Molecular dynamics simulations of D$_2$O ice photodesorption

C. Arasa, S. Andersson, H. M. Cuppen, E. F. van Dishoeck, and G. J. Kroes

Leiden Observatory, Leiden University, P. O. Box 9513, 2300 RA Leiden, The Netherlands, and Gorlaeus Laboratories, Leiden Institute of Chemistry, Leiden University, P. O. Box 9502, 2300 RA Leiden, The Netherlands

SINTEF Materials and Chemistry, P.O. Box 4760, 7465 Trondheim, Norway and Department of Chemistry, Physical Chemistry, University of Gothenburg, 41296 Gothenburg, Sweden

Leiden Observatory, Leiden University, P. O. Box 9513, 2300 RA Leiden, The Netherlands

Gorlaeus Laboratories, Leiden Institute of Chemistry, Leiden University, P. O. Box 9502, 2300 RA Leiden, The Netherlands

(Received 16 December 2010; accepted 7 April 2011; published online 27 April 2011)

Molecular dynamics (MD) calculations have been performed to study the ultraviolet (UV) photodissociation of D$_2$O in an amorphous D$_2$O ice surface at 10, 20, 60, and 90 K, in order to investigate the influence of isotope effects on the photodesorption processes. As for H$_2$O, the main processes after UV photodissociation are trapping and desorption of either fragments or D$_2$O molecules. Trapping mainly takes place in the deeper monolayers of the ice, whereas desorption occurs in the uppermost layers. There are three desorption processes: D atom, OD radical, and D$_2$O molecule photodesorption. D$_2$O desorption takes places either by direct desorption of a recombined D$_2$O molecule, or when an energetic D atom produced by photodissociation kicks a surrounding D$_2$O molecule out of the surface by transferring part of its momentum. Desorption probabilities are calculated for photoexcitation of D$_2$O in the top four monolayers and are compared quantitatively with those for H$_2$O obtained from previous MD simulations of UV photodissociation of amorphous water ice at different ice temperatures [Arasa et al., J. Chem. Phys. 132, 184510 (2010)]. The main conclusions are the same, but the average D atom photodesorption probability is smaller than that of the H atom (by about a factor of 0.9) because D has lower kinetic energy than H, whereas the average OD radical photodesorption probability is larger than that of OH (by about a factor of 2.5–2.9 depending on ice temperature) because OD has higher translational energy than OH for every ice temperature studied. The average D$_2$O photodesorption probability is larger than that of H$_2$O (by about a factor of 1.4–2.3 depending on ice temperature), and this is entirely due to a larger contribution of the D$_2$O kick-out mechanism. This is an isotope effect: the kick-out mechanism is more efficient for D$_2$O ice, because the D atom formed after D$_2$O photodissociation has a larger momentum than photogenerated H atoms from H$_2$O, and D transfers momentum more easily to D$_2$O than H to H$_2$O. The total (OD + D$_2$O) yield has been compared with experiments and the total (OH + H$_2$O) yield from previous simulations. We find better agreement when we compare experimental yields with calculated yields for D$_2$O ice than when we compare with calculated yields for H$_2$O ice.

I. INTRODUCTION

The formation of molecules in the interstellar medium (ISM) can proceed through several kinds of reactions. Surface reactions on nano- to micrometer sized particles are thought to play a key role in the formation of molecules in the ISM. Dust grains in the ISM consist of a core of silicates and carbonaceous components and, in dense clouds, these dust particles can be covered by icy mantles. The icy mantles contain mainly H$_2$O, and also traces of other molecules (e.g., CO, CO$_2$, NH$_3$, CH$_4$, among others). Observed infrared (IR) spectra reveal that H$_2$O and CO are the most abundant molecules in the icy mantles in the ISM. Recent ground and space based observations have also detected heavy water (D$_2$O) in the ISM.

Ultraviolet (UV) irradiation of an icy grain can photodissociate water molecules and cause desorption of the ice. The flux of UV photons in the ISM is low compared with the lamp UV photon flux used in the laboratories [which varies between, e.g., $(1.1–5.5) \times 10^{13}$ photons cm$^{-2}$ s$^{-1}$ (Ref. 15)], with photon fluxes of the order of $10^3$ photons cm$^{-2}$ s$^{-1}$, which is equivalent to roughly one incident photon per month per grain. The photodissociation dynamics is typically computed over a picosecond time scale, and hence, the photodissociation by one incident photon is completed before the next photon arrives at the ice. The energies of the incident photons ~6–13 eV (Refs. 16–18) cover the first absorption bands of water ice.

