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Biomineralization-inspired crystallization of monodisperse α -Mn₂O₃ octahedra and assembly of high-capacity lithium-ion battery anodes†

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Uniform colloidal building-blocks enable the creation of more stable, structurally sophisticated materials. Here we describe a simple polymer-mediated approach to generate grams of monodisperse, singlecrystal α-Mn₂O₃ nanocrystals bound by {111} facets. The technique is inspired in part by biomineralization, where organisms use macromolecular matrices or compartments to trigger the oriented nucleation and growth of crystalline phases. Polyvinylpyrrolidone (PVP) behaves as a polymeric nano-reactor by coordinating to the manganese (Mn) precursor while recruiting the NO_x oxidizing agent from solution to drive the co-precipitation of the manganese oxide. PVP also serves as a molecular template to guide the nucleation of trigonal bipyramids composed of Mn₃O₄. The porosity of the Mn₃O₄ particles indicates that they form non-classically via oriented attachment instead of atom-by-atom. The particles are further oxidized and transform into single-crystal α -Mn₂O₃ octahedra. This co-precipitation approach is advantageous because it can generate large amounts of monodisperse nanocrystals at low economic cost. α-Mn₂O₃ is an alternative lithium ion battery (LIB) anode material that is earth abundant and has \sim 2.7 times higher capacity than conventional graphite anodes. We assembled the monodisperse α -Mn₂O₃ octahedra into LIB anodes to examine their performance in a realistic device. The α -Mn₂O₃ octahedra exhibit good rate performance, cycling stability, coulombic efficiency and morphology retention during extended lithiation-delithiation cycles compared to previous reports for this material. We attribute the improved electrochemical performance of the α -Mn₂O₃ octahedra to the lack of agglomeration in the uniformly distributed electrode and improved lithiation of single crystalline α-Mn₂O₃ nanoparticles.

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Introduction

Inhomogeneities limit the performance of porous electrodes.¹ Inhomogeneity can thrive in porous materials when they are made from a mishmash of particles with a wide distribution in their sizes, shapes, porosities and crystallinities. Such lack of control negatively affects the transport, adsorption and intercalation of ions, which are important in technologies spanning from pH sensors to electrochemical reactors to lithium ion batteries (LIBs). Making homogenous particles is an important pursuit of the colloidal synthesis field, because uniform building blocks enable the creation of functional hierarchical nanostructured materials.²-⁵ In the context of porous electrodes

priority on material uniformity, even though nanoscale and microscale heterogeneities will impact performance and how material interfaces interact in an engineered system.^{6,7} The aim of the shape-controlled colloidal synthesis field is to develop new synthetic tools to create nanoparticles with uniform size, shape and phase. Most notably, understanding how different chemical and physical parameters control the shapes of metallic nanoparticles has led to extraordinary advances in diverse fields including heterogeneous catalysis, biochemical sensing and nano-optics.⁸⁻¹⁴ Achieving better uniformity is a key next step in the creation of more complex hierarchical nanostructured materials whose properties and functions depend on the ordering of materials at multiple length scales.¹⁵⁻¹⁸

for LIBs, new materials development does not place much

Manganese oxides are a promising class of alternative anode materials for use in LIBs because (1) they have higher theoretical specific capacities than conventional graphite anodes, (2) are less susceptible to short circuits due to Li dendrite formation that usually occurs on the graphitic surface at lower potentials, and (3) have lower operating potentials compared to other transition metal oxides based on Co, Ni and Fe. 19,20

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Moreover, recycling of Mn from rechargeable batteries is especially appealing when the battery cathode (LiMn $_2$ O $_4$) and anode (Mn $_2$ O $_3$) comprise Mn. Despite having an \sim 2.7 times better capacity than conventional graphite anodes, α -Mn $_2$ O $_3$ anodes have not been adopted in LIBs because current morphologies and electrode architectures suffer from serious capacity fading at higher charge–discharge rates, resulting in reduced capacity retention. 21

α-Mn₂O₃ is in thermodynamic equilibrium with spinel lithium manganese oxides (LMOs) within the limits of its stability.²² During the initial discharge of the battery, the α-Mn₂O₃ anode incorporates Li⁺ by the reduction of Mn³⁺ to Mn²⁺, which is accompanied by the irreversible formation of Li₂O and further reduction of Mn²⁺ into Mn metal.²³⁻²⁵ Charging causes the Mn metal to be oxidized to Mn²⁺. Like in most conversiontype anodes, the charge-discharge process is accompanied by a phase transition, which causes a huge volume change (≈200%) that can lead to pulverization of the material.26 Pulverization is a particular problem in large, micron-sized polycrystalline particles that can form electrically isolated or electrochemically inactive clusters that contribute to poor performance and device failure. Additionally, while irreversible capacity loss due to secondary electrolyte interface (SEI) formation is unavoidable in manganese oxides, the amount of material lost to the SEI depends on the crystallinity and the surface area to volume ratio of the particle.

In the analogous and well-studied case of spinel LMO cathodes, it is known that surface termination plays an important role in performance and stability.²⁷ Low index facets have the lowest surface energies, and the thermodynamic Wulff shape is predicted to be a truncated octahedron bound by large {111} and smaller {100} facets.28 In LIB cathodes, the stability of LMO is attributed to the growth of a uniform SEI layer, which leads to fewer lattice modifications, particularly on the (111) surface.²⁹ We hypothesize that α -Mn₂O₃ anodes, like LMO cathodes, will have improved performance and cycling stability if the nanoparticles are single crystalline and bound by low index {111} facets. Smaller {100} facets can still enable faster Li diffusion but they are less stable in LMO.27 Additionally, solid non-porous particles could still achieve high performance because the morphological nature of the pores leads to high index surface terminations that cause a non-uniform SEI layer and lower performance over time. Although highly porous materials reduce volume change during charging/discharging, in the case of metal oxides high surface area particles lead to more material being sacrificed to form the SEI layer.

