<table>
<thead>
<tr>
<th><strong>Title</strong></th>
<th>Single-crystal micro/nanostructures and thin films of lamellar molybdenum oxide by solid-state pyrolysis of organometallic derivatives of a cyclotriphosphazene</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Author(s)</strong></td>
<td>Díaz, Carlos; Lavayen, Vladimir; O'Dwyer, Colm</td>
</tr>
<tr>
<td><strong>Publication date</strong></td>
<td>2010-05-15</td>
</tr>
<tr>
<td><strong>Type of publication</strong></td>
<td>Article (peer-reviewed)</td>
</tr>
<tr>
<td><strong>Link to publisher's version</strong></td>
<td><a href="http://dx.doi.org/10.1016/j.jssc.2010.05.006">http://dx.doi.org/10.1016/j.jssc.2010.05.006</a></td>
</tr>
<tr>
<td>Access to the full text of the published version may require a subscription.</td>
<td></td>
</tr>
<tr>
<td><strong>Rights</strong></td>
<td>© 2010 Elsevier Inc. This manuscript version is made available under the CC-BY-NC-ND 4.0 license <a href="http://creativecommons.org/licenses/by-nc-nd/4.0/">http://creativecommons.org/licenses/by-nc-nd/4.0/</a></td>
</tr>
<tr>
<td><strong>Item downloaded from</strong></td>
<td><a href="http://hdl.handle.net/10468/2818">http://hdl.handle.net/10468/2818</a></td>
</tr>
</tbody>
</table>

Downloaded on 2019-08-03T02:19:20Z
Single-Crystal Micro/Nanostructures and Thin Films of Lamellar Molybdenum Oxide by Solid-State Pyrolysis of Organometallic Derivatives of a Cyclotriphosphazene

Carlos Díaz,a,1 Vladimir Lavayen,b and Colm O'Dwyer c,2

a Departamento de Química, Facultad de Ciencias, Universidad de Chile, Casilla 653, Santiago, Chile
b Departamento de Física, Universidad Técnica Federico Santa María, Valparaiso 2390123, Chile
c Department of Physics, and Materials & Surface Science Institute, University of Limerick, Limerick, Ireland

Abstract

The solid-state pyrolysis of organometallic derivatives of a cyclotriphosphazene is demonstrated to be a new, simple and versatile solid-sate templating method for obtaining single crystal micro- and nanocrystals of transition and valve metal oxides. The technique, when applied to Mo-containing organometallics N3P3[OC6H4CH2CN⋅Mo(CO)5]6 and N3P3[OC6H4CH2CN⋅Mo(CO)4 py]6, results in standalone and surface-deposited lamellar MoO3 single crystals, as determined by electron and atomic force microscopies and X-ray diffraction. The size and morphology of the resulting crystals can be tuned by the composition of the precursor. X-ray photoelectron and infrared spectroscopies indicate that the deposition of highly lamellar MoO3 directly on an oxidized (400 nm SiO2) surface or (100) single crystal silicon surfaces yields a layered uniphasic single-crystal film formed by cluster diffusion on the surface during pyrolysis of the metal carbonyl derivatives. For MoO3 in its layered form, this provides a new route to an important intercalation material for high energy density battery materials.

1 cdiaz@uchile.cl
2 colm.odwyer@ul.ie
Introduction

The method of solid-state pyrolysis of organophosphazene/organometallic as a synthetic templating method represents a new approach to the realization of micro- and nanocrystals of transition and valve metals and their respective oxides. Of these, MoO$_3$ is an extremely versatile material from many standpoints$^{[1]}$. This material exhibits useful properties that have already been applied to sensing, electrochromic coating materials, layered host materials for solid-state thin film battery cathodes,$^{[2,3]}$ solid-state lubricants$^{[4]}$ exhibiting load-independent friction$^{[5]}$, and even superconductivity$^{[6]}$ and thermoelectrics$^{[7]}$. A recent review$^{[8]}$ has poignantly highlighted the importance of micro- and nanostructured MoO$_3$ and related substoichiometric molybdenum oxides for charge storage/intercalation and particularly for electrochromics, which has been studied for a wide range of disordered partially crystalline or polycrystalline materials including amorphous MoO$_3$ films. Only brief studies of nanocrystalline MoO$_3$ materials for electrochromic applications have been reported to date$^{[9]}$ limited by the availability of single crystal MoO$_3$ synthesis with size and crystal structure control and suitable surface deposition methodologies.

