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Mechanistic understanding of microstructure formation during synthesis of metal oxide/carbon nanocomposites†

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Nanocomposite materials consisting of metal oxide and carbon are of interest as electrode materials for both high rate intercalation-type and high capacity conversion-type charge storage processes. Facile synthesis processes like the pyrolysis of an organic carbon-source can yield a well-dispersed carbon phase within the metal oxide structure. Detailed understanding of the carbon formation process is required to tailor the resulting material microstructure. Herein, both the formation and the final microstructure of a molybdenum oxide/carbon nanocomposite are studied in detail. Octylamine assembled in the interlayer space of layered MoO_3 serves as a carbon source. The structural changes during pyrolysis are characterized using a combination of *in situ* heating X-ray diffraction with simultaneous FTIR- and mass spectroscopy-coupled thermogravimetric analysis experiments. These reveal mobility and partial desorption of octylamine and interlayer water at low temperatures, octylamine decomposition and loss of long-range order at intermediate temperatures, and carbothermic reduction of molybdenum oxide at high temperatures during pyrolysis. The resulting nanocomposite mainly contains nanocrystalline MoO_2 domains surrounded by a well-dispersed carbon phase, as observed with scanning transmission electron microscopy of focus-ion beam prepared cross-sectional lamellae. The electrochemical behavior is evaluated in organic, lithium-containing electrolyte for both intercalation and conversion-type reactions, showing good intercalation kinetics and a high first cycle efficiency for the conversion-type reaction.

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1. Introduction

Advancing state-of-the-art electrochemical energy storage devices requires improvements in both specific power and energy.^{1,2} High power capability requires electrodes with both fast ionic and electronic transport properties, necessitating engineering across length scales, from the atomic scale to the macroscale. It is necessary to consider not only the active material properties (*e.g.*, composition, electronic conductivity, ion diffusivity, and particle size), but also the particle-to-particle interactions that contribute to the overall electrode

performance. For example, Luo *et al.* found that V_2O_5 electrodes with micron-scale platelets showed greater heterogeneity in the spatial variation of the state of charge than those with nano-scale platelets.³ Park *et al.* further showed that phase separation observed in layered metal oxides that undergo a solid-solution ion insertion is due to electro-autocatalytic reactions between particles with different states of charge.⁴ These recent results highlight the importance of engineering the particle-to-particle interactions and optimizing the electrode microstructure in energy storage materials.

Transition metal oxides (TMOs) can electrochemically react with Li^+ *via* intercalation⁵ and conversion reactions.⁶ The exact mechanism depends on the extent of electron transfer which determines the degree of structural distortion, with intercalation reactions favored for up to $\sim 1 \text{ Li}^+/\text{e}^-$ per transition metal and conversion reactions for $>1 \text{ Li}^+/\text{e}^-$ per transition metal. Many TMOs are semiconductors or insulators and require the addition of conductive carbon to the electrode for high-rate operation. Electrochemical conversion-type reactions in TMOs are further associated with large volumetric changes, leading to reduced efficiency and cycle life. One strategy to alleviate this issue is to confine TMO particles in a carbonaceous matrix to

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buffer volumetric expansion.^{7,8} Both functionalities – improved electronic percolation and volumetric buffering – can be achieved with TMO/carbon (nano)composite electrodes.^{9,10}

Among many synthesis approaches for TMO/carbon nanocomposites, one of the most widely adopted is the addition of an organic species to the TMO particles and its subsequent carbonization.^{8,11} Recent works have shown that the carbonization of amine molecules added into the interlayer of layered TMOs can produce TMO/C nanocomposites with a well-dispersed carbon phase.^{8,12–16} In some cases, it was described that this synthesis route (and similar ones using different sources of carbon¹⁷) yielded heterostructured TMO/C^{12,16} (and transition metal dichalcogenide/carbon¹⁸) nanocomposites, where layers of dissimilar phases alternate. This particular microstructure results in abundant heterointerfaces which may synergistically combine beneficial electrochemical properties of the individual building blocks (redox-activity, electrical conductivity).^{19,20}

Comparison of these studies^{8,12–16} reveals variations in structural properties of the resulting TMO/C materials, especially in terms of nanocomposite microstructure (*i.e.*, phase distribution), degree of TMO reduction, and carbon hybridization. A more generalized understanding of the synthesis process is required to be able to precisely tune the resulting nanocomposite properties. Therefore, it is important to develop mechanistic insights into the microstructure evolution during carbonization. Understanding the process *via in situ* characterization methods, as well as microscopic analysis representative of the bulk volume of the nanocomposites can give valuable insights into the microstructure evolution. This is essential to further optimize nanocomposite electrode materials for electrochemical energy storage applications and beyond.

In this study, the microstructural changes during pyrolysis of an octylamine-functionalized molybdenum oxide are systematically analyzed. Molybdenum trioxide (α -MoO₃) consists of layers of corner- and edge-sharing distorted MoO₆-octahedra in an orthorhombic crystal structure, where the individual layers are held together by van der Waals forces.²¹ MoO₃ is chosen for this study because organic molecules such as alkylamines can assemble in its interlayer space, serving as a confined carbon source.²² The resulting nanocomposite derived from pyrolysis has relevance for electrochemical applications, because molybdenum oxides can undergo intercalation and conversion reactions with lithium.^{23,24} The octylamine (OA) molecule is chosen for its model character, as alkylamines are linear organic molecules which are widely available with variable carbon chain length, allowing to customize the carbon-source molecule.

We employ a combination of *in situ* characterization and X-ray spectroscopy and electron microscopy techniques to analyze the structural and chemical changes of the molybdenum oxide/octylamine material during pyrolysis. We show that carbon formation occurs predominantly within the micron-sized product aggregates and coincides with molybdenum reduction that is accompanied by CO₂ & CO release. Simultaneously, nanosized oxide grains in the range of 5–10 nm are

formed within the aggregates that exhibit local variations in oxygen and carbon content. Scanning transmission electron microscopy (STEM) imaging of specimen prepared *via* focused ion beam (FIB) reveals the microstructure of the materials representative of their bulk volume. The obtained MoO_x–C nanocomposite exhibits improved kinetics for electrochemical intercalation and an enhanced initial coulombic efficiency for the conversion reaction, depending on the employed potential range, compared to pristine molybdenum oxide samples.

2. Experimental section

2.1 Materials synthesis

Molybdenum oxide dihydrate (MoO₃·2H₂O) was synthesized *via* a modified Freedman method.²⁵ 50 grams of sodium molybdate dihydrate Na₂MoO₄·2H₂O (VWR International) were dissolved in 100 mL of deionized H₂O and slowly added to 300 mL of 5 mol L^{−1} HNO₃ (VWR International) through a burette under continuous stirring. The solution was stirred at room temperature for about 2 weeks, then the stirring was stopped for another week, until a significant amount of yellow precipitate was visible. The precipitated MoO₃·2H₂O was recovered *via* filtration of the solution through a Büchner funnel (reaction yield around 20 g), followed by subsequent washing with about 3 L of deionized H₂O and drying at room temperature for 1 week.

Octylamine-functionalized molybdenum oxide (MoO_x–OA) was synthesized by dispersing 2 grams of MoO₃·2H₂O in 57 mL of ethanol under stirring, before adding 1.43 grams (1.825 mL) of octylamine (Sigma Aldrich) to obtain a molar ratio of Mo : OA of 1 : 1. The solution was stirred for 3 days at room temperature until a white precipitate formed. The precipitate was recovered by washing and centrifugation with ethanol at 2000 rpm for 5 minutes for 3 times and drying at 80 °C overnight. While in this study, a molar ratio of Mo : OA of 1 : 1 was chosen, variation of the ratio yields structures with similar crystallography (Fig. S1†).

Octylamine-derived molybdenum oxide–carbon composite (MoO_x–C) was synthesized by pyrolysis of MoO_x–OA in a tube furnace under flowing argon atmosphere at 700 °C for 2 hours, with a heating rate of 5 K min^{−1}, then allowed to cool to room temperature.

In order to compare the electrochemical properties to conventional molybdenum oxide, MoO₃·2H₂O underwent the same thermal treatment described above, but without assembly of OA in the interlayer (*i.e.*, in the absence of the carbon source). This led to a partially reduced form of molybdenum oxide (MoO_{3-x}).

