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## PAPER

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## 1 Introduction

In recent years, phosphates with sodium super ionic conductor (NASICON) structure have been a subject of great interest for their application as electrodes in sodium-ion batteries.<sup>1-3</sup> The most prominent example is  $Na_3V_2(PO_4)_3$  (NVP), which is used as a cathode because of its relatively high redox potential of about 3.4 V vs. Na<sup>+</sup>/Na, assigned to the  $V^{3+}/V^{4+}$  redox couple.<sup>2-4</sup> On the other hand, NaTi<sub>2</sub>(PO4)<sub>3</sub> (NTP) exhibits a relatively low redox potential of 2.1 V vs. Na<sup>+</sup>/Na attributed to the Ti<sup>3+</sup>/Ti<sup>4+</sup> redox couple, which makes it a suitable anode material.<sup>1,2,5</sup> The combination of NVP and NTP in a full cell is especially attractive because the overall cell voltage allows the use of aqueous electrolytes.6 Compared to organic electrolytes, such electrolytes are non-toxic, non-flammable, less expensive and more environmentally friendly, making aqueous sodium-ion batteries particularly appealing for future large-scale stationary energy storage systems.<sup>7,8</sup> However, the application of NVP in aqueous

# *Operando* monitoring the redox activity of sodium vanadium titanium phosphate electrodes in organic and aqueous electrolytes by magnetometry<sup>†</sup>

Benedikt Huemer,‡<sup>a</sup> Anna Jodlbauer,‡<sup>b</sup> Martin Wilkening, <sup>b</sup><sup>c</sup> Heinz Krenn,<sup>d</sup> Peter Knoll,<sup>d</sup> Roland Würschum, <sup>b</sup><sup>a</sup> Ilie Hanzu <sup>b</sup><sup>c</sup> and Stefan Topolovec <sup>\*a</sup>

A sol-gel method was used for the synthesis of composite electrodes consisting of sodium vanadium titanium phosphate (NVTP) and carbon. X-ray diffraction studies confirm that the NVTP possesses the NASICON crystalline structure, and cell tests indicate that the electrodes exhibit good electrochemical performance. Temperature-dependent magnetic susceptibility measurements reveal the presence of the V<sup>3+</sup> and Ti<sup>4+</sup> oxidation states in the pristine Na<sub>2</sub>VTi(PO<sub>4</sub>)<sub>3</sub>. The change in oxidation states during de-/ sodiation was monitored continuously by means of *operando* magnetometry measurements in both organic (1 M NaPF<sub>6</sub> in EC:PC) and aqueous electrolyte (1 M Na<sub>2</sub>SO<sub>4</sub>). The variation of the magnetic susceptibility demonstrates that in both types of electrolytes, the V<sup>3+</sup>/V<sup>4+</sup>redox couple is active in the high potential regime, whereas the Ti<sup>3+</sup>/Ti<sup>4+</sup> couple operates in the low potential regime, and that no significant side reactions occur. Furthermore, it is shown that the two redox couples are distinctly separated and occur in a consecutive manner. Additionally, the results of our *operando* study indicate that both redox couples contribute approximately equally to the total capacity of NVTP.

electrolytes is limited by the rapid degradation caused by the dissolution of vanadium.<sup>6,9</sup>

In 2015 Mason et al.<sup>10</sup> showed, that this degradation can be prevented by substituting half of the vanadium with titanium, and, that the resulting sodium vanadium titanium phosphate  $(Na_2VTi(PO_4)_3, NVTP)$  still exhibits a high potential redox couple. Further studies demonstrated, that multiple redox plateaus emerge for NVTP in organic11-16 as well as aqueous electrolytes,17-22 and, that symmetric full cells can be constructed using NVTP.11,13,14,17-21 Such symmetric NVTP cells have also recently been investigated for their potential use as electrochemical desalination devices.<sup>23,24</sup> In situ X-ray absorption spectroscopy measurements revealed that in organic electrolyte, the  $V^{3+}/V^{4+}$ redox couple is active in the high potential regime, while Ti<sup>3+</sup>/Ti<sup>4+</sup> is redox active at low potentials.11 Furthermore the redox activity of vanadium at high potentials and of titanium at low potentials was observed by ex situ X-ray photoelectron spectroscopy measurements in organic16 as well as aqueous electrolytes.17,19,21,23

One limitation of these spectroscopic techniques is that the spectra are recorded only at specific charging states. Continuous monitoring of the oxidation states by an *operando* technique would be particularly advantageous for electrodes containing multiple redox centers, such as NVTP, as it would allow for a more precise distinction between the various redox regimes. For instance, the behavior in the transition region could be resolved. Furthermore, in the individual studies mentioned above, these techniques were either applied solely in organic or aqueous electrolyte. A direct comparison of the redox