Photodissociation and photodesorption of water in ice are of interest to understand astronomical observations of gas-phase water in cold clouds, and also because the photoproducts (H and OH) can proceed to react with co-adsorbed species, which may lead to the formation of more complex
molecules. In addition, the process is interesting from a fundamental chemical physics point of view. Most studies of photodissociation processes of molecules on surfaces have focused on (sub)monolayers of species on mostly metallic surfaces, not on the thick (~100 monolayer (ML)) ices found in the interstellar space. Very different processes can occur in this case.

Several experiments on UV irradiation of amorphous and crystalline H$_2$O ice and D$_2$O ice have been carried out using different analysis techniques and different light sources.

In order to obtain insight into the basic molecular processes, the photodissociation of H$_2$O molecules in amorphous and crystalline ice in the temperature range of 10–90 K has been studied using molecular dynamics (MD) simulations. The most important photodesorption mechanism after photodissociation of water in the top three MLs of the ice surface is H atom photodesorption, followed by OH radical photodesorption, and H$_2$O molecule photodesorption. The calculated H$_2$O photodesorption probability is due to two mechanisms. (1) The direct mechanism: H and OH recombine after H$_2$O photodissociation to form H$_2$O, which eventually desorbs. (2) The kick-out mechanism: an energetic H atom released after photodissociation kicks out one of the surrounding water molecules by a transfer of momentum.

Many key experiments studying water ice photodesorption have been performed for D$_2$O rather than H$_2$O. In contrast, the MD simulations have so far been carried out only for H$_2$O. In order to better compare with experiments and to identify isotope effects on the photodesorption processes, we present here results of MD simulations of the UV photodissociation of amorphous D$_2$O ice at different temperatures.

In Sec. II, we present the methods used in this study, in Sec. III, the main results in comparison with previous UV photodissociation of amorphous H$_2$O ice results are presented, and in Sec. IV, the concluding remarks are given.

II. METHODS

A. Potentials

The total analytical potential energy surface (PES) for the ice in the photodissociation calculations is the same as in our previous studies and can be written as follows:

$$ V_{\text{tot}} = V_{\text{ice}} + V_{\text{H$_2$O$_2$}} + V_{\text{H$_2$O$_{\text{exc}}$}}. $$

The first term describes the intermolecular interactions between the H$_2$O molecules inside the ice excluding the H$_2$O molecule that is photoexcited. These interactions are described by the TIP4P potential with all molecules kept rigid.

The second term refers to the intermolecular interactions of the photoexcited molecule, which is treated as fully flexible with the rigid ice molecules, and the third term is the intramolecular potential of the photoexcited molecule. These potential terms also cover all interactions involved in the dissociation and possible recombination of the excited molecule. The potentials are exactly the same as previously used for photodissociation of H$_2$O ice. All details of the potentials and the functions used to switch between different potentials are given in Refs. 41 and 43.

B. Amorphous ice surface

To study the UV photodissociation of D$_2$O ice, we simply changed the mass of the H atom to that of the D atom. (Of course, all the interactions that take place in and on the ice during the photodissociation are described with the same potentials employed for the UV photodissociation of H$_2$O ice.)

Crystalline and amorphous D$_2$O ice surfaces were constructed using the MD method, and using the same procedure and cell parameters employed before to model H$_2$O ice.

Starting from the normal hexagonal ice (I$_h$) crystalline ice configuration (containing 8 bilayers (BLs) (16 MLs) with 60 (30) molecules in each ML), the amorphous ice surface was set up at 10, 20, 60, or 90 K using the “fast quenching” method. Further details can be found in our previous studies. Since the resulting amorphous ice surface has a more irregular bonding structure than the crystalline ice surface, assigning molecules to MLs is not straightforward. In our most recent study, a new definition of ML (binning method 2) was tested and shown to be a more realistic way to assign molecules to MLs. This binning (method 2) is used in this study and it consists of choosing a molecule and finding the first 23 closest molecules in terms of $(x, y)$ coordinates. This leads to 24 molecules that are divided in 12 bins of two molecules each, based on their $z$ center of mass coordinates. The 12 bins represent the top 12 MLs of the ice in which the molecules are allowed to move.

