Numerical and experimental studies of the carbon etching in EUV-induced plasma

D.I. Astakhov,1 W.J. Goedheer,2 C.J. Lee,1 C.J. Lee,1 V.V. Ivanov,3 V.M. Krivtsun,3 O. Yakushev,3 K.N. Koshelev,3 D.V. Lopaev,4 and F. Bijkerk1

1XUV Group, MESA+ Institute for Nanotechnology, University of Twente, P.O. Box 217, 7500 AE Enschede, The Netherlands
2FOM Institute DIFFER - Dutch Institute for Fundamental Energy Research, PO Box 6336, 5600HH Eindhoven, the Netherlands, The Netherlands
3Institute for Spectroscopy RAS (ISAN), Fizicheskaya 5, Troitsk 142190, Russian Federation
4Skobeltsyn Institute of Nuclear Physics, Lomonosov Moscow State University, Leninskie Gory, Moscow 119991, Russian Federation

We have used a combination of numerical modeling and experiments to study carbon etching in the presence of a hydrogen plasma. We model the evolution of a low density EUV-induced plasma during and after the EUV pulse to obtain the energy resolved ion fluxes from the plasma to the surface. By relating the computed ion fluxes to the experimentally observed etching rate at various pressures and ion energies, we show that at low pressure and energy, carbon etching is due to chemical sputtering, while at high pressure and energy a reactive ion etching process is likely to dominate.

I. INTRODUCTION

Many vacuum mirror optical tools suffer from the build-up of carbon contamination due to cracking of hydrocarbons under powerful vacuum ultraviolet radiation [1]. The large absorption of EUV radiation by carbon becomes significant in the case of multi-element optical systems, where throughput can be greatly reduced by even a very thin layer of carbon contamination on the top of each mirror.

The problem of EUV induced carbon contamination has been addressed in a series of publications [2–5]. The reduction of carbon films in a hydrogen atmosphere or hydrogen plasma has also been extensively studied [6–10]. Despite numerous studies, however, it is still difficult to predict the carbon removal rate, because there are many contributing factors. Several aspects that significantly affect the carbon removal rate include: many different allotropes and compounds of the carbon (e.g. soft black or hard graphite), many different contributing reaction paths (e.g. physical sputtering, chemical sputtering, reactive ion etching etc.), and, last but not least, small admixtures to the background gas, which, while residing on the carbon surface, can produce reactive species once irradiated by EUV. Despite this complexity, experiments have shown that carbon etching can be achieved under certain EUV-induced plasma conditions. Nevertheless, it has proven difficult to fully understand the etch process, because the characteristics of the EUV-induced plasma are poorly known, and the plasma-surface interaction has many contributing factors [10].

In this paper, we use a model of the EUV-induced plasma to numerically analyse the fluxes from the plasma to the sample surface. Our model is a self-consistent 2D Particle-in-Cell model of the weakly ionized low pressure hydrogen plasma that is formed during the EUV pulse due to ionization by EUV photons and secondary electrons from the surface. As described in Section III, our model provides an accurate estimate of the ion flux composition and energy distribution. However, a considerable number of parameters required for accurate simulations are not well known, therefore, the modeling results were combined with experimental observations. With the aid of simulations, we show that the shape of the energy distribution function of the ion fluxes in the considered experimental setup are mainly defined by the setup geometry, background pressure, and externally applied bias voltage. The ion dose, on the other hand is sensitive to the variations of many other parameters (e.g. EUV dose, secondary electron yield, etc.).

By combining the computed energy distribution function of the ion flux with the experimentally measured ion dose, insight into the mechanism for carbon removal was gained. By analysing the differences in yield between EUV-induced plasma and surface wave discharge plasma experiments in combination with numerical simulations, we show that chemical sputtering dominates for low pressures and energies. It was found that the carbon removal yield for both the surface wave discharge and EUV-induced plasmas was similar in the overlapping energy range. Hence, the effect of the EUV radiation on carbon removal is found to be significantly smaller than was estimated previously [10].

II. EXPERIMENTAL SETUP

The ISAN EUV experimental setup is based on a tin EUV radiation source, which is a Z-pinch discharge plasma with 1500 Hz repetition rate, which has been described in detail elsewhere [11]. This source is a good tool for exploring EUV-induced surface processes over a large number of pulses (> 1 MShot). In brief, EUV radiation is introduced into a so-called "clean" chamber (see Fig. 1), separated from the source and collector optics by a Si:Zr spectral purity filter (SPF). The clean chamber is differentially pumped to pressure of $3.5 \times 10^{-8}$ torr. Under vacuum conditions, the background hydrocarbon carbon
growth rate was measured to be 0.4±0.2 nm/10 Mshot.

The diameter of the EUV beam at the sample was 5 mm. In addition to the direct beam, some scattered EUV radiation was also incident on the sample. The EUV pulse duration is about 100 ns (FWHM), with a tail, as described in [12].