The recent interest in oxides, and in particular nano-oxides$^{[10]}$, as anodes for HEV lithium batteries is driven by the desire for an anode with a slightly more positive insertion voltage with respect to Li$^+$/Li to minimize any risks of high-surface-area Li plating while charging at high rates. Thus the ideal negative electrode for forthcoming vehicular applications that operate at higher potential than graphite are currently being investigated for future anode technologies$^{[11]}$. MoO$_3$ and methods for high yield and purity are under renewed scrutiny since it (and other transition metal oxides)$^{[12,13]}$ operates at higher potentials relative to Li$^+$/Li during intercalation/insertion reactions. The Li$^+$/intercalation cycling properties of $\alpha$-MoO$_3$ is proposed to be markedly improved upon nanostructuring, by minimizing the stability-reducing effect cause by unit cell volume increases during intercalation$^{[13]}$. The orthorhombic structure of $\alpha$-MoO$_3$ is characterized by narrow channels of ~2 nm.
Intercalation of lithium ions into a crystal of this form would be aided if the lithium ions can preferentially penetrate spaces between planes bound by van der Waals interactions.

Nanostructured MoO$_3$ in particular, is an attractive material due to its unusual chemistry and electrochromism produced by multiple valence states. Some metal oxides can spontaneously spread over supports forming self-assembled monolayers, which has been applied to heterogeneous catalysis$^{[14]}$, but nanostructured MoO$_3$ can exhibit greater activity, and the possibility of forming stable coordination environments could lead to improved selective oxidation$^{[15]}$ and isomerisation of hydrocarbons, and suitable supported intercalation cathodes. Lamellar or layered $\alpha$-MoO$_3$ in its thermodynamically stable orthorhombic form, based on the bilayered arrangement of corner-shared MoO$_6$ octahedra held together by covalent forces, is also used as an image rotation calibration standard for transmission electron microscopy, but currently only accessible in bulk form with a lamellar structure by tedious synthetic routes.

Despite its importance, success in obtaining nanoscale single crystal anhydrous MoO$_3$, compared to other metal oxides, is relatively rare.$^{[16,17]}$ Furthermore, the products are commonly obtained as irregularly agglomerated powders in which the polycrystalline domain size is typically 10-30 nm. Recent reports have shown the nanometer control of sheet thickness for MoO$_3$ nanobelts and nanoribbons, citing the importance of its properties as a lamellar material.$^{[18-20]}$ Ultimately, the possibility of synthesizing standalone micro- and nanocrystals of lamellar $\alpha$-MoO$_3$, as both standalone single crystal of micro- and nanoscale sizes in addition to monocrystalline deposits on amorphous and single crystal surfaces, is a considerable advance. Only localized islands are usually formed on related substrates making their removal and eventual application quite cumbersome. In addition, the van der Waals bound interatomic spacings are readily available for electrochemical lithium ion storage and release.$^{[21]}$
Our recent reports employing polyphosphazene as solid-state templates for the preparation of metallic nanostructured materials \[22\] inspired the search for a suitable methodology to produce nanostructured MoO\(_3\) with micro- and nanoscale morphological analogues of its bulk counterpart. This solid-state pyrolysis of organophosphazene/organometallic (SSPO) template method is outlined in Scheme 1. This methodology represents a new synthetic route for the realization of lamellar MoO\(_3\) micro- and nanocrystals with morphologies similar to multi-functional low-dimensional, layered MoO\(_3\).

Although several metallic nanostructured materials can be obtained by other synthetic methods, the relatively long and tedious synthetic procedures are major drawbacks to obtaining the precursors, and generally result in low yields. This is especially true also for the SSPO method using polymers as precursors. Some of these disadvantages are resolved by using organometallic derivatives of cyclophosphazenes as alternative precursors. These organometallic derivatives can be easily prepared by functionalization of N\(_3\)P\(_3\)Cl\(_6\) and subsequent coordination of organometallic fragments,\[23\] giving routinely high yields as either standalone crystalline product or as deposited films. Preliminary results with the trimer N\(_3\)P\(_3\)(OC\(_6\)H\(_5\))\(_5\)(OC\(_6\)H\(_4\)CH\(_2\)CN[RU])PF\(_6\) indicate similar results in comparison with the respective polymer.\[22c\]

Here, we report the first successful pyrolysis of the Mo-containing N\(_3\)P\(_3\)[OC\(_6\)H\(_4\)CH\(_2\)CN-Mo(CO)\(_5\)]\(_6\) (I) and N\(_3\)P\(_3\)[OC\(_6\)H\(_4\)CH\(_2\)CN-Mo(CO)\(_4\)py]\(_6\) (II) organometallics, and the characterization of their \(\alpha\)-MoO\(_3\) micro- and nanocrystalline products. As will be outlined, pyrolysis of (I) and (II) in air at 800 °C produces crystalline orthorhombic \(\alpha\)-MoO\(_3\) single crystals, determined using both electron and ion microscopy in addition to diffraction-based techniques, atomic force microscopy and surface spectroscopies. The size of the crystallites depends on the composition of the precursor and this uniphasic material can be deposited/prepared uniformly on either single crystal substrates or amorphous
supports. The probable mechanism involving the phosphazene as a solid-state template in the formation of MoO₃ is also discussed.