C. Initial conditions and dynamics

For each of the top four MLs, all the molecules were chosen to be photodissociated, and for each molecule, 200 different initial configurations were considered. To initialize the trajectories, a Wigner phase-space distribution function fitted to the ground-state vibrational wavefunction of gas-phase water is used. In the case of heavy water the trajectories are initialized by using the corresponding Wigner distribution of gas-phase heavy water, which has the same functional form as for gas-phase water (Eq. (5.13) in Ref. 50), but with $\alpha_{\text{D}_2\text{O}} = \sqrt{2} \alpha_{\text{H}_2\text{O}}$ (Eq. (5.16) in Ref. 50). The initial coordinates and momenta of the atoms from the dissociating molecule are sampled using a Monte Carlo procedure. Then, a Franck-Condon excitation is performed and the system is put on the first electronically excited state, on the DK $\tilde{A}^1B_1$ PES.

The excitation energies are calculated by computing the energy difference between a D$_2$O ice with an excited molecule and one with a ground state D$_2$O molecule (both molecules with the same coordinates). The calculated D$_2$O amorphous ice spectrum is shifted 0.02 eV with respect to that for H$_2$O amorphous ice.

To simulate the dynamics of a photodissociation event, Newton’s equations of motion are integrated in time with a time step of 0.02 fs and a maximum time of 20 ps. The stop
The average D atom photodesorption probability and the average H atom photodesorption probability\textsuperscript{43} taken over the top four monolayers (e.g., $P_{\text{H-des}} = \sum_{i=4}^{4} P_{\text{H-des}}$) are plotted in Fig. 1. The average H photodesorption probability is somewhat larger than that of D, by about 6\% for all $T_{\text{ice}}$ studied. The total deuterium (hydrogen) atom photodesorption probability is calculated by summing over two different processes: one in which the D (H) atom desorbs while the OD (OH) stays trapped in the surface, and one in which both photofragments desorb from the ice surface (outcomes 1 and 3).

The deuterium and hydrogen atom photodesorption probabilities are higher than the OD (OH) and D$_2$O (H$_2$O) photodesorption probabilities in the uppermost monolayers of the D$_2$O (H$_2$O) amorphous ice (see Fig. 2 where the H and D photodesorption probabilities are displayed versus ice temperature and ML). This is because D and H atoms are smaller and are formed with higher kinetic energies immediately after D$_2$O and H$_2$O photodissociation, which facilitates the desorption of these atoms.\textsuperscript{40-43}

The average photodesorption probability of D is smaller than that of H mostly because the probabilities of D atom photodesorption in the third and fourth monolayers (Fig. 2) are smaller than those for the H atom. This trend is expected because D is heavier than H. Therefore, the efficiency of energy transfer between D and D$_2$O molecules is larger than the corresponding efficiency between H and H$_2$O molecules. If the photoexcited molecule is isolated (i.e., in the absence of the surrounding ice), the initial kinetic energy in which D and H atoms are formed after D$_2$O and H$_2$O photodissociation should be similar in order to achieve energy conservation. But, in the presence of ice, D atoms lose more kinetic energy than H atoms when they interact with the surrounding molecules due to the larger efficiency of energy transfer, and they are therefore less able to penetrate the ice when moving through the upper layers.

The dependence on ice temperature is negligible. The average D atom photodesorption probability (Fig. 1) is almost constant at ~54\%. However, this probability depends on the ML, where the photoexcited molecule was initially located (Fig. 2). In the top two MLs the probability is high (~90\% to ~70\%), but it drops in the third ML and further below, because other processes such as trapping are in competition. Trapping becomes important because deeper in the ice the
FIG. 2. Total probability of D atom (dashed line) and H atom (solid line) (Ref. 43) photodesorption (per absorbed UV photon) versus temperature, for the uppermost four MLs.

structure is more closed and the molecules from the ice above can impede the D atom from reaching the ice surface.

D atoms travel through the D₂O ice at 90 K by an average distance of 8.4 Å before they become trapped, whereas H atoms travel around 9.1 Å. The OD and OH radicals travel 2.2 Å and 1.9 Å, respectively. The recombined D₂O and H₂O move on average a distance of 1.8 Å and 2.0 Å, respectively. Therefore, the mobility of the photofragments inside the ice is slightly affected by the mass of the photofragments: H atoms move further than D atoms until they become trapped, because H is lighter than D. The maximum distances travelled, which are about tens of angstroms should enable reaction with other species trapped in the ice. This could explain the formation of more complex molecules in the ISM.

