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## **Growth, characterization and performance of bulk and nanoengineered molybdenum oxides for electrochemical energy storage and conversion**

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#### **Abstract**

Molybdenum oxides (MoOy) exhibit quite interesting structural, chemical, electrical, optical and electrochemical properties, which are often dependent on the synthetic procedures and fabrication conditions. While these oxides show promise in a diverse applications ranging from electronics to energy storage and micromechanics, engineering the phase and microstructure is the key to achieving the desired properties and performance, especially in electrochemical energy storage and conversion technologies. In the course of development of lithium batteries delivering high-power and high-energy density for powering electric vehicles, here in this paper, we examine the performances of Mo-oxides, which are candidates as electrodes materials primarily for lithium-ion batteries (LIBs), while some aspects considered in sodium-ion batteries (SIBs) or electrochemical supercapacitors (ECs). Due to the wide range of oxidation states (from  $+6$  to  $+2$ ) they are promising as both positive (cathode) and negative (anode) electrodes of electrochemical cells. Based on their specific structural, chemical, electrical, and optical properties, which are dependent on the growth conditions and the fabrication technique, this review highlights the progress in improving and understanding the electrochemical performance of MoO<sub>y</sub> compounds. Various materials  $(2.0 \le y \le 3.0)$  including anhydrous, hydrates, nanorods, nanobelts, composites and thin films of  $MoO<sub>y</sub>$  are considered. Due to their higher oxidation states,  $MoO<sub>v</sub>$  have been shown to undergo reversible topotactic lithium intercalation reactions; however, electrochemical features appear strongly dependent on the crystal quality and structural arrangement in the host lattice. Several structural characteristics such as in-situ and ex-situ X-ray diffraction (XRD) patterns, Raman spectra are also presented. The approaches used for performance improvement are summarized, and the causes of first-cycle irreversible capacity loss are discussed in detail. Several sub-stoichiometric MoO<sup>y</sup> positive electrodes exhibit excellent cycle life (up to 300 cycles) with high initial coulombic efficiency (80–90%) and large reversible capacity (>300 mAh  $g^{-1}$ ). Molybdenum oxides also categorized as one of the conversion-type transition-metal oxides and applied as negative electrodes for LIBs and SIBs with a specific capacity approaching 1000 mAh  $g^{-1}$ . In addition to the discussion of the key aspects of crystal growth, characterization, and structureproperty relationships, the future prospects to design Mo-oxide materials to enhance the structural stability and electrochemical performance are presented.

#### **1. Introduction**

Nowadays, a great deal of efforts directed towards the green and sustainable energy development in view of the concerns about climate change and global warming. The major contribution comes from human activities for the energy conversion, especially the fossil fuel combustion for electricity, heat and transportation. The fossil fuels consumption for energy production and transporation presumably result in the emission of green-house gases and subsequently causes the global warming effects. Converting global energy consumption through electrification of alternative forms of transportation requires the use of efficient power sources, with an emphasis on hybrid electric (HEVs), plug-in hybrid electric (PHEVs) and fully electric vehicles (EVs) [1]. Other current applications concern batteries used to solve the intermittence problems of the electricity produced by wind or photovoltaic plants, more and more developed as renewable energy sources, and considered as the only solution to give access of electricity to the sub-Saharan countries in Africa, for example. Moreover, economics and environmental incentives, as well as advances in electricity production via renewable intermittent sources are reshaping the traditional distribution network (smart grid environment, vehicle-to-grid (V2G) infrastructure) [2]. High energy density lithium ion batteries (LIBs) and sodium ion batteries (SIBs) are suitable storage systems for such applications.

The concept of LIB (often called a 'rocking-chair battery') was first formulated by Armand in the late 1970s [3]. In a LIB, the lithium ion moved reversibly between the positive and the negative electrode as intercalation materials of different potentials. This important principle was experimentally established by Lazzari and Scrosati over 60 cycles and implemented with a lithiated WO<sub>2</sub> anode and a TiS<sub>2</sub> cathode, which operate in the potential range  $0.8-2.1$  volts [4]. In 1991, Sony Corporation announced the first commercialization of a LIB, which consisted of lithium cobaltate  $LiCoO<sub>2</sub> (LCO)$  as positive electrode and carbon (graphite) as negative electrode in between of which a separator soaked with an organic electrolyte (ionic conductor) [5]. Depending on the choice of the electrodes, the volumetric energy density of LIBs is in the range of 200–350 Wh L<sup>-1</sup> and on average the gravimetric density is 150 Wh kg<sup>-1</sup>. However, two critical issues need to be addressed: performance and safety [6]; also, significant improvements of LIBs must be achieved for vehicular applications in both energy density and rate capability. For an analysis of the LIBs deficiencies and possible remedies, see the report by Blomgren [7]. Briefly, the main deficiencies are: (i) low capacity density of the carbonaceous anode, (ii) formation of a resistive solid-electrolyte interphase (SEI), (iii) low specific capacity of the cathode and (iv) limiting cycle life due to the cathode-electrolyte interphase (CEI). Thus, the replacement with new anode and/or cathode materials is needed that could provide high

coulombic efficiency (CE), good power capability, low irreversible capacity, and low cost with little or no loss of capacity density or cell voltage. Due to its disadvantages of high price, cobalt toxicity and a rather small capacity ( $\approx$ 140 mAh g<sup>-1</sup>) LCO cannot be used as power supply for EVs. The energy density *E* must satisfy the relation  $E = QV$ , which imposes high operating cell voltage (*V*) and large stored capacity (*Q*) delivered to loads. According to the Faraday law  $(Q = nF/M_w)$ , the capacity is determined by the number of transferred electrons (*n*) and the molar weight of the active electrode material  $(M_w)$ . Since, for safety and energetic reasons discussed elsewhere [8], the most feasible methods involve an increase of the capacity and working voltage of the cathode and to avoid the use of graphite as an anode.

Among the transition-metal oxides (TMOs) of diverse compositions and morphological architectures, the various compounds with two- and tri-dimensional structures have been intensively investigated as cathode and anode materials. Molybdenum (Mo) oxides are among those TMOs. Micro- and nano-structured molybdenum oxides (MoO<sub>y</sub>) with  $2.0 \le y \le 3.0$  have shown to be attractive materials in early applications [8]. For example, in 1980s, taking advantages of the superior electrochromic response of  $MoO<sub>3</sub>$  to the other materials, General Motors introduced a rear-view electrochromic mirror in Cadillac cars based on MoO<sub>3</sub> [9,10]. MoO<sup>3</sup> is not only a candidate material for energy storage, batteries and supercapacitors [11-14], but also have potential applications in the fields of display devices [15], gas sensors [16-19], electrochromics [20-22], lubricants [23], field-emission-devices [24], organic solar cells [25], organic-light-emitting-diode (OLED) devices  $[26]$ . MoO<sub>3</sub> also serves as a precursor for preparing a wide variety of nanomaterials indetended for specific electronic and optoelectronic applications  $[27-29]$ . Moreover, MoO<sub>y</sub> are important compounds in selective oxidation catalysis [30-32]. MoO<sup>y</sup> materials are used industrially as catalysts for partial oxidation of hydrocarbons to carbon acids, selective catalytic reduction of NOy, production of acrylonitrile, fabrication of acrylic acid and for olefin metathesis reactions [33].

In addition to diverse crystallography and structural features, Mo-oxides posesses interesting electronic structure. With respect to electronic structure, the Mo-oxide phases can be arranged into five groups:

- (i) The strictly stoichiometric anhydrous MoO<sub>3</sub> polymorphs ( $\alpha$ -,  $\beta$  and  $\epsilon$ -phases), which are nearly insulators (with an unfilled 4d band).
- (ii) The nearly-stoichiometric  $\alpha$ -MoO<sub>3-δ</sub> (with  $\delta$  < 0.03), which retains the orthorhombic symmetry of α-MoO3. This set of materials are unintentionally *n*-type semiconductors at ambient conditions, partly due to intrinsic point defects related to oxygen vacancies  $(V<sub>O</sub>)$  and molybdenum interstitials  $(M<sub>Oi</sub>)$  [34].
- (iii) The defective  $MoO<sub>3-δ</sub>$  (with  $0.03 < δ < 0.11$ ).
- (iv) Mo<sub>n</sub>O<sub>3n-1</sub> suboxides (Magnéli phases,  $n = 4-9$ ), which collapse into ReO<sub>3</sub>-type structures with crystallographic shear planes due to the destabilization of the MoO<sub>3</sub> structure by the loss of oxygen [35-40].
- (v) The hydrated MoO3∙*n*H2O phases [41].

This review is focused on key issues and trends in research on the material science aspects of a variety of Mo-oxides (MoO3, MoOy, MonO3n-1, MoO3∙nH2O, MoO2) used in many advanced systems for energy storage and conversion with a special emphasis toward the compounds that are candidates as positive or negative electrodes in rechargeable lithium and sodium batteries. One of the most interesting properties of  $MoO<sub>v</sub>$  compounds is their intercalation ability, which is possible only if the bonding between the layers or tunnels of the host materials is weak. This technique has been applied to develop nanohybrids and widely used in electrochemical devices and displays because the layered structure facilitates the formation of  $Mo(VI)/Mo(V)$  couple [42]. With the realized property enhancements originating from the incorporation of conducting polymers into layered inorganic hosts, interest in  $MoO<sub>3</sub>$ nanocomposites have attracted great interest. Various combinations of electronic and/or ionic conductive polymers and  $MoO<sub>3</sub>$  have been explored, requiring a wide variety of synthetic methods to effect inclusion of the polymer in the interlamellar gap of the host material [43]. Thus, based on this brief overview, it is very clear that the fundamental understanding of the crystal growth, chemical bonding, structural quality, morphology, surface chemistry, electronic properties of Mo-oxides is critical to tune the synthetic conditions and procedures so as to obtain materials with enhanced electrochemical perfromance. Furthermore, a deeper understanding of the Mo-oxides from a fundamental scientific perspective will enable even design materials with unexpected properties and phenomena, which might be useful for integration into other future applications. Therefore, the synthesis, structural and electrochemical properties of Mo-oxides are thoroughly examined.

This paper is organized as follows. Section 2 is devoted to the properties of anhydrous MoO<sub>3</sub> phases including nanomaterials and MoO<sub>3</sub> used as cathode and anode materials in rechargeable batteries. In Section 3, we expose the properties of MoO3∙*n*H2O hydrates and their derivative products. The non-stoichiometric MoO<sup>y</sup> and oxygen deficient molybdenum oxides  $Mo<sub>n</sub>O<sub>3n-1</sub>$  are evaluated in Section 4. Finally, Section 5 is devoted to the electrochemical features of MoO<sup>2</sup> used as anode in lithium and sodium batteries. In the very section (conclusions), for completeness, a very short panel devoted to discuss the limit of performance of Mo oxides.

#### **2. Anhydrous molybdenum trioxides**

Anhydrous molybdenum trioxides have six polytypic forms. At ambient conditions the thermodynamically stable phase is orthorhombic  $(\alpha$ -MoO<sub>3</sub>), while hexagonal ( $h$ -MoO<sub>3</sub>) and monoclinic (β -MoO<sub>3</sub>, β'-MoO<sub>3</sub>, ε-MoO<sub>3</sub> (MoO<sub>3</sub>-II) and MoO<sub>3</sub>-III) are metastable phases [44]. The crystallographic parameters of anhydrous MoO<sup>3</sup> polymorphs are listed in **Table 1**.

Compound	Space group	Cell parameters (A)			Réf.
		$a/\alpha$	$b/\beta$	$c/\gamma$	
$\alpha$ -MoO <sub>3</sub>	Pbnm	3.962	13.850	3.696	[45]
$\beta$ -MoO <sub>3</sub>	P21/n	7.118	5.366/91.99°	5.568	$[46]$
$\beta$ '-MoO <sub>3</sub>	$P2_1/n$	7.424	7.478/90.09°	7.689	[47]
$h-MoO3$	P63/m	10.57		3.72	[48]
$\epsilon$ -MoO <sub>3</sub>	P21/m	3.954	$3.687/103.75^{\circ}$	7.095	[49]

**Table 1**. Crystallographic parameters of the anhydrous molybdenum trioxides.

#### 2.1. Orthorhombic α-MoO<sup>3</sup>

Molybdenum trioxide exists as mineral *molybdite* (molar mass 143.94 g mol<sup>-1</sup>, density 4.70  $g \text{ cm}^{-3}$ , melting point 802 °C).

#### *2.1.1. Brief history*

It was Carl Wilhem Scheele who, in 1778, identified molybdenum by separation from lead and isolated the oxide  $MoO<sub>3</sub>$  and the molybdenite,  $MoS<sub>2</sub>$ . In 1861, Adolf Nordenskiold synthesized a MoO<sub>3</sub> crystal by sublimation of ammonium molybdate and determined the crystal axial ratio as *a*:*b*:*c* = 0.3872:1:0.4792. In 1931, it was Haakon Bräkken and Nora Wooster, who reported the crystal structure of MoO<sub>3</sub>; the works carried out independently obtained fairly similar results [50,51]; a new axial ratio of 0.282:1:0.265 was determined by X-ray measurements. In 1940s, the Uppsala's group, by means of X-ray methods, revised the crystallographic structure of all forms of molybdenum oxides [52]. In 1950, Oskar Glemser published a series of papers on Mo oxides investigated by X-ray powder technique and electrical resistivity measurements [53]; they also formulated a series of molybdenum oxide hydroxides  $[54]$ . Since 1956, MoO<sub>3</sub> has been the subject of numerous studies as a host structure for the intercalation of various atoms or molecules, which lead in particular to molybdenum

bronzes  $H_xMOO_3$  [55]. Since then, a large number of works were devoted to the study of insertion in aqueous environments and in  $H_2SO_4$  [56]. In the 1960s, Morehouse Glicksman [57] investigated the discharge characteristics of  $MoO<sub>3</sub>$  in strongly alkaline aqueous electrolyte. In the 1970s, several attempts made to use  $MoO<sub>3</sub>$  as a battery electrode [58-60]. The electrochemical behavior of  $MoO<sub>3</sub>$  as positive electrode material in organic solvent lithium batteries was investigated for the first time by Campanella and Pistoia [58,59], who recognized the poor solubility of  $MoO<sub>3</sub>$  in aprotic solvent and reported the open circuit voltages (OCV) of the Li(s)|Li<sup>+</sup>(nonaqueous solvent|MoO<sub>3</sub>(s) of ~2.8 V. Further, during the early years of lithium battery research, MoO<sub>3</sub> was actively investigated as lithium insertion compound  $[42,61-65]$ . In 1974, the growing interest in lithium anode batteries has led to studies on the properties of  $MoO<sub>3</sub>$  in an aprotic medium [66,67]. The lithiated phase of  $MoO<sub>3</sub>$  was synthesized both by reaction of n-butyllithium and electrochemically. The maximum  $x$  value in  $Li_xMoO_3$  obtained by chemical lithiation is 1.55 [62]. Besenhard and Schöllhorn [61] investigated the charge and discharge process of Li//MoO<sub>3</sub> cells using different aprotic electrolytes and have shown the process to be a reversible "*topotactic*" redox reaction to form the ternary phase  $Li_xMoO_3$ , which exhibits an increase of the interlayer spacing of  $\sim$ 1 Å. The term "topotactic" refers to solid-state reactions, which are characterized by the fact that defined lattice matrix elements specific to the original solid phase remain unaltered in the course of the reaction with respect to their structural architecture [68]. The Hagenmuller's group reported that the high-temperature lithium molybdenum bronzes monoclinic  $Li_xMoO_3$  (0.31  $\leq x \leq 0.39$  at 560 °C) and  $Li_xMo_6O_{17}$  ( $x=0.9$ ) display structures different than that of  $\alpha$ -MoO<sub>3</sub> itself [69].

#### *2.1.2. Structure*

Of the anhydrous  $MoO<sub>3</sub>$  polymorphs, the orthorhombic α-Mo $O<sub>3</sub>$  phase is the stable form at room temperature and possesses a 2D layered structure  $[70]$ . The  $\alpha$ -phase MoO<sub>3</sub> can be represented as the double layers of linked distorted  $MoO<sub>6</sub>$  octahedra parallel to (010) planes (Fig. 1). In each double layer  $MoO<sub>6</sub>$  octahedra form edge sharing zigzag rows along the (001) direction and corner haring rows along the (100) direction. Successive layers are held together by weak interlayer forces with an easy cleavage along the (010) planes (**Fig. 1a**). The octahedral coordination of Mo with O characterizes the geometrical structure of MoO<sup>3</sup> with Mo surrounded by four close neighbor O atoms at distances of 1.95, 1.95, 1.73, and 1.67 Å and two O atoms at significantly longer distances of 2.25 and 2.33 Å. These values indicate that the MoO<sub>6</sub> octahedra are distorted and three different types of oxygen neighbors are present in the coordination sphere of Mo. They are termed as terminal (non-bridging)  $O(1)$ , bridging  $O(2)$ ,

and chain oxygens O(3) (see **Fig. 1c**). The terminal oxygen O(1) is doubly bound to a single Mo at a short bond length of 1.67 Å.  $O(2)$  is two-fold coordinated and the Mo–O bonds with bond lengths of 1.79 and 2.25 Å characterize the bridging oxygen located asymmetrically between two Mo centers. The symmetric bridging oxygen O(3) is bound to three Mo atoms, with two equivalent Mo–O bonds (1.94 Å) and a longer Mo–O bond (2.33 Å) [35]. The large work function of  $MoO<sub>3</sub>$  of 6.6 eV is due to its closed shell character and the dipole layer created by planes of terminal O(1) oxygen sites which lower the electrostatic potential of the inner Mo-O units. These  $O(1)$  sites arise from the high stoichiometry of MoO<sub>3</sub>. The oxygen vacancy is most stable at the 2-fold O(2) site. It is a shallow donor and has a small formation energy in the O poor limit so that MoO<sup>3</sup> easily becomes a degenerate semiconductor [71,72]. **Figure 1d** displays the XRD pattern of the layered  $\alpha$ -MoO<sub>3</sub> showing fingerprints at  $2\theta = 12.7^{\circ}$ , 23.3°, 25.6°, 27.2°, and 38.9°, which correspond to the  $(020)$ ,  $(110)$ ,  $(040)$ ,  $(0.21)$  and  $(060)$  planes (JCPDS card no. 05-0508, S.G.), and crystallizes in the unique orthorhombic crystal structure (space group *Pbnm*) with lattice unit-cell constants  $a = 3.962 \text{ Å}$ ,  $b = 13.858 \text{ Å}$  and  $c = 3.697 \text{ Å}$ . Unit cell parameters of  $\alpha$ -MoO<sub>3</sub> investigated through 80 years are given in **Table 2**. [45,52,73].

**Table 2**. Unit cell parameters of α-MoO<sup>3</sup> (*Pbnm* S.G.) through 80 years.

Authors (year)	a(A)	b(A)	c(A)	$V(A)^3$	Ref.
Bräkken (1931)	3.962	13.855	3.701	203.1	$[73]$
Wooster $(1931)$	3.93	13.97	3.67	201	[74]
Magnéli (1950)	3.966	13.85	3.696	203.0	$[52]$
Kilborg $(1963)$	3.9628	13.855	3.6964	202.9	[45]
Sitepu $(2005)$	3.9621(7)	13.855(1)	3.6986(4)	203.0	[75]





**Fig. 1**. Structural model for MoO<sup>3</sup> α-phase (a) The unit cell includes four formula units located on double-layer slabs along the (010) direction. (b) The (010) projection of an individual double-layer. (c) The three different types of oxygen neighbors present in the coordination sphere of molybdenum. The Mo-O framework consists of  $MoO<sub>6</sub>$  octahedra sharing edges (O2-O2'). (d) XRD pattern of orthorhombic  $\alpha$ -MoO<sub>3</sub> (white crystal). (e) Raman spectrum recorded with the excitation laser line  $\lambda = 532$  nm.

The local structure was investigated using Raman scattering (RS) and Fourier transform infrared (FTIR) spectroscopy, which are effective tools to study the structure of crystalline, disordered and amorphous substances. The vibrational spectrum of a molecule, as observed in Raman studies, is the direct manifestation of forces arising from mutual interactions of electrons and nuclei and the RS spectrum can be analysed in terms of internal and external mode. Thus, from the observed mode frequencies, we can glean information about the force constants which hopefully can then be correlated with electronic structure and bonding theories. Several reports have been devoted to the study of vibrational features of MoO<sup>y</sup> materials [41,76-80]. **Figure 1f** shows the RS spectrum of crystalline  $\alpha$ -MoO<sub>3</sub>. The observed bands are assigned according to the single crystal model of Py and Masche [76]. The Raman spectrum of α-MoO<sub>3</sub> single crystal has characteristic bands at 996 cm<sup>-1</sup> ( $A_g$ ,  $v_{as}$  M=O stretch), 823 cm<sup>-1</sup> ( $A_g$ ,  $v_s$  M=O stretch), 667 cm<sup>-1</sup> ( $B_{2g}$ ,  $B_{3g}$ ,  $v_{as}$  O–M–O stretch), 473 cm<sup>-1</sup> ( $A_g$ ,  $v_{as}$  O–M–O stretch and bend), 380 cm<sup>-1</sup> ( $B_{1g}$ ,  $\delta$ O–M–O scissor), 376 ( $B_{1g}$ ), 366 cm<sup>-1</sup> ( $A_{1g}$ ,  $\delta$  O–M–O scissor), 334 cm<sup>-1</sup> ( $A_g$ ,  $B_{1g}$ ,  $\delta$ O–M–O bend), 293 cm<sup>-1</sup> ( $B_{3g}$ ,  $\delta$  O=M=O wagging), 285 cm<sup>-1</sup> ( $B_{2g}$ ,  $\delta$  O=M=O wagging), 247 cm<sup>-1</sup> ( $B_{3g}$ ,  $\tau$  O=Mo=O twist), 216 cm<sup>-1</sup> (A<sub>g</sub>, rotational rigid MoO<sub>4</sub> chain mode, R<sub>c</sub>), 197 cm<sup>-1</sup> (B<sub>2g</sub>,  $\tau$ O=Mo=O twist), 159 cm<sup>-1</sup> ( $A_g/B_{Ig}$ , translational rigid MoO<sub>4</sub> chain mode,  $T_b$ ), 129 cm<sup>-1</sup> ( $B_{3g}$ , translational rigid MoO<sub>4</sub> chain mode,  $T_c$ ), 116 cm<sup>-1</sup> ( $B_{2g}$ , translational rigid MoO<sub>4</sub> chain mode,  $T_c$ ), 100 cm<sup>-1</sup> ( $B_{2g}$ , translational rigid MoO<sub>4</sub> chain mode,  $T_a$ ) and 89 cm<sup>-1</sup> ( $A_g$ , translational rigid MoO<sub>4</sub> chain mode,  $T_a$ ). The Raman bands at 995 and 823 cm<sup>-1</sup> are the fingerprints of the symmetrical stretching vibration of the terminal Mo=O bonds along the *a*- and *b*-axes (**Table** 

). The bridging oxygens (longer bond distances) along the *c*-axis are the weakly bound oxygens as shown by Mestl et al. [81]. Due to the stoichiometry deviation within the  $MoO<sub>v</sub>$ samples variations of the Raman bands of localized vibrations, e.g., the 666, 823 or 995 cm<sup>-1</sup> bands are expected as a displacement of the Mo atom toward the terminal oxygen in the *b*direction occurs upon the loss of the bridging oxygen, thus weakening the bond to the terminal O atom along the *a*-axis.

<b>Modes</b>	<b>Assignment</b>	Raman shift $(cm-1)$					
		$[82]$	$[44]$	$[78]$	[83]	[84]	$[41]$
$A_g$	$T_a$ Translation of rigid MoO <sub>4</sub> chain $(T_a)$	82		89	82	82	83
$B_{2g}$	$T_a$ Translation of rigid MoO <sub>4</sub> chain $(T_a)$	98	$\overline{\phantom{a}}$	100	$\overline{\phantom{a}}$	98	98
$B_{2g}$	$T_c$ Translation of rigid MoO <sub>4</sub> chain $(T_c)$	116	116	116	115	114	116
$B_{3g}$	$T_c$ Translation of rigid MoO <sub>4</sub> chain $(T_c)$	127	130	129	129	128	128
$A_g/B_{1g}$	$\delta$ (O <sub>2</sub> Mo <sub>2</sub> ) <sub>n</sub>	159	157	159	158	157	156
$B_{2g}$	$\delta O_2$ Mo $O_2$ scissoring	196	198	197	198	197	198
$A_{g}$	$\delta O_2$ Mo $O_2$ scissoring	218	217	216	216	216	218
$B_{3g}$	$\delta O_2$ Mo $O_2$ scissoring	245	246	247	246	244	246
$B_{2g}$	$\delta O_1$ = Mo = O <sub>1</sub> wagging	283	284	285	284	282	286
$B_{3g}$	$\delta O_1$ = Mo = O <sub>1</sub> wagging	292	291	293	÷,	÷,	291
$A_g/B_{1g}$	$\delta O_3$ Mo O <sub>3</sub> bending	338	338	334	336	336	338
$A_{g}$	$\delta O_2$ = Mo = O <sub>2</sub> scissoring	365	365	366	365	365	366
$B_{1g}$	$\delta O_2$ = Mo = O <sub>2</sub> scissoring	379	378	376	379	377	380
$A_{g}$	$v_{\text{as}}$ Mo__O <sub>2</sub> __Mo bending	471	471	473	÷,	469	472
$B_{2g}/B_{3g}$	$v_{\text{as}}$ Mo $\_\_O_2$ Mo stretching	664	667	667	666	666	666
$A_{g}$	$v_s$ Mo $\_\_O_3\_\_$ Mo stretching	816	817	823	819	818	820
$A_{g}$	$v_{\rm as}$ Mo $=$ O <sub>1</sub> stretching	992	996	996	996	994	996

**Table 3**. Raman vibration modes of nanostructured α-MoO3. Reproduced with permission from [82]. Copyright 2012 Elsevier.

#### *2.1.3. Synthesis*

A variety of synthesis routes have been developed for the preparation of  $MoO<sub>y</sub>$  materials to control their structure and morphology. They include solid state reaction [85], wet-chemical synthesis [86], sol-gel route [20], precipitation [87], thermal oxidation [88], thermal evaporation [83], flame synthesis process [89], hot-plate method [90], mechanochemical technique [91], electrodeposition [92], electrospinning technique [93,94], solvothermal method [95], spray pyrolysis [96-98], spin coating [99], and template synthesis [100]. Synthesis techniques are briefly reviewed in this section; for a complete view see Ref. [101]. Most of crystallized MoO<sup>y</sup> were easily fabricated by heat treatment of either molybdic acid powders  $(H_4MoO_4)$  or synthetic hydrated molybdenum trioxide  $(MoO_3 \cdot nH_2O)$  [102,103].

Different hydrothermal techniques were utilized to grow  $\alpha$ -MoO<sub>3</sub> crystals (see **Table 4**) [104-107]. Various raw materials are commonly used including Mo metal powders, acid molybdic (H4MoO4), ammonium molybdate ((NH4)2MoO4), ammonium molybdate tetrahydrate  $((NH_4)_6Mo_7O_{24} \cdot 4H_2O)$ , and sodium molybdate  $(Na_2MoO_4)$ . Zhou et al. [108] fabricated flexible single crystalline nanobelts with widths of 200-500 nm, lengths of 5-10 *μ*m, and thicknesses of ∼50 nm via hydrothermal treatment at 180 °C for 24 h of a peroxomolybdic acid solution prepared by reaction of  $H_2O_2$  with Mo metal powders. Li et al. [93] prepared MoO<sub>3</sub> nanofibers (diameter of 200 nm) by a combination method of sol–gel process and electrospinning technique using  $(NH_4)_6M_9T_9Q_{24} \cdot 4H_2O$  with the molar ratio of 1/1.28 dissolved in dilute nitric acid and then mixed with a water–ethanol  $(v/v 1:4)$  solution containing citric acid as a chelating agent for the metal ions. Subba-Reddy et al. [106] constructed nanobelts (1– 5  $\mu$ m in length and 100–600 nm in diameter) by hydrothermal method using MoO<sub>3</sub>·*n*H<sub>2</sub>O sols prepared by ion exchange of ammonium heptamolybdate tetrahydrate through a proton exchange resin. After ion exchange, a clear light-blue MoO<sub>3</sub> sol (pH  $\approx$  2.0) was obtained and modified with poly(ethylene glycol) (PEG). Nanocrystalline  $MoO<sub>3</sub>$  was synthesized by a solgel method using ammonium molybdate tetrahydrate citric acid (CA) as chelating agent with the metal/CA ratio of 1:1 under neutral pH  $[109]$ . Pure MoO<sub>3</sub> was obtained by calcination of dried precursor at 500 °C for 5 h in air, while  $Mo<sub>n</sub>O<sub>3n-1</sub>/C$  composite was formed under argon atmosphere (**Fig. 2**).



**Fig.** 2. Overall synthesis process of the crystalline MoO<sub>3</sub> and Mo<sub>m</sub>O<sub>3m−1</sub>/C composite. Copyright Hashem et al. [109]. Copyright 2020 Hashem et al. Under Creative Commons Attribution (CC BY) license.

Raw material	Additive	Conditions	Ref.
commercial MoO <sub>3</sub> bulk	CTAB <sup>a</sup>	180 $\degree$ C/4 days	$[110]$
Mo metal powders	$H_2C_2O_4.2H_2O$	$180 °C/5$ days	[111]
Na <sub>2</sub> MoO <sub>4</sub>	$0.2$ mmol HCl	160 °C/20 h	[112]
$(NH_4)_6Mo_7O_{24}\cdot 4H_2O$	$HNO3$ (pH=4.5-9-6)	140-220 °C/5-62 h	[105]
$(Mo(NO)(NH2O))^{2+}$	HNO <sub>3</sub>	$220 °C/7-24 h$	[113]
Mo metal powders	$H_2O_2$	180 °C/24 h	[108]
Na <sub>2</sub> MoO <sub>4</sub>	4 mmol $L^{-1}$ HClO <sub>4</sub>	140 °C/24 h	[114]
$Na2MoO4·2H2O$	3-diaminopropane	120 °C/48 h	[115]
$Na2MoO4·2H2O$	$NaCl + GO$	180 °C/24 h	[116]
$H_2MO_4$	$HNO3/H2O2$ (1:5 v/v)	120-180 °C/6 h	[117]
$MoO2$ powders	$H_2O_2$	180 °C/10 h	[118]
$MoO3·nH2O$ sols	poly(ethylene glycol)	$180 °C/4$ days	[106]

Table 4. Synthesis of α-MoO<sub>3</sub> through different hydrothermal routes.

a) cetyltrimethylammonium bromide.

The procedure developed by Jiao et al. [119] consists in the wet-chemical synthesis using an ammonium molybdate tetrahydrate solution adjusted at  $pH = 3.5$  with diluted glacial acetic acid with subsequent sintering at 600 °C for 24 h. The conversion of isopolymolybdate anions  $Mo<sub>7</sub>O<sub>24</sub><sup>6–</sup>$  to neutral  $Mo<sub>7</sub>O<sub>21</sub>$  (or α-MoO<sub>3</sub>) occurs by combining excess oxygen in Mo<sub>7</sub>O<sub>24</sub> with protons from the acidic medium:

$$
Mo_{7}O_{24}^{6-} + 6H^{+} \rightarrow 7MoO_{3} + 3H_{2}O. \tag{1}
$$

Chiang et al. [120] investigated the precipitation route using the mixture of an ammonium molybdate tetrahydrate solution with ethylene glycol (dark blue solution), which precipitates after centrifugation and heating at 80 °C for 24 h (white powder). In a typical solid-state reaction, α-MoO<sub>3</sub> crystals obtained from the decomposition of  $H_4MO_4$  raw powders loaded into porcelain crucibles and positioned in a multi-zone tube furnace with a  $300 \text{ cm}^3 \text{ min}^{-1}$  argon flow. After 2-h purge, the furnace temperature is ramped from room temperature to the desired heat-treatment temperature over a 1-h period. This thermal treatment maintained at 750 °C for 2 h provides well-crystallized specimens [121]. **Figure 3** shows the SEM images of samples produced by heat treatment of molybdic acid at temperature  $400 \leq T_a \leq 800$  °C. Hydrates contain  $MoO<sub>5</sub>(H<sub>2</sub>O)$  octahedral units, while they are intercalated by water molecules in  $MoO<sub>3</sub>·2H<sub>2</sub>O$ . During the topotactic dehydration process, the intercalated water is lost first, then neighbouring sheets condense giving the double sheets of the anhydrous  $MoO<sub>3</sub>$  phase as the final water is lost (see sketch of **Fig. 4a**). TG and DTA curves of the oxide-hydrate of molybdenum  $MoO_3 \cdot 2H_2O$ shown in Fig. 4b exhibit four endothermic peaks at annealing temperature  $(T_a)$  of 120, 198, 246, and 325 °C corresponding to the various steps for water removal via the formation of the intermediate α-MoO3·H2O. **Figure 4c** presents the XRD patterns of oxide-hydrates heat-treated

in the temperature range 75-800 °C. The phase  $\alpha$ -MoO<sub>3</sub> appears after a thermal treatment of the molybdic acid above 750 °C in air. FTIR spectra of  $MoO<sub>3</sub>·nH<sub>2</sub>O$  ( $n<1$ ) shows the spectral features of the layered α-MoO<sub>3</sub>·H<sub>2</sub>O phase for  $T_a \approx 280$  °C. Note the disappearance of the OH stretching mode (3400 cm<sup>-1</sup>) for  $T_a$ >250 °C (**Fig. 4d**). The low-temperature synthesis of MoO<sub>3</sub> nanobelts (15 nm thick) was based on the preparation of  $MoO<sub>3</sub>·H<sub>2</sub>O$  solution (i.e., colorless solution obtained by dissolution of 20 mmol  $Na<sub>2</sub>MoO<sub>4</sub>$  in distilled water added with 4 mol  $L^{-1}$  perchloric acid and subsequent treatment of a hydrothermal reaction [114].



**Fig.** 3. SEM images of samples produced by heat treatment of  $MoO<sub>3</sub>·2H<sub>2</sub>O$  hydrate at (a) 400, (b) 600, (c) 750 and (d) 800 °C. The latter sample is the well crystallized  $\alpha$ -MoO<sub>3</sub> phase, which displays the typical elongated shape the layered structure (edge of the platelet).



**Fig. 4**. (a) Sketch of the structural evolution upon heat-treatment of the hydrated MoO<sub>3</sub>·2H<sub>2</sub>O precursor to form α-MoO<sub>3</sub> via the intermediate α-MoO<sub>3</sub>·H<sub>2</sub>O phase. (b) TG and DTA curves for oxide-hydrate of molybdenum. The DTA curve exhibits four endothermic peaks at 120, 198, 246, and 325 °C corresponding to the various steps for water removal. (c) X-ray diffraction patterns of oxide-hydrates of molybdenum heat-treated at various temperature in the range 75- 600 °C. The phase α-MoO<sub>3</sub> appears after a thermal treatment of the molybdic acid above 750 <sup>o</sup>C in air. (d) FTIR spectra of dehydrated MoO<sub>3</sub>· $nH_2O$  ( $n<1$ ). Spectral responses are more complex than that of crystalline MoO3, but they can be discussed in terms of internal and external modes as usual for oxide lattices.

#### *2.1.4. MoO<sup>3</sup> nanostructures*

One major reason for the use of nanosized particles of materials for energy storage comes from their poor transport properties that imply poor rate performance of the electrochemical devices. This is the case for  $MoO<sub>3</sub>$  used as electrodes in batteries and supercapacitors. As the electronic conductivity of MoO<sub>3</sub> is  $\approx 10^{-8}$  S·cm<sup>-1</sup> at room temperature, it requires some sophisticated technology such as the use of slurry containing a huge quantity of carbon (carbon "Super P", acetylene black, amorphous carbon, etc.) or deposition at the surface of the grains for enhanced charge carrier transport [122-126]. Achieving high-rate capability depends ultimately on the geometry of the active objects building the positive and negative electrodes.

As the performance of an electrode is governed by the transport of both electrons and ions, the ionic and electronic conductivity of the materials must be considered. For a diffusion process, the characteristic time  $\tau$  for ionic species *i* (in practice Li<sup>+</sup> or Na<sup>+</sup> ions in the present case) to reach the surface of any active particle of dimension *L* is given by the second Fick's law that applies the chemical diffusion coefficient with  $D_{Li}^*$  of moving ions [127]. In the case when the chemical reaction proceeds by a single-phase process, i.e., within a solid solution, *τ* is given by  $\tau = L^2/4\pi D_{Li}^*$ . Moreover, decreasing the size of MoO<sub>3</sub> particles is an efficient strategy to mitigate the volume variation of the electrode during Li ions intercalation and deintercalation process. Nanosized MoO<sup>3</sup> particles with different morphologies (**Fig. 5**) include nanoparticles [128], nanobelts [108,129,130], nanowires [131-34], nanofibers [113,135], nanotubes [136-138], nanoflakes [90] and nanoribbons [18]. Designing of porous structure and carbon coating are another most popular strategy to improve the electrochemical performances of MoO<sub>3</sub> [139,140].



**Fig. 5**. SEM images of nanostructured a-MoO3. (a) nanofibers, (b) HRTEM image showing the lattice fringe of 0.38 nm which corresponds to the  $d_{110}$  spacing, (c) nanoplatelets, (d) nanospheres, (e) nanoflakes, (f) nanorods, (g) nanosheets, (h) nanoribbons, (i) nanotubes.

#### *2.1.5. MoO<sup>3</sup> thin films*

Due to their chromogenic, catalytic properties and ability to host foreign ions,  $MoO<sub>3</sub>$  thin films have been applied in electrochromic display devices (see  $[141]$  and Refs. therein), optical memory devices [142], gas sensors [17], and lithium microbatteries [143**-**146]. Currently, there are several approaches available for preparing  $MoO<sub>3</sub>$  films with either an amorphous or a polycrystalline structure using physical vapor deposition (PVD) and metal organic chemical vapor deposition (MOCVD) [147] are also used to fabricate MoO<sub>3</sub> in thin film forms. The different techniques include thermal evaporation [148], magnetron sputtering [144,149], chemical vapor deposition [150,151], electrodeposition [152], flash evaporation [153,154], pulsed-laser deposition (PLD) [155-157], electron-beam deposition [158,159], ion-beam deposition  $[17]$ , sol-gel spin-coating  $[160,161]$ , and spray pyrolysis  $[162,163]$ . Among these methods, magnetron sputtering and pulsed-laser deposition have received considerable attention for industrial purpose because the low temperatures process  $[164]$ . MoO<sub>3</sub> films were produced under varying conditions of growth temperature,  $T_s$ , and oxygen pressure  $p(O_2)$ . Analyses indicate that the microstructure of  $MoO<sub>v</sub>$  thin films is sensitive to growth conditions. For example, RHEED patterns of sputter-deposited Mo oxide films grown at 445 °C with various  $p(O_2)$  show that a secondary component phase, i.e., Mo<sub>8</sub>O<sub>23</sub>, is evidenced for films grown at lower  $p(O_2)$ . Well-textured MoO<sub>3</sub> film without secondary phases is obtained for  $p(O_2)$ =62.3% [165]. FTIR spectra of PLD MoO<sub>3</sub> films grown at different  $T_s$  show that the broad infrared absorption band in the region of  $500-1000$  cm<sup>-1</sup> starts to be resolved into strong absorption peaks at around 570, 625, 700, 840, and 985 cm<sup>-1</sup> at  $T_s > 200$  °C [157], indicating the pure  $\alpha$ -MoO<sub>3</sub> phase. In addition, the splitting of IR active mode at 900–1010 cm<sup>-1</sup> is evidenced for well-crystallized film. X-ray photoelectron spectroscopy (XPS) is also a sensitive tool to evaluated the local environment: (i) the XPS Mo 3d core level spectra evidence the reduction of  $Mo^{6+}$  ions to  $Mo^{5+}$  ions in non-stoichiometric film and (ii) the broadening of the Mo 3d core level peaks is assigned to the distortion of the oxide matrix [166].