<sup>&</sup>lt;sup>a</sup>Institute of Materials Physics, Graz University of Technology, NAWI Graz, Graz, Austria. E-mail: stefan.topolovec@tugraz.at

<sup>&</sup>lt;sup>b</sup>Institute for Chemistry and Technology of Materials, Graz University of Technology, NAWI Graz, Graz, Austria

<sup>&</sup>lt;sup>c</sup>ALISTORE – ERI European Research Institute, CNRS FR3104, Hub de l'Energie, Rue Baudelocque, F-80039 Amiens, France

<sup>&</sup>lt;sup>d</sup>Institute of Physics, University of Graz, NAWI Graz, Graz, Austria

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<sup>‡</sup> These authors contributed equally to this work.

activities in both aqueous and organic electrolytes is still lacking.

In this study we conducted operando magnetic susceptibility measurements on NVTP electrodes in organic and aqueous electrolytes. This technique enables the distinction of different redox couples based on the changing magnetic susceptibility, as well as a quantitative assessment if all electrons participate in these redox reactions.<sup>25-29</sup> This quantification is of prime interest in the context of NVTP in aqueous electrolytes, since several studies reported parasitic side reactions in the low potential regime.<sup>18,20</sup> We show in this work that the V<sup>3+</sup>/V<sup>4+</sup> and Ti<sup>3+</sup>/Ti<sup>4+</sup> redox regimes are well separated in both electrolytes, with both redox couples contributing approximately equally to the total capacity. Moreover, we elucidate that the charge compensation processes of NVTP are identical in both types of electrolyte, and that in both cases most of the charge is consumed for the oxidation/reduction of the NVTP and not for side reactions. As magnetometry is an integral method, our measurements also clearly show that the changes in the oxidation states occur uniformly throughout the bulk of the electrodes.

## 2 Experimental

#### 2.1 Material synthesis

Sodium vanadium titanium phosphate carbon composite (NVTP(a)C) was prepared by a sol-gel synthesis procedure.<sup>24</sup> Citric acid monohydrate (Sigma Aldrich, 99.5-102 %), acting as the gel forming agent, ammonium metavandate (Sigma Aldrich,  $\geq$ 99 %) and sodium dihydrogen phosphate (Sigma Aldrich,  $\geq$ 99%) were dissolved in 200 ml deionized water. The molar ratio between the reactants can be described by 1:1:3 (corresponding to 0.04 M: 0.04 M: 0.12 M in concentration units). The solution was heated up to 353 K and stirred for 7 hours at this temperature. While stirring the solution turned dark blue indicating a reduction process from V<sup>4+</sup> to V<sup>3+</sup>. A second solution was prepared where glacial acetic acid (Sigma Aldrich,  $\geq$ 99%) and titanium isopropoxide (Sigma Aldrich, >97 %) in a molar ratio of 1:1 (0.04 M: 0.04 M in concentration units) were dissolved in pure ethanol. This mixture was then stirred for 1 hour at room temperature. Both solutions were poured together and heated to 373 K until the solvents have fully evaporated, leaving a gel in the beaker. Drying the gel over night at 333 K yielded a powder which was grinded by hand using an agate mortar. Thermal treatment was performed in a two step program on the fine powder using a quartz synthesis tube inserted into a tube furnace. In the first step, the powder was heated to 623 K with a heating rate of 3 K min<sup>-1</sup> followed by a holding time of 5 hours, then a second heating step was carried out heating with the same rate up to 1223 K and holding this temperature for 12 hours. Finally, the sample was cooled down to room temperature by the natural cooling ramp of the oven. The whole thermal treatment was carried out strictly under nitrogen atmosphere. During the synthesis the citric acid converts into conductive carbon resulting in a NVTP@C composite. The obtained powder was again grounded by hand in an agate mortar before it was used for further analysis.

#### 2.2 XRD and TGA measurements

X-ray measurements were performed using a Rigaku Miniflex 600 diffractometer ( $2\theta$  Bragg–Brentano geometry, Cu K $\alpha$ ) with a D/teX Ultra silicon strip detector. The data was recorded in a  $2\theta$  range from 10 °–60° with a step size of 0.02.

For the thermogravimetric analysis (TGA) a Netzsch STA 449 F1 Jupiter thermal analyzer was used. The measurements were conducted in a temperature range from 303–1223 K.

