

OPEN ACCESS

Atomic Layer Etching at the Tipping Point: An Overview

To cite this article: G. S. Oehrlein et al 2015 ECS J. Solid State Sci. Technol. 4 N5041

View the article online for updates and enhancements.

You may also like

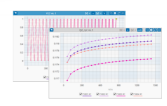
- Electrochemical Atomic Layer Etching (e-ALE) of Copper - Thermodynamic and Diffusion-Reaction Considerations of the Surface-Limited Sulfidization of Copper Yukun Gong and Rohan Akolkar
- Challenges of Tailoring Surface Chemistry and Plasma/Surface Interactions to Advance Atomic Layer Etching S. U. Engelmann, R. L. Bruce, M. Nakamura et al.
- (Invited) Atomic Layer Etching Using Thermal Reactions: Atomic Layer Deposition in Reverse Younghee Lee, Jaime W. DuMont and Steven M. George

PAT-Tester-x-8 Potentiostat: Modular Solution for Electrochemical Testing!



- ✓ Flexible Setup with up to 8 Independent Test Channels! Each with a fully equipped Potentiostat, Galvanostat and EIS!
- Perfect Choice for Small-Scale and Special Purpose Testing!
 Suited for all 3-electrode, optical, dilatometry or force test cells from EL-CELL.
- ✓ Complete Solution with Extensive Software! Plan, conduct and analyze experiments with EL-Software.
- Small Footprint, Easy to Setup and Operate!
 Usable inside a glove box. Full multi-user, multi-device control via LAN.







Contact us:

- +49 40 79012-734
- sales@el-cell.com
- www.el-cell.com





JSS Focus Issue on Atomic Layer Etching and Cleaning

Atomic Layer Etching at the Tipping Point: An Overview

G. S. Oehrlein, a,z D. Metzler, a and C. Lia,b

^aDepartment of Materials Science and Engineering, and Institute for Research in Electronics and Applied Physics, University of Maryland, College Park, Maryland 20742, USA

The ability to achieve near-atomic precision in etching different materials when transferring lithographically defined templates is a requirement of increasing importance for nanoscale structure fabrication in the semiconductor and related industries. The use of ultra-thin gate dielectrics, ultra thin channels, and sub-20 nm film thicknesses in field effect transistors and other devices requires near-atomic scale etching control and selectivity. There is an emerging consensus that as critical dimensions approach the sub-10 nm scale, the need for an etching method corresponding to *Atomic Layer Deposition* (ALD), i.e. *Atomic Layer Etching* (ALE), has become essential, and that the more than 30-year quest to complement/replace continuous directional plasma etching (PE) methods for critical applications by a sequence of individual, self-limited surface reaction steps has reached a crucial stage. A key advantage of this approach relative to continuous PE is that it enables optimization of the individual steps with regard to reactant adsorption, self-limited etching, selectivity relative to other materials, and damage of critical surface layers. In this overview we present basic approaches to ALE of materials, discuss similarities/crucial differences relative to thermal and plasma-enhanced ALD, and then review selected results on ALE of materials aimed at pattern transfer. The overview concludes with a discussion of opportunities and challenges ahead.

© The Author(s) 2015. Published by ECS. This is an open access article distributed under the terms of the Creative Commons Attribution Non-Commercial No Derivatives 4.0 License (CC BY-NC-ND, http://creativecommons.org/licenses/by-nc-nd/4.0/), which permits non-commercial reuse, distribution, and reproduction in any medium, provided the original work is not changed in any way and is properly cited. For permission for commercial reuse, please email: oa@electrochem.org. [DOI: 10.1149/2.0061506jss] All rights reserved.

Manuscript submitted February 2, 2015; revised manuscript received March 12, 2015. Published March 27, 2015. *This paper is part of the JSS Focus Issue on Atomic Layer Etching and Cleaning.*

A requirement of increasing importance for nanoscale device fabrication is the ability to achieve atomic scale etching control and materials selectivity during pattern transfer. ^{1–8} An etching method corresponding to *Atomic Layer Deposition* (ALD), i.e. *Atomic Layer Etching* (ALE), is expected to satisfy these needs as critical dimensions continue to shrink below the 10 nm scale.

Demonstrations of self-limited dry etching methods capable of near-atomic resolution have a long history in the dry etching community and a brief review will be presented below. Key challenges for these approaches have been specialized equipment, long process times and low throughput. 9,10 However, a recent demonstration using a commercial plasma etch tool¹⁰ and activities within the dry etching community¹¹ provide indications that the situation has changed, and that we may have reached for ALE the Tipping Point which Gladwell defined as the "the moment of critical mass, the threshold, the boiling point" when "ideas and products and messages and behaviors spread like viruses do". 12 This is due to several factors, including unprecedented demands on dry etching technology introduced by the semiconductor device evolution according to Moore's law that can be satisfied by atomic layer etching, advanced capabilities in plasma etch, and the existence of a critical level of information on plasma etch and ALD methods as applied in the semiconductor fabrication space.

In this article we will provide a review of background of these approaches, and focus on issues that have to be overcome for wide-spread implementation in manufacturing.

Technology Demands on Plasma Etch (PE) and Key Shortcomings of Continuous PE Methods

Technology demands.— Current semiconductor manufacturing is characterized by the need to mass produce features that are approaching 10 nm critical dimension (CD) and require CD variation of 0.5 nm or less. ^{1,3} The use of ultra-thin gate dielectrics, ultra thin channels, and overall decreasing film thicknesses in combination with more stringent demands on surface property control in field effect transistors, i.e. preventing materials damage, requires control over etching direction-

ality and materials selectivity that approaches the atomic scale. ¹³ Additionally, the material stacks making up devices are becoming more complex and exhibit higher aspect ratios. Examples are 3-dimensional gate etch applications which demand essentially infinite etch selectivity while avoiding introduction of materials damage as FinFETs, Trigates, nanowires and other 3D devices are produced. ^{14–16} For these applications the top of the fin/wire is exposed to the plasma and needs to withstand plasma exposure while the remaining gate is formed around the fin/wire.

The potential of graphene technology has introduced the challenges associated with fabrication of single atomic layer—based technologies. ^{17,18} Gate formation on single atomic layer materials such as graphene requires the ability to stop on a single layer with atomistic precision. Patterning of graphene sheets (ribbons) for digital/logic applications requires line edge roughness (LER) control of <1 nm so that the graphene exhibits semiconducting behavior. ¹⁹

Shortcomings of continuous plasma etching technology.— It will be difficult to meet the above demands of nanotechnological manufacturing using continuous plasma processing approaches for which complex process recipes are used to optimize the achievement of certain process objectives. Admitting all reactants simultaneously to the process chamber gives rise to large particle fluxes at certain surface locations through all phases of the plasma process and complex parallel reactions that can evolve with the long-time transients associated with plasma-chamber wall interactions. ^{3,6} For instance, etching selectivity for the prototypical case of fluorocarbon based etching of dielectrics is based on several parallel reactions, which for compositionally distinct materials can lead to different net overall reaction rates, i.e. fairly rapid etching for one material and slow etching or deposition for another material.²⁰⁻²⁶ However, the thicknesses of these steady-state surface layers can be of the order of several nm. During the time needed for these to form, significant material loss can take place.²⁷ Thick modified surface layers develop on semiconductor and dielectric surfaces that are simultaneously exposed to significant chemical reactant fluxes and ion bombardment. To achieve silicon etching directionality in plasma etching for these reactant-rich process conditions, often O2 is added to chlorine- or bromine-based discharges to enhance

^bDepartment of Physics, University of Maryland, College Park, Maryland 20742, USA

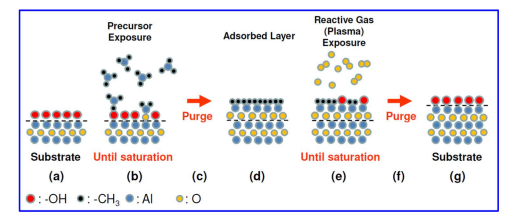


Figure 1. Schematic of one cycle of a typical ALD process to produce thin Al_2O_3 layers (redrawn after Potts and Kessels³⁷). The surface modification step ((b)-(d)) is based on self-limited adsorption of a reactive precursor, followed by pump-out. The steps ((e)-(f)) involve exposure to an oxidizing gas, e.g. water, O_2 plasma, etc, followed by pump-out and transform the precursor layer into roughly a monolayer of the final Al_2O_3 material. For the substrate (g) obtained after one ALD cycle, the steps (b) through (f) will be repeated. The overall deposited layer thickness is produced by n ALD cycles.

oxidation of vertical silicon surface features and prevent lateral attack which, however, changes critical dimensions of devices.²⁸ Additionally, profile imperfections, e.g. micro-trenching, can result.

These limitations have resulted in significant developments on pulsed discharges for plasma etching which are promising in overcoming some of these issues.³⁻⁶ An alternative and possibly complementary approach corresponds to the reverse of atomic layer deposition technology.

Atomic Layer Deposition and Atomic Layer Etching – General Principles

Atomic layer deposition (ALD).— Atomic layer deposition has become the method of choice for highly conformal coatings in many applications, including advanced semiconductor processing. ^{29–31} The ability to control the thickness of deposited films near one monolayer per process cycle is based on careful choice of chemical precursors which, once adsorbed at one monolayer, passivate the surface and prevent multi-layer adsorption (see Fig. 1). This is followed by a reaction step which transforms the precursor on the heated substrate into the desired material. An example is the deposition of an Al metallorganic monolayer which upon oxidation is transformed into Al₂O₃. ^{32–35} Deposition conformality is a key objective of many ALD processes. It can be successfully achieved even for very challenging geometrical situations, e.g. high-aspect ratio trench or via structures, since self-limited adsorption is insensitive to local variations in incident particle fluxes at surface locations.

For thermal ALD processes the activation energy for the chemical reaction that takes place during the reaction step is provided by substrate heating. Thermal ALD rates drop off at low substrate temperature when the thermal energy becomes insufficient to drive the chemical reaction. The possible substrate temperature range over which an *ALD window* exists is limited at low temperature by incomplete reaction during the reaction step or multi-layer condensation during precursor deposition²⁹ (see Fig. 2). The ALD window is limited at high substrate temperature by thermal decomposition of precursors, thermal desorption, and other loss processes.²⁹

Plasma-enhanced ALD.— Recently, there has been strong growth in plasma-enhanced ALD that uses plasma-generated radicals/energetic species during the reaction step to enable or speed up chemical reactions with the deposited precursor layer that are either too slow or not possible with just thermal energy and typical reactants. The reviews^{36–38} describe key features of plasma-enhanced ALD, and differences relative to thermal ALD. Plasma-assisted ALD offers greater processing flexibility relative to thermal ALD, including a larger number of precursors that may be used, the opportunity to use lower substrate temperatures than possible in thermal ALD

while maintaining growth per cycle, and an increased range of materials that may be deposited along with control of materials properties. One limitation of plasma-assisted ALD is reduced conformality (step coverage) relative to thermal ALD when coating substrates with pronounced surface topography, ³⁶ e.g. high aspect ratio trenches or holes. This is explained by the highly reactive species created in the plasma environment which make the achievement of self-limitation more difficult, i.e. it has been observed that deposition on passivated surfaces can take place more easily in the plasma environment. 36,39 The result is that process rates may become more strongly dependent on the locally available chemical precursor flux, and may be controlled by radical recombination loss probabilities in deep trenches, ³⁹ and/or energy flux, in contrast to ideal ALD processes. Profijt et al.³⁶ also point out additional concerns for plasma-based ALD processes, e.g. vacuum ultraviolet (VUV) induced electrical damage to insulators which is absent in a purely thermal environment. These features of plasma-enhanced ALD are expected to be also very important for plasma-based atomic layer etching methods.

Atomic layer etching (ALE).— Similar to ALD, realization of ALE has long been based on replacing the complex plasma-surface interactions of steady-state plasma etching by a sequence of individual, self-limited surface reactions (see Fig. 3). In a first reaction step, a chemical precursor is introduced into the reactor and adsorbed at the surface of the substrate. The precursor is chosen so that upon reaction with substrate atoms volatile products can be formed. Conditions must be chosen so that the precursor does not spontaneously etch the substrate, e.g. by lowering the substrate temperature. Ideally, reactants are present at the active surface at about a monolayer. Subsequently, the chamber is exhausted to remove remaining chemical reactants.