To fabricate nanoscopic MoO3, methods like template-directed reaction of molybdic acid and the subsequent leaching process [167], and templating against carbon nanotubes have been attempted  $[168]$ . Polycrystalline MoO<sub>3</sub> thin films have been prepared by oxidation at high temperature of molybdenum compound layers deposited by chemical vapour deposition (CVD) from molybdenum hexacarbonyl  $Mo(CO)_{6}$  [169].

The main parameter for a thin film microbattery is the delivered gravimetric capacity. Rather than being expressed as the conventional unit of mAh·g<sup>-1</sup>, due to the uncertainty in the film density, technologists prefer the stored charge, *Q* expressed in µAh or in coulomb, per film surface area and the film thickness, i.e.,  $\mu$ Ah·cm<sup>-2</sup>  $\mu$ m<sup>-1</sup> or mC·cm<sup>-2</sup>  $\mu$ m<sup>-1</sup>. The relation

between the gravimetric capacity, *Q*m, of the material and the volumetric capacity of a film, *Q*f, is given by:

$$
Q_{\rm f} = 0.36 \, d \, Q_{\rm m},\tag{2}
$$

where  $Q_f$  is expressed in mC·cm<sup>-2</sup>  $\mu$ m<sup>-1</sup>,  $Q_m$  in mAh·g<sup>-1</sup>, and *d* is the density of the material in g·cm<sup>-3</sup>. Considering the accommodation of 1.5Li per mole of MoO<sub>3</sub>, yielding a gravimetric capacity of 280 mAh·g<sup>-1</sup>, the capacity of a dense film can reach the value of ≈130  $\mu$ Ah·cm<sup>-2</sup>· $\mu$ m<sup>-1</sup>. For certain applications, high-quality films grown by PLD are required. Currently, PLD MoO<sub>3</sub> thin films are grown using a KrF excimer laser ( $\lambda = 248$  nm) with a fluence of 2 J cm<sup>-2</sup> (energy of 300 mJ per pulse) and deposited on various substrates heated in the range of  $25 \le T_s \le 500$  °C under an atmosphere of O<sub>2</sub> flow maintained at a pressure of  $0.1 \le$  $p(O_2) \leq 20$  Pa. In the prior report, Julien et al. showed that the structure analyzed by optical spectroscopy strongly depends on  $T_s$ . For  $T_s < 150$  °C, an amorphous phase is formed, the β-MoO<sup>3</sup> monoclinic phase grows at *T*<sup>s</sup> ≈ 200 °C, and the layered α-MoO<sup>3</sup> orthorhombic phase appears at  $T_s = 300 \degree C$  [170-172]. Al-Kuhaili et al. reported the growth of polycrystalline MoO<sub>3</sub> films on unheated substrates using both XeF and KrF excimer lasers. By tuning the annealing temperature in the range of 300 to 500 °C, both the grain size and surface roughness increased. Films formed using the XeF laser ( $\lambda = 351$  nm) and annealed at 400 °C have the best stoichiometry of  $MoO<sub>2.95</sub>$  [155]. Analyzing the growth mechanism, Ramana and Julien concluded that the thermochemical reaction during ablation strongly influences the structural characteristics of PLD MoO<sub>3</sub> films. Above  $T_s = 400 \degree C$ , the formation of compositional defects induces structural disorder, i.e.,  $\alpha$ -β-MoO<sub>3-δ</sub> phase mixture [165,173].

The applicability of  $MoO<sub>3</sub>$  films as cathode component in  $Li/M<sub>0</sub>O<sub>3</sub>$  microbatteries was demonstrated by the best electrochemical features. Early works have been reviewed by Julien et al. [174,175]. Assuming fully dense MoO<sub>3</sub> films, a theoretical specific capacity ~130  $\mu$ Ah cm<sup>-2</sup>  $\mu$ m<sup>-1</sup> is expected compared with 69  $\mu$ Ah cm<sup>-2</sup>  $\mu$ m<sup>-1</sup> for LiCoO<sub>2</sub>. In 1989, Ohtsuka and Yamaki [176] prepared a Li/solid-state electrolyte/MoO3-δ microbattery by thin film technology, in which the MoO<sub>3- $\delta$ </sub> cathode film thickness was 1.0  $\mu$ m. The Li<sub>2</sub>O–V<sub>2</sub>O<sub>5</sub>–SiO<sub>2</sub> thin film electrolyte exhibits a conductivity of  $1 \times 10^{-4}$  S cm<sup>-1</sup> at 25 °C. The Li//MoO<sub>3- $\delta$ </sub> cell was cycled over 240 times with a capacity of  $60 \mu$ Ah cm<sup>-2</sup>. In 1994, Julien et al. investigated the influence of the conditions of preparation of  $MoO<sub>3</sub>$  thin on their ability to intercalate  $Li<sup>+</sup>$  ions have been evaluated [177]. Microbatteries fabricated with cathode films formed at moderate temperature ( $T_s \leq 250$  °C) exhibit a monotonous discharge profile but show a strong dependence of the average voltage *vs*. deposition temperature. For  $T_s = 250$  °C, the average potential is 2.9 V and the Li uptake is close to 3 Li/Mo. The variation of the cell voltage is generally attributed

to (i) the presence of large crystallites surrounded by amorphous grain boundaries, (ii) the presence of a mixed  $\alpha$ -*β*-phase which may tune the standard potential, and/or (iii) high concentration of oxygen defects in the host structure involving a lower Fermi level in MoO<sup>3</sup> film. An in-depth study of polycrystalline thin films for cathode applications was reported [178]. It was found that the kinetically accessible discharge range was  $0 \le x \le 1.5$  in Li<sub>x</sub>MoO<sub>3</sub> and that the operating potential for the disordered phases was higher than that of the crystal. The OCVs ranged from  $3.2-2.5$  V and the cells were discharged to 1.5 V vs. Li<sup>+</sup>/Li. The chemical diffusion coefficient  $D_{Li}^*$  of the crystalline film ranged from  $10^{-11}$ – $10^{-12}$  cm<sup>2</sup> s<sup>-1</sup> for disordered polycrystalline films with a thermodynamic factor that was approximately two orders of magnitude higher than in MoO<sub>3</sub> crystals. Electrochemical properties of MoO<sub>3</sub> film grown by thermal and flash evaporation have been investigated in cell of 100 μAh capacity [179]. Thermodynamic data show that microbatteries fabricated with films formed at moderate temperature exhibit a monotonous discharge voltage indicating that the  $Li<sub>x</sub>MoO<sub>3</sub>$  material remains in the single phase even for large degree of intercalation, *x*. The lithium diffusivity increases with the temperature  $T_s$  of preparation of the film and reaches the value of  $10^{-11}$  cm<sup>2</sup>  $s^{-1}$  in Li<sub>x</sub>MoO<sub>3</sub> for  $T_s = 250$  °C. MoO<sub>y</sub> films have been deposited on Ni substrate by rf-sputtering using different flow rate of O<sub>2</sub>/Ar mixture (1.5  $\leq \Phi \leq 8.0$  sccm) as sputter gas [180]. Their electrochemical properties appear to be dependent of the oxygen flow rates and the nature of the substrate. At a flow rate of 8.0 sccm, the  $Li/MoO<sub>v</sub>$  microcell delivers a discharge capacity of 450  $\mu$ Ah cm<sup>-2</sup> at 5  $\mu$ A cm<sup>-2</sup> current density.  $\tilde{D}_{Li}$  increases at least by two orders of magnitude in films grown at high oxygen partial pressure. A value  $D_{Li}^* = 2 \times 10^{-12}$  cm<sup>2</sup> s<sup>-1</sup> is obtained for  $\Phi$  $= 8$  sccm. In 2000, Ferreira et al. [181] prepared MoO<sub>y</sub> thin films deposited by r.f. reactive sputtering of metallic Mo target in an  $Ar/O<sub>2</sub>$  atmosphere. The films with different compositions and crystal structures were obtained by varying the oxygen flow  $0.5 \le \Phi \le 4.0$ . Combined XRD and XPS analyses show that, with the increase of  $\Phi$ , the microstructures were obtained as: amorphous  $\rightarrow$  (MoO<sub>2.8</sub> + α-MoO<sub>3</sub>)  $\rightarrow$  β-MoO<sub>3</sub>  $\rightarrow$  (MoO<sub>2.8</sub> + β-MoO<sub>3</sub>). The film formed with the β-MoO<sub>3</sub> phase exhibits the best lithium insertion and electrochromism properties.

In 2001, Ohtsuka and Sakurai [144] fabricated an all-solid-state lithium microbattery with thicker cathode layer (4.66 µm) to increase the cell capacity per unit area. The thin film battery architecture deposited on stainless-steel substrate is composed of a thick  $MoO<sub>3-δ</sub>$  cathode film, Li film anode and  $Li_2O-V_2O_5-SiO_2$  thin film electrolyte. The cathode films (0.49 cm<sup>2</sup> area) with composition  $Mo<sub>9</sub>O<sub>26</sub>$  were grown at 100 °C by magnetron sputtering at rf-power of 200 W using a sintering-powdered MoO<sub>3</sub> target in Ar at pressure of Pa. The discharge curves are Sshaped with an average potential of 2.3 V. At discharge current density of 10 mA  $cm^{-2}$ , the battery has discharge capacity of 290 mAh cm<sup>-2</sup> (56.4  $\mu$ Ah cm<sup>-2</sup>  $\mu$ m<sup>-1</sup>) at the 40<sup>th</sup> cycle. Assuming a film of composition  $Mo<sub>9</sub>O<sub>26</sub>$  (determined by XRD) with density 4.7 g cm<sup>-3</sup>, the first discharge capacity is equivalent to 182 mAh  $g^{-1}$ .

In 2006, Ramana et al. reported a specific discharge capacity of 90  $\mu$ Ah cm<sup>-2</sup>  $\mu$ m<sup>-1</sup> for a PLD  $\alpha$ -MoO<sub>3</sub> film deposited at *T*<sub>s</sub> = 400 °C, while only 53  $\mu$ Ah·cm<sup>-2</sup>  $\mu$ m<sup>-1</sup> was delivered for  $T_s = 200 \degree C$  [157]. The superiority of MoO<sub>3</sub> cathode material that can be extend the upper limit of rechargeable lithium microbatteries was demonstrated in experiments on lithium thin-film cells fabricated on glass slides [182]. Each cell included a layer of Ti (for adhesion to the glass slide), a patterned layer of Pt that served as a cathode current collector, a cathode layer of MoO<sub>3</sub>, a solid electrolyte layer of  $Li_{3.3}PO_{3.8}N_{0.22}$  (LiPON), and an anode layer of Li. All the layers were deposited by magnetron sputtering except for the Li layer, which was deposited by thermal evaporation. After 5500 cycles of charge/discharge at a current density of 0.7 mA cm<sup>-2</sup> the cell operating at temperature 150 °C delivered specific capacity of 140  $\mu$ Ah cm<sup>-2</sup> $\mu$ m<sup>-1</sup>, while the cell containing  $LiCoO<sub>2</sub>$  failed. Mo $O<sub>3</sub>$  thin films were prepared by sol-gel process from molybdenum acetylacetonate sol [183]. The films were formed onto silica substrates by spincoating with subsequent heat treatment at 508 °C for 2 h. SEM images show grains shaping from oval to rod are distributed on the substrates in random orientation. It seems that the size and morphology of micrograins in  $MoO<sub>3</sub>$  films depend on the heat treatment temperature, ambient and type of substrates. Comini et al. [184] have obtained needle-like nanoparticles by rf-sputtering technique and continuous planar films with sol-gel method on alumina substrates. Yu and coworkers reported that a porous MoO<sub>3</sub> film prepared by a hydrothermal route and subsequent calcinations in air, delivers a high capacity of 750 mAh  $g^{-1}$  at 1C rate, and exhibits long cycle life of 120 cycles with 80 % capacity retention as the anode of LIB [140].

Recently, Moitzheim et al. [185] reviewed the progress toward solid-state 3D thin-film Liion microbatteries and discussed the status of the individual components: thin‐film cathodes, anodes, and thin‐film solid electrolyte. Julien et al. investigated the electrochemical performance of molybdenum-suboxide thin films as anode layers in planar lithium microbatteries [72]. Puppala et al. investigated the microstructure and morphology of PLD MoO<sub>3−x</sub> thin films' growth for catalytic applications using a femtosecond laser (f-PLD) and a nanosecond excimer-laser (n-PLD). Substantially textured films with a partially crystalline phase prior to annealing were obtained by the f-PDL laser, while the n-PLD-grown MoO<sub>3-x</sub> films were predominantly amorphous with a smooth surface [186]. Sunu et al. claimed that asdeposited PLD films ( $T_s = 400$  °C,  $\Phi = 4-5$  J·cm<sup>-2</sup>, repetition rate of 15 to 20 Hz, and  $p(O_2) =$ 500 Pa) are suboxide-like, i.e., mixture of η--Mo<sub>4</sub>O<sub>11</sub> and  $χ$ -Mo<sub>4</sub>O<sub>11</sub>, which transformed to

MoO<sub>3</sub> after annealing at 500 °C in air for 5 h [18]. Several works reported the PLD growth of films  $(MoO<sub>3</sub>)<sub>1-x</sub>(V<sub>2</sub>O<sub>5</sub>)<sub>x</sub>$  with  $0.0 \le x \le 0.3$  prepared at room temperature under an oxygen pressure of 13.3 Pa. The effect of the  $V_2O_5$  content on the coloring switching properties for thermochromic, gasochromic, photochromic, and electrochromic applications was investigated [187,188]. Contrary to pure  $MoO<sub>3</sub>$ , the electrochromism of  $MoO<sub>3</sub>-V<sub>2</sub>O<sub>5</sub>$  films showed that the Mo oxidation state  $(+6)$  did not change considerably upon Li<sup>+</sup> insertion, while  $V^{5+}$  was reduced considerably to  $V^{4+}$  [187]. A similar improvement of the gas-sensing properties, i.e., the shortest response time and highest transmittance change, was observed for  $V_2O_5$ -doped MoO<sub>3</sub> films under an H<sup>2</sup> atmosphere [188]. **Figure 6** presents the structural and electrochemical properties of MoO<sup>3</sup> thin films deposited on Si wafer using PLD technique. The average grain size was estimated to be 50 nm. The  $\alpha$ -MoO<sub>3</sub> thin films deposited at  $T_s$ =400 °C (500 nm thick) delivers a volumetric capacity of 90  $\mu$ Ah cm<sup>-2</sup>  $\mu$ m<sup>-1</sup>.



**Fig.** 6. Structural and electrochemical properties of  $\alpha$ -MoO<sub>3</sub> thin films deposited on Si wafer using PLD technique. (a) XRD patterns for different deposition temperature. Asterisks show the peaks due to the β-MoO<sub>3</sub> phase for film deposited at  $T_s=100$  °C. (b) Dependence of the growth temperature on the surface roughness determined by atomic force microscopy; (c) Influence of the growth temperature on the optical band gap. (d) The discharge-charge profiles

of Li//MoO<sub>3</sub> microcells cycled at a current density 10  $\mu$ A cm<sup>-2</sup>. A volumetric capacity of 90 μAh cm<sup>-2</sup> μm<sup>-1</sup> is obtained for film deposited at  $T_s$ =400 °C.

#### *2.1.6. Electronic properties*

The chemistry of molybdenum is dominated by the consequences of the considerable extension of its *d*-orbitals and the position of the *d*-electrons redox potential relative to the anion valence band edge. Even prepared at elevated temperatures under atmospheric oxygen pressure, molybdenum trioxide always contains some oxygen defects  $(M_0O_{3-\delta})$ . Deb estimated the oxygen vacancy concentration to be  $3\times10^{-19}$  mol<sup>-1</sup>, resulting in a formal stoichiometry of  $(3-\delta) = 2.999$  [189]. MoO<sub>3</sub> is an n-type semiconductor with indirect bandgap that has reported between 2.9 and 3.3 eV.  $\alpha$ -MoO<sub>3</sub> is found to be indirect insulator with the conduction band minimums (CBM) located at  $\Gamma$  point and the valence band maximum (VBM) at  $\overline{R}$  point [190]. The introduction of oxygen vacancies in  $MoO<sub>3</sub>$  gives rise to electrical conductivity. The valence band is generated by oxygen  $2p_{\pi}$  orbitals, while the conduction band is formed by overlapping metal 4d and 5s bands. Structural characterization of oxygen defects in MoO<sub>3-δ</sub> by Raman spectroscopy and X-ray diffraction has been reported by Dieterle et al. [78].

#### *2.1.7. Intercalation properties of MoO<sup>3</sup>*

The layered crystalline  $\alpha$ -MoO<sub>3</sub> is an attractive material for charge storage because of its ability to intercalate ions in a wide range of composition [108,191,192]. Both faradaic and nonfaradaic mechanisms can store charge. The latter is the dominant factor in electric double-layer capacitors, in which charge is stored electrostatically from the reversible adsorption of ions onto high-surface-area materials. The other option is to store charge in a faradaic process, which again can take place on a surface, as in pseudocapacitors [193,194], or in the bulk of the material, as in conventional batteries [195]. Due to their reversibility, fairly attractive energy densities, conductivities, and Li diffusion coefficients, and a good availability,  $MoO<sub>y</sub>$  materials are probably the most promising candidates for anodes in non-aqueous batteries among all second and third row transition metal oxides. A scheme of the modification of the structure of the layered lattice of  $\alpha$ -MoO<sub>3</sub> phase due to the intercalation of foreign ions (Li<sup>+</sup>, Na<sup>+</sup>, K<sup>+</sup>, etc) or molecules (polypyrrole) is shown in **Fig. 7**.



**Fig. 7**. Scheme of the modification of the structure of the layered lattice of α-MoO<sup>3</sup> phase due to the intercalation of foreign atoms or molecules.

#### 2.1.6.1. Chemical lithiation

Orthorhombic  $α$ -MoO<sub>3</sub> powders synthesized by thermal decomposition of molybdic acid containing ammonium ions,  $(H_2O)_y$  (NH<sub>4</sub>)<sub>z</sub> M<sub>o</sub>O<sub>3</sub>, in air at 600 °C for 6 h were lithiated by Hashem et al.  $[196]$ . Li<sub>x</sub>MoO<sub>3</sub> were prepared by reaction of various amounts of water-free lithium iodide (the molar ratios of Li/Mo 0.1≤*x*≤1.0 in organic solution; dispersions of LiI and MoO<sub>3</sub> powders were stirred for 24 h in dry n-hexane; the n-hexane was renewed two times during this period to remove the liberated iodine. The filtered powder was washed with excess of n-hexane until the filtrate was clear from iodine. The filtered powder was heated at 250 °C under vacuum for 8 h. The major amount of iodine evolved during this reaction step due to the reaction of LiI with  $MoO<sub>3</sub>$  and the subsequent liberation of  $I<sub>2</sub>$ . Structural studies show that a two-phase reaction occurs during chemical lithium insertion into  $α$ -MoO<sub>3</sub>. The XRD peaks related to the orthorhombic MoO<sub>3</sub> phase disappeared gradually until the formation of completely lithiated phase at  $x > 0.25$  in Li<sub>x</sub>MoO<sub>3</sub>, for which a new peak assigned to (010) refection appeared at about  $2\theta = 11.4^\circ$ . Mai et al. [197,198] reported the electroactivity of α-MoO<sub>3</sub> nanobelts after lithiation that show superior performance to non-lithiated α-MoO<sub>3</sub> nanobelts. The lithiated phase (light-blue color) was obtained by reaction of  $MoO<sub>3</sub>$  with LiCl in aqueous solution for 2 days followed by a hydrothermal process at 180 °C for 24 h. For the non-lithiated MoO<sub>3</sub>, the discharge capacity decreased from 301 to 180 mAh  $g^{-1}$  after 15 cycles, corresponding to a capacity retention of 60 %, whereas the discharge capacity of lithiated  $MoO<sub>3</sub>$ decreased from 240 to 220 mAh  $g^{-1}$  after 15 cycles, corresponding to a capacity retention of 92%, showing the stability and drastic improvement in the charge-storage capacity of the lithiated  $MoO<sub>3</sub>$  nanobelts. Villevieille et al. conducted similar studies on  $MoO<sub>3</sub>$  nanobelts

synthetized by hydrothermal route and chemically lithiated using a LiCl solution. The obtained nominal LiMoO<sub>3</sub>, shows enhanced electrochemical performance  $(>150 \text{ mA} \text{h g}^{-1})$  for a loading of 4 mg cm<sup>-2</sup> at 3C rate) and can be considered as a novel cathode with high specific charge (**Figs. 8a-b**) [199]. High resolution in situ XRD analysis reveals almost no damage to the nanobelt electrode after 50 cycles (**Fig. 8c**).



Fig. 8. (a) Galvanostatic curves of LiMoO<sub>3</sub> nanobelts prepared using 1 mol of LiCl per MoO<sub>3</sub> cycled at C/10 rate. (b) Cyclability at different C rate. (c) In situ XRD synchrotron pattern collected during the first cycle. In red the starting material before cycling, in green the end of the delithiation process, in blue the end of the lithiation process. Due to the small shift the changes in lattice parameter are less than 1%. Reproduced with permission from [199]. Copyright 2015 Elsevier.

#### 2.1.6.2. Electrochemical lithiation

In principle, many works have demonstrated the suitability of Li<sup>+</sup> ions to intercalate/deintercalate the interlayer spacing (van der Waals gap) of the  $\alpha$ -MoO<sub>3</sub> host structure by redox reaction with the accommodation up to 1.5 Li atom per Mo [11,64,135,196,200-203]. In the cut-off potential  $3.5$ -1.5 V vs. Li<sup>+</sup>/Li, the overall reversible intercalation/deintercalation reaction is:

$$
xLi^{+} + xe^{-} + [MoVI]O3 \leftrightarrow Lix+ [Mo(6-x)+]O3,
$$
\n(3)

assuming the reduction the transition-metal cations from  $Mo(VI)$  to  $Mo(V)$  and  $Mo(IV)$ oxidation states. Note that the  $Li<sup>+</sup>$  coupled electron transfer produces intense blue coloration  $(Lix+[Mo^{(6-x)+}]O_3)$  via strong light absorption of the pristine transparent  $[Mo<sup>VI</sup>]O_3$ . Figure 9a shows the cyclic voltammograms of the electrode made of crystalline  $\alpha$ -MoO<sub>3</sub> powders in the first five cycles at a sweep rate of  $0.05 \text{ mV s}^{-1}$ . In the first cycle, two sets of cathodic and anodic peaks appear at around the potential of 2.30, 2.73 V and 2.33, 2.48 V, which can be assigned to the insertion/extraction of  $Li^+$  ions between the MoO<sub>6</sub> octahedral interlayers and intralayers, respectively [11]. However, the set of peaks at (2.33, 2.75 V) completely disappeared in the second and subsequent cycles, indicating irreversible capacity losses due to the unrecoverable phase transformation. The first galvanostatic discharge charge profiles of a  $Li/\alpha$ -MoO<sub>3</sub> cell and those of subsequent cycles are shown in **Figs. 9b-c**. During the first cycle at C/10 current rate, the host can accommodate almost 1.6 Li/Mo, which corresponds to a specific capacity 265 mAh  $g^{-1}$ . A capacity retention of 152 mAh  $g^{-1}$  is remained after 100 cycles. The discharge capacities of anhydrous  $MoO<sub>3</sub>$  reported by several workers are in the range from 1.3 to 1.6 Li/Mo in the potential range above 1.2 V vs. Li<sup>+</sup>/Li  $[113,196,204]$ . The capacity of MoO<sub>3</sub> observed here is approximately in good agreement with the theoretical gravimetric capacity of 280 mAh  $g^{-1}$ . The disappearance of the irreversible peak (at ca. 2.75 V) in the CV and the absence of voltage plateau at ca. 2.78 V in the galvanostatic discharge curve is related to a structural modulation that occurs upon insertion/extraction of  $Li<sup>+</sup>$  ions between the MoO<sub>6</sub> octahedral interlayers and intralayers, respectively, and it is responsible for a capacity loss of ca. 20 mAh  $g^{-1}$ . This suggests that part of the Li<sup>+</sup> ions, first introduced during the reduction reaction (discharge), later remain in the lattice. Such an effect has also been observed in  $V_2O_5$  [205] The mechanism of Li-ions intercalation into the  $\alpha$ -MoO<sub>3</sub> has been described as the formation of three phases separated by bi-phase domains that are responsible for voltage plateaus in the first discharge curve [11,196,206]. It is believed that the  $Li_{0.25}MoO<sub>3</sub>$  phase formed at 2.75 V vs. Li<sup>+</sup>/Li is at the origin of the irreversible capacity at the end of the first charge. Thus, for subsequent cycles of discharge-charge, the cathode material shuttles from  $Li_{0.25}MoO<sub>3</sub>$  to  $Li_{1.5}MoO<sub>3</sub>$  without the appearance of the first plateau. Based on these results, an idealized scheme of the Li intercalation process into the MoO<sup>3</sup> framework is shown in **Fig. 10** [135].



**Fig. 9**. Electrochemical intercalation of Li ions in crystalline α-MoO3. (a) Five first cyclic voltammograms of Li//α-MoO<sub>3</sub> cell cycled at a sweep rate of 0.05 mV s<sup>-1</sup> and cut-off potential 2.0–3.3 V vs. Li<sup>+</sup>/Li. (b) First galvanostatic discharge-charge profile recorded at C/10 rate. (c) Electrochemical patterns at the  $2<sup>nd</sup>$  and  $100<sup>th</sup>$  cycle. (a) Variation of the chemical diffusion coefficient of  $Li^+$  ions in the  $\alpha$ -MoO<sub>3</sub> host network with the degree  $x(Li)$  of intercalation. (e) Experimental setup of *in situ* measurements of the electrical resistance of the  $\alpha$ -MoO<sub>3</sub> electrode upon Li extraction (charge). (f) Composition dependence of the electrode resistance. MoO<sub>3</sub> can be considered such as a good self-limiting over-charge material.



Fig. 10. Idealized scheme of the intercalation process in MoO<sub>3</sub> framework. At the end of the discharge, the chemical composition is  $Li<sub>1.5</sub>MoO<sub>3</sub>$ , while at the end of the charge, it is Li<sub>0.25</sub>MoO<sub>3</sub>. Reproduced with permission from [135]. Copyright 2012 Elsevier.

The chemical diffusion coefficient,  $D_{Li}^*$ , of Li<sup>+</sup> ions in the  $\alpha$ -MoO<sub>3</sub> host lattice depends on the composition  $x$  and the morphology of the material. **Figure 9d** shows the compositional dependence of  $D_{Li}^*$  determined using the current step method (plot of  $\Delta E$  vs  $t^{-1/2}$ ) given by the relation [207]:

$$
D_{Li}^* = \widetilde{D} \frac{RT}{xF} \frac{\Delta x}{\Delta E},\tag{4}
$$

where  $\tilde{D}$  is the self-diffusion coefficient, *R* and *F* are the usual constant and *T* is the absolute temperature. Dickens reported a chemical diffusion coefficient of  $3 \times 10^{-9}$  cm<sup>2</sup> s<sup>-1</sup> (self-diffusion of  $3 \times 10^{-10}$  cm<sup>2</sup> s<sup>-1</sup>) for Li<sub>0.3</sub>MoO<sub>3</sub> [200] A maximum value of ca.  $10^{-10}$  cm<sup>2</sup> s<sup>-1</sup> has been reported for *x*≈0.6. Li mobility was shown to slightly decrease upon subsequent discharge-charge cycles due to irreversible structural and morphological changes of the host matrix [64]. However, when the cell was recharged, reoxidation of Mo produces a resistive compound, which induces a large polarization of the cell at the end of the charge as shown in **Fig. 9a**. This effect could be beneficial for application, thus,  $MoO<sub>3</sub>$  can be considered such as a good self-limiting overcharge material. Upon lithiation of  $Li_xMoO_3$ , the electronic conductivity increases from  $10^{-4}$  S cm<sup>-1</sup> for  $x = 0$  to ca.  $10^{-1}$  S cm<sup>-1</sup> for  $0.3 \le x \le 0.9$  [64]. *In situ* measurements of the electrical resistance of the electrode has been reported by Yebka et al. [204]. The results shown in **Fig. 9f** are in good agreement with the variation of the overpotential at the end of charge. The structural changes occurring in the positive electrode materials during Li electrointercalation were followed by *in situ* XRD measurements (**Fig. 11a**) and Raman spectroscopy (**Fig. 11b**) [204]. **Figure 11a** presents the XRD diagrams of  $Li_xMoO_3$  during the discharge process (0.0  $\leq$  $x \le 1.5$ ). These patterns indicate the formation of a two-phase system in the range  $0.0 \le x \le 0.25$ followed by a single phase  $Li_xMoO_3$  for which the framework structure is almost unchanged during discharge reaction up to 1.5 Li/Mo. However, one can observe the vanishing of Bragg peak at ca.  $2\theta = 52^{\circ}$  for  $x > 0.5$ Li/Mo. This indicates a continuous structure reorganization corresponding to complete reversibility of the insertion/de-insertion process for the  $Li<sub>x</sub>MoO<sub>3</sub>$ material in the potential range  $3.5$ -1.2 V. The orthorhombic lattice parameters of  $Li<sub>0.5</sub>MoO<sub>3</sub>$ chemically intercalated by n-butyllithium have been reported by Dickens and Reynolds [200] as  $a = 3.64$  Å,  $b = 16.7$  Å,  $c = 3.77$  Å, which result in a slight contraction along the  $a$ - and  $c$ direction and an expansion of the *b*-axis of 20% due to the filling of the interlayer spacing. Such a structural evolution has been reported by Tsumura et al. [11] as shown in **Fig. 11c**. The initial formation of the lithiated phase results in expansion of the interlayer spacing from 0.69 to 1.175 nm.

The evolution of the local structure of  $\alpha$ -MoO<sub>3</sub> cathode materials during the first discharge has been studied by Raman spectroscopy. The electrode was washed with DMC solution to remove LiPF<sub>6</sub> salt and left in a glove box for 12 h to reach equilibrium. **Figure 11b** shows the Raman spectra of  $\alpha$ -Li<sub>x</sub>MoO<sub>3</sub> nanofibers as a function of the concentration of Li<sup>+</sup>ions inserted in the range  $0 \le x \le 1$ . At the first discharge, the Raman spectrum of MoO<sub>3</sub> nanofibers shows significant changes. As a general trend, we notice that all the main spectral features observed in  $MoO<sub>3</sub>$  are present in the spectrum of  $Li<sub>x</sub>MoO<sub>3</sub>$  and the broadening of the bands indicate that the  $Li<sub>x</sub>MoO<sub>3</sub>$  phase is less ordered than the  $\alpha$ -MoO<sub>3</sub> lattice. On comparing the Raman features of  $Li_xMoO_3$  and  $MoO_3$ , the following considerations can be established: (i) the intensity of the translational modes at low frequency, i.e., 83 and 113 cm<sup>-1</sup>, is progressively quenched, (ii) some bands are enhanced, i.e., the  $\delta$ (O<sub>2</sub>-Mo-O<sub>2</sub>) scissoring  $B_{3g}$  and  $B_{1g}$  modes at 244 and 377 cm<sup>-1</sup>, respectively, while some others vanish, i.e., the intense  $\delta(O_1-M_0-O_1)$ wagging  $B_{2g}$  mode at 282 cm<sup>-1</sup> and the  $\delta$ (O<sub>2</sub>-Mo-O<sub>2</sub>) scissoring  $A_g$  mode at 364 cm<sup>-1</sup>, (iii) the single line, corresponding to the Mo=O stretching mode (molybdyl mode) along the *c-*axis broaden and shifted up to 998 cm<sup>-1</sup> with respect to 994 cm<sup>-1</sup>, typically observed for the α-phase, and (iv) new spectral features are detected in the spectrum of  $Li<sub>x</sub>MoO<sub>3</sub>$  (bands at 440, 758, 967 and  $1013 \text{ cm}^{-1}$ ). At least two factors can account for the distinction between the spectra of these materials. First, the non-equivalent character of the transition metals in the lattice of  $Li_xMoO_3$ , meaning three kinds of molybdenum oxidation states from  $Mo^{6+}$  to  $Mo^{4+}$ , which implies the modification in the local environment of  $MoO<sub>6</sub>$  octahedra. Second, the Li atom oscillations may couple with some modes of the MoO<sup>3</sup> lattice. These results are consistent with the reported data in the literature, which describe that the lithium insertion process consists in rather moderate local distortions allowing the accommodation of 1.5 Li/mol of oxide without breaking of the orthorhombic symmetry [11]. The structural modifications in  $Li<sub>x</sub>MoO<sub>3</sub>$  can be also evidenced by the variation of the peak intensities of stretching modes; a decrease of the peak intensity ratio *I*994/*I*<sup>818</sup> with the increase of Li content is shown in **Fig. 11d**.



**Fig. 11**. (a) In-situ XRD patterns taken during the electrochemical lithium intercalation of α-MoO<sub>3</sub> crystal. Around  $2\theta = 55^{\circ}$  the diffraction line vanishes with the increase of Li concentration in the lattice. (b) Raman spectra of pristine and  $Li<sub>x</sub>MoO<sub>3</sub>$  ( $x = 0.5$  and 1.0). New vibrational modes are marked by asterisks. (c) Change of interlayer spacing between Mo–O octahedron layers of  $Li_xMoO_3$  with lithium composition. Reproduced with permission from [11]. Copyright 1997 Elsevier. (d) Evolution of the Raman peak intensity ratio *I*994/*I*<sup>818</sup> with *x*(Li).

The mechanism of a phase transition in electrochemical lithium insertion into  $\alpha$ -MoO<sub>3</sub> was also investigated by transmission electron microscopy (TEM), X-ray diffraction (XRD) and Xray photoelectron spectroscopy (XPS) [206,208]. The combined XPS, RBS, and NRA analysis of MoO<sup>3</sup> thin films electrochemically treated at selected potentials between 1.7 and 3.2 V vs. Li<sup>+</sup>/Li allowed the dosing of intercalated lithium and the determination of the composition of the lithiated phases.  $Li_{0.50}MoO<sub>3</sub>$ ,  $Li_{1.20}MoO<sub>3</sub>$ , and  $Li_{0.21}MoO<sub>3</sub>$  were obtained after intercalation at 2.58 and 1.73 V and deintercalation at 3.2 V, respectively, showing that ∼1.2 mol of Li can be initially intercalated in the potential range 1.7–3.2 V (capacity of 223 mAh  $g^{-1}$ ), and ~0.2 mol of Li per mol of  $MoO<sub>3</sub>$  is trapped in the oxide matrix after the initial stages of intercalation. The Li 1s core level (at  $E_B = 55.80$  eV) is most intense at 1.73 V and does not vanish at 3.2 V. Changes of the electronic structure after intercalation result from the occupation of the Mo 4d states (at  $E_B = 1.0$  eV) originally empty in the pristine oxide [166].

Recently, Guo et al. [209] investigated the electrochemical insertion of proton in  $\alpha$ -MoO<sub>3</sub> in acidic electrolyte (i.e., the  $4.4 \text{ mol L}^{-1}$   $H_2SO_4$  commonly used in lead-acid batteries). **Figure 12** presents the electrochemical patterns and operando structure changes of initial proton deintercalation in molybdenum bronzes  $H_xMoO_3$  ( $0 \le x \le 1.68$ ). Zhang et al. [210] reported how selective  $H_3O^+$  intercalation in a neutral  $ZnCl_2$  electrolyte can be achieved for water-proton co-intercalated  $\alpha$ -MoO<sub>3</sub>. H<sub>2</sub>O molecules located between MoO<sub>3</sub> interlayers block  $\text{Zn}^{2+}$ intercalation pathways while allowing smooth  $H_3O^+$  intercalation/diffusion through a Grotthuss proton-conduction mechanism. Thus, the water-proton insertion provides enhanced specific capacity of 356 mAh  $g^{-1}$ , a rate capability of 77.5% at 4 A  $g^{-1}$  and a cycling stability of 83% over 1000 cycles.



**Figure 12**. Electrochemistry and operando structure changes of initial proton intercalation (A) The initial galvanostatic charge-discharge profiles of  $MoO<sub>3</sub>$  at 1 A  $g<sup>-1</sup>$ . Typical redox processes are labeled as C (cathodic)/A (anodic) *n*, where *n* is the redox index. Note the initial cathodic processes are unique and labeled as  $Cl<sup>2</sup>/C2$ . (B) Structure diagrams of  $\alpha$ -MoO<sub>3</sub> and the hydrogen molybdenum bronzes (HMBs) with different hydrogen amounts. The purple, red, and white spheres refer to Mo, O, and H atoms, respectively. Note that these structures were defined by different authors; therefore, the coordinates are different. (C) The contour plot of operando

diffraction data of cell (in 4.4 mol  $L^{-1}$  H<sub>2</sub>SO<sub>4</sub> electrolyte, galvanostatically tested at 1C), correlated with the electrode potential as a function of time. Copyright Guo et al. [209]. Open access article under the CC BY-NC-ND license.

Several approaches have been conducted to improve the electrochemical performance of MoO<sup>3</sup> electrodes (i.e., capacity retention, long life cycling) including nanocrystallization [130,211-213] core-shell composite [214], carbon-based hybrid [215-217], carbon encapsulation [218], MWCNT-based hybrid [219-221], incorporation into graphene network [222-224], MoO<sub>2</sub> intergrowth [225,226], dual-ion intercalation [210,227], and coating [228-230]. Nanobelts of  $\alpha$ -MoO<sub>3</sub> have been modified to H<sub>x</sub>Mo(O,N)<sub>3</sub> through a moderate ammonolysis method, i.e., reaction at  $200-300$  °C with gaseous NH<sub>3</sub>, which diffuses inbetween the MoO<sub>3</sub> layers and reacts with terminal oxygen sites. As a consequence, hydrogen is bonded to terminal oxygen, and the unit cell volume significantly shrinks mostly along the *b*-axis. The modified H<sub>x</sub>Mo(O,N)<sub>3</sub> electrode exhibits a specific capacity of ~250 mAh g<sup>-1</sup> and shows no capacity fading over 300 cycles in the potential window 1.5–3.5 V [227].

#### *2.1.8. MoO<sup>3</sup> anodes for LIBs*

Molybdenum trioxide, which can be also categorized as one of the conversion type transition metal oxides, has additional advantageous features of low cost and environmental benignity as well as even higher capacity. However,  $MoO<sub>3</sub>$  suffers from serious capacity fading caused by large volume changes and poor rate capability due to low conductivity. Recently, these drawbacks have been mitigated by different strategies, i.e., reduction of particle size to nanoscale and fabrication of composites.