The homogenous oxidation of Mn precursors by O_2 is thermodynamically favorable, but it is kinetically limited without catalysts, alkaline conditions or high temperatures. Thus the conventional (*i.e.* anthropogenic) generation of α -Mn₂O₃ materials is typically done in two steps: co-precipitation of a poorly crystalline manganese adduct such as manganese carbonate, often in a polymer or surfactant matrix, followed by high temperature (>550 °C) annealing to crystallize polydisperse and multi-crystalline α -Mn₂O₃ products. There have been some reports of using solvothermal methods to generate highly

crystalline α -Mn₂O₃ particles, even α -Mn₂O₃ octahedra, but they are either highly polydisperse or porous. Thus the most common conventional approaches have limited control of crystallinity, shape and/or phase. And it is worth noting that, at least superficially, this procedure of making polycrystalline materials and then annealing them for greater crystallinity evokes the two-fold process of biomineralization.

In biomineralization systems, organisms use molecular matrices or compartmentalization to precipitate a poorly crystalline mineral phase, which interacts with ordered arrays of molecules that initiate the nucleation and growth of highquality crystals.36 The oxidation and reduction of manganese is ubiquitous in nature and is mediated by both biotic and abiotic processes. Although the complete role of biological organisms in the oxidation of manganese in the geochemical cycle is not well understood, there is strong evidence that many organisms can oxidize soluble Mn(II) ions with enzymatic superoxides to form Mn³⁺ and Mn⁴⁺ biogenic oxide minerals and even nanoparticles.37-39 The role of macromolecules and polymers in the nucleation and growth of metal oxide materials is fertile ground to seek inspiration for the design and synthesis of metal oxide nanocrystals.40 We hypothesized that a suitable molecular matrix might allow us to avoid the high temperature annealing procedure used in conventional manganese oxide synthesis methods, which would allow us to create nanocrystals with a high degree of monodispersity and crystallinity.

Polyvinylpyrrolidone (PVP) is a nonionic water-soluble polymer that is frequently used in the shape-controlled nanoparticle synthesis of noble metal nanoparticles and metal organic frameworks (MOFs).41-45 The pyrrolidone moiety has a large dipole moment in combination with dispersion forces that are responsible for its reversible interaction with polarizable anions, charged molecules and surfaces. 46-48 Additionally, PVP is known to form complexes with numerous metal ions including Mn.49 In the context of nucleation, these complexes reduce the difference in chemical potential between the precursor and solid,50 which decreases the oversaturation of metal precursors and leads to a slower rate of nucleation and ultimately generates large, monodisperse nanocrystals.51 PVP also has an affinity for oxidizing anions such as NO₃^{-.47} This combination of characteristics makes PVP a good candidate to serve as a polymeric nano-reactor for oxidation reactions and to sustain the growth of large facetted nanocrystals.

In this paper we describe an inexpensive and scalable method to make grams of monodisperse $\alpha\text{-Mn}_2O_3$ nanocrystals with a simple polymer-mediated, bio-inspired approach (Fig. 1). Only moderate temperatures are necessary to oxidize the Mn(NO₃)₂ precursor and trigger the co-precipitation of Mn₃O₄ tetragonal bipyramids within the PVP matrix. At lower temperatures the conditions are relatively non-oxidizing and we observed the formation of multiferroic Mn(HCOO)₃[(-CH₃)₂NH₂] metal organic frameworks (MOFs). Surprisingly, PVP was essential to the reaction, and its omission or exchange with different polymers resulted in different products. The Mn₃O₄ tetragonal bipyramidal nanocrystals appear to nucleate via a non-classical crystallization and then are further oxidized to transform into phase pure $\alpha\text{-Mn}_2O_3$ octahedra. The

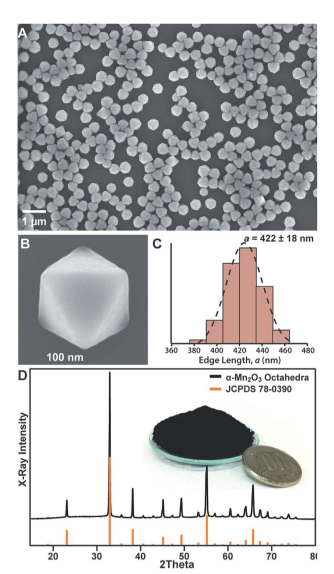


Fig. 1 Synthesis and characterization of $\alpha\text{-Mn}_2O_3$ octahedra. The SEM micrograph shows typically synthesised octahedral particles (A). The particles have smooth facets, and sharp edges and tips (B). The average edge length (a) of the octahedra was 422 \pm 18 nm (C). The synthesis method can yield grams of the material from a simple autoclave reactor ((D) inset). The material was measured using PXRD and the pattern obtained closely matches the reference pattern of $\alpha\text{-Mn}_2O_3$ (JSPDS #78-0390) (D).

oxidation step could serve as a secondary annealing step where the diffusion of Mn^{2+} atoms in the porous Mn_3O_4 framework fills in the gaps in the lattice to make the particles single-crystalline. In the synthesis method the PVP behaves as (i) a modifier to reduce the number of nuclei, (ii) as a shape-directing agent to guide the shape of the $\alpha\text{-M}n_2O_3$ octahedra, and (iii) as a steric barrier to prevent particle aggregation. The monodisperse $\alpha\text{-M}n_2O_3$ octahedra were then used to make uniform porous electrodes, which were assembled into anode half-cells (vs. lithium) for LIBs. We examine how monodisperse $\alpha\text{-M}n_2O_3$ octahedra bound by $\{111\}$ facets have improved electrochemical performance.