In a second surface reaction step, bombardment of the surface with energetic species, typically a beam of low energy ions, provides the necessary energy to induce chemical reactions between the adsorbed

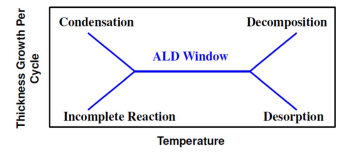


Figure 2. Schematic illustrating the thermal "ALD window" concept (redrawn after George²⁹).

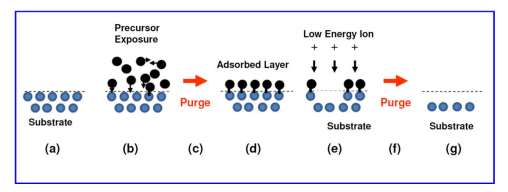


Figure 3. Schematic of one cycle of a typical ALE process. The surface modification step ((b)-(d)) may include self-limited adsorption, short deposition, etc, followed by pump-out. Low energy Ar ion bombardment is often used for selective removal of reacted region ((e),(f)). For the substrate (g) after one ALE cycle, the steps (b) through (f) will be repeated. The overall etch depth is produced by *n* ALE cycles.

species and the substrate. Other methods employed for inducing a reaction of adsorbed species with substrate atoms to produce volatile products are bombardment with fast neutral atoms, ^{40–42} electrons, ⁴³ or irradiation with photons. ⁴⁴ (Isotropic atomic layer etching based on thermal product desorption has been described, ⁴⁵ but these thermal approaches necessitating elevated substrate temperatures will not be discussed extensively in this review.) To minimize physical sputtering and surface damage, energies for ion or neutral beam bombardment are typically limited to 100 eV or lower. The reaction chamber may be evacuated again to complete one cycle. The ALE process ideally proceeds in a cyclic, self-limiting way, with a substrate thickness loss of about 1 monolayer per cycle.

ALE window.— To control the thickness loss (etch depth) per cycle, self-limited surface reactions are required. Both spontaneous chemical etching by the precursor and physical sputtering should be minimal. The concept of an ALE window located between the spontaneous chemical etching and the physical sputtering regimes is schematically illustrated in Fig. 4 for ion-induced etching. The key parameter for ALE that plays a similar role as substrate temperature in thermal ALD is ion energy. In a similar fashion as there exists an ALD window versus substrate temperature, there is an ALE window versus ion energy. In a plasma environment, ions with an energy distribution are incident on the substrate and control physical sputtering and substrate damage extent. Bombardment of the surface between chemical reactant exposures at a sufficiently high fluence is assumed in Fig. 4. The modification of the surface by the precursor allows the material to be etched with lower activation energy as compared to the underlying material without modification. The presence of chemical reactants at the substrate surface along with ion bombardment induce a chemical reaction between the surface atoms and the precursor which causes

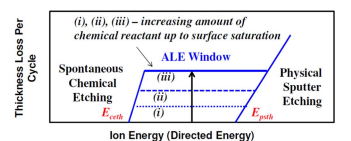


Figure 4. Schematic illustrating the concept of an "ALE window" located between the spontaneous chemical etching and the physical sputtering regimes. The presence of chemical reactant on the substrate surface enables directional etching of the material in a window of ion energies above the chemically enhanced etching energy threshold E_{ceth} and below the physical sputtering energy threshold E_{psth} , respectively. Spontaneous chemical etching of the reactant/substrate system interferes with this approach and is made negligible by reducing the temperature of the substrate sufficiently.

etch products to boil off or sputter from the surface. This process enables directional etching of the material at ion energies above the chemically enhanced etching energy threshold E_{ceth} and below the physical sputtering energy threshold E_{psth} , respectively. By carefully tailoring the energy of ion bombardment, it is possible to control the etching depth to about a monolayer. Spontaneous chemical etching of the reactant/substrate system interferes with this approach and the thermal energy required to drive these reactions is made negligible by reducing the temperature of the substrate sufficiently. While Fig. 4 illustrates this concept for ion bombardment, in the literature ALE windows have also been demonstrated by using electron, fast neutral and photon bombardment of surfaces for directed energy input.

An important question relates to the degree to which ion energy and the width of the distribution of ion energies must be controlled. While average ion energy has been shown to be useful, careful studies of the impact of ion energy distributions on ALE performance are required to answer the question how closely ion energies must be controlled to enable optimal exploitation of the ALE window. Shin et al. 6 investigated nearly mono-energetic ion energy distributions (IEDs) and observed novel phenomena (see below).

Critical neutral exposure and critical dose for energetic particle bombardment.— In order for the etching depth per cycle to be self-limited, the chemical reactant exposure of the substrate has to be sufficiently high to achieve surface saturation, and the dose of low energy particles received during a cycle has to be balanced with this and also be sufficiently high. Figure 5 schematically shows the impact of different chemical reactant exposure conditions, e.g. by changing reactant pressure, on the thickness loss depth per cycle. Ample energy

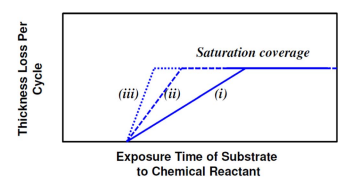


Figure 5. Schematic illustration of the dependence of etched thickness per cycle on exposure time for different pressures ((i) – lowest pressure, (ii) – intermediate and (iii) – highest) and assuming negligible spontaneous etching. Self-limitation requires saturation surface coverage which at the highest pressure (iii) is achieved after the shortest exposure time. Alternatively, a plasma and production of radicals may be used to reduce the time required to achieve saturation surface coverage.

input to the surface by low energy particle bombardment within the ALE energy window at a sufficiently high dose between chemical reactant exposures is assumed. The lower values of these have been termed critical exposure, i.e. pressure * time of chemical reactant, and critical dose (for energetic particle bombardment), respectively. For instance, self-limited etching of HfO₂ at 1 ML/cycle using BCl₃ and energetic Ar neutral beam bombardment required a BCl₃ pressure above a critical pressure of 0.22 mTorr for 20 s BCl₃ exposure and a critical dose for energetic particle bombardment of 1.48×10^{17} atoms/cm², respectively.⁴⁷ Corresponding information has been published for other reactant/materials systems.

One key need driving development of ALE approaches is the achievement of atomic scale etching selectivity with regard to a different material. More generally, owing to the complex nature of surfaces in advanced semiconductor devices, simultaneous etching control over multiple materials is typically required. By working close to the energy threshold for physical sputtering of one material, and exploiting energy threshold differences among different materials, etching selectivity can be optimized.⁹ The threshold energy for physical sputtering E_{psth} depends on the nature of the material and is higher in the case of SiO_2 than for Si, $E_{psth}(SiO_2) > E_{psth}(Si)$. By supplying chemical reactant to the substrate surface, chemically enhanced etching is possible, with an energy threshold $E_{ceth}(SiO_2)$ that is lower than for physical sputtering $E_{psth}(Si)$, which may enable selective etching of SiO₂ over Si. The amount of material etched per cycle will depend on the surface coverage of the chemical reactants up to a saturation coverage. In the schematic of Fig. 4 it is also assumed that in this ALE window between E_{ceth} and E_{psth} the substrate thickness loss per cycle is determined by the reactant coverage and essentially independent of ion energy, which does not need to be the case.

Maintaining etching directionality, achieving dimensional control approaching atomic scale, and leaving materials after ALE damage-free are other essential objectives.

An important advantage of a cyclic ALE process relative to continuous etching is that it provides the opportunity to decouple the reaction steps and through detailed study of each, establish how variations of incident particle parameters (chemistry, energies, etc.) enables product volatilization, self-limiting behavior and protection of lateral and vertical surfaces/underlayers that are consistent with the requirements on the overall process.

Although ALE shows some similarities to ALD with regard to physisorption/chemisorption requirements at the surface, the requirements with regard to "volatile product" removal are fundamentally different. Whereas for ALD, films are grown in a conformal fashion, for ALE the "etch product" removal ideally should take place in a directional fashion. Because of this fundamental difference from ALD, ALE provides profound surface chemistry challenges and energetic species/surface interaction problems that are special.

Atomic Layer Etching of Various Materials - A Brief Survey

As discussed, the study of ALE technology goes back more than 30 years. A multitude of different kinds of approaches have been applied to ALE of GaAs and silicon along with a significant number of other materials. In concert with the large number of approaches and methods, these approaches have been referred to by many different names, including atomic layer etching, molecular layer etching, digital etching, layer-by-layer etching and others. To make it easier to obtain an overview of this literature, we provide in Table I a brief listing of published work on materials and approaches for which atomic layer etching has been investigated using both experimental and theoretical methods. Essential information extracted from the publications is presented below. For halogen-based etching of GaAs, 48-50 silicon 51-56 and other materials there is a large body of atomistic etching work not aimed at pattern transfer and replacement of plasma etching methods. This literature provides important atomic layer etching and surface chemistry background, but is not reviewed. For instance, Maki and Ehrlich⁵⁷ described laser-induced bilayer etching of GaAs at the

Angstrom and sub-Angstrom level in 1989, but their work addressed a different application. Wet etching methods have also not been covered.

Table I lists materials in alphabetical order, and references for a given material are sorted primarily according to the year published. We apologize in advance for mistakes, omissions and redundancies of Table I.

Al₂O₃ and Be-Oxide.— ALE studies on Al₂O₃^{58,59} and Be-oxide on GaAs substrate⁶⁰ were performed by Yeom's group using a neutral beam system that is based on an ICP source, for which accelerated Ar⁺ ions are neutralized by low angle forward reflection from a surface. They used up to 30 s BCl₃ gas exposure for adsorbing chlorine reactants at the surface, followed by evacuation, striking a plasma and bombarding the passivated surface with a neutralized Ar beam at 100 eV. Measured etch depths per cycle were about 1 Angstrom. They also describe results of density functional theory for the interaction of BCl₃ with Al₂O₃ which they thought provided insights that could be exemplary for other ALE systems as well.⁵⁸ For BeO on GaAs negligible sputtering was seen for bombardment energies of less than 130 eV, and a self-limited etch depth of 0.75 Å/cycle at saturation.

III-V: GaAs, InP, InAlAs, InGaAs and others.— Atomic layer etching of III-V materials is among the oldest ALE demonstrations. Meguro et al. in 1990⁴³ used exposure of GaAs to Cl₂ gas and electron bombardment at 100 eV to demonstrate a self-limited iterative etching approach. They achieved about 1/3 ML etching per cycle. This work demonstrated clearly that the etching was limited by the adsorption of the chemical reactant. In related work they also evaluated surface activation using low energy Ar ion bombardment using an electron beam excited plasma. 61,62 They published subsequently additional work using Ar ion bombardment, 63 or photon irradiation of a chlorine-coated GaAs surface. 44,64 Aoyagi et al.65 studied GaAs ALE using alternating Cl₂ exposure and synchronized low energy Ar⁺ bombardment by applying a low bias voltage in an electron cyclotron resonance (ECR) plasma system. The ECR Ar discharge was continuously maintained, and Cl₂ was admitted for times up to 40 s to achieve adsorption of Cl atoms at the GaAs surface. They report self-limited etching of about a monolayer per cycle for a certain exposure time window of the substrate to the chlorine discharge and the low energy Ar ion beam. For extended exposure times (greater than 20 s), the etch depth per cycle decreased which they explained by multi-layer Cl adsorption. Ko et al. 66 also examined layer-by-layer etching of GaAs, InP, GaInAs, and AlInAs using Cl radicals produced by a low power discharge and Ar⁺ ion bombardment (with additional RF bias) produced sequentially in an ECR system. For complete chlorine surface coverage of GaAs achieved by 6 s exposure to an Ar/Cl₂ discharge at 1 mTorr they observed self-limited etching of about 5 A/cycle. For their approach they quote a typical cycle time of 45 s, which consists of 10 s reactive radical adsorption time, followed by pump out of excess radicals (30 s), and a desorption time with Ar ions of 5 s. This shows the strong impact of the purge cycle on cycling time. In another study, Lim et al.⁶⁷ used Cl₂ exposure of GaAs and a Ne neutral beam source and measured about one atomic layer/cycle.

