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Introduction

Batteries are not only part of our present lives in the form of portable energy sources for our numerous electronic devices, but also part of our future as the tools that will allow us to move further with the integration of clean renewable energy sources in the electricity grid. This will be achieved by using batteries, storing the excess energy generated from renewable sources and releasing the energy when necessary. However, to achieve widespread integration of batteries into the electricity grid,

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Structural evolution of mixed valent (V³⁺/V⁴⁺) and V⁴⁺ sodium vanadium fluorophosphates as cathodes in sodium-ion batteries: comparisons, overcharging and mid-term cycling⁺

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Sodium vanadium fluorophosphates belonging to the $Na_3V_2O_{2x}(PO_4)_2F_{3-2x}$ family of compounds have recently shown very good electrochemical performance versus Na/Na⁺ providing high working voltages (3.6 and 4.1 V) and good specific capacity values. In this work the electrochemical behaviour and structural evolution of two compositions, $Na_3V_2O_{16}(PO_4)_2F_{1.4}$ (V^{3.8+}) and $Na_3V_2O_2(PO_4)_2F$ (V⁴⁺), are detailed using time-resolved in situ synchrotron X-ray powder diffraction. For the first time in sodiumion batteries the effects of overcharging and mid-term cycling are analyzed using this technique. Differences in the composition of both materials lead to different combinations of biphasic and singlephase reaction mechanisms while charging up to 4.3 V and overcharging up to 4.8 V. Moreover, the analysis of particle size broadening of both samples reveals the higher stress suffered by the V^{4+} compared to the more disordered $V^{3.8+}$ sample. The more "flexible" structure of the $V^{3.8+}$ sample allows for maximum sodium extraction when overcharging up to 4.8 V while in the case of the V^{4+} sample no evidence is shown of more sodium extraction between 4.3 V and 4.8 V. Furthermore, the analysis of both materials after 10 cycles shows the appearance of secondary phases due to the degradation of the material or the battery itself (e.g. electrolyte degradation). This study shows examples of the possible degradation mechanisms (and phases) while overcharging and mid-term cycling which is in turn crucial to making better electrodes, either based on these materials or generally in cathodes for sodium-ion batteries.

essentially better, safer, longer lasting and cheaper batteries are required. In this context, research on battery materials has become broader, in the search for new ions that can reversibly insert/extract from a range of hosts, for example the use of sodium ions, instead of lithium ions.¹⁻⁴

The development of Na-ion based batteries demands research into new materials, electrodes and electrolytes, to make these devices a commercial reality. Many compounds have been examined as possible cathodes for more reliable, long-lasting and economic Na-ion batteries.5,6 Among them, sodium vanadium fluorophosphates show high voltage, high capacity and long cycle-life. The performance arises from the two reaction voltages, 3.6 and 4.1 V vs. Na/Na⁺, the high theoretical specific capacity of 130 mA h g⁻¹ and their 95% capacity retention over 200 cycles.⁷ In the last two years, it has been demonstrated that these sodium vanadium fluorophosphates belong to the $Na_3V_2O_{2x}(PO_4)_2F_{3-2x}$ family, where the substitution of F by O in the same crystallographic site controls the vanadium oxidation state from a V³⁺ compound when x = 0 (or Na₃V₂(PO₄)₂F₃), to a V⁴⁺ phase when x = 1 (or Na₃V₂O₂(PO₄)₂F). Mixed valent phases are readily evident for intermediate values of x (ref. 8–10) and Fig. 1 shows the structural model and the F/O substitution sites.

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Fig. 1 (a) Vanadium octahedra of Na₃V₂O_{2x}(PO₄)₂F_{3-2x} (0 < x < 1) compounds. Vanadium is represented in blue, oxygen in red and fluorine in green. The crystallographic structure of (b) Na₃V₂(PO₄)₂F₃ and (c) Na₃V₂O₂(PO₄)₂F in the tetragonal *P*4₂/mnm space-group. Phosphate tetrahedra are represented in blue.

In spite of belonging to the same family, there are slight differences between the V4+ compound and the thoroughlystudied mixed valent phase of the composition x = 0.8 with V^{3.8+}. First, they both are produced by hydrothermal syntheses but from different precursors. The choice of precursor provides the V^{3.8+} compound with a carbonaceous coating that the V⁴⁺ material lacks. Second, the electrochemical performance of the $V^{3.8+}$ material has been demonstrated to be better than the V^{4+} material, especially when cycled at high rates. This can, in part, be related to the carbon coating of the V^{3.8+} materials, or to the higher intrinsic conductivity of the mixed valent material (with V^{3+} and V^{4+}) compared to the V^{4+} materials (with only V^{4+}). It was recently demonstrated that these materials undergo different reaction mechanisms when extracting/inserting Na⁺ ions from/ into the structure.^{11,12} Recent calculations on the Na₃V₂O_{2x}- $(PO_4)_2F_{3-2x}$ family have suggested that the third Na can be extracted at 4.9 V for x = 0 and 5.3 V for x = 1;¹³ however for the x = 1 sample the oxidation state of all V atoms is V⁵⁺ when 2 Na atoms are removed so the removal of the third Na would need to involve another oxidation state or mechanism. Additionally, favourable high rate performance has also been shown with this

