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Solution Layer Deposition: A Technique for the Growth of Ultra-Pure Manganese Oxides on Silica at Room Temperature

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Abstract: With the ever increasing miniaturization in micro-electronic devices, new deposition techniques are required to form high-purity metal oxide layers. Herein, we report a liquid route to specifically produce thin and conformal amorphous manganese oxide layers on silicon substrate, which can be transformed into a manganese silicate layer. The undesired insertion of carbon into the functional layers is avoided through a solution metal–organic chemistry approach named Solution Layer Deposition (SLD). The growth of a pure manganese oxide film by SLD takes place through the decoordination of ligands from a metal–organic complex in mild conditions, and coordination of the resulting metal atoms on a silica surface. The mechanism of this chemical liquid route has been elucidated by solid-state 29Si MAS NMR, XPS, SIMS, and HRTEM.

The deposition of metal oxide thin films on supports is a real challenge in obtaining functional materials of high purity in several applications, such as microelectronics, photovoltaics, and energy storage. To deposit metal oxide layers, three main methods based on vapor deposition processes are commonly used: Physical Vapor Deposition (PVD), Chemical Vapor Deposition (CVD), and Atomic Layer Deposition (ALD). Among them, CVD involves the decomposition of organometallic precursors at high temperature (>250°C) and low pressure, but generates unwanted byproducts in the growing film. Conversely, ALD processes are considered as the most promising techniques to deposit thin layers in nanoscale patterns because it allows formation of various compositions of thin and conformal films. In ALD, the growth of metal oxides is obtained by the alternate introduction of water vapor (which forms surface hydroxy groups on the substrate), and metal precursors (in gas phase) which react on it. One of the key parameters of ALD relies on the choice of the metal precursors, which have to present high volatility, thermal and chemical stability, and efficient self-limited reactivity with surfaces.

Recently, Gordon and co-workers have developed a family of metal–organic amidinate precursors, which are efficient for CVD or ALD deposition. They have reported the conformal deposition of a thin Mn oxide layer on silica support by using bis(N,N′-disopropylpentylamidinato) manganese(II). They showed that CVD of the Mn metal layer allowed formation of a thin amorphous layer of MnSiOₓ on silica. Moreover, this layer presented good barrier properties against interdiffusion between Cu and SiOₓ, which is a crucial property for copper-based microtechnologies.

However, carbon (arising from the precursor ligands) trapped inside the deposited layers may be responsible for the loss of properties of the diffusion barrier layer. For instance, Koike and co-workers reported that the CVD of Mn oxide films induces the presence of an excess of inserted carbon atoms inside the layer, resulting in the progressive loss of adhesion between Cu and Mn oxide. To overcome this problem and decrease the cost of film deposition techniques, the requirement for new deposition processes has never been so crucial. Solution chemistry approaches are good candidates to meet these requirements.