When Ko et al. studied layer-by-layer etching of InP they found that sample heating to 150°C and a higher RF bias than for GaAs was required to observe etching. ⁶⁶ Etching of InP was studied by Otsuka et al. ⁶⁸ using brief exposures to tertiarybutylphosphine, pump-out followed by substrate heating using a halogen lamp for desorption of products. This resulted in etch depths of a fraction of an Angstrom per cycle.

Park et al.^{69,70} studied ALE of InP and InAlAs based on 20 s exposure to Cl₂ at 0.4 mTorr, followed by Ne neutral beam bombardment. They measured roughly 1 monolayer /cycle (1.47 A/cycle) for InP with high selectivity against InAlAs (0.02 A/cycle), and observed no significant surface compositional changes. This ALE approach was applied by Kim et al.⁷¹ for fabrication of InAlAs/InGaAs high electron mobility transistors. Changes in surface stoichiometry and surface roughness were investigated in several of the studies

Table I. Overview of materials and ALE investigations.

Material	Precursor Chemistries for Adsorption	Energy Source for Etching/Desorption	Reference
Al_2O_3	BCl ₃	Ar neutral beam	59
Al_2O_3	BCl_3	Ar neutral beam	58
BeO	BCl ₃	Ar neutral beam	60
III-V: GaAs	Cl_2	Electron bombardment	43
III-V: GaAs	Cl_2	low-energy Ar ⁺ ions	61–63
III-V: GaAs	Cl_2	248 nm KrF excimer laser and Ti sapphire laser	44,64
III-V: GaAs	Cl ₂ /Ar plasma	Ar ions from ECR plasma	65
III-V: GaAs,	Cl ₂ /Ar plasma	Ar ions from ECR plasma	66
III-V: GaAs	Cl_2	Ne neutral beam	67
III-V: InP	Tertiarybutylphosphine	Halogen lamp desorption	68
III-V: InP (and InAlAs, InGaAs)	Cl_2	Ne neutral beam	69-71
Ge	Cl_2	Ar ions from ECR plasma	72
Ge (and Si, SiGe)	Cl_2	Ar ions from ECR plasma	73
Graphene	O ₂ plasma	Ar neutral beam	74
Graphite	O ₂ plasma	Ar neutral beam	75
HfO_2	BCl ₃	Ar neutral beam	47,76
HfO_2	Cl_2	Ar neutral beam	77
Polymer (Polystyrene)	O_2	Ar ions from CCP plasma	78
Si	CF ₄ +O ₂ plasma	Ar ions from ECR plasma	79
Si	CF_4/O_2 , NF_3/N_2 , or F_2/He plasmas	Ar ions from ECR plasma	80
Si	Cl_2	Ar ions from ECR plasma	83
Si	Cl_2	Ar ions from ECR plasma	84
Si	Cl_2	50 eV Ar ions (MD simulation)	85
Si	Cl_2	Ar ions from helical resonator plasma	86
Si	Cl_2	Ar ions from helicon plasma	87,88
Si	Cl_2	Ar ions from ICP type ion gun	89
Si	Cl_2	Ar neutral beam	40,41,90,91
Si	Cl_2	Ar or He ions from ICP source	92
Si	Cl ₂ /Ar plasma	Ar ions from ICP source	9
Si	Cl ₂ /Ar plasma	Ar ions from ICP source	10
Si_3N_4	H ₂ plasma	Ion bombardment from Ar/H ₂ ECR plasma	93
Si_3N_4	Ion implantation from H ₂ plasma	Wet etching (aqueous HF solution)	94
SiO ₂ (and Si)	CF _x ⁺ ions or radical adsorption	Ar ions (MD simulation)	95
SiO_2	C ₄ F ₈ /Ar plasma	Ion bombardment from C ₄ F ₈ /Ar ICP plasma	9
SiO_2	C ₄ F ₈ /Ar plasma	Ar ions from magnetically enhanced plasma	9
SiO_2	C ₄ F ₈ /Ar plasma	Ar ions from ICP source	99
TiO_2	BCl_3	Ar neutral beam	105
ZrO_2	BCl ₃	Ar neutral beam	106

on compound semiconductors, and ALE methods generally appeared promising with regard to minimizing changes in these surface properties.

Germanium.— Sugiyama et al. 72 studied ALE of Ge based on alternating Cl_2 exposure and Ar^+ ion bombardment using an ECR system. Importantly, they found that they could not inject Cl_2 into an ECR plasma, since the plasma-generated Cl radicals etched the Ge spontaneously, whereas this was not the case for Si. They report 1.5 A/cycle for longer Ar^+ irradiation and higher microwave power, and stated that Ar^+ ions with an energy higher than ~ 13 eV were effective for etching. Similarly, Matsuura et al. 73 investigated ALE of Ge, Si, and SiGe using Cl_2 without plasma followed by low energy Ar^+ bombardment in an ECR system. For extended Ar^+ ion bombardment and chlorine surface saturation they measured an etch rate per cycle that approached an atomic-layer thickness. Ge was found to be more reactive than Si since it approaches more rapidly saturation of surface chlorination.

Graphene, graphite.— Lim et al.⁷⁴ performed ALE of graphene using an oxygen plasma for exposure of graphene to O radicals (5 min), followed by energetic Ar neutrals beam exposure (1 min). They measured removal of one graphene layer per cycle. They also discussed graphene damage issues, and damage annealing. Similarly, Kim et al.⁷⁵ applied this approach to etching of graphite, and measured removal of 1 monolayer per etching cycle.

 HfO_2 .— Park et al.^{47,76} studied ALE of HfO₂ using adsorption of BCl₃ followed by neutral beam bombardment. They observed self-limited etching of 1.2 A/cycle which required specific BCl₃ exposure and neutral beam dose. No etching was observed when using Cl₂ instead of BCl₃ for the pressures investigated (up to 0.35 mTorr). On the other hand, Min et al.⁷⁷ report HfO₂ etching using Cl₂ and neutral Ar beam etching at less than 1.0 A/cycle, with high selectivity against an SiO₂ underlayer. The reason for this difference was not discussed.

Polymer.— Vogli et al. 78 used a polystyrene-based photoresist material to demonstrate Angstrom layer removal of polymer in a capacitively coupled plasma system. One cycle consisted of O₂ exposure of the polymer material to adsorb species, O₂ exhaust from the chamber, Ar ion bombardment using low ion energies (~20 eV) to remove oxygen-associated carbon from the surface, followed by Ar exhaust. Molecular oxygen does not spontaneously react with polymers at room temperature and can be adsorbed on an activated polymer surface to form a monolayer of oxidized carbon material over unmodified polymer surface atoms. This work demonstrated that about 1.3 Å of unmodified material could be removed per step, but also illustrated the unexpected complexity of ALE processes. The key to the success of this process was the deposition of a thin (\sim 1 Å) reactive layer of polyimide-related film precursors inadvertently sputtered from a second electrode within the etching chamber. The polyimide-related deposition inhibited etching during the Ar ion bombardment step once

the oxygen-associated reactive layer had been removed. Additionally, the deposition of this ultra-thin layer increased O_2 adsorption during the O_2 gas exposure step.

Silicon.— For the **F/Si system** spontaneous etching that takes place at room temperature can be suppressed by using cryogenic temperatures. Horiike et al. 79 studied ALE of silicon and used exposure to a fluorine-based plasma for the formation of an SiF_x adsorbate layer. This was followed by bombardment with Ar ions of approximately 20 eV generated by an ECR plasma to desorb SiF_x reaction products. By careful choice of parameters, e.g. flow rate, atom mole fraction of halogen gases, exposure time, bias voltage, and cooling of the substrate to 113 K to decrease spontaneous chemical etching of silicon, they obtained an etching rate of approximately 1.5 A/cycle, close to a monolayer and indicative of approximately self-limiting etching. Sakaue et al.80 used a similar approach to achieve atomic layer etching of cooled silicon using exposures to different fluorine-based discharges. They observed that the amount of physisorbed fluorine molecules on Si surfaces controls the self-limited etching rate which varied between 2.5-8 A/cycle. They also found that the etching of Si with a 20 nm pattern width PMMA mask was anisotropic and the Si etching rate was five times larger than the etching rate of PMMA during this process, demonstrating that atomic layer etching can be a selective approach.

Cl₂/Si system.— A great deal of ALE work has focused on the Cl₂/Si system at room temperature since the saturation adsorption characteristics of Cl₂ on silicon followed by energetic ion bombardment are favorable to achieving self-limited ALE. For chlorine adsorption on a room temperature silicon substrate, Langmuir self-limited adsorption of about one monolayer of chlorine is typically seen. 81,82 Matsuura et al. 83 found a self-limited layer-by-layer etching mechanism with the substrate at room temperature using Cl₂ exposure followed by low energy (20 eV) Ar ion bombardment in an ECR system. An etch depth of 0.5 atomic layer per cycle was achieved which increased with Cl₂ exposure of the surface. Suzue et al. 84 also used Cl₂ exposure and low energy ion bombardment from an ECR plasma to examine the substrate orientation dependence of Si ALE. They found that the sticking probabilities of chlorine radicals were almost independent of the substrate orientation.

The group of Economou performed both modeling⁸⁵ and experimental work⁸⁶ on Cl₂/Ar⁺ based ALE of silicon. Molecular dynamics simulations of Si ALE by Athavale et al.85 using 50 eV argon ion bombardment of Si(100) passivated with a monolayer of adsorbed chlorine showed that 93% of etched Si originated from the top silicon layer and 7% from the underlayer. For 50 eV Ar⁺ ions the Si reaction yield was 0.172 Si atoms removed per ion, 84% in the form of SiCl, 8% elemental Si and 8% as SiCl₂. These results nicely demonstrate the concept of the ALE window, since this yield is higher than expected for physical sputtering. They also discussed introduction of structural damage in the top three silicon layers. In their experimental work⁸⁶ they used a helical resonator plasma source to achieve ALE of silicon by Cl₂ exposure and low energy Ar ion bombardment. They observed a self-limiting process with respect to both Cl₂ and ion dose, and concluded that control of the ion energy was the most important factor in realizing ALE.

Kim et al. 87,88 performed ALE of Si using Cl₂ and low energy Ar bombardment (\sim 30 eV) using a helicon plasma and employing a shutter for process control. They measured etch depths of about 0.7 A/cycle. Park et al. 89 performed Si ALE using Cl₂ exposure for 20 s followed by Ar ion bombardment at ion energies in the range of 70 to 90 eV to observe self-limited etching at 1.36 A/cycle. Subsequently, Yeom's group developed this approach to perform Si ALE using the same kind of Cl₂ exposure in combination with energetic Ar neutrals obtained by the low-angle forward reflection neutral beam technique. 40,41,90,91 They observed a self-limited Si etch rate of a monolayer per cycle for both Si (100) and Si (111) orientations when Cl₂ and Ar neutrals were supplied above the critical dose values, and sur-

face roughness that remained very low and comparable to a reference sample without ALE.

Yun et al.⁹² studied ALE of poly-Si using Cl₂ exposure and either Ar or He low energy bombardment. They observed self-limited etching of 0.8 A/cycle for Ar, and 0.6 A/cycle for He, with a process window that was much greater for He than Ar ion bombardment.

Agarwal and Kushner⁹ addressed the question if Si ALE can be performed in conventional plasma etching equipment, e.g. an ICP etching system for directional Si etching. In their computational study of Si ALE they modeled an inductively coupled plasma where Cl₂/Ar without bias of the Si substrate is used for chlorine passivation of the Si surface. During the passivation step, Cl⁺ is the dominant ion and ion energies are below 20 eV. The chlorination of the surface is due to neutral Cl atoms formed by the plasma, which at the 20 mTorr pressure used proceeds rapidly and does not produce multiple layers of passivation since diffusion of Cl into the Si is slow. This is followed by a second cycle in which a pure Ar plasma and biasing is used to etch the passivated Si. During the etching step, Ar⁺ ion energies are between 50 and 60 eV, and observed etch rates were about 1 ML/cycle. One conclusion of this work was that an essential prerequisite for achieving ALE performance of Si are control of the Ar/Cl₂ chemistry along with ion energy and angular distribution (IEAD) functions. This approach has significant potential to speed up processing relative to ALE based on halogenation of Si using simply gaseous halogens,

Recently, Kanarik et al.¹⁰ realized this ALE approach in an ICP reactor equipped with fast gas-switching capabilities to achieve short process cycles. Rapid surface chlorination using an Ar/Cl₂ plasma, rapid pump-out to establish a pure Ar plasma and RF biasing for efficient product desorption were discussed as key to achieve practical ALE cycle times. The ALE process was reported to consist of self-limiting cycles which yielded an etched Si surface that was smoother and showed no microtrenching as compared to Si etched using a continuous plasma etching process.

