

# Vanadium-based polyoxometalate as new material for sodium ion battery anodes

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# Abstract

Affordable energy storage is crucial for a variety of technologies. One option is sodium ion batteries (NIBs) for which, however, suitable anode materials are still a problem. We report on the application of a promising new class of materials, polyoxometallates (POMs), as an anode in NIBs. Specifically Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>]·16H<sub>2</sub>O has been synthesized and characterized. Galvanostatic tests revealed a reversible capacity of approximately 276 mA h g<sup>-1</sup> with an average discharge potential of 0.4 V, as well as a high cycling stability. The underlying mechanism was rationalized to be an insertion of Na<sup>+</sup> in between the [V<sub>10</sub>O<sub>28</sub>]<sup>6</sup>·anions rather than an intercalation into a crystal structure; the accompanying reduction of  $V^{+V}$  to  $V^{+IV}$  was confirmed by X-Ray Photoelectron Spectroscopy. Finally, a working full-cell set-up is presented with the POM as the anode, substantiating the claim that Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>]·16H<sub>2</sub>O is a promising option for future high-performing sodium ion batteries.

## Introduction

In recent years, research in the field of electrochemical energy storage has intensified as the need to reliably and economically store electricity has been identified as a crucial factor in a broad variety of fields. This includes portable electronic devices and electrically powered vehicles, which require high power and energy densities. Another area where energy storage will play a key role is the stabilization of the electricity grid. As green ways to harvest electricity from renewable sources like wind or solar power become increasingly popular, economical solutions to smoothen electricity peaks are paramount for these green technologies to play a major role in the energy mix. However, for such large-scale applications the crucial factor is neither energy nor power density, but rather cost. Thus, novel battery technologies that reduce the storage cost are being researched. One promising candidate is sodium ion batteries, due to the relative abundance of sodium in the earth's crust and, consequently, its lower cost.<sup>1–3</sup> Furthermore, in sodium ion batteries copper, current collectors can be replaced with the cheaper aluminium as sodium, as opposed to lithium, does not alloy with aluminium.<sup>4–6</sup>

However, for sodium ion batteries to become a viable alternative to lithium ion batteries, novel materials need to be developed that can compete with regard to capacity and cycle behaviour. For cathode materials, layered oxides seem to be the material of choice, with a variety of sodium metal oxides (metal: manganese, cobalt, vanadium, iron) being researched, including different compositions / doping of various metals.<sup>6–10</sup> So far, anode materials seem to be a big challenge, as graphite anodes commercially employed in lithium ion batteries are unsuitable for sodium ion batteries owing to the graphite interlayer spacing that is not sufficient to intercalate sodium ions.<sup>11</sup> Thus, several prospective anode materials are being researched

