ABSTRACT: We examine the initial differential sticking probability of CH₄ and CD₄ on CH₄ and CD₄ ices under nonequilibrium flow conditions using a combination of experimental methods and numerical simulations. The experimental methods include time-resolved in situ reflection–absorption infrared spectroscopy (RAIRS) for monitoring on-surface gaseous condensation and complementary King and Wells mass spectrometry techniques for monitoring sticking probabilities that provide confirmatory results via a second independent measurement method. Seeded supersonic beams are employed so that the entrained CH₄ and CD₄ have the same incident velocity but different kinetic energies and momenta. We found that as the incident velocity of CH₄ and CD₄ increases, the sticking probabilities for both molecules on a CH₄ condensed film decrease systematically, but that preferential sticking and condensation occur for CD₄. These observations differ when condensed CD₄ is used as the target interface, indicating that the film’s phonon and rovibrational densities of states, and collisional energy transfer cross sections, have a role in differential energy accommodation between isotopically substituted incident species. Lastly, we employed a mixed incident supersonic beam composed of both CH₄ and CD₄ in a 3:1 ratio and measured the condensate composition as well as the sticking probability. When doing so, we see the same effect in the condensed mixed film, supporting an isotopic enrichment of the heavier isotope. We propose that enhanced multi-phonon interactions and inelastic cross sections between the incident CD₄ projectile and the CH₄ film allow for more efficacious gas–surface energy transfer. VENUS code MD simulations show the same sticking probability differences between isotopologues as observed in the gas–surface scattering experiments. Ongoing analyses of these trajectories will provide additional insights into energy and momentum transfer between the incident species and the interface. These results offer a new route for isotope enrichment via preferential condensation of heavier isotopes and isotopologues during gas–surface collisions under specifically selected substrate, gas-mixture, and incident velocity conditions. They also yield valuable insights into gaseous condensation under non-equilibrium conditions such as occur in aircraft flight in low-temperature environments. Moreover, these results can help to explain the increased abundance of deuterium in solar system planets and can be incorporated into astrophysical models of interstellar icy dust grain surface processes.

INTRODUCTION

Adsorption is a key process in both astrophysical and terrestrial environments as it serves as the first step in many gas–surface interactions. In extraterrestrial environments where chemical species are scarce, adsorption onto an interstellar grain, planetesimal, or other larger body controls many combinatorial reactions. The formation of larger organic molecules becomes more feasible when species can engage on a surface rather than the void of space. In addition, the astrophysical environment is abundant with isotopes of many chemical species. In order to accurately model the chemical abundances, we need to better understand how differences in mass can influence the ability of a species to adsorb under specified conditions, and thus lead to observed relative isotope abundances.
Interstellar methane is the most common hydrocarbon, existing in both the gaseous and the solid form.13–19 Methane is commonly found in the gaseous planetary atmospheres or as molecular ices intermixed with water ice matrices.20,21 As the most basic hydrocarbon, CH4 serves as a base for addition reactions which form larger hydrocarbon species.22 Additionally, the isotopic twin of CH4, CD4, can serve as a model for understanding the effects and the abundance of deuterium within these environments.12,23 Theoretical methods and gas chromatography have found that the isotopic difference in CH4 and CD4 stems from the difference in polarizability and length of the C–H and C–D bonds; however, no studies have reported how this difference might translate into its sticking probability.12,24,25 Studying CH4 and CD4 adsorption is an excellent model system to determine how slight mass differences in the condensate and projectile can impact adsorption and surface abundance of isotopic species.

Here we present the first study of the isotopic sticking probability of CH4 and CD4 as a function of translational beam energy on CH4 and CD4 thin films under ultrahigh-vacuum (UHV) conditions at low temperatures using the King and Wells method,26 complemented by in situ infrared spectroscopic studies of gaseous condensation. VENUS code molecular dynamics (MD) simulations show the same sticking probability differences between isotopologues as were observed in the gas-surface scattering experiments. Taken together, these results accurately and independently determine the sticking probability, allowing us to explore how differences in isotopic composition of the surface and incident molecular mass can impact the overall energy accommodation, and thus adsorption of the gaseous species onto the film.