#### 2.3 Cell tests

For electrochemical cell tests slurries were prepared containing 80 wt% NVTP@C, 5 wt% carbon black (Super C, from TimCal) and 15 wt% PVDF binder dispersed in NMP. This composition of the slurry was taken over from the work of Shrivastava et al.24 The slurries were mixed in a planetary mill (Fritsch Pulverisette 7) for 15 minutes followed by a 5 minutes break for a total of four cycles. The dispersion was then blade coated (coating gap: 200  $\mu$ m) on Al foil (Goodfellow, 50  $\mu$ m thick) and the substrates were dried at 333 K for 12 hours. Prior to coating the slurry, the Al foil was etched by using 5 wt% KOH solution in water, followed by rinsing it with deionized water and drying. Electrodes were punched out with a diameter of 8.5 mm and their weight was determined. Prior to this, they were dried at 333 K in a vacuum oven (Büchi) for 24 hours to remove possible solvent residues. For the cell assembly three-electrode Swagelok cells were used stacking the dried electrodes, a glass fibre separator (Whatman GF/B) and sodium metal as counter and reference electrodes on top of each other. As an electrolyte a mixture of EC: PC 1: 1 (vol. ratio) containing 1 M of  $NaPF_6$  was used. The cells were electrochemically characterized by cyclic voltammetry as well as galvanostatic cycling with potential limitation (GCPL) using a Biologic MPG-2 potentiostat.



Fig. 1 XRD pattern of Na<sub>2</sub>VTi(PO<sub>4</sub>)<sub>3</sub> with reference patterns of NaTi<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub> (ICSD 29079) and Na<sub>3</sub>V<sub>2</sub>(PO4)<sub>3</sub> (ICSD 248140); the main reflexes are indexed according to the crystal structure  $R\bar{3}c$ .

#### 2.4 Ex situ magnetic characterization

Temperature dependent magnetic susceptibility measurements were performed with a SQUID magnetometer (Quantum Design, MPMPS-XL-7) for the pristine NVTP@C powder. The powder was filled into a polycarbonate capsule which was then mounted in a SQUID sample holder straw. The susceptibility measurements were performed at an applied magnetic field of 5000 Oe in a temperature range of 300–5 K. The background signal arising from the polycarbonate capsule was corrected.

#### 2.5 Operando SQUID magnetometry measurements

For the *operando* SQUID magnetometry measurements during electrochemical charging/discharging a cell design established in previous works<sup>25-27</sup> was applied. As described in detail in these references, this cell enables a continuous tracking of the changes of the magnetic susceptibility of the working electrode, which is in our case the NVTP@C electrode. A description and schematic of the general measurement principle of the SQUID magnetometer, as well as sketches of the electrochemical cell design used for the *operando* measurements, are provided in Section S1 of the ESI.<sup>†</sup>

The NVTP@C electrodes were prepared using the same slurry as for the cell tests (see above). Either Al (ChemPUR, 99.999%) or Au wires (ChemPUR, 99.9%) with a diameter of 0.25 mm were used as substrate for the electrode material. Approximately 0.5 cm of one end of the wires were pressed to form a flat surface and this flat part was then dipped into the slurry. After drying the slurry overnight at ambient conditions the coated part of the wires was pressed again slightly, to ensure a good contact between the substrate and the electrode material. Afterwards the electrodes were dried at 383 K in a vacuum oven (Pfeiffer Vacuum HiCube/Carbolite) for 24 hours to remove any residual solvent. The mass of the electrode material was determined by weighing the wires before and after the dip coating.

*Operando* magnetometry measurements were conducted for cells with organic electrolyte (1 M NaPF<sub>6</sub> in EC : PC 1 : 1 vol%) as well as for cells with aqueous electrolyte (1 M Na<sub>2</sub>SO<sub>4</sub>). For the

cells with organic electrolyte, the NVTP@C electrodes dipcoated on Al were used as working electrode. Metallic sodium pressed onto the tail ends of Cu wires (ChemPur, 99.9%, 0.25 mm diameter) served as counter and reference electrodes. For the aqueous electrolyte, symmetric cells were assembled with NVTP@C electrodes dip-coated on Au as working and counter electrodes. Here it was ensured, that the counter electrode contained more electrode material. A Ag wire (ChemPur, 99.995%, 0.25 mm diameter) coated on one end with AgCl was used as Ag/AgCl quasi reference electrode. Each electrode wire was inserted in a small polyethylene tube to ensure that only the parts with the electrode material were in contact with the electrolyte and to prevent short circuits in the cell. Long PE tubes (length: 17.5 cm, diameter: 5 mm) and NMR tubes (Rotilabo, length: 17.8 cm, diameter: 4.95 mm) were used as cell compartments for the organic and aqueous electrolyte cells, respectively. The electrodes were positioned in the tubes in such a way that the working electrode was approximately in the middle and the other two electrodes approximately 6 cm above. In this way it was ensured that only the working electrode causes a magnetic flux change in the SQUID pickup coils during the susceptibility measurements. The cells with organic electrolyte were assembled in an Ar filled glove box. The aqueous electrolyte was purged with Ar gas before the assembly. All cells were closed with a polypropylene plug which had feedthroughs for the wires and which were sealed with epoxy resin.

