Resonant infrared laser-induced desorption of methane condensed on NaCl(100): Isotope mixture experiments

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Resonantly enhanced infrared laser-induced desorption of methane condensed on a single-crystal NaCl(100) surface is observed after excitation with the widely tunable infrared laser output of the free-electron laser at the free-electron laser for infrared experiments facility using mass spectroscopic detection and time-of-flight analysis. Desorption of methane is observed only when the exciting light is in resonance with an internal vibrational mode of the molecule. Different intramolecular modes of the three methane isotopologues under study—CH4, CD4, and CD3H—are excited; the degenerate deformation mode v3 is observed for CH4 and CD4 at 7.69 and 10.11 μm, respectively, as well as the v2 and v4 modes of CD3H at 7.79, 9.75, and 9.98 μm. The desorption signals for the pure layers of these different methane isotopologues as well as for different mixtures of two of these are investigated as a function of the infrared wavelength and the laser fluence. The desorption behavior for pure and mixed layers is compared and the underlying desorption mechanism is discussed. © 2006 American Institute of Physics. [DOI: 10.1063/1.2159487]

INTRODUCTION

One of the goals of the studies on vibrational modes of adsorbates or condensates on surfaces is to gain a better understanding of their role in energy transfer and dissipation processes, as well as in chemical reactions on surfaces. In this field it has been shown that surface processes such as desorption from insulating or metal surfaces can be induced by excitation of internal vibrations of adsorbed molecules in the electronic ground state.1–8 For efficient resonant excitation of vibrational modes, a pulsed tunable infrared laser source has to be employed. Recently it has been shown that a free-electron laser (FEL) operating in the infrared is a well-suited radiation source for this kind of investigations.9–11 Theoretical studies of desorption by laser-adsorbate vibrational coupling and energy transfer have been carried out,12–17 particularly for the well-investigated physisorption system CO/NaCl(100).18,19

In the present work the infrared laser-induced desorption of different methane isotopologues condensed on a NaCl(100) single-crystal surface is investigated using the FEL of the free-electron laser for infrared experiments (FELIX) facility in the Netherlands as the excitation source. The present study is beginning to explore the field of mode and isotope selectivity after resonant infrared excitation of vibrational modes in condensed layer systems on dielectric surfaces. Considering different isotopologues the question arises whether the different IR-active modes such as bending and stretching modes can lead to a significantly different desorption behavior. The density of states accessible for further excitation is increased for larger systems after resonant excitation and one can ask how this influences the desorption process, e.g., due to coupling of intra- and intermolecular modes. Besides the mode selectivity also the isotope selectivity sheds light on the underlying mechanism of the desorption process. Isotope selectivity is only expected in cases of resonant excitation followed by a direct desorption step or when energy scrambling between different adsorbates is prohibited. The former is possible when the energy deposited in a specific mode exceeds the binding energy of the molecule within the layer or to the substrate and rupture of this bond occurs. In case no selectivity is observed for the desorption process, the desorption involves either a resonant heating mechanism leading to thermal desorption, a V-V transfer process among the species, followed by vibrational energy transfer from the internal vibrational mode to the desorption coordinate of the adsorbed molecule, or a collisional interaction within the adsorbate in dense systems. In general, the dynamics of the desorption process will depend on the coupling and the energy flow between the different degrees of freedom.

EXPERIMENT

The experiments are performed in a standard ultrahigh-vacuum (UHV) chamber using the infrared light output of the FEL of the FELIX facility at our Institute.20 The infrared radiation produced by the FEL is continuously tunable between 5 and 250 μm, and the wavelength ranges between 3 and 5 μm can be accessed by operation on the third har-
monic. For the desorption experiments presented here, the wavelength regimes from 7 to 12 $\mu$m and from 3.25 to 3.4 $\mu$m are used. The infrared light is pulsed and consists of so-called macropulses that are on the order of 5 $\mu$s long and that having a repetition rate of 5 Hz. A macropulse contains a series of micropulses that are equally spaced by 1, 20, or 40 ns. The experiments presented are performed in the 1 GHz operation mode corresponding to a micropulse spacing of 1 ns. The micropulse duration depends on the laser settings and is adjustable in duration between a few hundred femtoseconds and several picoseconds. For the desorption experiments the spectral bandwidth, which is Fourier transform limited, is an important parameter. For these measurements, it is kept constant to about 0.5% full width at half maximum (FWHM) of the central frequency of the laser light or to about 4–7 cm$^{-1}$, which corresponds to a pulse duration of about 2–3 ps. Macropulse energies can reach up to 100 mJ, but typical pulse energies at this wavelength range and bandwidth are on the order of 20 mJ at the experiment. Correspondingly, the energy in a micropulse is typically of the order of 4 $\mu$J. Values reported for the laser energy are measured directly in front of the KBr entrance window to the UHV chamber. The laser fluence is varied either using attenuators or by changing the size of the laser spot on the surface with a KBr lens ($f=470$ mm). The angle of incidence for the $p$-polarized laser light onto the surface is 45° with respect to the surface normal.

