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# Prussian blue analogues with $\text{Na}_2\text{Ni}_x\text{Co}_y\text{Mn}_z\text{Fe}(\text{CN})_6$ -multimetallic structures as positive and hydrogen vanadate as negative electrodes in aqueous Na-ion batteries for solar energy storage applications†

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Low-cost and environmentally benign aqueous Na-ion batteries are the suitable choice for large-scale solar energy storage applications. Na-ion batteries with Prussian blue analogues (PBAs) as positive and hydrogen vanadate ( $\text{H}_2\text{V}_3\text{O}_8/\text{HVO}$ ) as negative electrodes in hydrogel electrolytes exhibit excellent durability, good power capability and moderate energy density. In this work, three variants of PBAs, namely  $\text{Na}_2\text{Ni}_{0.33}\text{Co}_{0.33}\text{Mn}_{0.33}\text{Fe}(\text{CN})_6$  (PBA-1),  $\text{Na}_2\text{Ni}_{0.5}\text{Co}_{0.2}\text{Mn}_{0.3}\text{Fe}(\text{CN})_6$  (PBA-2) and  $\text{Na}_2\text{Ni}_{0.2}\text{Co}_{0.5}\text{Mn}_{0.3}\text{Fe}(\text{CN})_6$  (PBA-3), were synthesized by tuning the compositions of nitrogen-coordinated transition metals such as Ni, Co and Mn. The trend of capacity in PBAs is the function of the Ni content as  $\text{Ni}^{2+}$  is electrochemically inert. PBA-1, PBA-2 and PBA-3 exhibited specific capacities of 100, 75 and 120  $\text{mA h g}^{-1}$ , respectively, at 100  $\text{mA g}^{-1}$ . Electrochemical impedance spectroscopy analysis illustrated solid-state electronic conductivity, and PBA-1 exhibited the highest electronic conductivity of  $23 \times 10^{-4} \text{ S cm}^{-1}$  compared to PBA-2 ( $15 \times 10^{-4} \text{ S cm}^{-1}$ ) and PBA-3 ( $7.4 \times 10^{-4} \text{ S cm}^{-1}$ ). Although Ni does not contribute to the capacity, it improves the cycling life of the active material. PBA-3 demonstrated 86% capacity retention after 500 cycles at 300  $\text{mA g}^{-1}$ , whereas PBA-1 and PBA-2 respectively showed 91% and 92% retention. The lower Ni content in PBA-3 is accountable for the comparatively poor cycling performance. However, based on the optimum performances of PBAs, a 1.2 V rated PBA-1//HVO full cell was assembled, which delivered a specific capacity of 65  $\text{mA h g}^{-1}$  (at 100  $\text{mA g}^{-1}$ ), an energy density of 78  $\text{W h kg}^{-1}$  (at 121  $\text{W kg}^{-1}$ ), a power density of 520  $\text{W kg}^{-1}$  (at 31  $\text{W h kg}^{-1}$ ) and a good cycling life (81% capacity retention after 500 cycles at 300  $\text{mA g}^{-1}$ ). A 5 V/4 mA h prototype device was also developed for the practical demonstration of solar energy storage applications. Based on the energy storage performances, cost, safety and environmental issues, this Na-ion battery could be scaled up for large-scale production.

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## Introduction

Clean energy harvesting and storage are salient features for sustainable development, wherein fossil fuel-based economy is gradually phased out. Owing to the fluctuating weather conditions, harvesting of renewable energy, *i.e.*, solar, wind, *etc.*, is sporadic in nature, and therefore, large-scale energy storage devices are required. Secondary batteries are technologically mature enough to reversibly store and deliver energy on a large

scale.<sup>1</sup> Presently, lead-acid and Li-ion batteries are commercially successful products for large-scale energy storage, including load leveling and solar energy storage.<sup>2</sup> In general, the prerequisites of large-scale solar energy storage batteries are (i) low cost, (ii) environment friendliness, (iii) operational safety, (iv) long service life, and (v) moderate energy/power densities. Based on these criteria, lead-acid batteries satisfy only cost and safety issues, whereas short operational life, poor energy/power densities and lead-poisoning impede their full-fledged applications.<sup>3</sup> However, good energy/power densities, long cycle life and environment friendliness are the positive features of Li-ion batteries for large-scale energy storage; however, cost and safety concerns need to be addressed.<sup>4</sup> From the perspective of energy storage performances, Li-ion batteries are more advanced than lead-acid batteries. Hence, research interest should be directed towards Li-ion analogous chemistry, with

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safety and cost components optimally settled. In this light of foregoing, aqueous Na-ion batteries are the most suitable candidates for large-scale solar energy storage applications. Similar to Li-ion chemistry, the intercalation/deintercalation of Na ions in/from the positive and negative electrodes is the key operating principle of aqueous Na-ion batteries, whereas low-cost sodium and safe aqueous electrolytes are employed.<sup>5</sup>

Various classes of positive active materials are envisaged for aqueous Na-ion chemistries, while Prussian blue analogs (PBAs) are important materials because they have multiple redox centers and open framework structures.<sup>6</sup> PBAs belong to the family of coordination polymers with the generic structure of  $A_xM_y[M'(CN)_6]_z \cdot nH_2O$ , whereas A is an alkali metal (Li, Na, and K), and M and M' are transition metals coordinated with a CN ligand through respective nitrogen- and carbon-ends.<sup>7</sup> Most of the PBAs are synthesized for Na-ion battery applications by fixing Na as the alkali metal, Fe as the carbon-coordinated transition metal and Mn/Fe/Co/Ni, *etc.*, as the nitrogen-coordinated transition metal.<sup>8</sup> Among the various nitrogen-coordinated transition metals, Ni is electrochemically inert in the potential range of positive electrode operations, while all other transition metals are redox active.<sup>9</sup> However, bi-transition metallic compositions such as  $Na_2MnFe(CN)_6$ ,  $Na_2NiFe(CN)_6$ , and  $Na_2CoFe(CN)_6$  are widely reported in Na-ion-based energy storage systems.<sup>7</sup> Hou *et al.* investigated  $Na_2MnFe(CN)_6$  as the positive active material in aqueous Na-ion batteries, which delivers a specific capacity of 140 mA h g<sup>-1</sup> at 0.5C current load in the potential range of 0.2–1.2 V vs. standard hydrogen electrode (SHE).<sup>10</sup> Naskar *et al.* studied the electrochemical performances of  $Na_2CoFe(CN)_6$  as the positive electrode in aqueous Na-ion batteries, while it delivers specific capacities of 122, 100, 90, 84 and 80 mA h g<sup>-1</sup> respectively at 1, 2, 3, 4 and 5C rates within 0–1.2 V vs. Ag/AgCl.<sup>11</sup> The same group also reported a  $Na_2NiFe(CN)_6$ /polypyrrole composite as the positive active material in aqueous Na-ion hybrid batteries, but it exhibited a specific capacity of 73.6 mA h g<sup>-1</sup> at 2C current load within 0.05–1.2 V vs. Ag/AgCl.<sup>12</sup> Even though Ni does not contribute any storage capacity in PBAs, it improves the cycling stability by preserving the structural integrity of PBAs.<sup>9</sup> Therefore, modern research focuses on Ni-doping in PBAs at the sites of nitrogen-coordinated transition metals for achieving good cycle life. Therefore, several tri-transition metallic PBA compositions are obtained, which include  $Na_2Mn_{0.5}Ni_{0.5}Fe(CN)_6$ ,<sup>13</sup>  $Na_2Co_{0.5}Ni_{0.5}Fe(CN)_6$ ,<sup>14</sup>  $Na_2Ni_{0.17}Co_{0.83}Fe(CN)_6$ ,<sup>15</sup> and  $Na_{1.85}Ni_{0.40}Co_{0.31}Fe_{0.29}[Fe(CN)_6]_{0.97} \cdot 2.5H_2O$ .<sup>16</sup> Quan *et al.* studied  $Na_2Ni_{0.17}Co_{0.83}Fe(CN)_6$  composition as a positive electrode in a non-aqueous Na-ion system, and it delivers a specific capacity of 146 mA h g<sup>-1</sup> at 20 mA g<sup>-1</sup> with 93% capacity retention at 100 mA g<sup>-1</sup> over 100 cycles.<sup>15</sup> Naskar *et al.* studied a  $Na_2Co_{0.5}Ni_{0.5}Fe(CN)_6$  positive electrode in aqueous Na-ion batteries, which demonstrated a specific capacity of 49 mA g<sup>-1</sup> at 100 mA g<sup>-1</sup> and 90% capacity retention after 500 cycles at 200 mA g<sup>-1</sup> within 0–1.2 V vs. Ag/AgCl.<sup>14</sup> A comparative positive electrode performance study was conducted by Naskar *et al.* in  $Na_2MnFe(CN)_6$  and  $Na_2Mn_{0.5}Ni_{0.5}Fe(CN)_6$ , where the respective specific capacities were obtained as 36 and 47 mA h g<sup>-1</sup> at

