OVERVIEW

Studying chemical vapor deposition processes with theoretical chemistry

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Abstract In a chemical vapor deposition (CVD) process, a thin film of some material is deposited onto a surface via the chemical reactions of gaseous molecules that contain the atoms needed for the film material. These chemical reactions take place on the surface and in many cases also in the gas phase. To fully understand the chemistry in the process and thereby also have the best starting point for optimizing the process, theoretical chemical modeling is an invaluable tool for providing atomic-scale detail on surface and gas phase chemistry. This overview briefly introduces to the non-expert the main concepts, history and application of CVD, including the pulsed CVD variant known as atomic layer deposition, and put into perspective the use of theoretical chemistry in modeling these processes.

Keywords Chemical vapor deposition · Atomic layer deposition · Thin films · Surface chemistry · Gas phase chemistry · Theoretical chemistry

1 An introduction to vapor-phase deposition techniques

Thin films are layers of materials with thicknesses ranging from <1 nm (a few atomic layers) to hundreds of

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S. D. Elliott Tyndall National Institute, University College Cork, Lee Maltings, Cork, Ireland e-mail: Simon.Elliott@tyndall.ie micrometers (for reference, a human hair is about 75 µm thick) [1]. The importance of thin films in today's society is enormous, and thin films can be found everywhere; from low friction coatings in a car engine to the anti-reflecting coating on the lenses of spectacles, as well as the decorative coating on their frames. Most metal objects around us have been machined by cutting tools that are coated with a hard, wear-resistant thin film. Replacement parts for the human body, such as hip joints, are often coated with a thin film to make them more biocompatible. Furthermore, today's nanoelectronic devices are built up very precisely from stacks of thin films of various materials with different electrical properties, with some of the films as thin as one atomic layer. Technologically important thin films can be amorphous, polycrystalline or epitaxially grown single crystals, and the properties of the materials can often be tuned with great precision to suit various applications.

To coat an object (the "substrate") with a thin film, it is often preferred to start from atoms or molecules in a vapor phase and place the object(s) to be coated in that vapor, letting atoms and/or molecules from the vapor build up a thin film on the surface of the object. These vapor-based thin film synthesis methods are classified as either physical vapor deposition (PVD) or chemical vapor deposition (CVD), depending on whether the film deposition process is driven by physical impacts or by chemical reactions, respectively. Generating the vapor in the reactor is of course straightforward when the desired element is available in gaseous form, e.g., O₂, but this is not the case for most elements. Therefore, in PVD, a solid sample containing the target elements is subjected to substantial energy, often in the form of a plasma or an electric discharge, thereby ejecting atoms and producing a vapor, which can then condense onto the substrate [2]. In CVD, target elements are delivered in the form of volatile

molecules, denoted as precursors, and the film is built up via a series of chemical reactions between precursors, precursor fragments and the substrate. In the general case, such reactions can take place both in the gas phase and on the substrate surface. However, a form of CVD named atomic layer deposition (ALD) uses only surface chemical reactions to build up thin films with great precision [3, 4].

The precursor molecules are often diluted in a carrier gas that makes up the main part of the gas volume in the process, analogous to the solvent in liquid-phase chemical reactions. The carrier gas in CVD is most often hydrogen, nitrogen or argon, or mixtures of these. The majority of CVD processes are thermally activated by applying temperatures typically in the range 200-2,000 °C, although there are examples of CVD and ALD processes at lower temperatures, even down to room temperature, and at higher temperatures up to 2,500 °C. There are also CVD and ALD processes that use a plasma to activate the chemistry by opening up new reaction pathways, by electron impact collisions [5] and by the generation of ions and radicals, and these processes are referred to as plasma-enhanced CVD (PECVD) or alternatively plasma-assisted CVD (PACVD). The gas phase chemistry can also be activated by photons from a laser, referred to as laser-enhanced CVD (LECVD) or photo-assisted CVD [6].

CVD may indeed be regarded as a chemical process that spans many traditional disciplines: chemical physics of gases and plasmas, surface science, solid-state chemistry of inorganic materials and organometallic or organic chemistry for precursor synthesis. The field of CVD has been the subject of a number of books and book chapters [7–11], and several review articles on CVD [12–17] and ALD [3, 4, 18–22] have been published over the last years, providing both a detailed background to the processes and overviews of the current research frontier. The scope of this paper is limited to a brief overview of CVD, so as to put into perspective the use of theoretical studies, including those featured in this special collection of papers on theoretical chemistry for CVD.