which fulfil the requirements for commercial applications of sodium ion batteries<sup>12</sup>, a selection of which is presented in Figure 1. Research focuses on different storage mechanisms for Na<sup>+</sup>-ions in the active material. Recent works on intercalation materials include amorphous TiO<sub>2</sub><sup>13</sup>, and layered P2-Na<sub>0.66</sub>[Li<sub>0.22</sub>Ti<sub>0.78</sub>]O<sub>2</sub><sup>14</sup>. The first forms Na<sub>x</sub>TiO<sub>2</sub> in-situ with an approximate capacity of 140 mA h g<sup>-1</sup> at potentials between 0.9 and 2.0 V, while the latter provides 130 mAh  $q^{-1}$  (C/10) between 0.4 – 1.25 V. Phase change materials are mainly metal oxides; titanium-based materials seem promising with capacities of approximately 200 mA h  $g^{-1}$  for Na<sub>2</sub>Ti<sub>3</sub>O<sub>7</sub><sup>15</sup>. however, only five cycles are shown. Balaya et. al. reported 180 mA h g<sup>-1</sup> for this material, with 120 mA h g<sup>-1</sup> after 50 cycles.<sup>16</sup> Li<sub>4</sub>Ti<sub>5</sub>O<sub>12</sub> was found to provide 155 mA h g<sup>-1</sup> at an average potential of 0.75V.<sup>17</sup> Furthermore, phosphate-based NASICON materials have also been explored recently.<sup>18</sup> Metal-based anodes react with Na<sup>+</sup>ions by forming alloys; often, a carbon matrix is added to alleviate volume changes. Oh et. al., for example, described the application of tin-carbon in a sodium ion full cell<sup>19</sup>. Although alloy materials are considered to be safe anodes with a high capacity, cycle stability is usually poor, due to the large volume changes during (de-)insertion, which is often alleviated by introducing a carbon matrix.<sup>20,21</sup> Qian et. al. showed that Sb/C composites can deliver around 600 mA h g<sup>-1</sup> (100 mA g<sup>-1</sup>).<sup>22</sup> Work in the field of carbon-based materials focuses on hard-carbon, which consists of partially graphitized areas and irregular regions forming cavities. Stevens et. al. found that, for graphitic regions, an intercalation mechanism takes place up to a capacity of around 100 mA h g<sup>-1</sup>, followed by an insertion of Na<sup>+</sup> in the cavities, which contributes additional 150 mA h g<sup>-1</sup> (C/10).<sup>23</sup> While hard-carbons combine reasonable capacities with high stabilities, safety problems arise due to the low insertion voltage which facilitates Na-plating. Another class of materials that has

received some attention recently is organic materials. Zhao *et. al.* and Park *et. al.*, for example, recently reported 225 and 300 mA h g<sup>-1</sup>, respectively, for Na<sub>2</sub>C<sub>8</sub>H<sub>4</sub>O<sub>4</sub> at 0.4 V<sup>24,25</sup>; here, disodium terephthalate reacts to tetrasodium terephthalate through a radical intermediate.

Polyoxometalates (POM) are anionic metal-oxo clusters of early transition metal ions in high oxidation states and they exhibit unique structural and compositional properties leading to potential applications in various areas such as catalysis, magnetism, electro- and photochemistry, as well as in materials science<sup>26–31</sup>. It has been shown that POMs can also be used as electrode material in lithium ion batteries and that their multi-electron redox properties could provide a good capacity for lithium-based systems, especially for the Keggin-based POM structures<sup>32–37</sup>. A unique characteristic of POMs as electrode material in batteries is that they do not consist of extended crystal structures, but rather of isolated, discrete polyanions, making them an intermediate between intercalation electrodes and organic radical electrodes<sup>32,34,36</sup>. Thus, POMs tap into the same advantageous mechanism like sodium terephthalate and hard-carbons, i.e., Na<sup>+</sup>-insertion in intercluster cavities. Therefore, strain on crystal structures and resulting structural deterioration can be excluded as potential reasons for capacity fading. Moreover, safety features can be enhanced when this insertion occurs at a higher potential than the one at which Naplating occurs.

Even though polyoxometalates have been tested for lithium ion batteries, no reports of polyoxometalate-based electrodes for sodium ion batteries have been published. In this work, we report on the properties of the decavanadate ion in  $Na_6[V_{10}O_{28}]$ ·16H<sub>2</sub>O POM, and show its suitability as anode in sodium ion batteries.

We first describe its structural properties, and then discuss the electrochemical data, including cyclic voltammetry and charge / discharge performance.

## **Results and Discussion**

**Material characterization of Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>]-16H<sub>2</sub>O.** <sup>51</sup>V-NMR measurements revealed the characteristic signals of the decavanadate,  $[V_{10}O_{28}]^{6-}$  (-514, -500 and -424 ppm), indicating the successful synthesis of the targeted structure (Fig. 2a)<sup>38,39</sup>. This was confirmed by the FTIR spectrum, which showed the typical absorption bands of  $[V_{10}O_{28}]^{6-}$ , with asym. stretching at 847 and 746, and symmetric stretching at 521 cm<sup>-1</sup> being attributable to the V-O-V bridging, while the mode at 956 cm<sup>-1</sup> is attributed to the terminal V=O<sub>stretch</sub> bond (Fig. 2b)<sup>40,41</sup>. Inductively coupled plasma analysis revealed Na-V ratio of 0.59, confirming the expected stoichiometry within the POM.