The incident EUV power was measured using a sensitive thermo-couple attached to a thin copper disk. It was found that the radiation intensity was 0.13 W/cm² after the SPF and approximately 0.75 W/cm² without the SPF. The ratio of EUV intensities with and without the SPF corresponds to the calculated transmission of the 100 nm Si:Zr SPF filter over an energy range of 60 to 100 eV (see Fig 2).

The EUV intensity on the sample decreases with time because of carbon growth on the SPF filter and focusing optics. In later experiments, the EUV intensity was measured to be 0.1 W/cm² after the SPF.

During the experiments, the hydrogen pressure was set in the range of 2.8 Pa – 86.5 Pa. To ensure that each radiation pulse excited a plasma in an atmosphere dominated by hydrogen, hydrogen flowed through a liquid nitrogen trap (to remove water) and into the chamber at 100 liters×torr/minute.

To control the energy of the ion flux, the sample holder assembly was biased in the range of -200 – 0 V, while all other metallic electrodes were grounded. The samples consisted of carbon, deposited by magnetron sputtering on a silicon wafer to a thickness of ~30 nm. Each sample was exposed to the 10⁷ EUV pulses at a different combination of bias and hydrogen pressure. After exposure, the amount of carbon removed was measured by X-Ray fluorescence (XRF) (EDS) analysis and spectroscopic ellipsometry.

III. MODEL

A two dimensional particle in cell (PIC) model with rz geometry was used to model the experiment. Our model follows the general PIC scheme, described elsewhere [13].

Ionization is initiated by an EUV pulse, which directly ionizes the background gas, and produces electrons by photo emission from the SPF and the sample. This process leads to the formation of an EUV induced plasma in the chamber. The ionization and photo-emission process and their inclusion in the model is described below.

Although the plasma is continually re-ignited by the pulsed EUV radiation, we consider each EUV pulse to induce a plasma in a cold neutral gas that is at equilibrium, because the characteristic time of plasma decay in the given geometry is about ~ 20µs, which is much shorter than the 660 µs between pulses. The restoration of thermal equilibrium in the background gas is also much shorter than the time between pulses.

A. Chamber configuration

The configuration of vacuum chamber is presented in Fig.1. Because the chamber is axially symmetric, it is possible to include the structure of the internal chamber in the simulations. To accurately model the plasma dynamics when the sample is biased, the space between the SPF and the metal grid is included in our simulations. The grid is included in the model as a number of metallic rings. When a bias is applied to the sample, ions from the plasma, which is formed between SPF and metal grid, are pulled towards the sample.

B. Dielectric model

In the experiment, a dielectric mica diaphragm was used to prevent EUV radiation being incident on the sample holder structure, thereby decreasing electron emission from the sample holder structure. Therefore, allowing plasma parameters to be estimated from the discharge characteristics of the sample. In the model, this feature is included with a simple dielectric model: a dielectric is not conductive, it can accumulate charge and the secondary electron yield under EUV irradiation is order of magnitude lower than for the sample.

The electron reflection and secondary electron emission under electron impact (SEE) are included in the model. These two processes are combined into SEE with the probability \( P(E) \) defined as follows.

\[
P(E) = p_0 + \frac{E}{E_1}(1.0 - p_0) \tag{1}
\]

Here \( E \) is the energy of incoming electron. The parameters of the used mica are unknown, therefore, we choses \( E_1 = 45 \) eV and \( p_0 = 0.7 \), these parameters corresponds to linear fit for SEE yield for quarts in [14].
This approximation blends between significant slow electron backscattering for low energy electrons and secondary electron emission for higher impact energies. We do not use higher order approximations, since both effects are known to depend strongly on the surface conditions [14, 15].

This model for the dielectric is over simplified, but sufficient to allow us to to take the out of focus EUV radiation, which is incident on the diaphragm, into account in a consistent manner.

C. Length scales and grid resolution

The time evolution of one EUV pulse has two distinct stages: negative space-charge dominated during the beginning of the EUV pulse, and decay of the positively charged plasma after the pulse [16].

In order to model the negative space-charge dominated part of the plasma evolution, the potential well near the surface, which is formed by the electrons that have escaped to the volume, must be resolved. The length scale of the space-charge potential well depends on the energy distribution of photoelectrons from the surface and on the current which passes through the system.

For the purpose of estimation, it is possible to simplify the formulas from [17] and obtain

\[ z_m \simeq 0.1 \text{ cm} \times \frac{(T[\text{eV}])^{3/4}}{(I[\text{mA/cm}^2])^{1/2}} \]  

(2)

Here, \( T \) is the initial temperature of the emitted photoelectrons, \( I \) is the current density near the cathode, and \( z_m \) is the distance from the cathode to the bottom of the space charge potential well. For \( T \sim 1 \text{ eV} \) and \( I \sim 20 \text{ mA/cm}^2 \), one obtains \( z_m \sim 0.02 \text{ cm} \).

For the plasma dominated part of the plasma evolution we need to resolve the Debye length in the volume, and also have a finer grid near the surface to resolve the plasma sheath.