2 A brief history of CVD

In the history of CVD, J.M. Blocher is often mentioned as a father of modern CVD, since he suggested that thin film deposition processes based on chemical reactions should be distinguished from those based on physical processes. He proposed this at the symposium of the Electrochemical Society (ECS) 1960 in Houston and since then the terms CVD and PVD are used [23]. Blocher also significantly developed the understanding of CVD, notably by summarizing the factors for structure/property/process relationships for CVD films [24]. Refs. [23, 25] and [26] provide a more complete account of the history of CVD.

The first CVD processes were reported and patented in the late nineteenth century and were used for the production of carbon powder for color pigment and of carbon fibers for filaments in early versions of the electrical lamp. CVD of metals was also reported in the mid- and late nineteenth century; one of the earliest examples is the deposition of tungsten from WCl₆ in a hydrogen atmosphere reported by Wöhler in 1855 [27]. Some years later, the famous Mond process for deposition of Ni was reported [28, 29]. This process was developed to purify nickel ore by transforming it to nickel tetracarbonyl [Ni(CO)₄] at low temperatures and from this molecule deposit a film of nickel on a substrate at a higher temperature.

As early as 1909, CVD of silicon was reported from $SiCl_4$ in hydrogen [30]. This process is still used to produce pure silicon for industry, although somewhat refined to allow for greater control in the process. This is the first example of CVD being used to deposit a film of a semiconducting material and looking back in time it is clear that the microelectronics industry and CVD processes have developed hand-in-hand. The electronics boom, and associated drive toward miniaturization, has been pushing the development of CVD processes toward higher-quality films over larger areas with better run-to-run reproducibility and uniformity. An important advance for the CVD of electronic materials was the development in the late 1960s of metal-organic CVD (MOCVD), where a metal is rendered volatile by surrounding it with organic ligands. The first MOCVD process was used to deposit GaAs and was reported in 1968 [31]. The use of metal-organic precursors such as trimethylgallium [Ga(CH₃)₃, also known as TMG] has thus been a key factor in the development of GaAs and other III-V-based electronics and forms today a cornerstone of III-nitride technology.

The latest major breakthrough in the field of CVD is the development of ALD, which was first patented by Suntola in 1977 [32], although it should be pointed out that significant work was done prior to this in the former USSR (for more details on ALD history see ref [20]). The first commercial application of ALD was in thin film electroluminescent displays (TFELs), where ALD was shown to produce films for the luminescent and protective layers with superior quality compared to the state-of-the-art thin film synthesis techniques of the 1970s [33]. Nevertheless, research into ALD remained at a low level until the late 1990s, when it was identified as a possible process solution for high-permittivity ("high-k") thin films in the electronics industry, specifically in memory devices and transistors. Recently, Intel stated that the use of ALD was a key factor for the successful development of the high-k metal gate transistors that allowed further downscaling of the size of integrated circuit chips [34]. Spurred on by this success, ALD is now becoming a widespread nanofabrication

technique and being applied in a wide variety of industrial sectors.

3 Some applications of CVD

The applications of CVD are numerous, and their impact on today's society is enormous. Here, a few important examples of CVD applications are described to provide a flavor of the impact of CVD on our everyday lives.

3.1 Hard coatings

Metal objects can be found everywhere, and most of them are machined by some cutting operation, e.g., turning, milling or drilling. As an example, if one considers the amount of machined parts in an ordinary car and then considers how many cars are produced worldwide, it is obvious that metal cutting is of great importance for our society. Almost all cutting tools use exchangeable edges, referred to as inserts, made of cemented carbide. The great majority of inserts are coated with a hard, wear-resistant thin film that prolongs their lifetime by several orders of magnitude. Although PVD processes, especially for depositing hard nitride compounds, are emerging, CVD still is the work horse for coating cutting tools. A typical CVD coating for a cutting tool is a multilayer structure consisting mainly of some of the following materials: TiN, TiC, TiC_{1-x}N_x, α -Al₂O₃ and κ -Al₂O₃. These film stacks are deposited in a single deposition process, typically at around 50 mbar and 1,000 °C, and a coating batch consists of several thousands of cutting inserts. Importantly, the properties of the individual films can be controlled with great precision [35].