#### *2.1.8.1. MoO<sup>3</sup> nanoparticles*

The MoO<sup>3</sup> nanoparticles have been demonstrated as high-capacity Li-ion battery anodes for next-generation electric vehicles  $[231-235]$ . Specifically, the MoO<sub>3</sub> anodes have been shown to have approximately three times the Li-ion capacity of commercially employed graphite anodes in thick electrodes suitable for vehicular applications [128,193, 232,236-240]. Researches about MoO<sub>3</sub> as an anode in the potential range down to 0 V (*vs. Li<sup>+</sup>/Li*) have shown that MoO<sup>3</sup> can theoretically accommodate 6 Li/Mo providing a specific capacity of 1117 mAh  $g^{-1}$ . The overall first lithiation reaction for MoO<sub>3</sub> is described by two reactions: the lithium insertion (addition) at potential >1.5 V up to  $x \approx 1.2$  (Eq. 5) and the conversion (transformation) reaction at potential <0.5 V up to  $x \approx 6.0$  (Eq. 6) as follows:

$$
MoO3 + xLi+ + xe- \to LixMoO3,
$$
\n(5)

$$
\text{Li}_x\text{MoO}_3 + (6-x)\text{Li}^+ + (6-x)\text{e}^- \rightarrow \text{Mo} + 3\text{Li}_2\text{O}.\tag{6}
$$

Note that the amorphization of the material does not alter the overall electrochemical performance of the anode. The conversion reaction yields nanodispersed particles of Mo metal in the Li<sub>2</sub>O matrix  $[241]$ . **Figure 13** shows the typical first discharge-charge curve of MoO<sub>3</sub> nanofibers as anode materials in lithium battery. Nanofibers were synthesized by hydrothermal method from MoO3*∙*1.4H2O sols prepared by ion exchange of ammonium heptamolybdate tetrahydrate (NH4)6Mo7O24*∙*4H2O.



Fig. 13. The typical first discharge-charge curve of MoO<sub>3</sub> nanofibers as anode materials in lithium battery. The overall first lithiation reaction for  $MoO<sub>3</sub>$  is described by two reactions: the lithium insertion (addition) at potential >1.5 V up to  $x \approx 1.2$  and the conversion (transformation) reaction at potential <0.5 V up to  $x \approx 6.0$ .

Xia et al. [208] investigated the lithiation mechanism in  $α$ -MoO<sub>3</sub> nanobelts using in situ TEM experiments. Results reveal that  $\alpha$ -MoO<sub>3</sub> undergoes a two-stage delithiation process. Mo nanograins were first transformed into crystalline  $Li_{1.66}Mo_{0.66}O_2$  along with the disappearance of Li2O and size shrink, followed by the conversion to amorphous Li2MoO3. The electrochemical reactions of  $\alpha$ -MoO<sub>3</sub> during lithiation and delithiation processes can be expressed as:

\n
$$
MoO_3 + 6Li^+ + 6e^- \rightarrow Mo + 3Li_2O,
$$
\n
$$
Mo + 3Li_2O \rightarrow 1.5Li_{1.66}Mo_{0.66}O_2 + 3.5Li^+ + 3.5e^-,
$$
\n
$$
1.5Li_{1.66}Mo_{0.66}O_2 \rightarrow Li_2MoO_3 + 0.5Li^+ + 0.5e^-,
$$
\n
$$
(9)
$$
\n

where Eq. (7) represents the first lithiation and Eqs. (8) and (9) account for the two-stage phase of the subsequent delithiation/lithiation reactions (**Fig. 14**) [208]. This shows that the total amount of extracted Li is about 4 [132,236]. The theoretical lithium storage capacity of Li<sub>2</sub>MoO<sub>3</sub> is calculated to be about 734 mAh  $g^{-1}$  according to its weight and the ability of accommodating lithium ions, which is much lower than MoO<sub>3</sub> of 1117 mAh  $g^{-1}$ .



**Fig. 14**. HRTEM images and electron diffraction patterns (EDP) showing the microstructure evolution of  $MoO<sub>3</sub>$  nanobelt in lithiation and delithiation processes. (a) pristine  $MoO<sub>3</sub>$  nanobelt with a size of 56.1 nm. (b) The first lithiation. (c) Intermediate stage of the first delithiation. (d) Delithiated MoO<sub>3</sub> nanobelt. EDPs recorded from the nanobelt anode reveal the phase conversion during conversion. Reproduced with permission from [208]. Copyright 2016 American Chemical Society.

Ultralong  $\alpha$ -MoO<sub>3</sub> nanobelts (length of 200–300  $\mu$ m and uniform width of ~0.6–1.5  $\mu$ m) were hydrothermally synthesized using a molybdenum organic salt precursor. The electrodes fabricated with sodium carboxymethyl cellulose (Na-CMC) binder deliver the specific capacity of over 730 mAh  $g^{-1}$  for over 200 cycles at a 0.2C rate. Even cycled at high rates of 1–2C, high capacities of around 430–650 mAh  $g^{-1}$  can be still retained [129]. Sasidharan et al. [242] synthesized  $\alpha$ -MoO<sub>3</sub> hollow nanospheres (HNSs; outer diameter of 42 nm; hollow cavity of 23 nm) have been using a soft template of polymeric micelles with core–shell–corona architecture. The MoO<sub>3</sub> HNSs electrode delivers a discharge capacity of 263 mAh  $g^{-1}$  for the first cycle at a

rate of 0.3C and maintains structural integrity even after subjecting to a high current density 2000 mA  $g^{-1}$  (10C rate) The capacity retention is 2.2 times higher than that of dense MoO<sub>3</sub> particles. Ette et al. [243] prepared self-assembled lamellar  $\alpha$ -MoO<sub>3</sub> nanoflakes (20-30 nm) using facile polymer (polyvinyl pyrrolidone) assisted solvothermal method. After 50 cycles, a capacity value of 1027 mAh  $g^{-1}$  can be extracted at 0.2C rate while at 2C rate it shows capacity of 700 mAh  $g^{-1}$ . Riley and coworkers prepared  $\alpha$ -MoO<sub>3</sub> nanoparticles using a hot-wire chemical vapor deposition and reported that MoO<sub>3</sub> negative electrodes fashioned into thick electrodes containing 70% MoO<sup>3</sup> nanoparticles (10-40 nm in diameter), active material, 20% poly (vinylidenefluoride) (PVDF), binder, and 10% acetylene black (AB), conductive additive, can deliver a discharge capacity of  $\sim 1000$  mAh g<sup>-1</sup> at a reduced rate to C/10 [237]. Later, these workers showed that coating of the electrode with  $\sim 8$  Å of Al<sub>2</sub>O<sub>3</sub> by atomic layer deposition (ALD) achieves better rate capability [244]. It was suggested that the ALD coating enabled better adhesion of the high-volume expansion  $MoO<sub>3</sub> (≥100%)$  to the current collector. In a subsequent work,  $MoO<sub>3</sub>$  nanoparticles prepared by hot wire chemical vapor deposition (HWCVD) have been demonstrated as high-capacity Li-ion battery anodes for next-generation electric vehicles. The MoO<sub>3</sub> nanoparticles coated with ~8 Å thick  $Al_2O_3$  layer deliver a specific capacity of ~600 mAh  $g^{-1}$  at 5C rate [239]. The ALD-coated MoO<sub>3</sub> composite electrode (~15  $\mu$ m thick) prepared with an Al<sub>3</sub>O<sub>3</sub> coating of ~8 Å thick delivers a capacity of ~600 mAh g<sup>-1</sup> at 5C rate. Zhao and coworkers [245] synthesized MoO<sub>3</sub> hollow microspheres via a template-free solvothermal route and subsequent heat treatment in air. The MoO<sub>3</sub> hollow microspheres have a high specific surface area of  $48.2 \text{ m}^2 \text{ g}^{-1}$ , and with such a feature, when tested as a Li-storage anode material, they show a higher discharge capacity of 1377 mAh  $g^{-1}$  in the first discharge and a high reversible capacity of 780 mAh g<sup>-1</sup> after 100 cycles at a rate of 1C (**Fig. 15**). Varghese et al.  $[246]$  investigated the performance of full-cell Li-ion batteries based on MoO<sub>3</sub> nanobelts synthesized by hydrothermal process as cathode and prelithiated MoO<sub>3</sub> as anode. At 1 A  $g^{-1}$ current density, the Li<sub>x</sub>MoO<sub>3</sub> anode delivers a capacity of 700 mAh  $g^{-1}$  and at 2.5 A  $g^{-1}$  current density a capacity of 400 mAh  $g^{-1}$  is achieved. The MoO<sub>3</sub>//Li<sub>x</sub>MoO<sub>3</sub> full-cell delivers initial energy densities of 156 Wh kg<sup>-1</sup> and shows a CE of 99% over 100 cycles.



**Fig.** 15. (a) Discharge/charge voltage profile between 0.01 and 3 V at a rate of 1C of MoO<sub>3</sub> hollow microspheres; (b) cycling performance of  $MoO<sub>3</sub>$  hollow microspheres and bulk  $MoO<sub>3</sub>$ ; (c) cycling performance for the different rates  $(1-4C)$  of MoO<sub>3</sub> hollow microspheres; (d) electrochemical impedance spectroscopy of the MoO<sub>3</sub> hollow microspheres. Reproduced with permission from [245]. Copyright 2013 Elsevier.

The promising characteristics of both poly( $p$ -phenylene) (PPP) and  $Na<sub>0.25</sub>MoO<sub>3</sub>$  as anode materials in Li-ion cells, along with substantial amounts of polymer surface impurities resulting from the hydrothermal method, have led us to develop a new technique for the inclusion of PPP in MoO<sub>3</sub>  $[247]$ . Thus, the electrochemical reactivity of MoO<sub>3</sub> utilizing a conversion reaction has been widely investigated  $[129,236]$ . Reversible lithium-ion insertion in MoO<sub>3</sub> nanoparticles deposited by electrophoresis on stainless steel substrate have been reported by Lee et al. [128]. A stable capacity of 630 mAh  $g^{-1}$  was obtained for 150 cycles between 0.005 and 3.5 V. In contrast micron-sized particles of MoO<sub>3</sub> showed significant capacity fade after only several cycles. These thin films employed neither conductive additive nor binder as in traditional electrodes. Despite stable cycling at a rate of  $C/2$ , the reversible capacity was only 60% of the theoretical maximum of MoO<sub>3</sub>, 1117 mAh  $g^{-1}$ . Table 5 summarizes the electrochemical performance of MoO<sub>3</sub> nanoparticles as anode materials for LIBs prepared by various methods.

**Table 5**. Electrochemical performance of MoO<sub>3</sub> nanoparticles as anode materials for LIBs prepared by various methods. Relevant cycle number is given in brackets.



a) HMPs: hollow microspheres; NBs: nanobelts; NPs: nanoparticles; NFs: nanofibers; NSs: nanospheroids.

#### *2.1.8.2. MoO<sup>3</sup> nanocomposites*

The MoO<sub>3</sub> nanocomposites act as high capacity anode materials for lithium-ion batteries and exhibits good cyclic behavior. A variety of appealing strategies including carbon-based nanocomposites have been utilized to minimize the volume expansion/contraction and particle aggregation associated with the Li-insertion and extraction processes [232, 252,253]. MoO<sub>3</sub>based nanocomposites include: MoO3@carbon [232,252], MoO3@graphene [254-257], MoO<sub>3</sub>@rGO [223,258-260], MoO<sub>3</sub>@metal [261,262], and MoO<sub>3</sub>@metal oxides [238,263,264].

Tao and coworkers  $[252]$  synthesized a MoO<sub>3</sub>-carbon nanocomposite from a mixture of MoO<sub>3</sub> nanoparticles (2–180 nm) and graphite by a controlled ball milling procedure. The initial capacity of the MoO3/graphite (1:1 by weight) composite exceeds the theoretical capacity of 745 mAh  $g^{-1}$ , and the stable capacity of 700 mAh  $g^{-1}$  (94% of the theoretical capacity) still retains after 120 cycles. The electrode performance is linked with the unique nanoarchitecture of the composite and is compared with the performance of  $MoO<sub>3</sub>$ -based anode materials reported in the literature previously (nanoparticles, ball milled powders, and carbon-coated nanobelts). The high value of capacity and good cyclic stability of  $MoO<sub>3</sub>$ -carbon nanocomposite are attractive in respect to those of the reported  $MoO<sub>3</sub>$  electrodes [254]. Hussan et al. [232] synthesized MoO<sub>3</sub> nanobelts, diameter of 150 nm and a length of  $5-8$  µm, by a simple hydrothermal route followed by carbon coating. The  $C-MoO<sub>3</sub>$  nanobelts exhibit excellent cycling stability after being cycled at a current rate of C/10, maintaining their capacity at 1064 mAh g<sup>-1</sup> after 50 cycles. Highly uniform MoO<sub>3</sub> nanoparticles were prepared using hot-wire chemical vapor deposition (HWCVD) for use as active negative electrodes in high-energy LIBs [183]. An optimized reversible high capacity of  $\sim$ 1050 mAh g<sup>-1</sup> was obtained for an electrode fabricated from active material, acetylene black and PVDF (70:10:20) with a 250  $^{\circ}$ C pre-heat treatment.
Xue et al. [238] have synthesized  $\alpha$ -MoO<sub>3</sub> nanobelts and SnO<sub>2</sub>/ $\alpha$ -MoO<sub>3</sub> core-shell nanobelts, which show extraordinarily high reversible capacity as LIB anodes. The initial coulombic efficiency (ICE) of  $\alpha$ -MoO<sub>3</sub> nanobelts and SnO<sub>2</sub>/ $\alpha$ -MoO<sub>3</sub> core-shell nanobelts is 70.8% and 69.7%, respectively. The low ICE is probably attributed to the formation of solid electrolyte interphase (SEI) layer. In the following cycles, both  $\alpha$ -MoO<sub>3</sub> nanobelts and SnO<sub>2</sub>/ $\alpha$ - $MoO<sub>3</sub> core-shell nanobelts show high coulombic efficiency (>96.0%).$  The reversible capacity of SnO<sub>2</sub>/ $\alpha$ -MoO<sub>3</sub> core-shell nanobelts is ~2200 mAh g<sup>-1</sup>, much higher than that of pure  $\alpha$ -MoO<sub>3</sub> nanobelts ( $\sim$ 1400 mAh g<sup>-1</sup>). For pure  $\alpha$ -MoO<sub>3</sub> nanobelts, during the first lithiation process, three peaks can be observed at 2.25, 0.4 and 0.07 V. The peak at 2.25 V, only observed in the first discharge process, is probably ascribed to the formation of SEI layer. The other two peaks can be attributed to the lithiation process. In the subsequent lithiation processes, all the peaks (at 0.07, 0.3, 0.53, 1.29 and 1.48 V) are reversible, which can be attributed to the lithiation with different site energies and phase-transition process of  $\alpha$ -MoO<sub>3</sub> nanobelts. For SnO<sub>2</sub>/ $\alpha$ -MoO<sub>3</sub> core-shell nanobelts, during the first lithiation process, the irreversible peak, also observed at 2.25 V, can be ascribed to the formation of SEI layer. In the following lithiation processes, a new and reversible peak at  $\sim 0.7$  V is observed, corresponding to the formation of Li<sub>22</sub>Sn<sub>5</sub> alloys  $[238]$ . MoO<sub>3</sub> nanoplates were used as anode material for aqueous supercapacitors with higher energy density and relatively higher power density in comparison with traditional activated carbon [240]. MoO<sup>3</sup> nanowires were synthesized using a biomolecule-assisted hydrothermal approach and were demonstrated to be an excellent pseudocapacitor material. A specific capacitance 30 F  $g^{-1}$  was reported [193]. The nanocomposite MoO<sub>3</sub>/C was synthesized by a facile one-pot citric-nitrate method, where the carbon (from glucose – ratio  $MoO<sub>3</sub>:C = 10:3$  in weight corresponding to 23 wt.% carbon) was introduced *in situ*, and therefore uniformly dispersed in MoO<sup>3</sup> host, forming a homogeneous core-shell structure (thin carbon layer of 10- 15 nm). The MoO3/C anode exhibits superior electrochemical performance, a specific capacity of about 500 mAh  $g^{-1}$  in the voltage range of 0.01–3.0 V vs. Li<sup>+</sup>/Li can be maintained after 100 cycles  $[126]$ . Wang et al.  $[234]$  fabricated synergistic  $TiO<sub>2</sub>$ -MoO<sub>3</sub> core-shell nanowire array anode for high energy and high power lithium-ion batteries via a combination of hydrothermal and electrodeposition process. The  $TiO<sub>2</sub>-MoO<sub>3</sub>$  (1:1 mass ratio) hybrid exhibit a first specific capacity of 670 mAh  $g^{-1}$ , retention of ~400 mAh  $g^{-1}$  at 250 mA  $g^{-1}$  current rate after 200 cycles and good rate capability (up to 2 A  $g^{-1}$ ). The areal capacity is also as high as 3.98 mAh cm<sup>-2</sup>.

Martinez-Garcia et al. [265] presented the electrochemical performance of a low lithiation potential of 0.2 V vs. Li<sup>+</sup>/Li for pristine MoO<sub>3</sub> nanoplatelets and Si-decorated (0.3 wt.%). MoO<sub>3</sub> (Si@MoO<sub>3</sub>). The products of lithiation were identified as  $Li<sub>2</sub>MoO<sub>4</sub>$  and  $Li<sub>1.333</sub>Mo<sub>0.666</sub>O<sub>2</sub>$ , respectively. The initial cyclic voltammogram of  $Si@MoO<sub>3</sub>$  carried out in the 3.0 – 0.005 V potential range exhibits three cathodic peaks at 2.6, 2.18 and 0.02 V during (reduction process). Upon oxidation to 3.0 V, the anodic peaks are observed at 0.75, 1.23, 1.5, and 2.72 V. At subsequent cycle, the features over 2.0 V disappear and a new cathodic peak starts evolving at a potential of 1.45 V after the second polarization. Si-decorated  $MoO<sub>3</sub>$  nanoplatelets exhibit enhanced capacity of 1037 mAh  $g^{-1}$  with exceptional cyclability when charged/discharged at high current densities of 10 A  $g^{-1}$ . Mesoporous orthorhombic MoO<sub>3</sub> nanowire bundles with diameters 10-30 nm and lengths of up to 2 μm were prepared by topotactic chemical transformation from triclinic α-MoO3·H2O nanorods under vacuum condition at 260 °C The electrochemical measurements indicate that the high reversible capacities of 955 mAh  $g^{-1}$  can be retained over 150 cycles at 200 mA  $g^{-1}$  [266]. Single-crystalline  $\alpha$ -MoO<sub>3</sub> nanobelts with uniform width of  $\sim$ 200 nm and length at the micrometer level are first synthesized by a simple water-based hydrothermal route  $[248]$ . The  $\alpha$ -MoO<sub>3</sub> film electrode exhibits a reversible specific capacity of ~1000 mAh g<sup>-1</sup> at 50 mA g<sup>-1</sup> and a stable capacity retention of 387–443 mAh g<sup>-1</sup> at 2000 mA  $g^{-1}$ .

Graphene oxide-encapsulated  $\alpha$ -MoO<sub>3</sub> nanoribbons (GO/MoO<sub>3</sub>) composite was prepared by a self-assembly process. The great potential of this unique hybrid architecture as anode for Li-ion battery is revealed by a high specific capacity (up to 823 mAh  $g^{-1}$  after 70 cycles at 200 mA g<sup>-1</sup>), an excellent cycling performance (with more than 754 mAh g<sup>-1</sup> after 200 cycles at 1 A  $g^{-1}$ ) and a greatly-enhanced high-rate capability (displaying a high discharge capacity of 710 mAh g<sup>-1</sup> after 30 cycles at 3 A g<sup>-1</sup>) [267]. Ni et al. [268] fabricated the MoO<sub>3−*x*</sub>/CNTs composite via hydrothermal synthesis and controlled reduction in Ar/H<sup>2</sup> atmosphere, which consists of 63 mol%  $MoO<sub>3</sub>$  and 37 mol%  $MoO<sub>2</sub>$ . When evaluated as an anode in the 0.05–3.0 V voltage range, the MoO<sub>3</sub>/CNTs hybrid retains a capacity of 421 mAh  $g^{-1}$  after 100 cycles at 200 mA  $g^{-1}$ , and deliver 293 and 202 mAh  $g^{-1}$  at current densities of 2 and 4 A  $g^{-1}$ , respectively. MoO<sub>3</sub>@C nanofibers were synthesized by electrospinning the Mo precursor with polyvinyl pyrrolidone (PVP), followed by annealing. The as-prepared  $MoO<sub>3</sub>@C$  nanofibers deliver a discharge capacity of 623 mAh g<sup>-1</sup> after 100 cycles at 500 mA g<sup>-1</sup>, and 502 mAh g<sup>-1</sup> even at 1000 mA  $g^{-1}$  [124]. Ding et al. [122] adopted a facile two-step method in which pristine bulk MoO<sub>3</sub> is first converted into  $MoO<sub>3</sub>$  nanorods  $(MoO<sub>3</sub>$  NRs) through mechanical grinding, to buffer the continuous volume changes, and then coated with amorphous carbon (a-C) through simple stirring and heating. Electrochemical tests reveal that the a-C/MoO<sub>3</sub> NRs exhibits outstanding specific capacity of 856 mAh  $g^{-1}$  after 110 cycles at a current density of 0.1C and cycle life of 485 mAh  $g^{-1}$  after 300 cycles at 0.5C and 373 mAh  $g^{-1}$  after 400 cycles at 0.75C. Yang et al. [269] aimed to expanded the interlayer spacing and enhanced the electrical conductivity of α-MoO<sup>3</sup> by introducing glucose in the layered framework. It is found that glucose functioned as a cage to hold the pulverized  $MoO<sub>3-x</sub>$  pieces, while the subsequent conversion reaction of between  $Li_2O$ , Mo and  $Li<sub>x</sub>MoO<sub>3</sub>$  occurred in confined space, which maintained electrodes structural integrity. MoO<sub>3-x</sub>@G exhibits superior specific capacity of 607 mAh g<sup>-1</sup> after 200 cycles at 1 A  $g^{-1}$  and a coulombic efficiency kept stable near 100%. The exceptional performance of MoO<sub>3−x</sub>@G is attributed to the capacitive contribution and high reversible reaction. Cao et al. [270] hybridized nanostructured MoO<sub>3</sub> particles with foam-like carbon nanoflakes (CNFs) via the pyrolysis of molybdenum 2-ethtlhexanoate ( $C_{48}H_{90}MoO_{12}$ ) at a low temperature of 300  $^{\circ}$ C under ambient atmosphere. Electrochemical tests of the MoO<sub>3</sub>/CNFs composite as anode material for LIBs show a high discharge capacity of 791 mAh  $g^{-1}$  after 100 cycles at 500 mA  $g^{-1}$  and even ~600 mAh  $g^{-1}$  at a high rate of 2000 mA  $g^{-1}$  (**Fig. 16**).



Fig. 16. (a) CV curves and (b) discharge-charge profiles of the MoO<sub>3</sub>/CNFs. (c) Cycle performance at 500 mA  $g^{-1}$  and (d) rate performance at current densities ranging from 200 to 2000 mA  $g^{-1}$  for the MoO<sub>3</sub>/CNFs and MoO<sub>y</sub> electrodes. Reproduced with permission from [270]. Copyright 2018 Elsevier.

Ultrathin N-doped carbon layer encapsulated  $MoO<sub>3</sub>$  nanosheets have been synthesized by a simultaneous pyrolysis–reduction process of dodecylamine-intercalated  $MoO<sub>3</sub>$  composites at 600 °C under a nitrogen atmosphere. The as-fabricated nanosheets can reach high initial charge and discharge capacities of 1610 and 1359 mAh  $g^{-1}$ , respectively, and show remarkable cycle stability with a specific capacity of 1250 mAh  $g^{-1}$  after 60 cycles at 0.3C rate. High specific

discharge capacities are maintained at fast C rates, e.g., 1370, 1010, 940, 610, 490 and 370 mAh  $g^{-1}$  at 0.3C, 1C, 2C, 4C, 10C, and 20C, respectively [218]. A nanocomposite  $\alpha$ -MoO3/MWCNT was synthesized via a surfactant-assisted solvothermal process followed by low-temperature calcination [271]. When the discharge/charge current density is increased from 0.1 to 2 A  $g^{-1}$ , the reversible charge capacity is only decreased from 1138 to 941 mAh  $g^{-1}$ , giving a capacity retention of 82.7%. After 300 cycles at a current of 0.5 A  $g^{-1}$ , a high discharge capacity of 1350 mAh  $g^{-1}$  is maintained. Martinez-Garcia et al. [265] presented the electrochemical performance of a low lithiation potential of  $0.2$  V vs. Li<sup>+</sup>/Li for pristine MoO<sub>3</sub> nanoplatelets and Si-decorated (0.3 wt.%). MoO<sub>3</sub> (Si $@$ MoO<sub>3</sub>). The products of lithiation were identified as  $Li_2MoO_4$  and  $Li_{1,333}Mo_{0,666}O_2$ , respectively. The initial cyclic voltammogram of Si $@$ MoO<sub>3</sub> carried out in the 3.0 – 0.005 V potential range exhibits three cathodic peaks at 2.6, 2.18 and 0.02 V during (reduction process). Upon oxidation to 3.0 V, the anodic peaks are observed at 0.75, 1.23, 1.5, and 2.72 V. At subsequent cycle, the features over 2.0 V disappear and a new cathodic peak starts evolving at a potential of 1.45 V after the second polarization. Si-decorated MoO<sub>3</sub> nanoplatelets exhibit enhanced capacity of 1037 mAh  $g^{-1}$  with exceptional cyclability when charged/discharged at high current densities of 10 A  $g^{-1}$ . Mesoporous orthorhombic  $MoO<sub>3</sub>$  nanowire bundles with diameters 10-30 nm and lengths of up to 2  $\mu$ m were prepared by topotactic chemical transformation from triclinic  $α$ -MoO<sub>3</sub>·H<sub>2</sub>O nanorods under vacuum condition at 260 °C The electrochemical measurements indicate that the high reversible capacities of 955 mAh  $g^{-1}$  can be retained over 150 cycles at 200 mA  $g^{-1}$  [266]. Singlecrystalline  $\alpha$ -MoO<sub>3</sub> nanobelts with uniform width of  $\sim$ 200 nm and length at the micrometer level are first synthesized by a simple water-based hydrothermal route [248]. The α-MoO<sub>3</sub> film electrode exhibits a reversible specific capacity of  $\sim$ 1000 mAh g<sup>-1</sup> at 50 mA g<sup>-1</sup> and a stable capacity retention of 387–443 mAh  $g^{-1}$  at 2000 mA  $g^{-1}$ . A porous MoO<sub>3</sub> film was prepared by electrodeposition on Ni foam substrates and exhibits a capacity of 650 mAh  $g^{-1}$  at a current density of 3 A  $g^{-1}$  as anodes for LIBs [139]. A MoO<sub>3</sub>/C nanocomposite synthesized using the electrospinning method with 42% carbon takes the morphology of nanofibers, with the diameter of 50-200 nm. The MoO<sub>3</sub>/C anode cycled in a 3.0–0.01 V potential window at current density of 40 mA  $g^{-1}$  retains a discharge capacity of 710 mAh  $g^{-1}$  after 100 cycles. When the load current density is increased to 800 mAg<sup>-1</sup>, the composite delivers a discharge capacity of 300 mAh g<sup>-1</sup> after 100 cycles [125]. Sun et al. [272] reported on the synthesis of single wall carbon nanotubes (SWNTs)-bridged MoO<sub>3</sub> nanosheets as anode material for LIBs. The MoO<sub>3</sub>/SWNTs composite as fabricated via a liquid phase exfoliation of layered  $MoO<sub>3</sub>$  crystallites producing multilayer MoO<sup>3</sup> nanosheets dispersed in isopropanol and mixed with solution processed SWNTs in the same solvent. The MoO3/SWNTs hybrid leads to a specific capacity of 865 mAh  $g^{-1}$  at 100 mA  $g^{-1}$  after 100 cycles, with a coulombic efficiency approaching 100% and a capacity fading of 0.02% per cycle. The  $MoO<sub>3</sub>@CoMoO<sub>4</sub>$  hybrid was fabricated by a facile one-step hydrothermal method [273]. Galvanostatic charge–discharge tests show that the hybrid anode delivers a capacity of 586 mAh  $g^{-1}$  at the high current density of 1 Ag<sup>-1</sup> and an enhanced cyclic capacity of 887 mAh  $g^{-1}$  after 140 cycles at the current density of 200 mA  $g^{-1}$ . Reduced graphene oxide wrapped hollow  $MoO<sub>3</sub>$  nanorods  $(MoO<sub>3</sub>@rGO)$  were fabricated by freeze drying method [274]. The MoO<sub>3</sub>@rGO composite delivers a high reversible capacity of 842 mAh g<sup>-1</sup> at 0.1 A g<sup>-1</sup> rate, excellent cycling stability of 778 mAh g<sup>-1</sup> at 0.1 A g<sup>-1</sup> after 200 cycles and excellent rate capability (455 mAh  $g^{-1}$  at 2 A  $g^{-1}$ ).

Recently, a novel composite of  $\alpha$ -MoO<sub>3</sub> nanobelts and single-walled carbon nanohorns (SWCNHs) has been synthesized by a microwave hydrothermal method. Tested as a anode material for LIBs, the  $\alpha$ -MoO<sub>3</sub>/SWCNH composite displays a specific capacity of 654 mAh  $g^{-1}$ at 1C rate, excellent rate capability (275 mAh  $g^{-1}$  at 5 C), and outstanding cycle life (capacity retention of >99% after 3000 cycles at 1C) without any cracking of the electrode [275]. Feng et al.  $[276]$  produced  $SnO<sub>2</sub>/MoO<sub>3</sub>$  nanoparticles encapsulated in plate-like graphite via simple hydrothermal synthesis and dry ball milling. The  $SnO<sub>2</sub>–MoO<sub>3</sub>–C$  displays high ICE of 70%, high reversible specific capacity of 1338 mAhg<sup>-1</sup> at 0.2 A g<sup>-1</sup> current rate after 300 cycles (715 mAhg<sup>-1</sup> at 5 A g<sup>-1</sup>. Long-term cyclic tests show a capacity of 571 mAhg<sup>-1</sup> at 2 A g<sup>-1</sup> after 1000 cycles. Zhao et al. [277] designed a core–sheath structured  $MoO_3@MoS_2$  composite via in situ growth of few-layered  $MoS<sub>2</sub>$  nanoflakes on the surface of the biotemplated  $MoO<sub>3</sub>$ . The electrochemical tests show that the composite possesses a high specific capacity of 1545 mAh  $g^{-1}$  at 100 mA  $g^{-1}$  and coulombic efficiency (above 98%) after 150 cycles. Teng et al. [278] developed a hierarchically nanostructured composite of  $MoO<sub>3</sub>-NiO/graphene$ . Ultrafine  $MoO<sub>3</sub>$ nanosheets and NiO nanoparticles were homogeneously anchored on the graphene layers. Tested as anode material for LIB, this ternary composite exhibits specific capacities of 1164 mAh g<sup>-1</sup> at 100 mA g<sup>-1</sup> after 50 cycles and 946.9 mAh g<sup>-1</sup> at 1000 mA g<sup>-1</sup> after 180 cycles. A fibrous  $CNTs/MoO<sub>3</sub>$  composite, in which CNTs anchored with  $MoO<sub>3</sub>$  nanoplates was synthesized by electrospinning process of the  $MoO<sub>2</sub>(acac)<sub>2</sub>-PAN-CNTS$  complex in solution and subsequent annealing. The anode composite has high specific capacities of 972 mAh  $g^{-1}$ after 100 cycles at 1 A  $g^{-1}$  and 905 mAh  $g^{-1}$  after 800 long-term cycles at 2 A  $g^{-1}$ . Rate capability tests show discharge capacities of 599 and 374 mAh  $g^{-1}$  at 5 and 10 mAh  $g^{-1}$  rate, respectively [279]. The ternary composite  $MoO<sub>3</sub>/V<sub>2</sub>O<sub>5</sub>/C$  having the morphology of nanofibers with the diameter of 200~500 nm was synthesized by electrospinning combined with heat treatment.

Electrochemical tests of the composite containing ~40% carbon retains a specific capacity of 737 mAh  $g^{-1}$  after 200 cycles at 0.2 A  $g^{-1}$  current density, and its coulomb efficiency is 93% [280]. **Table 6** summarizes the electrochemical performance of MoO<sub>3</sub> nanocomposites as anode materials for LIBs prepared by various methods.

Table 6. Electrochemical performance of MoO<sub>3</sub> nanocomposites as anode materials for LIBs prepared by various methods. Relevant cycle number is given in brackets.

Material <sup>a)</sup>	Synthesis	Specific capacity $(mAh g-1)$	Current rate $(mA g^{-1})$	Ref.
MoO <sub>3</sub> NBs/CNTs	Hydrothermal	166	5C (500)	[281]
MoO <sub>3</sub> @GNS	Electrostatic	833	200 (70)	$[267]$
MoO <sub>3</sub> /C NBs	Hydrothermal	1000	100(50)	$[215]$
$MoO3-δ/CNTs$	Hydrothermal	421	200 (100)	$[268]$
				$[252]$
aC-MoO <sub>3</sub> NRs				[122]
$MoO3-δ/graphene$	Vacuum drying	607	1000 (200)	$[269]$
MoO <sub>3</sub> /CNFs	Pyrolysis	791	500 (100)	$[270]$
$MoO3/N-doped C$	Pyrolysis-reduction	1250	0.3C(60)	[218]
MoO <sub>3</sub> /MWCNT	Solvothermal	1350	500 (300)	$[271]$
$MoO3/C$ layer (10-15 nm)	Hydrosol-gel	500	100 (100)	[126]
$MoO3/C$ microspheres	Spray-pyrolysis	808	1000 (100)	$[282]$
$SnO_2/\alpha$ -MoO <sub>3</sub> NBs	Ultrasonication	1895	C/10(50)	[238]
MoO <sub>3</sub> /C	Electrospinning	300	800 (100)	$[125]$
$MoO3/29\%$ C microballs	Spray-pyrolysis	733	2000 (300)	[249]
MoO <sub>3</sub> /graphene	In situ hydrothermal	869	50(80)	[253]
MoO <sub>3</sub> /graphene	Spray-pyrolysis	1296	2000 (100)	[283]
MoO <sub>3</sub> /SWCNHs	Microwave hydrothermal	654	1000 (3000)	[275]
$MoO3/graphene$ (1:1 in wt.)	Sonication	574	500 (100)	[284]
MoO <sub>3</sub> /rGO	Freeze drying	778	100 (200)	[274]
SnO <sub>2</sub> /MoO <sub>3</sub> /C	Hydrothermal	1338	200 (300)	$[276]$
C/MoO <sub>3</sub> NBs MoO <sub>3</sub> @C NFs MoO <sub>3</sub> NPs/C $\Omega$ $\Omega$ $\overline{I}$	Hydrothermal Electrospinning <b>Ball milling</b> Mechanical grinding CITI CIRIT	1064 623 700 485	110(50) 500 (100) 150 (120) 0.5C(385)	$[232]$ $[124]$

a) CNFs: carbon nanofibers; SWCNHs: single-walled carbon nanohorns.

Xia et al. [215] designed a polyporous  $MoO<sub>3</sub>/C$  nanobelt anode material by a hydrothermal route with (NH4)6Mo7O24∙4H2O and activated carbon. The carbon component (8 wt.%) present in the composite serves as a physical barrier preventing the aggregation of the  $MoO<sub>3</sub>$ nanoparticles, and offers good electronic conductivity for electrode reaction. The MoO3/carbon nanobelts exhibit a high specific capacity of 1000 mAh  $g^{-1}$  after 50 cycles at a current density of 0.1 A  $g^{-1}$  and retain a discharge capacity of 675 mAh  $g^{-1}$  at a current density of 5 A  $g^{-1}$  and long-term cycle stability to 550 cycles. In situ XRD examination on  $MoO<sub>3</sub>/C$  electrode is

performed to understand the reaction mechanism of lithium with  $MoO<sub>3</sub>$  in initial process (**Fig. 17**).



**Fig.** 17. In situ XRD results recorded during the first discharge of the MoO<sub>3</sub>/C cell. During the initial lithiation process, the intensities of (040) and (060) peaks strongly decrease, indicating a phase transition from pristine  $MoO<sub>3</sub>$  to a lithium-inserted phase. The nanobelts anode delivers a specific capacity of 1014 mAh  $g^{-1}$  at 100 mA  $g^{-1}$  rate after 50 cycles in cut-off 0.01-3.0 V. Reproduced with permission from [215]. Copyright 2015 Elsevier.

### *2.1.9. Anodes for SIBs*

 $MoO<sub>3</sub>$  is considered as an appealing choice of anode for sodium-ion batteries (SIBs) because of its high theoretical capacity of 1117 mAh  $g^{-1}$  (i.e., considering the charge transfer of 6e- /Mo). However, the poor cycling stability and capacity fade results from the large volume change upon Na<sup>+</sup> storage. Hence, several investigations offer new possibilities to bypass multiple material challenges facing current sodium electrodes.

The first experimental evidence of the electrochemical insertion of sodium in the orthorhombic MoO<sup>3</sup> lattice has been reported in 1995 by Spahr et al. [202]. Specific capacity of up to 240 mAh  $g^{-1}$  (at 0.15 mA cm<sup>-2</sup> current density) was obtained using 0.5 mol L<sup>-1</sup> NaClO<sub>4</sub> solution in propylene carbonate (PC) as electrolyte. In the first cycle (**Fig. 18a**), a very sharp reduction peak found in the cyclic voltammogram at a potential of about  $2.4 \text{ V}$  vs. Na<sup>+</sup>/Na indicates the formation of a sodium bronze,  $Na_xMoO_3$ . It is further reduced to  $Na_{x+y}MoO_3$ . Subsequently, *y* moles of sodium are cycled rather reversibly. The galvanostatic dischargecharge curves (**Fig. 18b**) suggest the formation of the molybdenum bronze of sodium,  $Na_xMoO_3$ [285] with  $0 < x < 0.4$  at a voltage plateau of 2.5 V followed by a second plateau at about 1.8 V. A maximum Na uptake of  $\sim$ 1.5 mole is obtained at a potential limit of 1 V vs. Na<sup>+</sup>/Na. In

2013, Hariharan reported the sodium storage in commercial  $\alpha$ -MoO<sub>3</sub> powder (anode loading  $\sim$ 1.5 mg cm<sup>-2</sup>) using 2016-coin-type cell with 1 mol L<sup>-1</sup> NaClO<sub>4</sub> dissolved in ethylene carbonate (EC) and propylene carbonate (PC) as electrolyte (**Figs. 18c-e**) [233]. The sodiation and desodiation capacities of 771 and 410 mAh  $g^{-1}$  were delivered in the voltage window 0.04–3.0 V with average voltage profiles lying below 1.0 V. During sodiation (**Fig. 18c**), the voltage dropped from OCV (2.6 V) to reach a small plateau (i) at ca. 2.3 V. resulting the formation of sodium bronze  $Na<sub>x</sub>MoO<sub>3</sub>$ . Beyond the first  $Na<sup>+</sup>$  insertion plateau, the voltage profile in the region 1.8-1.2 V corresponds to additional insertion of Na in MoO<sub>3</sub> leading to the Na<sub>x+y</sub>MoO<sub>3</sub>, followed by a long sloping tail.  $MoO<sub>3</sub>$  anode also shows favorable rate performance and long cycle life of 117 mAh  $g^{-1}$  over 500 cycles at 0.2C (0.223 A  $g^{-1}$ ). The real-time sodiation/desodiaton behaviors of  $\alpha$ -MoO<sub>3</sub> during electrochemical cycling was observed by Xia et al.  $[286]$  investigating the *in situ* TEM and the electron diffraction patterns of  $\alpha$ -MoO<sub>3</sub> nanobelts. The EELS measurements show that upon the first sodiation,  $\alpha$ -MoO<sub>3</sub> initially forms an amorphous  $Na<sub>x</sub>MoO<sub>3</sub>$  phase and is subsequently sodiated into intermediate phase of crystalline NaMoO2, finally resulting in the crystallized Mo nanograins embedded within the Na<sub>2</sub>O matrix. During the first desodiation process, Mo nanograins are firstly re-oxidized into intermediate phase NaMoO<sub>2</sub> that is further transformed into amorphous Na<sub>2</sub>MoO<sub>3</sub>, resulting in an irreversible phase transformation. The overall electrochemical reactions can be expressed by the relations:

$$
MoO3 + 6Na+ + 6e- \rightarrow Mo + 3Na2O,
$$
\n(10)

$$
Mo + 3Na2O \leftrightarrow Na2MoO3 + 4Na+ + 4e+,
$$
\n(11)

where Eq. (10) states for the first sodiation and Eq. (11) for the subsequent desodiation/sodiation.



