

Water in star-forming regions with *Herschel*: highly excited molecular emission from the NGC 1333 IRAS 4B outflow ^{★,★★}

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Received 18 August 2011 / Accepted 17 January 2012

ABSTRACT

During the embedded phase of pre-main sequence stellar evolution, a disk forms from the dense envelope while an accretion-driven outflow carves out a cavity within the envelope. Highly excited ($E' = 1000\text{--}3000$ K) H_2O emission in spatially unresolved *Spitzer*/IRS spectra of a low-mass Class 0 object, NGC 1333 IRAS 4B, has previously been attributed to the envelope-disk accretion shock. However, the highly excited H_2O emission could instead be produced in an outflow. As part of the survey of low-mass sources in the *Water in Star Forming Regions with Herschel* (WISH-LM) program, we used *Herschel*/PACS to obtain a far-IR spectrum and several Nyquist-sampled spectral images to determine the origin of excited H_2O emission from NGC 1333 IRAS 4B. The spectrum has high signal-to-noise in a rich forest of H_2O , CO, and OH lines, providing a near-complete census of far-IR molecular emission from a Class 0 protostar. The excitation diagrams for the three molecules all require fits with two excitation temperatures. The highly excited component of H_2O emission is characterized by subthermal excitation of ~ 1500 K gas with a density of $\sim 3 \times 10^6 \text{ cm}^{-3}$, conditions that also reproduce the mid-IR H_2O emission detected by *Spitzer*. On the other hand, a high density, low temperature gas can reproduce the H_2O spectrum observed by *Spitzer* but underpredicts the H_2O lines seen by *Herschel*. Nyquist-sampled spectral maps of several lines show two spatial components of H_2O emission, one centered at $\sim 5''$ (1200 AU) south of the central source at the position of the blueshifted outflow lobe and a heavily extinguished component centered on-source. The redshifted outflow lobe is likely completely obscured, even in the far-IR, by the optically thick envelope. Both spatial components of the far-IR H_2O emission are consistent with emission from the outflow. In the blueshifted outflow lobe over 90% of the gas-phase O is molecular, with H_2O twice as abundant than CO and 10 times more abundant than OH. The gas cooling from the IRAS 4B envelope cavity walls is dominated by far-IR H_2O emission, in contrast to stronger [O I] and CO cooling from more evolved protostars. The high H_2O luminosity may indicate that the shock-heated outflow is shielded from UV radiation produced by the star and at the bow shock.

Key words. infrared: ISM – ISM: jets and outflows – stars: protostars – molecular processes – stars: individual: NGC 1333 IRAS 4B

1. Introduction

During the embedded phase of pre-main sequence stellar evolution, the protoplanetary disk forms out of a dense molecular envelope (e.g. Terebey et al. 1984; Adams et al. 1987). Meanwhile, as the protostar builds up most of its mass, it drives powerful, collimated outflows into the dense envelope (e.g. Bontemps et al. 1996). These processes together eventually cause the envelope to dissipate and set the initial conditions for disk evolution and planet formation.

At the interfaces between the outflow and envelope and between envelope and disk, shocks can heat the gas and potentially produce detectable emission. The well-studied outflow-envelope

interactions produce an outflow cavity with walls heated by shocks and energetic radiation from the central star (e.g. Snell et al. 1980; Spaans et al. 1995; Arce & Sargent 2006; van Kempen et al. 2009; Tobin et al. 2010). On the other hand, observational evidence for the disk-envelope interactions has been sparse. Velusamy et al. (2002) detected methanol emission from L1157 on scales of ~ 1000 AU, spatially-extended beyond the point-like continuum emission, and argued that the kinematics suggest that the emission is produced at the disk/envelope interface. Watson et al. (2007) detected emission in highly excited ($E' = 1000\text{--}3000$ K) H_2O lines from the NGC 1333 IRAS 4B system and attributed the heating to an envelope-disk accretion shock within 100 AU of the star. These observations offer two different interpretations for disk-envelope interactions, with material either entering the disk on large scales (Visser et al. 2009; Vorobyov 2011) or raining onto the disk at small radii (Whitney & Hartmann 1993). However, a persistent complication in interpreting spatially and spectrally unresolved emission as coming

* *Herschel* is an ESA space observatory with science instruments provided by European-led Principal Investigator consortia and with important participation from NASA.

** Appendices are only available in electronic form at <http://www.aanda.org>

from a compact disk-like structure is that outflows can also produce bright emission in highly excited lines. Molecular emission is a dominant coolant of outflow-envelope interactions, and H₂O emission is particularly sensitive to shocks in outflows (e.g. Nisini et al. 2002, 2010; van Kempen et al. 2010a).

The NGC 1333 IRAS 4B system (hereafter IRAS 4B) is a Class 0 YSO ($d = 235$ pc, Hirota et al. 2008) with a $0.24 M_{\odot}$ disk that is deeply embedded ($A_V \sim 1000$ mag) within a $2.9 M_{\odot}$ envelope (Jørgensen et al. 2002, 2009). Compact outflow emission is detected in many sub-millimeter (sub-mm) molecular lines (Di Francesco et al. 2001; Jørgensen et al. 2007). Near-IR emission in all four *Spitzer*/IRAC bands (3.5, 4.5, 5.8, and $8.0 \mu\text{m}$) is located in the blueshifted outflow lobe (Jørgensen et al. 2006, see also Choi et al. 2011), offset by $\sim 6''$ south of the peak of interferometric sub-mm continuum emission. The redshifted outflow is seen in sub-mm line emission (Jørgensen et al. 2007) but is not detected in the near- or mid-IR because the outflow is located behind IRAS 4B and hidden by the high extinction of the envelope. In the near-IR, the *Spitzer*/IRAC photometry is dominated by H₂ emission (Arnold et al. 2011; see also Neufeld et al. 2008), with some contribution of CO fundamental emission to the $4.5 \mu\text{m}$ bandpass (Tappe et al. 2011; see also Herczeg et al. 2011). Excited water emission from the NGC 1333 IRAS 4 system, including both IRAS 4A and 4B, was detected with ISO/LWS, but with too low spatial resolution to attribute the emission to any component in the system (Giannini et al. 2001). H₂O maser emission has also been seen from dense gas associated with the IRAS 4B outflow, although with a different position angle than the molecular outflow (Rodríguez et al. 2002; Furuya et al. 2003; Marvel et al. 2008; Desmurs et al. 2009).

In a sample of low-resolution *Spitzer*-IRS spectra of 30 Class 0 objects, Watson et al. (2007) found water emission from highly-excited levels in only IRAS 4B. The lines have upper levels with high excitations (1000–3000 K) and high critical densities ($\sim 10^{11} \text{ cm}^{-3}$). Watson et al. inferred that the emitting gas has a high density and argued that the high density indicates that the emission is produced in a $\sim 2 \text{ km s}^{-1}$ accretion shock at the envelope-disk interface. They also argued that the emission was detected only from IRAS 4B because the viewing angle may be well-aligned with the outflow, allowing a clear view of the embedded disk, and because the timescale for such high envelope-disk accretion rates may be short. H₂O emission has since been detected in at least two other components of the IRAS 4B system: (1) narrow ($\sim 1 \text{ km s}^{-1}$, with rotation signatures) p-H₂¹⁸O $3_{13-2_{20}}$ ($E' = 204$ K) emission in a spatially compact region, likely a (pseudo)-disk of ~ 25 AU in radius (Jørgensen & van Dishoeck 2010); and (2) broad (FWHM $\sim 24 \text{ km s}^{-1}$), spatially unresolved emission from low-excitation H₂O ($E' = 50\text{--}250$ K) lines, which are consistent with an outflow origin (Kristensen et al. 2010) but too broad for the $\sim 2 \text{ km s}^{-1}$ velocity expected of envelope gas in free-fall striking the disk at 25–100 AU (e.g. Shu et al. 1977).

The highly-excited mid-IR emission and the broad line profiles of lower-excitation lines could be reconciled if either (a) the high- and low-excitation H₂O emission lines originate in different locations, if (b) models for the envelope-disk accretion shock underpredict line widths, or if (c) both the high- and low-excitation H₂O lines are produced in the outflow. In this paper, we analyze a *Herschel*/PACS far-IR spectral survey and spectral imaging of IRAS 4B to resolve the discrepancy in the different possible origins of H₂O emission from IRAS 4B. The far-IR spectrum of IRAS 4B is as rich in lines as its mid-IR *Spitzer*/IRS spectrum. In Nyquist-sampled maps, the

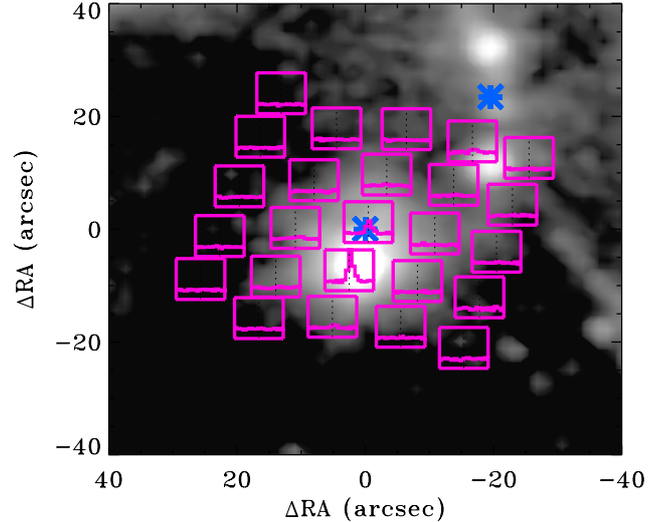


Fig. 1. The location of the 5×5 spaxel array (purple spectral map) against an $4.5 \mu\text{m}$ image obtained with *Spitzer*/IRAC (grayscale) from Jørgensen et al. (2006). The spectral map shows that the continuum-subtracted emission in the o-H₂O $6_{16-5_{05}}$ $82.03 \mu\text{m}$ ($E' = 643$ K) line is produced mostly in the blueshifted outflow lobe. The sub-mm positions of IRAS 4B and IRAS 4A are marked with blue asterisks.

H₂O emission is spatially offset from the continuum emission in the direction of the blueshifted outflow. The excitation of warm H₂O lines detected with *Herschel*/PACS and with *Spitzer*/IRS together can be explained by emission from an isothermal slab. An additional physical component(s) is located on-source and likely traces material closer to the base of the outflow. No evidence is seen for an envelope-disk accretion shock. This result is consistent with a conclusion by Tappe et al. (2012), obtained contemporaneous to the results in this paper, that the H₂O emission seen in the *Spitzer*/IRS spectra also coincides with the southern outflow position. We discuss the implications of these results for outflows and for the prospects of observationally studying disk formation with H₂O lines.

2. Observations and data reduction

We obtained far-IR spectra of NGC 1333 IRAS 4B ($03^{\text{h}}29^{\text{m}}12^{\text{s}}.0 +31^{\circ}13'08''.1$; Jørgensen et al. 2007) on 15–16 March 2011 with the PACS instrument on *Herschel* (Pilbratt et al. 2010; Poglitsch et al. 2010) as part of the WISH key program (van Dishoeck et al. 2011). The observations presented here consist of a complete scan of the $52\text{--}208 \mu\text{m}$ spectral range and deep, Nyquist-sampled spectral maps in four narrow ($\sim 0.5\text{--}1 \mu\text{m}$) wavelength regions. Each PACS spectrum includes observations of two different nod positions located at $\pm 3'$ from the science observation to subtract the instrumental background. All PACS data were reduced with HIPEv6.1 (Ott 2010). We supplemented the PACS observations with re-reduced archival *Spitzer*/IRS spectra that were previously analyzed by Watson et al. (2007). Details of the observations and reduction are described in the following subsections.

2.1. Complete *Herschel*/PACS far-IR spectrum

The $52\text{--}208 \mu\text{m}$ spectrum of IRAS 4B was obtained in 2.7 h of integration with PACS. PACS observed IRAS 4B simultaneously in the first order $>100 \mu\text{m}$ and in the second order at $<100 \mu\text{m}$. The grating resolution varies between $R = 1000\text{--}2000$

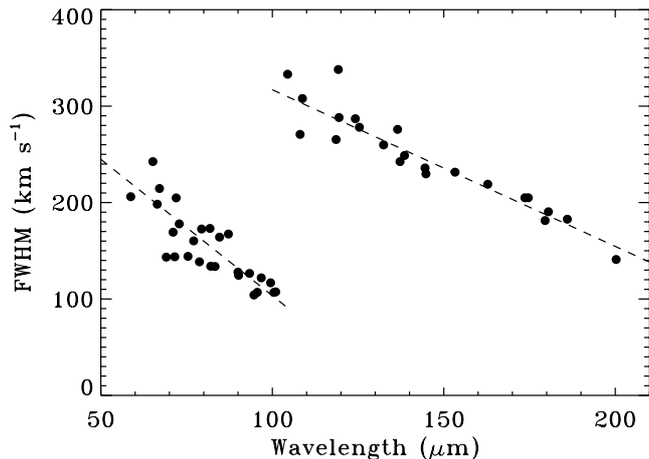


Fig. 2. The FWHM from initial fits to strong lines in our full scan of the 52–208 μm wavelength range. The lines are spectrally unresolved and provide a measure of the spectral resolution. The line widths for our final fits were set by linear fits to the FWHM at >100 and <100 μm .

at >100 μm and $R = 3000$ – 4000 at <100 μm . Large grating steps are used, so the spectrum is binned to ~ 2 pixels per resolution element. A spectral flatfielding was applied to the data to improve the S/N. The observed flux was normalized to the telescopic background and subsequently calibrated from observations of Neptune, which is used as a spectral standard. Appendix A describes a new method to calibrate the flux in PACS spectra between 97–103 μm and above >190 μm . The relative flux calibration is accurate to $\sim 20\%$ across most of the spectrum.

The spectral scan produced a single 5×5 spectral map over a $47'' \times 47''$ field-of-view. The central spaxel is centered at the location of the sub-mm continuum peak. An adjacent spaxel located $9'.4$ to the SE ($PA = 249^\circ$) is centered on the blueshifted outflow position (Fig. 1). Most of the line and continuum flux is located in these two spaxels. The southern outflow of NGC 1333 IRAS 4A is located in the NW corner of the array.

Extracting line fluxes requires an assessment of the distribution of flux on the detector caused by both the point-spread function of *Herschel* and the spatial extent of the emission. For a well-centered point source, the encircled energy in a single spaxel is $\sim 70\%$ at ≤ 100 μm wavelengths and declines to $\sim 40\%$ near 200 μm . However, the signal-to-noise decreases if the spectrum is extracted from many spaxels. The emission line spectrum is obtained by adding the flux from the central spaxel and the outflow spaxel. The line fluxes are subsequently corrected for light leakage and the spatial extent in the emission by comparing line fluxes from this 2-spaxel extraction with the line fluxes extracted from a 3×3 spaxel area centered on the central spaxel. The line flux ratio between the 2-spaxel and 3×3 spaxel extraction was calculated for strong lines of all detected molecules. The fluxes from the 2-spaxel extraction are then divided by a wavelength-dependent correction that is 0.76 at <100 μm and then decreases linearly to 0.6 at 180 μm . The wavelength dependence of this correction includes both the point-spread function of *Herschel* and the wavelength dependence in the spatial distribution of the detected emission. Finally, high signal-to-noise PACS spectra of the point source HD 100546 (Sturm et al. 2010) were then used to correct for emission leaked beyond the 3×3 spaxel area. This approach assumes a similar spatial distribution for all molecules and for lines at nearby wavelengths, and introduces a $\sim 10\%$ uncertainty in relative fluxes. The overall uncertainty in flux calibration is $\sim 30\%$. The typical noise level in

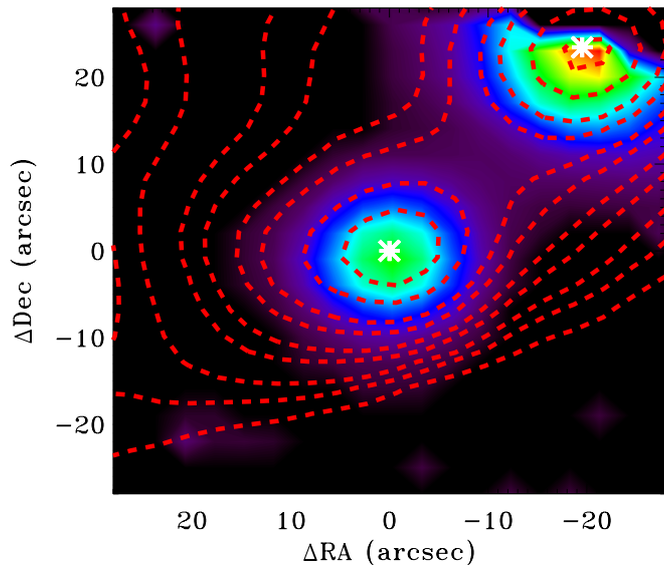


Fig. 3. The 63 μm continuum emission (colors) and the SCUBA 450 μm emission map (contours), with the location of IRAS 4A and IRAS 4B marked as white asterisks. The PACS maps are shifted so that the centroid of the 190 μm continuum emission is located at the position of IRAS 4B obtained from sub-mm continuum interferometry. The SCUBA map is shifted to the same position.

the two-spaxel extraction ($\sim 9.4 \times 18.8''$) is ~ 0.4 Jy per resolution element in the extracted continuum spectrum.

All lines in the complete spectral scan are spectrally unresolved. The lines were initially fit with Gaussian profiles, with the central wavelength, line width, and amplitude and a first-order continuum as free parameters. The instrumental line widths were then calculated from first-order fits to the wavelength-dependent spectral resolution, obtained from strong lines (Fig. 2). Our final fluxes are obtained from fitting Gaussian profiles to each line, with the line width set by the calculated instrumental resolution at the given wavelength. For strong lines, the median centroid velocity is 30 km s^{-1} with a standard deviation of 24 km s^{-1} . The absolute wavelength calibration is accurate to ~ 50 km s^{-1} and is limited by the spatial distribution of emission within each spaxel.

2.2. *Herschel*/PACS Nyquist-sampled spectral imaging

Nyquist-sampled spectral maps of narrow spectral regions at 54.5, 63.3, 108.5, and 190 μm were obtained in a total of 2.2 h of integration time. These maps were obtained from a 3×3 raster scan with $3''$ steps, yielding spatial resolutions of $\sim 5''$, $5''$, $8''$, and $10''$, respectively. Small grating steps were used to fully sample the spectral resolution. The final spectrum is rebinned onto a wavelength grid with ~ 4 pixels per resolution element.

The spectral maps include CO, H₂O, and [O I] lines listed in Table 1. In a $3 \times 3''$ area, the typical rms is ~ 0.02 Jy per resolution element at 63 μm and ~ 0.01 Jy per resolution element at 108 μm . This sensitivity level is better than that from the full spectral scan because of different integration areas and much longer integration times in each resolution element.

The data cubes from the Nyquist-sampled maps were reprojected onto a normal grid of right ascension and declination. In the automated calibration the 190 μm continuum emission

Table 1. Location of PACS line and continuum emission from the IRAS 4B system.