3.2 Functional coatings on glass

Considering total area of deposited film, the largest application of CVD is to coat window glass. One of the most important reasons for coating a window is to prevent heat passing through, reducing the need for cooling down or warming up buildings, and thus reducing energy consumption. Typically films of transparent SnO₂:F are used for this application. To alter the darkness of the window via electrochromism or thermochromism, films of WO3 or VO₂, respectively, are used. The coating can also reflect some of the incoming light, and for this, TiN films are employed. These films are deposited on the glass as a final production step by an atmospheric pressure CVD technique mounted on the float glass production line. The technique was developed by Pilkington in the mid 1980s. A recent development is to coat window glass with transparent TiO₂, making the window self-cleaning by breaking down dirt via photocatalysis with sunlight [36].

3.3 Microelectronics

It is fair to say that without CVD we would not have the electronics that we take for granted today. All sorts of electronic devices are constructed from stacks of thin layers with highly controlled electrical properties, and CVD is often the method of choice for depositing these thin layers. High process temperature is often not an issue for Si or III-V materials that form the bedrock of most of our everyday electronics devices, as well as for the emerging high-frequency and high-power electronics and light-emitting diodes based on silicon carbide (SiC) and III-nitrides. Therefore, CVD processes with process temperatures above 1,000 °C can be used; these processes are carried out close to thermodynamic equilibrium and do not suffer from particle bombardment. The film quality is therefore generally very high, with few defects in the films. The alternative to CVD would be PVD, which is done further from thermodynamic equilibrium and often with a substantial amount of particle bombardment which gives rise to crystal defects. As mentioned above, ALD-grown high-k dielectric films have proven to be vital for a new generation of nanometer-scale transistors [34], where standard CVD is unable to deliver the required quality and uniformity at the thickness scale of just a few nanometers. Now that the utility of ALD in the semiconductor industry has been proven, it is being targeted for the deposition of a variety of materials in ultra-thin layers, particularly as interface control and three-dimensional structures become more important with continued down-scaling.

3.4 Gas permeation barriers

Among all deposition techniques, ALD is unique in enabling nanometer-thin, pinhole-free films that are conformal over features at all length scales. By a happy coincidence, one of the most successful ALD processes across a wide temperature range is that of Al₂O₃, which is highly impermeable to oxygen gas and water vapor. It is therefore possible to use ALD to coat a variety of objects with an Al₂O₃ coating that is impermeable to these gases, while also being so thin (on the order of 10 nm) that the optical and mechanical properties of the object are almost unaffected. Examples include reduced CO₂ permeability through ALD-coated PET bottles [37] and moisture/oxygen diffusion barriers for organic light emitting diodes in flexible display technology [38].

4 Chemical processes in CVD

Thin film growth by CVD is the result of a complex sequence of chemical reactions. All CVD processes

Fig. 1 Schematic summary of the most important chemical reactions involved in thin film synthesis by CVD, after [41]. It should be noted that not all types of chemical reactions are desired in all types of CVD processes



involve surface chemical processes and most CVD processes, with the exception of pure ALD processes, involve also gas phase chemical reactions. The types of chemical reactions that have been recognized as playing a role in CVD are schematically summarized in Fig. 1. Such a conceptual scheme is generally the starting point for building theoretical models of the process. It is therefore important to test the relative importance of these chemical processes in the deposition of a given material system.

An important feature of CVD is that a boundary layer develops above the substrate surface, also known as stagnant boundary layer. The development of this layer is a consequence of the fluid dynamics when flowing a gas mixture above the surface [11]. In the boundary layer, the velocity of species in the flow direction is significantly lower than that in the main gas stream and the concentration of species differs substantially compared to the main body of the gas stream. It is generally considered that all chemistry of importance to the CVD process takes place in the boundary layer and on the substrate surface.

In most, but not all, CVD processes the precursors undergo gas phase chemical reactions that result in the formation of more reactive species. The reactions may be activated thermally or by an external source of energy, e.g., application of a plasma. Generally these more reactive species are smaller fragments of the original molecule and when a plasma is used to activate the chemistry the molecular fragments can even be radicals or ions. One should bear in mind here that many CVD processes are done at low pressures and high temperatures, allowing a significant lifetime of very reactive species, compared to many other chemical reaction environments in, e.g., a liquid solution. It should be mentioned that for some CVD chemistries, the gas phase chemistry produces larger complexes of several precursor molecules that after some molecular re-arrangements are the species active in film deposition [39]. When hydrogen is used as carrier gas, it often takes part in the gas phase reactions, but even relatively inert nitrogen gas can in some thermally activated CVD processes function as both carrier gas and film precursor. A mixture of hydrogen and nitrogen is then used as carrier gas, and only a very small fraction of the nitrogen molecules react to form the film.

