Sustainable Energy & Fuels

COMMUNICATION

View Article Online

Check for updates

Cite this: Sustainable Energy Fuels, 2018, 2, 941

Received 7th December 2017 Accepted 13th March 2018

DOI: 10.1039/c7se00587c

rsc.li/sustainable-energy

A carbon-coated pyrosilicate, $Na_2Mn_2Si_2O_7/C$, was synthesized and characterized for use as a new positive-electrode material for sodium ion batteries. The material consists of 20-80 nm primary particles embedded in a ≈ 10 nm-thick conductive carbon matrix. Reversible insertion of Na⁺ ions is clearly demonstrated with \approx 25% of its theoretical capacity (165 mA h g⁻¹) being accessible at room temperature at a low cycling rate. The material yields an average potential of 3.3 V vs. Na⁺/Na on charge and 2.2 V on discharge. DFT calculations predict an equilibrium potential for Na2Mn2Si2O7 in the range of 2.8-3.0 V vs. Na⁺/Na, with a possibility of a complete flip in the connectivity of neighboring Mn-polyhedra - from edge-sharing to disconnected and vice versa. This significant rearrangement in Mn coordination (~2 Å) and large volume contraction (>10%) could explain our inability to fully desodiate the material, and illustrates well the need for a new electrode design strategy beyond the conventional "down-sizing/coating" procedure.

Once overlooked in favour of high energy density Li-ion batteries (LIBs), research in the field of non-aqueous Na-ion batteries (NIBs) is now undergoing a revival by virtue of its economic advantages and the relative abundance of Na compared to Li.¹⁻³ Layered oxides (NaMO₂, with M = Co, Fe, Mn, *etc.*) are by far the most extensively investigated candidates as positive electrode materials.⁴ A range of Fe and Mn compounds combined with PO_4^{3-} , SO_4^{2-} , $P_2O_7^{4-}$, *etc.* polyanions have recently been reported and characterized as positive electrodes in NIBs with the ambition of overcoming the performance, cost and safety limitations of the layered oxides.⁵

Generally, polyanion-based Mn compounds exhibit a higher cell potential (>0.5 V) than their Fe-counterparts, yet pose

Manganese pyrosilicates as novel positive electrode materials for Na-ion batteries†

Viktor Renman, 跑 a Mario Valvo, a Cheuk-Wai Tai, 跑 b Cesar Pay Gómez, a Kristina Edströma and Anti Liivat ២ *a

greater challenges in terms of electrochemical performance, typically resulting from the Jahn–Teller (J–T) distortion of Mn^{3+} (3d⁴ valence electron configuration), leading to poor electronic conductivity and sluggish structural conversion between their (de)lithiated/(de)sodiated phases.6 Though nano-sizing of the active particles and the use of electronically conducting coatings7 can often resolve the first challenge, the structural issues still remain. If the crystal structure framework is not able to withstand the strain caused by the J-T distortion of Mn³⁺, this might even result in significant atomic rearrangement, making it energetically favourable for Mn to migrate into alkali ion sites and/or dissolve in the electrolyte.8 Although these possibilities are certainly discouraging, the potential rewards for taming Mn in these electrochemical applications remain huge - thereby justifying continued effort within the battery materials research community.

The structural stress on Na-ion (de)insertion could be greatly alleviated through the use of some polyhedral linkage in the structures. Recent studies have highlighted that the performance of pyrophosphates containing larger $P_2O_7^{4-}$ moieties can benefit from the introduction of this type of structural flexibility. For instance, corner-sharing [Mn–O] polyhedra in Na₂MnP₂O₇ are believed to give better performance than the corresponding edge-sharing connectivity in Li₂MnP₂O₇.^{9,10} The Na₂MnP₂O₇ structure can, in this way, better accommodate the strain introduced by the J–T distortion of Mn³⁺, resulting in only minor atomic rearrangement under charge/discharge.¹¹

