



Scalable Synthesis of Pre-Intercalated Manganese(III/IV) Oxide Nanostructures for Supercapacitor Electrodes: Electrochemical Comparison of Birnessite and Cryptomelane Products

Daniel R. Jones,^{*[a]} Haytham E. M. Hussein,^[b] Eleri A. Worsley,^[a] Sajad Kiani,^[a] Kittiwat Kamlungsua,^[c] Thomas M. Fone,^[a] Christopher O. Phillips,^[a] and Davide Deganello^[a]

Manganese(III/IV) oxide is a promising pseudocapacitive material for supercapacitor electrodes due to favorable attributes such as its chemical resilience, high earth abundance and low specific cost. Herein, the morphological, compositional and electrochemical characteristics of co-precipitated manganese(III/IV) oxide products, each described by the general formula Na_xK_yMnO_z, are investigated to establish how these properties are influenced by synthesis conditions. Na_xK_yMnO_z growths in low-temperature (< 100 °C) basic and acidic environments are shown to promote the formation of turbostratic birnessite and cryptomelane phases, respectively, with the latter polymorph containing a relatively low concentration of interstitial Na⁺ and K⁺ cations. It is demonstrated that K⁺ preinsertion during synthesis yields lower initial charge-transfer

Introduction

If inherently intermittent renewable energy technologies are to offer a practical alternative to conventional heat engines, it is critically important that efficient, cost-effective and sustainable energy storage solutions are developed to decouple supply and demand profiles. A promising strategy is to pair the competitive energy densities of batteries with the superior cycling lifetimes and power densities of supercapacitors,^[1] employing an appropriate maximum power point tracking (MPPT) system to

- [a] Dr. D. R. Jones, E. A. Worsley, Dr. S. Kiani, T. M. Fone, Dr. C. O. Phillips, Prof. D. Deganello Faculty of Science and Engineering Swansea University Swansea, SA1 8EN (UK) E-mail: d.r.jones@swansea.ac.uk [b] Dr. H. E. M. Hussein School of Natural and Computing Sciences University of Aberdeen Aberdeen, AB24 3UE (UK) E-mail: h.hussein@abdn.ac.uk [c] Dr. K. Kamlungsua Enserv Power Co. Ltd., Enserv Innovation Centre Bangkok, 10800 (Thailand) E-mail: kittiwat@enserv.co.th Supporting information for this article is available on the WWW under https://doi.org/10.1002/celc.202300210
- © 2023 The Authors. ChemElectroChem published by Wiley-VCH GmbH. This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

resistances than equivalent Na⁺ intercalation, and that this parameter correlates strongly with storage performance. Accordingly, Na-mediated storage initially delivers inferior specific capacitances and Coulombic efficiencies than K-based mechanisms, but K⁺ intercalation/deintercalation causes faster capacitance decay during prolonged galvanostatic cycling. Furthermore, whilst crystallographic phase is shown to have a weaker effect on Na_xK_yMnO_z storage properties than the choice of intercalating guest cations, cryptomelane electrodes are more susceptible to cycling-induced capacitance and efficiency losses than their birnessite counterparts. In combination, these insights provide an instructive foundation for the optimization of Na_xK_yMnO_z in high-power storage applications.

optimize energy exchange between these complementary devices.^[2] Mixed-valence manganese(III/IV) oxides (MnO₂, where z ranges between 1.5 and 2.0) have emerged as foremost candidates for the electroactive material in supercapacitor cathodes due to their low toxicity, minimal cost,^[3] ready availability (manganese appears at an average concentration of ca. 1,000 ppm in the Earth's crust^[4]), wide electrochemical stability window (ca. 1.0 V in aqueous electrolytes^[5]), and high maximum theoretical specific capacitance (ca. 1,370 F g⁻¹, based on the reduction of Mn⁴⁺ ions by a single electron^[6]). Charge storage in MnO, is predominantly mediated by the pseudocapacitive intercalation of cations from the surrounding electrolyte in each discharge phase, followed by deintercalation of these guest ions during charging; in the case of a monovalent cation (A⁺), this cyclical process is described by the reversible reaction scheme expressed in Equation 1,^{[7}]

$$MnO_z + A^+ + e^- \leftrightarrow MnO_z A$$
 (1)

where Mn^{4+} ions in MnO_z are reduced to the 3+ oxidation state by participating electrons (e⁻). Accordingly, effective storage performance is contingent on the accessibility of Mn^{4+} sites, and hence it is essential that cations can travel unimpeded throughout a charge-discharge cycle.

Of the various crystallographic phases of MnO_{zr} cryptomelane (α -MnO_z) and birnessite (δ -MnO_z) polymorphs are deemed two of the most promising candidates for storage applications because they possess characteristically wide channels for intercalation/deintercalation:^[8,9] α -MnO_z consists of ca. 4.6 Ådiameter one-dimensional tunnels enclosed between cornersharing MnO₆ octahedra, whereas δ -MnO₇ comprises twodimensional layers of MnO₆ octahedra separated by ca. 7.0 Å. In both cases, these spacious passages result in a theoretical specific integral capacitance, C_{int}, that compares favorably with the corresponding storage capabilities of alternative phases such as tetragonal β -MnO₂ and orthorhombic γ -MnO₂.^[8] In order to quantitatively assess α -MnO₂ and δ -MnO₂ products reported in the literature, it is instructive to collate the associated C_{int} values estimated from galvanostatic discharge measurements at a consistent current density. For this purpose, Figure 1 relates the discharge capacitances of selected α -MnO₂ and δ -MnO₂ materials to the temperature at which they were formulated, T_{synth} . The plotted data correspond to a gravimetric current density of 10 A per gram of electroactive material, with synthesis methods including the reduction of MnO₄⁻ anions, [10-18] the oxidation of Mn^{2+} cations, [19-21] and the coprecipitation of Mn(II) and Mn(VII) sources^[22-28] in acidic or alkaline conditions according to the multi-step reactions described in Equation 2,

$$(23 - 10z)Mn^{2+} + 2MnO_4^- + 2(11 - 5z)H_2O \rightarrow 5(2z - 3)MnO_2 + 10(2 - z)Mn_2O_3 + 4(11 - 5z)H^+$$
(2)

and Equation 3,

$$(23 - 10z)Mn^{2+} + 2MnO_4^- + 4(11 - 5z)OH^- \rightarrow 5(2z - 3)MnO_2 + 10(2 - z)Mn_2O_3 + 2(11 - 5z)H_2O$$
(3)

respectively.^[22] Whilst most of the products represented in Figure 1 were prepared at temperatures in excess of 100 °C, the weak correlation between C_{int} and T_{synth} suggests that lower synthesis temperatures do not necessarily yield inferior storage performance. Moreover, reducing T_{synth} below the boiling point of water provides a way to minimize the energy cost of MnO_z production whilst maximizing the scalability of the process.



Figure 1. Literature values^[10–28] of the specific integral capacitance, C_{intr} estimated from galvanostatic discharge measurements at 10 Ag⁻¹ and normalized with respect to the mass of electroactive material, plotted as a function of synthesis temperature, T_{synthr} for birnessite (δ -MnO_z) and cryptomelane (α -MnO_z) products.

Alongside specific capacitance and the electrochemical stability window, another key property of any supercapacitor electrode is its capacitance retention over many thousands of galvanostatic charge-discharge cycles.^[29] In the case of aqueous MnO₂ systems, capacitance fading is known to depend strongly on the ability of the material to recover from repeated sequences of contraction and expansion caused by electrostatic attractions between the negatively charged MnO₆ octahedra and the guest cations.^[30,31] Alternatively, deterioration may result from the disproportionation of Mn³⁺ ions at sufficiently low potentials (typically less than ca. -0.1 V vs Ag|AgCl (3 M KCI)),^[30] generating Mn(II) products that can subsequently dissolve in the aqueous electrolyte.^[29,30] Mass loss through dissolution can be circumvented by selecting an appropriate voltage range for electrochemical testing, whilst pre-inserting cations into the bulk material during synthesis also suppresses Mn reduction and enhances structural stability by reducing the propensity of the MnO₆ octahedra to shift during cycling.^[15,29,32-36] In addition, the form of the MnO₂ polymorph is known to play a critical role in capacitance decay; for instance, it has been shown that Jahn-Teller distortions of MnO₆ octahedra cause less disruption to the two-dimensional channels of δ -MnO_z than the corresponding ion pathways of α -MnO...[37]

Although numerous investigations into the electrochemical behaviors of MnO, materials have addressed how storage performance depends on factors such as crystallographic phase^[22,37-44] and the nature of intercalating cations,^[11,14,25,31,45-54] the effects of quest cation type and MnO₂ structure are rarely explored simultaneously within a single self-consistent study. Furthermore, researchers have hitherto adhered to a convention of measuring storage parameters as functions of current density and potential scan rate at just one stage during testing, either with the electrode in its as-prepared state or after a designated number of activation cycles.^[11] The variation of capacitance retention is typically evaluated at a single value of current density during prolonged galvanostatic cycling, but, to the present authors' knowledge, all previous works have neglected to consider the evolution of established pre-cycling relationships over the full range of current density and potential scan rate values. For example, Xiong et al.^[31] employed X-ray diffractometry (XRD) to link cyclical changes in the interlayer spacing of δ -MnO₇ to the intercalation/deintercalation of various cations, and it was shown that reducing the amplitude of lattice contraction and expansion yielded an improvement in capacitance retention over 5,000 galvanostatic cycles at 5 A g⁻¹. In turn, the authors reasoned that in comparison to mechanisms involving Li⁺, Na⁺ or Mg²⁺ ions, the relatively low structural deformation caused by K⁺ exchanges translated to superior cycling stability. However, prior to the cycling procedure, it was demonstrated that storage performance depended strongly on current density, with K-mediated storage delivering lower capacitance than corresponding Li⁺, Na⁺ and Mq^{2+} processes below ca. 1 Aq^{-1} , despite offering the highest capacitance when current density exceeded ca. 2 A g⁻¹. As the effects of current density on capacitance and Coulombic efficiency were not explored in the aftermath of prolonged cycling, it is unclear whether the inferred relationships between guest ion type and capacitance fading at 5 Ag^{-1} are applicable over the whole current density range.

The present treatment avoids the identified shortcomings of previous research by quantifying the storage properties of α - MnO_z and δ - MnO_z electrodes, each described by the general formula Na_xK_yMnO_z, across various operating regimes, both before and after subjecting the electrodes to many galvanostatic charge-discharge cycles. In contrast to existing methodologies, this rigorous and systematic approach enables the short- and long-term effects of cation intercalation/deintercalation mechanisms to be elucidated in enhanced detail over the full duration of electrochemical testing. Alongside these electrode measurements, a variety of other characterization techniques were implemented to explore the influence of synthesis pH and guest cation type on the morphology, composition and crystallography of the resulting Na_xK_yMnO_z product. By examining these relationships in greater detail than existing investigations, the present research serves as an instructive resource for the optimization of mixed-valence

 $Na_xK_yMnO_z$, providing a scalable basis to develop efficient, stable and charge-dense pseudocapacitive cathodes for use in high-power storage applications.