TGA of the POM shows a weight loss starting already at temperatures slightly above room temperature, which continues until 220 °C (Fig. 2c). The total weight loss of 21% is consistent with the removal of 16 water molecules from the structure, indicating the presence of 16 molecules of crystal water per formula unit. Thus, the overall chemical formula is  $Na_6[V_{10}O_{28}] \cdot 16H_2O$ . As the coated electrode were dried at 110 °C under vacuum for several hours, it can safely be assumed that no crystal water is present after the drying process, and the chemical composition of the POM on the electrode is hence  $Na_6[V_{10}O_{28}]$ .

Surface area of the material, as measured by BET, was 21 m<sup>2</sup> g<sup>-1</sup>; the corresponding adsorption-desorption isotherms can be seen in Figure 3a. Pore-size distribution, as elucidated by BJH, ranges from  $\sim$  3 - 5 nm, indicating mesoporosity,

for the majority of pores; for the remainder, pore size varies between ~ 5 - 160 nm (Fig. 3b).

According to FESEM and HRTEM measurements (Fig. 4), the Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>] 16H<sub>2</sub>O POM shows a rod-like morphology in the solid state; the micro-rods have a diameter of 0.5–1  $\mu$ m and a typical length of several micrometers, resulting in a typical aspect ratio of 5-10. According to TEM, the POM microrods are polycrystalline, and show a random orientation with d-spacings of 0.769 and 0.333 nm. These findings are consistent with the ICSD data file 66807. As discussed above, one characteristic of discrete POMs is that they do not consist of extended crystal structures, but rather consist of individual, well-separated polyanions. In the case of decavanadate examined in this work, the clusters consist of  $[V_{10}O_{28}]^{6-}$  units, comprising ten edgeshared VO<sub>6</sub> octahedra. As discussed in previous publications about the lithium salt, Li<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>], crystal water molecules and counter cations separate the individual  $[V_{10}O_{28}]^{6-}$  ions. Upon heating, the crystal water can be removed so that the material consists of individual  $[V_{10}O_{28}]^{6-}$  clusters separated only by the alkali ions. As the same  $[V_{10}O_{28}]^6$  units make up the structure of  $Na_6[V_{10}O_{28}]$  16H<sub>2</sub>O, and TGA shows that crystal water molecules can be removed from the structure, the same overall behavior is assumed. The lattice would then consist only of sodium ions separating the individual [V<sub>10</sub>O<sub>28</sub>]<sup>6-</sup> clusters. Thus, sodium is not incorporated into the crystal structure of the  $[V_{10}O_{28}]^{6-}$  clusters, but rather inserted into the space between different clusters.

**Electrochemical measurements.** In the cyclic voltammogram the half-cell is first discharged, and subsequently cycled between 0.01 V and 3 V. In the first discharge, i.e., in the first process of inserting Na<sup>+</sup> ions into the POM structure, various distinct

reduction peaks can be seen at 2.8, 2.2, 1.1, and 0.2 V in Figure 5a. This indicates distinct reduction or ordering processes. However, as the corresponding oxidation peaks are, if existent, significantly less pronounced, these processes are irreversible. In the subsequent cycles, no distinct peaks are occurring and the electrochemical processes do not occur at specific voltages. Hence, the cyclic voltammogram resembles that of a supercapacitor rather than that of a typical intercalation process for the major part of the voltage range. However, two minor peaks become visible at a potential of 0.01 V for the reduction and 0.09 for the oxidation, indicating a defined electrochemical process. This can potentially be attributed to the acetylene black in the electrode.

