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HIFISTARS Herschel/HIFI observations of VY Canis Majoris*,**

Molecular-line inventory of the envelope around the largest known star

J. Alcolea¹, V. Bujarrabal², P. Planesas¹, D. Teyssier³, J. Cernicharo⁴, E. De Beck^{5,6}, L. Decin^{5,7}, C. Dominik^{7,8}, K. Justtanont⁹, A. de Koter^{5,7}, A. P. Marston³, G. Melnick¹⁰, K. M. Menten⁶, D. A. Neufeld¹¹, H. Olofsson⁹, M. Schmidt¹², F. L. Schöier^{9,***}, R. Szczerba¹², and L. B. F. M. Waters^{7,13}

- Observatorio Astronómico Nacional (IGN), Alfonso XII, 3 y 5, 28014 Madrid, Spain e-mail: j.alcolea@oan.es
- ² Observatorio Astronómico Nacional (IGN), PO Box 112, 28803 Alcalá de Henares, Spain
- ³ European Space Astronomy Centre, Urb. Villafranca del Castillo, PO Box 50727, 28080 Madrid, Spain
- ⁴ CAB, INTA-CSIC, Ctra. de Torrejón a Ajalvir, km 4, 28850 Torrejón de Ardoz, Spain
- ⁵ Instituut voor Sterrenkunde, Katholieke Universiteit Leuven, Celestijnenlaan 200D BUS 2401, 3001 Leuven, Belgium
- Max-Planck-Institut für Radioastronomie, Auf dem Hügel 69, 53121 Bonn, Germany
- ⁷ Sterrenkundig Instituut Anton Pannekoek, University of Amsterdam, Science Park 904, 1098 Amsterdam, The Netherlands
- ⁸ Department of Astrophysics/IMAPP, Radboud University Nijmegen, Nijmegen, The Netherlands
- Onsala Space Observatory, Dept. of Earth and Space Sciences, Chalmers University of Technology, 43992 Onsala, Sweden
- ¹⁰ Harvard-Smithsonian Center for Astrophysics, Cambridge, MA 02138, USA
- ¹¹ The Johns Hopkins University, 3400 North Charles St, Baltimore, MD 21218, USA
- ¹² N. Copernicus Astronomical Center, Rabiańska 8, 87-100 Toruń, Poland
- Netherlands Institute for Space Research, Sorbonnelaan 2, 3584 CA Utrecht, The Netherlands

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ABSTRACT

Aims. The study of the molecular gas in the circumstellar envelopes of evolved stars is normally undertaken by observing lines of CO (and other species) in the millimetre-wave domain. In general, the excitation requirements of the observed lines are low at these wavelengths, and therefore these observations predominantly probe the cold outer envelope while studying the warm inner regions of the envelopes normally requires sub-millimetre (sub-mm) and far-infrared (FIR) observational data.

Methods. To gain insight into the physical conditions and kinematics of the warm (100–1000 K) gas around the red hyper-giant VY CMa, we performed sensitive high spectral resolution observations of molecular lines in the sub-mm/FIR using the HIFI instrument of the Herschel Space Observatory. We observed CO, H₂O, and other molecular species, sampling excitation energies from a few tens to a few thousand K. These observations are part of the Herschel guaranteed time key program HIFISTARS.

few tens to a few thousand K. These observations are part of the *Herschel* guaranteed time key program HIFISTARS. *Results.* We detected the J=6-5, J=10-9, and J=16-15 lines of ^{12}CO and ^{13}CO at ~ 100 , 300, and 750 K above the ground state (and the ^{13}CO J=9-8 line). These lines are crucial for improving the modelling of the internal layers of the envelope around VY CMa. We also detected 27 lines of H_2O and its isotopomers, and 96 lines of species such as NH_3 , SiO, SO, SO₂ HCN, OH and others, some of them originating from vibrationally excited levels. Three lines were not unambiguously assigned.

Conclusions. Our observations confirm that VY CMa's envelope must consist of two or more detached components. The molecular excitation in the outer layers is significantly lower than in the inner ones, resulting in strong self-absorbed profiles in molecular lines that are optically thick in this outer envelope, for instance, low-lying lines of H_2O . Except for the most abundant species, CO and H_2O , most of the molecular emission detected at these sub-mm/FIR wavelengths arise from the central parts of the envelope. The spectrum of VY CMa is very prominent in vibrationally excited lines, which are caused by the strong IR pumping present in the central regions. Compared with envelopes of other massive evolved stars, VY CMa's emission is particularly strong in these vibrationally excited lines, as well as in the emission from less abundant species such as $H^{13}CN$, SO, and NH_3 .

Key words. stars: AGB and post-AGB – stars: mass-loss – stars: individual: VY Canis Majoris – circumstellar matter

1. Introduction

The very luminous red evolved star VY CMa (M2.5 – M5e Ia; Wallerstein 1958; Humphreys 1974), also known as IRC –30087

and CRL 1111, is one of the most extreme stars in our galaxy. Thanks to trigonometric parallax measurements of $\rm H_2O$ and $^{28}\rm SiO$ masers, the distance to VY CMa is well known, 1.10–1.25 kpc (see Choi et al. 2008; Zhang et al. 2012), resulting in a total luminosity of $3\times 10^5\,L_\odot$ (Smith et al. 2001). This value is close to the empirical limit for cool evolved stars (Humphreys & Davidson 1994), and therefore VY CMa has been classified as a red hyper-giant (RHG), see de Jager (1998). These accurate distance estimates, in combination with VLTI/AMBER measurements of the angular size of VY CMa, result in a value for the stellar radius that is the highest among well-characterized

^{*} *Herschel* is an ESA space observatory with science instruments provided by European-led Principal Investigator consortia and with important participation from NASA. HIFI is the *Herschel* Heterodyne Instrument for the Far Infrared.

^{**} Appendices A and B are available in electronic form at http://www.aanda.org

^{***} Deceased 14 January 2011.

stars in our galaxy: $1420 \pm 120 R_{\odot}$ (Wittkowski et al. 2012). Values for its initial mass are more uncertain, ranging from 15 to $50 M_{\odot}$ (see Wittkowski et al. 1998; Knapp et al. 1993; Smith et al. 2001; Choi et al. 2008; Wittkowski et al. 2012). VY CMa is also characterized by displaying a huge mass loss, with rate values in the range of 5×10^{-5} to $10^{-4} M_{\odot} \text{ yr}^{-1}$ (Decin et al. 2006), and as high as 2×10^{-3} at least during the past 300 yr (see Smith et al. 2009, and references therein); this last value is so high that it is thought that VY CMa might explode as a supernova from its present state in about 10^5 yr (see Smith et al. 2009).