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family of compounds when combined with graphene and its derivatives.^{13,14}

A thorough understanding about these materials in terms of electrochemical performance and structure is required before they can be commercially used. In particular, the mechanisms that appear under cell stress or aging conditions, such as overcharging and mid-term cycling are of practical importance to manufacturers. To date, some *in situ* X-ray diffraction studies of materials in Na-ion batteries have been published, both from laboratory¹⁵⁻¹⁷ and synchrotron X-ray sources,¹⁸⁻²⁰ but none of them have focused on the degradation mechanisms of the electrode materials by overcharge or mid-term cycling, until now.

In this study, we detail two possible degradation mechanisms of these cathodic materials, overcharging and mid-term cycling. For this purpose, we have used time-resolved in situ synchrotron X-ray diffraction (XRD) data obtained from coincells designed to be analyzed during operation and without removal. In situ XRD provides insight into the reaction mechanism and crystal structure evolution of electrodes in a battery during charge/discharge and this technique has already proved its relevance.11,12,19-22 Such data are distinctly different from providing a snapshot of the equilibrated structure under equilibrium conditions as ex situ (post mortem) or pseudo-equilibrium in situ data provide. With sufficient time-resolution the in situ data show detailed evolution under non-equilibrium and real battery operation conditions. Using this technique, we determined the Na extraction mechanisms of both mixed valent and V^{4+} phases, when they are overcharged up to 4.8 V vs. Na/ Na⁺. Then, the structures and sodium content of both compounds at 4.8 V are compared, and the effect of a potentiostatic step at high voltages on the mixed valent sample is analyzed. For mid-term cycling analysis, both compounds have been examined using synchrotron XRD data after a sequence of offline cycles. The reaction mechanism, appearance of new secondary phases and the finally discharged cathode structure are illustrated.

Experimental

 $Na_3V_2O_2(PO_4)_2F$, the V⁴⁺ phase,⁹ was prepared under mild hydrothermal conditions (170 °C) and autogenous pressure, by reacting V₂O₃ (Sigma-Aldrich, 98% purity), $NaH_2PO_4 \cdot H_2O$ (Fluka, 99% purity) and NaF (Sigma-Aldrich, 99% purity) in a 1 : 2 : 1 molar proportion. The reaction mixture was sealed in a polytetrafluoroethylene (PTFE)-lined steel pressure vessel, which was maintained at 170 °C for 65 hours. The reaction product was washed repeatedly with distilled water and acetone.

Na₃V₂O_{2x}(PO₄)₂F_{3-2x} where x = 0.8, the V^{3.8+} phase,⁸ was synthesized by the hydrothermal method from a ceramic precursor. The synthesis process took place in two steps. First, the VPO₄/C composite was synthesized by the ceramic method. V₂O₅ (Sigma-Aldrich, 99.99% purity) and NH₄H₂PO₄ (Fluka, 99.5% purity) were mixed in an agate mortar in a stoichiometric ratio with a 25% molar excess of Kejten black. This mixture was annealed twice under a nitrogen atmosphere at 300 and 850 °C. Second, the sodium fluorophosphate was prepared under mild hydrothermal conditions (170 °C) and autogenous pressure by reacting NaF (Sigma-Aldrich, 99% purity) and the VPO₄/C composite in a 3.3/1 molar proportion. The reaction mixture was sealed in a polytetrafluoroethylene (PTFE)-lined steel pressure vessel, which was maintained at 170 °C for 65 hours.

The positive electrodes were manufactured by mixing 80% wt $Na_3V_2O_{2x}(PO_4)_2F_{3-2x}$ active material, 10% wt conductive carbon (Super C65, Timcal) and 10% wt polyvinylidenefluoride binder (PVDF 5130, Solvay). A few mL of N-methylpyrrolidone (NMP, Aldrich) were added and the resulting slurry was stirred for 1 h. This slurry was then coated on an aluminium foil using the "doctor blade" technique. The electrode film was dried at 80 °C in a vacuum oven for 24 h. The electrode sheets were pressed to 100 kN using a flat plate press (MTI corporation) and dried overnight at 100 °C before transfer to the Ar-filled glovebox. Coin cells with 3 mm diameter holes in the casing and 5 mm diameter holes in the stainless spacer were used for the construction of the coin cells for the *in situ* measurements. The coin cells contained Na metal (\sim 1 mm thickness), glass fibre separators with 1 M NaPF₆ in dimethyl carbonate and diethyl carbonate (1:1 wt%) electrolyte solution.