The capacitive behavior can be explained by the special structural and morphological characteristics of POMs discussed previously. It has been shown before for lithium ion batteries that POMs do not act as an intercalation material. In fact, the ions move between the polyanionic clusters and seem to be adsorbed at the surface.<sup>35,36</sup> Thus, the capacitive process for Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>] is based on a continuous insertion of Na<sup>+</sup>-ions into the cavities between the [V<sub>10</sub>O<sub>28</sub>]<sup>6</sup> clusters. X-Ray Photoelectron Spectroscopy was employed to examine whether the behavior is only physical adsorption, or accompanied by electrochemical redox reactions. As Figure 6 indicates, in the pristine POM material only vanadium with an oxidation state of +V is present. When discharged, however, vanadium with an oxidation state of +IV can be observed as well. Thus, it can be said that Na<sup>+</sup>-ions are not physically adsorbed on the surface of the [V<sub>10</sub>O<sub>28</sub>]<sup>6-</sup> clusters, but a reduction from  $V^{+V}$  to  $V^{+IV}$  is induced. Therefore, these clusters seem to act as redox centers in the sense that Na<sup>+</sup>-ions are taken up between them while vanadium ions are reduced to maintain the charge balance.

The hypothesis of a continuous insertion of Na<sup>+</sup>-ions is substantiated by the charge / discharge profile, in which the lack of plateaux indicates a steady insertion of Na<sup>+</sup> (Fig. 5b). Unlike phase change materials, in which Na<sup>+</sup> continuously occupies a defined lattice space and thus triggers a structural change at a specified voltage, POMs do not exhibit these defined insertion processes at a distinct voltage. This variety of insertion sites results in a variety of chemical potentials of the Na<sup>+</sup>-ions in the material, which in turn results in a virtually continuous distribution of insertion voltages.<sup>23</sup> This would explain the absence of peaks in the CV and the slopy charge / discharge curves, rather than a distinct plateau. In previous works, similar-looking curves for LIBs have been fitted using the Redlich-Kister relation.<sup>42,43</sup> However, as we do not yet have a full understanding of the Na<sup>+</sup>-insertion mechanism into our POM and have doubts if the theoretical background of the Redlich-Kister (thermodynamic of solutions<sup>42</sup>) derivation justify its application in this case, we refrain from performing this fit. The average discharge potential, as measured by the potential at which the area under the 10<sup>th</sup> discharge curve was cut in half, was found to be 0.4 V.

In the first discharge cycles, the capacities exceed 300 mA h g<sup>-1</sup>; these high values are not achieved in subsequent cycles and can probably be attributed to the formation of a solid-electrolyte interphase, based on the partial decomposition of electrolyte. This is a known phenomenon and is also substantiated by the charges transferred during cyclic voltammetry as found by integration. While for the first cycle the charge transferred during reduction is approximately 150% of the charge transferred during reduction, this number decreases for subsequent cycles. For the following cycles, discharge capacities stabilize and for the tenth cycle reach approximately 276 mA h g<sup>-1</sup> for a current rate of 20 mA g<sup>-1</sup>. As the rate test in Figure

5c shows, discharge capacities decrease for higher current rates, giving 221 mA h g<sup>-1</sup> for 50 mA g<sup>-1</sup> (cycle 15), 173 mA h g<sup>-1</sup> for 100 mA g<sup>-1</sup> (cycle 20) and 97 mA h g<sup>-1</sup> for 200 mA g<sup>-1</sup> (cycle 25). When changing the current rate again back to 50 mA g<sup>-1</sup>, discharge capacity is rather stable on a high level, with 220 mA h g<sup>-1</sup> for cycle 27, compared to 221 mA h g<sup>-1</sup> for cycle 15. This indicates that the fading, even though capacities are lower at higher current rates, is not exacerbated when increasing the currents. The reason for this remarkable cycle stability is probably the unique structural and morphological features of POMs. As the Na<sup>+</sup>-ions (de-)insert into / from cavities between the [V<sub>10</sub>O<sub>28</sub>]<sup>6-</sup> clusters and are not built into a defined crystal lattice, structural strain is minimized. Consequently, a major cause for capacity fading is possibly due to electrolyte decomposition; as reported in the literature, this can be remedied by adding fluorinated ethylene carbonate to the electrolyte.<sup>5</sup> Tests to examine the effect of this additive to the POM system are under way.