Experimental section

Synthesis of α-Mn₂O₃ octahedra

All chemicals were of analytical grade and were used as received without further purification. In a typical experiment, 1.648 g of polyvinylpyrrolidone (PVP; 55 000 M.W.; Sigma Aldrich) was dissolved in 35 mL of N,N-dimethylformamide (DMF; Sigma Aldrich). Then 4.81 mL (0.02 mole) of 50 wt% agueous Mn(NO₃)₂ solution (Alfa Aesar) was briefly mixed into the DMF solution using a magnetic stirrer. This solution was poured into a 100 mL Teflon-lined stainless steel autoclave and heated at 180 °C for 6 h, and then allowed to cool to room temperature and opened in a fume hood. Caution: care must be taken in opening the autoclave because some CO gas and dimethylamine is formed during the reaction. The precipitate obtained was centrifuged (9000 rpm; 10 minutes) and the pellet was resuspended in DMF to remove excess PVP polymer. This washing process was repeated twice. Finally the sample was dried in air and weighed, yielding ~1.1 grams from a 100 mL autoclave.

Structural characterization

SEM images were recorded using a Hitachi S-4800 field emission scanning electron microscope (FESEM). Powder XRD diffraction was carried out using an X-ray diffractometer (RIGAKU RINT 2000 ULTIMA 3) with CuK α radiation (λ = 1.54060 Å) as the incident beam, which is calibrated using SiO₂. TEM analysis was performed with a JEOL 2100-F1 (200 keV) electron microscope. Further details about the structure of the products were elucidated using selected area electron diffraction (SEAD) patterns and high-resolution transmission electron microscopy (HRTEM). Nitrogen adsorption-desorption isotherms were obtained with a high-precision Belsorp-mini apparatus (Bel Japan, Inc.) at 77 K, and the Brunauer-Emmett-Teller (BET) method was used to calculate the surface area of the particles.

Electrochemical testing

Monodisperse octahedral α-Mn₂O₃ single crystals were assembled into a composite anode by evenly mixing 70 wt% α -Mn₂O₃, 20 wt% Super-P carbon (conductive additive) and 10 wt% PVDF (binder) in N-methylpyrrolidone solvent. The obtained slurry was coated on a copper foil using a doctor blade and dried in a vacuum oven at 80 °C for 24 h. For electrochemical performance testing, coin-type 2032 half-cells were fabricated with an α-Mn₂O₃ single crystalline anode, a Celgard 2500 polypropylene separator and lithium metal as the cathode. The electrolyte was composed of 1 M LiPF₆ in a 1:1:1 mixture of ethylene carbonate/dimethyl carbonate/diethyl carbonate. A glovebox filled with high purity Ar (99.999%) and fitted with O_2 and H_2O absorbers was used for fabricating Li-ion half-cells. The oxygen and moisture contents in the glovebox were always maintained below 1 ppm. Cyclic voltammetry (CV) measurements were performed using a Gamry Reference-600 electrochemical workstation. A computer controlled Neware battery tester was used for recording the galvanostatic charge-discharge behavior

of α -Mn₂O₃ anodes in the 3–0 V voltage range. In order to investigate the effect of the lithiation–delithiation process on the morphology and microstructure of α -Mn₂O₃ single crystals, the composite electrodes after 100 galvanostatic cycles were examined by SEM and Raman spectroscopy analysis. To remove the electrolyte solution, the cycled electrodes were washed with dimethyl carbonate followed by drying at room temperature under vacuum. Galvanostatic rate performance and cycling tests were repeated three times and the results were within 5% error limit. All potentials mentioned in this paper are *versus* Li/Li⁺ unless specifically mentioned.

Results and discussion

Crystallizing monodisperse single-crystal $\alpha\text{-Mn}_2O_3$ octahedra with $\{111\}$ facets

Monodisperse α-Mn₂O₃ octahedra were generated via the coprecipitation of manganese oxide precursors into the Mn(III) oxide framework. The Mn precursor is composed of an aqueous 50% Mn(NO₃)₂ precursor mixed with 0.1% HNO₃ (w/w). 4.81 mL of this solution was mixed with 35 mL of 0.422 M PVP in N,Ndimethylformamide (DMF). It was typically heated in an autoclave at 180 °C for 6 hours, although in elaborating the mechanism we changed the temperature and duration of the reaction. Fig. 1A shows a scanning electron microscopy (SEM) micrograph of the octahedra, which conform closely to the ideal octahedral shape with relatively smooth facets and sharp edges and tips (Fig. 1B). The dispersity of these particles was calculated and they were found to have an edge length (a) of 422 nm \pm 18 nm, which is very good for this class of materials (Fig. 1C). The phase of the material was examined by powder X-ray diffraction (PXRD) and matches the known structure of α -Mn₂O₃ (JCPDS 78-0390) (Fig. 1D). Assessing whether a particle is a single crystal is challenging because a majority of crystalline domains can be crystallographically oriented and hide local disorder. Octahedra deposited on surfaces typically sit flat on one of their (111) facets. In high-resolution transmission electron microscopy (HRTEM) the particles could be tilted to align to the $\langle 110 \rangle$ zone axis. And the selected area electron diffraction (SAED) measurement of a single particle indeed matched the pattern for the $\langle 110 \rangle$ direction of crystalline α -Mn₂O₃ (Fig. S1†). There were no additional diffraction spots corresponding to other crystallites. HRTEM along this axis showed that the particle was composed of crystalline α-Mn₂O₃ up to the edge of the particle and no obvious amorphous phases are present. The lack of an amorphous phase or secondary phase is unusual because manganese has numerous oxidation states and the phase is usually determined by oxygen partial pressure. 52-54 To understand the monodispersity of the particles and their high level of crystallinity, a closer look at the chemical and structural mechanism of their formation is necessary.

