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Efficient Electron-promoted Desorption of Benzene from Water Ice Surfaces†

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Desorption of benzene (C₆H₆) from solid water surfaces [compact amorphous solid water (c-ASW) and crystalline ice (CI)] during irradiation of ultrathin solid films with low energy (250-300 eV) electrons has been investigated. The observed desorption behaviour is complex but typically two desorption components, with particularly large cross-sections, were present in the observed signal. A fast component, with a cross-section up to 10⁻¹⁵ cm², is attributed to desorption of isolated C₆H₆ molecules that are hydrogen-bonded to small clusters of water (H₂O) molecules on the solid water surface. A slower component, with a cross-section of ca. 10⁻¹⁷ cm², is attributed mainly to desorption from larger C₆H₆ islands on the solid water surface. Possible desorption mechanisms are proposed and astrophysical implications are discussed.

1 Introduction

The impact of ionising radiation (keV and above charged particles and electromagnetic radiation with wavelengths shorter than 120 nm) on condensed phase aqueous environments is of both considerable practical and fundamental interest. The key role played in the chemical evolution of the Universe by cosmic ray interactions with icy dust grains and the radiological effects of ionising radiation on biological systems are but two examples. Primarily, ionising radiation interacts with condensed phase water by generating a cascade of low energy secondary electrons, which provide the crucial energy input for promoting physical and chemical change. 1,2 Though much is known of the mechanisms by which these electrons are generated in the condensed phase, 3 there remain gaps in our knowledge specifically surrounding the role low energy electrons play in promoting physico-chemical change especially in the interfacial or selvedge regions of the aqueous environment. 1,2

Therefore, considerable effort has been devoted to investigating the processes underlying the low energy electron-promoted desorption (EPD) of neutral and ionic species from adsorbed layers of H₂O in recent years. A low threshold of ca. 5 eV, was found for the desorption of anionic H⁻ species from amorphous solid water (ASW) via the dissociative electron attachment (DEA) mechanism. 4 Frenkel-type excitons were implicated in desorption of H₂ at electron energies below 11 eV, 5 while the recombination of cationic species formed in the ice with quasi-free or trapped electrons was found to drive H₂ desorption at higher electron energies. Investigations of electron irradiation of thin ASW films deposited on Pr(111) 6,7 showed that H₂ formation is but one of the possible outcomes: O₂ production and desorption of intact H₂O molecules have also been observed. Frenkel-type excitons of 4A1 character lying near the bottom of the ice conduction band are thought to play an important role in the EPD of atomic fragments such as H (2S), O (3P), and O (1D) from ASW films. 8 Formation of H₂, O₂, and H₂O₂ has also been reported at higher energies (5 keV) although these species remain trapped within the ice selvedge, and hence, are not observed during the irradiation but detected during subsequent thermal desorption. 9 More recent work has illustrated the desorption, promoted by low energy electrons, of simple species adsorbed on, 10 and absorbed in, 11 water ice.

Benzene (C₆H₆) is efficiently produced by ion chemistry in the upper atmosphere of Titan, 12,13 and it has been detected in the protoplanetary nebula CRL-618. 14 It is thought to form in the interstellar medium (ISM) under single collision conditions via the gas phase reaction of ethynyl radicals with 1,3-butadiene. 15 C₆H₆ is also proposed to be the starting point for the generation of polycyclic aromatic hydrocarbons (PAHs) 16,17 which carry a significant proportion of the interstellar carbon budget, although the efficiency of such synthetic process is still debated. 18,19 C₆H₆ itself is a convenient system for investigating the interaction of the π-electron density with H₂O ices. For instance, Zwier and co-workers 20–22 have presented IR data using resonance enhanced ionization techniques and resonant ion-dip IR spec-
troscopy\textsuperscript{23} to probe the intermolecular binding between C\textsubscript{6}H\textsubscript{6} and (H\textsubscript{2}O)\textsubscript{n} clusters. Theoretical investigations on the same systems by Slipchenko et al.\textsuperscript{24} have highlighted that C\textsubscript{6}H\textsubscript{6} behaves as both a donor and an acceptor of a hydrogen bond with H\textsubscript{2}O. The interaction between C\textsubscript{6}H\textsubscript{6} molecules and an H\textsubscript{2}O ice film have also been considered. Previous temperature programmed desorption (TPD) studies of C\textsubscript{6}H\textsubscript{6} on ASW show island formation during film growth from the lowest exposures.\textsuperscript{25} This means that C\textsubscript{6}H\textsubscript{6}-C\textsubscript{6}H\textsubscript{6} interaction is favoured over that between C\textsubscript{6}H\textsubscript{6} and the ASW. Nonetheless, when the surface coverage is extremely low, deviation of the desorption kinetics from an ideal zero-order desorption, with evidence of a narrow range of desorption energies, at around 41.0 ± 0.5 kJ mol\textsuperscript{-1}, is compatible with the aromatic ring being \pi-hydrogen bonded in the sub-monolayer regime at the ice interface. Although not yet confirmed by theoretical studies,\textsuperscript{26} reflection-absorption infrared (RAIR) spectra for similar coverages support this interpretation with some evidence of small shifts in peak position of the aromatic C-C stretch from that of bulk C\textsubscript{6}H\textsubscript{6}.\textsuperscript{27, 29}