 Si_3N_4 . — Matsuura et al. 93 proposed layer by layer etching of Si_3N_4 by using a remote H_2 ECR plasma. The interaction of hydrogen atoms with the Si_3N_4 surface removed N atoms from the outermost surface of the Si_3N_4 . This was followed by bombardment of the modified surface using Ar and hydrogen ions. Posseme et al. 94 evaluated a thin layer etching method based on low energy ion implantation of the Si_3N_4 surface using an H_2 plasma. The modified Si_3N_4 surface layer could be selectively removed using wet etching. The modified surface layer had a typical extent of about 10 to 20 nm in this work, due to the projected range of hydrogen ions, and thus was significantly greater than typical layer thicknesses removed by ALE processes.

SiO₂.— For atomic layer etching of SiO₂ using fluorocarbon ions or precursors, computational work has been performed by Rauf et al.⁹⁵ and Agarwal and Kushner.⁹ The molecular dynamics simulation of Rauf et al.⁹⁵ first showed potential of a two-step etch process consisting of the formation of a nanometer-thick, self-limited fluorocarbon passivation layer on an SiO₂ or Si surface followed by etching with Ar⁺ ions with energies up to 50 eV using the deposited fluorocarbon as a source of etchant. A sequence of these steps enabled nanometer precise etching of SiO₂ and Si.

Agarwal and Kushner⁵ examined ALE of SiO₂ using C₄F₈/Ar in a capacitively coupled plasma (CCP) reactor. Their work was motivated by the observation that because of the use of different gas mixtures for the passivation and etching steps and the need to exhaust the reaction chamber, ALE results in an inherently slow etch rate. They argued that during actual device fabrication the switch to an ALE method would likely only be implemented after a conventional rapid plasma etching process had thinned the material to a few monolayers above the interface between materials. Such an approach could be realized using separate, dedicated plasma etching and ALE processing chambers. This is undesirable, since it is expensive from several points of view and requires additional wafer handling. Alternatively, if one could perform both main etch and atomic layer etching in the

same conventional plasma reactor, cost and processing time would be reduced. Since a limitation of these approaches with respect to throughput is the existence of purge steps, Agarwal and Kushner⁹ also examined the question if elimination of the purge steps is possible and self-limited etching can be achieved if the entire SiO₂ etching cycle is performed using a single gas mixture, and simply controlling ion bombardment energies during a cycle by changing RF bias. By utilizing a nonsinusoidal bias waveform, ^{96,97} they controlled ion energy distribution functions, and demonstrated self-limiting etching at 1 to several ML/cycle. This method is related to pulsed plasma approaches. ^{5,98}

Using a steady-state Ar plasma in conjunction with periodic injection of a defined number of C_4F_8 molecules and synchronized plasmabased Ar⁺ ion bombardment, Metzler et al.⁹⁹ evaluated an approach related to both simulations^{9,95} and demonstrated that in agreement with the simulations Angstrom level precision in etching of SiO₂ is possible. For low energy Ar⁺ ion bombardment conditions giving a maximum ion energy of about 20 eV, the physical sputter rate of SiO₂ vanishes. Conversely, for the same ion energies and a SiO₂ surface coated with several Angstroms of fluorocarbon (FC), SiO₂ etching is initiated, and stops once the FC supply is exhausted. Precise management of C_4F_8 supply enables control of the deposited FC layer thickness in the 1 to several Angstrom range. The temporal variation of FC deposition, FC and SiO₂ etching for Ar⁺ ion energies of 25 eV for this process during a number of cycles is shown in Fig. 6. As the fluorocarbon surface coverage decreases, the SiO₂ ER vanishes,

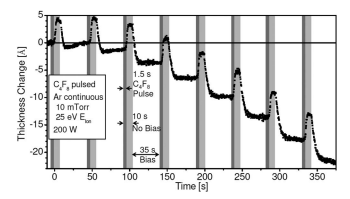


Figure 6. Example of thickness evolution during eight cycles of an SiO_2 ALE process. ⁹⁹

which enables controlled removal of Angstrom-thick SiO_2 layers per cycle.

Improved control of etching selectivity near the etching threshold energies is one motivation for ALE. In Figs. 7a and 7b thickness changes during a typical ALE cycle are shown for SiO₂ at maximum ion bombardment energies of 25 eV and 30 eV, respectively. ¹⁰⁰ Upon precursor injection a fast FC deposition is seen resulting in a film about

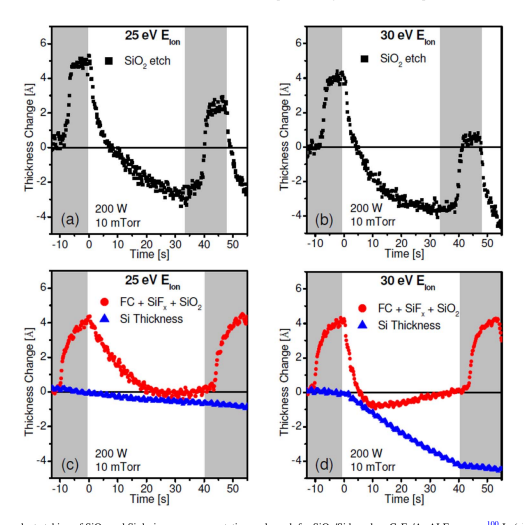


Figure 7. Time-dependent etching of SiO_2 and Si during one representative cycle each for SiO_2/Si based on C_4F_8/Ar ALE process. 100 In (a) and (b) deposition of FC on SiO_2 followed by etching at 25 eV and 30 eV maximum ion energy are shown, respectively, whereas in (c) and (d) deposition of FC on SiO_2/Si followed by etching at 25 eV and 30 eV maximum ion energy are shown, respectively. The most significant change is seen for the Si etching rate as the maximum ion energy is increased to 30 eV, leading to a very large increase of the Si etch rate, loss of self-limited etch rate and of SiO_2/Si etching selectivity.

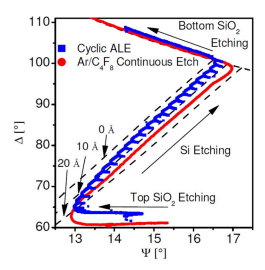


Figure 8. Real time ellipsometry trajectory showing key differences in surface processes between ALE and steady-state etching during continuous plasma etching of a SiO₂-Si-SiO₂ multi-layer stack. ¹⁰⁰ Each dashed line marks 10 Å of FC film.

4 Å thick. When the bias potential is applied at 0 s the ion energy is increased to the above values and the FC film is rapidly etched, followed by SiO_2 etching. Once the FC is depleted the etch rates cease before the next precursor injection starts another cycle. Figures 7c and 7d display the corresponding information for Si etching. While the variation of the SiO_2 etching rate with ion energy is fairly small, it is much larger for Si. The result is that the process exhibits SiO_2/Si etching selectivity for a maximum ion energy of 25 eV, whereas for a maximum ion energy of 30 eV the etching selectivity is reduced. The relative placement of the ALE windows for dissimilar materials, i.e. differences in the energy thresholds $E_{psth}(SiO_2) - E_{ceth}(SiO_2)$ versus $E_{psth}(Si) - E_{ceth}(Si)$ and precise placement of the ion energy distribution within the ALE window of the target material is important to maximize etching selectivity.

Figure 8 shows real time in situ ellipsometry data of a typical ALE process for a Si surface in comparison to a continuous Ar/C₄F₈ plasma etch of a multilayer stack sample. 100 The stack sample has a Si layer, about 10 nm thick, sandwiched between two SiO₂ layers. The transition from SiO₂ to Si etching and back to SiO₂ etching can clearly be seen as sharp turns in the trajectory (near descriptions "Top SiO₂ Etching" and "Bottom SiO₂ Etching"). During "Si Etching", each cycle of the ALE approach can clearly be seen by the FC deposition as an increase in Ψ . It is noticeable that the continuous etch is shifted to higher values of Ψ compared to the ALE process. The comparison shows that the continuous Si etch exhibits a significantly thicker steady-state FC film than the maximum FC film thickness deposited during each ALE cycle. These data demonstrate that ALE enables processing where surface conditions, including reactant supply, are highly controlled, strongly time-dependent and much closer to atomically abrupt interfaces.

Recently, Hudson et al. ¹⁰¹ reported a similar highly selective SiO₂ etch process based on repeated cycles of FC deposition and etch reaction activation using low energy ion bombardment.

While these FC deposition approaches are reminiscent of an approach called the "Bosch process" consisting of FC deposition during C_4F_8 passivation cycles followed by etching cycles using SF_6 , the Bosch process is primarily employed for deep reactive ion etching of silicon and the goal of the FC deposition is the achievement of sidewall passivation rather than as a source of etchant. 102,103 Roozeboom et al. 104 have proposed a method where this kind of process can be performed by horizontally moving the substrate back and forth during exposure to two chemically distinct gas discharges separated by inert gas curtains. This approach using ALD-based passivation may have potential as a basis of ALE processes.

 TiO_2 .— Park et al. ¹⁰⁵ studied ALE of TiO₂ using 20 s adsorption of BCl₃ followed by 60 eV Ar neutral beam bombardment, and measured 1.25 A/cycle. They report a critical BCl₃ pressure of 0.16 mTorr and neutral beam dose of 1.49×10^{17} atoms/cm² for these conditions, along with low surface roughness and no change in surface chemistry.

 ZrO_2 .— Lim et al. ¹⁰⁶ studied ALE of ZrO_2 using adsorption of BCl₃ followed by Ar neutral beam bombardment, and measured 1.07 A/cycle at the higher BCl₃ pressures investigated (0.15 mTorr).

Issues and Needs

In this section we attempt to summarize several considerations of developments and opportunities that will be helpful for realization of ALE processes in manufacturing environments applicable to a broad set of reactant/materials systems. Additionally, we will briefly discuss features and challenges that will have to be overcome.

Scientific and technological basis of ALE.— There has been a tremendous growth in knowledge in low temperature plasma materials processing/plasma etch science and technology since the first ALE studies. Examples are a) control of ionization, dissociation and uniformity across wafer, ^{1,3,107,108} b) ion energy control, ¹⁰⁹ including using shaped waveforms, ⁹⁶ c) pulsed plasma science and technology, ⁵ d) control of wafer charging and damage effects, ^{110–112} e) plasma characterization and metrology, ^{113,114} f) understanding of the control of plasma-polymer interactions, Line Edge Roughness, Line Width Roughness, CD and CD variation, ¹¹⁵ advanced modeling and simulations of all of the above, ¹¹⁶ and tremendous growth in advanced hardware engineering and capabilities.

Additionally, while true Atomic Layer Etching may be the ultimate goal for the most demanding applications, for many pattern transfer/dry etching applications dimensional control at the Angstrom level in combination with materials selectivity rather than true atomistic level control is sufficient. Indeed, the achievement of *self-limited* deposition and etching reactions by cyclic processing is deemed to be one of the key requirements on a practical ALE process, rather than achieving true atomistic resolution for each etching cycle. ¹⁰

The above advances place atomic layer etching on a much stronger scientific and technological basis than the early efforts. In particular, the great advances in computational modeling of plasma and plasma/surface interactions have been important in designing and evaluating potential ALE approaches prior to experimental validation, e.g. as seen in the case of SiO₂ ALE. ^{9,95,99} Additionally, the somewhat relaxed expectations relative to true atomistic level control make the prospect of broad implementation more realistic.

The potential of precursor chemistry.— Atomic layer etching also may be expected to have greater potential than conventional steady-state plasma etching to utilize the chemical nature of precursors and thus gain a new level of control over surface reactions. This ability is strongly reduced in continuous plasma processing, and has limited our possibilities of controlling surface reactions by choice of precursor molecular structure. Either exposure of a substrate to precursor gases without plasma or short plasma exposures offer the prospect of retaining a much larger proportion of the precursor molecular structure at the surface, and in this fashion impact etching reactions. The exposure parameters can be varied over a significant range, with steady-state behavior as a limit.

