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Hydrogen plasma induced photoelectron emission from low work function cesium covered metal surfaces

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Experimental results of hydrogen plasma induced photoelectron emission from cesium covered metal surfaces under ion source relevant conditions are reported. The transient photoelectron current during the Cs deposition process is measured from Mo, Al, Cu, Ta, Y, Ni, and stainless steel (SAE 304) surfaces. The photoelectron emission is 2–3.5 times higher at optimal Cs layer thickness in comparison to the clean substrate material. Emission from the thick layer of Cs is found to be 60%–80% lower than the emission from clean substrates. Published by AIP Publishing.

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I. INTRODUCTION

Negative hydrogen and deuterium ion sources are used in many applications, for example, neutral beam injection into fusion devices,1 charge exchange injection into storage rings, e.g., in spallation neutron sources2 and isotope production with cyclotrons.3 H+ ions are produced via the main reaction channels, dissociative electron attachment in the plasma volume and the surface production.4 Surface production of negative hydrogen ions is based on electron tunneling from the conduction band of a metal to the affinity level of a hydrogen atom.5 Cesium is commonly used in negative ion sources to enhance the surface production of negative ions by lowering the work function and thereby increasing the tunneling probability.6–8 However, lowering the work function can also enhance the photoelectron (PE) emission induced by the plasma light emission, the effect of which has not been studied as extensively.

Theoretical calculations based on fundamental conservation laws and reaction cross sections show that at least 10% of plasma heating power is dissipated via photon emission in low temperature hydrogen plasmas.9 The theoretical result is supported by experimental evidence showing that low temperature hydrogen plasmas are strong sources of vacuum ultraviolet (VUV) radiation with up to 15%–30% of the discharge power radiated at wavelengths of 120–250 nm in filament-driven arc discharge,10 up to 8% of the injected microwave power at 80–250 nm in ECR discharge,11 and up to 21% of RF power at 117–280 nm in RF discharge.12

Electrons in the conduction band of a metal follow the Fermi-Dirac distribution which can be used to derive the following relation for the quantum efficiency \( Y \) of the PE emission:13

\[
Y \propto \frac{(h\nu - \phi)^2}{(U_0 - h\nu)^{3/2}}, \quad \text{when } h\nu \geq \phi, \quad (1)
\]

\[
= 0, \quad \text{when } h\nu < \phi,
\]

where \( h\nu \) is the energy of the photon, \( \phi \) is the work function, and \( U_0 \) is the potential step at the surface–vacuum boundary. However, this relationship is accurate only for photon energies near the threshold, when electrons are emitted close to the surface and the quantum efficiency is not affected by the electron transport inside the metal lattice. Typical work functions of common metals are on the order of 4–5 eV. Adsorption of electropositive Cs atoms on a metal surface takes place by transfer of a valence electron to the conduction band of the metal.14 As the Cs coverage on the surface increases, the work function decreases reaching a minimum value at 0.5–0.7 monolayer thickness. Beyond this, repulsive forces between dipole bonds lead to an increasing work function saturating to a value corresponding to bulk Cs at one monolayer thickness. An area density of \( 5.5 \times 10^{13} \) Cs atoms per cm\(^2\) corresponds to one monolayer.15 The minimum work function \( \phi_{\min} \) of a cesiated metal surface can be estimated using a semiempirical expression14

\[
\phi_{\min} = 2.707 - 0.24\phi_0, \quad (2)
\]

where \( \phi_0 \) is the work function of uncesiated metal. Higher \( \phi_0 \) results to lower \( \phi_{\min} \), which is due to stronger electric field on the surface. The functional dependence of the work function on surface coverage \( \theta \) can be estimated using a semiempirical expression14

\[
\phi(\theta) \approx \phi_0 + \frac{6 \Delta\phi_{\min}}{(3 - \theta_{\min})\theta_{\min}} \theta - \frac{3 \Delta\phi_{\min}(\theta_{\min} + 1)\theta}{(3 - \theta_{\min})\theta_{\min}^2} \theta^2 + \frac{2 \Delta\phi_{\min}^2}{(3 - \theta_{\min})\theta_{\min}^3} \theta^3, \quad (3)
\]

where \( \Delta\phi = \phi_{\min} - \phi_0 \) is the work function change and \( \theta_{\min} \) is the relative coverage corresponding to the minimum work function. Work function values suggested by Eq. (3), which have been shown to be in good accuracy with experimental data,14 are plotted in Fig. 1 for different substrates (used in the experiments) with variable \( \phi_0 \)16,17 with a choice of \( \theta_{\min} = 0.5 \).