We report herein the synthesis of a novel carbon-coated, nanosized pyrosilicate $Na_2Mn_2Si_2O_7$ and a preliminary study of its electrochemical properties in Na half-cells. $Na_2Mn_2Si_2O_7$ has a theoretical capacity of 165 mA h g⁻¹, equal to that of other state-of-the-art Li-based compounds in this family of materials; typically, LiFePO₄.¹² To the best of our knowledge, pyrosilicates have remained unexplored in terms of electrochemical properties until very recently,¹³ unlike their orthosilicate counterparts (A_2MSiO_4 , where M = Mn, Fe, Co; A = Li,¹⁴⁻¹⁷ Na¹⁸⁻²⁰). A few relevant pyrosilicates, such as $Na_2Mn_2Si_2O_7$ and $Na_2Zn_2Si_2O_7$ were reported more than five decades ago; they were

^aDepartment of Chemistry – Ångström Laboratory, Ångström Advanced Battery Centre, Uppsala University, Box 538, SE-751 21 Uppsala, Sweden. E-mail: anti.liivat@kemi. uu.se

^bDepartment of Materials and Environmental Chemistry, Stockholm University, SE-106 91 Stockholm, Sweden

[†] Electronic supplementary information (ESI) available: Materials, methods, and supplementary results (PDF). See DOI: 10.1039/c7se00587c

synthesized hydrothermally under alkaline conditions at both high pressures and high temperatures.²¹ We speculate here that this family of compounds could provide new opportunities in terms of cost-effective and environmentally sustainable materials for both Li- and Na-ion secondary batteries through careful control of composition, particle size and conductive coating.

The target phase, carbon-coated Na₂Mn₂Si₂O₇ ("Na₂Mn₂Si₂-O₇/C") was synthesized through a solvothermally assisted sol–gel technique based on the well-known mechanism of acid-catalyzed hydrolysis and condensation of tetraethyl orthosilicate (TEOS); see the ESI.† The structure was refined successfully ($R_p = 3.87\%$, $R_{wp} = 5.00\%$, $\chi^2 = 3.73$) from powder XRD data (Fig. 1), using a monoclinic structural model derived earlier from single-crystal XRD data.²² A model involving mixed site occupation of Mn- and Na-sites²³ did not result in a significant improvement in the fit. The atomic parameters of the refined structural model are reported separately in Table S2 in the ESI.†

Impurity-free manganese silicates are known to be difficult to synthesize, especially in the presence of carbon.²⁴ Indeed, the phase purity of Na₂Mn₂Si₂O₇/C is here \approx 94 wt%, with \approx 6 wt% of Na₂MnSiO₄ impurity. The uncoated sample was free from impurity phases (Fig. S1 in the ESI†). An Electron Paramagnetic Resonance (EPR) spectrum of Na₂Mn₂Si₂O₇/C (Fig. S6 in the ESI†) displays a single Lorentzian absorption line (width of 21 mT and a *g*-factor of 2.01) indicating the presence of Mn²⁺ species – in good agreement with an earlier report.²⁵

The structure of $Na_2Mn_2Si_2O_7$ is made up of layers of Mn ions separated by $[SiO_3-O_b-SiO_3]^{6-}$ silicate dimers (illustrated as conjoined blue tetrahedra in Fig. 1 inset). These dimers serve as spacers, effectively opening up the structure and accommodating the Na ions. The conjoined silicate tetrahedra have

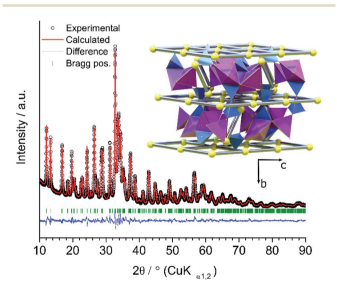


Fig. 1 Experimental (open black circles) and calculated (red line) X-ray diffractograms shown together with their difference (blue line) for the as-synthesized carbon-coated $Na_2Mn_2Si_2O_7$. The inset shows the refined structural model of the title compound; Na atoms (yellow spheres); Na–Na connectivity (rods) within the Si_2O_7 dimer (blue tetrahedra) gallery separating the [MnO_x] layers (purple polyhedra). Note: O atoms are situated at each polyhedral vertex, but have been omitted here for clarity.

a close to staggered configuration relative to one another, and have a non-straight Si–O_b–Si bond angle of 120.9°. The Si–O bond lengths are distributed over the range 1.46–1.74 Å with the longest bond to the bridging O_b as found in earlier studies of other pyrosilicates.²⁶ The Mn ions assume two distinct crystallographic positions within each Mn layer: tetrahedral in a corner-sharing configuration, and square-pyramidal (Mn–O < 2.5 Å) with both corner and edge-sharing connectivity to their neighbouring coordination polyhedra (Fig. 1 inset).

