Hydrides in Young Stellar Objects: Radiation tracers in a protostar-disk-outflow system*


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ABSTRACT

Context. Hydrides of the most abundant heavier elements are fundamental molecules in cosmic chemistry. Some of them trace gas irradiated by UV or X-rays.

Aims. We explore the abundances of major hydrides in W3 IRS5, a prototypical region of high-mass star formation.

Methods. W3 IRS5 was observed by HIIF on the Herschel Space Observatory with deep integration (≥ 2500 s) in 8 spectral regions.

Results. The target lines including CH, NH, H₂O⁺, and the new molecules SH⁺, H₂O⁺, and OH⁺ are detected. The H₂O⁺ and OH⁺ J = 1 − 0 lines are found mostly in absorption, but also appear to exhibit weak emission (P-Cyg-like). Emission requires high density, thus originates most likely near the protostar. This is corroborated by the absence of line shifts relative to the young stellar object (YSO). In addition, H₂O⁺ and OH⁺ also contain strong absorption components at a velocity shifted relative to W3 IRS5, which are attributed to foreground clouds.

Conclusions. The molecular column densities derived from observations correlate well with the predictions of a model that assumes the main emission region is in outflow walls, heated and irradiated by protostellar UV radiation.

Key words. Stars: formation - stars: high mass - ISM: molecules - ISM: individual objects: W3 IRS5 - Line: identification - Ultraviolet: ISM

1. Introduction

In interstellar clouds, chemical reactions with hydrogen molecules lead to an elementary class of molecules that represent key species in the chemical evolution to larger molecules. These fundamental molecules, known as hydrides, include OH, CH, NH, H₂O, and their ions, OH⁺, CH⁺, NH⁺, SH⁺, H₂O⁺, and H₂O⁺. The combination of hydrogen atoms with a heavier atom causes large dipole moments and large rotation constants, particularly in diatomic hydrides. This widely separates the excitation levels. Only low-J lines are excited at temperatures relevant to star and planet formation. These lines have now become observable with the Herschel Space Observatory (Pilbratt et al. 2010).

Many hydrides have a high activation energy in their formation paths. However, if high-energy photons - far UV (FUV) or X-rays - interact with the molecular gas and heat it, hydrides and particularly their ions are greatly enhanced in abundance (e.g., Hollenbach & Tielens 1999). Ionized hydrides are chemically even more active and can substantially drive chemical evolution. Many of the above hydrides have been observed from the ground in absorption in diffuse interstellar clouds (e.g. Swings & Rosenfeld 1937; Menten et al. 2010; Wyrowski et al. 2010a).

Here we focus on hydrides in dense star-forming regions surrounding protostars. We report first results of the ‘Radiation Diagnostics’ project within the Herschel key program ‘Water In Star-forming regions with Herschel’ (WISH, van Dishoeck et al. 2010). The ‘Radiation Diagnostics’ subprogram aims to trace high-energy radiation through the most abundant hydrides and their ions and relate them to the chemical network of water. Preparing for this project, Stäuber et al. (2005, 2006) showed that hydrides may be enhanced in star-forming regions affected by far UV and/or X-ray irradiation. In a second step, optimal
lines were selected using cataloged data in the available databases or computed from measured molecular constants (Bruderer 2006).

Spherically symmetric models including FUV and X-rays have found that most hydrides would be too weak for Herschel detection. However, evidence of extended high-energy irradiation (Doty et al. 2004) and in particular the observation of CO$^+$, with an abundance four orders of magnitude larger than predicted, in W3 IRS5 and other YSOs with the JCMT (Stäuber et al. 2007) raised the expectation that the effects of irradiation may be more dramatic in asymmetric reality. Bruderer et al. (2009) modeled and interpreted the CO$^+$ anomaly as FUV radiation originating in the YSOs and irradiating the walls of outflow cavities.

Based on these predictions, the ‘Radiation Diagnostics’ observations started with deep integrations of a large number of hydrides that was to be followed by a survey of a few species in many sources of various ages and masses. Here we present exploratory observations towards W3 IRS5, a species in many sources of various ages and masses. Here the YSOs and irradiating the walls of outflow cavities.