100 mA g<sup>-1</sup> current density within 0–1.2 V vs. Ag/AgCl in a 1 M  $Na_2SO_4$  electrolyte. Importantly,  $Na_2Mn_{0.5}Ni_{0.5}Fe(CN)_6$  exhibits far better capacity retention (80%) than that of  $Na_2MnFe(CN)_6$  (56%) over 250 cycles at 500 mA g<sup>-1</sup> current rate.<sup>13</sup> Zhang *et al.* reported a multi-centre nitrogen-coordinated PBA, namely,  $Na_{1.85}Ni_{0.40}Co_{0.31}Fe_{0.29}[Fe(CN)_6]_{0.97} \cdot 2.5H_2O$ , which delivered a discharge capacity of 120 mA h g<sup>-1</sup> (at 20 mA g<sup>-1</sup>) and excellent cycle life (0.0044% decay per cycle at 2 A g<sup>-1</sup>) in a non-aqueous electrolyte system.<sup>16</sup> Nonetheless, tetra-transition metallic PBAs as positive active materials in Na-ion chemistry are truly limited in the literature, which envisaged further improvements in charge storage capability and durability. In this study, we synthesized multiple tetra-transition metallic PBA compositions for aqueous Na-ion batteries and investigated their electrochemical performance.

However, the appropriate negative active materials are crucial in order to accomplish the desirable energy storage performances from aqueous Na-ion batteries. Different types of negative active materials such as carbonaceous matters, including hard carbons, soft carbons, and amorphous carbons<sup>17,18</sup> and metal oxides,<sup>13</sup> metal phosphates,<sup>14</sup> metal carbides, and metal chalcogenides<sup>19–21</sup> are extensively reported for Na-ion batteries in the literature. In the metallic oxide segment, vanadium-based polyoxometalates (POMs) are promising candidates for the negative electrode of aqueous Na-ion batteries due to the following reasons: (i) high storage capacity because of multielectronic redox activity and (ii) high power capability due to facile Na-ion diffusion through the open layered structure.<sup>22–24</sup> Deng *et al.* reported a  $Na_2V_6O_{16} \cdot nH_2O$  phase of Na-vanadate as the negative electrode in aqueous Na-ion batteries, which delivered a specific capacity of 42 mA h g<sup>-1</sup> at 40 mA g<sup>-1</sup> current density within 0 to –0.65 V vs. saturated calomel electrode (SCE) window.<sup>23</sup> Naskar *et al.* studied  $NaV_3O_8$  as a negative electrode in aqueous Na-ion batteries, and it exhibited specific capacities of 83, 65, 27, 28 and 23 mA h g<sup>-1</sup> at 1, 2, 3, 4 and 5C rates respectively within 0.2 to –0.8 V vs. Ag/AgCl window.<sup>11</sup> Not vividly, Na-vanadate is reported as the negative electrode in Na-ion systems,<sup>11</sup> but hydrogen vanadate has never been reported in the literature as the negative electrode for Na-ion chemistry. We have found very minimal reports on hydrogen vanadate as the negative electrode in Li-ion systems. For example, Li *et al.* studied  $H_2V_3O_8$  (HVO) as a potential negative electrode in aqueous Li-ion batteries, while it demonstrates a specific capacity of 234 mA h g<sup>-1</sup> at 100 mA g<sup>-1</sup> current density.<sup>25</sup> Nevertheless, here we selected  $H_2V_3O_8$  as the negative electrode in aqueous Na-ion batteries; which is the first ever report on the use of HVO as the negative electrode of Na-ion chemistry.

In this work, we synthesized three different tetra-transition metallic PBAs, namely,  $Na_2Ni_{0.33}Co_{0.33}Mn_{0.33}Fe(CN)_6$  (PBA-1),  $Na_2Ni_{0.5}Co_{0.2}Mn_{0.3}Fe(CN)_6$  (PBA-2) and  $Na_2Ni_{0.2}Co_{0.5}Mn_{0.3}Fe(CN)_6$  (PBA-3) as positive electrode materials and hydrogen vanadate ( $H_2V_3O_8$ /HVO) as a negative electrode material for hydrogel-based Na-ion batteries. The synthesized positive and negative active materials were thoroughly characterized by various physicochemical and electrochemical methods. Based on the comparative electrochemical performances of PBA-1, PBA-2 and PBA-3, PBA-1 was selected as the positive



active material in Na-ion full cells. Therefore, a 1.8 V PBA-1//HVO full cell was assembled in a 1 M Na<sub>2</sub>SO<sub>4</sub>-carboxymethylcellulose (CMC)-silica (SiO<sub>2</sub>)-based hydrogel electrolyte. The full cell demonstrated a specific capacity of 65 mA h g<sup>-1</sup> (at 100 mA g<sup>-1</sup>), an energy density of 118 W h kg<sup>-1</sup> (at 181 W kg<sup>-1</sup>), a power density of 780 W kg<sup>-1</sup> (at 46 W h kg<sup>-1</sup>) and a good cycle life (81% capacity retention after 500 cycles at 300 mA g<sup>-1</sup>). This study is culminated with the development of a 5 V/4 mA h prototype device for the practical demonstration of solar energy storage applications.

## Experimental

### Synthesis of active materials

Three variants of PBAs, namely, Na<sub>2</sub>Ni<sub>0.33</sub>Co<sub>0.33</sub>Mn<sub>0.33</sub>Fe(CN)<sub>6</sub> (PBA-1), Na<sub>2</sub>Ni<sub>0.5</sub>Co<sub>0.2</sub>Mn<sub>0.3</sub>Fe(CN)<sub>6</sub> (PBA-2) and Na<sub>2</sub>Ni<sub>0.2</sub>Co<sub>0.5</sub>Mn<sub>0.3</sub>Fe(CN)<sub>6</sub> (PBA-3) were synthesized using a citrate-assisted control crystallization technique *via* an *in situ* co-precipitation process.<sup>26</sup> The stoichiometric amounts of Ni, Co and Mn salts (NiCl<sub>2</sub>·6H<sub>2</sub>O, CoCl<sub>2</sub>·6H<sub>2</sub>O and MnCl<sub>2</sub>·4H<sub>2</sub>O) and 5 mmol tri-sodium citrate were taken in a beaker containing 50 ml distilled water, named A. In an another beaker (named B), 5 mmol of Na<sub>4</sub>Fe(CN)<sub>6</sub> was dissolved in 50 ml distilled water. Then, two solutions were mixed simultaneously in another beaker containing 100 ml distilled water and stirred for 12 h. The PBA-precipitates were obtained as suspensions, and the suspensions were centrifuged three times with water and ethanol. The resulting PBA-products were obtained after drying at 70 °C for 8 h in a hot air oven. Meantime, H<sub>2</sub>V<sub>3</sub>O<sub>8</sub> (HVO) was hydrothermally synthesized,<sup>27</sup> where 0.040 g of polyethylene glycol (PEG-600) and 0.237 g of V<sub>2</sub>O<sub>5</sub> were mixed in 40 ml distilled water. Then 10 ml of 30% H<sub>2</sub>O<sub>2</sub> was added to the resulting mixture and vigorously stirred for 30 min. Thereafter, the whole solution mixture was transferred into a 50 ml Teflon-lined autoclave and heated at 180 °C for 60 h in a hot air oven. The obtained product was filtered and washed thoroughly with distilled water. The washed product was dried in a hot air oven at 70 °C for 24 h and obtained.