The surface composition of plasma-deposited FC films using C_4F_8 and CHF $_3$ ALE processes for two thicknesses is shown in Fig. 9. 100 The C1s spectra were determined after the deposition step during the 10^{th} ALE cycle using X-ray photoelectron spectroscopy. Ultrathin FC films (~ 3 Å) deposited using CHF $_3$ exhibit a slightly higher F/C ratio composition than ultra-thin layers deposited using C_4F_8 (Fig. 9a). This is in contrast to continuous plasma etching, where the steady-state FC films formed from CHF $_3$ typically show a significantly lower F/C ratio than FC films deposited using C_4F_8 . This is due to both

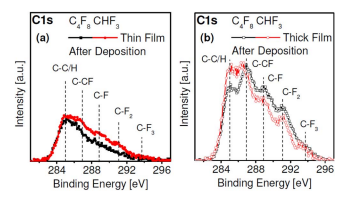


Figure 9. Enhanced potential of precursor choice relative to continuous plasma etching. 100 The steady-state surface composition is not reached during a short deposition cycle (\sim 3 Å in (a)), whereas for the thicker (\sim 15 Å) film deposited in (b) the carbon 1s photoemission spectrum has become similar to that seen for FC films produced using continuous plasma processes.

a high FC deposition rate for C_4F_8 , and lack of hydrogen in the discharge which during ion bombardment of the film enhances F loss from the FC surface by HF formation for CHF3. For very thin ALE depositions the FC films reflect better the relative fluorine to carbon ratio of the feedgas which is higher for CHF3 than for C_4F_8 . For thicker films of ≈ 15 Å (Fig. 9b), the F/C ratio of the film deposited by C_4F_8 is higher compared to the film deposited by CHF3. The surface chemistry differences between CHF3 and C_4F_8 seen for the thicker depositions are consistent with the findings of Standaert et al. 24

Besides these helpful developments and features, there is a number of considerations and observations which demonstrate the difficulty of transiting ALE from research and development to manufacturing.

As discussed, although ALE shows some similarities to ALD with regard to physisorption/chemisorption requirements at the surface, the requirements with regard to "volatile product" removal are fundamentally different. Whereas for ALD films are grown in a conformal fashion, for ALE the "etch product" removal ideally should take place in a directional fashion, and requires energetic bombardment. The ALE window is located within a range of ion energies, and depends very sensitively on minute changes in surface chemistry (of the order of a monolayer). Because of this fundamental difference from ALD, ALE provides profound surface chemistry challenges and energetic species/surface interaction problems that are unique.

Reproducing sputter etching energy thresholds.— To achieve self-limited etching in ALE, processing near the energy thresholds of physical sputtering is required. In Figs. 10 and 11 we have collected physical sputtering yield data by Arion bombardment with energies up to 400 eV for both Si and SiO₂, respectively. SRIM simulation results and physical sputtering yields by Ar ion bombardment in the ALE work performed at University of Maryland^{99,100} are also shown. The surface sputtering simulation was conducted for ion bombardment normal to the surface using the software http://www.srim.org/. The simulation parameters are presented below.¹¹⁷

The physical sputter yield of Si as reported in the literature 95,118-132 when plotted versus the square root of the Ar⁺ ion energy shows a great deal of scatter around the energy threshold for physical sputtering (see Fig. 10). Similarly, the physical sputter yield data of SiO₂ for Ar⁺ ion bombardment 95,122,124,128,133-137 show significant scatter (see Fig. 11). While the data span many years, the scatter reflects the difficulty to reproduce these experimental conditions. For instance, differences in materials, e.g. single crystal silicon versus silicon thin films deposited on quartz microbalances, along with vacuum quality and other factors may explain some of the differences. Additionally, surface modifications during sputtering can change observed sputter yield, e.g. surface roughness, surface impurities, and so forth. Figures 10 and 11 illustrate that controlling etching behavior near the energy threshold for physical sputtering, i.e. close to the ALE win-

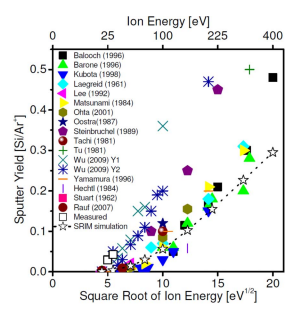


Figure 10. Review of literature data of physical sputter rates reported for Si versus the square-root of Ar ion energy up to energies of $400 \text{ eV}^{95,118-132}$ along with SRIM simulation results. The threshold for physical sputtering of Si is $\approx 20 \text{ eV}$. The large scatter with regard to the threshold energy of physical sputtering for Si shows the difficulty of controlling experimental conditions.

dow, is extremely challenging. On the other hand, physical sputtering will be most sensitive to small changes in residual impurities, and it is possible that for ALE the presence of chemical reactants at saturation coverage on a surface will overwhelm the factors leading to discrepancies and produce more stable and reproducible responses. Therefore, assuming that the energy of ions inducing physical sputtering can be precisely controlled, the ability to accurately control the flux of chemical reactants to the substrate is required.

Impact of surface reactions on plasma characteristics.— The role of reactor surfaces (heterogeneous reactions) on continuous plasma etching process stability is well-known.^{3,138–141} One important difference between ALD systems and ALE systems is the overall energy content of ALE systems, in particular if plasma is used continuously

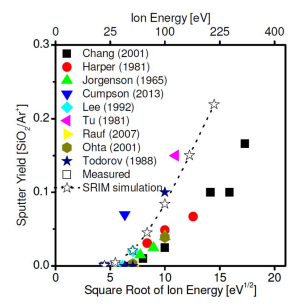


Figure 11. Review of literature data of physical sputter rates for SiO₂ versus the square-root of Ar ion energy up to energies of 400 eV. $^{95,122,124,128,133-137}$ The threshold for physical sputtering of SiO₂ is \approx 45 eV.

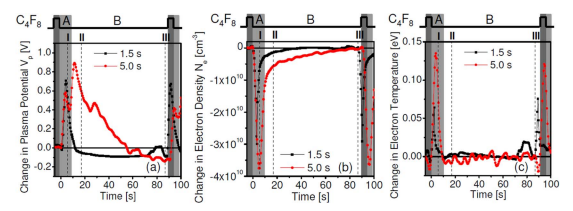


Figure 12. Evolution of plasma properties from cycle to cycle as measured using a Langmuir probe. 100 The cycle-to-cycle changes increase with C_4F_8 pulse length due to more residual FC in the system. This is explained by more deposition of FC on the walls of the apparatus, contributing residual FC during Ar cycles when ideally FC is only present on the active surface of the substrate. The overall behavior is consistent with the impact of continuous precursor addition on plasma electrical properties.

and for all surface reaction steps. The result will be a highly dynamic environment where species transport between different surfaces can easily take place. For atomic layer etching, changes in the state of reactor surfaces and enhanced supply of reactants from "passive" surfaces (e.g., walls) by plasma-wall interactions could potentially lead to loss of control over the supply of chemical precursors to "active" surfaces. While plasma-enhanced surface passivation is desirable when considering throughput, the interaction of chemical reactants with "passive" surfaces will be enhanced for this situation. The formation of reactive radicals required to speed up adsorption processes at "active" surfaces will lead to greater interactions with "passive" surfaces which can become a supply of chemical reactants. Assuming a partial pressure of 10^{-4} chemical reactants for a plasma system operating at 10 mTorr pressure leads to the arrival of about 1 monolayer per s of chemical reactants during the process step whereas during the surface activation step no chemical reactants should be present to achieve a self-limited etch per cycle. Work needs to address what may be a proper balance between achieving ultra-clean processing/process control and rapid processing required for enhanced throughput.

This enhanced feedback from "passive" surfaces on plasma properties has been seen in the FC-based ALE of SiO₂ and is demonstrated by Langmuir probe measurements during gas pulsing and shown in Fig. 12.¹⁰⁰ Figure 12 displays the change in plasma properties, i.e. plasma potential V_n (Fig. 12a), electron density n_e (Fig. 12b), and electron temperature T_e (Fig. 12c), respectively, during one ALE cycle for two different conditions. These measurements show rapid changes of the electrical discharge characteristics during gas pulsing and additionally slow long-term changes in plasma electrical properties due to FC film buildup on walls. The increase seen for V_p and T_e and the decrease of n_e upon C₄F₈ precursor injection agrees with the impact on electrical properties for experiments using continuous precursor addition. For short pulses, the plasma properties return to values similar to those measured before the pulse within 20 s. A stronger and longer impact can be seen when increasing the C₄F₈ pulse length from 1.5 s to 5.0 s, and are consistent with the presence of residual precursor in the Ar plasma long after the initial pulse has been pumped out. Similar effects may be expected for other ALE chemistries, and point out the need for ALE process chamber designs and compensation techniques that will minimize these effects.

Importance of photo-induced processes at low ion energies.— Irradiation of surfaces by vacuum ultraviolet (VUV) light generated in plasmas can strongly affect surface reactions, and also give rise to synergistic effects. Donnelly's and Economou's groups^{46,142} reported on the importance of photo-assisted etching of silicon in chlorine-and bromine-containing plasmas for very low ion bombardment energies using nearly mono-energetic ion energy distributions. At this

time the mechanistic origin of this observation is not well understood. The question of how important and generic photo-enhanced etching is for plasma-based ALE processes needs to be examined. This observation highlights the need to examine and understand the potential importance of simultaneous photon irradiation on ion-induced ALE processes in plasma environments in general.

Performance in 3-dimensional structures.— Little is known at this time on the application of basic ALE procedures to advanced structures and applications. Plasma-enhanced surface passivation could potentially face limitations when considering substrates with pronounced surface topography. Radicals may have to undergo several surface collisions in order to reach the bottom of contact holes or trenches. The reduction in radical flux to surface elements at the bottom of features by surface recombination has been discussed extensively, ${}^{3}6, {}^{3}9, {}^{107, 143}$ and depends strongly on the value of the recombination coefficient r. This potentially could lead to situations where one location and material may exhibit self-limited adsorption at a monolayer, whereas another material may show multi-layer adsorption. Additionally, the redeposition of etch product on the feature sidewalls has to be considered. These phenomena could complicate the application of certain ALE processes to high aspect ratio structures.

Conclusions

As critical dimensions of semiconductor devices approach the sub-10 nm scale, the industrial need for an etching method capable of near-atomic resolution has led to a large increase in activity on novel directional etching methods based on a sequence of individual, self-limited surface reaction steps. The ability to achieve controlled, self-limited removal of various films at the Angstrom-level has been demonstrated for a range of reactant/materials systems, although specialized equipment with low throughput has often been used. Since for many situations true atomistic layer removal is not yet required and self-limited processing at the Angstrom scale appears sufficient, recent efforts have focused on increasing the speed with which selflimited surface reaction steps can be realized using plasma etching systems with enhanced temporal control of power, substrate biasing and gas delivery/pumping. A key challenge will be the insertion of ALE approaches into manufacturing. Since the overall energy content of ALE systems is expected to be significantly higher than for typical ALD systems, in particular if plasma is used during all reaction steps, this will produce a highly dynamic environment where species transport between different surfaces can easily take place. Maintaining controlled etching behavior for such systems within the ALE window near the energy threshold for physical sputtering is extremely challenging and will require stringent control of system cleanliness and the state of "passive" surfaces. These efforts are expected to benefit from advances in ALE chemistries and approaches. Computational modeling has been important in designing and evaluating potential ALE approaches and is expected to play a crucial role in the future for pin-pointing/appraising opportunities.

Acknowledgments

The authors gratefully acknowledge the financial support of this work by the National Science Foundation (CBET-1134273), US Department of Energy (DE-SC0001939) and Lam Research Corporation. The authors also thank E. Vogli, M. Vollmer, N. Fox-Lyon, E. Bartis, A. Knoll, P. Luan, D. Kaplan, Th. Pitzel and M. Kawakami for contributions during phases of this project. We also thank R. L. Bruce, S. Engelmann, and E. A. Joseph of IBM Research, V. Godyak, and E. Hudson, S. Lai, M. Danek, and A. Dulkin of Lam Research Corporation for helpful discussions and collaborations on ALE.