The achieved capacities are in the same regions as the values reported for hard carbon (Fig. 1). However, as the insertion process takes place at higher voltages, the risk of Na-plating is eliminated, which leads to an increase in safety. When compared to phase-change materials like  $Li_4Ti_5O_{12}$  and  $Na_2Ti_3O_7$ ,  $Na_6[V_{10}O_{28}]$  reaches higher capacities in a voltage region that lies between the two aforementioned materials. The same classification can be made for intercalation materials, i.e., the POM lies between  $TiO_2^{13}$  and P2-Na<sub>0.66</sub>[Li<sub>0.22</sub>Ti<sub>0.78</sub>]O<sub>2</sub><sup>14</sup> with respect to voltage, and exceeds both materials regarding capacity. Alloy materials exceed  $Na_6[V_{10}O_{28}]$  with regard to capacity; however, they typically have the disadvantage of low cycle stabilities. Even though this can be improved by embedding the active material into a matrix (e.g., carbon nanotubes), this often requires complex synthesis methods, which is a barrier

to commercial applications. Sodium terephthalates<sup>24,25</sup> match and exceed Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>] with regard to capacities, while their plateau during discharge matches the average discharge voltage of the POM.

As discussed above, the chemical formula of the electrode material can be assumed to be Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>]. With respect to this composition, the capacity of 1 Na<sup>+</sup> ion is 24.5 mA h  $g^{-1}$ . Reduction of all vanadium +V to +IV would be accompanied by an insertion of 10 Na+-ions per formula unit Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>], i.e., of 245 mA h g<sup>-1</sup>. As discussed, in the first discharge cycles the current occurring with the formation of a SEI adds to the reversible capacity of the insertion of sodium ions, which expresses itself as a rather high fading during the initial cycles. Furthermore, this is also substantiated by a coulomb efficiency of > 100% in the first cycles. This would explain that, in case 10 Na<sup>+</sup> are inserted, the measured capacity exceeds theoretical capacity in the beginning. Another potential reason could be minor electrochemical activity of the acetylene black in the electrode. When increasing the current rate to 50 mA g<sup>-1</sup>, discharge capacities drop to values around 221 mA h g<sup>-1</sup>. The insertion of Na<sup>+</sup> would have to be accompanied by a reduction of  $V^{+V}$  to  $V^{+IV}$ . According to XPS measurements, as discussed before, this reduction indeed takes place. However,  $V^{+V}$  is still present in the discharged sample. This can either be attributed to an incomplete reduction of vanadium or to a re-oxidation during ex-situ sample handling. As the measured discharged sample was briefly exposed to air during the measurement, the oxygen in the air might have re-oxidized some from  $V^{+IV}$  to  $V^{+V}$ . However, based on the achieved capacity we assumed, from a mechanistic point of view, that ten additional Na<sup>+</sup>-ions are inserted per cluster, resulting in the following reaction:

$$Na_{6}[V_{10}^{+V}O_{28}] + 10Na^{+} + 10e^{-} \rightarrow Na_{16}[V_{10}^{+V}O_{28}]$$