Oxidation of the ${\rm Mn}^{2^+}$ precursor and sequestration of ${\rm Mn}^{3^+}$ in PVP polymer nano-reactors

In the materials field it is known that some divalent transition metal ions such as Mn²⁺ can form multiferroic Mn(HCOO)₃[(CH₃)₂NH₂] MOFs in DMF. ⁵⁵⁻⁵⁸ This reaction occurs when DMF is hydrolysed at temperatures above ~140 °C to form the formic acid (HCOO) and dimethylamine ((CH₃)₂NH₂) components of the MOF. Indeed, PXRD shows that our standard reaction heated at lower temperatures (t=147-149 °C) generated these multiferroic MOF particles (Fig. S2†). By increasing the temperature of the reaction above t=151 °C we observed α-Mn₂O₃ octahedra exclusively (Fig. S2†). Interestingly, Mn₃O₄ was observed at t=150 °C. The Mn atoms in Mn₃O₄ are in a mixture of +2 and +3 oxidation states, and it is reasonable to assume that the oxidizing power of the reagent/ solvent was not strong enough to completely oxidize the Mn²⁺ starting material. Clearly temperatures above 150 °C were important in triggering the oxidation reaction.

To go further and examine the importance of temperature, the standard reagent solution (*i.e.* 4.81 mL of aqueous 50% $Mn(NO_3)_2$ in 35 mL of 0.422 M PVP in DMF) was heated to $t=151\,^{\circ}\mathrm{C}$ in an open glass vial to observe colorimetric changes over time. Upon heating, the solution acquired a vivid yellow/orange hue within a few minutes (Fig. 2A). Ultraviolet-visible (UV-Vis) spectrophotometry showed that a peak appeared with an absorbance maximum at 468 nm that matches the literature

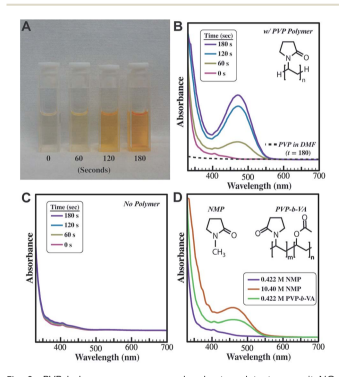


Fig. 2 PVP behaves as a macromolecular template to recruit NO_x from solution and drive the oxidation of the Mn^{2+} precursor. (A) An optical micrograph shows that the standard reaction solution quickly acquires a yellow/orange hue upon being heated at $t=151\,^{\circ}\mathrm{C}$ in an open vial. (B) UV-Vis measurements of the solution have a peak with an absorbance maximum at 468 nm, which matches the known values for Mn^{3+} and also NO_2 . When PVP was omitted from the reaction there was no change in the optical properties of the solution (C). PVP was exchanged for PVP-b-VA or different concentrations of NMP monomer and heated for 180 seconds (D). The pyrrolidone moiety was essential to drive the oxidation reaction, while the polymer microstructure assisted in cooperatively binding the Mn precursor and NO_x oxidizing agent.

values for Mn³⁺ (Fig. 2B).⁵⁹ PVP was critical in the reaction. Omission of PVP from the reagent solution resulted in no change in color (Fig. 2C), and omitting PVP from the solvothermal reaction resulted in the Mn(HCOO)₃[(CH₃)₂NH₂] MOF (Fig. S3†). Exchanging PVP with other DMF-soluble polymers such as poly(4-vinylpyridine) (PVPy) and polyethyloxazoline (PEOX) did not generate Mn³⁺ nor was there any color change indicative of a change in the oxidation state of Mn or generation of NO₂ (Fig. S4A and B†).

It is important to note that heating ($t = 151 \,^{\circ}\text{C}$) 1 mmol nitric acid (HNO₃) in 0.422 M PVP in DMF also led to a change in color that was at least qualitatively similar to the standard reagent solution containing Mn(NO₃)₂. However, the absorbance spectrum indicates that the nitrate ions thermally decompose into nitrous acid (HNO₂) and nitrogen dioxide (NO₂) (Fig. S4C†).^{60,61} Both NO₃ and NO₂ are good oxidizing agents, and NO₂ absorbs at nearly the same wavelength as the Mn3+ ion, so it is challenging to conclusively assign the absorbance change of our reagent solution to Mn³⁺ or NO₂. Oxidizing Mn²⁺ can generate NO₂ so the absorbance spectrum is likely a combination of the two compounds. DMF typically behaves as a reducing solvent so its direct participation in the oxidation reaction would be unlikely.62 Interestingly, when we added HNO3 to neat DMF at t = 151 °C, NO₂ was not observed (Fig. S4D†). This result further confirms that PVP strongly associates with NO3 and recruits and transforms polarizable species from solution to perform oxidation reactions.

In biomineralization systems the polymer creates supersaturation within its microstructure by cooperatively sequestering ions, which decreases the free ion concentration in solution and prevents nucleation outside of the polymer. To test the importance of the polymer microstructure we used 0.422 M N-methyl-2-pyrrolidone (NMP) instead of PVP so that there was an equivalent concentration of NMP to the pyrrolidone monomer residue in the reaction. With 0.422 M NMP in DMF we saw no formation of Mn³⁺ after several minutes (Fig. 2D and S5A†). However, by using NMP as the solvent (i.e. 10.4 M NMP) we could observe Mn3+ and NO2 although the absorbance intensity was lower than those of samples with PVP in DMF (Fig. 2D and S5B†). So the PVP microstructure and its pyrrolidone polymer monomer are critical in driving the reaction. Going further we used 0.422 M poly(1-vinylpyrrolidone-co-vinyl acetate) (PVP-VA) instead of PVP. This polymer contains half the concentration of pyrrolidone subunits separated by acetate groups. There was some generation of Mn³⁺ but the rate of generation was roughly half compared to that of PVP (Fig. 2D and S5C†).