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Quantified values for the work function of Cs covered metals are usually measured in ultra-high vacuum conditions. However, in ion sources the surface is unavoidably exposed to higher neutral particle fluxes, plasma bombardment, and UV radiation which can affect the formation of the Cs layer. Furthermore, the high reactivity of Cs leads to formation of compounds, e.g., CsOH and Cs₂O, with the impurities present in all low temperature ion source plasmas. The thickness of the Cs layer and the level of surface impurities are determined by the balance between their deposition and adsorption rates. Hence, the surface conditions can have significant temporal variation especially in pulsed high power discharges. Usually a continuous evaporation of Cs into the source is used to keep the surfaces “clean.” There are experimental results from a Penning type ion source showing notable transient lines of Cs compounds in the VUV spectra after plasma ignition. It has been suggested that the involvement of Cs compounds in the Cs dynamics of the ion source is significant or even dominating in ion sources for neutral beam injection. Therefore, work function values derived from the measurements performed at ultra-high vacuum conditions do not apply directly to ion source relevant conditions.

II. EXPERIMENTAL SETUP

The experimental setup is presented in Fig. 2. The measurements were performed with a filament-driven multi-cusp arc discharge plasma generator, which is a commonly used technology in volume production H⁻/C₀ ion sources, e.g., LIISA H⁻/C₀ ion source at JYFL. The vacuum chamber is evacuated down to 10⁻⁸ mbar background pressure with a turbomolecular pump before introducing 99.9999% purity hydrogen into the discharge volume. 70 V/4 A arc discharge voltage/current (280 W discharge power) and a plasma chamber (hydrogen) pressure of 2⁻²⁻ mbar were used in the measurements. It has been shown earlier that the PE current measured from clean metal samples increases linearly with the discharge power. A VUV emission spectrum measured from the hydrogen plasma with a spectrometer consisting of a monochromator (McPherson model 234/302), holographic grating, and photomultiplier tube (ET Enterprises 9406B) is presented in Fig. 3.

The PE emission is measured from a remote sample, which is illuminated by light emitted from the arc discharge.
It is not possible to measure the PE emission inside the plasma chamber, because the measured current would be affected by particle currents from various sources (plasma losses, secondary electron emission, etc.) making it impossible to determine the origin of electrons in the plasma. The PE current was measured with an updated version of a previous apparatus with an integrated Cs oven and a deposition monitor. The light emitted by the hydrogen plasma passes through the extraction aperture (Ø 4 mm) of the plasma chamber and travels through the collimator (Ø 6 mm) and illuminates the sample (4 cm behind the collimator, 90 cm away from the plasma generator). The PE current is measured from the sample with a SRS SR570 preamplifier and National Instruments data acquisition system. The emitted electrons are collected with an anode ring located approximately 10 mm from the target and biased to +150 V with respect to the grounded cathode. The vacuum system includes an optical filter revolver and Cs is supplied through a copper shaft. The oven is heated up to 200 °C and kept at a constant temperature as Cs evaporates onto the metal sample in the PE detector. An Inficon XTM/2 deposition monitor, placed next to the sample, was used to verify the accumulation of Cs, but absolute layer thickness on the sample was not measured.

III. RESULTS AND DISCUSSION

Figure 4 presents the PE current measured from the Mo sample while Cs is deposited on the sample surface. As Cs starts to evaporate, a significant increase in the PE current is observed. The thickness of the Cs coverage grows, the PE current exhibits a double hump structure before starting to decrease. The PE current eventually saturates to a value, lower than the signal from a clean substrate that corresponds to bulk Cs. The PE current measured with the Lyman-alpha filter shows similar behavior as a function of Cs coverage on the substrate compared to the current measured without a filter.

