The abundance of HOC$^+$ in diffuse clouds

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Abstract. We used the Plateau de Bure Interferometer to search for 13mm absorption lines of HOC$^+$ from local diffuse and translucent clouds occulting compact extragalactic mm-wave continuum sources. We detected HOC$^+$ in three directions with column densities only 70–120 times below that of the HCO$^+$ isomer, a factor 5–50 higher than typically found in dense dark gas but comparable to recent observations of dense photon-dominated regions. The observed amounts of HOC$^+$ $N$(HOC$^+$/N(H$_2$) = 3–6 x 10$^{-11}$) can be made in quiescent diffuse gas at thermal gas-kinetic rates if the H$_2$O/OH ratio is of order unity, in mild violation of extant observational limits.

Key words. ISM: molecules – astrochemistry

1. Introduction

The roster of polyatomic molecules known to exist in diffuse interstellar gas has been steadily enlarged over the last fifteen years and now numbers nearly one dozen. Two of these, C$_3$ (Maier et al. 2001; Roueff et al. 2002; Oka et al. 2003) and H$_3^+$ (McCall et al. 2003) were recently discovered in optical and near-infrared absorption spectra. The others have accumulated gradually over the last twenty or so years in cm-wave and mm-wave radiofrequency absorption data and now include H$_2$CO (Federman & Willson 1982; Nash 1990; Marscher et al. 1993; Liszt & Lucas 1995; Moore & Marscher 1995), HCO$^+$ (Liszt & Lucas 1996), NH$_3$ (Nash 1990), C$_2$H (Lucas & Liszt 2000), C$_3$H$_2$ (Cox et al. 1988; Lucas & Liszt 2000), HCN and HNC (Liszt & Lucas 2001) and H$_2$S and HCS$^+$ (Lucas & Liszt 2002).

Observationally, the polyatomics observed with radio techniques are often seen to be strongly and directly related to simpler species observed at optical wavelengths, i.e. HCO$^+$ to OH (Liszt & Lucas 1996; Lucas & Liszt 1996) HCN and HNC to CN (Liszt & Lucas 2001) etc. These connections point to fundamental lacunae in our understanding of the chemistry of even the simplest species in diffuse clouds: simple models which reproduce the abundances of diatomics, however easily, invariably fall orders of magnitude short of explaining the related polyatomics. Conversely, the high abundances of some polyatomics transform other mysteries: the poorly-understood abundances of CO and CS in diffuse clouds are readily understandable given the recombination of the observed columns of HCO$^+$ (Liszt & Lucas 2000) and HCS$^+$ (Lucas & Liszt 2002) respectively.

The work described here rounds out to an even dozen the number of polyatomics known to exist in diffuse and translucent clouds. HOC$^+$, the energetically-disfavored HCO$^+$ isomer whose 89.4874 GHz $J$ = 1–0 transition we sought, was indeed detected along several of the sightlines studied in the other molecular species mentioned in the preceding paragraphs. The HCO$^+$-HOC$^+$ comparison is especially interesting now because both species have recently been detected over a wide range of physical conditions in dense gas: our new spectra reinforce an emerging view which relates molecular abundances in dense and diffuse PDR. Section 2 gives details of the observations, which are discussed in Sect. 3 and summarized in Sect. 4.

2. Observations of HOC$^+$

The 89.4874 GHz $J$ = 1–0 transition of HOC$^+$ was observed at the Plateau de Bure at various times during the period 2000–2001 toward the five sources listed in Table 1. Profiles were taken with 39.1 kHz (0.13 km s$^{-1}$) channel separation and 70 kHz resolution and subsequently once hanning smoothed, yielding spectra with the line/continuum rms values as listed in Table 1.

The column density in the lowest ($J$ = 0) level of a simple linear molecule is related to the integrated optical depth of the $J$ = 1–0 transition as

$$N_0 = \frac{8.0 \times 10^{13} \text{ cm}^{-2} \int \tau_{10} \text{dv}}{\mu^2(1 - \exp(-h\nu_{10}/kT_{\text{exc}}))}$$

(1)
where $\mu$ is the permanent dipole moment of the molecule, $T_{\text{exc}}$ is the excitation temperature of the $J = 1 \rightarrow 0$ transition and $v_{10}$ is its frequency.

