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SPIRE spectroscopy of the prototypical Orion Bar photodissociation region*


(Affiliations are available in the online edition)

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ABSTRACT

Aims. We present observations of the Orion Bar photodissociation region (PDR) obtained with the SPIRE instrument on-board Herschel.

Methods. We obtained SPIRE Fourier-transform spectrometer (FTS) sparse sampled maps of the Orion bar.

Results. The FTS wavelength coverage and sensitivity allow us to detect a wealth of rotational lines of CO (and its isotopologues), fine structure lines of C and N$^+$, and emission lines from radicals and molecules such as CH, CH$_2$O or H$_2$S. For species detected from the ground, our estimates of the column densities agree with previously published values. The comparison between $^{12}$CO and $^{13}$CO maps shows particularly the effects of optical depth and excitation in the molecular cloud. The distribution of the $^{13}$CO and $^{12}$CO lines with upper energy levels indicates the presence of warm (~100–150 K) CO. This warm CO component is a significant fraction of the total molecular gas, confirming previous ground based studies.

Key words. infrared: ISM – ISM: lines and bands – ISM: molecules – evolution – submillimeter: ISM – ISM: general

1. Introduction

The Orion Bar located between the Orion molecular cloud and the HII region surrounding the Trapezium stars is one of the best-studied photodissociation regions (PDRs) in the Galaxy. Much of the emission from massive star-forming regions will originate from these interfaces, which are responsible for reprocessing the energy output from stars and reemitting this energy at infrared-millimetre wavelengths including a rich mixture of gas lines (i.e., Hollenbach & Tielens 1999). Visible-ultraviolet stellar radiation governs the chemical and thermal state of the gas in these regions. The impinging radiation field on the Bar is $\chi = (0.5-2.5) \times 10^5 \chi_0$ (Tielens & Hollenbach 1985; Marconi et al. 1998), where $\chi_0$ is the Solar neighbourhood far-UV interstellar radiation field as given by Draine (1978). The UV field varies as a function of depth within the cloud, providing a unique opportunity to study how the dust populations and the molecular content evolve with the excitation and physical conditions. This is important for the evolution of the cloud and its associated star formation.

The ESA Herschel Space Observatory (Pilbratt et al. 2010) offers a unique opportunity to observe continuously between 55 and 672 $\mu$m. This range includes most of the PDR gas lines and dust components emission and will provide a fundamental step in our understanding of the evolution of the interstellar matter. Here, we present a first analysis of Fourier-transform spectrometer (FTS) observations of the Orion Bar obtained with the SPIRE instrument (Griffin et al. 2010) on-board Herschel.

2. Observations with the FTS

The SPIRE FTS simultaneously measures the source spectrum across two wavebands: spectrometer long wavelength (SLW), covering 14.9–33.0 cm$^{-1}$ (303–671 $\mu$m) and spectrometer short wavelength (SSW) covering 32.0–51.5 cm$^{-1}$ (194–313 $\mu$m). Each band is imaged with a hexagonal bolometer array with pixel spacing of approximately twice the beam-width. The FWHM beam-widths of the SLW and SSW arrays vary between 29–42" and 17–21" respectively. The source spectrum, including the continuum, is obtained by taking the inverse transform of the observed interferogram. For more details on the FTS calibration and data reduction procedures, the reader is referred to the article by Swinyard et al. (2010).

Our observations are part of the “Evolution of Interstellar dust” key program of the SPIRE consortium (Abergel et al. 2010). The Orion Bar was observed with a single pointing in the high-resolution mode of the SPIRE FTS on 2009 September 13 (Herschel observation ID, 1342183819). Two scan repetitions were observed which gave an on-source integration time of 266.4 seconds. The pointing centre was at a right ascension and declination (J2000) of 05 h 35 m 22.83 s and -05°24′57.67″ (see Fig. 1). The unapodized spectral resolution was 0.04 cm$^{-1}$ (1.2 GHz). After apodization (using extended Norton-Beer function 1.5; Naylor & Tahir 2007) the FWHM of the resulting instrument line shape is 0.0724 cm$^{-1}$ (2.17 GHz).

