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Time- and space-resolved measurements of Ar(1s5) metastable density in a microplasma using diode laser absorption spectroscopy

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Abstract
Time- and space-resolved measurements of Ar(1s5) metastable (Ar*) density were carried out in a pulsed dc argon microplasma discharge, using diode laser absorption spectroscopy. The temporal behaviour of metastable density after discharge turn off (in the afterglow) depended on their spatial location in the microplasma. In the early afterglow, the Ar* density decayed monotonically with time in the region around the sheath edge, while in the bulk plasma the Ar* density showed a maximum with time. This behaviour was attributed to electron–ion dissociative recombination. Later in the afterglow, the Ar* decay was everywhere monotonic with time, mainly due to three-body collisional quenching by ground state argon atoms. The time evolution of the Ar* density in the afterglow predicted by a kinetic model is in good agreement with the experimental measurements.

1. Introduction

High pressure (hundreds of Torr) microdischarges have attracted much attention due to numerous applications in plasma display panels, excimer radiation sources, chemical and biological sensors, microreactors, materials treatment and modification, sterilization and plasma surgery [1–19].

Microdischarge diagnostics are important to better understand the physics of microdischarges and to optimize device performance. However, the application of conventional diagnostics (e.g. Langmuir probes) is complicated due to the small size (hundreds of micrometres), high operating pressure (hundreds of Torr) and high power density (ten to hundreds of kW cm−3) of microdischarges. In contrast, non-intrusive optical diagnostics are convenient for characterization of microdischarges.

Previously [18], diode laser absorption spectroscopy (DLAS) was used to obtain spatially resolved measurements of Ar(1s5) metastable density (Ar*) in a parallel plate direct current (dc) argon microdischarge. It was found that neutral gas heating resulted in decreasing Ar* density with increasing microdischarge current (i.e. power). This is in contrast to behaviour observed in low pressure (<100 s mTorr) plasmas, where the metastable density increases with input power. However, these measurements could not provide information about the mechanisms of metastable production and loss in high pressure microdischarges.

In this work time- and space-resolved measurements of Ar* density were performed in a pulsed dc argon high pressure microdischarge. Probing the Ar* density as a function of time...
in the afterglow (plasma power off) at different spatial locations in the microdischarge provided information about its formation and destruction kinetics.

**Experimental.** The experimental setup, shown in figure 1, was a modified version of the setup used by Belostotskiy et al [18]. A parallel plate slot-type microdischarge [12, 14], was sustained between two rectangular (5 mm × 0.5 mm) molybdenum electrodes, spaced 300 µm apart. An external cavity diode laser (ECDL) in the Littman configuration (LION—Sacher Lasertechnik, Marburg, Germany) was used as the light source. It was tuned to the \( \lambda = 801.48 \, \text{nm} \) argon line, \( \text{Ar}(2p_8) \leftarrow \text{Ar}(1s_5,3p_2) \) transition. The absorption spectral profile was recorded by scanning the laser frequency across the absorption line. The laser frequency was scanned by changing the voltage on the piezoelectric element that moved one of the cavity mirrors. The laser beam from the ECDL passed through a beam splitter guiding part of the beam to a Fabry–Perot interferometer (1 GHz free spectral range) to perform *in situ* calibration of the laser frequency.

Unlike the original DLAS setup which used continuous wave laser radiation [18], this system employed an external electro-optical modulator (Lithium Niobate Intensity Modulator, EOSPACE Inc., Redmond, WA), so that Ar metastables were probed with short (\( \tau_l = 200 \, \text{ns} \)) laser pulses. Furthermore, an intensified CCD (ICCD, Princeton Instruments model PI-MAX) was used for laser light detection. The ICCD operated in a gated mode, with a gate width of \( \tau_{\text{ICCD}} = 300 \, \text{ns} \), that was synchronized with the laser pulse. This arrangement allowed measurement of the absorption signal over a specified time window during the operation of the pulsed dc plasma.

After passing through the interelectrode space, the laser beam was directed through an aperture to suppress emission from the plasma by reducing the solid angle of light collection. A set of flat and concave mirrors magnified (11 ×) and imaged the laser light, through a red optical filter (2-64 Corning Glass filter), onto the photocathode of the ICCD camera. Pixel binning was applied in the direction parallel to the electrode surfaces to increase sensitivity and improve the signal-to-noise ratio. Each resulting macropixel corresponded to a certain position across the interelectrode gap, and spatially resolved measurements were obtained in a single acquisition.