### Physicochemical characterizations of materials

The synthesized active materials were physicochemically characterized by powder X-ray diffraction (PXRD), Fourier transformed infrared (FT-IR) spectroscopy, thermogravimetric analysis (TGA) and scanning electron microscopy (SEM) with energy-dispersive X-ray (EDX) analysis. PXRD patterns were acquired using a Malvern Panalytical Empyrean diffractometer (Ni-filtered Cu-K $\alpha$  radiation,  $\lambda = 1.5418$  Å). PXRD data for the Rietveld refinements were acquired at room temperature with a step size of 0.02° and a step duration of 50 s. The PXRD patterns were refined using the program GSAS-EXPGUI.<sup>28</sup> Lattice parameters, scale factors, background (Fourier polynomial background function), and pseudo-Voigt (U, V, W and X) were refined. The crystal structures were sketched using the software VESTA.<sup>29</sup> The theoretical PXRD patterns were simulated using the program POWDER-CELL.<sup>30</sup> The FT-IR spectroscopic studies

were carried out using a Bruker Optics ALPHA spectrometer with a Zn-Se ATR (attenuated total reflection) accessory. The TGA experiments were conducted using a TG 8120 Thermo Plus EvoII by Rigaku Corporation (Tokyo, Japan). The SEM imaging was performed using a Carl Zeiss Supra 55 field emission scanning electron microscope equipped with an EDX (Oxford INCA) facility. The Na-ion diffusion pathways in the PBAs and HVO were revealed using the softBV (bond valence) software.<sup>31,32</sup>

### Electrochemical characterizations and cell assembly

All the positive and negative electroactive materials were electrochemically characterized by cyclic voltammetry (CV), galvanostatic charge discharge (GCD) and electrochemical impedance spectroscopy (EIS). All the measurements were conducted in a three-electrode cell configuration in a 1 M Na<sub>2</sub>SO<sub>4</sub> electrolyte, where active material-coated current collectors, platinum foil and Ag/AgCl were used as the working, counter and reference electrodes respectively. A low-cost stainless steel (SS304) sheet was employed as the potential current collector, whereas a single-walled carbon nanotube (SWCNT) coating was applied as the corrosion protective layer.<sup>11</sup> The active material slurries were prepared by mixing 80% active materials, 10% conductive carbon and 10% polyvinylidene fluoride (PVDF) binder in a *N,N*-dimethylformamide (DMF) solvent. The slurry was coated onto the current collectors (geometric area: 1 cm<sup>2</sup> & mass loading: 1 mg cm<sup>-2</sup>) by the doctor blade method, and the same were dried in a hot air oven at 60 °C for 1 h. The dried working electrodes were pressed with 15 ton pressures to achieve uniform active material thickness. A 1.8 V full cell was assembled with a (+)PBA-1//HVO(-) configuration in the prismatic cell structure, whereas a hydrogel electrolyte was employed. The optimum hydrogel was prepared by mixing 2 wt% carboxymethylcellulose (CMC) and 2 wt% fumed silica (SiO<sub>2</sub>) as gelling agents in a 1 M Na<sub>2</sub>SO<sub>4</sub> (aq.) solution.<sup>13</sup> It is worthy to mention that the neutral electrolyte was preferred in order to minimize the current collector corrosion and active material disintegration. Despite superior electrochemical performances, highly concentrated 'water-in-salt' electrolyte systems were excluded in this work because of the cost constraint. However, the two-electrode full cell was configured with one positive and one negative electrodes (geometric area: 5 × 5 cm<sup>2</sup>), while the respective mass loading values were 1 and 1.12 mg cm<sup>-2</sup>. A low-cost polyvinyl chloride (PVC)-based fencing net was placed as the separator, which offered mechanical support to the thixotropic hydrogel in between two electrodes. A 5 V/4 mA h rated prototype device was developed by series connection of three 1.8 V cells to demonstrate the solar storage performance. A custom-made solar charging module with 3 V/270 mW light-emitting-diode (LED) load was used in field trials for monitoring the device performances. All the laboratory-based electrochemical measurements were carried out using a multichannel AUTOLAB potentiostat/galvanostat workstation (Model: M204). Until otherwise stated, all the chemicals/materials were purchased from Merck, India, and used without any further purification steps; and all the experiments were performed at ambient temperature (25 °C).



## Results and discussion

### Physicochemical characterizations of positive and negative active materials

The positive active materials, namely, PBA-1, PBA-2 and PBA-3 were structurally characterized by PXRD analysis through Rietveld refinement, which revealed that all the above-mentioned PBAs were crystallized in monoclinic phase with the space group  $P2_1/n$ . Fig. 1(a)–(c) depict the Rietveld refinement results with unit cell parameters and obtained reliability factors for PBA-1, PBA-2 and PBA-3, respectively. A representative crystal structure of the PBAs is presented in Fig. 1(d). It is noted that all the three structures are quite similar, where distorted Ni/Co/Mn(NC)<sub>6</sub> octahedra and Fe(CN)<sub>6</sub> octahedra are situated alternately along the crystallographic *a*-direction leading to the construction of a three-dimensional open-framework structure with one-dimensional passages for Na ion migration. Na ions are residing in a zigzag manner in the one-dimensional channels facilitating the movement of the Na-ions. Fig. S2 in the ESI† depicts the FT-IR spectra of all the synthesized PBAs, whereas a sharp peak is observed at 2070 cm<sup>-1</sup> assigned to the stretching vibration of CN coordinated in between Fe<sup>2+</sup> and Ni<sup>2+</sup>/Co<sup>2+</sup>/Mn<sup>2+</sup>. The observed peaks at 3614 and 3546 cm<sup>-1</sup> appear due to the stretching vibration of O–H, and a peak at 1600 cm<sup>-1</sup> is ascribed to the bending vibration of O–H. Such stretching and bending vibrations of O–H indicate the presence of surface-adsorbed and crystalline H<sub>2</sub>O molecules.<sup>33</sup> The SEM images of respective PBA-1, PBA-2 and PBA-3 are shown in Fig. 2(a)–(c), which represent homogeneous cubic-shaped morphologies with ~0.14, 0.12 and

0.13 μm particle sizes respectively. The non-aggregated block-shaped particles are developed in the crystallization process due to the presence of chelating agent; as it impedes the nucleation kinetics and prevents uncontrolled crystal growth.<sup>26</sup> The elemental analysis of the PBAs was performed by EDX (Fig. S3(a)–(c), ESI†), and the obtained metallic ratios provide a good synchronization between the compositions of PBA-1, PBA-2 and PBA-3 and the molar ratios of the precursors as per the synthetic routes (Table 1). The quantification of crystalline H<sub>2</sub>O in PBAs was carried out by TGA (Fig. 2(d)–(f)), whereas surface-adsorbed and crystalline H<sub>2</sub>O molecules escaped at ~110 and ~220 °C, respectively.<sup>34</sup> From the weight loss data in TGA, it is intended that nearly two crystalline H<sub>2</sub>O molecules are present in the molecular formula of synthesized PBAs (Table 1).

The negative active material, namely, HVO, was also structurally characterized by PXRD analysis through Rietveld refinement.<sup>35</sup> Fig. 3(a) depicts the Rietveld refinement of HVO with unit cell parameters and obtained reliability factors, and Fig. 3(b) shows the resulting crystal structure.<sup>35</sup> The layers of edge-shared VO<sub>6</sub> octahedra in the *b*–*c* plane are stacked along the crystallographic *a*-direction. The layers are interconnected by means of hydrogen bonding. These interlayer spacings are expected to promote Na-ion migration in the *b*–*c* plane under applied potential in an aqueous Na<sub>2</sub>SO<sub>4</sub> solution as the electrolyte. The FT-IR spectrum of HVO is shown in Fig. S4 (ESI†), where the peaks around 1005 and 970 cm<sup>-1</sup> correspond to the stretching vibrations of V=O bonds. The peaks at 3004 and 1600 cm<sup>-1</sup> are assigned to the respective stretching and bending vibrations of O–H, arising from surface-adsorbed and