With the exception of CO (Liszt & Lucas 1998), the molecules observed at mm-wavelengths in diffuse gas are well-described by assuming that the rotational ladder is thermalized at the temperature of the cosmic microwave background, in which case the total column density can be calculated straightforwardly from observations of one transition. We take the dipole moments of HOC$^+$ and HCO$^+$ as 2.8D and 3.93D, respectively, a slight departure from prior work in which we used 4.07D for HCO$^+$. Given these assumptions, one has $N(X)/\int \tau d\nu = 2.146 \times 10^{12}$ cm$^{-2}$ for $X = $ HOC$^+$, $1.093 \times 10^{12}$ cm$^{-2}$ for $X = $ HCO$^+$ and $1.128 \times 10^{12}$ cm$^{-2}$ for $X = $ H$^{13}$CO$^+$.

3. Observed and expected abundances of HOC$^+$ in diffuse gas

3.1. Observed abundance of HOC$^+$

The mm-wavelength data are summarized in Tables 1–3 and line profiles toward two sources are shown in Figs. 1 and 2. The HCO$^+$ and OH column densities are taken from our prior published (Lucas & Liszt 1996; Liszt & Lucas 1996) and unpublished work; for B0415+379 (see Fig. 1) we assert $N($HCO$^+$) = $59 \pm 2 N($H$^{13}$CO$^+$) (Lucas & Liszt 1998) because of the high optical depth in the main isotope. HOC$^+$ is found to have relative abundances HCO$^+$:HOC$^+$ = 70–120, HOC$^+$:OH = 3–6 $\times 10^{-3}$ toward three sources (see Table 3). The results obtained toward the other two, B0528 and B1730, are not of great significance. The relative abundance of OH, for which it is found that $X($OH$) = n($OH$)/n(H_2) = 1.0 \pm 0.2 \times 10^{-7}$ along the four sightlines where both $N$(H$_2$) and $N$(OH) have been measured (Liszt & Lucas 2002), sets the relative abundance scale for HOC$^+$, $X($HOC$^+)$ = $3\rightarrow 6 \times 10^{-11}$.

Note that the OH column density toward B2200+420 is only 20% larger than that seen along the archetypal diffuse line of sight toward $\zeta$ Oph, $N($OH$) = 4.7 \pm 0.7 \times 10^{13}$ cm$^{-2}$ (Van Dishoeck & Black 1986; Roueff 1996), while the reddening ($E_{B-V}$ = 0.33 mag toward B2200+420 from Schlegel et al. (1998)) is the same. The larger total OH column density seen toward B0355+508 at low galactic latitude ($-1.6^\circ$) is roughly evenly divided among 5 kinematically-separated diffuse clouds of modest OH and HCO$^+$ column density, only two of which have prominent CO emission (Lucas & Liszt 1996; Liszt & Lucas 1996). The line of sight toward B0415+379 is somewhat darker (see the discussion in Lucas & Liszt (1998)) but relatively little of the carbon along even this line of sight is in CO and the density is fairly low, as gauged by the weakness of emission from species beside CO (Lucas & Liszt 1996).

### Table 1. Background sources observed.

<table>
<thead>
<tr>
<th>Source</th>
<th>$l$</th>
<th>$b$</th>
<th>$\sigma_{S0}$</th>
<th>$\int \tau($HOC$^+)$d$\nu$</th>
</tr>
</thead>
<tbody>
<tr>
<td>B0355+508</td>
<td>150.38$^a$</td>
<td>-1.60$^a$</td>
<td>0.0022</td>
<td>0.042(0.005)</td>
</tr>
<tr>
<td>B0415+379</td>
<td>161.68$^a$</td>
<td>-8.82$^a$</td>
<td>0.0029</td>
<td>0.051(0.004)</td>
</tr>
<tr>
<td>B0528+134</td>
<td>191.37$^a$</td>
<td>-11.01$^a$</td>
<td>0.0101</td>
<td>&lt;0.036</td>
</tr>
<tr>
<td>B1730-130</td>
<td>12.03$^a$</td>
<td>+10.81$^a$</td>
<td>0.0081</td>
<td>&lt;0.026</td>
</tr>
<tr>
<td>B2200+420</td>
<td>92.13$^a$</td>
<td>-10.40$^a$</td>
<td>0.0041</td>
<td>0.017(0.006)</td>
</tr>
</tbody>
</table>

1 rms noise in line/continuum ratio at 140 kHz (0.46 km s$^{-1}$) resolution.

