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Characterization of energetic and thermalized sputtered tungsten atoms using tuneable diode-laser induced fluorescence in direct current magnetron discharge

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Abstract
In this study a tuneable diode-laser induced fluorescence (TD-LIF) technique (λ₀ = 407.4358 nm) is used to determine the atoms’ velocity distribution function (AVDF) of energetic and thermalized sputtered tungsten (W) atoms in direct current magnetron discharge. The AVDF is characterized by probing the plasma above the centre of the target racetrack along the magnetron cathode axis in an argon–helium (Ar–He) mixture. Quantitative absorption measurements corroborated by deposition on silicon substrates are used to calibrate the TD-LIF relative measurements. Density, flux, temperature, AVDF and the flux velocity distribution function are derived from fitting the TD-LIF signals with four Gaussians (thermalized atoms) and four (energetic atom) functions (Stepanova and Dew 2004 Nucl. Instrum. Methods Phys. Res. B 215 357) taking into account the natural abundance and resonance wavelength shifts of the four main isotopes. Measurements show transport improvement for W atoms and an increase of the ratio of Ar ions to Ar neutrals with the increase of the percentage of He. All measurements are performed at 0.4 Pa and 100 W. The mean velocity of energetic W atoms typically ranges from 1900 to 2200 m s⁻¹. The densities of thermalized and energetic atoms are in the same order of magnitude (∼10⁹ cm⁻³) and their corresponding fluxes are several tens of times higher for energetic atoms (∼10¹⁵ cm⁻² s⁻¹).

Keywords: tungsten sputtering, atoms energy distribution functions, laser-induced fluorescence, laser absorption spectroscopy

(Some figures may appear in colour only in the online journal)

1. Introduction
Sputtered particle sources such as magnetron discharges have been widely used in thin film deposition applications. In order to improve and control film properties, knowledge of the sputtered atoms’ velocity distribution function (AVDF) is important. In particular, for a high-Z material with low thermal diffusion such as tungsten (W), the impinging particles’ energy is determinant in controlling the mobility of condensing atoms during thin layer growth on the substrate [1], e.g. for WC (hard coating) [2] and WO₃ (electrochromic device technology) [3]. The low surface mobility of W atoms leads to porous structures with poor properties while, through an increase of the atoms’ mobility, dense coatings with good physical properties can...
be obtained [4]. Moreover, W is currently considered as a prime candidate for plasma-facing components of the ITER divertor because of its desirable properties [5, 6]. However, physical erosion results in the formation of W atoms and dust particles which are then subsequently carried into the plasma core inducing a severe degradation of the plasma properties required for the fusion reactions [7]. As the penetration depth into the plasma depends on the energy of the sputtered atoms, it is crucial to obtain experimentally the AVDF of neutral tungsten during the sputtering process. The ITER divertor will interact with He (a product of fusion reactions) and Ar (considered as one of the possible coolants). Therefore, the use of an Ar–He gas mixture is of great interest for divertor [8] and deposition studies.

Diagnostics using laser diodes are among the most powerful techniques for studying plasma as they are not intrusive and allow the acquisition of accurate experimental results. Only few studies using such techniques have been performed on sputtered atoms such as aluminum (Al) [9, 10] and titanium (Ti) atoms [11–17]. In this work, new measurements on W atoms are presented. The tuneable diode-laser induced florescence (TD-LIF) technique is used to determine the AVDF of energetic and thermalized sputtered W atoms in direct current (dc) magnetron discharges. The AVDF is measured by probing the plasma above the centre of the target racetrack along the cathode magnetron axis in an Ar–He gas mixture. Tuneable diode-laser absorption spectroscopy (TD-LAS) absolute measurements, corroborated with deposition on silicon substrates, were used to calibrate TD-LIF relative measurements. Density, flux, temperature, AVDF and flux velocity distribution function (FVDF) are determined taking into account the natural abundance and shifted resonance wavelength of the isotopes. This study is focused on the influence of the percentage of helium (%He) in the Ar–He gas mixture, the distance from the target, and on both energetic and thermalized W sputtered populations.

