

Title	Effect of reaction mechanism on precursor exposure time in atomic layer deposition of silicon oxide and silicon nitride
Authors	Murray, Ciarán A.;Elliott, Simon D.;Hausmann, Dennis;Henri, Jon;LaVoie, Adrian
Publication date	2014-01-10
Original Citation	MURRAY, C. A., ELLIOTT, S. D., HAUSMANN, D., HENRI, J. & LAVOIE, A. 2014. Effect of Reaction Mechanism on Precursor Exposure Time in Atomic Layer Deposition of Silicon Oxide and Silicon Nitride. ACS Applied Materials & Interfaces, 6, 10534-10541. http://dx.doi.org/10.1021/am5021167
Type of publication	Article (peer-reviewed)
Link to publisher's version	10.1021/am5021167
Rights	© 2014 American Chemical Society. This document is the Accepted Manuscript version of a Published Work that appeared in final form in ACS Applied Materials & Interfaces, copyright © American Chemical Society after peer review and technical editing by the publisher. To access the final edited and published work see http://pubs.acs.org/doi/abs/10.1021/am5021167
Download date	2025-07-04 09:02:53
Item downloaded from	https://hdl.handle.net/10468/2424



UCC

University College Cork, Ireland
Coláiste na hOllscoile Corcaigh

This document is confidential and is proprietary to the American Chemical Society and its authors. Do not copy or disclose without written permission. If you have received this item in error, notify the sender and delete all copies.

The Effect of Reaction Mechanism on Precursor Exposure Time in Atomic Layer Deposition of Silicon Oxide and Silicon Nitride

Journal:	<i>Chemistry of Materials</i>
Manuscript ID:	cm-2014-00998r
Manuscript Type:	Article
Date Submitted by the Author:	20-Mar-2014
Complete List of Authors:	Murray, Ciarán; Tyndall National Institute, Elliott, Simon; Tyndall National Institute, Hausmann, Dennis; Lam Research Corporation, Henri, Jon; Lam Research Corporation, LaVoie, Adrien; Lam Research Corporation,

SCHOLARONE™
Manuscripts

The Effect of Reaction Mechanism on Precursor Exposure Time in Atomic Layer Deposition of Silicon Oxide and Silicon Nitride

Ciaran A. Murray[†], Simon D. Elliott^{*†}, Dennis Hausmann[‡], Jon Henri[‡], Adrien LaVoie[‡]

[†]Tyndall National Institute, University College Cork, Lee Maltings, Cork, Ireland

[‡]Lam Research Corporation, 11155 SW Leveton Drive, Tualatin, OR 97062-8094, USA

KEYWORDS: Silicon nitride, Silicon Oxide, Atomic Layer Deposition, DFT, mechanisms.

ABSTRACT: Atomic layer deposition (ALD) of highly conformal, silicon-based dielectric thin films has become necessary due to the continuing decrease of feature size in microelectronic devices. The ALD of oxides and nitrides is usually thought to be mechanistically similar, but plasma-enhanced ALD of silicon nitride is found to be problematic, while that of silicon oxide is straightforward. To find why, the ALD of silicon nitride and silicon oxide dielectric films was studied by applying *ab initio* methods to theoretical models for proposed surface reaction mechanisms. The thermodynamic energies for the elimination of functional groups from different silicon precursors reacting with simple model molecules were calculated using density functional theory (DFT), explaining the lower reactivity of precursors towards the deposition of silicon nitride relative to silicon oxide seen in experiments, but not explaining the trends between precursors. Using more realistic cluster models of amine and hydroxyl covered surfaces, the structures and energies were calculated of reaction pathways for chemisorption of different silicon precursors via functional group elimination, with more success. DFT calculations identified the initial physisorption step as crucial towards deposition and this step was thus used to predict the ALD reactivity of a range of amino-silane precursors, yielding good agreement with experiment. The retention of hydrogen within silicon nitride films but not in silicon oxide observed in FTIR spectra was accounted for by the theoretical calculations and helped verify the application of the model.

INTRODUCTION

Conformal dielectric films based on silicon oxide or silicon nitride are used for liner and spacer applications in front-end-of-line (FEOL) semiconductor wafer processing. The traditional methods for depositing these films have been either plasma enhanced chemical vapor deposition (PECVD) or low pressure chemical vapor deposition (LPCVD). PECVD is capable of depositing semi-conformal films at low temperature (<400°C) and LPCVD is capable of perfectly conformal films at high temperature (>550°C). For sub-32 nm technology however, both low deposition temperature and perfect conformality is required, which has necessitated the move to atomic layer deposition (ALD). Additional applications such as Fin-FET conformal doping, double patterning, and through-Si-via (TSV) 3D integration are also enabled by ALD of silicon-based dielectrics.

Herein, we focus on two silicon dielectric films: SiO₂ and Si₃N₄. The successful deposition of these materials using ALD techniques is shown in Figure 1, demonstrating that high quality conformal films are possible. Plasma enhanced ALD techniques may be applied in the deposition of these materials where an oxygen plasma is used for SiO₂ and a nitrogen plasma for Si₃N₄^{1,2}. Both ALD silicon oxide and nitride films exhibit the properties desired of these materials with excellent step coverage, good dielectric properties and low wet etch rates. Deposition of these films has also demonstrated reasonable growth rates per ALD cycle if exposure is sufficient. Unfortunately, many experimental problems exist, making the application of the ALD of these silicon dielectrics in industrial processes unfeasible, in particular silicon nitride. These experi-

mental difficulties in the ALD of SiO₂ and Si₃N₄ therefore warrant further study.

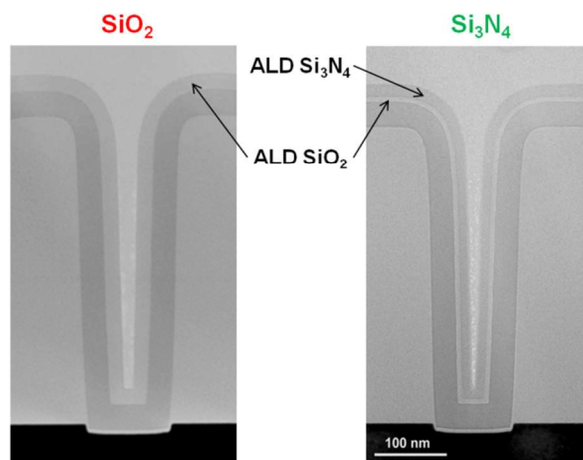


Figure 1: TEM images of SiO₂ and Si₃N₄ ALD thin films.