Species	Line	E_{up} (K)	λ (μm)	ΔRA ($''$)	ΔDec ($''$)	$\sigma(\text{RA})$ ($''$)	$\sigma(\text{Dec})$ ($''$)
CO	49–48	6457	53.9	3.4 ± 1.5	-2.4 ± 0.5	7 ± 2	9.9 ± 1.0
o-H ₂ O	5 ₃₂ –5 ₀₅	732	54.5	1.0 ± 1.0	-5.4 ± 0.5	9.9 ± 1.0	10.4 ± 1.0
[O I]	³ P ₁ – ³ P ₂	228	63.2	-1.3 ± 0.3	-3.9 ± 0.2	11.8 ± 0.7	12.2 ± 0.8
o-H ₂ O	8 ₁₈ –7 ₀₇	1070	63.3	0.2 ± 0.2	-5.1 ± 0.3	11.2 ± 0.3	12.2 ± 0.4
o-H ₂ O	8 ₀₈ –7 ₁₇	1070	63.5	0.2 ± 0.3	-5.2 ± 0.4	11.3 ± 0.6	14.4 ± 1.0
Continuum	–	–	63.3	0.4 ± 0.4	-0.7 ± 0.5	11.5 ± 0.5	10.1 ± 0.8
o-H ₂ O	2 ₂₁ –1 ₁₀	194	108.1	0.3 ± 0.3	-3.1 ± 0.2	12.5 ± 0.4	18.1 ± 0.4
CO	24–23	1524	108.8	0.0 ± 0.3	-3.7 ± 0.3	12.2 ± 0.5	17.0 ± 0.6
Continuum	–	–	108.5	-0.05 ± 0.5	-0.8 ± 0.1	12.5 ± 0.2	11.3 ± 0.4
Continuum	–	–	189.9	(0.0)	(0.0)	16.5 ± 0.5	14.4 ± 0.3

is offset from the object position¹, as measured from sub-mm interferometry (Jørgensen et al. 2007). Each map is shifted in position by 3'3 E and 1'5 N so that the location of the 190 μm continuum from our observations matches the peak location of the sub-mm emission. After the shift, the 63 and 108 μm continuum emission from IRAS 4B and IRAS 4A (located in the NW edge of the map) are well aligned with the *Spitzer*-MIPS 70 μm emission. No significant offset was measured in the staring observation of the full PACS SED, which was obtained with a new pointing one day after the Nyquist-sampled map. Because the same spatial shift is applied to both the line and far-IR continuum emission maps and because the shift moved both the line and continuum closer to the sub-mm continuum peak, the result that the line emission is spatially offset from the sub-mm continuum is robust to the pointing uncertainty.

The southern nod position in the map has spatially-extended [O I] emission in the southwest portion of the map. The [O I] emission is therefore measured from only the northern nod position. Inspection of *Spitzer*/MIPS 70 μm maps at the two nod positions does not indicate the presence of a strong, point-source continuum emission that could otherwise corrupt the continuum map for IRAS 4B. The 54 μm continuum map has low S/N and is not used.

The two H₂O and [O I] lines at 63 μm have spectral widths of 110 km s⁻¹, which places an upper limit of 65 km s⁻¹ on the intrinsic line width. That the line widths are broader than the instrumental resolution of ~ 3300 is not significant because emission that is spatially extended in the cross-dispersion direction can broaden the spectral line profile, as with any other slit spectrograph.

2.3. *Spitzer*/IRS spectrum

The *Spitzer*/IRS spectra of IRAS 4B were originally presented by Watson et al. (2007). We re-reduced the spectrum following the procedure described by Pontoppidan et al. (2010). The LH slit width is 11''. The flux extraction region of 5–10'' across the wavelength region was selected to optimize the final signal-to-noise in the limit of a point source (Horne 1986). The *Spitzer* and *Herschel* observations cover similar regions on the sky. The relative flux calibration between *Spitzer* and *Herschel* spectra is likely uncertain by $\sim 30\%$ flux.

¹ The optical guider observations onboard *Herschel* are typically accurate to $\sim 1''$. However, observations of nearby high-extinction regions, including NGC 1333 IRAS 4B, have few optical guide stars, which may introduce pointing offsets when the few guide stars are distributed asymmetrically in the field-of-view.

3. Results

3.1. Far-IR spectrum of IRAS 4B

The far-IR PACS spectrum of IRAS 4B is the richest far-IR spectrum of a YSO to date. A forest of high signal-to-noise CO, H₂O, and OH lines that provide a full census of far-IR molecular emission that can be detected from low-mass YSOs (Fig. 4; see also Figs. D.1, D.2 in the Appendix). Lines were discovered in the spectrum following a biased search for emission at the wavelengths for transitions of common species and an unbiased search for narrow features that peak above the noise level.

A total of 115 distinct emission lines are detected and identified from IRAS 4B (Table 3). All strong lines are identified. Several tentative detections of weak lines are unidentified and discussed in Appendix B. No H₂¹⁸O or ¹³CO emission is detected in the PACS observations, with typical flux limits in the strongest expected lines of ~ 0.03 times the observed flux of the main isotopologue. The [O I] 145.5 μm line is not detected, with a 2σ flux limit 1.2×10^{-21} W cm⁻². Lines of OH⁺ and CH⁺, HD 56 and 112 μm , [N II] 121.8 and 205.2 μm , [C II] 157.7 μm , and [O III] 88.7 μm are also not detected.

Figure 1 shows a spectral map of the o-H₂O 6₁₆–5₀₅ 82.03 μm ($E' = 643$ K) line emission overplotted on a *Spitzer*/IRAC 4.5 μm image of IRAS 4B (and IRAS 4A). The line emission is located at the position of the near-IR emission in the blueshifted outflow lobe, south of the central source of IRAS 4B. On the other hand, most of the continuum emission is located in the central spaxel, consistent with the location of the sub-mm continuum emission. Figure 5 demonstrates that the equivalent width of far-IR lines is much larger in the outflow spaxel than in the continuum spaxel, indicating a spatial offset between the line and continuum emission. The line emission in the central spaxel mostly disappears at <70 μm . All lines in the PACS spectrum are spatially offset from the continuum emission. The line fluxes are measured based on the summation of these two bright spaxels and correction for spatial extent (see Sect. 2.1).

3.2. PACS mapping of H₂O Emission from IRAS 4B

Figure 6 and Table 1 compare the spatial distribution of the continuum flux with H₂O, CO, and [O I] line fluxes obtained from the Nyquist-sampled spectral maps. The far-IR continuum emission is centered on-source, at the location of the sub-mm continuum, while line emission is centered to the south in the blueshifted outflow. Figure 7 shows a cartoon version of the approximate location of line and continuum emission and the morphology of IRAS 4B. In the following analysis, we simplify the analysis by assuming that emission consists of two unresolved

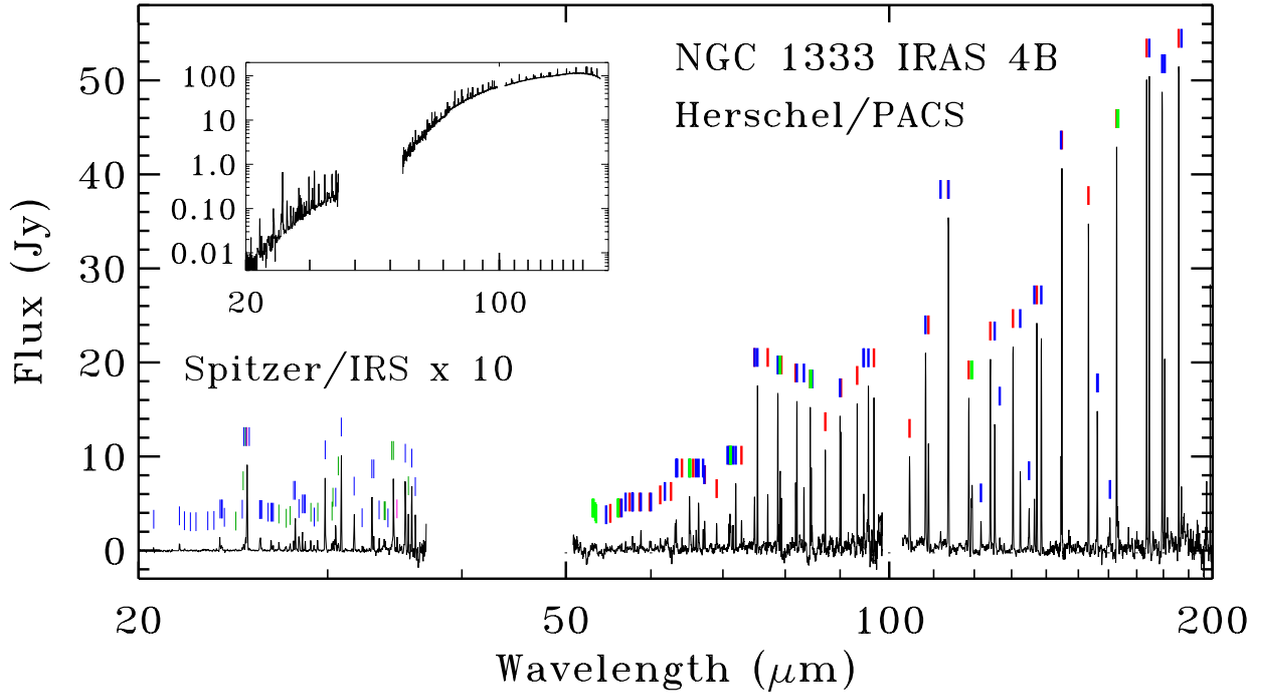


Fig. 4. The combined *Herschel*/PACS and *Spitzer*/IRS continuum-subtracted spectra of IRAS 4B, with bright emission in many H₂O (blue marks), CO (red marks), OH (green marks), and atomic or ionized lines (purple). The *Spitzer* spectrum is multiplied by a factor of 10 so that the lines are strong enough to be seen on the plot. The inset shows the combined spectrum including the continuum.

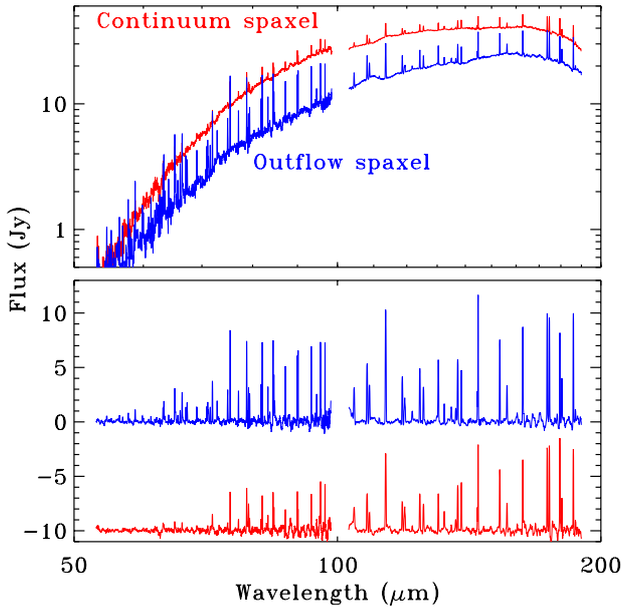


Fig. 5. *Top:* spectra extracted separately from a spaxel centered on the sub-mm continuum (red) and a spaxel offset by $9''.4$ to the S and centered on the outflow position (blue). The outflow position is dominated by line emission while the central object is dominated by continuum emission, indicating that the lines and continuum are spatially offset. *Bottom:* the same two spectra after continuum subtraction show that the on-source line emission gets much weaker to short wavelengths.

point sources, one at the outflow position and one at the sub-mm continuum peak, and are subsequently fit with 2D Gaussian profiles. More complicated spatial distributions would be unresolved in our maps.

The right panel of Fig. 6 demonstrates that the location of the warm H₂O coincides with the *Spitzer*/IRAC $4.5 \mu\text{m}$ imaging. The two H₂O lines near $63.4 \mu\text{m}$, o-H₂O $8_{18}-7_{07}$ and p-H₂O $8_{08}-7_{17}$, ($E' = 1070 \text{ K}$) are centered at $5.2 \pm 0.2''$ from the $63 \mu\text{m}$ continuum and are spatially extended relative to the continuum emission (assumed to be unresolved for simplicity) by $\text{FWHM} = 6.7 \pm 1.0''$. About 70% of the emission is produced at the southern outflow position (Fig. 8). The flux ratios for the two H₂O $63.4 \mu\text{m}$ lines are similar at both the on-source and off-source positions (Fig. 9). In the $108.5 \mu\text{m}$ spectral map, both the o-H₂O $2_{21}-1_{10}$ ($E' = 194 \text{ K}$) and CO $24-23$ ($E' = 1524 \text{ K}$) emission are centered $2.5 \pm 0.4''$ south of the $108.5 \mu\text{m}$ continuum emission and are spatially-extended in the outflow direction by $13.6 \pm 0.7''$, relative to the extent of the continuum emission. The larger spatial extent and smaller offset in the $108 \mu\text{m}$ lines both indicate that the outflow component contributes $\sim 40\%$ of the measured line flux. The spatial differences may be interpreted as differential extinction across the emission region, discussed in the next subsection.

The $54 \mu\text{m}$ maps are noisy because PACS has poor sensitivity at $<60 \mu\text{m}$. The o-H₂O $5_{32}-5_{05}$ $54.507 \mu\text{m}$ ($E' = 732 \text{ K}$) emission is offset by $5.9 \pm 0.4''$ south from the the peak of the sub-mm continuum emission. The CO $49-48$ $53.9 \mu\text{m}$ emission ($E' = 6457 \text{ K}$) is offset $2.9 \pm 0.4''$ south, between the peak of the sub-mm emission and the bright outflow location. The highly-excited CO emission is produced in a different location than the highly-excited H₂O emission.

The [O I] emission is offset by $3.7 \pm 0.3''$ at $\text{PA} = 168^\circ$, just west of the outflow, and is spatially extended by $\sim 7''.1 \pm 1.0$.

3.3. Extinction estimates to the central source and outflow position

The extinction to different physical structures within the IRAS 4B system depends on how much envelope material is

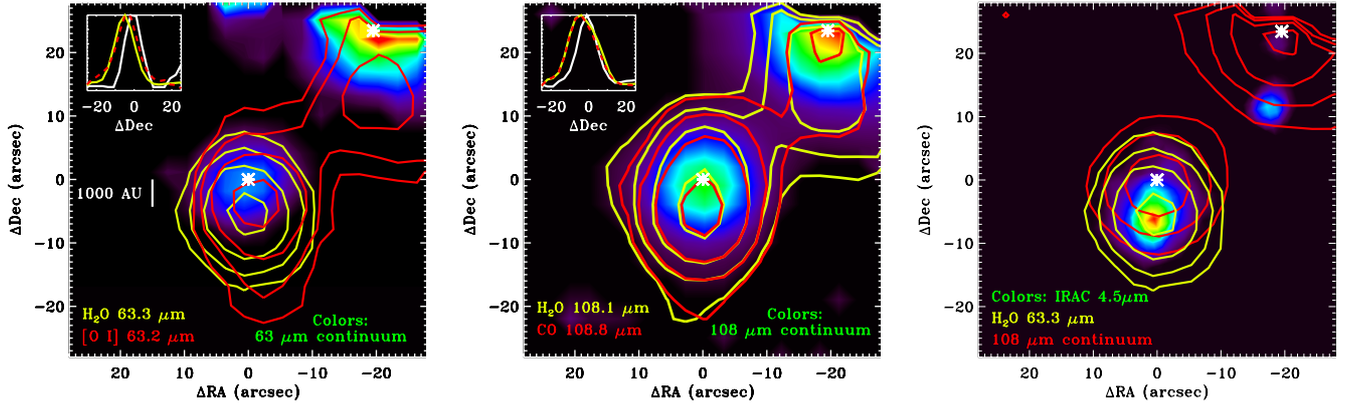


Fig. 6. *Left and Middle:* contour maps of H₂O (yellow), [O I] (red, *left*), and CO (red, *middle*) emission compared with continuum emission (color contours) at 63.3 μm (*left*) and 108.5 μm (*middle*). The insets show the spatial extent of continuum and line emission in the different components along the N-S outflow axis. *Right:* a comparison of the location of emission in H₂O 63.32 μm (blue), 108.5 μm continuum (red), *Spitzer*-IRAC 4.5 μm photometry (color contours; Jørgensen et al. 2006; Gutermuth et al. 2008). All contours have levels of 0.1, 0.2, 0.4, 0.8, and 1.6 times the peak flux near IRAS 4B. The asterisks show the sub-mm positions of IRAS 4B near the center of the field and IRAS 4A in the NW corner. Some noise in these maps is suppressed at empty locations far from IRAS 4A and IRAS 4B.

present in our line of sight. In this subsection, we discuss how these different extinctions affects the emission that is seen. The extinction law used here is obtained from Weingartner & Draine (2001) with a total-to-selective extinction parameter $R_V = 5.5$, typical of dense regions in molecular clouds (e.g. Indebetouw et al. 2005; Chapman et al. 2009). Appendix D includes a discussion of how extinctions may affect the molecular excitation diagrams.

The strength of the near-IR emission in the outflow (Jørgensen et al. 2006) indicates that extinction must be low to at least some of the outflow position. Spherical models of the dust continuum indicate that the central protostar is surrounded by $A_V = 1000$ mag (Jørgensen et al. 2002), so any emission from the redshifted outflow lobe may suffer from as much as $A_V \sim 2000$ mag of extinction. Any additional extinction would have likely introduced asymmetries in the H₂O line profiles that were presented in Kristensen et al. (2010). Depending on the wavelength and spatial location of the emission, the far-IR emission line fluxes may be severely affected by extinction.

The H₂O 54.5 and 63.4 μm lines have a different spatial distribution than the H₂O 108.1 μm line. If we assume that the on-source and off-source emission both have similar physical conditions, then the ratio of the H₂O 63.4 to 108.1 μm line luminosities should not change with position. In this scenario, the different locations for the detected flux is caused by differential extinction across the emission area. An average extinction to the on-source H₂O component of $A_V \sim 700$ mag would reduce the fractional contributions from the on-source and off-source locations observed values. This extinction may be the combination of a lightly-extinguished region on the front side of the protostar and a heavily-extinguished region on the back side of the protostar.