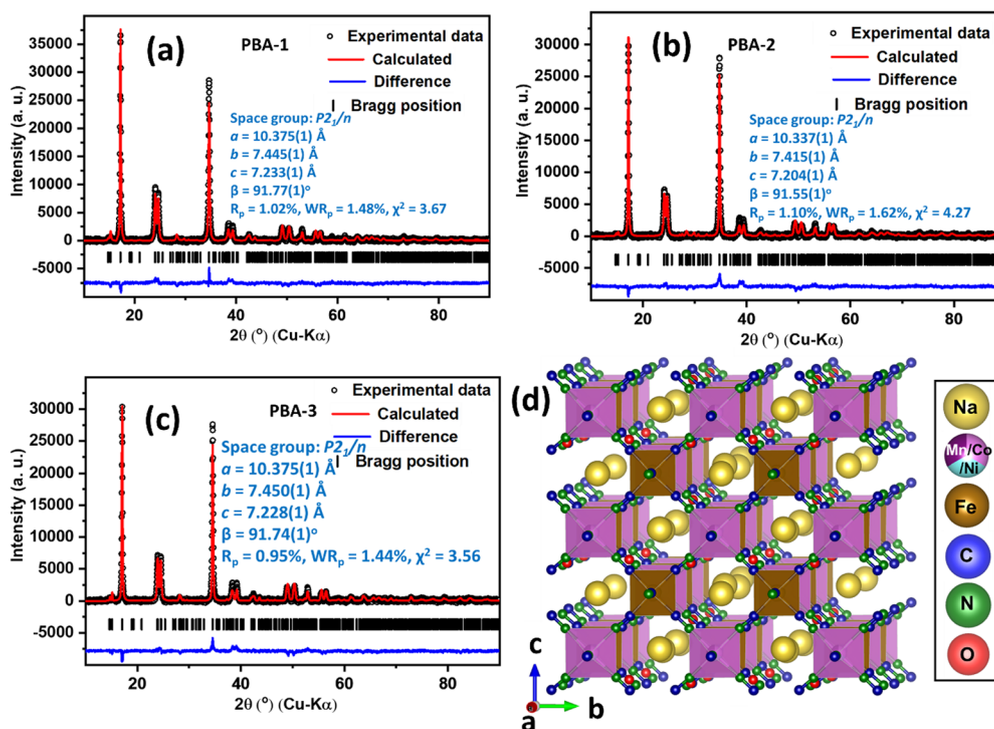


Fig. 1 Rietveld refinements of (a) PBA-1, (b) PBA-2, and (c) PBA-3 based on the monoclinic structure, unit cell parameters, and obtained reliability factors; (d) representative crystal structure of PBA (viewed along the crystallographic *a* direction).



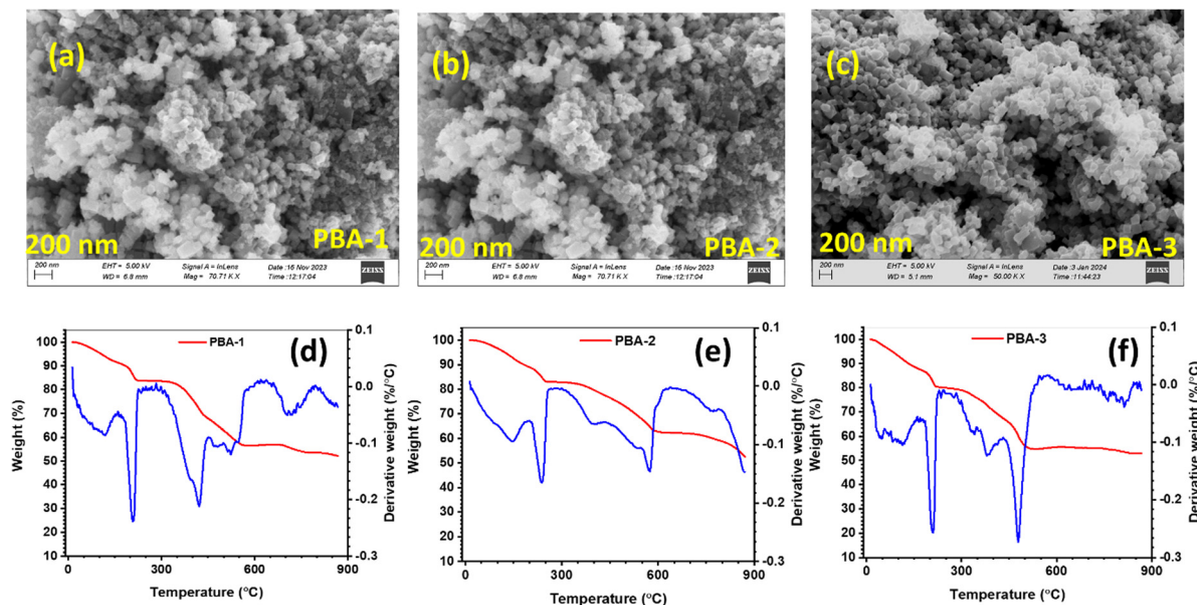


Fig. 2 SEM image of (a) PBA-1, (b) PBA-2, and (c) PBA-3; thermogravimetric profiles with derivative plots for (d) PBA-1, (e) PBA-2, and (f) PBA-3.

Table 1 Composition of PBAs with different nitrogen-coordinated transition metals

	Atomic %					No of crystalline water
	Na	Ni	Co	Mn	Fe	
PBA-1	1.0 ± 0.14	0.2 ± 0.16	0.2 ± 0.17	0.2 ± 0.21	0.5 ± 0.09	1.69
PBA-2	1.1 ± 0.14	0.4 ± 0.12	0.2 ± 0.17	0.2 ± 0.21	0.6 ± 0.09	1.76
PBA-3	4.6 ± 0.09	0.4 ± 0.06	0.9 ± 0.03	0.5 ± 0.04	1.6 ± 0.03	2.33

crystalline H<sub>2</sub>O molecules.<sup>36</sup> The SEM image and EDX profile of HVO are presented in Fig. 3(c) and Fig. S3(d) (ESI<sup>†</sup>), respectively, where rod-shaped particles are found with ~3.2 μm length and ~0.28 μm width. The rod-shaped particles accelerate the electrolyte diffusion into the bulk of the active materials, which improved the active material utilization and rate capability. The TGA experiment was also conducted for HVO (Fig. 3(d)), and it was found that crystalline waters escaped at 278 °C. The resulting weight loss indicates the number of crystalline water molecules (*i.e.*, 1.75) in the molecular structure of H<sub>2</sub>V<sub>3</sub>O<sub>8</sub>.

The bond valence energy landscape (BVEL) calculations on Na-ion diffusion pathways were carried out in all PBAs and HVO structures using the softBV software.<sup>31,32</sup> The Na-ion diffusion paths along the *b-c* and *a-b* planes in PBA-1 and HVO are demonstrated in Fig. 4(a)–(d), whereas the diffusion pathways in *b-c/a-b* planes for PBA-2 and PBA-3 are depicted in Fig. S5(a)–(d) (ESI<sup>†</sup>) respectively. Furthermore, the activation energy barriers under 3D, 2D and 1D routes are shown in Table 2. The Na-ion diffusions in all directions are probable in PBAs because of having open-framework structures. It is important to note that the activation energies and the diffusion pathways are quite comparable for all PBAs in 3D, 2D and 1D pathways, because of the resemblance in crystal structures. Besides, the activation energy barriers at 3D, 2D and 1D pathways were estimated for HVO, and are reported in Table 2. Due to the two-

dimensional layer structure of HVO, the Na-ion diffusion is limited in 2D and 1D pathways, while in the 3D pathway the activation energy is unattainable (*i.e.*, 999 eV). It is noteworthy that the BVEL calculations of Na-ion diffusion in the HVO structure were carried out by incorporating Na-ions into the crystal structure. It is expected to be close to reality, because of the use of an aqueous Na<sub>2</sub>SO<sub>4</sub> electrolyte in battery operations. However, the diffusion coefficient (*D*) values were also estimated from the EIS analysis, and the results showed  $3.09 \times 10^{-14}$ ,  $1.68 \times 10^{-14}$ ,  $1.29 \times 10^{-14}$  and  $2.50 \times 10^{-11}$  cm<sup>2</sup> s<sup>-1</sup> for PBA-1, PBA-2, PBA-3 and HVO, respectively. The higher *D*-value is expected in HVO due to the 2D layered structure; while PBAs depict a lower *D*-value because of the framework structure with one-dimensional Na-ion diffusion channels. However, the diffusion-related study confirms that no appreciable changes are observed in the diffusion properties of PBAs, when the compositions of nitrogen-coordinated transition metals are changed.