In a previous letter, we reported our initial studies on the effect of low energy electrons on adlayers of C\textsubscript{6}H\textsubscript{6} on ultrathin ASW films on amorphous silica.\textsuperscript{10} We demonstrated that a highly efficient channel exists for the desorption of isolated C\textsubscript{6}H\textsubscript{6} molecules from the ASW surface and a somewhat less efficient channel attributed to the desorption of C\textsubscript{6}H\textsubscript{6} from C\textsubscript{6}H\textsubscript{6} islands on the ASW surface. The cross-sections for both the processes were largely independent of incident electron energy in the range explored experimentally (100 eV - 350 eV); the observed small increase with primary electron energy is consistent with an increasing number of secondary electrons generated in the ASW film. In this paper, we more fully develop these studies to report on the effect of ASW film thickness and morphology in addition to the C\textsubscript{6}H\textsubscript{6} coverage dependence. We discuss possible mechanisms for the observed C\textsubscript{6}H\textsubscript{6} desorption and suggest that exciton-driven energy transfer from the bulk compact ASW to the selvedge plays an important role.

2 Experimental

The experiments described here were conducted in a stainless steel ultrahigh vacuum (UHV) chamber that has been described in detail elsewhere,\textsuperscript{28, 29} with a base pressure of approximately 2 × 10\textsuperscript{-10} Torr. The substrate is a polished stainless steel disk front-coated with ca. 200 nm of amorphous silica (SiO\textsubscript{2}) that mimics the core of siliceous interstellar dust grains.\textsuperscript{28} The disk was cooled by liquid nitrogen in a reservoir in thermal contact with the sample mount that gives a base temperature of 112 ± 2 K. Under these conditions, both exposure to an effusive molecular beam and background dosing results in growth of a compact, i.e. non-porous, ASW (c-ASW) film.\textsuperscript{30, 31} Deposition at 153 K results in a crystalline ice (CI) growth though a longer exposure (140 L) was necessary due to a non-negligible H\textsubscript{2}O thermal desorption rate. In addition, H\textsubscript{2}O was also deposited at base temperature and subsequently annealed to 138 K resulting in a film that has partially undergone the glass transition and it will be referred to as polycrystalline ice (PCI).\textsuperscript{32-35} C\textsubscript{6}H\textsubscript{6} vapour was subsequently deposited on top of the H\textsubscript{2}O ice, with the sample at base temperature in all cases. Film thickness (\textit{d}) is reported in Langmuir units (\textit{IL} = 1 × 10\textsuperscript{-6} Torr s) and can be converted to

\begin{equation}
\textit{d} = \frac{SP_{T}}{\sqrt{2\pi nk_{B}T}} \frac{1}{\rho_{s}}
\end{equation}

where S is the sticking coefficient assumed to be 1, \textit{k}_{B} is the Boltzmann constant, \textit{T} is the temperature of the dosed molecules, \rho_{s} is the molecular volume density, \textit{n} is the mass, \textit{P} is the pressure recorded on the hot cathode ion gauge corrected for the appropriate molecular ionisation efficiencies,\textsuperscript{36} and \textit{r} is the time of exposure. In the expression, we first define the number of molecules deposited onto the substrate (molecules per unit of surface area) during the dose and then divide this by the density (molecules per unit of volume). De-ionised H\textsubscript{2}O and spectroscopic grade C\textsubscript{6}H\textsubscript{6} (Fluka, ≥ 99.5%) were stored in glass vials and purified by repeated freeze-pump-thaw cycles before use.