**Experimental**

The precursors (I) and (II) were prepared using a previously reported methodology.[23] Briefly, the trimer N₃P₃[OC₆H₄CH₂CN]₆[23a] and the organometallic compounds Mo(CO)₅THF (generated from UV-irradiation of Mo(CO)₆ and THF) and Mo(CO)₄py₂ in stoichiometric quantities were stirred in THF and chloroform, respectively. The resulting solids were filtered and washed with a n-hexane/ether mixture.

(I) 0.5130 g, 1.94 mmol, of Mo(CO)₆ in 30 ml methanol was irradiated for 45 min. The yellow solution was transferred to a Schlenk and 0.299g, 0.32 mmol of N₃P₃[OC₆H₄CH₂CN]₆[23] was added and stirred for 24 h. After this step, the solvent was removed at reduced pressure and the solid washed with diethylether and n-hexane. Yield: 0.24 g, 32%. Elem. anal. Calcd for C₇₈H₃₆N₉O₃₆P₃Mo₆ (found): C, 40.01 (39.3); H, 1.53 (1.49); N, 5.38 (5.29). ¹H NMR (ppm, CDCl₃): 7.185 m, 7.01 M, [-C₆H₄-]. 1.28 s, 2H [-CH₂-]. ³¹P NMR (ppm, acetone-d₆) 8.75 s, ppm. IR(KBr, cm⁻¹): 2963, 2876, 2140 ν(CN), 2065, 1940 ν(CO), 1507, 1485, 1463, 1382, 1268, 1206, 1183, 1165 ν(PN), 957, 886, 850.

(II) To N₃P₃[OC₆H₄CH₂CN]₆, 0.0506g , 0.546 mmol in 80 ml chloroform 1.63 mmol Mo(CO)₄Py₂, 0.61 g, 1.63 mmol was added also dissolved in the same solvent (40 ml) and stirred at room temperature for 24 h. The solid was then decanted and the supernatant was extracted. The solid was washed twice with chloroform and dried under vacuum. Yield 0.08g, 63%. Elem. anal. Calcd for C₁₅₂H₆₀N₃₀P₃Mo₆ (found): C, 46.37 (45.5); H, 2.27 (2.3); N, 7.95 (7.85). ¹H NMR (ppm, CDCl₃): 8.29 m, [C₅H₄N]; 7.2 m, 7.1 m, [-C₆H₄-]. 1.3 s, 2H [-CH₂-]. ³¹P NMR (ppm, acetone-d₆) 8.93 s. IR(KBr, cm⁻¹): 2948, 2840, 2139 ν(CN), 2065, 1940 ν(CO), 1485, 1463, 1382, 1268, 1206, 1183, 1165 ν(PN), 957, 886, 850.
\( \nu(CO), 1660, 1638; 1585 \nu[CC, C_5H_4N], 1461, 1433, 1394; 1206, 1162 \nu(PN), 1029, 896, 842. \)

**Pyrolysis of the precursors:** The pyrolysis experiments were carried out by pouring a weighed portion (0.05-0.15 g) of the organometallic trimer into aluminum oxide boats placed in a tubular furnace (Lindberg/Blue Oven model STF55346C-1) under a flow of either air or nitrogen, heated from 25 to 300 °C and then to 800 °C, and annealed for 2 h. The heating rate was 10 °C min\(^{-1}\) under an air/N\(_2\) flow rate of 200 mL min\(^{-1}\).

**Characterization of the pyrolytic products:** X-ray diffraction (XRD) was conducted at room temperature on a Siemens D-5000 diffractometer with 0-20 geometry. The XRD data was collected using Cu-K\(\alpha\) radiation (40 kV, 30 mA). Scanning electron microscopy (SEM) and energy dispersive X-ray analysis (EDX) were acquired with a JEOL 5410 SEM with a NORAN Instrument micro-probe transmission microscope. Transmission electron microscopy (TEM) was carried out on a JEOL 2011 TEM operating at 200 kV. The finely powered samples were dispersed in \(n\)-hexane and dropped on a conventional carbon-coated copper grid and dried under a lamp. Atomic force microscopy (AFM) measurements were performed using a Digital Instruments Nanoscope IIIa and Veeco Enviroscope using both contact and tapping mode scans. Suitable depositions were obtained by dissolving the trimers (I), and (II) in dichloromethane, dropping on either an oxidized (400 nm SiO\(_2\)) surface or (100) single crystal silicon surfaces, followed by evaporation of the solvent at room temperature. Subsequent pyrolysis was conducted at 800 °C to remove surface bound water. AFM images were acquired with in-situ substrate heating to 110 °C. Optical microscopy imaging was performed using a Zeiss A1 Vario Axioscope. Fourier transform infrared (FTIR) characterization was performed in attenuated total reflection mode using a Perkin-Elmer Spectrum 100 spectrometer. X-ray photoelectron spectroscopy (XPS) was conducted on...
deposits from (I) and (II) on Si and SiO2 surfaces in normal emission geometry. A VG CLAM 2 spectrometer with an Al Kα X-ray source (1486.6 eV), operating at 250 W, were used several centimeters from the samples. The spectra were normalized to the total electron yield to correct for small differences in sample positions and X-ray source intensities. The resulting spectra were fitted by using a Shirley-type background and symmetric Voigt functions with variable Gauss and Lorentz contributions.