In situ synchrotron XRD experiments were performed within two days after the construction process for the overcharge investigation. Overcharge studies were performed by charging both starting phases up to 4.8 V at 0.3 mA. The mixed valent sample was held for 75 minutes at 4.8 V, whereas the V⁴⁺ phase discharged immediately. Although the experiment was designed to hold for both samples, unfortunately an error in the cell cycler during the experiment meant that for the V⁴⁺ sample the hold step finished rapidly. Both materials were subsequently discharged to 2.5 V vs. Na/Na⁺.

The cells for the mid-term cycling were started after construction and synchrotron XRD data were collected within a day of completion of the cycles. The V^{4+} material was cycled offline for 10 cycles between 2.5 and 4.3 V at 0.3 mA for the first cycle and 0.1 mA for the following cycles. The open circuit voltage (OCV) was 3.57 V, the cell was cycled *in situ* at 0.3 mA but for comparative purposes to the mixed valent electrode only the initial state after the offline cycling is considered. The mixed valent sample was only cycled 5 times, and then it failed.

In situ synchrotron XRD data were collected on the Powder Diffraction beamline²³ at the Australian Synchrotron with a wavelength (λ) of 0.68816(2) Å, determined using the NIST 660b LaB₆ standard reference material. Data were collected continuously in 4 minute acquisitions on the coin cell in transmission geometry throughout the charge/discharge cycles described above. Rietveld refinements were carried out using the GSAS²⁴ software suite with the EXPGUI²⁵ software interface. A similar Rietveld procedure to our previous work was used.^{11,12} Particle size distributions were determined using the lorentzian profile function as implemented in GSAS and the Scherrer equation.

Results and discussion

First, results related to the overcharge experiments will be described followed by the results of cathode degradation or mid-term cycling for both materials, $V^{3.8+}$ and V^{4+} . It should be

noted that the $Na_3V_2O_{2x}(PO_4)_2F_{3-2x}$ family of compounds have been studied recently where the V oxidation state has been allowed to vary. However, in order to achieve this variation the synthetic methods are typically very different, e.g. a one-step hydrothermal method for V⁴⁺⁹, a two-step hydrothermal method using a carbon containing precursor for the V^{3.8+8} or a ceramic method using a carbon containing precursor for the V3+20. These approaches lead to inherent differences in electrochemical performance, e.g. with or without carbon coating. In addition, for in situ experiments different rates have been used and reported.^{8,9,20} The phase evolution will in part be related to the current rate applied during charge and discharge, the current modifies the galvanostatic charge/discharge rates and these are manifested in the structural evolution. Bearing this in mind, future work can be targeted to understand the phase evolution as a function of the applied current. This is particularly pertinent due to the high rate performance recently observed with these materials.13,14,26

Overcharging of mixed valent and V⁴⁺ phases up to 4.8 V

The upper cutoff voltage is often determined by either the electrochemical curve increasing rapidly without the addition of significant amount of further capacity or by the limits of electrolyte stability. The cutoff voltage also corresponds to the maximum amount of Na that can be removed from the cathode. In our attempt to address what is the maximum amount of Na extraction from the Na₃V₂O_{2x}(PO₄)₂F_{3-2x} electrodes we charged the cells *in situ* to 4.8 V. Fig. 2a and b show the 2D diffraction plots of the V⁴⁺ and the mixed valent samples focused on 220 and 113 reflections (13.1° to 13.7° 2 θ range) corresponding to charge up to 4.8 V.

Similar to our previous report,¹² charging of the V⁴⁺ sample proceeds via a short initial solid solution mechanism followed by two sets of two-phase transitions (P to P' to P''12) around 3.6 V linked to the lower potential plateau-like feature in the charge curve (Fig. 2a), then a single phase region for the majority of the higher potential plateau until close to 4.3 V where an abrupt two-phase transition occurs (see Fig. 2a). After this transition and to higher voltages a solid solution mechanism continues until the overcharge cutoff voltage. In the case of the mixed valent sample, again this is similar to the literature as expected, beginning with a solid solution region, followed by a two-phase region and then a solid solution region. The solid solution region continues until the end of overcharge, at 4.8 V. The detailed structural evolution of both these materials when overcharging is parallel to that reported earlier without an overcharge step.11,12