Sample irradiation was performed using an electron gun (Kimball Physics, ELG-2) incident at 60° with respect to the substrate normal and over an area of 1 mm\textsuperscript{2}. Desorption of the species was followed using a quadrupole mass spectrometer (VG Microtech PC300D, further modified by ESS) with a homemade line-of-sight tube facing the front of the sample. Several control experiments revealed evidence of substrate charging during the irradiation of the ices with electrons. The incident electron current decreased by up to 30% after 600 s of exposure to the beam giving a typical average flux of (9 ± 2) × 10\textsuperscript{13} electron cm\textsuperscript{-2} s\textsuperscript{-1}. However this had little impact on the quality of the reported measurements as the relevant events occur either in the first 50 s, when the flux can be assumed to be constant at around (1.1 ± 0.2) × 10\textsuperscript{14} electron cm\textsuperscript{-2} s\textsuperscript{-1} (\textit{\phi}_{1}), or at longer times, when the flux has reached a limiting value of (7.5 ± 0.5) × 10\textsuperscript{13} electron cm\textsuperscript{-2} s\textsuperscript{-1} (\textit{\phi}_{2}). No significant difference was noted between flux measurements made on the SiO\textsubscript{2} coated and the uncoated side of the substrate.

Monte Carlo simulations were performed to determine the electron penetration depth in pure C\textsubscript{6}H\textsubscript{6} and ASW films using version 2.48 of the CASINO code.\textsuperscript{37, 38} Calculations for an incident beam at 60°, over an area of 1 mm\textsuperscript{2}, consistent with the experimental conditions (e.g. electron energy), showed that all the binary layered systems investigated have an overall larger thickness than the calculated electron maximum penetration depth. For example, assuming that the densities of the target molecular solids are 2.91 × 10\textsuperscript{22} molecule cm\textsuperscript{-3} and 8.57 × 10\textsuperscript{21} molecule cm\textsuperscript{-3} for H\textsubscript{2}O and C\textsubscript{6}H\textsubscript{6} respectively,\textsuperscript{30, 39} it is found that, for an energy of 250 eV, most of the incident electrons (~ 85%) are stopped within the uppermost 5 nm of the ices. Distributions of electrons within the film as a function of the ASW depth for each experimental energy are reported in the ESI\textsuperscript{10}.

3 Results and Discussion

3.1 Effect of Benzene Coverage

In order to investigate the mechanisms underlying the EPD of C\textsubscript{6}H\textsubscript{6} from an underlying ASW film, we have re-investigated the dependence of the desorption signal on the thickness of the C\textsubscript{6}H\textsubscript{6} overlayer. Previously we reported that desorption signal is dominated by a prompt rise immediately following the onset of electron irradiation.\textsuperscript{10} This is shown in Figure 1, where for a small
exposure of 0.5 L C₆H₆ on a thick ASW film the desorption signal clearly displays a single component that decays rapidly within the first 50 s of irradiation. We will subsequently refer to this as the fast desorption component. For small exposures, the result of 200 L the fast desorption peak is no longer present, consistent with the formation of a thick C₆H₆ film that completely covers the ASW substrate. The significantly reduced intensity of this desorption signal and much smaller desorption cross-section is consistent with the dominant electron-induced process in the C₆H₆ film being dehydrogenation and the formation of a graphitic deposit.⁺ Previously we showed the desorption cross-section for the fast component to be of the order of 10⁻¹⁵ cm² and that this large cross-section can be related to the transfer of excitations within the ASW film to the interface region where C₆H₆ desorption from surface bound C₆H₆-(H₂O)ₙ cluster is induced.

3.2 Effect of Water Ice Film Thickness and Morphology

Since the efficient EPD of C₆H₆ requires the interaction of electrons with the underlying ASW underlayer, it follows that the role of the H₂O has to be carefully investigated. In principle, neglecting the direct interaction with the C₆H₆ molecules, the impinging electrons can promote excitations at the SiO₂/H₂O interface, within the bulk H₂O and/or at the H₂O/C₆H₆ interface. The excitation is subsequently transferred to the H₂O/C₆H₆ interface and ultimately the aromatic ring leading to non-thermal desorption. In order to elucidate some of the aspects of this process, a relatively thin film of C₆H₆ was deposited onto the H₂O ice films of varying thickness and phase (c-ASW, PCI, CI) and then irradiated with 250 eV electrons. Changing the H₂O thickness allows the investigation of the role of excitation within the bulk and at the SiO₂/H₂O interface while changing the phase probes the contribution made by dangling OH bonds at the H₂O/C₆H₆ interface in transferring the excitation itself.

3.2.1 ASW Film Thickness.

Irradiation experiments with 250 eV electrons were performed on layered binary ices comprising 5 L of C₆H₆ on ASW of variable thickness: 70 L (7.7 nm), 100 L (11 nm), 200 L (22 nm) and 500 L (55 nm). Given the results of the CASINO simulations (see the ESI†), ~ 85% of the electrons at this energy are stopped within the top 5 nm of the ASW film. Thus, for ASW films thicker than 5 nm, no change should be expected in the EPD cross-sections as the bulk thickness of the ASW ice is increased and if the desorption process is specific to the H₂O/C₆H₆ interface.