References

- V. M. Donnelly and A. Kornblit, "Plasma etching: Yesterday, today, and tomorrow," *Journal of Vacuum Science & Technology A*, 31, 050825 (2013).
- N. Marchack and J. P. Chang, "Perspectives in nanoscale plasma etching: what are the ultimate limits?," *Journal of Physics D-Applied Physics*, 44, 174011 (2011).
- C. G. N. Lee, K. J. Kanarik, and R. A. Gottscho, "The grand challenges of plasma etching: a manufacturing perspective," *Journal of Physics D-Applied Physics*, 47, 273001 (2014).
- S. Banna, A. Agarwal, G. Cunge, M. Darnon, E. Pargon, and O. Joubert, "Pulsed high-density plasmas for advanced dry etching processes," *Journal of Vacuum Science & Technology A.* 30, 040801 (2012).
- ence & Technology A, 30, 040801 (2012).
 5. D. J. Economou, "Pulsed plasma etching for semiconductor manufacturing," *Journal of Physics D-Applied Physics*, 47, 303001 (2014).
- C. Petit-Étienne, M. Darnon, P. Bodart, M. Fouchier, G. Cunge, E. Pargon, L. Vallier, O. Joubert, and S. Banna, "Atomic-scale silicon etching control using pulsed Cl₂ plasma," *Journal of Vacuum Science & Technology B*, 31, 011201 (2013).
- S. Samukawa, M. Hori, S. Rauf, K. Tachibana, P. Bruggeman, G. Kroesen, J. C. Whitehead, A. B. Murphy, A. F. Gutsol, S. Starikovskaia, U. Kortshagen, J. P. Boeuf, T. J. Sommerer, M. J. Kushner, U. Czarnetzki, and N. Mason, "The 2012 Plasma Roadmap," *Journal of Physics D-Applied Physics*, 45, 37 (2012).
- M. Guillorn, J. Chang, N. Fuller, J. Patel, M. Darnon, A. Pyzyna, E. Joseph, S. Engelmann, J. Ott, J. Newbury, D. Klaus, J. Bucchignano, P. Joshi, C. Scerbo, E. Kratschmer, W. Graham, B. To, J. Parisi, Y. Zhang, and W. Haensch, "Hydrogen silsesquioxane-based hybrid electron beam and optical lithography for high density circuit prototyping", J. Vac. Sci. Technol. B., 27, 2588 (2009).
- A. Agarwal and M. J. Kushner, "Plasma atomic layer etching using conventional plasma equipment," *Journal of Vacuum Science & Technology A*, 27(1), 37 (2009).
- K. J. Kanarik, S. Tan, J. Holland, A. Eppler, V. Vahedi, J. Marks, and R. A. Gottscho, "Moving atomic layer etch from lab to fab," *Solid State Technology*, 56, 14 (2013).
- 11. "First Workshop on Atomic-Layer-ETch (ALET) and -Clean (ALC) technology", April 2014, San Francisco (organized by Craig Huffman, Int. SEMAT-ECH); http://www.sematech.org/meetings/archives/fep/10605/index.htm; In conjunction with ALD 2015, AVS Plasma Science and Technology Division will be hosting a "Workshop on Atomic Layer Etching", July 1–2, 2015, Portland (Chair, Eric A. Joseph).
- 12. http://en.wikipedia.org/wiki/The_Tipping_Point.
- P. D. Agnello and P. D. "Process requirements for continued scaling of CMOS the need and prospects for atomic-level manipulation," *IBM J. Res. Dev.*, 46(2–3), 317 (2002).
- S. Bangsaruntip, G. M. Cohen, A. Majumdar, Y. Zhang, S. U. Engelmann, N. C. M. Fuller, L. M. Gignac, S. Mittal, J. S. Newbury, M. Guillorn, T. Barwicz, L. Sekaric, M. M. Frank, and J. W. Sleight, "High Performance and Highly Uniform Gate-All-Around Silicon Nanowire MOSFETs with Wire Size Dependent Scaling", IEDM Baltimore (2009).
- M. Guillorn, J. Chang, A. Pyzyna, S. Engelmann, E. Joseph, B. Fletcher, C. Cabral, Jr., C.-H. Lin, A. Bryant, M. Darnon, J. Ott, C. Lavoie, M. Frank, L. Gignac, J. Newbury, C. Wang, D. Klaus, E. Kratschmer, J. Bucchignano, B. To, W. Graham, I. Lauer, E. Sikorski, S. Carter, V. Narayanan, N. Fuller, Y. Zhang, and W. Haensch, "Trigate 6T SRAM Scaling to 0.06 m²", IEDM Baltimore (2009).
- K. J. Kanarik, G. Kamarthy, and R. A. Gottscho, "Plasma etch challenges for Fin-FET transistors," Solid State Technology, 55(3), 15 (2012).
- 17. F. Schwierz, "Graphene transistors," Nature Nanotechnology, 5(7), 487 (2010).
- D. Akinwande, N. Petrone, and J. Hone, "Two-dimensional flexible nanoelectronics," *Nature Communications* 5 (2014).
- J. L. Zhang, X. Wang, G. Dinakov, and H. Dai, "Narrow graphene nanoribbons from carbon nanotubes," *Nature*, 458, 877 (2009).
- G. S. Oehrlein and H. L. Williams, "Silicon Etching Mechanisms in a CF₄/H₂ Glow Discharge", J. Appl. Phys., 62, 662 (1987).
- T. E. F. M. Standaert, M. Schaepkens, N. R. Rueger, P. G. M. Sebel, G. S. Oehrlein, and J. M. Cook, "High-Density Fluorocarbon Etching of Silicon in an Inductively Coupled Plasma: Mechanism of Etching through a Thick Steady-State Fluorocarbon Layer," J. Vac. Sci. & Technol. A, 16, 239 (1998).

- N. R. Rueger, J. J. Beulens, M. Schaepkens, M. F. Doemling, J. M. Mirza, T. E. F. M. Standaert, and G. S. Oehrlein, "The role of steady-state fluorocarbon films in the etching of silicon dioxide using CHF₃ in an inductively coupled plasma reactor," *J. Vac. Sci. & Technol. A*, 15, 1881 (1997).
- M. Schaepkens, T. E. F. M. Standaert, N. R. Rueger, P. G. M. Sebel, and G. S. Oehrlein, "Study of the SiO₂-to-Si₃N₄ etch selectivity mechanism in inductively coupled fluorocarbon plasmas and a comparison with the SiO₂-to-Si mechanism," *J. Vac. Sci. & Technol. A*, 17, 26 (1999).
- 24. T. E. F. M. Standaert, C. Hedlund, E. A. Joseph, G. S. Oehrlein, and T. J. Dalton, "On the role of the fluorocarbon film formation in the etching of silicon, silicon dioxide, silicon nitride, and amorphous hydrogenated silicon carbide," *J. Vac. Sci. & Technol. A*, 22, 53 (2004).
- A. Sankaran and M. J. Kushner, "Integrated feature scale modeling of plasma processing of porous and solids SiO₂. I. Fluorocarbon etching," *Journal of Vacuum Science & Technology A*, 22(4), 1242 (2004).
- A. Sankaran and M. J. Kushner, "Etching of porous and solid SiO₂ in Ar/c-C₄F₈, O₂/c-C₄F₈ and Ar/O₂/c-C₄F₈ plasmas," *Journal of Applied Physics*, 97(2) (2005).
- 27. G. S. Oehrlein and R. Kalish, "Silicon loss and transient etch rate tech rate is selective reactive
- ion etching of oxide overlayers," *App. Phys. Lett.*, **54**, 2698 (1989).
 28. G. S. Oehrlein and Y. Kurogi, "Sidewall Chemistry in Plasma Etching," *Materials Science and Engineering Reports R*, **24**, 153 (1998).
- S. M. George, "Atomic Layer Deposition: An Overview," *Chemical Reviews*, 110(1), 111 (2010).
- M. Leskelä and M. Ritala, "Atomic layer deposition (ALD): from precursors to thin film structures," *Thin Solid Films*, 409, 138 (2002).
- M. Leskelä and M. Ritala, "Atomic layer deposition chemistry: Recent developments and future challenges," Angewandte Chemie-International Edition, 42, 5548 (2003).
- S. M. George, A. W. Ott, and J. W. Klaus, "Surface chemistry for atomic layer growth," *Journal of Physical Chemistry*, 100, 13121 (1996).
- A. W. Ott, J. W. Klaus, J. M. Johnson, and S. M. George, "Al₂O₃ thin film growth on Si(100) using binary reaction sequence chemistry," *Thin Solid Films*, 292, 135 (1997).
- 34. M. D. Groner, J. W. Elam, F. H. Fabreguette, and S. M. George, "Electrical characterization of thin Al_2O_3 films grown by atomic layer deposition on silicon and various metal substrates," *Thin Solid Films*, **413**, 186 (2002).
- M. D. Groner, F. H. Fabreguette, J. W. Elam, and S. M. George, "Low-temperature Al₂O₃ atomic layer deposition," *Chemistry of Materials*, 16, 639 (2004).
- H. B. Profijt, S. E. Potts, M. C. M. van de Sanden, and W. M. M. Kessels, "Plasma-Assisted Atomic Layer Deposition: Basics, Opportunities, and Challenges," *Journal of Vacuum Science & Technology A*, 29(5), 050801 (2011).
- S. E. Potts and W. M. M. Kessels, "Energy-enhanced atomic layer deposition for more process and precursor versatility," *Coord. Chem. Rev.*, 257(23–24), 3254 (2013).
- H. Kim and I. K. Oh, "Review of plasma-enhanced atomic layer deposition: Technical enabler of nanoscale device fabrication," *Jpn. J. Appl. Phys.*, 53(3) 03DA01 (2014).
- H. C. M. Knoops, E. Langereis, M. C. M. van de Sanden, and W. M. M. Kessels, "Conformality of Plasma-Assisted ALD: Physical Processes and Modeling," *Journal of the Electrochemical Society*, 157(12), G241 (2010).
- S. D. Park, D. H. Lee, and G. Y. Yeom, "Atomic layer etching of Cl-adsorbed silicon by using a low-angle forward reflected Ar neutral beam," *Journal of the* Korean Physical Society, 47(3), 469 (2005).
- S. D. Park, D. H. Lee, and G. Y. Yeom, "Atomic layer etching of Si(100) and Si(111) using Cl₂ and Ar neutral beam," *Electrochemical and Solid State Letters*, 8(8), C106 (2005).
- D. J. Economou, "Fast (tens to hundreds of eV) neutral beams for materials processing," *Journal of Physics D-Applied Physics*, 41, 024001 (2008).
 T. Meguro, M. Hamagaki, S. Modaressi, T. Hara, Y. Aoyagi, M. Ishii, and
- T. Meguro, M. Hamagaki, S. Modaressi, T. Hara, Y. Aoyagi, M. Ishii, and Y. Yamamoto, "Digital Etching of GaAs - New Approach of Dry Etching to Atomic Ordered Processing," *Applied Physics Letters*, 56(16), 1552 (1990).
- M. Ishii, T. Meguro, K. Gamo, T. Sugano, and Y. Aoyagi, "Digital Etching Using Krf Excimer-Laser - Approach to Atomic-Order-Controlled Etching by Photo Induced Reaction," *Japanese Journal of Applied Physics Part 1-Regular Papers Short Notes* & Review Papers, 32(12B), 6178 (1993).
- E.g. see, K. Nakayama, C. M. Aldao, and J. H. Weaver, "Halogen etching of Si(100)-2×1: Dependence on surface concentration," *Physical Review B*, 59, 15893 (1999).
- H. Shin, W. Y. Zhu, V. M. Donnelly, and D. J. Economou, "Surprising importance of photo-assisted etching of silicon in chlorine-containing plasmas," *Journal of Vacuum Science & Technology A*, 30(2) (2012).
- S. D. Park, W. S. Lim, B. J. Park, H. C. Lee, J. W. Bae, and G. Y. Yeom, "Precise depth control and low-damage atomic-layer etching of HfO₂ using BCl₃ and Ar neutral beam," *Electrochemical and Solid State Letters*, 11(4), H71 (2008).
- A. Ludviksson, M. D. Xu, and R. M. Martin, "Atomic Layer Etching Chemistry of Cl₂ on GaAs(100)," Surface Science, 277, 282 (1992).
- J. C. Patrin, Y. Z. Li, M. Chander, and J. H. Weaver, "Atomic Layer Etching of GaAs(110) with Br₂ Studied by Scanning Tunneling Microscopy," *Applied Physics Letters*, 62, 1277 (1993).
- J. C. Patrin and J. H. Weaver, "Br₂ and Cl₂ Adsorption and Etching of GaAs(110) Studied by Use of Scanning-Tunneling-Microscopy," *Physical Review B*, 48, 17913 (1993)
- H. F. Winters and J. W. Coburn, "Surface Science Aspects of Etching Reactions," Surf. Sci. Rep., 14, 161 (1992).
- D. Rioux, F. Stepniak, R. J. Pechman, and J. H. Weaver, "Chemisorption and Thermally Activated Etching of Si(100)-2×1 by Iodine," *Physical Review B*, 51(16), 10981 (1995).