Both materials present a different structural evolution with charge, probably induced by the subtle changes in the vanadium oxidation state (and the F : O ratio) in Na₃V₂O_{2x}(PO₄)₂- F_{3-2x} (x = 0.8) and Na₃(VO)₂(PO₄)₂F phases (and possibly due to the synthetic approach, *e.g.* with or without carbon coating). The compositional changes slightly modify the structure of the compound. It must be recalled that the mixed valent phase has a crystallographic site occupied by either F or O, so the V^{3.8+} compound is more disordered than the V⁴⁺ one. Thus,



Fig. 2 2D diffraction plots of the charge process of (a) V^{4+} and (b) mixed valent fluorophosphate electrodes focused on the 220 and 113 reflections (from 13.1° to 13.7° 2 θ range).

structural evolution is strongly affected by this phenomenon and each phase does not evolve in the same way when extracting Na from the structure.

Overcharging: what happens at high voltage charge?

As the structural evolution during charge has been established, the behaviour of the V4+ compound under overcharge conditions, in the 3.8-4.8 V range (104 to 120 minutes) is now discussed. The structural details of the overcharged V⁴⁺ phase are shown in the ESI, Table S1.† Fig. 2 shows the lattice and Na site occupation evolution in this high voltage region. Very small fluctuations are detected in the lattice, which is consistent with the small variations in sodium content from 4.3 V onwards. An average total sodium content of 1.21(10) with a clear preference for the Na(1) site was found in the high voltage region, which indicates that the vanadium oxidation state is \sim 4.90. This value, sodium content and preferential occupation are very similar to the ones obtained when applying a potentiostatic step at 4.3 V to this phase,¹² so it can be concluded that no extra specific capacity is obtained from this phase when going to voltages above 4.3 V. The fact that the phase reaches 4.8 V very fast, with a sharp rise in the voltage curve is also in good agreement with the minimal addition of extra capacity during overcharge.

The mixed valent material undergoes subtle structural evolution during overcharge that can be analyzed in terms of diffraction intensity, position and full width at half maximum (FWHM) as shown in Fig. 3. Between 4.3 and 4.8 V the intensity of the 002/111 reflection significantly increases, whereas 220 and 113 intensities slightly decrease and the 222/311 reflection intensity shows a minor increase. In the constant potential region all the reflection intensities tend to stabilize, with the exception of the 220 reflection, which appears to increase at the later stages of the 4.8 V constant potential.

In terms of reflection positions, 222/311 and 220 reach their highest position at 4.3 V and stabilize during the constant voltage region. On the other hand, the 113 reflection position decreases continuously up to 4.8 V and 002/111 reflections reach their lowest positions at 4.3 V and slightly increase until 4.8 V and stabilize during the constant potential region. When the FWHMs of these reflections are compared, the 220 and 222/311 reflections remain quite stable, while the 002/111 FWHM



Fig. 3 (a) Lattice parameter and cell volume evolution and, (b) sodium site occupation in the high voltage range (3.8-4.8 V) for the V⁴⁺ electrode. Purple lines represent the voltage curves and, in (b), nominal 4.3 V voltage.

decreases between 4.3 and 4.8 V, and then continues to slightly decrease during the potentiostatic step. It must be also remarked that, at the end of the potentiostatic step when a wait step is applied before discharge, from 203 to 207 minutes, the structure changes abruptly. This happens because the structure, which is at a high potential, starts to relax quickly when the potential/current is not applied.

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Summarizing, two regions must be distinguished for the mixed valent sample at high voltages; the one that corresponds to the potential rise up to 4.8 V, and the one corresponding to the constant potential step. In the first section, structural changes coming from the charge process are more evident, and these changes are minimized during the potentiostatic step.

Finally, all the changes that are observed in the diffraction intensity turn into modifications of the lattice and sodium occupation in the material (Fig. 4). The lattice (volume) shrinks



Fig. 4 (a) Integrated intensity, (b) 2θ position and (c) FWHM of 002, 111, 220, 222, 311 and 112 reflections of the mixed valent electrode between 100 and 200 minutes, which corresponds to the overcharge and potentiostatic step at 4.8 V.

during the voltage rise up to 4.8 V and then stabilizes with slight fluctuations of *a* and *c* lattice parameters. In order to refine sodium occupation, a variety of models were attempted. Finally, it was necessary to fix or sequentially refine most atomic parameters (positions/ADPs) and set Na(1) occupation as 0. The constraints were necessary due to the presence of some correlations when using more refinable parameters. However, it must be noted that the Na(2) occupation results obtained when constraining Na(1) occupation to 0 and when allowed to freely refine were very similar. The structural details of the overcharged mixed valent phase are shown in the ESI, Table S2.†

Structural refinements of the mixed valent model with the diffraction data at the end of charge at 4.8 V (before the potentiostatic step) indicate an average total sodium content of ~0.51. There is a clear preference for the Na(2) site, leaving Na(1) site empty (with or without including this site into the model). This implies that the vanadium oxidation state at the end of the charge process and before the potentiostatic step is V^{5+} , which is the maximum possible oxidation state of this element. Thus the mixed valent material would have reached the maximum possible specific capacity for this phase, since the structure is at the electrochemical V^{5+} extreme.