The observations can be qualitatively explained by the decreasing work function of the surface, and by comparison of the photon penetration depth within the Cs layer and the metal substrate to the escape depth of the PEs. It is clearly noticeable that the work function change, illustrated in Fig. 1, alone does not explain the observed behavior of the PE emission. The mean free path for 10 eV VUV photons, which is the average energy for the predominant part of the spectrum, is 10 nm in Mo while the penetration depth in Cs is 13 μm.04 This means that photons interact predominantly with the Mo substrate even at a considerable Cs layer thickness. With a thin layer of Cs, the work function is lowered and thus the PE yield is higher, as long as the emitted electrons are able to propagate through the Cs layer with sufficient energy. Typical escape depth of PEs with energies of a few eV is 1–3 nm.05 If the Cs layer is too thick, the PE current is limited by the short escape depth of the electrons, although the work function is lower for bulk Cs in comparison to Mo. The weaker PE emission from bulk Cs compared

![Figure 3. VUV emission spectrum of the hydrogen plasma generator. The spectrum is not calibrated for spectral response.](image-url)
to clean Mo can be attributed to at least two factors. One is the long mean free path of photons in Cs versus the short escape depth of PEs and the other is the lower density of conduction band electrons in Cs (Fermi energy 1.59 eV) in comparison to Mo (Fermi energy 5.32 eV). It was confirmed with the deposition monitor that the observed transition was indeed caused by a thin layer of Cs, since a change in the PE signal was observed before measuring any detectable signal with the deposition monitor, which has a resolution of 0.01 nm. The measurements are repeatable with less than 12% variation in the maximum gain with optimal Cs layer thickness as measured with Mo and Al samples in four independent measurements per sample. The repeatability includes variable evaporation rates, which presumably affects the concentration of Cs compounds on the sample surface. Uneven Cs distribution on the substrate can create patches with different PE quantum efficiency causing nonhomogeneity of the PE emission. These patches can be caused by local nonuniformity of the substrate due to different crystal faces, roughness, contamination, etc. However, the characteristic dimension of the irradiated surface area (Ø 6 mm) is much larger than the surface roughness (peak-to-peak over 100 nm) and the dimensions of the possible patches. Thus, it can be expected that the patch effect is averaged over multiple measurements and included in the uncertainty probed by repeated measurements.

Figure 5 presents the measured absolute PE currents from different substrates [Mo, Al, Cu, Ta, SS (SAE 304), Y, and Ni] as Cs is deposited on the surface. The horizontal axes have been normalized with the Cs deposition rate for easier comparison, because even a small change in the oven temperature has a significant impact on the deposition rate. Clean substrate materials have variable work functions in the range of 3.1–5.15 eV, while the mean free path for 10 eV VUV photons varies in the range of 8–13 nm. The observed behavior in the PE current transition is similar for all the substrate materials. The PE current is 2–3.5 times higher at optimal Cs layer thickness compared to the clean substrate material. Y has the lowest work function of all the substrates (3.1 eV) and, thus, has the smallest change in the work function due to Cs deposition as suggested by Eq. (2). Y also has the smallest increase in PE current by Cs deposition. Ni,

**FIG. 5.** Measured photoelectron currents from different substrates as a function of Cs coverage by exposing the sample to the whole emission spectrum of the hydrogen plasma (left axes) and by applying the Lyman-alpha filter (right axes).
which has the highest work function (5.15 eV), behaves similar to the other substrate materials, which have work functions between 4 and 5 eV. With all the substrates the PE current saturates to the same value, which is 60%–80% lower than the starting value, supporting the conclusion that the emission is from bulk Cs. The relative difference in the PE current measured with the Lyman-alpha filter in proportion to the PE current measured without a filter has somewhat different values for different metals. The ratio also changes as a function of the Cs coverage. This is believed to be due to the functional dependence of the PE emission quantum efficiency on the photon energy with varying work function.\(^{30,46}\)