**Fig. 18**. Electrochemical patterns of the sodium intercalation in commercial  $\alpha$ -MoO<sub>3</sub> powders. (a) Cyclic voltammogram in 0.5 mol  $L^{-1}$  NaClO<sub>4</sub>/PC at 20  $\mu$ V s<sup>-1</sup> sweep rate. (b) Galvanostatic discharge charge curves in 0.5 mol  $L^{-1}$  NaClO<sub>4</sub>/PC at 0.15 mA cm<sup>-2</sup> (first three cycles). Reproduced with permission from [202]. Copyright 1995 Elsevier. Electrochemical patterns of the sodium intercalation in  $\alpha$ -MoO<sub>3</sub> powders prepared by precipitation of phosphomolybdic acid. (c) Cyclic voltammogram of  $MoO<sub>3</sub>$  vs. Na<sup>+</sup>/Na in the window 0.04– 3.0 V recorded at 0.058 mV s<sup>-1</sup>. (d) First and second galvanostatic voltage profiles of MoO<sub>3</sub> vs. Na<sup>+</sup>/Na at 0.1C (0.1117 A  $g^{-1}$ ) in the window 0.04–3.0 V. (e) Voltage profiles showing hysteresis at selected cycles. Reproduced with permission from [233]. Copyright 2013 Elsevier.

Sreedhara et al. [287] fabricated a 3D MoO<sub>3</sub>-rGO composite from MoO<sub>3</sub> nanosheets chemically tagged with optimum amounts of rGO. The  $MoO<sub>3</sub>(10 wt. %)-rGO$  composite exhibits remarkable electrochemical stability (first discharge capacity of 1061 mAh g<sup>-1</sup> at C/20 rate; ICE of 88%; capacity retention of 665 mAh  $g^{-1}$  after 100 cycles), cyclability and high rate capability over a wide range of operating currents  $(0.05-1C)$ . At 1 C, the MoO<sub>3</sub>– $(10 \text{ wt.}\%)$ -rGO dislays a capacity of 330 mAh  $g^{-1}$ . Jiang et al. [288] demonstrated a surface phosphorylation strategy to mitigate the degradation of  $3D$  MoO<sub>3</sub> array electrodes, which allows to sustain a capacity of 265 mAh  $g^{-1}$ , or ~90% of the initial value, at a rate of 2 A  $g^{-1}$  over 1500 cycles. Kinetic analysis unveils a capacitance-dominated Na<sup>+</sup> storage behavior, owing to the enhanced electron mobility imparted by oxygen vacancies that are simultaneously introduced by phosphorylation. Liu et al.  $[289]$  prepared a nanocomposite of MoO<sub>3</sub> nanobelts coated with polypyrrole (PPy) as anode material and investigated the electrochemical performance of the full cell formed by PPy@MoO<sub>3</sub> as anode and Na<sub>0.35</sub>MnO<sub>2</sub> as cathode in 0.5 mol L<sup>-1</sup> Na<sub>2</sub>SO<sub>4</sub> aqueous electrolyte, which delivers an energy density of 20 Wh  $kg^{-1}$  at 80 Wkg<sup>-1</sup> and even maintain 18 Wh  $kg^{-1}$  at 2.6 kWkg<sup>-1</sup>.

Yang et al. reported the synthesis of  $MoO<sub>3</sub>$  nanoplates ( $MoO<sub>3</sub>$  NPs) from Mo-based metal– organic frameworks via a facile heating treatment [235]. When used as an anode in SIB, the material showed a discharge capacity of 154 mAh  $g^{-1}$  at 50 mA  $g^{-1}$  after 1200 cycles. Even at 500 mA  $g^{-1}$ , it exhibits a specific capacity 217 mAh  $g^{-1}$  after 500 cycles. It is believed that the nanoplate morphology prevents the pulverization caused by volume expansion in SIBs. Li et al. [290] reported that MoO<sub>3−*x*</sub> grown on flexible carbon cloth, with a well-controlled oxygen vacancy, exhibits a specific capacity of  $\sim$ 150 mAh g<sup>-1</sup> after 200 cycles at 0.1 A g<sup>-1</sup> current rate in the voltage range 4.0-1.0 V vs. Na<sup>+</sup>/Na, and long-life performance of 79 mAh  $g^{-1}$  after 2000 cycles at 1 A  $g^{-1}$ . The oxygen vacancy in MoO<sub>3- $\delta$ </sub> is responsible for the two-plateau voltage profile, in contrast to the sloping feature observed in  $\alpha$ -MoO<sub>3</sub>. Xu et al. [291] demonstrated the beneficial effect of oxygen vacancies (OVs) in MoO3−δ nanosheet anode for SIBs. A series of measurements show that the OVs increase the electric conductivity and Na‐ion diffusion coefficient, and that the ultrathin  $Al_2O_3$  coating promotes the effective reduction of cyclinginduced solid-electrolyte interphase. The Al<sub>2</sub>O<sub>3</sub> coated MoO<sub>3</sub><sup>-</sup> $\delta$ </sup> nanosheets exhibited high reversible capacity and great rate capability with the capacities of 283.9 (50 mA  $g^{-1}$ ) and 179.3 mAh  $g^{-1}$  (1 A  $g^{-1}$ ) after 100 cycles. Rod-like  $\alpha$ -MoO<sub>3</sub> is successfully designed by alternating voltage induced electrochemical synthesis (AVIES) approach with subsequent control of the morphology and texture of MoO<sub>3</sub> by applying various calcination temperature of 300–600  $^{\circ}$ C [292]. When utilized for SIBs, it presents the high charge capacities of 305 mAh  $g^{-1}$  at 1C rate and 143 mAh  $g^{-1}$  at 10C. Even after 3000 cycles at 10C, a significant capacity of 108 mAh  $g^{-1}$ is still delivered. Zhu et al. [293] fabricated an amorphous  $MoS<sub>2</sub>/MO<sub>3</sub>/nitrogen-doped carbon$ composite using a reaction that partially sulfurizes organic–inorganic hybrid material  $Mo<sub>3</sub>O<sub>10</sub>$  $(C_2H_{10}N_2)$  (named as MoO<sub>y</sub>/ethylenediamine) nanowire precursors at low temperature of 300 °C. Tested in sodium cell, the composite retains 539 mAh  $g^{-1}$  after 200 cycles at 300 mA  $g^{-1}$ and maintains 340 mA h g<sup>-1</sup> at 1000 mA g<sup>-1</sup> after 220 cycles. The  $\alpha$ -MoO<sub>3</sub> nanosheets vertically grown on activated carbon fiber cloth exhibit a discharge capacity of 2.5 mAh cm<sup>-2</sup> (1621 mAh  $g^{-1}$ ) at 0.1 mA cm<sup>-2</sup> and capacity retention of 90% after 200 cycles at 0.2 mA cm<sup>-2</sup> [294]. Zhang et al.  $[259]$  prepared MoO<sub>3</sub>/reduced graphene oxide (MoO<sub>3</sub>/rGO) composites via one-step hydrothermal method, and evaluated as anode materials for SIBs. The  $MoO<sub>3</sub>/rGO$  composite

with 6 wt.% rGO delivers the highest reversible capacity of ~208 mAh  $g^{-1}$  at 50 mA  $g^{-1}$  after 50 cycles. **Table 7** summarizes the electrochemical performance of MoO<sup>3</sup> composites as anode materials for SIBs prepared by various methods.



**Table 7.** Electrochemical performance of MoO<sub>3</sub> composites as anode materials for SIBs prepared by various methods. Relevant cycle number is given in brackets.

#### 2.2. Monoclinic β-MoO<sup>3</sup>

 $MoO<sub>3</sub>$  in its β-modification presents a great interest in many applications such as catalysts [297,298], gas sensors [299] and batteries [300]. It was revealed that β- MoO<sub>3</sub> exhibits better response than that of α-MoO<sub>3</sub> in some catalysis reactions due to the high acidity of Lewis acid sites on the surface of β-MoO<sub>3</sub> [301]. Therefore, a metastable β-MoO<sub>3</sub> which has been reported to have high catalytic property in methanol oxidation to formaldehyde [302] due to the overall symmetry of this phase.

# *2.2.1. Structure*

Since 1986, the β-phase MoO<sub>3</sub> with distorted 3D ReO<sub>3</sub>-related structure similar to WO<sub>3</sub> is known from the work of McCarron [46,303,304]. This novel metastable phase with monoclinic symmetry results from the thermal treatment of spray-dried powders of aqueous molybdic acid solutions [46]. The β-MoO<sub>3</sub> phase (*P*2<sub>1</sub>/c S.G., *a* = 7.118 Å, *b* = 5.366 Å, *c* = 5.568 Å, β = 91.99°, and JCPDS: 00-047-1081) has a crystal structure similar to  $\text{ReO}_3$  (cubic,  $Pm\overline{3}m$ ), which possesses a three-dimensional network of  $\text{ReO}_6$  octahedra sharing the six corners. Its structure

is markedly different from the crystal structure of  $\alpha$ -MoO<sub>3</sub>, in which the MoO<sub>6</sub> octahedral units share corner oxygen atoms in the direction of the *c*-direction, and edge sharing occurs in the direction of the *c*-axis (inset **Fig. 19a**). The absence of van der Waal forces is due to adjacent MoO<sup>6</sup> octahedra sharing corners, which form a 3D monoclinic network. As shown in (**Fig. 19a**) the XRD pattern of β-MoO<sub>3</sub> is characterized with two dominant peaks at  $2\theta = 22.9^\circ$  and 24.9° corresponding to the (001) and (200) planes, respectively. **Fig. 19b** shows the Raman spectrum of β-MoO<sub>3</sub>, which differs than the α-MoO<sub>3</sub> spectral features. The peaks at 775, 840 and 904 cm<sup>-1</sup> are the Raman profiles corresponding to the Mo-O stretching modes of β-MoO<sub>3</sub>. The absence of peak in the high-frequency range ( $v > 904$  cm<sup>-1</sup>) indicates that no oxygen atoms are singly coordinated [305].

Phuc et al. [306] reported the existence of a new metastable β-MoO<sub>3</sub> green phase synthesized from MoO<sub>2</sub>Cl<sub>2</sub> solution derived from H<sub>2</sub>MoO<sub>4</sub> powder and concentrated HCl solution. This new phase crystallizes in a triclinic structure similar to that of  $WO<sub>3</sub>$  and is stable up to 435 °C. The XRD pattern exhibits three main reflection at 22.9°, 24.0°, and 24.9°, which are indexed as (002), (020) and (200) of the triclinic symmetry ( $P\overline{1}$  space group) [307]. The Raman spectrum of the β-MoO<sub>3</sub> green phase displays Mo—O—Mo vibration modes located at 767, 842 and 898 cm<sup>-1</sup> along with a weak peak at 708 cm<sup>-1</sup>, which does not exist in the spectrum of the β-MoO<sub>3</sub> yellow phase. The β'-MoO<sub>3</sub> phase was prepared by heating the D<sub>0.99</sub>MoO<sub>3</sub> intercalate in oxygen at 200 °C, driving off D<sub>2</sub>O. The structure of  $\beta$ '-MoO<sub>3</sub> was determined from neutron powder diffraction as monoclinic  $(P2_1/n \text{ S.G.}, a = 7.4245(1) \text{ Å}, b = 7.4783(1) \text{ Å},$  $c = 7.6897(1)$  Å,  $\beta = 90.090(2)°$  is isostructural with the room temperature monoclinic modification of  $WO<sub>3</sub> [47]$ .



**Fig.** 19. (a) XRD pattern of the β-MoO<sub>3</sub> yellow phase ( $P2<sub>1</sub>/c$  space group). (b) Raman spectrum recorded with the excitation line at  $\lambda = 532$  nm. (c) GCD profile carried out at C/10 rate. (d) Differential capacity plot (-d*Q*/d*V*) vs. *V*.

### *2.2.2. Synthesis*

The synthesis of the metastable  $\beta$ -MoO<sub>3</sub> phase is limited due to the difficulty in finding suitable molybdenum source such as ammonium molybdate, molybdic acid, polyoxomolybdate. The first successful preparation of  $\beta$ -MoO<sub>3</sub> was reported by McCarron [46] who employed a soft-chemical route. The synthesis consists of the dehydration of blue molybdic acid ( $Mo_{36}O_{112}(H_2O)_{16}$ )<sub>8</sub> solution, which is obtained via cation exchange of Na<sub>2</sub>MoO<sub>4</sub> solution (1 mol  $L^{-1}$  of Na<sub>2</sub>MoO<sub>4</sub> only gives 0.1 mol  $L^{-1}$  of blue molybdic acid solution). Mizushima et al. [301] reported that this method consumes a lot of water and also has very low yield, i.e., the Mo content in the final solution was only 3.3% of the initial value. In 2011, Mizushima et al. synthesized the  $\beta$ -MoO<sub>3</sub> phase (yellow powders) by a soft chemical transformation of  $α$ -MoO<sub>3</sub> via the addition of HNO<sub>3</sub> to an aqueous solution of  $α$ -MoO<sub>3</sub>, followed by drying and calcination [308]. Yao et al. synthesized  $\beta$ -MoO<sub>3</sub> from sodium molybdate solution [299] but α-MoO<sub>3</sub> was detected in the final product. β-MoO<sub>3</sub> crystal has a yellow color [46,306,309-311]. Moreover, β-MoO<sub>3</sub> can reportedly be obtained by a novel method of freeze drying the same molybdic acid solution [312]. These methods yield a mixture of α-MoO<sup>3</sup> and β-MoO3. Juárez Ramírez and Martínez-de-la-Cruz [310] produced pure β-MoO<sup>3</sup> free from the α phase through vacuum drying. This method, however, requires a very long drying time. In 2005, Mizushima et al. [311] reported a new preparation method for pure

β-MoO<sup>3</sup> using a generalized rotary evaporator. The simple evaporation of a cation-exchanged molybdic acid solution forms pure  $\alpha$ -MoO<sub>3</sub> crystallites (bright yellow powder) after calcination at 573 K, whereas the addition of a small amount of nitric acid to the solution resulted in selective formation of the β-phase. The preparation of  $\beta$ -MoO<sub>3</sub> thin films has reported by several workers. Porous high surface area thin films of nanosheet-shaped (50-100 nm thick) monoclinic β-MoO<sub>3</sub> were deposited onto platinized Si substrates using patch antenna-based atmospheric microplasma processing [313]. β**-**MoO<sup>3</sup> thin films were prepared by reactive rf sputtering deposition at low substrate temperature of 200  $^{\circ}$ C under a 65%Ar+35%O<sub>2</sub> gas mixture at the pressure of 0.13 Pa [314]. Nearly a pure β-phase was formed (i.e., well-resolved XRD reflections at ca.  $2\theta = 23.0, 25.0, 47.5$  and  $51.8^{\circ}$  corresponding to the (011), (200), (022) and (400) of the  $P2<sub>1</sub>/c$  structure, respectively), which exhibits a preferred *a*-axis orientation when the annealing temperature is in the range of  $300-350$  °C. MoO<sub>3</sub> films were electrodeposited on fluorine-doped tin oxide (FTO) glass substrates from sodium-molybdate (Na<sub>2</sub>MoO<sub>4</sub>) solutions. The monoclinic β-phase was obtained by manipulating the cyclic voltammetry conditions  $[299]$ . Characterization of the MoO<sub>3</sub> films grown by molecular beam epitaxy on *c*-plane sapphire substrates was conducted by Koike et al. [315]. XRD and Raman measurements revealed that amorphous, (100) β-phase, and (010) α-phase MoO3 films were preferentially grown at 150, 200, and 350 °C, respectively. Their optical bandgap energies were estimated to be ~3.5 eV for the amorphous, ~3.7 eV for the β-phase, and ~4.1 eV for the  $\alpha$ -phase films. The β-phase films were completely transformed into stable α-phase films at 600 °C.

A transformation from the  $\beta$  to  $\alpha$  phase took place spontaneously at the temperature ranging from 387 to 450 °C, according to several reports [304,316-318]. Topotactic transition of metastable β-MoO<sub>3</sub> to orthorhombic  $\alpha$ -MoO<sub>3</sub> has been reported as a result of thermal treatments at temperature 350 °C and above. The presence of the β-phase of MoO<sub>3</sub> at low temperature ~250 °C is supported by Kuzmin [316], Julien [317] and McEvoy [318], who have shown that sintering in air of hydrated, amorphous molybdenum oxide at relatively low temperatures produces a morphological change from an amorphous phase to a metastable, monoclinic β**-**MoO3. Raman microprobe spectroscopy experiments have revealed that two phases of MoO<sup>3</sup> coexist. Some regions exhibit characteristic Raman shifts of 997, 821, and 668 cm $^{-1}$  that are unique to  $\alpha$ -MoO<sub>3</sub> while other regions exhibit Raman stretches at 850 and 776 cm<sup>-1</sup> that are specific to  $\beta$ -MoO<sub>3</sub> [319].

#### *2.2.3. Li intercalation*

Pristine β-MoO<sub>3</sub> exhibits an electrical conductivity below  $10^{-7}$  S cm<sup>-1</sup> [320], which implies to build an electrode using of carbon additive. In 2003, Juárez Ramírez and Martínez-de-la-Cruz demonstrated the ability of  $β$ -MoO<sub>3</sub> obtained through soft chemistry methods to accommodate electrochemically Li<sup>+</sup> ions [300,310]. Electrochemical experiments were carried out in Swagelok-type cells using Li foil as negative electrode and a 1 mol  $L^{-1}$  LiPF<sub>6</sub> in ethylene carbonate (EC)/dimethyl carbonate (DMC) as electrolyte. The cathode electrode was made of β-MoO<sup>3</sup> powders, carbon black and ethylene–propylene–dieneterpolymer (EPDT) in a 90:9:1 ratio. Lithium insertion in β-MoO<sub>3</sub> proceeds at least in a two-step reduction process. The maximum amount of lithium inserted leads to a specific capacity of 370 mAh  $g^{-1}$  (gravimetric energy density of 666 Wh  $kg^{-1}$ ). However, a total loss of 25% occurs after the first chargedischarge cycle due to structural transformations with variable  $Li_xMoO_3$  composition. For several discharge/charge cycles, the Li/ $\beta$ -MoO<sub>3</sub> cell is able to maintain 280 mAh g<sup>-1</sup> at 400  $\mu$ A cm<sup>-2</sup> current density. Porous β-MoO<sub>3</sub> nanosheets (50-100 nm thick) produced by atmospheric microplasma were applied to LIBs [313]. The insertion/de-insertion of Li<sup>+</sup> ions was performed in 1 mol  $L^{-1}$  LiN(SO<sub>2</sub>CF<sub>3</sub>)<sub>2</sub> in EC:DEC (1/1, v/v) electrolyte. The cyclic voltammogram displays cathodic peaks at 2.2 and 1.6 V and the anodic peaks at 1.7 and 2.1 V. At galvanostatic current of 10  $\mu$ A g<sup>-1</sup>, the discharge process proceeds through several steps and two-phase regions are observed at 1.7 and 2.7 V, respectively. The charge capacity is 370 mAh  $g^{-1}$  and corresponds to a mole fraction of 2 Li atoms per Mo atom. McEvoy et al. [321] investigated the Li-ion insertion energetics and kinetics of β-MoO<sub>3</sub> films prepared by electrodeposition from acidic peroxo-polymolybdate solutions. Films deposited on ITO-coated glass substrates and sintered in the temperature range 250-350 °C display an intermixed  $\alpha$ -β-MoO<sub>3</sub> structure. In LiClO<sub>4</sub>/propylene carbonate solution, the electrochemical response for films sintered at  $T \ge 250$ °C exhibits three sets of distinguishable redox peaks. This response is assigned to the faradaic electron transfer and to a large capacitive charge during Li<sup>+</sup> insertion. The smaller anodic peak area after the first cycle suggests that the insertion/deinsertion process is only 60% reversible. It is more likely due to some fraction of  $Li<sup>+</sup>$  becoming trapped in the film due to the irreversible formation of a molybdenum bronze (i.e.,  $Li_{x+2}Mo_{1-x}^{VI}Mo_{x-z}^{V}Mo_{z}^{IV}O_{3-y-z}$ ). **Figure 19c** shows the typical first discharge charge curve of a β-MoO<sub>3</sub> film prepared by pulsed-laser deposition on silicon wafer at  $T_s = 250$  °C. The OCV of the Li/ $\beta$ -MoO<sub>3</sub> cell is 3.20 V vs. Li<sup>+</sup>/Li. In the potential range 3.2-1.0 V, the transfer of 1.92 e<sup>-</sup>/Mo occurs during the discharge at C/10 rate (36 mA  $g^{-1}$ ) yielding a specific capacity of 357 mAh  $g^{-1}$ . The first charge occurs with a coulombic efficiency of 83%. The two distinct sets of redox reaction are determined by the plot

of the differential capacity (-d*Q*/d*V*) vs. *V* at 2.66/1.72 V and 2.90/2.04 V for the anodic and cathodic peaks, respectively (**Fig. 19d**).

### 2.3. Hexagonal h-MoO<sup>3</sup>

Hexagonal molybdenum trioxide (h-MoO<sub>3</sub>) is known for one century when, in 1906, Rosenheim described the preparation of a molybdic acid hydrate by precipitation of an aqueous solution of ammonium paramolybdate,  $(NH_4)_6[M_07O_{24}]_3$  4H<sub>2</sub>O, acidified with HNO<sub>3</sub> [322]. In 1969, the Roseheim's process was used by Peters for the XRD analysis of the isolated precipitates [323]. In 1980s, several workers revised the structure of hexagonal molybdates [324,325]. Caiger et al. [325] reported that the "hexagonal MoO<sub>3</sub>" series corresponds to the partially dehydrated and de-ammoniated ammonium decamolybdate  $NH<sub>4</sub>Mo<sub>5</sub>O<sub>15</sub>OH·2H<sub>2</sub>O$ . Mc Carron et al. investigated the crystal structure of h-MoO<sub>3</sub> with incorporation of sodium and proton in the tunnels  $[324]$ . Functional modifications and diversified applications of h-MoO<sub>3</sub> can be found in Refs. [325-331]

# *2.3.1. Structure*

The framework of hexagonal h-MoO<sub>3</sub> is constructed from zigzag chains of MoO<sub>6</sub> octahedra linked to each other by corner sharing along the *c* direction giving large one-dimensional tunnels (**Fig. 20a**). The h-MoO<sub>3</sub> phase can generally be formulated as  $(A_2O)_x \cdot \text{MoO}_3 \cdot (H_2O)_y$ , where *A* is an alkali-metal ion or ammonium ion, and the exact values of *x* and *y* depend on the details of the preparation and subsequent treatment [101,332-334]. The salient structural feature for crystalline h-MoO<sub>3</sub> is the presence of tunnel ( $\sim$ 3 Å in diameter) running along the *c* axis, in which cations or water molecules can be sitting. Each tunnel is enclosed by  $12 \text{ MoO}_6$  octahedra linked by sharing corners along the *a* and *b* directions and by edges along the *c* direction (**Fig. 20a**). The tunnel structure in h-MoO<sub>3</sub> exhibits better intrinsic properties than α-MoO<sub>3</sub>, such as high sensitivity, coloration efficiency, and faster response. The sample prepared via the precipitation of molybdenum oxide from an ammonium paramolybdate solution, by the addition of nitric acid has a "sea-urchin" shape composed of  $h-MoO<sub>3</sub>$  nanorods (SEM images in Fig. 20b) [335]. Figure 20c depicts the XRD pattern of crystalline h-MoO<sub>3</sub>. The diffraction peaks appeared at  $2\theta = 9.62^{\circ}$ , 16.72°, 19.34°, 25.68°, 29.26°, 33.86° and 35.32° can be attributed to the planes of  $(100) (110) (200) (210) (300) (220)$  and  $(310)$ , respectively of the hexagonal structure (space group  $P6_3/m$ ) with lattice parameters  $a=10.53 \text{ Å}$  and  $c=14.97 \text{ Å}$  (JCPDS card No. 21-0569) [13,27,28,332,335]. However, using Rietveld refinements, XRD reflections were successfully indexed with  $a=10.55$  Å and  $c=3.72$  Å [48]. A structural transformation from hexagonal to stable orthorhombic α-MoO<sub>3</sub> phase was observed at 350 °C [107]. The publication by Lunk et al. [334] aimed to dispel the confusion concerning the identity of the hexagonal molybdenum trioxide (HEMO) phases variously described in the literature. The reported HEMO composition are characterized by virtually identical XRD patterns with unit cell parameters a  $\approx$  10.60 Å and c  $\approx$  3.72 Å. As occurred in the case of MnO<sub>2</sub> hollandite, the presence of large cations into the tunnel can stabilize the h-MoO<sub>3</sub> structure  $[337]$ . However, the growth of crystalline h-MoO<sup>3</sup> without large cations incorporated into the tunnels can be obtained from two methods: one is the hot liquid-phase process [335], and the other is the liquid-phase process followed by autoclaving and vacuum heat-treatment [27,332]. The crystallinity and phase purity of as-synthesized nanorods were evidenced by micro-Raman spectrum (**Figure 20d**), which exhibits sharp high-frequency peaks at 980, 915, 902 and 888 cm<sup>-1</sup> assigned to the symmetrical and asymmetrical stretching modes of Mo=O double bond. The peak at 818 cm<sup>-1</sup> is assigned to the stretching mode of Mo<sub>2</sub>-O and the band at 694 cm<sup>-1</sup> is the scissoring vibration of the O-Mo-O bonds. Raman peaks at 395, 412 and 495 cm<sup>-1</sup> are attributed to the bending modes of O-Mo-O bonds, while the peaks at 247 and 220 cm<sup>-1</sup> are assigned to the twisting vibration of O-Mo-O bonds  $[48,338,339]$ . Raman peaks in the low-frequency (174 and 145 cm<sup>-1</sup>) belong to the external lattice modes.



**Fig. 20**. (a) Crystallographic structure of h-MoO3. The unit cell is shown by a thin black line. (b) SEM images of h-MoO<sup>3</sup> nanorods: projected view (top) and side view (bottom). Reproduced with permission from [335]. Copyright 2009 Elsevier. (c) XRD pattern of h-MoO<sub>3</sub> with

reflection indexed with *P*6<sub>3</sub>/*m* S.G (JCPDS card No. 21-0569). (d) Raman spectrum of h-MoO<sub>3</sub> recorded with the laser excitation  $\lambda_{\text{exc}} = 532$  nm. (e) Galvanostatic discharge-charge profiles recorded at 50  $\mu$ A cm<sup>-2</sup> current density for h-MoO<sub>3</sub> electrode with empty tunnels and h-Ag0.2MoO3.1 electrode. Reproduced with permission from [340]. Copyright 1995 Elsevier.

### *2.3.2. Synthesis*

The metastable hexagonal form of  $h-M<sub>0</sub>O<sub>3</sub>$  has been successfully prepared starting from the ammonium molybdate, using a "chimie douce" reaction [107,332,341-345]. In the precipitation method, a mixture of ammonium heptamolybdate tetrahydrate  $(NH_4)_6M_0T_2^4$  4H<sub>2</sub>O and concentrated nitric acid (37% HNO<sub>3</sub>) heated to a temperature in the range 55-85 °C was used [13,107,346]. Hydrothermal technique has been widely used for the synthesis of h-MoO<sub>3</sub> [332,333,347]. Zheng et al. [332] prepared h-MoO<sub>3</sub> nanobelts by a simple hydrothermal route from peroxomolybdate solution with the presence of sodium nitrate as a mineralizer. It was demonstrated that the high openness degree in the tunnel structure of h-MoO<sub>3</sub> provides larger spatial locations for cation insertion/extraction and diffusion. Song et al. prepared the high purity h-MoO<sub>3</sub> phase via chemical precipitation, hydrothermal treatment and calcination. In the procedure of precipitation, the concentrated HCl played a key role for the formation of the metastable h-MoO<sub>3</sub> [13]. Atuchin et al. [28] prepared h-MoO<sub>3</sub> nanocrystals in the form of well-faceted straight hexagonal rods with an aspect ratio of  $1/d \approx 60$ . Han et al. [348] demonstrated that rod-shaped h-MoO<sub>3</sub> were obtained by hydrothermal method using a low concentration of nitric acid  $(0.1-1.0 \text{ mol L}^{-1} \text{ HNO}_3)$ , while higher concentration leads the nanostructured α-MoO<sub>3</sub> phase. Hexagonal and truncated hexagonal shaped MoO<sub>3</sub> nanoplates (MoO<sup>3</sup> HNP) were synthesized through a simple vapor-deposition method in Ar atmosphere under ambient pressure without the assistant of any catalysts. The nanoplates have a large area surface. The Raman spectrum shows a significant size effect on the vibrational property of MoO<sub>3</sub> HNP [344]. Hexagonal MoO<sub>3</sub> nanoparticles were synthesized by hydrothermal method  $(T = 90$  °C for 12 h) from a solution of 0.2 mol L<sup>-1</sup> ammonium molybdate in 10 ml of distilled water added with 5 ml of concentrate HNO<sub>3</sub>. When heated to  $T > 150$  °C, h-MoO<sub>3</sub> transforms to the layered α-MoO<sub>3</sub> phase [107,326]. Using chemical precipitation, the h-MoO<sub>3</sub> phase with unique rod-like morphology was synthesized by Song et al. [13]. Guan et al. [349] obtained a nanodisk of hexagonal h-MoO<sub>3</sub> by heat treatment of rhodamine B/molybdic acid aggregates (RBMA). Dhage et al.  $[29]$  fabricated *h*-MoO<sub>3</sub> nanorods via the probe sonication route using ammonium molybdate, urea and sodium dodecyl sulphate at 70 °C. Irmawati and Shafizah  $[344]$  produced pure h-MoO<sub>3</sub> nanocrystals through the acid washing method, which consists in the reduction of the basic aqueous solution of ammonium heptamolybdate tetrahydrate and ammonium carbonate from pH of 8.85 to 1.95-2.10 by addition of 5 mol  $L^{-1}$  HNO<sub>3</sub> (lattice parameters  $a = 10.55 \text{ Å}$  and  $c = 14.89 \text{ Å}$ ). Pan and coworkers [48] prepared nano- and microrods h-MoO<sub>3</sub> using modified liquid-phase processes with concentrated  $HNO<sub>3</sub>$  and  $H<sub>2</sub>SO<sub>4</sub>$  and determine the hexagonal structural patterns as  $a \approx 10.57$  Å and  $c \approx 3.72$  Å. The fabricated rods have a BET total surface area of  $\sim$ 3 m<sup>2</sup> g<sup>-1</sup>, with crystallite size ranging from 33 to 46 nm. Zheng et al.  $[332]$  synthesized h-MoO<sub>3</sub> nanobelts by a simple hydrothermal route from peroxomolybdate solution with the presence of sodium nitrate as a mineralizer.

#### *2.3.3. Li intercalation*

Due to the many empty sites in the tunnel structure, i.e., hexagonal tunnel and trigonal cavity (Fig. 20e), the h-MoO<sub>3</sub> electrode exhibits a high capacity on Li<sup>+</sup> battery application [324,325,332,340,350-351]. In the early work by Gao et al. [340], the lithium intercalation/deintercalation reaction was studied on  $h-MoO<sub>3</sub>$  with empty tunnels and Agcontaining tunnels. In the discharge curves carried out at  $30 \mu A \text{ cm}^{-2}$  in the potential range 1.7-3.5 V vs. Li<sup>+</sup>/Li (Fig. 20e), the voltage plateau at  $x = 0.7$  indicates a two-phase region with lattice parameters  $a = 10.559(5)$  Å and  $c = 3.725(3)$  Å (i.e., 0.97% volume change). For the hexagonal  $Ag_{0.2}MoO_{3.1}$  electrode, the ICE was reduced to ~60%. Electrochemical tests, carried out at current density of 0.1 mA  $cm^{-2}$  in the voltage range 1.2–4.0 V vs. Li<sup>+</sup>/Li, show excellent performances with an initial reversible discharge specific capacity of  $402 \text{ mA} \text{h} \text{g}^{-1}$ , i.e., insertion of 2.2 Li/Mo. Xu et al. reported a first discharge voltage plateau at ~1.8 V for hydrothermally synthesized h-MoO<sub>3</sub> [352]. Tang et al. [353] fabricated h-MoO<sub>3</sub>/graphene composites using a facile hydrothermal method via a template effect of GO on catalyzing the phase transition from α-MoO<sup>3</sup> to h-MoO3. The galvanostatic charge/discharge tests of this composite as anode of Li cell exhibit a first reversible capacity of 987 and 739 mAh  $g^{-1}$  after 30 cycles under the current density of 100 mA  $g^{-1}$  in the voltage range of 0.01–3 V vs. Li<sup>+</sup>/Li. Recently, Joseph et al. [354] fabricated MoO<sub>3</sub> microrods with hexagonal structure as an Al-ion intercalation host material and show its use as an electrode for an aqueous Al-ion battery. h- $MoO<sub>3</sub>$  microrods yield a discharge capacity of ~300 mAh  $g^{-1}$  for 150 cycles at a current density of 3 A  $g^{-1}$  and 90% capacity retention after 400 cycles. Xiao et al. [355] used a solution-based method to synthesized 2D h-MoO<sub>3</sub> nanosheets Mo was dissolved in  $H_2O_2$  to produce the molecular peroxomolybdate precursor solution mixed with NaCl microcrystals as lattice-matched templates. 2D h-MoO<sub>3</sub> was tested in a 1 mol  $L^{-1}$  LiClO<sub>4</sub> solution with a 1:1 mixture of ethylene carbonate and dimethyl carbonate. Intriguingly, a gravimetric capacity of 996 C  $g^{-1}$  (277 mAh

 $g^{-1}$ , 1.49 Li/Mo) at 2 mV s<sup>-1</sup> was much larger than that previously reported for mesoporous  $\alpha$ - $MoO<sub>3</sub>$  in 1 mol L<sup>-1</sup> LiClO<sub>4</sub> with propylene carbonate.

The coloration properties from facile Li<sup>+</sup> ion insertion/extraction in electrochromic devices was demonstrated by several workers. Zheng et al. [332] showed that the performances concerning photochromism on two types of  $MoO<sub>3</sub>$  nanobelt suspensions show that the photochemical efficiency of h-MoO<sub>3</sub> is more excellent than that of α-MoO<sub>3</sub>. Electrochromic performance of the h-MoO<sup>3</sup> nanobelts coated film was tested in a three-electrode cell cycled between -1.0 and -2.0 V vs. SCE with 1 mol  $L^{-1}$  LiClO<sub>4</sub> in propylene carbonate as electrolyte and graphite plate as the counter electrode. Two strong and wide transmittance peaks at ∼620 and 590 nm imply that the  $MoO<sub>3</sub>$  nanobelt film can display from green-blue to dark blue color at different voltages. Huang et al.  $[211]$  reported the synthesis of h-MoO<sub>3</sub> nanorods (average length of 20-50 mm and a width of 1-10 mm) using a hydrothermal treatment at 180 °C for 24 h of a solution of  $(NH_4)_6M_0T_2^4$ -4H<sub>2</sub>O adjusted at pH~1 by addition of 8 mol L<sup>-1</sup> HNO<sub>3</sub>. Electrochemical properties of h-MoO<sub>3</sub> (Fig. 21) were examined as anode for LIB over 100 cycles at 0.1C rate. The discharge capacity of 780 mAh  $g^{-1}$  (2nd cycle) rapidly drops to 132 mAh  $g^{-1}$  after 100 cycles, merely 14% of the 2nd cycle. The structure of h-MoO<sub>3</sub> leads to poor capacity stability. After 20 cycles, the hexagonal framework transforms to agglomerated bulk structure with slow Li<sup>+</sup>/e<sup>-</sup> transfer kinetic and finally nanorods are pulverized after 100 cycles (**Fig. 21d**). **Table 8** lists the electrochemical performance of some h-MoO<sup>3</sup> anode materials for LIBs. Electrochemical properties of h-MoO<sub>3</sub> have been also investigated by Song et al. [350]. The first reversible discharge specific capacity can reach  $402 \text{ mA}$ h g<sup>-1</sup> (2.2 Li/Mo uptake) at 0.1 mA cm<sup>-2</sup> current density in the voltage range  $1.2-4.0$  V *vs.* Li<sup>+</sup>/Li.



**Fig. 21**. (a) Galvanostatic charge-discharge curves in different cycles carried out at 0.1C rate of h-MoO<sub>3</sub>. (b) Differential capacity vs; voltage curves. (c) Rate capability of h-MoO<sub>3</sub> at rates between 0.1 and 2C. (d) Schematic illustration of the structural changes upon cycling. Reproduced with permission from [211]. Copyright 2016 Elsevier.