As an additional substantiation to the hypothesis of reversible redox reactions, a sodium full cell with Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>] as anode and spherical sodium manganese oxide<sup>44</sup> as cathode was built. It is important to note that for this full cell the target was not to optimize performance, but rather to conduct a proof-of-concept that Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>] can be used for full-cells and undergoes reversible redox reactions. As can be seen in the cyclic voltammogram in Figure 5a, this is the case. The system exhibits several reversible electrochemical processes, even though at this point the variety of peaks cannot be attributed to defined electrochemical processes, as the resulting CV is a superposition of anodic and cathodic reactions as well as complex sodium ordering processes. However, the peaks clearly indicate reversible redox processes, which substantiate the assumption that the POM undergoes electrochemical reactions. Moreover, as batteries with metallic sodium electrodes pose significant safety risks, full cells without metallic sodium are essential for application-based problems. With this full-cell set-up demonstrated here, a capacity of 140 mA h g<sup>-1</sup> (with regard to the mass of  $Na_{6}[V_{10}O_{28}]$ ) was achieved for a voltage range of 0.1 – 4.0 V. As the sodium manganese oxide was the limiting factor in this set-up, a higher mass of this material had to be used, so that the capacity with regard to the mass of NMO, would be inherently lower (as it would obviously be for the total mass of active material, i.e., anode + cathode, in the cell). However, this shows that  $Na_{6}[V_{10}O_{28}] \cdot 16H_{2}O$  not only shows promising half-cell data, but is also an interesting option for sodium ion full cells.

## Conclusion

We have introduced POMs as a new class of materials for sodium ion batteries and demonstrated their capability as anode material with good cycle stability.

 $Na_6[V_{10}O_{28}] \cdot 16H_2O$  was synthesized via a known procedure and characterized by <sup>51</sup>V-NMR and FTIR. Cyclic voltammetry and charge / discharge profiles showed no distinct peaks or plateaux respectively, but continuous slopes. This can be attributed to the insertion of  $Na^+$ -ions between the  $[V_{10}O_{28}]^{6-}$  anions instead of into defined crystal interstices. We suggest that this working principle of vanadium POM electrodes is also responsible for the good cycle stability and the negligible "damage" caused by cycling at higher current rates. Finally, we demonstrated that vanadium POM electrodes function as anodes in sodium ion full cells, using sodium manganese oxide as cathode material.

#### Methods

**Synthesis.** Sodium decavanadate, Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>]·16H<sub>2</sub>O, was synthesized according to Domaille *et. al.*<sup>38</sup>. Typically, 3 g of NaVO<sub>3</sub> is dissolved in de-ionized water (100 mL); then, 4 M HCl is added to acidify the solution until a pH of 4.8 is reached. The solution is filtered and addidtional HCl is added to maintain a pH of around 4.5. Subsequently, ethanol (95%, 200 mL) was added to precipitate Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>]·16H<sub>2</sub>O as an orange bulk product, which is then filtrated and air-dried.

**Characterization.** The product was characterized by Vanadium Nuclear Magnetic Resonance spectroscopy (<sup>51</sup>V NMR) and Fourier Transform Infrared Spectroscopy (FTIR). NMR measurements were performed on a 400-MHz JEOL-ECX instrument in 5 mm tubes at 105.4 MHz. FTIR was measured with a PerkinElmer Frontier FT-IR Spectrometer using KBr pellets. For inductively coupled plasma spectroscopy (ICP) measurements, a Dual-view Optima 5300 DV ICP-OES system was used. Field Emission Scanning Electron Microscopy (FESEM, JEOL JSM 7600F) and High Resolution Transmission Electron Microscopy (HRTEM, JOEL-JSM 2100F) were used to elucidate information on the morphology of Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>]-16H<sub>2</sub>O. N<sub>2</sub> adsorption-desorption isotherms were used to determine the Brunau-Emmet-Teller (BET) surface area and Barrett-Joyner-Halenda (BJH) pore size distribution. Thermogravimetric Analysis (TGA) was performed on a TA Instruments SDT Q600 thermobalance to test the thermal stability of the product. The flow rate of nitrogen was 100 mL min<sup>-1</sup>, and the temperature range was 20 to 800 °C with a heating rate of 5 °C min<sup>-1</sup>.