B. OD radical photodesorption

The second main photodesorption mechanism in the uppermost MLs of the ice is OD photodesorption. Figure 3 shows that the average of the OD photodesorption probabilities taken over the top four MLs is larger than that for OH for all ice temperatures studied. To calculate the OD (OH) photodesorption probabilities we have summed over the probabilities of two pathways: the probability of the channel in which the OD (OH) radical desorbs while the D (H) atom remains trapped in the ice surface, and the probability of the channel in which both photofragments leave the ice surface (outcome 2 and 3, respectively).

The probabilities per monolayer are plotted in Fig. 4 versus ice temperature. Those for OD are much larger than those for OH (Ref. 43) in the top two monolayers, which gives rise to a larger total average OD photodesorption probability (Fig. 3). In the absence of the surrounding ice, the water fragments (D and OD, H and OH) have to obey momentum conservation \( p_X = -p_{OX}, X = H, D \) and energy conservation \( E_X + E_{OX} = E_{X,O} = E \), where \( E \) being the initial available energy \( E_{exc} - E_{diss}(X_2O), X = H, D \); the excitation energy \( E_{exc} \) is in the range 7.5–9.5 eV with a peak at 8.6 eV, and the dissociation energy \( E_{diss}(X_2O) \approx 5.4 \text{ eV} \) (Ref. 53), leading to the following equation:

\[
\frac{1}{2}m_{OX}v_{OX}^2 = \frac{E}{(1 + \frac{m_X}{m_O})}.
\]

Thus, if the molecule is isolated and dissociates, the OD radicals will be formed with a higher initial translational energy than the OH radicals \( E_{OD} < E_{10} \text{ and } E_{OH} < E/18 \), which leads to \( E_{OD} \approx 1.8 \times E_{OH} \) according to Eq. (3). Because the OD radicals have a higher initial translational energy than the OH radicals in the uppermost monolayers, they leave the ice surface more easily.

The OD photodesorption probability decreases with an increasing depth, similar to the OH photodesorption probability. In the third and fourth MLs, the OD and OH photodesorption probabilities drop to less than \( 1 \times 10^{-2} \) because the OD and OH radicals do not have enough translational energy to escape from the ice surface.

FIG. 3. The probability of OD radical (dashed line) and OH radical (solid line) photodesorption averaged over the top four MLs per absorbed UV photon is shown as a function of ice temperature. OH radical results from Ref. 43.

FIG. 4. Total probability of OD radical (dashed line) and OH radical (solid line) (Ref. 43) photodesorption (per absorbed UV photon) versus ice temperature, for the uppermost four MLs.
An oscillatory effect is observed when the OD and OH photodesorption probabilities are plotted in MLs 1–4 versus ice temperature (Fig. 4). We attributed these oscillations in our previous paper to the irregular nature of the amorphous ice surface, which makes it very complicated to assign molecules to specific MLs, and also to the finite sample size of about 30 molecules per ML.

The average OD and OH photodesorption probabilities increase with ice temperature by ~24% and ~25%, respectively, from 10 to 90 K. If longer time scales in our simulations could be considered, a stronger dependence on ice temperature would be expected, because processes like thermal diffusion and thermal desorption are more efficient at higher \( T_{\text{ice}} \).\(^{43} \)