**Results and Discussion**

The new compounds (I) and (II) are brown or yellow solids, soluble in common organic solvents. IR spectra of (I) clearly exhibit the presence of the Mo(CO)5 moiety. Absorption at 2056 cm\(^{-1}\) and a broad band between 1946 and 1908 cm\(^{-1}\), corresponding to \(\nu(CO)\) vibrations typical of M(CO)5 were observed.\(^{[22a]}\) For (II), the respective \(\nu(CO)\) vibrations are centered at 1985 cm\(^{-1}\) and a weak vibration at 2025 cm\(^{-1}\), were also observed. The presence of the phosphazene is clearly indicated by the typical \(\nu(CN)\)\(^{[23]}\) bands: at 2140 cm\(^{-1}\) for (I) and 2139 cm\(^{-1}\) for (II) of the ligands, as well as by the 1206 cm\(^{-1}\), 1183 cm\(^{-1}\), and 1165 cm\(^{-1}\) bands for (I) and 1206 cm\(^{-1}\), 1162 cm\(^{-1}\) for (II) corresponding to the \(\nu(PN)\) band of the ring.\(^{[23,24]}\) In the case of the pyridine complex (II) the emerging of a band at 1638 cm\(^{-1}\) typical of coordination of pyridine ligand\(^{[22a]}\), was also observed. In addition, NMR spectra corroborate the presence of the ligand i.e. the signal corresponding to the C6H4 and CH2 groups were observed as expected (see Experimental section). The total coordination of the Mo(CO)5 or Mo(CO)4Py fragments in the six sites of the phosphazene ring is clearly established by the solitary signal in the \(^{31}\)P NMR spectra of both (I) and (II)\(^{[22c]}\).

Pyrolysis of the trimers (I) and (II) in air at 800 °C results in a white crystalline solid. The X-ray diffraction (XRD) patterns of the pyrolytic product from (I) and (II), shown in Figures 1a,b, exhibit excellent correspondence with the standard XRD data for orthorhombic
α-MoO₃ (JCPDS 00-005-0508). The pattern in Fig. 1a was taken from large scale pyrolysis of the organometallic and shows the pure α-MoO₃ phase in addition to small quantities of residual P₄O₇ which acts as a solid-stage stabilizer.[22] The inset to Fig. 1a shows an expanded view of the region 14° < 2θ < 20°, indexed to monoclinic pyrophosphate (JCPDS 01-072-0880). The (020), (040) and (060) peaks (see Fig. 1a,b) are predominantly more intense than those from (110), (021), (130), i.e. (hk0) reflections, indicating a two-dimensional growth of the crystallites along the {010} basal planes, as shown schematically by the extended unit cell in Fig. 1b (inset).[2,25] For the as-obtained pyrolytic product from precursor (II), similar XRD patterns to those for (I) are observed; the relative predominance of the (0k0) reflections is more pronounced and the three-dimensional crystal diffraction peaks have a reduced relative intensity. A dominant presence of (hk0) would indicate the presence of monoclinic α-MoO₃, but is never observed. Additionally, the presence of metastable β-MoO₃ is not detectable and both precursors are almost entirely orthorhombic α-MoO₃, either as crystalline powder or deposited films.

Both precursors result in micro- and nanoscale crystals exhibiting a high degree of crystallinity; the pronounced (0k0) reflection intensities confirm a layered structure packed in the direction of its crystallographic b-axis, parallel to the {010} plane as shown in crystal structure inset to Fig. 1b. Within each MoO₆ octahedron, one oxygen atom is essentially unshared forming a terminal Mo=O bond. The layers themselves are in a staggered arrangement along the c axis and are bound by van der Waals’ forces resulting in individual layers of this material forming as a turbostratic arrangement. Analysis of the degree of grain orientation and lamellar stacking was determined from XRD data using the Lotgering method[26] using the intensities of the (0k0) and (hk0) reflections for the synthesized samples compared to pure MoO₃. The lamellar orientation degree, F, is given by

$$F = \frac{(P - P_0)}{(1 - P_0)}; P = \frac{\sum (0k0)}{\sum (hk0)} I$$
where $P$ is the ratio of the sums of the $(0k0)$ and $(hk0)$ reflection intensities, respectively, and $P_0$ represents similar relative intensities from pure MoO$_3$. The products from precursor (I) and (II) have lamellar orientation degrees (Lotgering factors) of 0.78 and 0.92 respectively, indicating pronounced turbostratic layering of the products, which are uniphasic, orthorhombic MoO$_3$ micro- and nanocrystals.