The use of plasma to activate the gas phase chemistry, as in PECVD, opens up several new reaction paths at significantly lower temperatures, mainly by electron impact collisions but also by collisions between plasma ions and precursor molecules, and by ions or radicals reaching the surface [5]. The effect of the plasma is therefore often controlled by the distance between plasma and substrate in the reactor; if the substrate is placed in the plasma or very close to the plasma ("direct PECVD"), ionic species will significantly contribute to the chemistry, both in the gas phase and on the surface. A longer distance between plasma and substrate ("remote PECVD") leads to more significant contribution from radical species rather than ions. It should be noted also that the power supplied to the plasma discharge will change the amount of ions in the plasma and thereby also the ionic contribution to the film growth chemistry [40]. In PECVD, dinitrogen gas is often used as nitrogen precursor since the molecule rather easily dissociates in the plasma.

The precursor molecules or the more reactive fragments of the precursor molecules are transported via gas diffusion through the boundary layer to the substrate surface. The adsorption mode depends critically on the chemical properties of the incident precursor or fragment and of the surface. All precursor molecules can be expected to weakly physisorb at least for a short time, followed by desorption, since CVD reagents are volatile by design. If there is sufficient chemical complementarity between precursor or reactive fragment and surface, stronger chemisorption is possible, leading to longer lifetimes of the adsorbate. Some adsorbates react rapidly on the surface (e.g., dissociative chemisorption), either because bonds within the adsorbate are weakened or because of co-reagents on the surface. ALD is based on the elimination of ligands from the adsorbate through reaction with co-adsorbed remnants from a prior co-reagent pulse.

Adsorbate molecules or fragments may be mobile on the surface and may sample a variety of surface sites via diffusion, which again may lead to reactions. In general, surface diffusion is the process that leads ultimately to formation of a film of the target material. Finding the optimum adsorption site via diffusion is particularly important in the growth of epitaxial thin films for electronics since the crystalline quality of the film determines the performance of the final device. Finding optimum sites is facilitated by a strong energetic driving force toward forming the target material from the reagents. In ALD, this has been described as "densification," whereby the coordination number of the constituent atoms increases from the low value of molecular precursors to the high value of the solid product [42, 43].

An important aspect of thin film deposition is the growth mode of the film on a given substrate. Fragments may aggregate into nuclei or islands, or may attach to an atomic step, or may favor a uniform coating. Roughness will increase if subsequent growth is favored at islands. By contrast, well-behaved ALD processes give conformal films that exactly follow the roughness of the substrate. Ultimately the growth processes combine and dictate the larger-scale morphology of the as-grown film (epitaxial crystal, polycrystalline or amorphous), but the factors determining this are not in general well understood [4].

The parts of the precursor molecule (e.g., the ligands) that do not constitute the target material should then desorb from the film surface as by-products. These by-products can be simple molecules formed by the chemical reactions on the surface, e.g., H_2 , CO_2 or Cl_2 , or can be large molecules derived from the ligands. For example, a protonated ligand is the by-product in thermal ALD of metal oxides using H_2O as co-reagent. In any case, it is of great importance to all CVD processes that these by-products desorb cleanly from the film surface and are not incorporated as impurities in the film.

5 The use of theoretical studies development and understanding of CVD processes

The main motivation for theoretical chemical studies of CVD processes is obtaining a detailed understanding of the chemical reactions summarized in Fig. 1. There are various outcomes from obtaining this understanding, as shown in the following examples.

5.1 A better understanding of established CVD processes

Thermochemical modeling of the gas phase chemistry in CVD is a convenient approach for understanding the gas chemistry, given that not many experimental techniques are available. Spectroscopic technique, e.g., Raman or FT-IR spectroscopy, could in principle be used but this requires that all molecules in the gas phase can be detected by these techniques and that the CVD reactor can be fitted with viewports of a material that is both transparent for the excitation light and at the same time not affected by the CVD gases and process temperatures. A mass spectrometer measuring the gas flow could also be used to study the process chemistry; the problem with this is that only the stable molecules are detected while all unstable intermediates are lost. An example of a CVD process where much understanding has been provided by thermochemical modeling is CVD of electronic grade SiC. Significant effort has been directed toward understanding the process, and a gas phase and surface reaction model has been proposed [44]. Using this model, the CVD process can be discussed based on the gas phase and surface chemistry, rather than susceptor design and gas flow patterns. Modeling can then be used to predict growth rate, etching rate, surface morphology and doping, all key aspects when doing CVD of electronic device structures. Detailed gas phase chemistry models can also be obtained by quantum chemical calculations, an example is the gas chemistry model for CVD of boron carbide from BCl₃ and C₃H₈ involving 16 intermediates presented in ref [45].