From the EUV intensity (\( \sim 0.02 \text{ mJ per pulse} \)) and background hydrogen pressure (2.8 Pa, 11.2 Pa), the plasma density can be estimated to be about \( 10^9 \text{ cm}^{-3} \). For \( T_e \sim 0.5 \text{ eV} \) and \( N_e \sim 10^9 \text{ 1/cm}^3 \) one obtains \( R_D \sim 0.015 \text{ cm} \), which is comparable to the space charge length scale estimated above. To achieve this, the grid is refined near the sample surface and SPF. The minimum grid cell size is \( 0.05 \times R_D \), and is gradually increased in steps of 5% until the bulk cell size of \( 0.5 \times R_D \) is reached. Several tests were performed to ensure that the chosen grid resolution does not affect the plasma dynamics.

D. photo-electron emission

The energy spectrum of photoelectrons emitted from the SPF and sample surface are calculated from [18]:

\[ P(E) \sim \frac{E}{(E + W)^4} \]  

(3)

Here \( P(E) \) is the probability of emitting an electron with energy, \( E \), from a surface with a work function, \( W \). For carbon \( W = 5 \text{ eV} \) [19]. For electrons emitted from the surface, we assume an angular dependence given by a cosine emission law [18]. For purpose of estimation, we used the same approximation for the spectrum of photoelectrons emitted from the dielectric.

E. EUV spectrum and photoionization

To calculate the distribution of ion species, the spectrum of the EUV radiation must be included in the model, along with the energy-dependent cross sections for each photoionization process. Direct photoionisation of hydrogen by EUV photons is included as two processes:

\[ \text{H}_2 + h\nu \rightarrow \text{H}_2^+ + e \]

\[ \text{H}_2 + h\nu \rightarrow \text{H}^+ + \text{H} + e \]

The photoionisation cross-sections were taken from [20].

In the experiment, the EUV spectrum was measured (green curve Fig. 2) and found to be in agreement with data from [12]. The EUV spectrum, however, was measured before reflection from the Zr collector mirrors, and transmission through the SPF.

The calculated transparency curve that was provided with the Si:Zr SPF was used to calculate the EUV spectrum after transmission through the SPF (See Fig. 2 blue curve).

The SPF transmission curve has a transmission of about 1% in the range of 20 – 30 eV, which is important because the photoabsorption cross-section is very large in this energy range (see Fig. 2 red curve).

The contribution due to radiation in the 20 – 30 eV range is difficult to quantify for two reasons: the intensity of the radiation varies significantly due to carbon growth on the SPF, and the accuracy of the SPF transmission curve in this energy range is unknown. Nevertheless, we include it because, even a small transmission will result in additional photoionization, which leads to a reduced space-charge potential barrier at low pressures (e.g. 2.8 Pa, 11.2 Pa).

Although the photoionization due to the 20 – 30 eV radiation range is significant compared to the 13 nm band, it has significance only for cases where no bias is applied to the sample. For cases with bias, the presence of this radiation leads to a decrease of the space-charge potential.

F. Cross-sections set

The cross-sections set used in the model consists of electron collisions with hydrogen and ion collisions with hydrogen. The collisions between plasma species and three body processes are neglected due to their low probability under the conditions considered here. To accu-
We use an approach similar to that described in [21]. The reaction probability is sampled using the integrated reaction cross-section. The differential cross-section data is used to determine the collision kinematics and energy redistribution between products. Electron elastic scattering and hydrogen electronic excitations, and angular scattering data is taken from [22]. For electron impact ionization of hydrogen, we use an experimentally determined doubly differential cross-section [23, 24]. The set of cross-sections for collisions between ions and hydrogen is based on [25] because this set provides good agreement with swarm data for ions in hydrogen. We neglect the formation of H\(^{-}\), because the cross-section of dissociative electron attachment is very low, and the density of vibrationally exited hydrogen molecules too low to make a significant contribution to the production of H\(^{-}\).

We make use of the procedure described in [26] to perform Monte Carlo collisions with the background gas. We tested the consistency of our implementation by modeling swarm experiments and found good agreement with experimental values [27] for the first Townsend electron ionization coefficient, the electron mobility, and for H\(^{+}\) and H\(_2\)\(^{+}\) mobility in hydrogen [28].

Let us begin with the analysis of the charge — bias voltage characteristics (Q — V), which were measured during the experiments. The ion dose on the sample per pulse for the range of pressures and bias voltages considered here was estimated from the experimental results as a difference between the collected charge for given pressure and bias voltage and the collected charge for vacuum conditions [10]. The measured curves are presented in Fig. 3. It is worth noting that these Q — V characteristics have some peculiarities.