The argon dc microdischarge was operated in a pulsed power mode. The laser probe was synchronized with the plasma pulse using a Delay Generator (DG 535, Stanford Research Systems Inc., Sunnyvale, CA), which was externally triggered by a personal computer (PC). Time-resolved measurements were performed by sweeping the plasma pulse with respect to the probing laser pulse. The experiment was controlled by LabVIEW (National Instruments, Austin, TX) from the PC via a GPIB interface and a National Instruments data acquisition board (NI PCI-6064E). To obtain the space-dependent Ar(1s5) metastable densities at a given delay time, the Beer–Lambert law was applied to the recorded transmitted laser signals on the ICCD camera with and without the presence of microdischarge (see [18] for more detail).

### 2. Results and discussion

Profiles of \( \text{Ar}^* \) density across the microdischarge as a function of time after the discharge was turned off (i.e. in the afterglow) are shown in figure 2 for a discharge pressure of \( P = 100 \, \text{Torr} \) and current of \( I = 5 \, \text{mA} \). The metastable density near the sheath edge (*sheath edge density*) and that in the bulk plasma (*bulk density*) show different temporal behaviour. The sheath edge density decays in the first \( \sim 2 \, \mu s \), while the bulk density...
increases during this early afterglow time. Later in the afterglow, both the sheath edge and bulk metastable densities decay at a similar rate. The time evolution of the sheath edge (\(\sim 25 \mu m\) from the cathode) and bulk (\(\sim 210 \mu m\) from the cathode) metastable densities are shown in figure 3.

The observed spatiotemporal behaviour of Ar* density can be understood by considering the production and loss mechanisms of argon metastables. At steady state, Ar* production and loss are determined mainly by electron impact excitation and electron impact (stepwise) ionization, respectively [18]:

\[
\text{Ar}^*(1s_0) + e \rightarrow \text{Ar}^*(1s_2, 1s_5) + e, \quad (R1)
\]

\[
\text{Ar}^*(1s_2, 1s_5) + e (\geq 4.2 \text{eV}) \rightarrow \text{Ar}^+ + 2e. \quad (R2)
\]

In the afterglow, the electron temperature decays rapidly (\(\sim 100 \text{ns}\)), resulting in drastic decrease in the rates of high threshold energy reactions such as (R1) and (R2). Under these conditions, other production and loss mechanisms come into play, such as electron–ion dissociative recombination ((R3a), (R3b)), electron impact collisional transfer to the resonant Ar*(1s_1, 1s_4) state followed by radiative decay to the ground state (R4a), (R4b), the reverse of reaction (R4a) (R4c), metastable pooling (R5), collisional quenching by ground state atoms (R6) and (R7) and collisional quenching by electrons (R8):

\[
\text{Ar}^*_2 + e \rightarrow \text{Ar}^*(1s_2, 1s_5) + \text{Ar}^*(1s_0), \quad (R3a)
\]

\[
\text{Ar}^*_2 + e \rightarrow \text{Ar}(2p) + \text{Ar}^*(1s_0) \rightarrow \text{Ar}^*(1s_2, 1s_5) \quad + \text{Ar}^*(1s_0) \quad + hv, \quad (R3b)
\]

\[
\text{Ar}^*(1s_2, 1s_5) + e \rightarrow \text{Ar}^*(1s_1, 1s_4) + e, \quad (R4a)
\]

\[
\text{Ar}^*(1s_2, 1s_5) \rightarrow \text{Ar}^*(1s_0) + hv(106.7 \text{nm}), \quad (R4b)
\]

\[
\text{Ar}^*(1s_1, 1s_4) + e \rightarrow \text{Ar}^*(1s_2, 1s_5) + e, \quad (R4c)
\]

\[
\text{Ar}^*(1s_2, 1s_5) + \text{Ar}^*(1s_2, 1s_5) \rightarrow \text{Ar}^*(1s_0) + \text{Ar}^+ + e, \quad (R5)
\]

\[
\text{Ar}^*(1s_2, 1s_5) + \text{Ar}^*(1s_0) \rightarrow 2\text{Ar}^*(1s_0), \quad (R6)
\]

\[
\text{Ar}^*(1s_2, 1s_5) + 2\text{Ar}^*(1s_0) \rightarrow 3\text{Ar}^*(1s_0), \quad (R7)
\]

\[
\text{Ar}^*(1s_2, 1s_5) + e \rightarrow \text{Ar}^*(1s_0) + e (\geq 11.55 \text{eV}). \quad (R8)
\]