2.2 Structural and morphological characterization

Powder X-ray diffraction (XRD) patterns of all the samples were obtained using a Bruker D8 Advance equipped with a Cu K α_1 radiation source ($\lambda = 0.15406$ nm). The patterns were recorded in Bragg–Brentano mode in the range between 2° and 60° 2 θ with 0.02° step size. X-ray photoelectron spectroscopy (XPS) of MoO_x–C powder was performed in a fixed analyzer transmission

mode, using a monochromatic Al K α ($h\nu = 1.487$ eV) X-ray and a Phoibos 150 XPS spectrometer (Surface Concept) equipped with a micro-channel plate and Delay Line Detector (DLD). High resolution Mo 2p, C 1s, and O 1s regions were acquired with an X-ray power source of 200 W (12 kV), pass energy of 30 eV, and 0.1 eV energy steps. The spectra analysis was carried out by CasaXPS software, using a nonlinear Shirley-type background and 70% Gaussian and 30% Lorentzian profile functions.²⁶ Scanning electron microscopy (SEM) imaging was performed using a ZEISS Crossbeam 340 electron microscope operated at 5 kV and working distance of 7.7 mm. Raman spectra were obtained using a Renishaw inVia Raman spectrometer equipped with an Nd:YAG laser with a wavelength of 532 nm using 0.25 to 5 mW laser power. *In situ* heating XRD was carried out using a PANalytical Empyrean diffractometer equipped with a HTK 1200N heating stage (Anton Paar) operating under nitrogen atmosphere and replicating the heat treatment conditions during pyrolysis synthesis, while recording a diffractogram every 6 minutes. Thermogravimetric analysis coupled with simultaneous mass spectroscopy (TGA-MS) and Fourier transform infra-red spectrometry (TGA-FTIR) of the pyrolysis gas was performed using a NETZSCH TG 209 F1 Libra thermal analyzer under Ar flow at a heating rate of 10 K min⁻¹ from 40 to 700 °C in Al₂O₃ crucibles loaded with *ca.* 20 mg of material. MS data were acquired with a QMS 403 Aëlos Quadro mass spectrometer in the 10–300 *m/z* range, while FTIR spectra were recorded using a Bruker Invenio spectrophotometer in absorption mode, in the range of 4500–650 cm⁻¹ and at a resolution of 2 cm⁻¹. Scanning transmission electron microscopy (STEM) was carried out at 200 kV and with a semi-convergence angle of 24.7 mrad in an aberration-corrected Titan G2 60-200 CREWLEY microscope (ThermoFisher Scientific, formerly FEI, Netherlands) equipped with a high-brightness field emission electron gun and a Super-X energy-dispersive X-ray spectroscopy (EDS) system. High-angle annular dark-field (HAADF) images were acquired to make use of the associated Z-number contrast with a semi-collection angle of 69–200 mrad.

Samples for STEM were prepared using a dual-beam Helios NanoLab400S (ThermoFisher Scientific, formerly FEI, Netherlands) focused ion beam (FIB). Powder particles were fixated on a conventional scanning electron microscope stub using glue. Afterwards, regions of high particle density were identified, and Pt was deposited on top to make a good connection between the particles. A conventional lift-out technique was applied to transfer the thick TEM lamella onto a TEM grid. Before thinning the lamella, the side of it was again glued with Pt to maintain a good particle connection during thinning. Finally, the sample was thinned down to electron transparency with a final thinning step consisting of 5 kV ion polishing to reduce beam damage and Ga contamination as much as possible.

2.3 Electrode and electrochemical cell preparation

The electrode slurries were prepared by mixing 80 wt% active material and 10 wt% conductive carbon black (Super C65, C-NERGY) in an agate mortar, followed by the addition of

10 wt% polyvinylidene fluoride (PVDF, Solef 6020, Arkema Group) in *N*-methyl-2-pyrrolidone solvent (NMP, anhydrous, Sigma-Aldrich, 2 wt% of PVDF in NMP). The slurries were homogenized using a planetary mixer (Thinky, ARE-250) and then cast onto aluminum and copper foil current collectors (Wellcos Corporation) using a laboratory scale doctor blade (wet film thickness set to 60 μ m). The cast electrodes were further dried in an oven set at 80 °C overnight. Electrode discs 12 mm in diameter were then punched with active material loading ranging between 0.7 to 0.9 mg cm⁻², and dry coating thickness ranging between 12.0 to 13.8 μ m.

2.4 Electrochemical characterization

All cyclic voltammetry and galvanostatic cycling measurements were done in 2-electrode configuration coin cells, where 12 mm diameter metallic Li discs were used as a counter and reference electrode and glass microfiber (Whatman grade GF/A) as separator. Standard LP30 (1 M LiPF₆ in 1 : 1 volume mixture of ethylene carbonate and dimethyl carbonate, Solvionic) was used as the electrolyte. Electrochemical measurements were performed in temperature-controlled chambers (Binder) at 20 °C using a potentiostat/galvanostat (Bio-Logic VMP3). The potential window was between 1.1 and 3.1 V vs. Li⁺/Li (for MoO_x-C) or 1.5 and 3.5 V vs. Li⁺/Li (for MoO_{3-x}) to study the Li⁺ intercalation reaction. To study the conversion reactions at more negative potentials, the electrodes were cycled between 0.01 and 3 V vs. Li⁺/Li.

Electrochemical impedance spectroscopy (EIS) measurements were performed in custom-made 3-electrode cells, where the components are sandwiched between two spring-loaded titanium pistons, and Li foils were used as separate counter and reference electrodes. Detailed description of the cell is found in ref. 27. Impedance spectra were recorded under open circuit potential conditions with sinusoidal potential amplitude of 10 mV, in the frequency range between 200 kHz and 100 mHz.

3. Results and discussion

3.1 Structural characterization

The synthesis of molybdenum oxide dihydrate (MoO₃·2H₂O, also referred to as molybdic acid) was performed *via* the method described by Freedman²⁵ and the chemical self-assembly reaction of octylamine (OA) in the interlayer of molybdenum oxide was achieved by a wet chemistry synthesis to obtain MoO_x-OA. This material was subsequently pyrolyzed at 700 °C for 2 hours under argon flow to form an OA-derived carbonaceous phase. The final product of this synthesis is a nanocomposite MoO_x-C, as described in the Experimental section and shown in the schematic in Fig. 1A. In the following, we provide mechanistic insight into the synthesis/pyrolysis process and final product structure, pursuant of the overarching goal to understand where a carbonaceous phase forms and how the microstructure evolves in the molybdenum oxide/carbon nanocomposite material. In particular, we aim to answer whether a heterostructured microstructure forms, where in an ideal case, 2D



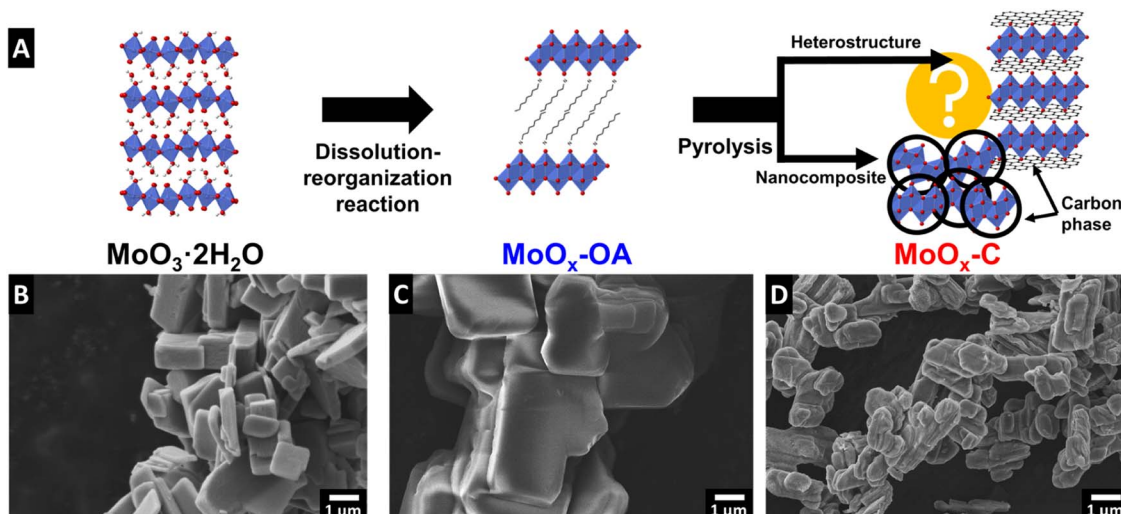


Fig. 1 (A) Schematic illustration of the synthesis process: octylamine is assembled in the MoO_3 interlayer space via a dissolution–reorganization reaction; subsequent pyrolysis yields the carbon-containing nanocomposite. SEM images of (B) molybdenum oxide dihydrate ($\text{MoO}_3 \cdot 2\text{H}_2\text{O}$), (C) MoO_x -OA, and (D) MoO_x -C after pyrolysis in argon at 700 °C.