Firstly, all the characteristics reach saturation, and the turning point for the almost all curves is approximately -75 V. If there is a significant contribution from ionization by the accelerated secondary electrons from the sample in the drift regime, one would expect a rapid increase of the collected charge for biases in the range of -200 .. -50 V. Secondly, the increase of the collected charge for -200 V bias does not depend linearly on pressure in the range of 2.8 Pa to 22.4 Pa, i.e. the collected charge increases less than two-fold while the pressure increases eight times. But, one would expect at least linear growth of the collected charge for biases in the range of -200 .. -50 V.

Secondly, the increase of the collected charge for -200 V bias does not depend linearly on pressure in the range of 2.8 Pa to 22.4 Pa, i.e. the collected charge increases less than two-fold while the pressure increases eight times. But, one would expect at least linear growth of the collected charge with pressure if the volume ionization provides a significant contribution to the total collected charge. Thus, the overall contribution of photoionization in the volume is small compared with the other factors.

Thirdly, for -200 V bias and 22.4 Pa background pressure, due to the photoionization process, all the external bias would be applied over a very small layer near the sample, thus, all the secondary electrons will gain 200 eV energy. The H\(_2\) ionization cross-section for this energy is
approximately $6 \cdot 10^{-17}$ cm$^2$, thus, in the space between the sample and the grid ($l \sim 2.5$ cm), these electrons will, on average, have 0.8 ionization inducing collisions. If we assume that, under vacuum conditions and -200 V bias, the SE current from the sample saturates, and we obtain that, just due to the direct electron induced ionization, without cascade process and photoionization, the collected charge should be greater than 1.8 nC. In the experiment, however, the collected charge is approximately 1.9 nC.

A. Average secondary electron yield

The above analysis suggests that the combination of EUV power per pulse, and effective secondary electron yield (SEY) produces approximately 1 nC of electrons from the sample.

It is instructive to estimate the total SE charge from the experimental parameters. From the EUV intensity, repetition rate and spot size, the average dose per pulse was approximately $\sim 0.017$ mJ. The secondary electron yield for carbon under EUV radiation is estimated to be approximately $\gamma_{SE} \sim 0.01$ for the photon energy range of 60 eV – 100 eV [2, 29], which leads to a saturation charge of 2 nC, which would lead to a significant disagreement between the simulated and measured charge bias characteristics. It is also worth noting that effective SEY from mica was small, since, in the experiment, the charge collected from a sample made from mica was measured to be an order of magnitude smaller than for the carbon sample.

However, the measured charge bias characteristics are the only experimental data that provides a reference point for the simulations of the experiment dynamics. Since the spatial distribution of EUV intensity is subject to an unknown systematic measurement error, and the SEY is known to vary widely, depending on the surface conditions, we chose to keep the product of incident EUV and SEY a constant, chosen to provide 1 nC total SE charge from the sample.

B. Role of dielectric ring

During the simulations we found that the mica accumulated charge over many EUV pulses, significantly effecting the local field distribution, and, hence changing the flux incident on the sample. If charging is neglected, the Q – V response under vacuum and 2.8 Pa conditions cannot be reproduced for any reasonable parameter values. The discrepancy is caused by the potential barrier near the sample surface, which is created by the negative space charge generated during the EUV pulse. Although the potential is low, it produces a significant effect because the SE energy spectrum (3) is strongly peaked at rather low energies: approximately $W/3 = 1.3$ eV.

The charge accumulation on the mica significantly increases the local field strength near the sample surface (see Fig. 4). This removes the space charge potential barrier for biases of -100V bias and higher under the conditions that we consider.

The charging of the mica should saturate, leading to an unchanging charge density distribution on the mica. To estimate the charge density, we simulated one hundred pulses under vacuum conditions for all experimentally applied bias values. In these calculations, the effective SEY from the mica is assumed to be about 0.001, which corresponds to the experimentally measured value.

This approach corresponds to the experimental procedure, since the Q – V characteristics were measured via averaging a large number of pulses for every combination of bias and pressure. No special means were used in the experiment to remove the accumulated charge from the mica between pulses.

For negative biases, simulations show that the mica potential rapidly reaches approximately 0 V potential. For these conditions, further charging is very slow, because (a) the electric field strength near the mica is very small, therefore, even a small amount of emitted charge creates a space charge potential barrier, and, (b), because the mica potential is close to zero, there is a small flux of electrons from the sample to the mica.

The continued slow variation in charge distribution is impractical to simulate, however. The final charge distri-
is also a non-negligible contribution from fast H$_2^+$ pressure, H$_3^+$ transition significantly on the pressure. For the 3 Pa case, the main ion is H$_3^+$, because the characteristic time for H$_2^+$ to H$_3^+$ conversion in 3 Pa H$_2$ is approximately 0.5 µs. With increasing pressure, H$_3^+$ becomes the main ion, as expected. There is also a non-negligible contribution from fast H$_2$ which is produced due to a resonant charge exchange reaction.

The maximum ion energy is larger than the applied bias voltage due to the build-up of the plasma potential. But the number of such fast ions is small, due to the fast plasma decay. For the same reason, the most energetic ion is H$_3^+$ because its small mass allows it to accelerate during the decay of the plasma potential.