An independent estimate of the extinction can also be made from the flux ratio of [O I] 63.18 to 145.5 μm lines, which is typically observed to be about 10 (Giannini et al. 2001; Liseau et al. 2006). The undetected [O I] 145.5 μm line flux is less than 10% of the 1.8×10^{-20} W cm⁻² flux in the [O I] 63.18 μm line. If we conservatively assume that the true ratio is 30, then we estimate $A_V < 200$ mag to the [O I] emission region. Some additional [O I] emission could only be hidden behind a high enough extinction ($A_V \sim 4000$ mag) to attenuate emission in both the 63.18 and 145.5 μm lines. Therefore, the effect of extinction on

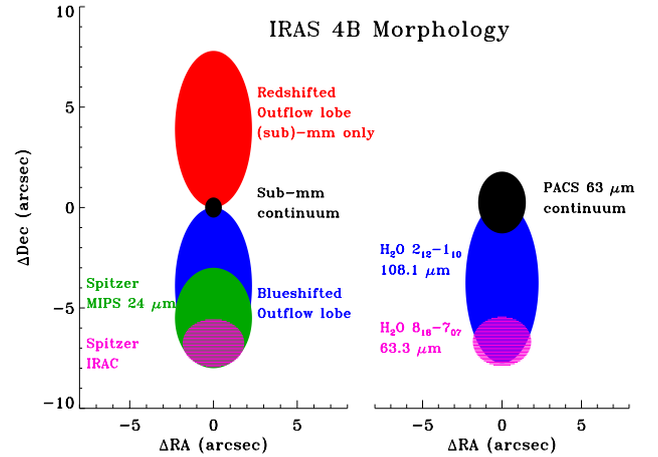


Fig. 7. A cartoon showing the location of different emission components from IRAS 4B, based in part on Fig. 6.

the [O I] luminosity is likely not too significant for the southern outflow lobe, where [O I] emission is seen.

3.4. *Spitzer*/IRS spectrum and broadband images of IRAS 4B

The *Spitzer*/IRS spectrum of IRAS 4B includes emission in lines of highly-excited H₂O and OH, plus [S I] and [Si II]. Our line identification mostly agrees with that of Watson et al. (2007) for H₂O lines, with some modifications to account for the identifications of OH lines (Table 2 and Fig. 10, see also Tappe et al. 2008). The H₂O line identification was informed from line intensities predicted by RADEX modelling (see Sect. 4.1) of both the low-density case discussed here and the high density-case of Watson et al. (2007). All lines with significant detections are identified.

An analysis of the *Spitzer*/MIPS 24 μm image, with sensitivity from 20–31 μm, helps us place limits on the amount of H₂O emission that could arise on-source. Convolution of the filter transmission curve with the *Spitzer*/IRS spectrum from Watson et al. (2007) indicates that ~24% of the light in the MIPS 24 μm

Table 2. OH lines detected in the *Spitzer*/IRS spectrum of IRAS 4B^a.

Line ID	E_{up} (K)	$\log A_{\text{ul}}$ (s ⁻¹)	λ_{vac} (μm)	Flux ^b	Err ^b
${}^2\Pi_{1/2} - {}^2\Pi_{3/2} J = 9/2 - 7/2^c$	875	-1.40	24.62	1.6	0.2
${}^2\Pi_{3/2} J = 21/2 - 19/2^+$	2905	1.29	27.39	6.0	0.4
${}^2\Pi_{3/2} J = 21/2^+ - 19/2^-$	2899	1.29	27.45	8.7	0.4
${}^2\Pi_{1/2} J = 19/2 - 17/2^c$	2957	1.28	27.67	8.3	0.2
${}^2\Pi_{1/2} - {}^2\Pi_{3/2} J = 7/2 - 5/2^c$	617	-1.50	28.94	8.3	0.2
${}^2\Pi_{3/2} J = 19/2^+ - 17/2^-$	2381	1.16	30.28	11.9	0.3
${}^2\Pi_{3/2} J = 19/2^- - 17/2^+$	2375	1.15	30.35	13.1	0.3
${}^2\Pi_{1/2} J = 17/2^+ - 15/2^-$	2439	1.14	30.66	7.2	0.6
${}^2\Pi_{1/2} J = 17/2^- - 15/2^+$	2436	1.14	30.71	10.9	0.6
${}^2\Pi_{3/2} J = 17/2^- - 15/2^+$	1905	1.01	33.86	11.3	0.8
${}^2\Pi_{3/2} J = 17/2^+ - 15/2^-$	1901	1.00	33.95	8.4	0.8
${}^2\Pi_{1/2} J = 15/2 - 13/2^c$	1969	0.98	34.61	9.3	1.0

Notes. ^(a) Listed OH lines are the sum of unresolved triplet hyperfine structure transitions. ^(b) 10^{-22} W cm⁻², with $1 - \sigma$ error bars. ^(c) Includes two sets of unresolved triplets with different parities.

bandpass is in molecular emission (mostly H₂O), 17% in the [S I] 25.24 μm line, and 59% in the continuum.

Jørgensen & van Dishoeck (2010) found that the emission from IRAS 4B in the *Spitzer*/MIPS 24 μm images is offset from the peak emission of the sub-mm continuum (see also Choi & Lee 2011). We measure that the emission is centered at 5'2 S and 0'9 E from the central source, consistent with the location of the outflow emission, and is spatially extended by $\sim 5'5$ in the north-south direction, along the outflow axis. An additional component is present at the location of the sub-mm continuum peak. The *Spitzer*/MIPS emission is assumed here to be a combination of emission from two unresolved sources, one at the sub-mm continuum peak and one at the position of the *Spitzer* IRAC 4.5 μm emission located 6'2 S and 0'4 E. From fitting two dimensional Gaussian profiles to the image, the component at the blueshifted outflow lobe accounts for 78% of the *Spitzer*/MIPS 24 μm emission and the sub-mm point source accounts for the remaining 22% of the emission (see Fig. 8 for the fit to the image collapsed onto the outflow direction). Some additional *Spitzer*/MIPS emission is located at 20'' S of the sub-mm continuum peak and is ignored here.

In the *Spitzer*/IRAC images of emission between 3.8–8 μm , the emission is located entirely at the outflow position. In contrast, the 63 μm continuum emission is located mostly on the central source at the sub-mm continuum position. Much of the 20–31 μm continuum emission must be located at the outflow position, but some continuum emission could also be located on the central source. Given the fraction of emission located on-source (22%) and the relative contributions of molecular lines (24%) and continuum to the *Spitzer*/MIPS photometry, the MIPS map could be consistent with an on-source location of H₂O emission only if the continuum emission is located entirely at the outflow position.

4. Excitation of molecular emission from IRAS 4B

The spatial distribution of the highly excited H₂O emission in the PACS observations places the bulk of the highly excited far-IR emission at the outflow position. In this section, we analyze the excitation of the H₂O lines in detail to demonstrate that the highly excited H₂O emission in both the *Herschel*/PACS and *Spitzer*/IRS spectra can be explained with emission from a single isothermal, plane-parallel slab. We subsequently analyze CO, OH, and [O I] emission from IRAS 4B. Although the

H₂O emission region is likely complicated and includes multiple spatial and excitation temperature components, our simplified approach is able to reproduce the highly excited H₂O lines. The properties of this slab are the combination of the spatially offset outflow component and the on-source component. We lack sufficient spatial resolution throughout most of the spectrum to analyze the excitation of the two components separately.

Figure 11 shows excitation diagrams for H₂O, CO, and OH emission². Without considering sub-thermal excitation, each molecule requires two temperature components to reproduce the measured fluxes. For convenience, the two components for each molecule are called “warm” and “cool”, however this terminology applies separately to each molecule³. The warmer component of CO may not be related to the warmer component of H₂O or OH. For the temperature and density derived below, the two apparent excitation temperatures for OH and H₂O could even be produced by a single component, with the warm and cool regimes resulting from subthermal excitation. Table 3 describes the excitation temperatures, molecular column density, and luminosity for fits to these diagrams. In the following subsections we discuss the excitation of the H₂O in detail, and briefly describe the excitation of CO, OH, and O. RADEX models of H₂O are used to fit only the higher excitation H₂O lines because the lower excitation lines are optically thick and difficult to use to infer physical conditions of the emitting gas. The emitting area, temperature, and density derived from the fits to the observed H₂O lines are assumed to also apply to CO, OH, and O for simplicity.

4.1. RADEX models of H₂O emission

The H₂O line emission extends to high energy levels, with level populations indicating an excitation temperature of 220 K but with significant scatter. Among the highly excited lines, the H₂O excitation diagram does not show any break at high energies that would indicate multiple excitation components.

² Throughout the paper all logarithms are base 10, all units for column density are in cm⁻², and all units for density are cm⁻³. The units in excitation diagrams are in number of detected molecules rather than column density.

³ The terminology “warm” and “cool” components, as defined here, corresponds to “hot” and “warm” gas, respectively, in other works, including Visser et al. (2011), which considers colder gas, usually observed in the sub-mm, than the gas studied here.

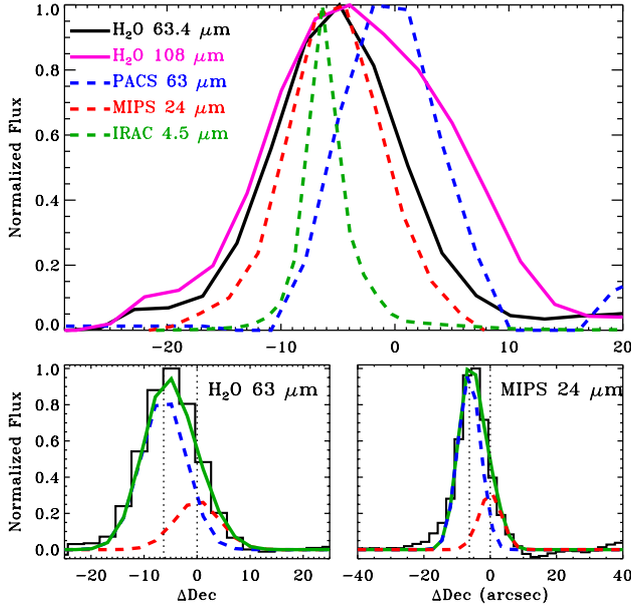


Fig. 8. *Top:* a cross cut of the flux at several wavelengths along the N-S outflow axis. The 63 μm continuum is centered at the peak of the sub-mm continuum while the 4.5 μm and 24 μm photometry are centered in the outflow. The H₂O 63.3 μm lines are produced primarily at the outflow location while the H₂O 108.1 μm line has similar on- and off-source contributions. The (0,0) position is defined here by the peak of the sub-mm continuum emission from Jørgensen et al. (2007). *Bottom:* the spatial cross cut of H₂O 63.3 μm and MIPS 24 μm emission, shown as the combination of two unresolved Gaussian profiles located at the on-source position (red dashed line) and the blueshifted outflow position (blue dashed line).

The highly-excited H₂O levels have high critical densities ($\sim 10^{11} \text{ cm}^{-3}$). The bottom left panel in Fig. 11 demonstrates that a large amount of scatter in excitation diagrams may be explained by sub-thermal excitation. The excitation temperature could therefore be the kinetic temperature of dense ($> 10^{11} \text{ cm}^{-3}$) gas or could result from subthermal excitation of warmer gas with lower density. Line opacities also increase the scatter in observed fluxes.

We calculate synthetic H₂O spectra from RADEX⁴ models of a plane-parallel slab (van der Tak et al. 2007) characterized by a single temperature T , density $n(\text{H}_2)$, and H₂O column density $N(\text{H}_2\text{O})$ with an emitting surface area A . RADEX is a radiative transfer code that simultaneously calculates non-LTE level populations and line optical depths for a plane-parallel slab to produce line fluxes. A large grid was calculated using molecular data obtained from LAMDA (Schöier et al. 2005; Faure et al. 2007). Since this molecular data file lacks the most highly excited lines detected with *Spitzer*, individual RADEX models were calculated at specific gridpoints using a much larger and more complete database with energy levels obtained from Tennyson et al. (2001), radiative rates from the HITRAN database (Rothman et al. 2009), and collisional rates with H₂ from Faure et al. (2008).

The RADEX models are calculated to obtain a rough idea of the physical properties of the emitting gas. Radiative pumping is not included in the model but is likely important, especially at low densities. Including radiative pumping would require detailed physical and chemical modeling of the envelope and is

⁴ <http://www.strw.leidenuniv.nl/~simoldata/radex.html>

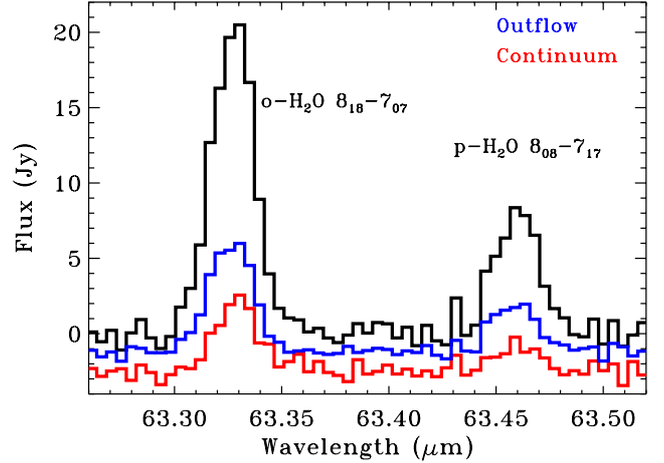


Fig. 9. The H₂O 63.4 μm spectral region extracted from the on-source position (red) and from the blueshifted outflow position (blue), and over the entire spectral map (black). No significant differences are detected in the ratio of the two lines, which indicate that the two lines are optically thin at both locations.

beyond the scope of this work. The line profile is assumed to be a Gaussian profile with a FWHM of 25 km s^{-1} , based on the FWHM of low-excitation H₂O lines observed with HIFI (Kristensen et al. 2010)⁵. The extinction to the warm H₂O gas is highly uncertain and is mostly ignored (see Sect. 3.3 and Appendix D for a discussion of extinction estimates and their implications). The ortho-to-para ratio is assumed to be 3, based on flux ratios of optically-thin lines that range from 2.8–3.5.

A χ^2 fit (upper left panel of Fig. 12) to the measured fluxes with PACS and IRS lines with upper energy level above 400 K (the warm component) yields acceptable solutions with high temperature ($T > 1000 \text{ K}$) and H₂ densities of $\log n < 7.5$. Appendix C provides a detailed description of the line ratios and the non-detections of H₂¹⁸O emission that constrain the best-fit parameters. The size of the emission region further limits the range of acceptable parameter space (Fig. 12). The emitting surface area A is equivalent to the area of a circle with radius from 25–500 AU, which is reasonably close to the projected length of the outflow on the sky ($\sim 1000 \text{ AU}$ or 0.005 pc). If we assume that $N(\text{H}_2\text{O}) < 10^{-4} N(\text{H}_2)$, then the given column density and H₂ density yields the length scales (depth along our line of sight) for the outflow that range from $> 10^{-4} \text{ pc}$ (for the acceptable model with the highest H₂ density and lowest H₂O column density) to $> 130 \text{ pc}$ (for the model with the lowest H₂ density and highest H₂O column density). Given the projected size of the outflow of 0.005 pc, a length scale greater than 0.1 pc is uncomfortably large and rules out solutions with $\log n(\text{H}_2) < 5$. The lack of obvious vibrational excitation in spectra at 6 μm (Maret et al. 2009; Arnold et al. 2011) limits the kinetic temperature to $\lesssim 2000 \text{ K}$. The number of H₂O molecules scales with density.

Combining these analyses, we adopt the parameters $T = 1500_{-600}^{+2000} \text{ K}$, $\log n = 6.5 \pm 1.5$, and $\log N(\text{H}_2\text{O}) = 17.6 \pm 1.2$ over an emitting area equivalent to a circle with radius 25–300 AU and length scale 0.003 pc. Figure 13 shows that the PACS H₂O spectrum is well fit with model fluxes obtained with these parameters.

The temperature and column density are both inversely correlated with the density, so the acceptable parameter space is

⁵ The HIFI H₂O lines are much more optically-thick and likely have larger emitting areas than most H₂O lines in the PACS spectrum, which may lead to differences in line profile.

Table 3. Cooling budget with *Herschel*/PACS^a.

Species	Obs. ^b log <i>L</i>	Fraction of ^b gas cooling	Cool component ^c			Warm component ^c			RADEX model ^d		
			<i>T</i> (K)	log ₁₀ <i>N</i>	log <i>L</i>	<i>T</i> (K)	log ₁₀ <i>N</i>	log <i>L</i>	<i>T</i> ^e	<i>N</i>	log <i>L</i>
H ₂ O	-1.6	0.45	110	47.0	-1.1	220	46.0	-1.6	(220)	48.2	-1.5
CO	-1.6	0.45	280	49.6	-1.8	880	48.5	-1.9	(880)	48.5	-2.0
OH	-2.3	0.09	60	47.1	-1.6	425	44.2	-3.2	(60)	47.6	-2.4
[O I] ^f	-3.5	0.005	–	–	–	–	–	–	–	47.9	-3.5

Notes. ^(a) All luminosities in units of L_{\odot} . ^(b) PACS lines from 53–200 μm , uncorrected for extinction. ^(c) From fits to excitation diagrams. The warm and cool components are not necessarily the same region for each molecule. ^(d) RADEX model with temperature of 1500 K and $\log n(\text{H}_2) = 6.5$. ^(e) The component explained by RADEX model, not the excitation temperature calculated from RADEX model. ^(f) [O I] emission is detected from a different physical location than the molecular emission.

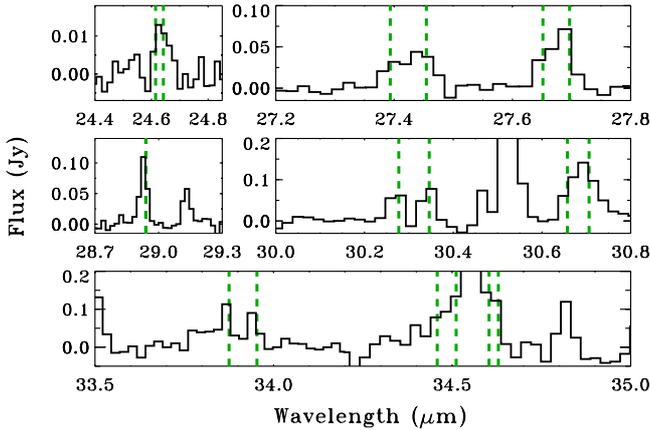


Fig. 10. OH emission lines (vertical dashed lines) in the *Spitzer*/IRS spectrum of IRAS 4B. The lines expected to be strongest are either detected (Table 2) or are blended with lines of other species.

tighter than implied by the large error bars. The choice of line width scales the optical depth. A broader line width would require the same factor increase in column density $N(\text{H}_2\text{O})$ and decrease in total emitting surface area. The widths of the far-IR lines may differ from the optically thick low-excitation H₂O lines analyzed by Kristensen et al. (2010). The far-IR lines are primarily seen from the offset outflow location, while the longer-wavelength lines are dominated by on-source emission and include both red- and blue-shifted outflow lobes. Very broad lines ($>65 \text{ km s}^{-1}$) are ruled out from the widths of lines in the Nyquist sampled spectral maps.

4.2. Comparing H₂O spectra for warm, subthermal excitation and cool, thermalized gas

The high temperature, low density solution presented here (hereafter H12, with properties obtained from the χ^2 fit listed above) produces sub-thermal excitation of H₂O, in contrast to the high-density (thermalized, with $\log n \sim 11$, $\log N(\text{H}_2\text{O}) = 17.0$, and $T = 170 \text{ K}$), low-temperature solution from Watson et al. (2007, hereafter W07). The W07 slab has many optically-thick lines, which leads to a spectrum where the mid-IR H₂O lines observed with *Spitzer* are brighter than those in the PACS wavelength range (Fig. 14). In contrast, many of the far-IR lines in the H12 parameters are optically-thin, so that most H₂O emission escapes in the PACS wavelength range. As a consequence, in principle the far-IR H₂O emission could trace a high temperature, low density region (H12) while the mid-IR H₂O emission traces a high density, low temperature component (W07).

