Modeling of Reactive Ion Etching for Si/SiO₂ Systems

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Abstract
Molecular dynamics (MD) simulations have been used to study surface reaction dynamics of Si and SiO₂ etching by halogens. To perform classical MD simulations for SiO₂ etching reaction by halogens (Cl or F), we have constructed new sets of two- and three-body interatomic potential functions based on potential energy data obtained from ab initio quantum mechanical calculations of the electronic states. Etching yields for Si and SiO₂ targets are obtained from MD simulations for both beam etching and reactive ion etching (RIE). The obtained yields are in good agreement with experimental observations.

1 Introduction
Patterned etching of Si is one of the most important fabrication processes employed in the semiconductor chip manufacturing. To transfer desired submicron patterns to the Si surface efficiently, one needs to use highly selective etching processes. As the minimum dimensions of required etched patterns are now close to 0.1 µ under the current state-of-the-art manufacturing technologies and are diminishing rapidly, development of more reliable, high-throughput processes with good critical-dimension control is pursued aggressively in the semiconductor industry.

One of the most important issues for the control of critical dimensions in such processes is to understand the surface reaction mechanisms during actual processing conditions. In the present paper, we focus on Si and SiO₂ etching processes by plasmas containing halogen (Cl or F) atoms. Using molecular dynamics (MD) simulations, we attempt to elucidate details of etching processes under realistic etching conditions.

Classical MD simulations with empirical potentials have been used by various authors to analyze details of surface reaction dynamics [1]–[9]. Data obtained from such simulations can compensate for scarcity of some experimental data. As to beam etching and/or reactive ion etching (RIE) of Si by halogens, Stillinger-Weber (SW) type interatomic potentials [1, 2] are known to represent many aspects of surface reactions during etching processes with reasonable accuracy.

To extend such MD simulations to etching of SiO₂ surfaces by halogens, we have developed new potential sets for Si-O-F and Si-O-Cl systems [10], which are consistent with previously known interatomic potentials of SW type for Si.
[1], Si-Cl [2], Si-F [3], and Si-O [11] systems. As the SW-type potential functions include three-body interactions, we have determined ternary interactions by fitting a functional form to potential energy data obtained from *ab initio* calculations of electronic states of such systems.

## 2 Molecular Dynamics Simulation

The simulation method is now presented briefly. Atoms are normally injected to a target which is placed in a simulation cell with periodic boundary conditions imposed in the horizontal directions. The atoms in the bottom layer are rigidly fixed to prevent the drift of the entire simulation cell. Since energetic ions impinging on the surface are expected to be neutralized near the target surface due to an Auger emission process, we only consider charge-neutral atoms as impinging species. It is also assumed that the target surface is kept charge-neutral during the process.

After injection of each atom, we let the system evolve for 0.7 ps under the constant total-energy conditions. Most transient processes such as rapid release of the kinetic energy of the injected atom to the target are almost completed during this period. We then artificially cool the entire system (by decreasing the velocities of all atoms) for 0.3 ps to reduce the target temperature to the initial temperature (i.e., 300° K) in order to prevent the system from being unrealistically heated up.

![Fig. 1. Etching yields of the silicon target for Cl (○), F (□), and Ar (△) beam etching. The filled circles represent etching yields for a Cl RIE process with Ar bombardment.](image)

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