It was observed, that contrary to the ion dose, the shape of the ion flux EDF was mainly defined by bias, pressure and the accumulated charge on the dielectric. However, as discussed in previous section, we expect that the equilibrium mica potential does not significantly differ from 0 V. For small variations of the mica potential (if all other parameters are the same), the variations of the EDF shape are small for all simulations where a sample bias was applied. Moreover, the EDF was barely sensitive to reasonable variations of other parameters.

This insensitivity is a consequence of the experimental procedure. The applied bias allows ions to be collected from the entire chamber volume. Hence, the characteristics of the plasma are important only for the short time when the sheath between plasma and sample is small. As ions are collected at the sample, the sheath size becomes larger and larger. Therefore, as time progresses, the instantaneous ion flux EDF starts to depend only on the bias potential, mica potential, and the distance that the ions travel between the plasma and the sample. This is because these parameters determine the maximum possible energy of the ions, while the background pressure determines the number of collisions (e.g. energy loss), and, hence, the average energy of the ion at the sample. Therefore, for the considered conditions, the shape of the ion flux EDF is mostly determined by the bias, mica potential, pressure, and the chamber geometry.

It is notable, that the mica charge, due to exposure to EUV radiation and plasma, leads to the ion flux being focused on the sample (see Fig. 7). The ion flux to the mica itself becomes negligible compared to the simulation with zero charge on the mica. Therefore, the measured ion doses should represent accurately the ion doses on the sample.

Hence, for the analysis of the experiments, it is reasonable to take the computed EDF shape, normed to the experimentally measured ion dose (if it is available). This approach allows us to compare the experimental results at different pressures.

V. ION FLUXES TO THE SAMPLE SURFACE

The removal of carbon from the surface material should be directly dependent on the energy distribution function (EDF) of the ion flux incident on the surface. Conditions of the experimental study of carbon cleaning in the ISAN EUV experiment are summarized in Table I. We computed the plasma conditions, ion fluxes, and ion EDF for these conditions. The simulated time for all cases was 10 µs. For 2.8 Pa – 11.2 Pa and biases -200 V – -100 V, the plasma had completely decayed and all ions were collected during the simulation time. But, for the 60 Pa and 86.5 Pa cases some plasma was still left in the simulation domain.

The EDFs, integrated over the simulation window, and averaged over the sample surface, are presented in Fig. 5. The composition of the computed ion flux depends significantly on the pressure. For the 3 Pa case, the main ion is H$_3^+$, because the characteristic time for H$_2^+$ to H$_3^+$ conversion in 3 Pa H$_2$ is approximately 0.5 µs. With increasing pressure, H$_3^+$ becomes the main ion, as expected. There is also a non-negligible contribution from fast H$_2$ which is produced due to a resonant charge exchange reaction.

The maximum etched depths and total etched volumes per exposure of $10^7$ pulses for different pressures and biases are presented in Table I. The etched volumes are the integrals over etch profiles, which were determined by XRF measurement (see Fig. 8). In the cases of 60 Pa and 86.5 Pa pressure, the etch profile was only measured at the center of the EUV spot. We assumed, for these
two cases, a uniform etching profile to estimate the carbon removal yield. Hence, for these cases, the amount of removed carbon is most probably overestimated.

As was discussed in the previous section, we used the experimentally measured ion doses for yield estimation. However, for some experiments, the ion dose was not measured. In these cases, we used the ion doses obtained from simulation results. This approach is valid at high pressures because the uncertainty in the simulations is on the order of 30% (see Fig. 3).

For all samples, the average carbon removal yield was larger than 0.2 carbon atoms per ion. Thus, physical

Figure 5: Energy distribution function of ion and fast neutrals flux to the surface for 3 Pa (top) and 11.2 Pa (bottom) for -100 V bias. The fluxes were integrated over time and over sample surface and sum of fluxes normalized to unity. Note that $H_3^+$ spike is on the same position over energy in spite of the increase of pressure.

Figure 6: Comparison of the shape of the ion flux EDF obtained with pre-charged dielectric mica for different pressures. The ion fluxes were integrated over time and over sample surface, summed over ion types and normalized to unity. The carbon removal yield curve corresponds to surface wave discharge plasma (SWD) from [10]. The dotted part of the yield curve corresponds to linear extrapolation.

Figure 7: Comparison of the ion flux spatial distribution obtained with pre-charged mica and with zero charge on the mica. The ion fluxes were integrated over time and energy. Both distribution are normed to one if integrated over sample and dielectric mica surface areas. The conditions are 3 Pa background pressure and -100 V bias on the sample. The charge on the mica focuses the ion flux on the sample. Other combinations of pressure and bias show similar behavior.
In the case of experiments with -50 V and -100 V biases, the results can be compared with carbon etching in a surface wave discharge plasma [10], where similar magnetron deposited carbon on silicon substrate samples were used. The carbon removal yield for experiments with biases of -50 V and -100 V was estimated by convolving the simulated EDF with the energy dependent yield, as measured in the SWD experiments (see Fig. 6 and column "SWD recomputed" in Table I). In spite of \( \text{H}_4^+ \) being the main ion in SWD plasma, we used the same yield for estimation, since we do not expect dramatic difference between \( \text{H}_4^+ \) and \( \text{H}_2^+ \) for carbon removal. These estimates show that the yields found in EUV experiments with low biases agree with the recomputed yield from the SWD experiment. The margin of error is, however, large enough that we cannot completely exclude direct influence from EUV radiation.