* Herschel is an ESA space observatory with science instruments provided by European-led Principal Investigator consortia and with important participation from NASA.
While unapodized FTS spectra provide the highest spectral resolution, the instrument line shape, which for an ideal FTS is the classical sinc function, is characterized by relatively large secondary oscillations with negative lobes. An iterative spectral-line fitting routine was developed to extract line parameters from unapodized FTS spectra (Jones et al. 2009). This algorithm fits a continuum (either a low order polynomial or a blackbody variant) and a series of lines with the Levenberg-Marquardt least-squares method. The fitting procedure weights the spectral intensity at a given frequency of an averaged spectrum by the statistical uncertainty at that frequency. The fitting routine returns the line centres, intensities, and line widths, together with their associated errors.

3. Results

3.1. Detected gas lines

The averaged apodized FTS spectra over the three SLW/SSW detectors aligned on the Bar and corrected for obliquity effects1 are presented in Figs. 1 and 2. The FTS wavelength coverage allows us to detect a wealth of rotational lines of CO (and its isotopologues), fine structure lines of C and N, and emission lines from several radicals and molecules. The expected line positions for detected species are marked in Figs. 1 and 2. The 13CO transitions, which appear as the bright narrow lines, are here seen for the first time together from J = 4–3 to 13–12 in a single spectrum. The 13CO lines are clearly detected from J = 5–4 to 13–12. Most of the C18O lines are visible but blended with the 13CO lines; some C17O lines are detected. One emission line at about 359 μm lies at the position of the fundamental rotational transition of CH+ (Naylor et al. 2010). This detection can be related to the observation of the CH lambda doublet transitions at about 556.5 μm and 560.7 μm, although it is possibly blended with an HCO+ J = 6 → 5 line. The ortho-H2O 1_00 → 1_01 line at ~538 and para-H2 2_11 → 2_02 at 398 μm are clearly detected. The ~269 μm para-H2O 1_11 → 0_00 line was also detected, but the signal-to-noise ratio is low. Some other H2O lines may be blended. The H2S 2_10 → 1_01 line at ~407 μm is detected, while other fainter H2S lines at shorter wavelengths are only marginally detected. Some features related to the emission of HCO+, HCN, CN and C2H are observed as expected (e.g., Hogerheijde et al. 1995; Simon et al. 1997; Young Owl et al. 2000; Teysier et al. 2004; van Der Wiel et al. 2009), but to help distinguish the spectral confusion for fainter lines or unresolved k-ladder transitions from species such as methanol, the actual signal-to-noise ratio will be improved as the SPIRE FTS response is better understood and, scheduled deeper observations will also help.

3.2. Mapping gas lines

Figure 3 presents sparse sampled2 maps of nearly the complete CO and 13CO band measured. Off-axis calibrations are not guaranteed because both detector arrays have not yet been fully characterised. The comparison between these maps shows the effects of optical depth and excitation in the molecular cloud particularly well. The emission of the less abundant 13CO isotopologue

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1 The obliquity effect is important at the highest frequencies, where a significant error in the line position is introduced.

2 The present FTS science demonstration phase observations sparsely samples the field of view and do not allow us to present fully sampled maps.
probes the denser shielded regions, while the $^{12}$CO optically thick emission likely comes from the less dense surface layers (Lis et al. 1998). The highest rotational lines, which are very sensitive to both gas densities and temperatures, show strong and peaked emission on the Bar, while they are not visible in the off Bar positions. Emission lines of species such as C, N$^+$ or CH$^+$ show spatially extended emission.

3.3. Molecular column densities

We used the observed line intensities and the CASSIS software\(^3\) to estimate the beam-averaged molecular column densities. We list in Table 1 column densities estimated for a volume density of $10^5$ cm$^{-3}$ as applicable to the extended molecular gas in the Bar (Hogerheijde et al. 1995) and in the high-density limit, because some of the line emission may originate from dense clumps\(^4\). We adopt the mean molecular gas temperature towards the Bar: $T = 85$ K across the bar, consistent with many observed transitions from the ground. However, the $^{12}$CO $J = 12-11$ and $J = 13-12$ at 209 $\mu$m lines. Scales are in $10^5$ erg s$^{-1}$ cm$^{-2}$ sr$^{-1}$. Contour levels are drawn at 10, 20, ..., 90% of the maximum intensity.