**Electrochemistry.** The composite electrodes were prepared by mixing the POM with acetylene black (Alfa Aesar, > 99%) and polyvinylidene fluoride (PVDF, Arkema, Kynar HSV 900) binder in the weight ratio 60:20:20 with N-Methyl-2-pyrrolidone (NMP) to form a homogeneous slurry. This mixture was coated on an Al foil using a doctor blade; the coating was dried in air at 80 °C to remove the NMP. The coating was punched into pieces with a diameter of 16 mm, which were then roll-pressed. Subsequently, they were dried at 110 °C under vacuum. These electrodes were then assembled in 2016 coin cells with circular metallic sodium pieces with a diameter of 16 mm as the anode, and glass fibre (Whatman) as the separator. A 1M solution of NaClO<sub>4</sub> (Sigma Aldrich,  $\geq$  98 %) in a mixture of ethylene carbonate and propylene carbonate (1:1 weight %, EC: 99%, PC: Sigma Aldrich,  $\geq$  99.7%) was employed as electrolyte. Cyclic voltammetry was measured using a BioLogic potentiostat, and a Neware battery tester was used for galvanostatic charge / discharge tests.





Illustrates the voltage profiles of different sodium ion battery anode materials. The circles represent potentials at which the respective materials have a plateau; for the other materials, the voltage profile is indicated. The capacities shown here represent approximate values after initial stabilization, i.e., after irreversible reactions in the first cycles.



Figure 2 | Different characterization methods to identify the compound. (a) Shows that the <sup>51</sup>V NMR spectrum of our POM similar to what would be expected from literature data. (b) Shows the absorption bands of the FTIR spectrum, and as an inset the polyhedral representation of  $[V_{10}O_{28}]^{6-}$ . (c) Illustrates the weight loss of the material during a TGA experiment with Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>]·16H<sub>2</sub>O.



**Figure 3 | Determination of the surface characteristics of the compound. (a)** Shows the graph resulting from BET measurements. **(b)** Illustrates the corresponding BJH pore size distribution for Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>]·16H<sub>2</sub>O.

![](_page_18_Figure_0.jpeg)

**Figure 4 | Morphological analysis of the POM. (a)**, **(b)** & **(c)** Illustrates the rod-like morphology of  $Na_6[V_{10}O_{28}] \cdot 16H_2O$  by SEM (a, b) and bright-field TEM (c) measurements. **(d)** Shows the high-resolution TEM image of the POM.

![](_page_19_Figure_0.jpeg)

Figure 5 | Electrochemical behaviour and peformance of the POM electrode in a sodium cell. (a) Shows the cyclic voltammogram (cycles 1-5) of  $Na_6[V_{10}O_{28}]$  in a half-cell set-up with 1M NaClO<sub>4</sub> in EC:PC (1:1 % weight) as electrolyte at a scan rate of 0.01 mV s<sup>-1</sup> (b) Illustrates the galvanostatic charge / discharge profile in a half-cell set-up with 1M NaClO<sub>4</sub> in EC:PC (1:1 % weight) as electrolyte at 50 mA g<sup>-1</sup>. (c) Illustrates the rate capability and stability of  $Na_6[V_{10}O_{28}]$  in a half-cell set-up with 1M NaClO<sub>4</sub> in EC:PC (1:1 % weight) as electrolyte.

![](_page_20_Figure_0.jpeg)

Figure 6 | Determination of the oxidation state of vanadium before and after cycling. Illustrates the oxidation state of vanadium in the POM by XPS measurements of (left) a pristine  $Na_6[V_{10}O_{28}]$  electrode and (right) discharged  $Na_6[V_{10}O_{28}]$  electrode. This shows that vanadium is reduced during Na<sup>+</sup>-insertion.