The *operando* magnetometry measurements were performed at 300 K and an applied magnetic field of 5000 Oe. The NVTP@C electrodes were charged and discharged with a constant current density of  $\pm 12$  mA g<sup>-1</sup>.

### 3 Results and discussion

# 3.1 Characterization of pristine NVTP@C powder by XRD and TGA

In Fig. 1, the XRD pattern of the synthesized powder and a comparison with the reference patterns of  $Na_3V_2(PO_4)_3$  and  $NaTi_2(PO_4)_3$  are shown. The measurement results indicate that the reflexions of our sample match very well with the reflexions



Fig. 2 (a) Cyclic voltammetry of the NVTP@C electrode at a scan rate of  $0.1 \text{ mV s}^{-1}$ . (b) Cyclic voltammetry of the same electrode at different scan rates between  $0.1 \text{ mV s}^{-1}$  and  $5 \text{ mV s}^{-1}$ . Note the different scales of the ordinate axis.

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of the Na<sub>3</sub>V<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub> reference sample. This is expected as we wanted to incorporate Ti into the Na<sub>3</sub>V<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub> host structure. The structure of the compound can be described by space group  $R\bar{3}c$  which belongs to the NASICON structure class. Accordingly the reflexion positions can be indexed at 14.3°, 20.1°, 20.6°, 23.9°, 28.9°, 32.2°, and 36.0° to certain lattice planes (see Fig. 1).<sup>11</sup> The crystal structure can be described by [TiO<sub>6</sub>]/[VO<sub>6</sub>] octahedra and [PO<sub>4</sub>] tetrahedra which are corner-sharing to build up a 3-dimensional framework where the Na+ ions are located at interstitial sites 6b and 18b.<sup>19,30</sup>

The carbon content in the pristine NVTP@C powder was determined to be about 5 wt% by TGA measurements (see Section S2 in ESI† for details). This value coincides quite well with literature values for other NVTP@C composites prepared by sol-gel synthesis.<sup>11,13,19,23</sup> Since the TGA revealed a water content of about 3.5 wt%, the NVTP fraction in the pristine powder can be estimated to be about 91.5 wt%.

#### 3.2 Cell test

To gain insight on the redox activity of the synthesized NVTP@C material, cyclic voltammetry was performed. In Fig. 2(a) the cyclic voltammogram between 1.75 V and 4 V at a scan rate of 0.1 mV s<sup>-1</sup> is shown. As expected, the measurement yields two reversible redox peak couples, the one in the range of 2.1–2.3 V can be attributed to the redox reaction of  $\text{Ti}^{3+}/\text{Ti}^{4+}$  while the other peak between 3.3–3.4 V stands for the V<sup>3+</sup>/V<sup>4+</sup> reaction.<sup>1-5</sup> In Fig. 2(b) the influence of different scan rates is presented. With increasing scan rate the oxidation and reduction peaks shift slightly towards higher potentials, however even at scan rates as high as 5 mV s<sup>-1</sup> the two different peaks are still clearly visible.

We conducted galvanostatic cycling with potential limitation in order to characterize the cycling performance of our material. Initially the potential range 1.75-4 V was examined (see Fig. 3(a)). The coulombic efficiency reaches nearly 100 % for all cycles and without any visible decay. Cycling the cell at 0.1C resulted in a capacity as high as 105.7 mA h g<sup>-1</sup> for the first discharge. This value coincides within a few percent with literature values of NVTP@C obtained under similar conditions11,13 and therefore shows that the electrochemical properties of our NVTP@C are consistent with literature. The discrepancy between the obtained capacity and the theoretical one of NVTP (124.8 mA h  $g^{-1}$ ) can be explained partly by the reduced NVTP content in our NVTP@C sample. However, the lower value also indicates that not all active material particles can be accessed and that there are some kinetic limitations. Increasing the cycling rates leads to smaller capacities, but even at a cycling rate as high as 10 C still 5.5 mA h g<sup>-1</sup> could be reached. Capacity retention can be checked by cycling again at 0.1 C, this results in a value of 86 mA h  $g^{-1}$  for cycle number 69. This capacity value at the end of the measurement indicates only a slow capacity fade. However, the fact that the capacity at each cycling rate is not totally constant can be a hint towards slow decay of the material.