Rb was used as an alternative alkali metal to confirm the results obtained with Cs. The work function of Rb (2.16 eV) is almost as low as the work function of Cs (2.14 eV).\(^{16}\) A comparison of PE emission from Mo and Al substrates as a function of Cs and Rb coverage is presented in Fig. 6. In four independent measurements for both alkali metals on Mo substrate, the increase factor in PE current from the clean substrate to maximum current was 3.4 for Cs and 2.6 for Rb on average. Based on two independent measurements with Rb on Al, it is concluded that the PE emission from Rb covered Al is practically the same as for Cs covered Al. The PE current saturates to the same value with both alkali metals, i.e., low temperature hydrogen plasma induced PE emission from bulk Cs and bulk Rb are identical. The comparison of the two alkali metals also suggests that their impurity compounds with different chemical properties play a minor role in determining the PE current.

The role of PEs in plasmas is not well known as they might have a considerable effect on various plasma processes depending on the intensity and the energy distribution of the emitted electrons. In addition to the energy of the absorbed photon, the final energy distribution of the emitted electrons is determined by the plasma sheath structure, i.e., (positive) plasma potential and a possible virtual cathode. A virtual cathode can be formed, if the emission of electrons (and negative ions) from the wall is high enough to prevent the compensation of the space charge by incoming positive ions.\(^{47,48}\) If the virtual cathode exists, it limits the transport of the low energy PEs to the plasma, but does not change the energy of the PEs that reach the plasma as the sheath potential forms between the conducting chamber wall and bulk plasma. The emitted electrons may have an impact on the volumetric rates of the dissociative electron attachment \((e_{\text{cold}} + \text{H}_2(X^1\Sigma_g^+: \nu'') \rightarrow \text{H}_2(^3\Sigma_u^+ \rightarrow \text{H}(1s) + \text{H}^-)\) which has a large cross section for vibrationally excited molecules at low electron energies (around 1 eV), electron detachment \((e + \text{H}^- \rightarrow 2e + \text{H})\) which has a large cross section for energies higher than \(\sim 2\ eV\),\(^{49}\) molecular excitation from the ground state to \(a^3\Sigma_g^+\) and repulsive \(b^3\Sigma_u^+\) triplet states (threshold of about 12 and 8 eV, respectively)\(^{50}\) excitation from the ground state \((\chi^1\Sigma_u^+: \nu' = 0)\) to \(B^1\Sigma_u^+\) and \(C^1\Pi_u^+\) singlet states (threshold approximately 12 eV)\(^{50}\) and molecular ionization (threshold 16 eV)\(^{50}\). Nevertheless, processes with threshold energies higher than the difference between the energy of the absorbed photon and the surface work function are possible only if the emitted electrons with energies ranging from 0 to \(h\nu - \phi\) are accelerated to sufficient energy across the plasma sheath by the plasma potential.

In the filament-driven arc discharge, the total PE current from the plasma chamber walls has been estimated to be in the order of 1 A per kW of discharge power, which corresponds to almost 10% of the arc current of 14 A at 70 V discharge voltage.\(^{29}\) A thin layer of alkali metal can increase the PE emission by a factor of 2–3.5 at optimal layer thickness. The results obtained in this study suggest that the PE emission especially from cesiated surfaces should be included in plasma simulations, in which the plasma surface interaction is taken into account.\(^{51}\) This is especially true for surface production ion sources used for neutral beam injection, in which the Cs covered multiparticle extraction grid covers a large area exposed to VUV photons.\(^{52,53}\) The formation of the virtual cathode requires few hundred \(\text{Am}^{-2}\) of negative ion current density and, due to lower space charge of electrons, it can be assumed that the PE current density needs to be higher by an order of magnitude to impact the sheath structure. Nevertheless, the effect of PEs on the plasma sheath structure may be significant for example in Penning type ion sources,\(^{54}\) where a few kW discharge power is deposited into a small plasma volume (<1 cm\(^3\)), which can lead to high PE emission density from the walls as a significant part of the injected power is dissipated via photon emission.\(^{7,12}\) On the contrary, in neutral beam injection sources the current density can be expected to be smaller, but this does not exclude the PEs effect on the reaction rates.

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