Owing to its high mass-loss rate, the star is surrounded by a thick circumstellar envelope, which was detected as an optical reflection nebulosity almost a century ago (Perrine 1923), and which has been studied in great detail using the *Hubble* Space Telescope and ground-based adaptive optics (see Smith et al. 2001; Humphreys et al. 2007, and references therein). This nebulosity is relatively compact, $8'' \times 8''$ at most (even in the mid-IR), but displays much sub-structure. At short wavelengths, the nebula is highly asymmetric, with a much brighter lobe located south of the assumed position of the star. This suggests the existence of a bipolar structure oriented in the north-south direction, the south lobe pointing towards us (see Herbig 1970, for example). Due to the large amount of dust in the envelope, the star itself is hardly visible in the optical, and most of its luminosity is re-radiated in the mid-IR and at longer wavelengths. This and the high luminosity of the source makes VY CMa one of the brightest IR sources in the sky.

The envelope around VY CMa has a very rich molecular variety. In the centimetre- and millimetre-wave range, VY CMa has been known for many years to exhibit strong maser emission in the three classical circumstellar maser species: OH, H₂O, and ²⁸SiO. In particular, VYCMa is the strongest emitter of highly excited ${}^{28}SiO$ (v > 2) and H₂O ($v_2 \ge 1$) masers (see e.g. Cernicharo et al. 1993; Menten et al. 2006). At these wavelengths, however, the strength of the thermal molecular emission of VY CMa's envelope is not outstanding, probably due to its large distance and small extent, which led to little interest in the object. This situation has changed drastically in the past half decade, when the results of sensitive spectral surveys in millimetre, sub-millimetre (sub-mm), and far-IR (FIR) ranges were published, from which a wealth of molecular lines from about two dozen species have been identified. Since then, this source has become the main target of the search of new molecular species in O-rich circumstellar environments.

Using the ISO Short Wavelength Spectrometer, Neufeld et al. (1999) discovered that the FIR spectrum of VY CMa is rich in spectral features that are caused by water vapour. Subsequently, Polehampton et al. (2010) studied VY CMa at longer wavelengths with the ISO Long Wavelength Spectrometer, finding that it is also dominated by strong lines of H₂O, and concluding that this is the most abundant molecular species after H₂ in its circumstellar envelope. In addition to H₂O, CO, and OH, these authors also reported the identification of lines from less common species such as NH₃, and tentatively H₃O⁺. In the same year, Royer et al. (2010) published the Herschel PACS and SPIRE full spectral scan (55 to 672 μ m wavelength range). These more sensitive observations confirmed that the FIR and sub-mm spectrum of VY CMa is dominated by H₂O lines, which are responsible for nearly half of the 930 emission lines identified in this work. In addition to H₂O, lines of CO, CN, CS, SO, NH₃, OH, SiO, SiS, SO₂, H₂S, and H₃O⁺ (and of some of their isotopomers) are also detected in the spectrum, in spite of the relatively poor spectral resolution. From a preliminary analysis of the H₂O spectrum, Royer and collaborators concluded that the fractional abundance of this species (w.r.t. the total abundance of hydrogen) is very high, 3×10^{-4} . They also found a low ortho-topara abundance ratio of ~1.3:1, which would support the hypothesis that non-equilibrium chemical processes control the formation of H₂O in this envelope. Meanwhile, Tenenbaum et al. (2010b; but see also Ziurys et al. 2007 and 2009; Tenenbaum & Ziurys 2009 and 2010; Tenenbaum et al. 2010a) presented the results of a full-scan survey in the 215-285 GHz range, conducted with the Arizona Radio Obseratory's 10 m diameter Submillimeter Telescope¹ on Mt. Graham (ARO-10 m SMT). This survey yielded the detection of ten more new species in the envelope of VY CMa, namely HCN, HNC, HCO⁺, CN, NS, PN, PO, NaCl, AlO, and AlOH. These studies will soon be complemented with the results from the full spectral surveys that have been or are being performed with Herschel/HIFI and the IRAM 30 m telescope in all the available bands of these two instruments, and with the Submillimeter Array (SMA) in the 280–355 GHz (~0.9 mm) frequency range (see Kamiński et al. 2013). Altogether, these works have revealed, and will continue to reveal, the chemical richness and complexity that can be present in the circumstellar envelopes of cool, high mass-loss O-rich stars.

A proper understanding of the chemical characteristics of the circumstellar envelope of this unique source necessarily requires a prior good knowledge of the main physical conditions in the envelope. For the cool layers, this can be attained from the ground by means of low-J rotational lines of CO and other abundant species. However, to gain insight into the deep warm layers where most of the molecular species are formed, we need to observe high-J lines that in general are not or not easily accessible for ground-based telescopes. Moreover, for O-rich environments, the expected high H₂O abundances make this species the major coolant for the gas phase, and therefore knowing the distribution and excitation of H₂O is crucial for determining the temperature of the molecular gas in general. Yet, it is important to observe these molecular lines with very high spectral resolution to adequately probe the velocity field in the envelope. All these observational needs are fully met by the HIFI instrument on-board the *Herschel* Space Observatory. Here we present new Herschel/HIFI observations of VY CMa. Observational and data reduction procedures are detailed in Sect. 2. In Sect. 3 and the appendices, we discuss the main observational results. The conclusions are presented in Sect. 4.

2. Observations and data reduction

The observations we present in this paper were obtained with the Herschel Space Observatory (Pilbratt et al. 2010), using the Heterodyne Instrument for the Far Infrared (HIFI, de Graauw et al. 2010). This data set is part of the results obtained by the guaranteed time key program HIFISTARS, which is devoted to the study of the warm gas and water vapour contents of the molecular envelopes around evolved stars. The main target lines of the HIFISTARS project were the J = 6-5, 10-9, and 16-15 transitions of ¹²CO and ¹³CO, and several lines of ortho- and para-H₂O sampling a wide range of line-strengths and excitation energies, including vibrationally excited states. In addition, some other lines were observed simultaneously, thanks to the large instantaneous bandwidth coverage of the HIFI receivers. We observed 16 different frequency settings for VY CMa. In the naming adopted in the project, these spectral setups are, in order of increasing local oscillator (LO) frequency,

¹ Formerly known as the *Heinrich Hertz* telescope.

Table 1. Telescope characteristics and main observational parameters.