Herein, we report the deposition of a carbon-free Mn oxide layer by a metal–organic solution approach: Solution Layer Deposition (SLD). This method involves the reaction in organic solvent between a hydroxylated silica surface and a metal–organic precursor: bis(N,N′-disopropylpentylamidinato) manganese(II), hereafter named MnAmd₂. The surface of silica is functionalized first by an O₂ plasma treatment, and exposed to ambient air to form surface hydroxy species. The substrate is then immersed at room temperature into a toluene solution of MnAmd₂, which spontaneously reacts with the hydroxy groups of the substrate. This process can be repeated several times, leading to the growth of the Mn oxide layer (Scheme 1). A final annealing ensures the full conversion of Mn oxide into manganese silicate MnSiOₓ. This method is simple and inexpensive, because the oxide layer is grown in very mild conditions.
To examine, by solid state NMR, the nature of the interactions between the silica and Mn oxide layers, one single cycle of the grafting process was performed on silica nanoparticles (NPs), which present a similarly high hydroxy group density compared to plasma-treated SiO$_2$ planar substrate.$^{[10]}$ A first $^{29}$Si MAS NMR analysis, derived from the Bakhmutov procedure, was achieved with pure silica NPs as reference sample 1.$^{[11]}$ Then, the same analysis was performed on silica NPs after contact in MnAmd solution (sample 2). Sample 3 consisted of the mechanical mixture of silica NPs and Mn oxide powder (10% by weight) prepared by a simple air oxidation of MnAmd. The paramagnetic nature of the Mn oxide powder was confirmed by magnetic measurements (Supporting Information, Figure S5). $^{29}$Si magic angle spinning (MAS) NMR spectrum of 1 is presented in Figure 1a. Overlapping lines centered at $-92$, $-100$, and $-109$ ppm were attributed to different Si atoms, respectively $\text{Si}$(Q$_1$), $\text{Si}$(Q$_2$), and $\text{Si}$(Q$_3$) in the silica matrix, where units $\text{Si}$(Q$_4$) dominate. The $^{29}$Si MAS spectrum of 2 mostly presents an isotropic signal at $-109$ ppm ($\text{Si}$(Q$_3$) moieties) displaying sideband patterns (Figure 1b). The signals corresponding to $\text{Si}$(Q$_2$) and $\text{Si}$(Q$_3$) moieties show notable intensity decrease; the $(\text{Si}$(Q$_2$))/$(\text{Si}$(Q$_3$) + $\text{Si}$(Q$_2$)) ratio is 89:11 for 2 compared to 78:22 for 1 (Supporting Information, Figure S6). These signals decrease are most probably due to strong Fermi-contact interactions for nuclei located at the surface of silica, and with paramagnetic Mn centers in close proximity, which renders part of them spectrally invisible.$^{[12]}$ Indeed, the $^{29}$Si nuclei in the first coordination spheres of the Mn paramagnetic centers, which should present paramagnetic isotropic shifts, are not observed. The intense spinning sideband patterns could be attributed to electron–nucleus dipolar interactions, but also to bulk magnetic susceptibility (BMS) effects.$^{[12]}$ It is noteworthy that dipolar electron–nucleus interactions are a through-space effect that decreases as a function of $1/r^3$, and therefore have a significant influence only over short distances. The BMS field is a demagnetization field, arising from larger macroscopic phenomena, that leads to resonance broadenings in solid-state NMR spectra.$^{[12]}$ Since the dipolar coupling and BMS terms depend on the molecular orientations relative to the external magnetic field, they can both lead to spinning sidebands in the MAS NMR spectra.$^{[12]}$ Therefore, these NMR experiments reveal the presence of paramagnetic Mn centers, but are not yet conclusive for chemical interactions between manganese and the silica lattice.$^{[13]}$ To discriminate between both phenomena, a $^{29}$Si MAS NMR spectrum of a physical mixture of silica NPs and Mn oxide powder was recorded (sample 3; Figure 1c). In this case, the Fermi contact and dipolar interactions between $^{29}$Si nuclei and the paramagnetic center are excluded, as the electron–nucleus distances become too important, and only BMS effects should impact the NMR spectra. The $^{29}$Si MAS spectra of 3 did not show the spinning side patterns observed for 2, strongly suggesting that the spinning sidebands are associated to the electron–nucleus dipolar interactions. This confirmed the close spatial proximity between $^{29}$Si nuclei and paramagnetic Mn centers in MnSiO$_3$. This was obtained after a single grafting cycle of MnAmd$_2$ on SiO$_2$ NPs in solution at room temperature. This result is also strongly supported by $T_1$ relaxation time measurements, because the $T_1$ value decreased from $81 \pm 10$ s for 1 to $2.0 \pm 0.3$ s for 2 (Supporting Information, Table S1).

Recent studies indicate that MnSiO$_3$ is the most thermodynamically stable compound formed through the interaction of MnO$_2$ and SiO$_2$.$^{[14,15]}$ MnSiO$_3$ forms an interfacial and covalently bound layer on silica, on which a MnO$_2$ layer grows during further cycles of the process (Supporting Information, Figure S7). As for ALD, the SLD process is self-limited: when MnAmd$_2$ molecules have reacted with all the surface hydroxy moieties, the grafting stops. A new O$_2$ plasma treatment
reactivates the surface, allowing further growth of a Mn oxide layer on the substrate (Figure 2). A STEM-EDS analysis of the cross section of samples prepared with 8 SLD cycles revealed an interfacial zone where both Mn and Si atoms were present between MnO$_x$ and SiO$_2$ layers (Figure 2e,d). The Mn oxide layer deposited on silica can be fully converted into MnSi$_x$O$_y$ by annealing. This reaction was followed by X-Ray Photoelectron Spectroscopy (XPS).\textsuperscript{16,17} Annealing at different temperatures and durations were carried out on 40 nm thick MnO$_x$ layers (Figure 3). The assignment of the O 1s, Mn 2p, and Si 2p peaks from the XPS spectra are summarized in Table S3 (Supporting Information). Sample 4 corresponds to an unannealed MnO$_x$ layer on silica. The O 1s and Si 2p signals, respectively located at 531.1 and 102.2 eV, are characteristic of a MnSi$_x$O$_y$ compound (Figure 3a). This oxide lies at the MnO$_x$/SiO$_2$ interface, in agreement with the MAS NMR conclusions. A component corresponding to MnO$_x$ compound was observed at 529.7 eV for O 1s. When the annealing temperature was increased from 200 to 300°C (samples 5 and 6), the envelope related to the MnO$_x$ compound decreased (Figure 3b,c). Conversely, O 1s and Si 2p components related to MnSi$_x$O$_y$ (531.1 and 102.2 eV) increased. In sample 6 (300°C annealing), the initial 40 nm thick MnO$_x$ layer is fully converted into MnSi$_x$O$_y$ layer. This Mn diffusion into SiO$_2$ is confirmed by the STEM-EDS analysis of cross-sectioned sample 6 (Supporting Information, Figure S8) where the elementary distributions of Mn and Si atoms clearly overlap in the same region.