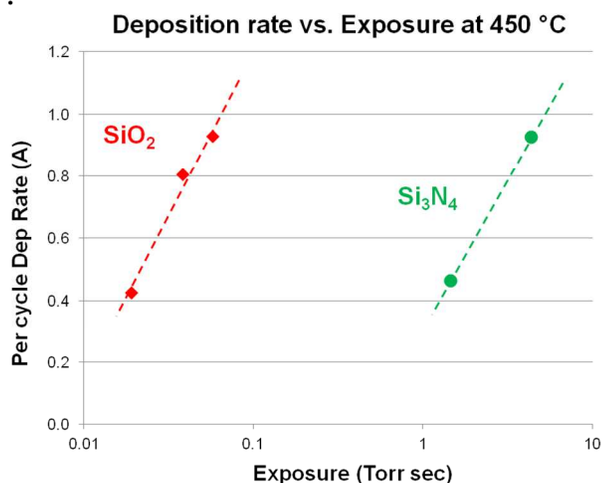


Figure 2: Deposition rate versus exposure of SiO₂ (red diamonds) and Si₃N₄ (green circles) films deposited from chloro-silane precursors by ALD at 450 °C.

Of specific interest is the experimental observation that the required silicon precursor exposure is significantly (>100 ×) higher for Si₃N₄ than for SiO₂. Figure 2 compares the relative reactivity for simple chloro-silane precursors (e.g. DCS, HCDS, etc.) while similar data (not shown) has been obtained using amino-silane precursors (e.g. BTBAS, BDEAS, etc.). The very long precursor exposure for deposition of silicon nitride makes this process economically unviable, due to both the excessive throughput time per film deposited and the unacceptably high volume of silicon precursors consumed. It is the goal of this work to explain the difference in deposition efficiency for a given exposure between SiO₂ and Si₃N₄ and to examine the effect of different silicon precursor on deposition efficiency.

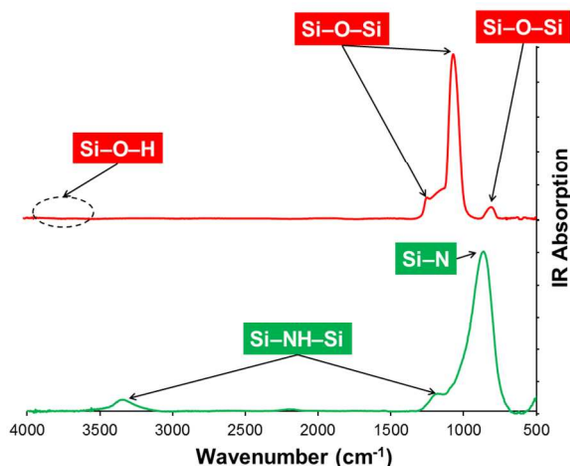


Figure 3: FTIR absorption spectra of SiO₂ (upper red plot) and Si₃N₄ (lower green plot) thin films both deposited by ALD at >400°C.

Representative FTIR data are shown in Figure 3. One of the marked differences between these spectra is the lack of bands associated with hydrogen (Si–O–H) in SiO₂. In Si₃N₄ a peak assigned to NH stretching modes can be clearly observed at 3350 cm⁻¹, whereas the equivalent OH stretching modes in SiO₂ are non-existent. The incorporation of hydrogen in ALD

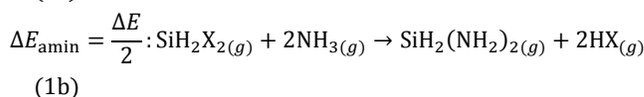
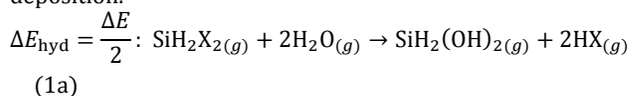
of silicon nitride but not in ALD of silicon oxide suggests different surface chemistries during deposition. In order to investigate the surface chemistry of these materials, models for the deposition of silicon oxide/nitride have been developed. The prediction of hydrogen incorporation can be used as a way of verifying the mechanistic models.

In this paper, we use first principles density functional theory to probe the reasons behind the differences between the ALD of SiO₂ and Si₃N₄. Various theoretical approaches are used including model reaction pathways, acidity/basicity of the oxide vs. nitride surfaces and overall energetics as a function of precursor functional group. A variety of silicon precursors will be taken into account with particular consideration of amino-silane precursors. Amino-silane precursors would be preferred in the ALD of silicon nitride due to the detrimental incorporation of chlorine in films deposited using chloro-silanes. It is assumed that the NH₃ plasma produces an amine-terminated surface, in analogy with the hydroxyl-terminated surface that is produced by oxygen plasma³.

METHOD

All species in this work were modeled as isolated molecules in vacuum in their ground state using the TURBOMOLE suite of programs^{4,5}. All optimized structures and energies (including those of the transition states) were calculated using the generalized gradient approximation Becke-Perdew density functional BP86^{6,7} with the resolution of identity (RI) approximation⁸⁻¹⁰. Atom-centered basis sets were used for all atoms in this work: the large def2-TZVPP basis set for amination/hydrolysis thermodynamic calculations and the smaller def-SV(P) basis set for both cluster models^{11,12}. Transition state structures were optimized by following a vector with a negative eigenvalue (or imaginary frequency) corresponding to the reaction pathway of interest¹³. These transition vectors were determined by performing a vibrational analysis.

The elimination of H–L from a surface has been successfully modeled previously for the ALD of metal oxides using a gas phase hydrolysis model¹⁴. ΔE_{hyd} is the computed energy change of the following model reaction: $\text{ML}_{q(g)} + q\text{H}_2\text{O}_{(g)} \rightarrow \text{M}(\text{OH})_{q(g)} + q\text{HL}_{(g)}$, where M is a metal of valence q and L is a monodentate ligand. In this model gas-phase H₂O represents the source of hydroxyl groups on the surface of the metal oxide while the substitution of OH groups for the ligands represents the formation of new M–O bonds in the solid. Here, in order to model the elimination of groups from difunctionalized silane precursors SiH₂X₂, the hydrolysis model is modified so that only elimination of two functional groups (X) is considered during the ALD of silicon oxide (Equation 1a). An equation to represent surface reactivity on silicon nitride is proposed where gas-phase ammonia (NH₃) represents NH bonds on the surface (analogous to H₂O representing surface hydroxyl groups) and the substitution of amide groups (NH₂) for the functional groups, X, represents the formation of new Si–N bonds (Equation 1b). This model reaction is referred to in this work as “hydrolysis” for the deposition of silicon oxide and “amination” when considering silicon nitride deposition.



