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Deuterium retention in tungsten exposed to low-energy, high-flux clean and carbon-seeded deuterium plasmas

V.Kh. Alimov^{a,*}, J. Roth^b, R.A. Causey^c, D.A. Komarov^a, Ch. Linsmeier^b, A. Wiltner^b, F. Kost^b, S. Lindig^b

^a Institute of Physical Chemistry and Electrochemistry, Russian Academy of Sciences, Leninsky Prospect 31, 119991 Moscow, Russia ^b Max-Planck-Institut für Plasmaphysik, EURATOM Association, Boltzmannstrasse 2, D-85748 Garching, Germany ^c Sandia National Laboratories, P.O. Box 969, Livermore, CA 94550, USA

Abstract

Depth profiles of deuterium trapped in tungsten exposed to a low-energy ($\approx 200 \text{ eV/D}$) and high deuterium ion flux (about 1×10^{21} D/m² s) in clean (We use the term 'clean' in quotation marks having in mind the impossibility to obtain absolutely clean plasma. In our case the conception 'clean' D plasma means the plasma without intentionally introduced carbon impurities.) and carbon-seeded D plasmas at an ion fluence of about 2×10^{24} D/m² and various temperatures have been measured up to a depth of 7 µm using the D(³He,p)⁴He nuclear reaction at a ³He energy varied from 0.69 to 4.0 MeV. The deuterium retention in single-crystalline and polycrystalline W increases with the exposure temperature, reaching its maximum value at about 500 K (for 'clean' plasma) or about 600 K (for carbon-seeded D plasmas, and then decreases as the temperature grows further. It is assumed that tungsten carbide formed on the W surface under exposure to the carbon-seeded D plasmas serves as a barrier layer for diffusion and prevents the outward transport of deuterium, thus increasing the D retention in the bulk of tungsten.

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1. Introduction

Tungsten (W) and carbon fibre composites (CFC) are the candidate materials in the divertor region in ITER where the armors will be subjected to high-flux $(10^{22} \text{ m}^{-2} \text{ s}^{-1} \text{ or higher})$ deuterium–tritium plasmas [1]. Because of high sputtering yield and chemical erosion of the CFC materials, the sputtered carbon particles and hydrocarbon species can migrate into the plasma. They will be ionized by the energetic electrons and, together with hydrogen isotope ions, implanted into tungsten. Carbon and hydrocarbon impurities in the hydrogen plasma will not only govern the process of tungsten erosion and carbon deposition as reported in Refs. [2–8], but will create nearsurface layers of tungsten carbides, W₂C and WC, depending on the conditions as the supply of thermal energy or

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particle bombardment [9-13], and also affect the surface deformation such as blistering [11,14-16].

Despite of the fact that a great deal of publications was devoted to the hydrogen isotope retention in W materials [17–47] the influence of the carbon impurities in a hydrogen (deuterium, tritium) plasma or ion beam on the hydrogen isotope retention in W materials has not been examined yet. There are only indications that the vacuum conditions during D ion irradiation could influence deuterium retention in single crystal tungsten: a higher level of background impurities leads to a higher D retention [32,43].

Studies of carbon re-deposition on polycrystalline W in a controlled carbon-contaminated deuterium plasma environment were performed in the linear magnetized plasma facility PISCES-B [23]. Tungsten substrates were exposed to a high ion flux $(10^{22} \text{ m}^{-2} \text{ s}^{-1})$ and low-energy ($\leq 100 \text{ eV/D}$) deuterium plasma for a total fluence of the order of 10^{26} D/m^2 . Downstream CD₄ puffing was used to produce carbon impurities in the deuterium plasma at

^{*} Corresponding author. Tel.: +7 495 330 2192; fax: +7 495 334 8531. *E-mail address:* vkahome@mail.ru (V.Kh. Alimov).

various substrate temperatures. At an impurity level of about 1% CD₄, no carbon deposition was observed on the plasma-exposed substrates at temperatures below 700 K. While the substrate temperature was increased to 850 K, carbon particles started to form on the tungsten surface. Deuterium retention measurements were done using thermal desorption spectrometry. The dominant factors influencing the hydrogen isotope retention in plasmaexposed tungsten were the substrate temperature, T_{exp} , and whether carbon was deposited on the tungsten surface. For uncontaminated tungsten surfaces, the total isotope retention at $T_{exp} \leq 500$ K was in the range from about 5×10^{20} to about 5×10^{21} D/m². These values were about one order of magnitude higher than those after exposure at higher temperatures (>700 K). The deposited carbon increased the retention of deuterium, which was mainly residing within the carbon film.

The retention of deuterium implanted at room temperature in a pure W single crystal being pre-irradiated with 40 keV C ions was studied by Alimov et al. [48]. D ion implantation was performed at energies of 10 keV, being confined to the carbon-modified layer, and of 100 keV exceeding the carbon-modified layer. The carbon preimplantation influences the deuterium retention only if the range of the D ions is confined to the carbon-modified surface layer. In this case, deuterium diffusion beyond the ion range does not occur and the retained amount of deuterium is smaller than in the pure W crystal. Regarding a D ion energy where the deuterium range exceeds the carbonmodified layer, the deuterium retention is the same for carbon-implanted and pure W samples.

