Hydrogenation at low temperatures does not always lead to saturation: the case of HNCO

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ABSTRACT

Context. It is generally agreed that hydrogenation reactions dominate chemistry on grain surfaces in cold, dense molecular cores, saturating the molecules present in ice mantles.

Aims. We present a study of the low temperature reactivity of solid phase isocyanic acid (HNCO) with hydrogen atoms, with the aim of elucidating its reaction network.

Methods. Fourier transform infrared spectroscopy and mass spectrometry were employed to follow the evolution of pure HNCO ice during bombardment with H atoms. Both multilayer and monolayer regimes were investigated.

Results. The hydrogenation of HNCO does not produce detectable amounts of formamide (NH₂CHO) as the major product. Experiments using deuterium reveal that deuteration of solid HNCO occurs rapidly, probably via cyclic reaction paths regenerating HNCO. Chemical desorption during these reaction cycles leads to loss of HNCO from the surface.

Conclusions. It is unlikely that significant quantities of NH₂CHO form from HNCO. In dense regions, however, deuteration of HNCO will occur. HNCO and DNCO will be introduced into the gas phase, even at low temperatures, as a result of chemical desorption.

Key words. astrochemistry – ISM: molecules – molecular processes – molecular data

1. Introduction

Isocyanic acid, HNCO, is the simplest molecule containing the four most abundant elements: hydrogen, carbon, nitrogen, and oxygen. It is an important interstellar molecule, with gas phase abundances of around $10^{-9}$ to $10^{-8}$ with respect to H₂ in molecular clouds, where it is believed to trace dense, cold gas (Jackson et al. 1984). Since its first detection in the Sgr B molecular cloud complex (Snyder & Buhl 1972), HNCO has been detected in multiple environments, including hot cores (e.g. Helmich & van Dishoeck 1997), high mass young stellar objects (Bischop et al. 2007a), molecular outflows (Rodríguez-Fernández et al. 2010), comets (Lis et al. 1997), and other galaxies (Nguyen-Q-Rieu et al. 1991). It has been shown that HNCO is also a tracer of warm gas, and that its formation origins are likely to be predominantly in icy grain mantles (Bischop et al. 2007a).

Isocyanic acid has not yet been observed in the solid state, but it is believed to be responsible for the formation of the abundant cyanate ion OCN⁻ (e.g. Soifer et al. 1979; Demyk et al. 1998; Lowenthal et al. 2002; van Broekhuizen et al. 2005) via reaction of the acidic HNCO with bases such as NH₃ (Raunier et al. 2003a; van Broekhuizen et al. 2004; Mispelaer et al. 2012) and H₂O (Raunier et al. 2003b; Theule et al. 2011), or by the irradiation of ices with ultraviolet photons (Lacy et al. 1984) or protons (Moore et al. 1983). Moreover, its solid phase chemical reactions have been shown to give rise to isomerisation, with cyanic acid, HOCCN, formed thermally in mixtures of HNCO and H₂O (Theule et al. 2011).

The chemical network surrounding HNCO has not been fully studied experimentally. Early theoretical models assumed that HNCO formed only in the gas phase (e.g. Iglesias 1977), but more recent studies contend that HNCO forms on grain surfaces via the thermal reaction NH + CO or by the hydrogenation of OCN⁻ (e.g. Garrod et al. 2008; Tideswell et al. 2010). Its presence in the gas phase is explained by the subsequent desorption of HNCO from grains, or by the destruction of larger species such as urea, (NH₂)₂CO. Abundances of HNCO are enhanced in shocked regions, and thus it is likely that sputtering or additional gas phase formation routes are active in post-shock gases (Zinchenko et al. 2000).

The abundance of solid HNCO is predicted to remain relatively low, at $\sim 10^{-4}$ with respect to H₂O, and thus is not detectable in infrared spectra. If the dominant formation route of HNCO (NH + CO) were efficient (Garrod et al. 2008), destruction routes would be required to explain this. As mentioned above, the formation of OCN⁻ by reaction of HNCO with H₂O and NH₃ are favourable routes, the reactions having activation barriers of 26 ± 2 kJ mol⁻¹ (3130 ± 240 K; Theule et al. 2011) and 0.4 ± 0.1 kJ mol⁻¹ (48 ± 12 K; Mispelaer et al. 2012), respectively. Irradiation of HNCO with vacuum ultraviolet (VUV) radiation has been shown to produce formaldehyde (H₂CO), formamide (NH₂CHO), and urea (H₂NCONH₂) (Raunier et al. 2004).

