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In situ infrared absorption spectroscopy of dusty plasmas

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In situ, time-resolved Fourier transform infrared spectroscopy was used to study particulate formation in rf discharges in mixtures of silane, argon, and nitrogen. The spectra were taken at a maximum rate of 20 Hz. The discharge conditions were chosen such that previous calibrations of the time evolutions of particle size and density could be used. The measurements indicate that the onset of the solid-state vibrational absorptions of the SiH and SiH2 bands only takes place after the nucleation and coagulation phase have finished; it coincides with the previously predicted start of the deposition of amorphic hydrogenated silicon on the particles. The dissociation of the silane feed gas is found to be in the range of 30%, and its time development suggests that also the large-scale dissociation of silane only starts after the coagulation phase. This is in agreement with previously observed trends for the electron temperature. If silicon particles are grown in the plasma, and the silane flow is stopped, the Si particles stay trapped in the glow. The infrared measurements, however, show that they almost completely oxidize: the SiH/SiH2 vibrations disappear and a strong SiO vibration appears. If nitrogen gas is allowed into the plasma, the SiO vibration is replaced by a SiN vibration. © 1996 American Vacuum Society.

I. INTRODUCTION

A great effort has been made lately to clarify the formation mechanism of macroscopic particulates in a rf plasma because of their destructive role in surface processing like deposition, etching, or sputtering. More recently, interest in the possibility to generate particles with a well defined size distribution and controllable chemical composition has emerged. Examples of application of those plasma generated particles can be found within the field of electronics (crystalline silicon particles), catalysis and ceramics (particles with a core of silicon nitride, oxide, or carbide covered with a metal or metal oxide film), and lighting (covering fluorescent powders with a protective coating). A better understanding of the particle formation and growth process will make it possible to enhance process and product control.

In order to elucidate the actual formation process in a given gas mixture and configuration it is necessary to monitor the particulate composition during the growth process. If the clusters contain infrared active bonds, this analysis can be performed using Fourier transform infrared (FTIR) absorption spectroscopy. In this work we apply this technique to particulates formed in rf discharges containing argon, silane, methane, and nitrogen.

II. EXPERIMENT

The experiments were performed in a 13.56 MHz capacitively coupled plasma in a planar 12-cm-diam parallel plate configuration. The distance between the aluminum electrodes is 5 cm. The input power can be varied between 0 and 100 W. The gas pressure and flow can be varied independently from 5 to 500 mTorr and from 0 to 10 sccm, respectively, using a throttle valve in the pumping line. The gases are fed through MKS mass flow controllers and introduced homogeneously through a shower head in the rf electrode. The opposite, grounded electrode consists of a grid to allow the particles to be removed from the “plasma box” with the gas flow. In order to obtain a good base pressure an oil diffusion pump is used. During plasma operation just a rotary pump is applied. The discharge and the growth of silicon particles in it are well characterized. For certain parameter settings, the time evolution of the particle size, the particle density, and the electron density and temperature have been determined previously (see Fig. 1). In short, one can distinguish three phases: (1) nucleation: almost immediately after plasma ignition the particle density is very high (10⁹ cm⁻³) and the average size is very small (a few nm); (2) coagulation: in the next 5 s these precursors coagulate to form larger particles (up to 50 nm), meanwhile, the density decreases (down to 10⁷ cm⁻³); (3) a-Si:H deposition: after 5 s there is a steady growth of the particles, presumably caused by the deposition of amorphic hydrogenated silicon (a-Si:H), without the density changing much. In this work we have used those calibrated settings (120 mTorr total pressure, 4 mass percent of silane in argon, 3 W of rf power).

For the infrared measurements NaCl windows have been mounted. The spectra have been collected using a Bruker IFS-66 Fourier interferometer with a glowbar as a continuous light source. The FTIR is basically a Michelson interferometer, supplied with a fixed and a moving mirror. This provides a time varying interference signal for all wavelengths. The parallel beam exiting the spectrometer is directed
through the plasma (single pass, see Fig. 2). Then it is focused with an off-axis paraboloid mirror onto a mercury–cadmium–telluride (MCT) detector. A Fourier transform of the detector signal yields a spectrum in the wave number range of 600–5000 cm\(^{-1}\) with a spectral resolution of 4 cm\(^{-1}\). In order to avoid too strong an appearance of the absorption bands of water and \(\text{CO}_2\), that are always present in the ambient, the infrared (IR) light beam was enclosed in plastic tubing that was continuously flushed with nitrogen gas. Nevertheless, sometimes the (clearly recognizable) absorptions of \(\text{H}_2\text{O}\) and \(\text{CO}_2\) could not be completely avoided. In order to avoid arcing of the discharge in the slits that were made in the plasma box to allow access of the IR beam, in some cases (especially at high power) those slits were covered with a grid.