Results and Discussion

Phase Identification

Through the co-precipitation of Mn^{2+} and MnO_4^{-} ions, $Na_xK_yMnO_z$ samples were prepared under either basic (B) or acidic (A) conditions; for concision, materials produced in the presence of $Na_2SO_{4(aq)}$ are henceforth assigned the prefix "Na", whilst the prefix "K" is used to reference products obtained when $K_2SO_{4(aq)}$ was instead used as a synthesis precursor. X-ray diffraction (XRD) measurements acquired from the four $Na_xK_yMnO_z$ products (Figure 2) indicate that the choice of cation in the sulfate salt had a seemingly negligible effect on crystallographic structure, although the predominant polymorph was altered by changing the pH of the synthesis solution: the XRD



Figure 2. Transmission-mode powder X-ray diffractograms acquired from samples of Na-B (a), K-B (b), Na-A (c) and K-A (d); crystallographic peaks associated with turbostratic birnessite δ -MnO₂ and cryptomelane α -MnO₂ are indexed to reference patterns provided by JCPDS cards 23-1046 and 20-0908, respectively.



patterns of Na-B and K-B (Figure 2a and b, respectively) are consistent with the lamellar structure of birnessite MnO_z (δ-MnO₂), as indexed by JCPDS card 23-1046,^[55] whereas the diffractograms of samples Na-A and K-A (Figure 2c and d) accord with tetragonal cryptomelane MnO_{7} (α -MnO₇), corresponding to JCPDS card 20-0908.^[56] A notable feature of the birnessite systems is the apparent absence of the (002) peak, which characteristically appears at a 2θ value of ca. 12.5°; this omission suggests that Na-B and K-B did not adopt the archetypal form of pristine δ -MnO₂. Indeed, similar suppression of the (002) peak has been witnessed in the X-ray diffractograms of other Na- and K-intercalated δ -MnO_z products, ^[14,25,57,58] and the phenomenon may be attributed to poor alignment between lattice planes due to turbostratic stacking of the δ -MnO, nanosheets.^[59] Conversely, the Na-A and K-A diffractograms both contain sharp peaks matching the full reference pattern of α -MnO₂, which shows that as well as promoting formation of the cryptomelane phase, acidic conditions also enhanced lattice ordering.

The relationship between Na_vK_vMnO_z crystallography and synthesis pH has been considered in previous research,^[22,38,60] wherein it has been demonstrated that the principal polymorph depends on the relative concentrations of hydroxonium and alkali metal cations: cryptomelane structures preferentially form when there is a sufficient abundance of hydroxonium ions to competitively intercalate into the one-dimensional channels, whilst lower hydroxonium availability causes MnO₆ octahedra to instead assemble as two-dimensional layers. As the relatively narrow tunnel configuration of α -MnO, renders it less vulnerable than δ -MnO_z to infiltration by electrolyte species, the formation of well-ordered crystallites in δ -MnO₂ is more strongly inhibited by pre-insertion processes. Consequently, literature comparisons between the X-ray diffractograms of α -MnO, and turbostratic δ -MnO_z^[8,22,38,61,62] typically reveal that a much higher degree of crystallinity exists in the cryptomelane case.

By examining each sample using a combination of fieldemission scanning electron microscopy (SEM), transmission electron microscopy (TEM) and scanning transmission electron microscopy (STEM), the relationship between lattice structure and synthesis pH was further explored (Figure 3). Images of the samples prepared under basic conditions, Na-B (Figure 3a-c) and K-B (Figure 3d-f), show that they consisted of microscale particles with nanoscopic surface features, which are representative of agglomerated δ -MnO₇ nanoflakes^[8] exemplified elsewhere in the literature.^[13,25,61] In TEM images of Na-B and K-B (Figure 3c and f, respectively), no lattice planes were discernible amongst the turbostratic δ -MnO₂ matrix; this finding is consistent with the low signal-to-noise ratio of the corresponding Xray diffractograms (Figure 2a and b, respectively) and the absence of (002) peaks in these measurements. By contrast, SEM and TEM images acquired from the products of acidic synthesis conditions, Na-A (Figure 3g-i) and K-A (Figure 3j-I), indicate that they possessed a high degree of crystallinity, evidenced by an abundance of nanorods consisting of welldefined lattice planes and atomic columns (Figure 3i and I, and Figure S1 of the Supporting Information). Lattice d-spacing measurements obtained from the high-magnification TEM images of Na-A (Figure 3i, and Figure S1a and b) and K-A (Figure 3l, and Figure S1c and d) may be indexed to characteristic peaks in the XRD patterns of Na-A (Figure 2c) and K-A (Figure 2d), with examples of (110), (200) and (310) plane arrangements identified in the images. Nevertheless, it should be noted that both Na-A and K-A also contained high quantities of seemingly amorphous particles, akin to the microstructures of Na-B and K-B. It is thus apparent that whilst the acidic growth environments of Na-A and K-A encouraged nanorod formation, this process was accompanied by alternative co-precipitation mechanisms that culminated in a more disordered product.

Composition Analysis

The chemical compositions of the four Na_xK_yMnO_z products were investigated by X-ray photoelectron spectroscopy (XPS); Mn 2p, O 1s, C 1s, Na 1s, and K 2p spectra were acquired from four separate locations on the surface of each pelletized sample. XPS data obtained from one region of the Na-A pellet are provided in Figure 4, whilst the Supporting Information displays corresponding measurements from Na-B (Figure S2), K-B (Figure S3), and K-A (Figure S4). To illustrate the magnitude of composition variations across each pellet surface, the Supporting Information also includes overlaid presentations of the survey spectra (Figure S5) and the Mn 2p (Figure S6), O 1s (Figure S7), C 1s and K 2p (Figure S8), and Na 1s (Figure S9) measurements from the four sample locations, normalizing the measured count rate with respect to the relevant Mn $2p_{3/2}$ peak maximum in each case; in all of these comparisons, it is clear that the shape and intensity of the core peaks varied little between the scanned areas. By deconvolving the core spectra into constituent components associated with distinct electronic environments, it is possible to estimate the relative proportions of chemical species; the mean and standard deviation of atomic ratios from the four pellet locations were calculated in order to evaluate the significance of chemical differences between samples.

In general, MnO₆ octahedra generated by the co-precipitation of Mn^{2+} and MnO_4^{-} ions are composed of Mn(III) and Mn(IV) oxidation states; consequently, the Mn 2p spectra of Na-B (Figure S2a), K-B (Figure S3a), Na-A (Figure 4a) and K-A (Figure S4a) can be resolved into two pairs of doublets, with the 2p_{3/2} components of the Mn(III) and Mn(IV) species centred at characteristic binding energies of ca. 640.6-641.0 eV and ca. 642.1-642.3 eV, respectively.^[14,15,63-68] From the relative areas of these doublets it is evident that Na-A and K-A comprised a higher proportion of Mn³⁺ environments than Na-B and K-B, implying that basic synthesis conditions suppressed the reduction of Mn⁴⁺ ions; this deduction concurs with previous research on co-precipitated cryptomelane and turbostratic birnessite MnO₇, which has demonstrated that α -MnO₇ inherently possesses a lower average oxidation state than δ -MnO₇.^[62,69] By comparing the relative areas of the Mn(III) and Mn(IV) components from each set of four sample locations, Mn^{3+}/Mn^{4+} ratios of 0.07 \pm 0.03 and 0.42 \pm 0.05 are obtained for Na-B and K-B, respectively, which are considerably lower





Figure 3. Field-emission SEM (a, b, d, e, g, h, j and k) and TEM/STEM (c, f, i and I) images acquired from Na-B (a–c), K-B (d–f), Na-A (g–i) and K-A (j–I); STEM images corresponding to Na-B (c) and K-B (f) suggest that these samples consisted of randomly orientated agglomerates of birnessite (δ -MnO₂) nanoflakes, whereas distinctive collections of well-aligned crystallographic lattice planes corresponding to the cryptomelane (α -MnO₂) phase are discernible within the TEM images of Na-A (i, and Figure S1a and b of the Supporting Information) and K-A (I, and Figure S1c and d of the Supporting Information), as indicated in the images. In the lowest magnification SEM images (a, d, g and j), the regions surveyed at higher magnification (b, e, h and k) are marked by dashed red boxes.



Figure 4. X-ray photoelectron measurements from Na-A, including the Mn 2p (a), O 1s (b), K 2p and C1s (c), and Na 1s (d) core peaks. Deconvolution of these spectra was achieved by fitting Gaussian-Lorentzian (O 1s, K 2p, C 1s and Na 1s) or asymmetric Lorentzian (Mn 2p) functions to the data following subtraction of a Shirley-type background function. For clarity, components associated with separate electronic environments are represented by different colors, whilst the 2p_{3/2} and 2p_{1/2} contributions of a given 2p doublet are assigned the same color. The proposed species responsible for the plotted spectral components are annotated accordingly, and the peak position of each component is indicated by a vertical dashed line.

than corresponding values of 0.97 ± 0.05 and 0.88 ± 0.05 yielded, respectively, by the Mn 2p spectra of Na-A and K-A.