In dense molecular clouds the secondary photon field is weak and hydrogen atoms have a long residence time on grain surfaces because of the low temperature (Tielens & Hagen 1982; Amiaud et al. 2007). Hydrogen atoms are mobile on the surface at 10 K,
and thus hydrogenation reactions dominate low temperature ice chemistry. Experimentally, there still remain many hydrogenation reactions to characterise, although studies have been carried out, particularly on simple molecules. The hydrogenation of CO to form H₂CO and CH₂OH has been extensively studied (e.g. Hiraoka et al. 1994; Watanabe & Kouchi 2002; Fuchs et al. 2009), as have the reaction pathways to the formation of H₂O from atomic oxygen (Hiraoka et al. 1998; Dulieu et al. 2010), molecular oxygen (Miyauchi et al. 2008; Ioppolo et al. 2008; Chaabouni et al. 2012), and ozone (Mokrane et al. 2009). The formation of C₂H₅OH, CH₄, H₂CO, and CH₃OH from CH₃CHO has been studied by Bisschop et al. (2007b), while the formation of NH₂OH by the hydrogenation of NO (Congiu et al. 2012) and the formation of CH₂NH and CH₃NH₂ from HCN (Theule et al. 2011) have recently been demonstrated.

Given that atomic hydrogen is present at fractional abundances of [H₄]/[H₂] ~ 10⁻³ in molecular clouds (Li & Goldsmith 2003), hydrogenation is likely to dominate the destruction pathways of HNCO at low temperatures. Theoretical studies may suggest that the radical intermediates HNCHO or NH₂CO form rapidly from HNCO, and the stable molecule formamide forms on further hydrogenation of these intermediate species (Garrod et al. 2008), giving overall

\[
HNCO + 2H \rightarrow H₂NCHO. \tag{1}
\]

This work focuses on the reaction of HNCO with H, which has not yet been examined experimentally. The experiments performed are introduced in Sect. 2, the results of these experiments are presented and discussed in Sect. 3, while the astrophysical implications of the results are considered in Sect. 4.

2. Experimental

Experiments were performed using two different experimental set-ups: RING, as described in Theule et al. (2011), and FORMOLISM, as described in Amiaud et al. (2006).

The RING set-up was used to perform multilayer, bulk ice experiments. Briefly, RING consists of a gold-plated copper surface within a high vacuum chamber (a few 10⁻⁹ mbar). Molecular species in the form of room temperature gas are dosed onto the surface held at 90 K via a molecular beam. This ensured that the deposited HNCO was not mixed with the byproduct CO₂. The HNCO was cooled to ~10 K, then bombarded with H or D atoms (at approximately 300 K, with a flux of 1 ± 0.3 × 10¹⁴ cm⁻² s⁻¹) produced in a hydrogen or deuterium plasma in the second molecular beam. After bombardment, the ices were probed using mass spectrometry during temperature programmed desorption (TPD) experiments.

The spectra are as follows: a) pure HNCO deposited at 17 K; b) HNCO bombarded with H atoms for 140 min; and c) the difference spectrum of traces a and b. Trace d) is a reference spectrum discussed in full in the text.

The HNCO ice was bombarded with H atoms (at approximately 300 K, with a flux of ~10¹⁵ cm⁻² s⁻¹) produced in a molecular hydrogen plasma generated by a 2.45 GHz microwave discharge. The plasma source and its calibration are fully detailed in Theule et al. (2011). The hydrogenation was monitored at regular time intervals using IR spectroscopy. In monolayer and sub-monolayer experiments, HNCO was dosed onto the surface held at 90 K via a molecular beam. This ensured that the deposited HNCO was not mixed with the byproduct CO₂. The HNCO was cooled to ~10 K, then bombarded with H or D atoms (at approximately 300 K, with a flux of 1 ± 0.3 × 10¹⁴ cm⁻² s⁻¹) produced in a hydrogen or deuterium plasma in the second molecular beam. After bombardment, the ices were probed using mass spectrometry during temperature programmed desorption (TPD) experiments.

3. Results and discussion

3.1. The multilayer, bulk HNCO regime

3.1.1. H bombardment of HNCO

The spectrum of pure multilayer HNCO deposited at 17 K is presented in Fig. 1, curve a. The molecular species HNCO is identified via its characteristic absorption bands, as listed in Table 1. The most intense bands are the N–H stretching mode absorption infrared spectroscopy (FT-RAIRS) using a MCT detector in a Vertex 70 spectrometer. A typical spectrum has a 1 cm⁻¹ resolution and is averaged over a few tens of interferograms.

The FORMALISM set-up was used to perform experiments in the monolayer and sub-monolayer regime. The experimental set-up consists of an ultra high vacuum (UHV) chamber (base pressure ~1 × 10⁻¹⁰ mbar), containing a previously oxidised graphite HOPG sample (7–400 K, controlled by a closed-cycle He cryostat). Molecules are dosed onto the surface via two triply differentially pumped beam lines. Desorption of molecules from the surface is monitored using a quadrupole mass spectrometer (QMS, Hiden HAL-3F), positioned directly in front of the surface.

Isocyanic acid was prepared in the gas phase from cyanoacetic acid (HNCO)₃ via thermal decomposition of the commercially available trimer (Aldrich, 98%) at 650 °C under primary vacuum (Raumier et al. 2003b). Small quantities of CO₂ and traces of CO are always present in the HNCO as a residual of the synthesis method.