Material	Synthesis	Specific capacity $(mAh g-1)$	Current rate $(mA g^{-1})$	Ref.
$h-MoO3 NRs$	Hydrothermal	132	0.1C(100)	[211]
$h-MoO3/graphene$	Hydrothermal	739	100(30)	[353]
$h-MoO3 NRs$	Solvothermal	780	0.1C(150)	[356]

**Table 8.** Electrochemical performance of h-MoO<sub>3</sub> anode materials for LIBs.

### 2.4. Monoclinic  $MoO<sub>3</sub>$ -II phase

The metastable  $MoO<sub>3</sub>-II$  ( $\varepsilon$ -MoO<sub>3</sub>) phase was highlighted by Mc Carron and Calabrese [357] from the combination of high pressure and high temperature (60 kbar and 700  $^{\circ}$ C), using a tetrahedral anvil apparatus at elevated temperature. Using a soft-chemical synthesis of the high-pressure phase of molybdenum trioxide,  $MoO<sub>3</sub>-II$ , has been described by Baker et al. [49]. A topotactic dehydration of either the white molybdenum trioxide monohydrate MoO3∙H2O, or the hemihydrate MoO3∙½H2O provides a convenient synthetic route to the high-pressure phase. The structural filiations between the various molybdenum trioxide and trioxide hydrate phases are delineated, and simple mechanistic models for the transformations have been proposed, which involves a change in the stacking sequence of the Mo-O layers. MoO<sub>3</sub>-II is the highpressure modification of  $\alpha$ -MoO<sub>3</sub>. Like the  $\alpha$ -MoO<sub>3</sub> layered structure, the individual MoO<sub>3/3</sub>O<sub>2/2</sub>O<sub>1/1</sub> layers of MoO<sub>3</sub>-II and  $\alpha$ -MoO<sub>3</sub> are virtually identical. The stacking sequence of the layers of MoO3-II (*AAA* stacking) differs from that of α-MoO<sup>3</sup> (*ABA* stacking), which is equated with an improved packing efficiency for the layers of MoO<sub>3</sub>-II versus those of  $\alpha$ -MoO<sub>3</sub> (**Fig. 22**). Its structure is monoclinic,  $P2_1/m$  S.G., with unit cell parameters  $a = 3.954(1)$  Å,  $b =$ 3.687(2) Å,  $c = 7.095(4)$  Å and  $\beta = 103.75(4)^\circ$  (4.75 g cm<sup>-3</sup>). In contrast, Åsbrink et al. [358] reported the lattice parameters of  $α$ -MoO<sub>3</sub> as a function of pressure up to 7.4 GPa at ambient temperature, but did not find any high-pressure phase transition. Liu et al. [44] demonstrated that the MoO3-II phase can be achieved at 12 GPa and room temperature. The lattice parameters refined within the  $P2_1/m$  space group for the new phase were  $a=3.860(3)$  Å,  $b=3.711(6)$  Å,  $c$  $=6.329(4)$  Å,  $\beta = 104.811^{\circ}$ , and unit cell volume *V*=87.66(5) Å<sup>3</sup> (*Z*=2). The metastable MoO<sub>3</sub>-II phase is relatively more stable than  $h-MoO<sub>3</sub>$  but can convert rapidly to stable orthorhombic phase α-MoO<sub>3</sub> at temperatures above 200 °C [8,357]. Another high-pressure phase, the monoclinic MoO<sub>3</sub>-III phase  $(P21/c$  S.G.), has been observed at pressure of 25 GPa at room temperature. The lattice parameters refined within the  $P2<sub>1</sub>/c$  space group for the MoO<sub>3</sub>-III phase were  $a=5.749(2)$  Å,  $b = 4.804(6)$  Å,  $c=6.095(3)$  Å,  $\beta =116.73(5)$ °, and unit cell volume *V*=150.36(5)  $\AA$ <sup>3</sup> (*Z*=4). The Raman spectrum of MoO<sub>3</sub>-III did not display the peak associated with the unique molybdyl bond ( $M_0=O$  stetch) confirming this phase is not a layered structure. The stretching modes at 786, 834 and 893 cm<sup>-1</sup> reveal the Mo-O bonds in the corner-sharing  $MoO<sub>6</sub> octahedra.$  The  $MoO<sub>3</sub>$ -III phase is likely a distortion of the  $ReO<sub>3</sub>$ -type structure [44].



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**Fig.** 22. Crystallographic structures of molybdenum trioxides. (a) Orthorhombic  $\alpha$ -MoO<sub>3</sub> structure with *ABA* stacking. (b) Monoclinic  $P2_1/m$  MoO<sub>3</sub>-II structure with *AAA* stacking. Mo atoms of the *B* layer within MoO<sub>6</sub> a mirror plane compared with  $\alpha$ -MoO<sub>3</sub>. (c) Monoclinic *P*2<sub>1</sub>/*c* structure of the high-pressure phase MoO3-III. Purple circles represent the Mo atoms and red circles the O atoms. Reproduced with permission from [44]. Copyright 2009

#### 2.5. Bronzes of Molybdenum

In 1895, the first molybdenum bronze was synthesized by Stavenhagen and Engels [359]. The sodium molybdenum bronze Na<sub>6</sub>Mo<sub>7</sub>O<sub>24</sub> (3NaO∙7MoO<sub>3</sub>) was produced by electrolytic reduction of fused sodium molybdate. Under water, the bronze appears as indigo blue, shiny metal needles that lost some of their metallic sheen when drying and turned a dark blue color. In 1970, Réau et al. [69] investigated the monoclinic  $Li<sub>x</sub>MoO<sub>3</sub>$  phases, which refer to violetblue bronzes prepared in a range of stoichiometry with  $0.31 \le x \le 0.39$ . The main interest of molybdenum bronzes was motivated by their transport anomalies (charge-density wave instabilities). In the 1980s, molybdenum bronzes were studied as low-dimensional metallic conductors, with the generic formula  $A_xMo_yO_z$  where *A* is hydrogen or alkali metal. They are distinguished by their color: red bronzes ( $Li_{0.33}MoO<sub>3</sub>$ ,  $K_{0.33}MoO<sub>3</sub>$ ), blue bronzes ( $K_{0.3}MoO<sub>3</sub>$ ), and purple bronzes  $(Li<sub>0.9</sub>Mo<sub>6</sub>O<sub>17</sub>, Na<sub>0.9</sub>Mo<sub>6</sub>O<sub>17</sub>)$  [360-363]. In 1984, McCarroll and Greenblatt reported the synthesis of several single crystals of  $Li<sub>x</sub>MoO<sub>3</sub>$  bronze with different compositions: violet-red Li<sub>0.9</sub>Mo<sub>6</sub>O<sub>17</sub>, violet-blue Li<sub>0.32</sub>MoO<sub>3</sub> and blue Li<sub>0.04</sub>MoO<sub>3</sub> [364]. A lithium-ion battery was fabricated with a lithium molybdenum bronze as anode [365]. The molybdenum bronze Li<sub>x</sub>MoO<sub>3</sub>⋅εH<sub>2</sub>O with  $0.20 \le x \le 0.30$  and  $\varepsilon \le 0.1$  was prepared in a preferred embodiment, using a method that prevent the co-intercalation of proton. For instance, LixMoO3∙εH2O is formed from MoO<sup>3</sup> suspended in aqueous solution in argon atmosphere, by simultaneous addition of sodium dithionite and hydrated lithium molybdate. In the voltage range 3.5-0.005 V vs. Li, the composite electrode exhibits a capacity retention of 72% after 20 cycles.

### **3. Hydrated Mo-O phases**

## 3.1. Structure

A rich family of crystalline molybdenum(VI) oxide hydrates (MOHs), MoO3·*n*H2O, are known for  $1 \le n \le 2$ , which include the monoclinic dihydrate, MoO<sub>3</sub>·2H<sub>2</sub>O, two monohydrates, the yellow monoclinic  $MoO<sub>3</sub>·H<sub>2</sub>O$ , and the white triclinic  $MoO<sub>3</sub>·H<sub>2</sub>O$ , a white hemihydrate, MoO3·½H2O, and a white orthorhombic MoO3·⅓H2O (**Table 9**). The MoO3·⅔H2O hydrate has been also reported [366]. The structure of  $MoO<sub>3</sub>·2H<sub>2</sub>O$  is made of layers of  $MoO<sub>5</sub>(OH<sub>2</sub>)$  octahedra exchanging their oxygens in the equatorial plane and stacking along the *b*-direction [367]. Yellow monohydrate  $(\beta-MoO_3 \cdot H_2O)$  displays a structure with equatorial sharing oxygens and axial Mo-OH<sub>2</sub> and Mo=O bonds [368]. White monohydrate ( $\alpha$ -MoO<sub>3</sub>·H<sub>2</sub>O) has a triclinic structure (space group  $\overline{P1}$ , JCPDS card No. 16-1449) closely related to that  $\alpha$ -MoO<sub>3</sub> and crystallizes in well-shaped needles. Its structure is formed by edge-sharing octahedra parallel to the *b*-direction and includes  $Mo-OH_2$ ,  $Mo=O$  and  $Mo-O_3$  bonds [369]. The  $MoO<sub>3</sub>·½H<sub>2</sub>O$  structure is built of layers parallel to the b-direction formed of edge-sharing distorted MoO<sub>6</sub> octahedra and MoO<sub>5</sub>(OH<sub>2</sub>) octahedra [370]. MoO<sub>3</sub>·1/<sub>3</sub>H<sub>2</sub>O presents slightly distorted MoO<sub>6</sub> octahedra leading to a base-centered orthorhombic structure [41,309,371].

Compound	Space group	Cell parameters (A)		
		$a/\alpha$	$b/\beta$	$c/\gamma$
MoO <sub>3</sub> ·H <sub>2</sub> O	$P2\sqrt{c}$	7.55	$10.69/91.0^{\circ}$	7.28
$\alpha$ -MoO <sub>3</sub> ·H <sub>2</sub> O	PI	7.388	$3.70/113.6^{\circ}$	$6.673/91.6^{\circ}$
MoO <sub>3</sub> ·2H <sub>2</sub> O	P2/n	10.476	$13.822/91.6^{\circ}$	10.606
$MoO_3·\frac{1}{3}H_2O$	Phnm	7.697	12.647	7.338
$MoO3·\frac{1}{2}H2O$	P2/m	9.658	$3.71/102.4^{\circ}$	7.087
$h-MoO3·1/2H2O$	$P6_3/m$	10.584		3.728

**Table 9**. Crystallographic parameters of the various molybdenum oxide hydrates.

During the dehydration process of molybdic acid, several hydrated phases are formed. Yellow monoclinic mono-hydrate MoO<sub>3</sub>·H<sub>2</sub>O is formed by the topotactic loss of the intralayer water molecule on the dehydration of  $MoO<sub>3</sub>·H<sub>2</sub>O$  [368]. The white monoclinic hemi-hydrate  $(n=1/2)$  was prepared by precipitation from nitric acid solution [368], and by ion exchange method from sodium molybdate [366]. White orthorhombic MoO3·⅓H2O was grown by hydrothermal treatment at 110 °C of aqueous solution of molybdic acid [309]. By dehydration of molybdic acid, the monoclinic  $\beta$ -MoO<sub>3</sub> with ReO<sub>3</sub>-type structure is formed which transforms to the orthorhombic  $\alpha$ -MoO<sub>3</sub> phase above 400 °C [70,154].

The different steps are observed as endothermic peaks in TGA curves indicating multistep dehydration and structural rearrangements of the MOH framework (**Fig. 4**). Above 325 °C,  $\beta$ -MoO<sub>3</sub> and  $\alpha$ -MoO<sub>3</sub> phases are formed. A broad exothermic peak at 480 °C originates from the irreversible phase transformation from monoclinic to orthorhombic structure. The phase transformation is associated with loss of oxygen observed particularly for the samples heat

treated above 750°C leading sub-oxide such as  $MoO_{2.8}$  ( $Mo<sub>5</sub>O_{14}$ ) [121]. The DSC curve of MoO<sub>3</sub>·H<sub>2</sub>O shows three broad endothermic peaks at 198, 246, and 325 °C on heating in argon atmosphere at the rate  $2^{\circ}$ C min<sup>-1</sup>, which is correlated with the weight losses. At the first stage 33% of the water molecules was removed forming the product MoO3·⅓H2O.

Günter [368] reported that the dehydration process is a topotactic-like reaction with a slight expansion in both axial direction and an increase in symmetry; in molybdenum trioxide, the coordination octahedra partly share edges instead of corners only, which explains the change in the lattice constants and in symmetry. The low-temperature phase  $\beta$ -MoO<sub>3</sub> (bright yellow colour) is formed around 450 °C, while the layered  $\alpha$ -MoO<sub>3</sub> phase (white transparent) is formed at 750 °C. Some samples with black colour are formed at 750 °C, indicating an oxygen-deficient  $MoO<sub>3-δ</sub> structure. The dehydration mechanism of Mo mono hydrate (molybdic acid) can be$ summarized by the relation

$$
MoO3·H2O → MoO3·2⁄3H2O + 1⁄3H2O1 → MoO3·1⁄3H2O + 1⁄3H2O1 \n→ β-MoO3 + 1⁄3H2O1 → α-MoO3 → MoO3. (12)
$$

Vibrational spectra of MoO3∙H2O and dehydrated products obtained after heat treatment were studied by Raman spectroscopy, which appears to be a sensitive technique to distinguish unambiguously inter-lamellar  $MoO<sub>6</sub>$  octahedra from octahedra sharing two common edges. The Raman spectrum of the product heated above 470 °C exhibits superposition the spectral features of β-MoO<sub>3</sub> and α-MoO<sub>3</sub> crystal. Moreover, FTIR spectra of dehydrated MoO<sub>3</sub>⋅*n*H<sub>2</sub>O (*n*<l) shown in Fig. 4c are also more complex than that of crystalline MoO<sub>3</sub>, but spectral features can be discussed in terms of internal and external modes as usual for oxide lattices. The highest frequency bands are due to the symmetric stretching mode of the shortest Mo=O bond (the socalled molybdyl-mode), whereas the next highest bands are assigned to the stretch of intermediate bridging Mo-O bonds (longer Mo-O distances). The bending modes due to O-Mo-O bonds are located in the medium frequency range [77], while the external modes appear below 300 cm-1 . FTIR results (**Fig. 4c**) clearly show the change in the local structure of the MOs for annealing treatment *T*>250 °C of starting MoO<sub>3</sub>⋅H<sub>2</sub>O in good agreement with XRD patterns (**Fig. 4b**). Here, the modifications of the crystallinity can be deduced using the shape and frequency of the four groups of peaks. At intermediate heat treatment we observe the bands attributed to the mixed phase during the transformation from  $\beta$ -MoO<sub>3</sub> to  $\alpha$ -MoO<sub>3</sub> [309].

Camacho-López et al. [372] reported that the MoO<sub>3</sub>·1.4H<sub>2</sub>O glassy phase transforms to the amorphous MoO<sub>3</sub> $\cdot$ 0.7H<sub>2</sub>O phase prior to its crystallization, while the sample heated at 500  $\degree$ C crystallizes into the orthorhombic  $\alpha$ -MoO<sub>3</sub> with micro-crystallites having an average size of 6.8 µm. The local structure of products obtained by dehydration was investigated by Raman spectroscopy as shown in **Fig. 23** with the correspond SEM images of the as-prepared samples. In the Mo-O stretching mode region  $(100-600 \text{ cm}^{-1})$  the broad bands are due to the highly disordered  $MoO<sub>5</sub>(H<sub>2</sub>O)$  octahedral units in the amorphous hydrated sample (a-Mo $O<sub>3</sub>·1.4H<sub>2</sub>O$ ). These bands are centered on those of the Raman spectrum of the polycrystalline  $MoO<sub>3</sub>·1H<sub>2</sub>O$ phase, so the patterns of  $MoO_3 \cdot 1.4H_2O$  are the features envelop of  $MoO_3 \cdot 1H_2O$ . The sample a-MoO<sub>3</sub>·1.4H<sub>2</sub>O heated at 120 °C for 1 h transforms to MoO<sub>3</sub>·0.7H<sub>2</sub>O. The overall decrease of the Raman signal is due to the enhancement of the electrical conductivity (decrease of the penetration depth of the light) of the partially dehydrated sample [319]. The Raman spectrum exhibits the spectral features at 705, 850 and 948 cm<sup>-1</sup> of the MoO<sub>3</sub>·1.4H<sub>2</sub>O with additional vibrations at 665 and 983 cm<sup>-1</sup> assigned to the Mo-O3 stretch ( $B_{2g}$ ,  $B_{3g}$  mode) and M=O stretch  $(A<sub>g</sub>$  mode) of the MoO<sub>3</sub>·0.7H<sub>2</sub>O phase, respectively. These structural changes are related to water removal, which induces shorter Mo=O bonds and the less octahedral distortion. For wellcrystallized  $\alpha$ -MoO<sub>3</sub>, the peak at 983 cm<sup>-1</sup> shows a red shift to 995 cm<sup>-1</sup> and becomes a narrow line due to its non-polar character. It is worth noting that the line shape of the stretching modes is a function of the particle size growing with the annealing temperature of the glassy phase. Thus, the phonon confinement is observed in the case of  $MoO<sub>3</sub>$  crystallization, i.e., the Raman peaks broadened and little asymmetry is observed at 819 cm<sup>-1</sup>. When  $MoO<sub>3</sub>$  nano-particles formed were large in diameter (*L*≥40 nm) no significant downshift can be observed. The frequency of the stretch of terminal Mo=O shows a step at ca.  $T_a=120$  °C.



**Fig. 23**. Raman spectra for four different molybdenum trioxide phases: the  $MoO<sub>3</sub>·1H<sub>2</sub>O$ polycrystalline powders, the MoO<sub>3</sub> $\cdot$ 1.4H<sub>2</sub>O hydrated phase, the partially MoO<sub>3</sub> $\cdot$ 0.7H<sub>2</sub>O

dehydrated phase, and the orthorhombic  $\alpha$ -MoO<sub>3</sub> phase with their corresponding SEM images.

Haro-Poniatowski et al. [102] have reported the preparation of hydrated  $MoO<sub>3</sub>$  by acidification of sodium molybdate (Na2MoO4) solution and have also revealed the dependence of the morphology and color of the resulting product on the drying procedure. Using Raman scattering techniques, they demonstrated the evolution of the sample from a low temperature glass phase to a high temperature glass phase then crystallization under laser radiation. The thermally induced structural transformations in hydrated MoO<sub>3</sub> samples prepared by acidification of Na2MoO<sup>4</sup> solution have been investigated dynamically by combined analytic techniques such as thermo-Raman spectroscopy, thermogravimetry, differential thermal calorimetry and X-ray diffraction [103]. Results reveal that the structural evolution proceeds through an amorphous phase after dehydration processes, formation of sub-microcrystals, the possible MoO<sub>3</sub>-II phase and the  $\alpha$ -MoO<sub>3</sub> phase. The structural evolution under a flow of nitrogen gas indicated that the sample formed after dehydration could be oxygen deficient MoO<sub>3- $\delta$ </sub>. Du et al. [373] prepared the MoO<sub>3</sub>⋅0.55H<sub>2</sub>O phase by sonication of ammonium molybdate solution added to a mixture of 3-mercaptopropyltrimethoxysilane (MPTS) with anhydrous toluene. The light-blue sample obtained after vacuum drying at 70 °C for 6 h has a hexagonal structure with lattice parameters  $\alpha = 10.581$  Å,  $c = 3.720$  Å (JCPDS card No. 76-1003). MoO<sub>3</sub>⋅0.55H<sub>2</sub>O was converted into α-MoO<sub>3</sub> till the temperature reached 400 °C.

#### *3.2. Electrical conductivity*

**Figure 24** shows the temperature dependence of the electrical conductivity,  $\sigma_{dc}$ , of MOHs heat-treated at various temperatures in the range 140-458 °C. It is well known that transitionmetal oxides can be non-stoichiometric. MoO<sub>3</sub> is such a material. Mo<sup>5+</sup> ions can be produced as a consequence of the formation of oxygen vacancies. The conduction process has been described as a carrier hopping mechanism between localized states [18]. Small polarons are formed around these ions, due to the highly polar structure of this oxide. A thermally activated electronic hopping between lower and higher valence states is responsible for the electric transport. Also, it has been pointed out that the number of vacancies in  $MoO<sub>3</sub>$  films is indicated by the colour of the film [177]. The electrical conductivity curves shown in **Fig. 24** are typical of a hopping mechanism. The activation energies of the high-temperature regime are in the range 0.5-0.7 eV. The increase in  $\sigma_{dc}$  suggests a decrease of the O/Mo ratio due to the oxygen loss in annealed materials. The highly treated samples  $(T_a=458 \degree C)$  have the highest electrical conductivity values.



**Fig. 24.** Arrhenius plots of the electrical conductivity of molybdenum monohydrate MoO<sub>3</sub>⋅1H<sub>2</sub>O and products heat-treated at various temperatures in the range 140-458 °C.

## *3.3. Intercalation in MoO3∙nH2O*

Molybdate-based layered hybrids derived by intercalating organic guest species into the interlayer spaces of  $MoO<sub>6</sub>$  layers have also been a hot research topic, and the as-obtained organic–inorganic hybrids can be used as the precursors to prepare  $MoO<sub>3</sub>$  nanostructures [167,374-376]. Afsharpour et al. [374] synthesized molybdenum oxide nanohybrids using yellow molybdic acid and amino-carboxylates as the starting materials in an ethanol-water (1:3) mixed solution, and  $\alpha$ -MoO<sub>3</sub> nanostructures with high surface areas were obtained by calcining the as-obtained nanohybrids at 600 °C. Niederberger et al. [167] prepared molybdenum oxideamine hybrid composites by hydrothermally treating the ethanol-water mixed suspension of yellow MoO3∙2H2O and amines at 120 °C, and the molybdenum oxide-amine hybrid composites were used as the precursor to synthesize a-MoO3∙H2O nanofibers. Shukoor et al. [375] systematically investigated the intercalation behaviors of amines with various n-alkyl lengths into yellow molybdic acid (MoO3∙2H2O) in ethanol/water (1/3, v/v) solutions, and the as-obtained hybrids were transformed to  $MoO<sub>3</sub>$  rods, scrolls and disks by 33% HNO<sub>3</sub> treatment. In these intercalation reactions, elevated temperatures are usually necessary. Chen et al. [374] investigated molybdate-based inorganic–organic hybrids, which are formed from the reaction of white molybdic acid (MoO3∙H2O) powders and n-octylamine at room temperature, for sensing applications.

The suitability of MOHs as 3-volt electrode materials for non-aqueous lithium batteries has been assessed by several workers [121,319,366,377-379]. In early studies of electrochemical lithium insertion in MOHs Kumagai et al. [201,380,381]. Discharge profiles and kinetics are dependent on the amount of "structural water" into the host lattice. Compositional range was reported to be more than 2.5 Li/Mo reversibly inserted into crystalline MoO3⋅2H<sub>2</sub>O, which is higher than the 1.5 Li/Mo obtained for anhydrous  $\alpha$ -MoO3. Numerous hydrated forms of MoO<sub>3</sub>⋅y(H<sub>2</sub>O) (y = 0.3∼0.6) were prepared using the Freedman's method by hydrothermal acidification of Na<sub>2</sub>MoO<sub>4</sub> solution with 6 mol L<sup>-1</sup> HCl. The orthorhombic structure was obtained at HCl/Na<sub>2</sub>MoO<sub>4</sub> ratio of  $2.0 \sim 1.5$  and  $150 \sim 180$  °C, while a hexagonal network was formed at lower ratio. The as-prepared MoO<sub>3</sub>⋅0.58(H<sub>2</sub>O) has crystallographic parameters  $a = 3.955$  Å,  $b = 13.805$  Å,  $c = 3.684$  Å and  $V = 201.1$  Å<sup>3</sup>. As shown in the XRD patterns (**Fig. 25a**), the (021) reflection intensity is sensitive with the annealing temperature. The initial discharge-charge curves in 1 mol  $L^{-1}$  LiClO<sub>4</sub> in propylene carbonate as electrolyte at 0.2 mA cm-2 current density and cycling performance are shown in **Figs 25b-c**. The maximum first discharge capacity of 335 mAh  $g^{-1}$  is delivered by the MoO<sub>3</sub>⋅0.58(H<sub>2</sub>O) electrode heat-treated at 200 °C. However, a rapid decrease in specific capacity is observed after few cycles (**Fig. 25c**) [382]. The same group of workers reported the suitability of MoO<sub>3</sub>·H<sub>2</sub>O to deliver a discharge capacity of about 400 mAh  $g^{-1}$  with a voltage plateau around 2.5 V vs. Li<sup>+</sup>/Li. The MoO<sub>3</sub>·H<sub>2</sub>O crystal was found to be changed from a monoclinic to orthorhombic structure with lattice parameters of  $a = 5.285$  Å,  $b = 10.824$  Å,  $c = 5.237$  Å on discharge to 0.5 e<sup>−</sup> /Mo [201]. Similar synthesis process was used to prepared yellow MoO<sub>3</sub>·2H<sub>2</sub>O powders in 1 mol L<sup>-1</sup> Na<sub>2</sub>MoO<sub>4</sub> aqueous solution with 3 mol L<sup>-1</sup> HCl at 100 °C. Starting from an initial value of about 340 mAh  $g^{-1}$  obtained at a discharge exchange rate of C/10, the specific capacity continuously decreases with cycling. However, after the tenth cycle in the potential limits 3.8-2.0 V about 300 mAh  $g^{-1}$  are still available against only 240 mAh  $g^{-1}$ in the case of conventional  $MoO<sub>3</sub>$  [383]. Komaba et al. [336] showed that the orthorhombic *x*(Li<sub>2</sub>O)⋅MoO<sub>3</sub>⋅y(H<sub>2</sub>O) electrode which was hydrothermally formed from Li<sub>2</sub>MoO<sub>4</sub> system underwent electrochemical lithium intercalation up to Li/Mo  $\approx 1.6$  (>300 mAh ( $g_{\text{oxide}}$ )<sup>-1</sup>) on electroreduction until 1.3 V vs. Li<sup>+</sup>/Li.



**Fig. 25**. (a) XRD patterns of orthorhombic MoO3∙0.58(H2O) electrode prepared at 150 °C with H<sup>+</sup>/Na<sup>+</sup> ratio of 2.0 and its heat-treated products (o: orthorhombic phase). (b) Initial dischargecharge curves in 1 mol  $L^{-1}$  LiClO<sub>4</sub> in propylene carbonate as electrolyte at 0.2 mA cm<sup>-2</sup> current density. (c) cycling performance. On line at https://books.google.fr/books? [hl=fr&lr=&](https://books.google.fr/books?%20hl=fr&lr=)id=Ju5viChENAwC&oi=fnd&pg=PR6&dq=intercalation+compounds+for+batter y+materials.

It was made clear as a general trend that the discharge capacity of MOHs increases with decreasing water content, but cycle life increases with increasing water content in the compositional range 0.33<*n*<1.0. Julien et al. [121,366,377] reported the discharge curves performed on MOHs from open circuit potential down to 2.5 V vs.  $Li<sup>+</sup>/Li$  (-0.77 V vs. Ag/AgCl). This general trend has been also observed for  $MoO<sub>3</sub>$  thin films prepared by electrodeposition from peroxo-polymolybdate solutions [318]. Amorphous oxide-hydrate of molybdenum MoO<sub>3</sub>·2H<sub>2</sub>O was obtained during the synthesis of  $\beta$ -MoO<sub>3</sub> by cation exchange of an aqueous solution of Na2MoO4·2H2O passed through a cation exchange resin (Dowex 50WX8-200). As a product of this process, a solution with a pale green color and a pH of 2 was obtained. This solution was placed in a 50 ml vessel that was connected to a vacuum pump for 36 h. In this time, all water was removed slowly and a green powder was obtained as a residue [378]. Electrochemical lithium insertion in amorphous  $MoO<sub>3</sub>·2H<sub>2</sub>O$  led to a specific capacity of the cell of 490 mAh  $g^{-1}$ . This value exceeds notably the capacity developed by anhydrous β-MoO<sub>3</sub> (370 mAh g<sup>-1</sup>). Kumagai et al. [380] investigated *x*((NH<sub>4</sub>)<sub>2</sub>O)·MoO<sub>3</sub>·*y*(H<sub>2</sub>O) (with *x* = 0.075-0.042,  $v = 0.40$ -0.043) as cathode materials in Li cells. At current densities of 0.5 mA  $\text{cm}^{-2}$  and an average discharge voltage of 2 V, theoretical energy densities of ca. 600 Wh kg<sup>-1</sup>

have been predicted, which are considerably higher than in the case of completely dehydrated molybdenum oxides. However, the performance of *x*((NH<sub>4</sub>)<sub>2</sub>O)⋅MoO<sub>3</sub>⋅*y*(H<sub>2</sub>O) decreases drastically with subsequent cycling. The differing discharge mechanisms for the latter compounds (heterogeneous reaction) and MoO<sub>3</sub> (homogeneous reaction) has been discussed on the grounds of discharge characteristics, XRD data, and structural models. The maximum amount of lithium (~3.3 Li/Mo) incorporated in amorphous MoO<sub>3</sub>⋅2H<sub>2</sub>O leads to a specific capacity of 490 mAh  $g^{-1}$ . The charge–discharge curve showed a good reversibility in the potential range from 3.2 to 1.1 V *vs*. Li<sup>+</sup>/Li, where the cell voltage decreased monotonously as a function of the degree of lithium inserted [378].

Monoclinic monohydrate, MoO<sub>3</sub>·1H<sub>2</sub>O, having only one coordinated water molecule, shows a discharge capacity of about 200 mAh  $g^{-1}$  of acid weight equivalent and a discharge potential around 2.5 V vs.  $Li^{+}/Li$ . This electrode material displays a good charge-discharge cyclic behaviour at a capacity below 1 Li/Mo, while keeping the original layered lattice on cycling. Guzman et al. [366] studied a crystalline MoO<sub>3</sub>⋅1H<sub>2</sub>O electrode prepared by sol-gel method, while Yebka et al. used commercial molybdebum-trioxide hydrate powders [319]. **Figure 27** presents the electrochemical behavior of the crystalline MoO3∙1H2O investigated at the current rate of C/10 (cathode loading of 2.3 mg  $cm^{-2}$ ). In the potential voltage 1.5-3.0 V vs. Li<sup>+</sup>/Li, the first discharge occurs a Li uptake identical to that of anhydrous  $MoO<sub>3</sub>$  (1.5Li/Mo). The electroinsertion of Li ions occurs mainly in two steps in the potential range between 3.0 and 1.5 V (compositional range  $0.0 \le x \le 1.5$ ): a first plateau at ca. 2.85 V in the range  $0 \le x \le 1.5$ 0.25 and at wide plateau at ca. 2.35 V with a sloppy region in between, which corresponds to a single-phase insertion mechanism (**Fig. 26a**). The MoO<sub>3</sub>·1H<sub>2</sub>O electrode delivers a first discharge capacity of 344 mAh  $g^{-1}$ , which maintains at  $\sim$ 200 mAh  $g^{-1}$  after 100 cycles (**Fig. 26b**). The discharge/charge curves have been modelled using the Armand's model with two adjacent domains including an ion-ion interaction term for each voltage region [3]. The electrochemical behaviour of anhydrous sample shows higher discharge capacity than that of hydrated MoO<sub>3</sub> (262 mAh g<sup>-1</sup>). MoO<sub>3</sub>⋅1H<sub>2</sub>O has only a coordinated water molecule in the structure. It is obvious that this open structure is favourable to the lithium insertion. The temperature dependence of the discharge-charge shows that Li//MoO<sub>3</sub>⋅1H<sub>2</sub>O cells can operate efficiently at different temperatures in the range of 25-50 °C. However, in the case of Li//MoO3∙1H2O operating at 50 °C, the cell voltage range is about 5% less than for the material operating at 25 °C. Amorphous MoO3∙1H2O electrode exhibits stepwise discharge behaviour, including two plateaus: the first step up to 0.3e<sup>-</sup>/Mo and the second one up to 1.0e<sup>-</sup>/Mo. A

discharge capacity of 260 mAh  $g^{-1}$  is delivered in the potential range corresponding to 1.5 e<sup>-</sup>/Mo [366].



**Fig. 26**. (a) First galvanostatic discharge-charge curve of the MoO3•1H2O positive electrode recorded at  $C/10$  rate in the potential range 2.0-3.5 V vs.  $Li<sup>+</sup>/Li$ . (b) Electrochemical patterns of  $MoO<sub>3</sub>•1H<sub>2</sub>O$  at the 2<sup>nd</sup> and 100<sup>th</sup> cycle.

The electrochemical features of the crystalline  $MoO<sub>3</sub>$ <sup>1</sup>/<sub>2</sub>H<sub>2</sub>O electrode are rather similar with the appearance of three plateaus at. ca. 2.5, 1.75 and 1.5 V (**Fig. 27a**). This electrochemical behaviour is attributed to the large cavities available for Li ions, which prevent repulsive forces between inserted ions. The structure of MoO3∙½H2O may be described as an alteration of linear double rows of edge-sharing distorted  $MoO<sub>6</sub>$  and  $MoO<sub>5</sub>(H<sub>2</sub>O)$  octahedral [309]. In the potential range 3.0-1.0 V, the MoO<sub>3</sub> hemihydrate electrode delivers a first discharge capacity of 353 mAh g<sup>-1</sup> for a charge transfer of ~1.5e<sup>-</sup>/Mo. The cyclability of MoO<sub>3</sub>⋅<sup>1</sup>/<sub>2</sub>H<sub>2</sub>O (**Fig. 27b**) shows the electrode cycled 100 times at C/10 rate retains a capacity of 201 mAh g<sup>-1</sup>. Figure 27c presents the XRD patterns of  $Li<sub>x</sub>MoO<sub>3</sub>·<sup>1</sup>/<sub>2</sub>H<sub>2</sub>O$  as a function of the amount of Li electrochemically inserted. The monoclinic structure remains largely unchanged upon lithiation with the strongest (001) reflection line shifted toward lower angles (inset **Fig. 27c**). This result suggests that, on discharging, Li<sup>+</sup> ions are inserted between the layers leading to a small increase of the interlayer spacing, which can be explained by a model of rigid hydrated-MoO<sub>3</sub> layers in their original state separated by intercalated Li cations. **Figure 27d** displays the FTIR absorption spectra of pristine and Li intercalated products. The FTIR spectra of Li0.15MoO3∙½H2O and Li<sub>0.15</sub>MoO<sub>3</sub>⋅<sup>1</sup>/<sub>2</sub>H<sub>2</sub>O samples show new bands at 1780, 1200, 1148, 1117 and 1087 cm<sup>-1</sup>. The band at 1780 cm<sup>-1</sup> is assigned to the stretching vibration  $v_{C=0}$  of propylene carbonate molecules. Its position is shifted towards lower energies compared with pure propylene carbonate suggesting that solvent molecules are bonded at the surface of the oxide network. The bands situated in the region 1200-1080 cm<sup>-1</sup> are attributed to the stretching mode of Li atoms inserted between layers vibrating against the nearest neighboring oxygen atoms [366].



**Fig. 27**. (a) Typical discharge curves of amorphous MoO3∙½H2O powders cycled at C/10 rate in the potential window3.0-1.5 V vs. Li<sup>+</sup>/Li. (b) Electrochemical patterns of MoO<sub>3</sub>⋅1/2H<sub>2</sub>O at the 5<sup>th</sup> and 100<sup>th</sup> cycle. (c) XRD patterns of lithiated MoO<sub>3</sub>⋅½H<sub>2</sub>O powders using electrochemical titration. The insert shows the variation of the d-spacing vs.  $x(Li)$  (reflection at  $2\theta = 12.8^{\circ}$ ). (d) FTIR absorption spectra of pristine MoO3∙½H2O and electrochemically Li inserted Li<sub>x</sub>MoO<sub>3</sub>⋅<sup>1</sup>/<sub>2</sub>H<sub>2</sub>O ( $x = 0.15$  and 0.3).

MoO3∙0.66H2O powders were prepared by heating molybdic acid (MoO3∙H2O) at *T*a=216 °C [121,317,319]. **Figures 28a-b** show the typical discharge-charge curves of Li<sub>x</sub>MoO<sub>3</sub>⋅<sup>2</sup>⁄<sub>3</sub>H<sub>2</sub>O//Li cells cycled at C/10 rate in the potential range 3.5-1.5 V vs. Li<sup>+</sup>/Li, which exhibit electrochemical patterns as follows: (i) an initial OCV of ~3.5 V, (i) a Li uptake 1.5 Li per Mo atom providing a discharge capacity of 260 mAh  $g^{-1}$ , (iii) a large voltage plateau at ca. 2.2 V, and (iv) the discharge capacity remains at  $162 \text{ mA}$ h g<sup>-1</sup> after 100 cycles. Note the absence of the first discharge voltage plateau in the 0.0 < *x* < 0.25 observed in discharge profiles of other MOHs and of anhydrous MoO3, which show the strong dependence of the water content on the Li insertion process in MOH frameworks. The S-shaped behaviour of the discharge-charge curves of the MoO3∙⅔H2O electrode suggests single-phase reaction LixMoO3∙⅔H2O in the range  $0.0 \le x \le 1.5$ . Structural changes in Li-intercalated were analysed using in situ XRD in MoO3∙⅔H2O/LiClO4-PC electrolyte/Li cell (**Fig. 28c**) and Raman spectroscopy (**Fig. 28d**). The Raman spectrum of MoO<sub>3</sub>⋅<sup>2</sup>/<sub>3</sub>H<sub>2</sub>O is in fair agreement with the spectrum of MoO<sub>3</sub> [77]. Bands located at 120 and 150 cm<sup>-1</sup> are assigned to vibrations of the  $(O_2Mo_2)_n$  chains while bands

located at 335, 440 and  $657 \text{ cm}^{-1}$  are attributed to the vibration of the O- Mo<sub>3</sub>. After lithium insertion, we observed features as follows. (i) The intensities of original Raman-active vibrations decrease drastically and the background increases. (ii) The background also increases as the frequency shift increases, probably owing to the modification of the electronic band structure upon intercalation. (iii) The low-frequency part of the spectrum remains almost unchanged, indicating that the layered structure of  $MoO<sub>3</sub>·<sup>2</sup>/<sub>3</sub>H<sub>2</sub>O$  is maintained. (iv) New peaks appear at 562, 546, and  $278 \text{ cm}^{-1}$ , which are assigned to a longitudinal and transverse intercalation mode [384], in which intercalant species vibrate strongly against the host lattice. Moreover, the reversible insertion behaviour has been also observed by Raman measurements. The original Raman spectrum has been recovered on discharge-charge cycling. Note that the change in colour with increasing Li content seems to start from the physical edges of the sample proceeding along the crystal imperfections (grain boundaries, micro cracks, planer defects, etc.) to a uniform coloration of the crystal [385]. **Figure 28e** shows the interplanar distance, *d*int, deduced from the position of the (200) reflection peak as a function of the Li concentration within the MoO<sub>3</sub>⋅<sup>2</sup><sub>3</sub>H<sub>2</sub>O framework. *In situ* XRD patterns were recorded during charge and discharge in the potential range 3.4-1.2 V. The shift toward lower  $2\theta$  position of the (200) line reveals that Li ions are inserted between layers upon discharge the cell, resulting in a significant increase in the interlayer spacing of about 6%. Layer separation in spite of the reduced dimensions of Li ions (0.69 Å, *CN*=6) is constantly reported for layered compounds. Besenhard and Schollhorn demonstrated an interlayer separation of 1  $\AA$  for  $x = 1$  for the parent oxide Li<sub>x</sub>MoO<sub>3</sub> [61]. An expansion of 12% for  $d_{\text{int}}$  in the basic structure of Mo<sub>18</sub>O<sub>52</sub> has been observed by Pasquali et al [386]. The lattice expansion in  $Li<sub>x</sub>MoO<sub>3</sub>$  does not increase continuously with the degree of intercalation, *x*, but shows a maximum in the vicinity of  $x = 0.1$ . Similar situation has been observed for molybdenum bronzes  $H_xMOO_3$  [387].





**Fig. 28**. (a) Galvanostatic discharge-charge curve of the MoO3∙⅔H2O positive electrode. Electrochemical measurements were performed at a rate of C/5. (b) Electrochemical patterns of MoO<sub>3</sub>⋅<sup>2</sup><sub>3</sub>H<sub>2</sub>O at the 2<sup>nd</sup> and 100<sup>th</sup> cycle. (c) In situ XRD patterns of Li-intercalated MoO<sub>3</sub>⋅<sup>2</sup><sub>3</sub>H<sub>2</sub>O in LiClO<sub>4</sub>-PC electrolyte. (d) Raman spectra of molybdenum hydrate Li<sub>x</sub>MoO<sub>3</sub>⋅<sup>2</sup><sub>3</sub>H<sub>2</sub>O and its lithium intercalated product. New peaks marked by stars are the fingerprints of Li ions located in the interlayer space. (e) Variation of the interplanar distance deduced from the (200) Bragg peak with the Li concentration within the MoO3∙⅔H2O framework. In-situ XRD patterns were recorded during charge (■) and discharge (●) in the potential range 3.4-1.2 V.