In equations 1a and 1b, ΔE is quoted per functional group X. The more negative a ΔE_{hyd} or ΔE_{amin} value is, the more exothermic the hydrolysis/amination reaction and the greater the possibility of HX elimination. As this model is concerned solely with the changes in bonding, temperature effects are neglected.

In many cases, the activation energies and surface geometries are important in determining the reactivity of a molecule at a surface and so the model proposed in Section 2.1 is not adequate. The proposed mechanism for the reactive chemisorption of a SiH_2X_2 precursor is (i) physisorption, (ii) proton transfer via a transition state followed by (iii) HX elimination¹⁵. The various steps required to model this process are described in Figure 4.

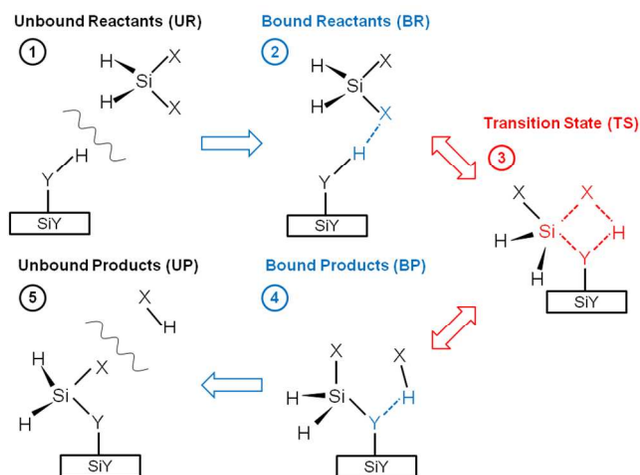


Figure 4: The proposed mechanistic steps for reactive chemisorption of SiH_2X_2 precursor and desorption of functional group X via proton transfer from a $\text{Si}-\text{Y}-\text{H}$ functionalized surface ($\text{Y} = \text{O}, \text{NH}$ or N).

The first step in Figure 4 shows the “Unbound Reactants” (UR) where the precursor SiH_2X_2 is isolated, by an effectively infinite distance, from a surface group $\text{Y}-\text{H}$ (where $\text{Y} = \text{O}, \text{NH}$ or N). In the second “Bound Reactants” (BR) step, a loosely bound complex is formed between a group X on the precursor and a surface hydrogen atom. Direct coordination of the Si precursor to surface-Y via dative bonding is not expected in this mechanism because Si does not tend to increase its coordination number beyond four in stable structures. This BR structure leads into a four-membered ring “Transition State” (TS) involving the Si and X on the precursor and Y and H on the surface. With five-fold coordination about Si in TS, this structure is expected to be unstable and transient. Step 4 describes the “Bound Products” (BP) where the precursor is now chemisorbed to the surface but the newly formed molecule $\text{H}-\text{X}$ is still hydrogen bonded to Y. Step 5 in Figure 4 shows the final “Unbound Products” (UP) where $\text{H}-\text{X}$ has been fully eliminated from the surface and the SiH_2X group bound to the surface has relaxed to its most stable structure.

RESULTS AND DISCUSSION

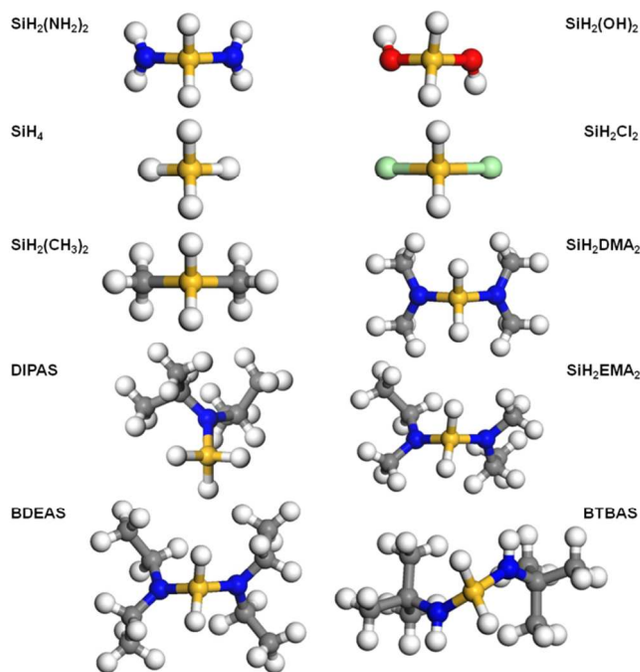


Figure 5: Molecular structures of some of the silicon precursors considered in this work. Structures were optimized using BP86/TZVPP. The silicon atoms are represented by yellow spheres, hydrogen by white, nitrogen by blue, oxygen by red, carbon by grey and chlorine by green.

In the following sections, results from the application of the theoretical models described in above will be given. Multiple di-alkylamide silanes with the general formula $\text{SiH}_2(\text{NR}'\text{R}'')_2$ were considered including: SiH_2DMA_2 [$\text{R}'=\text{R}''=\text{CH}_3$, bis(dimethylamino)silane], $\text{SiH}_2\text{HFMA}_2$ [$\text{R}'=\text{R}''=\text{CF}_3$, bis(hexafluorodimethylamino)silane], SiH_2EMA_2 [$\text{R}'=\text{CH}_3$, $\text{R}''=\text{CH}_2\text{CH}_3$, bis(ethylmethylamino)silane], BDEAS [$\text{R}'=\text{R}''=\text{CH}_2\text{CH}_3$ bis(diethylamino)silane], and BTBAS [$\text{R}'=\text{C}(\text{CH}_3)_3$, $\text{R}''=\text{H}$, bis(tert-butylamino)silane] as well as $\text{SiH}_2(\text{NH}_2)_2$ [diamino-silane] and DIPAS [diisopropylamino-silane] with only one amide functional group with $\text{R}'=\text{R}''=\text{CH}(\text{CH}_3)_2$ attached to the silyl SiH_3 group. Some results for other silanes including SiH_2Cl_2 [dichlorosilane], $\text{SiH}_2(\text{CH}_3)_2$ [dimethylsilane] and SiH_4 [silane] will also be given. The molecular structures of these precursors are shown in Figure 5.