Key to these studies is the use of essentially monoenergetic seeded supersonic beams so that the CH4 and CD4 have the same incident velocity but different kinetic energies and momenta. It is shown that as the incident velocity of CH4 and CD4 increases, the sticking probabilities for both molecules on a CH4 condensed film decrease systematically, but that preferential sticking and condensation occur for CD4. These observations differ when condensed CD4 is used as the target interface, indicating that the film’s phonon and rovibrational densities of states, and collisional energy transfer cross sections, play a role in differential energy accommodation between isotopically substituted incident species. In addition, a mixture of gaseous CH4 and CD4 was grown on a methane thin film. While both species adsorbed creating a mixed isotopologue condensate, we saw an increased abundance of CD4 versus CH4 within the film as opposed to initial beam concentration. We demonstrate an isotopic enrichment for CD4 in our mixed surface based on the difference in sticking probabilities between CH4 and CD4.

This experiment builds on previous work by our group where CH4 sticking was investigated on the surfaces of D2O of varying morphologies and where H2O and D2O sticking on their own films were studied.20–28 In particular, we consider a similar isotopic experiment as the H2O and D2O sticking but expand to study CH4 and CD4 sticking on both films rather than only their own films.27,28 Additionally, previous work examined the sticking of only CH4 on H2O which we expand to include CH4 and CD4 sticking onto both CH4 and CD4 ices to examine how the mass difference can affect the overall sticking.28 We examine how these differences in mass, energy, and surface composition can affect the ability of the film to absorb and dissipate energy from the impinging molecules to allow adsorption onto the film structure.

Our work demonstrates differential condensation between methane isotopologues under specifically selected substrate, gas-mixture, and incident velocity conditions. The demonstrated outcomes have obvious implications for the development of novel isotopic enrichment and separation techniques. These results also provide new insights into gaseous condensation under non-equilibrium conditions such as occur in aircraft flight in low-temperature environments. More broadly, this work is critical to understanding the nature of methane adsorption within astrophysical environments. Our sticking probability differences can be incorporated into astrophysical models to explain molecular abundances and increased deuterium abundance in cometary ices and outer solar system planets. Aside from astrophysical environments, adsorption has implications into fields such as heterogeneous catalysis or thin film growth where the adsorption process serves as the first step in film formation.28

## EXPERIMENTAL METHODS

All experiments were conducted in a molecular beam scattering instrument previously discussed in full detail.28 Briefly, this instrument consists of a UHV chamber with base pressures of 10−10 Torr connected to a triply differentially pumped molecular beamline. In the main chamber, a state-of-the-art closed-cycle helium-cooled sample manipulator (Advanced Research Systems) enables precise and accurate temperature control of the Au(111) sample substrate between 16 and 800 K. The crystal is exposed to the impinging beam at normal incident angle and monitored in real time with optics for in situ reflection absorption infrared spectroscopy (RAIRS). Gas scattering and incident flux monitoring occur with a residual gas analyzer (RGA).

All RAIR spectra are analyzed with Gaussian peak fitting atop cubic baselines. Spectra were acquired with a Nicolet 6700 infrared spectrophotometer (Thermo Fisher) using incident p-polarized IR radiation at an angle of 75° to the Au(111) crystal and a liquid-nitrogen-cooled mercury cadmium telluride (MCT/A) detector. Each RAIR spectrum is an average of 25–200 scans taken by using 4 cm−1 resolution with a clean Au(111) sample used for the background correction.