Herschel	Obs. date	Durat.	HIFIST	ARS setting	Sky frequen	icy coverage	$T_{ m sys}^{ m DSB}$	HIFI	HPBV	$W T_{\rm mb}$	/T _a Cal.§
OBSID	yr:mo:day	(s)	Name	LO (GHz)	LSB (GHz)			band	(")		uncer.
1342192528	2010:03:21	1575	14	564.56	556.55-560.69	568.54-572.68	93	1B	37.5	1.3	15%
1342192529	2010:03:21	1575	13	614.86	606.85-610.99	618.84-622.98	91	1B	34.5	1.3	15%
1342192533	2010:03:21	1623	12	653.55	645.55-649.69	657.54-661.68	131	2A	32.0	1.3	15%
1342192534	2010:03:21	617	17	686.42	678.42-682.56	690.41-694.55	142	2A	30.8	1.3	15%
1342194680†	2010:04:13	1618	11	758.89	750.90-755.04	762.89-767.03	196	2B	27.8	1.3	15%
1342195039^{\dagger}	2010:04:18	1505	10	975.23	967.27-971.41	979.26-983.40	352	4A	21.7	1.3	20%
1342195040†	2010:04:18	1505	09	995.63	987.67-991.81	999.66-1003.80	364	4A	21.3	1.3	20%
1342194786	2010:04:17	1487	08	1102.92	1094.98-1099.11	1106.97-1111.10	403	4B	19.2	1.3	20%
1342194787	2010:04:17	1487	07	1106.90	1098.95-1103.09	1110.94-1115.08	416	4B	19.1	1.3	20%
1342192610	2010:03:22	1618	06	1157.67	1149.72-1153.85	1161.71-1165.84	900	5A	18.3	1.5	20%
1342192611	2010:03:22	1538	05	1200.90	1192.95-1197.08	1204.94-1209.07	1015	5A	17.6	1.5	20%
1342195105‡	2010:04:19	3547	04	1713.85	1709.11-1711.68	1716.37-1718.97	1238	7A	12.4	1.4	30%
1342195106 [‡]	2010:04:19	3851	03	1757.68	1752.95-1755.52	1760.20-1762.77	1580	7A	12.0	1.4	30%
1342230403‡	2011:10:09	3103	19^{\flat}	1766.89	1762.23-1764.78	1769.48-1772.04	1248	7A	12.0	1.4	30%
1342194782‡	2010:04:17	2992	16^{\sharp}	1838.31	1833.59-1836.16	1840.84-1843.41	1300	7B	11.6	1.4	30%
1342196570‡	2010:05:15	3207	16 [#]	1838.47	1833.73-1836.30	1840.99-1843.55	1415	7B	11.6	1.4	30%
1342194781‡	2010:04:17	1535	01	1864.82	1860.10-1862.67	1867.36–1869.92	1330	7B	11.4	1.4	30%

Notes. ^(§) Following Teyssier et al. (2012), we based our absolute-calibration accuracy estimates on the error budget reported by Roelfsema et al. (2012), plus an additional contribution arising from the ripples in the baseline for bands 7A and 7B. ^(†) Instabilities in the baseline that simultaneously affect both receivers were removed by discarding several sub-scans. ^(‡) Ripples in the baseline, which mainly affected the V-receiver, were removed by discarding several sub-scans. ^(b) Setting 19 was added later in the program. ^(‡) Setting 16 was observed twice using a slightly different value for the systemic velocity.

settings 14, 13, 12, 17, 11, 10, 09, 08, 07, 06, 05, 04, 03, 19, 16, and 01; their main observational parameters are listed in Table 1.

The observations were all performed using the two orthogonal linearly polarized receivers available for each band, named H and V (horizontal and vertical) for their mutually perpendicular orientations. These receivers work in double side-band mode (DSB), which doubles the instantaneous sky frequency coverage of the HIFI instrument: 4 plus 4 GHz for the superconductorinsulator-superconductor (SIS) receivers of bands 1 to 5, and 2.6 plus 2.6 GHz for the hot-electron bolometer (HEB) receivers of band 7. The observations were all performed in the dual-beam switching (DBS) mode. In this mode, the HIFI internal steering mirror chops between the source position and two (expected to be) emission-free positions located 3' at either side of the science target. The telescope alternately locates the source in either of the two chopped beams (a and b), providing a double-difference calibration scheme, $(ON_a-OFF_b)-(OFF_a-ON_b)$, which allows a more efficient cancellation of the residual baseline and optical standing waves in the spectra (see Roelfsema et al. 2012, for additional details). In our program, the DBS procedure worked well except for band 7, where strong ripples (generated by electrical standing waves) are often found in the averaged spectra, especially in the case of the V-receiver. The HIFI data shown here were taken using the wide-band spectrometer (WBS), an acousto-optical spectrometer that provides full coverage of the whole instantaneous IF band in the two available orthogonal receivers, with an effective spectral resolution slightly varying across the band, with a mean value of 1.1 MHz. This spectrometer is made of units with bandwidths slightly wider than 1 GHz, and therefore four/three units per SIS/HEB receiver are needed to cover the full band.

The data were retrieved from the *Herschel* Science Archive and were reprocessed using a modified version of the standard

HIFI pipeline using HIPE². This customized level-2 pipeline provides as final result individual double-difference elementary integrations without performing the final time-averaging, but stitching the three or four used WBS sub-bands together³. Later on, these spectra were exported to CLASS⁴ using the hiClass tool within HIPE for a more detailed inspection and *flagging* of data with noticeable ripple residuals. Time-averaging was also performed in CLASS, as well as baseline removal. Finally, spectra from the H and V receivers were compared and averaged together, as the differences found between the two receivers were always smaller than the calibration uncertainties.

In general, the data presented no problems and did not need a lot of *flagging*, except for the settings observed using band 7 (see Table 1). All these settings presented severe residual ripples whose intensity varied from sub-scan to sub-scan. A semi-automated procedure was designed in CLASS to detect and remove the sub-scans in which the ripples were more severe. The application of this procedure normally results in the rejection of 30% to 50% of the non-averaged spectra, which produces a final spectrum slightly noisier, but with a more reliable baseline, since the standing waves are largely suppressed. Some instrumental features affecting the baseline were also detected in

² HIPE is a joint development by the *Herschel* Science Ground Segment Consortium, consisting of ESA, the NASA *Herschel* Science Center, and the HIFI, PACS, and SPIRE consortia. Visit http://herschel.esac.esa.int/HIPE_download.shtml for additional information.

³ The standard HIPE pipeline does perform this time average, but does not perform sub-band stitching by default, providing just a single spectrum per receiver and WBS unit.

⁴ CLASS is part of the GILDAS software package, developed and maintained by IRAM, LAOG/Univ. de Grenoble, LAB/Obs. de Bordeaux, and LERMA/Obs. de Paris. For more details, see http://www.iram.fr/IRAMFR/GILDAS

settings 09, 10, and 11, which are also largely suppressed by removing the most affected sub-scans.

The original data were oversampled to a uniform channel spacing of 0.5 MHz, but we smoothed all spectra down to a velocity resolution of about $1\,\mathrm{km\,s^{-1}}$. The data were re-calibrated into (Rayleigh-Jeans equivalent) main-beam temperatures (T_{mb}) adopting the latest available values for the telescope and main beam efficiencies (Roelfsema et al. 2012). In all cases we assumed a side-band gain ratio of one. A summary of these telescope characteristics, including the total observational uncertainty budget, is given in Table 1.

3. HIFI results

The final results of the HIFISTARS observations of VY CMa are presented in Figs. B.1 and B.2, where we show the full bandwidth observed in each of the settings using the WBS. The original spectral resolution was degraded down to about 1 km s⁻¹ by averaging to the nearest integer number of channels, and a baseline was removed. Because the receivers work in DSB mode, each spectral feature has two possible values for its rest frequency, one if the emission enters the receiver via the LSB, and another if it does via the USB. In Figs. B.1 and B.2 we therefore plotted both frequency axes. In all cases, frequency scales and line identifications are made assuming a systemic velocity for the source of 22 km s⁻¹ w.r.t. the LSR frame. Lines from the LSB and USB, marked in red and blue, are labelled on the lower/upper x-axis of each panel. For the three lines that remain unidentified or unassigned, marked in green, we give labels for the two (LSB/USB) possible rest frequencies.