In the multilayer experiments presented here, HNCO was dosed onto the surface held at 17 K via an injection line.

The pure solid HNCO at 17 K was bombarded with H atoms for a total of 140 min (corresponding to a dose of ~8 × 10¹⁷ cm⁻²). The temperature of the deposition and H bombardment was chosen to be low enough to allow the H atoms to have a relatively long residency time on the surface, while being high enough to allow the H atoms to have a high mobility on the surface, to penetrate the bulk ice as deeply as possible, and to aid in overcoming any potential activation barrier to the hydrogenation of HNCO. The spectrum of the H-bombarded HNCO is presented in Fig. 1, curve b, while the difference spectrum of pure and H-bombarded HNCO is presented in Fig. 1, curve c. The positions of the absorption bands of the observed products are given in Table 1.

There are a number of crucial points to note with regard to the difference spectrum of H-bombarded HNCO (Fig. 1, curve c). First, in the wavelength region 2200–2300 cm⁻¹...
the peak absorption at 2240 cm$^{-1}$ has diminished, indicative of a decrease of approximately 1 to 2 monolayers (of 580 total monolayers, calculated assuming a band strength of $7.8 \times 10^{-17}$ cm molec$^{-1}$ van Broekhuizen et al. 2004). Bands appearing at $\sim 3300$ cm$^{-1}$ and 1680 cm$^{-1}$ are attributed to H$_2$O, and the band at 2342 cm$^{-1}$ is attributed to CO$_2$; both of these species are contaminants introduced via the H plasma, confirmed by blank experiments where the bare gold surface was bombarded. Figure 1, curve d is a pure H$_2$O spectrum at 15 K fitted to the 1680 cm$^{-1}$ band of the difference spectrum. Thus, H$_2$O accounts for most of the main feature in the difference spectrum (at $\sim 3300$ cm$^{-1}$) as well as the full 1680 cm$^{-1}$ band. Remaining differences between the spectra of pure and H-bombarded HNCO are relatively minor.

After inspection of the difference spectrum it appears that the expected hydrogenation product, methanamide (formamide, NH$_2$CHO), has not been produced in detectable quantities. Arguably the simplest potential product, formamide would be an ion of the environment, with a range of around 2145–2305 cm$^{-1}$ (Falk & Giguère 1957), and much lower than that of OCN$^-$ (1.3 $\times$ 10$^{-16}$ cm molec$^{-1}$ van Broekhuizen et al. 2004), so the absorption features would be vanishingly weak in our ice. If OCN$^-$ is present in very small quantities: approximately 0.1 monolayers account for the peak produced after 140 min of H bombardment. However, given the quantity of H$_2$O deposited over the course of the H bombardment, it is not unreasonable that HNCO could react as in Eq. (2), as the rate of reaction was found to be 26 kJ mol$^{-1}$ (Theule et al. 2011), which could be delivered to the ice surface by room temperature H atoms.

Another potential, but rather unlikely, explanation for the absorption feature is that the radical species NCO is formed by reaction with the deposited H$_2$O:

$$\text{HNCO} + \text{H}_2\text{O} \rightarrow \text{OCN}^- + \text{H}_3\text{O}^+. \quad (2)$$

The counterion H$_3$O$^+$ was not observed in the H-bombarded HNCO ice, but it should be noted that the band strength is very weak (Falk & Giguère 1957), and much lower than that of OCN$^-$(1.3 $\times$ 10$^{-16}$ cm molec$^{-1}$ van Broekhuizen et al. 2004), so the absorption features would be vanishingly weak in our ice. If OCN$^-$ is the species responsible for the absorption band at 2163 cm$^{-1}$, it is present in very small quantities: approximately 0.1 monolayers account for the peak produced after 140 min of H bombardment. However, given the quantity of H$_2$O deposited over the course of the H bombardment, it is not unreasonable that HNCO could react as in Eq. (2), as the rate of reaction was found to be 26 kJ mol$^{-1}$ (Theule et al. 2011), which could be delivered to the ice surface by room temperature H atoms.

Another potential, but rather unlikely, explanation for the absorption feature is that the radical species NCO is formed by decomposition of HNCO, probably by H abstraction:

$$\text{HNCO} + \text{H} \rightarrow \text{NCO} + \text{H}_2. \quad (3)$$

The spontaneous decomposition of HNCO adsorbed on metal surfaces has been extensively reported in the surface science literature (e.g. Kiss & Solymosi 1983; Celio et al. 1997; Jones & Trenary 2008). The NCO radical formed by this decomposition is relatively stable, being observed on the surface at temperatures up to 600 K under UHV conditions (Celio et al. 1997). The position of the adsorption band associated with the asymmetric stretching mode absorption of NCO is highly dependent on the environment, with a range of around 2145–2305 cm$^{-1}$ (e.g. Németh et al. 2007).