The Na/Mn and K/Mn atomic ratios, respectively denoted x and y in the general formula Na_xK_yMnO_z, may be similarly estimated from the XPS measurements. K 2p spectra recorded from Na-B (Figure S2c), K-B (Figure S3c), Na-A (Figure 4c) and K-A (Figure S4c) all exhibit a doublet that is well-represented by a single electronic environment, located at binding energies in the vicinity of C 1s components from the adventitious organic surface layer.^[70] Similarly, the Na 1s spectra of Na-B (Figure S2d) and Na-A (Figure 4d) consist of one symmetric peak, indicative of a single Na species. From the enclosed areas of the Na 1s and Mn 2p spectra, a mean Na/Mn atomic ratio of 0.30 ± 0.03 is obtained for Na-B, which compares with a corresponding value of 0.044 ± 0.006 from the Na-A data. No Na 1s signal was observed in the case of K-B (Figure S3d) or K-A (Figure S4d), as no Na-containing precursors were involved during formulation of these materials. It is also apparent that basic synthesis conditions facilitated intercalation of K⁺ ions, as K-B possessed a mean K/Mn atomic ratio of 0.39 ± 0.02 , which is three times greater than a corresponding estimate of 0.130 ± 0.001 yielded by the K 2p and Mn 2p doublets of K-A. As anticipated from the low concentration of K-based precursors employed in their syntheses, Na-B and Na-A comprised far lower quantities of preinserted K⁺ ions: the relative magnitudes of associated K 2p and Mn 2p spectra are consistent with mean K/Mn atomic ratios of 0.040 ± 0.002 and 0.043 ± 0.001 for Na-B and Na-A, respectively. A qualitative rationale for the differing levels of preintercalation within the acidic and basic samples may be developed by recalling the crystallographic structures of the α -MnO_z and δ -MnO_z polymorphs: the planar configuration of MnO₆ octahedra in δ -MnO_z provides wide interlayer passages for ionic transport, in contrast to the comparatively narrow onedimensional channels of α -MnO_z.^[8]

To determine the mean O/Mn atomic ratio within each sample, denoted z in the general formula $Na_xK_yMnO_z$, it is necessary to analyse the contributions of Mn-containing species to corresponding O 1s spectra. Due to the high surface-



sensitivity of the XPS technique, each O 1s spectrum contains components associated with C–O and C=O environments from the thin adventitious organic layer that enveloped the particles, in addition to interstitial water.^[71-73] These species were accompanied by O²⁻ anions situated in the bulk MnO₇ lattice and Mn-bound hydroxide groups with characteristic O 1s peaks centred at binding energies of ca. 529.6-530.0 eV and ca. 531.1-531.4 eV, respectively,^[64,65,72,73] which must be summed to determine the overall quantity of O that was affiliated with the octahedral MnO₆ units. By comparing these combined contributions to the total atomic percentages of Mn, respective z estimates of 1.92 ± 0.02 and 1.91 ± 0.04 are computed from the O 1s spectra of Na-B (Figure S2b) and K-B (Figure S2c), which far exceed corresponding values of 1.604 ± 0.008 and 1.639 ± 0.008 yielded by the O 1s spectra of Na-A (Figure 4b) and K-A (Figure S4b), respectively.

A summary of key results from the XRD and XPS characterizations is provided in Table 1, which collates the identified crystallographic phases of Na-B, K-B, Na-A and K-A alongside the estimated mean atomic ratios Na/Mn (x), K/Mn (y) and O/ Mn (z). Also included in Table 1 are the Brunauer-Emmett-Teller (BET) specific surface areas of the four products estimated via the Supporting Information (Figure S10), which depicts the measured Type IV isotherms^[74,75] of Na-B (Figure S10a), K-B (Figure S10c), Na-A (Figure S10e) and K-A (Figure S10g) in conjunction with BET plots (Figure S10b, d, f and h, respectively) constructed from the nitrogen adsorption curves. As observed in previous $Na_x K_v MnO_z$ research,^[14,38,65,68] each pair of adsorption and desorption isotherms is separated by a H3 hysteresis loop,^[74,75] which is indicative of condensation at mesopores present within the material. The BET specific surface areas of Na-A and K-A were found to be an order of magnitude greater than those of their basic counterparts, reflecting the nanoscopic dimensions of the α -MnO₂ crystallites. The high abundance of mesoporous adsorption sites in Na-A and K-A is also evident from their hysteresis loops, which are appreciably more pronounced than the corresponding hysteresis behaviors of Na-B and K-B.

Electrochemical Performance Testing

To assess the electrochemical stability of the $Na_xK_yMnO_z$ products, and to explore the nature of charge storage mechanisms in the presence of alkali metal cations, $Na_xK_yMnO_z$ electrodes were subjected to cyclic voltammetry (CV) sweeps at potential scan rates, ν , in the ranges 10–100 mVs⁻¹ (Figure 5) and 100–1,000 mV s⁻¹ (Figure S11 of the Supporting Information). Prior to galvanostatic charge-discharge (GCD) cycling, the as-prepared electrodes of Na-B (Figure 5a and Figure S11a), K-B (Figure 5c and Figure S11c), Na-A (Figure 5e and Figure S11e) and K-A (Figure 5g and Figure S11g) all exhibited nearrectangular relationships between potential, E, and current density, J, at scan rates ranging from 10 mV s^{-1} to 100 mV s^{-1} (Figure 5a, c, e and g). However, after performing 2,000 successive galvanostatic cycles at 28.6 Ag⁻¹ between 0.0 V and 1.0 V vs Ag | AgCl (3 M KCl), the corresponding cyclic voltammograms (Figure 5b, d, f and h) diverged significantly from the rectangular form of an ideal supercapacitor,^[76] especially at the upper limit of the scanned potential window. A notable aspect of these post-cycling CV measurements is the emergence of redox peaks at potentials close to 0.5 V vs Ag AgCl (3 M KCl), which are especially evident in the cases of Na-B (Figure 5b) and Na-A (Figure 5f); similar redox features observed in the voltammograms of other Na_xK_yMnO_z electrodes have been assigned to the Mn⁴⁺/Mn³⁺ redox couple, accompanied by reversible alkali metal cation intercalation/ deintercalation.[15,31,34,44,77-79] Increasing the potential scan rate to 100–1,000 mV s⁻¹ caused the J vs ν relationships of Na-B (Figure S11a and b), K-B (Figure S11c and d), Na-A (Figure S11e and f) and K-A (Figure S11g and h) to deviate from the linear J vs ν trend witnessed during slower CV scans, both in the asprepared state (Figure S11a, c, e and g) and after prolonged galvanostatic cycling (Figure S11b, d, f and h).

Further analysis of the pre- and post-cycling voltammograms was achieved via application of the Dunn formula (Equation 4), $^{[80]}$

$$J = k_{\text{surf}}\nu + k_{\text{diff}}\nu^{1/2} \tag{4}$$

where the term containing the constant coefficient k_{surf} represents the linear J vs ν trend yielded by fast pseudocapacitive processes at the electrode surface, and the term possessing a square-root dependence on ν , with a constant coefficient k_{diffr} , corresponds to diffusion-controlled ion exchange mechanisms. In conjunction with Equation 4, it is possible to assess the relative contributions of surface- and diffusion-mediated storage interactions by plotting the quotient $|J|/\nu^{1/2}$ as a function of $\nu^{1/2}$ (Figure S12 of the Supporting Information), as the coefficients k_{surf} and k_{diff} are given, respectively, by the gradient and *y*-intercept of the relationship between these quantities. Applying this approach to pre- and post-cycling *J* measurements from the CV data of Na-B (Figure S12a), K-B (Figure S12b), Na-A (Figure S12c) and K-A (Figure S12d) at 0.5 V vs Ag|AgCl

Table 1. Compiled mean atomic ratio estimates determined from the deconvolved XPS spectra of Na-B (Figure S2), K-B (Figure S3), Na-A (Figure 4) and K-A (Figure S4), alongside the principal crystallographic phases identified from XRD measurements (Figure 2) and specific surface areas calculated through BET analysis of nitrogen adsorption isotherms (Figure S10).

Na _x K _y MnO _z sample	Mean atomic ratios Na/Mn (x)	K/Mn (y)	O/Mn (<i>z</i>)	Principal phase	BET specific surface area [m ² g ⁻¹]
Na-B K-B Na-A K-A	$\begin{array}{c} 0.30 \pm 0.03 \\ 0.000 \\ 0.044 \pm 0.006 \\ 0.000 \end{array}$	$\begin{array}{c} 0.040 \pm 0.002 \\ 0.39 \pm 0.02 \\ 0.043 \pm 0.001 \\ 0.130 \pm 0.001 \end{array}$	$\begin{array}{c} 1.92 \pm 0.02 \\ 1.91 \pm 0.04 \\ 1.604 \pm 0.008 \\ 1.639 \pm 0.008 \end{array}$	Birnessite Birnessite Cryptomelane Cryptomelane	21.4 29.0 181.0 151.0

Research Article doi.org/10.1002/celc.202300210



Figure 5. Cyclic voltammograms measured at potential scan rates, v, between 10 mV s⁻¹ and 100 mV s⁻¹, both before (a, c, e and g) and after (b, d, f and h) subjecting the Na-B (a and b), K-B (c and d), Na-A (e and f) and K-A (g and h) electrodes to 2,000 GCD cycles at a current density of 28.6 A g⁻¹.

(3 M KCl), it is clear that the displayed relationships adhere to the Dunn equation up to ν values of ca. 100 mV s⁻¹, whereupon they diverge from the predicted linearity between $|J|/\nu^{1/2}$ and $\nu^{1/2}$. Na-B and Na-A were most weakly affected by the cycling procedure, whilst more marked mechanistic changes in the K-based products, K-B and K-A, are evident from the greater discrepancies between their pre- and post-cycling $|J(0.5 \text{ V})|/\nu^{1/2}$ vs $\nu^{1/2}$ trends.

Repeating Dunn's protocol over the entirety of the cyclic voltammogram at a given value of ν enables estimation of the surface-mediated storage contributions. As shown in the Supporting Information (Figure S13), these mechanisms accounted for most of the transferred power within the central plateau of CV measurements acquired at 50 mV s⁻¹, but diffusion-controlled processes delivered an increasing proportion of the total current density close to the extrema of the spanned potential range. In agreement with the $|J(0.5 \text{ V})|/\nu^{1/2}$ vs $v^{1/2}$ relationships depicted in Figure S12, prolonged galvanostatic cycling had a seemingly weak effect on surface contributions in Na-B (Figure S13a and b) and Na-A (Figure S13e and f): integration of the pre-cycling surface current curves reveals that pseudocapacitive charge exchange delivered 88% (Figure S13a) and 83% (Figure S13e), respectively, of total current in these systems, compared with 91% (Figure S13b) and 86% (Figure S13f), respectively, in the post-cycling case. Conversely, precycling surface contributions of 90% in both K-B (Figure S13c) and K-A (Figure S13g) decreased to post-cycling values of 87% (Figure S13d) and 79% (Figure S13h), respectively.

The specific integral capacitances, C_{int}, of the Na_xK_yMnO_z electrodes were quantified for charging and discharging between 0.0 V and 1.0 V vs Ag AgCI (3 M KCI) by considering GCD measurements acquired over the current density ranges 1.43–14.3 A g^{-1} (Figure 6) and 14.3–143 A g^{-1} (Figure S14 of the Supporting Information), both before (Figure 6a, c, e and g, and Figure S14a, c, e and g) and after (Figure 6b, d, f and h, and Figure S14b, d, f and h) conducting 2,000 galvanostatic cycles at 28.6 Ag^{-1} . To map the evolution of GCD behaviors, the Supporting Information (Figure S15) also provides overlaid depictions of the 2,000 cycles performed by each system (Figure S15a, c, e and g), alongside the measured variations in discharge capacitance retention (Figure S15b, d, f and h). It was observed in all cases that the durations of the charge and discharge phases diminished gradually during prolonged cycling, equilibrating at a decreased value of C_{int} and increased equivalent series resistance (ESR); the latter property is directly proportional to the instantaneous potential change at a given charge-discharge transition,^[76,81] and hence it is measurable at both the upper and lower threshold of the spanned potential window (0.0 V and 1.0 V). Of the four samples, Na-B (Figure S15a and b) experienced the lowest rate of capacitance decay, delivering a capacitance retention of 89% after 2,000 cycles. By comparison, K-B (Figure S15c and d) retained 75% of its initial discharge capacitance by the end of the cycling process, whilst the cryptomelane samples, Na-A (Figure S15e and f) and K-A (Figure S15g and h), exhibited markedly inferior retention values of 60% and 46%, respectively. It should be noted that since C_{int} effectively averages the capacitance of a given system over the measured *E* range, its application as a storage metric should be restricted to cases where differential capacitance, which correlates with the instantaneous rate of potential variation, dE/dt, at a given location on the GCD curve, exhibits negligible potential-dependence.^[76] In the present case, every dataset depicted in Figure 6 and Figure S14 is characterized by a strongly linear relationship between *E* and time *t*, indicating that differential capacitance indeed remained approximately constant for the duration of each GCD experiment.