We detected 34 lines of CO and H₂O (including their isotopomers), 96 additional lines of 14 other species, and 3 unidentified lines. Therefore, in the 114 GHz covered by the observations (in both upper and lower side-bands), we detected a total 133 spectral lines, resulting in a mean line density of 1.17 lines per GHz. In Tables 2, 3, A.1, and A.2, we give the main observational parameters of the detected lines. We list the species and transition names, the energy of the upper state above the ground level of the species, the rest frequency found in the spectral line catalogues, the name of the HIFISTARS setting and side-band in which the line was observed, the root mean square (rms) value for the given spectral resolution, the peak intensity, the integrated area, and the velocity range over which the line's emission was integrated (in the cases where this applies). For unblended well-detected lines, peak and area were computed directly from the observational data. For blended lines or spectral features for which the signal-to-noise ratio (S/N) is modest, we fitted Gaussian profiles to the spectra, and we give the peak and area resulting from these fits. In the case of blended lines, we simultaneously fitted as many Gaussian profiles – usually two, but sometimes up to four – as evident lines appeared in the blended profile. The results from these fits, as well as the resulting total composite profile, can be seen in the figures of the individual lines (see next paragraph). Parameters for unidentified features are listed in Table A.3.

Individual plots for the different-well detected lines are presented in Figs. 3–10 and A.2–A.5. In these individual line plots, the horizontal scale is given in LSR velocities assuming the rest frequency quoted in the tables. In many cases, some other lines, close in frequency or from the other side-band, can be seen in the plots. To avoid confusion, the spectral line we refer to is plotted in the centre of the panel, and the grey area below the histogram shows the velocity range considered when computing its integrated intensity as given in Tables 2, 3, A.1, and A.2. In the case

of line blends, we also plot the results of the multi-line Gaussian fit we used to separate the emission from the different lines, as well as the resulting composite spectrum. Here the grey area also shows the contribution to the profile of the spectral line shown in the panel. Species and line names, rest frequency, setting and side-band, as well as the HPBW of the telescope at the corresponding frequency are also given in each panel. To facilitate the comparison between the different line profiles, vertical red dashed lines mark the assumed systemic velocity and the central velocities of the two high expansion velocity components (see next paragraphs).

We only made observations pointed at the central position of the star: J2000 RA $07^{h}22^{m}58.33$, Dec $-25^{\circ}46'03''.2$ (see Zhang et al. 2012). Therefore, for emitting sizes larger than or comparable with the beam size of the telescope, some flux is missed. In the optical and IR, the nebula around VYCMa is not larger than 8" in diameter (see Sect. 1). From the interferometric maps of lines of CO and other molecules (Muller et al. 2007; Verheyen 2011; Fu et al. 2012), it is found that the molecular emission is also smaller than 8". In these cases, however, it can be argued that any more extended component could be resolved out. We investigated the existence of such an extended component (10" in diameter or larger) by comparing single-dish data from radio telescopes of different beam sizes. We compared the $^{12}CO\ J = 2-1$ spectrum from VY CMa observed at the IRAM 30 m (unpublished observations by some of us) and JCMT 15 m (Kemper et al. 2003) telescopes, with HPBWs of 12" and 19".7, respectively. When comparing these two spectra, we found an IRAM 30 m-to-JCMT 15 m flux ratio of 2.1, except for the velocity range 12–22 km s⁻¹, where we obtained values between 1.8 and 2.1. If we assume that these ratios are only due to the different beam dilution factors, we derive a FWHM size for the emitting region of $\sim 8''$ for the 2.1 ratio, while the 1.8 value can be explained if the size of the source is a large as $\sim 12''.5$. We note that this analysis is affected by the uncertainties in the absolute calibration of both telescopes, but these sizes are compatible with existing images of the nebula. Still, the existence of a structure larger than $\sim 8''$ at moderate blue-shifted velocities (between $12-22\,\mathrm{km\,s^{-1}}$) agrees with the lost flux at these velocities noted by Muller et al. (2007) in their interferometric observations. All these estimates agree with other results obtained from single-dish studies from groundbased telescopes such as those by Ziurys et al. (2007, 2009). Our very crude size estimates should be taken as upper limits, because they are based on a low-energy transition, with E_{upp}/k of just 17 K, of a very abundant and easily thermalizable molecule. For the observations presented here, we note that the HPBW of the telescope is larger than these estimated sizes, and/or the excitation energies of the transitions are considerably higher (see Tables 1 to 3, and A.1 to A.3). Hence no significant amount of lost flux is expected in our data.

Because the envelope around VY CMa is peculiar (it is not dominated by a constant mass-loss wind that isotropically expands at constant velocity), the observed line profiles differ from those typical of circumstellar envelopes. VY CMa's molecular lines are very often characterized by up to three components: a central component close to the systemic velocity of $22\,\mathrm{km\,s^{-1}}$ and two high expansion velocity winds (HEVWs) placed symmetrically w.r.t. the central one at $\sim\!22\pm26.2\,\mathrm{km\,s^{-1}}$, that is, approximately at -4.2 and $48.2\,\mathrm{km\,s^{-1}}$. This profile decomposition was first noted by Muller et al. (2007) and Ziurys et al. (2007) and has been systematically used for the analysis of the spectral lines of VY CMa since, although some authors have identified up to four components in some cases (Fu et al. 2012). We

Table 2. Spectral-line results for carbon monoxide and water vapour.