2. Experimental apparatus

2.1. Reactor

The experiments were carried out in a conventional dc magnetron sputtering discharge. The reactor illustrated in figure 1 is a cylindrical stainless steel chamber, 35 cm in height and 30 cm in diameter. We used a standard planar balanced magnetron cathode which operates with a tungsten target of 50 mm diameter, 3 mm thickness and 99.6% purity. Two positions of the magnetron are possible depending on the laser diagnostic used (detailed in the following subsection). The magnetic field flux density is 100 mT at the centre of the target and lower elsewhere. The cathode is moveable in the direction perpendicular to the target surface to allow one to perform space-resolved measurements along this z-axis without modifying the optical arrangement set-up. The chamber is equipped with long cylindrical pipes in order to use optical diagnostic methods and to avoid W deposition on the windows. The chamber also has two other fused quartz windows, facing each other; through one (the one protected by a manual shutter) light is collected with an optical fibre for TD-LIF measurements. The vacuum system consists of two vacuum pumps. A low vacuum of 0.1 Pa is reached using a primary vacuum pump and a high vacuum of $7 \times 10^{-5}$ Pa is achieved using a secondary turbomolecular pump (VAT). The gas is introduced through Ar and He flowmeters injecting up to 20 sccm and 50 sccm, respectively. The measurements are performed with different gas mixtures starting from pure Ar and then increasing the %He in steps of about 10%. The pressure is fixed at 0.4 Pa and the magnetron power at 100 W for all the results presented in this paper.

2.2. TD-LIF, TD-LAS and deposition arrangements

The laser source is a tuneable diode blue-laser system, Toptica Phototonics DL 100 (15 mW), single mode operated with...
resonance transition centred at a λ₀(W) = 407.4358 nm wavelength, which corresponds to the neutral W transition from the low energy (0.37 eV) metastable level 5d⁵(S)6s to radiative excited level 5d⁵(S)6p (figure 2). In this study, as shown in section 3.2, the TD-LAS, TD-LIF and deposition rate measurements suggest a low sensitivity to the electron density (nₑ) variations of the metastable level 5d⁵(S)6s. Consequently, this may be considered as being representative of the W neutral ground state.

The diode laser spectral resolution is 0.005 pm which is much narrower than that of an OPO laser (~2 pm) or a dye Nd:YAG laser (~0.5 pm). The typical scanning frequency is 60 Hz. Keeping the diode temperature constant and systematically controlling the current, the scanning range available, without cavity mode jump, is about 8 pm. The laser beam is separated via a beam splitter into two beams. One probes the plasma volume while the other ones pass through a Fabry–Pérot interferometer to calibrate the wavelength (Δλ = λ激光 − λ₀) [12].

Figure 3 presents a schematic view of the optical arrangements used for TD-LAS (figure 3(a)) and TD-LIF (figure 3(b)) measurements. The absorption and fluorescence wavelengths are the same around λ₀ (a two energy level system, figure 2). To perform TD-LAS measurements, the laser beam is positioned in the direction parallel to the target at z = 9 cm (the distance required for calibration, see section 3.2) corresponding to position ① of the magnetron (figure 1). The light passing through the quartz windows mounted on long pipes is collected by a photodiode. The characteristic absorption curve is determined by measuring the intensity I₀ emitted by the laser diode and the light beam intensity I₁ transmitted after crossing the plasma, and calculating the absorption profile resulting from the Beer–Lambert law (1), where L is the absorption length and k(λ, n) (a product of the attenuation cross section σ(λ) and the atoms’ density n) is the attenuation coefficient at the wavelength λ.

\[ k(\lambda, n) \cdot L = \sigma(\lambda) \cdot n \cdot L = -\ln \left( \frac{I_1}{I_0} \right). \]  

(1)

To perform the TD-LIF measurements, the position of the magnetron is switched from position ① to position ② to avoid changing the position of the laser diode. In this set-up, the target is placed in front of the laser beam. The latter is positioned in the direction perpendicular to the target surface and placed in the centre of the racetrack where W atoms are sputtered. The fluorescence emitted from the probed volume (1 × 1 × 3 mm³) for several distances z from the target is focused by a lens (focal length of 30 cm) into an optical fibre. The TD-LIF signal is carried into a monochromator (Chromex 500 IS/IM) and converted into an electrical signal using a photomultiplier tube (Hamamatsu R928). The tube output is connected to a current amplifier (SR570) and recorded with a digital oscilloscope (Lecroy Wavesurfer no 104Xi).