By constructing linear fits over the charge and discharge regions of the GCD measurements, the variation of C_{int} was estimated as a function of J (Figure 7) for Na-B (Figure 7a and b), K-B (Figure 7 c and d), Na-A (Figure 7e and f) and K-A (Figure 7g and h), both before (Figure 7a, c, e and g) and after (Figure 7b, d, f and h) prolonged galvanostatic cycling. Additionally, the Coulombic efficiencies, η_{0} , of the pre- and postcycled systems were estimated at each J value (Figure S16 of the Supporting Information) by computing the ratio of the cumulative charge transferred to and from the electrodes during the relevant charge-discharge cycle. Even though the four Na_xK_vMnO_z products comprised two different polymorphs and displayed varying levels of cation pre-insertion, the asprepared electrodes delivered remarkably similar storage characteristics up to current densities on the order of 10 Ag^{-1} ; indeed, differences between C_{int} estimates associated with galvanostatic discharge at 10 Ag⁻¹ (Table 2) are of comparable magnitude to the corresponding experimental uncertainties. At higher current densities, pre-cycling pseudocapacitance was more significantly influenced by the intrinsic properties of guest cations, with K-B and K-A initially delivering greater values of C_{int} and η_0 at 100 A g⁻¹ than the Na-based electrodes. However, K-B and K-A subsequently experienced a faster cycling-induced decline in storage performance than Na-B and Na-A, evidenced by the relatively large separations between their pre- and postcycling the η_0 vs J trends. The post-cycling inferiority of the Kintercalated systems is most apparent from capacitance measurements acquired at high current density; for instance, the C_{int} values exhibited by K-B and K-A at 100 Ag^{-1} (Table 2) are an order of magnitude lower than the corresponding estimates yielded by Na-B and Na-A. The comparatively pronounced destabilization caused by K⁺ intercalation/deintercalation is reflective of the substantial discrepancies between the pre- and post-cycling $|J(0.5 \text{ V})|/\nu^{1/2}$ vs $\nu^{1/2}$ behaviors of K-B (Figure S12b) and K-A (Figure S12d), which are much greater than corresponding differences between the $|J(0.5 \text{ V})|/\nu^{1/2}$ vs $\nu^{1/2}$ relationships of Na-B (Figure S12a) and Na-A (Figure S12c).

Table 2. Pre- and post-cycling estimates of the specific integral capaci- tance, C_{intr} exhibited by the four $Na_xK_yMnO_z$ samples under discharge at current densities of 10 A g ⁻¹ and 100 A g ⁻¹ .					
Na _x K _y MnO _z sample	Pre-cycling C _{int} (discharge) [F g ⁻¹] 10 A g ⁻¹ 100 A g ⁻¹		Post-cycling C_{int} (discharge) [F g ⁻¹] 10 A g ⁻¹ 100 A g ⁻¹		
Na-B K-B Na-A K-A	155 ± 6 149 ± 6 141 ± 5 146 ± 5	70 ± 3 96 \pm 4 71 \pm 3 94 \pm 4	146 ± 5 116 ± 5 138 ± 5 112 ± 4	$50 \pm 3 \\ 4.6 \pm 0.6 \\ 22.0 \pm 1.3 \\ 5.6 \pm 0.6$	







Figure 6. Pre- (a, c, e and g) and post-cycling (b, d, f and h) GCD behaviors of Na-B (a and b), K-B (c and d), Na-A (e and f) and K-A (g and h) measured at current densities between 1.43 A g⁻¹ and 14.3 A g⁻¹.

21960216, 202



Figure 7. Influence of current density, J, on the estimated specific integral capacitance, C_{intr} of Na-B (a and b), K-B (c and d), Na-A (e and f) and K-A (g and h) during charging and discharging, both before (a, c, e and g) and after (b, d, f and h) subjecting the systems to 2,000 successive GCD cycles at 28.6 Ag⁻¹.



The effects of guest cation type on Na_xK_yMnO_z storage characteristics depend on a complex interplay of factors including ion-transport kinetics, the thermodynamics of ion interactions with the host material, and the influence of quest ions on the dimensions of interstitial channels. In order to better understand differences between the C_{int} characteristics displayed in Figure 7, it is therefore prudent to address each stage of the pseudocapacitance mechanism in turn. In the hydrated state, K⁺ ions exhibit greater conductivity $(73.5 \text{ S} \text{ cm}^2 \text{mol}^{-1} \text{ at } 25 \,^{\circ}\text{C})$ than their Na⁺ counterparts (50.1 S cm² mol⁻¹ at 25 °C),^[13,48,51,82] enabling more rapid diffusion towards the Na_xK_yMnO_z surface. Upon entering the host framework during a discharge cycle, K⁺ ions can more readily shed the solvation shell of coordinating water molecules as they possess a relatively low enthalpy of hydration compared with the Na⁺ case.^[48-50] The resulting bare K⁺ ions have a lower charge density than Na⁺ ions and consequently experience weaker electrostatic attraction to the MnO₆ octahedra, resulting in a smaller activation barrier for deintercalation and less contraction of the diffusion pathways.^[26,30,31,48-50] Furthermore, pre-insertion of alkali metal ions improves the electronic conductivity of Na_xK_yMnO_z with respect to pristine MnO₂ by enhancing the Mn 3d and O 2p density of states close to the Fermi level,^[14,33,45,50] and density functional theory (DFT) predicts that K⁺ insertion promotes this phenomenon to a greater extent than Na⁺ intercalation.^[14,45,50] Although the pseudocapacitive behaviors of as-prepared K-B and K-A electrodes were well-matched by Na-B and Na-A at current densities on the order of 10 Ag^{-1} , these considerations provide a plausible explanation for the superior performance of K-based systems at higher current densities, where storage processes are more significantly hindered by underlying kinetic limitations.

Whilst undergoing the sequence of 2,000 galvanostatic cycles, K-B and K-A experienced more rapid capacitance decay than their Na-based equivalents. Given the comparatively high mobility of K⁺ ions, it is ostensibly paradoxical that K-mediated storage mechanisms were more susceptible to cycling-induced disruption than corresponding processes involving Na⁺ ions; however, several factors could be responsible for this finding. For instance, the radius of a bare K^+ ion (1.4 Å) is significantly larger than in the Na^+ case (1.0 Å), $^{\scriptscriptstyle [30,45,51,52,54,83]}$ causing it to encounter greater steric hindrance and a larger energy barrier for diffusion^[50,54] after removal of the hydration shell. Charge exchange with the MnO₆ octahedra is similarly impeded by the size of K⁺ ions, resulting in fewer accessible electroactive sites for intercalation,^[47] again, this effect is likely to be exacerbated by degradation of the Na_xK_yMnO₂ lattice, with electroactive sites depleted by the collapse of interstitial channels or material loss through disproportionation and subsequent dissolution of Mn-(III) species.^[29,30] Finally, since the electrostatic forces between guest K⁺ ions and MnO₆ octahedra are inherently weaker than corresponding Na⁺ interactions, it follows that K⁺ pre-insertion cannot counteract structural disintegration as effectively as the pre-intercalation of Na⁺ ions.^[15,29,32-36]

Due to competition between the properties that caused enhanced capacitance fading in K-B and K-A and the compensating factors that enabled these materials to outperform Na-B and Na-A in their as-prepared states, the relative storage characteristics of Na- and K-based electrodes are liable to vary between studies due to disparate synthesis and testing protocols. For example, whilst various investigations have demonstrated that K⁺ insertion/extraction affords greater capacitance and Coulombic efficiency than intercalation/deintercalation of Na⁺ ions,^[14,31,45,48-50] others showcase the opposite trend.^[11,25,46,47,51-53] Similarly, the superior capacitance retention of K-mediated systems in some reports^[31,45,48] is countered by alternative publications wherein K⁺ intercalation yielded similar or inferior cycling stability to the Na⁺ case.^[14,30,46,50,53] These literature disagreements highlight the clear need for research that rigorously addresses the benefits and pitfalls of Na_xK_yMnO_z products using a self-consistent approach, rather than relying on comparisons between potentially incompatible studies.

Returning to the relationships depicted in Figure 7 and Figure S16, it is apparent that the crystallographic structure of the as-prepared materials had a lesser impact on storage behaviors than the nature of intercalating cations: the C_{int} vs J and $\eta_{\rm O}$ vs J trends of the as-prepared Na-B and K-B electrodes closely resemble those of Na-A and K-A, respectively, whilst the post-cycling storage characteristics of the acid-synthesized products are only fractionally inferior to corresponding relationships from their basic counterparts. This similarity between the electrochemical properties of the birnessite and cryptomelane systems is potentially surprising given that the BET specific surface areas of Na-A and K-A were an order of magnitude greater than in the birnessite samples (Table 1). However, it should be recognized that specific surface area estimates derived from nitrogen adsorption isotherms are not representative of the area that is accessible to intercalating cations, which experience greater steric hindrance than nitrogen molecules and are also obstructed by electrode additives such as the ethyl cellulose binder layer. Indeed, research has shown that there is little correlation between the specific capacitance and surface area of Na,K,MnO, systems, even when comparing products with the same crystallographic phase.[41-43] Analogous to the contradictions between literature comparisons of Na- and Kintercalated materials, existing studies deliver varying conclusions regarding the relative electrochemical attributes of comparable α -MnO_z and δ -MnO_z products: some works indicate that the superior specific surface area afforded by cryptomelane structures is the primary determining factor in overall storage performance, [40,41,44] whereas others suggest that the birnessite phase offers superior capacitance, [22,37-39,42] likely due to the availability of wider ion passages than in the cryptomelane case.^[8]

Equivalent Circuit Modelling

When discussing the GCD behaviors of the electrodes, it was noted that the instantaneous step-change in potential at a charge-discharge transition scales in direct proportion to the ESR of the pseudocapacitive system. However, the true magnitude of this resistance is difficult to determine accurately due to the finite time required for electrochemical instrumenta-



tion to reverse the direction of the applied current. In the case of a linear GCD relationship, an alternative strategy is to estimate the ESR by extrapolating the *E* vs *t* trend back to the charge-discharge transition,^[81] but this approach is complicated by a transient non-linear phase associated with the "equivalent distributed resistance" (EDR) caused by charge redistribution through surface electrode pores.^[76,81] Another problem with estimating the ESR from GCD measurements is that it fundamentally represents an intricate combination of Ohmic, charge-transfer and diffusion-borne resistances; hence, the practice of determining a single characteristic resistance value is implicitly predicated on an unrealistic assumption that the system can be approximated as a series configuration of a resistor and an ideal capacitor.