Since the growth of the particles is very reproducible, the following measurement sequence was performed. First, the gas flows and pressure are set and allowed to stabilize. Then the discharge is ignited, and IR absorption spectra are taken at a rate of 20 Hz for several seconds. The first spectrum serves as reference spectrum and is taken before the discharge is ignited. The rf power is switched on and off by a digital port of the FTIR in order to synchronize the experiment. After the rf power is switched off, a few seconds are taken to allow the grown particles to be blown out of the plasma box. Then the rf power is switched on again, and the sequence is repeated. The series of IR spectra are co-added each time. By repeating this sequence a few thousand times, the signal to noise ratio is brought up to a level that allows absorptions of \(10^{-3}\) to be detected. The experiments typically take a total time of a few hours and result in one sequence of averaged IR spectra. Since the spectra are taken at intervals of 50 ms, the IR absorption by molecules as well as particles in the plasma can be monitored with the same time resolution.

### III. RESULTS AND DISCUSSION

First, a note concerning the labels of all vertical axes of displayed spectra. They all read “transmission ratio” because they are obtained by dividing two (transmission) spectra as directly measured by the FTIR. For each figure it is then indicated separately in the text how to interpret the direction (upwards or downwards) of the various peak structures (visible).

Figure 3 shows the absorption spectrum of pure silane gas. It was obtained by dividing the spectrum with silane gas (measured last) on the spectrum without silane gas (measured first). Clearly recognizable are the two \(\text{SiH}_4\) bands around 900 and around 2100 cm\(^{-1}\). The intensity of the \(Q\)-branch of \(\text{SiH}_4\) at 2187 cm\(^{-1}\) can be used as a measure for the concentration of silane gas. An absorption of 3% of this peak corresponds to the partial pressure of silane without plasma (4.8 mTorr). The strong bands around 1500 and 3700...
cm\(^{-1}\) are caused by water. Apparently the amount of water in the optical system or in the plasma chamber has decreased slightly between the two measurements. In Fig. 4 the absorption spectrum of a silane discharge (5 min) with powders is given. This spectrum is obtained by dividing the spectrum with plasma (and powders) on the spectrum without plasma (but with the silane gas). Bands that point upwards indicate an increase in absorption; bands that point downwards indicate a decrease of absorption upon plasma ignition. Apart from the already known silane absorptions (which now point downwards with about 1% absorption in the \(Q\)-band, indicating a decrease in density when the plasma is switched on, corresponding to a dissociation degree of 30%), one can recognize the solid-state Si–H vibration around 2010 and Si–H\(_2\) around 2110 cm\(^{-1}\), a band around 3000 cm\(^{-1}\) (maybe, C–H, could be carbon contamination coming from pump oil and incorporated in the particles), and a band around 650 cm\(^{-1}\) (Si–H). The spectra of solid-state SiH/SiH\(_2\) and gas phase SiH\(_4\) overlap partially.

When the spectra are taken time resolved (one spectrum each 2 s, first spectrum without plasma, then each 2 s with plasma on; then the ratio of the spectrum without plasma and the spectra with plasma are calculated), the time evolution of the SiH/SiH\(_2\) solid-state and SiH\(_4\) gas phase absorptions as a function of time can be obtained. Immediately after switch on (Fig. 5) no solid-state absorption and only a small change in the silane concentration are visible. After 40 s (Fig. 6) the Si–H\(_2\) bands at 650 and 2110 cm\(^{-1}\) become visible. Also one again sees the change in the SiH\(_4\) absorption. Furthermore, a continuous increase towards larger wave numbers (shorter wavelength) is observed. This is caused by Mie–Rayleigh scattering of the IR beam by the particles. In Fig. 7 the silane gas contribution in the spectrum of Fig. 5 is eliminated by subtracting the (scaled) silane spectrum (Fig. 3); now a clean Si–H/Si–H\(_2\) solid-state vibrational band is visible. The deformations around 1500 and 3700 cm\(^{-1}\) in Fig. 7 are caused by the contributions of H\(_2\)O (which is present in the optical path outside the plasma reactor) which are also visible in the used silane spectrum. In Figs. 8 and 9 the time evolution of the directly measured intensity at 2110 (Si–H\(_2\)) and 2187 cm\(^{-1}\) (silane \(Q\)-branch absorption) is given. It has been verified that the displayed time dependencies are not caused by changes in the IR background signal or the scattered light by comparing the time dependencies with results obtained at frequencies that are close to but not in the Si–H\(_2\) or the SiH\(_4\) absorption regions. The time evolution of the 2110 cm\(^{-1}\) SiH vibration is similar to the one of SiH\(_2\) and is not displayed separately. For 2110 cm\(^{-1}\), a decrease of the measured intensity corresponds to an increase in SiH\(_2\) absorption. For 2187 cm\(^{-1}\) an increase of the mea-