### Electrochemical characterizations of positive active materials

The electrochemical charge storage mechanisms of the synthesized PBAs were characterized by performing CV analysis at variable scan rates (5–25 mV s<sup>-1</sup>) within the 0–1.2 V window. The predominating diffusion-controlled faradaic and surface-controlled non-faradaic/pseudo-faradaic charge storage phenomena in electroactive materials could be predicted from the following power law equation:<sup>37</sup>



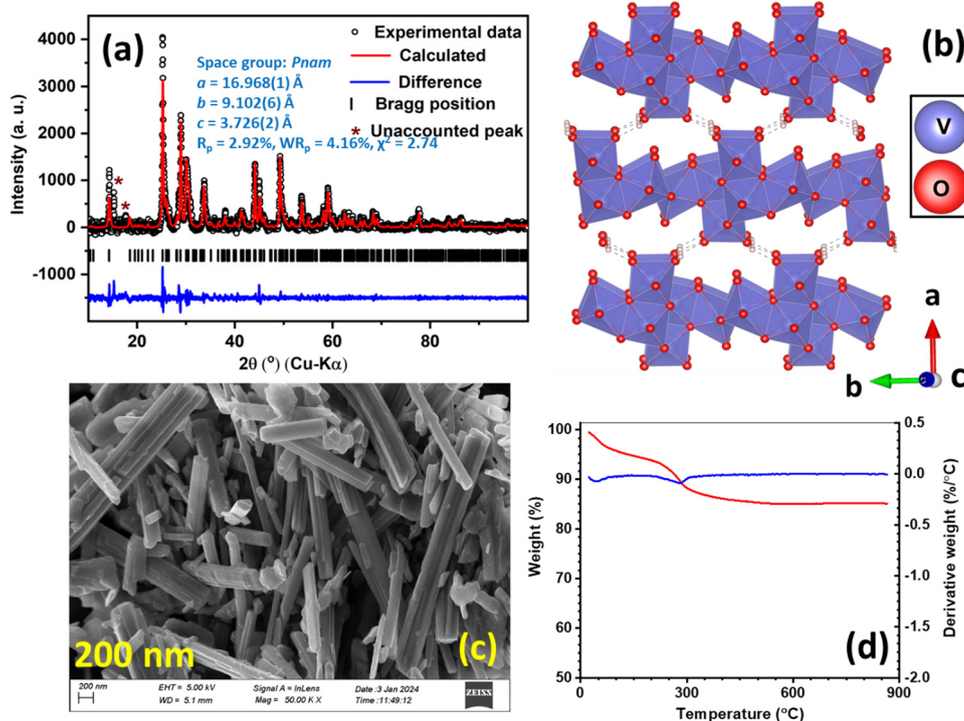


Fig. 3 (a) Rietveld refinements of HVO based on the orthorhombic structure together with the unit cell parameters and obtained reliability factors, (b) crystal structure of HVO (viewed along the crystallographic *c* direction), (c) SEM image of HVO, and (d) thermogravimetric profile with the derivative plot for HVO.

$$i = a \times \nu^b \quad (1)$$

where  $i$  and  $\nu$  are the current response and scan rate in CV, and  $a$  and  $b$  are constant values. As per the Randles–Sevcik equation,  $b \sim 0.5$  designates the faradaic mechanism, whereas  $b \sim 1$  implies the non-faradaic/pseudo-faradaic mechanisms. However, the variable scan rate CV profiles of PBAs are shown in Fig. S6(a)–(c) (ESI<sup>†</sup>), whereas prominent anodic/cathodic redox pairs are noted at 0.60/0.40, 0.67/0.56 and 0.61/0.42 V in PBA-1, PBA-2, and PBA-3, respectively. The redox activities arise in PBAs from the  $\text{Co}^{2+/3+}$ ,  $\text{Mn}^{2+/3+}$  and  $\text{Fe}^{2+/3+}$  redox pairs, while  $\text{Ni}^{2+}$  is electrochemically inert and does not contribute to the charge storage capacity of the materials.<sup>9</sup> The electrochemical responses of PBA-1, PBA-2 and PBA-3 are depicted in eqn (2)–(4), respectively:

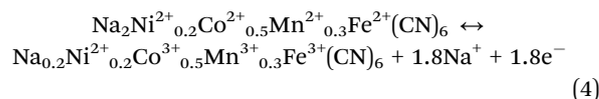
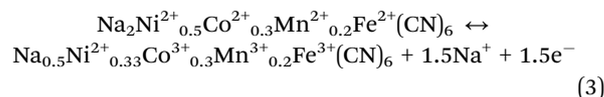
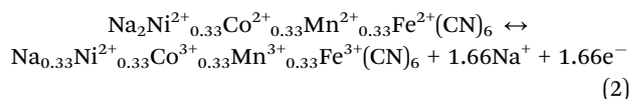


Fig. 4 Na-ion diffusion pathways found using BVEL calculations for (a) PBA-1 in the *b*–*c* plane, (b) PBA-1 in the *a*–*b* plane, (c) HVO in the *b*–*c* plane, and (d) HVO in the *a*–*b* plane.

Table 2 Activation energies for Na-ion diffusion in PBA-1, PBA-2, PBA-3 and HVO

	PBA-1 (in eV)	PBA-2 (in eV)	PBA-3 (in eV)	HVO (in eV)
3D Pathway	0.220	0.221	0.219	999
2D Pathway	0.168	0.165	0.168	1.253
1D Pathway	0.168	0.165	0.168	0.704

The separated peak pairs for  $\text{Co}^{2+/3+}$ ,  $\text{Mn}^{2+/3+}$  and  $\text{Fe}^{2+/3+}$  couples are not distinctive in the experimental CV profiles, rather the broad overlapped redox pairs are obtained.<sup>38</sup> The



calculations of 'b' values are shown in Fig. S6(d)–(f) (ESI<sup>†</sup>), while PBA-1, PBA-2 and PBA-3 exhibit 'b' values of 0.61/0.67, 0.71/0.74 and 0.60/0.64 at the respective anodic/cathodic peak positions. In brief, the 'b' values are obtained in the range of 0.60 to 0.75 that strongly correlates with the predominant diffusion-controlled faradaic charge storage mechanism. However, the both anodic and cathodic peak positions are not appreciably shifted towards the higher over potentials upon increasing the scan rate from 5 to 25 mV s<sup>-1</sup>, which signifies that the minimal mass transfer resistances are associated with the faradaic charge storage processes in PBAs. Fig. 5(a)–(c) show the GCD profiles of PBA-1, PBA-2 and PBA-3, respectively, at different current densities from 100 to 500 mA g<sup>-1</sup> within 0–1.2 V range. The prominent voltage plateaus are observed in both charge and discharge profiles for all PBAs, which further confirms the diffusion-controlled phase transformations in the faradaic charge storage mechanism.<sup>37</sup> The estimated specific capacities and coulombic efficiencies (CE) for PBA-1 are 100 (CE = 90%), 84 (CE = 93%), 76 (CE = 97%), 69 (CE = 98%) and 63 (CE = 99%) mA h g<sup>-1</sup> at 100, 200, 300, 400, and 500 mA g<sup>-1</sup> current densities respectively, while PBA-2 depicts 74 (CE = 93%), 54 (CE = 89%), 43 (CE = 93%), 41 (CE = 94%) and 39 (CE = 96%) mA h g<sup>-1</sup> and PBA-3 demonstrates 120 (CE = 85%), 73 (CE = 84%), 57 (CE = 85%), 48 (CE = 87%) and 42 (CE = 89%) mA h g<sup>-1</sup> under similar experimental conditions. The capacity trend exactly matches with the content of Ni-doping, as Ni remains electrochemically inactive during the charge/discharge processes of PBAs. The rate capability data are presented in Fig. 5(d), whereas 32, 47 and 65% capacity losses are observed for PBA-1, PBA-2 and PBA-3, respectively, when the

current density increases from 100 to 500 mA g<sup>-1</sup>. The rate capability data are also interpreted by the capacity retention values after 25 cycles by returning the current at 100 mA g<sup>-1</sup>, whereas PBA-1 (81%) and PBA-2 (84%) exhibit better retention than PBA-3 (60%). The minimal capacity losses upon higher current densities are accountable for higher electronic conductivity of the materials, while good structural stability promotes good capacity retention after multiple cycling at multiple current loads. The EIS studies were carried out on PBAs, and the resulting Nyquist plots are shown in Fig. 5(e). The solution resistance (*R<sub>s</sub>*) and charge transfer resistance (*R<sub>ct</sub>*) were obtained from the equivalent circuit fitting, and the same are presented in Table 3. The solid-state electronic conductivities ( $\rho$ ) of the PBAs were estimated from the *R<sub>ct</sub>* values using eqn (5):<sup>39</sup>