A more reliable way to investigate the impedance properties of a storage electrode is electrochemical impedance spectroscopy (EIS), which enables the various contributions of resistance and capacitance phenomena to be distinguished from relationships between the real and imaginary components of impedance, Z_{i} and the frequency, f_{i} of applied potential perturbations.^[84] EIS characterization is typically performed at the measured open-circuit potential of the working electrode, as this value corresponds to equilibrium between oxidation and reduction reactions of electrolyte species at the electrode surface, where the Butler-Volmer equation for redox processes is approximately linear.^[85] However, open-circuit potential measurements recorded prior to every CV experiment reveal that this equilibrium scenario did not occur at a consistent potential value over the course of electrochemical testing: open-circuit potential values in the range 0.21–0.76 V vs Ag AqCI (3 M KCI) were exhibited by the four electrodes, possibly due to cyclical variations in the average Mn oxidation state and the changing concentrations of intercalated Na⁺ and K⁺ ions.^[30,86] To overcome this difficulty, EIS was instead conducted at two widely separated bias potentials within the electrochemical stability window, 0.0 V and 0.5 V vs Ag AgCI (3 M KCI), thereby enabling impedance behaviors of each system to be assessed as a function of the applied potential. Indeed, it has been acknowledged elsewhere that since the equivalent circuit components of supercapacitor systems are commonly voltagedependent, the practice of conducting EIS exclusively at the open-circuit potential is liable to restrict the information that can be gleaned from the technique.^[87]

Using EIS data acquired at a base potential of 0.5 V vs Ag | AgCl (3 M KCl) (Figure 8), Nyquist (Figure 8a and b) and Bodemagnitude (Figure 8c and d) plots were constructed to develop equivalent circuits for the systems both before (Figure 8a and c) and after (Figure 8b and d) they underwent 2,000 galvanostatic cycles at 28.6 Ag⁻¹; corresponding plots yielded by EIS measurements obtained at 0.0 V vs Ag |AgCl (3 M KCl) are supplied in the Supporting Information (Figure S17). Each dataset was modelled using a modified Randles equivalent circuit^[88–94] (Figure 8e and Figure S17e) composed of an Ohmic series resistance, R_{sr} equal to the sum of electrode and electrolyte resistance, a resistance R_{ct} and capacitance C_{ct} to account for pseudocapacitive charge transfer resulting from cation intercalation/deintercalation, and a constant phase element, Q_{dlr} to approximate the non-ideal capacitive behavior of electrostatic double-layers at the electrode surface. The latter contribution has a complex impedance given by Equation 5,^[88]

$$Z_{dl} = \frac{1}{Q_{dl}(i\omega)^n} \tag{5}$$

where ω denotes the angular frequency of potential perturbations, and *n* is a constant that equals one in the case of an ideal capacitor. Finally, a Warburg impedance element, $Z_{\rm W}$, describes slow diffusion-controlled ion transport to electro-active sites via passivating surface layers such as the ethyl cellulose polymer binder. Over the practical frequency range of the EIS experiments, $Z_{\rm W}$ obeys the Cottrell equation (Equation 6),^[26,88]

$$Z_W = \frac{\sigma_W}{\sqrt{i\omega}} = \frac{\sigma_W}{\sqrt{2\omega}} (1 - i)$$
(6)

where $\sigma_{\rm W}$ is a constant known as the Warburg coefficient. It can be shown that $\sigma_{\rm W}$ is inversely proportional to the square root of the diffusion constant for charged species travelling to and from the electrode surface, and hence it serves as a useful indicator of ion mobility. Estimates of the equivalent circuit parameters for the as-prepared systems are compiled in Table 3 and Table S1 of the Supporting Information, which correspond to bias potentials of 0.5 V and 0.0 V vs Ag|AgCl (3 M KCl), respectively. Similarly, post-cycling fitting values obtained from EIS data acquired at 0.5 V and 0.0 V vs Ag|AgCl (3 M KCl) are collated in Table 4 and Table S2 of the Supporting Information, respectively.

A noteworthy feature of the applied equivalent circuit model is that it uses a single ideal capacitance, C_{ctr} to simulate the pseudocapacitive characteristics of a given $Na_xK_yMnO_z$ system. Nevertheless, the generated fitting functions provide an

Table 3. Pre-cycling estimates of the Ohmic series resistance, R_s , charge-transfer resistance, R_{cr} , charge-transfer capacitance, C_{cr} , Warburg coefficient, σ_{wr} , electric double-layer capacitance, Q_{dr} and the constant phase element parameter n, obtained by modelling EIS measurements acquired at a bias potential of 0.5 V vs Ag | AgCl (3 M KCl); all gravimetric variables are normalized with respect to the estimated mass of Na_xK_yMnO_z.

Parameter	Na-B	K-B	Na-A	K-A
$R_{\rm s} [\Omega {\rm mg}]$	0.51	0.49	0.54	0.38
$R_{\rm ct} [\Omega {\rm mg}]$	2.4	1.5	2.4	1.1
$C_{\rm ct} [{\rm F} {\rm g}^{-1}]$	150	140	120	130
$\sigma_{\rm W} \left[\Omega {\rm mg}{ m s}^{-1/2} ight]$	3.8	3.2	3.0	2.6
Q_{dl} [F s ^{<i>n</i>-1} g ⁻¹]	6.9	1.6	2.8	1.5
n	0.70	0.80	0.76	0.83

Research Article doi.org/10.1002/celc.202300210



Figure 8. Nyquist (a and b) and Bode-magnitude (c and d) plots constructed from EIS measurements at a bias potential of 0.5 V vs Ag AgCl (3 M KCl), acquired before (a and c) and after (b and d) subjecting the electrodes to 2,000 GCD cycles at 28.6 A g^{-1} . Fitting functions were generated using a modified Randles equivalent circuit model (e) comprising impedance contributions from: Ohmic series resistances, R_s, in the electrodes and electrolyte; charge-transfer resistances, R_{dt}, at the interface between electrode and electrolyte; a capacitance, C_{ct}, resulting from intercalation/deintercalation processes; a constant phase element, Q_{dt} corresponding to electric double-layer capacitance; and a Warburg impedance element, Z_w, describing diffusion-controlled ion transport.

accurate representation of pre-cycling measurements, indicating that this simplification is indeed appropriate due to the fast kinetics of intercalation/deintercalation mechanisms. It is instructive to recall that the rapidity of cation insertion/extraction is also evident from corresponding pre-cycling low- ν CV (Figure 5a, c, e and g) and low-J GCD (Figure 6a, c, e and g) trends, which bear a close resemblance to the ideal rectangular CV and linear GCD relationships of a classic capacitor. By contrast, slow, diffusion-controlled storage phenomena would manifest as distinct peaks in the cyclic voltammograms and plateaus in the GCD traces,^[76] but no such features appear in any of the pre-cycling CV or GCD datasets. The model is further

Table 4. Post-cycling estimates of the Ohmic series resistance, $R_{s'}$ charge-transfer resistance, $R_{ct'}$ charge-transfer capacitance, $C_{ct'}$ Warburg coefficient, $\sigma_{W'}$ electric double-layer capacitance, Q_{dv} and the constant phase element parameter n, obtained by modelling EIS measurements acquired at a bias potential of 0.5 V vs Ag | AgCl (3 M KCl); all gravimetric variables are normalized with respect to the estimated mass of Na_xK_vMnO_z.

Parameter	Na-B	K-B	Na-A	K-A
$R_{\rm s} [\Omega {\rm mg}]$	0.31	0.35	0.33	0.32
$R_{\rm ct} [\Omega {\rm mg}]$	3.6	8.7	5.2	16.0
$C_{ct} [Fg^{-1}]$	160	130	170	110
$\sigma_{\rm W} \left[\Omega {\rm mg}{ m s}^{-1/2} ight]$	2.5	4.0	5.5	7.9
$Q_{\rm dl} [{\rm F} {\rm s}^{n-1} {\rm g}^{-1}]$	4.9	2.0	5.3	2.5
n	0.70	0.77	0.70	0.75

validated by the parameter C_{ctr} which, in line with the magnitudes of C_{int} estimates derived from GCD measurements (Figure 7), mostly adopts values in the range 100–200 Fg⁻¹. When comparing C_{ct} and C_{int} in this way, it is important to acknowledge that the form of each EIS fitting function is not significantly sensitive to small changes (on the order of 10%) in the variables $C_{ctr} Q_{dl}$ and σ_{W} , and hence these quantities are constrained to an accuracy of two significant figures in Table 3 and Table 4, and in Table S1 and Table S2 of the Supporting Information. Whilst the equivalent circuit model generates marginally inferior descriptions of the post-cycling EIS data, most noticeably in the case of Na-A, it nevertheless provides useful indicative parameter estimates for the purpose of investigating the origins of capacitance decay.

From the near-convergence of the high-frequency x-intercepts in the Nyquist plots (Figure 8a and b, and Figure S17a and b), Ohmic resistance was evidently affected little by compositional and crystallographic differences between the four Na_xK_yMnO_z materials. However, at both 0.0 V and 0.5 V vs Ag AgCl (3 M KCl), R_s was found to decrease over the course of electrochemical testing, which is possibly attributable to gradual penetration of the electrode films by surrounding electrolyte. As apparent from the relative diameters of the semicircular sections of the Nyquist plots, K-B and K-A offered significantly less resistance to charge transfer processes than Na-B and Na-A in the as-prepared state, but they also experienced greater increases in $R_{\rm ct}$ during prolonged testing. Charge transfer was similarly influenced by crystallographic phase: the birnessite materials, Na-B and K-B, exhibited higher initial R_{ct} values than their cryptomelane counterparts at both bias potentials, but they delivered lower R_{ct} values after the GCD cycling process. With the notable exception of Na-B, enhanced charge-transfer resistance during GCD cycling was accompanied by a corresponding increase in $\sigma_{\rm W}$, which signifies a diminished rate of ion diffusion between the electrolyte and electroactive surface sites.