Consider the CVD of Al₂O₃ from AlCl₃ and CO₂, which has mainly been developed by research departments in the hard coatings industry. CO2 forms H2O and CO together with the hydrogen carrier gas in a gas phase reaction in the reactor, and H₂O then acts as the oxygen precursor in the formation of Al₂O₃. The catalytic effect of H₂S in this process has been known for many years. The addition of H₂S leads to a higher deposition rate and more surfacecontrolled deposition process as evidenced by more uniform coating thickness on complex substrate geometries [46]. But since this process was mainly developed in industry, where detailed understanding often has low priority, it was only lately that a surface chemical model supported by careful theoretical chemistry modeling could be presented [47]. It was suggested that H_2S acts as a true surface catalyst by facilitating first the removal of adsorbed chlorine from the surface and then the adsorption of H₂O. Another example where modeling has been able to explain details in CVD of hard coatings is the observation that thin films of titanium carbide grow along the (111) direction when aromatic hydrocarbons such as benzene are used, while aliphatic hydrocarbons, e.g., methane, produce growth in the (100) direction. This is of interest since (111)TiC performs better in cutting operations [48] and since control of the preferred TiC growth direction enables control of the preferred growth direction of alumina deposited on top of the TiC layer [49]. The explanation for the change in preferred growth direction with aromatic hydrocarbons, provided by quantum chemical calculations, was that benzene chemisorbs significantly more strongly on the (111) surface compared to the (100) surface and reduces the surface energy of the (111) surface [50]. Methane did not show any significant preference for either surface, so the lower surface energy of the (100) surface controls the growth direction when aliphatic hydrocarbons are used. Although these findings on the surface chemistry are not likely to change the well-established and optimized CVD process for industrially used hard coatings, such deeper understanding of the chemistry can serve as a guide for design of future CVD processes.

5.2 Using modeling for designing new CVD processes

Theoretical studies of CVD surface chemistry that is more directed toward designing a new CVD process for a material are the theoretical studies of CVD of cubic boron nitride (c-BN) onto diamond surfaces. Models are developed of both the initial growth stage onto a diamond surface [51] and the subsequent growth of c-BN onto c-BN [52, 53], including surface reconstructions induced by adsorption of precursor molecules onto c-BN [54, 55]. The authors of these studies boil down their results to a suggested ALD chemistry for deposition of c-BN [52], although it remains to be seen whether it will be tested given the large amount of HF produced as by-product.

When developing precursors for new CVD processes, modeling can give a better understanding of the behavior of the molecules. It is especially important to understand how the molecules behave upon vaporization and in the vapor phase. One such example is the development of CVD precursors for metal boride films of the actinides and lanthanides [56, 57], where DFT modeling provided a better understanding of why the same type of organometallic molecule works as CVD precursor for Sm, Pr and Er but decomposes instead of subliming for U [58].

Approaches exist for the simulation of gas–liquid equilibria for small molecules [59], which allow the vapor pressure to be predicted. Extending these to metalorganic molecules requires accurate estimation of both metal– ligand bonding and intermolecular interaction, and therefore requires an approach beyond standard density functional theory. Simulating hundreds of molecules at such a high level is currently beyond routine computational capacity.

5.3 ALD development and understanding

Given its great importance for the electronics industry, ALD is today the field of CVD where perhaps the largest amount of effort is being directed toward increased understanding of the process chemistry. Research on ALD is somewhat simplified by the surface-controlled nature of the process—at least in the ideal case—so that growth should be independent of other factors such as gas phase chemistry and gas flow in the reactor. However, the stringent and partially contradictory requirements for achieving ALD make it very difficult to design a new process, even when theory and experiment work closely together.