CH4 or CD4 was dosed on the Au(111) substrate via beam deposition at 18 K prior to measurements at 20 K. Dosing conditions resulted in a deposition rate of 0.5 layers per second. CH4 and CD4 beams were produced by expanding 1% CH4 in H2 or 1% CD4 in H2 at stagnation pressures of 150–400 psi through a 15 μm platinum pinhole. Resistively heating the beam nozzle from room temperature up to 1100 K resulted in beam velocities of up to 4600 m/s and rotationally cold molecules resulting from the seeded expansion. The translational energy distribution widths (∆v/v) ranged from 5 to 24%. We note that the velocity slip between the two isotopologues varied between 0 and at most 100 m/s with velocities spanning 2400 to 4600 m/s; therefore, the incident velocities of the two isotopologues were essentially identical for the purposes of a given experiment. Incident velocities were measured by time-of-flight methods using a mechanical chopper to modulate the beam prior to detection with an in-line quadrupole mass spectrometer. To confirm all the results and further understand phonon interactions at cold temperatures, a mixed beam was produced by expanding 1% CD4 and 3% CH4 in H2.
The sticking probability was determined by using the King and Wells technique. This was previously described in more detail for our system, and a typical King and Wells experiment conducted on a CH4 surface temperature at 20 K is shown in Figure 1 where m/z = 20 for CD4 is monitored as a function of time by using an RGA out of line with the beam. The experiment involves monitoring the background signal (P1), indirect flux (P2) with a flag in front of the substrate and removal of that flag where molecules start to stick (P3) to calculate the initial sticking probability:

\[ S = \frac{P_3 - P_2}{P_2 - 1} \]

Figure 1. CD4 signal (m/z = 20) monitored with the RGA during a representative King and Wells experiment conducted on a CH4 surface at 20 K. P1 (pink) is the background CD4 signal, P2 (blue) is the full CD4 flux with the flag blocking the Au substrate, and P3 (orange) is the initial CD4 adsorption without the flag.

King and Wells measurements were performed at 20 K for all results presented in this study. This temperature was carefully chosen due to the methane surface interaction and the King and Wells method itself. UHV conditions at 20 K accurately model astrophysical chemistry rich environments such as dense molecular clouds. Additionally, at 20 K, multilayer CH4 is stable on a gold substrate and frozen ice films which enables measuring the condensate via RAIRS. As mentioned in He et al., and detailed in our previous work examining the initial sticking probability of CH4 on D2O ices, the liquid helium cooling of the sample manipulator could impact the pumping speed and thus the reflected portion of the beam; therefore, we take all measurements at a single sample temperature. This ensures that the unknown pumping speed remains consistent across measurements. We also calculate sticking probability by using the initial CH4 indirect flux instead of the value at saturation.

### RESULTS AND DISCUSSION

To fully understand the role that mass matching and preadsorbed hydrocarbons play in trapping dynamics for CH4 and CD4, we examined sticking probability on top of amorphous CH4 and CD4. Although the sticking probability was previously found to be independent of ice film thickness, we choose to grow films for ~80 layers to achieve self-similarity in film structure. The measured sticking probabilities for CH4 and CD4 on a CH4 substrate are shown in Figure 2. For physiosorption trapping to occur, the CH4 or CD4 molecule must lose some initial kinetic energy when impinging upon the surface. If the energy loss is not efficient enough, the impactor molecule just bounces back. As expected, the sticking probability decreases with an increase in energy as more energy must be lost in the initial condensation in order for sticking to occur. The corrugation of the gas–surface potential for CH4 and CD4 is greater on the alkane-covered surface than it is on a bare metal substrate. Although our films are thicker than one monolayer, previous rare gas and alkene studies demonstrate that sticking probabilities are enhanced by such adlayers that allow for enhanced energy accommodation. Sticking probabilities are close to unity at low incident velocities for both incident isotopologues before decaying down to 0.85 for both CH4 and CD4. There was no strong variation in sticking probabilities between the CH4 and CD4 projectiles indicating, overall, very similar energy accommodation. This suggests that both phonon creation and translational to intramolecular energy transfer are essentially the same for both CH4 and CD4 on the condensed CD4 film.