Fig. 1 C₆H₆ EPD traces for (i) 0.5, (ii) 5, (iii) 50 and (iv) 200 L of C₆H₆ adsorbed on a thick c-ASW film. The ices were irradiated with 300 eV electrons with a beam current of 100 nA. Traces are offset for clarity with the dashed lines showing the zero lines for each. The desorption signal for 200 L is clearly significantly reduced compared to the low coverage experiments. This highlights the role played by the c-ASW film in the desorption of C₆H₆ molecules.

Fig. 2 shows the C₆H₆ desorption signal for the four ices. Consistently in all cases the intensity is highest as soon as the irradiation begins and quickly decays within the first 20 s. After that, a slow decrease culminates in a rather constant desorption rate that endures throughout the experiment. The irradiation was stopped before the signal reached zero. Since the beam spot (1 mm²) is significantly smaller than the substrate itself (1 cm²), diffusion from unirradiated areas occurs at 110 K, slowly repopulating with C₆H₆ the probed region of the sample. This explains the non-zero residual obtained in the fit. The observed trend, composed of fast and slow components, is consistent with the results presented in section 3.1. In order to quantitatively estimate the EPD cross-section for C₆H₆ on ASW it is necessary to know both (i) the decay time constant, τᵣ, which is given by the fit, and (ii) the current density, φ. The
A bi-exponential decay function that explicitly contains two desorption components:

\[ I(t) = I_1 e^{-\sigma_1 t} + I_2 e^{-\sigma_2 t} + I_\infty \]  

where \( \sigma_i \) are the desorption cross-sections in \( \text{cm}^2 \) for the fast \((i=1)\) and slow \((i=2)\) component, \( I_i \) is the corresponding amplitude, while \( I_\infty \) is the residual.

The values of the decay time constants, reported in Figure 2, are found to range from 13 to 18 seconds (mean of \( 14.5 \pm 1.7 \) s) and from 213 to 299 seconds (mean of \( 241 \pm 40 \) s) for the fast and slow process respectively. Table 1 lists the electron fluxes employed in the measurements and the cross-sections obtained for this series of experiments. Note that due to the small amount of substrate charging observed, a lower current \((120 \pm 20 \text{ nA})\), and hence current density, was considered for the slow component. When both the statistical and experimental errors are taken into account, both \( \sigma_1 \) and \( \sigma_2 \) are independent of the thickness of the bulk H\(_2\)O films.

![Figure 2](image)

**Fig. 2** C\(_6\)H\(_6\) EPD signal obtained for 5 L of C\(_6\)H\(_6\) on c-ASW of different thicknesses: a) 7.7 nm (70 L), b) 11 nm (100 L), c) 22 nm (200 L), and d) 55 nm (500 L). The experimental data are represented by black closed circles while the solid lines are the results of fitting a two component decay model to the data as described in the text. Irradiation was conducted with 250 eV electrons and initial beam current of 180 nA.

latter is the experimental electron flux in electron \( \text{cm}^{-2} \text{ s}^{-1} \) and allow us to express \( \tau_i \) in terms of desorption cross-section \((\sigma_i)\) as it follows:

\[ \sigma_i = \frac{1}{\tau_i \phi_i} \]  

where the subscript \( i \) refers to the \( i \)th-component in the multi-exponential decay describing the experimental data. In fact, by assuming first-order kinetics for the EPD process\(^{40-42}\), we obtain \( \sigma_i \) and note that all the curves are well fitted by a bi-exponential desorption kinetics are independent of the thickness of the bulk H\(_2\)O films. This suggests that the energy which promotes desorption is deposited in the ASW film within 5 to 10 nm of the ASW/C\(_6\)H\(_6\) ice and is transported to the interface initiating C\(_6\)H\(_6\) desorption.

**3.2.2 Effect of Water Ice Morphology.**

Base temperature conditions in our apparatus (110 K) are compatible with the formation of a c-ASW layer. Annealing this later to 138 K allows the formation of an ice layer that is not yet fully crystalline and, hence, for the sake of clarity will be referred to as polycrystalline ice (PCI). Dosing at 153 K results in the growth of larger crystallites with a much reduced heterogeneity, and hence a CI film. However, at this temperature, the H\(_2\)O accommodation
Table 1 Table listing values of electron fluxes, and cross-sections obtained from the experiments for the EPD process changing the c-ASW thickness.