- 53. R. J. Pechman, X. S. Wang, and J. H. Weaver, "Interactions of Br with Si(111)-7×7 - Chemisorption, Step Retreat, and Terrace Etching," Physical Review B, 52, 11412 (1995)
- 54. C. M. Aldao and J. H. Weaver, "Halogen etching of Si via atomic-scale processes," Progress in Surface Science, 68(4-6), 189 (2001).
- 55. C. M. Aldao, Abhishek Agrawal, R. E. Butera, and J. H. Weaver, "Atomic processes during Cl supersaturation etching of Si(100)-(2×1)," Physical Review B, 79, 125303 (2009).
- 56. J. J. Boland and J. H. Weaver, "A surface view of etching," Phys. Today, 51(8), 34 (1998).
- 57. P. A. Maki and D. J. Ehrlich, "Laser Bilayer Etching of Gaas-Surfaces," Applied Physics Letters, 55(2), 91 (1989).
- 58. Young I. Jhon, Kyung S. Min, G. Y. Yeom, and Young Min Jhon, "Understanding time-resolved processes in atomic-layer etching of ultra-thin Al2O3 film using BCl3 and Ar neutral beam," Applied Physics Letters, 105(9), 093104 (2014).
- 59. K. S. Min, S. H. Kang, J. K. Kim, Y. I. Jhon, M. S. Jhon, and G. Y. Yeom, "Atomic layer etching of Al2O3 using BCl3/Ar for the interface passivation layer of III-V MOS devices," Microelectronic Engineering, 110, 457 (2013).
- 60. K. S. Min, S. H. Kang, J. K. Kim, J. H. Yum, Y. I. Jhon, W. Todd Hudnall, C. W. Bielawski, S. K. Banerjee, G. Bersuker, M. S. Jhon, and G. Y. Yeom, "Atomic layer etching of BeO using BCl₃/Ar for the interface passivation layer of III-V MOS devices," Microelectronic Engineering, 114, 121 (2014).
- T. Meguro, M. Ishii, H. Kodama, M. Hamagaki, T. Hara, Y. Yamamoto, and Y. Aoyagi, "Layer-by-Layer Controlled Digital Etching by Means of an Electron-Beam-Excited Plasma System," Japanese Journal of Applied Physics Part 1-Regular Papers Short Notes & Review Papers, 29, 2216 (1990).
- 62. T. Meguro, M. Ishii, K. Kodama, Y. Yamamoto, K. Gamo, and Y. Aoyagi, "Surface Processes in Digital Etching of GaAs," Thin Solid Films, 225(1-2), 136 (1993).
- 63. M. Ishii, T. Meguro, H. Kodama, Y. Yamamoto, and Y. Aoyagi, "Study of Surface Processes in the Digital Etching of GaAs," Japanese Journal of Applied Physics Part 1-Regular Papers Short Notes & Review Papers, 31(7), 2212 (1992).
- 64. M. Ishii, T. Meguro, T. Sugano, K. Gamo, and Y. Aoyagi, "Digital Etching by Using a Laser-Beam - on the Control of Digital Etching Products," Applied Surface Science, 80, 104 (1994).
- 65. Y. Aoyagi, K. Shinmura, K. Kawasaki, T. Tanaka, K. Gamo, S. Namba, and I. Nakamoto, "Molecular Layer Etching of GaAs," Applied Physics Letters, 60(8), 968 (1992).
- 66. K. K. Ko and S. W. Pang, "Controllable Layer-by-Layer Etching of III-V Compound Semiconductors with an Electron-Cyclotron-Resonance Source," Journal of Vacuum Science & Technology B, 11(6), 2275 (1993).
- 67. W. S. Lim, S. D. Park, B. J. Park, and G. Y. Yeom, "Atomic layer etching of (100)/(111) GaAs with chlorine and low angle forward reflected Ne neutral beam," Surface & Coatings Technology, 202, 5701 (2008).
- 68. N. Otsuka, Y. Oyama, H. Kikuchi, J. Nishizawa, and K. Suto, "Digital etching of (001) InP substrate by intermittent injection of tertiarybutylphosphine in ultrahigh vacuum," Japanese Journal of Applied Physics Part 2-Letters, 37(12B), L1509
- 69. S. D. Park, C. K. Oh, J. W. Bae, G. Y. Yeom, T. W. Kim, J. I. Song, and J. H. Jang, "Atomic layer etching of InP using a low angle forward reflected Ne neutral beam," Applied Physics Letters, 89, 043109 (2006).
- 70. S. D. Park, C. K. Oh, W. S. Lim, H. C. Lee, J. W. Bae, G. Y. Yeom, T. W. Kim, J. I. Song, and J. H. Jang, "Highly selective and low damage atomic layer etching of InP/InAIAs heterostructures for high electron mobility transistor fabrication, Applied Physics Letters, **91**, 013110 (2007).
- T. W. Kim, D. H. Kim, S. D. Park, S. H. Shin, S. J. Jo, H. J. Song, Y. M. Park, J. O. Bae, Y. W. Kim, G. Y. Yeom, J. H. Jang, and J. I. Song, "A two-step-recess process based on atomic-layer etching for high-performance In_{0.52}Al_{0.48}As/In_{0.53}Ga_{0.47}As p-HEMTs," *IEEE Transactions on Electron Devices*, 55(7), 1577 (2008).
- 72. T. Sugiyama, T. Matsuura, and J. Murota, "Atomic-layer etching of Ge using an
- ultraclean ECR plasma," *Applied Surface Science*, **112**, 187 (1997).

 73. T. Matsuura, T. Sugiyama, and J. Murota, "Atomic-layer surface reaction of chlorine on Si and Ge assisted by an ultraclean ECR plasma," Surface Science, 402, 202 (1998).
- 74. W. S. Lim, Y. Y. Kim, H. Kim, S. Jang, N. Kwon, B. J. Park, J. H. Ahn, I. Chung, B. H. Hong, and G. Y. Yeom, "Atomic layer etching of graphene for full graphene device fabrication," *Carbon*, **50**(2), 429 (2012).
- 75. Y. Y. Kim, W. S. Lim, J. B. Park, and G. Y. Yeom, "Layer by Layer Etching of the Highly Oriented Pyrolythic Graphite by Using Atomic Layer Etching," Journal of the Electrochemical Society, 158(12), D710 (2011).
- 76. J. B. Park, W. S. Lim, B. J. Park, I. H. Park, Y. W. Kim, and G. Y. Yeom, "Atomic layer etching of ultra-thin HfO2 film for gate oxide in MOSFET devices," Journal of Physics D-Applied Physics, 42, 055202 (2009).
- 77. K. S. Min, C. Y. Kang, C. Park, C. S. Park, B. J. Park, J. B. Park, M. M. Hussain, J. C. Lee, B. H. Lee, P. Kirsch, H. H. Tseng, R. Jammy, and G. Y. Yeom, "A Novel Damage-Free High-k Etch Technique Using Neutral Beam-Assisted Atomic Layer Etching (NBALE) for Sub-32nm Technology Node Low Power Metal Gate/Highk Dielectric CMOSFETs," in 2009 IEEE International Electron Devices Meeting (IEEE, New York, 2009), pp. 406.
- 78. E. Vogli, D. Metzler, and G. S. Oehrlein, "Feasibility of atomic layer etching of polymer material based on sequential O2 exposure and Ar low-pressure plasmaetching," Applied Physics Letters, 102, 253105 (2013).
- Y. Horiike, T. Tanaka, M. Nakano, S. Iseda, H. Sakaue, A. Nagata, H. Shindo, S. Miyazaki, and M. Hirose, "Digital Chemical Vapor-Deposition and Etching Tech-

- nologies for Semiconductor Processing," Journal of Vacuum Science & Technology a-Vacuum Surfaces and Films, 8(3), 1844 (1990).
- 80. H. Sakaue, S. Iseda, K. Asami, J. Yamamoto, M. Hirose, and Y. Horiike, "Atomic Layer Controlled Digital Etching of Silicon," *Japanese Journal of Applied Physics Part 1-Regular Papers Short Notes & Review Papers*, **29**(11), 2648 (1990).
- 81. T. Engel, "Fundamental aspects of the reactions of thermal and hyperthermal F, F2. Cl, and Cl₂ with Si surfaces," Japanese Journal of Applied Physics Part 1, 35, 2403 (1996).
- 82. A. Szabo, P.D. Farrall, and T. Engel, "Reactions of Chlorine with Si(100) and Si(111) - Adsorption and Desorption-Kinetics," Surface Science, 312, 284
- 83. T. Matsuura, J. Murota, Y. Sawada, and T. Ohmi, "Self-limited layer-by-layer etching of Si by alternated chlorine adsorption and Ar+ ion irradiation," Applied Physics Letters, 63(20), 2803 (1993).
- 84. K. Suzue, T. Matsuura, J. Murota, Y. Sawada, and T. Ohmi, "Substrate Orientation Dependence of Self-Limited Atomic-Layer Etching of Si with Chlorine Adsorption and Low-Energy Ar⁺ Irradiation," Applied Surface Science, 82-3, 422 (1994).
- 85. S. D. Athavale and D. J. Economou, "Molecular-Dynamics Simulation of Atomic Layer Etching of Silicon," Journal of Vacuum Science & Technology a-Vacuum Surfaces and Films, 13(3), 966 (1995).
- 86. S. D. Athavale and D. J. Economou, "Realization of atomic layer etching of silicon,"
- Journal of Vacuum Science & Technology B, 14(6), 3702 (1996).
 B. J. Kim, S. H. Chung, and S. M. Cho, "Atomic layer etching of Si (100) for reducing etching damage," in Thin Film Materials, Processes, and Reliability (ELEC-TROCHEMICAL SOCIETY INC, Pennington, 2001), Vol. 2001, pp. 54.
- 88. B. J. Kim, S. Chung, and S. M. Cho, "Layer-by-layer etching of Cl-adsorbed silicon surfaces by low energy Ar⁺ ion irradiation," Applied Surface Science, 187(1-2), 124 (2002).
- 89. S. D. Park, K. S. Min, B. Y. Yoon, D. H. Lee, and G. Y. Yeom, "Precise depth control of silicon etching using chlorine atomic layer etching," Japanese Journal of Applied Physics Part 1-Regular Papers Short Notes & Review Papers, 44(1A), 389 (2005).
- 90. S. D. Park, C. K. Oh, D. H. Lee, and G. Y. Yeom, "Surface roughness variation during Si atomic layer etching by chlorine adsorption followed by an Ar neutral beam irradiation," Electrochemical and Solid State Letters, 8, C177 (2005).
- 91. C. K. Oh, S. D. Park, H. C. Lee, J. W. Bae, and G. Y. Yeom, "Surface analysis of atomic-layer-etched silicon by chlorine," Electrochemical and Solid State Letters, 10, H94 (2007).
- 92. H. J. Yun, T. H. Kim, C. B. Shin, C. K. Kim, J. H. Min, and S. H. Moon, "Comparison of atomic scale etching of poly-Si in inductively coupled Ar and He plasmas," Korean Journal of Chemical Engineering, 24(4), 670 (2007).
 93. T. Matsuura, Y. Honda, and J. Murota, "Atomic-order layer-by-layer role-share
- etching of silicon nitride using an electron cyclotron resonance plasma," Applied Physics Letters, 74, 3573 (1999).
- 94. N. Posseme, O. Pollet, and S. Barnola, "Alternative process for thin layer etching: Application to nitride spacer etching stopping on silicon germanium," Applied Physics Letters, 105, 051605 (2014).
- 95. S. Rauf, T. Sparks, P. L. G. Ventzek, V. V. Smirnov, A. V. Stengach, K. G. Gaynullin, and V. A. Pavlovsky, "A molecular dynamics investigation of fluorocarbon based layer-by-layer etching of silicon and SiO2," Journal of Applied Physics, 101, 033308
- 96. S. B. Wang and A. E. Wendt, "Control of ion energy distribution at substrates during plasma processing," Journal of Applied Physics, 88(2), 643 (2000).
- 97. R. Silapunt, A. E. Wendt, and K. H. R. Kirmse, "Ion energy control at substrates during plasma etching of patterned structures," Journal of Vacuum Science & Technology B, 25(6), 1882 (2007).
- 98. N. Schaepkens, G. S. Oehrlein, and J. M. Cook, "Effects of radio frequency bias frequency and radio frequency bias pulsing on SiO2 feature etching in inductively coupled fluorocarbon plasmas," Journal of Vacuum Science & Technology B, 18(2), 856 (2000).
- 99. D. Metzler, R. L. Bruce, S. Engelmann, E. A. Joseph, and G. S. Oehrlein, "Fluorocarbon assisted atomic layer etching of SiO2 using cyclic Ar/C₄F₈ plasma," Journal of Vacuum Science & Technology A, 32, 020603 (2014).
- 100. D. Metzler, C. Li, R. L. Bruce, S. Engelmann, E. A. Joseph, V. Godyak, and G. S. Oehrlein (to be published).
- 101. E. Hudson, V. Vidyarthi, R. Bhowmick, R. Bise, H. J. Shin, G. Delgadino, B. Jariwala, D. Lambert, and S. Deshmukh, "Highly selective etching of Silicon Dioxide Using Fluorocarbons"; AVS 61st International Symposium & Exhibition $(2014); http://www2.avs.org/symposium2014/Papers/Paper_PS2+TF-ThM2.html.\\$
- 102. F. Laermer and A. Schilp, "Method for anisotropic plasma etching of substrates," US Pat. 5498312, March-12 (1996).
- F. Laermer and A. Urban, "Challenges, developments and applications of silicon deep reactive ion etching," *Microelectronic Engineering*, 67–8, 349 (2003).
- 104. F. Roozeboom, B. Kniknie, A. M. Lankhorst, G. Winands, R. Knaapen, M. Smets, P. Poodt, G. Dingemans, W. Keuning, and W. M. M. Kessels, "A new concept for spatially divided Deep Reactive Ion Etching with ALD-based passivation," IOP Conf. Ser.: Mater. Sci. Eng., 41, 012001 (2012).
- 105. J. B. Park, W. S. Lim, S. D. Park, Y. J. Park, and G. Y. Yeom, "Etch Characteristics of TiO₂ Etched by Using an Atomic Layer Etching technique with BCl3 Gas and an Ar Neutral Beam," *Journal of the Korean Physical Society*, **54**(3), 976 (2009). 106. W. S. Lim, J. B. Park, J. Y. Park, B. J. Park, and G. Y. Yeom, "Low Damage Atomic
- Layer Etching of ZrO2 by Using BCl3 Gas and Ar Neutral Beam," Journal of Nanoscience and Nanotechnology, 9(12), 7379 (2009).
- R. A. Gottscho, C. W. Jurgensen, and D. J. Vitkavage, "Microscopic Uniformity in Plasma-Etching," Journal of Vacuum Science & Technology B, 10(5), 2133 (1992).