Fig. 5 (a) Lattice parameter and unit cell volume evolution of the mixed valent sample, and (b) Na(2) site occupation which is the total sodium content evolution, as Na(1) = 0, during the high voltage range and potentiostatic step (3.8-4.8 V). Purple lines represent the voltage curves.

Table 1 Key crystallographic data of sodium in both overcharged structures

Atom	x	у	z	Site occupancy factor
V ⁴⁺ overcl	harged material	l		
Na(1)	0.5091	0.252	0	0.60(6)
Na(2)	0.794	0.035	0	0.01(8)
Mixed val	ent overcharge	d material		
Na(1)	0.5495	0.2155	0	0
Na(2)	0.7403	-0.0353	0	0.254(12)

The comparison between the two 4.8 V overcharged electrode materials, V^{4+} and mixed valent, is shown in Fig. 5 and key crystallographic data for sodium in both materials are displayed in Table 1.

The first feature that must be noted is that only Na(2) sites are represented in the mixed valent overcharged material, since Na(1) site occupation is zero; whereas both Na(1) and Na(2) are partially occupied for the V⁴⁺ sample. There is an extremely low but not insignificant Na(2) content in the V^{4+} compound. Notably, the sodium distribution amongst the crystallographic sites is surprisingly inverse in both oxidized materials, with overcharged V⁴⁺ showing preferential Na(1) site occupation and the oxidized mixed valent electrode showing preferential Na(2) site occupation. This exchange can be due to the fact that both sodium sites are located inside the tunnels of the structure and these can be filled/extracted alternatively. There is likely to exist a close relationship between these sites, such that cooperative insertion/extraction of sodium in the channels is possible or speculatively a concerted filling/removing into/from the Na sites. It must be also remarked that, in spite of having a lower overall sodium content, the cell volume of the mixed valent electrode at the oxidized state is greater than the V⁴⁺ volume. A smaller volume may influence the probability of preferential occupation on a particular sodium site.

The difference in volume is also found in the raw materials,^{8,9} possibly due to the greater disorder that exists in the mixed valent sample, with the partial substitution of O by F in the vanadium coordination octahedra and the lower overall vanadium oxidation state. Thus the different sodium distributions between the V4+ and mixed valent electrodes at the overcharged state can be related to the different structural transitions leading to the overcharged state, the ease of transport between the two sodium sites and the influence the cell volume has on the site-specific occupancy.

Electrochemically the performance of V⁴⁺ compared to the mixed valent electrode can be related in part to the structural differences observed in these overcharged structures. Table 2 shows the total sodium content and V oxidation state of V4+ and mixed valent electrodes at 4.3 V (normal charge) and 4.8 V (overcharge). It appears that the V^{4+} electrode approaches its Na extraction limit at 4.3 V and it is not possible to extract more, even going up to 4.8 V, whereas the mixed valent electrode can go beyond the Na extraction at conventional cut-off voltages of 4.3 V. This can partially explain the inferior performance of the V^{4+} relative to the mixed valent electrode. While V^{4+} is pushed up to the conventional Na extraction limit in normal charge, the mixed valent 4.3 V charged structure suffers less structural stress because it doesn't reach its limit and can be cycled to a higher potential. This fact can be also considered as a consequence of the initial structural differences in both materials. In addition, calculations suggest that further desodiation might be possible.13

Overcharging - particle size broadening

The changes observed in the FWHM during overcharge, Fig. 3, shows a potential anisotropic change of crystallite size, e.g. preference in the 002 direction, for the mixed valent sample at the overcharged state. This indicates that the crystallite and particle size may play a role in conventional and overcharge regimes.

Crystallinity evolution of these fluorophosphates can be evaluated by the fitting of selected reflections or in whole pattern refinements in the discharged materials before and after the overcharge. In the case of the V⁴⁺ sample two sets of reflections are depicted in Fig. 6. 111 at 7.7° and 404 at ca. 24.6° were selected for comparison and a SEM image of the pristine material is also shown in Fig. 6. Two phases coexist in the V^{4+}



Structures of the overcharged (a) V^{4+} phase, $Na_{1,21(10)}V_2O_2$ -Fig. 6 $(PO_4)_2F$ and (b) mixed valent phase, $Na_{0.508(12)}V_2O_{1.6}(PO_4)_2F_{1.4}$.