Deposition on silicon substrates (with preliminary cleaning in acetone and a methanol ultrasound bath) is performed making use of a substrate holder positioned in front of the magnetron cathode (figure 1, position ③). The thickness of the deposition profile was measured using scanning electron microscopy.

3. Results and discussion

3.1. TD-LIF experimental and data analysis procedures

In this study, the particularities of W such as isotopes with their respective resonance wavelength shifts have to be taken into account. In the available scanning range, the high spectral resolution of the laser diode (5 × 10⁻⁵ pm) allows one to clearly distinguish two well-defined populations of W corresponding to thermalized and energetic sputtered atoms and leading to the study of the general behaviour of heavy sputtered species transport. In this section the method used for the analysis of the TD-LIF signal is presented.

At low pressure, i.e. with limited quenching processes, and with conditions for which we checked that the absorption saturation does not occur, the LIF signal intensity can be considered to be directly proportional to the probed particles’ density. The wavelength scale of the laser diode scanning range is converted into velocity scale using \( v = c \cdot \Delta \lambda / \lambda \). The TD-LIF signal then represents the AVDF of the W sputtered atoms. Figure 4 shows a typical shape of the TD-LIF signal recorded at z = 0.8 cm in the axial direction in front of the target racetrack.

The isotope shifted resonance wavelengths and the hyperfine structures of the transitions at 384.8620, 400.8750 and 407.4358 nm were measured in [18]. W has five isotopes whose natural abundances and shifts are given in table 1. The isotopes ¹⁸⁰W, ¹⁸²W, ¹⁸⁴W and ¹⁸⁶W have a nuclear spin of 0 while ¹⁸³W has a nuclear spin of 1/2 giving rise to hyperfine structure. For the considered transition at λ₀ = 407.4358 nm, the contributions of ¹⁸⁰W (with 0.12% natural abundance) and the hyperfine state of ¹⁸³W are negligible. Another transition such the one at 400.8750 nm would make the data analysis more challenging because of line blending issues (singly ionized tungsten lines being coincidentally at nearly the same wavelength as the neutral lines). Also, the transition at 384.8620 nm shows a significant contribution of the hyperfine state of ¹⁸³W masking the ¹⁸⁰W peaks. That is why we chose the transition at λ₀ = 407.4358 nm.
Then, the symmetrical part of the signal corresponding to thermalized W atoms is fitted with a Gaussian distribution function for each isotope with its corresponding shift and natural abundance but the same full width at half maximum (FWHM) as they have the same temperature. The latter was assumed considering the slight difference of mass between W isotopes (≈2%). In the fitting procedure the isotope shifts and the height ratios (accounting for isotope natural abundance) are fixed.

Note that in our conditions, Doppler broadening is typically of 1.5 pm which is much higher than the laser spectral resolution of $5 \times 10^{-3}$ pm. In figure 4 the origin of the velocity scale is arbitrary fixed with the maximum of the thermalized population of the $^{183}$W isotope with the corresponding isotropic Doppler profile in the laser beam direction. The $^{183}$W atoms with negative velocities mean that they travel in the laser beam direction (towards the target) and the opposite (towards the substrate) for positive values.

The asymmetrical part of the signal with positive velocities corresponds to energetic W sputtered atoms (anisotropic distribution function). Considering the physics of sputtering, the energetic distribution can be fitted by a Thompson’s distribution function. The initial energy of the sputtered atoms immediately leaving the target surface obeys Thompson’s formula (equation (2) [19]).

$$J_T(v) \propto \frac{1}{2} M v^2 \left( \frac{1}{2} M v^2 + U \right)^{-3+2m}$$

where $m$ is the binary interaction parameter, $M$ the mass of the sputtered atoms, $v$ the velocity of the sputtered atoms and $U$ is the value of the surface energy barrier. To improve the sputtered atoms’ fitting, the following empirical modification of a previous distribution function proposed by Stephanova et al [20] is required:

$$J_{TM}(v) = J_T(v) \left[ 1 - \left( \frac{\frac{1}{2} M v^2 + U}{\frac{1}{2} M v_{max}^2 + U} \right)^{3+2m} \right].$$
Figure 5. TD-LIF signals with total fits at \( z = 0.8 \text{ cm} \) (a) and \( z = 3 \text{ cm} \) (b) (four Gaussians + four Stephanova et al functions) for several Ar–He mixtures. The velocity scale is arbitrarily centred on the maximum of thermalized population of \(^{183}\text{W}\). The measurements were performed at \( R_0 = 1.3 \text{ cm} \); pressure = 0.4 Pa; magnetron power = 100 W.