The rate coefficient of dissociative recombination is given by \(k_3 = 9.1 \times 10^{-7}(T_e/T_g)^{0.61} \text{cm}^3 \text{s}^{-1}\) for \(T_g\) in the range 300–8500 K. Here, \(T_g\) (in K) is the gas temperature, assumed equal to the ion temperature [24]. According to Royal and Orel [20] the dissociative recombination reaction (R3) yields mainly an Ar atom in the ground state and an Ar atom in the 2p (\(\sim 25\%\)) or 1s (\(\sim 75\%\)) state. Ar(2p) may in turn radiatively decay to the Ar(1s_0) state. The overall yield of Ar(1s_0) is not known. A rough estimation based on the statistical weights of the 1s manifold suggests that the yield of Ar(1s_0) should be around \(5/(5 + 3 + 3 + 1) \approx 0.4\). Thus, in the absence of experimental data, the overall yield was taken equal to 40%.

Reaction (R4a) has a very low threshold (\(\sim 0.075 \text{eV}\)) compared with electron temperatures prevailing in the active discharge, so its rate coefficient \(k_{4a} = 2 \times 10^{-7} \text{cm}^3 \text{s}^{-1}\) is taken to be independent of electron temperature [21]. However, in the afterglow (when the electron temperature plummets) this threshold is no longer neglected. For this purpose, an Arrhenius type expression was assumed \(k_{4a} = 2 \times 10^{-7} \exp(-0.075/T_e) \text{cm}^3 \text{s}^{-1}\). The rate coefficient of the reverse reaction was calculated using detailed balancing (taking into account the statistical weights of the levels) as \(k_{4c} = \frac{2}{\pi} \frac{1}{k_{4a} L} \exp(-0.075/T_e) \approx 3.3 \times 10^{-7} \text{cm}^3 \text{s}^{-1}\) [26].

The resonant state Ar*(1s_1, 1s_4) decays very fast to the ground state (Einstein’s coefficient \(A_{bg} = 1.19 \times 10^8 \text{s}^{-1}\) [22]). However, at high pressures, resonant radiation is re-absorbed by ground state atoms resulting in so-called radiation trapping. The effective loss rate of Ar*(1s_1, 1s_4) atoms through reaction (R4b) is characterized by an escape factor, estimated using Holstein’s theory [23, 24]. For slab geometry and Lorentzian lineshape, the escape factor is given by

\[
g = \frac{1.15}{\sqrt{\pi k_0 L}}, \quad (1)
\]
where \( k_0 \) is the absorption coefficient at the centre of the spectral profile and \( L \) is the interelectrode gap. Under the present conditions, the spectral line of the resonant radiation (\( \lambda = 106.7 \text{ nm} \)) is not necessarily a pure Lorentzian. In fact, the Doppler width (FWHM), for the typical conditions of \( P \sim 100 \text{ Torr} \) and \( T_e \sim 600 \text{ K} \), is ~5 times larger than the Lorentzian FWHM. Nevertheless, at high opacity (\( k_0L \gg 1 \)), radiation mainly ‘escapes’ through the wings of the spectral distribution. Thus, the Lorentzian component of the spectral lineshape dominates the ‘escape’ of radiation, even though the bulk of the spectral line is still governed by Doppler broadening.

The absorption coefficient can be calculated from [24]

\[
k_0 = \frac{\lambda_0^2}{2\pi} \frac{g_r}{g_0} \frac{A_{tg}}{N} (\frac{Y_0}{N})^{-1},
\]

where \( \lambda_0 = 106.7 \text{ nm} \) is the central wavelength, \( g_r/g_0 = 3 \) is the statistical weight ratio and \( \gamma_0/N \) is the reduced width of the Lorentzian profile (in units of angular frequency), which is governed by resonant broadening. The reduced width is given by [25]

\[
\frac{Y_0}{N} \approx 2\pi \frac{1.45}{8} \lambda_0 c n_e r_e (\text{cm s}^{-1}),
\]

where \( c \) is the speed of light in vacuum, \( f_{tg} = 0.066 \) is the oscillator strength of the transition and \( r_e = e^2/m_e c^2 = 2.818 \times 10^{-13} \text{ cm} \) is a classical electron radius. In the high pressure limit, when the Lorentzian component of the spectral line dominates the ‘escape’ of radiation, the escape factor is independent of pressure. For the Ar resonant transition \( \text{Ar}^{(1s^4 P_1, 1s_4)} \rightarrow \text{Ar}^{(1s^3 S_0)} \) the escape factor, calculated using equations (1)–(3), was found to be \( g = 3.8 \times 10^{-3} \), resulting in a loss frequency of \( \text{Ar}^{(1s^4 P_1, 1s_4)} \) given by \( \nu_{tg} = g A_{tg} = 4.6 \times 10^8 \text{ s}^{-1} \).