The molecular structures for a selection of silicon precursors and the molecules ($\text{SiH}_2(\text{NH}_2)_2$ and $\text{SiH}_2(\text{OH})_2$ respectively) employed to model the silicon nitride and oxide surfaces were optimized using BP86/TZVPP. The resulting geometries are depicted in Figure 5. The functional groups are coordinated to the silicon atom in a quasi-tetrahedral fashion, exemplified by the SiH_4 parent molecule. Some distortion from this idealized coordination is seen for $\text{SiH}_2\text{HFMA}_2$, SiH_2EMA_2 , BDEAS and BTBAS molecules due to the relatively large size of the functional group and for $\text{SiH}_2(\text{OH})_2$ due to intra-molecular interactions between the two OH groups. Optimization of the same molecules using the SV(P) basis set resulted in similar geometries.

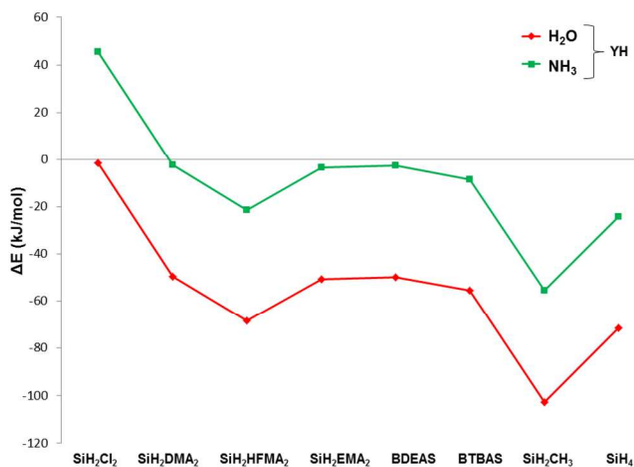


Figure 6: ΔE for the hydrolysis (YH = H_2O) and amination (YH = NH_3) of various silicon precursors calculated using equations 1a and 1b from BP86/TZVPP total energies.

The ΔE values for both the hydrolysis and amination of a selection of silicon precursors were determined from Equations 1a and 1b using DFT calculated energies (BP86/TZVPP) and are presented in Figure 6. ΔE_{hyd} was significantly more negative than ΔE_{amin} by 47.14 kJ/mol (this value is actually $|\Delta E|$ of $\text{SiH}_2(\text{NH}_2)_2 + 2\text{H}_2\text{O} \leftrightarrow \text{SiH}_2(\text{OH})_2 + 2\text{NH}_3$). This result predicts that functional group elimination from di-functionalized silane precursors is thermodynamically more favorable on OH-covered SiO_2 surfaces than on NH_2/NH -covered Si_3N_4 surfaces. This appears to correlate with the much slower ALD growth rates for silicon nitride compared with silicon dioxide.

Unfortunately, the limitations of this thermodynamic model become apparent when the trends between the different precursors are considered. $\text{SiH}_2(\text{CH}_3)_2$ and SiH_4 are predicted here to be the most reactive and SiH_2Cl_2 the least reactive. These theoretical predictions are contradicted experimentally where $\text{SiH}_2(\text{CH}_3)_2$ and SiH_4 are not precursors for the ALD of either SiO_2 or Si_3N_4 due to slow or negligible growth rates. SiH_2Cl_2 is one of the more promising precursors, in particular for the deposition of Si_3N_4 , and has demonstrated significant ALD growth rates in experiment. Even for the alkyl amides (SiH_2DMA_2 , SiH_2EMA_2 , BDEAS and BTBAS), greater variation in deposition rates is seen in experiments than compared to the theoretical results presented in Figure 6. A kinetic model, considering energy barriers and surface geometries, is therefore needed to explain the differences between silicon amide precursors in the deposition of silicon nitride and dioxide.

Simple surface models consisting of one functional group of interest were investigated. The silyl group SiH_3 was selected to represent both the SiO_2 and Si_3N_4 bulk material. To model hydroxyl groups on SiO_2 , an OH group was added to the SiH_3 fragment resulting in a $\text{SiH}_3\text{-OH}$ (silanol) surface model. For the nitride models, either NH_2 was added to SiH_3 to represent a primary amine, resulting in a $\text{SiH}_3\text{-NH}_2$ (silylamine) surface model or NH-SiH_3 to model secondary amides, a $\text{SiH}_3\text{-NH-SiH}_3$ (disilylamine) surface model. Two precursors were initially considered, $\text{SiH}_2(\text{NH}_2)_2$ and SiH_2DMA_2 , due to their small size, reducing the computation resources required and allowing easy analysis of the resulting structures. All $\text{SiH}_3\text{-YH}$ results presented here were calculated using the BP86 GGA density functional and SV(P) basis set.

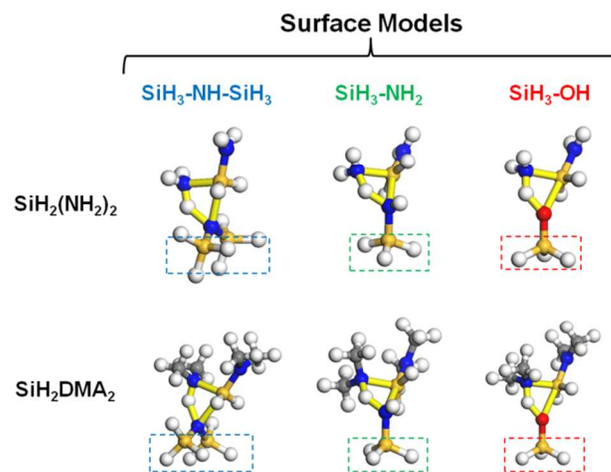


Figure 7: DFT optimized structures of the transition states for the reactions of $\text{SiH}_2(\text{NH}_2)_2$ and SiH_2DMA_2 silicon precursors with $\text{SiH}_3\text{-NH-SiH}_3$, $\text{SiH}_3\text{-NH}_2$ and $\text{SiH}_3\text{-OH}$ surface models. The silicon atoms are represented by yellow spheres, hydrogen by white, nitrogen by blue, oxygen by red and carbon by grey. The bonds of the 4-membered ring transition state are highlighted in yellow.

The optimized geometries and energies for the proton transfer steps (outlined in Figure 4) from $\text{SiH}_3\text{-NH-SiH}_3$, $\text{SiH}_3\text{-NH}_2$ and $\text{SiH}_3\text{-OH}$ to one of the functional groups of $\text{SiH}_2(\text{NH}_2)_2$ and SiH_2DMA_2 were calculated. The transition state (TS) geometries from these calculations are shown in Figure 7. For these transition states a planar, “kite” shaped 4-membered ring is found between the O/N and H of the surface model molecule and the N and Si of the incoming precursors. The orientation of the precursors towards the surface models is different for each group. While the precursors may approach $\text{SiH}_3\text{-OH}$ vertically to form a transition state, a side-on approach of the precursors is necessary for the $\text{SiH}_3\text{-NH}_2$ due to the orientation of the amine hydrogen atoms. This seems to reflect the direction of the lone pair on O and N respectively.