$$\rho = \frac{1}{R_{ct}} \times \frac{l}{A} \quad (5)$$

where *l* (75 × 10<sup>-4</sup> cm) is the thickness of the active material layer and *A* (1 cm<sup>2</sup>) is the geometrical surface area of the electrode. The calculated conductivities are 23 × 10<sup>-4</sup>, 15 × 10<sup>-4</sup> and 7.4 × 10<sup>-4</sup> S cm<sup>-1</sup> for PBA-1, PBA-2 and PBA-3, respectively. The compositions of Ni, Co and Mn are greatly varied among three different PBAs, which definitely differentiate their electronic band structures, while the obtained electronic conductivity harmonizes with the band structure of the material.<sup>13,16</sup> However, the trend in electronic conductivity well synchronizes with the capacity loss data of PBAs under 100 to 500 mA g<sup>-1</sup> current loads. The Ni<sup>2+</sup>-content in the PBA structure reduces the charge storage capacity, but simultaneously improves the structural integrity because of the absence of the

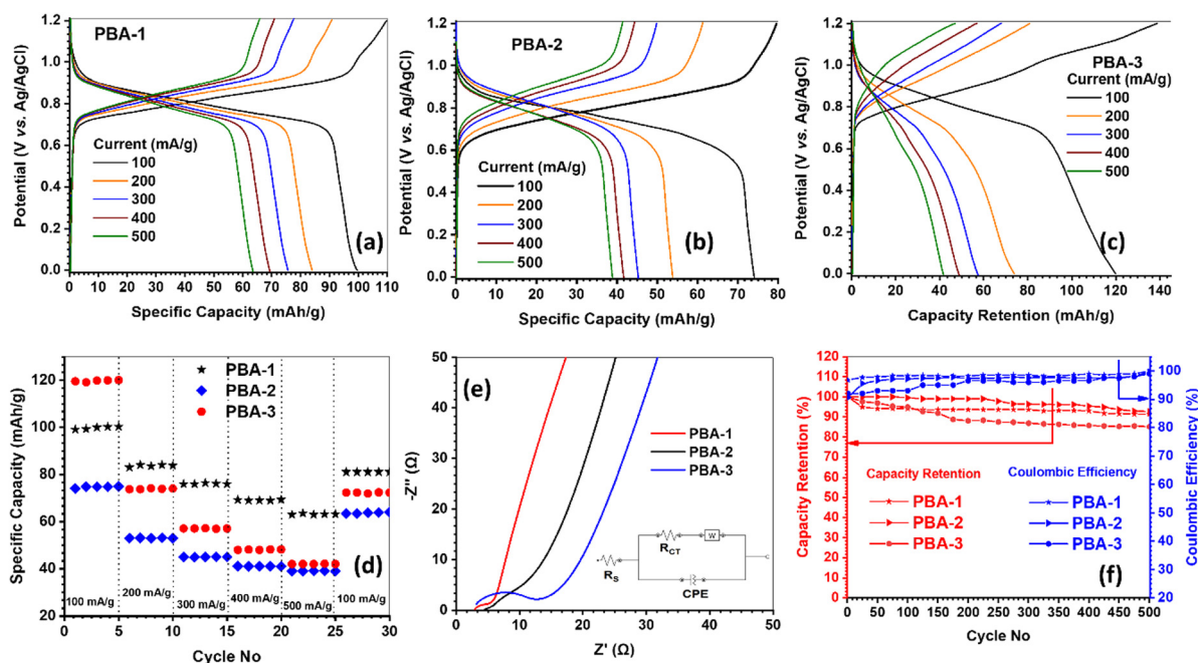


Fig. 5 Electrochemical characterizations of positive active materials: GCD profile of (a) PBA-1, (b) PBA-2, and (c) PBA-3; (d) comparative rate capability data for 30 cycles; (e) comparative Nyquist plots at discharged state (inset: equivalent circuit; W = Warburg impedance and CPE = constant phase element); and (f) comparative cycling life data at 300 mA g<sup>-1</sup> current density.



Table 3 EIS analysis of PBAs before and after the cycling test

	PBA-1		PBA-2		PBA-3	
	Before cycling	After cycling	Before cycling	After cycling	Before cycling	After cycling
$R_s$ ( $\Omega$ )	2.5	2.9	4.3	4.7	3.0	3.0
$R_{ct}$ ( $\Omega$ )	3.2	3.3	5.2	5.4	10	12.9

Jahn–Teller distortion between  $Ni^{2+}$  and NC bonds.<sup>40</sup> Therefore, PBA-1 and PBA-2 demonstrate a better capacity retention than that of PBA-3 after 25 cycles by returning the current at  $100 \text{ mA g}^{-1}$ . Long-term cycling stability was also tested for PBAs under 500 charge/discharge cycles at  $300 \text{ mA g}^{-1}$  current density within 0–1.2 V potential window (Fig. 5(f)), while PBA-1 and PBA-2 demonstrate 91 and 92% capacity retentions respectively. However, PBA-3 depicts 86% retention due to the lower level of Ni content. Nevertheless, the increase in CE values in few initial cycles is mainly due to the formation of passivation layers on the electrode surfaces, which minimize the further parasitic reactions in cells. The EIS analysis was also conducted after cycling experiment to elucidate the increment in resistive components ( $R_s$  and  $R_{ct}$ ) in PBA electrodes. Fig. S7(a)–(c) (ESI<sup>†</sup>) present the Nyquist plots before and after cycling, and the relevant  $R_s$  and  $R_{ct}$  data are summarized in Table 3. It is worthy to mention that there are no appreciable changes in  $R_s$  and  $R_{ct}$  for PBA-1 and PBA-2 electrodes; but PBA-3 depicts 29% increments in  $R_{ct}$  and upshifted  $R_s$  upon cycling experiments. The  $R_{ct}$  enhancement indicates the degradation of the PBA-3 electrode, and the same is reflected in the capacity retention data after 500 cycles. However,

from the high-throughput electrochemical characterizations of PBAs, it is clear that PBA-3 exhibits the best storage capacity and worst rate capability cum durability among the three PBAs. PBA-2 shows minimum capacity, but exhibits moderate rate capability and good durability. The PBA-1 provides moderate capacity, but demonstrates the best rate capability and good cycle life. Based on the above-mentioned comparative data analysis, PBA-1 was selected for the positive electrode in a full-cell assembly.

### Electrochemical characterizations of the negative active material

The variable scan rate CV measurements were carried out within 0.4 to  $-1.0 \text{ V}$  potential window at  $5\text{--}25 \text{ mV s}^{-1}$ , while the voltammograms are shown in Fig. S8(a), (ESI<sup>†</sup>). There are two distinct cathodic/anodic redox peak pairs found at around  $-0.26/0.86$  and  $-0.41/-0.12 \text{ V}$ , which correspond to  $V^{5+}/4^{+}$  and  $V^{4+}/3^{+}$  redox couples respectively.<sup>41,42</sup> The ‘*b*’ values were also calculated at the cathodic and anodic peak positions, whereas the same were reported as 0.36/0.42 (at  $-0.26/0.86 \text{ V}$ ) and 0.40/0.52 (at  $-0.41/-0.12 \text{ V}$ ) (Fig. S8(b), (ESI<sup>†</sup>)). The ‘*b*’ values near 0.5 and even lower signifies the diffusion-controlled faradaic charge storage mechanism, which is associated with the charge/discharge of HVO.<sup>43</sup> Here, no mass transfer limitations are observed in higher scan rates, which is manifested by the non-shifting of peak positions towards large overpotentials at higher rates. The GCD profiles of HVO at variable current loads ( $100\text{--}500 \text{ mA g}^{-1}$ ) within 0.2 to  $-0.7 \text{ V}$  range are shown in Fig. 6(a). The HVO exhibits respective specific capacities of 82 (CE = 91%), 67 (CE = 91%), 61 (CE = 99%), 54 (CE = 98%) and 46