For polycrystalline tungsten pre-irradiated with 1 keV C ions and subsequently implanted with 500 eV D ions, Poon et al. [49] observed that deuterium retention in carbonimplanted W strongly depends on the C ion fluence and the implantation temperature. With a low C ion fluence, tungsten carbide WC was formed at the surface, leading to a decreased D retention at low D ion fluence, compared to the pure W case. With a high carbon ion fluence and low D ion fluence, the presence of a graphitic surface layer increases the D retention to values larger that the pure W case. With increasing D ion fluence, preferential sputtering removes both the graphitic surface layer and the implanted C from the bulk, such that the D retention behaviour of the specimen tends towards that of pure W (with no carbon ion pre-implantation) at high D ion fluencies. The identical results were obtained by Ogorodnikova et al. [36] for polycrystalline W pre-irradiated with 1 keV C ions and afterward implanted with 200 eV D ions.

The purpose of this work is to study deuterium retention in monocrystalline and polycrystalline tungsten exposed to 'clean' and carbon-seeded D plasmas at exposure temperatures in the range from 313 to 853 K. Information about the total amount of deuterium retained in the W materials is obtained from thermal desorption measurements. Moreover, measurements of deuterium depth profiles can give detailed information about the fractions of deuterium retained both in the near-surface layers and in the bulk of the investigated materials. Therefore, the impact of carbon surface impurities on the D retention in the bulk of the plasma-exposed tungsten can be revealed. In our study the $D({}^{3}\text{He}, p){}^{4}\text{He}$ nuclear reaction at different ${}^{3}\text{He}$ energies was used for quantitative depth profiling.

2. Experimental

Two types of W materials were investigated:

(i) Single crystal tungsten specimens produced by double electron-beam zone melting. The manufacturer, the State Institute of Rare Metals (Moscow), quoted the purity as about 99.9 at.% with the main impurities being H (0.02 at.%), C (0.05 at.%) and O (0.05 at.%). The macrocrystallite sizes were in the range of 10-20 mm. The specimens were cut from a W rod by spark cutting and were 0.8–0.9 mm in thickness. The sample surface was parallel to the (100) crystallographic plane.

(ii) A hot-rolled polycrystalline tungsten foil (powdermetallurgy product, 99 at.% W, 0.03 at.% H, 0.1 at.% C, 0.2 at.% O, 0.1 at.% N, 0.5 at.% other impurities – Fe, Al, Si, Ni, Mg) has been investigated. The grain sizes were $1-5 \mu m$. The samples were cut from the 0.5 mm thick foil by spark cutting.

The single crystal and polycrystalline W samples, $6 \times 8 \text{ mm}^2$ in size, were mechanically and electrochemically polished.

The W samples were exposed to 'clean' and carbonseeded deuterium plasmas at various exposure temperatures T_{exp} in the range from 313 to 853 K at Institute of Physical Chemistry and Electrochemistry, Russian Academy of Sciences, Moscow.

The 'clean' plasma was generated in a planar dc magnetron operated with D_2 gas at a pressure of about 1 Pa. The samples placed on the cathode surface and covered with a tantalum mask with an aperture of 5 mm in diameter were bombarded with plasma ions accelerated in the cathode sheath of the magnetron discharge with a discharge voltage of 450 V. The spatial distribution of the plasma potential showed that the cathode-drop voltage was about 0.85 of the discharge voltage. Assumed that D_2^+ ions dominate in the plasma, the mean energy of D ions was estimated to be about 200 eV. The ion flux was determined from ion current measurements and was about $1 \times 10^{21} \text{ D/(m}^2 \text{ s})$.

The carbon-seeded D plasmas were generated by placing plates of pyrolytic graphite within the sputter-dominated area of the cathode surface at a distance of about 8 cm from the W sample. The plasma generated by using one graphite plate with a surface area of about 7.5 cm² is denoted as 'D + C(I)' plasma. The plasma generated by using two graphite plates with a total surface area of about 15 cm² is denoted as 'D + C(II)'. Only polycrystalline W samples were exposed to the carbon-seeded D plasmas.

The W samples were exposed to the 'clean' and carbonseeded D plasmas for 30 min resulting in an ion fluence of about $2 \times 10^{24} \text{ D/m}^2$. During plasma exposure the sample was heated by plasma and the sample temperature mounted to steady-state level for 1-2 min. The temperature was modified by varying heat conductivity between the sample and cathode and was measured by a chromel-alumel thermocouple welded to the front surface of the sample outside the irradiation area.

The deuterium profiles were determined by nuclear reaction analysis (NRA) at Max-Planck-Institut für Plasmaphysik, Garching (IPP). The D concentration within the near-surface layer (at depths up to about 0.5 µm) was measured by means of the $D({}^{3}He, \alpha)H$ reaction at a ${}^{3}He$ energy of 0.69 MeV, and the α particles were energy-analyzed with a small-angle surface barrier detector with a solid angle of 9.17 msr at the laboratory scattering angle of 102°. The α spectrum was transformed into a D depth profile using the program SIMRNA [50]. To determine the D concentration at larger depths, an analyzing beam of ³He ions with energies being varied from 0.69 to 4.0 MeV was used. The protons from the $D({}^{3}He, p){}^{4}He$ nuclear reaction were counted using a wide-angle proton detector with a solid angle of 0.15 sr placed at a scattering angle of 135°. The $D(^{3}He, p)^{4}He$ nuclear reaction has a broad maximum at around 0.63 MeV, which can be used for resonant depth profiling. In order to determine the D concentration profile in deeper layers, the computer program SIMNRA was used for the deconvolution of the proton yields measured at different ³He ion energies. A deuterium depth distribution was assumed taking into account the near-surface depth profile obtained from the α particle spectrum, and the proton yield as a function of incident ³He energy was calculated. The form of the D depth profile was then varied using an iterative technique until the calculated curve matched the measured proton yields [51].