The W07 model produces optically-thick emission in the o-H₂O 8₁₈–7₀₇ and p-H₂O 8₀₈–7₁₇ lines at 63.4 μm , with a flux in the p-H₂O 8₀₈–7₁₇ line similar to the observed flux. However, the two lines are observed in an optically-thin flux ratio both on-source and at the blueshifted outflow lobe. Therefore, the W07 model cannot explain the PACS H₂O emission located at the outflow position. If the extinction in W07 is reduced to $A_V \sim 0 \text{ mag}$, then the two 63.4 μm lines both become two times weaker relative to the mid-IR H₂O lines.

A comparison between the *Spitzer*/IRS spectrum and the synthetic H₂O spectrum (see Fig. 15 and a further discussion of line ratios in Appendix C) shows that both the W07 and H12 models could explain the mid-IR H₂O emission alone. Both models are able to accurately reproduce the emission in most detected lines, with a few notable exceptions. The p-H₂O 8₃₅–7₂₆ 28.9 μm line flux is well reproduced in W07 but not H12. However, the wavelength of this line is more consistent with an OH line than with the H₂O line. The OH rotational diagram (Fig. 11) shows that the line flux is also consistent with fluxes in other OH lines with similar excitations. The inability of H12 to reproduce this line flux with an H₂O model is therefore not significant. An OH line at 24.6 μm was also misidentified as o-H₂O 8₆₃–8₃₆ despite neither W07 nor H12 being able to produce flux in the H₂O line. Because the synthetic fluxes of these lines are faint, the misidentification of this emission as H₂O can help to drive a best fit physical parameters to an optically thick solution.

In models with high $N(\text{H}_2\text{O})$ and $n(\text{H}_2)$, including W07, the line blend of o-H₂O 6₅₂–5₂₃ and 5₅₀–4₂₃ at 22.4 μm and the p-H₂O 6₄₂–5₁₅ 23.2 μm line are predicted to be strong but are not detected. Both W07 and H12 overpredict the flux in the H₂O 21.15 μm line. The H12 model does not significantly overpredict any other line in the IRS spectrum, even if the best fit H12 fluxes are scaled to the level of the strongest IRS lines rather than to the far-IR PACS lines.

This analysis demonstrates that a high temperature, low density model (H12) of the H₂O emitting region can reasonably reproduce both the PACS and IRS spectra. On the other hand, a low temperature, high density model (W07) cannot reproduce the PACS spectrum, is inconsistent with the spatial distribution of the H₂O 63.4 μm line emission, and overpredicts the emission in several lines in the IRS spectrum.

4.3. CO excitation

In the CO excitation diagram, a cool ($280 \pm 30 \text{ K}$) component dominates mid-*J* lines and a warm ($880 \pm 100 \text{ K}$) component dominates high-*J* lines. The uncertainty in temperature includes the choice of energy levels to fit for the high and cool

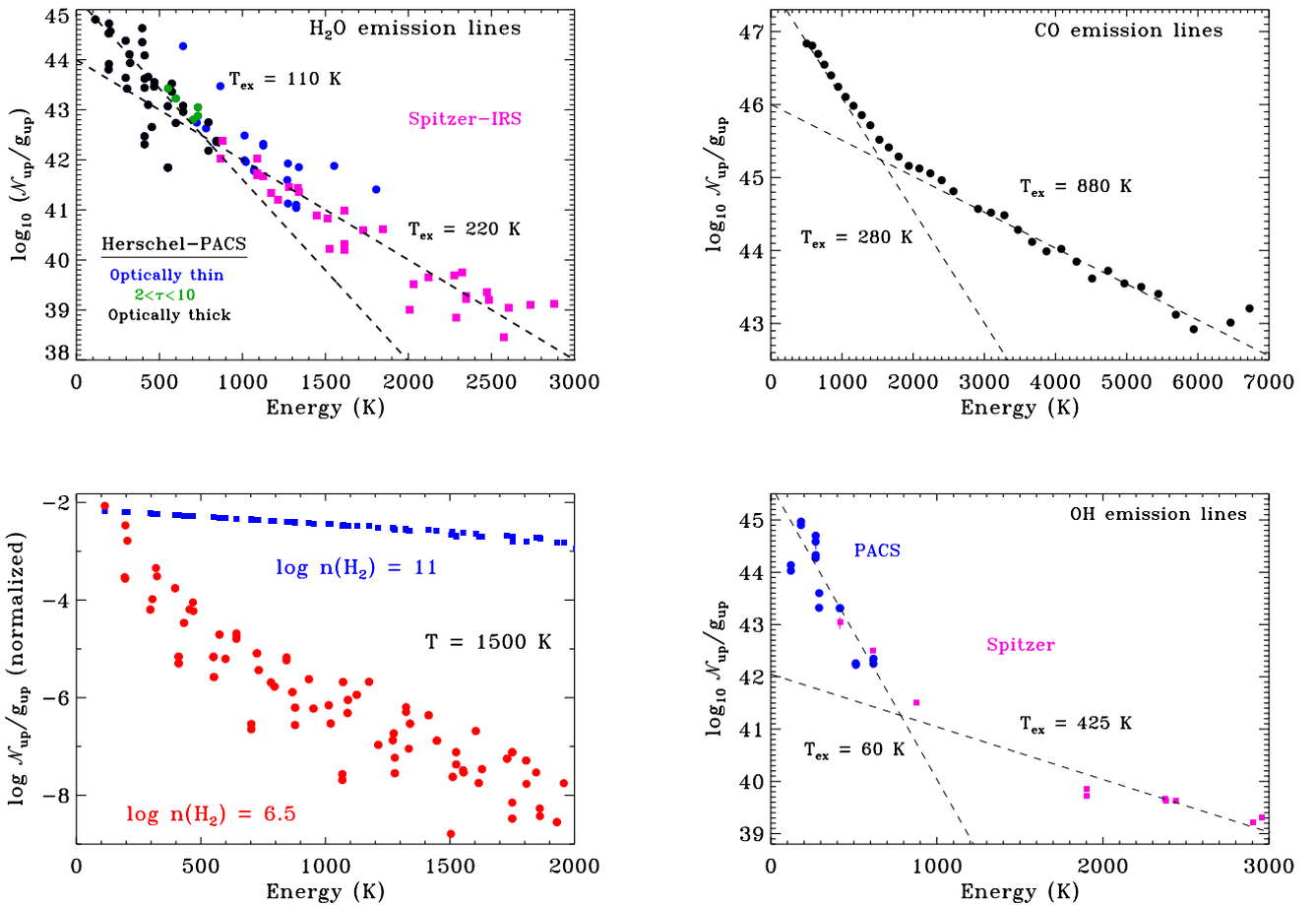


Fig. 11. Excitation diagrams, in units of total number of detected molecules N divided by degeneracy g , for H_2O (upper left), CO (upper right), and OH (lower right) emission lines detected with *Herschel*-PACS (circles) and *Spitzer*-IRS (purple squares). For the H_2O excitation diagram, PACS data are subdivided into lines that are optically-thin (blue), moderately optically-thick (green), and optically-thick (black), for the H12 model. Most of the *Spitzer* lines are optically-thin. Lower left: H_2O excitation diagrams obtained from RADEX models for optically thin models with $T = 1500$ K at high (blue) and low (red) density. At low density, subthermal excitation leads to cooler measured excitation temperatures and significant scatter in the level populations.

components. The two excitation components could relate to regions with different kinetic temperatures or with different densities. RADEX models of CO were run using molecular data obtained from LAMDA (Schöier et al. 2005; Yang et al. 2010) with an extrapolation of collision rates up to $J = 80$ by (Neufeld 2012).

The detection of high- J CO lines with an excitation temperature of ~ 880 K requires $\log n(\text{H}_2) > 6$ for reasonable kinetic temperatures (< 4000 K). For $T = 1500$ K, $\log n(\text{H}_2) = 6.5$, and an emitting area with radius 100 AU, the physical parameters adopted to explain the water emission, produces an excitation temperature of 950 K for lines with $J = 30\text{--}45$. This temperature is sensitive to the density, with $\log n(\text{H}_2) = 6.0$ leading to an excitation temperature of 640 K. The total number of CO molecules, N , for $\log n(\text{H}_2) = 6.5$ is $\log N = 48.4$.

In principle the two temperature components could relate to a single region with high temperature (~ 4000 K) and $\log n(\text{H}_2) < 4$ (Neufeld 2012). In this case, the CO emission would be physically unrelated to the highly excited H_2O emission, and the CO abundance in the highly excited H_2O emission region would be much smaller than that measured here.

4.4. OH excitation

The OH excitation diagram shows cool (60 ± 15 K) and warm (425 ± 100 K) components. RADEX models of the low excitation levels of OH ($E_{\text{up}} < 1000$ K) were run, using collisional rate coefficients from (Offer et al. 1994) and energy levels and Einstein A values from (Pickett et al. 1998). Molecular data for higher excitation levels were obtained from HITRAN (Rothman et al. 2009). A 1500 K gas with $\log n(\text{H}_2) = 6.5$ and emitting area of radius 100 AU roughly reproduces the emission in the cool component OH emission, with $\log N(\text{OH}) = 17.3$. Whether these parameters could also reproduce the highly excited OH emission is not clear.

The RADEX model fluxes are also somewhat discrepant with the observed fluxes, The $24.6 \mu\text{m}$ line flux is much lower than predicted. In addition, all detected OH doublets have similar line fluxes but the RADEX model predicts different fluxes in several transitions. A lower opacity, caused either by broader lines or a lower column density and larger emitting area, would alleviate some of these discrepancies.

The OH molecule has energy levels with high critical densities and with strong far-IR transitions to low energy levels

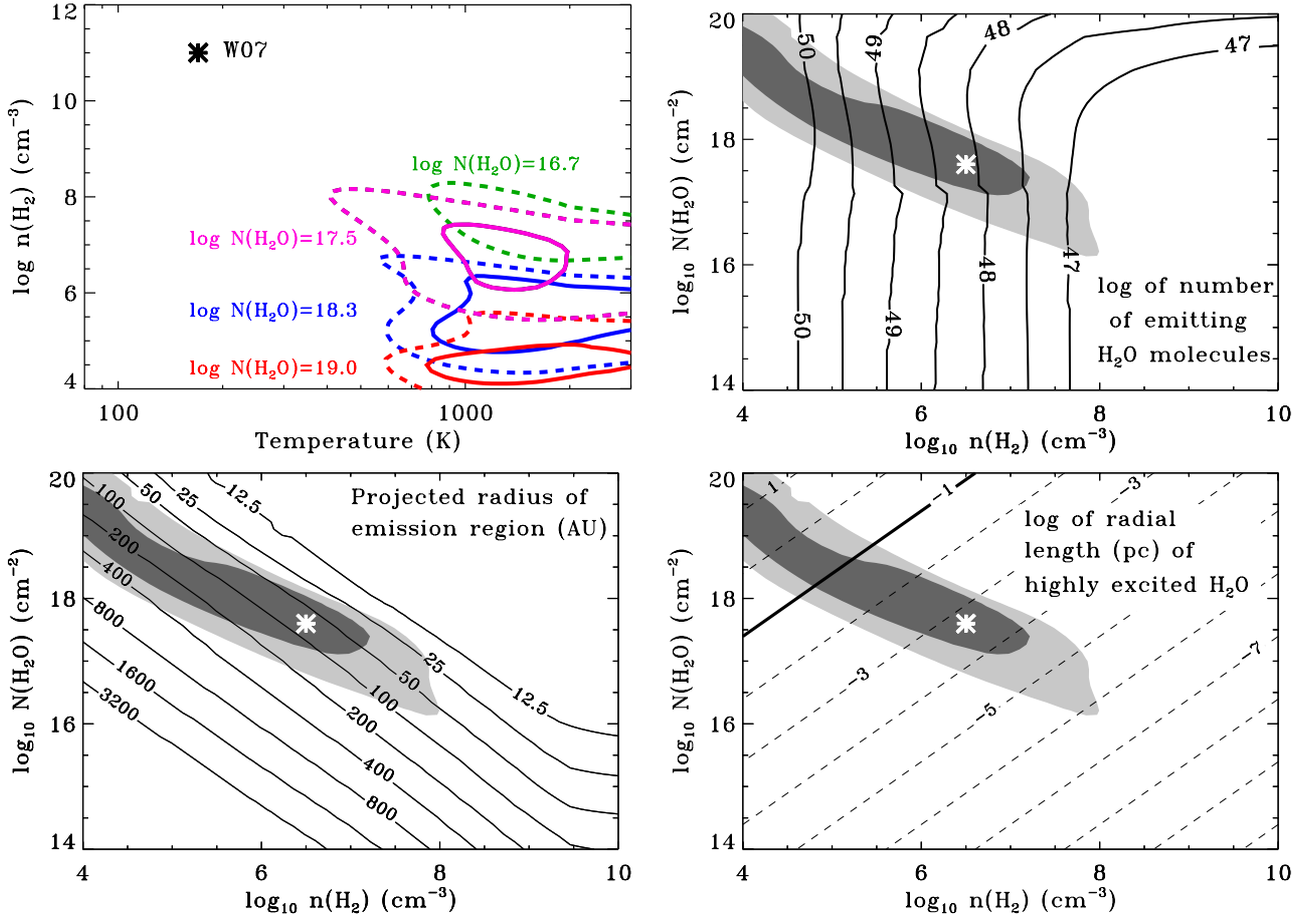


Fig. 12. *Upper left:* contours of χ^2 versus temperature and H_2 density for several different H_2O column densities (the different colors), with the minimum reduced $\chi^2 \sim 4$. The contours show where $\chi_{\text{red}}^2/4 = 1.2$ (solid lines) and 1.5 (dashed lines), which roughly indicate the acceptable parameter space. The contours shown here are calculated to fits of lines with $E' > 400$ K. The asterisks show the parameters of the sub-thermal model presented here (H12) and the high density envelope-disk accretion shock model (W07). *Upper right and lower panels:* contours of total number of H_2O molecules (*upper right*), radius (AU) of a circular emission area for H_2O emission (*lower left*), and length scale (log pc) of H_2O emission (*lower right*) compared with the best fit contours of $n(\text{H}_2)$ and $N(\text{H}_2\text{O})$ for $T = 1500$ K. The asterisks indicate the parameters adopted for this paper. In the lower right panel, the solid line at 0.1 pc shows the conservative upper limit to the length scale of the emission. The parameters adopted for this paper are located in the lower right of the acceptable parameter space so that the length scale for the emission is small.

that are favorable to IR photoexcitation. Radiative pumping may therefore severely alter the level populations (e.g. [Wampfler et al. 2010](#)). The IR pumping would increase the populations in excited levels, which may cause us to overestimate the total number of OH molecules for the given temperature and density. As with H_2O , a rigorous assessment of the IR pumping requires a full physical model of the envelope and is beyond the scope of this work.

4.5. O excitation

From the $63.18 \mu\text{m}$ line flux and assuming $T = 1500$ K, the total number of neutral O atoms is $\log N = 47.9$. This number is robust to changes in temperature and density within the parameter space discussed here, based on RADEX models of [O I] lines. If the [O I] emission is spread out over a circle of 100 AU in radius, the column density $\log N(\text{O}) = 16.5$ is much less than the column density required for the line to become optically thick ($\log N(\text{O}) \sim 19$ for a Gaussian profile with a FWHM of 25 km s^{-1}). These physical properties are adopted from the H_2O emission for simplicity but are likely incorrect

because the location of [O I] emission is spatially different than the H_2O emission (left panel of Fig. 6).

5. Discussion

5.1. The origin of highly-excited H_2O emission

Prior to this work, H_2O emission has been attributed to three different regions in or near the IRAS 4B environment:

- (1) *A compact disk:* spectrally narrow (FWHM $\sim 1 \text{ km s}^{-1}$) H_2^{18}O emission is produced in a (pseudo)-disk with a radius ~ 25 AU around IRAS 4B ([Jørgensen & van Dishoeck 2010](#)). From the inferred H_2^{18}O column density, the compact disk is optically-thick in most H_2^{16}O rotational transitions. The disk covers only a small area on the sky and therefore contributes very little emission to the broad lines detected with HIFI and to the mid- and far-IR H_2O emission. From the assumed $T = 170$ K and resulting column density $\log N(\text{p-H}_2\text{O}) \sim 18.4$ and $b = 1 \text{ km s}^{-1}$, the $\text{p-H}_2^{16}\text{O } 3_{31}-2_{02}$ $138.5 \mu\text{m}$ line, from the same upper level as the observed H_2^{18}O line, would have a flux of $3 \times 10^{-22} \text{ W m}^{-2}$, 50 times

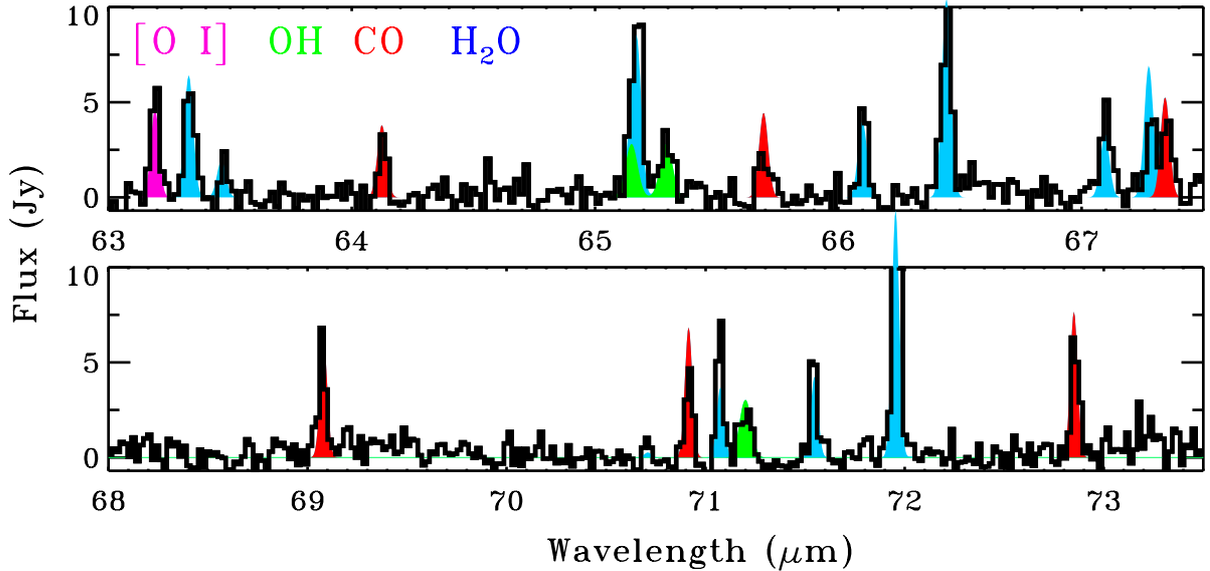


Fig. 13. Segments of the PACS spectrum (solid black line). The H_2O line fluxes (shaded in blue) are obtained from RADEX, CO line fluxes (red) from the thermal distribution in Fig. 11, and OH (green) and [O I] line fluxes (purple) from Gaussian fits.