After the 70 cycles seen in Fig. 3(a) the potential range was changed to 4-2.5 V for five cycles to obtain the capacity caused by

the redox reaction of  $V^{3+}/V^{4+}$  (see Fig. 3(b)). Lastly the same was done for Ti<sup>3+</sup>/Ti<sup>4+</sup> in a range of 2.5–1.75 V. The cycling was performed with a cycling rate of 0.1 C. For the higher potential range, namely the  $V^{3+}/V^{4+}$  redox couple we obtained capacities of 43 mA h g<sup>-1</sup>, the cycling of the Ti<sup>3+</sup>/Ti<sup>4+</sup> redox couple yielded 30 mA h g<sup>-1</sup>. An overview of the different capacities is given in Table 1.

#### 3.3 Ex situ magnetic characterization

Fig. 4(a) depicts the temperature dependence of the magnetic susceptibility of the pristine NVTP@C powder. Since carbon exhibits a small diamagnetic susceptibility,<sup>31</sup> its contribution to the total magnetic susceptibility can be deemed negligible. The molar susceptibility normalized to the amount of NVTP in the composite was obtained by considering the NVTP weight fraction of about 91.5% determined by the TGA analysis (see above).

The molar susceptibility in Fig. 4(a) exhibits the typical temperature dependence of localized paramagnetic magnetic moments, like it is expected for the 3d transition metal ions in NVTP. Therefore, the susceptibility can be described by the Curie-Weiss law:<sup>32</sup>

$$\chi_{\rm mol} = \frac{C}{T - \Theta_{\rm CW}} \text{ with } C = \mu_0 \frac{N_{\rm A} \mu_{\rm eff}^2}{3k_{\rm B}}.$$
 (1)

To determine the effective magnetic moment  $\mu_{\rm eff}$  and the Curie–Weiss temperature  $\Theta_{\rm CW}$ , the inverse susceptibility was plotted as a function of temperature (see Fig. 4(b)). Thereafter, a straight line was fitted to the linear regime of this plot (150–300 K). This procedure yielded values of  $\Theta_{\rm CW} = -36$  K and  $\mu_{\rm eff} = 2.71 \mu_{\rm B}$ .

Considering that the paramagnetic susceptibility arises from the 3d transition metal ions in NVTP, the theoretical expected value of  $\mu_{\text{eff}}$  can be calculated by

$$\mu_{\text{eff}}^{2} = \sum_{i} c_{i} \mu_{i}^{2} \text{ with } \mu_{i} = 2\sqrt{S(S+1)},$$
 (2)

where  $c_i$  is the concentration of the individual transition metal ions and  $\mu_i$  the corresponding ionic magnetic moments, which can be calculated from the total spin moment S. The total spin moments and the corresponding ionic moments for the different possible formal oxidation states of the transition metal ions in NVTP are given in Table 2. The table additionally includes their presumed concentrations for different charging states of NVTP, and the resulting values of  $\mu_{eff}$ . As can be seen from this table, for the pristine material, *i.e.*,  $Na_2VTi(PO_4)_3$ , a value of  $\mu_{\rm eff} = 2.83 \mu_{\rm B}$  is expected for the oxidation states V<sup>3+</sup> and Ti<sup>4+</sup>. This value is in good agreement with the experimentally obtained result of  $2.71\mu_{\rm B}$ . Thus, it can be concluded that the initial state of the NVTP can be described quite well by this formal oxidation states and the corresponding ionic magnetic moments. The discrepancy between the two values is most likely due to experimental uncertainties or a slight off-stoichiometry in the NVTP. Another possibility to explain the discrepancy in  $\mu_{\rm eff}$ , would be a deviation from the formal oxidation states and thus of the corresponding ionic magnetic moments. However, different values of  $\mu_i$  would also affect the theoretically expected



**Fig. 3** Cycling protocol of the NVTP@C electrode. (a) Electrode cycled at different C-rates in a potential range of 1.75–4 V: specific capacity and coulombic efficiency are plotted against cycle number. (b) The same electrode cycled with a C-rate of 0.1 C in different potential ranges to gain insight into the capacities for the two different redox couples. The potential ranges chosen for the experiment are labeled in the figure. Note that the first five cycles shown in (b) correspond to the last cycles shown in (a), and that the deviations of the coulombic efficiency from around 100 % are due to a change of the C-rate or the potential range.