Our values for the column densities agree for species detected from the ground with previously published values to within a factor of 2--3: Hogerheijde et al. (1995) for C$^{18}$O and HCO$^+$; Johnstone et al. (2003) for C$^{17}$O; and Leurini et al. (2006) for H$_2$S. Beam dilution effects could introduce a significant factor. To convert the observed C$^{18}$O $J = 8-7$ and C$^{17}$O $J = 8-7$ line intensities to a total H$_2$ column density, we assume isotopic ratios $^{18}$O/$^{16}$O $= 560$, $^{17}$O/$^{16}$O $= 1800$ (Wilson & Rood 1994) and a relative CO abundance of H$_2$ of $1.1 \times 10^{-4}$ as applicable for the Orion Bar PDR (Johnstone et al. 2003). We find $N$(H$_2$) $\sim 9 \times 10^{22}$ cm$^{-2}$ assuming $T$ $= 85$ K, which implies the following molecular abundances on the Bar: $x$(ortho-H$_2$O) $\leq 3.3^{+3.3}_{-1.7} \times 10^{-7}$, $x$(para-H$_2$O) $\leq 5^{+11.7}_{-3.1} \times 10^{-7}$, $x$(HCO$^+$) $\leq 3.9^{+8.3}_{-2.1} \times 10^{-9}$, $x$(CH$^+$) $= 7.5^{+12.6}_{-0.7} \times 10^{-11}$, $x$(H$_2$S) $= 3.4^{+2.3}_{-1.1} \times 10^{-10}$.

H$_2$O is extremely sensitive to the local physical conditions in molecular clouds: close to the surface, molecules are photodissociated, while deeper into the cloud molecules freeze onto grain surfaces (i.e., Hollenbach et al. 2009). Desorption of ices (Westley et al. 1995; Seperuelo Duarte et al. 2009) could supply gas-phase species. The high abundances of sulphur species remain an interesting puzzle for interstellar chemistry (i.e., Goicoechea et al. 2006). The observed abundance of species such as H$_2$S are difficult to interpret in models. H$_2$S results from a mixed chemistry involving gas-phase reactions and grain-related processes.

3.4. CO excitation

Figure 4 shows the distribution of the $^{12}$CO and $^{13}$CO line intensities as a function of the upper energy levels. The observed intensities of the optically thick $^{12}$CO lines provide an estimate of the temperature of about $85$ K across the bar, consistent with many observed transitions from the ground. However, the $^{12}$CO $J = 12-11$ and $J = 13-12$ transitions do not agree with that temperature (intensities higher by a factor of 2 and 4 respectively), which is consistent with ground data of the $J = 14-13$ transition dominated by warmer CO (Stacey et al. 1993; Tauber et al. 1994). Similarly, the distribution of the $^{13}$CO lines cannot be described by a single temperature. A dense and warm ($\sim 100–150$ K) component with a significant column density (about 15% of the total column) is required to fit the observed line intensities with $J_{\text{LO}} \geq 9$. We find that the $^{13}$CO lines become optically thin for $J_{\text{LO}} \geq 9$, while all the $^{12}$CO lines are optically thick.

We computed PDR models for the Bar with an updated version of the Meudon PDR code described in Le Petit et al. (2006) solving in an iterative way the chemical and thermal balances at each point of the cloud. Adopting $\chi = 10^7 \chi_0$, a constant gas density of $10^5$ cm$^{-3}$ or constant thermal gas pressure of $8 \times 10^7$ K cm$^{-3}$ (Allers et al. 2005), the models predict a gas temperature of about $50–80$ K for the CO emitting gas. Consequently, it cannot explain the observed warm CO. One explanation is that the warm CO originates from dense clumps at the PDR surface (Tauber et al. 1994). An alternative solution are additional heating mechanisms for the interior, like shocks (or turbulence) or cosmic ray heating (Pellegreni et al. 2009). Out-of-equilibrium effects such as advection of molecular gas from the shielded cloud interior to the warm surface could also enhance the column densities of warm CO. Progress is expected from the spectroscopy of additional cooling lines, to be obtained from Herschel PACS and HIFI instruments. In particular, observations of gas cooling lines at high spectral resolution with HIFI will provide missing information about the gas velocity within

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\(^3\) Based on analysis carried out with the CASSIS software and CDMS, JPL spectroscopic databases and RADEX (van Der Tak et al. 2007) molecular databases. CASSIS has been developed by CESR-UPS/CNRS (http://cassis.cesr.fr).

\(^4\) The clumpiness of the PDR inferred by Hogerheijde et al. (1995) was confirmed by interferometric data of Young Owl et al. (2000); Lis & Schilke (2003). Clump densities up to $10^6$ cm$^{-3}$ were derived by Lis & Schilke (2003), while the density of the interclump medium should fall between a few $10^5$ cm$^{-3}$ (Young Owl et al. 2000) and $2 \times 10^5$ cm$^{-3}$ (Simon et al. 1997).
Table 1. Beam-averaged molecular column densities.