- 108. P. L. G. Ventzek, R. J. Hoekstra, and M. J. Kushner, "2-Dimensional Modeling of High Plasma-Density Inductively-Coupled Sources for Materials Processing, Journal of Vacuum Science & Technology B. 12(1), 461 (1994).
- 109. R. J. Hoekstra and M. J. Kushner, "Predictions of ion energy distributions and radical fluxes in radio frequency biased inductively coupled plasma etching reactors," Journal of Applied Physics, 79(5), 2275 (1996).
- 110. C. T. Gabriel and Y. Melaku, "Gate Oxide Damage in a High-Density Inductively-Coupled Plasma," Journal of Vacuum Science & Technology B, 12(1), 454 (1994)
- 111. P. K. Aum, R. Brandshaft, D. Brandshaft, and T. B. Dao, "Controlling plasma charge damage in advanced semiconductor manufacturing-challenge of small feature size device, large chip size, and large wafer size," IEEE Transactions on Electron Deces, 45(3), 722 (1998).
- 112. S. Rauf, A. Haggag, M. Moosa, and P. L. G. Ventzek, "Computational modeling of process induced damage during plasma clean," Journal of Applied Physics, 100,
- 113. J. P. Chang and J. W. Coburn, "Plasma-surface interactions," Journal of Vacuum Science & Technology A, 21(5), S145 (2003).
- 114. V. M. Donnelly, J. Guha, and L. Stafford, "Critical review: Plasma-surface reactions and the spinning wall method," Journal of Vacuum Science & Technology A, 29, 010801 (2011).
- 115. G. S. Oehrlein, R. J. Phaneuf, and D. B. Graves, "Plasma-polymer interactions: A review of progress in understanding polymer resist mask durability during plasma etching for nanoscale fabrication," *J. Vac. Sci. Technol. B*, **29**, 010801 (2011).
- 116. M. J. Kushner, "Hybrid modelling of low temperature plasmas for fundamental investigations and equipment design," Journal of Physics D-Applied Physics, 42, 194013 (2009).
- 117. SRIM simulation parameters used for simulations

99999 ions 0 ^{deg} injection	Density		Lattice Binding Energy	Surface Binding Energy	Displacement Energy
Si SiO ₂	2.321 2.65	Si (28.086) Si (28.086)	2 2	4.7 4.7	15 15
		O(15.999)	3	2.	28

Most of these values were obtained from http://www.srim.org/. The SiO₂ density was obtained from Greenwood, Norman N.; Earnshaw, Alan (1984). Chemistry of the Elements. Oxford: Pergamon Press. pp. 393-99. ISBN 0-08-022057-6.

- 118. M. E. Barone and D. B. Graves, "Chemical and physical sputtering of fluorinated silicon," J. Appl. Phys., 77, 3 (1995).
- 119. M. E. Barone and D. B. Graves, "Molecular dynamics simulations of plasma-surface chemistry," Plasma Sources Sci. Technol., 5, 2 (1996).
- 120. N. A. Kubota, D. J. Economou, and S. J. Plimpton, "Molecular dynamics simulations of low-energy (25-200 eV) argon ion interactions with silicon surfaces: Sputter yields and product formation pathways," J. Appl. Phys., 83, 8 (1998).
- 121. N. Laegreid and G. K. Wehner, "Sputtering Yields of Metals for Ar+ and Ne+ Ions with Energies from 50 to 600 ev," J. Appl. Phys., 32, 3 (1961).
- 122. Y. H. Lee, "Surface damage threshold of Si and SiO2 in electron-cyclotronresonance plasmas," J. Vac. Sci. & Technol. A, 10, 4 (1992).
- 123. N. Matsunami, Y. Yamamura, Y. Itikawa, N. Itoh, Y. Kazumata, S. Miyagawa, K. Morita, R. Shimizu, and H. Tawara, "Energy-Dependence of the Ion-Induced Sputtering Yields of Monatomic Solids," At. Data Nucl. Data Tables, 31, 1 (1984).

- 124. H. Ohta and S. Hamaguchi, "Molecular dynamics simulation of silicon and silicon dioxide etching by energetic halogen beams," J. Vac. Sci. & Technol. A, 19, 5 (2001).
- 125. D. J. Oostra, R. P. van Ingen, A. Haring, A. E. de Vries, and G. N. A. van Veen, 'Near threshold sputtering of Si and SiO2 in a Cl2 environment," Appl. Phys. Lett., 50, 21 (1987).
- 126. C. Steinbruüchel, "Universal energy dependence of physical and ion-enhanced
- chemical etch yields at low ion energy," *Appl. Phys. Lett.*, **55**, 19 (1989). S. Tachi, K. Miyake, and T. Tokuyama, "Chemical and Physical Sputtering in F⁺ Ion-Beam Etching of Si," *Jpn. J. Appl. Phys.*, **20**, 6 (1981).
- 128. Y.-Y. Tu, T. Chuang, and H. Winters, "Chemical sputtering of fluorinated silicon," Phys. Rev. B: Condens. Matter, 23, 2 (1981).
- S.-M. Wu, R. van de Kruijs, E. Zoethout, and F. Bijkerk, "Sputtering yields of Ru, Mo, and Si under low energy Ar⁺ bombardment," J. Appl. Phys., 106, 5 (2009).
- Y. Yamamura and H. Tawara, "Energy dependence of ion-induced sputtering yields from monatomic solids at normal incidence," At. Data Nucl. Data Tables, 62, 2
- E. Hechtl and J. Bohdansky, "Sputtering Behavior of Graphite and Molybdenum at Low Bombarding Energies," *J. Nucl. Mater.*, **123**, 1 (1984).
- 132. R. V. Stuart and G. K. Wehner, "Sputtering Yields at Very Low Bombarding Ion Energies," J. Appl. Phys., 33, 7 (1962).
- 133. J. P. Chang and H. H. Sawin, "Molecular-beam study of the plasma-surface kinetics of silicon dioxide and photoresist etching with chlorine," J. Vac. Sci. & Technol. B, 19.4 (2001).
- 134. J. M. E. Harper, J. J. Cuomo, P. A. Leary, G. M. Summa, H. R. Kaufman, and F. J. Bresnock, "Low-Energy Ion-Beam Etching," J. Electrochem. Soc., 128, 5 (1981)
- 135. G. V. Jorgenson and G. K. Wehner, "Sputtering Studies of Insulators by Means of Langmuir Probes," J. Appl. Phys., 36, 9 (1965).
- 136. P. J. Cumpson, J. F. Portoles, A. J. Barlow, and N. Sano, "Accurate argon clusterion sputter yields: Measured yields and effect of the sputter threshold in practical depth-profiling by X-ray photoelectron spectroscopy and secondary ion mass spectrometry," J. Appl. Phys., 114, 12 (2013).
- 137. S. S. Todorov and E. R. Fossum, "Sputtering of Silicon Dioxide near Threshold," Appl. Phys. Lett., 52, 5 (1988).
- 138. M. Schaepkens, R. C. M. Bosch, T. E. F. M. Standaert, G. S. Oehrlein, and J. M. Cook, "Influence of reactor wall conditions on etch processes in inductively coupled fluorocarbon plasmas," Journal of Vacuum Science & Technology A-Vacuum Surfaces And Films, 16, 2099 (1998).
- 139. S. J. Ullal, H. Singh, J. Daugherty, V. Vahedi, and E. S. Aydil, "Maintaining reproducible plasma reactor wall conditions: SF6 plasma cleaning of films deposited on chamber walls during Cl₂/O₂ plasma etching of Si," Journal of Vacuum Science & Technology A-Vacuum Surfaces And Films, 20(4), 1195 (2002).
- 140. G. Cunge, M. Kogelsehatz, and N. Sadeghi, "Influence of reactor walls on plasma chemistry and on silicon etch product densities during silicon etching in halogenbased plasmas," Plasma Sources Science & Technology, 13(3), 522 (2004).
- 141. G. Cunge, M. Kogelschatz, O. Joubert, and N. Sadeghi, "Plasma-wall interactions during silicon etching processes in high-density HBr/Cl₂/O₂ plasmas," Plasma Sources Science & Technology, 14(2), S42 (2005)
- 142. W. Y. Zhu, S. Sridhar, L. Liu, E. Hernandez, V. M. Donnelly, and D. J. Economou, "Photo-assisted etching of silicon in chlorine- and bromine-containing plasmas," Journal of Applied Physics, 115, 203303 (2014).
- 143. M. F. Doemling, N. R. Rueger, and G. S. Oehrlein, "Observation of Inverse Reactive Ion Etching (RIE) Lag for Silicon Dioxide Etching in Inductively Coupled Plasmas (ICP)," Appl. Phys. Lett., 68, 10 (1996).