However, we note a higher sticking coefficient for CD4 on CH4 ice than for CH4 on CH4 ice particularly at high incident translational energies (Figure 3). We monitored the amount of adsorbed CH4 and CD4 via the intensity of the degenerate 4 bending mode to calculate the initial growth rate. To ensure that these measurements were taken during an essentially constant film thickness regime, the IR measurements were completed by adding no more than an additional 0.75 MLs of condensate over <4 min. As shown in Figure 4 for CH4 and CD4 beams at 4600 m/s, sticking probability differences between the CH4 and CD4 result in a larger amount of CD4 stuck on the surface after exposure and therefore a higher initial growth rate. Based on the total spectral intensity vs time and, thus, condensed projectile on the surface, we calculated the initial growth rate for each incident velocity. As a consistency check, at the end of the growth exposure, we took an additional King and Wells measurement, which matched the initial sticking probability at the beginning. Taken together, this indicates that the coverage following the

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**Figure 2.** Sticking probabilities for CH4 and CD4 on a CD4 film at 20 K. The sticking probability decreases with increasing velocity. Error bars represent the standard deviation of at least three measurements on at least three different days.
growth rate is not enough to change the underlying film structure and that RAIRS allows us to determine the amount of CH$_4$ or CD$_4$ on the surface. When overlaying this with the initial sticking probability (Figure 4), we confirm that both the King and Wells measurements and infrared spectroscopy of the condensate demonstrate an increased condensation of CD$_4$ on CH$_4$ compared to CH$_4$ on CH$_4$.

This isotopic effect increases with increasing translational energy. To understand this, we start by examining the Baule model, which predicts that a more efficient collision occurs when the gas and surface masses match due to the singularity in the momentum case. An incident molecule containing mass $m$ and energy $E$ encounters a square well potential of depth $D$ and a surface species of mass $M$, resulting in an energy transfer ($\Delta$) in the collision:

$$\Delta = \frac{4mM(E + D)}{(M + m)^2}$$

Figure 3. Sticking probabilities for CH$_4$ and CD$_4$ on a CH$_4$ film at 20 K. The sticking probability decreases with increasing velocity, but remains higher for the CD$_4$ projectile. Error bars represent the standard deviation of at least three measurements on at least three different days.

Figure 4. Representative RAIR spectra of CD$_4$ (a) and CH$_4$ (b) $\nu_4$ bending mode as a function of exposure time for the highest energy beam (4600 m/s) on a CH$_4$ surface at 20 K. Spectra were taken approximately every 25 s (c) as a function of intensity to get the initial growth rate. Differences in sticking probability result in an increased amount of CD$_4$ on the surface and thus a higher growth rate.

Figure 5. Confirmation of increased condensation of CD$_4$ on CH$_4$ in comparison with CH$_4$ on CH$_4$ as a function of incident methane velocity. Monitoring of the amount of adsorbed methane via the intensity of the $\nu_4$ bending mode for CH$_4$ and CD$_4$ by RAIRS, we calculate the initial growth rate to overlay with the sticking probability. Error bars represent the standard deviation of at least three measurements on at least three different days.
For this, we assume that the energy of the incoming molecule is much greater than the well depth of the potential. Thus, for the CH₄ film, the energy transfer for the CD₄ projectile is greater than that of the CH₄ which would generally indicate a higher sticking probability. This model does not account for the density of states of the film or the internal modes of the molecule, as discussed later. These contributions can influence sticking probabilities. Because of the role of these molecular degrees of freedom, complex and multi-phonon interactions between the surface and the incident projectiles CH₄ film and CD₄ clearly need to be taken into consideration, as they are in the MD simulations shown herein.