- I	D 4 4' 1	F /1†	D 4 C	C 11, 0	+	D 1		X7.1	
Species and elec./vibr. state	Rotational quantum nums.	$E_{\mathrm{upp}}/k^{\dagger}$ (K)	Rest freq. (GHz)	Setting & sideband	rms [‡] (mK)	Peak (mK)	Area $(K \text{ km s}^{-1})$	Veloc. range LSR (km s ⁻¹)	Comments
$\frac{^{12}\text{CO }v = 0}{^{12}\text{CO }v = 0}$	J = 6-5	116	691.473	17 USB	13.4 (1.08)	1085	57.6	[-50; +120]	Comments
v = 0	J = 6-3 J = 10-9	304		06 LSB			80.6		
	J = 10-9 J = 16-15	752	1151.985 1841.346	16 USB	59.5 (1.04) 55.5 (0.98)	1554 1352	52.8	[-60; +100] [-30; +80]	
v = 1	J = 10-13 J = 5-4	3166	571.021	10 USB 14 USB	6.6 (1.05)	£20	\$2.8 \$0.15	[-30, +60]	
v = 1	J = 3 - 4 J = 15 - 14	3741	1710.861	04 LSB	66.5 (1.05)	≈200 ≲200	≈0.13 ≲7.00		
	J = 13-14	3741	1710.801	04 LSD	00.5 (1.05)	≈200	≈7.00		
13 CO $v = 0$	J = 6-5	111	661.067	12 USB	7.8 (0.91)	85	5.02	[-45; +80]	
	J = 9 - 8	238	991.329	09 LSB	30.5 (1.06)	131	6.49	g-fitted	
	J = 10-9	291	1101.350	07 LSB	22.0 (1.09)	173	7.20	g-fitted	
	J = 16-15	719	1760.486	03 USB	78.3 (1.02)	350	5.19	[-10; +50]	
$C^{18}O v = 0$	J = 6-5	111	658.553	12 USB	7.8 (0.91)	≲23	≲0.50		
ortho- $H_2O v = 0$	I = 1 1	27	556.936	14 LSB	6.6 (1.08)	1169	49.3	[-55; +88]	
$011110-\Pi_2O v = 0$	$J_{K_{a},K_{c}} = 1_{1,0} - 1_{0,1}$ $J_{K_{a},K_{c}} = 3_{0,3} - 2_{1,2}$	163	1716.770	04 USB	64.5 (1.05)	4685	193.8	[-30; +80]	
	$J_{K_a,K_c} = 3_{0,3} - 2_{1,2}$ $J_{K_a,K_c} = 3_{1,2} - 3_{0,3}$	215	1097.365	04 USB	23.1 (1.10)	1904	85.0	[-60; +115]	
	$J_{K_a,K_c} = 3_{1,2} - 3_{0,3}$ $J_{K_a,K_c} = 3_{1,2} - 2_{2,1}$	215	1153.127	06 LSB	59.5 (1.04)	2993	141.7	[-65; +85]	
	$J_{K_{\rm a},K_{\rm c}} = 3_{1,2} - 2_{2,1}$ $J_{K_{\rm a},K_{\rm c}} = 3_{2,1} - 3_{1,2}$	271	1162.912	06 USB	59.5 (1.04)	2140	96.8	[-50; +87]	
	$J_{K_a,K_c} = 5_{2,1} - 5_{1,2}$ $J_{K_a,K_c} = 5_{3,2} - 4_{4,1}$	698	620.701	13 USB	5.4 (0.97)	1716	39.5	[-55; +120]	maser
	$J_{K_a,K_c} = 5_{3,2} - 5_{2,3}$ $J_{K_a,K_c} = 5_{3,2} - 5_{2,3}$	698	1867.749	01 USB	90.9 (0.96)	1802	60.2	g-fitted	masci
$v_2 = 1$	$J_{K_{\rm a},K_{\rm c}} = J_{3,2} - J_{2,3}$ $J_{K_{\rm a},K_{\rm c}} = 1_{1,0} - 1_{0,1}$	2326	658.004	12 USB	7.8 (0.91)	5654	66.9	[-30; +80]	maser
02 1	$J_{K_a,K_c} = 2_{1,2} - 1_{0,1}$	2379	1753.914	03 USB	78.3 (1.02)	1007	34.8	g-fitted	maser
	$J_{K_a,K_c} = 8_{2,7} - 8_{3,4}$	3556	968.049	10 LSB	19.8 (1.08)	46	0.80	g-fitted	
$v_{1,2} = 1,1$	$J_{K_a,K_c} = 3_{1,2} - 3_{0,3}$	7749	1194.829	05 LSB	88.8 (1.01)	366	2.12	[-10:+50]	maser?
$para-H_2O v = 0$	$J_{K_{\rm a},K_{\rm c}} = 1_{1,1} - 0_{0,0}$	53	1113.343	07 USB	22.0 (1.08)	3141	136.5	[-50; +100]	
	$J_{K_a,K_c} = 2_{0,2} - 1_{1,1}$	101	987.923	09 LSB	33.1 (1.06)	3368	139.4	[-30; +75]	
	$J_{K_a,K_c} = 2_{1,1} - 2_{0,2}$	137	752.033	11 LSB	13.6 (1.00)	1637	79.4	[-50; +115]	
	$J_{K_a,K_c} = 4_{2,2} - 4_{1,3}$	454	1207.639	05 USB	88.8 (0.99)	1299	66.7	[-60; +100]	
	$J_{K_a,K_c} = 5_{2,4} - 4_{3,1}$	599	970.315	10 LSB	19.8 (1.08)	3188	72.3	[-30; +95]	maser
	$J_{K_a,K_c} = 6_{3,3} - 6_{2,4}$	952	1762.043	03 USB	78.3 (1.02)	1011	33.8	g-fitted	
$v_2 = 1$	$J_{K_{\rm a},K_{\rm c}} = 1_{1,1} - 0_{0,0}$	2352	1205.788	05 USB	88.8 (0.99)	244	7.36	g-fitted	
$v_2 = 2$	$J_{K_{\rm a},K_{\rm c}} = 2_{1,1} - 2_{0,2}$	4684	1000.853	09 USB	33.1 (1.05)	114	3.32	g-fitted	
$v_1 = 1$	$J_{K_a,K_c} = 3_{2,2} - 3_{1,3}$	5552	1868.783	01 USB	90.9 (0.96)	217	2.01	g-fitted	tent. detec.
$v_3 = 1$	$J_{K_{\rm a},K_{\rm c}} = 6_{3,3} - 6_{2,4}$	6342	1718.694	04 USB	64.5 (1.05)	169	4.79	g-fitted	tent. detec.
ortho- H_2^{18} O $v = 0$	$J_{K_{\rm a},K_{\rm c}} = 3_{1,2} - 3_{0,3}$	215	1095.627	08 LSB	23.1 (1.10)	229	5.68	[-10; +100]	
$para-H_2^{18}O v = 0$	$J_{K_{\rm a},K_{\rm c}} = 1_{1,1} - 0_{0,0}$	53	1101.698	07 LSB	22.0 (1.09)	161	3.53	g-fitted	
ortho- $H_2^{17}Ov = 0$	$J_{K_a,K_c} = 3_{0,3} - 2_{1,2}$	162	1718.119	04 USB	64.5 (1.05)	350	9.59	g-fitted	
	$J_{K_a,K_c} = 3_{1,2} - 3_{0,3}$	215	1096.414	08 LSB	23.1 (1.10)	281	7.59	[+0; +60]	
para- $H_2^{17}O v = 0$	$J_{K_{\rm a},K_{\rm c}} = 1_{1,1} - 0_{0,0}$	53	1107.167	08 USB	23.1 (1.08)	169	4.16	[-20; +60]	
Para 112 0 0 1 0	$J_{K_a,K_c} = 1_{1,1} 0_{0,0}$ $J_{K_a,K_c} = 2_{0,2} - 1_{1,1}$	101	991.520	09 LSB	33.1 (1.06)	72	2.72	g-fitted	tent. detec.
IIDO O					` ,			Č	
HDO v = 0	$J_{K_{a},K_{c}} = 3_{1,2} - 3_{0,3}$	168	753.411	11 LSB	13.6 (1.00)	≲40	≲1.00		

Notes. (†) Since the ortho and para spin isomer variants of water behave as different chemical species, for all ortho-water isotopomers the level excitation energies are relative to their lowest (ortho) energy level, the $J_{K_a,K_c}=1_{0,1}$, while for the para species they are given w.r.t. the $J_{K_a,K_c}=0_{0,0}$ level. (‡) For the spectral resolution in km s⁻¹ given in parentheses.

refer to these three components as the blue-HEVW, the central or systemic component, and the red-HEVW. Some lines exhibit all three components, others only the central one, and in some others the emission in the HEVWs is much stronger than that of the central component. No lines match this latter case in our survey, but some good examples can be found in the literature, for instance, the SO, SO₂, and HNC lines presented in Ziurys et al. (2007), Tenenbaum et al. (2010b), and references therein, many of which are emitted from levels with lower energies above the ground than the lines detected by us.