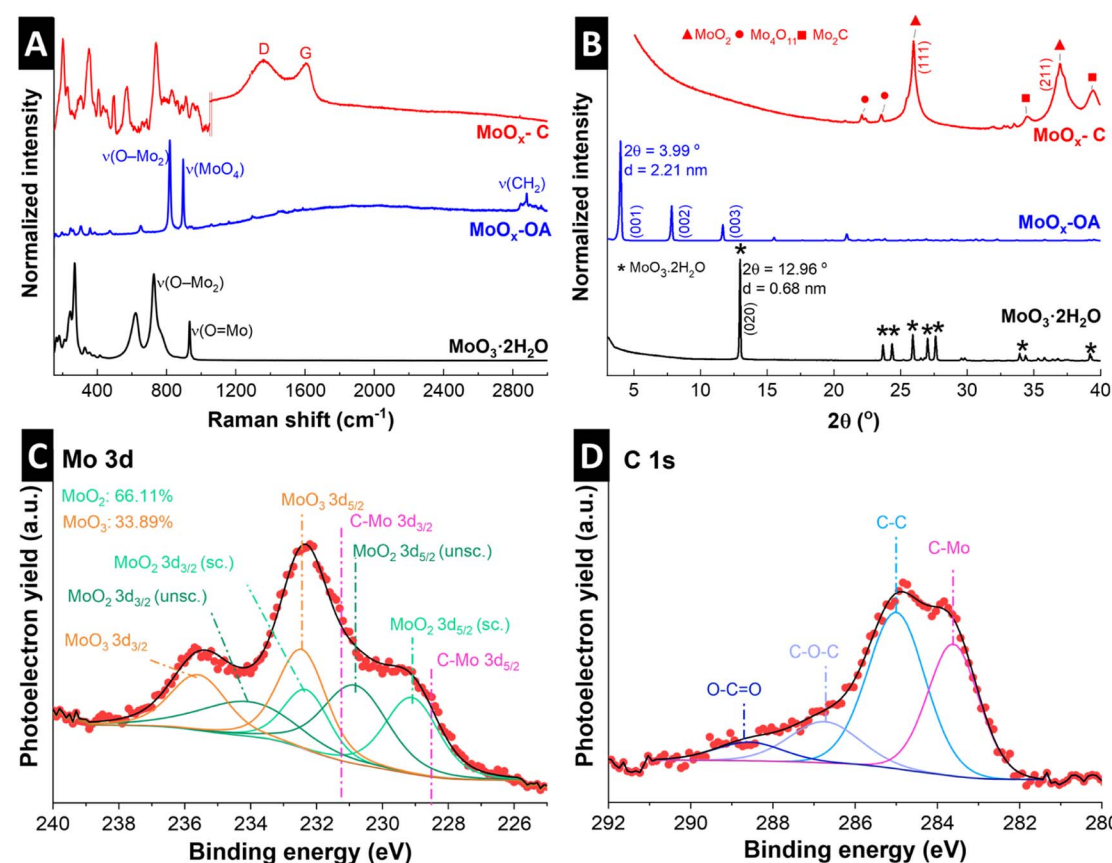


Fig. 2 Structural characterization of hydrous molybdenum oxide ($\text{MoO}_3 \cdot 2\text{H}_2\text{O}$), molybdenum oxide with octylamines assembled in the interlayer space (MoO_x -OA) and pyrolyzed MoO_x -C: (A) Raman spectra of all samples. Raman spectrum of MoO_x -C consists of two individual measurements with different laser power: 0.25 mW between 100–1050 cm^{-1} to avoid oxidation of MoO_2 to MoO_3 ; 5 mW between 1050–3000 cm^{-1} for better resolution of carbon signals (individual full spectra given in Fig. S4†). (B) X-ray diffractograms with referenced peak positions of $\text{MoO}_3 \cdot 2\text{H}_2\text{O}$ (PDF 00-016-0497, asterisk), monoclinic MoO_2 (PDF 00-032-0671, triangle), Mo_2C (PDF 00-011-0680, square), and Mo_4O_{11} (PDF 00-005-0337, circle). (C) Mo 3d, and (D) C 1s X-ray photoelectron spectrum regions of pristine powder MoO_x -C.

layers of molybdenum oxide and carbon alternate throughout the bulk volume of the structure.

The reaction mechanism of alkylamines with molybdic acids can be likened to that in isostructural tungstic acids, which has been described in literature to occur *via* a dissolution–reorganization mechanism that leads to a change in the morphology and particle size.^{28,29} Scanning electron micrographs of $\text{MoO}_3 \cdot 2\text{H}_2\text{O}$ and $\text{MoO}_x\text{-OA}$ (Fig. 1B and C) show an increased particle size after reaction with octylamine. Therefore, we hypothesize that, like in tungstic acids,²⁸ the alkaline conditions upon octylamine addition lead to an initial dissolution of $\text{MoO}_3 \cdot 2\text{H}_2\text{O}$ and a subsequent reassembly of the structure with octylamine in the interlayer space of MoO_3 .

Subsequent pyrolysis of $\text{MoO}_x\text{-OA}$ to form $\text{MoO}_x\text{-C}$ leads to a change in particle morphology (Fig. 1D), most notably, a reduced average particle size from 7.0 ± 2.3 to $1.7 \pm 0.6 \mu\text{m}$ (Fig. S2†), indicating significant densification after pyrolysis.

The Raman spectrum of $\text{MoO}_3 \cdot 2\text{H}_2\text{O}$ (Fig. 2A) shows the typical Raman signals of $\alpha\text{-MoO}_3 \cdot 2\text{H}_2\text{O}$, where the high intensity peaks at 934 cm^{-1} and 727 cm^{-1} are assigned to the stretching modes of $\text{O}=\text{Mo}$ and $\text{O}-\text{Mo}_2$, respectively.³⁰ Peaks in

lower wavenumber ranges, from 400 to 200 cm^{-1} , are assigned to lattice and deformation modes.³⁰ After assembly of OA in the interlayer, a sharp peak appears at 896 cm^{-1} , corresponding to $\text{Mo}-\text{O}$ stretching in $[\text{MoO}_4]^{2-}$,³¹ suggesting the formation of molybdate as a result of OA functionalization.³² The other strong peak at 819 cm^{-1} is characteristic of the doubly-coordinated oxygen ($\text{O}-\text{Mo}_2$) stretching mode.³³ The Raman peak around 2882 cm^{-1} could be assigned to the asymmetric stretching of the methylene (CH_2) group in octylamine.³⁴ The Raman spectrum of the pyrolyzed $\text{MoO}_x\text{-C}$ shows characteristic peaks for MoO_2 according to previous reports (note that a reduced laser power has to be chosen to avoid (re-)oxidation of MoO_2 to MoO_3 during Raman measurement³⁵).³⁶ This suggests that the $\alpha\text{-MoO}_3$ is reduced to MoO_2 during pyrolysis. Successful carbon formation is confirmed by two peaks appearing in $\text{MoO}_x\text{-C}$ at 1604 and 1357 cm^{-1} , corresponding to bond stretching of pairs of sp^2 -carbon (G-mode) and defect-induced breathing modes of sp^2 -carbon (D-mode), respectively.³⁷ In the 2000 to 3000 cm^{-1} range, the absence of the G' band, also called the 2D band, indicates that the formed carbon is highly defective.³⁸