We performed chemical dynamics simulations using the VENUS MD computer program. Classical trajectories simulated collisions of a beam of CH₄ or CD₄ with the CH₄ or CD₄ surface at a surface temperature of 20 K. Initial conditions for the trajectories were selected to sample the experimental beam’s translational and vibrational energy. After collision the trajectories were terminated at 50 ps; CH₄ and CD₄ remaining on the surface were considered trapped. The scattered trajectories are dominated at the level of ~99% by direct scattering rather than those that trap and then desorb.

## COMPUTATIONAL DETAILS

The potential energy function for the (CH₄)beam and (CH₄)surface on top of a Au(111) crystal is given by

\[ V = V_{\text{beam}} + V_{\text{surface}} + V_{\text{beam+surface}} \]

where \( V_{\text{beam}} \) is the beam CH₄ intramolecular potential. \( V_{\text{surface}} \) is composed of intramolecular CH₄ potentials (same as \( V_{\text{beam}} \)) as well as the intermolecular CH₄---CH₄ and Au---CH₄ potentials using the 6-12 Lennard-Jones fashion. Lastly, \( V_{\text{beam+surface}} \) is the intermolecular CH₄---CH₄ potential. Each intramolecular CH₄ potential is expressed as a sum of Morse potentials for the C–H stretches and quadratic potentials for the H–C–H bends: Morse parameters are \( D = 112.5 \text{ kcal/mol}, \beta = 1.86 \text{ Å}^{-1}, \) and \( r_0 = 1.086 \text{ Å}. \) Each HCH quadratic bend has \( f = 0.585 \text{ mdyn Å}^{-1}\text{rad}^1 \) and \( \theta = 109.47^\circ. \)

The methane harmonic frequencies are 3193, 3021, 1583, and 1413 cm⁻¹.

The surface model consists of six methane layers stacked in an AB sequence on top of a layer of gold to form a cubic close-packed structure. There are 789 CH₄ molecules in alternating layers of 120/143 molecules to so that \( x \) and \( y \) are each 40 Å for an area of 800 Å for each layer. The total surface height of all the stacked layers is 18 Å, including the gold layer on the bottom. All intermolecular potentials are written as sums of Lennard-Jones two-body potentials with a cutoff distance of 10 Å and are summarized in Table 1. For the Au (111) base, \( \epsilon_0 = 5.29 \text{ kcal/mol} \) and \( \sigma_0 = 2.951 \text{ Å} \) were used to give an atomic spacing of 2.93 Å, closely matching that determined from STM images of the reconstructed (111) surface. Our surface contains CH₄ spaced by 3.8 Å, which is comparable to calculated CH₄ intermolecular potentials. CH₄---CH₄ intermolecular potentials among all methane molecules (including those in different layers) are written as sums of 6-12 Lennard-Jones two-body potentials and include interactions between carbons and hydrogens. To calculate the Au---CH₄ interaction, we employ standard mixing rules and assume a geometric mean between C and Au to get a \( \epsilon_0 = 0.7336 \text{ kcal/mol} \) and \( \sigma_0 = 2.99 \text{ Å}. \)

Geometry optimization of the surface occurred prior to trajectory simulations to obtain a potential energy minima configuration. Additionally, we note that this is a flat crystalline surface, which is a model representation of a local section in the experimental surface topology which in reality may contain domains of small, imperfect crystallites. However, even with this difference, there is qualitatively similar energy-transfer dynamics and thus is appropriate to use for our study.

A microcanonical ensemble averaged intermolecular potential curve for CH₄ approaching to the surface is obtained by averaging the potential energies of randomly oriented CH₄ as a function of CH₄-surface center-of-mass separation parallel to the surface. Such a potential energy minimum is ~0.07 eV at a center-of-mass separation of 4.25 Å.