Nucleation of Mn_3O_4 tetragonal bipyramids and further oxidation into Mn_2O_3 octahedra

Fig. 3 shows a general mechanism for the reaction. It is clear from the previous section that the Mn^{2+} precursor and NO_3 associate with the PVP polymer via the pyrrolidone subunit. The polymer microstructure cooperatively binds Mn^{2+} and NO_3 , while assisting in the formation of the NO_x oxidizing agent. PVP serves as a local site for the oxidation of Mn^{2+} to Mn^{3+} thus serving as a nucleation point for the co-precipitation of the

MOF: Mn(HCOO)₃[(CH₃)₂NH₂]

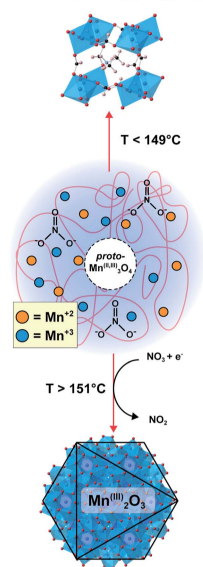


Fig. 3 A scheme describing the polymer-mediated oxidation of the Mn precursor and growth of $\alpha\text{-Mn}_2\text{O}_3$ octahedra. Reactions between 145 and 149 °C yielded the multiferroic Mn(HCOO)_3[(CH_3)_2NH_2] MOF by decomposition of DMF solvent (ref. 52–55). Above 151 °C the Mn precursor is oxidized inside the PVP polymer, generating $\alpha\text{-Mn}_2\text{O}_3$ octahedra. Both the pyrrolidone moiety and PVP microstructure were important in the reaction.

manganese oxide. 47,63,64 The solvothermal reaction is conducted at $t=180\,^{\circ}\text{C}$ (above the boiling point of DMF at 1 atm), so it is difficult to halt the reaction to observe any intermediates. Another approach is to decrease the amount of Mn(NO₃)₂·6H₂O starting material, but the simultaneous change in Mn²⁺ and water concentration will affect the chemistry of the solvent. 35 In the previous section we learned that the starting material begins to oxidize at $t\sim150\,^{\circ}\text{C}$; thus we ran several reactions at this temperature but changed the reaction time to observe any intermediates on the path to α -Mn₂O₃ octahedra. Fig. 4A shows the PXRD measurements of samples heated at 151 $^{\circ}\text{C}$ for 1.0

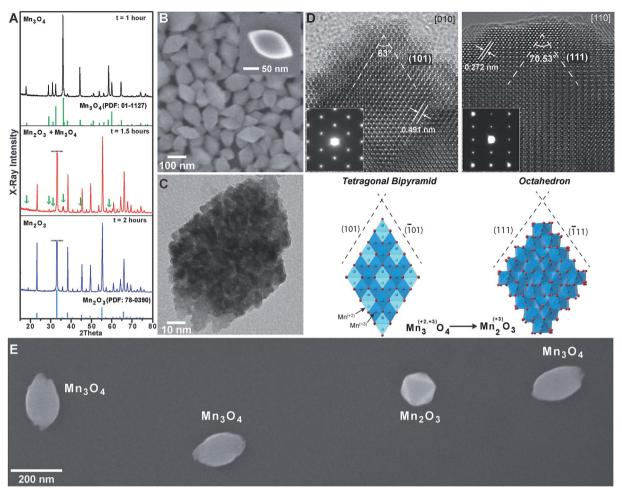


Fig. 4 The solvothermal reaction was conducted at a lower temperature ($t=151\,^{\circ}\text{C}$) to observe any intermediate structures and further elaborate the mechanism. (A) PXRD spectra taken from reactions synthesized for 1.0 hour, 1.5 hours and 2.0 hours. Within the first hour the Mn²⁺ starting material was converted into pure Mn₃O₄, which has a mixed oxidation state of Mn^{II,III}. (B) SEM image of the Mn₃O₄ taken at t=1 hour shows that the particles are tetragonal bipyramids. (C) TEM of a single Mn₃O₄ particle shows that the Mn₃O₄ particles are initially quite porous. ((D) top left) HRTEM image of the tetragonal bipyramid matches the spacing of the {101} planes of Mn₃O₄ and the angle between adjacent planes is 63° ((D) bottom left). After 2 hours the material is oxidized to α -Mn₂O₃. ((D) top right) HRTEM image of the octahedra matches the spacing of the {111} planes of α -Mn₂O₃ and the angle between adjacent planes is 70° ((D) bottom right). The inset selected area electron diffraction patterns (SAED) indicate that each individual particle is a single crystal and no other phases are present ((D) top insets). To view the transition from Mn₃O₄ to α -Mn₂O₃, the reaction vessel was heated at 151 °C for 1.5 hours and then rapidly cooled in water. (E) An SEM image of this sample was composed primarily of Mn₃O₄ tetragonal bipyramids and small amounts of α -Mn₂O₃ octahedra (E).

hour, 1.5 hours or 2.0 hours. The PXRD patterns show that the reaction initially forms Mn_3O_4 which then quickly transforms into α -Mn₂O₃ by the progressive oxidation of manganese:

$$3Mn^{2+} \rightarrow Mn_3O_4 + e^- \rightarrow Mn_2O_3 + 2e^-$$
 (1)

This result again confirms that Mn_3O_4 is the partially oxidized intermediate in the reaction. Initially Mn_3O_4 forms crystals that conform to the pseudo-octahedral shape of a tetragonal bipyramid (Fig. 4B).^{65,66}