Table 2 Total sodium content and the average V oxidation state of V⁴⁺ and mixed valent electrodes at 4.3 V (normal charge) and overcharge conditions (4.8 V)

	4.3 V normal charge	Ref.	4.8 V overcharge	Ref.
V ⁴⁺	Na content = $1.24(13)$, V ^{4.88+}	12	Na content = $1.21(10)$, V ^{4.90+}	This work
Mixed valent	Na content = $0.77(4)$, V ^{4.92+}	11	Na content = $0.508(12)$, V ⁵⁺	This work

Table 3 Particle size broadening of the V⁴⁺ material before cycling and in the discharged state after the 4.3 V potentiostatic step, with conventional charge up to 4.3 V and overcharge conditions up to 4.8 V vs. Na/Na⁺

	Initial	Conventional CH/DIS (4.3 V)	% change	4.3 V pot. step	% change	Overcharge 4.8 V	% change
V ⁴⁺ [ref. 12] V ⁴⁺ [this work]	210 nm ^a 1440 nm	200 nm	+5%	100 nm	-52%	50 nm	-96%
^{<i>a</i>} Indicates that th	e electrode was	s pre-cycled.					

discharged electrode after 4.8 V overcharge, which is unlike what was observed in our previous investigation of this material when cycled up to 4.3 V, in normal charge conditions where a single phase electrode is found on discharge.¹² Thus, whereas the V⁴⁺ material comes back to its original (single phase) state when charging and discharging under normal conditions, the structure does not reset when overcharge conditions are used. This can be considered as an adverse influence of the high voltage used. Moreover, diffraction reflection broadening can be observed when the overcharged electrode is discharged, indicating the crystallinity loss. The Scherrer equation was used to semi-quantitatively study the changes in the crystallite size in the material based on whole pattern refinement. The obtained data have been compared to the crystallite size obtained from our previous *in situ* cycling of this material.¹² Table 3 displays the obtained particle size broadening of the V⁴⁺ material before cycling and in the discharged state after the 4.3 V potentiostatic step, with conventional charge up to 4.3 V and overcharge conditions up to 4.8 V vs. Na/Na⁺.

Note that the particle size analysis is relevant in terms of relative changes, not necessarily in absolute particle size values. This is because calculated sizes are affected by the peak shape function used in all cases. A significant size difference can be observed for the initial V⁴⁺ phase in the material analyzed in ref. 12 and the one from this work. This could be due to the fact that the V^{4+} electrode in ref. 12 had been precycled offline (1 cycle), so this could account for the differences. When comparing the initial particle size to the size of the discharged material after conventional charge/discharge up to 4.3 V, no significant changes are observed, whereas, when subjecting the material to a potentiostatic step at 4.3 V there is a large 52% reduction in particle size distributions. The use of an overcharge potential of 4.8 V leads to an even more massive reduction in particle size. Thus the overcharge conditions induce significant structural changes and stresses on the V⁴⁺ phase, as it causes the existence of two phases when discharged, with a concerted reduction in particle size. This can lead to a premature failure of the material, shortening its cycle life.

In the case of the mixed valent sample, the material is monophasic after overcharge and a potentiostatic step at 4.8 V, as it can be seen in Fig. 7. Furthermore, no significant diffraction peak broadening is observed, which indicates that crystallinity is preserved after overcharge, see Table 4. SEM images of the pristine mixed valent electrode are also shown in Fig. 8. As it can be seen in Table 4, changes in the particle size distributions for the mixed valent electrode are of a lower magnitude under the three conditions analysed compared to the V⁴⁺ electrode. The three conditions are normal charge up to 4.3 V, after a potentiostatic step at 4.3 V, and after overcharge at 4.8 V. This can be due to the slight structural disorder characteristic of the mixed valent phase that allows the material to adapt better to structural changes. Overcharge at 4.8 V gives the largest change in size (65%) relative to the conventional and potentiostatic charge (~25%).

It should be noted that one sample is carbon coated $(V^{3.8^+})$ while the other is not (V^{4^+}) and this may also contribute to the observed changes in the particle size. The coating layer may act



Fig. 7 Selected diffraction peaks before and after overcharge of the V⁴⁺ material. 7.7° corresponds to 111, 7.9° to 002 and 24.6° to 404 reflections. The inset shows an SEM image of the sample.