Figure 6. TD-LIF signals with total fits at \( z = 5 \text{ cm} \) (a) and \( z = 9 \text{ cm} \) (b) (only four Gaussians) for several Ar–He mixtures. The velocity scale is arbitrarily centred on the maximum of the thermalized population of \(^{183}\text{W}\). The measurements were performed at \( R_0 = 1.3 \text{ cm} \); pressure = 0.4 Pa; magnetron power = 100 W.

This provides a cut off velocity \( v_{\text{max}} \) which represents the limit of the velocity distribution function. In our data analysis, \( v_{\text{max}} \) is fixed at \( 1 \times 10^4 \text{ m s}^{-1} \) to take into account the whole energetic tails, \( U = 1.4 \times 10^{-18} \text{ J} \) (8.8 eV) [21], \( m = 0.2 \) [20]. The AVDF is then fitted with a group of four Gaussians and a group of four Stephanova et al [20] distribution functions. An additional constraint is that, for each isotope, the maximum of the Gaussian distribution function \( (v = 0) \) corresponds to the starting point of the Stephanova et al one since energetic particles only have positive velocities (towards the substrate). Figures 5 and 6 show the raw signals (black lines) with their corresponding total fits (red lines) for several Ar–He mixtures at \( z = 0.8 \text{ cm} \) (figure 5(a)), \( z = 3 \text{ cm} \) (figure 5(b)), \( z = 5 \text{ cm} \) (figure 6(a)) and \( z = 9 \text{ cm} \) (figure 6(b)).

As expected, the intensity of the TD-LIF signal drastically decreases with the increase of \%He due to a lower sputtering as \( M_{\text{He}} \ll M_{\text{Ar}} \). At \( z = 0.8 \) and \( 3 \text{ cm} \), the two populations can be clearly distinguished from \( z = 5 \text{ cm} \) where only a small fraction of energetic sputtered atoms remains (the discrepancy between the black and red lines in the inset). Nevertheless at this distance, the information about energetic atoms cannot be extracted because their density is too low to be fitted with the
Figure 7. (a) A TD-LIF signal recorded at $z = 3$ cm and $R_0 = 1.3$ cm fitted with four Gaussians (thermalized atoms) and four Stephanova et al. [20] functions (energetic atoms). (b) The same TD-LIF signal fitted with eight Gaussians (four for thermalized and four for energetic atoms). The velocity scale is arbitrarily centred on the maximum of the thermalized population of $^{183}$W. Pressure = 0.4 Pa; magnetron power = 100 W; pure Ar.

Figure 8. $[W]_{th}$, measured by means of TD-LIF (relative values) and TD-LAS (absolute values) in addition to the deposition rate as a function of %He. The measurements were performed at $z = 9$ cm and $R_0 = 1.3$ cm; pressure = 0.4 Pa; magnetron power = 100 W.

3.2. Calibration of TD-LIF using TD-LAS

The main advantage of TD-LIF calibrated with TD-LAS is to obtain local quantitative measurements on both energetic and thermalized W sputtered atoms. The use of TD-LAS is, however, restricted to conditions for which the absorption can be assumed to be homogeneous along the laser beam pathway; therefore in the volume where the absorbing particles are thermalized, i.e. far from the target or at high pressure. Since the TD-LAS method integrates the local information along the laser beam pathway, the laser beam is positioned in the direction parallel to the target at $z = 9$ cm, where only thermalized W atoms are measured (as shown in the previous sub-section, figure 6). Figure 8 represents the density of W thermalized atoms ($[W]_{th}$) measured above the centre of the racetrack (at $R_0 = 1.3$ cm) by means of TD-LIF (relative density without calibration) and TD-LAS as well as the deposition rate as a function of %He.