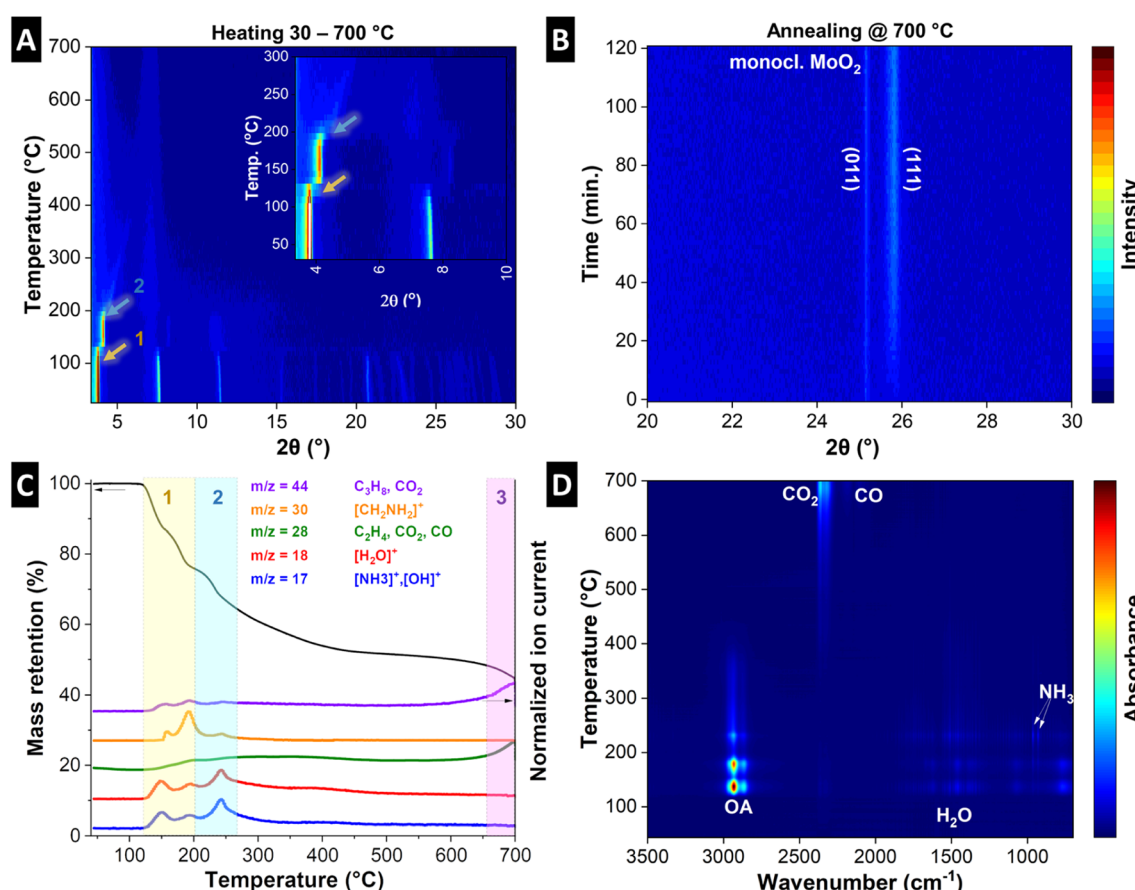


Fig. 3 (A) *In situ* heating XRD patterns of $\text{MoO}_x\text{-OA}$ during the heating step of pyrolysis, with inset showing magnified view of the 2θ range $2\text{--}10^\circ$ and temperature range $30\text{--}300 \text{ }^\circ\text{C}$, (B) *In situ* heating XRD patterns of $\text{MoO}_x\text{-OA}$ during the two-hour annealing step. (C) Thermogravimetric analysis with mass spectroscopy of $\text{MoO}_x\text{-OA}$ during the heating step of pyrolysis, detecting gaseous decomposition products with various mass-to-charge ratios (m/z). Three distinct processes occurring during pyrolysis are graphically indicated by colored mass loss regions (yellow, blue, purple), also highlighted by the arrows in the *In situ* heating XRD diffractograms. (D) TGA with FTIR analysis of gaseous decomposition products during the heating step of pyrolysis.



Fig. 2B shows powder XRD patterns of $\text{MoO}_3 \cdot 2\text{H}_2\text{O}$, $\text{MoO}_x\text{-OA}$, and $\text{MoO}_x\text{-C}$. Diffraction peaks of the as-synthesized molybdcic acid are referenced to $\text{MoO}_3 \cdot 2\text{H}_2\text{O}$ according to PDF 00-016-0497. Upon assembly of OA in the interlayer, a shift of the (020) peak at 12.96° to $3.99^\circ 2\theta$ is observed, corresponding to an expansion of the interlayer, with the *d*-spacing increasing from the initial 0.68 nm in $\text{MoO}_3 \cdot 2\text{H}_2\text{O}$ to 2.21 nm in $\text{MoO}_x\text{-OA}$. Since the octylamine molecule has a chain length of *ca.* 1.2 nm ,³⁹ the large increase in interlayer spacing in $\text{MoO}_x\text{-OA}$ suggests a tilted bilayer arrangement of OA molecules between the MoO_x sheets.²²

After pyrolysis, the XRD pattern of $\text{MoO}_x\text{-C}$ shows peaks matching with monoclinic MoO_2 reflections according to PDF 00-032-0671, in agreement with Raman measurements. Furthermore, low-intensity signals at *ca.* $34^\circ 2\theta$ and $39.4^\circ 2\theta$ suggest that traces of Mo_2C and Mo_4O_{11} were also formed, according to PDF 00-011-0680 and PDF 00-005-0337, respectively.

The oxidation states of Mo in the resulting nanocomposite $\text{MoO}_x\text{-C}$ were investigated by means of XPS. The analysis of Mo 3d is rather complex, and the accurate interpretation is often not well reported. In this case, the Mo 3d (Fig. 2C) region was analyzed following Scanlon *et al.* work, in which the Mo(iv)O₂ consisted of two doublets due to the presence of both screened and unscreened electronic environment of Mo (for more detail see ref. 40).⁴⁰ The Mo 3d region indicates the presence of Mo(iv) and Mo(vi) on the $\text{MoO}_x\text{-C}$, in agreement with XRD. The formation of the latter species is highly common on Mo(iv) oxides.⁴¹ Nevertheless, the $\text{MoO}_x\text{-C}$ consists mainly of MoO_2 (66% *vs.* 33% of Mo(vi)), as also shown in the O 1s spectrum (Fig. S3†). In addition, the C 1s region (Fig. 2D) displays the presence of Mo_2C and C-C, C-O, and O-C=O species, at 283.6, 285.0, and 286.7 and 288.6 eV in agreement with Raman and XRD.^{42,43}

To understand the structural evolution of the material during the pyrolysis, we combined *in situ* heating XRD of $\text{MoO}_x\text{-OA}$ in an inert atmosphere (Fig. 3A and B) with thermogravimetric analysis with simultaneous mass spectroscopy (TGA-MS, Fig. 3C) and Fourier-transform infrared spectroscopy (TGA-FTIR, Fig. 3D). The results indicate that $\text{MoO}_x\text{-OA}$ undergoes three major structural transitions during pyrolysis, which can be conclusively explained in combination with the analysis of the evolving gaseous products *via* MS and FTIR at different temperatures:

Stage 1: a sudden shift of the diffraction peak at *ca.* $3.8^\circ 2\theta$, representative of the interlayer spacing, to a higher diffraction angle (*ca.* $4.1^\circ 2\theta$) can be observed at a temperature of *ca.* 120°C , indicating an initial shrinking of the interlayer spacing. In the same temperature range ($120\text{--}200^\circ\text{C}$), TGA reveals a two-step mass loss of around 24 wt%, coinciding with the evolution of gaseous species. MS signals in profiles of five mass-to-charge ratio (*m/z*) signals are detected, namely *m/z* = 17, 18, 28, 30, and 44. The signals of *m/z* = 17 (OH^+) and 18 (H_2O^+) suggest the desorption of surface and loosely-bound interlayer water molecules.⁴⁴ The *m/z* = 17 signal can also be assigned to ammonia (NH_3^+), which can result from the decomposition of the OA molecule. The peaks in the profile of *m/z* = 30 can be

assigned to the base peak ion $[\text{CH}_2\text{NH}_2]^+$, characteristic of β -cleavage of OA molecule.^{45,46} Finally, the signal at *m/z* = 28 can be attributed to hydrocarbons, CO_2 , or N_2 , and *m/z* = 44 can be attributed to hydrocarbons (C_3H_8) or CO_2 .⁴⁴ FTIR results in Fig. 3D confirm the evolution of mainly molecular OA and water, and small traces of CO_2 and NH_3 as likely OA decomposition product in this temperature range (see the selected FTIR spectra of the pyrolysis gas compared with the typical FTIR spectra of OA, water, NH_3 , and CO_2 in Fig. S5A and B†).⁴⁷ Hence it can be concluded that loss of loosely bonded OA and water likely by evaporation are the main causes of initial interlayer shrinking, while a first onset of OA decomposition can be detected.