We adopted this component splitting in our work and tried to identify the contribution from these three components by fitting the spectra of some lines with three Gaussian shapes at fixed LSR velocities of -4.2, 22, and $48.2 \,\mathrm{km \, s^{-1}}$. To facilitate

the fitting, we also imposed that the widths of the two HEVWs were the same. We note that at lower frequencies, particularly in lines that are dominated by the HEVW components like those of the species mentioned before, the width of the red-HEVW is noticeably broader than that of the blue counterpart (see e.g. Figs. 1 and 3 in Tenenbaum et al. 2010b). However, this seems not to be our case. We see no indications of such a large difference between the red- and blue-HEVW widths in our spectra. For the few cases in which our lines have a high S/N and are free from blending of other lines, namely the three 12 CO lines and the J = 6-5 line of 13 CO in Fig. 3, the $J_K = 1_0^+ - 0_0^-$ of NH₃ and the OH lines in Fig. 8, and the J = 7-6 line of HCN in Fig. A.5, we tried to fit our data by allowing the widths of the two HEVWs to vary independently, which resulted in small differences, and the

Table 3. Spectral-line results for ammonia, hydrogen sulphide, hydroxyl, and silicon monoxide.

Species and elec./vibr. state	Rotational quantum nums.	$E_{ m upp}/k^{\dagger}$ (K)	Rest freq. (GHz)	Setting & sideband	rms [‡] (mK)	Peak (mK)	Area (K km s ⁻¹)	Veloc. range LSR (km s ⁻¹)	Comments
ortho-NH ₃ $v = 0$	$J_K = 1_0^+ - 0_0^-$ $J_K = 3_0^+ - 2_0^-$	27 170	572.498 1763.525	14 USB 19 LSB	6.6 (1.05) 77.8 (1.02)	308 1441	15.5 68.9	[-50; +65] g-fitted	
para-NH ₃ $v = 0$	$J_K = 3_2^+ - 2_2^- J_K = 3_1^ 2_1^+$	127 143	1763.821 1763.602	19 LSB 19 LSB	77.8 (1.02) 77.8 (1.02)	450 ≲100	21.0 ≲4.5	g-fitted	
ortho- $H_2S v = 0$	$J_{K_{\rm a},K_{\rm c}} = 3_{1,2} - 2_{2,1}$	117	1196.012	05 LSB	88.8 (1.01)	177	10.3	g-fitted	
para- $H_2S v = 0$	$J_{K_{\rm a},K_{\rm c}} = 3_{1,3} - 2_{0,2}$ $J_{K_{\rm a},K_{\rm c}} = 5_{2,4} - 5_{1,3}$	103 303	1002.779 1862.436	09 USB 01 LSB	33.1 (1.05) 90.9 (0.97)	97 590	4.22 11.8	g-fitted g-fitted	tent. detec.
OH $^{2}\Pi_{1/2} v = 0$	J = 3/2 - 1/2	270	1834.747	16 LSB	55.5 (0.98)	1768	85.5	[-40; +90]	
28 SiO $v = 0$	J = 14-13 J = 16-15	219 283	607.608 694.294	13 LSB 17 USB	5.4 (0.99) 13.4 (1.08)	370 384	15.5 15.1	[-40; +105] [-35; +85]	
v = 1	J = 10-13 J = 13-12 J = 15-14	1957 2017	560.325 646.429	14 LSB 12 LSB	6.6 (1.08) 7.8 (0.93)	76 108	2.13 3.01	[-35, +65] [-15; +67] [-20; +60]	
	J = 23-22 J = 27-26	2340 2550	990.355 1161.945	09 LSB 06 USB	33.1 (1.06) 59.5 (1.03)	178 190	5.12 5.36	g-fitted g-fitted	
v = 2	J = 40-39 J = 26-25	3462 4282	1717.236 1111.235	04 USB 07 USB	64.5 (1.05) 22.0 (1.08)	160 140	4.98 3.43	g-fitted g-fitted	tent. detec.
	J = 27-26 J = 28-27	4297 4355	1153.804 1196.353	06 LSB 05 LSB	59.5 (1.04) 88.8 (1.01)	925 50	25.3 2.50	g-fitted g-fitted	maser? tent. detec.
29 SiO $v = 0$	J = 13-12 J = 26-25	187 722	557.179 1112.798	14 LSB 07 USB	6.6 (1.08) 22.0 (1.08)	138 203	5.73 7.07	[-15; +75] g-fitted	
v = 1	J = 20-25 J = 16-15	2036	680.865	17 LSB	13.4 (1.10)	35	0.97	g-fitted	
30 SiO $v = 0$	J = 26-25 J = 42-41	713 1832	1099.708 1771.251	07 LSB 19 USB	22.0 (1.09) 77.8 (1.02)	231 153	6.47 3.68	[-20; +70] g-fitted	tent. detec.

Notes. (†) Since ortho- and para-NH₃ behave as different chemical species, for para-NH₃ the level excitation energies are relative to that of the lowest para level ($J_K = 1_1^-$), while for ortho-NH₃ are given w.r.t. the $J_K = 0_0^-$ level. Since the ortho and para spin isomer variants of hydrogen sulphide behave as different chemical species, for ortho-H₂S the level excitation energy is relative to its lowest (ortho) energy level, the $J_{K_a,K_c} = 1_{0,1}$, while for the para species they are given w.r.t. the $J_{K_a,K_c} = 0_{0,0}$ ground level. (‡) For the spectral resolution in km s⁻¹ given in parentheses.