### VII. CONCLUSION

The evolution of an EUV induced hydrogen plasma was simulated. The simulations, due to their close coupling to experimental conditions, allowed the magnitude, composition, and energy spectrum of the flux from the plasma to the surface to be estimated. Our model successfully computes the charge bias characteristics of the sample to an accuracy of a factor of two. This was used to quantitatively compare carbon etch rates between different EUV plasma conditions, as well as compare carbon etching under EUV-induced plasma to etching under SWD plasmas. In addition, the model describes the focusing effect of the dielectric surround, which allows the measured carbon etch profiles to be better understood in a quantitative manner.

It was observed that the carbon etching mechanism at low bias and pressure was different than that at high bias and pressure. For the higher energy range, the carbon re-

---

**Table I: Experimental conditions and carbon removal rate**

| Pressure (Pa) | Bias (V) | EUV Power (W/cm²) | Max Etched Depth (nm/10⁷ pulse) | Etched Volume (10⁻⁶ cm³/10⁷ pulse) | Ion Dose (nC/pulse) | Average Yield (C atom/ion) |
|--------------|----------|-------------------|-------------------------------|---------------------------------|--------------------|--------------------------|
| 3.0          | -50      | 0.11              | 1.7 ± 0.5                     | ~ 0.03                          | ~ 0.1              | 0.5                      | 0.4                      |
| 3.0          | -100     | 0.11              | 5.7 ± 0.5                     | ~ 0.1                           | ~ 0.25             | 0.6                      | 0.5                      |
| 3.0          | -200     | 0.13              | 14 ± 0.5                      | ~ 0.3                           | ~ 0.25             | 1.9                      | —                        |
| 11.2         | -200     | 0.13              | 30 ± 0.5                      | ~ 0.5                           | ~ 0.5              | > 1.6                    | —                        |
| 60           | -100     | 0.13              | 23 ± 0.5                      | ~ 1.2                           | > 2.2              | < 0.9                    | 0.5                      |
| 60           | -200     | 0.1               | 27 ± 0.5                      | ~ 1.4                           | > 2.3              | ~ 1.0                    | —                        |
| 86.5         | -100     | 0.2               | 19 ± 0.5                      | ~ 1                             | > 3.0              | < 0.5                    | 0.4                      |

a) The substrate was exposed, thus only lower boundary for cleaning rate could be estimated.
b) Estimated from measured Q – V characteristics (see Fig. 3) as the difference between the collected charge for a given pressure and bias voltage, and the collected charge for vacuum conditions.
c) The carbon removal was measured only in the center of the exposed area. However, an SEM image of the EUV exposed area showed no features. Therefore, for the purposes of estimating the carbon removal rate, we assume that the carbon was etched uniformly over the exposed area of the sample.
d) Experimental Q – V values were missing, therefore the simulated ion dose were used for estimation. Only ions which hit the sample during the simulated period are included. No corrections were made for the plasma which was present in the chamber after the end of the simulation.
e) This value is only an order of magnitude estimation, because all carbon was removed from the sample.
moval yield in EUV-induced plasma was larger than one carbon atom per ion. Most probably, etching is dominated by a reactive ion etching process, which may be due to the production of methane radicals (e.g. CH$_2$) that can desorb under energetic ion flux.

However, at low bias energies, the etch rates found in EUV-induced plasma agree with those found in SWD plasma to within model and measurement uncertainties. However, our study does not exclude EUV enhancing etch rates at low biases or having a small (< 0.2 C/ion) enhancing effect over the range of biases and pressures that we studied. Therefore, experiments to determine the effect of EUV in the limit of low-to-no bias are required.

References

[1] K. Boller, R. P. Haelbich, H. Hogrefe, W. Jark, and C. Kunz. Investigation of carbon contamination of mirror surfaces exposed to synchrotron radiation. Nuclear Instruments and Methods in Physics Research, 208(1-3): 273–279, 1983. ISSN 0167-5087. doi:10.1016/0167-5087(83)91134-1.

[2] Jeremy Hollenshead and Leonard Klebanoff. Modeling extreme ultraviolet/H2O oxidation of ruthenium optic coatings. Journal of Vacuum Science & Technology B: Microelectronics and Nanometer Structures, 24(1):118–130, 2006. doi:10.1116/1.2150225.