The impurity concentration and chemical state in the outermost atomic layers on the tungsten surface was determined at IPP by X-ray photoelectron spectroscopy (XPS), which probes core-level transitions giving information on the specific element including its chemical environment (Table 1). All photoelectron spectra were measured using a monochromatic Al K α source ($E_{k\alpha} = 1486.6$ eV). For survey spectra, a pass energy of 93.9 eV was used. To achieve the highest energy resolution, the pass energy was reduced yielding an energy resolution of 0.3 eV. The binding energy was calibrated with the Au 4f_{7/2} peak at 84.0 eV and Cu 2p and Ag 3d peaks for linearity. Depth profiles were acquired by sputtering with a scanned 3 keV Ar ion

Table 1

Binding energies for C 1s in carbon films and tungsten carbides [52] (preliminary data were published in [9])

Substance	Binding energy for C 1s (eV)
Graphitic carbon	284.2
Disordered graphitic carbon	285.1
W ₂ C	283.4
WC	282.9

beam. The sputtered area was about $2 \times 2 \text{ mm}^2$, and ion beam current on the target was about 0.4 μ A.

To determine the binding energies and different fractions of binding states in the high-resolution XPS spectra, curvefitting procedures were carried out using the MultiPak software package [53]. For the non-linear least-squares optimization procedure, a Gaussian–Lorentzian line shape was used and a Shirley background was applied [54].

The surface morphology of the plasma-exposed W samples was investigated by scanning electron microscope (SEM).

3. Results

3.1. Surface composition

The carbon, oxygen, and tungsten concentration on the surface of the polycrystalline W exposed to the 'clean' and carbon-seeded D plasmas was determined from the survey XPS spectra (an example is shown in Fig. 1). The elemental concentrations were calculated from the survey spectra using tabulated sensitivity factors. Since no elemental depth profile within the XPS information depth (several nm) is available, the concentrations were calculated assuming a homogeneous distribution over this range. In the survey spectra the signals from Na, N and Cu were detected in addition to the expected peaks of C, O and W. Cu was presumably sputtered from copper parts in the magnetron device and co-deposited during exposure to the D plasmas. After the plasma exposure, the samples were kept in air until the XPS analysis was carried out and Na, N and C impurity accumulation at the top surface were attributed to the air exposure. The high C concentrations (around 50 at.%) measured directly after sample loading (before Ar sputtering) are typically observed on samples which were carried through the air. These adsorbed layers can

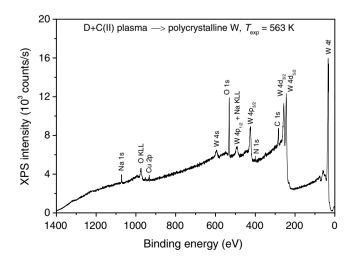


Fig. 1. Survey XPS spectrum of polycrystalline W exposed to a lowenergy and high ion flux 'D + C(II)' plasma at $T_{exp} = 563$ K to an ion fluence of about 2×10^{24} D/m². The XPS measurement was performed after surface sputtering with 3 keV Ar ions at a fluence of 1×10^{19} Ar/m².

be easily removed by sputtering with low Ar fluences. Therefore, only results obtained after surface sputtering with fluences above $1 \times 10^{19} \text{ Ar/m}^2$ were considered. In all cases, the remaining concentrations of the detected impurities (besides carbon and oxygen) are below 1 at.% and are not expected to influence the deuterium inventory.

Although all samples contain carbon in the surface layer, the concentration is lowest for the samples exposed to the 'clean' D plasma. The carbon concentration in the near-surface layers in these cases does not exceed 18 at.% (Table 2). After exposure to the carbon-seeded plasmas, however, the C concentration always exceeds 25 at.%.

The chemical state of the carbon presented at the surface is determined in high-resolution C 1s spectra. Disordered graphitic carbon (peak at binding energy of 285.1 eV) is found on the W samples exposed both to the 'clean' and carbon-seeded D plasmas. Contaminations from the air usually also show peaks at higher binding energies in the C 1s region (around 289 eV) and are not any more present after the lowest applied sputtering fluence of 1×10^{19} Ar/ m², therefore we concluded that contaminations from the transport through the air have largely been removed by a short Ar sputtering procedure.

For the polycrystalline W exposed to the 'clean' D plasma, no indications for carbide formation are visible in the C 1s core-level XPS spectra measured after sputtering with Ar ions at fluences below 7.5×10^{19} Ar/m². Since Ar ion bombardment can also lead to the formation of carbides already at 300 K [55], the absence of carbide intensity in these spectra indicates that no tungsten carbides are present in the surface after the exposure to the 'clean' D plasma. In the spectra measured after the surface sputtering with fluences above 7.5×10^{19} Ar/m², small peaks at the carbide positions are detected (not shown here) and can be explained by the influence of the Ar ion beam.