Stage 2: above 200°C up to around 350°C , a continuous shrinking of the interlayer spacing to about 1.7 nm (*ca.* $5^\circ 2\theta$) is observed with the intensity strongly reducing before fully disappearing, suggesting complete loss of long-range crystalline order above *ca.* 350°C during pyrolysis. The corresponding TGA in this region up to 350°C detects an accumulated mass loss of *ca.* 43 wt% with simultaneous MS signals at *m/z* = 17, 18, and 44. This indicates further release of water and decomposition products of OA (NH_3 and hydrocarbon fragments). From the FTIR spectra of the pyrolysis gas in this temperature range, the evolution of hydrocarbons is evident from the strong absorption between 3000 and 2800 cm^{-1} , but it cannot be assigned univocally to a specific molecule like in the previous temperature range, due to the lack of significant fingerprints in the $1500\text{--}650\text{ cm}^{-1}$ region of the spectra (see the selected FTIR spectra of the pyrolysis gas in Fig. S5C and D†). The detection of a stronger absorption pattern typical of NH_3 in the same temperature range confirms the decomposition of OA, hence the hydrocarbons detected through FTIR could be regarded as decomposition products. Simultaneously, an increase in the CO_2 signal is detected in FTIR above 350°C (Fig. S5D†), suggesting the onset of (elemental) carbon formation in the material.

Stage 3: towards the end of the heating step (above 650°C) and during the two hours holding step at 700°C in argon (Fig. 3B), the emergence and intensification of two diffraction peaks at 25.8 and $25.2^\circ 2\theta$ can be detected, indicative of the (111) and (011) signals of monoclinic MoO_2 . This can be conclusively linked to the last mass loss step observed above *ca.* 650°C (highlighted in purple, accumulated mass loss of 58 wt%), and an MS signal at *m/z* = 44, which is attributed to CO_2 and indicative of carbothermic reduction of MoO_3 to MoO_2 with carbon acting as reducing agent.⁴⁸ Another strong peak at *m/z* = 28 can be attributed both to CO_2 and CO , and the FTIR spectra at this temperature confirm the simultaneous presence of these two gases (Fig. 3D and S5E†). This is further supported by the detection of MoO_2 after the pyrolysis by Raman results (Fig. 2A), XRD (Fig. 2B), and further confirmed by the electrochemical characterization (*vide infra*).

The stepwise transformations of $\text{MoO}_x\text{-OA}$ during pyrolysis, primarily during the heating step with several structural changes and loss of initial long range order, indicate the formation of an intricate, intraparticle microstructure. Due to the apparent mobility of OA and OA-fragments (*i.e.*, the carbon source) during the heating step, the intraparticle arrangement



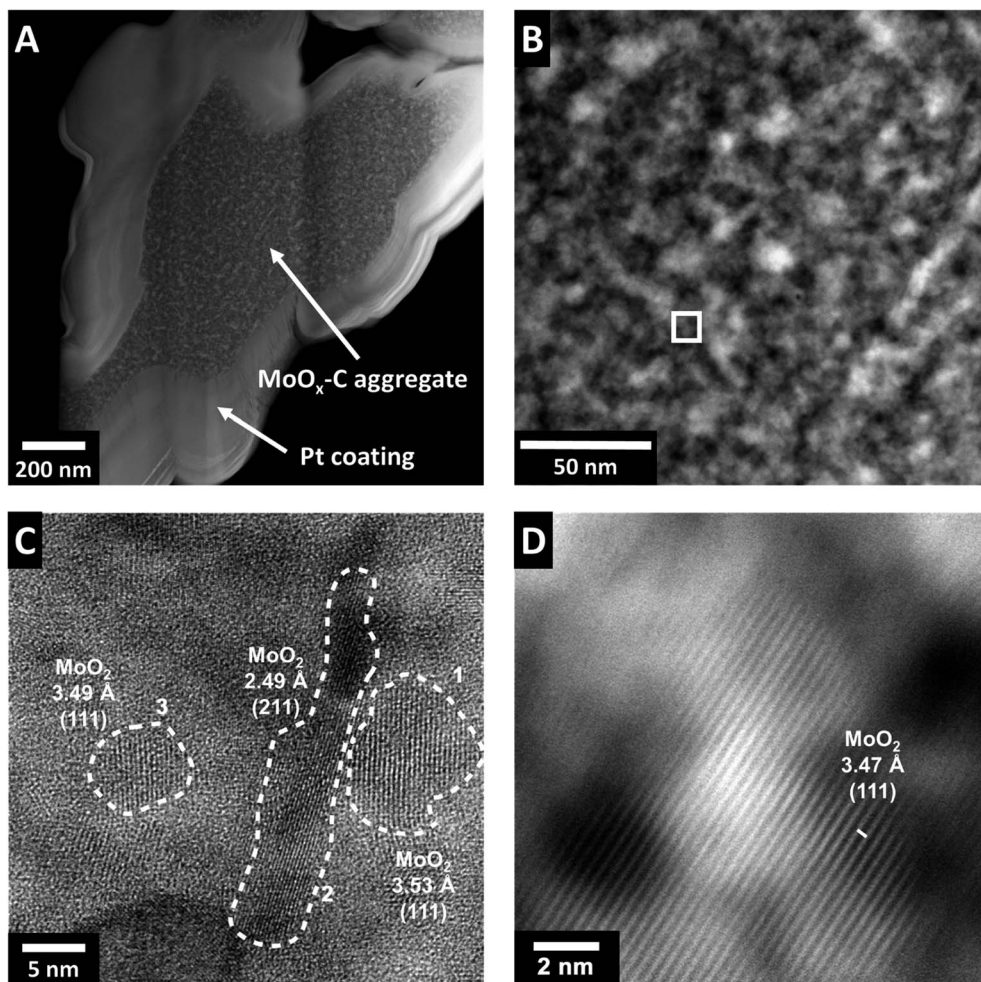


Fig. 4 Cross-sectional overview of the MoO_x-C hybrid nanostructure. (A) HAADF-STEM micrograph of MoO_x-C showing microstructure with non-homogeneous particle composition. (B) Magnified micrograph showing phase Z-contrast variations in MoO_x-C, (C) bright field STEM micrograph taken at the area indicated in (B). The crystalline domains are highlighted and measurements of planar spacings are indicated on the image. Lattice spacings in regions 1, 2, and 3 are in good agreement with the (111), (211), and (111) planes of MoO₂, respectively. (D) High resolution HAADF-STEM image of monoclinic MoO₂ nanodomain.

of the carbon in the nanocomposite product has to be investigated carefully. For this purpose, the microstructural features of MoO_x-C were analyzed using scanning transmission electron microscopy (STEM). Electron-transparent lamellae were prepared *via* focused ion beam (FIB) to observe regions of interest taken from the bulk of MoO_x-C particles (Fig. 4A) to reveal a representative intraparticle structure. STEM images show a difference in the Z-contrast for features with a size of 5–10 nm. This indicates the formation of domains or grains in a size range of 5–10 nm with varying elemental composition, as shown in the zoomed-in view in Fig. 4B. High-resolution STEM micrographs (Fig. 4C and D) show the presence of nanocrystalline domains homogeneously distributed throughout the sample with a size comparable to the sizes of differing Z-contrast. The observation suggests that carbon formation occurred primarily outside the crystalline nanodomains, which contradicts the initially hypothesized formation of a heterostructured, *i.e.*, alternating layer-by-layer arrangement of

molybdenum oxide and the carbon phase. Rather, pyrolysis of MoO_x-OA yields a carbon-containing molybdenum oxide nanocomposite. The lattice fringes in Fig. 4C and D (corresponding HAADF-STEM image in Fig. S6†) show two characteristic distances with slight local variation. The larger one is around 3.5 Å and the smaller around 2.5 Å, in good agreement with the averaged values obtained from XRD for the *d*-spacings of the (111) plane (3.43 Å) and the (211) plane (2.43 Å) of monoclinic MoO₂. Both of these planes account for the most intense reflections on the XRD diffractogram of MoO_x-C (Fig. 2B), hence, the nanodomains observed in Fig. 4C and D can be assigned to monoclinic MoO₂.