The total TD-LAS signal is fitted with four Gaussians. The experimental maxima of the four absorbance profiles...
(equation (1)), $k_{\exp} (\lambda = \lambda_i)$ and the $\lambda_{\text{FWHM}}$ are determined to calculate the total absolute value of $[W_{\text{abs}}]_{\text{th}}$ (cm$^{-3}$) (4). The corresponding calculated temperature (5) is in the range of 400–500 K (respectively 230–260 m s$^{-1}$). Thus, at such a long distance, the variation of the atoms’ velocity is not significant.

$$\lambda_{\text{FWHM}} = \sum_{i=1}^{4} k_{\exp} (\lambda_i) \cdot \Delta\lambda_{\text{FWHM}} / 8.25 \times 10^{-13} \cdot f \cdot \lambda_i^2$$

$$T_W = \left( \frac{\Delta\lambda_{\text{FWHM}}}{\lambda_i} \cdot \frac{1}{7.16 \times 10^{-7}} \right)^2 \cdot M_W,$$  

where $\lambda_{\text{FWHM}}$ (cm$^{-1}$) is the FWHM of each Gaussian distribution fitting the absorbing profile, $k_{\exp}$ (cm$^{-1}$) is the attenuation coefficient given by the Lambert–Beer law, $f$ is the oscillator strength of the transition ($f = 2.5 \times 10^{-2}$), $\lambda_i$ (cm$^{-1}$) is the resonance wavelength with $i$ the index of the corresponding isotope (see table 1), and $M_W$ (uma) and $T_W$ (K) are the mass and the temperature of W, respectively. From the TD-LIF signal the relative $[W_{\text{rel}}]$ is calculated by determining the area of the AVDF

$$[W_{\text{rel}}] = \int_{-\infty}^{+\infty} \text{AVDF} \, dv.$$  

Measurements by TD-LIF and TD-LAS are in very good agreement $([W_{\text{rel}}]_{\text{th}} = [W_{\text{abs}}]_{\text{th}} = [W]_{\text{th}})$ and are supported by the deposition rate on silicon substrates making the calibration possible. Additional measurements on $n_e$ (not shown here) indicate an increase at $z = 9$ cm by a factor of 3 between %He = 0 and 80 while the metastable level measurements are in good agreement with deposition rate. In fact, $n_e$ seems to not impact the measured metastable densities. This suggests a low sensitivity to $n_e$ of the metastable level $5d^6(6S)6s$. Thus, the latter may be produced close to the target (where the electron density is high) and then diffuses up to the measurement location. Consequently, measurements are considered as being representative of the W neutral ground state. The TD-LIF relative measurements $([W_{\text{rel}}]_{\text{th}})$ and $[W_{\text{abs}}]_{\text{th}}$ are then rescaled for each condition using the consistency at $z = 9$ cm between the TD-LIF signal and $[W_{\text{abs}}]_{\text{th}}$. Thus it becomes possible to obtain a local quantitative estimate of the AVDF and the density of both energetic and thermalized W atoms along the magnetron cathode axis.

The observed maximum of $[W]_{\text{th}}$ and the deposition rate at 20–40%He and the deposition rate at 20–40%He is not a consequence of the variation of the current and the voltage with %He. Indeed, as shown in figure 9, the voltage decreases and the current increases only by 10% from 0 to 90%He and both are quasi-constant up to 40–50%. As detailed in the following sub-section, this behaviour could be the consequence of an improved transport and an increase of the ratio of argon ions to argon neutrals (Ar$^+$/Ar) with the increase of %He. Thus this effect would overcome the lower sputtering of W with higher He content.

### 3.3. Velocities and fluxes of thermalized and energetic W sputtered atoms

In this section we report absolute measurements along the axial magnetron cathode axis of AVDF, FVDF, density ([W]), flux ($\Phi$) as well as the mean velocity ($v_z$) of energetic atoms. The absolute FVDF was obtained by multiplying the absolute AVDF with the corresponding velocity ($\text{FVDF}(v) = v \cdot \text{AVDF}(v)$). Then, the fluxes of thermalized ($\Phi_{\text{th}}$, directed towards the substrate) and energetic ($\Phi_{\text{en}}$) W atoms were calculated by determining the area of the absolute FVDF for each population (7).

$$\Phi = \int_{0}^{+\infty} \text{FVDF} \, dv$$

$$\langle v_z \rangle = \int_{0}^{+\infty} \frac{\text{FVDF} \, dv}{\int_{0}^{+\infty} \text{AVDF} \, dv}.$$  

The purpose is the characterization of the gas mixture effect on the transport of thermalized and energetic W atoms. Figure 10 shows the absolute AVDF ($\Phi$) and FVDF ($\langle v_z \rangle$) obtained by TD-LIF at $z = 0.8$ cm ($R_0 = 1.3$ cm) as a function of Ar–He gas mixtures ranging from 0 to 96% He, by fitting the raw signals of figure 5(a).