The NaCl(100) single crystal ($20 \times 20 \times 3$ mm$^3$, purity of 99.99%) is cleaved $ex$ $situ$ under dry nitrogen atmosphere and is immediately transferred to the vacuum chamber. It is mounted to a copper sample holder, which is connected to a manipulator constructed for low-temperature application. The sample can be cooled to 25 K using liquid helium. Between the experiments, the sample is maintained at 450 K to avoid contamination, especially adsorption of water onto the surface. The sample temperature is controlled using two thermocouple pairs (NiCr–Ni and AuFe–Cr) attached to the sample holder. The sample can be manipulated in $x$, $y$, and $z$ directions, rotated polarly by 360°, and tilted azimuthally, which enables precise positioning with respect to the laser beam.

The cryostat and manipulator are mounted in a standard UHV chamber operated at a base pressure of $2 \times 10^{-10}$ mbar. Both the entrance and the exit windows are made of KBr to minimize the influence of light scattered inside the chamber. The UHV apparatus is equipped with a sensitive, differentially pumped quadrupole mass spectrometer (Extrel Q50). The neutral species desorbing from the surface are detected mass selectively using electron-impact ionization and a channeltron detector. The direction of the detection is normal to the surface plane. The distance between the surface and the electron-impact ionizer is 16 cm. The mass-selected signal is recorded as a function of time after the laser pulse.

The condensed layers of the different methane isotopes (CH$_4$ from Messer Griesheim 99.995%, CD$_3$H and CD$_4$ from Cambridge Isotope Laboratories Inc. D-98% and D-99%, respectively) are prepared at low temperature under nonequilibrium conditions via background dosing using a leak valve. The adsorption temperature is chosen low enough to ensure that on the time scale of the experiment no significant thermal desorption occurs. The gas inlet is made of stainless steel and operates at a base pressure of $1 \times 10^{-6}$ mbar; no further purification of the gas is carried out. Typically a few hundred monolayers (ML) are condensed using gas pressures on the order of $1 \times 10^{-5}$ mbar. At these coverages a fairly homogeneous irradiation of the layer is still ensured. For a 500-ML-thick methane film on resonance, about 55% of the IR radiation is transmitted. The coverage is estimated using a sticking coefficient of 1. In addition, a Fourier transform infrared (FTIR) spectrometer (Bruker IFS66v) is attached to the UHV chamber, which allows to record FTIR spectra in transmission geometry with a liquid-nitrogen-cooled mercury-cadmium-telluride (MCT) detector. Analysis of the integrated absorption enables a cross-check of the coverage.

**RESULTS**

The feasibility to study infrared laser-induced desorption of small molecules from insulator surfaces using the FEL at the FELIX facility as an infrared radiation source was shown for CD$_3$F and N$_2$O condensed on NaCl(100). In the present investigation, we report results on desorption experiments of methane condensed on NaCl(100). Experiments are performed with three methane isotopologues—CH$_4$, CD$_4$, and CD$_3$H—as well as with mixed layers of these isotopes.

