. S. Whittingham, C. P. Grey, G. G. Amatucci, Y. Zhu, and J. Graetz, *J. Am. Chem. Soc.*, 2011, **133**, 18828–36.
- F. Wang, H.-C. Yu, M.-H. Chen, L. Wu, N. Pereira, K.
 Thornton, A. Van der Ven, Y. Zhu, G. G. Amatucci, and J. Graetz, *Nature Comm.*, 2012, 3, 1201.
- Z.-W. Fu, C.-L. Li, W.-Y. Liu, J. Ma, Y. Wang, and Q.-Z. Qin, J. Electrochem. Soc., 2005, 152, E50.
- 23. C. Wall, R. Prakash, C. Kübel, H. Hahn, and M. Fichtner, *J. Alloys Compd.*, 2012, **530**, 121–126.
- G. Malandrino, S. T. Finocchiaro, P. Rossi, P. Dapporto, and I. L. Fragalà, *Chem. Comm.*, 2005, 8, 5681–3.
- 25. S. De Dea and D. Graziani, *The Journal of Supercritical Fluids*, 2007.

45 26. J. T. Berg, Eugene W.; Truemper, Journal of Physical Chemistry, 1960, 2948.

50

60

- H. Soling, A. Jonsson, S. Westman, K. J. Watson, and M. Sandström, *Acta Chemica Scandinavica*, 1976, **30a**, 163–170.
- G. Malandrino, M. Blandino, L. M. S. Perdicaro, I. L. Fragalà, P. Rossi, and P. Dapporto, *Inor. Chem.*, 2005, 44, 9684–9.
- 29. J. Haisma, P. H. Joosten, H. J. P. Nabben, T. J. A. Popma, and H. A. M. Van Hal, .
- H. Li, P. Balaya, and J. Maier, J. Electrochem. Soc., 2004, 151, A1878.
- 55 31. Y. Zhou, W. Liu, M. Xue, L. Yu, C. Wu, X. Wu, and Z. Fu, *Electrochem. Solid-State Lett.*, 2006, 9, A147.