In sample 7 (450°C annealing), O 1s and Si 2p XP spectra show a decrease of the components related to MnSi$_x$O$_y$. This suggests a deeper diffusion of Mn atoms inside the silica layer. Indeed, STEM-EDS observations reveal a progressive diffusion of MnSi$_x$O$_y$ inside the silica lattice, leading to a higher apparent thickness (25 nm) of the mixed oxide layer (Supporting Information, Figure S8). Consequently, a silicon oxide-rich layer appears over the top of the mixed oxide layer. Too high annealing temperatures lead to a deeper diffusion of the functional oxide layer. For many applications, the adhesion strength between layers is a key parameter, but the presence of carbon atoms inside the deposited layer can induce a loss of adhesion properties.\textsuperscript{19} To quantify the carbon composition into a manganese oxide layer deposited by SLD, Secondary Ion Mass Spectrometry (SIMS) analyses were carried out on samples before and after annealing. The diffusion of Mn in silica was confirmed in the annealed sample, as the depletion of O and Si signals was simultaneously observed with the maximum of Mn content (Figure 4b). The presence of C atoms only appears at the outermost surface of the sample and not inside the deposited layer (Figure 4; Supporting Information, Figure S9).

The relative intensity of C to Mn is about 1:100 in both cases, and appears to be independent of the annealing. This surface carbon may be due to the adsorption of carbon species from the ambient air environment. This low carbon content in the growing film is related to the very mild grafting process in SLD, where the Mn precursor is gently de-coordinated on hydroxy groups on the substrate, and byproducts are removed in the solvent medium.

In conclusion, we have presented an inexpensive and efficient solution deposition technique (SLD) that allows preparation of a Mn oxide layer on a silica substrate. During the first SLD deposition cycle, O$_2$ plasma treatment of the substrate, followed by air exposure, generates hydroxy moieties that react with the MnAm$_2$ precursor in solution. Owing to the high reactivity of MnAm$_2$ to silicon hydroxy groups, the Mn atoms are covalently bound to silica and spontaneously form MnSi$_x$O$_y$, as confirmed by MAS NMR and XPS studies. The repetitive steps of plasma treatment followed by dipping the substrate in solution precursor lead to the growth of a Mn oxide layer. It is noteworthy that SIMS analyses indicated the lack of carbon incorporation in the Mn
oxide layer obtained by this mild solution deposition technique. A controlled annealing (300°C) of the Mn oxide layer induces the diffusion of Mn atoms into the silica lattice, leading to a full conversion into MnSiOx. Such layers can be used in microelectronics applications as a diffusion barrier layer between copper and silica. A copper layer, prepared by decomposition of a copper amidinate precursor according to a previously described method, has been deposited on a silica substrate functionalized with one cycle of the SLD method. After a subsequent annealing at 300°C, no evidence for copper diffusion was observed, thus demonstrating the very efficient barrier property of our MnSiOx layer (Supporting Information, Figure S10). The SLD method allows for a much larger choice of precursors compared with ALD. The amidinate family are good candidates for SLD deposition. We also showed, in other experiments, that zinc amidinate allows deposition of ZnO layers by using the same process. Additionally, many other metal complexes that are reactive towards hydroxy groups can be applied using this method, such as metal alkyls which are highly reactive and would be easily implemented in a liquid phase and low temperature process. Indeed, thermal stability and high volatility are not required properties for precursors employed at room temperature and atmospheric pressure in solution processes. The SLD is therefore a facile and very promising method for the deposition of functional metal oxide layers on various surfaces.

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