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2. Observations

The Heterodyne Instrument for the Far Infrared (HIFI, de Graauw et al. 2010) on Herschel observed W3 IRS5 between 1 and 8 March 2010 in the Science Demonstration Phase in eight 4 GHz frequency bands for about 2500 s each. One of them includes [C II] at 1090.5369 GHz. We used the Wide Band Spectrometer which has a spectral resolution of 1.1 MHz, and HIPE 3.0 for pipeline and data analysis. The data were taken by double beam switching (DBS), the high-frequency (HEB) bands in fast DBS mode. The off-source position was at a distance of 3 arcmin in the NE and SW without remarkable IR sources.

The current accuracy of the velocity calibration is estimated to be better than 2 km s$^{-1}$. The antenna temperature was converted to main beam temperature, using pre-flight antenna efficiencies. After visual inspection and defringing, the V polarization was shifted linearly in flux to match the H polarization. The two polarizations were then added. A second observation of equal length was made using a local oscillator frequency shifted by 10 km s$^{-1}$. The two data sets were plotted in both upper and lower sideband presentation. If a line matched velocities in one sideband and was double with 20 km s$^{-1}$ separation in the other, the frequency of the former was assumed. All lines of interest could be attributed to a sideband without ambiguity. The continuum was divided by 2 for double sideband observations, assuming that it is the same in both sidebands, of equal sensitivity.

3. Results

Most lines (except H$_3$O$^+$) are split by fine or hyperfine interaction as indicated in Fig. 1. Table 1 lists the observed lines (strongest only for multiples) and summarizes the quantitative observational results. Molecules here detected for the first time in star-forming regions include H$_2$O$^+$, OH$^+$, and SH$^-$. Having the most prominent line near the H$_2$O para ground-state line at 1113.3 GHz, H$_2$O$^+$ is serendipitously detected in many Herschel observations. H$_2$O$^+$ and OH$^+$ are detected in absorption by the interstellar medium (Bruderer et al. 2010b; Ceccarelli et al. 2010; Gerin et al. 2010; Palgarone et al. 2010; Neufeld et al. 2010; Ossenkopf et al. 2010; Schiike et al. 2010), but also near the systemic velocity of other high-mass YSOs (Wyrowski et al. 2010b).

Several lines are found in absorption, indicated in Table 1 by negative peak values relative to the continuum. All lines predominantly in absorption (NH, OH$^+$, H$_2$O$^+$) originate in molecules in the ground state with a J = 1 level energy exceeding 47 K. Lines of CH and SH$^+$, observed in emission, are transitions from a J = 1 level of energy less than 26 K. We do not detect SH and NH$^+$ above the 5σ limit.

The observed line shifts are within ±2 km s$^{-1}$ of the YSO, thus within the accuracy of the frequency calibration, spectral resolution, and molecular data. The only deviations in Table 1 are OH$^+$ and H$_2$O$^+$. Their strongest peaks, both in absorption, are shifted by 34 - 39 km s$^{-1}$ relative to the YSO, similar to an absorption feature in [C II].

Figure 2 illustrates a possible explanation of the exceptional line shifts of OH$^+$ and H$_2$O$^+$. In both cases, a theoretically strong hyperfine structure line corresponds to a smaller absorption dip within less than 5 km s$^{-1}$ of the YSO. For OH$^+$, the low-frequency line (a doublet) is blended with the high-frequency line of another, red-shifted OH$^+$ component. We thus interpret the spectra of OH$^+$ and H$_2$O$^+$ as the superimposition of a component originating in the star-forming region and another component at lower frequency related to an outflow or the foreground interstellar medium near zero velocity in the LSR.

The line widths in Table 1 are between 4 and 7 km s$^{-1}$. Measured at the component with the systemic velocity of the YSO. Two lines attract attention: (i) OH has an absorption feature near the YSO velocity. This absorption reduces the peak flux and widens the line profile at half power. (ii) The line width of NH is extremely narrow in all hyperfine structure components.

The lines are generally symmetric. A remarkable exception is OH$^+$, showing blue tails. We note that the shifted lines of OH$^+$ (unblended line) and H$_2$O$^+$ are 17.7 and 11.1 km s$^{-1}$ wide, respectively, indicating that they have a different origin from the component at zero systemic velocity.

Several lines in Fig. 1 show slightly blue-shifted absorption and red-shifted emission, thus a P-Cyg-like behavior. Figure 2 compares them with the [C II] line. CH (strongest component) exhibits similar absorption to [C II], their peak absorptions being at -1.6 km s$^{-1}$ and -1.7 km s$^{-1}$, respectively, relative to the YSO motion. OH$^+$ and H$_2$O$^+$ differ, having absorption peaks at -2.5 and -4.8 km s$^{-1}$, respectively, and emission peaks at 4.2 and 3.5 km s$^{-1}$.