In this range of He content, population densities are $4 \times 10^8 \leq [W]_{\text{th}} \leq 7 \times 10^9$ cm$^{-3}$ and $2 \times 10^8 \leq [W]_{\text{en}} \leq 1 \times 10^{10}$ cm$^{-3}$ with corresponding fluxes $\Phi_{\text{th}}$ and $\Phi_{\text{en}}$ of $1 \times 10^{13}$–$3 \times 10^{14}$ atoms cm$^{-2}$ s$^{-1}$ and $4 \times 10^{14}$–$3 \times 10^{15}$ atoms cm$^{-2}$ s$^{-1}$, respectively. The densities of thermalized and energetic atoms are in the same order of magnitude while their fluxes are about ten times higher for $\Phi_{\text{en}}$. Indeed, close to the target and for all mixtures, $\Phi_{\text{en}}$ prevails compared to $\Phi_{\text{th}}$ due to the higher velocity of energetic W atoms. Far away from the target where sputtered atoms have the same energy (only a thermalized population with slow variation of temperature), the behaviour of the density is comparable to the flux one. Thus, it justifies the comparison between the densities and the deposition rate in figure 8. However, close to the target, where thermalized...
and energetic atoms have very different transport properties or behaviour in terms of velocities and directions (anisotropic and isotropic), the important parameter to consider is the flux (and not only the particle density). The flux governs the transport, in which the thermalization process takes part as well as the deposition properties. Also, the flux of energetic atoms $\Phi_{en}$ at the vicinity of the target where the collisional process is limited (here at 0.8 cm which is the closest distance accessible because of the magnetron top protection) leads to the proportion of W sputtering with respect to the gas mixture. Figure 11 presents $\Phi_{en}$ at $z = 0.8$ cm ($R_0 = 1.3$ cm) as a function of the %He. One can observe the non-linear decrease of $\Phi_{en}$ which is lower than the expected behaviour (dashed line). Considering the mass of He to be extremely low and so likewise the sputtering yield ($Y$) of W by He, the ions participating in the sputtering are mainly Ar ions since $Y_{Ar}/Y_{He} \approx 300$ (incident ions’ energy $\approx 250$ eV (figure 9), $Y_{He} = 3 \times 10^{-3}$ atoms/ion, $Y_{Ar} = 9 \times 10^{-1}$ atoms/ion [23]). Consequently, the decrease of $\Phi_{en}$ should follow the decrease of %Ar. In our plasma conditions, it is clear that the populations of ions (He+, Ar+) do not reflect the Ar–He gas mixture variations. Moreover, $\Phi_{en}$ remains significant and so the sputtering of W, even at 80%He ($\Phi_{en} = 1.3 \times 10^{14}$ atoms cm$^{-2}$ s$^{-1}$). This could be explained by a more important Ar$^+/Ar$ ratio in the sputtering plasma than expected. In fact, it is well known that the He excitation and ionization energies are higher than the Ar ones (see table 2).

Additionally, the He ionization cross section ($\sigma_{He^+}$) is about six times lower than the Ar one ($\sigma_{Ar^+}$) in our experiment’s secondary electron energy range (secondary electron energy $\approx 250$ eV, $\sigma_{Ar^+} = 2 \times 10^{-20}$ m$^2$, $\sigma_{He^+} = 0.3 \times 10^{-20}$ m$^2$ [24]). Also, the secondary emission $\gamma_{se}$ of W by He is two and a half times more important than by Ar (incident ions’ energy $\approx 250$ eV, $\gamma_{se-He} = 0.25$ electrons/ion and $\gamma_{se-Ar} = 0.10$ electrons/ion [25]). Thus, as seen in figure 11 in the change of curvature around 80%He, it seems that there is a change in the Ar$^+/Ar$ ratio. Similarly, it is well known that if a small amount of nitrogen or hydrogen is injected in a pure He or Ar plasma, the electron energy is greatly consumed in the low levels of rotation and vibration of molecules implying significant changes of plasma properties (even with weak quantities). In the same manner, the electron energy is preferentially allocated to the lower excitation and ionization of Ar. This means that up to 80%He the Ar$^+/Ar$ ratio increases with respect to %He in the gas mixture since $\Phi_{en,20%Ar}$ is only two times lower than $\Phi_{en,100%Ar}$ instead of the expected five times. It seems this ratio increases by a factor of 2.5. Beyond 80%He the depletion of Ar atoms means that more
electron energy is available to the excitation and ionization of He leading back to the expected drop of the sputtering rate.