The carbon content of the Na₂Mn₂Si₂O₇/C compound was estimated by thermogravimetric analysis (TGA) to be ~13 wt% (Fig. S3 in the ESI†). Raman spectroscopy confirmed the presence of typical pyrolytic carbon on the surface of Na₂Mn₂Si₂O₇/C with maxima centred at 1594 cm⁻¹ and 1345 cm⁻¹, attributed to the so-called G ("Graphitic") and D ("Disorder") bands^{27,28} (Fig. 2a).

The spectrum of pristine Na₂Mn₂Si₂O₇ shows only weak G and D bands, apparently due to minimal amounts of pyrolysis by-products from the acetate precursor. The additional peaks and bands below 1000 cm⁻¹ are assigned to the pyrosilicate phase.²⁹ The fact that these characteristic peaks and bands are not visible in the Na₂Mn₂Si₂O₇/C sample demonstrates complete coverage of Na₂Mn₂Si₂O₇ surface with highly Raman active carbon. The main peak at 691 cm⁻¹ is ascribed to a symmetric stretching vibration ν_s (Si–O_b–Si) of the pyrosilicate phase and serves as a fingerprint for the presence of the Si₂O₇ groups.^{30,31}

The presence of the pyrosilicate bands is also confirmed by IR spectroscopy, where the bands related to the various symmetric stretching modes within the Si₂O₇ groups appear at 689, 895, 927, 964 and 997 cm⁻¹ for both C-coated and pristine specimens (Fig. 2b). An additional IR peak centred at 830 cm⁻¹ can be assigned to an antisymmetric stretching vibration v_{as} (Si–O_b–Si) of the Si₂O₇ groups, which is typical of non-straight Si–O_b–Si bridges (Si–O_b–Si angle < 180°) and appears to be inactive in Raman.³² The characteristic separation (Δ) between the symmetric (v_s) and antisymmetric (v_{as}) Si–O_b–Si bridge stretching modes in Fig. 2b is then 141 cm⁻¹. As suggested by Lazarev,³³ this separation typically increases as the Si–O_b–Si angle increases. Finally, the purity (within the detection limit) of both products is confirmed by the absence of IR absorption bands at higher wavenumbers, which might have arisen

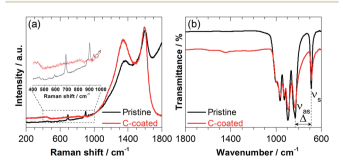


Fig. 2 Raman (a) and FT-IR spectra (b) for uncoated ("pristine") and carbon-coated powders of $Na_2Mn_2Si_2O_7$. Note the domination of the D and G bands in the Raman spectrum of the C-coated powder in (a).

from residues of their xerogel precursors and/or water in the final powders.

Scanning electron microscopy (SEM) shows that Na₂Mn₂-Si₂O₇/C is made up of micron-sized agglomerates (>1 μ m) of primary particles with typical dimensions ~100 nm (Fig. S4†). More accurate determination of the size of the primary particles is hindered by the presence of the surrounding carbon.

Using volume-weighted integral breadth (LVol-IB) in the XRD data, a crude evaluation of the average crystallite size gives ${\sim}46$ nm.

High resolution transmission electron microscopy (HRTEM) of the Na₂Mn₂Si₂O₇/C sample confirms that the secondary particles comprise agglomerated crystallites with typical size 20-80 nm (Fig. 3b and c). Characteristic d-spacings for Na₂- $Mn_2Si_2O_7$ are $d_{110} \approx 7.3$ Å, $d_{111} \approx 4.5$ Å and $d_{230} \approx 3.2$ Å (Fig. 3b and c). It is also evident that the crystallites are effectively embedded in a matrix of conductive carbon, which apparently inhibits the growth of the crystallites beyond nano-size under these synthesis conditions. The carbon is mainly amorphous (dominant sp³ hybridization), although turbostratic ordering (sp² hybridized) is also observed in parts of the Na₂Mn₂Si₂O₇/C composite, in line with our Raman observations. A Selected Area Electron Diffraction (SAED) pattern originating from a larger area of the composite material shows that both the main phase and the minor impurity product are monoclinic. As such, they contain a large number of observable (and partly overlapping) diffraction lines, as evidenced by the numerous reciprocal lattice points in Fig. 3d. Totally correct indexing of the diffraction pattern is inhibited by instrument resolution,