The XRD findings are in agreement with SEM imaging of the products, shown in Fig. 2 clearly showing the typical rectangular morphology of the resulting crystals. Figure 2a shows the typical morphology of the surface of a MoO$_3$ crystal formed by pyrolysis of (I) on silicon. The surface, as in all samples in this work, is coated with smaller nanocrystals of MoO$_3$ of similar morphology and crystallinity. Additionally, the EDAX spectrum of all crystallites (from (I) and (II)) exhibited only peaks corresponding to the presence of Mo and O atoms; the spectrum in Fig. 2b is from the large crystal in Fig. 2c where contributions from Au arise from the coating for SEM imaging. The product from precursor (II) is noted to be very similar to that from (I), except that the resulting crystal structures vary in size from micro- to nanocrystals. These pure MoO$_3$ single crystals are observed with a range of sizes and aspect ratios, all with a characteristic high aspect ratio rectangular morphology. Thus the resulting crystals are distinct and all crystals exhibit identical crystallographic phase and space group, and apart from absolute size and aspect ratio, are self-similar. The shapes of these crystals can be classified into 4 general morphological groups: nano and micro rods, flakes, monoliths and lamina, all of which have length to width aspect ratios typically between 2 and 50. Each of these crystals comprises crystallographically stacked layers of MoO$_3$, corroborating the XRD findings. Some clear examples of laminar MoO$_3$ crystals in SEM images are shown in Fig. 2d.

AFM and TEM images were also acquired of individual flakes of the lamellar products from pyrolysis of (I) and (II), and are shown in Fig. 3. After pyrolysis, the crystalline flakes have typical dimensions in the range 200-1000 nm for (I) but 150-300 $\times$ 300-700 nm
for (II), similar to those in Fig. 2. All flakes are taken from two sources: the surfaces of larger crystals and also from film deposits on Si by mechanical exfoliation. Kalantar-zadeh *et al.*[20] have suggested that the mechanical exfoliation of MoO$_3$ sheets delaminates at their weakest bond points, where they are held together by weak van der Waals forces. Microscale thickness flakes exhibited characteristics layering at their edges, as shown in the high resolution contact AFM image in Fig. 3a. The surface profile shown in Fig. 3b exhibits typical layering, where the repeat step height is consistently in the range 2.7 – 3.1 nm. This value is close to twice the magnitude of $b$ lattice parameter for $\alpha$-MoO$_3$ ($b = 1.38$ nm) based on the structure depicted in Fig. 1b. In some cases, step heights of roughly half this value (similar to $b$) are observed (see * in Fig. 3b), but is not a common observation.

TEM analysis in Figs. 3c and d show the individual crystals and also the exfoliated flakes. The corresponding electron diffraction patterns and FFT can be indexed to non-hydrated orthorhombic MoO$_3$ with reconstructed lattice parameters of $a_0 = 0.395$ nm, $b_0 = 1.383$ nm, $c_0 = 0.369$ nm; Space Group: $Pnma$. The HRTEM image in Fig. 3d acquired perpendicular to the growth direction shows MoO$_3$ lattice fringes spaced by 0.685 nm, corresponding to the spacing between the (020) planes. The flakes for TEM measurement actually have thicknesses of the order of 2-6 molecular layers of MoO$_3$ with excellent layer matching. Predominant intensity (020) reflections are observed typical of lamellar (010) stacking along the $c$-axis, in agreement with indications of lamellar two-dimensional growth from XRD data. The crystalline structures obtained as product from precursor (II) are routinely observed to be larger in length and width than from (I), according to the SEM and TEM data, and correspondingly, exhibit higher relative $(0k0)$ reflections in their XRD spectra (Fig. 1b). The planar or lamellar topology of these turbostratic orthorhombic crystals, which consist of bilayers parallel to the (010) plane weakly bound by covalent interactions, would allow exfoliation of nanoscale flake crystals from microscale single-crystal structures, which is what is usually observed experimentally.
In order to apply these nanostructured lamellar materials to surfaces or supports, synthesis and subsequent pyrolysis on suitable substrates was carried out to characterize the as-deposited morphology and investigate the mechanism of surface spreading and long-range uniformity in crystallinity and coverage/thickness. Suitable depositions were obtained by dissolving the trimers (I) and (II) in dichloromethane, dropping on either an oxidized (400 nm SiO₂) surface or (100) single crystal silicon surfaces, followed by evaporation of the solvent at room temperature. Subsequent pyrolysis was conducted at 800 °C. Surprisingly few AFM studies of MoO₃ have been reported in the literature [4,16] for uniform MoO₃ deposition on surfaces directly and this is the first report of single crystal deposition of this material on monocrystalline and amorphous surfaces. Scanning probe and scanning tunneling microscopy (SPM and STM) techniques have been principally used for the atomic scale study of MoO₃ island growth on both Au{111} and MoS₂ along with the friction-reducing benefits of a Mo/MoS₂/MoO₃/MoS₂ sandwich layer system. [4] Atomic force microscopy (AFM) imaging of pyrolyzed substrates, shown in Fig. 4, reveals a surface morphological dependence of the coverage on the composition of the precursor: from (I) craters of typically 10-20 nm in depth and two principal diameters of either ~500 nm or ~60-100 nm are routinely observed (Fig. 4a-c) whereas from (II) encrustations or islands of 23-50 nm in height and diameters of 0.6-1.2 μm are formed (Fig. 4e-g), on surfaces that are otherwise covered with a relatively uniform layer of MoO₃ with rms roughness values of 2.6-4 nm for both surfaces over 100 μm² (including the surface features). High resolution contact AFM images shown for precursor (I) in Fig. 4d and precursor (II) in Fig. 4h show that the deposition using precursor (II) results in an extremely smooth surface. This is also the deposition that gives the lowest density of crater formation, but also a higher density of excess MoO₃ in the form of islands on the surface. Although both islands and craters are found on the surface deposits from both precursors, the general trend is a predominance of craters from (I) and islands of MoO₃ from pyrolysis of (II)
whether deposited on Si or SiO₂. As the pyrolysis was conducted in air, the presence of O₂ promotes the spreading of MoO₃.[27]