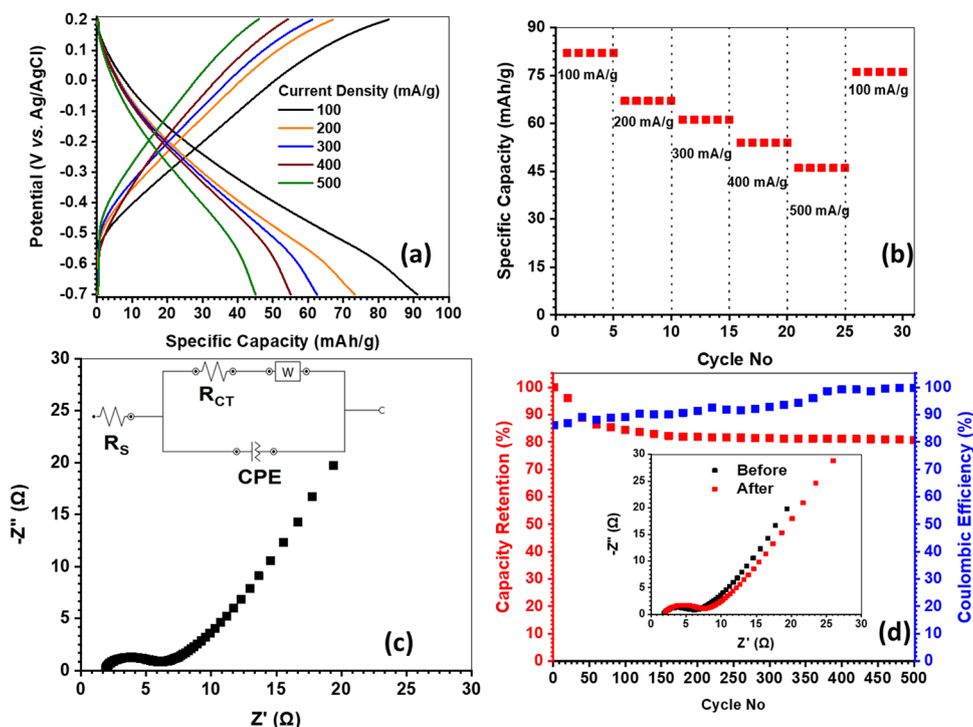


Fig. 6 Electrochemical characterizations of HVO: (a) GCD profiles at different current densities; (b) rate capability data for 30 cycles; (c) Nyquist plot at discharge state (inset: equivalent circuit); and (d) cycling life data at  $300 \text{ mA g}^{-1}$  current density (inset: Nyquist plots before and after cycling).



(CE = 99%) mA h g<sup>-1</sup> at 100, 200, 300, 400 and 500 mA g<sup>-1</sup> current density. Fig. 6(b) depicts the rate capability data, while 43% capacity loss is noted under shifting the current density from 100 to 500 mA g<sup>-1</sup>. Nonetheless, 92% of initial capacity is retained after 25 rate capability cycling, when the current reverts to 100 mA g<sup>-1</sup>. The Nyquist plot of HVO is presented in Fig. 6(c), and the related  $R_s$  and  $R_{ct}$  values are reported as 1.9 and 4.4  $\Omega$ , respectively. The electronic conductivity was estimated from the  $R_{ct}$  value to be  $17 \times 10^{-4}$  S cm<sup>-1</sup>. The cycle life data are presented in Fig. 6(d), where 80% capacity retention is observed under 500 charge/discharge cycles operated at 300 mA g<sup>-1</sup> current load. The Nyquist plots recorded before and after cycling are shown in the inset of Fig. 6(d), while unaltered  $R_s$  and 28% increments of  $R_{ct}$  are noted after cycling. The detailed electrochemical characterizations of HVO provide good insights into the potential negative active material for aqueous Na-ion batteries.

### Performance testing of Na<sub>2</sub>Ni<sub>0.33</sub>Co<sub>0.33</sub>Mn<sub>0.33</sub>Fe(CN)<sub>6</sub>/H<sub>2</sub>V<sub>3</sub>O<sub>8</sub> full cells

The full cell was assembled with positive limited configuration in the CMC-SiO<sub>2</sub>-Na<sub>2</sub>SO<sub>4</sub> hydrogel electrolyte. The gel electrolytes enhance the operating voltage window of aqueous batteries by entrapping the free water molecules into the extensive H-bonded gelling networks.<sup>44</sup> Therefore, the present full cell could be operated within 0–1.8 V window without excessive gassing. Fig. 7(a) demonstrates the GCD profiles of full cells, while 65 (CE = 75%), 47 (CE = 85%), 40 (CE = 89%), 35 (CE = 89%), and 26 (CE = 92%) mA h g<sup>-1</sup> specific capacities are manifested at 100, 200, 300, 400 and 500 mA g<sup>-1</sup> current densities respectively. The rate capability data are presented

in Fig. 7(b), whereas the full cell depicts 60% capacity decay from 100 to 500 mA g<sup>-1</sup>, and 82% capacity retention after 25 cycles upon reverting the current density at 100 mA g<sup>-1</sup>. The Ragone plot is shown in Fig. 7(c), where the energy/power densities of PBA-1//HVO are presented along with the literature data for various PBA based cells.<sup>11–14,45–50</sup> The PBA-1//HVO full cell demonstrates an energy density of 118 W h kg<sup>-1</sup> (at 181 W kg<sup>-1</sup>) and a power density of 780 W kg<sup>-1</sup> (at 46 W h kg<sup>-1</sup>). The effect of temperature on the full cell capacity is demonstrated in Fig. 7(d). The specific capacity is marginally increased from ambient condition to 50 °C, and it is due to the improved ionic conductivity in the gel electrolyte. However, drastically low ionic conductivities are obvious at sub-zero temperatures, while resultantly the capacity values are decreased appreciably. However, based on this temperature-dependent study, one can predict the operating temperature range to be –10 °C to 50 °C. Fig. 7(d) demonstrates the cycle life data of the full cell, which shows 81% capacity retention with 91% CE after 500 cycles at 300 mA g<sup>-1</sup> current density. The inset of Fig. 7(d) shows the Nyquist plots before and after cycling, and an increment in internal resistance is observed upon cycling. To demonstrate the solar energy storage performances, a 5 V/4 mA h rated prototype device was developed by series connection of three 1.8 V cells, where each 1.8 cell was fabricated using two positive and two negative electrodes in parallel connections. The GCD profile of the 5 V prototype is presented in Fig. 8(a), which justifies the capacity rating of the prototype. The EIS analysis was carried out in the 5 V integrated device along with individual cells, and the respective Nyquist plots are depicted in Fig. S9 (ESI†). However, the ESI† study signifies that the internal resistance of 5 V prototype is the summation of the resistances from three