For the polycrystalline W exposed to the carbon-seeded plasmas, however, the C 1s core-level XPS spectra measured after surface sputtering with lowest Ar ion fluence $(1 \times 10^{19} \text{ Ar/m}^2)$ clearly indicate peaks for tungsten subcarbide W₂C (at binding energy of 283.4 eV). The appearance of this peak can be explained by the implantation of carbon from the carbon-seeded plasmas into the tungsten surface and the formation of carbide by the energetic C atoms. Assuming formation of tungsten carbide with a stoichiometry of W₂C and analyzing data on the impurity

concentration in the near-surface layer of polycrystalline W exposed to the 'pure' and carbon-seeded D plasmas (Table 2), the coverage of W surface with carbide phase can be estimated to be 20–40% for the D + C(I) plasma and 55–60% for the D + C(I) plasma.

Oxygen atoms are present on the W surfaces both after 'clean' and carbon-contaminated D plasma exposures. The oxygen concentration shows no dependence on the type of the plasma.

3.2. Deuterium depth profiles

The retention of deuterium in tungsten develops differently at different depths. The depth at which deuterium is retained in the W materials irradiated with low-energy D ions or exposed to a D plasma is divided into three zones: (i) the near-surface layer (up to a depth of $0.2-0.5 \mu m$, depending on the ion energy), (ii) a sub-surface layer (from about 0.5 to about 3 μm) and (iii) the bulk (>6 μm) [39].

In the single-crystalline and polycrystalline W materials exposed to the 'clean' low-energy ($\approx 200 \text{ eV/D}$) and highflux (about $1 \times 10^{21} \text{ D/m}^2 \text{ s}$) magnetron D plasma at exposure temperatures of $T_{\text{exp}} = 303-313$ K, the D depth profiles are characterized by a sharp near-surface concentration maximum of about 1 at.% and a decreasing concentration tail into the bulk (Fig. 2). At $T_{\text{exp}} = 413-493$ K, the D profiles demonstrate, in addition to the near-surface peak, a relatively high D concentration (0.3–0.4 at.%) in the sub-surface layer. The main difference between single-crystalline and polycrystalline W materials is the constant concentration level in the bulk for polycrystalline material which is absent in single crystals.

The concentration of D_2 molecules (in arbitrary units) within the near-surface layer of the W crystal was determined by measuring the D_2 release rate in the course of surface sputtering with Ar ions (residual gas analysis, RGA, by quadrupole mass spectrometer, for details see [38]). A characteristic of the D_2 release rate after exposure to the 'clean' D plasma at $T_{exp} = 373$ K is that repetitive D_2 gas bursts occur throughout the whole analysis time corresponding to a depth of \approx 450 nm (Fig. 3). The appearance of D_2 molecules is related to the formation of voids filled by gaseous deuterium. Based on the D depth profiles (Fig. 2), one may assume the void formation at depths at least up to 1–2 µm.

Table 2

The impurity concentration in the near-surface layer of polycrystalline W exposed to the 'pure' and carbon-seeded D plasmas

Type of plasma exposure temperature	W (at.%)	O (at.%)	C (total) (at.%)	C (in carbide phase) (at.%)	C (in carbide phase) to W atomic ratio
'clean' D plasma $T_{exp} = 418 \text{ K}$	52.0	35.9	12.1	1.6	0.03
'clean' D plasma $T_{exp} = 493 \text{ K}$	34.5	47.6	17.9	0	0
'D + C(I)' plasma $\parallel T_{exp} = 413 \text{ K}$	41.8	32.6	25.6	8.0	0.19
'D + C(I)' plasma $\parallel T_{exp} = 603 \text{ K}$	46.9	24.4	28.7	5.0	0.11
'D + C(II)' plasma $T_{exp} = 383 \text{ K}$	30.4	25.3	44.3	8.9	0.29
'D + C(II)' plasma $T_{exp} = 563 \text{ K}$	37.2	35.2	27.6	10.2	0.27

The XPS measurements were carried out after surface sputtering with 3 keV Ar ions at a fluence of 1×10^{19} Ar/m².

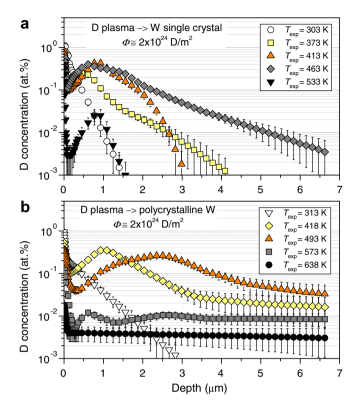


Fig. 2. Depth profiles of deuterium trapped in W single crystal (a) and polycrystalline W (b) exposed to a low-energy and high ion flux 'clean' deuterium plasma at various temperatures.