Using a vacuum drying technique Ramirez et al. have prepared an amorphous hydrate molybdenum oxide a-MoO3∙2H2O [310]. Electrochemical lithium insertion in a-MoO3∙2H2O shows a maximum amount of Li incorporated of ∼3.3 Li/Mo, which leads to a specific capacity of 490 mAh  $g^{-1}$  [378]. This value exceeds by more than 20% the specific capacity developed by its homolog crystalline form (400 mAh  $g^{-1}$ ). The charge–discharge curves show a good reversibility in the potential range from 3.2 to 1.1V vs. Li<sup>+</sup>/Li, where the cell voltage decreased monotonously as a function of the degree of lithium inserted. Yuan and Si prepared a triclinic

 $\alpha$ -MoO<sub>3</sub>·H<sub>2</sub>O sample by direct chemical reaction of Mo powders and H<sub>2</sub>O<sub>2</sub> with the formation of precursor MoO<sub>2</sub>(OH)(OOH). Partly dehydrated  $\alpha$ -MoO<sub>3</sub>·*x*H<sub>2</sub>O with *x*=0.936, 0874 and 0.752 were prepared by vacuum dehydration at 110, 150 and 165 °C, respectively (**Fig. 29**) [379]. **Figure 29c** presents the first two galvanostatic discharge–charge curves for the triclinic  $\alpha$ -MoO<sub>3</sub>·H<sub>2</sub>O and its dehydrated products (loading  $\sim$ 2 mg) tested in 2016-type coin cell with Li metal anode cycled at 0.1 A  $g^{-1}$  current density in the potential range 3.0-0.001 V vs Li<sup>+</sup>/Li. These profiles show a broad plateau at 0.2 V ascribed to the conversion reaction involving the transformation of  $Li_xMoO_3·H_2O$  to Mo and  $Li_2O$  matrix.

Recently, Yu et al. [388] demonstrated a water-incorporation strategy to expand the interlayer gap of  $\alpha$ -MoO<sub>3</sub>, in which water molecules take the place of the non-bridging oxygen of layered lattice. The incorporated H2O molecules expand the interlayer channel dimension, i.e., increase of the b-lattice parameter to 15.02 Å compared to  $\alpha$ -MoO<sub>3</sub> (13.85 Å). The composition MoO<sub>2.92</sub>⋅0.07H<sub>2</sub>O was determined by TGA. The modified  $α$ -MoO<sub>3</sub> electrode exhibits specific capacity of 963 C  $g^{-1}$  at 0.1 mV s<sup>-1</sup>, and boosted cycling stability, i.e., a capacity of 455 C  $g^{-1}$  is maintained after 600 cycles at 100 mA  $g^{-1}$ . A fast-kinetics dual-ionintercalation energy storage device was assembled by combining the  $Li<sup>+</sup>$  intercalation α-MoO<sub>3</sub>⋅nH<sub>2</sub>O anode with an anion-intercalation graphite cathode in 2 mol L<sup>-1</sup> LiPF<sub>6</sub> electrolyte. Operating within a 1.0–3.5 V voltage window an energy density of 44 Wh L−1 was delivered (based on the whole device). The significant role of the incorporated  $H_2O$  molecules in accelerating the diffusion of Li<sup>+</sup> within the MoO<sub>2.92</sub>⋅0.07H<sub>2</sub>O lattice was studied by galvanostatic titration. The  $D_{Li}^*$  values  $2.4 \times 10^{-12} - 3.5 \times 10^{-10}$  cm<sup>2</sup> s<sup>-1</sup>) are substantially higher than those in  $\alpha$ -MoO<sub>3</sub> (3.9 × 10<sup>-13</sup>–6.0 × 10<sup>-11</sup> cm<sup>2</sup> s<sup>-1</sup>).




**Fig. 29**. (a) The crystal structure of the triclinic monohydrated molybdenum oxide  $\alpha$ -MoO<sub>3</sub>·1H<sub>2</sub>O. (b) XRD patterns of pristine  $\alpha$ -MoO<sub>3</sub>·1H<sub>2</sub>O and its partially dehydrates  $\alpha$ -MoO<sub>3</sub>·0.936H<sub>2</sub>O,  $\alpha$ -MoO<sub>3</sub>·0.874H<sub>2</sub>O, and  $\alpha$ -MoO<sub>3</sub>·0.752H<sub>2</sub>O. (c) The first two dischargecharge profiles of  $\alpha$ -MoO<sub>3</sub>· $xH_2O$  cycled at 0.1 A  $g^{-1}$  current density between 3.0 and 0.001 V vs. Li<sup>+</sup>/Li. Reproduced with permission from [379]. Copyright 2013 The Royal Society of Chemistry.

## **4. Oxygen deficient MoO3-**

## 4.1. MoO<sub>3- $\delta$ </sub> suboxides

MoO<sub>3- $\delta$ </sub> suboxides with stoichiometric deviation  $\delta$  < 1 belong to the class of materials that offer a tunability of their intrinsic electronic properties from wide bandgap semiconductor MoO<sub>3</sub> to semi-metallic MoO<sub>2</sub> [8]. Upon inducing oxygen defects, Mo<sup>6+</sup> ions can be reduced to  $Mo^{5+}$  and finally  $Mo^{4+}$ , the oxygen vacancies not only increase the interlayer spacing and the electrical conductivity of  $MoO<sub>3</sub>$ , but also enhance charge storage kinetics [389]. The varieties of oxidation states in oxygen-deficient  $MoO<sub>3-δ</sub>$  compounds make them highly attractive in the field of energy storage and conversion and they have found applications in LIBs [236,390], solid state thin-film microbatteries [72,173,], electrochemical supercapacitors [194,391], gas sensors [392], electrochromic devices [231], solar cells [393,394], anode interlayers for photovoltaic devices [395], smart windows [396], light-emitting diodes [397], etc. These devices take advantage of the changes in the degree of crystallinity, cationic environment, stoichiometry deviation, band gap energy and electronic conductivity, which can be controlled by the growth conditions.

## 4.1.1. Structure

Regarding the electronic structure, the defective oxygen generates gap states, which change the optical appearance, i.e., the oxides exhibit color transitions from transparent to yellow and eventually grayish/blue. Thus, the molybdenum suboxides phases can be classified in three groups: (i) the nearly-stoichiometric MoO<sub>3-δ</sub> (with  $\delta$  < 0.03), which retains the orthorhombic symmetry, is unintentionally an *n*-type semiconductor at ambient conditions, partly due to intrinsic point defects related to oxygen vacancies  $(V<sub>O</sub>)$  and molybdenum interstitials  $(M<sub>Oi</sub>)$ [34], (ii) the defective MoO<sub>3- $\delta$ </sub> (with 0.03 <  $\delta$  < 0.11) and (iii) Mo<sub>n</sub>O<sub>3n</sub>-1 suboxides (Magnéli phases,  $n = 4-9$ ), which collapse into  $\text{ReO}_3$ -type structures with crystallographic shear planes due to the destabilization of the  $MoO<sub>3</sub>$  structure by the loss of oxygen [35-40]. The schematic representation of the crystal structure of the layered  $α$ -MoO<sub>3</sub> phase and the vacancies of the three inequivalent oxygen positions O1 (apical), O2 (corner-sharing) and O3 (edge-sharing) are shown in Fig. 30 [398]. An O1 vacancy  $(V_{O(1)})$  leads to two localized electrons on the neighboring Mo that are thus in the  $Mo^{4+}$  (4d<sup>2</sup>5s<sup>0</sup>) configuration; O2 and O3 vacancies create a bipolaron with electrons localized on two separate  $Mo^{5+}$  sites in the first and second coordination spheres for  $V_{O(2)}$  and on the opposite side of the vacancy for  $V_{O(3)}$  [399]. At higher vacancy concentration, the shear structure is energetically more favorable than the point defects [400]. Inzami et al. accurately explored the effect of oxygen nonstoichiometry on the electronic structure and properties of the oxygen-deficient  $MoO<sub>3-δ</sub>$  material using a van der Waals density functional calculation [398]. It is shown that: (i) the electronic structure of layered  $MoO<sub>3-d</sub>$  is highly sensitive to changes in oxygen stoichiometry as  $Mo<sup>6+</sup>$  has an empty 4d shell; (ii) the layered structure efficiently screens the structural perturbations caused by oxygen vacancies, and (iii) the oxygen deficiency gives rise to Mo 4d gap states with energy levels dependent on the type of oxygen vacancy.



**Fig.** 30. Crystal structure of the layered α-MoO<sub>3</sub> phase (a) and the vacancies of the three inequivalent oxygen positions O1 (apical) (b), O2 (corner-sharing) (c) and O3 (edge-sharing) (d). Adapted from [398]. Copyright 2016 American Chemical Society.

**Figure 31** shows a schematic diagram of the electronic states of molybdenum oxides as a function of the oxygen vacancies. The electronic character varies from insulator  $(M_0O_3)$  to semiconductor ( $MoO<sub>3-δ</sub>$ ) and finally to metal-like ( $Mo<sub>9</sub>O<sub>26</sub>$ ), in agreement with conductivity measurements [399]. It is well established that increasing  $\delta$  in MoO<sub>3- $\delta$ </sub> can lower the bandgap [401]. The electron distribution in pure MoO<sub>3</sub> follows the ionic model (Mo<sup>6+</sup> and O<sup>2−</sup>) corresponding to the Mo  $4d^0$  configuration. For non-stoichiometric MoO<sub>3</sub> oxides, extended Mo 4*d* states, which then lie in the bandgap as gap states due to the occurrence of  $Mo^{5+}$  and  $Mo^{4+}$ ions, are filled by electrons donated from oxygen vacancies. The formation of oxygen deficiency (termed also as sub-stoichiometric) not only produces an increase of the electrical conductivity  $\sigma_e$  [402,403] owing to the additional gap states, but also increases the surface energy of the particles and promotes electrochemical reactions [397,399]. Magnetic susceptibility measurements showed that  $MoO<sub>3</sub>$ ,  $MoO<sub>2</sub>$  and  $Mo<sub>n</sub>O<sub>3n-1</sub>$  shear suboxides in between are all feebly paramagnetic.



**Fig. 31.** Schematic diagram of the electronic states of molybdenum oxides as a function of the oxygen vacancies. The electronic character varies from insulator (MoO<sub>3</sub>,  $\sigma_e > 10^{-9}$  S cm<sup>-1</sup>) to semiconductor (defective Mo<sub>9</sub>O<sub>26</sub>, σ<sub>e</sub> ≈ 10<sup>-1</sup> S cm<sup>-1</sup>) to metal-like (γ-Mo<sub>4</sub>O<sub>11</sub>, σ<sub>e</sub> ≈ 10<sup>2</sup> S cm<sup>-1</sup>). Copyright Lakshmi-Narayana et al. [72]. Open access article under the CC BY-NC-ND license.

#### 4.1.2. Determination of  $\delta$

X-ray photoelectron spectroscopy (XPS) is an appropriate tool to evaluate the stoichiometry of oxygen deficient  $MoO<sub>3-δ</sub>$  compounds. From the Mo 3d core level spectra, the presence of Mo in low valence states, i.e.,  $Mo^{5+}$  and  $Mo^{4+}$ , can be evidenced by fitting the XPS 3d5/2 and 3d3/2 peaks at binding energies around 235 and 232 eV, respectively (**Table 10**) [85,404].

**Table 10**. Standard binding energies of Mo 3d levels for molybdenum at different oxidation states [405].

Peak assignment	Binding energy (eV)		
	$3d_{5/2}$	$3d_{3/2}$	
$Mo^{6+}$	232.5	235.7	
$Mo^{5+}$	231.5	234.7	
$Mo^{4+}$	230.1	233.3	
$Mo^{3+}$	229.3	232.5	
$Mo^{2+}$	228.4	232.6	
$Mo^{0}$	227.7	230.9	

Based on the deconvolution of the Mo 3d core-level peaks by using equal-width Gaussian peaks  $3d_{5/2}$  and  $3d_{3/2}$  spin-orbit components, the ratio of  $Mo^{6+}/Mo^{5+}$  in a sub-stoichiometric  $MoO<sub>3-δ</sub>$  oxide is calculated by the following equation [406]:

$$
k = \frac{Mo^{6+}}{Mo^{5+}} = \frac{R(Mo^{6+}3d_{5/2}) + R(Mo^{6+}3d_{3/2})}{R(Mo^{5+}3d_{5/2}) + R(Mo^{5+}3d_{3/2})},
$$
\n(13)

where k is the ratio of the quantity of  $Mo^{6+}/Mo^{5+}$  calculated from the ratio of the integrated peak area  $(R)$ , the valence state for molybdenum can be calculated by the relation  $[407]$ :

$$
\phi = \frac{6}{1+k^{-1}} + \frac{5}{1+k},\tag{14}
$$

and the stoichiometric deviation is given by:

$$
\delta = 3 - \frac{\phi}{2} \,. \tag{15}
$$

As an example, **Figure 32a** shows the XPS spectra of oxygen vacancies-rich  $\alpha$ -MoO<sub>3-δ</sub> nanobelts synthesized by hydrothermal process. Blue  $MoO<sub>3-δ</sub>$  nanobelts were obtained by calcining the MoO<sub>3</sub> powder NaH<sub>2</sub>PO<sub>2</sub>·H<sub>2</sub>O at 250 °C for 1 h in N<sub>2</sub> atmosphere. A graphene nanomesh carbon nanotube/MoO<sub>3- $\delta$ </sub> (GC/MoO<sub>3- $\delta$ </sub>) nanocomposite with more O-vacancy defects was also characterized [408]. For the pristine MoO<sub>3</sub> nanobelts, the atomic ratio of Mo<sup>5+</sup>/Mo<sup>6+</sup> is calculated to be 0.132 giving  $\delta = 0.02$ , whereas is greatly increased to 0.308 for the GC/MoO<sub>3</sub> $δ$  nanocomposite, indicating the partial reduction of MoO<sub>3</sub>,  $δ = 0.12$ . Consequently, the electrons occupied in 4d band increase with  $\delta$  improving the electrical conductivity to  $0.83 \times 10^{-5}$  S cm<sup>-1</sup>

(two orders of magnitude higher than that of pristine pure  $MoO<sub>3</sub>$ ). Diffuse reflectance spectra show that the bandgap is reduced largely from 3.25 to 2.95 eV with the increase of O-vacancy concentration.



**Fig. 32**. (a) XPS core-level spectra of Mo 3d for the oxygen vacancies-rich *α*-MoO3-*<sup>δ</sup>* nanobelts. Reproduced with permission from [408]. Copyright 2019 Elsevier. (b) UV-Vis absorption spectra of flash-deposited  $\alpha$ -MoO<sub>3-δ</sub> thin films at different substrate temperatures: (1) 30 °C, (2) 120 °C, and (3) 250 °C. (c) Plot of  $[(α-α<sub>0</sub>)\hbarω]<sup>2</sup>$  vs. photon energy for thin films and MoO<sub>3</sub> crystal. The intercepts at  $\alpha = 0$  (full lines) give the bandgap energy. (d) variation of the bandgap with the stoichiometric deviation. Reproduced with permission from [153]. Copyright 1995 Elsevier.

Ultraviolet-visible spectroscopy can be also used to determine the stoichiometric deviation in MoO3-δ (**Figs. 32b-c**) [153]. The increase of O-vacancy concentration results in the enhanced absorbance from 300 to 1000 nm. The fundamental absorption edge occurs at about 4 eV with a high value of the absorption coefficient  $3 \times 10^4$  cm<sup>-1</sup>. A broad absorption band in the red region, i.e., 1.5 eV, corresponds to the excitation of trapped electrons into the conduction band. As a result, various degrees of coloration occurred. It is worth noting that the coloration of the films is associated with the absorption edge shift. For example, films grown at room temperature are white and transparent, whereas films deposited at higher substrate temperatures

 $(T<sub>s</sub> > 120$ °C) are deeply colored. The intrinsic absorption edge of the films was evaluated in terms of the direct transition. The optical absorption coefficient  $\alpha$  is calculated by the expression:

$$
\alpha = \frac{1}{d} \ln \left\{ \frac{(1 - R)^2}{2T} + \sqrt{\left[ \frac{(1 - R^2)}{2T} \right]^2 + R^2} \right\},\tag{16}
$$

where *d* is the sample thickness, *T* the transmittivity and *R* the reflectivity determined from the optical transmission and reflection spectra, respectively (**Fig. 32b**). The theory of interband absorption near the threshold shows that  $\alpha$  vary according the equation:

$$
\alpha h v = B(hv - E_g)^n,\tag{17}
$$

where *h* is the Planck constant, v the frequency, *B* the probability parameter,  $E<sub>g</sub>$  is the optical bandgap and *n* an exponent, which characterizes the transition process ( $n = \frac{1}{2}$  for direct allowed transitions). Thus, the Tauc plot  $[(\alpha-\alpha_0)\hbar\omega]^2$  vs. photon energy yields a linear behavior in the region of strong absorption near the absorption edge, as shown in **Fig. 32c**. Extrapolating the linear portion of this straight portion to  $\alpha = 0$  gives the optical bandgap. The optical bandgap of crystalline MoO<sub>3</sub> is  $E_g = 3.05$  eV. As shown in Fig. 32d, the bandgap of MoO<sub>3-δ</sub> suboxides decreases linearly with the increase of the stoichiometric deviation in the range  $0 \le \delta \le 0.15$ .

### 4.1.3. Li insertion in  $MoO<sub>3-δ</sub>$

Generally, the oxygen deficient  $MoO<sub>3-δ</sub>$  oxides, as cathode or anode materials, exhibits several appealing characteristics for lithium-ion storage, including high specific capacity, good stability against cycling and fast charge transport kinetics [132]. The bulk structure of  $MoO<sub>3-δ</sub>$ under reductive and oxidative reaction conditions was investigated in situ with time-resolved X-ray absorption spectroscopy (XAS) [409]. The MoO<sub>3-δ</sub> suboxide phases (with  $0 < \delta < 0.25$ ) exhibit Li-insertion capacity much higher than that of stoichiometric  $MoO<sub>3</sub>$  phase, which justifies the efforts to fabricate them under the form of thin films. For the development of these technologies, oxygen vacancies are easily generated in the  $MoO<sub>3-δ</sub>$  materials prepared in a thin film architecture by optimizing the deposition conditions  $[410]$ . Jung et al.  $[236]$  investigated the electrochemical reactivity of ball-milled MoO3−*<sup>δ</sup>* as anode materials for lithium-ion batteries. High-energy ball-milling converts highly-crystalline  $MoO<sub>3</sub>$  bulk powders into partially reduced low crystalline MoO3−*<sup>δ</sup>* materials with a reduced particle size. Oxygen deficiency gradually increases as the ball milling time increases. Bulk  $MoO<sub>3</sub>$  powders were partially reduced to give nanostructured  $MoO<sub>2.929</sub>$ ,  $MoO<sub>2.903</sub>$ , and  $MoO<sub>2.895</sub>$  by ball-milling for 4, 6, and 8 h, respectively. When tested in the  $0.01-3$  V *vs.* Li<sup>+</sup>/Li range the first discharge

capacity is beyond 1100 mAh  $g^{-1}$ , which is indicative of a complete conversion reaction. The lithiation in MoO<sub>3−δ</sub> suboxides occurs with two steps: intercalation (>1.5 V) and conversion  $(<0.5 V$ ), although their voltage profiles become smother than that of MoO<sub>3</sub> bulk. Additionally, incomplete re-oxidation during subsequent charge results in the formation of  $MoO<sub>2</sub>$  instead of  $MoO<sub>3</sub>$ , which in turn affects the reactivity in subsequent cycles. As compared to bulk  $MoO<sub>3</sub>$ , ball-milled MoO<sub>3−δ</sub> showed significantly enhanced cycle performance. The anode material ballmilled for 8 h has 64.4% discharge efficiency at the  $35<sup>th</sup>$  cycle against 27.6% for bulk at the  $10<sup>th</sup>$ cycle, which has been attributed to the nano-texture wherein nanometer-sized particles aggregate to form secondary ones. Jegal et al.  $[390]$  prepared a mixed-valence MoO<sub>3- $\delta$ </sub>/CNT nanocomposite  $(x<0.1)$  using a one-pot microwave-assisted hydrothermal reaction in which 5 nm MoO<sub>3-δ</sub> nanoparticles are coated on the surfaces of CNT. The nanocomposite had a reversible discharge capacity of could deliver discharge capacities of 770, 700, and 570 mAh  $g^{-1}$  at current densities of 450, 900, and 1800 mA  $g^{-1}$ , respectively.

Highly dispersed ultra-small  $MoO<sub>y</sub>$  nanoparticles (1.5-3.5 nm) anchored on N-doped 3D hierarchically porous carbon (3D-MoO<sub>y</sub>@CN) were prepared using an efficient in-situ chelating and hard-templating strategy. An optimized 3D-MoO<sub>y</sub>@CN heat-treated at 700  $^{\circ}$ C delivers specific capacities of 742 and 431 mAh  $g^{-1}$  at current density of 100 and 1000 mA  $g^{-1}$ after 1000 cycles, respectively. This excellent performance is attributed to the unique hierarchical porous morphology (specific BET surface area of 109  $m^2$  g<sup>-1</sup> and pore volume of 0.30 cm<sup>3</sup>  $g^{-1}$ ) structure with strong binding of the ultra-small MoO<sub>y</sub> nanoparticles onto N-doped carbon surface, which can avoid the agglomeration and alleviate the volume expansion of  $MoO<sub>v</sub>$ nanoparticles in the charge-discharge process  $[406]$ . MoO<sub>3– $\delta$ </sub> nanowires were synthesized in a hot-filament chemical vapor deposition reactor in which molybdenum filaments were resistively heated to 775 °C in 10 sccm of oxygen at a pressure of  $\sim$ 150 Pa [134]. Structural characterization revealed the Mo<sub>17</sub>O<sub>47</sub> phase. It has been shown that Mo<sub>17</sub>O<sub>47</sub> nanowire arrays can retain a capacity of 630 mAh g<sup>-1</sup> for up to 20 cycles at a current density of 50 mA g<sup>-1</sup>. Also, it has been reported that a 10-nm silicon coated  $MoO<sub>3-δ</sub>$  hybrid architectures, deposited by microwave plasma CVD have shown a capacity retention of 780 mAh  $g^{-1}$ . Sun et al. [411] designed vertically aligned oxygen-deficient α-MoO3-*<sup>δ</sup>* nanoflake arrays as 3D cathode for advanced all-solid-state thin film lithium batteries (TFLIBs). The α-MoO<sub>3-δ</sub> electrode were fabricated by magnetron sputtering of metal Mo target. The Li/LIPON/MoO<sub>3- $\delta$ </sub> 3D-TFLIB exhibits high specific capacity of 266 mAh  $g^{-1}$  at 50 mA  $g^{-1}$  current density, good rate performance (110 mAh  $g^{-1}$  at 1000 mA  $g^{-1}$ ), and excellent cycle performance (92.7% capacity retention after 1000 cycles).

Sun et al. prepared  $MoO<sub>y</sub>$  nanobelts with different quantities of oxygen vacancies through a "sauna reaction" and tested their performance in LIBs [412]. The number of oxygen vacancies depends on the duration for the production of  $H_2$  by the reaction of the carbon cloth and water vapor to reduce  $MoO<sub>3</sub>$ . The as-obtained  $MoO<sub>v</sub>$  nanobelts deliver a reversible capacity of 400 mAh·g<sup>-1</sup> at 1 A·g<sup>-1</sup> after 200 cycles without obvious capacity loss. Furthermore, MoO<sub>y</sub> exhibits an excellent rate capability (400 and 267 mAh·g<sup>-1</sup> at 100 and 200 mA·g<sup>-1</sup> current rate, respectively. Wang et al.  $[225]$  investigated the MoO<sub>3</sub>-MnO<sub>2</sub> intergrown nanoparticles (6-8 nm in size) composite prepared by one-step hydrothermal synthesis as anode for LIBs. The composite attains an initial specific capacity of 2034 mAh  $g^{-1}$  and stays 1446 mAh  $g^{-1}$  after 50 cycles at a rate of 0.5C in the voltage range of 0.01-2.0 V. The synergistic effect of  $MoO<sub>3</sub>$  and  $MnO<sub>2</sub>$  during intergrowth is attributed to the incorporation of  $Mn<sup>4+</sup>$  is incorporated into the MoO<sup>3</sup> lattice to modify its crystal structure, i.e., creation of oxygen deficiency. Carbon nanotube-wired and oxygen-deficient MoO<sub>3-δ</sub> nanobelts (MoO<sub>3-δ</sub>/CNTs) were constructed via hydrothermal synthesis with subsequent controlled reduction in  $H_2/Ar$  gas mixture [268]. The MoO3/CNTs composite, evaluated as an anode in the 0.05–3.0 V potential window, is able to retain a specific capacity of 421 mAh  $g^{-1}$  after 100 cycles at 200 mA  $g^{-1}$ , and delivers 293 and 202 mAh  $g^{-1}$  at current densities of 2 and 4 A  $g^{-1}$ , respectively. Zhang et al. [413] prepared several oxygen deficient  $\alpha$ -MoO<sub>3-δ</sub> cathode materials using H<sub>2</sub> plasma etching process. From XPS measurements, the composition is calculated to be  $MoO<sub>2.912</sub>$  and  $MoO<sub>2.870</sub>$  for  $MoO<sub>3</sub>$  etched for 10 and 20 min, respectively, which is confirmed by the appearance of the Raman peak at  $1008$  cm<sup>-1</sup> as a result of oxygen vacancy. The MoO<sub>2.912</sub> electrode shows the best performance delivering a first specific capacity of 224 mAh  $g^{-1}$  at 1 A  $g^{-1}$  rate, which declines slowly to 67 mAh  $g^{-1}$  over 1000 cycles.

In a SIBs, the oxygen vacancies of  $MoO<sub>3-δ</sub>$  play also an important role. They can enhance the electric conductivity and Na-ion diffusion coefficient [291]. Li and coworkers revealed how oxygen vacancies of  $MoO<sub>y</sub>$  influenced the performance of the SIBs [290]. Mo $O<sub>2.97</sub>$ , which possesses only a few oxygen vacancies exhibits the best specific capacity (176.6 mAh·g<sup>-1</sup> at 50 mA·g<sup>-1</sup>) and a ~8% capacity loss over 2000 cycles compared with the highest capacity at the  $500<sup>th</sup>$  cycle. The voltage profile of MoO<sub>2.5</sub> displays a two-plateau behavior suggesting the stabilized function of oxygen vacancies during the charge/discharge cycling. Yi et al. reported that oxygen-deficient MoO<sup>3</sup> demonstrates excellent performances in Li-S batteries. The cells with MoO<sub>3-δ</sub> cathode deliver a capacity of 690 mAh g<sup>-1</sup> after 200 cycles at 0.2C, demonstrating a significant improvement of the capacity retention. Furthermore, at the current rate of 1C,  $MoO<sub>3-δ</sub>$  cell displays a high initial capacity of 775 mAh  $g<sup>-1</sup>$  and an enhanced surplus capacity

of 480 mAh  $g^{-1}$  after 600 cycles [414]. Wu et al. [226] adopted a carbon-free strategy for an advanced LIB anode material, which consists of a mixture of  $MoO<sub>3</sub>$  and  $MoO<sub>2</sub>$ . Mo $O<sub>2</sub>$ nanoparticles treated with a smaller fraction of  $O_2$  (0.2%  $O_2$  in N<sub>2</sub>) show mixed phases which can be attributed to  $MoO<sub>2</sub>$ ,  $MoO<sub>3</sub>$ , as well as a small fraction of  $Mo<sub>4</sub>O<sub>11</sub>$ . A high specific capacity up to 930.6 mAh·g<sup>-1</sup>, long cycle-life (>200 cycles) and high rate capability. MoO<sub>3-δ</sub> nanobelts with different concentrations of oxygen vacancies were synthesized by a one-step hydrothermal process using different concentrations of ethanol as the reductant [415]. With the increasing amount of ethyl alcohol in the hydrothermal reaction, the sample color gradually changes from white (pure  $\alpha$ -MoO<sub>3</sub>) to dark (MoO<sub>2</sub>+Mo<sub>4</sub>O<sub>11</sub>). Mo<sup>5+</sup> in MoO<sub>3- $\delta$ </sub> appears blue and MoO<sub>2</sub> looks black. MoO<sub>3-δ</sub> nanobelts, tested as anode for LIBs in a voltage window of 0.01–3.0 V vs. Li<sup>+</sup>/Li, can maintain a high specific capacity of ~500 mAh  $g^{-1}$  at a high current density of 1 A  $g^{-1}$  and can retain a high specific capacity of 641 mAh  $g^{-1}$  after 50 cycles at 100 mA  $g^{-1}$  and 420 mAh  $g^{-1}$  after 100 cycles at 500 mA  $g^{-1}$ .

## *4.2. MoO3-δ thin films*

MoO3-δ thin films have been easily prepared using a broad variety of physical vapor deposition (PVD) methods, including thermal evaporation [194,416,417], magnetron sputtering [418,419], dip-coating [394], pulsed layer deposition (PLD) [115,173,420,421]. However, few works report the spin-coating deposition of  $MoO<sub>3-δ</sub>$  via a low-temperature solution process  $[422]$ . α-MoO<sub>3-δ</sub> thin films with low resistivity were fabricated on glass substrate via conducting a two-step annealing treatment on the thin film deposited by thermal evaporation method. A heat-treatment in air enables the formation of the  $\alpha$ -phase layered structure (resistivity of  $10^6$ )  $\Omega$ ⋅cm), whereas an annealing in N<sub>2</sub> gives rise to the lowest resistivity of 5.4  $\Omega$ ⋅cm without deteriorating crystal structure [417]. The increase in carrier concentration was assigned to the presence of ionized oxygen vacancy during the  $N_2$  annealing as illustrated by the following equation:

$$
O_0^x + 2Mo^{6+} \rightleftharpoons V_0 + 2Mo^{5+} + \frac{1}{2}O_2(g).
$$
 (18)

Like stoichiometric MoO<sub>3</sub>, sub-stoichiometric MoO<sub>3- $\delta$ </sub> thin films can be obtained by evaporation in a vacuum at relatively low temperature (*T* < 500 °C) or in reducing atmosphere. However, few works explore the effect of oxygen deficiency and electrical conduction enhancement on energy storage properties [423]. Parashar demonstrated that the deposition temperature plays a crucial role on the composition of  $MoO<sub>3-δ</sub>$  thin film deposited by magnetron sputtering in argon plasma. At working pressure of 0.5 Pa and RF power of 75 W, the film

deposited at  $T_s = 250$  °C is a mixture of  $\alpha$ -MoO<sub>3</sub> and Mo<sub>4</sub>O<sub>11</sub> phases, while a pure  $\alpha$ -MoO<sub>3</sub> phase is grown at  $T_s = 350$  °C [418]. Sub-stoichiometric MoO<sub>3- $\delta$ </sub> thin films were deposited on  $SiO<sub>2</sub>$  glass and stainless-steel substrate using conventional RF sputtering technique [144]. The films fabricated in an Ar gas atmosphere at a sputtering gas pressure of 4 Pa, a radio-frequency power of 200 W, a substrate temperature of 100  $^{\circ}$ C and a deposition rate of 1.16  $\mu$ m h<sup>-1</sup> exhibit the  $Mo<sub>9</sub>O<sub>26</sub>$  structure (dark-blue color) with the crystallographic (010) plane oriented parallel to the substrate surface. The OCV of the  $Mo<sub>9</sub>O<sub>26</sub>$  thin-film electrode (4.6 µm thick) was 2.7 V. The discharge capacities of the 1<sup>st</sup>, 10<sup>th</sup> and 21<sup>st</sup> cycles were 398, 318 and 289  $\mu$ Ah cm<sup>-2</sup>, respectively, at a current density of 10 mA cm<sup>-2</sup>. Recently, He and coworkers [419] prepared sputtered  $MoO<sub>y</sub>$  thin films with different oxygen contents (fixed Ar flow rate of 10 sccm and variable  $O_2$  flow rate from 0.4 to 1.5 sccm corresponding to 3.8%-13%). Mo $O_{2.95}$ , Mo $O_{2.43}$ ,  $MoO<sub>2.7</sub>$ , and  $MoO<sub>1.45</sub>$  thin films were obtained with  $O<sub>2</sub>$  concentration of 13.0%, 8.6%, 6.5% and 3.8%, respectively, showing that the decrease of the oxygen content pushes the Fermi level very close to the conduction band. Puppala et al. investigated the microstructure and morphology of PLD MoO3-*<sup>δ</sup>* thin films growth for catalytic applications using a femtosecond laser (f-PLD) and a nanosecond excimer-laser (n-PLD). Substantially textured films with a partially crystalline phase prior to annealing were obtained by the f-PDL laser, while the n-PLD-grown  $MoO<sub>3-δ</sub>$ films were predominantly amorphous with a smooth surface [186]. Sunu et al. claimed that asdeposited PLD films ( $T_s = 400 \degree C$ ,  $\Phi = 4-5$  J cm<sup>-2</sup>, repetition rate of 15 to 20 Hz, and  $p(O_2) =$ 500 Pa) are suboxide-like, i.e., mixture of  $n-Mo<sub>4</sub>O<sub>11</sub>$  and  $\gamma-Mo<sub>4</sub>O<sub>11</sub>$ , which transformed to  $MoO<sub>3</sub>$ after annealing at 500 °C in air for 5 h [392].

Recently, the effects of operational conditions on structural, electronic and electrochemical properties on molybdenum suboxides  $(M_0O_{3-\delta})$  thin films were investigated [72]. Three classes of samples are obtained with different degrees of stoichiometric deviation without posttreatment: (i) amorphous  $Mo_{3-\delta}$  ( $\delta$  < 0.05) (ii) nearly-stoichiometric samples ( $\delta \approx 0$ ) and (iii) suboxides MoO<sub>3-δ</sub> ( $\delta > 0.05$ ). The suboxide films  $0.05 \leq \delta \leq 0.25$  deposited on Au/Ti/SiO2/flexible-Si substrates with appropriate processing conditions show high electrochemical performance as an anode layer for lithium planar microbatteries. In the realm of simple synthesis, the MoO<sub>3-δ</sub> film deposited at 450 °C under oxygen pressure of 13 Pa is a mixture of  $\alpha$ -MoO<sub>3</sub> and Mo<sub>8</sub>O<sub>23</sub> phases (15:85). The electrochemical test of the 0.15MoO<sub>3</sub>-0.85Mo<sub>8</sub>O<sub>23</sub> (MoO<sub>2.894</sub>) film shows a specific capacity of 484  $\mu$ Ah cm<sup>-2</sup>  $\mu$ m<sup>-1</sup> after 100 cycles of charge-discharge at a constant current of 0.5 A  $cm^{-2}$  in the potential range 3.0-0.05 V. This capacity value corresponds to the insertion of  $\sim 5.78 \mathrm{Li}^{\dagger}/\mathrm{Mo}$ , is close to the theoretical maximum (5.79 Li<sup>+</sup> /Mo). **Figure 33** presents the electrochemical performance of the MoO3-δ deposited at

450 °C under oxygen pressure of 13 Pa [72]. Amorphous  $MoO<sub>v</sub>$  thin films prepared by reactive magnetron sputtering deposition (total Ar+ $O_2$  gas flow kept at 40 sccm at 0.8 Pa using 20%  $O_2$ ) consists of a mixture of amorphous  $MoO<sub>3</sub>$  and  $MoO<sub>2</sub>$  phases [424]. The a-Mo $O<sub>y</sub>$  thin films (1.1)  $\mu$ m thick) exhibit a high capacity of 338  $\mu$ Ah·cm<sup>-2</sup> (about 845 mAh·g<sup>-1</sup>) at a current density of 90 µA cm<sup>-2</sup> even after 100 discharge/charge cycles, and have a rate capacity of 158 µAh cm<sup>-2</sup> at a high current density of 2000  $\mu$ A cm<sup>-2</sup>.



**Fig.** 33. Electrochemical tests of the MoO<sub>3</sub>-Mo<sub>8</sub>O<sub>23</sub> (15:85) thin film in Li cells with 1 mol L<sup>-1</sup> LiPF<sup>6</sup> dissolved in ethylene carbonate and dimethyl carbonate (EC:DMC; 1:1 *w*/*w*) as aprotic electrolyte. (a) Galvanostatic charge discharge profiles of a cell cycled at 1 A  $g^{-1}$  current density in the voltage range 3.0–0.05 V. (c) Differential capacity (-d*Q*/d*V* vs. *V*) of the first lithiationdelithiation cycle. (c) Rate capability. (d) Cyclability over 100 cycles at 1 A  $g^{-1}$  compared with amorphous MoO<sub>3-δ</sub> and α-MoO<sub>3-δ</sub> film deposited at 300 °C. Copyright Lakshmi-Narayana et al. [72]. Open access article under the CC BY-NC-ND license.

# **5. Magnéli-type Mo oxides**

In the Mo-O binary system, there is the series  $Mo<sub>n</sub>O<sub>3n-1</sub>$ , which crystallizes in the so-called Magnéli phases described in details by Kihlborg [37] (see crystallographic parameters in **Table 11**). They are oxygen deficient relative to MoO3. The removal of oxygen and accompanied partial reduction of Mo(VI) cations introduce crystallographic shear maintaining the layered framework. These oxides have been identified as derived from  $MoO<sub>3</sub>$  by shear mechanism with networks of  $MoO<sub>6</sub> octahedra$ . These units are connected in three dimensions with empty parallel tunnels involving the presence of potential sites for hosting Li<sup>+</sup> ions. The crystal structures of

the MonO3n-1 molybdenum oxides, with their respective single unit cells are shown in **Fig. 34**. Since 1980s, quasi-2D layered  $Mo<sub>n</sub>O<sub>3n-1</sub>$  compounds with commensurate/incommensurate periodic lattice distortion (PLD) have attracted great attention due to their fascinating property of charge-density-wave (CDW) [425,426]. These intermediate molybdenum oxides are believed to be also catalytically active [427]. Furthermore, they exhibit a rich phase polymorphism and a variety of morphologies.

Compound	Space group	Cell parameters (A)			Ref.
		$a/\alpha$	b/B	$c/\gamma$	
Mo <sub>17</sub> O <sub>47</sub>	Pha <sub>2</sub>	$21.61/90^{\circ}$	$19.63/90^{\circ}$	$3.951/90^{\circ}$	$\lceil 38 \rceil$
$Mo_9O_{26}$	P2/c	14.45	$4.00/96.0^{\circ}$	16.750	$\left[36\right]$
$Mo_8O_{23}$	P2/a	16.88	$4.052/106.19^{\circ}$	13.39	[428]
$\eta$ -Mo <sub>4</sub> O <sub>11</sub>	$P2\sqrt{a}$	$6.695/90^{\circ}$	5.440/94.31°	324.573/90°	[429]
$\gamma$ -Mo <sub>4</sub> O <sub>11</sub>	Pnam	$24.49/90^{\circ}$	$6.752/90^{\circ}$	5.457/90°	[430]
Mo <sub>5</sub> O <sub>14</sub>	P4/nmm	23.00		3.937	[431]
Mo <sub>18</sub> O <sub>52</sub>	P1	8.145/102.67	11.89/67.82°	21.23/109.97°	$[35]$

**Table 11.** Structural parameters of  $Mo<sub>n</sub>O<sub>3n-1</sub>$  molybdenum oxides.



**Fig.** 34. Crystal structures of Magnéli Mo<sub>n</sub>O<sub>3n-1</sub> phases. Reproduced with permission from [400]. Copyright 2017 Royal Society of Chemistry.