Metal oxides in solution are thought to grow either classically atom-by-atom or non-classically *via* the oriented aggregation of prenucleation clusters. Interestingly, the smallest single particles in the 1.0 hour sample were initially

quite porous according to high-resolution TEM (HRTEM) micrographs (Fig. 4C). The $\{110\}$ reflections from the Fourier transform of Fig. 4C were used to construct moiré images, which show that the lattice extended across the entire particle forming a coherent crystalline unit (Fig. 86†). Porosity here indicates that nucleation is likely not homogenous but that the particle grows via the crystallization of primary units that undergo oriented aggregation into some kind of proto-Mn $_3$ O $_4$ particle inside the PVP matrix. This seems reasonable given that the formation of iron oxide minerals in the environment is thought to occur in part via the oriented aggregation of prenucleation Keggin clusters, rather than classical growth. Manganese-based Keggin clusters have been synthesized in the lab, so it is intriguing to

speculate that the Mn₃O₄ particles may form *via* a nonclassical growth mechanism much like iron oxides.⁶⁹

In terms of geometry a tetragonal bipyramid is simply an octahedron that is elongated along one axis. Comparing both crystal structures in HRTEM, it is clear that the surfaces of the Mn₃O₄ particles (growth time t=1.0 hours) are roughly bound by the {101} planes, which have characteristic spacings of 0.491 nm that intersect at 63° (Fig. 4D; left). The selected area electron diffraction (SAED) pattern shows that these intermediate Mn₃O₄ particles are single crystals and no other phases are present (Fig. 4D; top left inset). For the α -Mn₂O₃ octahedra (t=2 hours) viewed along the [110] direction, we can see the {111} planes with spacings of 0.272 nm and facets that intersect at 70.53° (Fig. 4D; right). The SAED pattern of this sample is a distinct diffractogram that matches that of α -Mn₂O₃ along the [110] direction (Fig. 4D; top left inset).

The size of the Mn_3O_4 bipyramids and α - Mn_2O_3 octahedra was roughly the same; thus it is likely that the particles transform via a topochemical oxidation reaction. The overall structural elements of the particles are maintained, but the conversion is very fast and thus it is challenging to observe an initial site of crystallographic rearrangement. So in situ studies are required to prove such a claim. To transform Mn₃O₄ into α-Mn₂O₃ the manganese cations in the particles must diffuse to the surface. The radius of O^{2-} is almost double the radius of Mn(II) or Mn(III), so the diffusion of oxygen should be small in comparison to that of manganese. Additionally, the transformation (topochemical or otherwise) from Mn₃O₄ into α-Mn₂O₃ would result in a great reorganization in the connectivity and bonding of the Mn atoms inside the particles. Again, solute-induced phase transformation in single-crystals tends to happen quickly so it is difficult to capture the observed transition outside of in situ measurements.70 Regardless we heated a reaction at 151 °C for 1.5 hours and then rapidly cooled it in water to room temperature. A typical SEM image of this sample showed that it was composed primarily of Mn₃O₄ tetragonal bipyramids and small amounts of α-Mn₂O₃ octahedra (Fig. 4E). Their similarity in size indicates that the Mn₃O₄ particles must transform into monodisperse α-Mn₂O₃ octahedra. Interestingly, the diffusion of the Mn atoms could act as a secondary annealing step to assist in the formation of single crystal nanoparticles filling the pores and stitching together the lattice. Nitrogen adsorption-desorption isotherms were used to calculate the porosity of the particles compared to hard (i.e. nonporous) octahedra. The specific surface area ($S_{\rm BET}$) of the particles was ~ 2.0 cm³ g⁻¹ which was actually smaller than the theoretical surface area of hard α-Mn₂O₃ octahedra with an edge length of 420 nm (Fig. S7†). This indicates that the particles are non-porous and are essentially hard particles.

Assembly of α-Mn₂O₃ octahedra into LIB anodes

We assembled the α -Mn₂O₃ octahedra into porous electrodes and tested them as anodes in Li-ion half-cells. The Li-ion storage electrochemical performance of single crystalline Mn₂O₃ nanocrystals is first evaluated by cyclic voltammetry. The first five consecutive cycles in the voltage range of 0.0–3.0 V at a scan rate of 1 mV s⁻¹ are presented in Fig. 5A. In the first cathodic sweep

a broad signal between 1.1 and 0.50 V corresponds to the irreversible reduction of $\mathrm{Mn^{3+}}$ to $\mathrm{Mn^{2+}}$ and SEI formation. 23,25,71,72 Another prominent peak centered at 0.10 V matches the reduction of $\mathrm{Mn^{2+}}$ to $\mathrm{Mn^{0}}$. During the subsequent cycles, the main cathodic peak becomes wider and is shifted to \sim 0.12 V. Nevertheless, the first anodic peak was observed at \sim 1.3 V, characteristic of the oxidation of $\mathrm{Mn^{0}}$ to $\mathrm{Mn^{2+}}$. $^{73-75}$ The following anodic signals appeared at 1.4 V, which is a relatively high potential. These potential shifts during consecutive cycles indicate the formation of $\mathrm{Li_2O}$, which increases impedance. Interestingly, the anodic peak at 2.35 V reported in the literature is absent in our α - $\mathrm{Mn_2O_3}$ samples, demonstrating that $\mathrm{Mn^{2+}}$ does not get oxidized back to $\mathrm{Mn^{3+}}$. Consequently, the reduction of the original $\mathrm{Mn^{3+}}$ ion in α - $\mathrm{Mn_2O_3}$ into $\mathrm{Mn^{2+}}$ was irreversible; thus the dominant Li-ion storage mechanism proceeds by an $\mathrm{Mn^{2+}/Mn^{0}}$ couple.