The rate coefficients for reactions (R5)–(R7) were taken from [26] \( k_6 = 6.4 \times 10^{-10} \text{ cm}^3 \text{ s}^{-1} \), \( k_8 = 2.3 \times 10^{-15} \text{ cm}^3 \text{ s}^{-1} \) and \( k_7 = 1.4 \times 10^{-32} \text{ cm}^6 \text{ s}^{-1} \), and the rate coefficient for reaction (R8) was taken from [27] \( k_5 = 1 \times 10^{-9} \text{ cm}^3 \text{ s}^{-1} \).

Analysis of the metastable density evolution in the argon afterglow was based on a kinetic model accounting for five species: electrons, atomic ions \( \text{Ar}^+ \), molecular ions \( \text{Ar}_2^+ \) and atoms in metastable \( \text{Ar}(1s_4) \) and resonant \( \text{Ar}(1s_3) \) states. The electron temperature in the afterglow was assumed to be \( T_e \approx 0.1 \text{ eV} \). The initial conditions for bulk electron density \((2 \times 10^{13} \text{ cm}^{-3})\) and gas temperature \((600 \text{ K})\) were taken from [19]. The initial \( (t = 0) \) densities of \( \text{Ar}^+ + \text{Ar}_2^+ \) were calculated from the mass balance of \( \text{Ar}_2^+ \) ions coupled to the electroneutrality constraint:

\[
k_0 N^2 \text{Ar}^+ - (k_3) T_e n_0^3 \text{Ar}_2^+ = \frac{d\text{Ar}_2^+}{dt} = 0,
\]

where \( (k_3) T_e \) is \( k_3 \) evaluated at the steady-state temperature \( T_e^0 \), and \( k_0 = 2.5 \times 10^{-31} \text{ cm}^6 \text{ s}^{-1} \) is the rate coefficient for the production of molecular ions by reaction (R9).

\[
\text{Ar}^+ + 2\text{Ar}(1s_4) \rightarrow \text{Ar}_2^+ + \text{Ar}(1s_5).
\]

Solving the system of equations (4) one obtains the following expressions for the ion densities used as initial conditions for the afterglow calculation:

\[
\text{Ar}^+ = n_0^0 \left(1 - \frac{1}{1 + ((k_3) T_e n_0^0/k_9 N^2)}\right) \quad at \ t = 0,
\]

\[
\text{Ar}_2^+ = \frac{n_0^0}{1 + ((k_3) T_e n_0^0/k_9 N^2)} \quad at \ t = 0.
\]

The bulk electron temperature in the steady-state microdischarge (before the voltage was turned off) was taken as \( T_e^0 \approx 1 \text{ eV} \) [14]. For a \( P = 100 \text{ Torr}, \ (k_3) T_e n_0^0/k_9 N^2 \approx 5 \), so \( \text{Ar}^+ \) is the dominant ion (~82%). The ratio \( \text{Ar}^+ / \text{Ar}_2^+ \) increases in the early afterglow, since the dissociative \( e-\text{Ar}_2^+ \) recombination becomes faster as the electron temperature decreases. However, in the late afterglow, after the electron density has decayed enough \( n_e < (k_9 N^2/k_3) \), \( \text{Ar}_2^+ \) takes over.

The initial density of resonant species \( \text{Ar}(1s_4) \) was found using the corresponding mass balance equation,

\[
(k_{4a}) T_e n_0^0 \text{Ar}(1s_5) - k_4 n_0^0 \text{Ar}(1s_4) - g A_{tg} \text{Ar}(1s_4)
\]

\[
= \frac{d\text{Ar}(1s_4)}{dt} = 0
\]

yielding

\[
\text{Ar}(1s_4) = \text{Ar}(1s_5)^0 \frac{(k_{4a}) T_e n_0^0}{k_4 n_0^0 + g A_{tg}} \quad at \ t = 0.
\]

The species densities as a function of time in the afterglow (calculated with the equations given above) are presented in figure 4. The rate of reactions involved in the \( \text{Ar}(1s_5) \) production and loss are presented in figure 5. The evolution of \( \text{Ar}(1s_5) \) density can be divided into two stages:

(1) early afterglow (0 to a few \( \mu \text{ s} \)): the electron density is still high and electron reactions (including generation of metastables through the electron–ion recombination reaction (R3)) are important.
(2) late afterglow (beyond a few μs): the electron density has decayed to the point that electron reactions are unimportant and metastables are mainly quenched by three-body collisions (R7).