![Fig. 5. The absorption spectrum of a silane plasma immediately after plasma ignition.](image)

![Fig. 6. The absorption spectrum of a silane plasma after 40 s of plasma operation.](image)

![Fig. 7. The absorption spectrum of a silane plasma after 40 s of plasma operation.](image)

![Fig. 8. The absorption spectrum of a silane plasma after 40 s of plasma operation.](image)

![Fig. 8. Evolution of the measured intensity at the SiH\(_2\) solid-state absorption wave number (2110 cm\(^{-1}\)) as a function of time after plasma ignition.](image)
sured intensity corresponds to a decreased SiH₄ absorption and hence to an increased SiH₄ dissociation. One can see that the Si–H₂ solid-state absorption only appears after about 5 s, and then increases rapidly to an equilibrium value. If one compares this to the time evolutions of particle size and density given in Fig. 1, one can conclude that the solid-state absorption of SiH/SiH₂ only sets in after the coagulation of the clusters into 50 nm particulates has finished. Apparently, before that point there is no amorphous silicon deposition on the particles. One can also argue that it may not be appropriate to speak about a “solid-state phase” for particles smaller than 50 nm. Figure 9 indicates that the dissociation degree of silane only becomes substantial after 5 s. This is consistent with previous observations using Langmuir probe techniques and emission spectroscopy, which indicate that the electron temperature and electron density also only show a substantial increase and decrease respectively at that point.

In Fig. 10 the spectrum of Si powders trapped in an argon plasma is shown. The powders were produced in a 5 min Ar/SiH₄ discharge, after which the silane flow was switched off. The powders remain trapped in the glow. The displayed spectrum is the ratio of the spectrum of the vacuum vessel filled with only argon and the spectrum of the trapped powders. If one looks closely, one can see that the SiH/SiH₂ band is reduced by a factor of 2, and that a strong Si–O₅ band (1100–1200 cm⁻¹) appears. Apparently, if they are trapped in argon, the powders rapidly oxidize by the impact of the small amount of residual water which is always present in a vacuum system. Figure 11 illustrates the difference of powders trapped in argon and the subsequent treatment by a nitrogen plasma. The spectrum was obtained by subtracting the spectrum after N₂ treatment from the spectrum before N₂ treatment. If peaks point downward, the corresponding absorption reduces, and if they point upwards, the corresponding absorption increases at nitrogen introduction. Disappearing or at least reducing are the Si–O band at 1100–1200 cm⁻¹ and the Si–H bands at 650 and 2000–2100 cm⁻¹. Appearing are a weak Si–N band (1400 cm⁻¹) and a Rayleigh scattering continuum (3000–5000 cm⁻¹). This indicates that hydrogen and oxygen in the particles are replaced by nitrogen. The increase of the Mie–Rayleigh continuum points to either an increasing particle size (improbable) or a changing refractive index (more likely, SiO₂ has a refractive index of about 1.46, Si₃N₄ of 2.0). Also an extra band around 3000 cm⁻¹ appears (probably CH from pump oil contamination).

IV. CONCLUSIONS

In situ infrared absorption spectroscopy during the growth of particulates in discharges of silane/argon mixtures provides a wealth of information. Even at the applied low discharge powers (3 W) the silane dissociation is large (30%). The solid-state absorptions of SiH and SiH₂ only appear after the coagulation phase. Silicon powders trapped in an argon plasma rapidly oxidize. After nitrogen introduction in the discharge, the hydrogen and oxygen atoms which are present in the trapped powders are exchanged for nitrogen.

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