C. \( \text{D}_2\text{O} \) molecule photodesorption

1. Kick-out vs. direct mechanism

The third photodesorption channel upon UV photodissociation of \( \text{D}_2\text{O} \) amorphous ice is \( \text{D}_2\text{O} \) molecule photodesorption. This is due to two mechanisms: the direct and the kick-out mechanism. The direct mechanism consists of the recombination of the D atom and the OD radical that leads to the formation of an energetic \( \text{D}_2\text{O} \) molecule that eventually desorbs (outcome 5). In this situation the \( \text{D}_2\text{O} \) molecule has a high probability of desorbing from the ice in a vibrationally excited state.\(^{42} \) The kick-out mechanism takes place after the photodissociation of a \( \text{D}_2\text{O} \) molecule when an energetic D atom transfers momentum to one of the surrounding \( \text{D}_2\text{O} \) molecules, which is then likely to desorb vibrationally cold.\(^{29,38,43} \) Since in our model we cannot quantify the energy transfer from X to internal modes of \( \text{X}_2\text{O} \), because the kicked out molecule is treated as internally rigid, we have carried out quasi-classical trajectory calculations on the isolated X–\( \text{X}_2\text{O} \) system at different incident X atom kinetic energies (X = H, D). Here we only report the results for \( E_X = 1.5 \text{ eV} \) because it is the average kinetic energy with which H and D atoms kick a surrounding molecule out of the ice at \( T_{\text{ice}} = 10 \text{ K} \). Further details on these calculations will be reported in the paper we are preparing.\(^{54} \) Gas phase collisions at \( E_X = 1.5 \text{ eV} \) lead to final average vibrational energies of ~0.16 eV for \( \text{H}_2\text{O} \) and ~0.77 eV for \( \text{D}_2\text{O} \), whereas in the ice system, recombined \( \text{H}_2\text{O} \) and \( \text{D}_2\text{O} \) molecules desorb from the ice surface with average ro-vibrational energies of 5.3 and 5.4 eV, respectively. Thus, the kicked out molecules are much more likely to be formed in states with lower vibrational energies than the molecules formed after recombination of the photofragments. Yabushita et al.\(^{29} \) and Hama et al.\(^{38} \) observed photodesorbed \( \text{X}_2\text{O} \) in the ground vibrational state by using resonance-enhanced multiphoton ionization detection methods and classified them as kicked out. They speculated that when the X atom kicks out an \( \text{X}_2\text{O} \) molecule, most of the energy is transferred into translation and much less into internal energy because the X atom hits the \( \text{X}_2\text{O} \) molecule close to the center of mass. This is supported by our simulations (e.g., ~89% of the trajectories classified as kicked out occurs when the X atom kicks the \( \text{X}_2\text{O} \) molecule close to the oxygen atom at \( T_{\text{ice}} = 10 \text{ K} \) (X = H, D). Isotope effects are also observed in our calculations: the efficiency of energy transfer from D to intramolecular vibrational modes of \( \text{D}_2\text{O} \) molecule in gas phase collisions is larger than the corresponding efficiency for collisions of H with \( \text{H}_2\text{O} \), because D is heavier than H and because the vibrational frequencies of \( \text{D}_2\text{O} \) are lower than those of \( \text{H}_2\text{O} \).

Figure 5(a) shows the average of the \( \text{D}_2\text{O} \) photodesorption probabilities compared with those for \( \text{H}_2\text{O} \) (Ref. 43) over the top four monolayers versus ice temperature. The average values for \( \text{D}_2\text{O} \) and \( \text{H}_2\text{O} \) (Ref. 43) due to the kick-out mechanism are displayed in Fig. 5(b), and those due to the direct mechanism are displayed in Fig. 5(c).

The total average \( \text{D}_2\text{O} \) photodesorption probabilities are larger than those for \( \text{H}_2\text{O} \) [Fig. 5(a)] because of the large contribution of the \( \text{D}_2\text{O} \) kick-out mechanism [Fig. 5(b)]. For every ice temperature studied, the \( \text{D}_2\text{O} \) photodesorption probability due to the kick-out mechanism is larger than that for \( \text{H}_2\text{O} \) [Fig. 5(b)], because in the case of heavy water, the D atom formed after \( \text{D}_2\text{O} \) photodissociation has a larger momentum (by about a factor \( \sqrt{2} \)) than the H atom formed after \( \text{H}_2\text{O} \) photodissociation. In addition, the efficiency of

![Figure 5](image-url)
momentum transfer from D to D$_2$O is larger than that from H to H$_2$O, so that the kick-out mechanism is much more successful for D$_2$O. The effect occurs mainly after photoexcitation in the second and third monolayers [see Figs. 6(b) and 6(c)], because a D atom produced in these MLs is more likely to kick out a molecule located above it. The kick-out photodesorption probabilities are much lower in the first and fourth MLs [Figs. 6(a) and 6(d)], where there are no important differences between D$_2$O and H$_2$O.

We have calculated the probabilities of the parallel outcomes that take place in coincidence with the kicking out of a D$_2$O molecule by an energetic D atom. These probabilities are summarized in Table I. We do not observe differences with those calculated for the H$_2$O kicked out molecules from H$_2$O ice (Table I in Ref. 43). The most dominant simultaneous process is that where the D atom that kicks out the D$_2$O molecule also desorbs, while the OD fragment is trapped (Table I). The next most important parallel processes are those in which the photofragments recombine and form a D$_2$O molecule that remains trapped in the ice, and those in which both photofragments are trapped inside the ice at separate locations. It is also possible that two molecules desorb at the same time, i.e., the kicked out molecule and a recombined molecule, but this process occurs with a very low probability.