The early afterglow is defined by the following condition:

\[ n_e \gg \frac{k_9 N^2}{k_3}. \] (9)

For 100 Torr argon, this condition implies \( n_e \gg 10^{12} \text{ cm}^{-3} \). Furthermore, for these levels of electron density, decay of electrons through e–Ar\(^+\) recombination (R3) is much faster than electron generation through metastable pooling reaction (R5). Making the pseudo steady-state (PSS) approximation for Ar\(^+\) (which is valid in view of equation (9)) yields

\[ \text{Ar}^+ = \frac{n_e}{1 + (k_3 n_e/k_9 N^2)} \approx \frac{k_9 N^2}{k_3}, \] (10)

where \( k_3 \) is now evaluated at the afterglow \( T_e \). One can then rewrite the electron balance equation as

\[ \frac{dn_e}{dt} = -k_3 n_e \text{Ar}^+ \approx -k_9 N^2 n_e, \] (11)

and solve for the electron density evolution,

\[ n_e = n_e^0 \exp(-k_9 N^2 t). \] (12)

The equation for metastable density can also be simplified. As seen in figure 5, in the early afterglow, one may neglect metastable pooling (R5), two-body collisional quenching by neutrals (R6) and quenching by electrons (R8). Then, \( \frac{d\text{Ar}}{dt} \) becomes

\[ \frac{d\text{Ar}}{dt} = -k_7 N^2 \text{Ar} + 0.4 k_9 N^2 n_e - k_{a\text{Ar}} n_e \text{Ar}(1s_4), \] (13)

where \( k_{a\text{Ar}} \) is evaluated at the afterglow \( T_e \). The factor of 0.4 multiplying \( k_9 \) accounts for the 40% yield of reaction (R9) in producing metastables. In addition, if

\[ n_e \gg \frac{g A_{\text{Ar}}}{k_{a\text{Ar}}} \approx 10^{12} \text{ cm}^{-3}, \] (14)

the radiative state is strongly coupled with the metastable state, and the PSS approximation can also be applied to \( \text{Ar}(1s_5) \). Equation (14) is satisfied in the early afterglow in view of the condition imposed by equation (9). Applying the PSS approximation,

\[ \text{Ar}(1s_5) = \text{Ar}(1s_5) \frac{k_{a\text{Ar}} n_e}{k_{a\text{Ar}} n_e + g A_{\text{Ar}}} \] (15)

and

\[ \frac{d\text{Ar}(1s_5)}{dt} \approx -k_7 N^2 \text{Ar}(1s_5) + 0.4 k_9 N^2 n_e - \frac{k_{a\text{Ar}}}{k_{a\text{Ar}} + g A_{\text{Ar}}} g A_{\text{Ar}} \text{Ar}(1s_5). \] (16)

Using the electron density equation (12), equation (16) can be solved analytically,

\[ \text{Ar}(1s_5) = \text{Ar}(1s_5)^0 \exp(-vt) + 0.4 \cdot k_9 N^2 n_e^0 \exp(-vt) \]

\[ - \exp(-k_9 N^2 t), \] (17)

where \( v = (k_{a\text{Ar}}/k_{a\text{Ar}} + g A_{\text{Ar}} + k_7 N^2) \). This equation provides the metastable density evolution in the early afterglow.

The characteristic time for the transition from the early afterglow to the late afterglow is found by combining equations (9) and (12)

\[ n_e = \frac{k_9 N^2}{k_3} = n_e^0 \exp(-k_9 N^2 t_{\text{EL}}), \] (18)

which gives \( t_{\text{EL}} \approx 4.7 \mu s \) at 100 Torr. In the late afterglow, the electron and ion densities have decayed to the point that reaction (R3) does not contribute to metastable production significantly, resulting in net metastable loss mainly through three-body collisional quenching by neutrals (R7).