The formation of islands and craters in pyrolyzed precursor functionalized surfaces is reminiscent of the ramified-cluster-diffusion mechanism [16,28], which can explain the process of spreading on the nanoscale in most studies thus far, i.e. bulk MoO₃ spread in a ramified manner in the form of agglomerated cluster islands. The overall uniformity of the deposit (from either precursor) depends on the nature of that precursor and the mechanism of pyrolysis. Investigation of the surface by optical microscopy was also performed over larger deposition areas. Deposits on SiO₂ were examined since the large refractive index contrast afforded by the SiO₂ allows imaging of very thin several-molecule-thick layers in a similar manner to optical microscopy of graphene.[29] Figure 5a shows the typical MoO₃ film deposition prior to full coverage. The red regions are MoO₃ and the deposit is interspersed with craters. The blue colour comes from the SiO₂ under white light. The boxed region in Fig. 5a shows how the quantity or thickness of MoO₃ varies along the surface. In this case, it is always observed that the higher the density of craters, the lower the film thickness in accord with a cluster diffusion spreading of a finite local amount of product. For greater quantities of precursor, we observe the same type of film coverage as was seen by AFM in Fig. 4e, i.e. a uniform deposit with some evidence for cratering (but the craters do not extend down to the SiO₂ layer) and the presence of MoO₃ islands (shown in red). The contrast is less effective in Fig. 5b compared to Fig. 5a due solely to the reduced depth of the craters. Since the deposit can transfer to the Si or SiO₂ in single-crystal form, optical imaging of individual and several monolayers was possible. Figure 5c shows an optical micrograph of the coated SiO₂ surface, a further single layer on top of this, marked by (1), and also a double layer (2), which appears as the brightest, green region. Similar to Fig. 3 for single crystals, the surface deposit is also layered which was confirmed by AFM.
As with polyphosphazenes,[22,23] insight into the possible mechanism of the pyrolysis was obtained from thermogravimetric analysis and differential scanning calorimetric (TGA/DSC) studies obtained during the pyrolysis of precursors (I) and (II) in both air and nitrogen atmospheres, which are shown in Fig. 6. The deposits were heated at a rate of 10 °C min⁻¹ from ambient (25 to 800 °C under a constant air flow of 200 mL min⁻¹). Conditions similar to those of the preparative pyrolysis experiments were used. From samples of precursor (I), the TGA curves shown in Fig. 6a (in air and nitrogen) indicate a gradual loss of CO₂ and H₂O from 150 °C to 350 °C which corresponds to oxidation of the carbon atoms of the organic matter. The data indicate that the complete oxidation of carbon is much more efficient in air, even though under N₂ significant oxidation of carbon as CO₂ and H₂O still does occur. This is crucial for ensuring a pristine MoO₃ surface during pyrolysis, for example, and not routinely found by other general approaches.[30] This agrees with the intense exothermic peaks in the DSC response shown in Fig. 6b, at 284 °C and 494 °C, which are normally observed after combustion of the organic part of organometallic compounds[31,32] and crystallization of the MoO₃, which is also represented schematically in Fig. 6c.