Column densities are given in Table 1 for the upper energy level of transitions in emission, and for the lower
Molecular data are taken from: \(^a\) CDMS (Müller et al. 2001), \(^b\)Mürtz et al. (1998), \(^c\)Bruderer (2006), \(^d\)Hübers et al. (2009), \(^e\)JPL catalogue

Table 1. Frequency, upper level energy, and Einstein coefficient of molecules and lines observed by Herschel/HIFI towards W3 IRS5. The numbers in parentheses give the decimal power. Negative peak fluxes signify background minus line temperature of lines in absorption. Line widths refer to the FWHP value of the most intense line peak or absorption and its line shift to the systemic velocity of -38.4 km s\(^{-1}\). Non-detected peak fluxes are 5 \(^a\) upper limits at 1 km s\(^{-1}\) resolution, and non-detected line fluxes assume a 5 km s\(^{-1}\) line width. The column densities refer to the upper (emission) or lower (absorption) state.

Fig. 1. Spectral lines observed towards W3 IRS5 in main beam temperature. The positions of the theoretical fine/hyperfine structure lines are shown shifted by -38.4 km s\(^{-1}\), the systemic motion of the YSO. The length of the vertical bars indicates the theoretical intensities in arbitrary scale. The frequency binning is 1 km s\(^{-1}\).
The ground-state column densities of the component moving with the YSO are $8.1 \times 10^{11} \text{cm}^{-2}$ for H$_2$O$^+$, and $9.7 \times 10^{12} \text{cm}^{-2}$ for OH$^+$. The values for the red-shifted components are $3.8 \times 10^{12} \text{cm}^{-2}$ and $6.1 \times 10^{13} \text{cm}^{-2}$, respectively. Both components yield larger OH$^+$/H$_2$O$^+$ ratios than the other observations reported in this volume.

The measured line widths are generally small ($< 7 \text{ km s}^{-1}$) and show no anisotropies. We thus find no evidence of shocks except possibly in OH$^+$, which needs to be studied in combination with shock tracers.

In Fig. 3, column densities are derived from integrated line fluxes neglecting re-emission or reabsorption of the final state. The derivation is based on Table 1, except for OH$^+$ and H$_2$O$^+$ where only the unshifted component attributed to the YSO is used. For H$_2$O$^+$, the value of the $J = 4$ level as well as the one from the rotational diagram, integrated over all levels, are shown. The column densities are compared with abundances predicted by Bruderer et al. (2010a) in a two-dimensional ‘standard’ YSO model used here as a template, assuming UV and X-ray irradiation by a central high-mass YSO. It enhances the abundance of diatomic hydrides in the outflow walls by many orders of magnitude, such that the beam-averaged abundance is significantly changed. It is averaged over a radius of 20 000 AU, or 10.9$''$ at the distance of W3 IRS5.

The similarity of observations and model abundances in Fig. 3 support the scenario of hydride enrichment in outflow walls heated and irradiated by protostellar far UV.

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References

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**Appendix A:**

The rotational diagram in Fig. A1 is complemented with ground-based data for the frequency range 300 - 400 GHz observed at the CSO with comparable beam size by Phillips, van Dishoeck & Keene (1992). The non-detection at 307 MHz is surprising, but possibly an effect of optical depth. The data in Fig. A1, except 307 GHz, are well fitted by a single rotational temperature of 239 K, suggesting that the observed levels are populated according to an exponential distribution. The derived temperature and column density infer an optical depth of $\tau < 0.1$ for all lines except at 307 GHz. The fitted line (dashed) corresponds to a column density of $8.5(\pm2) \times 10^{13}$ cm$^{-2}$, consistent with the value derived by Phillips, van Dishoeck & Keene (1992). This leads to a beam averaged $\text{H}_3\text{O}^+$ abundance of $4.2(\pm1) \times 10^{-10}$ relative to H, to be compared with the theoretical value of $4 \times 10^{-10}$ reported by Bruderer et al. (2010a).

**Fig. A.1.** Rotational diagram of $\text{H}_3\text{O}^+$. Numbers indicate the frequencies in GHz of the observed lines.