To investigate the elemental distribution, we performed energy dispersive X-ray spectroscopy (EDS). HAADF-STEM (Fig. 5A) with EDS elemental mapping of molybdenum, oxygen, and carbon (Fig. 5B–D) were recorded for FIB-prepared MoO_x-C sample cross-sections. The EDS maps reveal an elemental separation with regions enriched in Mo and C and

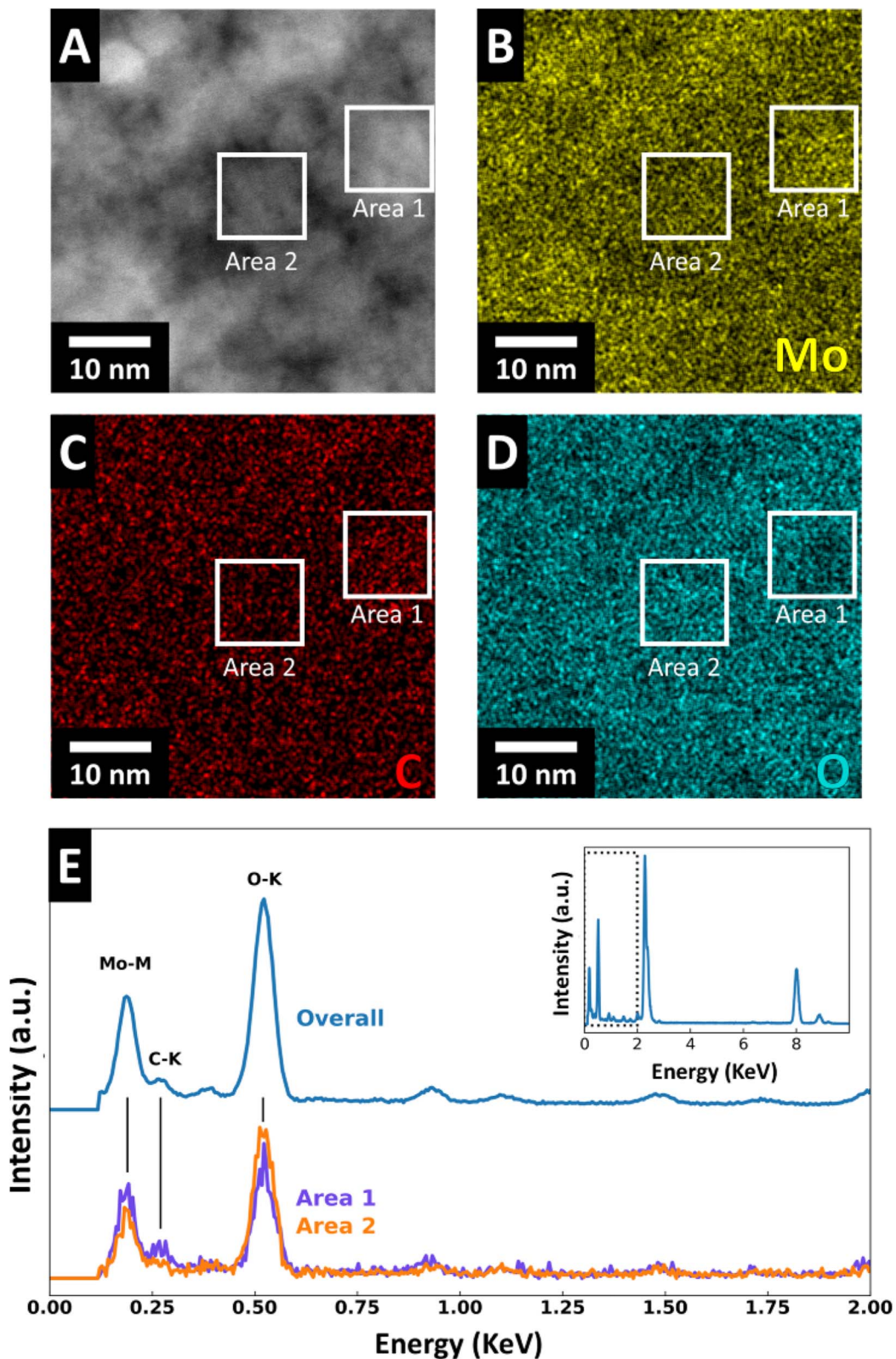


Fig. 5 (A) HAADF-STEM micrograph showing Z-number contrast and (B–D) EDS mapping with elemental distribution of Mo, C, and O in MoO_x –C. (E) Chemical analysis of the two marked areas by quantitative STEM-EDS spectral imaging analysis.

regions enriched in O. The size of these regions corresponds well with the domain/grain size identified in HAADF and high-resolution STEM micrographs (Fig. 4 and S6[†]). Furthermore,

the EDS maps show variations of oxygen- and molybdenum-concentration in the nanosized grains of molybdenum oxide suggesting localized carbothermic reduction of the



molybdenum oxide during pyrolysis. EDS spectra indicate that the carbon-concentration is higher in the areas with lower oxygen concentration (Fig. 5E), suggesting that the carbon formation process and MoO_3 reduction to MoO_2 are correlated. EDS line scan profiles of Mo, O, and C in $\text{MoO}_x\text{-C}$ are shown in Fig. S7,† where local variations in the concentrations of the three elements are shown. TGA measurements in oxygen reveal that the bulk carbon content of $\text{MoO}_x\text{-C}$ nanocomposites is *ca.* 2.7 wt% (Fig. S8†).

3.2 Electrochemical characterization

To understand the implications of the nanocomposite microstructure in $\text{MoO}_x\text{-C}$ on electrochemical reactivity, we tested the nanocomposite as an electrode material in standard organic, lithium-containing electrolyte (LP30). Two different potential windows are chosen to establish a complete picture of the electrochemical performance of the electrode material: according to literature, cycling of molybdenum oxides in a potential range of 1–3 V *vs.* Li^+/Li leads to reversible lithium intercalation according to:²³

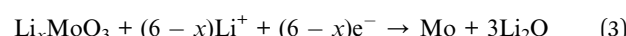


and for MoO_2 :⁸



resulting in a theoretical capacity for the intercalation reaction of 279 and 209 mA h g^{-1} for MoO_3 and MoO_2 , respectively.

After complete lithium intercalation, further extension of the potential range to 0.01 V leads to the conversion-type reaction during the first full reduction according to:



and for MoO_2 :



yielding a theoretical capacity of 1117 and 832 mA h g^{-1} for the combined intercalation and conversion reaction during the first full reduction of MoO_3 and MoO_2 , respectively.^{6,24} We highlight the corresponding processes known from literature in the cyclic voltammograms obtained in the first cycle for the studied materials (Fig. S9†).