The pre- and post-cycling variations of $R_{\rm ct}$ are qualitatively consistent with results from the GCD tests, wherein it was shown that the as-prepared Na-B and Na-A systems delivered fractionally lower values of $C_{\rm int}$ at given J than K-B and K-A (Figure 7a, c, e and g, and Table 2), despite displaying greater post-cycling capacitances (Figure 7b, d, f and h, and Table 2). The $R_{\rm ct}$ trends also align with the $|J(0.5 \text{ V})|/\nu^{1/2}$ vs $\nu^{1/2}$ relationships acquired from CV data (Figure S12): in comparison to K-B and K-A, the post-cycling $|J(0.5 \text{ V})|/\nu^{1/2}$ vs $\nu^{1/2}$ variations of Na-B

and Na-A more closely resemble their pre-cycling behaviors, implying that they were less significantly impacted by the cycling procedure. From these arguments it follows that changes in $R_{\rm ct}$ are strongly linked to performance deterioration during GCD cycling, characterized by decreases in $C_{\rm int}$ and $\eta_{\rm Q\nu}$ and diminished linearity of the J vs ν relationship. In this regard, Na⁺ insertion/extraction ostensibly had a weaker destabilizing influence than the intercalation/deintercalation of K⁺ ions, whilst δ -MnO₂ showcased greater cycling stability than α -MnO₂.

Conclusions

To explore the effects of synthesis precursor concentrations on the morphology, composition and electrochemical properties of co-precipitated Na_xK_yMnO_z products, sample materials formulated at two different pH values with an abundance of either Na⁺ or K⁺ cations were studied using a combination of characterization techniques. Through these investigations it was demonstrated that:

- Acidic synthesis conditions (pH 1.7) favor the formation of cryptomelane (α -MnO₂) nanorods, whereas basic solutions (pH 12.3) preferentially yield a turbostratic birnessite (δ -MnO₂) product composed of agglomerated nanoflakes with randomly ordered lattice planes.
- Hydroxide ions in the synthesis solution suppress reduction of Mn⁴⁺ ions to the 3+ oxidation state, with δ -MnO_z materials exhibiting an O/Mn atomic ratio close to two, as expected for pristine MnO₂.
- Intercalation/deintercalation of alkali metal cations occurs more readily in the interlayer channels of δ -MnO_z than the one-dimensional tunnel structure of α -MnO_z, enabling the former to achieve higher Na/Mn and K/Mn atomic ratios than their α -MnO_z equivalents.
- Variations in specific integral capacitance and cycling stability correlate with charge-transfer resistance: materials with no Na content initially exhibited higher capacitances and lower charge-transfer resistances than Na_xK_yMnO_z products from a Na-rich environment, yet the K-only electrodes became more resistive to charge transfer than Na-based systems during prolonged GCD cycling, culminating in inferior post-cycling capacitance and Coulombic efficiency.
- Crystallographic phase has a weaker influence than cation type on the electrochemical characteristics of as-prepared Na_xK_yMnO_z electrodes. Comparing α -MnO_z and δ -MnO_z

products intercalated by the same type of principal guest cation, pre-cycling differences between the specific integral capacitances of the two polymorphs were of similar magnitude to the experimental uncertainties.

• In comparison to equivalent α -MnO_z materials, the δ -MnO_z phase is less vulnerable to electrochemical performance deterioration during prolonged GCD cycling.

Combining these findings, charge-transfer resistance is identified as the primary source of performance loss during prolonged cycling, and it is reasoned that K⁺ intercalation/ deintercalation has a stronger destabilizing effect on capacitance and Coulombic efficiency than the insertion/extraction of Na⁺ ions. In addition, the kinetics of cation intercalation/ deintercalation into birnessite δ -MnO_z were shown to be less sensitive to GCD cycling than in cryptomelane α -MnO_z, as evidenced by the evolution of charge-transfer resistance in these materials. It therefore follows that the long-term storage performance and cycling stability of a Na_xK_yMnO_z electrode may be maximized by selecting a birnessite product with Na⁺ as the predominant guest cation, despite K-based systems offering marginally superior storage capabilities in the initial stages of electrochemical testing.

Whilst a plethora of research has addressed the effects of quest cations and crystallographic phase on the electrochemical properties of Na_xK_yMnO_z, studies have hitherto provided insufficient detail regarding the effects of prolonged testing on storage performance. By evaluating the relationships between storage parameters both before and after subjecting Na,K,MnO, electrodes to thousands of GCD cycles, the present work overcomes this limitation of existing investigations by providing a more comprehensive understanding of how storage performance is influenced by crystallography, morphology and the nature of intercalating species. Furthermore, competitive electrode characteristics have been attained via a scalable, lowtemperature synthesis strategy and environmentally sustainable, low-toxicity materials such as an ethyl cellulose (an inexpensive, non-toxic and environmentally sustainable polymer binder^[95,96]). In these ways, the present investigation provides a useful platform for additional development and optimization of Na_xK_yMnO_z materials, as well as informing further research into areas such as the interplay between crystallographic phase and cation pre-insertion, and the physical origins of capacitance decay during cycling.

Experimental Section

Reagents and Materials: All chemicals employed in this work were of analytical grade and were used as received. Manganese(II) sulfate monohydrate (MnSO₄.H₂O, \geq 99%), potassium permanganate (KMnO₄, \geq 99.0%), sodium sulfate (Na₂SO₄, \geq 99.0%), potassium sulfate (K₂SO₄, \geq 99.0%), sodium hydroxide (NaOH, \geq 98%), potassium hydroxide (KOH, \geq 85%) and 4-hydroxy-4-methyl-2-pentanone (99%) were acquired from Sigma-Aldrich Ltd., whilst ethyl cellulose (48.0–49.5% ethoxyl content), Super P carbon black (\geq 99%), nickel foil (99.5%, 0.25 mm thickness) and platinum wire (\geq 99.99%, 0.5 mm diameter) were obtained from Fisher Scientific Ltd.

Synthesis of Na_xK_yMnO₂: For the gram-scale synthesis of Na_xK_yMnO₂, a 400 mL aqueous solution of Mn₂SO₄ (100 mM), KMnO₄ (67 mM), and either Na₂SO₄ or K₂SO₄ (400 mM) was prepared by sequentially combining separate solutions of each precursor, with the KMnO₄ solution introduced as the final reagent. In an alkaline synthesis process, the pH of the solution was subsequently adjusted to 12.3 under constant stirring through the dropwise addition of aqueous NaOH or KOH (1.0 M), yielding a final added volume of between 104 mL and 113 mL; the cation in the hydroxide base was chosen to match the cation present in the sulfate salt. To obtain a pH 1.7 precursor solution, 104 mL of deionized water was instead added to the original 400 mL volume in place of the NaOH or KOH solution. In both synthesis routes, the final pH of the solution was measured using a Mettler Toledo FiveGo F2 pH probe. With continuous stirring, the combined solution was then heated under reflux for 12 hours at 75 °C in a thermocouple-controlled silicone oil bath. The resulting suspension was allowed to cool naturally to room temperature before being centrifuged several times with deionized water in order to remove residual reagents from the Na_xK_yMnO_z precipitate. Finally, the product was redispersed in acetone and dried in a muffle furnace at 60 °C.

Phase Identification: The crystallographic structures of the Na_xK_yMnO_z products were determined by transmission-mode powder X-ray diffractometry (XRD) using a Bruker D8 Advance X-ray diffractometer. The instrument was equipped with Cu–K α irradiation and a nickel filter, and it was set to operate at an X-ray power of 1,600 W (40 V×40 A); measurements were recorded over a 2 θ range of 5–90°, with a dwell time of 15 s and 2 θ increments of 0.025° between steps, and each sample powder was rotated at 30 rpm inside a special glass capillary with a 0.7 mm outside-diameter.

Morphology Investigations: Examinations of product morphology were carried out via a combination of field-emission scanning electron microscopy (SEM), transmission electron microscopy (TEM) and scanning transmission electron microscopy (STEM). For SEM, a Hitachi S-4800 instrument was employed with emission voltage and current settings of 10 kV and 10 µA, respectively, and each sample powder was pressed onto an adhesive carbon tab. A FEI Talos F200X system was used to perform TEM and STEM at an accelerating voltage of 200 kV, for which each sample was prepared by first introducing a small quantity of the product (less than 0.1 mg) into a 1:1 v/v mixture of isopropanol and methanol, before agitating the suspension in an ultrasonic bath and then transferring it dropwise onto a 300-mesh lacey carbon-coated copper grid. Where possible, lattice d-spacings were estimated from TEM images by applying ImageJ software to obtain distance measurements over a minimum of ten lattice planes.

Composition Analysis: The surface chemical composition of each material was estimated through deconvolution of X-ray photoelectron spectroscopy (XPS) measurements acquired using a Kratos Axis Supra system, which was fitted with a monochromated Al–K α source operating at an emission current of 15 mA. For these characterizations, a Retsch PP 25 pellet press was employed in conjunction with a Specac pellet die to compress the Na_xK_yMnO_z products into pellets of 5 mm diameter, and data were collected from four 700 µm×300 µm rectangular locations across the pelletized sample surface. In each case, survey spectra were recorded over binding energies between -5 eV and 1,350 eV, with a dwell time of 100 ms, 160 eV pass energy, and 1 eV energy increments. Higher resolution scans of core photoelectron peaks were subsequently obtained using a dwell time of 1 s, 20 eV pass energy and 50 meV energy increments. Throughout all measurements, electronic charge neutralization was applied with a charge balance potential of 3.3 V, and a filament bias and current of 1.0 V and



0.4 A, respectively. Additional information regarding XPS data interpretation is supplied in the Supporting Information.

Surface Area Measurement: The Brunauer-Emmett-Teller (BET) surface area, $S_{\text{BET}'}$ of each product was estimated from the nitrogen adsorption curve of a 0.27–0.66 g powder sample, as measured using a TriStar II 3020 surface area and porosity analyser; prior to BET analysis, the powders were degassed under vacuum by the instrument at 120 °C for one day. A comprehensive overview of the BET analysis procedure is included in the Supporting Information.

Electrode Preparation: In order to improve electron conductivity between Na_xK_yMnO_z particles, viscous slurries composed of the synthesized Na_xK_yMnO_z powder, Super P carbon black and ethyl cellulose in 4-hydroxy-4-methyl-2-pentanone were prepared, with the solid constituents present in a mass-ratio of 7:2:1, respectively; each slurry had a ca. 5:1 mass-ratio of liquid to solid constituents. Mixing was carried out using a Thinky ARE-250 planetary mixer, operated in five-minute steps at a rotational velocity of 2,000 rpm; this blending process was conducted over a minimum of two steps after combining the slurry components, with a further mixing step performed immediately prior to coating. All mixtures were allowed to age overnight before use. Each slurry was deposited over a rectangular 40 mm × 15 mm area of nickel foil (100 mm × 20 mm × 0.25 mm) by tape-casting with a glass rod, before allowing the film to dry naturally in air.