**Pure isotopes**

At a temperature of 25 K, layers of CH$_4$, CD$_4$, or CD$_3$H with a thickness of about 500 ML are condensed onto a NaCl(100) single-crystal surface. The mass-selected ion signal for the respective isotope is recorded as a function of the FELIX wavelength in the range between 7 and 12 $\mu$m. For the pure isotopes the primary peak is used for mass-selective detection (CH$_4$: 16 amu, CD$_4$: 20 amu, and CD$_3$H: 19 amu). The mass spectra of the desorbing species show a fragmentation pattern similar to the ones known for gas-phase molecules, indicating desorption of neutral and intact molecules. Figure 1(a) shows the desorption spectra of the pure isotope systems for an IR fluence between 1.5 and 2.2 J/cm$^2$ per macropulse. At these fluences between a fraction and a few monolayers are desorbed per macropulse. For each isotope desorption is observed only at specific infrared wavelengths: CH$_4$ at 7.69 $\mu$m (1300 cm$^{-1}$), CD$_4$ at 10.11 $\mu$m (989 cm$^{-1}$), and CD$_3$H at 7.79 (1284 cm$^{-1}$), 9.75 (1026 cm$^{-1}$), and 9.98 $\mu$m (1002 cm$^{-1}$). These wavelengths show that resonant excitation of the first internal vibrational modes in the molecules occurs. In the spectral range studied here there is no detectable desorption signal at frequencies other than the ones mentioned, even at the highest laser fluences applied. The corresponding linear absorption spectra recorded with a FTIR spectrometer are shown in Fig. 1(b) for comparison. The spectra are taken under comparable experimental conditions at a sample temperature of 25 K and for a coverage of 1000 ML. The spectral resolution is 0.5 cm$^{-1}$. The absorption spectra are obtained by division of a sample spectrum by a reference spectrum, which was recorded for the uncovered
surface at the measurement temperature of 25 K. The spectra exhibit the absorption bands of the condensates CH$_4$, CD$_4$, and CD$_3$H, which are labeled according to convention.$^{21}$

Comparing the desorption spectra to the linear absorption spectra, a clear correspondence is found. Frequencies and bandwidths of the peaks are observed to be identical in both sets of spectra within the experimental error. The only exception is the $\nu_4$ mode of CD$_3$H, which splits in both cases into two components, but the observed splitting is slightly different. In the desorption case the modes are observed at 9.75 $\mu$m (1026 cm$^{-1}$) and 9.98 $\mu$m (1002 cm$^{-1}$), while for the linear absorption spectrum the peak maxima are found at 9.73 $\mu$m (1028 cm$^{-1}$) and 10.02 $\mu$m (998 cm$^{-1}$). A systematic error is unlikely since the frequency shift is opposite for the two components. However, since the spectra are not recorded for the same layer and coverage, small differences in the preparation procedure might cause this change in splitting. Nevertheless, it can be concluded that desorption is observed only if the infrared laser excitation is in resonance with an internal vibrational mode of the methane isotope. Nonresonant desorption is not observed.

For CH$_4$ and CD$_4$ the relative intensities compare well for absorption and desorption. In linear absorption the integrated intensities of the $\nu_4$ bands yield ratios of 0.97:1.00 for CH$_4$:CD$_4$ [Fig. 1(b)]. In desorption the symmetric isotopes yield ratios of 0.99:1.00. The situation is very different for the asymmetric isotope CD$_3$H. The integrated intensity of the splitted $\nu_4$ band amounts in desorption to only 9% of that of CD$_4$ while in absorption a relative intensity of 82% is measured. From the data one could conclude that the two components of the $\nu_4$ band show different efficacies for desorption, but as discussed in more detail later on the effect can mainly be attributed to the way of data acquisition. Finally, the widths of the spectral features may be mentioned. For CH$_4$ and CD$_4$ again similar results are obtained in desorption and linear absorption with respective values of 8.5 and 7.2 cm$^{-1}$ for CH$_4$ and correspondingly of 6.8 and 5.3 cm$^{-1}$ for CD$_4$. CD$_3$H behaves now similar to the other isotopes. In desorption the spectral widths are 6.6 cm$^{-1}$ for the $\nu_4$ band and 12.6 and 10.0 cm$^{-1}$ for the high- and low-frequency components of the $\nu_4$ band. In absorption the corresponding values are 9.2, 10.5, and 6.0 cm$^{-1}$, respectively.