### 2. Trends with ice temperature

The total D$_2$O photodesorption probability increases faster with ice temperature than that for H$_2$O [Fig. 5(a)]: by 130% vs 30%, going from 10 to 90 K. The average desorption probability due to the direct mechanism [Fig. 5(c)] is relatively small, and there are no differences between the average D$_2$O and H$_2$O photodesorption probabilities: both rise with ice temperature by only ~30% going from 10 to 90 K. However, the D$_2$O average kick-out photodesorption probability increases strongly with ice temperature (by ~180% from 10 to 90 K), whereas the H$_2$O kick-out average photodesorption probability shows a much weaker increase with ice temperature (by ~50% from 10 to 90 K). Thus, the stronger trend with ice temperature for D$_2$O results from the increase of the D$_2$O kick-out photodesorption probability [Fig. 5(b)]. This probability increases with ice temperature because the molecules have higher initial kinetic energies at higher ice temperatures, which promotes the desorption of the surrounding molecules through the kick-out mechanism.

At higher ice temperatures the D$_2$O kick-out mechanism has a high probability (per absorbed UV photon): 2.8% at 60 K after photoexcitation in the 3rd ML [Fig. 6(c)] and 2.2% at 90 K after photoexcitation in the 2nd ML [Fig. 6(b)]. The probabilities for D$_2$O and H$_2$O desorption through the kick-out mechanism in the top four MLs (Fig. 6) show an oscillatory dependence on ice temperature, as also seen before for OD and OH photodesorption probabilities versus ice temperature and monolayer (Fig. 4). These oscillations are attributed to the corrugation of the amorphous ice surface (for more details see Ref. 43).

### D. Energies of the kicked out molecules

The average translational and rotational energies taken over the top four MLs of the D$_2$O and H$_2$O (Ref. 43) kicked out molecules are plotted versus ice temperature in Fig. 7, together with the corresponding experimental values for H$_2$O photodesorbed molecules ($v = 0$) at 90 K. The translational energies tend to increase with ice temperature, because the energy of the ice also rises. The final rotational energies are low.
and do not display any dependence on ice temperature. The calculated translational and rotational energies do not show a significant isotope effect. Our calculations cannot say anything about the vibrational state of the kicked out molecules, because the molecules that are not photoexcited are kept rigid in our model. However, it seems unlikely that the kicked out molecules would emerge highly vibrationally excited. The average translational and rotational energies at 90 K of the H$_2$O kicked out molecules are 0.29 and 0.044 eV, respectively, and of the D$_2$O kicked out molecules 0.27 and 0.021 eV, respectively. These results are in good agreement with the experimental translational and rotational energies of the H$_2$O desorbed molecules observed by Hama et al. and experimental average translational and rotational energies of H$_2$O molecules desorbed in their ground vibrational state at $T_{\text{ice}} = 90$ K.

The agreement between theory and experiment is better at low ice temperatures, (e.g., $\xi = 1.3$ at 10 K and $\xi = 2.3$ at 90 K). This trend strengthens our explanation that the difference between theory and experiments can be due to long time scale processes promoted by prolonged irradiation effects leading to an accumulation of radicals, thermal desorption, and thermal diffusion. At higher ice temperatures, the photofragments, possibly formed by different photodissociation events, become more mobile, allowing them to recombine and eventually desorb as a consequence of the excess energy. Some of the OD photofragments that are trapped deeper in the ice, could probably desorb at long time scales due to a higher diffusion rate at higher $T_{\text{ice}}$.

The calculated average of the total (O + H$_2$O, for X = H or D) photodesorption probability per absorbed UV photon is larger for D$_2$O ice than for H$_2$O ice at every ice temperature (Fig. 8).

In our previous study, we compared the total photodesorption yield of the O containing species (H$_2$O$_{\text{des}}$ + OH$_{\text{des}}$) with the total experimental photodesorption yield (Eq. (4) in Ref. 15). In the calculation of this yield, the deuterium (or hydrogen) atom photodesorption is not included (outcomes 2, 3, 5, and the kick-out), because D (or H) was not detected in the experiments by Öberg et al. The photodesorption yields (i.e., photodesorption probabilities per incident photon) have been calculated through Eq. (2) (see Sec. II C).

Although the actual experimental photodesorption yields (Table II) (Ref. 15) were for D$_2$O ice, Öberg et al. applied the results for D$_2$O ice to H$_2$O ice (Eq. (4) in Ref. 15), because they found that the total photodesorption yields from D$_2$O and H$_2$O were indistinguishable (i.e., no isotope effects) within the experimental uncertainties (60%) at 18 and 100 K.