Since the earliest days of ALD research, atomic-scale models have been made of the underlying chemistry. Pioneering calculations were carried out on the zinc sulfide system by Pakkanen and Lindblad in the 1980s [60, 61]. Over 150 publications modeling ALD have followed since then, mostly carrying out quantum chemistry on the reactions of precursors with model substrates and many of these concerned with highpermittivity dielectrics on silicon. The state of the art to 2012 is summarized in ref [62]. Since then, our most recent atomicscale simulations have revealed new details of the mechanism of oxide ALD [43], have accounted for the stoichiometry of ternary oxides [63], have assessed precursor ligands for alkaline earth metal oxides [64] and for copper metal [65] and have developed kinetic Monte Carlo models of film growth [66].

The ALD process is manifest across many length scales. The pulsed flow of gases into meter-scale reactors, around millimeter-scaled geometries, leads to chemical reactions between atoms, which grow into nanometer-thick films and coat micronscaled pores or particles. It is clearly impossible to describe explicitly all of these length scales in one model, and most simulations are "multi-scale" insofar as they involve coupling between selected length scales according to the property of interest. The problem of timescale in ALD is perhaps even more acute than that of length scale, since a combination of fast (psns) and slow reactions (µs-ms) contribute to film growth, and gases are pulsed and purged over second-long timescales. Wellbehaved ALD processes are independent of reactor conditions, and so modeling ALD chemistry exclusively at the atomic scale can in fact explain many features of film growth, without reference to reactor conditions. Nevertheless, explaining ALD's unique selling point-conformal and uniform growth regardless of the size and shape of the substrate-requires modeling at higher length scales, ideally incorporating information about the reaction chemistry from the atomic scale.

6 An overview of the special collection of papers on CVD and ALD modeling

This overview strives to aid readers with little or no experience in CVD and ALD and to put this special

collection of papers on CVD and ALD modeling into a broader perspective. The special collection has been edited by the authors of this overview who also invited wellknown researchers in the field of CVD/ALD modeling to submit original research manuscripts for the special collection. The collection consists of nine papers on both CVD and ALD modeling, summarized below.

Two papers strive to shine light on the CVD process for epitaxial film growth of SiC for manufacturing of highpower electronic devices. The first paper by Danielsson et al. [67] points to several weaknesses of the thermochemical modeling of SiC CVD heavily used in process development today. One of the main weaknesses is the suggested underestimation of the surface reactivity of hydrocarbons in today's modeling. Also, the great influence the quality of the molecular data used has on the continuous flow dynamics (CFD) simulations is pointed out by using state-of-the-art quantum chemical calculations to provide more accurate molecular data. The other paper on SiC CVD is a quantum chemical approach by Kalered et al. [68] to better understand the effect of using a chlorinated growth chemistry in SiC CVD. The use of chlorinated growth chemistry also for electronic grade SiC has enabled growth rates of 100 μ m/h or higher [69], but the effect on the surface chemistry when adding the chlorine has not been investigated before. It is shown in the paper that SiCl₂, the main Si species for a chlorinated chemistry, once chemisorbed has a higher energy barrier for desorption compared to SiH₂, the main Si species for the standard chemistry. It is therefore suggested that the chlorinated chemistry can bring a more favorable surface chemistry to SiC CVD.

An industrial CVD process where chlorinated chemistry is well established is CVD of α -Al₂O₃ for hard coatings. Århammar et al. [70] used a combination of thermochemistry and first principles modeling to study possible point defects incorporated in the α -alumina films from the CVD gas mixture. Besides Al and O, the gas mixture for alumina CVD contains H, C, Cl and S, but little is known on the solubility of the possible point defects that can be created from these atoms. It is found that especially interstitial hydrogen and chlorine have the lowest formation energies and are thus likely to form, whereas sulfur from H₂S, which act as catalyst in the process [47], is not likely to incorporate in α -alumina.

Boichot et al. [71] used thermochemical modeling to better understand a newly developed high-temperature CVD process for aluminum nitride (AlN). The focus of their modeling were the chemical conditions in the boundary layer and close to the surface—parameters very hard to extract from experiments. The modeling results were used to design validation experiments presented in the paper. Even though the authors can deduce some experimental parameters needed for deposition of highquality AlN and further reactor development, they also state that the modeling used fail to give the whole picture since surface diffusion mechanisms are not included.