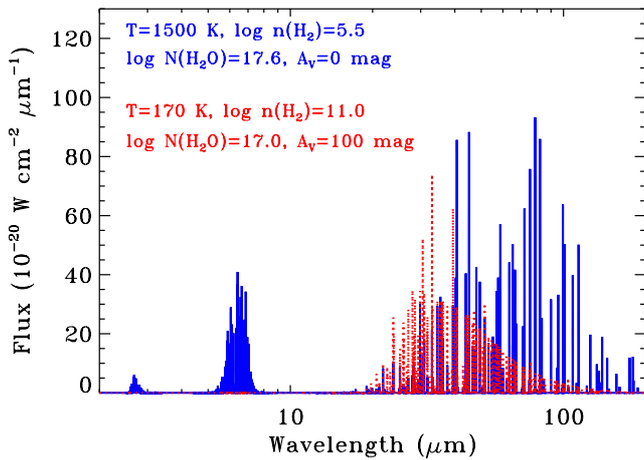


Fig. 14. The synthetic spectra for the H12 (blue) and W07 (red) models at a spectral resolution $R = 1500$. Because the W07 models are optically-thick in many highly-excited levels, the strongest H_2O lines peak in the mid-IR rather than the far-IR. The H12 model predicts some rovibrational emission at $6 \mu\text{m}$ because of the high temperature, with a strength that could further constrain the temperature and density of the highly excited H_2O .

smaller than the measured line flux. Our PACS observations are unable to probe this disk component.

- (2) *Outflow emission, (a) in low-excitation lines with a large beam:* spectrally-broad emission in low-excitation H_2O lines was observed in spatially-unresolved observations over a $\sim 20\text{--}40''$ beam and was attributed to outflows based on the line widths (Kristensen et al. 2010). The outflow is compact on the sky, so that both the red- and blue-shifted outflow lobes are located within this aperture. And (b), *in masers:* spectrally narrow ($\sim 1 \text{ km s}^{-1}$) and spatially compact H_2O maser emission is seen from dense gas in an outflow (e.g. Desmurs et al. 2009). The relationship between the maser emission and the molecular outflow is unclear.
- (3) *Disk-envelope shock:* H_2O emission in highly-excited mid-IR lines, which are spectrally and spatially-unresolved at

low spectral and spatial resolution, was attributed to the disk-envelope accretion shock (Watson et al. 2007). The primary argument for the accretion shock is that high H_2 densities ($>10^{10} \text{ cm}^{-3}$) are needed to populate the highly excited levels, which have high critical densities. Such high densities are expected in an envelope-disk accretion shock but are physically unrealistic for an outflow-envelope accretion shock because envelope densities are much lower than disk densities.

Our primary goals in this work are to use the spatial distribution and excitation of warm H_2O emission to test (3), the envelope-disk accretion shock interpretation proposed by Watson et al. (2007), and to subsequently use the far-IR emission to probe the heating and cooling where the emission is produced. In the following subsections, we discuss the outflow origin of the highly-excited H_2O emission and subsequently discuss the implications for the envelope-disk accretion shock and outflows.

5.2. Outflow origin of highly excited H_2O emission

The H_2O emission detected with PACS is spatially offset from the peak of the far-IR continuum emission to the south, the direction of the blueshifted outflow. Of the mapped lines, the H_2O $8_{18-7_{07}}$ and $8_{08-7_{17}}$ lines at $63.4 \mu\text{m}$ are closest in excitation to the mid-IR *Spitzer* lines. The location of the emission in both $63.4 \mu\text{m}$ H_2O lines is consistent with the location of near-IR emission from IRAS 4B imaged with *Spitzer*/IRAC. The full PACS spectrum from $50\text{--}200 \mu\text{m}$ demonstrates that all other molecular lines are also offset from the location of the sub-mm continuum peak (Fig. 5). These lines are produced in the southern, blueshifted outflow lobe of IRAS 4B, as shown in the plots and cartoon of Figs. 6, 7. Contemporaneous to our work, Tappe et al. (2012) found that the spatial distribution of H_2O emission in *Spitzer*/IRS spectra is consistent with an outflow origin. The images of CO 24–23 and 49–48 also demonstrate that highly excited CO emission is produced in the blueshifted outflow lobe. The CO 49–48 emission is located closer to the central source than the highly excited H_2O emission.

The combined *Herschel*/PACS and *Spitzer*/IRS excitation diagram does not show any indication of multiple components in

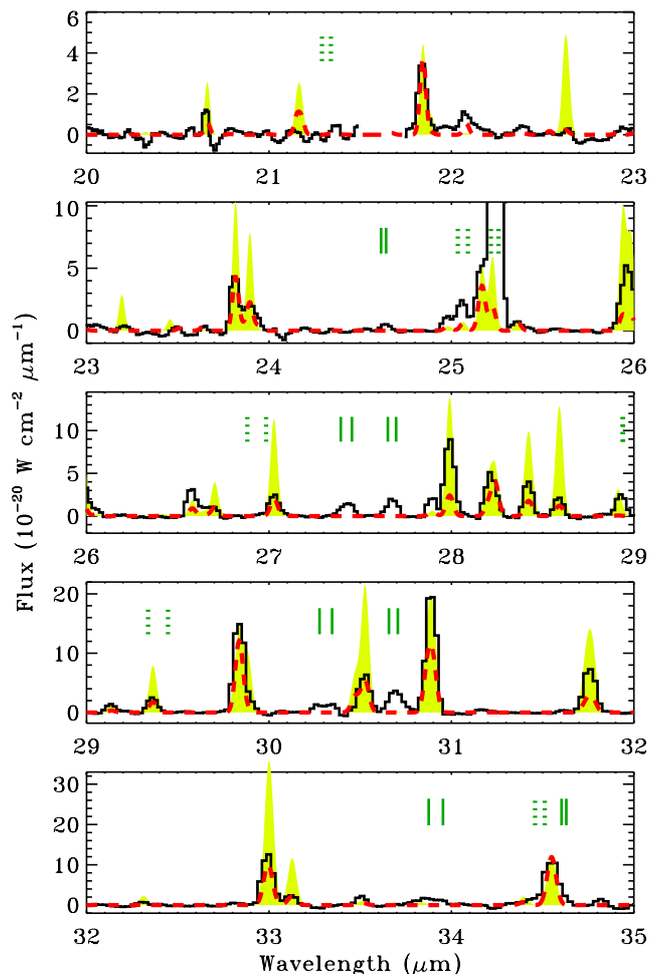


Fig. 15. A comparison of H12 (red dashed line, scaled to the flux in the 63.4 μm lines) and W07 (yellow filled regions, scaled to the flux of the 35.5 μm line) model spectra to the *Spitzer* IRS spectrum. Green vertical lines mark the wavelengths of OH lines that are detected (solid lines) or are too weak or blended to be detected (dotted lines). Both models reproduce most *Spitzer*/IRS lines reasonably well, with a few exceptions described in detail in the text.

the highly-excited H_2O lines. RADEX models indicate that the highly excited PACS lines are consistent with emission from a single slab of gas with $T \sim 1500$ K, $\log N(\text{H}_2\text{O}) \sim 17.6$, $\log n \sim 6.5$, and an emitting area equivalent to a circle with radius ~ 100 AU. The same parameters reproduce the mid-IR H_2O emission lines detected in the *Spitzer*/IRS spectrum of IRAS 4B. These same physical conditions may also produce CO fundamental emission, which could explain the bright IRAC 4.5 μm emission from the IRAS 4B outflow (Tappe et al. 2012).

The on-source component of the far-IR H_2O is not as well described as the outflow component because this component is faint in the short wavelength PACS lines, which are able to constrain the properties of the emission at the offset outflow position. The on-source emission suffers from higher extinction, so the brightest lines are at longer wavelengths, have low excitation energies, and are optically thick. Our RADEX modeling is restricted to the higher excitation component of the H_2O emission. However, the similarity of the ratio of the two 63.4 μm lines at the sub-mm continuum location and at the outflow location suggests similar excitations. The non-detection of H_2^{18}O lines (see Appendix C) place a strict limit on the optical depth of the H_2O lines. The H_2O 108.1 μm line, of which 70% is located

on-source, likely traces the same material as the HIFI spectra of low-excitation H_2O lines (Kristensen et al. 2010). The width of the HIFI emission (FWHM ~ 24 km s^{-1} , with wings that extend out to ~ 80 km s^{-1}) is consistent with an outflow origin and inconsistent with a slow (~ 2 km s^{-1}) envelope disk accretion shock that would be expected for infalling gas.

In sub-mm line emission, both the red- and blueshifted outflows are detected and spatially separated (Jørgensen et al. 2007; Yıldız et al. submitted), which indicates that the outflow is not aligned exactly along our line of sight to the central object. In previous near- and mid-IR imaging, only the blueshifted outflow is detected and the redshifted outflow is invisible. The extinction to the redshifted outflow is so high that even the far-IR emission is obscured by the envelope. The outflow therefore cannot be aligned in the plane of the sky and likely is aligned to within 45° of our line of sight to the central star. This large extinction is consistent with the outflow angle of $\sim 15^\circ$ relative to our line of sight, as inferred from assuming that the IRAS 4B outflow has the same age as the IRAS 4A outflow (Yıldız et al. 2012).

The relationship between the outflow and the maser emission is uncertain. The maser emission has a position angle of 151° from IRAS 4B (Park & Choi 2007; Marvel et al. 2008; Desmurs et al. 2009), in contrast to the $\sim 174^\circ$ position angle of the molecular outflow (Jørgensen et al. 2007). Unlike the molecular outflow, the maser emission is located in the plane of the sky ($\sim 77^\circ$ relative to our line of sight), based on the radial velocity and projected velocity on the sky. In this case, the outflow would have a dynamical age of only 100 yr. However, the high extinction to the redshifted outflow and low extinction to the blueshifted outflow together suggest that the outflow is aligned closer to our line of sight than in the plane of the sky. Jet precession, perhaps a result of binarity of the central object, has been suggested as a possible explanation for the difference between the maser emission and molecular outflow (Desmurs et al. 2009).

H_2O maser emission is produced in gas with densities of 10^7 – 10^9 cm^{-3} (Kaufman & Neufeld 1996). The density of the far-IR H_2O emission could be as high as $\sim 10^7$ cm^{-3} . However, the maser emission is produced in very small spots within the outflow, while the H_2O emission detected here is likely spread over a projected area equivalent to a circle with a radius ~ 100 AU. In principle, the maser emission could simply be the smallest, densest regions within the same outflow that produces the far-IR H_2O emission, although the maser and far-IR H_2 emission may be unrelated.

In summary, we conclude that most of the H_2O emission from IRAS 4B is produced in the blueshifted outflow. The highly excited H_2O emission is located at the blueshifted outflow position. The redshifted outflow is likely not detected in these lines because any emission is obscured by the envelope. The optically thick lower-excitation H_2O lines have an additional on-source component and are likely similar to the spectrally broad outflow seen by Kristensen et al. (2010). Similarly, the cool components of OH and CO emission are also likely produced in the outflow rather than a quiescent envelope.

5.3. Implications for envelope-disk accretion

The primary motivation for invoking the envelope-disk accretion shock to interpret the mid-IR H_2O lines was the high critical densities for the mid-IR H_2O lines. We have demonstrated that these lines may also be produced in gas with sub-thermal excitation. An additional component besides the outflow does not need to be invoked at present to explain the presence of the mid-IR H_2O emission.

Table 4. Summary of components for H₂O emission from IRAS 4B.

Emission source	Probe	Line profile	Opacity	Location
Compact disk	sub-mm H ₂ ¹⁸ O	Narrow, symmetric	optically thick	on-source
Off-source outflow	mid/far-IR H ₂ O	Unresolved	optically thin	off-source
Maser emission	mm H ₂ O	Narrow	high density	off-source
On-source outflow	far-IR H ₂ O	Symmetric	optically thin	on-source
Disk-envelope shock?	(sub)-mm?	Narrow (if present)	optically thick	on-source

The envelope-disk accretion shock may still have some undetected contribution to the on-source H₂O emission seen with PACS, in which case the mid-IR H₂O emission could in principle be the combination of the blueshifted outflow lobe and the envelope-disk accretion shock. The [Watson et al. \(2007\)](#) interpretation of the H₂O emission requires high line optical depths, including in the 63.4 μm H₂O lines. However, the 63.4 μm H₂O ortho/para lines are observed at the optically-thin flux ratio, with an upper limit that no more than 30% of the observed emission may be produced in optically thick gas. Since the envelope-disk accretion shock would only contribute to the on-source emission, the maximum contribution to the total flux in the 63.4 μm lines is $\sim 10\%$. In this case, we find that the luminosity of the disk-envelope accretion shock would be $1.5 \times 10^{-3} L_{\odot}$, 5% of that calculated by [Watson et al. \(2007\)](#)⁶. The accretion rate onto the disk would correspond with $\sim 3 \times 10^{-6} M_{\odot} \text{ yr}^{-1}$, which is likely too low to represent the main phase of disk growth.

An alternate and likely explanation is that any disk-envelope accretion shock is buried inside the envelope, with an extinction that prevents the detection of any mid-IR emission produced by such a shock. The on-source H₂O and CO emission is likely produced in an outflow that is seen behind $A_V \sim 700$ mag of extinction. In this case, whether the envelope-disk accretion shock exists is uncertain and the accretion rate is completely unconstrained. The prospects for studying envelope-disk accretion likely require a high-density ($>10^{11} \text{ cm}^{-3}$) tracer ([Blake et al. 1994](#)) observed at high spatial and spectral resolution in the sub-mm, where emission is not affected by extinction.

5.4. Gas line cooling budget and abundances for IRAS 4B

The bulk of the far-IR line emission is produced in the outflow. The emission is smoothly distributed within the outflow and is consistent with emission from two different positions, one located on-source and one located off-source.

Table 3 lists the total cooling budget attributed to the molecular and atomic emission in the far-IR, as extracted from the on-source position and the blueshifted outflow lobe. Most of the cooling is in molecular lines rather than atomic lines, which is consistent with previous estimates of far-IR cooling from a larger sample of Class 0 objects that were observed with ISO ([Nisini et al. 2002](#)).

In contrast to the luminous molecular lines, the [O I] emission from IRAS 4B is surprisingly faint. The [O I] 63 μm line is typically the brightest far-IR emission line from low-mass protostars, with an average luminosity of $10^{-3} L_{\text{bol}}$ for Class 0 stars and $10^{-2} L_{\text{bol}}$ for Class I stars ([Giannini et al. 2001](#); [Nisini et al. 2002](#)). For IRAS 4B, the [O I] flux is $10^{-3.5} L_{\text{bol}}$. The weak [O I] emission may be a signature of the youngest protostars with outflows that have not yet escaped the dense envelope, as may be the case for IRAS 4B. Although some [O I] emission could be

located behind the envelope for IRAS 4B and other Class 0 objects, the non-detection of the [O I] 145.5 μm line suggests that neglecting extinction does not lead to a serious underestimate in [O I] line luminosity. The PACS spectrum also does not show any evidence for high-velocity [O I] emission from the jet, unlike the more evolved embedded object HH 46 ([van Kempen et al. 2010a](#)).

The abundance ratio⁷ OH/H₂O ~ 0.2 is consistent with the abundance ratio OH/H₂O > 0.03 for the outflow from the high-mass YSO W3 IRS 5 ([Wampfler et al. 2011](#)), the only other YSO with such a measurement. The H₂O/CO abundance is ~ 1 , on the high end of the value of 0.1–1 found from emission in lower-excitation lines of CO and H₂O ([Kristensen et al. 2010](#)). The ratio of H₂O/H₂ is $\sim 10^{-4}$, based on the total number of H₂ molecules ($\log \mathcal{N}(\text{H}_2) = 52.2$) calculated from the extinction corrected fluxes of pure-rotational H₂ lines from ([Tappe et al. 2012](#)). The H₂ and H₂O emission trace the same projected region on the sky (right panel of Fig. 6, with the *Spitzer*/IRAC imaging dominated by H₂ emission).

The abundance ratio of O/H₂O is ~ 0.1 for the detected emission in the blueshifted outflow. The [O I] emission is predominantly located alongside the outflow axis and is offset from the location of the H₂O emission. Within the shock that produces the highly excited H₂O emission, the O/H₂O abundance ratio must be much lower than 0.1. The non-detection of [C II] emission suggests that the ionization fraction in the shock is low. At least 90% of the gas-phase O is in H₂O, CO, and OH, which is consistent with the high H₂O/H₂ ratio measured above.

Relative to other sources, the listed H₂O column density is $\sim 10^2$ – 10^4 times larger than that found from L1157 ([Nisini et al. 2010](#); [Vasta et al. 2012](#)). Some of this discrepancy might be attributed to methodology. The H₂O column density calculated here is measured directly from the opacities in many different lines, which are spectrally unresolved. In contrast, the H₂O column density is calculated from optically thick lines, many of which are spectrally resolved but measured with a range of beam sizes. However, the H₂O abundance from IRAS 4B may be much larger than that of most other young stellar objects, possibly because of some physical difference in the outflow properties.

5.5. Shock properties for the highly excited molecular outflow

The H₂O line cooling is qualitatively consistent with excitation of molecular gas predicted from C-shocks by [Kaufman & Neufeld \(1996\)](#). The OH emission also likely traces shocks because OH has a high critical density and is an intermediate in the high temperature gas phase chemistry that produces H₂O. The two-temperature shape of the CO excitation diagram is expected

⁶ The rate is also adjusted from [Watson et al. \(2007\)](#) for updated distance of 235 pc.

⁷ As calculated from the highly excited H₂O emission and the low temperature component of OH emission, both modeled with RADEX with the same temperature and density. It is unclear whether the warmer component of OH emission can be reproduced by these same parameters. The OH abundance may be severely affected by infrared pumping ([Wampfler et al. 2010](#)).

from models of shock and UV-excitation of outflow cavity walls (Visser et al. 2011) and is qualitatively consistent with similar observations of the Class I sources HH 46 and DK Cha (van Kempen et al. 2010ab). The ratio of H₂O to CO luminosity is higher for IRAS 4B than for the Class I source HH 46 and for the borderline Class I/II source DK Cha. These differences may indicate that shock heating of the envelope plays a more important role than UV heating during the Class 0 stage. Indeed, Visser et al. (2011) suggested an evolutionary trend that for more massive and denser (younger) envelopes the shock heating should dominate. The different evolutionary stages may not change the shape of the CO ladder, as the relative contributions of the mid-*J* (15–25) and high-*J* (25–50) CO lines are similar for DK Cha and IRAS 4B. The highly excited H₂O emission may also be especially strong from IRAS 4B because the outflow covers only a very compact area when projected on the sky.