Reaction (R3b) yields a photon in the \( \sim 800 \text{ nm} \) spectral range, giving rise to the emission in the near-IR, as observed in figure 6. In figure 6, the initial fast drop of emission intensity reflects the fast decay of electron temperature. After the initial drop, plasma emission decays with a characteristic time that
corresponds to the metastable density decay timescale in the late afterglow (~5 μs). The spatial profile of plasma emission becomes nearly symmetric with respect to the central plane of the microdischarge, resembling a diffusion profile. Since diffusion of the radiating neutrals can be neglected under the conditions of the experiment (their diffusion length before radiating is \( \sqrt{D_{\text{rad}}\tau_{\text{rad}}/\alpha} < 10 \mu\text{m} \)), the emission profile should reflect the spatial distribution of Ar\(^+\) density.

To describe the late afterglow one should consider gas cooling with a concomitant increase in gas density, \( N \). For a 100 Torr afterglow, the characteristic heat conduction time is \( \tau_c = \Lambda^2/4\alpha \approx 7.5\mu\text{s} \), where \( \Lambda \approx 100 \mu\text{m} \) is the microplasma length scale and \( \alpha \) is the thermal diffusivity of the gas. For 100 Torr and 5–10 mA, the spatially averaged gas temperature decreased from ~600 K in the steady-state microdischarge [19] to ~300 K in the late afterglow. This corresponds to an increase in the rate of reaction (R7) from \( \sim 10^{-4} \times 10^3\text{s}^{-1} \) at 600 K to \( \sim 1.4 \times 10^3\text{s}^{-1} \) at 300 K. The experimental loss rate, extracted from the slope of \( \ln(N(1s^5)) \) decay of figure 8, gives \( \nu_{\text{loss}} \approx 2.4 \times 10^2\text{s}^{-1} \), consistent with the estimated value.

Therefore, in the late afterglow, the metastable density can be described by

\[
\text{Ar}(1s^5) = \text{Ar}(1s^5)^* \exp \left(-k_7 \int_{\mu}^{\nu} N(z)^2 dz\right), \tag{19}
\]

where \( \nu^* \) is the beginning of the late afterglow and Ar(1s5)* is the corresponding metastable density. Figure 7 shows a comparison of the metastable density calculated using the full five-species kinetic model with that predicted by the simplified model (equations (17) and (19)). The simplified model provides a reasonable account of the metastable density.

The time evolution of the spatially averaged Ar(1s5) density measured experimentally is compared in figure 8 with the results of the full five-species kinetic model. The model captures the experimentally observed features in a quantitative manner. In the early afterglow, the average metastable density goes through a slight maximum because of the relatively high production rate through electron–ion recombination. Later in the afterglow, when the electron density has decayed sufficiently, the evolution of metastable density is governed by the three-body collisional quenching (R7).

### 3. Conclusions

Time- and space-resolved measurements of Ar(1s5) metastable (Ar*) density were carried out in a pulsed argon dc microdischarge, using DLAS. The time evolution of Ar* density in the afterglow (plasma off) depended on position in the microdischarge. Around the sheath edge, the Ar* density decayed monotonically with time, while in the bulk plasma the Ar* density went through a maximum in the early (0 to few μs) afterglow.

This behaviour was attributed to dissociative recombination of Ar\(_2^+\) with electrons, yielding Ar* as one of the products. In the bulk, during the early afterglow, Ar* production by dissociative recombination exceeded Ar* loss by pooling reactions, and by ground state atom and electron quenching. Thus, the Ar* density exhibited a maximum with time. The opposite situation prevailed in the sheath edge region.

Later in the afterglow, the Ar* decay was everywhere monotonic with time. During this period, the charge density was too low for dissociative recombination to be a significant source of metastables, and the Ar* loss rate was dominated by three-body collisional quenching. Due to gas cooling in the afterglow, the neutral gas density in the microdischarge increased, causing an increase in the metastable loss rate.

A five-species kinetic model of the argon afterglow was developed. The time evolution of the Ar* density predicted by the model was in good agreement with the experimental measurements. Approximate analytical expressions were also provided for the bulk Ar* density as well as for the transition time from the early afterglow to the late afterglow.
Acknowledgments

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References

[21] Ferreira C M, Loureiro J and Ricard A 1985 Populations in the metastable and the resonance levels of argon and stepwise ionization effects in a low-pressure argon positive-column J. Appl. Phys. 57 82–90