Figure 12 reports both the absolute $\Phi_{en}$ and $\Phi_{th}$ on the same plot by using a semi-log scale as a function of the distance for 0, 40 and 80%He. It can be observed that $\Phi_{en}$ strongly decreases while $\Phi_{th}$ grows slightly with the distance from the target. The drastic decrease of $\Phi_{en}$ is not the result of the thermalization process since only a small fraction of $\Phi_{en}$ loss passes to $\Phi_{th}$. Indeed, between $z = 0.8$ cm and $z = 3$ cm, $\Delta \Phi_{en} = \Phi_{en,0.8\text{ cm}} - \Phi_{en,3\text{ cm}} \sim 1 \times 10^{15}$ atoms cm$^{-2}$ s$^{-1}$ while $\Delta \Phi_{th} = \Phi_{th,3\text{ cm}} - \Phi_{th,0.8\text{ cm}} \sim 5 \times 10^{13}$ atoms cm$^{-2}$ s$^{-1}$. To explain the behaviour of $\Phi_{en}$, one has to take into account the geometry of the energetic sputtered cone emission [22] as represented in figure 13. In fact, the further away from the target the probed volume is, the less likely it is to contain energetic W atoms since the section of the sputtered emission cone increases (figure 13: $S_1 < S_2$). Considering the hypothesis of a distribution in cosine of the sputtered atoms, 50–65% of the sputtered vapour is globally emitted in a $\alpha = 30^\circ$–$40^\circ$ emission cone with respect to the normal to the target surface [22]. The section at the basis of the cone is $\pi R_0^2$ with $R_0 = 1.3$ cm being the racetrack radius, the section is equal to $\pi [R_0 + z \cdot \tan (\alpha)]^2$ at distance $z$. In this range of $\alpha$, the ratio of sections between $z = 3$ cm and $z = 0.8$ cm is $S_2,3\text{ cm}/S_1,0.8\text{ cm} \sim 3–4$ which is in agreement with the ratio $\Phi_{en,0.8\text{ cm}}/\Phi_{en,3\text{ cm}} \sim 2–3$. This indicates once again that the measurements on the metastable level $5d^5(6S)6s$ seem to be representative of the W neutral ground state. However, the slight increase of $\Phi_{th}$ is the result of the conversion of a small part (~5%) of the energetic atoms flux into the thermalized one through elastic collisions with the buffer gas. Of course, the thermalization process depends on the Ar–He gas mixture.

The mean value of the ratio of sputtered W atom energies $E_f/E_i$ before ($E_i$) and after ($E_f$) elastic collision with the buffer gas is given by:

$$
\frac{E_f}{E_i} = 1 - \frac{2M_WM_S}{(M_W + M_S)^2} \left( R_0 + z \cdot \tan (\alpha) \right)^2
$$

where $M_W$ is the atomic mass of W atoms and $M_S$ is the atomic mass of the gas species. The ratio $E_f/E_i$ gives 0.71 for a W/Ar collision and 0.98 for a W/He collision. Consequently, W atoms need about 15 times more collisions to thermalize with He. The slower energy transfer in the collisional process than with Ar leads to a slower thermalization and better transport given by the increase of $(v_z)$ with %He (figure 14).