Highly excited H₂O and OH lines were previously detected in mid-IR *Spitzer*/IRS spectra of the HH 211 bow shock, which is strong enough ($\sim 200 \text{ km s}^{-1}$) to dissociate molecules at the location of direct impact. Such a high velocity shock also produces strong UV radiation (see models by Neufeld & Dalgarno 1989 and observations by, e.g., Raymond et al. 1997 and Walter et al. 2003) that photodissociates molecules both upstream and downstream of the shock. In Fig. 2 of Tappe et al. (2008), many mid-IR OH emission lines are stronger than the H₂O lines from HH 211. In contrast, the mid-IR H₂O lines are much stronger than the mid-IR OH lines from IRAS 4B. In addition, the OH emission from HH 211 is detected from levels with much higher excitation energies than detected here, in a pattern that is consistent with prompt emission following production of OH in excited levels through photodissociation of H₂O by far-UV radiation (presumably Lyman α , Tappe et al., 2008). The atomic fine-structure lines are also brighter from the HH 211 outflow than from IRAS 4B (Giannini et al. 2001; Tappe et al. 2008). Thus, UV radiation likely plays a large role in both the chemistry and excitation of OH at the HH 211 shock position, which is well separated from the YSO itself and is outside the densest part of the envelope.

In the case of IRAS 4B, the combination of bright H₂O emission and faint [O I] emission suggests that the H₂O dissociation rate is low. Moreover, the OH emission from IRAS 4B is not seen from the very highly excited levels detected from HH 211. Thus, OH likely forms through the traditional route of high temperature ($>230 \text{ K}$) chemistry of reactions between O and H₂ to form OH. The OH can then collide with H₂ to form H₂O. These reactions control the oxygen chemistry in dense C-type shocks (Draine et al. 1983; Kaufman & Neufeld 1996). The balance between O, OH and H₂O in well-shielded regions depends primarily on the H/H₂ ratio of the gas. In dense photo-dissociation regions, these same high temperature reactions are effective in forming H₂O, but the strong UV field can drive H₂O back into O and H₂O (Sternberg et al. 1995). Indeed, for the Orion Bar, the very strong UV field leads to a higher OH/H₂O abundance ratio (OH/H₂O > 1) than detected here (OH/H₂O ~ 0.1), in addition to emission in other diagnostics of strong UV radiation (strong [C II] and CH⁺ emission) that are detected from the Orion Bar (Goicoechea et al. 2011) but are not detected from IRAS 4B. Thus, the C-shock that produces the OH and H₂O emission from IRAS 4B is likely not irradiated by UV emission. The shocked gas may be shielded from any UV emission produced by the central source and internal shocks within the jet.

This shielding lends support to a C-type shock explanation rather than molecular formation downstream of a dissociative J-type shock because such a shock would produce UV emission

(see discussion above). A non-dissociative shock is also consistent with the presence of H₂ emission from IRAS 4B (Arnold et al. 2011; Tappe et al. 2012). This scenario is different than that postulated for HH 46 by (van Kempen et al. 2010a), where the on-source O and OH emission was thought to be produced by a fast dissociative J-type shock based on the different spatial extents of OH and H₂O. This scenario also differs from the interpretation of Wampfler et al. (2011) that the OH and H₂O emission from the high-mass YSO W3 IRS 5 arises either in a J-type shock or in a UV-irradiated C-shock. While the J-type shock or UV-irradiated C-type shock is an unlikely explanation for the H₂O emission, the OH emission could be produced in a different location than the H₂O emission, as is the case for [O I].

6. Conclusions

We have analyzed *Herschel*/PACS spectral images of far-IR H₂O emission from the prototypical Class 0 YSO IRAS 4B. Table 5 summarizes the different components of H₂O emission from IRAS 4B. We obtained the following results:

- 1) A rich forest of highly-excited H₂O, OH, and CO emission lines is detected in the blueshifted outflow from IRAS 4B. The spectrum is more line-rich than any other low-mass YSO that has been previously published.
- 2) Nyquist-sampled spectral maps place the highly-excited 63.4 μm lines at a average distance of $5''.2$ (projected distance of 1130 AU) south of IRAS 4B. The lower-excitation H₂O 108.1 μm line has a centroid closer to the peak of the sub-mm continuum emission and has a larger spatial extent than the 63.4 μm emission along the outflow axis. The far-IR H₂O emission can be interpreted as one component located at the blueshifted outflow position and a second component at the peak position in the mass distribution (sub-mm continuum peak). The redshifted outflow lobe is not detected in highly-excited H₂O emission, likely because of a high extinction ($>1500 \text{ mag}$) through to the back side of the envelope.
- 3) The highest excitation lines detected with PACS are optically-thin, indicating an ortho-to-para ratio of ~ 3 . RADEX models of the highly excited H₂O lines indicate that the emission is produced in gas described by $T \sim 1500 \text{ K}$, $\log n(\text{H}_2) \sim 6.5 \text{ cm}^{-3}$, and $\log N(\text{H}_2\text{O}) \sim 17.6 \text{ cm}^{-2}$ over an emitting area equivalent to a circle with radius $\sim 100 \text{ AU}$. These same physical parameters can reproduce the mid-IR H₂O emission seen with *Spitzer*/IRS and the CO and OH emission seen with PACS. The total mass of warm H₂O in the IRAS 4B outflow is about 140 times the amount of water on Earth.
- 4) From results (2) and (3), we conclude that the bulk of the far-IR H₂O emission is produced in outflows. Any contribution of the envelope-disk accretion shock to highly-excited H₂O lines is minimal. Moreover, at present the mid-IR H₂O emission does not offer any support for the presence of an envelope-disk accretion shock, as had been previously suggested by Watson et al. (2007). Any mid-IR H₂O emission produced by the envelope-disk accretion shock is likely deeply embedded in the envelope.
- 5) In the blueshifted outflow lobe over 90% of the gas phase O is in the H₂O, CO, and OH molecules rather than in neutral O. The H₂O is twice as abundant as CO and 10 times more abundant than OH. The cooling budget for gas in the envelope around IRAS 4B is dominated by H₂O emission.

- 6) The H₂O emission traces high densities in non-dissociative C-shocks. In contrast, much of the heating of lower mass envelopes occurs by energetic radiation. The highly excited H₂O in the shock-heated gas is likely shielded from UV radiation produced by both the central star and the bow shock. The OH likely forms through reactions between O and H₂ and provides a pathway to form H₂O.

Acknowledgements. G.J.H. thanks Achim Tappe for interesting and extensive discussions regarding the location of H₂O in the *Spitzer*/IRS spectra and the outflow inclination, and for detailed comments on the manuscript. G.J.H. also thanks Tom Megeath for discussions regarding the comparison of *Spitzer*/IRS and *Herschel*/PACS spectra, and Mario Tafalla, Javier Goicoechea, the anonymous referee, and the editor, Malcolm Walmsley for careful reads of the manuscript and insightful comments, Per Bjerkelli, Jeong-Eun Lee, and Doug Johnstone for some useful comments, and Stefani Germanotta for help in preparing the manuscript. We also thank Javier Goicoechea for providing us with the SPIRE spectrum of Ser SMM1 and for use of some DIGIT data to help calibrate the PACS spectrum at long wavelengths. Astrochemistry in Leiden is supported by NOVA, by a Spinoza grant and grant 614.001.008 from NWO, and by EU FP7 grant 238258. The research of KJ is supported by a Lundbeck Foundation Junior Group Leader Fellowship and by the Danish Research Council through the Centre for Star and Planet Formation.

References

- Adams, F. C., Lada, C. J., & Shu, F. H. 1987, *ApJ*, 312, 788
 André, P., Ward-Thompson, D., & Barsony, M. 1993, *ApJ*, 406, 122
 Arce, H., & Sargent, A. I. 2006, *ApJ*, 646, 1070
 Arnold, L. A., Watson, D. M., Kim, K. H., et al. 2011, *ApJ*, submitted [arXiv:1107.3261v2]
 Bergin, E. A., Kaufman, M. J., Melnick, G. J., Snell, R. L., & Howe, J. E. 2003, *ApJ*, 582, 830
 Blake, G. A., van Dishoeck, E. F., Jansen, D. J., Groesbeck, T. D., & Mundy, L. G. 1994, *ApJ*, 428, 680
 Bontemps, S., Andre, P., Terebey, S., & Cabrit, S. 1996, *A&A*, 311, 858
 Chapman, N. L., Mundy, L. G., Lai, S.-P., & Evans, N. J. 2009, *ApJ*, 690, 496
 Choi, M., & Lee, J.-E. 2011, *JKAS*, 44, 201
 Curiel, S., Raymond, J. C., Wolfire, M., et al. 1995, *ApJ*, 453, 322
 Desmurs, J.-F., Codella, C., Santiago-Garcia, J., Tafalla, M., & Bachiller, R. 2009, *A&A*, 498, 753
 Di Francesco, J., Myers, P. C., Wilner, D. J., Ohashi, N., & Mardones, D. 2001, *ApJ*, 562, 770
 Draine, B. T., Roberge, W. G., & Dalgarno, A. 1983, *ApJ*, 264, 485
 Faure, A., & Josselin, E. 2008, *A&A*, 492, 257
 Faure, A., Crimier, N., Ceccarelli, C., et al. 2007, *A&A*, 472, 1029
 Furuya, R. S., Kitamura, Y., Wootten, A., Claussen, M. J., & Kawabe, R. 2003, *ApJS*, 144, 71
 Giannini, T., Nisini, B., & Lorenzetti, D. 2001, *ApJ*, 555, 40
 Goicoechea, J. R., Joblin, C., Contursi, A., et al. 2011, *A&A*, 530, L16
 Gutermuth, R. A., Myers, P. C., Megeath, S. T., et al. 2008, *ApJ*, 674, 336
 Herczeg, G. J., Brown, J. M., van Dishoeck, E. F., & Pontoppidan, K. M. 2011, *A&A*, 533, A112
 Hirota, T., Bushimata, T., Choi, Y. K., et al. 2008, *PASJ*, 60, 37
 Horne, K. 1986, *PASP*, 98, 609
 Indebetouw, R., Mathis, J. S., Babler, B. L., et al. 2005, *ApJ*, 619, 931
 Jørgensen, J. K., & van Dishoeck, E. F. 2010, *ApJ*, 710, L72
 Jørgensen, J. K., Schöier, F. L., & van Dishoeck, E. F. 2002, *A&A*, 389, 908
 Jørgensen, J. K., Harvey, P. M., Evans, N. J., et al. 2006, *ApJ*, 645, 1246
 Jørgensen, J. K., Bourke, T. L., Myers, P. C., et al. 2007, *ApJ*, 659, 479
 Jørgensen, J. K., van Dishoeck, E. F., Visser, R., et al. 2009, *A&A*, 507, 861
 Kaufman, M. J., & Neufeld, D. A. 1996, *ApJ*, 456, 611
 Kirk, H., Johnstone, D., & Di Francesco, J. 2006, *ApJ*, 646, 1009
 Kristensen, L. E., Visser, R., van Dishoeck, E. F., et al. 2010, *A&A*, 521, L30
 Liseau, R., Justtanont, K., & Tielens, A. G. G. M. 2006, *A&A*, 446, 561
 Maret, S., Bergin, E. A., Neufeld, D. A., et al. 2009, *ApJ*, 698, 1244
 Marvel, K. B., Wilking, B. A., Claussen, M. J., & Wootten, A. 2008, *ApJ*, 685, 285
 Neufeld, D. A. 2010, *ApJ*, 708, 635
 Neufeld, D. A. 2012, *ApJ*, submitted
 Neufeld, D. A., & Dalgarno, A. 1989, *ApJ*, 344, 251
 Neufeld, D. A., & Hollenbach, D. J. 1994, *ApJ*, 428, 170
 Neufeld, D. A., & Yuan, Y. 2008, *ApJ*, 678, 974
 Nisini, B., Benedettini, M., Giannini, T., et al. 2000, *A&A*, 360, 297
 Nisini, B., Giannini, T., & Lorenzetti, D. 2002, *ApJ*, 574, 246
 Nisini, B., Benedettini, M., Codella, C., et al. 2010, *A&A*, 518, L120
 Offer, A. R., van Hemert, M. C., & van Dishoeck, E. F. 1994, *J. Chem. Phys.* 100, 362
 Ott, S. 2010, *ASPC*, 434, 139
 Park, G., & Choi, M. 2007, *ApJ*, 664, L99
 Pickett, H. M., Poynter, I. R. L., Cohen, E. A., et al. 1998, *JQSRT*, 60, 883
 Pilbratt, G. L., Riedinger, J. R., Passvogel, T., et al. 2010, *A&A*, 518, L1
 Poglitsch, A., Waelkens, C., Geis, N., et al. 2010, *A&A*, 518, L2
 Pontoppidan, K. M., Saylk, C., Blake, G. A., et al. 2010, *ApJ*, 720, 887
 Raymond, J. C., Blair, W. P., & Long, K. S. 1997, *ApJ*, 489, 314
 Rodriguez, L. F., Anglada, G., Torrelles, J. M., et al. 2002, *A&A*, 389, 572
 Rothman, L. S., Gordon, I. E., Barbe, A., et al. 2009, *JQSRT*, 110, 533
 Schöier, F. L., van der Tak, F. F. S., van Dishoeck, E. F., & Black, J. H. 2005, *A&A*, 432, 369
 Shu, F. H. 1977, *ApJ*, 214, 488
 Snell, R. L., Loren, R. B., & Plambeck, R. L. 1980, *ApJ*, 239, L17
 Spaans, M., Hogerheijde, M. R., Mundy, L. G., & van Dishoeck, E. F. 1995, *ApJ*, 455, L167
 Sternberg, A., & Dalgarno, A. 1995, *ApJS*, 99, 565
 Sturm, B., Bouwman, J., Henning, Th., et al. 2010, *A&A*, 518, L129
 Tappe, A., Lada, C. J., Black, J. H., & Muench, A. A. 2008, *ApJ*, 680, L117
 Tappe, A., Forbrich, J., Martin, S., Yuan, Y., & Lada, C. J. 2012, *ApJ*, submitted
 Terebey, S., Shu, F. H., Cassen, P. 1984, *ApJ*, 286, 529
 Tennyson, J., Zobov, N. F., Williamson, R., Polyansky, O. L., & Bernath, P. F. 2001, *Phys. Chem. Ref. Data*, 30, 735
 Tobin, J. J., Hartmann, L., Looney, L. W., & Chiang, H.-F. 2010, *ApJ*, 712, 1010
 van der Tak, F. F. S., Black, J. H., Schöier, F. L., Jansen, D. J., & van Dishoeck, E. F. 2007, *A&A*, 468, 627
 van Dishoeck, E. F., Kristensen, L. E., Benz, A. O., et al. 2011, *PASP*, 123, 138
 van Kempen, T. A., van Dishoeck, E. F., Hogerheijde, M. R., & Güsten, R. 2009, *A&A*, 508, 259
 van Kempen, T. A., Kristensen, L. E., Herczeg, G. J., et al. 2010a, *A&A*, 518, L121
 van Kempen, T. A., Green, J. D., Evans, N. J., et al. 2010b, *A&A*, 518, L128
 Vasta, M., Codella, C., Lorenzani, A., et al. 2012, *A&A*, 537, A98
 Velusamy, T., Langer, W. D., & Goldsmith, P. F. 2002, *ApJ*, 565, L43
 Visser, R., van Dishoeck, E. F., Doty, S. D., & Dullemond, C. P. 2009, *A&A*, 495, 881
 Visser, R., Kristensen, L. E., Bruderer, S., et al. 2012, *A&A*, 537, A55
 Vorobyov, E. I. 2011, *ApJ*, 729, 146
 Walter, F. M., Herczeg, G. J., Brown, A., et al. 2003, *AJ*, 126, 3076
 Wampfler, S. F., Herczeg, G. J., Bruderer, S., et al. 2010, *A&A*, 521, L36
 Wampfler, S. F., Bruderer, S., Kristensen, L. E., et al. 2011, *A&A*, 531, L16
 Watson, D. M., Bohac, C. J., Hull, C., Forrest, W. J., et al. 2007, *Nature*, 448, 1026
 Weingartner, J. C., & Draine, B. T. 2001, *ApJ*, 548, 296
 Whitney, B. A., & Hartmann, L. 1993, *ApJ*, 402, 605
 Yang, B., Stancil, P. C., Balakrishnan, N., Forrey, R. C. 2010, *ApJ* 718, 1062
 Yıldız, U. A., van Dishoeck, E. F., Kristensen, L. E., et al. 2010, *A&A*, 521, L40
 Yıldız, U. A., et al. 2012, submitted

Appendix A: Calibration of first and second order light longward of 190 μm

PACS spectra are poorly calibrated in regions where light from different orders are recorded at the same physical location on the detector, especially between 97–103 μm and longward of 190 μm . Most photons between 97–105 μm get dispersed into the second order and contaminate the flux at $>190 \mu\text{m}$ because of a mismatch between the grating and the filter transmission. Similarly, the light at 97–103 μm can be contaminated by third-order emission at $\sim 69 \mu\text{m}$. As a consequence, the first order light at 97–103 μm has low S/N and that light and the light at $>190 \mu\text{m}$ has not previously been flux calibrated. In this appendix, we use PACS spectra of Serpens SMM 1 (Goicoechea et al., in prep.) and HD 100546 (Sturm et al. 2010), both reduced in the same method as IRAS 4B, to calibrate the first and second order emission from PACS at $\lambda > 190 \mu\text{m}$. The continuum emission from HD 100546 is produced by a disk and peaks (in Jy) at $\sim 60 \mu\text{m}$. The continuum emission from Serpens SMM 1 is produced in an envelope and peaks at $\sim 150 \mu\text{m}$.

First and second order light can be separated because the diffraction-limited point spread function is twice as large at 200 μm as at 100 μm . Figure A.1 shows the fraction of flux in the central spaxel divided by the flux in the central 3×3 spaxels. This fraction decreases smoothly at $>100 \mu\text{m}$. The fraction starts to rise at $\sim 190 \mu\text{m}$ because of a contribution from second order emission. At each wavelength, this fraction directly leads to a ratio of first order photons to second order photons recorded by PACS. The second order light is then calibrated by measuring the first order light between 90–110 μm for HD 100546. Most of the light recorded at $>190 \mu\text{m}$ from HD 100546 is second order light.

This calibration is then applied in reverse to the spectrum of Serpens SMM 1 to subtract the second order emission, leaving only first order emission at $>190 \mu\text{m}$. A SPIRE spectrum of Serpens SMM 1 (Goicoechea et al., in prep.) is then used to flux calibrate the PACS spectrum from 190–210 μm . Despite the red spectrum of Serpens SMM 1, $\sim 50\%$ of the photons recorded at 202 μm are second-order 101 μm photons.

To calibrate the PACS spectrum of IRAS 4B, the first and second order light are initially separated based on the point spread function for both wavelengths (Fig. A.1). The separate first and second order spectra are then flux calibrated using the relationships calculated above. Lines are identified as first or second order emission after searching for the correct line identification at λ and $\lambda/2$.

Figure A.2 shows the resulting spectrum of IRAS 4B at 95–105 μm . Analysis of a large sample of PACS spectra from the DIGIT program (P. I. N. Evans) suggest that our flux calibration has an uncertainty of $\sim 20\%$ between 98–103 μm , in addition to other sources of uncertainty in the standard PACS flux calibration. The line fluxes measured from second order light at $>190 \mu\text{m}$ are consistent to 20% of the fluxes measured directly at 97–103 μm but with smaller error bars. The accuracy of the flux calibration at $>190 \mu\text{m}$ has not been evaluated but is likely uncertain to $\sim 40\%$.