This loss of the organic matter produces vacancies in the previously formed cyclomatrix which allows agglomeration of the metallic particles followed by growth of the nanoclustered structures. The organometallic fragments Mo(CO)₅ and Mo(CO)₄py linked to the phosphazene precursors (I) and (II) respectively, as determined from previous studies,[33] loose carbon monoxide upon heating leading to Mo centers that migrate toward the vacancies generated in the cyclomatrix. Similarly, the phosphazenic phosphorus and the nitrogen polymeric chain evolve as the respective oxides upon oxidation, and do so as single crystals. As noted during the pyrolysis of organometallic derivatives of polyphosphazenes, the remaining phosphorus oxide, as shown by XRD, acts as a structural stabilizer.

Infrared vibrational and X-ray photoelectron spectroscopies were also conducted on few-monolayer coverage MoO₃ deposits after pyrolysis of (II). The FTIR spectrum in Fig. 7a
exhibits three distinct vibrations. The secondary electron image from a focused Ga\(^{+}\) beam is shown in Fig. 7b and the corresponding AFM image in Fig. 7c. The bridging Mo–O–Mo stretching vibration observed at 809 cm\(^{-1}\) is characteristic of most molybdenum oxides, where the stoichiometric ratio of oxygen is reduced, e.g. for Mo\(^{4+}\) as MoO\(_2\) and Mo\(^{6+}\) as MoO\(_3\). The \(\alpha\)-Mo–O vibration at 855 cm\(^{-1}\) is distinctly related to \(\alpha\)-MoO\(_3\) which is most typically found in several crystal structures. Corroboration of the orthorhombic structure deduced from diffraction measurements is found with the observation of the terminal molybdyl bond vibration (M=O) measured at 981 cm\(^{-1}\); this stretching vibration is unique to \(\alpha\)-MoO\(_3\) in an orthorhombic structure. In contrast to the Mo–O–Mo and M=O vibrations, the \(\alpha\)-Mo–O vibrations are comparatively broadband. This is most likely linked to the varied terminal surface morphology from various crystal facets, since FIB and AFM imaging indicate sub-nanoscale features on the most pristine (smoothest) regions of deposit.

XPS measurements conducted at various take-off angles conclude that the deposited layers (and uppermost atomic layers) are pure MoO\(_3\), since the wavenumber limit of the FTIR spectrometer prevented determination of true lattice vibrational modes, rather the IR active stretching modes. The Mo3d, Mo3p, O1s and C 1s core-level peaks are shown in Fig. 8a-d, respectively. The Mo3d\(_{5/2}\) and Mo3d\(_{3/2}\) doublet shown in Fig. 8a are observed at binding energies of 232.5 eV and 235.6 eV, respectively, with the corresponding Mo3p\(_{3/2}\) and Mo3p\(_{1/2}\) measured at 399.2 eV and 416.6 eV (Fig. 8b). These doublets are consistent with Mo only in its hexavalent(VI) oxidation state as MoO\(_3\). For all samples studied from (I) or (II) on Si or SiO\(_2\), deconvolution of the spectra showed there was no appreciable shift in binding energy of Mo3d core level emission towards lower energy, confirming the complete stoichiometry of the product, i.e. we never observe sub-stoichiometric molybdenum oxides. The O1s peak, for pyrolytic products from (I) and (II) is observed only at 530.8 eV with no dominant overlapping contributions from other oxygen species.\([34]\) Examination of the C 1s
core-level spectrum confirms that there are no C-O bonds and that the detected carbon is adventitious. Quantification of the O 1s core-level spectrum in $\alpha$-MoO$_3$ formed by other methods often includes contributions from two other species of O. The presence of OH groups would give a shoulder peak at 531.8 eV (scaled to the data presented here) and another at 532.7 eV, associated with adsorbed water. The shoulder peak in the deconvoluted O 1s spectrum in Fig. 8c corresponds to adsorbed moisture, most probably from exposure to air prior to chamber loading as no indicative vibrations from hydrated MoO$_3$-$n$H$_2$O are observed in the IR data. Thus, dedicated surface studies of the deposited product confirms a self-similar crystal structure to the individual crystals formed during pyrolysis of the organometallic Mo-containing derivatives of cyclotriphosphazenes directly. This readily accessible technique is promising for the synthesis of important lamellar materials and their incorporation onto surfaces and supports.

Conclusions

The pyrolysis of organometallic Mo-containing derivatives of cyclotriphosphazenes is found to be a simple, versatile and useful method to grow monocrystalline $\alpha$-MoO$_3$. The products can be micro- and nanocrystals with a high degree of phase purity and laminarity, and also smooth, single crystal deposits (monolayers and thin films) of $\alpha$-MoO$_3$ on crystalline silicon and amorphous silicon dioxide surfaces by cluster diffusion during pyrolysis. The size and the morphology can be tailored by compositional control of the organometallic precursor for a given surface. Transmission electron microscopy and electron diffraction measurements confirm the nanoscale crystallinity observed in larger deposits by XRD where the laminar single crystals are orthorhombic $\alpha$-MoO$_3$. The same crystalline phase is found for layer deposits on either silicon or silicon dioxide after pyrolysis. AFM and ion-beam morphology measurements indicate a smooth film deposit which IR and XPS analysis confirm stoichiometric MoO$_3$ and successful combustion of the organic matter from the precursor.
The technique is shown to be effective in realizing stand-alone micro- and nanostructures of an important electrochromic and intercalation host materials (MoO$_3$), but can also be directly applied to the formation of other valve and transition metals, leading to a variety of micro- and nanostructured materials and their oxides through a very clean and quick synthetic methodology involving pyrolysis of metal-carbonyl derivatives.