For direct comparison, the electrochemical behavior of MoO_{3-x} , (*i.e.*, partially reduced molybdenum oxide that

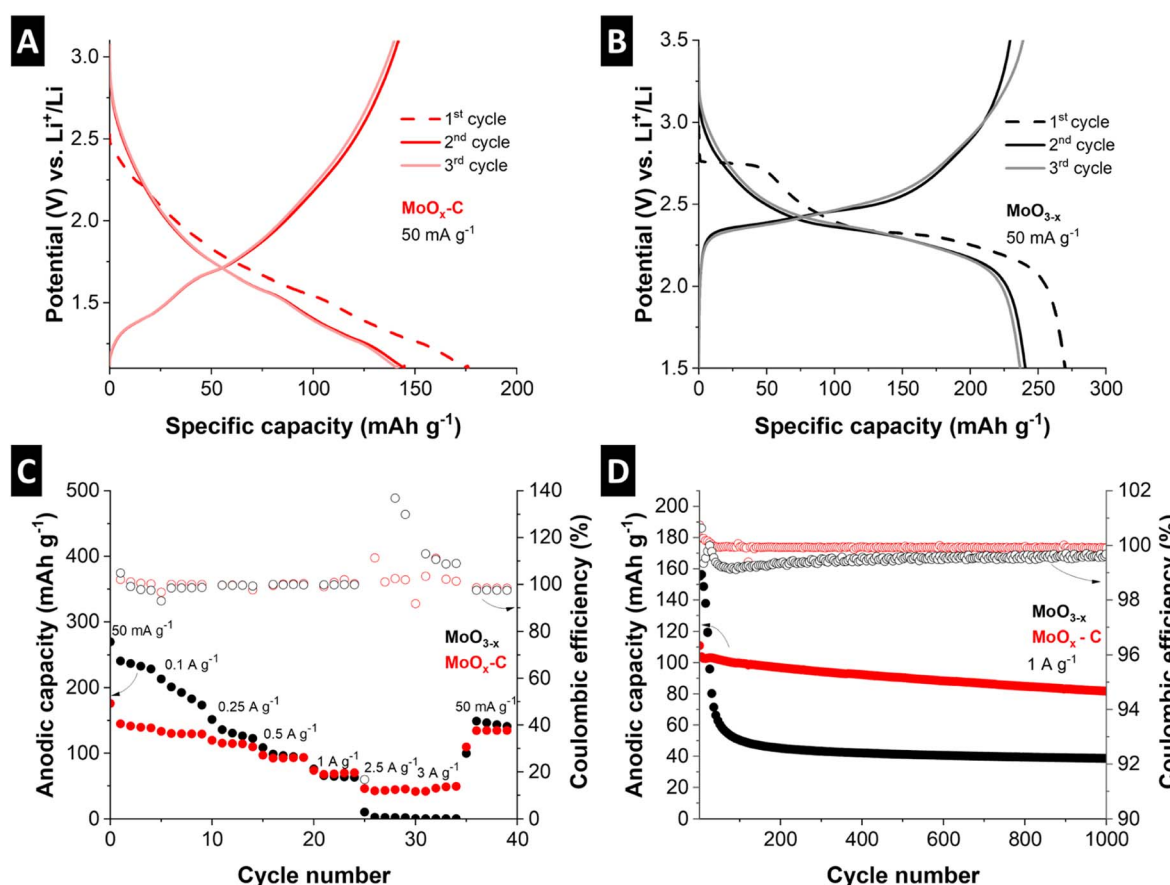


Fig. 6 First three galvanostatic charge–discharge cycles of (A) $\text{MoO}_x\text{-C}$ and (B) MoO_{3-x} at 50 mA g^{-1} , (C) rate capability tests at various specific current values from 50 mA g^{-1} to 3 A g^{-1} , then back to 50 mA g^{-1} , in the voltage range 1.5 V to 3.5 V for MoO_{3-x} and 1.1 V to 3.1 V for $\text{MoO}_x\text{-C}$, (D) cycling stability test for MoO_{3-x} and $\text{MoO}_x\text{-C}$ at 1 A g^{-1} for 1000 cycles.



underwent the same pyrolysis treatment, but without a carbon source, Fig. S10†) was studied alongside $\text{MoO}_x\text{-C}$. This allowed for insights into the influence of the nanocomposite microstructure resulting from the pyrolysis treatment of $\text{MoO}_x\text{-OA}$ on both, the electrochemical lithium intercalation and conversion reactions.

3.2.1 Electrochemical lithium intercalation. The galvanostatic charge–discharge profiles of $\text{MoO}_x\text{-C}$ (Fig. 6A) show Li^+ intercalation with a sloping potential profile below *ca.* 2 V vs. Li^+/Li , characteristic of intercalation in nanosized, monoclinic MoO_2 .⁵ The specific capacity of 145 mA h g^{-1} indicates the intercalation of 0.69 Li per MoO_2 . We hypothesize that the full theoretical capacity of MoO_2 is not reached as a consequence of partial side product formation (Mo_2C , Mo_4O_{11}), as observed by XRD (Fig. 2B). Hence, future work should focus on eliminating side product formation during pyrolysis to achieve specific capacities closer to the theoretical capacity of MoO_2 . For comparison, galvanostatic profiles of MoO_{3-x} (Fig. 6B) show the characteristic plateau of Li^+ (de-)intercalation into the orthorhombic structure of MoO_3 at 2.67 V (anodic) and *ca.* 2.15 V (cathodic),⁴⁹ with a specific capacity of 240 mA h g^{-1} . Electrodes based on MoO_3 often fail to reach full theoretical capacity due to their limited ionic and electronic conductivities.⁵⁰

Fig. 6C shows a comparison of the rate capability of $\text{MoO}_x\text{-C}$ and MoO_{3-x} at current densities ranging from 50 mA g^{-1} to

3 A g^{-1} . Despite the higher initial capacity of MoO_{3-x} , the cycling stability is poor (decrease from 213 mA h g^{-1} after 5 cycles at 100 mA g^{-1}). The anodic capacity continues to deteriorate at high currents. Meanwhile, $\text{MoO}_x\text{-C}$ outperforms MoO_{3-x} at high currents, retaining 41 mA h g^{-1} discharge capacity at 3 A g^{-1} , compared to the negligible 0.6 mA h g^{-1} capacity for MoO_{3-x} at the same specific current.

Fig. 6D shows cycling stability for the two materials cycled at a fixed specific current of 1 A g^{-1} for 1000 cycles. $\text{MoO}_x\text{-C}$ shows an initial anodic capacity of 110 mA h g^{-1} , which stabilizes to 82 mA h g^{-1} after 1000 cycles, retaining *ca.* 75% of the initial capacity. On the other hand, MoO_{3-x} undergoes a rapid capacity decay, with the anodic capacity dropping from 155 mA h g^{-1} after 1000 cycles, a loss of more than 74% of the initial anodic capacity. The averaged coulombic efficiency of MoO_{3-x} , and $\text{MoO}_x\text{-C}$ over 1000 cycles is 99.5%, and 99.9%, respectively, demonstrating the absence of significant side reactions specifically in the nanocomposite electrode $\text{MoO}_x\text{-C}$.

The improved rate handling behavior, as well as the improved cycling stability and coulombic efficiency of $\text{MoO}_x\text{-C}$, can be attributed to two factors. First, the nanocomposite approach leads to a well-dispersed conductive carbon phase. This improves the electronic conductivity, as demonstrated by a significantly lower impedance as compared to MoO_{3-x} electrodes (Fig. S11†). Second, the carbothermic reduction during