One final comment to make about the multilayer reactivity of HNCO is that the VUV irradiation of pure HNCO has previously been shown to produce formaldehyde, formamide, and urea (Rauzier et al. 2004). In this work, urea was not observed in the IR spectra after H bombardment of HNCO. As the N=C bond must be broken to form urea, this route is unlikely. The only clear product of H bombardment of HNCO is the species responsible for the absorption band at 2163 cm$^{-1}$; probably OCN$^-$. The other absorption features remain ambiguous.

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**Table 1.** Fundamental infrared band positions (cm$^{-1}$), for the species identified in the ice during the bombardment of HNCO$^{ab}$.  

<table>
<thead>
<tr>
<th>Species</th>
<th>$\nu$N-H</th>
<th>$\nu$C-H</th>
<th>$\nu$=C=O$_{as}$</th>
<th>$\nu$C=O</th>
<th>$\delta$N-H$_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>HNCO</td>
<td>3554/3562/3231</td>
<td>-</td>
<td>2240</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>NH$_2$CHO</td>
<td>3313/3169$^c$</td>
<td>2886$^c$</td>
<td>-</td>
<td>1685$^c$</td>
<td>1385$^c$</td>
</tr>
<tr>
<td>OCN$^-$/</td>
<td>-</td>
<td>2163$^a$</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>CO</td>
<td>-</td>
<td>-</td>
<td>2136$^b$</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>CO$_2$</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>2344$^a$</td>
<td>-</td>
</tr>
</tbody>
</table>

**Notes.** $\nu$ represents a stretching vibration, and $\delta$ a bending vibration.  
(a) Positively identified in the H-bombarded HNCO ice.  
(b) Tentatively identified in the H-bombarded HNCO ice.  
(c) Not identified in the H-bombarded HNCO ice.

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**Fig. 2.** Observed products of H + HNCO in the multilayer regime. In the left panel, the spectra are as follows: a) the new band at 1680 cm$^{-1}$ (magnification of the difference spectrum in Fig. 1c); b) a spectrum of pure H$_2$O, scaled to the band at 1680 cm$^{-1}$ (magnification of Fig. 1d); and c) a spectrum of pure NH$_2$CHO, scaled to the band at 1680 cm$^{-1}$. In the right panel, the spectra are the difference spectra of H-bombarded HNCO and pure HNCO are shown for bombardment times of d) 20 min; e) 80 min; and f) 140 min (magnification of the difference spectrum in Fig. 1c).
3.1.2. H bombardment of NH$_2$CHO

As the results for the hydrogenation of HNCO were inconclusive, the reactivity of the expected product, formamide, was examined. The hydrogenation of formamide could lead to the production of aminomethanol according to

$$\text{NH}_2\text{CHO} + 2\text{H} \rightarrow \text{NH}_2\text{CH}_2\text{OH}. \quad (4)$$

The molecule NH$_2$CH$_2$OH is the most saturated form of the NCO moiety, so is the logical end-point for the hydrogenation chemistry of HNCO.

Gas phase NH$_2$CHO was dosed onto the surface held at 15 K, then bombarded with H atoms for 190 min using the RING experimental set-up. The spectrum of pure NH$_2$CHO at 15 K is presented in Fig. 3, curve a, with the H-bombarded sample spectrum as curve b. The difference spectrum of pure and H-bombarded NH$_2$CHO is presented in Fig. 3, curve c.

Pure formamide exhibits peaks at 3313, 3169, 2886, 1683, 1385, and 1327 cm$^{-1}$, as seen in Fig. 3, curve a. The CO$_2$ molecule was present as a pollutant in the deposited NH$_2$CHO, as shown by the absorption feature in curve a (at 2341 cm$^{-1}$), and in the H plasma, resulting in absorptions in curves b and c. There is no clear decrease observed in the peak of the νC=O absorption band at 1681 cm$^{-1}$, and thus we conclude that NH$_2$CHO has not reacted.

Similarly to the bombarded HNCO, the difference spectrum of bombarded NH$_2$CHO is dominated by absorption in the range 3600–3000 cm$^{-1}$. As discussed above, this might have multiple sources, but a large contribution is due to H$_2$O. The four small peaks on the red wing between 2961 and 2859 cm$^{-1}$ are again attributed to contamination of the H plasma due to primary vacuum. Such contamination is only observed because of the very long irradiation times. The peak observed in the difference spectrum at 1676 cm$^{-1}$ is interpreted as further evidence of the deposition of H$_2$O, with a potential contribution from the crystallisation of the NH$_2$CHO sample during H bombardment. We confirmed that, after heating a sample of pure NH$_2$CHO at 2 K min$^{-1}$, the crystallisation of NH$_2$CHO results in the shifting of the νC=O absorption band by 4–5 cm$^{-1}$ towards the blue, as illustrated in the inset of Fig. 3. After H bombardment of NH$_2$CHO, no formation of OCN$^-$ was observed. We conclude that formamide is unreactive to H under our experimental conditions, and aminomethanol is not formed via this route.