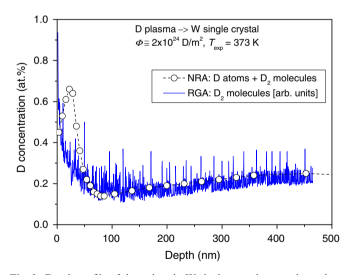


Fig. 3. Depth profile of deuterium in W single crystal exposed to a lowenergy and high ion flux 'clean' deuterium plasma at 373 K as measured with the $D(^{3}He, \alpha)H$ nuclear reaction (NRA) and with the sputter-residual gas analysis technique (RGA). Note that the concentration of D_2 molecules (RGA measurement) is given in arbitrary units.

In the W single crystal exposed to the 'clean' D plasma, the deuterium retention is about $1 \times 10^{20} \text{ D/m}^2$ at $T_{\text{exp}} = 303 \text{ K}$ and, as exposure temperature further increases, rises to its maximum of about $4 \times 10^{20} \text{ D/m}^2$ at $T_{\text{exp}} = 463 \text{ K}$ and then decreases down to about $2 \times 10^{19} \text{ D/m}^2$ at $T_{\text{exp}} = 533 \text{ K}$ (Fig. 4). In polycrystalline W exposed to the 'clean' D plasma, the D retention demonstrates practically the same temperature dependence as that in W single crystal, but the maximum retention in polycrys-

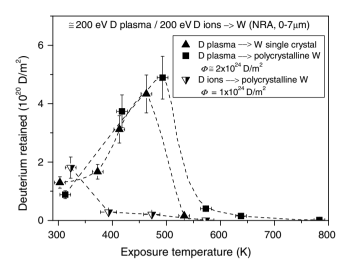


Fig. 4. Deuterium retention in single-crystalline and polycrystalline tungsten-exposed to a low-energy ($\approx 200 \text{ eV/D}$) and high ion flux (about $1 \times 10^{21} \text{ m}^{-2} \text{ s}^{-1}$) 'clean' deuterium plasma to an ion fluence of about $2 \times 10^{24} \text{ D/m}^2$ as a function of the exposure temperature. For comparison, the temperature dependence of the D retention in polycrystalline W irradiated with 200 eV D ions and flux of $4 \times 10^{19} \text{ D/m}^2$ s to a fluence of $1 \times 10^{24} \text{ D/m}^2$ [56] is also shown. The deuterium retention was calculated from deuterium depth profiles measured up to a depth of 7 µm.

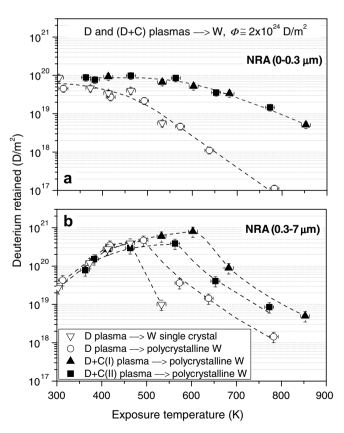


Fig. 5. Deuterium retention at depths of $0-0.3 \,\mu\text{m}$ (a) and at depths of $0.3-7 \,\mu\text{m}$ (b) in single-crystalline and polycrystalline tungsten-exposed to low-energy and high ion flux 'clean' and carbon-seeded deuterium plasmas as a function of the exposure temperature.

talline W is about 5×10^{20} D/m² at $T_{exp} = 493$ K (Fig. 4). Both in single-crystalline and polycrystalline W exposed to the 'clean' D plasma at $T_{exp} = 303-313$ K, the retention in the near-surface (at depths from 0 to 0.3 µm) is higher than that in the sub-surface layer and in the bulk (at depths from 0.3 to 7 µm); whereas at T_{exp} above 350 K, the D retention at depths above 0.3 µm starts to be dominating and reaches about 90% of the total D retention at $T_{exp} = 450-500$ K (Fig. 5).

At first sight the D depth profiles in the polycrystalline W exposed to the carbon-seeded D plasmas ('D + C(I)' and 'D + C(II)' plasmas) (Fig. 6) are similar to those measured after 'clean' D plasma exposure (Fig. 2). However, after exposure to carbon-seeded plasmas, the near-surface concentration maximum is about 3 at.% (compared with about 1 at.% for the 'clean' D plasma exposure), and the exposure temperature at which the D profiles demonstrate a maximum D concentration at depths of 2–3 µm is higher than that for the 'clean' D plasma exposure. It is significant, that in the W samples exposed to the carbon-seeded D plasmas at temperatures above 600 K, the D concentration in the bulk (at depths beyond 6 µm) is about one order of magnitude higher than that after exposure to the 'clean' D plasma (compare Figs. 2 and 6).

After exposure to carbon-seeded D plasmas, the maximum D retention is observed at the exposure temperatures of 603 K ('D + C(I)' plasma) and 563 K ('D + C(II)'

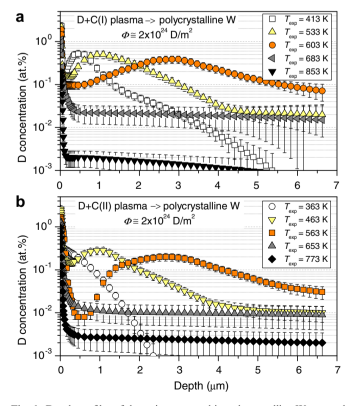


Fig. 6. Depth profiles of deuterium trapped in polycrystalline W exposed to low-energy and high ion flux carbon-seeded deuterium plasmas at various temperatures.

plasma), whereas for 'clean' D plasma exposure the maximum retention is at $T_{\rm exp} = 493$ K (Fig. 5). Generally, maximum D retention after exposure to carbon-seeded plasmas at elevated temperatures is $(5-9) \times 10^{20}$ D/m² and is comparable to the maximum retention after exposure to a 'clean' D plasma.