### Table 2. Column densities.

<table>
<thead>
<tr>
<th>Source</th>
<th>$N($HOC$^+$)</th>
<th>$N($HCO$^+$)</th>
<th>$N($OH$)$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10$^{10}$ cm$^{-2}$</td>
<td>10$^{12}$ cm$^{-2}$</td>
<td>10$^{13}$ cm$^{-2}$</td>
</tr>
<tr>
<td>B0355+508</td>
<td>9.01(1.07)</td>
<td>6.51(0.02)</td>
<td>14.87(0.67)</td>
</tr>
<tr>
<td>B0415+379</td>
<td>10.9(0.90)</td>
<td>12.8(0.40)$^a$</td>
<td>42.50(0.10)</td>
</tr>
<tr>
<td>B0528+134</td>
<td>&lt;7.7$^a$</td>
<td>2.28(0.02)</td>
<td>4.72(0.31)</td>
</tr>
<tr>
<td>B1730-130</td>
<td>&lt;5.7$^a$</td>
<td>1.25(0.02)</td>
<td>1.34(0.08)</td>
</tr>
<tr>
<td>B2200+420</td>
<td>3.7(1.3)</td>
<td>2.58(0.03)</td>
<td>5.73(0.21)</td>
</tr>
</tbody>
</table>

1 Limits are $2\sigma$.

2 Entry is $59 \pm 2 N($H$^{13}$CO$^+$).

### Table 3. Column density ratios$^a$.

<table>
<thead>
<tr>
<th>Source</th>
<th>$N($HOC$^+$/N($HCO^+$)</th>
<th>$N($HOC$^+$/N($OH$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B0355+508</td>
<td>72(9)</td>
<td>6.1(0.7) $\times 10^{-4}$</td>
</tr>
<tr>
<td>B0415+379</td>
<td>117(12)</td>
<td>2.6(0.3) $\times 10^{-4}$</td>
</tr>
<tr>
<td>B0528+134</td>
<td>&gt;30</td>
<td>&lt;1.6 $\times 10^{-3}$</td>
</tr>
<tr>
<td>B1730-130</td>
<td>&gt;18</td>
<td>&lt;4.3 $\times 10^{-3}$</td>
</tr>
<tr>
<td>B2200+420</td>
<td>70(24)</td>
<td>6.5(2.1) $\times 10^{-4}$</td>
</tr>
</tbody>
</table>

1 Limits are $2\sigma$. 

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Fig. 1. HOC$^+$ and H$^{13}$CO$^+$ observed toward B0415+379 (aka 3C 111).
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3.2. Expected abundance of HOC$^+$

The most obvious path to HOC$^+$ in diffuse gas is via the reaction C$^+$ + H$_2$O → HOC$^+$ + H, which proceeds with a rate constant $1.8 \times 10^{-9}$ cm$^3$ s$^{-1}$ in the UMIST reaction database: these two reactants also produce HCO$^+$ at half the rate, but this is not an important source of HOC$^+$. We parametrize the unknown abundance of water relative to OH and assume that all free carbon is once ionized in a fully molecular gas having a free carbon abundance $N$(C)/$N$(H) = $1.4 \times 10^{-4}$ (Savage & Sembach 1996). In these terms, the volume formation rate of HOC$^+$ from the reaction of C$^+$ and water can be expressed as $\frac{dn(\text{HOC}^+)}{dt} = 5.04 \times 10^{-20} n(\text{H}_2)^2 X(\text{H}_2\text{O})/X(\text{OH})$ cm$^{-3}$ s$^{-1}$.