The best place to observe lines between 98–103 μm with PACS is in the second order, despite overlap with first-order light because of sensitivity and higher spectral resolution. The PACS sensitivity to first order photons at $>200 \mu\text{m}$ is low. This new calibration allows us to improve the accuracy of flux measurements for lines between 98–103 μm and at $>190 \mu\text{m}$.

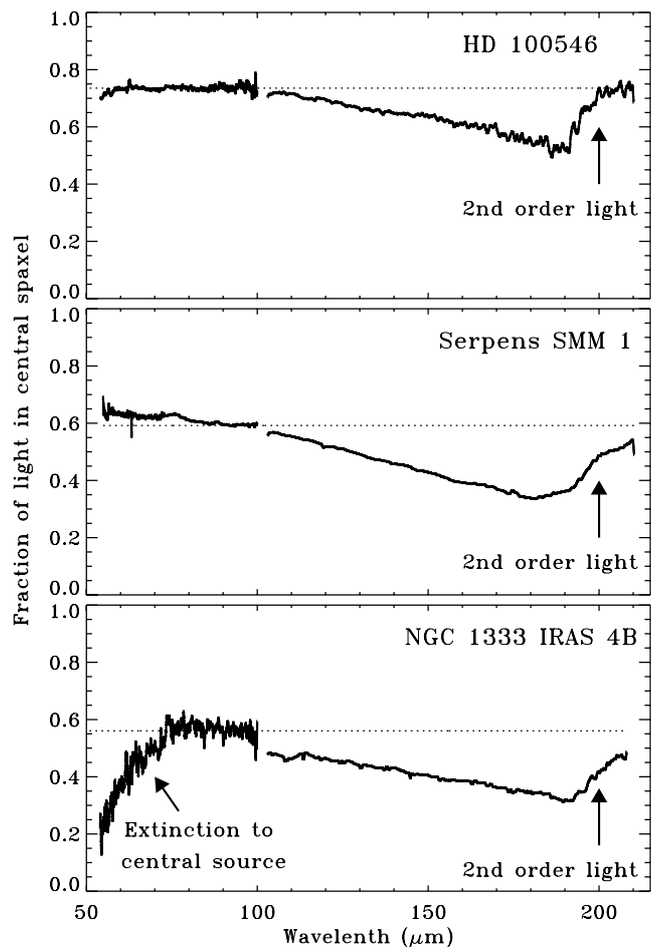


Fig. A.1. The ratio of flux in the central spaxel to the flux in the central 3×3 spaxels for HD 100546, Serpens SMM 1, and NGC 1333 IRAS 4B. The ratio falls linearly above $>100 \mu\text{m}$ until 190 μm . Second order light contaminates the spectrum at $>190 \mu\text{m}$. Because the first and second order light have different point-spread functions, the plotted ratio determines the fraction of first and second order photons versus wavelength. For NGC 1333 IRAS 4B, the continuum flux in the central spaxel falls at $<70 \mu\text{m}$, likely because of extinction.

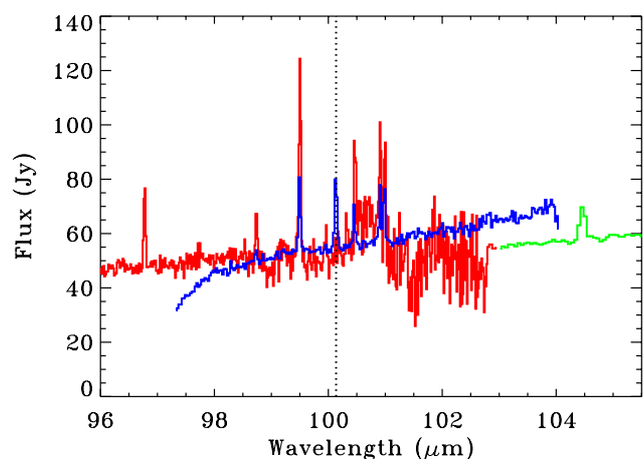


Fig. A.2. The first and second order spectrum of IRAS 4B at $\sim 100 \mu\text{m}$. The second order light at 100 μm (blue), calibrated from the $\sim 200 \mu\text{m}$ spectrum, matches the standard first and second order spectra (red and green). The lines in the blue spectrum are artificially weak, by a factor equivalent to the ratio of first to second order light, because the lines and continuum are treated separately. The dotted vertical line shows the location of the CO 13–12 line at 200.27 μm .

Table B.1. Unidentified lines^a.

λ_{obs} (μm)	Flux ^b	Error ^b	Possible ID		λ_{vac} (μm)
57.930	0.38	0.10	–	–	–
60.863	0.27	0.14	–	–	–
71.809	0.34	0.09	p-H ₂ O	5 ₅₁ –6 ₂₄	71.787
77.729	0.23	0.08	o-H ₂ O	7 ₅₂ –7 ₄₃	77.761
95.799	0.54	0.13	–	–	–
109.301	0.39	0.13	o-H ₂ ¹⁸ O	2 ₁₂ –1 ₁₀	109.346
112.846	0.24	0.12	o-H ₂ O	4 ₄₁ –5 ₁₄	112.802
127.164	0.21	0.08	–	–	–
129.885	0.25	0.08	–	–	–
147.380	0.28	0.08	–	–	–
170.017	0.26	0.07	p-H ₂ O	6 ₃₃ –6 ₂₄	170.138
181.595	0.32	0.14	–	–	–

Notes. ^(a) These lines may not be real. ^(b) 10^{-20} W cm⁻² s⁻¹.

Appendix B: Unidentified lines

All strong lines in the IRAS 4B spectrum are identified. Table B.1 lists several possible lines that are detected at the $\sim 3\sigma$ significance level. The position of several of these tentative detections do not correspond to expected emission lines. Although no o-H₂¹⁸O are clearly detected, the o-H₂¹⁸O 2₁₂–1₁₀ 109.346 μm line is expected to be among the strongest H₂¹⁸O lines and is tentatively detected. The centroid of the detected emission is -170 km s⁻¹ from the expected centroid, based on the measured wavelengths of other nearby lines. Emission is detected consistent with the location of the p-H₂O 5₅₁–6₂₄ 71.787 μm line, but the line flux is expected to be very weak.

These lines are all very weak and may be statistical fluctuations in the spectrum rather than significant detections of unidentified lines.

Appendix C: Exploring the parameter space in model fits to H₂O line fluxes

In Sect. 4, we characterized the physical properties of the H₂O emission region by comparing the observed line fluxes to synthetic fluxes calculated in RADEX models of a plane-parallel slab. In this appendix, we describe how the measured line fluxes constrain the parameters of the highly excited gas.

The backbone o-H₂O lines 9₀₉–8₁₈, 8₁₈–7₀₇, 7₀₇–6₁₆, 6₁₆–5₀₅, 5₀₅–4₁₄ and their p-H₂O counterparts are amongst the best diagnostics for the excitation and optical depth of the H₂O emitting region. The relative flux calibration between these ortho/para pairs should be accurate to better than 5% because the lines are located very near each other. The flux ratio ranges from 2.8–3.5 in the three of the backbone line ratios with highest excitation, indicating that the ortho-to-para ratio of ~ 3 is thermalized and that these lines are optically-thin. The 6₁₆–5₀₅, 5₀₅–4₁₄, and 4₁₄–3₀₃ and their p-H₂O counterparts have flux ratios of ~ 2.3 –2.5, suggesting that either the modeled emission is moderately optically thick in these lines, or that a second, optically-thick component contributes some emission to the flux in lower-excitation, longer wavelength lines. The top panels of Fig. C.1 show the acceptable space of $n(\text{H}_2)$ and T for two different values of $N(\text{H}_2\text{O})$ based on two of these line ratios.

The χ^2 statistic automates this type of analysis over all of the applicable lines to find the best-fit parameters for a single isothermal slab. Limiting the χ^2 calculation to lines with $E' > 600$ K yields two acceptable parameter spaces, one at the H12 location ($T \sim 1500$ K, $\log n(\text{H}_2) \sim 6.5$, and $\log N(\text{H}_2\text{O}) \sim 17.6$)

and one at low-temperature, high density, and low column density (hereafter X11, with $T \sim 200$ K, $\log n(\text{H}_2) \sim 11$, and $\log N(\text{H}_2\text{O}) \sim 14.0$). The X11 parameters differ from W07 because W07 includes a large H₂O column density to produce optically thick lines. The X11 parameters are disqualified because several lines, such as o-H₂O 3₃₀–2₂₁ 66.44 μm , have a synthetic flux that is much stronger than the observed emission⁸. As a consequence, the χ^2 statistic for all lines with $E' > 400$ K yields acceptable line fluxes only around the H12 parameters, adopted in this paper, and disqualifies the X11 parameter space.

The o-H₂O transitions 5₅₀–5₄₁ at 75.91 μm and 6₅₂–6₄₃ at 75.83 μm have a 2σ upper limit on the combined flux of 3×10^{-21} W cm⁻². Although these upper limits are not included in the χ^2 calculations, the non-detections confirm that the parameter space with high H₂ density and high H₂O column density cannot explain the bulk of the far-IR H₂O emission (lower left panel of Fig. C.1). Similarly, the line ratio o-H₂O 8₄₅–7₃₄ 35.67 μm to o-H₂O 7₀₇–6₁₆ 71.95 yields physical parameters consistent with the best-fit solution (lower right panel of Fig. C.1).

The non-detection of emission in H₂¹⁸O lines (with the possible exception of o-H₂¹⁸O 2₁₂–1₁₀ 109.35 μm) also places a limit on the opacity of H₂¹⁶O lines. The bottom right panel of Fig. C.1 shows where the flux ratio of o-H₂¹⁸O 3₂₁–2₁₂ 75.87 μm to o-H₂O 6₁₆–5₀₅ 82.03 μm becomes > 50 , large enough that the H₂¹⁸O line would be detected. This upper limit rules out the parameter space of high density and high column density.

In the H12 model, the strength of the 6 μm continuum emission (Maret et al. 2009) is similar to the strength of the 6 μm rovibrational H₂O lines. The non-detection with $R = 60$ spectra is marginally consistent with the predicted emission and rules out temperatures higher than ~ 2000 K. An $A_V = 20$ mag. would reduce the predicted 6 μm emission by 35%. Decreasing the temperature from 1500 K to 1000 K would reduce the predicted 6 μm emission by a factor of 4.8. Measurements of emission in H₂O vibrational lines would place significant additional constraints on the properties of the H₂O emission.

Appendix D: Effect of extinction estimates on rotational diagrams

Extinction can affect the derivation of excitation temperatures and line luminosities, especially at shorter wavelengths where lines typically have upper levels with higher energies. In this analysis, we assume that for all H₂O lines, 70% of the luminosity is produced on source behind $A_V = 700$ mag. and 30% is produced off-source at the blueshifted outflow lobe, with $A_V = 0$ mag. (see also Sect. 4.3). This description is consistent with the different observed spatial distributions of the H₂O 108.1 and 63.4 μm lines. The excitation conditions and emission line ratios may instead vary smoothly with distance in the outflow.

Applying an extinction correction under these assumptions increases the excitation temperature of the warm (cool) CO component from 850 to 950 K (250 to 280 K). If all the CO emission were located behind the $A_V = 700$ mag., then the temperature difference would become much larger, but the short wavelength CO lines would be very faint. The H₂O temperature does not change significantly because the fits are based mostly

⁸ The opposite case, where the synthetic emission is weaker than observed in a low-excitation line, would not disqualify a model because low-excitation line emission may include an additional contribution from cooler gas.

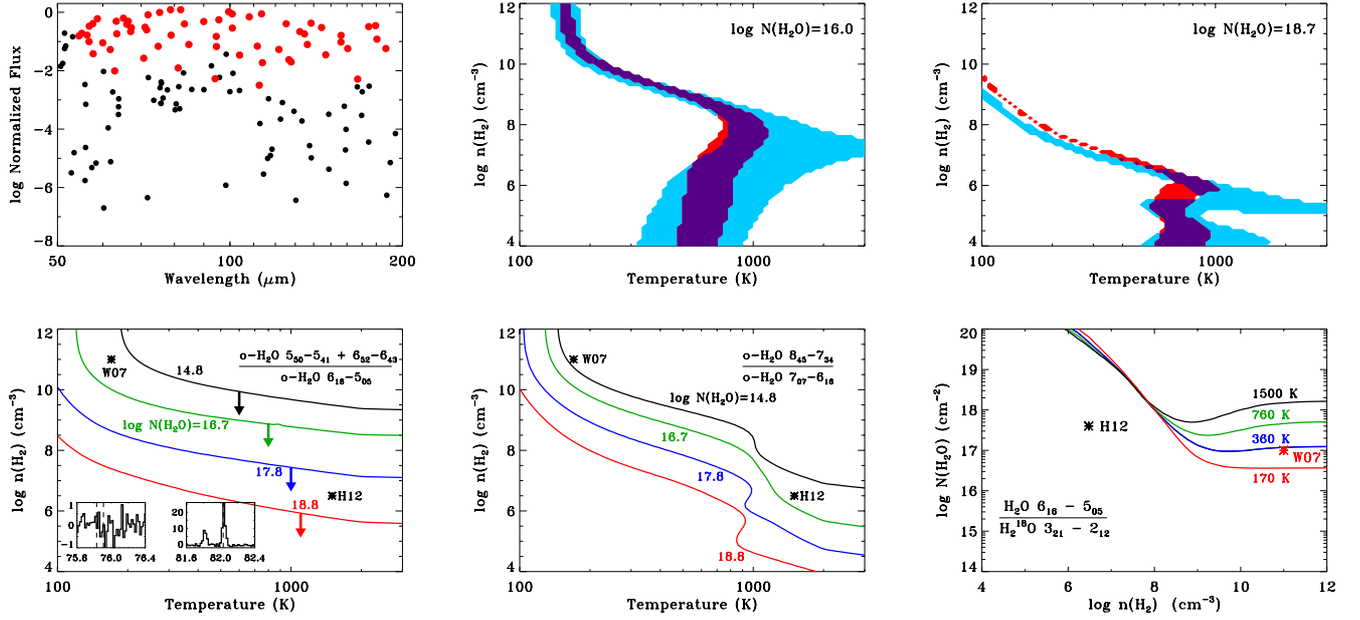


Fig. C.1. H₂O fluxes and acceptable parameter space of temperature and H₂ density. The *upper left panel* shows that in the H12 model, all far-IR H₂O lines expected to be strong are detected (red dots), while those expected to be weak are undetected (black dots). Several lines at $\sim 100 \mu\text{m}$ were likely undetected because the S/N is degraded in that wavelength region. The *upper middle panel* shows the acceptable contours for flux ratios of $o\text{-H}_2\text{O } 8_{18-7_{07}}/7_{07-6_{16}}$ (blue), $o\text{-H}_2\text{O } 7_{07-6_{16}}/6_{16-5_{05}}$ (red), and both (purple), for two different column densities. The acceptable parameter space shows the 1σ error bars with a 20% relative flux calibration uncertainty. The *lower left panel* shows that the non-detection of the $o\text{-H}_2\text{O } 5_{50-5_{41}}$ and $6_{52-6_{43}}$ lines relative to $o\text{-H}_2\text{O } 6_{16-5_{05}}$ line places a limit on the optical depth of the slab. The *lower middle panel* shows where the flux ratio of $o\text{-H}_2\text{O } 8_{45-7_{34}}$ $35.669 \mu\text{m}$ to $o\text{-H}_2\text{O } 7_{07-6_{16}}$ $71.947 \mu\text{m}$ is equal to the observed value of 0.07. The *lower right panel* shows the parameter space (high density and high column density) ruled out by the non-detection of the $o\text{-H}_2^{18}\text{O } 3_{21-2_{12}}$ line, for four different temperatures.

on lines at short wavelengths, so the emission from the heavily embedded region is mostly extinguished. An $A_V = 100$ mag to the blue outflow lobe would increase the relative luminosity of lines at $25 \mu\text{m}$ compared with those at $100 \mu\text{m}$ by a factor of four,

thereby increasing the warm H₂O excitation temperature from 220 K to 240 K. An $A_V > 200$ mag to the outflow is ruled out by the significant increase in scatter in the excitation diagram.

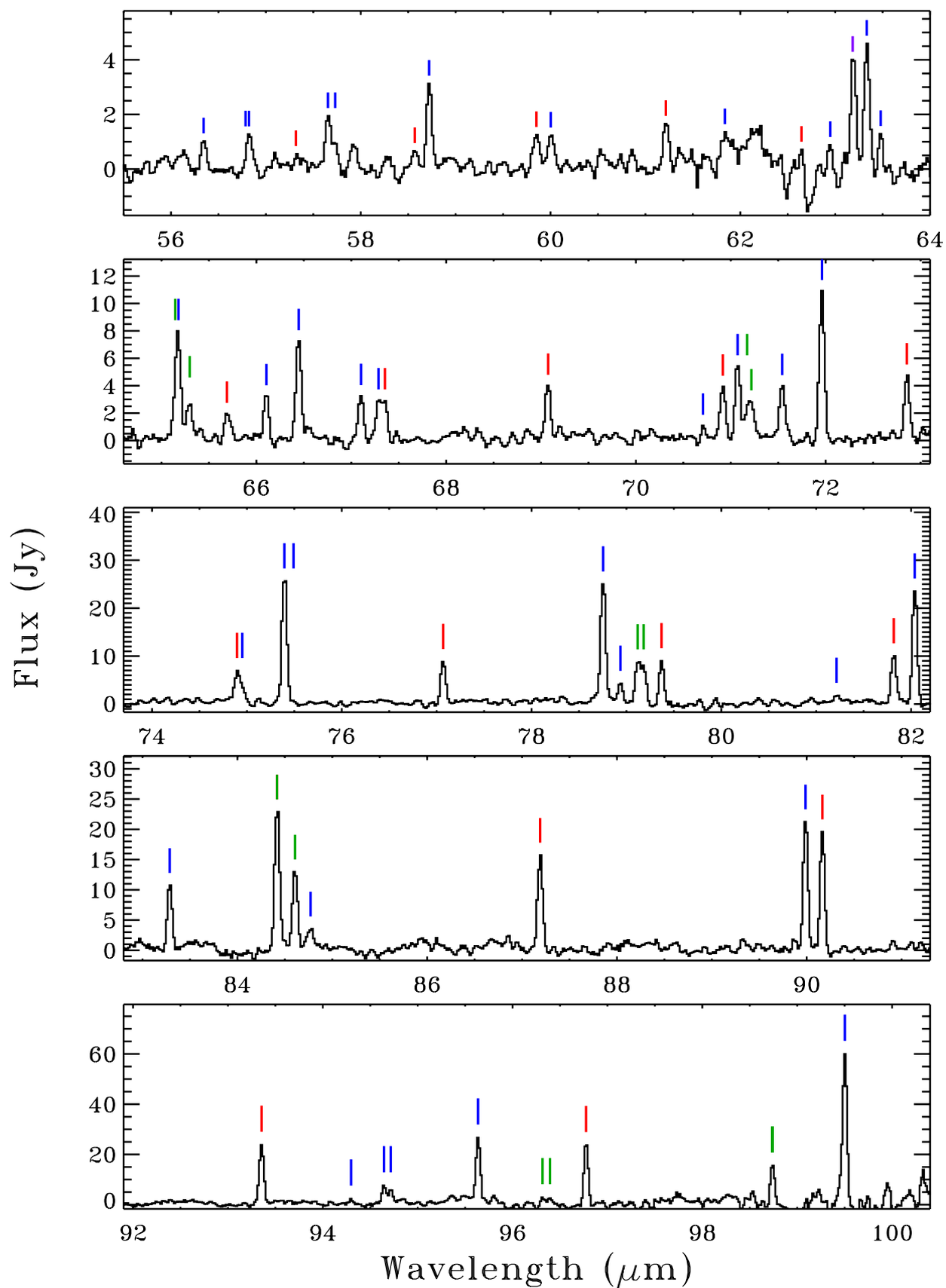


Fig. D.1. The continuum-subtracted PACS spectrum of NGC 1333 IRAS 4B from 55–100 μm . The marks identify lines of H_2O (blue), CO (red), OH (green), and [O I] (purple).