Figure 2 shows the dependence of the desorption signal on the applied laser fluence after resonant excitation of the $\nu_4$ mode of CH$_4$, CD$_4$, and CD$_3$H ($\lambda = 7.69$, 10.11, and 9.75 $\mu$m), respectively. Day-to-day differences in the operating conditions of the FEL cause the variability in the laser fluences applied to study the desorption yield of the individual isotopes. The overall dependence of the desorption signal for this mode is similar for the different isotopes. For CH$_4$ and CD$_4$ a measurable desorption yield can be observed only at fluences above 0.5 J/cm$^2$, while for CD$_3$H signifi-

![Desorption and Absorption Spectra](image)
significantly higher laser fluences of more than 1.5 J/cm² have to be applied. For all three isotopes the desorption signal shows a slightly nonlinear dependence on the fluence.

The time-of-flight (TOF) spectra of the different isotopes excited at the resonance frequency of the ν₄ mode are shown in Fig. 3. The resulting velocity distribution has therefore been fitted with a modified Maxwell distribution of the form

\[ f(v) = \frac{a}{\pi} \frac{d}{v^3} \exp \left( -\frac{d}{2kT} (\frac{d}{v} - u)^2 \right), \]

where \( d \) denotes the distance from the surface to the detector, \( u \) is the stream velocity, and \( a \) is a normalization factor. For clarity the results of these fits to the TOF distributions are not included in the figure. The fits show that the leading slope, the maximum of the flight-time distribution, and the beginning of the falling tail can be well described with a thermal velocity distribution; due to insufficient differential pumping capabilities in the setup, the signal decreases slower than expected for a Maxwellian distribution for later times. The fits yield the maxima of the TOF distributions at the times indicated in Fig. 3. These times scale with the square root of the mass as expected for the most probable velocity resulting from a Maxwell-Boltzmann distribution. The resulting translational temperatures are on the order of 40–50 K, i.e., well above the surface temperature of 25 K.

**Randomly mixed isotopes**

In the following, experiments performed on different randomly mixed (50:50) multilayers of the three isotopes are described. During the series of measurements the experimental parameters such as thickness (300 ML), temperature (25 K), and the parameters of the FEL such as bandwidth and intensity are kept constant as much as possible. In the case of CH₄ and CD₄ both molecules exhibit only one infrared-active mode in the spectral range under investigation, the ν₄ vibrations, which are spectrally well separated from each other by about 300 cm⁻¹. The molecules are detected on mass 15 for CH₄ and on mass 20 for CD₄.

The desorption yields are shown in Fig. 4(a) for the detection of CH₄ and in Fig. 4(b) for the detection of CD₄ at three different laser fluences, as indicated in the inset of Fig. 4. The laser fluence for the longer wavelength is higher by about 50% and the laser linewidth at those frequencies is slightly larger. Also for the mixed layers a desorption signal was observed only at the resonance frequencies independent of the exciting laser fluence. The observed frequencies are almost unchanged as compared to the pure systems. It is

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**FIG. 2.** Dependence of the desorption signal on the laser fluence for multilayers of CH₄, CD₄, and CD₃H condensed on NaCl(100) after resonant excitation of the ν₂ mode at 7.69, 10.11, and 9.75 μm, respectively.

**FIG. 3.** Time-of-flight spectra for the desorption of CH₄, CD₄, and CD₃H recorded at the resonance frequencies of the ν₄ mode. The maxima of the flight-time distribution are indicated for each time-of-flight spectrum in the figure. The spectra are offset for clarity.

**FIG. 4.** Desorption spectra of isotopically mixed layers (CH₄:CD₄ = 50:50) resonantly excited via the ν₄ bending modes of the isotopes at 7.69 and 10.11 μm. The desorption yield for CH₄ (a) is monitored on mass 15 and for CD₄ (b) on mass 20 for different laser energies. Layer thickness and temperature are kept comparable during the experiments; the experimental parameters of FELIX are indicated.
evident that the excitation of the internal deformation vibration of one isotope also causes the other isotope to desorb. With increasing laser fluences a broadening of the desorption bands is observed.