Anodes composed of α-Mn₂O₃ single crystals had a high firstcycle discharge capacity of 1202 mA h g⁻¹ (Fig. S8†). This can be divided into four distinct regions corresponding to different electrochemical processes. The reduction of Mn₂O₃ to Mn₃O₄ $(2Li^{+} + 3Mn_{2}O_{3} + 2e^{-} \rightarrow 2Mn_{3}O_{4} + Li_{2}O)$ during lithiation causes a sudden voltage fade and a plateau around 1.2 V. Further reduction of Mn₃O₄ resulted in the formation of MnO (2Li⁺ + $Mn_3O_4 + 2e^- \rightarrow 3MnO + Li_2O$, causing a corresponding voltage fade of \sim 0.4 V in the discharge profile. Finally, the complete reduction of MnO to Mn ($2Li^+ + MnO + 2e^- \rightarrow Mn + Li_2O$) caused a voltage plateau at 0.4 V. All transition metal oxide anodes form Li₂O during the lithiation step. The fourth region consists of a gradual voltage fade up to 0 V, which is characteristic of electrolyte decomposition to form a solid electrolyte interface (SEI) on the electrode surface. The first charge profile with a total specific capacity of 831 mA h g⁻¹ (Fig. S8†) displayed a voltage plateau around 1.25 V and a shoulder at 2.25 V corresponding to the oxidation of Mn to MnO and MnO_x, respectively by reaction with Li_2O (Mn + $x\text{Li}_2\text{O} \leftrightarrow \text{MnO}_x + 2x\text{Li}^+ + 2x\text{e}^-$) (1.0 < x < 1.5). This Mn²⁺/Mn⁰ redox coupling mechanism limits the practical specific capacity to 832 mA h g^{-1} rather than the theoretical capacity of 1018 mA h g^{-1} for α -Mn₂O₃. A first cycle capacity loss of 44% resulted from irreversible SEI formation and the partial reversibility of Li2O, which is in good agreement with the cathodic CV signals centered at 0.04 and 0.5 V, respectively.

Previous reports describing α-Mn₂O₃ anodes show capacity/ voltage fading with increasing current densities. 23,71,76-78 Fig. 5B shows the second charge-discharge voltage profiles of our octahedral α-Mn₂O₃ anodes at current densities ranging from 100 to 3200 mA g^{-1} . Their profiles were similar to the first charge-discharge curve, and we observed negligible capacity/ voltage fading. During subsequent cycles, the slope between 1.5 and 0.4 V and plateau at 0.4 V can be ascribed to the reduction of Mn3+ to Mn2+ and Mn2+ to Mn0, respectively. There are two slopes in the charge curves, one between 0.0 and 2.1 and the other between 2.1 and 3.0 V. These match the complete oxidation of Mn⁰ to Mn²⁺ and partial oxidation of Mn²⁺ to Mn³⁺. The α-Mn₂O₃ electrodes were then subjected to chargedischarge at increasing current densities (100, 200, 400, 800, 1600 and 3200 mA g^{-1}) to investigate their electrochemical rate capability (Fig. 5C). We observed reversible discharge capacities of 791, 682, 646, 618, 560 and 435 mA h g^{-1} , respectively. These

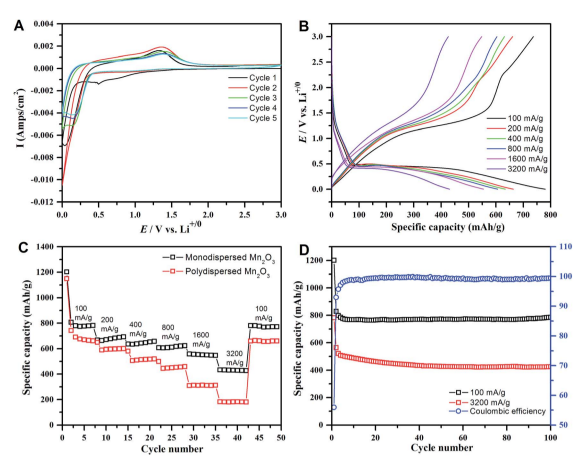


Fig. 5 (A) Cyclic voltammogram, (B) charge–discharge profiles of single crystalline monodisperse α -Mn₂O₃ vs. lithium at various current densities, (C) electrochemical rate performance of monodisperse and polydisperse Mn₂O₃, and (D) galvanostatic cycling performance at various current densities of single crystalline monodisperse octahedral α -Mn₂O₃ single crystals.

rate performances are significantly better than previous reports of $\alpha\text{-M}n_2O_3$ anodes composed of hierarchical structures, nanowires and hollow microspheres (Table S1†), even though our particles had a markedly lower surface area. In order to understand the effect of single-crystallinity and the exposed facets, their electrochemical performances were compared with those of the polydisperse $\text{M}n_2O_3$ sample. The superior rate performance of the monodisperse sample especially at higher current densities can be attributed to the exposed {111} facets.

Additionally, the long-term cycling stabilities of the α -Mn₂O₃ octahedra are superior to those of several Mn₂O₃ nanostructures reported in the literature (Fig. 5D). ^{23,71,77,9,80} A stable specific capacity of 780 mA h g⁻¹ was achieved after 100 galvanostatic charge–discharge cycles at a current density of 100 mA g⁻¹. Even after extended cycling at a high current density of 3.2 A g⁻¹, the α -Mn₂O₃ single crystal anodes retained a capacity of 425 mA h g⁻¹, which is still higher than the theoretical capacity of graphite anodes, which has only been achieved at very low current densities. Our α -Mn₂O₃ anodes experienced no capacity loss after 5 cycles at a low current density of 100 mA g⁻¹, whereas 30 cycles were required for capacity stabilization at a higher current density of 3.2 A g⁻¹. This capacity fading during the initial charge–discharge cycles resulted from the irreversible formation of a SEI and Li₂O. The coulombic

efficiency of these α -Mn₂O₃ electrodes reached 98.8% after 5 charge–discharge cycles, and maintained 99.5% on prolonged cycling even at a high current density of 3.2 A g⁻¹.