Table II contains the total experimental photodesorption yield, the computed total (H$_2$O$_{\text{des}}$ + OH$_{\text{des}}$), and the computed total (D$_2$O$_{\text{des}}$ + OD$_{\text{des}}$) photodesorption yield per incident photon, the ratio $\xi$ between the experimental yield and the calculated (H$_2$O$_{\text{des}}$ + OH$_{\text{des}}$) yield, and the same for the (D$_2$O$_{\text{des}}$ + OD$_{\text{des}}$) yield, at all ice temperatures. $\xi$ increases from 3.0 to 5.9 for H$_2$O ice and from 1.3 to 2.3 for D$_2$O ice. Thus, the computed (D$_2$O$_{\text{des}}$ + OD$_{\text{des}}$) photodesorption yield is significantly larger than the computed (H$_2$O$_{\text{des}}$ + OH$_{\text{des}}$) yield at all ice temperatures, as also illustrated in Fig. 8, and compares better with the experimental D$_2$O photodesorption yield than the yield calculated for amorphous H$_2$O ice.

The agreement between theory and experiment is better at low ice temperatures, (e.g., $\xi = 1.3$ at 10 K and $\xi = 2.3$ at 90 K). This trend strengthens our explanation that the difference between theory and experiments can be due to long time scale processes promoted by prolonged irradiation effects leading to an accumulation of radicals, thermal desorption, and thermal diffusion. At higher ice temperatures, the photofragments, possibly formed by different photodissociation events, become more mobile, allowing them to recombine and eventually desorb as a consequence of the excess energy. Some of the OD photofragments that are trapped deeper in the ice, could probably desorb at long time scales due to a higher diffusion rate at higher $T_{\text{ice}}$. In our simulations we can only reach the picosecond time scale, therefore these kind of secondary processes are beyond the scope of our simulations. Another difference is the UV wavelength covered by the lamp used in the experiments. This UV lamp includes Lyman-α photons which can excite H$_2$O to the Α state whereas our calculations consider only the Α state. Given the experimental uncertainties and our approximations (such as the use of a gas phase PESs for the H$_2$O intramolecular interactions).
interactions, the freezing of the intramolecular degrees of freedom of the surrounding molecules, and the short time scale of our simulations,
interactions may be considered to be in reasonable agreement. An important result for astrochemists is that the computational results fall within the range of the photodesorption probability per incident photon ($1 \times 10^{-3} - 3.5 \times 10^{-3}$)
(Refs. 55–59) used to model astrophysical environments. However, our calculations suggest that the computed total photodesorption yield can be different for H$_2$O and D$_2$O ice, in contrast to the experimental results.

IV. CONCLUSIONS

In this work we have studied the processes following UV photodissociation of D$_2$O in amorphous heavy water ice and compared them with previous UV photodissociation results in H$_2$O ice at different ice temperatures, in order to investigate isotope effects in photodesorption.

D atom photodesorption is the most important desorption process in the uppermost MLs of the ice, like H atom photodesorption. The average D atom photodesorption probability is smaller than that of the H atom, because in the top two MLs of the ice both the H and D atom can easily escape from the ice surface, but if the atoms are located in the third and fourth MLs, the D atom is less likely to penetrate the upper ice layers due to the more efficient collision energy transfer to D$_2$O. Therefore, the D atom photodesorption probabilities in these MLs decrease, and the same is then for the average D atom photodesorption probability. The D and H atom photodesorption probabilities do not show any dependence on ice temperature.

OD and OH radical photodesorption constitute the second most important desorption channel in the top two MLs of the ice. Deeper into the ice the probabilities of these processes drop because OD and OH radicals do not have enough translational energy to desorb from the surface. The average OD photodesorption probability is higher than that of OH. This trend can be explained by the initial translational energy of OD being higher by about a factor of 1.8, a result obtained if the photoexcited molecule is considered to be isolated (i.e., in the absence of the surrounding ice) and the laws of momentum and energy conservation are applied. The average OD photodesorption probabilities increase smoothly with ice temperature, by about 24% from 10 to 90 K.