Both molecules show a similar but nontrivial dependence of the desorption yield on the laser fluence. At low intensity the efficacy of the CD$_4$ $\nu_2$ vibration of desorbing either CD$_4$ or CH$_4$ seems to be higher than the corresponding CH$_4$ vibration despite its lower energy content (989 cm$^{-1}$ vs 1300 cm$^{-1}$). This behavior is reversed at high laser intensities. Moreover, excitation of the CD$_4$ vibration results in a nearly linear increase of the desorption yield with laser fluence irrespective of the desorbing molecule. On the other hand, excitation of the CH$_4$ vibration clearly shows a non-linear increase of the desorption yield, again irrespective of the desorbing molecule. Finally, it can be noted that the yield of CD$_4$ is significantly higher—by a factor of 2—than for CH$_4$ when the system is excited on the CH$_4$ vibration. Looking carefully at all data it is also observed that the maxima of the desorption peaks shift with increasing laser fluence to shorter wavelength and that the peaks become asymmetric.

In another set of experiments isotope mixtures consisting of CH$_4$ and CD$_3$H as well as mixtures of CD$_4$ and CD$_2$H are investigated. These two systems are different from the mixture of CH$_4$ and CD$_4$ discussed before, because the absorption bands are not well separated anymore. For the mixture containing CH$_4$ they are almost overlapping around 7.7–7.8 $\mu$m and for the one with CD$_4$ around 10.0–10.1 $\mu$m. In Fig. 5 the desorption spectra are shown for different laser fluences for the mixture CH$_4$:CD$_3$H =50:50 (a) detected on mass 15 for CH$_4$ and (b) detected on mass 19 for CD$_3$H. The mass channels for detection are selected such that the contribution from the other isotope is not existing or minimal. Also for this mixed system it is found that desorption of both isotopes occurs only after resonant excitation of one of the isotopes. For the CH$_4$ line again a blueshifted broadening is observed at high laser intensities. The two spectra measured do not depend on the detection channel. The efficacy of the $\nu_2$ vibration of CD$_4$H is low, as can be expected from the pure isotopic spectrum. More striking is the complete absence of the $\nu_2$ vibration of CD$_3$H, for desorption of CH$_4$ as well as for CD$_2$H molecules, although the band is spectrally well separated (by 17 cm$^{-1}$) from the CH$_4$ line.

Figure 6 shows corresponding studies for the mixture CD$_4$:CD$_3$H =50:50 (a) detected on mass 16 for CD$_4$ and (b) detected on mass 19 for CD$_3$H. Mass channel 16 instead of 20 for the detection of CD$_4$ was an unfortunate choice. However, the only consequence is that the detection is less efficient by more than one order of magnitude, because mass 16 is a minor fragmentation channel of CD$_4$; the data in Fig. 6 are corrected for this difference. Now the $\nu_2$ vibration of CD$_3$H is again active and is compared to the $\nu_2$ vibration with similar efficacy as in the pure isotope case. In all spectra shown in Figs. 4–6 the desorption yield shows a clear non-linear dependence on laser-pulse energy, as is also expected from the results shown in Fig. 2.

Layered isotopes

In a last set of measurements in experiment (1) 300 ML CD$_4$ was condensed on top of 300 ML CH$_4$ and in experiment (2) the order is reversed and 300 ML CH$_4$ was condensed on top of 300 ML CD$_4$. Figure 7(a) shows the corresponding FTIR spectra for experiments (1) and (2) for the first and second adsorption step in the spectral range of the $\nu_3$ and $\nu_4$ modes. This time, for the laser-induced desorption experiments, the FEL was operated on the third harmonic to enable excitation of the $\nu_4$ vibration of CH$_4$ at 3.33 $\mu$m (3003 cm$^{-1}$). In this mode of operation the tuning range is very restricted and it was only possible to excite the $\nu_4$ vibration of CH$_4$. The results are shown in Fig. 7(b). In experiment (1), where the CH$_4$ layer was covered by a CD$_4$ layer, desorption of CH$_4$ as well as of CD$_4$ was observed after excitation of CH$_4$. In contrast, in the case of experiment (2) where the CH$_4$ layer was condensed on top of the CD$_4$ layer, only desorption of CH$_4$ was found.

DISCUSSION

The desorption spectra recorded for CH$_4$, CD$_4$, and CD$_3$H condensed on NaCl(100) show desorption peaks only when the infrared wavelength is in resonance with an internal vibrational mode of the molecules. Therefore, it can be unambiguously concluded that the first step leading to the
Desorption process is resonant in character. A comparison of the desorption spectra to the linear absorption spectra shows identical behavior for the systems CH$_4$ and CD$_4$ concerning frequencies, linewidths, and relative and absolute intensities, suggesting therefore for both systems a very similar energy-flow and desorption processes after resonant excitation of the $\nu_4$ mode.