3.2. The monolayer HNCO regime

Because of the uncertainty in interpretation of our multilayer HNCO data, related to the low level contamination introduced by the H plasma of RING, we decided to perform experiments in the monolayer regime using FORMOLISM. Both experimental set-ups are described in Sect. 2, but the major advantage of FORMOLISM for this study is a base pressure of $\sim 10^{-8}$ mbar in the last stage of the molecular jets, which reduces the level of contamination of the H introduced into the chamber. Working in the monolayer regime allows us to bombard all deposited HNCO with H, unlike the multilayer regime where hydrogen penetrates the first $\sim 1$–3 monolayers, and the signal of the bulk HNCO ice overlows that of product molecules in both the Fourier transform infrared (FTIR) and mass spectra. However, the monolayer regime also limits the signal-to-noise ratio of potential products observed using mass spectrometry and, in these experiments, we were unable to observe a clear signal for our HNCO reactant using FTIR spectroscopy.

3.2.1. Desorption energy of HNCO

Using FORMOLISM, we performed experiments to determine the deposition conditions necessary to deposit 1 ML of HNCO on the graphite surface; HNCO was deposited for a series of fixed times, and TPDs were performed after each deposition. The TPD spectra are shown in Fig. 4. The characteristic monolayer deposition time was determined by visual inspection of the leading edges of these spectra. All HNCO depositions were subsequently quantified by comparison with the identified monolayer deposition (Fig. 4, curve b). The low flux in the molecular beam allows us to deposit very reproducible quantities of molecular species. It takes almost 13 min to deposit 1 ML of HNCO, so the deposition uncertainty on coverages of 0.5 ML and 1 ML (used in this work) is vanishingly small and the depositions are highly reproducible.

The multilayer desorption energy of HNCO was calculated via two different methods. The rate of desorption by unit surface, $r$, can be expressed by the Polanyi-Wigner equation (Redhead 1962; Carter 1962; King 1975), where the desorption rate constant $k_{des}$ is described in terms of an Arrhenius law,

$$r = \frac{dN}{dT} = \frac{A}{\beta} e^{-E_{des}/RT} N^n,$$  \quad (5)
where $A$ is the pre-exponential factor, $\beta = \frac{dT}{dt} = 12 \text{ K min}^{-1}$ is the heating rate, $E_{\text{des}}$ is the energy of desorption of a molecule from the surface (J mol$^{-1}$), $R$ is the gas constant (J K$^{-1}$ mol$^{-1}$), $T$ is the temperature of the surface (K), $N$ is the number of adsorbed molecules on the surface (molecules cm$^{-2}$), and $n$ is the order of the reaction. The units of $A$ depend on $n$: molecules$^{1-n}$ cm$^{-2}$ s$^{1-n}$.

By fixing the pre-exponential factor, $A$, at a value of $10^{28}$ molecules cm$^{-2}$ s$^{-1}$ (assuming that the lattice vibrational frequency of the solid is $10^{13}$ s$^{-1}$ and the number of molecules in a monolayer is approximately $10^{15}$ cm$^{-2}$), the desorption energy is calculated as the only free parameter in the fit of Eq. (5) to the leading edge of the experimental data (Collings et al. 2003). This fit is plotted as a dotted line in Fig. 4. Using this method, the calculated desorption energy was $E_{\text{ads, HNCO}} = 32.9 \pm 1.7$ kJ mol$^{-1}$ ($3957 \pm 204$ K).

A second fitting method, proposed by Hasegawa et al. (1992), assumes that the pre-exponential factor in Eq. (5) is a function of $E_{\text{ads}}$ approximated by

$$A = N_{\text{ML}} \nu = N_{\text{ML}} \sqrt{\frac{2N_{\text{ML}} E_{\text{ads}}}{\pi^2 M}},$$

where $M$ is the mass of the adsorbate molecule, and $N_{\text{ML}} \sim 10^{15}$ cm$^{-2}$. The advantage of this method is that the fit requires only one variable, $E_{\text{ads}}$, rather than assuming or fitting the pre-exponential factor and fitting $E_{\text{des}}$ (Acharyya et al. 2007; Noble et al. 2012). Using the second method, the derived desorption energy was $E_{\text{ads, HNCO}} = 31.0 \pm 1.6$ kJ mol$^{-1}$ ($3729 \pm 192$ K).

### 3.2.2. H and D bombardment of HNCO

A series of experiments were performed on the H or D bombardment of monolayer and sub-monolayer quantities of solid HNCO at 10–15 K. In this temperature range, H atoms do not form a bulk solid because of the rapid self recombination on the surface. A steady state, with an H$_2$ coverage of $\sim 10\%$ (Amiaud et al. 2007; Kristensen et al. 2011) due to desorption, is reached within the first ten seconds of H bombardment (Congiu et al. 2009).