![](_page_21_Figure_0.jpeg)

**Figure 7 | Proof of concept for a POM –based full cell set-up. (a)** Shows the cyclic voltrammogram of the full cell at 0.2 mV/s. Anode: POM, Cathode: Spherical Na<sub>x</sub>MnO<sub>2+y</sub> (not weight balanced). Cycles 7 – 9 after cycling six times in a narrower voltage ranges. **(b)** Illustrates the full cell charge / discharge capacities with respect to the POM as well as the corresponding coulomb efficiencies. Anode: POM, Cathode: Spherical Na<sub>x</sub>MnO<sub>2+y</sub> (not weight balanced). Potential range: 0 – 3.8 V; previous cycles were cycled in narrower voltage ranges

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# Author contributions

M.S. conceived the idea of studying POMs in NIBs. S.H. and N.B. performed the electrochemical tests and wrote the manuscript. H.C. provided helpful input for the POM part of the manuscript. R. A.-O. and U.K. provided the POM as well as their expertise in this field. U.S. established the contact. H.E.H. helped with the electrochemistry part of the manuscript. S.S., P.B. and Y.Z. conducted and analysed the XPS measurements.

# Competing financial interests.

The authors declare no competing financial interests.

#### Figure legends

#### Figure 1 | Overview of different anode materials for sodium ion batteries.

Illustrates the voltage profiles of different sodium ion battery anode materials. The circles represent potentials at which the respective materials have a plateau; for the other materials, the voltage profile is indicated. The capacities shown here represent approximate values after initial stabilization, i.e., after irreversible reactions in the first cycles.

## Figure 2 | Different characterization methods to identify the compound. (a)

Shows that the <sup>51</sup>V NMR spectrum of our POM is similar to what would be expected from literature data. **(b)** Shows the absorption bands of the FTIR spectrum, and as an inset the polyhedral representation of  $[V_{10}O_{28}]^{6-}$ . **(c)** Illustrates the weight loss of the material during a TGA experiment with Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>]·16H<sub>2</sub>O.

**Figure 3** | **Determination of the surface characteristics of the compound. (a)** Shows the graph resulting from BET measurements. **(b)** Illustrates the corresponding BJH pore size distribution for  $Na_6[V_{10}O_{28}] \cdot 16H_2O$ .

**Figure 4 | Morphological analysis of the POM. (a)**, **(b)** & **(c)** Illustrates the rod-like morphology of  $Na_6[V_{10}O_{28}] \cdot 16H_2O$  by SEM (a, b) and bright-field TEM (c) measurements. **(d)** Shows the high-resolution TEM image of the POM.

Figure 5 | Electrochemical behaviour and performance of the POM electrode in a sodium cell. (a) Shows the cyclic voltammogram (cycles 1-5) of  $Na_6[V_{10}O_{28}]$  in a

half-cell set-up with 1M NaClO<sub>4</sub> in EC:PC (1:1 % weight) as electrolyte at a scan rate of 0.01 mV s<sup>-1</sup> (b) Illustrates the galvanostatic charge / discharge profile in a half-cell set-up with 1M NaClO<sub>4</sub> in EC:PC (1:1 % weight) as electrolyte at 50 mA g<sup>-1</sup>. (c) Illustrates the rate capability and stability of Na<sub>6</sub>[V<sub>10</sub>O<sub>28</sub>] in a half-cell set-up with 1M NaClO<sub>4</sub> in EC:PC (1:1 % weight) as electrolyte.

Figure 6 | Determination of the oxidation state of vanadium before and after cycling. Illustrates the oxidation state of vanadium in the POM by XPS measurements of (left) a pristine  $Na_6[V_{10}O_{28}]$  electrode and (right) discharged  $Na_6[V_{10}O_{28}]$  electrode. This shows that vanadium is reduced during  $Na^+$ -insertion.

**Figure 7 | Proof of concept for a POM –based full cell set-up. (a)** Shows the cyclic voltrammogram of the full cell at 0.2 mV/s. Anode: POM, Cathode: Spherical Na<sub>x</sub>MnO<sub>2+y</sub> (not weight balanced). Cycles 7 – 9 after cycling six times in a narrower voltage ranges. **(b)** Illustrates the full cell charge / discharge capacities with respect to the POM as well as the corresponding coulomb efficiencies. Anode: POM, Cathode: Spherical Na<sub>x</sub>MnO<sub>2+y</sub> (not weight balanced). Potential range: 0 – 3.8 V; previous cycles were cycled in narrower voltage ranges