To decrease the risk of delamination during electrochemical testing, the cathode films were pressed between PTFE-coated heating sheets using a Devil Press S15 hydraulic press for 15 minutes at an estimated pressure of 7.5 MPa and a temperature of 120°C; a piezoelectric CBES button load cell was employed to construct a linear calibration plot for the press. The total film mass after pressing was measured by subtracting the mass of the uncoated nickel substrate from the final coated mass, using a mass balance with a stated precision of 0.1 mg. The electrodes investigated herein had a measured film mass in the range 2.6-2.8 mg, which translates to 0.30–0.33 mg cm⁻² of electroactive material, assuming that the film components were distributed homogeneously throughout the coated slurry. Finally, uncoated areas of the nickel substrate were masked with a UV-curing Solarez epoxy resin, which was hardened at a peak wavelength of 302 nm by a Thermo Scientific Pierce 3UV lamp with a stated power density of 2.8 mW cm⁻². A schematic illustrating the electrode preparation sequence is provided in the Supporting Information (Figure S18).

Electrochemical Testing: An Ivium-n-Stat electrochemical station was used to characterize each Na,K,MnO, working electrode in a threeelectrode configuration containing a platinum coil counter electrode, an Ag AgCI (3 M KCI) reference electrode, and an electrolyte of aqueous Na₂SO₄ or K₂SO₄ (0.5 M), selected so that the cation matched the predominant alkali metal ion present during synthesis. Prior to testing, dissolved oxygen was expelled from the electrolyte by sparging with nitrogen gas for two hours at a flow rate of 0.5 Lmin⁻¹, and the electrode was allowed to stabilize in the deaerated electrolyte for a minimum of one hour at open-circuit. Electrochemical storage properties were subsequently investigated using a combination of cyclic voltammetry (CV), galvanostatic charge-discharge (GCD) cycling and electrochemical impedance spectroscopy (EIS). Throughout the present study, gravimetric variables such as specific capacitance and specific current density are normalized with respect to the estimated mass of Na_xK_vMnO_z within the electrode film, invoking an assumption of slurry homogeneity to approximate the mass of Na_xK_yMnO_z as 70% of the measured film mass.

CV experiments were conducted over five cycles between 0.0 V and 1.0 V vs Ag|AgCl (3 M KCl) at potential scan rates in the range 10– $\,$

1,000 mVs⁻¹; the first cycle of each CV dataset was commenced at the open-circuit potential of the system, which was measured prior to cycling for a period of five minutes. Analysis of the CV data was performed by addressing measurements from the fifth cycle of each CV procedure. When utilizing Dunn's formula (Equation 4), the magnitude of current J, |J|, was taken as the mean of the two |J|values yielded by the oxidative and reductive sections of a given cyclic voltammogram, and the uncertainty in |J| was approximated as half the difference between these values. GCD measurements were carried out over two consecutive cycles between 0.0 V and 1.0 V vs Ag AgCI (3 M KCI) at current densities ranging from 1.43 A g⁻¹ to 143 A g⁻¹, with all electrochemical parameters estimated from the second charge-discharge cycle. In both CV and GCD sequences, tests were carried out in staggered order so that any systematic drift could be quantified from step-changes within the measured trends. EIS data were generated over a frequency range of 100 kHz to 0.1 Hz at DC bias potentials of 0.0 V and 0.5 V vs Ag AgCl (3 M KCl), applying AC potential oscillations with a peak-to-peak amplitude of 10 mV. Electrode durability was evaluated by repeating the same sequence of CV, GCD and EIS procedures after performing 2,000 GCD cycles at 28.6 A g^{-1} ; prior to commencing these post-cycling CV, GCD and EIS experiments, the electrolyte was replaced with a fresh batch of $Na_2SO_{4(aq)}$ or $K_2SO_{4(aq)}$ (0.5 M) solution, which was again sparged with a 0.5 Lmin⁻¹ flow of nitrogen gas for two hours prior to use. The Supporting Information contains further details about the protocol used to obtain specific integral capacitance estimates from the GCD measurements.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author. Additional references are cited within the Supporting Information.^[66,70,74-76,97]

Acknowledgements

This study is partially funded by Enserv Power Co. Ltd., and the research contributes to the IMPACT operation which has been part-funded by the European Regional Development Fund through the Welsh Government and Swansea University. The authors are grateful for access to XPS and TEM instrumentation provided by the Advanced Imaging of Materials (AIM) facility, which is funded in part by EPSRC (EP/M028267/1), the European Regional Development Fund via the Welsh Government (80708), the Ser Solar project via Welsh Government, and Carl Zeiss Microscopy.

Competing Interests

Author Kittiwat Kamlungsua is employed by Enserv Power Co. Ltd, which part-funded the research. The authors have no other potential conflicts of interest to declare.

Data Availability Statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Keywords: Birnessite · Cryptomelane · Intercalation · MnO₂ · **Pseudocapacitance**

- [1] B. Babu, P. Simon, A. Balducci, Adv. Energy Mater. 2020, 10, e2001128.
- [2] W. Jing, C. H. Lai, W. S. H. Wong, M. L. D. Wong, Sustain. Energy Technol. Assess. 2017, 22, 55-64.
- [3] T. M. Narayanan, Y. G. Zhu, E. Gençer, G. McKinley, Y. Shao-Horn, Joule 2021, 5, 2934-2954.
- [4] A. J. Hurd, R. L. Kelley, R. G. Eggert, M. H. Lee, MRS Bull. 2012, 37, 405-410
- [5] J. Xie, Z. Liang, Y. C. Lu, Nat. Mater. 2020, 19, 1006–1011.
- [6] M. Toupin, T. Brousse, D. Bélanger, Chem. Mater. 2004, 16, 3184–3190. [7] W. Guo, C. Yu, S. Li, Z. Wang, J. Yu, H. Huang, J. Qiu, Nano Energy 2019,
- 57, 459-472. [8] R. Samal, B. Chakraborty, C. S. Rout, J. Appl. Phys. 2019, 126, e045112.
- [9] S. K. Ghosh, ACS Omega 2020, 5, 25493-25504.
- [10] H. Bai, S. Liang, T. Wei, Q. Zhou, M. Shi, Z. Jiang, J. Feng, M. Zhang, Z. Fan, J. Power Sources 2022, 523, e231032.
- [11] Q. Ye, R. Dong, Z. Xia, G. Chen, H. Wang, G. Tan, L. Jiang, F. Wang, Electrochim. Acta 2014, 141, 286-293.
- [12] H. Zhang, L. Wu, J. Electrochem. Soc. 2018, 165, 2815-2823.
- [13] N. Zarshad, A. U. Rahman, J. Wu, A. Ali, F. Raziq, L. Han, P. Wang, G. Li, H. Ni, Chem. Eng. J. 2021, 415, e128967.
- [14] L. Niu, L. Yan, Z. Lu, Y. Gong, T. Chen, C. Li, X. Liu, S. Xu, J. Energy Chem. 2021, 56, 245-258.
- [15] N. Jabeen, Q. Xia, S. V. Savilov, S. M. Aldoshin, Y. Yu, H. Xia, ACS Appl. Mater. Interfaces 2016, 8, 33732-33740.
- [16] Q. Chen, J. Chen, Y. Zhou, C. Song, Q. Tian, J. Xu, C. P. Wong, Appl. Surf. Sci. 2018, 440, 1027-1036.
- [17] B. Xu, L. Yu, M. Sun, F. Ye, Y. Zhong, G. Cheng, H. Wang, Y. Mai, RSC Adv. 2017, 7, 14910-14916.
- [18] M. Huang, Y. Zhang, F. Li, L. Zhang, R. S. Ruoff, Z. Wen, Q. Liu, Sci. Rep. 2014, 4, e3878.
- [19] A. Xia, W. Yu, J. Yi, G. Tan, H. Ren, C. Liu, J. Electroanal. Chem. 2019, 839, 25-31.
- [20] Y. Qian, C. Huang, R. Chen, S. Dai, C. Wang, Int. J. Electrochem. Sci. 2016, 11.7453-7460.
- [21] R. Su, H. Wang, Y. Sun, P. Guo, Colloids Surf. A 2021, 609, e125579.
- [22] R. A. Aziz, R. Jose, J. Electroanal. Chem. 2017, 799, 538-546.
- [23] Z. Hu, X. Xiao, L. Huang, C. Chen, T. Li, T. Su, X. Cheng, L. Miao, Y. Zhang, J. Zhou, Nanoscale 2015, 7, 16094-16099.
- [24] A. Adomkevicius, L. Cabo-Fernandez, T. H. Wu, T. M. Ou, M. G. Chen, Y. Andreev, C. C. Hu, L. J. Hardwick, J. Mater. Chem. A 2017, 5, 10021-10026.
- [25] P.Y. Chen, A. Adomkevicius, Y.T. Lu, S.C. Lin, Y.H. Tu, C.C. Hu, J. Electrochem. Soc. 2019, 166, 1875-1883.
- [26] C. J. Raj, R. Manikandan, P. Sivakumar, D. O. Opar, A. D. Savariraj, W. J. Cho, H. Jung, B. C. Kim, J. Alloys Compd. 2022, 892, e162199.
- [27] X. Wang, Y. Wang, X. Zhao, Crystals 2022, 12, e115. [28] R. Teimuri-Mofrad, E. Payami, A. Piriniya, R. Hadi, Appl. Organomet.
- Chem. 2022, 36, e6620.
- [29] T. Liu, Y. Li, InfoMat 2020, 2, 807-842.
- [30] D. Chen, D. Ding, X. Li, G. H. Waller, X. Xiong, M. A. El-Sayed, M. Liu, Chem. Mater. 2015, 27, 6608-6619.
- [31] P. Xiong, R. Ma, N. Sakai, X. Bai, S. Li, T. Sasaki, ACS Appl. Mater. Interfaces 2017, 9, 6282-6291.
- [32] Y. Liu, J. Xu, J. Li, Z. Yang, C. Huang, H. Yu, L. Zhang, J. Shu, Coord. Chem. Rev. 2022, 460, e214477.
- [33] S. Zhu, Y. Wang, J. Zhang, J. Sheng, F. Yang, M. Wang, J. Ni, H. Jiang, Y. Li, Energy Environ. Mater. 2022, 6, e12382.
- [34] C. Y. Tsai, T. S. Su, T. C. Wei, M. S. Wu, Electrochim. Acta 2022, 430, e141107.
- [35] H. Dai, R. Zhou, Z. Zhang, J. Zhou, G. Sun, Energy Mater. 2022, 2, e200040.
- [36] D. A. Kitchaev, S. T. Dacek, W. Sun, G. Ceder, J. Am. Chem. Soc. 2017, 139, 2672-2681.
- [37] H. A. Pan, O. Ghodbane, Y. T. Weng, H. S. Sheu, J. F. Lee, F. Favier, N. L. Wu, J. Electrochem. Soc. 2015, 162, 5106-5114.
- [38] A. A. Radhiyah, M. I. Izwan, V. Baiju, C. Kwok Feng, I. Jamil, R. Jose, RSC Adv. 2015, 5, 9667-9673.
- [39] E. A. Arkhipova, A. S. Ivanov, K. I. Maslakov, R. Y. Novotortsev, S. V. Savilov, H. Xia, A. V. Desyatov, S. M. Aldoshin, Ionics 2022, 28, 3501-3509