Table 1Overview of the different specific capacities with respect tothe potential ranges used during the GCPL experiment shown inFig. 3(b). The capacity values given always correspond to the firstdischarge cycle in a new potential range shown in the figure

Potential range/V	Redox couple	Capacity/mA h g <sup>-1</sup>		
1.75-4 2.5-4 1.75-2.5	$\begin{array}{c} {\rm Ti}^{3+}/{\rm Ti}^{4+}, {\sf V}^{3+}/{\sf V}^{4+} \\ {\sf V}^{3+}/{\sf V}^{4+} \\ {\rm Ti}^{3+}/{\rm Ti}^{4+} \end{array}$	88 43 30		

variations of the magnetic susceptibility during oxidation/ reduction. Since, as will be shown below, the observed susceptibility variations in our *operando* measurements can be well described by the theoretical slopes assuming the formal oxidation states and the corresponding values of  $\mu_i$ , it can be concluded that these deviations from the formal oxidation states, if any, are only small.§

#### 3.4 Operando SQUID magnetometry measurements

Fig. 5 summarizes all the results of the operando SQUID magnetometry measurements. It is evident from this figure that, for all experiments, a systematic variation of the magnetic susceptibility  $\chi$  (gray dots) occurred while charging and discharging the NVTP@C with constant current (see green lines for potential profiles). Prior to examining the individual measurements, we will initially discuss which changes of the magnetic susceptibility can be theoretically expected with sodiation and desodiation of the NVTP@C electrodes. As the magnetic signal of the carbon is negligible small and the de-/sodiation will exclusively occur at the NVTP, it is sufficient to consider the magnetic susceptibility of NVTP alone. As can be seen from the eqn (1) the susceptibility of NVTP is proportional to  $\mu_{eff}^2$ , which depends on the concentration of the transition metal ions and their corresponding ionic magnetic moments (see eqn (2)). Supposing now that the  $V^{3+}/V^{4+}$  redox couple is active for the desodiation of Na<sub>2</sub>VTi(PO<sub>4</sub>)<sub>3</sub> to NaVTi(PO<sub>4</sub>)<sub>3</sub>, the corresponding change of  $V^{3+}$  to  $V^{4+}$  would result in a decrease of the susceptibility  $\chi_{mol}$  (see Table 2). Conversely, the sodiation of Na<sub>2</sub>- $VTi(PO_4)_3$  to  $Na_3VTi(PO_4)_3$  with the  $Ti^{4+}$  to  $Ti^{3+}$  redox transition active, is expected to result in an increase of the magnetic susceptibility. Due to the different ionic magnetic moments of the vanadium and titanium oxidation states, the second reaction results in a smaller susceptibility change. If we consider now that the operando cells are charged/discharged with a constant current, thus that the change of the concentrations of the oxidation states  $c_i$  occurs continuously, linear variations of  $\chi_{mol}$  are awaited, whose slopes will be characteristic for the active redox couple.¶ These theoretical expected changes of  $\chi_{mol}$  are indicated in Fig. 5 by orange dotted lines for the redox couple of  $V^{3+}/V^{4+}$  and by blue ones for the  $Ti^{3+}/Ti^{4+}$  redox couple.

Fig. 5(a) shows the results for a cell with organic electrolyte. In this experiment, the cell was first cycled four times in the potential range of 2.5-4 V, followed by a cycle encompassing the entire potential range of 1.75-4 V. Subsequently, three cycles were conducted within the potential range of 1.75-2.5 V.|| In the initial potential range the magnetic susceptibility exhibits a linear decrease during charging and a subsequent increase

<sup>§</sup> Note that theoretically one could construct a case where two simultaneous redox reactions could yield the same slope as the formal oxidation states. However, as pointed out in Section S3 of the ESI, it is highly unlikely that this case actually occurs.

 $<sup>\</sup>P$  Note that slight deviations from the linear behavior would occur in the case that the Curie–Weiss temperature  $\varTheta_{\rm CW}$  changes with the charging state.

 $<sup>\</sup>parallel$  It should be noted that due to the higher internal resistance of the *operando* cells in comparison to conventional cells typically used for cell tests (*e.g.* Swagelok cells), minor potential discrepancies may arise, due to higher ohmic drops.



Fig. 4 (a) Molar and (b) inverse molar magnetic susceptibility of pristine NVTP@C powder as a function of temperature. The straight line in (b) shows a linear fit for the temperature range of 150–300 K according to the Curie–Weiss law (see eqn (1)). Note that as described in the text the magnetic susceptibility is normalized to the amount of NVTP in the NVTP@C composite, since the magnetic contribution of carbon is negligible.