The temperature dependences of the D content in the near-surface layer (at depths from 0 to 0.3 µm) for exposure to the 'clean' and carbon-seeded D plasmas are different (Fig. 5(a)). For the 'clean' D plasma exposure, the D content is about 5×10^{19} D/m² at $T_{exp} = 313$ K and starts to fall drastically at the exposure temperature of about 500 K. For the carbon-seeded D plasma exposure, the D content in the near-surface layer is 1×10^{20} D/m² at low exposure temperatures and starts to decrease slowly at the exposure temperature of about 500 K. As evident from XPS spectra (see paragraph 3.1), the fraction of the W surface covered with an amorphous deuterated carbon film is higher for the carbon-seeded plasma than that for the 'clean' D plasma.

As for the 'clean' D plasma exposure, the D content at depths from 0.3 to 7 μ m in polycrystalline W exposed to carbon-seeded D plasmas with $T_{exp} = 400-700$ K is higher than that in the near-surface W layer (Fig. 5(b)).

3.3. Formation of blisters

SEM micrographs of the single crystal and polycrystalline W exposed to the 'clean' and carbon-seeded D plasmas to an ion fluence of about 2×10^{24} D/m² have shown that dome-shaped blisters are formed on the metal surfaces at elevated temperatures. The shapes of most blisters are almost spherical with sizes ranging from about 2 to 50 µm, depending on the W structure and conditions of the D plasma exposure.

After exposure to the 'clean' D plasma at $T_{exp} = 303$ K, the blisters are not formed on the surface of the single-crystalline W sample. However, after exposure at temperatures in the range from 373 to 533 K, blisters appear on the surface of the W crystals, and the size (diameter) of blisters is 1–5 µm (Fig. 7(a)). The maximum areal density of blisters is observed at $T_{exp} = 413-463$ K, whereas at $T_{exp} = 533$ K only solitary blisters are present on the surface.

For the polycrystalline W exposed both to the 'clean' D plasma at $T_{exp} = 418-493$ K and to the carbon-seeded D plasma at $T_{exp} = 463-603$ K, cracks are observed on the surface of blister lids (Fig. 7(b)). Note that after exposure to the D plasmas at these temperatures, the D depth profiles demonstrate a concentration minimum between the near-surface and bulk maxima indicating D release due to blister cracking (Figs. 2(b) and 6). The maximum areal density of blisters is observed at $T_{exp} = 493$ K for the 'clean' D plasma and at $T_{exp} = 533-603$ K for the carbon-seeded D plasmas. At these exposure temperatures, the D content at depths beyond 0.3 µm reaches its maximum (Fig. 5(b)).

The dependence of blister mean size (diameter) on the exposure temperature for the polycrystalline W is shown

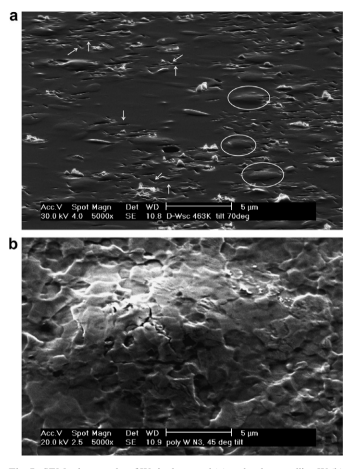


Fig. 7. SEM micrographs of W single crystal (a) and polycrystalline W (b) exposed to a low-energy and high ion flux 'clean' D plasma to an ion fluence of about $2 \times 10^{24} \text{ D/m}^2$ at 463 K (a) and 493 K (b). The surfaces were analyzed at tilt angles of 70 (a) and 45 (b) degrees. Some of the blisters on the surface of W single crystal (a) are marked by ovals, some of the etching pits are indicated by arrows.

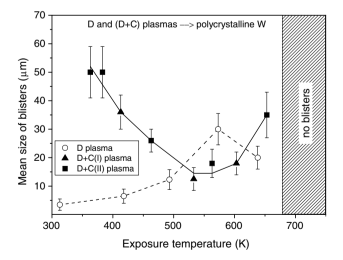


Fig. 8. Mean size (diameter) of blisters formed on the surface of polycrystalline W exposed to low-energy and high ion flux 'clean' and carbon-seeded D plasmas to an ion fluence of about 2×10^{24} D/m², as a function of the exposure temperature.

in Fig. 8. For the 'clean' D plasma exposure, the mean blister size is about 3 μ m at $T_{exp} = 313$ K and grows to the maximum value of about 30 μ m at $T_{exp} = 573$ K. At a

higher exposure temperature, the blister size starts to decrease. For the carbon-seeded D plasma exposure, the blister size demonstrates the opposite temperature dependence. At $T_{exp} = 363-413$ K, the mean blister size is around 50 µm and decreases to about 20 µm at the exposure temperature of 563 K. Further increase of the exposure temperature leads to the growth of the blisters in size. It should be noted that a tungsten carbide layer is formed on the W surface during exposure to the carbon-seeded D plasmas and is not formed during exposure to the 'clean' D plasma.