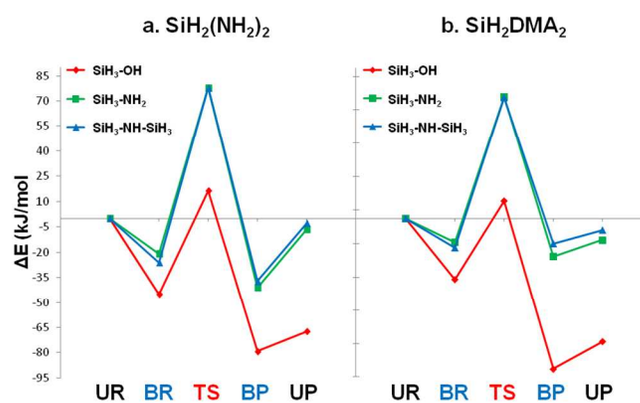


Figure 8: Energetics for the Bound Reactants (BR), Transition State (TS), Bound Products (BP) and Unbound Products (UP) relative to the Unbound Reactants (UR) for the reaction of a. $\text{SiH}_2(\text{NH}_2)_2$, and b. SiH_2DMA_2 precursor with the surface group models $\text{SiH}_3\text{-OH}$ (red diamonds), $\text{SiH}_3\text{-NH}_2$ (green squares) and

SiH₃-NH-SiH₃ (blue triangles). ΔE values are given in kJ/mol and were determined using BP86/SV(P) DFT calculations.

The energetics calculated for SiH₂(NH₂)₂ and SiH₂DMA₂ precursors are very similar (Figure 8). The reactions leading to BR, BP and UP (optimized to local minima) on the SiH₃-OH model surface were calculated to be more exothermic than those on the SiH₃-NH₂ and SiH₃-NH-SiH₃ systems, consistent with the results of the thermodynamic model (c.f. Section 4.1). The ΔE values for the SiH₃-NH₂ and SiH₃-NH-SiH₃ models were almost identical, indicating the similar chemistry of primary (NH₂) and secondary (NH) amine groups. The transition state energy barriers or activation energies, E_{act} , (i.e. $E_{act} = E(TS) - E(BR)$) are presented in Table 1. Again the SiH₂(NH₂)₂ and SiH₂DMA₂ results are quite similar with E_{act} for both precursors with the SiH₃-OH substrate lower than with SiH₃-NH₂/SiH₃-NH-SiH₃ substrates. Although E_{act} values (as well as energies of the local minima BR, BP and UP) are lower for SiH₂(NH₂)₂ and SiH₂DMA₂ with SiH₃-OH than on either SiH₃-NH₂ or SiH₃-NH-SiH₃, it must be remembered that these calculations ignored temperature effects that may reduce the significance of these energy differences.

Table 1: Activation energies, E_{act} , determined from $E(TS) - E(BR)$ in the reaction pathway for SiH₂(NH₂)₂ and SiH₂DMA₂ precursors with SiH₃-OH, SiH₃-NH₂ and SiH₃-NH-SiH₃ substrate models. Energies are in kJ/mol and were determined from BP86/SV(P) DFT calculations.

E_{act} (kJ/mol)	SiH ₂ (NH ₂) ₂	SiH ₂ DMA ₂
SiH ₃ -OH	61.3	46.7
SiH ₃ -NH ₂	98.8	87.1
SiH ₃ -NH-SiH ₃	104.1	89.8

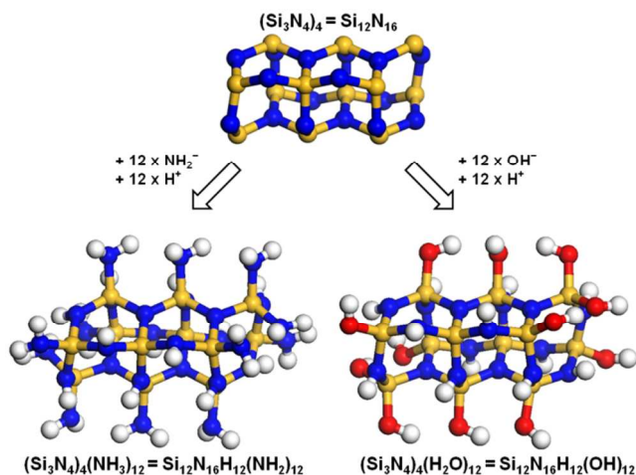


Figure 9: $(Si_3N_4)_4$ cluster model (upper center) with NH₂/H terminated $(Si_3N_4)_4(NH_3)_{12}$ cluster (lower left) and OH/H terminated $(Si_3N_4)_4(H_2O)_{12}$ cluster (lower right).

A cylindrical silicon nitride cluster consisting of 4 Si_3N_4 stoichiometric units ($Si_{12}N_{16}$ with 28 atoms) was constructed,

based on the chemically stable, β -phase crystal structure with hexagonal symmetry¹⁶. To this bare cluster, 12 NH₂⁻ anions and 12 H⁺ cations were added, terminating the uncoordinated Si and N atoms respectively on the outside of the cluster. This had the effect of adding 12 NH₃ molecules to the cluster, retaining the neutral charge of the cluster. This $(Si_3N_4)_4(NH_3)_{12}$ cluster ($Si_{12}N_{16}H_{12}(NH_2)_{12}$ with 76 atoms) was used to model ALD reactions of silicon precursors at a Si_3N_4 surface. The model used for reactions at a silicon oxide surface was constructed in a similar fashion to the silicon nitride, where 12 OH⁻ and 12 H⁺ fragments (i.e. 12 H₂O molecules) were added to the $Si_{12}N_{16}$ cluster resulting in a $(Si_3N_4)_4(H_2O)_{12}$ cluster ($Si_{12}N_{16}H_{12}(OH)_{12}$ with 64 atoms). Although SiO₂ has a different crystal structure than Si_3N_4 , the use of the same $(Si_3N_4)_4$ core for both silicon dioxide and nitride models allowed direct comparison of many calculated properties e.g. changes in geometry between reaction steps.