- [40] S. Devaraj, N. Munichandraiah, J. Phys. Chem. C 2008, 112, 4406-4417.
- [41] T. Brousse, M. Toupin, R. Dugas, L. Athouël, O. Crosnier, D. Bélanger, J. Electrochem. Soc. 2006, 153, 2171-2180.
- [42] X. Bai, X. Tong, Y. Gao, W. Zhu, C. Fu, J. Ma, T. Tan, C. Wang, Y. Luo, H. Sun, Electrochim. Acta 2018, 281, 525-533.
- [43] N. Li, X. Zhu, C. Zhang, L. Lai, R. Jiang, J. Zhu, J. Alloys Compd. 2017, 692, 26-33.
- [44] A. Boisset, L. Athouël, J. Jacquemin, P. Porion, T. Brousse, M. Anouti, J. Phys. Chem. C 2013, 117, 7408-7422.
- [45] Q. Xie, G. Cheng, T. Xue, L. Huang, S. Chen, Y. Sun, M. Sun, H. Wang, L. Yu, Mater. Today Energy 2022, 24, e100934.
- [46] M. A. Desai, A. Kulkarni, G. Gund, S. D. Sartale, Energy Fuels 2021, 35, 4577-4586.
- [47] N. Ma, S. Kosasang, A. Krittayavathananon, N. Phattharasupakun, S. Sethuraman, M. Sawangphruk, Chem. Commun. 2019, 55, 1213-1216.
- [48] J. Shao, X. Li, Q. Qu, Y. Wu, J. Power Sources 2013, 223, 56-61.
- [49] Q. Qu, P. Zhang, B. Wang, Y. Chen, S. Tian, Y. Wu, R. Holze, J. Phys. Chem. C 2009, 113, 14020-14027.
- [50] J. Wan, X. Gu, P. Ji, J. Li, J. Lu, S. Luo, B. Li, L. Huang, M. He, C. Hu, Appl. Phys. Lett. 2021, 119, e163901.
- [51] C. Xu, B. Li, H. Du, F. Kang, Y. Zeng, J. Power Sources 2008, 184, 691–694.
- [52] F. Yi, Y. Huang, A. Gao, F. Zhang, D. Shu, W. Chen, H. Cheng, X. Zhou, R. Zeng, Ionics 2019, 25, 2393-2399.
- [53] Y. Shi, M. Zhang, J. Zhao, L. Zhang, X. Cui, X. Zhu, D. Jin, J. Gong, D. Yang, J. Li, Electron. Mater. Lett. 2022, 18, 475-488.
- [54] M. J. Young, A. M. Holder, S. M. George, C. B. Musgrave, Chem. Mater. 2015, 27, 1172-1180.
- [55] N. Qiu, H. Chen, Z. Yang, S. Sun, Y. Wang, Electrochim. Acta 2018, 272, 154-160.
- [56] M. F. L. Haris, C. Y. Yin, Z. T. Jiang, B. M. Goh, X. Chen, W. A. Al-Masry, A. M. Abukhalaf, M. El-Harbawi, N. M. Huang, H. N. Lim, Ceram. Int. 2014, 40, 1245-1250.
- [57] J. Zhou, L. Yu, W. Liu, X. Zhang, W. Mu, X. Du, Z. Zhang, Y. Deng, Sci. Rep. 2015, 5, e17858.
- [58] Y. Munaiah, B. G. S. Raj, T. P. Kumar, P. Ragupathy, J. Mater. Chem. A 2013, 1, 4300-4306.
- [59] M. Villalobos, B. Lanson, A. Manceau, B. Toner, G. Sposito, Am. Mineral. 2006, 91, 489-502.
- [60] B. Yin, S. Zhang, H. Jiang, F. Qu, X. Wu, J. Mater. Chem. A 2015, 3, 5722-5729
- [61] A. Zahoor, H. S. Jang, J. S. Jeong, M. Christy, Y. J. Hwang, K. S. Nahm, RSC Adv. 2014, 4, 8973-8977.
- [62] A. Iyer, J. Del-Pilar, C. K. King'ondu, E. Kissel, H. F. Garces, H. Huang, A. M. El-Sawy, P. K. Dutta, S. L. Suib, J. Phys. Chem. C 2012, 116, 6474-6483.
- [63] K. Chen, M. Wang, G. Li, Q. He, J. Liu, F. Li, Materials 2018, 11, e601.
- [64] R. A. Davoglio, G. Cabello, J. F. Marco, S. R. Biaggio, Electrochim. Acta 2018, 261, 428-435.
- [65] C. Tanggarnjanavalukul, N. Phattharasupakun, J. Wutthiprom, P. Kidkhunthod, M. Sawangphruk, Electrochim. Acta 2018, 273, 17–25.
- [66] M. C. Biesinger, B. P. Payne, A. P. Grosvenor, L. W. M. Lau, A. R. Gerson, R. S. C. Smart, Appl. Surf. Sci. 2011, 257, 2717-2730.
- [67] M. Wang, K. Chen, J. Liu, Q. He, G. Li, F. Li, Catalysts 2018, 8, e138.
- [68] L. Yan, C. Shen, L. Niu, M. C. Liu, J. Lin, T. Chen, Y. Gong, C. Li, X. Liu, S. Xu, ChemSusChem 2019, 12, 3571-3581.
- [69] J. Wang, D. Zhang, F. Nie, R. Zhang, X. Fang, Y. Wang, Environ. Sci. Pollut. Res. Int. 2023, 30, 15377-15391.
- [70] M. C. Biesinger, Appl. Surf. Sci. 2022, 597, e153681.
- [71] C. Zhu, X. Dong, X. Mei, M. Gao, K. Wang, D. Zhao, J. Mater. Sci. 2020, 55, 17108-17119.
- [72] Z. Li, J. Wang, X. Liu, S. Liu, J. Ou, S. Yang, J. Mater. Chem. 2011, 21, 3397-3403.
- [73] D. Wang, K. Wang, L. Sun, H. Wu, J. Wang, Y. Zhao, L. Yan, Y. Luo, K. Jiang, Q. Li, S. Fan, J. Li, J. Wang, Carbon 2018, 139, 145-155.
- [74] R. Bardestani, G. S. Patience, S. Kaliaguine, Can. J. Chem. Eng. 2019, 97, 2781-2791.
- [75] M. Thommes, K. Kaneko, A.V. Neimark, J.P. Olivier, F. Rodriguez-Reinoso, J. Rouquerol, K. S. W. Sing, Pure Appl. Chem. 2015, 87, 1051-1069
- [76] A. Noori, M. F. El-Kady, M. S. Rahmanifar, R. B. Kaner, M. F. Mousavi, Chem. Soc. Rev. 2019, 48, 1272-1341.
- [77] N. Jabeen, A. Hussain, Q. Xia, S. Sun, J. Zhu, H. Xia, Adv. Mater. 2017, 29, e1700804
- [78] L. Athouël, F. Moser, R. Dugas, O. Crosnier, D. Bélanger, T. Brousse, J. Phys. Chem. C 2008, 112, 7270-7277.

Chemistry Europe European Chemical Societies Publishing



European Chemical Societies Publishing

1960216, 202

- [79] H. Jung, M. Taillefert, J. Sun, Q. Wang, O. J. Borkiewicz, P. Liu, L. Yang, S. Chen, H. Chen, Y. Tang, J. Am. Chem. Soc. 2020, 142, 2506-2513.
- [80] J. Wang, J. Polleux, J. Lim, B. Dunn, J. Phys. Chem. C 2007, 111, 14925-14931
- [81] J. Zhao, A. F. Burke, Adv. Energy Mater. 2020, 11, e2002192.
- [82] S. Paul, K. S. Choi, D. J. Lee, P. Sudhagar, Y. S. Kang, Electrochim. Acta 2012, 78, 649-655.
- [83] A. S. Vedpathak, M. A. Desai, S. Bhagwat, S. D. Sartale, Energy Fuels 2022, 36, 4596-4608.
- [84] A. R. C. Bredar, A. L. Chown, A. R. Burton, B. H. Farnum, ACS Appl. Energ. Mater. 2020, 3, 66-98.
- [85] D. Qu, W. Ji, H. Qu, Commun. Mater. 2022, 3, e61.
- [86] P. Scheitenberger, S. Brimaud, M. Lindén, Mater Adv 2021, 2, 3940–3953. [87] A. Adán-Más, T. M. Silva, L. Guerlou-Demourgues, M. F. Montemor,
- Electrochim. Acta 2018, 289, 47-55. [88] Q. A. Huang, Y. Li, K. C. Tsay, C. Sun, C. Yang, L. Zhang, J. Zhang, J. Power
- Sources 2018, 400, 69-86. [89] B. Yan, M. Li, X. Li, Z. Bai, L. Dong, D. Li, Electrochim. Acta 2015, 164, 55-61.

- [90] Z. Wei, Z. Zhang, Y. Ren, H. Zhao, Front. Chem. 2021, 9, e646218.
- X. Li, J. Liu, M. N. Banis, A. Lushington, R. Li, M. Cai, X. Sun, Energy [91] Environ. Sci. 2014, 7, 768-778.
- [92] L. Feng, Z. Xuan, H. Zhao, Y. Bai, J. Guo, C. W. Su, X. Chen, Nanoscale Res. Lett. 2014, 9, e290.
- [93] E. Umeshbabu, M. Satyanarayana, G. Karkera, A. Pullamsetty, P. Justin, Mater Adv 2022, 3, 1642-1651.
- [94] J. Kang, J. Wen, S. H. Jayaram, A. Yu, X. Wang, Electrochim. Acta 2014, 115, 587-598.
- [95] D. Bresser, D. Buchholz, A. Moretti, A. Varzi, S. Passerini, Energy Environ. Sci. 2018, 11, 3096-3127.
- N. Böckenfeld, S. S. Jeong, M. Winter, S. Passerini, A. Balducci, J. Power [96] Sources 2013, 221, 14-20.
- [97] G. Greczynski, L. Hultman, Prog. Mater. Sci. 2020, 107, e100591.

Manuscript received: May 15, 2023 Revised manuscript received: June 9, 2023 Version of record online: June 22, 2023