Our initial experiments into the monolayer H or D bombardment of HNCO were carried out on short timescales. The results of 10 min (i.e. 1.7 ML) of H or D bombardment of 0.5 ML HNCO are shown in Fig. 5. The left panel includes two curves: the TPD spectrum of m/z 43 (HNCO) for 0.5 ML pure HNCO (curve a) and the corresponding spectrum after 10 min of H bombardment (curve b). The H-bombarded HNCO sample contains only 0.42 ML HNCO, which represents a 16% decrease compared to the pure HNCO sample. No m/z 45 (formamide) was observed to desorb during the TPD experiment, suggesting that no reaction occurred during H bombardment. However, the difference of 0.08 ML HNCO could be, at least partly, due to the low signal-to-noise ratio of the spectra (which is limited by the simultaneous measurement of multiple m/z).

The results of D bombardment, shown in the right panel of Fig. 5, are more revealing. After D bombardment we observe the species DNCO in our ice sample. In Fig. 5 the TPD spectrum of m/z 43 (HNCO) for 0.5 ML pure HNCO is again plotted as curve a. The remaining curves, corresponding to 10 min of D bombardment (1.7 ML), are m/z 43 (HNCO, curve c) and m/z 44 (DNCO, curve d), respectively. In this experiment, only 0.33 ML of HNCO remain after 10 min of D bombardment, representing a decrease of 34%. Assuming that the mass spectrometer is equally sensitive to HNCO and DNCO, we calculate the quantity of DNCO on the surface after D bombardment to be $\sim 0.18$ ML.

Thus, the total quantity of HNCO and DNCO on the surface represents 0.51 ML, i.e. equivalent to the quantity of HNCO originally deposited. We will discuss the implications of the deuteration of HNCO in Sect. 3.2.4.

### 3.2.4. Proposed reaction mechanism

In order to determine the origin of the HNCO loss, we searched for the presence of formamide in the D-bombarded HNCO sample. The TPD method is typically sensitive to $\sim 0.01$ ML (Noble et al. 2011). In Fig. 7, curve a, we show the TPD spectrum of 1 ML pure HNCO (m/z 43) as a reference. All other curves are m/z 47 (NHDCDO, formamide) desorbed during the TPD, therefore we observe a conversion efficiency from HNCO to DNCO of approximately 100%.
hydrogenation of HNCO to NH₂CHO (Eq. (1)) does not readily occur and thus has a very high barrier. However, it is clear that D bombardment of HNCO produces DNCO and, additionally, that for longer bombardment times there is some loss of HNCO that is not accounted for by this DNCO formation. We provide a list of typical experiments, including the percentage loss of HNCO, in Table 2. By comparing the loss of initial HNCO to the doses of H and D, we see that there is a general trend in the data, whereby one molecule of HNCO is lost from the sample for approximately 1–1.5% of the incident H and D atoms. Although there was fluctuation in the absolute quantity of HNCO lost from the sample in our experiments, a loss was consistently seen for all H and D exposures greater than ~2 ML. These losses are likely due to chemical desorption from the surface (Dulieu et al. 2013).

Chemical desorption is the stimulated desorption of reactant or product molecules due to an exothermic surface reaction. Recent experimental studies using FORMOLISM have highlighted the significance of the chemical desorption mechanism in the monolayer regime at low temperatures (Dulieu et al. 2013; Minissale et al. 2014; Noble et al. 2011). It has been shown that up to 90% of molecules (on a graphite surface) formed by radical-molecule or radical-radical surface reactions sublimate from the surface after formation because of the inability of the surface to quench the energy released during the reaction.

On ice surfaces, more astrophysically relevant for the study of molecules such as HNCO (which will not be present on bare grains), this number is typically much lower.

To account for both the formation of DNCO (after D bombardment) and the loss of HNCO from the surface (after both H and D bombardment), the chemistry is required to proceed via a cyclic pathway, such as the two proposed here:

$$\text{HNCO} + \text{H} \rightarrow \text{H}_3\text{NCO} \rightarrow \text{HNCO} + \text{H}_2, \quad (7a)$$

$$\text{HNCO} + \text{D} \rightarrow \text{OCN} + \text{H} + \text{H} \rightarrow \text{HNCO}. \quad (7b)$$

Cycle 7a corresponds to a classical hydrogenation pathway, i.e. H addition, but at the second step the branching ratio between the products NH₂CHO (Eq. (1)) and HNCO is fully dominated by the reverse reaction to the initial HNCO population. Cycle 7b corresponds to the abstraction of a hydrogen atom by the incident H atom. The reaction intermediates H₂NCO and/or OCN are not expected to be observed in the TPDs as these radicals would either react with incident H atoms at a much faster rate than HNCO or, ultimately, recombine during the heating phase. It is also possible that, if cycle 7b occurs, the second step could result in the isomerisation of HNCO to HOCN (as previously observed during the heating of mixtures of HNCO and H₂O (Theule et al. 2011)). However, under current sub-monolayer experimental conditions it is not possible to differentiate between HNCO and HOCN, and no evidence of HOCN was observed in the multilayer experiments.