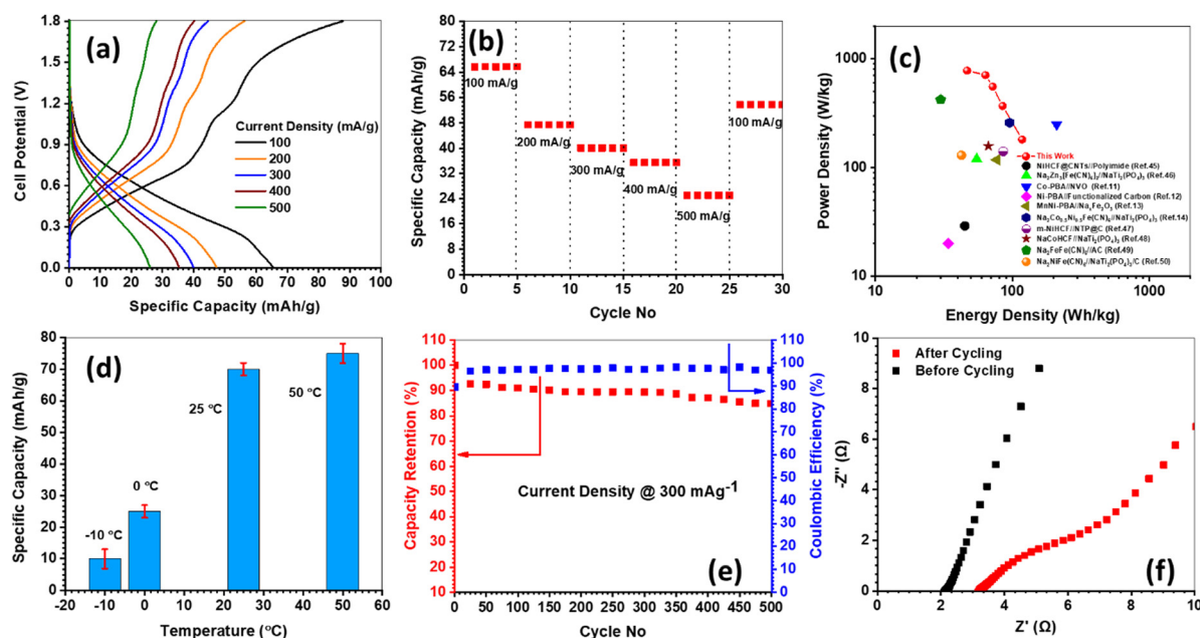


Fig. 7 Electrochemical performance testing data for a 1.8 V PBA-1//HVO full cell: (a) GCD profiles at different current densities; (b) rate capability data for 30 cycles; (c) Ragone plot; (d) effect of temperature on cell capacity; the capacity is estimated at 100 mA g<sup>-1</sup> current density. (e) Cycling life data at 300 mA g<sup>-1</sup> current density. (f) Nyquist plots before and after cycling.



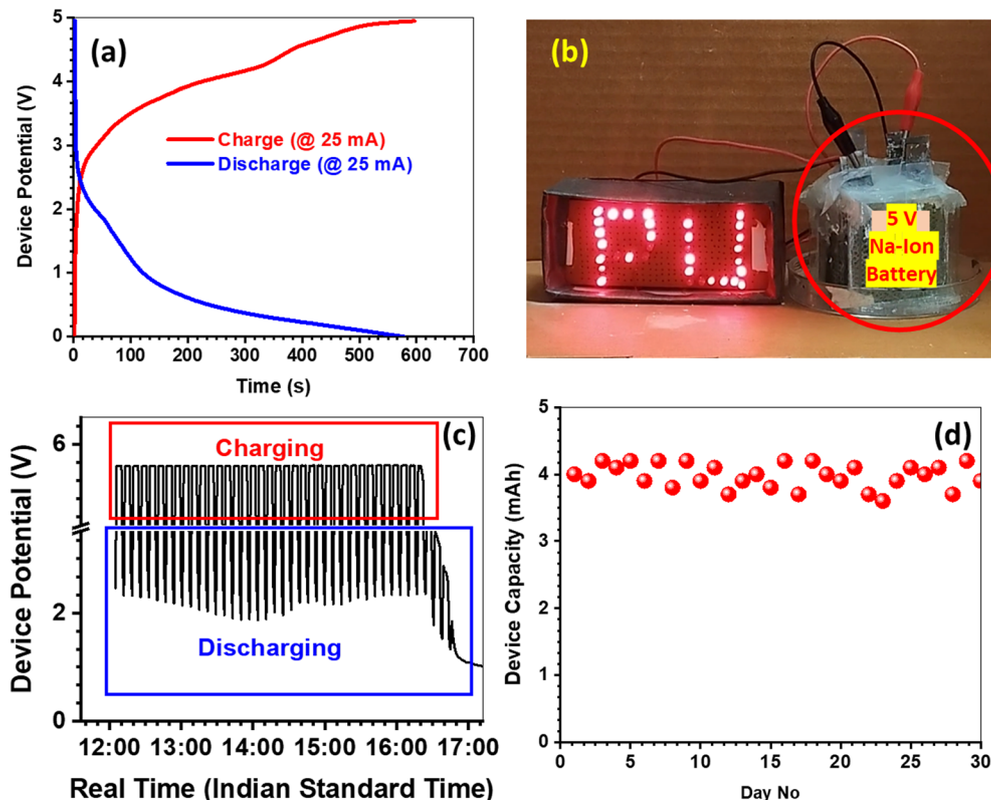


Fig. 8 Performance testing data for a 5 V/4 mA h prototype device: (a) GCD profile at 25 mA current load; (b) illumination of a 3 V/270 mW LED by a 5 V device; (c) voltage profile of the 5 V device under 5 min solar-charging and 2 min LED-discharging for a single day from around 12:00 noon to 05:00 pm; and (d) device capacity measured at 25 mA load after each day solar-charge/LED-discharge experiment during the 30-day field trial.

constituent cells. This result clearly demonstrates that there are no additional resistances associated in the prototype device during the cell integration process. Moreover, the patterns of three Nyquist plots from the three constituent cells are similar, and it signifies that the individual cell is equally performing. Otherwise, the cycle life of the integrated device will be drastically reduced due to the uneven performance distributions among the constituent cells. However, the 5 V/4 mA h rated prototype device is able to illuminate a custom-made 3 V/270 mW LED load, and the photograph is presented in Fig. 8(b). Fig. 8(c) demonstrates the representative profile of one-day charge/discharge data for a 5 V/4 mA h device under 5 min solar-charge and 2 min LED-discharge schedule. The custom-built solar-charging cum LED-discharging module is schematically demonstrated in Fig. S10 (ESI<sup>†</sup>). However, we also carried out a field trial for 30 days, while the capacity of the device was estimated by 25 mA charge/discharge current load after completion of each day solar-charge/LED-discharge experiment. Fig. 8(d) depicts the variation in the device capacity during 30 days' field trial, where 4 mA h average capacity is maintained with legitimate fluctuations.

## Conclusions

In this work, we demonstrated the synthesis and characterizations of three tetra-transition metallic PBAs with multiple redox centres for the positive electrode of aqueous Na-ion batteries.

All three PBAs crystallize in the monoclinic phase and demonstrate almost identical crystal structures. The elemental composition determination along with the quantification of crystalline water was carried out by EDX spectroscopy and TGA, which provide the chemical structures of the PBAs. All the PBAs were electrochemically characterized by CV and EIS studies, and their performance was investigated through GCD profiles, rate capability and cycle life. The CV analysis indicates predominating faradaic charge storage mechanisms in PBAs and the relevant charge transfer resistances were estimated by EIS. The Na-ion diffusion barriers are almost similar in all three PBAs, due to the resemblance of the crystal structure; but the electrochemical performances markedly deviate from each other because of the electronic conductivity of the materials. The EIS study demonstrated solid-state electronic conductivities of PBA-1, PBA-2 and PBA-3 as  $23 \times 10^{-4}$ ,  $15 \times 10^{-4}$  and  $7.4 \times 10^{-4}$  S cm<sup>-1</sup>, respectively. PBA-3 showed the highest capacity (120 mA h g<sup>-1</sup>) compared to those of PBA-1 (100 mA h g<sup>-1</sup>) and PBA-2 (75 mA h g<sup>-1</sup>) at 100 mA g<sup>-1</sup> current load, while it demonstrated poor rate capability (65% capacity loss from 100 to 500 mA g<sup>-1</sup>) and durability (86% capacity retention after 500 cycles at 300 mA g<sup>-1</sup>). Besides, PBA-1 and PBA-2 depicted better performances on rate capability (32 and 47%, respectively) and durability (91 and 92%, respectively). However, this performance matrix identified PBA-1 as the optimally suitable candidate for the positive electrode of aqueous Na-ion full cells.



In this work, we have synthesized HVO as the negative active material, and the same was thoroughly characterized by physicochemical and electrochemical means. Thereafter, we have assembled a 1.8 V PBA-1//HVO full cell in a hydrogel electrolyte, and this cell exhibited a specific capacity of 65 mA h g<sup>-1</sup> (at 100 mA g<sup>-1</sup>), an energy density of 118 W h kg<sup>-1</sup> (at 181 W kg<sup>-1</sup>), a power density of 780 W kg<sup>-1</sup> (at 46 W h kg<sup>-1</sup>) and a good cycle life (81% capacity retention after 500 cycles at 300 mA g<sup>-1</sup>). Finally, we have validated the integration of a 5 V/4 mA h prototype device for the practical demonstration of solar energy storage applications. It is noteworthy that the combination of tetra-transition metallic PBA as the positive electrode and HVO as the negative electrode in Na-ion-based energy storage devices is unique, and this unique combination is truly prospective towards industrial transformation.

## Conflicts of interest

There are no conflicts to declare.

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