<table>
<thead>
<tr>
<th>Coverage/L</th>
<th>$\phi_1 \times 10^{14}$ electron cm$^{-2}$ s$^{-1}$</th>
<th>$\sigma_1 \times 10^{-16}$ cm$^2$</th>
<th>$\phi_2 \times 10^{14}$ electron cm$^{-2}$ s$^{-1}$</th>
<th>$\sigma_2 \times 10^{-16}$ cm$^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>70</td>
<td>1.1 ± 0.2</td>
<td>5 ± 2</td>
<td>7.5 ± 0.2</td>
<td>5.6 ± 0.7</td>
</tr>
<tr>
<td>100</td>
<td>1.1 ± 0.2</td>
<td>6 ± 2</td>
<td>7.5 ± 0.7</td>
<td>4.4 ± 0.6</td>
</tr>
<tr>
<td>200</td>
<td>1.1 ± 0.2</td>
<td>6 ± 2</td>
<td>7.5 ± 0.7</td>
<td>6.2 ± 0.6</td>
</tr>
<tr>
<td>500</td>
<td>1.1 ± 0.2</td>
<td>7 ± 2</td>
<td>7.5 ± 0.7</td>
<td>6.2 ± 0.6</td>
</tr>
</tbody>
</table>

as irradiation begins; (2) from ca. 50 s up to 250 s, the C$_6$H$_6$ EPD signal is greater than on c-ASW; and (3) after ca. 350 s, the decay is faster than on the other substrates. The observed trends appear to correlate with the degree of crystallinity within the H$_2$O ice layer. There is a progressive change in the desorption behaviour with increasing H$_2$O film growth temperature. Besides the decrease in the initial desorption intensity, the gradual appearance of a second increase between 50 and 100 s giving rise to a bump is noted: this is strongest for the CI, but is clearly absent for the c-ASW, while it appears to be present for the PCI substrate, where the rapid change in slope at ca. 25 s could be associated with the appearance of the delayed desorption component. Certainly, the remainder of the PCI curve is somewhat similar to that obtained for CI. Any differences observed in the EPD traces in the first instants of irradiation are likely to be linked to the impact that the phase has on the H$_2$O/C$_6$H$_6$ interface. At longer times, ca. 50 s onwards, the effects that the electron bombardment has on the ice structure itself should be taken into account. Previous studies have demonstrated that continuous exposure of CI to ionising sources (H$^+$, VUV, and especially electrons) leads to generation of an amorphous phase. 43–46 This might be reflected by the appearance of the bump in the EPD signal around 70 s as well as the faster decay after 350 s due to gradual electron induced changes to the ice morphology in the interface region. In order to quantitatively evaluate the effect of the H$_2$O ice morphology on the C$_6$H$_6$ desorption, the EPD data were fitted with multi exponential functions. The bi-exponential expression in Equation 3 was used for the curves resulting from deposition of H$_2$O as c-ASW (110 K) and PCI (138 K). However, this function is not sufficient to describe the more complex behaviour shown by the third curve in Figure 4. As in previous work, 10 the phenomenological equation that successfully replicates the appearance of a bump peaking around 70 s is:

$$I(t) = I_1 e^{-\sigma_1 \phi_1 t} + I_2 \left( e^{-\sigma_2 \phi_1 t} - e^{-\sigma_3 \phi_1 t} \right) + I_0$$

(4)