Acknowledgements

Financial support from FONDECYT Grants No. 1085011, 1095135, 1090683, and 1090282 are acknowledged. V. L. acknowledges PBCT Grant ACT027. Part of this work was conducted under the framework of the INSPIRE programme, funded by the Irish Government's Programme for Research in Third Level Institutions, Cycle 4, National Development Plan 2007-2013.

References


**Figure Captions**

**Scheme 1.** Schematic representation of the solid-state pyrolysis of organophosphazene/organometallic (SSPO) template method.

**Figure 1.** X-Ray diffraction patterns of the pyrolytic products from (a) (I) and (b) (II). The patterns were indexed to JCPDS 00-001-0706 and JCPDS 00-005-0508 (stick pattern overlaid) for non-hydrated orthorhombic $\alpha$-MoO$_3$. The inset to (a) shows the region $14^\circ < 2\theta < 20^\circ$, indexed to monoclinic pyrophosphate (JCPDS 01-072-0880). The inset to (b) shows layered $\alpha$-MoO$_3$ unit cell within the repeating crystallographic unit of length $2b = 27.60$ Å.

**Figure 2.** (a) Scanning electron microscopy images of the pyrolytic products from (I) at beam voltages between 1 and 10 kV. (b) Energy dispersive X-ray analysis of the products showing their bi-elemental (Mo and O) composition. (c) Micro and nanoscale rods, flakes, monoliths and lamina of MoO$_3$ obtained after pyrolysis of (II). (d) Individual and stacked lamina of MoO$_3$ obtained from pyrolysis of (II).

**Figure 3.** (a) contact mode AFM image of crystallographic step features at the edge of a MoO$_3$ crystal from (I). (b) Line scan profile showing the repeat unit height of 2.7-3.1 nm. The * indicates a step of unusually low thickness. Transmission electron micrographs of the
pyrolytic products from (c) (I) and (d) (II) evidencing highly turbostratic flakes and the presence of rod-like lamellar monocrystals of MoO₃. The FFT pattern (inset (c)) taken from HRTEM images acquired along the [010] zone axis confirms single-crystal orthorhombic MoO₃, also evidence by the HRTEM image inset to (d).

**Figure 4.** Atomic force microscopy images of the products from (a-d) (I) and (e-h) (II) pyrolyzed on a silicon substrate. High resolution images from (I) in (d) and from (II) in (h) show very smooth surfaces. The corresponding line scans highlight the typical feature dimensions resulting from spreading during pyrolysis.

**Figure 5.** Optical microscopy of MoO₃ films at (a) low quantity coverage where cratering of the deposit is observed (b) smooth deposition for thicker deposits and (c) several monolayer thick lamina. In (a), the blue regions in craters come from the SiO₂ under white light, the red regions from MoO₃. Variations in MoO₃ thickness are shown in the boxed region. In (b) the red crystals are MoO₃. In (c) single and double layer MoO₃ lamina are marked (1) and (2) respectively.

**Figure 6.** (a) Differential scanning calorimetric analysis of the pyrolysis of the product from (I) in air and N₂. Pyrolysis in air exhibits intense exothermic peaks during organic combustion. Total weight = 4.000 mg. (b) Thermogravimetric analysis of the pyrolytic product from (I); 100% = 9.000 mg. (c) Pyrolysis condition and schematic of organometallic metal carbonyl to yield single crystal MoO₃.

**Figure 7.** (a) Fourier transform infra-red spectroscopy (absorption) of the pyrolytic product from (I) deposited on (100) silicon. The spectrum evidences the terminal molybdyl (M=O) vibration characteristic of orthorhombic α-MoO₃ and this is highlighted by the hatched
region. (b) Secondary electron image from a focused Ga⁺ beam scan of the morphology of the deposited product after pyrolysis. (c) AFM image of the surface morphology of the post-pyrolytic product of (I) acquired in contact mode in vacuum after in-situ heating to 110 °C to remove surface bound water.

**Figure 8.** High resolution XPS spectra acquired from the deposit in Fig. 7 showing (a) the Mo 3d core-level doublet (b) Mo 3p doublet peaks (c) the O1s photoelectron emission and (d) the C 1s binding energy.
Scheme 1
Figure 1
Figure 2
Figure 5
Figure 6
Figure 7
Figure 8