Both HCO$^+$ and HOC$^+$ are formed from the reaction of H$_3^+$ + CO → (HOC$^+$ or HCO$^+$) + H$_2$. The HCO$^+$/HOC$^+$ branching ratio is typically taken as 20 (Illies et al. 1982) or more (the current UMIST reaction database has a branching ratio of 62). The volume formation rate from this reaction may be written as $\frac{dn(\text{HOC}^+)}{dt} = 2.7 \times 10^{-22} n(\text{H}_2)^2 (X(\text{H}_3^+)) X(\text{CO})/10^{12}$ cm$^{-3}$ s$^{-1}$. The relative abundance of CO is known along these lines of sight to be 1–5% of the free carbon abundance (Liszt & Lucas 1998), i.e. 1–5 $\times$ 10$^{-6}$. The observed amounts of H$_3^+$, while surprisingly large (McCall et al. 2003) nevertheless still yield $X(\text{H}_3^+) < 10^{-6}$. Thus the only circumstance in which the reaction of H$_3^+$ and CO could be competitive in forming HOC$^+$ would be for very small water abundances, in which case neither formation route would suffice to form the observed HOC$^+$.

HOC$^+$ is likely destroyed by two processes which proceed with very nearly equal rates in diffuse gas. The HOC$^+$ isomer lies some 17 000 K higher in energy compared to HCO$^+$ and interconversion by molecular hydrogen (HOC$^+$ + H$_2$ → HOC$^+$ + H$_2$) has recently been shown to proceed with a rate constant $k_1 = 4 \times 10^{-10}$ cm$^3$ s$^{-1}$ nearly independent of temperature between 25 K and 300 K (Smith et al. 2002). This is 70% smaller than in the UMIST database, enhancing the prospects for reproducing $X$(HOC$^+$). The destruction rate for isomerization can be written as $\frac{dn(\text{HOC}^+)}{dt} = -4 \times 10^{-10} n(\text{H}_2) n(\text{HOC}^+) cm^{-3}$ s$^{-1}$.

Dissociative recombination, HOC$^+$ + e$^-$ → CO + H, proceeds with a rate constant $1.1 \times 10^{-9}(30/$T$_K$) cm$^3$ s$^{-1}$. Assuming that free carbon is once-ionized and expressing the electron fraction in terms of the ratio of free carbon to H$_2$ expected for a fully molecular diffuse gas (i.e. $N$(C)/$N$(H$_2$) = $2 \times 1.4 \times 10^{-4}$ (Savage & Sembach 1996)). The destruction rate from recombination can be written as $\frac{dn(\text{HOC}^+)}{dt} = -3.1 \times 10^{-10} n(\text{H}_2)n(\text{HOC}^+)(30/$T$_K$) (X(e)/X(C$^+$)) cm$^{-3}$ s$^{-1}$.

The expected electron abundance is a small multiple of the free carbon abundance (Liszt 2003) so dissociative recombination and interconversion probably compete quite effectively to destroy HOC$^+$.

The net destruction rate from both processes is of order $-10^{-9}$ n(He) cm$^{-3}$ s$^{-1}$ and the expected abundance of HOC$^+$ is $X$($\text{HOC}^+$) = $5 \times 10^{-11}$ X(He) X(OH) or $X$($\text{HOC}^+$)/X(OH) = $5 \times 10^{-4}$ X(He) X(OH). The observed amounts of HOC$^+$ can be made in diffuse gas if the abundance of water is comparable to that of OH. Limits on the OH/water ratio in diffuse clouds $N$(OH)/$N$(H$_2$O) $> 2$ – 3 were derived by Van Dishoeck & Black (1986) toward $\zeta$ Oph and $\zeta$ Per.

Given the weakness of the limits on the water abundance in diffuse clouds, straightforward gas-phase chemistry fails only weakly to produce the observed amount of HOC$^+$. This situation can be compared to the case of HCO$^+$ where the thermal reaction of C$^+$ upon the known quantities of OH falls 30–50 times short of producing the observed values of $N$(HCO$^+$) via the reaction chain C$^+$ + OH → CO$^+$ + H, H$_2$ + CO$^+$ → HOC$^+$ + H, HOC$^+$ + e$^-$ → CO + H. At thermal rates, this sequence of reactions (which predicts an HCO$^+$/OH ratio dependent only on temperature) produces $N$(HCO$^+$) scarcely larger than the observed $N$(HOC$^+$). The predicted abundance of HCO$^+$ could be increased in quiescent gas by hypothesizing that it recombines slowly, but only at the cost of understanding CO (Liszt & Lucas 2000).