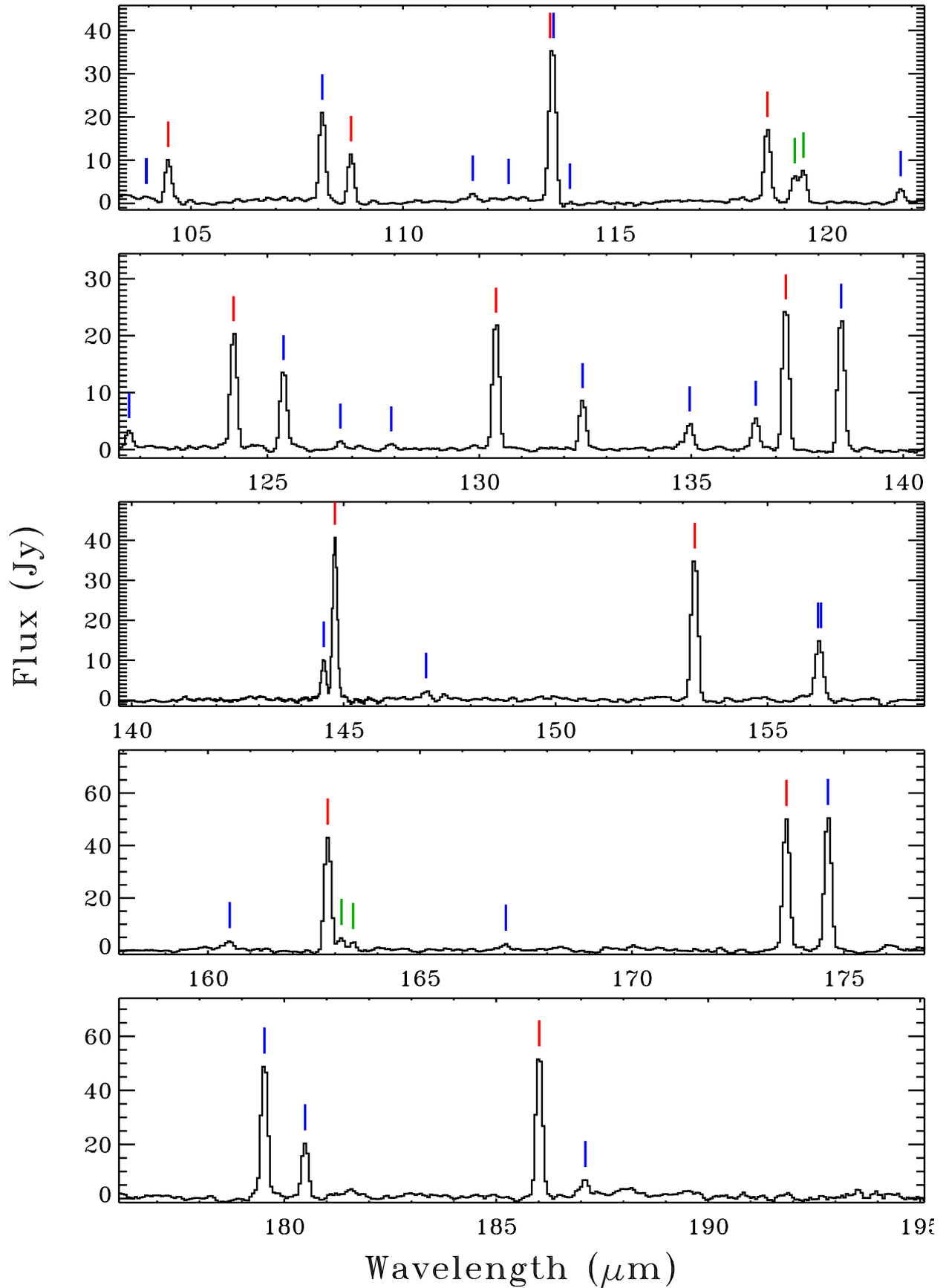


Fig. D.2. The continuum-subtracted PACS spectrum of NGC 1333 IRAS 4B from 103–195 μm . The marks identify lines of H₂O (blue), CO (red), and OH (green).

Table D.1. Detected lines in *Herschel*/PACS spectrum of IRAS 4B.

Species	Line	E' (K)	$\log A_{ul}$ (s^{-1})	λ_{vac} (μm)	λ_{obs} (μm)	Flux ^a	Error ^a
CO	49–48	6724	–2.17	53.897	53.893	0.66 ^c	0.11
o-H ₂ O	5 ₃₂ –5 ₀₅	732	–1.42	54.506	54.495	0.61	0.12
CO	48–47	6457	–2.18	54.986	54.965	0.39	0.07
o-H ₂ O	8 ₂₇ –7 ₁₆	1274	0.28	55.131	55.146	0.53 ^b	0.13
p-H ₂ O	4 ₃₁ –3 ₂₂	552	0.16	56.325	56.344	0.59	0.08
p-H ₂ O	9 ₁₉ –8 ₀₈	1324	0.40	56.771	56.786	0.21 ^b	0.07
o-H ₂ O	9 ₀₉ –8 ₁₈	1323	0.39	56.816	56.822	0.74	0.09
CO	46–45	5939	–2.22	57.308	57.317	0.27 ^b	0.09
p-H ₂ O	4 ₂₂ –3 ₁₃	454	–0.42	57.636	57.655	0.97	0.11
p-H ₂ O	8 ₁₇ –7 ₂₆	1270	0.21	57.709	57.730	0.43	0.08
CO	45–44	5688	–2.23	58.547	58.570	0.39	0.09
o-H ₂ O	4 ₃₂ –3 ₂₁	550	0.14	58.699	58.721	1.60	0.09
CO	44–43	5442	–2.25	59.843	59.851	0.70	0.08
p-H ₂ O	7 ₂₆ –6 ₁₅	1021	0.13	59.987	60.003	0.66	0.07
CO	43–42	4202	–2.27	61.201	61.215	0.79	0.10
p-H ₂ O	4 ₃₁ –4 ₀₄	552	–1.61	61.808	61.838	0.40	0.18
CO	42–41	4967	–2.29	62.624	62.644	0.80 ^b	0.23
o-H ₂ O	9 ₁₈ –9 ₀₉	1552	–0.38	62.928	62.949	0.56	0.20
[O I]	³ P ₁ – ³ P ₂	228	–4.05	63.185	63.193	1.83	0.08
o-H ₂ O	8 ₁₈ –7 ₀₇	1070	0.24	63.323	63.333	1.92	0.14
p-H ₂ O	8 ₀₈ –7 ₁₇	1070	0.24	63.457	63.480	0.68	0.13
CO	41–40	4737	–2.31	64.117	64.127	1.08	0.13
OH	² Π _{3/2} $J = 9/2^- - 7/2^+$	512	0.11	65.131	65.145	0.99	0.12
o-H ₂ O	6 ₂₅ –5 ₁₄	795	–0.03	65.166	65.180	2.94	0.13
OH	² Π _{3/2} $J = 9/2^+ - 7/2^-$	510	0.10	65.278	65.297	1.05	0.12
CO	40–39	4513	–2.34	65.686	65.689	0.76	0.10
o-H ₂ O	7 ₁₆ –6 ₂₅	1013	–0.02	66.092	66.105	1.38	0.10
o-H ₂ O	3 ₃₀ –2 ₂₁	410	0.09	66.437	66.446	3.07	0.13
p-H ₂ O	3 ₃₁ –2 ₂₀	410	0.09	67.089	67.102	1.44	0.09
o-H ₂ O	3 ₃₀ –3 ₀₃	410	–2.07	67.268	67.287	1.24	0.10
CO	39–38	4294	–2.36	67.336	67.355	1.18	0.10
CO	38–37	4080	–2.39	69.074	69.075	1.57	0.12
o-H ₂ O	8 ₂₇ –8 ₁₈	1274	–0.48	70.702	70.707	0.45 ^b	0.12
CO	37–36	3872	–2.41	70.907	70.916	1.29	0.10
p-H ₂ O	5 ₂₄ –4 ₁₃	598	–0.18	71.067	71.072	1.98	0.09
OH	² Π _{1/2} $J = 7/2^- - 5/2^+$	617	0.01	71.170	71.172	0.60	0.07
OH	² Π _{1/2} $J = 7/2^+ - 5/2^-$	617	0.01	71.215	71.217	0.75	0.07
p-H ₂ O	7 ₁₇ –6 ₀₆	843	0.07	71.539	71.542	1.44	0.10
o-H ₂ O	7 ₀₇ –6 ₁₆	843	0.06	71.946	71.961	3.79	0.12
CO	36–35	3669	–2.44	72.843	72.855	1.53	0.11
CO	35–34	3471	–2.47	74.890	74.896	2.01	0.11
o-H ₂ O	7 ₂₅ –6 ₃₄	1125	–0.59	74.944	74.951	0.89	0.11
o-H ₂ O	3 ₂₁ –2 ₁₂	305	–0.48	75.380	75.395	9.35	0.25
o-H ₂ O	8 ₅₄ –8 ₄₅	1805	–0.24	75.495	75.491	0.33	0.10
CO	34–33	3279	–2.50	77.059	77.068	2.76	0.12
o-H ₂ O	4 ₂₃ –3 ₁₂	432	–0.32	78.742	78.755	7.57	0.23
p-H ₂ O	6 ₁₅ –5 ₂₄	781	–0.34	78.928	78.936	1.09	0.11
OH	² Π _{1/2} – ² Π _{3/2} $J = 1/2^- - 3/2^+$	181	–1.44	79.115	79.118	2.53	0.12
OH	² Π _{1/2} – ² Π _{3/2} $J = 1/2^+ - 3/2^-$	181	–1.44	79.178	79.182	2.16	0.12
CO	33–32	3092	–2.53	79.360	79.370	2.66	0.12
p-H ₂ O	7 ₂₆ –7 ₁₇	1021	–0.61	81.215	81.215	0.38	0.07
CO	32–31	2911	–2.56	81.806	81.817	2.64	0.09
o-H ₂ O	6 ₁₆ –5 ₀₅	643	–0.13	82.031	82.038	6.67	0.16
p-H ₂ O	6 ₀₆ –5 ₁₅	642	–0.15	83.283	83.287	2.75	0.11
OH	² Π _{3/2} $J = 7/2^+ - 5/2^-$	291	–0.28	84.420	84.418	5.90	0.22
CO	31–30	2735	–2.60	84.410	blend with OH ² Π _{3/2} $J = 7/2^+ - 5/2^-$		
OH	² Π _{3/2} $J = 7/2^- - 5/2^+$	290	–0.28	84.596	84.607	3.06	0.11
o-H ₂ O	7 ₁₆ –7 ₀₇	1013	–0.67	84.766	84.771	0.76	0.09

Notes. ^(a) 10^{-20} W cm⁻², extracted from two spaxels as discussed in Sect. 2. Listed errors are 1σ and do not include calibration uncertainty. ^(b) Flux measured from outflow spaxel only. ^(c) Flux measured from Nyquist-sampled map.

Table D.1. continued.

Species	Line	E' (K)	$\log A_{ul}$ (s^{-1})	λ_{vac} (μm)	λ_{obs} (μm)	Flux ^a	Error ^d
CO	30–29	2565	–2.63	87.190	87.190	3.38	0.13
p-H ₂ O	3 ₂₂ –2 ₁₁	296	–0.45	89.988	89.988	4.56	0.10
CO	29–28	2400	–2.67	90.163	90.165	4.12	0.10
CO	28–27	2240	–2.71	93.349	93.355	4.36	0.12
p-H ₂ O	5 ₄₂ –5 ₃₃	878	–0.62	94.209	94.297	0.32	0.10
o-H ₂ O	6 ₂₅ –6 ₁₆	795	–0.76	94.643	94.645	1.38	0.11
o-H ₂ O	4 ₄₁ –4 ₃₂	702	–0.82	94.704	94.716	1.00	0.10
p-H ₂ O	5 ₁₅ –4 ₀₄	469	–0.35	95.626	95.637	4.44	0.12
OH	${}^2\Pi_{1/2} - {}^2\Pi_{3/2} J = 3/2^+ - 5/2^-$	270	–2.03	96.271	96.316	0.57	0.11
OH	${}^2\Pi_{1/2} - {}^2\Pi_{3/2} J = 3/2^- - 5/2^+$	269	–2.03	96.362	96.394	0.44	0.12
CO	27–26	2086	–2.75	96.773	96.775	4.28	0.13
OH	${}^2\Pi_{1/2} J = 5/2^- - 3/2^+$	415	–0.45	98.736	98.728 ^c	2.46	0.43
OH	${}^2\Pi_{1/2} J = 5/2^+ - 3/2^-$	415	–0.45	98.764	blend with OH ${}^2\Pi_{1/2} J = 5/2^- - 3/2^+$		
o-H ₂ O	5 ₀₅ –4 ₁₄	468	–0.41	99.492	99.496 ^c	9.85	0.26
CO	26–25	1937	–2.80	100.460	100.464 ^c	3.86	0.20
o-H ₂ O	5 ₁₄ –4 ₂₃	574	–0.81	100.912	100.910 ^c	4.56	0.22
p-H ₂ O	2 ₂₀ –1 ₁₁	195	–0.58	100.982	100.981 ^c	5.42	0.17
p-H ₂ O	6 ₁₅ –6 ₀₆	781	–0.87	103.916	103.944	3.50	0.80
CO	25–24	1794	–2.84	104.445	104.464	4.22	0.18
o-H ₂ O	2 ₂₁ –1 ₁₀	194	–0.59	108.072	108.097	9.47	0.19
CO	24–23	1656	–2.89	108.763	108.774	4.86	0.15
p-H ₂ O	5 ₂₄ –5 ₁₅	598	–0.92	111.627	111.646	0.71	0.10
o-H ₂ O	7 ₄₃ –7 ₃₄	1339	–0.67	112.510	112.492	0.21 ^b	0.09
CO	23–22	1524	–2.94	113.458	113.470	4.88	0.15
o-H ₂ O	4 ₁₄ –3 ₀₃	323	–0.61	113.536	113.549	11.70	0.15
p-H ₂ O	5 ₃₃ –5 ₂₄	725	–0.78	113.947	113.937	0.31	0.15
CO	22–21	1397	–3.00	118.581	118.595	6.46	0.11
OH	${}^2\Pi_{3/2} J = 5/2^- - 3/2^+$	120	–0.86	119.233	119.241	2.23	0.12
OH	${}^2\Pi_{3/2} J = 5/2^+ - 3/2^-$	120	–0.86	119.441	119.447	2.83	0.11
o-H ₂ O	4 ₃₂ –4 ₂₃	550	–0.91	121.721	121.740	1.17	0.08
CO	21–20	1276	–3.05	124.193	124.204	7.02	0.10
p-H ₂ O	4 ₀₄ –3 ₁₃	319	–0.76	125.353	125.383	4.73	0.10
p-H ₂ O	3 ₃₁ –3 ₂₂	410	–1.11	126.713	126.724	0.46	0.08
o-H ₂ O	7 ₂₅ –7 ₁₆	1125	–0.87	127.883	127.922	0.30	0.05
CO	20–19	1160	–3.11	130.369	130.393	7.38	0.05
o-H ₂ O	4 ₂₃ –4 ₁₄	432	–1.09	132.407	132.437	2.73	0.05
o-H ₂ O	5 ₁₄ –5 ₀₅	574	–1.12	134.934	134.964	1.27	0.07
o-H ₂ O	3 ₃₀ –3 ₂₁	410	–1.18	136.495	136.520	1.63	0.06
CO	19–18	1050	–3.18	137.196	137.230	7.66	0.07
p-H ₂ O	3 ₁₃ –2 ₀₂	204	–0.90	138.527	138.536	6.96	0.07
p-H ₂ O	4 ₁₃ –3 ₂₂	396	–1.48	144.517	144.533	2.04	0.07
CO	18–17	945	–3.24	144.784	144.794	8.13	0.09
p-H ₂ O	4 ₃₁ –4 ₂₂	552	–1.09	146.923	146.945	0.49	0.09
CO	17–16	846	–3.32	153.267	153.282	8.87	0.06
p-H ₂ O	3 ₂₂ –3 ₁₃	296	–1.28	156.193	156.190	2.16	0.06
o-H ₂ O	5 ₂₃ –4 ₃₂	642	–1.95	156.264	156.261	1.55	0.06
o-H ₂ O	5 ₃₂ –5 ₂₃	732	–1.09	160.509	160.513	0.66	0.07
CO	16–15	752	–3.39	162.812	162.825	9.13	0.20
OH	${}^2\Pi_{3/2} J = 3/2^+ - 1/2^-$	270	–1.19	163.123	163.149	1.01	0.08
OH	${}^2\Pi_{3/2} J = 3/2^- - 1/2^+$	269	–1.19	163.395	163.423	0.88	0.04
p-H ₂ O	6 ₂₄ –6 ₁₅	867	–1.12	167.034	167.027	0.44	0.06
CO	15–14	663	–3.47	173.631	173.643	9.45	0.11
o-H ₂ O	3 ₀₃ –2 ₁₂	196	–1.30	174.624	174.625	9.53	0.11
o-H ₂ O	2 ₁₂ –1 ₀₁	114	–1.25	179.525	179.532	8.86	0.10
o-H ₂ O	2 ₂₁ –2 ₁₂	194	–1.51	180.487	180.491	3.56	0.09
CO	14–13	580	–3.56	185.998	186.014	8.64	0.11
p-H ₂ O	4 ₁₃ –4 ₀₄	396	–1.43	187.109	187.102	0.93	0.10
CO	13–12	503	–3.66	200.271	200.262 ^c	6.33	0.10

Notes. ^(a) 10^{-20} W cm⁻², extracted from two spaxels as discussed in Sect. 2. Listed errors are 1σ and do not include calibration uncertainty. ^(b) Flux measured from outflow spaxel only. ^(c) Measured from first or second order emission at $>190 \mu m$.