To understand the stability of these α -Mn₂O₃ anodes, we opened up an electrode that had been subjected to 100 galvanostatic lithiation–delithiation cycles and examined it using SEM (Fig. 6A–D). SEM images of the pre- and post-cycled particles appear similar in shape and show no evidence of pulverization. This is surprising because strain caused by the lithiation–delithiation process induces non-uniform expansion/contraction. The octahedra are able to accommodate the huge volume change (~200%) without fracturing, even though they are relatively large and non-porous. Uniform {111} surface termination and monodispersity helps ensure that the lithiation–delithiation process proceeds uniformly throughout the electrode and each particle has a similar state of charge (SOC). As a result, the homogeneity should decrease the frequency of hotspots and other heterogeneities that might lead to electrode failure.

The cycled α -Mn₂O₃ octahedra in Fig. 6D have a faint halo around the particles, indicating that a thin and highly uniform SEI layer forms on the {111} facets. A thinner and more uniform SEI layer is beneficial because it consumes less of the liquid electrolyte for charge–discharge cycling, but still protects the Mn³⁺ cations from acidic species that promote the dissolution

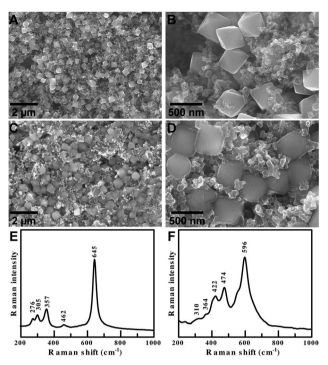


Fig. 6 SEM images of single crystalline octahedral α-Mn₂O₃ electrodes (A and B) before cycling and (C and D) after 100 galvanostatic cycles. Raman spectra of α-Mn₂O₃ composite electrodes (E) before cycling and (F) after cycling of a Li-ion half-cell.

of the anode.29 A thinner SEI has lower interfacial impedance, which is beneficial for stable long term cycling and high rate performance. Surface termination, polycrystallinity, monodispersity and agglomeration are all key factors that reduce electrochemical performance. The cycled anodes do not suffer from any of these problems. Their shape and monodispersity facilitate superior contact of the electrode with the electrolyte solution, resulting in reduced charge transfer resistance and improved performance at high current densities.

Finally, the pristine and cycled electrodes were examined by Raman spectroscopy (Fig. 6E and F). The pristine electrodes displayed peaks that are characteristic of pure α-Mn₂O₃. The cycled electrodes only displayed peaks from Li₂O (596, 474 and 422 cm⁻¹) and MnO (364 and 310 cm⁻¹). This clearly demonstrates that Li₂O is formed during the lithiation of α-Mn₂O₃ octahedra, and that delithiation during the charge process is incomplete. This observation helps confirm our hypothesis that the Li-ion storage mechanism of our α-Mn₂O₃ anodes proceeds by an Mn²⁺/Mn⁰ couple. The Raman measurements show that Li is stored by reversible redox reactions between metallic Mn and a stable fraction of Li₂O to form MnO and MnO_x (1.0 < x < 1.5). The shape and monodispersity confirm less aggregation and good electrode/ electrolyte contact, helping ensure that the oxidation of Mn²⁺ to Mn³⁺ is minimal.

Conclusions

Monodisperse single crystal α-Mn₂O₃ octahedra were synthesized inside PVP polymer nano-reactors. This method

is inspired by a simple biomineralization approach where the initial mineral phase is precipitated in a macromolecular matrix, which then interacts with ordered arrays of molecular ligands to initiate the nucleation and growth of single crystals. The pyrrolidone moiety played an essential role in the reaction by associating with the Mn precursor and recruiting NO₃ from solution. The PVP polymer microstructure enhanced the cooperative binding of precursors and generation of the NOx oxidizing agent locally so that there was a slower rate of nucleation via a decrease in oversaturation in solution. The omission of PVP from the reaction resulted in the generation of multiferroic Mn(HCOO)₃[(CH₃)₂NH₂] MOF microcrystals.

The progressive oxidation of the Mn precursor in the PVP matrix initiated the nucleation of Mn₃O₄. Initially the Mn₃O₄ tetragonal bipyramids were quite porous, indicating that the reaction nucleates via a non-classical mechanism at first, which is similar to some iron oxide mineralization schemes. Others have observed the oriented aggregation of Mn oxide clusters into octahedral nanoparticles but the mechanism was unclear in part because it omitted the Mn₃O₄ bipyramidal intermediate.35 As the reaction was heated longer the Mn3O4 tetragonal bipyramids were transformed into α-Mn₂O₃ octahedra. It is not yet known if the octahedral shape of α-Mn₂O₃ at this length scale is in thermodynamic equilibrium at room temperature, or guided by some kinetic factor determined by the reaction conditions. But simulations of bixybite phase of In2O3 indicate the Wulff shape is an octahedron.81 Finally, other researchers have noted that understanding shape-controlled colloidal synthesis in noble metals is like "making sense of the mayhem". We think extending this idea further to include manganese oxides and the labile nature of the manganese oxidation state adds a new layer of complexity and beauty to colloidal reactions.82

This co-precipitation method enables us to synthesize grams of monodisperse α-Mn₂O₃ octahedra for technological applications. Surface termination and particle-size dispersity are known to play an outsized role in the performance and stability of LIBs. Anodes composed of α-Mn₂O₃ octahedra demonstrated outstanding rate performance, cycling stability and coulombic efficiency compared to previous reports. At the microscale level, monodispersity ensures the spatially homogenous porosity of the composite anode, lending structural stability that is maintained even after extended galvanostatic cycling. Microscale homogeneity increases the maximum current density and limits damage due to hot spots.1 At the nanoscale level, uniform surface terminations ensure similar Li+ insertion and extraction, resulting in a uniform SOC from particle to particle. Nanoscale homogeneity increases power density and improves the response of the cell. Beyond LIBs the material α -Mn₂O₃ is also useful as a bifunctional non-precious metal catalyst in fuel cells.33,83 The synthetic method described here is scalable and inexpensive, and can be extended for the shape-controlled synthesis of other metal oxides to create a new class of hierarchically organized nanostructured materials that are completely assembled from the bottom up.

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