The third most important desorption mechanism is D$_2$O and H$_2$O photodesorption. This process takes place either by direct desorption of the photoexcited molecule after the recombination of D (H) and OD (OH) or by indirect desorption due to an energetic D (H) atom which transfers a part of its momentum to a surrounding molecule that is kicked out from the ice surface. The average photodesorption probability is higher for D$_2$O than for H$_2$O at all ice temperatures considered. This trend is due to the contribution of the kick-out mechanism, which is much more important for D$_2$O than for H$_2$O. This result is expected because the photoproduced D atoms have a higher average momentum (by about a factor of $\sqrt{2}$) than the H atoms, and because energy transfer in D–D$_2$O collisions is more efficient than energy transfer in H–H$_2$O collisions. The kick-out mechanism mainly takes place when the photoexcited molecule is initially located in the second and third MLs of the ice. Photodissociation leads to an energetic D (H) atom that can transfer its momentum to a molecule located above it, which will desorb from the ice if it has enough kinetic energy. The average direct photodesorption probability (which involves recombination) does not show any isotope effect. The average total D$_2$O photodesorption probability tends to increase with ice temperature faster than that of H$_2$O: by $\sim$130% vs $\sim$30%, from 10 to 90 K.

Experiments show, and a consideration of the mechanism suggests, that the kicked out molecules leave the surface vibrationally cold. In contrast, the molecules that desorb due to the direct mechanism are formed vibrationally excited. The average translational and rotational energies in which the D$_2$O and H$_2$O molecules desorb due to the kick-out mechanism have been calculated and compared with the corresponding experimental values at 90 K. The agreement between our MD calculations and the experimental measurements is good, and leads to the conclusion that the final energies with which the kicked out molecules are formed do not display an isotope effect.

We have also estimated the total photodesorption probability (OD$_{\text{des}} +$ D$_2$O$_{\text{des}}$) per incident photon from the total photodesorption probabilities per absorbed UV photon, and compared this quantity with the previously calculated values for (OH$_{\text{des}} +$ H$_2$O$_{\text{des}}$), and with the available experimental yields. Our total photodesorption probability for D$_2$O compares better with the experimental photodesorption yield than that for H$_2$O, and also better at low ice temperatures. Presumably at higher ice temperatures long time scale processes become increasingly important, such as diffusion and thermal desorption, which are not covered in our picosecond simulations.

---

**TABLE II.** Experimental $^a$, theoretical (OH$_{\text{des}} +$ H$_2$O$_{\text{des}}$), and theoretical (OD$_{\text{des}} +$ D$_2$O$_{\text{des}}$) photodesorption yields per incident photon, the experimental yield/theoretical yield $\xi$ (H$_2$O), and the experimental yield/theoretical yield $\xi$ (D$_2$O) at all ice temperatures.

<table>
<thead>
<tr>
<th>$T_{\text{ice}}$ / K</th>
<th>Exp. (OH$_{\text{des}} +$ H$<em>2$O$</em>{\text{des}}$), X = H or D $\times 10^{-3}$</th>
<th>(OH$_{\text{des}} +$ H$<em>2$O$</em>{\text{des}}$) $\times 10^{-3}$</th>
<th>(OD$_{\text{des}} +$ D$<em>2$O$</em>{\text{des}}$) $\times 10^{-3}$</th>
<th>$\xi$ (H$_2$O)</th>
<th>$\xi$ (D$_2$O)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>1.62 ± 0.48</td>
<td>0.54 ± 0.04</td>
<td>1.27 ± 0.063</td>
<td>3.0</td>
<td>1.3</td>
</tr>
<tr>
<td>20</td>
<td>1.94 ± 0.56</td>
<td>0.57 ± 0.04</td>
<td>1.46 ± 0.066</td>
<td>3.4</td>
<td>1.3</td>
</tr>
<tr>
<td>30</td>
<td>2.26 ± 0.64</td>
<td>0.71 ± 0.09</td>
<td>1.57 ± 0.068</td>
<td>3.2</td>
<td></td>
</tr>
<tr>
<td>60</td>
<td>3.22 ± 0.88</td>
<td>0.71 ± 0.05</td>
<td>1.83 ± 0.074</td>
<td>5.9</td>
<td>2.3</td>
</tr>
<tr>
<td>90</td>
<td>4.18 ± 1.1</td>
<td>0.71 ± 0.05</td>
<td>1.83 ± 0.074</td>
<td>5.9</td>
<td>2.3</td>
</tr>
</tbody>
</table>

$^a$Calculated from the empirical fit of the total photodesorption yield, Eq. (4) in Ref. 15.
Current experiments cannot distinguish between (OH + H₂O) and (OD + D₂O) yields within the experimental uncertainties of 60%. More accurate future experiments may reveal the isotope effects predicted here.

ACKNOWLEDGMENTS

The authors would like to thank M. C. van Hemert and T. P. M. Goumans for valuable discussions. This project was funded with computer time by NCF/NWO, and with TOP Grant No. 700.56.321 by CW/NWO.