<table>
<thead>
<tr>
<th>Species</th>
<th>Transition</th>
<th>Wavelength (microns)</th>
<th>E_J (K)</th>
<th>Intensity (10^{-6} erg s^{-1} cm^{-2} sr^{-1})</th>
<th>Column density (10^{15} cm^{-2})</th>
</tr>
</thead>
<tbody>
<tr>
<td>C^{18}O</td>
<td>J = 7 → 6</td>
<td>390.2</td>
<td>147.5</td>
<td>2.6[1.1]</td>
<td>7.7[24.3, 8]</td>
</tr>
<tr>
<td>C^{18}O</td>
<td>J = 8 → 7</td>
<td>341.5</td>
<td>189.6</td>
<td>3.3[1.1]</td>
<td>15[72, 5.5]</td>
</tr>
<tr>
<td>C^{18}O</td>
<td>J = 9 → 8</td>
<td>303.6</td>
<td>237.0</td>
<td>1[1.3]</td>
<td>9[720, 25]</td>
</tr>
<tr>
<td>C^{13}O</td>
<td>J = 7 → 8</td>
<td>333.7</td>
<td>194.1</td>
<td>1.4[1.1]</td>
<td>6.6[33, 2.3]</td>
</tr>
<tr>
<td>CH^+</td>
<td>J = 1 → 0</td>
<td>359.0</td>
<td>40.1</td>
<td>6.7[1.1]</td>
<td>0.006[0.0058, 0.0088]</td>
</tr>
<tr>
<td>ortho-H_2O</td>
<td>J_{0} → 1_{0}</td>
<td>538.3</td>
<td>61.0</td>
<td>2.6[1.1]</td>
<td>≤30[60, 15]</td>
</tr>
<tr>
<td>para-H_2O</td>
<td>J_{2} → 2_{0}</td>
<td>398.6</td>
<td>136.9</td>
<td>2.0[1.1]</td>
<td>≤45[150, 17]</td>
</tr>
<tr>
<td>HCO^+</td>
<td>J = 8 → 7</td>
<td>420.3</td>
<td>154.1</td>
<td>0.7[1.1]</td>
<td>≤0.35[1.0, 16]</td>
</tr>
<tr>
<td>H_2S</td>
<td>J_{2} → 1_{0}</td>
<td>407.5</td>
<td>55.1</td>
<td>1.4[1.1]</td>
<td>0.031[0.023, 0.052]</td>
</tr>
</tbody>
</table>

Notes. (a) Intensities with uncertainty in between brackets. (b) Beam-averaged column densities using RADEX with n = 10^5 cm^{-3} and T = 85 K, with estimations for 50 and 150 K in between brackets. (c) Beam-averaged column densities in the high-density (LTE) limit for 85 K, with estimations for 50 and 150 K in between brackets. (d) Note that the density (or temperature) could be higher than assumed, which would decrease the column density, and the para-H_2O 2_{1} → 2_{0} line could be affected by IR pumping. (e) We used the HCO^+ J = 8 → 7 line because the HCO^+ J = 7 → 6 line is possibly blended with an HCl J = 1 → 0 line. Column densities from ^{12}CO and ^{13}CO lines are uncertain due to high optical depth and are not listed.

Fig. 4. Distribution of the ^{12}CO (black) and ^{13}CO (red) line intensities as a function of the upper energy levels. The squares show the FTS data complemented by ground-based measurements shown by triangles (Stacey et al. 1993; Tauber et al. 1994; White & Sandell 1995; Hogerheijde et al. 1995; van Der Wiel et al. 2009). Error bars are small compared to the symbol size. The arrows indicate upper limits. The empty circles show the RADEX calculation for T = 85 K, n = 10^5 cm^{-3}, N({^{12}CO}) = 10^{18} cm^{-2} and N({^{13}CO}) = 1.3 × 10^{17} cm^{-2}. The full circles show the LTE calculation for T = 120 K and N({^{13}CO}) = 2 × 10^{16} cm^{-2}.

the PDR and allow us to assign some lines that could be merged in the lower resolution SPIRE spectra.

4. Conclusions
We have analysed the first spectral survey taken in the Orion Bar by the FTS of SPIRE. A wealth of rotational lines of CO (and its isotopologues), fine structure lines of C and N^+, and emission lines from radicals and molecules were found. We present the first sparse sampled maps, which illustrate FTS line mapping capabilities. We discussed the CO excitation and emphasized the need for complementary spectroscopic data.

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