Table 4Particle size broadening for the mixed valent electrode before cycling and in the discharged state after conventional charge up to 4.3 V,4.3 V potentiostatic step (pre-cycled cell), and overcharge conditions up to 4.8 V vs. Na/Na⁺

	Initial	Conventional CH/DIS (4.3 V)	(% change)	After pot. step at 4.3 V	(% change)	After overcharge (4.8 V)	(% change)
Mixed valent "fresh" ¹¹ Mixed valent cycled ¹¹	250 nm	180 nm 320 nm	-28%	240 nm	-25%		
Mixed valent [this work]	500 nm					175 nm	-65%



Fig. 8 111 and 002 reflections before (top) and after overcharge (bottom) of the mixed valent material. Example of a single peak fit is shown but the whole pattern fits are used to derive the particle size distributions in Tables 3 and 4. The inset shows a SEM image of the pristine electrode material prior to electrochemical cycling.

to enhance or restrict the dynamics of ion insertion/extraction. However, there is no method at the moment to synthesize both V^{4+} and $V^{3.8+}$ phases under the same conditions.

Mid-term cycling

The cathodes in selected batteries were not extracted, they were cycled in *in situ* cells and data were collected after 10 cycles while the batteries remained in the cell. *In situ* or *ex situ* analysis of a battery after ten cycles offline can shed light on the processes that take place in the cathode, that can be related to the degradation of the active material. This knowledge is of great value to evaluate the stability of the cathode in the battery system and, in case it is possible, to propose ways to avoid this degradation.

V⁴⁺ material: fresh cell *vs.* start of the 11th cycle

The V⁴⁺ electrode structure has been analyzed after 10 charge/ discharge cycles. Fig. 9 shows the XRD patterns of the fresh cell and at the start of charge corresponding to the 11th cycle. It is worth noting that the amorphous broad feature that is observed in the starting material lowers in intensity and stabilises at the beginning and over the course of the 11th cycle (which was conducted *in situ*). Usually this broad background feature at low angles is attributed to the liquid structure factor of the electrolyte,²⁷ thus it is consistent that, after 10 cycles, some electrolyte has been consumed or even lost during cell operation.

In terms of diffraction reflections, the main reflections associated with the V^{4+} phase are maintained, although some new peaks are observed. These new reflections can be assigned to degradation products generated during material cycling. Two kinds of new reflections are marked in Fig. 9b, broad and sharp peaks which could be attributed to a crystalline secondary or by-product phase and to a secondary or by-product with a marked amorphous or nanocrystalline character. In addition, the V⁴⁺ electrode component appears to be of two-phases at this state of charge. Thus, we can observe that the electrode contains broad and sharp reflections associated with possibly at least two phases and the V⁴⁺ phases.

V⁴⁺ material: start and end of the 11th cycle

When comparing the beginning and end points of the 11th cycle it can be seen that they are very similar (Fig. 10). The two-phase mixture at the beginning of the 11th charge mentioned above is not obvious in the discharged state, similar to the monophasic phase when fresh. This suggests that the electrode relaxed after the offline cycling at the beginning of the 11th cycle which is consistent with the lattice parameters being different from the initial fresh electrode found in ref. 12.

The analysis of the 7–8.6° range in the XRD patterns provides important clues about the extra peaks found at the start and end of the 11th cycle. It can be seen that there are small differences in the diffraction reflections related to the V⁴⁺ phase (at about 7.7 and 7.9° characteristic of two-phase reactions), whereas the 7.4° reflection does not change (even in the *in situ* data, not shown). This indicates that the phase this peak belongs to is not electroactive and it can be formed by degradation over time or slow crystallization during electrochemical reactions.

Mixed valent material: fresh cell vs. cycled

The mixed valent electrode was cycled offline for 5 times and then failed, so the analysis of the XRD pattern for the failed, discharged material will comprise both secondary phases produced during cycle life and degradation products from the failure. Fig. 11 shows, in the lower part, the XRD pattern of the failed mixed valent cell fitted to the mixed valent structural model. There are a number of extra reflections, which indicates the variety of by-products present in the cathode. The upper part



Fig. 9 (a) XRD patterns of the fresh V^{4+} electrode and the electrode at the start of charge in the 11^{th} cycle. (b) XRD pattern of the material at the start of the charge in the 11^{th} cycle fitted to that of the initial V^{4+} material. Red arrows mark the broad reflections and blue arrows the sharp ones, showing that two species have been formed. Upper diagram is a zoom of the wider one below.

of the figure is an enlargement of the 7 to 14° range, where the arrows indicate some reflections that are observed in the cycled V^{4+} electrode above. Surprisingly most of these are also observed in the mixed valent failed electrode in addition to some new reflections.



Fig. 10 (a) XRD patterns at the start of charge (red), and the discharged state (black) of the 11th cycle and (b) detail of XRD patterns in the 7– 8.6° range with the multiple phases seen at the beginning of the 11th cycle by the split reflection at ~7.9°.