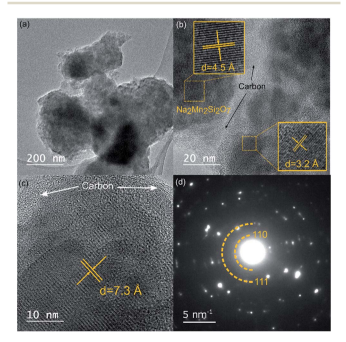


Fig. 3 (a) Low magnification TEM micrograph of the as-prepared carbon-coated $Na_2Mn_2Si_2O_7/C$ compound showing the general features of a typical agglomerate. (b, c) Higher magnification TEM micrographs revealing distinct lattice fringes from the nanocrystals embedded in a matrix of conductive carbon. (d) SAED pattern originating from a larger agglomerate irradiated by the electron beam.

though the largest *d*-spacings (those closest to the primary beam) are uniquely attributed to the primary phase, Na_2Mn_2 -Si₂O₇, by virtue of its significantly larger unit cell (see dashed semicircles in Fig. 3d).

The typical size of the primary particles – spanning tens of nanometers – along with the limited thickness (≈ 10 nm) of the conductive carbon coating should ensure short diffusion pathways for the mobile Na⁺ ions and thereby enhance the overall electronic conductivity of the entire composite electrode. Both factors are deemed necessary in order to mitigate the adverse effects of low conductivity in such polyanionic materials. Note that Na-ion conductivity in Na₂Mn₂Si₂O₇ is not expected to be a performance limiting factor considering the similarity of the Bond Valence Sum (BVS) based Na-ion migration pathways to those in Na₂Fe₂Si₂O₇,¹³ see Fig. S9.†

Na half-cells with $Na_2Mn_2Si_2O_7/C$ and $Na_2Mn_2Si_2O_7$ cathodes both exhibit an open circuit voltage (OCV) of about 2.5–2.6 V vs. Na^+/Na .

However, $Na_2Mn_2Si_2O_7$ shows poor charging performance, as expected in the absence of a proper conductive coating. It was therefore excluded from our further studies.

The cells with $Na_2Mn_2Si_2O_7/C$ clearly cycled better providing approximately 25% of the theoretical capacity at a low cycling rate (C/100); see the inset in Fig. 4.

The differential capacity plot in Fig. 4 reveals two oxidation maxima at \approx 3.2 and \approx 3.4 V in the first charge and a broadened peak around 3.4 V in subsequent cycles. Only one maximum near 2.8 V is observed during discharge, together with a sloping voltage profile below 2.8 V (see the inset). Increasing the upper voltage cut-off beyond 4.0 V makes it possible to access \approx 50% of the theoretical capacity (Fig. S2 inset in the ESI†). However, this was detrimental in terms of higher polarization and could well cause the onset of electrolyte oxidation. Such sluggish cycling behaviour cannot be ascribed to poor electronic contact

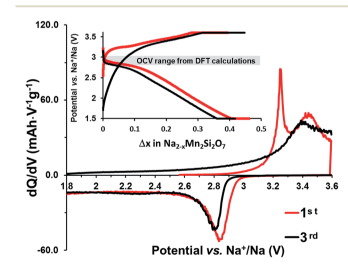


Fig. 4 Differential capacity curve for the 1st (red line) and the 3rd (black line) cycles of the carbon-coated $Na_2Mn_2Si_2O_7$ material cycled galvanostatically in a Na half-cell at RT at C/100 and held at constant upper voltage (3.6 V) and lower voltage (1.5 V) cut-offs for 10 h. The inset shows the corresponding voltage profiles together with the OCV range (shaded) from DFT calculations.

in the active material or to long diffusion times for the Na⁺ ions, since our HRTEM data show the opposite. Alternatively, structural rearrangement, occurring in the desodiation-induced amorphization of Na₂MnSiO₄,¹⁶ could lead to such electrochemical behavior.