The results of our D bombardment experiments are critical to explaining the reaction mechanism. At a low dose of D (Table 2 and Fig. 5), we do not observe any loss of reactants or products from the surface, but deuteration appears to be very efficient. Around 34% of the initial HNCO (0.17 ML) is transformed into DNCO. Of the total D atoms incident on the surface, approximately 10% have been included in the product. This represents a high efficiency, particularly when we consider that the reaction D + HNCO is in competition with D + D. For comparison, the CO + H system described by Fuchs et al. (2009) or Watanabe & Kouchi (2002) uses only 1% of the incident H atoms in the formation of H₂CO and CH₃OH. Although we perform our experiments under different conditions, we are able to place the reaction HNCO + H between that of H + H₂CO (which is faster) and H + CO (which is slower). It is clear that the barrier to the reaction HNCO + D is not very high, and is probably in the region of 250–700 K. A specific study investigating varying doses and surface temperatures would be required to calculate this value; this represents a large amount of work and is beyond the scope of this paper.

When we extend D bombardment to longer times, we observe a loss of material from the surface in addition to the formation of DNCO (Table 2 and Fig. 6). For an initial deposition of 1 ML HNCO subjected to 25 ML of D, we recover 43% of the HNCO and produce 22% DNCO i.e. a loss of 35% of the original material. Assuming that the proportion of D atoms reacting is the same as for short timescales (~10%), we approximate that HNCO + D occurs 2.5 × 10¹⁵ cm⁻² times. The observed loss represents 3.5 × 10¹⁴ molec cm⁻², and we can make a rough estimation that the efficiency of chemical desorption is ~14%. Unlike in some previous studies (Dulieu et al. 2013), the relative efficiency of the desorption mechanism must be low, as only 0.35 ML is lost from the surface over a bombardment period of 150 min. This is equivalent to a desorption rate of ~4 × 10⁻³ ML s⁻¹, and as such any desorbing molecules would not be observable using the mass spectrometer. The result
Table 2. Results of experiments carried out on monolayer and sub-monolayer quantities of HNCO.

<table>
<thead>
<tr>
<th>Initial HNCO (ML)</th>
<th>Exposed to</th>
<th>Temperature (K)</th>
<th>HNCO reduction (%)</th>
<th>DNCO produced (%)</th>
<th>Loss (to CD) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>1.7 ML D</td>
<td>10</td>
<td>34 ± 5</td>
<td>36 ± 5</td>
<td>−2 ± 5</td>
</tr>
<tr>
<td>1.0</td>
<td>10 ML H</td>
<td>7</td>
<td>11 ± 5</td>
<td>none</td>
<td>11 ± 5</td>
</tr>
<tr>
<td>1.0</td>
<td>25 ML D</td>
<td>15</td>
<td>57 ± 5</td>
<td>22 ± 5</td>
<td>35 ± 5</td>
</tr>
</tbody>
</table>

Fig. 8. Proposed reaction scheme for HNCO with H.

Figures indicate that the reaction driving the chemical desorption mechanism is not highly exothermic.

Each of the two reaction scenarios that we proposed above proceeds via two steps. The first step (H-abstraction or H-addition) limits the kinetics of the overall reaction and is also less likely to provoke chemical desorption since less energy will be released than in the second radical-radical step. Thus most of the desorption should occur at the second reaction step. If the reactivity of both the HNCO and DNCO species was equivalent, we would assume that, with the longer exposures, over 90% of the initial HNCO should be transformed, which is not the case. The low deuteration yield could be due to the presence of a second reactant (DNCO) and/or steric limitation of the HNCO desorption. However, we propose that the most likely limiting factor is the stabilisation of the HNCO by its formation of dimers or larger polymers, preventing its reaction with H. Experiments performed on HNCO in a H2O ice, or isolated in a low temperature matricial gas, would reduce the intermolecular bonding and may allow hydrogenation to occur. Because of the inability to observe desorbing HNCO/DNCO (owing to the low rate of chemical desorption) and the hypothesised intermediates (OCN and/or H2NCO) by mass spectrometry, we are unable to determine whether one reaction cycle dominates the reactivity of HNCO + H. In order to differentiate between the abstraction and addition mechanisms, a full ab initio quantum mechanical calculation treatment including surface effects would be required.

By comparing our monolayer experiments with those in bulk ice, we confirm that formamide is not produced in detectable quantities after H bombardment of HNCO. It is also plausible that the inconsistency between the loss of HNCO (~1–2 ML) and the only confirmed product OCN−/OCN (~0.1 ML) in our bulk experiments can be explained by the proposed cyclic reactivity of HNCO combined with chemical desorption. The efficient H/D cyclic substitution suggests that the N-H bond in HNCO is relatively weak, which strengthens the argument for a low level of HNCO decomposition into OCN. We show our proposed overall reaction scheme in Fig. 8.