### Procedure for the Chemical Dynamics Simulations

Chemical dynamics simulations were performed by using the VENUS general chemical dynamics computer program. Classical trajectories were used to simulate collisions of a beam of CH₄ or CD₄ with the CH₄ or CD₄ surface. Simulations at each collision energy were performed by using a surface temperature of 20 K. Initial conditions for the trajectories were selected to sample the beam’s translational and vibrational energy at the experimental conditions. The selection of initial conditions follows from previous VENUS studies. For each simulation, a beam of colliding molecules was aimed within a circular area. Each trajectory was initialized with a separation of 10 Å between the center of the beam and surface aiming point. For each beam, the initial vibrational quantum states were sampled from Boltzmann distributions at 300, 700, 900, or 1100 K, and the translational energies were determined from the molecular beam velocity distributions (Figure 2). Using the experimental velocities, the CH₄ translational energies were 0.49, 1.16, 1.48, and 1.79 eV and the CD₄ translation energies were 0.67, 1.41, 1.74, and 2.19 eV. Zero-point energy was included in these samplings, and the rotational energy was set to 0 K to match the experimental supersonic molecular beam conditions.

For each trajectory, the gold and bottom three layers were held rigid and acted as anchor layers. Additionally, the mass of carbon atoms in rim CH₄ molecules was artificially increased by 10000 to truncate the surface. Initial conditions for this surface were selected by assigning velocities to the carbon atoms of these layers, sampled from a Maxwell–Boltzmann distribution at 20 K. The surface was equilibrated by a 50 ps molecular dynamics simulation with velocity scaling every 1000 steps and another equilibration without velocity scaling. The trajectories were propagated with a Velocity-Verlet integrator, with a time step of 0.01 fs. Trajectories were terminated when either the distance between the central methane molecule and outgoing product exceeds 30 Å or the total integration exceeds 50 ps. Typically, 750–2000 trajectories were calculated for each ensemble of initial conditions including the surface composition and beam conditions.

### Simulation Results

Overall, we find that there is nice agreement between the chemical trajectory simulation results and the experimentally determined sticking probabilities. The
VENUS calculations demonstrate a decrease in sticking probability with increasing incident velocity as well as a difference between CH₄ and CD₄ on a CH₄ surface (Figure 6). Additionally, for the simulated collisions on a CD₄ surface (Figure 7), there is no difference in sticking probability, again in agreement with our experimental results. In a more careful comparison to the results shown in Figure 3, the theoretical sticking probability for the CD₄ on the CH₄ surface is slightly lower than the experimental value. This could arise from various effects; for example, Lennard-Jones potentials are not optimized for the repulsive region.

Full details of the energy transfer and chemical dynamics simulations will be discussed in a forthcoming paper to provide a molecular-level understanding of the mechanisms occurring between the methane projectile and the methane surface. When examining phonon dispersion curves for CH₄ and CD₄, not only are the CD₄ phonon modes at a lower energy, but there is enhanced translational–rotational coupling. In addition to this coupling, local corrugation of the surface can also influence trajectory paths and therefore energy flow. Full analysis of our molecular dynamics studies will provide necessary insight into lattice vibrations and how energy is efficiently dissipated to trap the methane isotopologues.

To further explore and confirm our experimental results, we consider a beam composed of both CH₄ and CD₄ in a 3:1 ratio; this ratio was not selected to optimize condensation differences, but rather to demonstrate the robust nature of differential sticking. This allows us to quantify the sticking probability as well as condensate composition. While dosing a multilayer film of both CH₄ and CD₄ at 20 K, the integrated area of the degenerate ν₄ mode was tracked over time using RAIRS. Once the condensate reached a self-similar structural steady state of at least 100 layers, at least 10 spectra per experiment on at least three different days were averaged to determine the film composition. As depicted in Figure 8, the condensate composition for the room temperature beam (2200 m/s) is 74.5% CH₄ and 25.5% CD₄. However, as the beam velocities increase, the heavier isotope (CD₄) becomes preferentially adsorbed into the film. Because of increased adsorption into the film, the condensate film structure changed to 73.7% CH₄ and 23.6% CD₄. Overall, by measuring the condensate with RAIRS, we confirm that due to the increased sticking probability of CD₄ on a CH₄ film, we see an increased affinity for CD₄. We demonstrate for our fastest beam (4400 m/s) that there is a 3.12 ± 0.06% enrichment of CD₄ compared to the room temperature beam (2200 m/s). When taking the individual King and Wells values (Figures 2 and 3) and combining that with the film composition determined from the RAIR spectra, we calculate the sticking probabilities for CH₄ and CD₄ on the mixed film. For the highest velocity beam, these sticking probabilities also result in a CD₄ enrichment of 3.9 ± 0.02%, indicating excellent agreement with the observed condensate enrichment.