Density Functional Theory (DFT) calculations were used to probe in more depth the possible structural origins of such a slow reaction kinetics and the local structural rearrangement occurring under sodium removal from Na₂Mn₂Si₂O₇. In a fully sodiated structure (Fig. 5a), the tetrahedrally coordinated Mn1 sites are ≈ 4 Å apart, while the five-fold coordinated Mn2 sites form an edge-sharing dimer. However, a completely desodiated structure (Fig. 5d) shows an inversion of the Mn1-Mn1 and Mn2-Mn2 connectivity: the Mn1 sites switch to five-coordination and edge-sharing, whereas the Mn2 sites break up their edge-sharing configuration through a rearrangement of the coordinating oxygen positions. This is highlighted by the change in the Mn2–O3 distance from ≈ 4.3 Å in the fully sodiated (Fig. 5a) to ≈ 2 Å in the fully desodiated structure (Fig. 5d). Note that large changes in the metal-oxygen distance (up to 1.2 Å) were reported on the extraction of guest ions for structurally related Li₂MnP₂O₇. This was argued to be the main cause of the sluggish cycling performance in this system.¹¹ Here, we observe even larger changes in the bonding environment of the Mn ions; displacements over a range of ≈ 2 Å occur on full extraction of Na (Fig. S8 in the ESI⁺). Accordingly, it is not surprising that large overpotentials occur to overcome the excessive stress built-up in the structure, especially considering the substantial volume contraction ($\Delta vol = -12\%$) and monoclinic distortion ($\Delta\beta = +15^{\circ}$) accompanying this conversion (Fig. S7 in the ESI[†]).

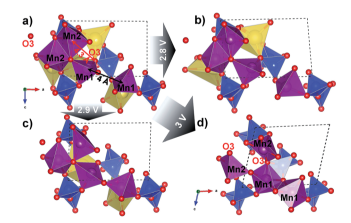


Fig. 5 Series of DFT-optimised Na₂Mn₂Si₂O₇ structures shown along their *b*-direction in the vicinity of two neighbouring Mn1 and Mn2 sites (purple) in sodiated (a), half-desodiated with only Na2 (b) or Na1 (c) sites (yellow) occupied, and fully desodiated (d) states. The marked distances undergo large contraction upon desodiation with the connectivity of Mn polyhedra flipping from edge-sharing to disconnected and *vice versa* for Mn2 and Mn1, respectively. The arrows departing from the initial structure show the calculated desodiation voltages for the states (b)–(d). Si (blue) and O (red) of the Si₂O₇ dimers in the vicinity are also shown. The structure pictures are prepared in VESTA.³⁴

Structural features of this kind could also account for similarly modest electrochemical behaviour in the isostructural $Na_2Fe_2Si_2O_7$ system.¹³

Simulated XRD patterns from the DFT models for x = 2, 1(Na2), 1(Na1) and x = 0 suggest a gradual increase of the (020) reflection intensity relative to the other peaks (Fig. S2c†). This is consistent with *ex situ* XRD data collected for a cycled electrode recovered after charging to 4.2 V using constant current followed by holding the voltage at this constant cut-off value (Fig. S2b†). Such an increase in the intensity of (020) reflection originates from the increase in the ordering of the layer structure of Na₂Mn₂Si₂O₇ under desodiation (Fig. 6).

The calculated OCV values shown for the extraction of Na1 (with the Na2 site occupied, Fig. 5b) and Na2 (with the Na1 site occupied, Fig. 5c) are 2.8 V and 2.9 V vs. Na⁺/Na, respectively. We see here that the configuration in which the Na2 site is still occupied (Fig. 5b) corresponds to a smaller structural rearrangement (judging from the changes in the Mn–Mn and Mn–O distances) compared to the initial structure and is therefore the more probable de-sodiation path in the first charge half-cycle (see S2.7 in the ESI† for further details).