For exposure to both the 'clean' and carbon-seeded D plasmas at T_{exp} above 680 K, no blister formation is observed on all W surfaces.

4. Discussion

The high deuterium concentration in the sub-surface layer and the formation of blisters allows the conclusion that exposure to a low-energy and high ion flux D plasma modifies the surface to depths up to $2-4 \,\mu\text{m}$, both for W single crystals and polycrystalline W. The sputter-RGA measurements show that a part of deuterium is accumulated in the form of D_2 molecules in the near-surface layer (Fig. 3). It could be supposed that D_2 molecules (or rather D₂-filled voids) are formed in the sub-surface layer as well (up to depths of several micrometers). According to van Veen et al. [57], the D_2 gas inside the voids is expected to be released during annealing at 400-600 K (detrapping energy ≈ 1.0 eV), whereas the D atoms bound on the inner surface of voids (binding energy $\approx 1.7 \text{ eV}$) is expected to be released at 700–900 K. Actually, the D depth profiles (Figs. 2 and 6) show that the D concentration at depths of $1-3 \,\mu m$ starts to decrease significantly at the exposure temperatures above 500 K confirming the assumption that D_2 -filled voids are formed.

The depths of D accumulation (several micrometers) are much larger than the deuterium implantation range (several nanometers). Moreover, deuterium ions with energies less than 800 eV are not able to produce displacement damage in tungsten [22]. A review of hydrogen bubbles in metals by Condon and Schober [58] outlines a possible mechanism of void formation - the mechanism of nearsurface plastic deformation caused by the deuterium super-saturation within the near-surface layer. During D ion irradiation or D plasma exposure, the D concentration in the implantation zone greatly exceeds the solubility limit and stresses the matrix lattice until plastic deformation with formation of voids and vacancy clusters occurs to alleviate these tensions. This deformation is assumed to be responsible for the sudden increase in trapping sites for deuterium and the concurrent accumulation of deuterium, both in the form of D_2 molecules and D atoms. Traces of the plastic deformation within the sub-surface layer of the W single crystal are observed on the surface as etching pits (Lüders bands) oriented along a certain crystallographic plane (Fig. 7(a)). These etching pits are created due to preferential

sputtering of dislocation lines intersecting the surface. According to Savitsky and Burkhanov [59], at plastic deformations of 6-10% the dislocation lines in the bcc lattice are parallel to the direction of the closest packing. But at higher degree of the deformation, a texture is formed.

It has been shown recently [39] that after irradiation with 200 eV D ions at a low-flux of about 4×10^{19} D/m² s, the maximum concentration of deuterium (rather D2 molecules) at depths of 1-3 µm in the polycrystalline W specimen is at least one order of magnitude higher than that in the W single crystals. This suggests that the initial structure of the W material plays a major role in the stress-induced void formation. Possibly, the grain boundaries serve as preferential sites for void nucleation and growth. However, after exposure to the high-flux (about 1×10^{21} D/m² s) D plasma, the maximum D concentration in the sub-surface layer (at depths of $1-3 \mu m$) is the same for both the W single crystal and polycrystalline W (Fig. 2). Apparently, for high-flux D ion implantation formation, the stress-induced defect depends on the W structure to a lesser degree than for low-flux implantation.

Interestingly, in polycrystalline W irradiated with 200 eV D ions at low-flux (about 4×10^{19} D/m² s), the D retention decreases drastically as the irradiation temperature increases from 323 to 393 K [56] (Fig. 4). In our experiments with the ion flux of about 1×10^{21} m⁻² s⁻¹, the decrease in the D retention begins at about 500 K. Evidently, the ion flux affects not only the D retention in W materials, as it was reported in [27,32], but also the temperature where the D retention reaches its maximum. The possible reason of the temperature dependence of the D retention on the ion flux is a balance between the incident flux and temperature dependent D atom diffusion rate out of the implantation zone [27].

In polycrystalline W exposed to the carbon-seeded D plasma, the temperature of maximum retention of deuterium (rather in the form of D_2 molecules) is shifted by 100 K towards higher temperatures as compared to the 'clean' D plasma exposure. Additionally, for the carbonseeded D plasma exposure, the D concentration at intrinsic defects (beyond $6 \,\mu$ m) at the exposure temperatures above 600 K is about one order of magnitude higher than that for the 'clean' D plasma exposure. We expect that the tungsten carbide layer on tungsten surfaces formed by simultaneous deuterium and carbon implantation could prevent the implanted deuterium from leaving the tungsten [15]. As a result, the flux of deuterium atoms diffusing into the bulk increases. In doing so, the temperature of the maximum deuterium retention in the zone of the stress-induced defects is shifted from about 500 to about 600 K (Fig. 5(b)), and the concentration of deuterium trapped at intrinsic defects at higher exposure temperatures significantly increases.