The geometries of both $(Si_3N_4)_4(H_2O)_{12}$ and $(Si_3N_4)_4(NH_3)_{12}$ clusters were optimized using the BP86 DFT functional and SV(P) basis set. The resulting structures are shown in Figure 9 with only minor changes to the underlying $(Si_3N_4)_4$ cluster seen during the optimizing process. Optimization of the various structures corresponding to the reaction steps outlined in Figure 4, was attempted for the SiH₂(NH₂)₂ and SiH₂DMA₂ precursors with both the $(Si_3N_4)_4(H_2O)_{12}$ and $(Si_3N_4)_4(NH_3)_{12}$ surface models, using the same method as for the smaller SiH₃-XH models (BP86/SV(P)). Only reactions of precursors with NH₂ and OH groups were considered due to the relative inaccessibility of the secondary, NH amine groups on the cluster.

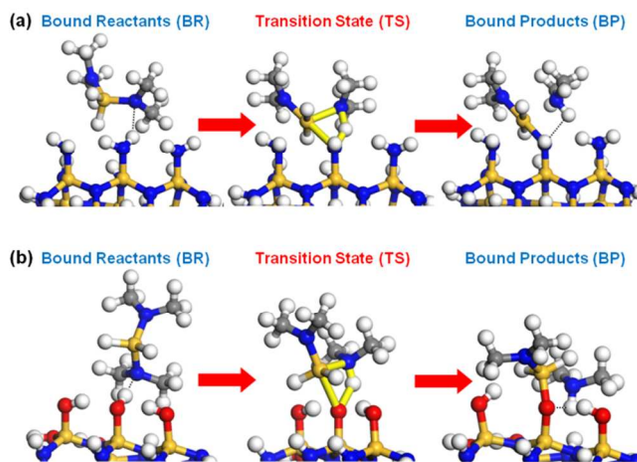


Figure 10: BP86/SV(P) optimized geometries of Bound Reactants, Transition State and Bound Products structures for SiH₂DMA₂ with a. $(Si_3N_4)_4(NH_3)_{12}$ as a model for silicon nitride growth and b. $(Si_3N_4)_4(H_2O)_{12}$ as a model for silicon oxide growth.

The structures determined for both SiH₂DMA₂ and SiH₂(NH₂)₂ amide precursors were quite similar. Figure 10 depicts the geometries determined for the BR, TS and BP steps of the SiH₂DMA₂ reaction with cluster model surfaces. The structures calculated for the reaction steps are qualitatively similar to those determined using the SiH₃-YH models with a few notable differences. In the BR and TS steps of SiH₂DMA₂ on $(Si_3N_4)_4(NH_3)_{12}$, there is an increased steric interaction between one of the CH₃ groups on the reacting DMA fragment and the cluster surface. A side-on orientation of the reacting precursor alkyl amide group with respect to the surface

means that only one CH₃ group of DMA interacts strongly with the surface while the other points away. For reactions involving SiH₂(NH₂)₂, these steric interactions are reduced due to the smaller NH₂ precursor groups. The favorable orientation of hydroxyl groups on the (Si₃N₄)₄(H₂O)₁₂ cluster also reduces the steric interactions.

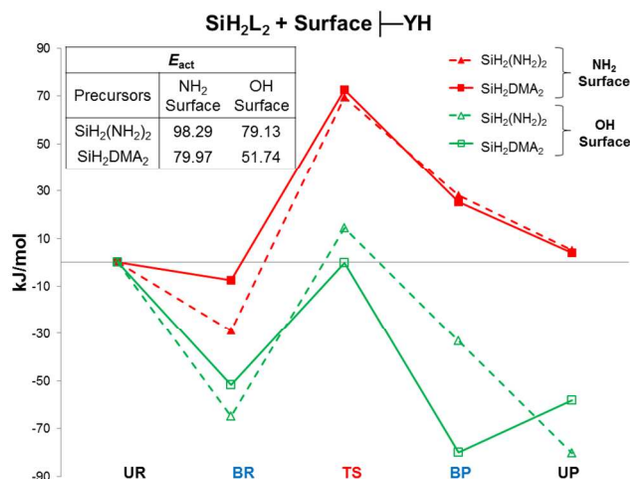


Figure 11: Energetics for the Bound Reactants (BR), Transition State (TS), Bound Products (BP) and Unbound Products (UP) relative to the Unbound Reactants (UR) for the reaction of SiH₂(NH₂)₂ (triangles, dashed lines), SiH₂DMA₂ (squares, solid lines) with the cluster models (Si₃N₄)₄(NH₃)₁₂ (red) and (Si₃N₄)₄(H₂O)₁₂ (green). The table inset presents the E_{act} values for the precursors with the NH₂ and OH covered surfaces. ΔE values are given in kJ/mol and were determined from BP86/SV(P) DFT calculations.

The relative energies of the mechanism steps for the three model precursors with the NH₂ and OH covered clusters are plotted in Figure 11. The E_{act} barriers are comparable to those determined using the smaller SiH₃-NH₂ and SiH₃-OH models. The reduction in energy from BP to UP steps for SiH₂(NH₂)₂ reactions with both surfaces and SiH₂DMA₂ with (Si₃N₄)₄(H₂O)₁₂ is due to the removal of the hydrogen bonded NH₃ and H-DMA molecules. This allows the newly formed surface SiH₂NH₂/SiH₂DMA groups to relax to more stable structures that were prohibited by the presence of the H bonded amine molecules in the Bound Products structures.

One of the most striking differences between these larger cluster calculations and those of the SiH₃-YH models, is the wide spread now apparent in the BR energies relative to UR (Figure 11). This energy difference is the adsorption energy. The larger steric interactions experienced by SiH₂DMA₂ with the amine-covered surface compared with the hydroxyl-covered surface destabilize BR. The same steric interactions explain the weaker adsorption by SiH₂DMA₂ relative to SiH₂(NH₂)₂ on both substrates. The higher energy of BR reflects a reduced bond strength of the precursor to the surface. This may increase the probability of the precursor returning unreacted to the gas phase at ALD temperatures and may prevent the remaining reaction steps occurring. This initial BR step is therefore crucial in determining the deposition rate of a precursor and was chosen as a metric for ALD reactivity of other potential precursors.

Table 2: Energy difference calculated between Unbound Reactants (UR) and Bound Reactants (BR) structures, $\Delta E[BR] =$

$E(UR) - E(BR)$, for SiH₂(NH₂)₂, SiH₂DMA₂, BTBAS, BDEAS and DIPAS precursors and (Si₃N₄)₄(NH₃)₁₂ and (Si₃N₄)₄(H₂O)₁₂ cluster models. Energy values (in kJ/mol) were determined using BP86/SV(P) calculations.