Figure 6. Sticking probabilities calculated from the number of CH₄ and CD₄ direct and physisorption scattering trajectories on a CH₄ layered surface at 20 K. Error bars represent the standard error of at least 750 trajectories for each velocity.

Figure 7. Sticking probabilities calculated from the number of CH₄ and CD₄ direct and physisorption scattering trajectories on a CD₄ layered surface at 20 K. Error bars represent the standard error of at least 750 trajectories for velocity.

Figure 8. Integrated area of the ν₄ mode for CD₄ and CH₄ demonstrates an enrichment of the heavier isotope (CD₄) into the mixed condensed film at higher beam velocities. Error bars represent the standard deviation of at least 35 steady-state films on at least three different days.
CONCLUSION

We examined the differential sticking probability of CH₄ and CD₄ on CH₄ and CD₄ ices using RAIRS for measuring on-surface gaseous condensation and complementary King and Wells mass spectrometry techniques for monitoring sticking probabilities. We found that as the incident velocity of CH₄ and CD₄ increases (up to 4600 m/s), the sticking probability decreases for both films. Interestingly, we conclude that preferential sticking and condensation occurs for CD₄ when striking the surface in comparison to the outcome for CH₄. This observation was confirmed both experimentally from infrared spectroscopy of the condensation and via mass spectrometric detection of the reflected molecules and theoretically from the gas–surface chemical trajectory simulations. This theoretical model system will be explored in more detail to provide insight into energy transfer and lattice vibrations. Next, we employed a mixed incident supersonic beam composed of both CH₄ and CD₄ in a 3:1 ratio to measure the condensate as well as the sticking probability. When doing so, we see the same effect in the condensed mixed film, supporting an isotopic enrichment of the heavier isotope. Because the Baule model does not accurately represent this condensed phase system due to its molecular complexity, we propose that enhanced multi-phonon interactions attributable to the film’s phonon and rovibrational densities of states and inelastic cross sections including intermolecular energy exchange between the incident CD₄ projectile and the CH₄ film allow for more efficacious gas–surface energy transfer.

In general, these results indicate the importance of understanding gas–surface energy exchange under non-equilibrium conditions at cold substrate temperatures and have important astrophysical and terrestrial implications. Our work demonstrates the importance of film structure and surface lattice coupling to allow for efficient energy transfer and an isotopic enrichment of the heavier isotope (CD₄). The insights gained from gaseous condensation under nonequilibrium conditions are also important for understanding aircraft flight in low-temperature environments. These results also offer a new route for isotopic enrichment via the preferential condensation of heavier isotopes and isotopologues during gas–surface collisions under carefully selected substrate, gas-mixture, and incident velocity conditions.

Importantly, our experiments are conducted at low-temperature astrophysical conditions. By experimentally determining initial sticking probability differences between methane and its heavier isotopologue as a function of incident energy, we find that the film composition is important, especially for high-energy projectiles bombarding icy dust grains. Because adsorption is often a first step for many cold temperature reactions occurring on these grains, differences in sticking probabilities have notable implications for allowed reaction probabilities and follow-on events leading to increased molecular complexity. Our work, therefore, not only can explain increased abundance of deuterium in solar system planets but also can be incorporated into astrochemical models of the icy dust grain processes including those in the interstellar region.

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Notes

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