**Table 2** Upper part: total spin moment *S* and associated ionic magnetic moment  $\mu_i$  for the various formal oxidation states of the 3d transition metals in NVTP. Lower part: presumed concentrations  $c_i$  of the transition metal oxidation states and the associate effective magnetic moment  $\mu_{eff}$  for different charging states of NVTP, along with the resulting molar magnetic susceptibility  $\chi_{mol}$  calculated according eqn (1) for 300 K and  $\Theta_{CW} = -36$  K

	v	3+			$V^{4+}$	Ti <sup>3+</sup>	Ti <sup>4+</sup>
S	1				1/2	1/2	0
$\mu_i/\mu_{\rm B}$	2.	83			1.73	1.73	0
		C	Ĉi		$\mu_{\mathrm{eff}}/\mu_{\mathrm{B}}$	$\chi_{mol}/10^{-3}$ emu mol	$l^{-1}$ Oe <sup>-1</sup>
NaVTi(PO <sub>4</sub> ) <sub>3</sub>	0	1	0	1	1.73	1.11	
$Na_2VTi(PO_4)_3$	1	0	0	1	2.83	2.98	
Na <sub>3</sub> VTi(PO <sub>4</sub> ) <sub>3</sub>	1	0	1	0	3.32	4.09	

with the same slope during discharging. As shown by the orange dotted lines the slopes align with the theoretically expected slopes of the  $V^{3+}/V^{4+}$  redox reaction, indicating that this redox couple is indeed active within this potential range and that no significant side reactions occur. The fact that the slope of the susceptibility change is constant during the whole charging/ discharging process, further reveals that the oxidation/ reduction takes places uniformly throughout the bulk of the NVTP. Upon extending the lower potential limit to 1.75 V after the fourth cycle, a transition to a reduced slope of the magnetic susceptibility can be observed in the discharge cycle once the potential is below 2.5 V. In the subsequent charging cycle, the magnetic susceptibility initially decreases with this reduced slope, followed by a transition to the higher slope. The ratio of the reduced slope to the total susceptibility change is approximately equal in the discharging and charging cycles. As shown by the blue dotted lines, the reduced slope in the low potential region coincide well with the expected slope for the Ti<sup>3+</sup>/Ti<sup>4+</sup> redox couple. This proves that this redox couple becomes active

in this potential range, and again that no considerable side reactions are occurring. In the charging cycle the transition between the two regimes occurs at a higher potential than in the discharging cycles, as indicated by the black dashed lines. The activity of the  $Ti^{3+}/Ti^{4+}$  redox couple within the low potential range is further demonstrated by the final cycles in the potential range of 1.75–2.5 V. In these cycles, the susceptibility consistently exhibits the slope expected for this redox transition.

The final cycles in the low potential range of 1.75-2.5 V where the Ti<sup>3+</sup>/Ti<sup>4+</sup> redox couple is active, are notably shorter in comparison to the cycles with the active  $V^{3+}/V^{4+}$  redox couple in the high potential range of 2.5-4 V, which were performed at the beginning of the measurement. The same outcome was observed in the cell tests, where a diminished capacity was determined for the low potential range in comparison to the high potential range (see Table 1). Since, as discussed above, the transition between the two redox regimes occurs at higher potentials in the charging cycle than in the discharging cycle (see black dashed lines in Fig. 5(a)), it can be concluded that the assumed lower capacity in the low potential range arises, because in the charging cycle the oxidation of Ti<sup>3+</sup> to Ti<sup>4+</sup> is only partially completed when the cut-off potential of 2.5 V is reached. To further investigate this charge-discharge asymmetry of the potential-capacity profiles additional measurements were performed with a second organic cell, where we always cycled in the entire potential range of 1.75-4 V (see Fig. 5(b)). Once more, a distinct separation between the  $V^{3+}/V^{4+}$  redox activity within the high potential range and the Ti<sup>3+</sup>/Ti<sup>4+</sup> activity within the low potential range is evident, as indicated by the different slopes of the susceptibility change. At the boundary between the two regimes a distinct kink appears in the susceptibility data, which reveals that even in the transition region the two reactions occur consecutively. This kink between the two slopes is observed in all charging and discharging cycles at approximately  $\Delta \chi = 0$ , which demonstrates a high reversibility for both redox couples. For each charge/discharge halfcycle the susceptibility undergoes a change with both