$\Delta E[BR]$ (kJ/mol)	SiH ₂ (NH ₂) ₂	SiH ₂ DMA ₂	BTBAS	BDEAS	DIPAS
(Si ₃ N ₄) ₄ (NH ₃) ₁₂	-28.8	-7.5	-21.4	-5.8	-10.6
(Si ₃ N ₄) ₄ (H ₂ O) ₁₂	-64.8	-51.8	-47.4	-41.9	-32.4

The $\Delta E[BR]$ values calculated for a series of precursors bound to either (Si₃N₄)₄(NH₃)₁₂ or (Si₃N₄)₄(H₂O)₁₂ surface models are tabulated in Table 2. In all cases $\Delta E[BR]$ for adsorption onto the oxide model surface (Si₃N₄)₄(H₂O)₁₂ is significantly lower than that onto the nitride model (Si₃N₄)₄(NH₃)₁₂. The most exothermic $\Delta E[BR]$ value determined for (Si₃N₄)₄(H₂O)₁₂ was for the SiH₂(NH₂)₂ precursor, gradually increasing to the least reactive, BDEAS. A greater variation in relative reactivity was determined for the same amino-silane precursor with (Si₃N₄)₄(NH₃)₁₂ than seen for (Si₃N₄)₄(H₂O)₁₂. As described above in Section 4.2, the difference in $\Delta E[BR]$ between SiH₂(NH₂)₂ and SiH₂DMA₂ was primarily due to the difference in size between the functional groups, the smaller hydrogen atom in SiH₂(NH₂)₂ reduced steric interactions compared with the larger methyl groups of SiH₂(DMA)₂. The substantially different $\Delta E[BR]$ values of -21.4 and -5.8 kJ/mol determined for BTBAS and BDEAS on the nitride substrate are, at first glance, surprising considering that they have the same number of constituent atoms. Examining the molecular structure of these precursors (see Figure 5), the presence of NH in the BTBAS allows the alkyl amide functional group to approach surface NH₂ groups with relatively low steric hindrance, comparable to the much smaller SiH₂(NH₂)₂. It is therefore possible to combine large R groups for volatility with accessible NH groups for adsorption in precursors of the type SiH₂(NHR)₂. BDEAS, unlike BTBAS, has two ethyl groups on each of its alkyl amide functional groups, resulting in steric interactions similar to SiH₂DMA₂, decreasing the stability of this BR complex. Although DIPAS has only one alkyl amide attached to its central silicon atom, the large isopropyl groups interact in a similar manner to the alkyl groups of SiH₂DMA₂ and BDEAS creating steric hindrance around the reacting site.

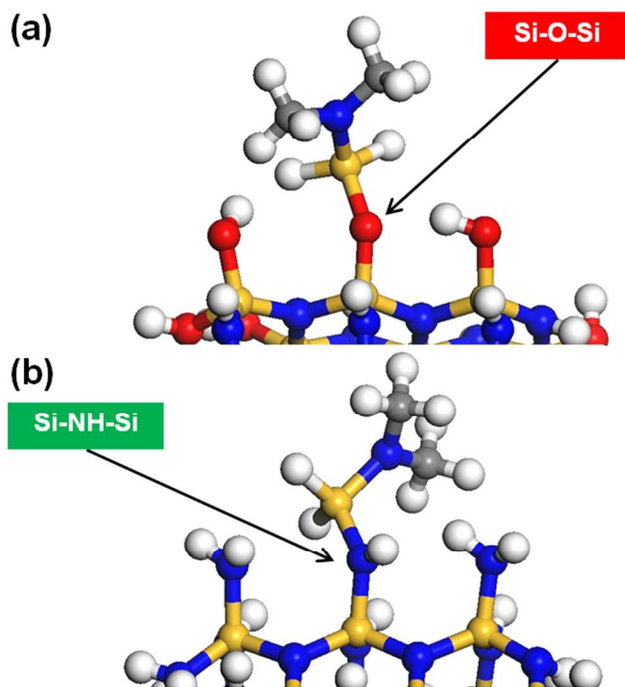


Figure 12: Structures of chemisorbed SiH_2DMA_2 product (UP) on (a) $(\text{Si}_3\text{N}_4)_4(\text{H}_2\text{O})_{12}$ and (b) $(\text{Si}_3\text{N}_4)_4(\text{NH}_3)_{12}$ clusters.

The presence of intense absorption bands, assigned to NH stretching and bending modes, in FTIR spectra of Si_3N_4 compared to the absence of OH bands in SiO_2 is one of the notable differences observed between ALD-grown silicon nitride and silicon dioxide. This retention of hydrogen in silicon nitride films may be explained by looking at the final unbound products (UP) calculated using the proton transfer mechanism described in this work. The molecular structure shown in Figure 12a depicts a SiH_2DMA group chemisorbed to a $(\text{Si}_3\text{N}_4)_4(\text{H}_2\text{O})_{12}$ surface after the elimination of an H-DMA molecule. The new silicon atom from the precursor is bonded to the surface via an oxygen atom which in turn is bonded to a Si atom of the bulk substrate (Si–O–Si), leaving no H atoms coordinated to the oxygen. Elimination of the other DMA group as H-DMA may further deplete the surface of hydrogen, forming a second Si–O–Si bridge.

Considering the product from the same precursor onto the $(\text{Si}_3\text{N}_4)_4(\text{NH}_3)_{12}$ cluster in Figure 12b, the incoming silicon atom is bonded to the surface via a nitrogen and then to a substrate silicon atom. However, unlike the SiO_2 model, the nitrogen atom bridging the new silicon atom with the underlying surface has a remaining hydrogen atom (Si–NH–Si) where elimination of the other DMA group would form another Si–NH–Si bridge. Further adsorption of a second precursor to bridging NH group and removal of second hydrogen during the silicon precursor ALD pulse is prevented by steric hindrance from the first precursor fragment, particularly in attempting to form the four-membered ring TS (Figure 10). In this way, despite removal of DMA by the nitrogenation pulse (e.g. by treatment with NH_3 plasma), this NH group will be buried within the film and will remain even after annealing.