[3] Takahiro Nakayama, Akira Miyake, Hiromitsu Takase, Shigeru Terashima, Takashi Sudo, Yutaka Watanabe, and Yasuaki Fukuda. Analysis of carbon deposition on multilayer mirrors by using two different beamlines. In Frank M. Schellenberg and Bruno M. La Fontaine, editors, Alternative Lithographic Technologies, volume 7271, pages 72713P–8, San Jose, CA, USA, 2009. SPIE.

[4] H Shin, J Sporre, R Raju, and D Ruzic. Reflectivity degradation of grazing-incident EUV mirrors by EUV exposure and carbon contamination. Microelectronic Engineering, 86(1):99–105, January 2009. ISSN 01679317. doi:10.1016/j.mee.2008.10.009.

[5] David J. Davis, Georgios Kyriakou, Robert B. Grant, Mintcho S. Tikhov, and Richard M. Lambert. To-ward the in situ remediation of carbon deposition on ru-capped multilayer mirrors intended for EUV lithography: exploiting the electron-induced chemistry. The Journal of Physical Chemistry C, 111(33):12165–12168, 2007. doi:10.1021/jp074766y.

[6] C. Hopf, A. von Keudell, and W. Jacob. Chemical sputtering of hydrocarbon films. Journal of Applied Physics, 94(4):2373–2380, August 2003. ISSN 00218979. doi:10.1063/1.1594273.

[7] Jürgen Kiippers. The hydrogen surface chemistry of carbon as a plasma facing material. Surface Science Reports, 22(7-8):249–321, 1995. ISSN 0167-5729. doi:10.1016/0167-5729(95)80002-1.

[8] Shengguang Liu, Jizhong Sun, Shuyu Dai, Thomas Stirner, and Dezhen Wang. A general model for chemical erosion of carbon materials due to low-energy h+ impact. Journal of Applied Physics, 108(7):073302, 2010. ISSN 00218979. doi:10.1063/1.3485821.

[9] Bhavin N. Jariwala, Cristian V. Ciobanu, and Sumit Agarwal. Atomic hydrogen interactions with amorphous carbon thin films. Journal of Applied Physics, 106(7): 073305, 2009. ISSN 00218979. doi:10.1063/1.3228305.

[10] A. Dolgov, D. Lopaev, T. Rachimova, A. Kovalev, A. Vasilyeva, C. J. Lee, V. M. Krivtson, O. Yakushev, and F. Bijkerr. Comparison of h2 and he carbon cleaning mechanisms in extreme ultraviolet induced and surface wave discharge plasmas. Journal of Physics D: Applied Physics, 47(6):065205, 2014. doi:10.1088/0022-3772/47/6/065205.

[11] A Dolgov, O Yakushev, A Abrikosov, E Snegirev, V M Krivtson, C. J. Lee, and F. Bijkerr. Extreme ultraviolet (EUV) source and ultra-high vacuum chamber for studying EUV-induced processes. Plasma Sources Science and Technology, 24(3):035003, June 2015. ISSN 0963-0252, 1361-6595. doi:10.1088/0963-0252/24/3/035003.

[12] Vivek Bakshi. EUV sources for lithography. SPIE Press, Bellingham, Wash., 2006. ISBN 978-1-61538-714-9 1-61538-716-7 978-0-8194-5845-2 0-8194-5845-7.

[13] Charles K Birdsal and A. Bruce Landgon. Plasma physics via computer simulation. McGraw-Hill, New York, 1985. ISBN 0-07-003571-5 978-0-07-003571-7.

[14] A. Dunaevsky, Y. Raitses, and N. J. Fisch. Secondary electron emission from dielectric materials of a hall thruster with segmented electrodes. Physics of Plasmas, 10(6):2574, 2003. ISSN 1070664X. doi:10.1063/1.1568344.

[15] H. A. Fowler and H. E. Farnsworth. Reflection of very slow electrons. Physical Review, 111(1):103–112, 1958. doi:10.1103/PhysRev.111.103.

[16] M. H. L. van der Velden, W. J. M. Brok, J. J. A. M. van der Mullen, W. J. Goedheer, and V. Banine. Particle- and field-driven plasma in-cell monte carlo simulations of an extreme ultraviolet light source. Physical Review E, 73(3):036406, March 2006. doi:10.1103/PhysRevE.73.036406.

[17] Irving Langmuir. The effect of space charge and initial velocities on the potential distribution and thermionic current between parallel plane electrodes. Physical Review, 21(4):419–435, April 1923. doi:10.1103/PhysRev.21.419.

[18] Burton L. Henke, Jerel A. Smith, and David T. Acknowledgments

The authors would like to thank Tatiana Rakhimova (Skobeltsyn Institute of Nuclear Physics) for helpful discussion.

This work is part of the research program “Controlling photon and plasma induced processes at EUV optical surfaces (CP3E)” of the “Stichting voor Fundamenteel Onderzoek der Materie (FOM)” which is financially supported by the Nederlandse Organisatie voor Wetenschappelijk Onderzoek (NWO). The CP3E programme is co-financed by industrial partners, including ASML (Veldhoven), and the AgentschapNL through the Catrene EX-EPT program.
Attwood. 0.1–10-keV x-ray-induced electron emissions from solids—models and secondary electron measurements. *Journal of Applied Physics*, 48:1852, 1977. ISSN 00218979. doi:10.1063/1.323938.