4. Astrophysical implications

Our results contradict theoretical studies which proposed that the initial hydrogenation step to form either HNCHO or NH2CHO is very important to HNCO chemistry and that subsequent hydrogenation leads to NH2CHO (e.g. Garrod et al. 2008; Tideswell et al. 2010). However, ab initio calculations determine an activation barrier of 1390 K to the first hydrogenation step (Nguyen et al. 1996). A recent experimental study of the formation of HNCO by addition of the radical NH to CO revealed no formation of NH2CHO after co-deposition of N, H, and CO, despite formation of HNCO (Fedoseev et al. 2015).

The NH2CHO molecule has been tentatively identified in ices towards the objects W 33A (Schutte et al. 1999) and NGC 7538 IRS9 based on comparison with laboratory spectra (Raunier et al. 2004), but the identification of molecules in the 6–8 μm region is complicated by the overlap of multiple absorption bands, and the results of these two studies are not conclusive (Boogert et al. 2008). What are the reactions that would drive the chemistry of NH2CHO in the interstellar medium (ISM)? Although gas phase formation routes to complex molecules exist, it is generally accepted that grain surface chemistry is the dominant formation mechanism for such species (Bisschop et al. 2007a). The NH2CHO molecule can form by recombination between the radicals NH2 and HCO following energetic processing of ices by, for example, UV photons (Allamandola et al. 1999; Muñoz Caro & Schutte 2003; van Broekhuizen et al. 2004).

It has recently been determined that NH2CHO is the most energetically stable CH3NO isomer that can be formed, and that the amide bond is the most stable bond possible (Lattanzi et al. 2010). Jones et al. (2011) contend that the hydrogenation of HNCO is insignificant as a route to formamide formation because of the low barrier to the reaction HNCO + H (Raunier et al. 2003a; Mispelaer et al. 2012). While it is true that NH3 does react rapidly with HNCO, the flux of hydrogen atoms onto a grain will be superior to the quantity of NH3 molecules in the ice mantle. This is particularly true before formation of NH3 in the mantle. In order to reproduce observed abundances of HNCO and OCN− in dense molecular clouds, it has been speculated that hydrogenation of HNCO must dominate its destruction mechanisms (Theule et al. 2011). However, our results suggest that hydrogenation is not an efficient process and thus thermal reactions, such as the reaction with NH3 or H2O, should dominate HNCO grain surface chemistry.

It has already been demonstrated that CH3CHO undergoes hydrogenation after H bombardment, but that it also forms CH4, H2CO, and CH3OH (Bisschop et al. 2007b). The hydrogenation product, ethanol, represents approximately 20% of the total products formed. The relative strength of the C=O bond prevents hydrogenation being the most prevalent reaction, and the C-C bond breaks to form products that are chemically simpler and of lower mass than the original CH3CHO. Our results confirm that hydrogenation of the C=O bond is not favourable under low temperature H bombardment conditions.

Our results show that the OCN moiety is formed in its radical or anionic form during H bombardment. The OCN− ion has been positively identified in interstellar ices (Soifer et al. 1979; Grim & Greenberg 1987) and its rotational spectrum has been measured (Lattanzi et al. 2010), so searches in the gas phase are possible. The neutral OCN (Kawaguchi et al. 1985) has not yet
been identified in the ISM; it has, however, long been predicted in dense clouds (Prasad & Huntress 1978). If OCN/OCN$^-$ can be formed by the reaction of H with HNCO, this could help explain the destruction of HNCO, while confirming that for- 
mamide formation does not occur via this route. Additionally, in dense regions such as pre-stellar cores, H/D exchange is poten- 
tially the dominant destruction mechanism for HNCO (Roberts et al. 2003).

We have also shown that formamide does not react with hy- 
drogen to produce aminomethanol, the logical saturated end-
point for HNCO hydrogenation. Aminomethanol has, however,
been already been shown to form under interstellar conditions by the 
purely thermal reaction between H$_2$CO and NH$_3$, with an activ- 
ization barrier of 4.5 ± 0.5 kJ mol$^{-1}$ (541 ± 60 K, Bossa et al. 2009). Theoretical studies suggest that, in the presence of acids,
aminomethanol can undergo spontaneous dehydration to form methyleneimine (CH$_2$=NH) and water (Walch et al. 2001) experi-
ments have shown that aminomethanol yields hexamethylenete-
tramine (HMT) from the polymerisation of the dehydration product methyleneimine (Bernstein et al. 1995; Vinogradoff et al. 2011). Although it has not yet been observed in the ISM, in
the laboratory HMT has a sublimation temperature of ∼2011). Although it has not yet been

References