In order to compare the similarities and differences between the cycled V^{4+} and failed mixed valent electrode they are plotted together in Fig. 12. The zones highlighted in blue correspond to mixed valent or V^{4+} phases, black arrows mark new reflections only seen in the failed mixed valent electrode and the purple arrows point out new reflections observed in both cases. It must be remarked that the relative intensity of these latter reflections are higher in the failed mixed valent sample than in the cycled V^{4+} material.

The reflections that are common for both materials can be divided into sharp and broad peaks, so the composition of the V^{4+} cycled material consists of the V^{4+} phases and at least two unidentified by-products, whereas the failed mixed valent electrode is made up of the mixed valent phases, at least two unidentified by-products that are related to the aging of these materials with cycle life, common with the V^{4+} electrode, and at least one other phase produced by the cell failure. This failure can be due to different reasons, such as exposure to air, formation of dendrites, *etc.* The next step for the analysis of mid-term cycling-related degradation mechanisms will be undertaken on full cells with an anode, so the possibility of dendrites would be eliminated.

Conclusions

In this work, the electrochemical mechanisms involved in overcharge and repeated cycling-related degradation have been illustrated. The structural evolution during cycling is affected by the compositional differences between V^{4+} and $V^{3.8+}$ phases, leading to different combinations of biphasic and single-phase stages for both, with normal charge up to 4.3 V and overcharge up to 4.8 V.

The charge limits of both compounds have been explored. Whereas the V⁴⁺ phase reaches the Na⁺ extraction limit at the usual cutoff voltage of 4.3 V; the V^{3.8+} phase is able to reach the maximum theoretical Na⁺ extraction limit when charging up to 4.8 V. Based on the conditions of our work, *e.g.*, V⁴⁺ not carboncoated and V^{3.8+} carbon coated, the V⁴⁺ sample appears to reach a point where the structure does not easily provide further sodium when charged to 4.3 V while $V^{3.8+}$ appears to be more flexible at this voltage. It should be noted that the computational calculations suggest that the V⁴⁺ can be further desodiated to 5.3 V (ref. 13) but these voltages require specialised electrolytes and are not easily accessible at this stage. It may be possible to try using highly concentrated electrolyte solutions²⁷⁻²⁹ to achieve these voltages in sodium-ion batteries in order to determine the mechanism at 5.3 V in the future. The other experimental variable to be considered is the use of carbon coating which was applied to the V^{3.8+} sample but not the V⁴⁺ sample. Overall, the mixed valent electrode is not at the extreme limit during the normal 4.3 V operation which positively affects the cycle-life performance of this compound compared to the V^{4+} phase.

The higher stress suffered by the more ordered V⁴⁺ phase is also revealed when analyzing particle size broadening. During conventional 4.3 V charging the particle size distribution is not significantly affected, but there is a large 52% reduction when using a potentiostatic step at 4.3 V and an even larger reduction in particle size distribution when charging up to 4.8 V. On the other hand, the mixed-valent phase exhibits a significantly smaller variation in particle size, even when overcharging.



Fig. 12 Comparison of XRD patterns of the cycled V⁴⁺ electrode (in red) and failed mixed valent material (in black). Zones highlighted in blue are those corresponding to mixed valent or V⁴⁺ phases, purple arrows point out new reflections observed in both cases and black arrows mark new reflections only seen in the failed mixed valent electrode

Fig. 11 Fit of the mixed valent model to the failed cell synchrotron XRD data (red crosses are the data, the black line is the calculated model, the purple line is the difference between the data and model and the vertical black lines are the reflection markers for the mixed valent model). Some reflections found in Fig. 10 are very similar to those found here, thus there are distinct similarities between mixed valent and V⁴⁺ cells. Red arrows mark the broad reflections and blue arrows mark the sharp ones, showing that at least two species have been formed. Upper diagram is a zoom of the wider one below.

9

8

10

12

14

 2θ (degrees)

16

18

10

11

12

20

22

24

13

Moreover, the V^{4+} electrode shows two phases when discharged after overcharge, while the $V^{3.8+}$ material remains monophasic.

The higher tolerance of the $V^{3.8+}$ material to stress can be related to the structural disorder characteristic of this phase, where some O are substituted by F atoms in the vanadium coordination octahedra, conferring to this material the "flexibility" to adapt better to the structural changes induced by continuous Na⁺ extraction up to different cutoff limits analysed in this work.

Examination of medium-term cycling for the V^{4+} phase indicates that secondary products are already present after 10 cycles, and these phases can be classified into crystalline and more amorphous or nanocrystalline phases. Analysis of the failed cathode of the $V^{3.8+}$ material showed that the by-products observed in the cycled V^{4+} phase were also present in this case, together with another set of diffraction peaks, probably linked to the cell exposure to air or the failure mechanism.

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