This expression is similar to Equation 3, but introduces a delay in the appearance of the second (slow) component with time decay constant $\tau_d$ from which we can derive a cross-section $\sigma_d$ as per Equation 2. Table 2 lists the values of cross-section found for the three curves distinguishing between the fast component ($\sigma_1$), the slow component ($\sigma_2$), and the delayed component ($\sigma_d$) when the latter clearly occurs. The $\sigma_1$ cross-sections for the experiments at 110 K and 153 K are consistent within the error bars. While $\sigma_1$ is larger, reaching a value of ca. $10^{-15}$ cm$^2$, for the PCI, this increase could be related to the presence of the underlying bump which is not intense enough to be revealed by the experimental curve, in such a way to allow the fit employing Equation 4, but might yet indirectly impact on the observed decay. In conclusion, there is no clear evidence for an increase in the cross-section for the fast component with the phase of the underlying ice. The slow component, $\sigma_2$, is almost halved going from c-ASW to CI; although overall this is a rather modest variation when the error bars are considered. It is important to stress that the C$_6$H$_6$ desorption from CI follows more complex
We now propose a qualitative model-mechanism to explain the findings: The results presented in this and a previous 3.3 non-thermal desorption at the H₂ and 670 nm hinting to a common fundamental mechanism for proposed for the photodesorption (PD) of H₂ beam. An identical interpretation to similar findings could be confirmed by further EPD experiments using thinner to accommodate the energy of the impinging electrons in the first clearly there needs to be a sufficiently thick H₂ molecule is exclusively four in a perfect crystalline film, while amorphous phases display two-, three-, four- and even five-fold coordination. This, combined with a less ordered structure, results in a larger number of H₂O/vacuum (hence H₂O/C₆H₆) interface dangling OH bonds for compact or porous ASW than for CI. It follows that a larger number of C₆H₆ molecules can be found directly bound to dangling bonds when the substrate is amorphous, and hence increasing the intensity of the rapid desorption signal. In conclusion, changing the phase of the H₂O ice underlying the C₆H₆ film strongly suggests that the mechanism is sensitive to the H₂O/C₆H₆ interface, although clearly there needs to be a sufficiently thick H₂O ice (bulk) film to accommodate the energy of the impinging electrons in the first instance before transferring that energy to the adsorbate. This could be confirmed by further EPD experiments using thinner ASW films which do not completely attenuate the electron beam. An identical interpretation to similar findings has been proposed for the photodesorption (PD) of H₂O between 275 and 670 nm hinting to a common fundamental mechanism for non-thermal desorption at the H₂O surface.

3.3 Key Findings and Proposed Mechanism
The results presented in this and a previous work on EPD of C₆H₆ from H₂O ices are summarised below, highlighting the key findings:

- EPD is dependent on the C₆H₆ coverage on the H₂O surface. At low exposures only one efficient process is observed (σ₁ ≈ 10⁻¹⁵ cm²), while this is quenched at large C₆H₆ exposures when the H₂O film is completely covered by a C₆H₆ multilayer. At intermediate coverages, desorption can be described using a multi-exponential decay that contains at least a fast and slow process.

- H₂O ice is a necessary substrate to observe the efficient desorption of the aromatic molecules.

- The process is independent of the thickness of the underlying H₂O ice in the range explored experimentally for thick films.

- The nature of the C₆H₆/H₂O interface is significant as it impacts on the intensity of the desorption trace and delays the slow desorption.

We now propose a qualitative model-mechanism to explain the large cross-section for non-thermal desorption of C₆H₆ from H₂O ices compatible with the key findings listed above. The starting point lies in the vast literature of previous studies on electron irradiation of icy films and on optical-absorption and electron-energy-loss spectroscopy (EELS) of solid and liquid water. These will likely induce further multiple excitations, lying in the 8.7 eV - 21 eV range, and further ionisation events. In the work of Kimmel and Petrik, 5,6,58,59 molecular hydrogen (H₂) formation is observed following the irradiation of ASW with 100 eV electrons and was also observed during our measurements (results not shown). The formation and observation of H₂ is explained in terms of electronic excitation of the H₂O (exciton formation) and electronic energy transport to the H₂O/vacuum interface where H₂ formation occurs. Diffusion and long-range transport of H atoms through the ASW or dissociative electron attachment (DEA) reactions should not contribute significantly to the H₂ yield since the mobility of hydronium ions is very limited at 100 K and because DEA would be efficient only at the H₂O/vacuum interface in a smaller energy range (~ 10 eV) than the 100 eV employed in the experiment. 61 Based on this mechanism, the energy deposited in the ice in the form of long-lived excitons during irradiation, would be transported to the H₂O/C₆H₆ interface and promote non-thermal desorption of the aromatic molecules (see upper panel in Figure 1) competitively with H₂ formation. The energy is passed via π-hydrogen bond to the adsorbed aromatic molecule promoting desorption (see the cartoon in Figure 5). It follows that the observed EPD is peculiar for adsorbates interacting with the H₂O surface via such a weak, directed interaction. Although further experiments are clearly necessary to confirm the role of
the hydrogen bond at the interface and the hydrogen bonding network in the H$_2$O film. In contrast, as displayed in the lower panel in Figure 1, increasing the C$_6$H$_6$ coverages to form thick films (e.g. 200 L) reduces significantly the number of electrons reaching the underlaying H$_2$O. EPD promoted by interaction with H$_2$O is therefore quenched, while desorption of the adsorbates is still observed as it is activated by less efficient processes. The value of $\sigma_2$, $(6 \pm 4) \times 10^{-17}$ cm$^2$, is consistent with previous studies of irradiation of both chemisorbed and multilayer C$_6$H$_6$ on W(110) and of amorphous silica, and hence, can be attributed to desorption from C$_6$H$_6$ multilayer on the solid H$_2$O surface. Although at intermediate exposures C$_6$H$_6$ forms islands, some molecules will still be H-bonded to the icy substrate leading to observation of both the fast and the slow desorption components. The latter does not simply contain the contribution to the signal from C$_6$H$_6$ desorbing from pure C$_6$H$_6$, but might be enhanced as a result of the islands disrupting, morphological changes within the ice, and non-thermal diffusion of islanded C$_6$H$_6$ towards the dangling OHs at the interface intensifying the desorption yield. Increasing the thickness of the H$_2$O film has no effect on the kinetics of the desorption, since if excitons were to form within the bulk and then migrate isotropically in all directions, those travelling toward the H$_2$O/C$_6$H$_6$ would not be affected by a larger spacing below 7 - 8 nm of depth. However, in Figure 2 the curve for 500 L of c-ASW, the initial rise is less intense compared to thinner H$_2$O films (70 L, 100 L, and 200 L). This is interpreted as due to the available surface area at the interface; a thinner H$_2$O film would reflect peaks and troughs of the underlying substrate, while these would be smoothed out in thicker films. A smaller degree of surface roughness means a reduced surface area at the interface, and hence fewer sites where C$_6$H$_6$ molecules could be hydrogen-bonded and accept the energy deposited in the bulk during the irradiation. The same interpretation can be extended to the bump appearing in Figure 4 and in previous work. The latter case this feature was observed also for C$_6$H$_6$ on c-ASW when high electron energies were used (e.g. 350 eV). Irradiation dynamically changes (roughens) the H$_2$O surface and generates additional dangling bonds that coordinate to the aromatic rings. This leads to a further increase of the desorption rate in the form of the aforementioned bump. This process can be particularly efficient, with a cross-section of $\sim 10^{-16}$ cm$^2$, and depends on the phase of the H$_2$O ice and on the energy of the impinging electrons. The exact mechanism is not yet clear, however the data are consistent with the idea that the entity in which the icy surface roughens correlates with a more or less efficient rearrangement of the H$_2$O/vacuum interface. This is due to amorphisation of the H$_2$O/vacuum interface accompanied by disruption of the C$_6$H$_6$ islands when the electron energy is highest and the morphology changes most. Hence, it follows that diffusion of the aromatic molecules from the edges of the islands themselves to the newly generated dangling OHs promotes the second increase (bump) in the EPD traces.