[19] David R Lide. *CRC handbook of chemistry and physics, 2003-2004: a ready-reference book of chemical and physical data.* CRC Press, Boca Raton, Fla., 2003. ISBN 0-8493-0484-9 978-0-8493-0484-2.

[20] Y. M Chung, E. M Lee, T. Masuoka, and James A. R Samson. Dissociative photoionization of h2 from 18 to 124 eV. *The Journal of Chemical Physics*, 99(2):885–889, July 1993. ISSN 00219606. doi:10.1063/1.465352.

[21] M. S. Mokrov and Yu P. Raizer. Monte carlo method for finding the ionization and secondary emission coefficients and i–v characteristic of a townsend discharge in hydrogen. *Technical Physics*, 53(4):436–444, April 2008. ISSN 1063-7842, 1090-6525. doi:10.1134/S1063784208040075.

[22] M. J. Brunger and S. J. Buckman. Electron–molecule scattering cross-sections. i. experimental techniques and data for diatomic molecules. *Physics reports*, 357(3):215–458, 2002.

[23] T. W. Shyn, W. E. Sharp, and Y.-K. Kim. Doubly differential cross sections of secondary electrons ejected from gases by electron impact: 25-250 eV on h2. *Physical Review A*, 24(1):79–88, July 1981. doi:10.1103/PhysRevA.24.79.

[24] M. E. Rudd, K. W. Hollman, J. K. Lewis, R. R. Porter, and E. L. Fagerquist. Doubly differential electron-production cross sections for 200–1500-eV e+H2 collisions. *Physical Review A*, 47(3):1866–1873, March 1993. doi:10.1103/PhysRevA.47.1866.

[25] T. Šimko, V. Martičovský, J. Bretagne, and G. Gousset. Computer simulations of h+ and h3+ transport parameters in hydrogen drift tubes. *Physical Review E*, 56(5):5908-5919, November 1997. doi:10.1103/PhysRevE.56.5908.

[26] Kenichi Nanbu. Simple method to determine collisional event in monte carlo simulation of electron-molecule collision. *Japanese Journal of Applied Physics*, 33(Part 1, No. 8):4752–4753, August 1994. ISSN 0021-4922. doi:10.1143/JJAP.33.4752.

[27] J. Dutton. A survey of electron swarm data. *Journal of Physical and Chemical Reference Data*, 4(3): 577–856, July 1975. ISSN 0047-2689, 1529-7845. doi:10.1063/1.555525.

[28] E. Graham, D. R. James, W. C. Keever, D. L. Albritton, and E. W. McDaniel. Mobilities and longitudinal diffusion coefficients of mass-identified hydrogen ions in h2 and deuterium ions in d2 gas. *The Journal of Chemical Physics*, 59(7):3477–3481, October 1973. ISSN 00219606. doi:10.1063/1.1680505.

[29] B. V. Yakshinskiy, R. Wasielewski, E. Loginova, and Theodore E. Madey. Carbon accumulation and mitigation processes, and secondary electron yields of ruthenium surfaces. In *Proceedings of SPIE*, pages 65172Z–65172Z–11, San Jose, CA, USA, 2007. doi:10.1117/12.711785.

[30] YASUNORI YAMAMURA and HIRO TAWARA. ENERGY DEPENDENCE OF ION-INDUCED SPUTTERING YIELDS FROM MONATOMIC SOLIDS AT NORMAL INCIDENCE. *Atomic Data and Nuclear Data Tables*, 62(2):149–253, March 1996. ISSN 0092-640X. doi:10.1016/0092-640X(95)00005.

[31] A. Gao, E. Zoethout, J.M. Sturm, C.J. Lee, and F. Bijkerk. Defect formation in single layer graphene under extreme ultraviolet irradiation. *Applied Surface Science*, 317:745–751, October 2014. ISSN 01694332. doi:10.1016/j.apsusc.2014.08.177.
Figure 8: Experimentally observed and simulated carbon removal as function of pressure, bias and radius. For the conditions of 11.2 Pa, -200V the carbon layer was completely removed, exposing the substrate, at the center of the EUV spot. The etch yield for this case was set to the 1.6 C atom/ion as in table I. For the conditions of 60 Pa, -200V, the carbon layer was completely removed from entire area exposed to plasma, hence the case is not shown. The simulated profiles for -50 V and -100 V bias were computed via convolution of the simulated ions fluxes with the yield from SWD experiments Fig. 6. The profiles were normed to the ion dose per pulse from table I. The EUV dose was $10^7$ pulses per sample in all cases, which is approximately 170 J. The EUV radiation was concentrated in a spot with radius approximately 2 mm. The detailed intensity distribution of the EUV spot is not accurately known.