Clearly, knowledge of the abundance of water is the limiting factor in assessment of the diffuse cloud chemistry of HOC$^+$.

3.3. Comparison with HCO$^+$ and HOC$^+$ in dense dark and bright gas

The work presented here is the first evidence of HOC$^+$ in diffuse gas, but even the case for its existence in the dense molecular ISM was marginal before the pioneering work of Aponi and his collaborators during a brief period in the mid-1990’s while the NRAO 12 m telescope was still operating.

Fig. 2. HOC$^+$ and HCO$^+$ observed toward B2200+420 (aka BL Lac).

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1 available at http://www.rate99.co.uk/
To summarize the prior results we note that the HCO⁺/HOC⁺ ratio in dense dark gas is typically as large as 1000–9000 (Ziurys & Apponi 1995; Apponi & Ziurys 1997), but is found to be much smaller (50–400) in dense PDR like the Orion Bar (Apponi et al. 1999) and/or NGC 7023 (Fuente et al. 2003). In these PDR the relative abundance of HOC⁺ is \( X(\text{HOC}^+) = 1\times10^{-11} \), quite similar to the diffuse clouds studied here.

The difference between the HCO⁺/HOC⁺ ratios seen in diffuse and dark dense gas is largely the result of the low ionization fraction in the latter. Whereas HOC⁺ is destroyed only slightly faster than HCO⁺ in diffuse gas, the isomerization reaction undergone by HOC⁺ is much faster than recombination when the electron fraction is \( 10^{-7} \) instead of \( 10^{-4} \). In dense dark gas HCO⁺ and HCO⁺ are formed by \( \text{H}_2^+ + \text{CO} \rightarrow \text{H}_2 + (\text{HCO}^+ \text{ or } \text{HOC}^+) \) with a branching ratio 20–60:1 in favor of HCO⁺ and this ratio is increased by another factor of 100 by the unequal destruction rates. Under favorable conditions, measured HCO⁺/HOC⁺ ratios could be good diagnostics of the ionization fraction in dense gas.

Reasoning by analogy from the discussion of diffuse and dark gas, we infer that the relatively low HCO⁺/HOC⁺ ratios in dense PDR also reflect the relative weakness of isomerization. In cases where isomerization is weak both isomers are destroyed by recombination at roughly equal rates, and their abundance ratio is more directly indicative of the relative rates at which the two isomers are formed.

4. Summary

We have added HOC⁺ to the roster of polyatomic molecules known to exist in diffuse clouds. The HCO⁺/HOC⁺ ratio, 70–120, is much smaller than in dense dark clouds but quite comparable to the most extreme values seen in dense PDR. The inferred relative abundance \( X(\text{HOC}^+) = 3\times6 \times 10^{-11} \), similar to that seen in dense PDR, can be synthesized at thermal rates in quiescent diffuse gas if the water/OH ratio is of order unity. This is a mild discrepancy, much less severe than is inferred for HCO⁺, as the water/OH ratio is currently known only to be less than 0.3–0.5.

Several authors have recently noted that hydrocarbons like \( \text{C}_3\text{H}_2 \) and \( \text{C}_2\text{H} \) have unexpectedly high abundances in dense PDR (Fosse et al. 2002; Fuente et al. 2003; Teyssier et al. 2004), very much as has been observed in diffuse clouds (Lucas & Liszt 2000). The HCO⁺/HOC⁺ ratios found here in diffuse clouds, which are PDR of lower density and presumably weaker illumination (hence similar ionization rates per \( \text{H}_2 \)), constitute another example of chemical similarities between dense and diffuse PDR. It will be interesting to see how much more complete a parallel can be drawn between the two regimes.

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References

Fosse, D., Teyssier, D., Gerin, M., et al. 2002, in Chemistry as a Diagnostic of Star Formation, 104