In 1951, Glemser and Lutz [54] investigated the structure of several molybdenum blue oxides, the crystallized samples  $Mo_8O_{23}$  and  $Mo_9O_{26}$ , amorphous blue oxide hydrates MoO2.75∙*n*H2O and MoO2.80∙*n*H2O, and crystallized blue hydroxides Mo4O10(OH)<sup>2</sup> and  $M_0<sub>2</sub>O<sub>4</sub>(OH)<sub>2</sub>$ , which are closely related to the starting material  $Mo<sub>3</sub>$ . The  $Mo<sub>n</sub>O<sub>3n-1</sub>$  phases include Mo<sub>4</sub>O<sub>11</sub> [432,433], Mo<sub>5</sub>O<sub>14</sub> [434], Mo<sub>6</sub>O<sub>17</sub>, Mo<sub>8</sub>O<sub>23</sub> [36], Mo<sub>9</sub>O<sub>26</sub> [40], Mo<sub>18</sub>O<sub>52</sub> [35,435] and Mo<sub>17</sub>O<sub>47</sub>, which are formed by a mixture of Mo oxidation states between +6 and +4. As experimental results, the topotactic reaction of lithium into the MoO<sup>y</sup> framework and reversibility have been demonstrated [66,67], and extent of lithiation depend on the stoichiometry of the  $Mo<sub>n</sub>O<sub>3n-1</sub>$  phases. Due to their tunneling structure with empty cavities, the MonO3n-1 phases are performant cathode materials in non-aqueous lithium cells [61,436-439]. Christian and co-workers [440] shown that of the oxides  $Mo_4O_{11}$  ( $MoO_{2.75}$ ),  $Mo_{17}O_{47}$  $(MoO<sub>2.765</sub>)$ ,  $Mo<sub>8</sub>O<sub>23</sub>$  (Mo $O<sub>2.889</sub>$ ), and  $Mo<sub>9</sub>O<sub>26</sub>$  (Mo $O<sub>2.875</sub>$ ) the greatest reversible capacity (1.5 Li/Mo) after several deep discharge-charge cycles was exhibited by  $Mo<sub>17</sub>O<sub>47</sub>$  in cells with LiClO4/propylene carbonate electrolyte in the potential range 2.9-1.4 V (**Table 12**).

Compound	$\rho$ ( $\Omega$ cm)	Li uptake (Li/Mo)		
		1st cycle	20th cycle	
$Mo_{18}O_{52}$	10	0.7	0.3	
$Mo_8O_{23}$		1.0	0.4	
$Mo_4O_{11}$	0.01	1.5	0.75	
$Mo_{17}O_{47}$	0.05	1.5	0.8	
Mo <sub>5</sub> O <sub>14</sub>	0.02	1.45	-	

Table 12. Electrical and electrochemical characteristics of  $Mo<sub>n</sub>O<sub>3n-1</sub>$  oxides.

## *5.1. Mo4O<sup>11</sup> (MoO2.75)*

The Mo<sub>4</sub>O<sub>11</sub> phase has two 2D modifications (metal-like), orthorhombic γ-Mo<sub>4</sub>O<sub>11</sub> [441] and monoclinic  $\eta$ -Mo<sub>4</sub>O<sub>11</sub> [442]; in the metallic  $\eta$ -Mo<sub>4</sub>O<sub>11</sub> phase the layers with the ReO<sub>3</sub>-type are ordered parallel to each other, whereas in  $\gamma$ -Mo<sub>4</sub>O<sub>11</sub> they are arranged like the mirror image of the former ones [443]. Cignini et al. [436] investigated the electrochemical behavior of the orthorhombic  $Mo<sub>4</sub>O<sub>11</sub>$  phases in Li cells with different electrolytes. The higher specific capacity of 340 mAh  $g^{-1}$  was delivered in LiClO<sub>4</sub>/PC. A Li//Mo<sub>4</sub>O<sub>11</sub> cell, discharged at 0.5 mA cm<sup>-2</sup>, has given 610 Wh kg<sup>-1</sup>, based on the weight of Li and  $Mo_4O_{11}$ . When the discharge for  $Mo_4O_{11}$  is stopped above 2 V corresponding to  $x < 0.7$  e/Mo), the lattice of this oxide is essentially retained, while at 1.5 V, the  $Mo<sub>4</sub>O<sub>11</sub>$  structure is greatly altered with the appearance of new phases.

## *5.2. Mo5O<sup>14</sup> (MoO2.8)*

The crystal structure of  $Mo<sub>5</sub>O<sub>14</sub>$  (denoted as  $\theta$ -Mo oxide) is tetragonal (*P4/mbm* S.G.). The lattice consists of either distorted  $MoO<sub>6</sub>$  octahedra and  $MoO<sub>7</sub>$  pentagonal bi-pyramids which are coupled together by edge- and corner-sharing [431]. As shown in **Fig. 34e**, the pentagonal bipyramids are edge sharing with the neighbored octahedra perpendicular to the fivefold axis

and tunnels are parallel to the *c*-axis. Few works reporting the lithiation of  $Mo<sub>5</sub>O<sub>14</sub>$  are available in the literature  $[121,436,410,444]$ . Preliminary patterns of the fully lithiated Mo<sub>5</sub>O<sub>14</sub> crystal suggest that the unit cell expands in the *b*-direction and contracts in both the *a-* and *c*-directions upon insertion of lithium [440]. Among the  $Mo<sub>n</sub>O<sub>3n-1</sub>$  oxides,  $Mo<sub>4</sub>O<sub>11</sub>$ ,  $Mo<sub>17</sub>O<sub>47</sub>$ , and  $Mo<sub>8</sub>O<sub>23</sub>$ , the compound  $Mo_{17}O_{47}$  has the highest intercalation cyclability. Its average discharge voltage is 1.85 V, and its reversible discharge capacity is 1.5 Li/Mo. Its theoretical specific energy is 490 Wh kg<sup>-1</sup>. Cignini et al. reported a specific capacity of 310 mA  $g^{-1}$  for Mo<sub>5</sub>O<sub>14</sub> as the positive electrode of a primary Li cell discharged at  $0.5$  mA cm<sup>-2</sup> and  $1.0$  V cut-off. [436]. Nazri and Julien  $[121]$  showed that MoO<sub>2.8</sub>, an oxygen-deficient Mo<sub>5</sub>O<sub>14</sub> prepared by dehydration and annealing treatment of molybdic acid powder (MoO<sub>3</sub>·1H<sub>2</sub>O) at 750 °C, exhibits a high electrical conductivity of  $10^{-2}$  S cm<sup>-1</sup>. Such a material displays an excellent reversible capacity (1.45 e<sup>-</sup> /Mo) in cells with 1 mol  $L^{-1}$  LiClO<sub>4</sub> in propylene carbonate (PC) electrolyte in the potential range 3.3–1.3 V vs. Li<sup>+</sup>/Li. The open-circuit voltage is 3.1 V vs. Li/Li<sup>+</sup>. The first discharge displays a stepped behavior with a voltage plateau at ca. 2.2 V followed by a potential decline for  $x > 0.7$  e/Mo. This compound can be assigned to the  $Mo<sub>3</sub><sup>6+</sup>Mo<sub>2</sub><sup>5+</sup>$  valence distribution. If reversible lithium incorporation produced all  $Mo^{4+}$ , the expected lithium uptake would be 8/5(1.6) e<sup>-</sup>/Mo; a value close to that measured by electrochemical titration. The gravimetric capacity of 326 mAh  $g^{-1}$  was much higher than the 280 mAh  $g^{-1}$  of anhydrous MoO<sub>3</sub>. Shembel et al.  $[445]$  reported the electrochemical performance of e-Mo<sub>4</sub>O<sub>11</sub> synthesized from ammonium molybdate electrolyte in thin-layer binder-free electrodes. A first discharge capacity of 900 mAh  $g^{-1}$  is delivered at 30 mA  $g^{-1}$  current density. The reversible discharge capacity can exceed 225 mAh  $g^{-1}$  for more than 170 cycles.

Oxygen deficient  $MoO<sub>3-δ</sub>$  nanoparticles (200 nm thick) were prepared by a citrate sol-gel using ammonium heptamolybdate tetrahydrate as a source of Mo and heat treated with a small fraction of zirconia under reducing atmosphere [446]. The structural analyses reveal that the composite is a blend formed by layered α-MoO<sub>3</sub>, orthorhombic oxygen deficient phases MoO<sub>3</sub>-δ with  $\delta = 0.25$  (γ-Mo<sub>4</sub>O<sub>11</sub>) and α-ZrMo<sub>2</sub>O<sub>8</sub>. At C/10 rate, the composite delivered a stable reversible capacity of 135 mAh  $g^{-1}$  after the 50<sup>th</sup> cycle. At a rate of 2C the reversible capacity is maintained at 118 mAh  $g^{-1}$ . An amorphous Mo<sub>n</sub>O<sub>3n-1</sub>/carbon nanocomposite (*n*  $\approx$  5) was fabricated from a citrate–gel precursor heated at moderate temperature (500 °C) in inert (argon) atmosphere. The as-prepared  $Mo<sub>5</sub>O<sub>14</sub>$ -type/C material is compared to  $\alpha$ -Mo $O<sub>3</sub>$  synthesized from the same precursor in air. Thermal gravimetry and elemental analysis indicate the presence of  $25.8 \pm 0.2$ % of carbon in the composite. The RS spectrum of Mo<sub>5</sub>O<sub>14</sub>/C demonstrates an oxygen deficiency in the molybdenum oxide and the presence of a partially graphitized carbon.

Outstanding improvement in electrochemical performance is obtained for the  $M_{0.5}O_{14}$ encapsulated by carbon (325 mAh  $g^{-1}$  at the 50<sup>th</sup> cycle at 70 mA  $g^{-1}$ ) in comparison with the carbon-free MoO<sub>3</sub> (95 mAh  $g^{-1}$ ). The high performance of the Mo<sub>5</sub>O<sub>14</sub>/C composite is clearly revealed with a specific discharge capacity of 155 mAh g<sup>-1</sup> at 10C rate (7 A g<sup>-1</sup>) [109]. Mo<sub>4</sub>O<sub>11</sub> nanoparticles were decorated onto ultralight graphene sheets (HRG) self-assembled with sulfur homogeneously dispersed onto the surface were used as the cathode material for a lithium sulfur battery [447]. The Mo<sub>4</sub>O<sub>11</sub>-HRG/S cathode deliver a specific capacity of ~900 mAh g<sup>-1</sup> at 0.1C rate after 80 cycles at cut-off of 1.5 V. Tysyachny et al. [448] investigated the transport properties of the electrolytic oxide  $Mo_4O_{11}$  as cathode material of lithium battery. The average values of  $D_{Li}^*$  in Mo<sub>4</sub>O<sub>11</sub> determined by cyclic voltammetry are in the range from 3.0 to 6.4×10<sup>-12</sup>  $\text{cm}^2 \text{ s}^{-1}$ .

## *5.3. Mo8O<sup>23</sup> (MoO2.875)*

 $M_{08}O_{23}$  has a monoclinic crystal structure ( $P_{2}/a$  S.G.), which derived from that of ReO<sub>3</sub> by a shear mechanism, is formed by slabs,  $8 \text{ MoO}_6$  octahedra thick, mutually connected by edge-sharing between octahedra. It is a 3D framework characterized by intersecting channels and of cavities (**Fig. 34a**). Besenhard and Schollhorn [61] reported that cathodes containing  $M<sub>08</sub>O<sub>23</sub>$  can be galvanostatically reduced resulting in the first uptake of  $\sim$ 1 e<sup>-</sup> per Mo in 0.35 mol  $L^{-1}$  LiAlCl<sub>4</sub>/DME electrolyte, which reduced to 0.4 Li after 20 cycles. Pistoia and coworkers examined the electrochemical properties of  $Mo_8O_{23}$  as cathode in lithium cells. Fiordiponti et al. [439] reported that Li cells containing the monoclinic  $Mo<sub>8</sub>O<sub>23</sub>$  cathode show a stepwise discharge curve with particularly noticeable knees for 0.2<*x*<0.3, 0.6<*x*<0.7, and 0.8<*x*<0.9, which does not prevent a complete reversibility. For high values of Li content, more serious structure alterations start to take place to which the sharp fall of potential may be attributed.  $Mo_8O_{23}$  exhibits at first a shrinkage of the unit cell until 0.2Li/Mo, which is attributed to the large empty cavities with at least 12 coordinating sites, followed by a lattice re-expansion. At 1.0 mA cm<sup>-2</sup> and 2.0 V cut-off, 85 mAh  $g^{-1}$  (0.89 e<sup>-</sup>/Mo) and 200 Wh  $kg^{-1}$  were obtained. Mo<sub>8</sub>O<sub>23</sub> maintains good recharge efficiencies for over 15 cycles. The chemical diffusion coefficient Li ions in Li<sub>x</sub>Mo<sub>8</sub>O<sub>23</sub> was measured in the range 0.0< $x$ <0.8.  $D_{Li}^*$  reaches a maximum value of  $5 \times 10^{-9}$  cm<sup>2</sup> s<sup>-1</sup> at *x* near to 0.3.

#### *5.4. Mo18O<sup>52</sup> (MoO2.889)*

 $M_{018}O_{52}$  crystallized with the triclinic symmetry ( $P\overline{1}$  S.G.). Its network maintains the layered structure of MoO<sup>3</sup> (**Fig. 34f**). Ressler et al. [449] investigated the bulk structural modification of Mo-O oxides by the reduction of  $MoO<sub>3</sub>$  with propene and the oxidation of  $MoO<sub>2</sub>$  with oxygen. The formation of a  $Mo<sub>18</sub>O<sub>52</sub>$ -type shear structure consists of generation of oxygen vacancies at the (100) or (001) facets by reaction with propene and a vacancy diffusion in the MoO<sub>3</sub> bulk. Kaiser et al. [450] prepared  $Mo<sub>n</sub>O<sub>3n-1</sub>$  compounds by the solid-state reaction of powdered α-MoO<sup>3</sup> with Mo in the spark-plasma synthesis (SPS) process at temperatures up to 973 K:  $M_{018}O_{52}$  ( $x = 2.889$ ),  $M_{017}O_{47}$  ( $x = 2.760$ ), and  $\gamma$ - $M_{04}O_{11}$  ( $x = 2.750$ ), which show different thermoelectric properties.  $Mo_{18}O_{52}$ , which appears to be an intrinsic narrow-gap semiconductor (activation energy  $\approx 0.3$  eV), exceptionally low thermal conductivity over the full temperature range (0.5–0.9 W m<sup>-1</sup> K<sup>-1</sup>), while γ-Mo<sub>4</sub>O<sub>11</sub>, and Mo<sub>17</sub>O<sub>47</sub> show poor metallic conductivity. The highest power factor is achieved by  $Mo_{17}O_{47}$  (30 W m<sup>-1</sup> K<sup>-2</sup> for  $440 \le T \le 610$  K). Preliminary electrochemical tests of Mo<sub>18</sub>O<sub>52</sub> in 0.35 mol L<sup>-1</sup> LaAlCl<sub>4</sub>/DME were conducted at  $0.3 \text{ mA cm}^{-2}$  current density. The galvanostatic reduction in the potential window from -0.1 to -2.8 V vs. SCE yields 1.5 e<sup>-</sup> per Mo [61]. Investigations of Mo<sub>18</sub>O<sub>52</sub> by Pistoia et al. [438] show that this non-stoichiometric  $Mo_{18}O_{52}$  oxide displays a high-rate capability as cathode for secondary Li cells. In a cell with 1 mol  $L^{-1}$  LiClO<sub>4</sub>/PC electrolyte, it can intercalate reversibly up to 1.35 Li/Mo (250 mAh  $g^{-1}$ ) at current density in the range 1.0 mA cm<sup>-2</sup> and 1 V cut-off, forming ternary phases. In a cell with 2 mol  $L^{-1}$  LiAsF<sub>6</sub>/methyl formate electrolyte, it delivers 340 mAh  $g^{-1}$  (1.81 e<sup>-</sup>/Mo charge transfer). Li<sup>+</sup>-ion diffusion is relatively fast in the Mo<sub>18</sub>O<sub>52</sub> channels ( $D_{Li}^* = 10^{-9}$  cm<sup>2</sup> s<sup>-1</sup>) and gives rise to a structure, in which Li<sup>+</sup> is tetrahedrally coordinated by oxygen.

## *5.5. Mo9O<sup>26</sup> (MoO2.889)*

 $M_{09}O_{26}$  has the same composition than  $M_{018}O_{52}$ , but crystallizes with a monoclinic structure ( $P2/c$  S.G.), which derives from the metastable ReO<sub>3</sub>-type modification β-MoO<sub>3</sub>.

## *5.6. Mo17O<sup>47</sup> (MoO2.765)*

Christian et al. [440] obtained their best results with  $Mo_{17}O_{47}$ . Li/ $Mo_{17}O_{47}$  cells could be cycled over 35 times at current densities of  $0.5$  mA cm<sup>-2</sup> and average discharge voltages of 1.85 V. The  $M_{017}O_{47}$  framework accommodates 1.5 electrons per Mo during the first cycle, which reduces to 0.8e<sup>-</sup> at the 20<sup>th</sup> cycle. Among the  $Mo<sub>n</sub>O<sub>3n-1</sub>$  oxides,  $Mo<sub>17</sub>O<sub>47</sub>$  has the highest intercalation cyclability. From these figures, the theoretical specific energy is 400 Wh  $kg^{-1}$ . Meduri et al. [132] prepared MoO<sub>3−*x*</sub> nanowire arrays with diameters of ∼90 nm in a hotfilament chemical vapor deposition reactor at 775 °C in 10 sccm of oxygen at a pressure of 146 Pa. XRD and Raman characterization indicate an oxygen deficient  $Mo_{17}O_{47}$  orthorhombic

phase. The initial discharge capacity retention of  $\sim$ 770 mAhg<sup>-1</sup> at a current density of 25 mA g<sup>-1</sup> in the range 0.1-3.5 V corresponds to 4.1 Li<sup>+</sup> ions per MoO<sub>3−*x*</sub>. A capacity of ~630 mAh g<sup>-1</sup> is retained for up to 20 cycles at a current density of 50 mA  $g^{-1}$ . Hugues et al. [449] considered that, among Mo oxides with formulae between  $MoO<sub>2</sub>$  and  $MoO<sub>3</sub>$ ,  $Mo<sub>17</sub>O<sub>47</sub>$  is the best cathode materials, which can supply current densities of 15 mA cm<sup>-2</sup>.

## **6. Molybdenum Dioxide MoO<sup>2</sup>**

Molybdenum dioxide occurs as the mineral *tugarinovite* (molar mass  $127.94$  g mol<sup>-1</sup>, density 6.47 g cm<sup>-3</sup>, insoluble in water, melting point 1100 °C).

## 6.1. Structure

MoO<sub>2</sub> crystallizes with the monoclinic structure with space group  $P2_1/c$  ( $C_{2h}^5$ ), which can be viewed as a distorted rutile phase ( $\text{Fig. 35a}$ ). This structure is composed of MoO<sub>6</sub> octahedra joined by edge-sharing, which form a  $(1\times1)$ -tunnelling network (**Fig. 35b**). The lattice parameters of MoO<sub>2</sub> are  $a = 5.6109$  Å,  $b = 4.8562$  Å,  $c = 5.6285$  Å, and  $\beta = 120.95^{\circ}$  (JCPDS card No. 66-6787) [452,453]. The unit cell comprises four formula units. The metal atoms as well as the two inequivalent oxygen atoms occupy the general Wyckoff positions (4*e*). An interesting feature of MoO<sub>2</sub> is that its metallic ground state (electrical resistivity  $6.9 \times 10^{-4} \Omega$ cm) (**Fig. 35c**) [454]. The *d* orbital can be separated into two sub-bands, i.e., *t*2g and *e*<sup>g</sup> bands by the crystal field. The *t*2g band has a lower energy level and is partially filled by the two electrons, and the *e*<sup>g</sup> band is empty and has a higher energy level located above the Fermi energy (Fig.  $35d$ ). This metallicity indeed differentiates  $MoO<sub>2</sub>$  from insulating  $MoO<sub>3</sub>$ .



**Fig.** 35. (a) The distorted rutile structure of  $MoO<sub>2</sub>$  (monoclinic, P2<sub>1</sub>/c S.G.). (b) View of the  $(1 \times 1)$  tunnels in the rutile network. (c) Temperature-dependent resistivity. It shows a typical metallic behavior. The lowest resistivity is around  $6.45 \times 10^{-4} \Omega$  cm. reproduced with permission from [454]. Copyright 2014 Royal society of Chemistry. (d) Schematic band

diagram. The *d* orbital can be separated into two sub-bands, i.e., *t*2g and *e*<sup>g</sup> bands by the crystal field. The *t*2g band has a lower energy level and is partially filled by the two electrons, and the *e*<sup>g</sup> band is empty and has a higher energy level located above the Fermi energy.

## 6.2. Lithium insertion

Importantly,  $MoO<sub>2</sub>$  is considered to be one of the most studied anode systems for lithiumion batteries [455,456]. The lithium ions are inserted in the octahedral interstitial sites that are located in the tunnels between chains to form  $Li_xMoO_2$  as reported 30 years ago [457,458]. MoO<sup>2</sup> as Li-ion battery anode was systematically studied by Auborn and Barberio in 1987 for the first time [459] who demonstrated a relatively large capacity of 400-600 mAh  $g^{-1}$ .

The lithiation of  $MoO<sub>2</sub>$  is a subject of debate since different mechanisms have been described, i.e., insertion reaction and conversion reaction delivering a theoretical gravimetric capacity of 838 mAh  $g^{-1}$  and a theoretical volumetric capacity of 1357 mAh cc<sup>-1</sup> [460,461]. The first lithiation mechanism is an insertion-type reaction, which took place in the bulk and amorphous  $MoO<sub>2</sub>$  electrodes with only one-electron reduction as described by the equation [457]:

$$
MoO2 + xLi+ + xe- \leftrightarrow LixMoO2,
$$
 (19)

with 0≤*x*≤0.98. The second mechanism is a conversion reaction that gradually resolves  $Li<sub>x</sub>MoO<sub>2</sub>$  as described by the equation [462]:

$$
LixMoO2 + 3.02Li \leftrightarrow 2Li2O + Mo.
$$
 (20)

that evidences the formation of metallic Mo and Li2O. In-situ XRD structural investigations of the lithium ions insertion into  $MoO<sub>2</sub>$  have shown that the original monoclinic lattice  $(P2<sub>1</sub>/c)$ S.G.) transforms to another monoclinic structure (*P*21/*c* S.G.) via an orthorhombic phase (*Pnnm* S.G.). **Figure 36** presents the crystal structure of monoclinic MoO<sub>2</sub>, orthorhombic Li<sub>x</sub>MoO<sub>2</sub> and Li0.98MoO2. [463].



Fig. 36. The crystal structure of monoclinic MoO<sub>2</sub>, orthorhombic Li<sub>x</sub>MoO<sub>2</sub> and Li<sub>0.98</sub>MoO<sub>2</sub>. reproduced with permission from [463]. Copyright 2016 Elsevier.

The volume change associated with the Li insertion is only 11%. Based on Li<sup>+</sup> insertion and extraction reaction, the phase transition  $Mo_{2}$  to  $LiMo_{2}$  occurs in the potential range 2.5-1.0 V on discharge (reduction) and vice versa on charge (oxidation) [455,457,464]. In the first cycle, two pronounced reduction peaks were observed at 1.49 and 1.20 V, which may arise from the phase transition from orthorhombic to the monoclinic phase due the Li insertion [465]. Ku et al. [466] demonstrated that structural defects in amorphous  $MoO<sub>2</sub>$  electrodes can serve as a reversible Li<sup>+</sup> storage site for lithium secondary batteries. An unexpectedly high Li<sup>+</sup> storage capacity (up to four Li per  $MoO<sub>2</sub>$  unit), which is larger by a factor of four than that for the crystalline counterpart. The conversion-type lithiation is discarded for this electrode from the absence of Mo metal and Li<sub>2</sub>O in the lithiated  $\alpha$ -MoO<sub>2</sub> electrode with high specific capacity up to 810 mAh  $g^{-1}$ . In comparison to MoO<sub>3</sub>, MoO<sub>2</sub> has superior electronic conductivity, less charging potential, and higher volumetric capacity, which are highly important for application as an anode material in LIBs [467,468]. **Figure 37** presents the typical electrochemical behavior of a  $MoO<sub>2</sub>$  composite anode for LIB [468].



**Fig.** 37. Typical electrochemical properties of a  $MoO<sub>2</sub>$  composite anode for LIB. (a) Cyclic voltammograms conducted at a sweep rate of 58  $\mu$ V s<sup>-1</sup> in the voltage range of 0.005–3.0 V. CV curves contain strong redox couples centered at 1.25/1.47 and 1.54/1.72 V, which corresponds to Li intercalation and deintercalation into  $MoO<sub>2</sub>$  according to Eq. 1. (b) Galvanostatic discharge-charge profiles of  $Li_xMoO_2$  recorded at 100 mA  $g^{-1}$  current rate. (c) Chemical diffusion coefficient of Li ion in  $MoO<sub>2</sub>$  anode measured after 100 discharging/charging cycles. (d) Cyclability and coulombic efficiency over 100 cycles at 100

mA g<sup>-1</sup> current rate. Reproduced with permission from [468]. Copyright 2016 American Chemical Society.

MoO<sup>2</sup> has been synthesized using various techniques [469]: template method [455,470- 473], hydrothermal method  $[467,474-478]$ , reduction of MoO<sub>3</sub>  $[479-481]$ , soft chemistry route [464,465,482], spray pyrolysis [483-485], carbothermal reaction [486], ion-exchange route [487], and so on. Traditionally, molybdenum dioxide is prepared by reducing  $MoO<sub>3</sub>$  with hydrogen at high temperature and the product grows as micrometer size [479,480]. Manthiram et al. [488] and Liu et al. [489] reported the preparation of  $MoO<sub>2</sub>$  powders by a reduction process in solution reaction routes, but their products showed a limited success because their lowdimensional nanosized powders. Liang et al.  $[477]$  prepared  $MoO<sub>2</sub>$  nano-particles by a hydrothermal reaction route. The results show the resulting spherical, sheet-like and bar-shaped samples present a single phase  $MoO<sub>2</sub>$  with monoclinic structure.  $SnO<sub>2</sub>/MoO<sub>3</sub>$  nanowires were synthesized by a combination of hydrothermal (preparation of a white  $MoO<sub>3</sub>$  precipitate from the yellowish solution of  $MoO<sub>2</sub>$  powders dissolved in  $H<sub>2</sub>O<sub>2</sub>$ ) and wet chemical (decoration of  $MoO<sub>3</sub>$  nanowires by  $SnO<sub>2</sub>$  nanoparticles by mixing  $MoO<sub>3</sub>$  with  $SnCl<sub>2</sub>$  with addition of HCl) routes [228].

#### 6.3. MoO<sup>2</sup> anode for LIBs

The characteristic lithium-intercalation properties of spherical MoO<sup>2</sup> powders imply there is a correlation between their nano-sized structure and an irreversible phase transition after the initial discharge process.  $MoO<sub>2</sub>$  anode materials were made with various morphologies: nanobelts [490], nanorods [474,491], nanoporous powders [455,492], nanospheres [473,477] and nanosheets  $[493]$ . Rod type-MoO<sub>2</sub> was synthesized by two steps: preparation of hexagonal MoO<sub>3</sub> powders with rod shape followed by an ethanol-assisted reduction in 5% H<sub>2</sub>/Ar atmosphere [335]. Zhang et al. [474] have prepared uniform  $MoO<sub>2</sub>$  nanorods by a facile hydrothermal method, and the resultant samples were lithiated by subsequent treatment with LiCl solution. After lithiation, the  $MoO<sub>2</sub>$  nanorods possessed a similar crystalline structure to the unlithiated sample. At a current density of 20 mA  $g^{-1}$ , the lithiated specimen with better coulombic efficiency displayed a preferable cycling performance and exhibited excellent reversible charge-discharge capacity, while the unlithiated  $MoO<sub>2</sub>$  exhibited more irreversible behavior. MoO<sub>2</sub> nanosheets were synthesized for the first time by a solvothermal route using an ionic liquid [494]. When used as anode, the  $MoO<sub>2</sub>/C$  nanosheets exhibit a high reversible capacity of 620 mAh  $g^{-1}$  at 100 mA  $g^{-1}$  for 20 cycles.

Since 2000, many works achieve improved electrochemical performance of  $MoO<sub>2</sub>$  anode materials. Different architectures have been proposed including 2D [495,496] and 3D assemblies [455,460,493,497]. Spherical type-MoO<sub>2</sub> was prepared by ultrasonic spray pyrolysis of aqueous ammonium molybdate and  $5{\text -}20\%$  sucrose in 550-650 °C under 5% H<sub>2</sub>/Ar atmosphere [484]. Shi et al. [455] prepared highly ordered mesoporous  $MoO<sub>2</sub>$  by using phosphomobyldic acid as precursor and silica KIT-6 as hard template via nanocasting strategy. The mesoporous  $MoO<sub>2</sub>$  electrode shows a first discharge capacity of 960 mAh  $g<sup>-1</sup>$  (Li<sub>4.6</sub>MoO<sub>2</sub>) and a charge capacity of 630 mAh  $g^{-1}$  (Li<sub>3</sub>MoO<sub>2</sub>) in the potential range of 0.01-3.0 V. Sun et al. [498] fabricated MoO<sub>2</sub> monolith using cotton cloth that is utilized as a binder-free alternative anode that exhibits a high reversible Li-storage capacity of 719 mAh  $g^{-1}$  at 200 mA  $g^{-1}$  after 30 cycles. Manthiram and Tsang [464] synthesized amorphous  $MoO<sub>2+δ</sub>$  ( $\delta=0.3$ ) at ambient temperatures by a reduction of aqueous potassium molybdate solution with potassium borohydride. The amorphous sample exhibits excellent cyclability with a capacity of over 200 mAh g<sup>-1</sup> in the range 3 to 1 V in lithium cells with electrochemical behavior distinctly different from that of crystalline  $MoO<sub>2</sub>$ . Mo $O<sub>2</sub>$  nanoparticles (~100 nm) were synthesized by reduction of MoO<sup>3</sup> with ethanol vapor. This anode material exhibits good rate capability for the LIBs It displays an initial charge capacity of 318 mAh  $g^{-1}$ , which maintains 100% after 20 cycles in the range of 0.01–3.00 V at a current density of 5 mA cm<sup>-2</sup>, and around 85% of the retrievable capacity is in the range of 1.0–2.0 V [467].

Tremella-like  $MoO<sub>2</sub>$  consisting of nanosheets (NSs) were prepared using a Fe<sub>2</sub>O<sub>3</sub>-assisted hydrothermal reduction of  $MoO<sub>3</sub>$  in ethylenediamine aqueous solution [493]. As an anode material for LIB,  $MoO<sub>2</sub> NSs$  deliver a reversible capacity of 538 mAh  $g<sup>-1</sup>$  with the coulombic efficiency of 70% at the current density of 0.5 mA cm<sup>-2</sup> in the first cycle and sustains  $\sim$  300 mAh  $g^{-1}$  at 5.0 mA cm<sup>-2</sup> after 22 cycles. Ultrafine MoO<sub>2</sub> nanorods (~5 nm diameter) were successfully synthesized by a nanocasting method using mesoporous silica SBA-15 as hard template. The ultrafine  $MoO<sub>2</sub>$  nanorod electrode exhibits a first discharge capacity of 775 mAh  $g^{-1}$  (Li<sub>3.7</sub>MoO<sub>2</sub>) at C/20 rate in the voltage range 3.0-0.01 V. The capacity even reaches 830 mAh g<sup>-1</sup> after 29 cycles [491]. MoO<sub>2</sub> powder electrode was electrochemically activated [499] at high temperature. The lithiation behavior occurs by a one-electron reduction (by addition reaction) at room temperature. At elevated temperatures, this electrode is lithiated with fourelectron reduction by addition and continued conversion reaction. This phenomenon is named "thermoelectrochemical activation" because the extension from one- to four-electron reduction is achieved by a simple charge–discharge cycling made at elevated temperatures. This  $MoO<sub>2</sub>$ electrode delivers a reversible specific capacity that is close to the theoretical four-electron

capacity (838 mAh  $g^{-1}$ ) with an excellent cycle performance at room temperature. A templatefree solvothermal method has been developed to synthesize yolk–shell  $MoO<sub>2</sub>$  microspheres [500]. They are composed of both a uniform porous shell of 80 nm in thickness and a porous core constructed from primary  $MoO<sub>2</sub>$  nanocrystal clusters of 20 nm in size. Yolk–shell  $MoO<sub>2</sub>$ microspheres can deliver a high specific capacity of 955 mAh  $g^{-1}$  in the first discharge at a current density of 50 mA  $g^{-1}$ , and retain a reversible capacity of 847 mA  $g^{-1}$  after 50 cycles.

#### 6.4. MoO<sup>2</sup> composites

Several strategies have demonstrated the ability to mitigate the particle pulverization as a consequence of Li insertion and extraction and to improve the  $MoO<sub>2</sub>$  electrochemical performance via the fabrication of nanocomposites including carbonaceous materials. By scaling down to the particle size, it results in shortening the transport paths for both ions and electrons, while the carbonaceous matrix maintains high conductivity, large surface area and chemical stability. The composites studied as lithium battery anodes include various carbon forms, such as carbon cloth  $[501]$ , carbon coating  $[490,502-504]$ , MoO<sub>2</sub>/C spheres  $[460]$ , MoO<sub>2</sub>/carbon hybrid nanowires [482], MoO<sub>2</sub>/Mo<sub>2</sub>C composite [461,487], MoO<sub>2</sub>/ordered mesoporous carbon [471,505]. Nitride-coating [506] and Mo<sub>2</sub>N nanolayer-coating [507] are also attractive synthetic approaches.

Uniform carbon-coated MoO<sub>2</sub> nanospheres synthesized a one-pot hydrothermal method [502] exhibit significant improved electrochemical performance for high-rate lithium storage. Carbon-coated MoO<sup>2</sup> lengthened a few micrometers produced by carbothermal reduction of  $M_0O_3$  nanobelts were further ball-milled to reduce the particle size below 100 nm.  $C/M_0O_2$ nanoparticles demonstrate a discharge capacity of 248 mAh  $(g_{\text{MoO2}})^{-1}$  at 150 mA  $g^{-1}$  (0.5C rate) with an average operation voltage of 1.3 V. This anode material is also suitable for high rate charge and discharge, reaching 129 mAh  $g^{-1}$  for charge and 146 mAh  $g^{-1}$  for discharge even at 30C-rates (9 A  $g^{-1}$ ) [486]. Luo et al. [503] fabricated carbon-coated MoO<sub>2</sub> nanofibers (with a uniform carbonaceous shell of ∼3 nm in thickness and a hierarchical core made of primary MoO<sub>2</sub> nanocrystal clusters of ∼20 nm in size) through a controlled route based on single-nozzle electrospinning, air stabilization, and reduction/carbonization processes. These nanofibers exhibit a highly reversible capacity as high as  $762.7$  mAh  $g^{-1}$  over 100 cycles. Carbon coated MoO<sup>2</sup> nanobelts were synthesized via a hydrothermal method followed by calcination under inert atmosphere, using  $\alpha$ -MoO<sub>3</sub> nanobelts as the precursor and self-template, ethanol as the reducer and glucose as the carbon source [490]. Electrochemical tests of this anode material show a reversible capacity of 769 mAh  $g^{-1}$  at a current density of 100 mA  $g^{-1}$  in the first cycle, and retain 80.2% of the capacity after 30 cycles. Zhou et al. [462] have evaluated C-coated  $MoO<sub>2</sub>$  nanocomposite, i.e., interconnected  $MoO<sub>2</sub>$  nanocrystals with carbon nanocoating, prepared by one-pot hydrothermal route that show high specific capacities of ~640 mAh  $g^{-1}$  at 200 mA g<sup>-1</sup> and ~575 mAh g<sup>-1</sup> at 400 mA g<sup>-1</sup>. Chen et al. [470] designed a novel MoO<sub>2</sub>-ordered mesoporous carbon (MoO2–OMC) hybrid through a two-step solvothermal chemical reaction route. Electrochemical tests reveal a first discharge capacity of 1642 mAh g−1 with an ICE of 63.6%, and a reversible capacity as high as 1049 mAh  $g^{-1}$  even after 50 cycles at a current density of 100 mA  $g^{-1}$ . The MoO<sub>2</sub>–OMC hybrid demonstrates an excellent high rate capability with capacity of ~600 mAh g<sup>-1</sup> even at a charge current density of 1600 mA g<sup>-1</sup> after 50 cycles. A nano-sized  $MoO<sub>2</sub>/C$  composite was synthesized using a spray pyrolysis technique, and investigated as an anode material for Li-ion batteries. Spherical  $MoO<sub>2</sub>/C$  particles with the monoclinic phase  $(P2)/c$  S.G.) were obtained without any impurities, and with a primary particle size in the range 30–50 nm [483]. Liu et al. [475] synthesized MoO<sub>2</sub>/C composite nanospheres with cage-like structure by a simple hydrothermal reduction process using ammonium molybdate tetrahydrate  $((NH_4)_6Mo_7O_{24}·4H_2O)$  and ascorbic acid as a structure directing agent, reductive agent and carbon source. These porous structured electrodes can deliver a capacity of 692.5 mAh  $g^{-1}$  after 80 charge–discharge cycles at a current density of 200 mA g<sup>-1</sup> and can maintain excellent cycling stability (about 550 mAh g<sup>-1</sup> after 475 cycles). Nanostructured monolithic hybrid mat made of C/MoO2+*<sup>δ</sup>* nanofibers were fabricated through an electrospinning process by thermal treatment in N<sub>2</sub>. The free-standing anode  $C/MoO<sub>2+\delta</sub>$ composite exhibits a reversible capacity as high as 877 mAh  $g^{-1}$  after 250 cycles at a current density of 200 mA  $g^{-1}$ . A capacity of ~448 mAh  $g^{-1}$  could be maintained after 1000 cycles at 2000 mA  $g^{-1}$  rate [508]. Recently, Qing et al. [509] fabricated the composite MoO2/VC via a facile hydrothermal route, where vitamin C (VC) is a source of carbon. This composite LIB anode displays a discharge capacity of  $\sim$ 300 mAh g<sup>-1</sup> at 0.5 A g<sup>-1</sup> after 120 cycles.

Several MoO2/Mo2C heterostructures have been fabricated as anode materials for LIBs [461,487,510]. MoO<sub>2</sub>/Mo<sub>2</sub>C heteronanotubes were fabricated by a one-step carbothermal reduction by using a mesoporous carbon CMK-3 acting as template and reactant  $[461]$ . Mo<sub>2</sub>C is an electrochemical inactive compound, which exhibits a high conduction of  $\sim 10^{-2}$  S cm<sup>-1</sup> and minimizes the charge transfer resistance of the composite. Cycled at 200 and 1000 mA  $g^{-1}$  for 140 cycles, the discharge capacities of the MoO2/Mo2C heteronanotubes remain to be 790 and 510 mAh g<sup>-1</sup>, respectively. Ihsan et al. [510] synthesized MoO<sub>2</sub>/Mo<sub>2</sub>C/C spheres through hydrothermal and calcination processes that demonstrates not only high cycling performance, but also good rate capability when they are used as anode materials for LIBs. After 100 cycles

at 100 mA  $g^{-1}$ , the discharge capacities of the MoO<sub>2</sub>/Mo<sub>2</sub>C/C spheres as anode materials remain at 800 mAh  $g^{-1}$ .