Nevertheless, it is likely that Na1 site occupation (Fig. 5c) can occur at higher overpotentials, and this can also lead to structural perturbation, as reflected in experimental differential capacity data (Fig. 4) which shows a smearing out of the two distinct oxidation processes during subsequent cycling. We speculate, however, that this fully desodiated structure (Fig. 5d) cannot be formed under the test conditions because the available capacity is <50% of the theoretical value, despite the calculated OCV value of \approx 3 V vs. Na⁺/Na (Fig. 5a–d). In practice, only one sodium per formula unit is extracted from Na₂Mn₂Si₂O₇.

To conclude, manganese pyrosilicate was successfully prepared for the use as a new positive electrode material for sodium ion batteries. Despite its very attractive working voltage of \sim 2.8 V and high theoretical capacity of 165 mA h g⁻¹, the

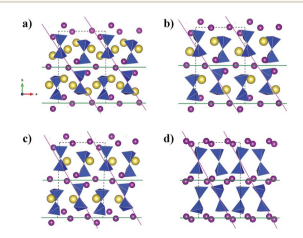


Fig. 6 A layer arrangement of Mn ions (purple) and the Si₂O₇-dimers (blue) together with Na (yellow) is shown for a series of DFT-optimised Na₂Mn₂Si₂O₇ structures in sodiated (a), half-desodiated (b and c) and completely desodiated (d) states as in Fig. 5. The lattice planes corresponding to the reflections (020) (green) and (110) (pink) are shown.

electrochemical activity of this compound, as exhibited in Na half-cells, appears rather moderate. DFT calculations reveal the possibility for complete flip in the connectivity of neighbouring Mn-polyhedra – from edge-sharing to disconnected and *vice versa*. Such a large rearrangement of the Mn coordination (≈ 2 Å) accompanied by a substantial volume contraction (>10%) could well explain our difficulties in completely desodiating the material. Clearly, some alternative to a conventional "down-sizing/ coating" strategy is needed in these and also other polyanion structures exhibiting similar structural freedom. Further studies of the pyrosilicate family using, for example, structure stabilizing dopants, could open new opportunities for cost-effective and environmentally friendly materials for Li- and Na-ion batteries.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was made possible through the financial support from the Swedish Research Council (Grants 2011-6512, 2010-4824), the Röntgen-Ångström Cluster, StandUp for Energy and Swedish Electromobility Centre. The authors would like to thank Henrik Eriksson for his valuable technical support. Mario Valvo gratefully acknowledges the contribution from the Swedish Research Council (FORMAS) through a personal grant no. 245-2014-668. The Knut and Alice Wallenberg (KAW) Foundation is acknowledged for its financial support of the electron microscopy facilities at Stockholm University. The computations were performed using resources provided by the Swedish National Infrastructure for Computing (SNIC) at Linköping University. Fikret Mamedov is also gratefully acknowledged for preliminary EPR measurements.

References

- 1 N. Yabuuchi, K. Kubota, M. Dahbi and S. Komaba, *Chem. Rev.*, 2014, **114**, 11636–11682.
- 2 H. Kim, H. Kim, Z. Ding, M. H. Lee, K. Lim, G. Yoon and K. Kang, *Adv. Energy Mater.*, 2016, **6**, 1600943.
- 3 M. Valvo, F. Lindgren, U. Lafont, F. Björefors and K. Edström, *J. Power Sources*, 2014, **245**, 967–978.
- 4 N. Yabuuchi, M. Kajiyama, J. Iwatate, H. Nishikawa, S. Hitomi, R. Okuyama, R. Usui, Y. Yamada and S. Komaba, *Nat. Mater.*, 2012, **11**, 512–517.
- 5 Q. Ni, Y. Bai, F. Wu and C. Wu, Adv. Sci., 2017, 4, 1600275.
- 6 G. Chen, A. K. Shukla, X. Song and T. J. Richardson, *J. Mater. Chem.*, 2011, **21**, 10126.
- 7 N. Ravet, J. B. Goodenough, S. Besner, M. Simoneau,
 P. Hovington and M. Armand, in *Abstract no. 127, 196th Meeting of the Electrochemical Society*, HI, 1999.
- 8 T. Nordh, R. Younesi, M. Hahlin, R. F. Duarte, C. Tengstedt,
 D. Brandell and K. Edström, *J. Phys. Chem. C*, 2016, 120, 3206–3213.
- 9 M. Tamaru, P. Barpanda, Y. Yamada, S. Nishimura and A. Yamada, J. Mater. Chem., 2012, 22, 24526.