Ueda et al. [15] proposed two mechanisms to control deuterium migration through a tungsten carbide layer. One is to decrease deuterium diffusion through the tungsten carbide layer, the other is to reduce the recombination rate of deuterium atoms on the surface. Regarding the hydrogen diffusion in tungsten carbide, Atsumi and Tanabe [60] have reported data indicating that the diffusivity of hydrogen isotopes in tungsten carbide layer is very different from that in pure W. In these experiments, hydrogen absorption was measured under a hydrogen atmosphere of about 10 kPa. They showed that coexisting carbon with tungsten strongly suppressed hydrogen retention and diffusion in tungsten. Deuterium depth profiles in chemical vapor deposited tungsten carbides W₂C and WC [56] and in polycrystalline W exposed to the low-energy and high ion flux D plasma to the same ion fluence at temperatures below 500 K show that D atoms penetrate in the polycrystalline W much deeper than in tungsten carbides. Thus, these results suggest that tungsten carbide layers formed on the W surface during exposure to the carbon-seeded D plasmas act as a diffusion barrier to deuterium.

Concerning the recombination of hydrogen atoms on the surface, there are no reliable data so far. According to Ogorodnikova et al. [36] and Ueda et al. [15], the presence of carbon atoms on the W surface, however, can reduce the recombination rate of hydrogen atoms.

Both the carbide barrier layer for D diffusion and the reduced recombination rates lead to the build-up of high D concentrations in the nearest-surface layer, to a larger in-depth migration of deuterium, and finally to a larger probability of vacancy cluster and void formation in deeper layers. In doing so, formation of deuterium-defects complexes/voids may occur at higher exposure temperature (at about 600 K) than that in the case of the W surface without the tungsten carbide layer (at about 500 K).

Recent observation of modification of tungsten surfaces exposed to low-energy and high-flux plasmas and ion beam bombardment have shown the formation of blisters [22,23,27,37,42,45–47,61–63]. In our experiments at the exposure temperatures between 350 and 500 K, in spite of the same D retention, the mean size of blisters after the carbon-seeded D plasma exposure is by about one order of magnitude larger than that after the 'clean' D plasma exposure. Mechanical properties of tungsten carbides (the Poisson ratio, the shear modulus) are thought to differ from those for pure W. Possibly, these parameters are the reason for the different sizes of the blisters formed at the same temperature under exposure to the 'clean' and carbon-seeded D plasmas.

There are mainly two different models which describe the blistering process. One is the gas driven model [64– 66], in which bubbles are assumed to grow by coalescence or the loop punching mechanism. As more deuterium is deposited and bubbles grow, cooperative fracture between the bubbles suddenly becomes an easy way of relieving their overpressure, thus initiating cracks, allowing internal gas release, and lifting the surface off into dome-shaped blisters. Another is the lateral stress model [67,68] in which stresses are assumed to be introduced in the implanted layer and the surface will buckle if the induced stress becomes too high. After exposure to the 'clean' D plasma at $T_{\rm exp} = 573$ K to an ion fluence of about 2×10^{24} D/m², the blister size (diameter) reaches 30 µm (Fig. 8), while the concentration of deuterium in the sub-surface layer and bulk is about 10^{-2} at.% (Fig. 2(b)). This fact points to the lateral stress model to describe the blister formation mechanism.

5. Conclusions

Depth profiles of deuterium trapped in single-crystalline and polycrystalline W exposed to a low-energy ($\approx 200 \text{ eV}/\text{D}$) and high deuterium ion flux (about $1 \times 10^{21} \text{ D/m}^2 \text{ s}$) in 'clean' and carbon-seeded D plasmas at an ion fluence of about $2 \times 10^{24} \text{ D/m}^2$ and temperatures from 313 to 853 K have been measured using the D(³He, p)⁴He nuclear reaction, analyzing both the α particles and protons. To determine the D concentration at depths up to 7 μ m, an analyzing beam of ³He ions with energies from 0.69 to 4.0 MeV was used.

In the case of the 'clean' D plasma exposure, the total deuterium retention in single-crystalline and polycrystalline W increases with the exposure temperature, reaching its maximum value of 5×10^{20} D/m² at about 500 K, and then decreases as the temperature grows further. Seeding of carbon into the D plasma does not significantly change the maximum D retention in the polycrystalline W, but shifts the temperature at which the D retention peaks are at maximum, to about 600 K. The high D concentration (about 0.3 at.%) at depths of $1-3 \mu m$, observed after exposure to the 'clean' and carbon-seeded D plasma at temperatures of the maximum D retention, is due to accumulation of D₂ molecules in voids created during the D plasma exposure. At exposure temperatures above 600 K, deuterium is retained in the polycrystalline W mainly as D atoms chemisorbed at the walls of microvoids present initially in the hot-rolled powder-metallurgy W matrix.

It can be assumed that tungsten carbide is formed on the W surface under exposure to the carbon-seeded D plasmas, it serves as a barrier layer for diffusion and prevents the outward transport of deuterium, thus increasing the D retention in the bulk of polycrystalline W matrix.

D plasma exposure with ion energies well below the displacement threshold modifies the W structure to depths of up to about 5 μ m, both in W single crystals and polycrystalline W. Plastic deformation of the W surface caused by deuterium super-saturation within the near-surface layer is suggested as a mechanism for D₂-filled void formation.

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