Fig. 5 Potential (green line) and change in magnetic susceptibility  $\Delta \chi$  (gray dots) of NVTP@C charged and discharged with 12 mA g<sup>-1</sup>. (a) Organic electrolyte (1 M NaPF<sub>6</sub> in EC:PC) cycled first between 2.5–4 V, then between 1.75–4 V, and, finally between 1.75–2.5 V. (b) Organic electrolyte (1 M NaPF<sub>6</sub> in EC:PC) cycled between 1.75–4 V. (c) Aqueous electrolyte (1 M Na<sub>2</sub>SO<sub>4</sub>) cycled first between -0.18–0.38 V, then between -1.18–0.38 V, and, finally between -1.18 to -0.18 V. The dotted lines indicate the theoretically expected change of  $\chi_{mol}$  for the V<sup>3+</sup>/V<sup>4+</sup> (orange) and the Ti<sup>3+</sup>/Ti<sup>4+</sup> (blue) redox couples in NVTP. The black dashed lines in (a) exemplary show the correlation between the kink in the susceptibility data and the potentials.

individual slopes for almost the same duration. This suggests that both redox couples contribute about equally to the total capacity when cycling within the entire potential range, and supports the notion that the different capacities for the  $V^{3+}/V^{4+}$  and  $Ti^{3+}/Ti^{4+}$  regimes observed in the cell tests are primarily due to the selected cut-off potentials.

Finally, the same measurement procedure as for the first cell was applied to a cell with aqueous electrolyte (see Fig. 5(c)). Here the potential limits of the entire potential range (-1.18-0.38 V) were adapted to the different electrolyte and reference electrode. In addition, the cut-off potential separating the high (-0.18-0.38 V) and the low potential (-1.18 to -0.18 V) ranges was adjusted, to ensure a better adaptation to the two different redox regimes. As can be seen from the susceptibility variation, the same behavior as for the organic electrolyte occurs here. The V<sup>3+</sup>/V<sup>4+</sup> redox couple is active in the high potential regime, while the Ti<sup>3+</sup>/Ti<sup>4+</sup> couple operates in the low potential regime. Again, the two redox couples are well separated from each other, as indicated by the rather sharp transition between the different slopes, and, both of them contribute about equally to the total

capacity. The good agreement between the measured and the theoretically expected susceptibility variations, implying that most of the electrons are involved in the oxidation/reduction of the NVTP and not in parasitic side reactions, is particularly noteworthy in the low potential regime, where the Ti<sup>3+</sup>/Ti<sup>4+</sup> redox couple is active. This is because a parasitic side reaction, which was assumed to be mainly the oxygen reduction reaction, has been reported in several studies in this regime.<sup>18,20</sup> Since we found indications for the occurrence of this side reaction when we did not purge the aqueous electrolyte with argon (not shown), we can assume that the careful argon purging performed for the aqueous *operando* cells has largely suppressed this unwanted reaction.

## 4 Conclusions

In conclusion, magnetometry measurements have been used to gain a detailed insight into the bulk oxidation states of the transition metals in NVTP@C electrodes and their variation during de-/sodiation in both organic and aqueous electrolytes.

#### Paper

Temperature-dependent magnetic susceptibility measurements revealed that in the pristine state the  $Na_2VTi(PO_4)_3$  in the NVTP@C powder prepared by a sol-gel synthesis procedure exhibits a good stoichiometry and that the V<sup>3+</sup> and Ti<sup>4+</sup> oxidation states are present. The systematic variations of the magnetic susceptibility occurring during charging/discharging of the electrodes showed that the  $V^{3+}/V^{4+}$  redox couple, which is active in the high potential regime, and the Ti<sup>3+</sup>/Ti<sup>4+</sup> redox couple, which becomes active at low potentials, are well separated from each other and occur one after the other when cycling over the whole potential range. In addition, these measurements revealed that there were no significant side reactions in either type of electrolyte and provided compelling evidence that the analogous charge compensation processes take place in both. Furthermore, it was demonstrated that both redox couples contribute about equally to the total capacity of NVTP and that the potential ranges must be carefully chosen to allow the individual capacities to be determined by electrochemical experiments alone. Overall, it was shown that operando magnetometry studies can provide additional insights into the charge compensation processes of electrode materials comprising multiple redox centers, and are therefore a very valuable complement to spectroscopic techniques.

## Data availability

All data supporting the findings of this study are available within the paper or the ESI.†

## Author contributions

Benedikt Huemer: investigation, formal analysis, writing – review & editing. Anna Jodlbauer: investigation, writing – original draft, visualization. Martin Wilkening: supervision. Heinz Krenn: resources. Peter Knoll: resources. Roland Würschum: writing – review & editing, supervision. Ilie Hanzu: conceptualization, writing – review & editing, supervision. Stefan Topolovec: conceptualization, formal analysis, writing – original draft, visualization, supervision.

## Conflicts of interest

There are no conflicts to declare.

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