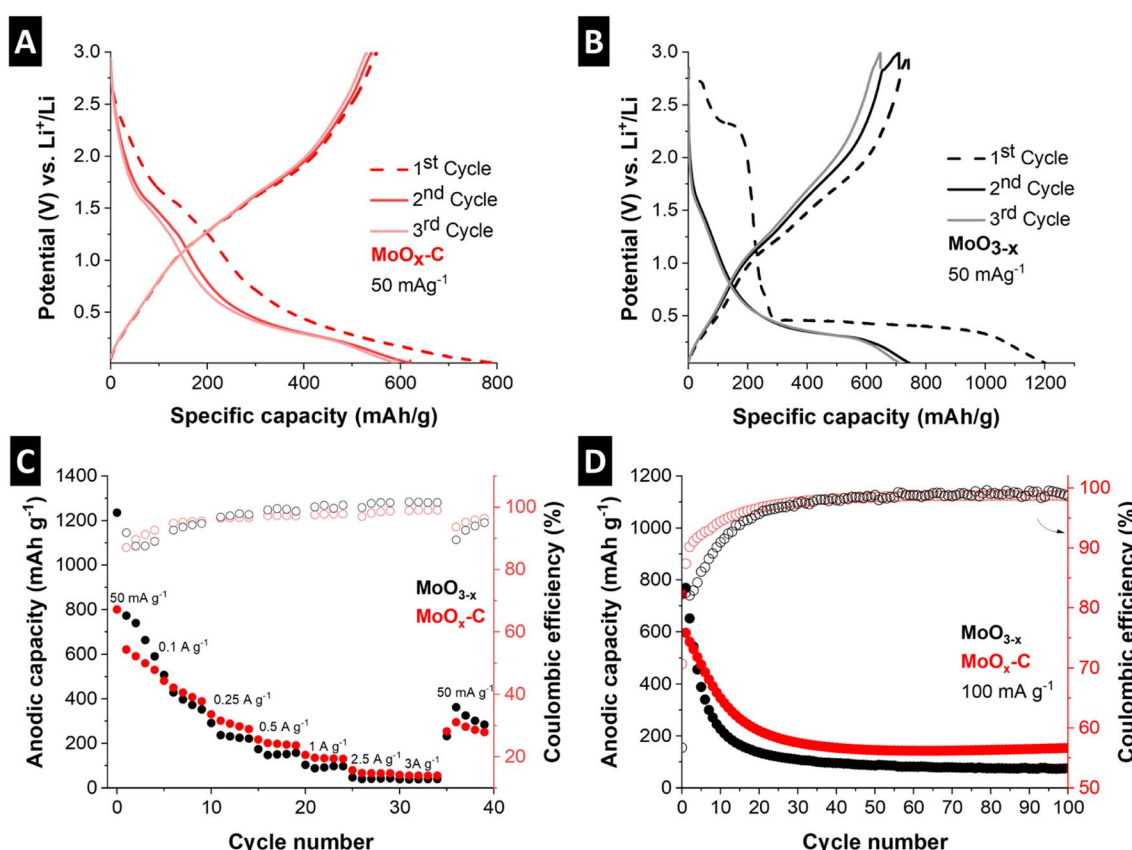


Fig. 7 (A) Galvanostatic charge–discharge curves of $\text{MoO}_x\text{-C}$ and (B) MoO_{3-x} at a specific current of 50 mA g^{-1} in the potential range of 0.01 V to 3 V vs. Li^+/Li , (C) rate performance of MoO_{3-x} and $\text{MoO}_x\text{-C}$ at rates from 50 mA g^{-1} to 3 A g^{-1} , (D) stability test of MoO_{3-x} and $\text{MoO}_x\text{-C}$ cycled at 100 mA g^{-1} .



the pyrolysis process in the nanocomposite led to the formation of MoO_2 , which was shown to exhibit improved cycling stability compared to MoO_3 .⁵¹

3.2.2 Electrochemical conversion reaction. Electrochemical cycling of molybdenum oxides to lower cathodic potential limits can lead to an electrochemical conversion reaction, as described in eqn (3) and (4) and experimentally verified in previous studies,⁶ yielding a significant increase in the specific charge storage capacity. To evaluate the electrochemical behavior, $\text{MoO}_x\text{-C}$ and MoO_{3-x} were galvanostatically cycled between 0.01–3 V vs. Li^+/Li . The occurrence of a conversion reactions forming Li_2O and metallic Mo during the first electrochemical reduction is indicated by the high capacity in the first cathodic cycle (Fig. 7A), according to eqn (3) and (4). The nanocomposite $\text{MoO}_x\text{-C}$ displays a sloping potential profile below 1 V, whereas the MoO_{3-x} sample exhibits a flat plateau around 0.45 V (Fig. 7B). The electrochemical profile of $\text{MoO}_x\text{-C}$ is analogous to previous studies that confirmed a conversion-type reaction of $\text{MoO}_2\text{-C}$ nanohybrids with advanced analytical methods.⁶ Therefore, we hypothesize that the identical conversion-type charge storage mechanism is present in $\text{MoO}_x\text{-C}$ as noted in eqn (4). Notably, the initial coulombic efficiency (ICE) of the nanocomposite $\text{MoO}_x\text{-C}$ (69%) is slightly improved compared to MoO_{3-x} (60%). The anodic capacity in the first three cycles for $\text{MoO}_x\text{-C}$ drops from 800 to 620 mA h g^{-1} , which indicates ~ 3 electron transfer per MoO_2 and amounts for *ca.* 75% of the total theoretical conversion capacity of MoO_2 . For MoO_{3-x} , the anodic capacity drops from 1200 at the initial cycle to *ca.* 700 mA h g^{-1} at subsequent cycles, corresponding to the transfer of 3.76 Li and achieving only 63% of the theoretical conversion capacity of MoO_3 .

The rate handling performance and cycling stability of the electrochemical conversion reaction in $\text{MoO}_x\text{-C}$ and MoO_{3-x} were evaluated by galvanostatic charge–discharge at rates ranging from 50 mA g^{-1} to 3 A g^{-1} and at 100 mA g^{-1} for 100 cycles, respectively (Fig. 7C and D). While the nanocomposite $\text{MoO}_x\text{-C}$ outperforms the reference material, the rate handling and cycling stability performance of both materials is limited. For $\text{MoO}_x\text{-C}$, significant capacity fading can be observed from 768 mA h g^{-1} to 192 mA h g^{-1} after 25 cycles, where the performance stabilizes. For comparison, the capacity of MoO_{3-x} drops to 165 mA h g^{-1} after about 15 cycles. To analyze the improved electrochemical conversion-type charge storage performance of the $\text{MoO}_x\text{-C}$ nanocomposite, the morphology of both electrodes was evaluated *via ex situ* SEM after the first cathodic cycle at 0.01 V vs. Li^+/Li and post mortem SEM after 39 cycles (Fig. S12†). The morphological changes of MoO_{3-x} were more severe compared to $\text{MoO}_x\text{-C}$. This is especially visible by the extensive particle cracking in MoO_{3-x} already occurring after the initial reductive cycle.

Future work optimizing $\text{MoO}_x\text{-C}$ nanocomposite electrodes for conversion-type reactions needs to involve a careful investigation of the electrode microstructure after the initial cathodic cycle, for example *via ex situ* TEM of a FIB-prepared lamella, to gain further insights on the morphological changes occurring during the electrochemical conversion reaction.

4. Conclusion

In this work, layered molybdenum oxide with octylamine molecules assembled in its interlayer was pyrolyzed to convert the organic molecules to a carbon phase, yielding a nanocomposite $\text{MoO}_x\text{-C}$ material. The present work set its main focus on analyzing the carbon formation and microstructure evolution during pyrolysis and the precise phase distribution in the nanocomposite product.

Processes occurring during pyrolysis were examined with a combination of *in situ* XRD, TGA-MS, and TGA-FTIR experiments. It was found that during the heating step, some loosely bound molecular OA and OA decomposition products escape from the interlayer space before being converted to elemental carbon, and a strong carbothermic reduction of the oxide takes place above 650 °C. With this knowledge, future directions towards improving the synthesis route can focus the use of more strongly bonded and/or less volatile organic molecules together with faster heating of the material during pyrolysis to obtain more localized carbon formation or a higher carbon content.

Cross-sectional specimens of the resulting $\text{MoO}_x\text{-C}$ nanocomposite prepared *via* FIB were analyzed to reveal the intricate intraparticle microstructure. It was found that nanoscopic domains of MoO_2 are surrounded by a carbon phase, and in addition, trace amounts of Mo_2C and Mo_4O_{11} formation could be observed by XRD and XPS. Further tuning the synthesis temperature and time could thus be employed to avoid carbide formation (or to support carbide formation, if desirable).

The electrochemical properties of the resulting $\text{MoO}_x\text{-C}$ nanocomposite electrode were evaluated in detail for both, intercalation-type and conversion-type charge storage in lithium-containing, organic electrolyte. The electrode significantly outperformed the MoO_{3-x} reference sample in terms of rate handling and stability for lithium intercalation and showed an increase in initial coulombic efficiency for the conversion reaction.

The work can be used as a guideline to obtain transition metal oxide/carbon nanocomposites with desired structural and electrochemical properties. It showcases pitfalls and opportunities arising from the pyrolysis route (organic molecule mobility, carbothermic reduction, carbide formation, *etc.*) which should be considered in future works.

Conflicts of interest

There are no conflicts of interest to declare.

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