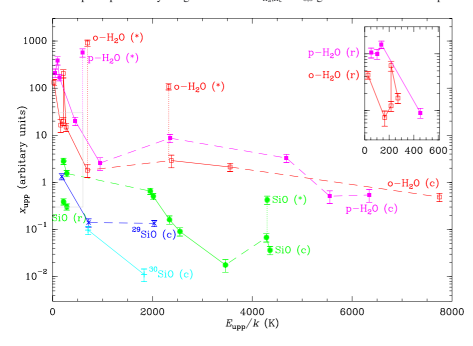


Fig. 1. Rotational diagrams for species discussed in Sect. 3 with emission from vibrationally excited levels: for H₂O and ²⁸SiO, central (c) and red-HEVW (r) components; for ²⁹SiO and ³⁰SiO, central component (c) only. Solid lines join upper levels in the same vibrational state, while dashed lines join upper levels in different vibrational states. ²⁸SiO and H₂O maser lines, indicated with an asterisk (*), are connected to their respective diagrams by vertical dotted lines.

red-HEVW was not always the widest of the two. Therefore, we decided to leave just five free parameters in the fitting procedure: the intensities of the three components, the width of the central component, and the width of the two HEVW components. We did not use this multi-component approach in the cases where

the S/N was low, when there was severe line blending (as the fitting of more than three/four components would have become very uncertain), and when the observed line profile clearly deviated from this triple-peaked template, as is the case of maser lines. Separating the molecular emission in these components

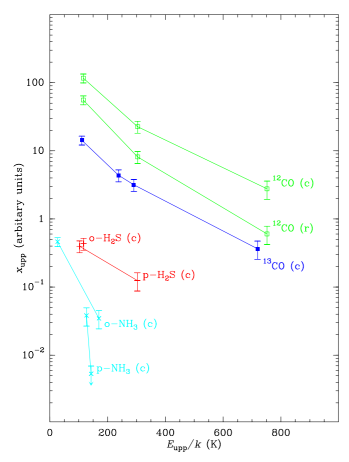


Fig. 2. Rotational diagrams for species discussed in Sect. 3 with no emission from vibrationally excited levels: for ¹²CO, central (c) and red-HEVW (r) components; for ¹³CO, NH₃, and H₂S, central component (c) only. Upper limits are marked with downward arrows.

allowed us to investigate their origin by studying their differing excitations. Some authors have argued that the HEVWs, and in general the high-velocity emission, arise from a wide-opening bipolar flow, whereas the emission from the central midvelocity component is due to a slower expanding isotropic envelope (Muller et al. 2007; Ziurys et al. 2007; Fu et al. 2012). However, other authors have tried to reproduce VY CMa's profiles using fully isotropic models consisting of multiple shells with different mass-loss rates (Decin et al. 2006).

We built rotation diagrams to readily obtain estimates for the excitation conditions of the different molecules and line components, but also to help in the line identification process. In these diagrams we computed the relative population of the upper levels, $x_{\rm upp}$, assuming that the lines are optically thin as well as a constant small size ($\ll 12''$) for the emitting region; see Eq. (23) in Goldsmith & Langer (1999). Although these assumptions are not always valid, the resulting diagrams still provided some useful information on the general trend of the excitation of a species. We inspected these rotational diagrams for the three components separately. Plots of these rotational diagrams can be seen in Figs. 1, 2, and A.1. From these diagrams, it is clear that the rotational temperature varies with the excitation energy of the lines (more highly excited lines tend to show higher rotational temperatures), as is expected in a medium with a steep temperature gradient

In the following subsections we describe the main observational results for the most relevant species: CO, H₂O, NH₃, H₂S, and OH, and SiO. The other detected species and unassigned features are discussed in detail in Appendix A.

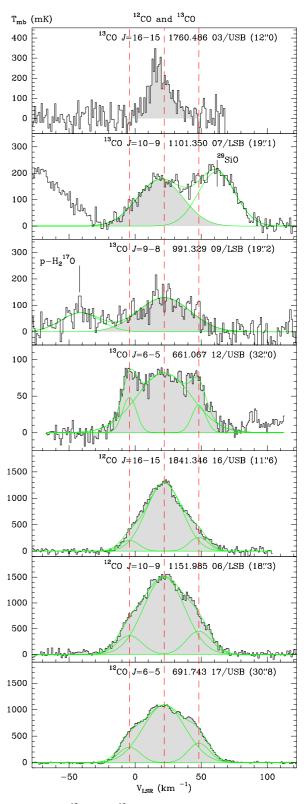


Fig. 3. Results for 12 CO and 13 CO from HIFISTARS observations. Line name, rest frequency (GHz), setting and side-band in which the line has been observed, and HPBW of the telescope at the frequency are also given. The grey area below the intensity histogram indicates the integrated line intensity quoted in Table 2. The horizontal axis gives the Doppler velocity for the adopted rest frequency. The intensities are given in $T_{\rm mb}$ scale. The vertical dashed lines mark the velocities adopted for the central and HEVW components. The green lines show Gaussian fits either for the HEVW and main components or for blended lines, and their composite profile. See Sects. 3 and 3.1 for more details. Labels for other lines in the spectra are also given in the cases of blended lines.

3.1. Carbon monoxide (CO)

The ^{12}CO and ^{13}CO results are presented in Fig. 3 and Tables 2 and A.4. We detected all the v=0 lines that we observed: the ^{12}CO J=6-5, J=10-9, and J=16-15, and the ^{13}CO J=6-5, J=9-8, J=10-9, and J=16-15. Two vibrationally excited lines of ^{12}CO , the J=5-4 and J=15-14 of v=1 were also observed, but were not detected; the corresponding upper limits provide no significant constraints on the excitation of these v=1 states. We did not detect the J=6-5 line of C^{18} O, which has been detected in other O-rich sources in HIFISTARS (see Bujarrabal et al. 2012; Justtanont et al. 2012); the obtained upper limit for the C^{18} O-to- 13 CO J=6-5 intensity ratio is consistent with the values for the O-rich stars detected in C^{18} O (≤ 0.1 , see Justtanont et al. 2012).

The ¹²CO spectra seen by HIFI are very different from those at lower frequencies and excitation energies (J = 3-2 and below, see e.g. Kemper et al. 2003; De Beck et al. 2010). Although we were able to identify the three *classical components* in the spectra, even in the J = 6-5 line the emission is highly dominated by the central component. This pre-eminence increases as we move up the rotational ladder: while in the J = 6-5 line the central-to-HEVW peak ratio is about 3, and the two HEVWs represent 24% of the total emission, in the J = 16-15 transition this peak ratio is about 5, and the HEVW emission only accounts for 16% of the total emission. The only spectrum in which the triple-peaked shape is clearly seen is the $^{13}CO\ J = 6-5$, where the central-to-HEVW peak ratio is only 2. In fact, this spectrum resembles very much those of the J = 2-1 and J = 3-2 ¹²CO lines. Although we did not try to separate the emission from the three components in the other ¹³CO spectra (because the J = 9-8 and J = 10-9 lines are blended and the J = 16-15 line is noisy), the contribution of the HEVWs in these lines is minor. In summary, it is evident that the excitation of CO in the HEVWs is lower than in the central component, especially for ¹³CO.

In spite of the different opacities expected for the two CO isotopic substitutions, the rotational temperature diagrams (see Fig. 2) give a similar excitation for the two species. Data for the main central component yield values between 120 and 210 K for ¹²CO, and just 10 K less for ¹³CO. As expected, when we examine the rotational diagram for the HEVW components of ¹²CO, we derive lower (but not very different) temperatures, 90 to 170 K. All the CO profiles are quite symmetric; for example, the blue-to-red HEVW intensity ratio is between 0.77 and 0.81 for ¹²CO, while for ¹³CO we obtain a value of 1.29. This means that the opacities of the CO lines in the HEVW components cannot be very high, since otherwise we should have detected some self-absorption in the red part of the spectra, because the temperature and velocity gradients in the envelope.