CONCLUSIONS

Silicon nitride films deposited using ALD with silicon precursors and NH_3 plasma require a precursor exposure more than 100 times greater than that for silicon oxide films deposited with oxygen plasma. Experiments also show significantly

different deposition rates between the precursors employed in the deposition of silicon nitride. In order to explain these differences, various theoretical models employing DFT calculations have been applied. Thermodynamic models using DFT calculated energies correctly predict the lower reactivity of silicon precursors with amine-terminated surfaces compared to hydroxylated surfaces, but failed to predict the trends in reactivity between precursors. A mechanistic pathway for growth involving the elimination of a precursor functional group via proton transfer mechanism was applied first to small molecules representing surface groups and then to larger cluster models of silicon nitride and silicon oxide surfaces. A significant difference in reactivity is observed due to the orientation of the hydrogen atoms attached to the hydroxyl and amine groups, where in-coming precursors approach the OH group vertically and approach the NH_2 group side-on. The nitride surface is therefore considerably more sensitive to precursor bulk.

The significance of this difference in approach of the precursors towards the surface groups becomes apparent when reactivity with the larger cluster models is considered. For the OH covered surface little interaction is observed between the precursor and surrounding surface, but for the NH_2 covered surface, the side-on approach of the precursor causes the precursor groups to be oriented towards the surface. This has a substantial effect on the strength of the H-bonding between precursor and surface (“Bound Reactants”) and thus on the lifetime of the adsorbed state and the probability of further reaction before desorption. Their relative adsorption energetics are therefore used to estimate the ALD kinetics and exposure required. With regard to the silicon dioxide surface model, all the amino-silane precursors considered in this work were determined to have reasonably strongly bound reactants (adsorption energies between -32 and -65 kJ/mol) and therefore a reasonable ALD growth rate is predicted. The steric bulk of the amine functional groups attached to the precursors was found to have a greater effect on ALD growth of silicon nitride.

For the precursors where one or more smaller R groups were attached to the amine functional groups (e.g. $\text{SiH}_2(\text{NH}_2)_2$, $\Delta E[\text{BR}] = -28.2$ kJ/mol), more stable bound reactants structures were found than those with larger R groups (e.g. BDEAS, $\Delta E[\text{BR}] = -5.9$ kJ/mol). DFT calculations for the bound reactants of the larger SiH_2DMA_2 and BDEAS precursors yielded the lowest adsorption energies -7.5 and -5.8 kJ/mol. Despite the same number of atoms as BDEAS and a large t-butyl group attached to one position of the amine functional groups, adsorption of BTBAS (-21.4 kJ/mol) was determined to be significantly more exothermic than that of BDEAS (-5.8 kJ/mol). In fact BTBAS adsorbs as easily as the model precursor $\text{SiH}_2(\text{NH}_2)_2$ (-28.8 kJ/mol) where the presence of the small hydrogen on the amine functional group allows the precursors to form bound reactant structures with reduced steric interactions with the surrounding surface compared to larger alkyl groups.

The mechanism presented in this work for the adsorption of silicon precursors via functional group elimination predicts hydrogen atoms to be present both on the surface and embedded within the growing silicon nitride film. By contrast, the same mechanism predicts that hydrogen is only present on the surface of silicon oxide film. In the experimental FTIR absorption spectra, vibrational bands associated with NH bonds in silicon nitride are indeed detected, in contrast with a lack of OH vibrational bands in silicon oxide. This helps validate the proposed growth mechanism and theoretical approach. Due the relative unreactivity of NH

1 groups towards functional group elimination, the plasma assisted
2 ALD silicon nitride is much slower compared to that of silicon
3 oxide systems and requires longer precursor exposure. Targeted
4 reduction in precursor bulk may improve the situation, but the
5 main reason is the inflexible orientation of amine groups at the
6 surface, which is an intrinsic property of the silicon nitride material being deposited.

7 AUTHOR INFORMATION

8 Corresponding Author

9 * E-mail: simon.elliott@tyndall.ie

10 Author Contributions

11 All authors have given approval to the final version of the manu-
12 script.
13

14 ACKNOWLEDGEMENTS

15 The authors would like to thank Lam Research Corporation for
16 the financial support of this work.
17

18 REFERENCES

- 19 (1) Koehler, F.; Triyoso, D. H.; Hussain, I.; Mutas, S.; Bern-
20 hardt, H. *IOP Conference Series: Materials Science and Engineering*
21 **2012**, *41*, 012006.
22 (2) Triyoso, D. H.; Jaschke, V.; Shu, J.; Mutas, S.; Hempel, K.;
23 Schaeffer, J. K.; Lenski, M. In *IC Design & Technology (ICICDT),*
24 *2012 IEEE International Conference on 2012*, p 1.
25 (3) Elliott, S. D.; Scarel, G.; Wiemer, C.; Fanciulli, M.; Pavia,
26 G. *Chem. Mater.* **2006**, *18*, 3764.
27 (4) Ahlrichs, R.; Bär, M.; Häser, M.; Horn, H.; Kölmel, C.
28 *Chem. Phys. Lett.* **1989**, *162*, 165.
29 (5) TURBOMOLE V6.1 2009, a development of University of
30 Karlsruhe and Forschungszentrum Karlsruhe GmbH, 1989-2007,
31 TURBOMOLE GmbH, since 2007
32 (6) Becke, A. D. *Phys. Rev. A* **1988**, *38*, 3098.
33 (7) Perdew, J. P. *Phys. Rev. B* **1986**, *33*, 8822.
34 (8) Eichkorn, K.; Treutler, O.; Öhm, H.; Häser, M.; Ahlrichs,
35 R. *Chem. Phys. Lett.* **1995**, *242*, 652.
36 (9) Eichkorn, K.; Weigend, F.; Treutler, O.; Ahlrichs, R. *The-*
37 *or. Chem. Acc.* **1997**, *97*, 119.
38 (10) Sierka, M.; Hogeckamp, A.; Ahlrichs, R. *J. Chem. Phys.*
39 **2003**, *118*, 9136.
40 (11) Schäfer, A.; Horn, H.; Ahlrichs, R. *The Journal of Chemi-*
41 *cal Physics* **1992**, *97*, 2571.
42 (12) Weigend, F.; Ahlrichs, R. *Phys. Chem. Chem. Phys.* **2005**,
43 *7*, 3297.
44 (13) Helgaker, T. *Chem. Phys. Lett.* **1991**, *182*, 503.
45 (14) Murray, C.; Elliott, S. D. *ACS Applied Materials & Inter-*
46 *faces* **2013**, *5*, 3704.
47 (15) O'Neill, M. L.; Bowen, H. R.; Derecskei-Kovacs, A.;
48 Cuthill, K. S.; Han, B.; Xiao, M. *Interface-Electrochemical Society*
49 **2011**, *20*, 33.
50 (16) Grün, R. *Acta Crystallographica Section B* **1979**, *35*, 800.

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