CVD is the standard technique for synthesizing thin films of diamond, a topic studied by Yiming et al. [72] using DFT. Particularly, the authors study the experimentally reported significant increase in growth rate of diamond when adding nitrogen to the CVD gas mixture; however, from experiments, it is not clear whether it is nitrogen incorporated into the diamond lattice or nitrogen, in the form of N, NH or NH2, chemisorbed to the diamond surface that leads to the higher growth rate. The authors conclude from their studies that nitrogen incorporated into the diamond lattice one or two atomic layers beneath the surface gives a higher surface reactivity leading to a higher growth rate. The authors estimate that the growth rate on the (100) surface of diamond will increase with a factor of 3.7, which is well in agreement with experimental findings.

Lin and Teplyakov [73] used density functional theory to explore the reactions of two representative precursors with silicon surfaces terminated with various functionalities (Si-H, Si-N, Si-O), yielding an impressive comparison across 32 reaction pathways. The use of small Si clusters as substrate models is discussed. The results for the trimethylaluminum (TMA) precursor illustrate how relatively simple C-Al and C-H dissociation processes are affected by the silicon termination. Extending this to tetrakis(dimethylamido)titanium introduces more complex chemistry and provides an opportunity to decouple electronic and steric factors. Molecular adsorption is seen to depend strongly on the electron-donating properties of the substrate, consistent with frontier orbital analysis. For proton transfer from substrate to ligand (the dominant ALD reaction), all activation energies relative to molecular adsorption are computed to be >50 kJ/mol, with the exception of three pathways involving the Ti precursor and an O-terminated silicon surface.

The reaction kinetics of ALD are described analytically and numerically in this collection by Travis and Adomaitis [74]. Taking alumina deposition from TMA as the sample system, a model describing the reaction kinetics and surface species dynamics for the TMA half-reaction is developed. Four TMA reactions are included, accounting for TMA adsorption and subsequent reaction on a range of growth surfaces spanning bare to fully hydroxylated states. Published reaction energetics for these reactions from quantum chemical studies are converted into kinetic parameters using statistical thermodynamics and absolute reaction rate theory. The cyclic nature of ALD can be exploited to determine the overall growth rate from the concentrations of surface species. Interestingly, the model reveals that substantial growth of alumina is possible regardless of the level of hydroxylation of the substrate, as there are two competing pathways with favorable kinetics. The reaction rate expressions developed here can be implemented into reactor-scale simulations, which can guide the optimization of commercial ALD processes.

An approach toward precursor design from first principles is outlined by Dey and Elliott [75] in this collection. The subject is the ALD of metallic copper, where of course the precursor combination must provide both copper cations and electrons for reduction to the metal. Here, it is proposed that the electron source should also contain copper, thus removing the chance of contamination by other metals, but that this is pulsed alternately with a "standard" copper source in 2-step or 3-step ALD. The hydride anion is proposed as the reducing agent, as this may provide both electrons for reduction and protons for elimination of ligands from the standard copper precursor. A copper carbene complex is computed to yield maximal activation of the hydride and lead to volatile by-products, and so the proposed precursor is 1,3-diphenyl-4,5-imidazolidinedithione copper hydride.

The paper in this collection by Yangas-Gil and Elam [76] highlights that simple first-order Langmuir kinetics is sometimes insufficient to capture the complexities of ALD surface chemistry, especially when precursor exposure is high, but that understanding such non-ideal cases is crucial for optimization and reliability of commercial ALD processes. The authors' approach is to cast gas transport inside nanostructures as a single particle discrete Markov chain process, decoupled from reaction mechanisms that can include branching and non-ideal surface kinetics as needed. This description of surface chemistry is applicable generally to all types of CVD, not just ALD. A link is established between ballistic models, kinetic Monte Carlo simulations, and continuous models based on the use of the diffusion equation under Knudsen conditions. The calculations confirm that ALD coverage depends solely on total exposure as long as reaction probabilities are pressureindependent.

7 Concluding remarks

There is substantial variation in our level of understanding of CVD and ALD processes: some processes are investigated in great detail and are well understood, while others have been tuned to perfection over many years of development and/or commercialization, without much understanding of the details. In this overview, we have tried to point out the effectiveness and versatility of theoretical chemistry as a tool to obtain a deeper understanding without experiments that require highly complex equipment and conditions far from what is used in the actual deposition process. The importance of CVD and ALD processes for our everyday lives is enormous, and the continuing development of, e.g., electronics requires constantly better CVD and ALD processes. Theoretical chemistry modeling will play a key part in this development.

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