In contrast, the desorption spectra of CD$_3$H show clear differences: The integrated desorption yield of the $\nu_4$ band is almost an order of magnitude lower than the corresponding bands in CH$_4$ and CD$_4$. The two peaks of the $\nu_4$ band show differences in the desorption efficacies as well as in their spectral separation when compared to the linear absorption spectrum. These observations are most likely explained by a desorption process requiring several photons for desorption. Given the fact that there is already, in the linear absorption spectrum, a considerable intensity difference of the two $\nu_4$ bands, this can easily lead to a difference of an order of magnitude for the nonlinear case. While the slight differences in the splitting of the $\nu_4$ band for the linear and the desorption case can be readily explained with small differences in the layer, the different efficacies of the two peaks have a different origin. The scanning programs for the FEL permitted only to scan from shorter to longer wavelength and not reversed. This might, for thicker layers and at high intensities, lead to a situation where on the blue side of the band a significant fraction of the layer is desorbed before the red part of the band is reached. Since at higher fluences the bands appear to be broadened, the increased desorption on the blue wing appears like a blueshift of the peak and can also introduce an asymmetry in the line shape. However, besides this technical reason, additional contributions can also be discussed as, e.g., a different energy-loss mechanism not operative in the highly symmetric molecules or a different desorption mechanism for the CD$_3$H molecule.
separated but almost overlapping. In those cases it is observed that the weaker components in the overlapping regime, i.e., the $v_2$ mode of CD$_3$H in the mixture with CH$_4$ and the lower-frequency component of the $v_3$ mode of CD$_4$ in the mixture with CD$_4$, are not detectable as compared to the pure layer. The explanation for this observation is not readily clear. It is known from linear absorption spectroscopy that within mixed layers, intensity transfer can occur from the lower- to the higher-frequency mode due to shielding effects. Such an effect would result in a reduction of the absorption coefficient and therefore it may be that the energy stored in the respective vibrational mode is not sufficient anymore to overcome the threshold value for desorption. To clarify this point unambiguously more detailed investigations are required.

The results observed for the layered methane isotopes yield yet another insight in the desorption mechanism. In experiment (1) the lower layer is resonantly excited and species from both layers are observed in the desorption flux. On the other hand when just the upper layer is excited, as in experiment (2), only molecules of this layer can be detected in the desorption flux. Thus the energy deposited in the layer via the resonant excitation remains locally confined.

Energy does not just flow to the top most layer causing these molecules then to desorb, because in that case one would observe only CD$_4$ molecules in experiment (1). Rather, in the laser-excited area the molecular constituents acquire so much energy in the translational coordinate that they boil off from the solid. On their way out these molecules collide with other, not excited molecules and transfer in these collisions enough energy to cause the desorption of also these primarily nonexcited species. This mechanism can also explain the lack of isotopic selectivity in the randomly

![Graphs showing FTIR spectra and desorption signals for different experiments.](image-url)
mixed layers. Also the equilibrated kinetic energies of different isotopes can be established by such collisions.

CONCLUSIONS

Infrared laser-induced desorption of multilayers of different methane isotopes and different mixtures of these isotopes condensed on NaCl(100) has been investigated. Desorption is observed only if the infrared excitation wavelength produced by the FEL is in resonance with one of the vibrational modes of the systems under study. Investigations of the fluence dependence and the time-of-flight spectra of the pure layers of CH₄, CD₄, and CD₃H favor a resonant heating mechanism over a direct desorption process. This finding is confirmed by the studies on mixed isotope layers of CH₄ and CD₄, CH₄ and CD₃H, as well as CD₄ and CD₃H. In these systems desorption of both components is observed after resonant excitation of one of the isotopes in the layer, which cannot be explained by a direct desorption mechanism as this would lead to a selectivity in the desorption product. In addition, the dependence of the laser fluence for the mixed layers can be explained by the existence of a threshold value for the laser energy required to introduce desorption, in agreement with a resonant heating process. The energy is, however, converted locally into kinetic energy of the molecular constituents. These hot molecules then collide with neighboring unexcited molecules causing also their desorption. This in turn causes the isotopic scrambling in randomly mixed layers.

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