- 10 P. Barpanda, T. Ye, M. Avdeev, S.-C. Chung and A. Yamada, *J. Mater. Chem. A*, 2013, 1, 4194.
- 11 C. S. Park, H. Kim, R. A. Shakoor, E. Yang, S. Y. Lim, R. Kahraman, Y. Jung and J. W. Choi, *J. Am. Chem. Soc.*, 2013, **135**, 2787–2792.
- 12 A. K. Padhi, K. S. Nanjundaswamy and J. B. Goodenough, *J. Electrochem. Soc.*, 1997, **144**, 1188.
- 13 A. Panigrahi, S. Nishimura, T. Shimada, E. Watanabe,
 W. Zhao, G. Oyama and A. Yamada, *Chem. Mater.*, 2017, 29, 4361–4366.
- 14 A. Nytén, A. Abouimrane, M. Armand, T. Gustafsson and J. O. Thomas, *Electrochem. Commun.*, 2005, 7, 156–160.
- 15 R. Dominko, M. Bele, M. Gaberšček, A. Meden, M. Remškar and J. Jamnik, *Electrochem. Commun.*, 2006, **8**, 217–222.
- 16 M. S. Islam, R. Dominko, C. Masquelier, C. Sirisopanaporn, A. R. Armstrong and P. G. Bruce, *J. Mater. Chem.*, 2011, 21, 9811.
- A. Boulineau, C. Sirisopanaporn, R. Dominko,
 A. R. Armstrong, P. G. Bruce and C. Masquelier, *Dalton Trans.*, 2010, **39**, 6310.
- 18 C.-Y. Chen, K. Matsumoto, T. Nohira and R. Hagiwara, *Electrochem. Commun.*, 2014, **45**, 63–66.
- 19 Y. Kee, N. Dimov, A. Staykov and S. Okada, *Mater. Chem. Phys.*, 2016, **171**, 45-49.
- 20 J. C. Treacher, S. M. Wood, M. S. Islam and E. Kendrick, *Phys. Chem. Chem. Phys.*, 2016, **18**, 32744–32752.
- 21 B. N. Litvin, I. M. Dianova and L. A. Kogan, *Kristallografiya*, 1964, **9**, 571–574.
- 22 L. P. Otroshchenko, V. I. Simonov and N. V. Belov, *Dokl. Akad. Nauk SSSR*, 1982, **265**, 76–79.
- 23 R. Tripathi, S. M. Wood, M. S. Islam and L. F. Nazar, *Energy Environ. Sci.*, 2013, 6, 2257.
- 24 R. J. Gummow and Y. He, J. Power Sources, 2014, 253, 315– 331.
- 25 O. V. Kachalov and N. M. Kreines, *J. Exp. Theor. Phys.*, 1968, 26, 522–525.
- 26 V. Kahlenberg, E. Brunello, C. Hejny, H. Krüger, D. Schmidmair, M. Tribus and D. M. Többens, *J. Solid State Chem.*, 2015, 225, 155–167.
- 27 A. C. Ferrari and J. Robertson, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2000, **61**, 14095–14107.
- 28 A. Cuesta, P. Dhamelincourt, J. Laureyns, A. Martínez-Alonso and J. M. D. Tascón, *Carbon*, 1994, 32, 1523–1532.
- 29 M. Wierzbicka-Wieczorek, U. Kolitsch and E. Tillmanns, *Can. Mineral.*, 2010, **48**, 51–68.
- 30 G. Clowry, Z. Molnár and P. Rakic, J. Anat., 2010, 217, 276– 288.
- 31 M. E. Fleet and G. S. Henderson, *Phys. Chem. Miner.*, 1997, 24, 345–355.
- 32 A. N. Lazarev, T. F. Tenisheva, I. A. Bondar' and L. N. Koroleva, Bull. Acad. Sci. USSR, Div. Chem. Sci., 1962, 11, 514–516.
- 33 A. N. Lazarev and T. F. Tenisheva, *Bull. Acad. Sci. USSR, Div. Chem. Sci.*, 1961, **10**, 894–901.
- 34 K. Momma and F. Izumi, *J. Appl. Crystallogr.*, 2008, **41**, 653–658.