4 Astrophyiscal Implications and Conclusions

The experiments discussed in this work simulate the secondary electron flux forming subsequent to ionising radiation interaction with interstellar ices. There are several sources that could lead to ionisation and, hence, to secondary electrons: X-rays, $\gamma$-rays, cosmic rays and to some extent also energetic VUV photons. Particularly noteworthy is the case of cosmic ray protons where the differential flux exhibits a relatively flat distribution between 1 - 100 MeV, with a maximum at 50 MeV. The integrated flux is $\sim 1$ cm$^{-2}$ s$^{-1}$ in this energy range. The distribution of secondary electrons produced by primary ions ($^1$H, $^6$He) in liquid water is only slightly dependent on the energy of the ions themselves between 100 keV and 100 MeV. These energy distribution curves, with a mean electron energy between 38.6 eV and 65.1 eV, overlap with the electronic excitations of solid H$_2$O at 8.7 eV, 10.4 eV and 14.5 eV, and 21 eV. Although these distributions were obtained from liquid water, they are not expected to be significantly different in ASW, which has often been used in the past as a model of liquid phase. Therefore, it can be concluded that cosmic rays impacting the icy ISM grain mantles, mostly made of H$_2$O, produce an ionisation track within the mantle which is rich in secondary electrons having energies resonant with valence electronic excitations of solid H$_2$O. The primary electron beam used in the experiments reported herein would likely be inelastically scattered, ionising molecules and forming a similar flux of secondary electrons to that associated with cosmic rays; thought somewhat more localised in the selvedge that would be the case for cosmic ray irradiation. Therefore, the reported results are appropriate for considerations in astrophysical context. It should be stressed that the relative abundance of C$_6$H$_6$ in dense clouds is not sufficient for the formation of thick layers, however, solid H$_2$O is certainly the main component of the ice mantles and presents peculiarities (long-lived exciton formation) that could favour EPD of any small molecule, including carbon monoxide (CO) which is known to be weakly bound to the ice surface via a dangling OH bond. The key point is that the efficiency of the desorption process observed in the present work results from the deposition of energy in the underlying H$_2$O film rather than the nature of the adsorbate film. Although dedicated measurements are clearly required, it is therefore reasonable to consider that a similar efficient desorption may occur for other adsorbates bound on a H$_2$O substrate. Therefore, the surface abundance of a molecule such as CO on the ASW would be the result of several processes: adsorption from the gas phase; thermal and non-thermal desorption; and reactions on the surface. It is noteworthy that hydrogenation of CO is the first and crucial step to the reactive accretion of complex organic molecule (COM) on the ISM grain surface. Clearly an additional efficient desorption step might potentially delay the mantle enrichment in CO, and hence methanol (CH$_3$OH) formation, and thence other organic molecules.