Zhang et al.  $[460]$  synthesized uniform hierarchical MoO<sub>2</sub>/C spheres through calcination of a MoO3/resin precursor generated via a hydrothermal method in the presence of resorcinol, formaldehyde and  $(NH_4)_6M_07O_{24}$  4H<sub>2</sub>O. Used as an anode material, these spheres exhibit an initial capacity of  $\sim$ 400 mAh g<sup>-1</sup> at 0.5 A g<sup>-1</sup> and high cycling performance after 400 cycles with capacities of ~600 and ~500 mAh  $g^{-1}$  at 0.5 and 1.0 A  $g^{-1}$  rate, respectively. MoO2/carbon nanowires were prepared by a method based on organic–inorganic hybrid nanocomposites with nanometer periodic structures  $[482]$ . The MoO<sub>2</sub>/C hybrid nanowires show good rate capability even at the high current density of 1000 mA  $g^{-1}$  as anode material for LIBs.

MoO2-loaded porous carbon hollow sphere composite materials were synthesized using MoO<sub>2</sub> nanoparticles with an average size of ∼12 nm uniformly dispersed in the shells of the porous carbon hollow spheres (PCHS) [511]. The  $MoO<sub>2</sub>/PCHS$  composite containing 44.2% of MoO<sub>2</sub> revealed a reversible capacity of 574 mAh  $g^{-1}$  at a current density of 50 mA  $g^{-1}$ . After 80 cycles, this composite still retained a capacity of 640 mAh  $g^{-1}$  due to the presence of porous carbon, which prevents volume change or aggregation of the MoO<sub>2</sub> nanoparticles during the Li ion insertion/extraction process. Zeng et al. [471] synthesized the nanocomposite of  $MoO<sub>2</sub>$ ordered mesoporous carbon ( $MoO<sub>2</sub>-OMC$ ) using a carbon thermal reduction route. This nanocomposite exhibits a large reversible capacity of 689 mAh  $g^{-1}$  after 50 cycles at a current density of 50 mA  $g^{-1}$ . This nanocomposite offered a better accommodation of the strain and volume changes and a shorter path for Li-ion and electron transport, leading to the improved capacity and enhanced rate capability of the anode material for LIBs.

Carbon-coated MoO<sub>2</sub> nanobelts were fabricated via a hydrothermal route followed by calcination under inert atmosphere, using  $\alpha$ -MoO<sub>3</sub> nanobelts as the precursor and self-template, ethanol as the reducer and glucose as the carbon source. The  $C/MnO<sub>2</sub> NBs$  anode materials exhibit a reversible capacity of 769 mAh  $g^{-1}$  at a current density of 100 mA  $g^{-1}$  in the first cycle, and retain 80.2% of the capacity after 30 cycles [490]. Nanorods (250 nm in diameter), which consist of α-MoO<sub>3</sub> core and β-MnO<sub>2</sub> shell (46 nm thick, tetragonal phase) were synthesized by a two-step hydrothermal process [214]. The α-MoO<sub>3</sub>@β-MnO<sub>2</sub> nanorods show a reversible capacity of 1475 mAh  $g^{-1}$  at 0.1C rate and retains at 1127 mAh  $g^{-1}$  after 50 cycles. Even at high current rate of 6C, the reversible capacity is 394 mAh  $g^{-1}$  and retains at 286 mAh  $g^{-1}$  after 50 cycles. A heterostructure, which consists of MoO<sub>2</sub> nanoparticles anchored and embedded in the 1D N-doped carbon matrix were fabricated using the organic–inorganic  $M_{03}O_{10}(C_6H_8N)_2.2H_2O$  hybrid nanowires as a precursor and self-template. The nanowires were

calcined at 650 °C for 5 h under an Ar flow  $(50 \text{ mL min}^{-1})$  to yield the MoO<sub>2</sub>/N-C heteronanowires. Tested at a current density of 2 A  $g^{-1}$ , the MoO<sub>2</sub>/N-C composite delivers a reversible capacity of 700 mAh g<sup>-1</sup> after 400 cycles, which still remains 570 mAh g<sup>-1</sup> even after 1500 cycles [512]. Nanoporous 3D MoO<sup>2</sup> single crystals were partially covered with a few atomic layers of  $MoS<sub>2</sub>$  to form  $MoS<sub>2</sub>/MoO<sub>2</sub>$  nanonetworks, which exhibit a reversible discharge specific capacity of 1233 mAh  $g^{-1}$  at a current density of 100 mA  $g^{-1}$ . When discharged at 200 and 500 mA  $g^{-1}$ , the capacities are 1158 and 826 mhA  $g^{-1}$  [513]. Wang et al. [514] prepared a mesoporous MoO2@C nanocomposite using oleic acid to reduce the MoO<sup>3</sup> precursor and to simultaneously coat the resultant 1D MoO<sub>2</sub> nanorods with carbon layers (BET surface area of 45.7 m<sup>2</sup> g<sup>-1</sup>, and pore size of 3.8 nm). The MoO<sub>2</sub>@C electrode exhibits a high capacity of 1034 mAh g<sup>-1</sup> at 0.1 A g<sup>-1</sup>, and a capacity of 155 mAh g<sup>-1</sup> at a super-high current rate of 22 A g<sup>-1</sup>. When, the Li/MoO<sub>2</sub>@C half cells cycled at 0.5 and 10 A  $g^{-1}$  retain 861 and 312 mAh  $g^{-1}$ capacity after 140 and 268 cycles, respectively. Recently, Zhang et al. [515] reported the design and fabrication of a composite, which consists of encapsulated MoO<sub>2</sub> nanocrystals into flexible carbon nanofibers, using a needle-free electrospinning method combined with the subsequent carbonization process. The  $MoO<sub>2</sub>/C$  nanofiber membrane as self-supporting anode exhibits a discharge capacity of 450 mAh g−1 after 500 cycles at 2000 mA g−1 . **Table 13** summarizes the electrochemical performance of various MoO<sub>2</sub> composites as anode materials for LIBs.

		Reversible	Current rate	Ref.
Material	Synthesis	capacity	$(mA g^{-1})$	
		$(mAh g-1)$		
Nano $MoO2$	Rheology	402	100(40)	[516]
MoO <sub>2</sub> /Mo <sub>2</sub> N	Reduction of $MoO3$	815	100(150)	[517]
MoO <sub>2</sub> /graphene	Chemical vapor deposition	986	50 (150)	$[518]$
MoO <sub>2</sub> /C	Ion exchange	574	100(100)	[519]
MoO <sub>2</sub> /C	Carbothermal reduction	500	100(50)	[471]
$MoO2/C$ hollow spheres	Solvothermal	580	200 (200)	$[520]$
Mesoporous $MoO2$	Template casting	750	42(30)	[455]
Activated $MoO2$	Electrochemical activation	850	100(30)	[521]
MoO <sub>2</sub> HCSMSs	<b>Hydrolysis</b>	420	50(30)	$[522]$
$W$ -doped $MoO2$	Nanocasting	670	75 (20)	[492]
$C/WO_x/MoO_2$	Hydrothermal	670	90(50)	[523]
MoO <sub>2</sub> /C NWs	Solvothermal	500	200 (20)	[482]
$C/MoO2$ NSs	Hydrothermal+annealing	675	838 (30)	$[502]$
$MoS_2/M_0O_2$	Sulfur assisted	654	500 (80)	[513]
C/MoO <sub>2</sub> NBs	Hydrothermal+annealing	617	100(30)	[490]
$MoO2$ monolith	Morphosynthesis	719	200 (20)	[498]

**Table 13**. Electrochemical performance of various  $MoO<sub>2</sub>$  composites as anode materials for Li-ion batteries. The relevant cycle number is displayed in brackets.



6.5. MoO2/graphene composites

To overcome the sluggish lithiation/delithiation kinetics of micron-scale particles, several graphite-based nanocomposites have been created such as graphite oxide [524], and graphene  $(GR)$  [465,525,526], graphene oxide  $(GO)$  [465,527-529], and reduced graphene oxide (rGO) [530-534]. Using a microwave-assisted hydrothermal process and subsequent thermal annealing, Palanisamy et al.  $[535]$  prepared 3D porous self-assembled MoO2/graphene microspheres. Such hybrid is built from interconnected  $MoO<sub>2</sub>$  nanoparticles (3-5 nm) in the graphene matrix. The  $MoO<sub>2</sub>/graph$ ene composite anode delivers a high reversible capacity of ~1300 mAh  $g^{-1}$  after 80 cycles at C/10 rate and excellent rate capability (913 and 390 mAh  $g^{-1}$ at 2C and 5C rates, respectively). Sun et al. [465] have investigated self-assembled hierarchical MoO2/graphene nanoarchitectures fabricated on a large scale through a facile solution-phase process and subsequent reduction of the Mo-precursor/graphene composite. The as-formed MoO<sub>2</sub>/graphene nanohybrid (containing 11.2 wt.% graphene) as an anode material for LIBs exhibits not only a highly reversible capacity but also an excellent cycling performance as well as good rate capability, a specific capacity of 597 mAh  $g^{-1}$  is delivered after 70 cycles at current density of 1000 mA  $g^{-1}$ . Tang et al. [528] reported the electrochemical performance of MoO2/graphene nanocomposite synthesized via a two-step of hydrothermal-calcination method. When used as anode material for LIBs, the MoO2/graphene composite shows a first discharge capacity of 674 mAh  $g^{-1}$  with a reversible capacity of 430 mAh  $g^{-1}$ . Significantly, the composite can also deliver a reversible capacity of as high as1009 mAh  $g^{-1}$  after 60 charge/discharge cycles. Chen et al. [481] reported the electrochemical properties of hierarchical nanocomposites including MoO<sub>2</sub> nanocrystal-functionalized graphene synthesized by in-situ reduction process. The discharge capacity is still up to 997 mAh  $g^{-1}$  after 50 cycles and 370 mAh  $g^{-1}$  after 200 cycles at 10C rate.

Nanostructured MoO<sub>2</sub>/GO composites, which consists of a mixture of MoO<sub>2</sub> nanorods and nanoparticles uniformly distributed at the surface of GO sheets, were synthesized by a simple solvothermal method of a GO suspension with  $(NH<sub>4</sub>)<sub>6</sub>Mo<sub>7</sub>O<sub>24</sub>$  and HNO<sub>3</sub> in ethanol at 160 °C for 16 h  $[524]$ . The MoO<sub>2</sub>/GO with 10 wt.% GO tested in lithium half-cell delivered an initial capacity of 800 mAh  $g^{-1}$  at current density of 0.8 A  $g^{-1}$  corresponding to Li uptake of 3.43 moles and a reversible capacity of 720 mAh  $g^{-1}$  at a current density of 100 mA  $g^{-1}$  and 560 mAh  $g^{-1}$  at

a high current density of 800 mA  $g^{-1}$  after 30 cycles. The improved reversible capacity, rate capacity, and cycling performance of the composites are attributed to synergistic reaction between MoO<sub>2</sub> and GO. MoO<sub>2</sub>/rGO composites fabricated by hydrothermal route followed by a heat treatment at 500 °C for 2 h [530]. The MoO<sub>2</sub>-based materials have monoclinic crystal structure, and worm like shape with average dimensions of 100-200 nm width and 0.5-1 μm length. There are two steps of Li ion intercalation/de-intercalation for the  $MoO<sub>2</sub>/rGO$  anode at the potential ranging from 1.0 to 3.5 V vs. Li<sup>+</sup>/Li, locating at  $1.60/1.75$  and  $1.25/1.40$  V, respectively with a first discharge capacity of 221 mAh  $g^{-1}$ and  $R_{\text{SEI}}$  and  $R_{\text{CT}}$  resistances of 2-4 Ω and below 5 Ω, respectively. Petnikota et al. [468] studied the insertion of Li and conversion mechanism of MoO<sub>2</sub> (below 0.8 V) in exfoliated graphene oxide/MoO<sub>2</sub> composites as anode materials. Graphene oxide was used as a reducing agent of  $MoO<sub>3</sub>$  (molar ratio of 4:1) and as a source for exfoliated graphene oxide (EGO) to fabricate EGO/MoO<sub>2</sub> composites with 46 wt.% graphene. The small particle size of  $\sim$ 33 nm indicates that agglomeration of MoO<sub>2</sub> has been controlled with the increased amount of EGO. The initial capacity of 713 mAh  $g^{-1}$  increased slightly to 878 mAh g<sup>-1</sup> after 100 cycles at 100 mA g<sup>-1</sup> current rate in the voltage window 0.005-3.0 V. Kinetics investigated by cyclic voltammetry revealed high values of the diffusion coefficients of Li ions in the range  $6 \times 10^{-10}$ -1  $\times 10^{-11}$  cm<sup>2</sup> s<sup>-1</sup> during Li<sub>x</sub>MoO<sub>2</sub> lithiation (0 < *x*  $<$  4). Hu et al. [478] prepared MoO<sub>2</sub>/GO composites via hydrothermal route at 200 °C for 24 h using Mo powder and exfoliated GO produced by sonication. MoO<sup>2</sup> nanoparticles (5−15 nm in size) were homogeneously dispersed on GO. When tested as an anode material for lithium, the MoO<sub>2</sub>/GO composite displayed an initial specific capacity of 780 and 500 mAh  $g^{-1}$  at 0.1 and  $0.5$  A  $g^{-1}$ , respectively. The beneficial contribution of exfoliated GO was evidenced by EIS measurements, as the charge transfer resistance of 44 W for  $MoO<sub>2</sub>/GO$  was much lower than 300 W for pure MoO<sup>2</sup> nanoparticles, due to more conductive pathway for Li-ion transportation. Huang and coworkers [536] fabricated a composite of  $MoO<sub>2</sub>$  nanoparticles (~100 nm) anchored on GO for sodium-ion battery anodes. This MoO2/GO hybrid (15 wt.% GO) yielded a discharge capacity of 483 mAh  $g^{-1}$  (~2318 mAh cm<sup>-3</sup>) at the current density of 100 mA  $g^{-1}$  and a capacity fading of 0.019% per cycle over the first 1000 cycles. Tang et al. reported the electrochemical properties of rGO-wrapped MoO<sup>2</sup> porous nanobelts (rGO/MoO<sup>2</sup> NBs) synthesized using a reduction process of the GO-MoO<sub>3</sub> hybrids with Ar/ethanol vapor at 400 °C for 12 h [537]. The full-scale multi-electron conversion reaction in rGO/MoO<sup>2</sup> NBs (with 8 wt.% rGO) led to a high reversible capacity of 974 mAh g<sup>-1</sup> at the current density of 60 mA g<sup>-1</sup> after 6<sup>th</sup> cycle and, after initial activation, a specific capacity of  $420$  mAh  $g^{-1}$  was retained at the end of 1900 cycles at 5 A  $g^{-1}$  rate. Zhu et al. [216] demonstrated that rGO/MoO<sub>2</sub> nanobelt composite (with 19.4

wt.% rGO) synthesized by a simple thermal reduction of  $MoO<sub>3</sub>$  and GO at 550 °C for 2 h in a 10% H<sub>2</sub>/Ar atmosphere had a BET specific surface area and pore volume of 17.68 m<sup>2</sup> g<sup>-1</sup> and 0.09 cm<sup>3</sup>  $g^{-1}$ , respectively. This self-assembled hybrid shows attractive electrochemical performance as LIB anodes with specific discharge capacity reaching 584 mAh g<sup>-1</sup> after 100 cycles at  $1 \text{ A g}^{-1}$ .

In 2017, Ju and coworkers [538] designed a  $TiO<sub>2</sub>/rGO/MoO<sub>2</sub>@Mo$  composite electrode for ESC. This architectural nanostructure with 3D walnut-shape synthesized by one-step hydrothermal method was composed of amorphous TiO<sub>2</sub> decorating the rGO/MoO<sub>2</sub> surface grown on Mo foam. It exhibited a specific capacitance of 1636 F  $g^{-1}$  at 1.25 A  $g^{-1}$  with only 3.5% capacitance loss after 5000 cycles. Tightly anchored MoO<sup>2</sup> nanoparticles (10-20 nm in size) deposited on the basal plane of rGO were fabricated using a green supercritical methanol (scMeOH) route. When tested as an anode in lithium cells, the composite with  $37$  wt.% MoO<sub>2</sub> loading (BET surface area of 14.3 m<sup>2</sup> g<sup>-1</sup>, 15% porosity) delivered a reversible capacity of 793 mAh  $g^{-1}$  at 50 mA  $g^{-1}$  current rate [531]. Graphene oxide flexibly supported MoO<sub>2</sub> porous hybrids were constructed by decomposition of (NH4)2MoO<sup>4</sup> /GO preforms with the percentage of GO at 4.3, 15.2, and 20.8 wt.%. The lithium storage performance was investigated at the current density of 0.1 A  $g^{-1}$  using various MoO<sub>2</sub>/GO architectures showing specific capacities of 901, 1127, and 967 mAh  $g^{-1}$  after 100 cycles for micrometer MoO2/GO, MoO2/GO nanohoneycomb, and layered  $MoO<sub>2</sub>/GO$ , respectively. The  $MoO<sub>2</sub>/GO$  nanohoneycomb delivered the capacity of 461 mAh  $g^{-1}$  at 5 A  $g^{-1}$  current rate with CE of 99.3% [539]. Wang et al. [540] synthesized 2D layered mesoporous-MoO2/rGO composites (m-MoO2/rGO) through a novel strategy using silica KIT-6/rGO as a template and ammonium molybdate as a precursor via a nanocasting method with subsequent heat treatment heated up to 600 °C for 6 h at a rate of 2 °C min<sup>-1</sup> in a mixed gas flow of H<sub>2</sub>/Ar (10:90). Analyses of the morphology validated the ordered mesoporous structure with a BET specific surface area of 86  $m^2 g^{-1}$ , a pore size of 3-4 nm in a large domain and a pore volume of 0.19 cm<sup>3</sup>  $g^{-1}$ . The m-MoO<sub>2</sub>/rGO(1:1) electrode delivered an initial discharge capacity of 1160 mAh  $g^{-1}$  at current density of 0.1 A  $g^{-1}$  with a capacity loss of 31.9% during the first cycle and a reversible capacity is 801 mAh  $g^{-1}$  after 50 cycles.

In 2018, Chen et al. [541] investigated a hierarchical MoO<sub>2</sub>@rGO (~10 nm MoO<sub>2</sub> particle size and 21.1 wt.% rGO) composite prepared by hydrothermal route at 200 °C for 2 days, which exhibited outstanding electrochemical performance with a reversible capacity of 708 mAh g<sup>-1</sup> at a current density of 0.5 A  $g^{-1}$  after 50 cycles. A capacity of 473 mAh  $g^{-1}$  was maintained when cycled at  $2 \text{ A } g^{-1}$  current rate. Li et al. [542] utilized Mo particles as both the reductant for GO and as the Mo source to prepare  $MoO<sub>2</sub>/rGO$  hybrid anodes. The  $MoO<sub>2</sub>/rGO$  composite with Mo:GO of 3:1 was used as an efficient cathode catalyst for the oxygen reduction reaction (ORR). Recently, Liu et al. [532] reported the direct growth of  $MoO<sub>2</sub>/rGO$  hollow sphere composites through hydrothermal technique. This advanced anode materials tested in potassium-ion batteries delivered a reversible specific capacity of 219 mAh  $g^{-1}$  after 200 cycles at 50 mA  $g^{-1}$  current rate. When cycled at 0.5 A  $g^{-1}$ , a high charge specific capacity of 104 mAh g<sup>-1</sup> was achieved after 500 cycles. Li et al. [543] prepared the MoO2/Mo-GO hybrids using a freeze-drying technique, which exhibited a high ICE of 97%. A capacity of 550 mAh  $g^{-1}$  ws delivered by the MoO<sub>2</sub>/Mo-GO anode materials after 150 cycles at 0.1 A  $g^{-1}$  rate. Devina et al. [544] used the same technique to prepare  $MoO<sub>2</sub>/rGO$  composite attached with  $Mo<sub>2</sub>C$  during carbothermal hydrogen reduction. The  $MoO<sub>2</sub>/Mo<sub>2</sub>C/rGO$  product displayed an increased ICE (77%), long-term cyclability (500 mAh  $g^{-1}$  at 50 mA  $g^{-1}$  after 150 cycles), and high-rate performance (200 mAh  $g^{-1}$  at 1 A  $g^{-1}$ ) compared with those of the MoO2/rGO composite (53%; 280 mAh g<sup>-1</sup> at 50 mA g<sup>-1</sup> after 150 cycles; 120 mAh g<sup>-1</sup> at 1 A g<sup>-1</sup>).

Recently, the fabrication of submicrometer-sized  $MoO<sub>2</sub>@MoS<sub>2</sub>/rGO$  composite with a hierarchical core-shell structure was suggested by Zheng et al. [545]. The three-step synthesis includes: (i) the preparation of the  $MoO_2@MoS_2$  precursor using a solid-state reaction (i.e., MoO<sub>3</sub> and sulfur heated at 600 °C for 1 h in a gas stream of Ar/H<sub>2</sub> (90:10)), (ii) a hydrothermal process at 180 °C for 4 h using a GO suspension, and (iii) a final heat treatment at 400 °C for 2 h in Ar/H<sub>2</sub> flowing gas. The electrochemical tests of the  $MoO<sub>2</sub>@MoS<sub>2</sub>/rGO$  anode showed an initial reversible capacity of 833 mAh  $g^{-1}$  with ICE of 80.6% and improved cycling stability (733 mAh g<sup>-1</sup> after 80 cycles at 0.2 mA g<sup>-1</sup>). The good rate capability with a capacity of 645 mAh g<sup>-1</sup> at 4 A g<sup>-1</sup> was also evidenced. An assembled MoO<sub>2</sub>@MoS<sub>2</sub>/rGO LiCoO<sub>2</sub> full cell maintained a reversible capacity of 694 mAh  $g^{-1}$  (based on the mass of MoO<sub>2</sub>@MoS<sub>2</sub>/rGO) at 0.2 mA g−1 after 40 cycles. Electrochemical performance of MoO2/rGO composite anodes for LIBs from the recent literature are summarized in **Table 14**.







a) graphene content in wt.%.

#### 6.6. Lithium molybdate  $LiMoO<sub>2</sub>$

Molybdenum also forms several stable phases with lithium including  $LiMoO<sub>2</sub>$  [551], Li<sub>2</sub>MoO<sub>3</sub> [552], Li<sub>4</sub>Mo<sub>3</sub>O<sub>8</sub> [553] and non-stoichiometric compounds such as Li<sub>0.74</sub>MoO<sub>2</sub> [551]. There were few reports on the structure, physico-chemical properties and electrochemistry of LiMoO<sub>2</sub> [554,555]. However, its structure has been the subject of debate. Several groups have claimed that  $LiMoO<sub>2</sub>$  crystallizes with the rock-salt-like structure isomorphs to the layered  $\alpha$ -NaFeO<sub>2</sub> [551,554,556,557]. These groups did not consider the accepted theoretical predictions for 3D-systems according to which the structure would distort, while Hibble et al. [558] have determined that LiMoO<sub>2</sub> adopts the monoclinic *C2/m* space group by refinement of by neutron scattering data. The structure is best described with  $a = 10.543(6)$  Å,  $b = 2.8626(5)$ Å,  $c = 10.899(6)$  Å, and  $β = 153.29°$ . The schematic representation is given in Fig. 38a. Ben-Kamel et al.  $[555]$  synthesized LiMoO<sub>2</sub> by a carbothermal method using citric acid as a chelating agent with a subsequent thermal treatment at 800 °C for 72 h in air. Analysis of the crystal structure by Rietveld refinements shows that lithium molybdate crystallizes in the monoclinic system ( $R_{\text{Bragg}} = 1.24\%$ ) instead of the rhombohedral structure ( $R_{\text{Bragg}} = 8.53\%$ ). This result is confirmed by the Raman spectroscopy experiment, which displays twelve bands instead of two for the rhombohedral  $(R\overline{3}m)$  structure. **Figure 38b** represents the charge– discharge profile of the  $Li/LiMoO<sub>2</sub>$  cell cycled in the potential range 2–4 V at current density of  $C/24$ . Almost 0.9Li can be extracted when the cell is charged at 4.0 V vs. Li<sup>+</sup>/Li. A good reversibility at the first cycle with the capacity retention of 168 mAh  $g^{-1}$ , which can be

compared with the theoretical value 198 mAh  $g^{-1}$ . Measurements reveal the presence of multiple phase transitions during the lithium extraction process: the S-shaped potential corresponds to the extraction of  $0.5Li$  in  $Li<sub>x</sub>MoO<sub>2</sub>$ , the formation of an ordered composition close to  $x=0.5$  and a plateau at 3.6 V attributed to the existence of a two-phase system, i.e., Li0.1MoO<sup>2</sup> and Li0.5MoO2. The differential capacity curve (inset **Fig. 38b**) displays a sharp peak at ca. 3.6 V corresponding to the voltage plateau observed in the charge–discharge curves and redox contributions of the single-phase regime appearing at 2.9 and 3.1 V vs. Li<sup>+</sup>/Li. Measurements of Li ion kinetics reveal that diffusion coefficients for the LiMoO<sub>2</sub> electrode are in the range of  $10^{-12}$ – $10^{-10}$  cm<sup>2</sup> s<sup>-1</sup> (Fig. 38c). The rate capability of LiMoO<sub>2</sub> compared with that of LiNi<sub>0.8</sub>Co<sub>0.2</sub>O<sub>2</sub> is shown in Fig. 38d.



**Fig.** 38. (a) Schematic representation of the rhombohedral  $(R\overline{3}m)$  and monoclinic  $(C2/m)$ structures of LiMoO<sub>2</sub>. (b) Charge–discharge profile of the Li// LiMoO<sub>2</sub> cell cycled in the potential range 2–4 V at C/24 rate. The electrode was LiMoO<sub>2</sub> powders synthesized at 600 °C for 72 h ( $C2/m$  S.G.) and the electrolyte was 1 mol  $L^{-1}$  LiPF<sub>6</sub> in 1:1 EC/DMC. The insert shows the differential capacity d*Q*/d*V* vs. cell voltage obtained at the second cycle. (c) Evolution of the chemical diffusion coefficient of  $Li^+$  ions in the  $Li_xMoO_2$  cathode material as a function of the composition *x*. (d) Rate capability of the LiMoO<sub>2</sub> electrode recorded at the  $5<sup>th</sup>$  cycle. Data

obtained for the layered  $LiNi<sub>0.8</sub>Co<sub>0.2</sub>O<sub>2</sub>$  cathode are shown for comparison. Reproduced with permission from [555]. Copyright 2012 Elsevier.

### 6.7. Lithium molybdate  $Li<sub>2</sub>MoO<sub>3</sub>$

Because of high lithium storage capacity, high first coulombic efficiency, low cost and  $Mo(IV)/Mo(VI)$  redox couple,  $Li_2MoO_3$ , can serve as anodes [559] as well as cathodes for LIBs [560,561]. Li<sub>2</sub>MoO<sub>3</sub> has a theoretical Li<sup>+</sup> extraction/insertion capacity of 339 mAh  $g^{-1}$  simply by a single  $Mo^{4+}/Mo^{6+}$  redox reaction. Li<sub>2</sub>MoO<sub>3</sub> synthesized through ball milling and thermal reduction presents a first discharge capacity of about 835 mAh g−1 with an ICE of 97.6% at a current density of 100 mA  $g^{-1}$ ; a discharge capacity of 902 mAh  $g^{-1}$  is preserved after 150 cycles [559]. Phase-pure  $Li<sub>2</sub>MoO<sub>3</sub>$  powders were prepared by reducing commercial  $Li<sub>2</sub>MoO<sub>4</sub>$ at 650 °C for 24 h in flowing H<sub>2</sub>/Ar (10:90 v/v) [562]. A fresh electrode delivers ~80 mAh g<sup>-1</sup> after 20 cycles at 10 mA  $g^{-1}$  with ICE of 98.9%, whereas the capacity of aged Li<sub>2</sub>MoO<sub>3</sub> (obtained after 120 days in relative humidity <10%) decreases drastically (75% capacity loss at the  $2<sup>nd</sup>$  cycle). Kumakura et al. [563] prepared a  $Li<sub>2</sub>MoO<sub>3</sub>/C$  composite with the simple addition of acetylene black. As a cathode material tested in the range 1.5-4.3 V, it delivers a first specific capacity of 230 mAh  $g^{-1}$ . The polycrystalline Li<sub>2</sub>MoO<sub>3</sub> synthesized via simple liquid chemical reaction, followed by thermal reduction in H<sub>2</sub>/Ar (5:95 v/v) atmosphere exhibits a first discharge capacity of 836 mAh  $g^{-1}$  (ICE = = 94.5%) at a current density of 100 mA  $g^{-1}$  over a voltage window of 0.01–3.0V. After 200 cycles at 300 mA  $g^{-1}$ , a discharge capacity of 654 mAh g<sup>-1</sup> is preserved and the composite still keeps a discharge capacity of 489 mAh g<sup>-1</sup> at 1600 mA  $g^{-1}$  current rate [564]. Recently, Mao et al. [565] investigated the electrochemical performance of layered  $Li<sub>2</sub>MoO<sub>3</sub>$  microspheres prepared from  $MoO<sub>2</sub>$  microsphere as a selftemplate. Due to their enhanced mesoporous structure,  $Li_2MO_3$  microspheres deliver specific capacities of 247 and 76 mAh  $g^{-1}$  at current densities of 34 and 1000 mA  $g^{-1}$ , respectively. **Table 15** list the electrochemical performance of some Li<sub>2</sub>MoO<sub>3</sub> used as cathode and anode of LIBs.

Table 15. Electrochemical performance of Li<sub>2</sub>MoO<sub>3</sub> cathode/anode materials for LIBs. Relevant cycle number is given in brackets.

Material	Synthesis	Specific capacity $(mAh g-1)$	Current rate $(mA g^{-1})$	Ref.
Cathode				
$Li2Mo0.9Co0.1O3$	Solid phase	140	5(50)	[566]
C/Li <sub>2</sub> MoO <sub>3</sub>	<b>Ball milling</b>	180	20(50)	$[567]$
C/Li <sub>2</sub> MoO <sub>3</sub>	Solid state reaction	150	16(35)	$[563]$



6.8. Other lithium molybdates

Monoclinic  $Li_{13}MO_{2}O_{4}$  was prepared by low temperature ion exchange of lithium for sodium in  $\text{Na}_{x}\text{Mo}_{2}\text{O}_{4}$  [568]. The study of the  $\text{Li}/\text{Li}_{0.3}\text{Mo}_{2}\text{O}_{4}$  electrochemical cells shows that the electrode material can reversibly accommodate up to 1.7 Li per  $Mo<sub>2</sub>O<sub>4</sub>$  unit formula at an average of 3.0 V resulting in a theoretical energy density of 530 Wh kg<sup>-1</sup> for the  $Li_2Mo_2O_4$ cathode material. Structural studies indicate that  $Li_xMo_2O_4$  is a multiphase intercalation system over the range of composition  $0.3 < x < 2.0$ . Li<sub>2</sub>MoO<sub>4</sub> nanoparticles (hexagonal phenacite-like structure) have investigated as anode materials for LIBs [569,570]. Liu and coworkers [570] synthesized 1D Li<sub>2</sub>MoO<sub>4</sub> nanostructures (nanorods and nanotubes) via a sol-gel method adding  $Li<sub>2</sub>CO<sub>3</sub>$  and MoO<sub>3</sub> powders into distilled water with citric acid as an assistant agent and carbon source. The materials display initial discharge/charge capacities of  $688/319$  mAh g<sup>-1</sup> (carbonfree powders), 750/420 mAh  $g^{-1}$  (carbon-free nanorods), 850/476 mAh  $g^{-1}$  (carbon-free nanotubes), 1016/648 mAh  $g^{-1}$  (carbon-coated nanotubes). After 50 cycles at 90 mA  $g^{-1}$  current rate, the carbon-coated Li<sub>2</sub>MoO<sub>4</sub> nanotubes deliver a specific capacity of 550 mAh  $g^{-1}$ . Crystallographic parameters of lithium molybdates are listed in **Table 16**.

Compound	Structure	Cell parameters (A)		
		$a/\alpha$	$b/\beta$	$C/\gamma$
$Li1.3Mo2O4$	monoclinic	10.666	2.885/99.5	4.953
$\beta$ -Li <sub>2</sub> MoO <sub>3</sub>	monoclinic	4.990	$8.642/99.5^{\circ}$	10.092

**Table 16**. Crystallographic parameters of lithiated Mo oxides.

#### 6.9. MoO<sup>2</sup> anodes for SIBs

The application of MoO<sub>2</sub> as anode material for SIBs has made rapid progress, and the energy density has been improved greatly [571,572]. Several approaches have been conducted to improve the electrochemical performance of  $MoO<sub>2</sub>$  anodes (i.e., capacity retention, long life cycling) including core-shell composite [573,574], carbon-based hybrid [536,575-577], and element doping [578]. Bao et al. [575] fabricated nanosized  $MoO<sub>2</sub>$  on 3D porous carbon using a simple hydrothermal method with subsequent calcination. The porous  $MoO<sub>2</sub>/C$  composite

(with surface area of 171.6  $m^2 g^{-1}$ ) exhibits the specific charge capacity of ~463 mAh  $g^{-1}$  in the first cycle at 0.1 A  $g^{-1}$ . A reversible capacity of ~367 mAh  $g^{-1}$  can be maintained after 200 cycles. Jiang et al. [577] prepared a  $MoO<sub>2</sub>/C$  nanocomposite using a simple hydrothermal method. This composite delivers a charge capacity of  $557 \text{ mA} \text{h} \text{g}^{-1}$  at 0.1C in the first cycle and has a good cycling stability for Na storage. Cui et al. [579] synthesized  $MoO<sub>2</sub>@C$  nanoflowers through a grinding method followed by an annealing process. The  $MoO<sub>2</sub>@C$  hybrid was tested at 0.1 A  $g^{-1}$  and showed a reversible charge capacity of 172 mAh  $g^{-1}$ . In the long-term cycling test (1 A  $g^{-1}$ , 1000 cycles), a reversible charge capacity of 166 mAh  $g^{-1}$  remained. He et al. [580] reported a MoO2-C composite with MoO<sup>2</sup> nanosheets embedded in carbon matrix. The charge capacity vs. Na<sup>+</sup>/Na after 100 cycles is ~367 mAh  $g^{-1}$ . Zhao et al. [581] synthesized a MoO<sub>2</sub> nanoparticles anchored graphene oxide, and the material exhibited a reversible charge capacity of 345 mAh  $g^{-1}$  after 100 cycles at 0.1 A  $g^{-1}$ . MoO<sub>2</sub> combined carbon dots (CDs) with nanoparticle stacking structure were fabricated via a facile hydrothermal route [509]. The MoO<sub>2</sub>/CDs granules (50 nm in size) exhibit a discharge capacity of 236 mAh  $g^{-1}$  at 0.5 A  $g^{-1}$ after 300 cycles in 1 mol  $L^{-1}$  NaClO<sub>4</sub> in ethylene carbonate (EC) and dimethyl carbonate (DMC) (1:1 in volume) as SIB electrolyte. MoO<sub>2</sub> nanoparticles ( $\approx$ 100 nm) anchored on graphene oxide (MoO2/GO) were fabricated as SIB for anodes. The discharge gravimetric capacity of the MoO<sub>2</sub>/GO hybrid is stabilized at 345 mAh  $g^{-1}$  (volumetric capacity  $\approx$ 1656 mAh cm<sup>-3</sup>) at the current density of 100 mA  $g^{-1}$  after 100 cycles. After 1000 cycles, the capacity loss is only 19% and the electrode is still able to deliver 276 mAh  $g^{-1}$  [536]. Valdez et al. [582] prepared MoO<sub>2</sub>/C composite fiber anodes using precursors solutions with different concentrations of MoO2/PAN, which were force-spun into microfibrous mats and subsequently stabilized in air at 280ºC and calcined at 700 $^{\circ}$ C under Ar atmosphere. The MoO<sub>2</sub>/C composite exhibits a discharge capacity of 300-400 mAhg<sup>-1</sup> after 100 cycles at 100 mAg<sup>-1</sup>. Recently, Zhang et al. proposed the used of 2 wt.% vinylene carbonate (VC) additive to improve the cyclic stability of  $MoO<sub>2</sub>$  microspheres anode for SIBs. After 1000 cycles at 300 mA  $g^{-1}$  current rate, the MoO<sub>2</sub> electrode exhibits a capacity retention of 96.5%  $(\sim 110 \text{ mA} \text{h s}^{-1})$  [583].

## **7. Concluding remarks**

This review article shed light on the promising design strategies of molybdenum oxides for high kinetic energy storage. The technology of the molybdenum oxides used in energy storage devices from the 1980s to the recent progress was discussed. Due to their low molecular weight and their favourable electrochemical and solid-state properties,  $MoO<sub>v</sub>$  compounds appeared attractive as cathode and anode materials in electrochemical energy storage systems. Three

major classes of Mo oxides, which exhibit various structural properties, have be distinguished: (i) anhydrous Mo oxides  $(MoO<sub>3</sub>, Mo<sub>n</sub>O<sub>3n-1</sub>, Mo<sub>2</sub>)$ , (ii) molybdenum-oxide hydrates (MoO3∙*n*H2O), and (iii) nanostructured MoOy. A comparison between the electrochemical features of the molybdenum-oxide hydrates with different degrees of dehydration and their anhydrous products shows that oxide-hydrates materials have a good cyclability with a discharge capacity of 160 mAh  $g^{-1}$  at the 100<sup>th</sup> cycle. This value was greater than that of anhydrous MoO3. In a series of MoO3∙*n*H2O including sol-gel compounds, the main factor affecting the performance is expected to be the water content, which influencing the electrical conductivity and structural properties. Thermodynamics and kinetics of lithium-ion insertion are functions of the structural arrangement of the  $MoO<sub>6</sub>$  octahedra in the host lattice. Diffusion coefficients can be modelled using either a simple model for site occupancy or an ideal lattice gas model. A raw MoO<sup>3</sup> powder prepared by heat-treatment of molybdic acid in proper conditions exhibits the highest conductivity and displays the best electrochemical features for Li-ion insertion. Despite the attractive energy and power densities of the molybdenum oxides, these cathode materials belong to the class of 3-volt systems. The new development of nanostructured oxygen deficient  $MoO<sub>v</sub>$ , and  $MoO<sub>v</sub>$ -based composites as well, are promising electrodes materials for electrochemical energy storage devices due to their enhanced electronic conductivity, particularly supercapacitors and microbatteries, because they can instantaneously provide a high energy density with long cycle life.

Existing LIBs utilise graphite as anode with moderate theoretical capacity  $(372 \text{ mA} \text{h} \text{ g}^{-1})$ and limited fast-charging capabilities. Hence, the attractive electrochemical performances of the  $MoO<sub>3</sub>$  and  $MoO<sub>2</sub>$  composite networks benefit from the synergistic effects of the unique nanoarchitectures and the integrity of these anode materials. Monodispersed  $MoO<sub>y</sub>$  nanocrystals encapsulated in carbon matrix not only provide interfacial storage but also improve the transport kinetics of electrons and lithium ions. Extensive efforts have been made to overcome the defect of capacity degradation in  $MoO<sub>2</sub>$  anode, and the preparation of  $MoO<sub>2</sub>/carbon$  composite materials has proven to be an effective way. The mesoporous nature of the  $MoO<sub>2</sub>/C$ nanocomposite and the thin-layer carbon coating are believed to contribute to the enhanced electrochemical performance, which not only feature the efficient four-electron conversion reaction for Li<sup>+</sup> storage, but also effectively tolerate volume expansion during the cycling. In this review, we have also seen that with many distinctive properties, stoichiometric  $MoO<sub>3</sub>$  and MoO<sub>2</sub> can sustain high-rate capability as high-power battery electrodes to power electric vehicles. It is worthy to note that  $MO<sub>3</sub>$  is a material used in various devices such as gas sensors, catalysts, recording materials, electrochromic devices, and so on.

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