Note that the cathode voltage cannot explain this increase since it decreases. Even close to the target (at $z = 0.8$ cm), the $(v_z)$ increases slightly from 1900 m s$^{-1}$ in pure Ar to 2200 m s$^{-1}$ in 90% He. Note that the measured $(v_z)$ agrees well with the literature [26]. One may notice that this slight increase of about 10% cannot be responsible for the discrepancy between $\Phi_{en}$ and the expected results (dashed line) in figure 11 which can rise up to 200%. The slower energy transfer is also observed on the evolution of $(v_z)$ with the distance to the target (figure 15). For 80% He, $(v_z)$ does not vary much with the distance indicating again limited energy transfers due to collisions during the transport. In accordance with the fact that the energy transfer is more efficient with the increase of Ar content, the measured $(v_z)$ values decrease more rapidly with the distance $z$. In fact, W atoms with an initial velocity of about 1900 m s$^{-1}$ (~3.5 eV) thermalize (note that an atom is said to be thermalized when its velocity falls below three times the thermal speed which here is $\sim 260$ m s$^{-1}$ (~0.04 eV)) in a 500 K background gas [27] after about 10 collisions in Ar gas and after about 100 collisions in He gas.

Using the previous conclusions, it becomes possible to give an interpretation of the observed behaviour of thermalized [W]$_{th}$ atoms far from the target (figure 8, $z \approx 9$ cm). The [W]$_{th}$ (similarly to the flux at this distance) and the deposition rate as functions of %He exhibit a maximum at about 20–40%He instead of a proportional decrease (dashed lines). At first, the sputtering of W was expected to be in proportion to the gas mixture (as $Y_{Ar}/Y_{He} \approx 300$) which turned out to be an underestimate as the Ar$^+/Ar$ ratio increases up to 80%He. Moreover, the transport is enhanced with the increase of He content as the energy transfer during collisions is less efficient. This implies different properties of the sputtered vapour in terms of velocity and direction, i.e. atom flux is less isotropic leading to lower losses to the wall. Thus with the increase of %He, the lower sputtering of W possibly ends up being overcome by the improved transport and the increase of the Ar$^+/Ar$ ratio.

### Table 2. Excitation and ionization energies of W, Ar and He.

<table>
<thead>
<tr>
<th></th>
<th>W</th>
<th>Ar</th>
<th>He</th>
</tr>
</thead>
<tbody>
<tr>
<td>Excitation energy (eV)</td>
<td>3.41</td>
<td>13.1</td>
<td>23.0</td>
</tr>
<tr>
<td>Ionization energy (eV)</td>
<td>7.86</td>
<td>15.8</td>
<td>24.6</td>
</tr>
</tbody>
</table>
4. Conclusion

This work consisted in the development of a method to perform original measurements on thermalized and energetic W atoms using tuneable laser-diode fluorescence (TD-LIF) and absorption (TD-LAS) spectrosopies. Those measurements were carried out in Ar/He DC magnetron plasmas. The atoms’ velocity distribution function (AVDF), flux velocity distribution function (FVDF), density, flux as well as the mean velocity ($v_z$) of the energetic atoms were determined from the measured signals (TD-LAS and TD-LIF) by taking into account the particularities of W in the isotopes’ hyperfine structure and resonance shifts for the resonance transition centred at $\lambda_0$ (W) = 407.4358 nm. The TD-LIF calibration by TD-LAS allowed for the compensation of its main drawback (relative measurements). The TD-LIF data analysis highlighted that even at $z = 3$ cm the distribution of energetic W atoms has to be fitted with Stephanova et al’s function [20]. Even though energetic W atoms undergo collisions with the buffer gas, due to the great difference in mass (with Ar and even more with He), the energetic W atoms conserve a memory of the local information of the ejection (ejection angles and velocity distribution at the target surface $z = 0$).

The study showed that with the increase of the percentage of He in an Ar–He mixture, the expected lower sputtering is overcome by, on one hand, improved transport due to the poorer energy transfer implying different properties of the sputtered vapour in terms of velocity and direction and, on the other hand, an increase of the Ar$^+$/Ar ratio leading to a significant sputtering yield and deposition rate even at 80%He. This was corroborated by measurements on the deposition rate exhibiting a maximum at 20%He (2.8 $\mu$hm$^{-1}$) and a significant value at 80%He (1.6 $\mu$hm$^{-1}$). This study reports absolute measurements, along the magnetron cathode axis, of AVDF, FVDF, density, flux as well as the mean energy of energetic atoms which can be of great interest both for fusion and deposition applications.

References


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