In order to ascertain the impact of EPD from ASW on an astrophysically relevant time-scale, it is desirable to quantitatively compare the effectiveness of EPD and photodesorption (PD) of
just CO on ASW at sub-monolayer, and monolayer level. Fayolle et al.\textsuperscript{57} have reported an average value of $1.8 \times 10^{-2}$ molecule photon$^{-1}$ for PD yields of CO from CO multilayers in the investigated range (120 - 177 nm). This process was shown to be independent of the thickness of the CO ice confirming that only the top few layers are directly involved. No direct measurements for the PD of CO from ASW are available to date, thus we will assume that, in contrast with the EPD process, the H$_2$O substrate has no effect on the PD yield of the adsorbate (CO). In order to compare these findings with the EPD results, the CO PD rate, $k_{PD}^{CO}$, can be estimated under astrophysical conditions using the following equation:

$$ k_{PD}^{CO} = \Phi \sigma_{abs} F_{UV} $$

(5)

where $\Phi$ is the yield of the process ($1.8 \times 10^{-2}$); $\sigma_{abs}$ is the photon absorption cross-section for CO in the VUV range between 120 nm and 160 nm ($4.7 \times 10^{-18}$ cm$^2$);\textsuperscript{68} and $F_{UV}$ is the estimated VUV flux in dense clouds. The latter, is dominated by the emission from H$_2$ excited by cosmic rays and is typically\textsuperscript{63,69} about $5 \times 10^3$ photon cm$^{-2}$ s$^{-1}$. Therefore a rate of $4.2 \times 10^{-16}$ s$^{-1}$ is obtained.

To estimate the EPD rate for CO from ASW is less straightforward and requires that a few assumptions to be made:

1. CO bonds weakly to the ASW surface via a hydrogen bond like interactions through dangling OH on the ASW surface. EPD occurs via the same mechanism for CO as for C$_6$H$_6$.

2. Only the fast desorption process will be considered as this relates to isolated molecules on the ASW surface and it might be more relevant to astrophysical irradiation time-scales.

3. The cross-section is assumed to be the same as for C$_6$H$_6$ ($10^{-15}$ cm$^2$).

4. The flux of secondary electrons, $F_{e-}$, produced by cosmic rays in 5 nm of c-ASW can be estimated using the SRIM code\textsuperscript{70,71} for a flux of 1 cm$^{-2}$ s$^{-1}$ 1 MeV H$^+$ ions and this is 4.5 electron cm$^{-2}$ s$^{-1}$ (see ESI\textsuperscript{t}).

5. The energy deposited per ionisation event is assumed to be twice the ionisation potential of the target molecule being ionised (11.0 eV).\textsuperscript{72,73}

6. Few isolated CO molecules are adsorbed on the ASW surface. Thus, no direct effect of the incoming ions will be considered on CO, which will be assumed to be “transparent”.

The EPD rate constant, $k_{EPD}^{CO}$, is defined as:

$$ k_{EPD}^{CO} = \sigma F_{e-} $$

(6)

and its value is $4.5 \times 10^{-15}$ s$^{-1}$. The ratio between the two non-thermal desorption rates is

$$ \frac{k_{EPD}^{CO}}{k_{PD}^{CO}} = 11 $$

(7)

The fast EPD process is thus about one order of magnitude more efficient than PD in dense clouds. Thus, the presence of this efficient desorption channel means that the EPD of an adsorbate, such as CO, on ASW cannot be ignored and should be taken into account in more complete astrochemical models as much as other non-thermal processes. Future works are in preparation to further expand on the astrophysical relevance of these results.

In conclusion, our results confirm the already reported high EPD cross-section of C$_6$H$_6$ from H$_2$O ices and provide additional support for the proposed mechanism involving exciton transport to the H$_2$O/vacuum interface. In addition we have discussed the possible astrophysical impact of our observations on CO accretion and subsequent hydrogenation on the icy ISM dust grains. CO depletion would be faster than currently believed which in turn will slow the formation of hydrogenated CO derivatives and, hence, slow the radiation-driven formation of COM.

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References