3.2. Water vapour (H₂O)

VY CMa HIFISTARS results for water vapour are shown in Figs. 4 to 7, and in Tables 2 and A.4. In total we have detected 27 water lines. Seven lines of ortho- H_2O , six lines of para- H_2O , two lines of both ortho- $H_2^{17}O$ and para- $H_2^{17}O$, and one line of ortho- $H_2^{18}O$ and para- $H_2^{18}O$ each, all from the ground-vibrational state, and eight lines of H_2O from several vibrationally excited states. Frequencies of several HDO transitions were also within the observed bands but, as expected, no HDO line was detected. We note the upper limit obtained the for $J_{K_a,K_c} = 3_{1,2}-3_{0,3}$ of HDO, which is about five to seven times lower than the intensities measured for the same transition of $H_2^{17}O$ and $H_2^{18}O$ (see Table 2).

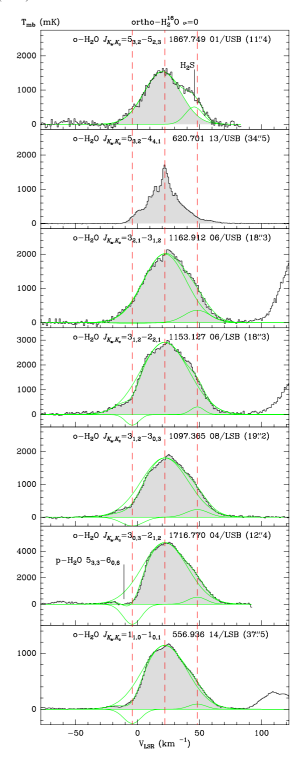


Fig. 4. Same as Fig. 3 for the detected v=0 lines of ortho-H₂O. Note the distinctive profile of the $J_{K_a,K_c}=5_{3,2}-4_{4,1}$ maser line. The location of the undetected $J_{K_a,K_c}=5_{3,3}-6_{0,6}$ line of para-H₂O is also indicated.

For ortho- and para- H_2O , we detected all the observed transitions with high line-strengths ($\Delta J=0,\pm 1$ with $\Delta K_a,\Delta K_c=\pm 1$) from the ground-vibrational state. We also detected the intrinsically weaker ($\Delta K_c=\pm 3$) $5_{2,4}$ – $4_{3,1}$ transition of para- H_2O , but its intensity and profile show that it is a maser (Figs. 5 and 1), as is normally the case for the other low line-strength v=0 transitions of water detected so far. We also detected maser emission from the (intrinsically strong) $J_{K_a,K_c}=5_{3,2}$ – $4_{4,1}$ transition of ortho- H_2O (Harwit et al. 2010). The highest excitation energy

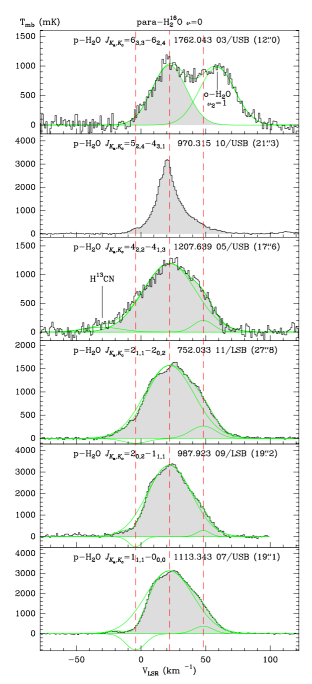


Fig. 5. Same as Fig. 3 for the detected v = 0 lines of para-H₂O. Note the distinctive profile of the $J_{K_a,K_c} = 5_{2,4}-4_{3,1}$ maser line.

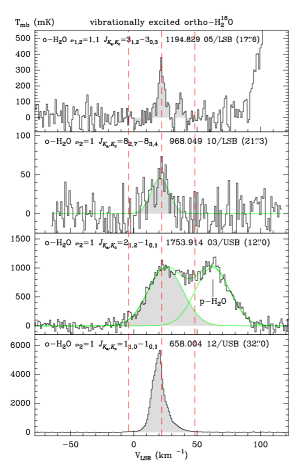
for these ground-vibrational transitions is the nearly 1000 K of the upper level of the $J_{K_a,K_c}=6_{3,3}-6_{2,4}$ line of para-H₂O (see Table. 2). We note that we did not detect the $J_{K_a,K_c}=5_{3,3}-6_{0,6}$ line of para-H₂O, which has been observed in other O-rich envelopes in HIFISTARS (Justtanont et al. 2012), the reason being that for VY CMa this transition blends with the much stronger $J_{K_a,K_c}=3_{0,3}-2_{1,2}$ line due to the broad linewidths (see Fig. 4).

The profiles of the lines of ortho- and para- H_2O , shown in Figs. 4 and 5, are not triple-peaked, but they can be decomposed into the same three components. We did not attempt this decomposition for the $5_{3,2}$ – $4_{4,1}$ (at $620\,\text{GHz}$) and $5_{2,4}$ – $4_{3,1}$ (at $970\,\text{GHz}$) lines because they are masers, nor for the $5_{3,2}$ – $5_{2,3}$ and $6_{3,3}$ – $6_{2,4}$ lines because they are blended. As for CO, the importance of the HEVW components decreases with the excitation of the line. The blue-HEVW component is always found

in absorption, whereas the red one is always found in emission. This asymmetry reflects the high opacity of the water lines in general, and that the H₂O excitation in the HEVWs is significantly lower than in the central component. If we inspect the rotational diagrams for the central component of the two spin isomers (Fig. 1), we realize that interpreting this plot is more complex than in the case of linear molecules, because the linestrength, level multiplicity, beam dilution, and frequency of the transitions do not monotonically increase with the excitation energy of the levels. However, we can estimate a rotation temperature of about 200 K for both ortho- and para-H₂O. Comparing the plots for the two spin isomers, the para-H₂O points lie about a factor 3-5 above the ortho points if we assume an ortho-topara abundance ratio of 3:1. Although this value is affected by the probably high opacity of the lines and the complex excitation of the molecule, the result suggests that the true ortho-topara ratio must be lower than 3 (see also Royer et al. 2010), and hence that water vapour is formed under non-equilibrium chemical conditions. The result of inspecting the rotational diagram for the red-HEVW component is less clear, but suggests a temperature as low as 100 K.

We systematically searched for a detection of water lines from its four lowest vibrationally excited states $v_2 = 1$, $v_2 = 2$, $v_1 = 1$, and $v_3 = 1$ at 2294, 4598, 5262, and 5404 K above the ground; more than 30 of these lines lie within the observed frequency ranges. We were only able to identify the emission from seven lines, see Table 2: four lines from the $v_2 = 1$ state, and one line for each of the other three vibrationally excited states listed before. We also tentatively assigned the feature at 1194.829 GHz (setting 05 USB, see Fig. B.2) to the $3_{1,2}-3_{0,3}$ line of the $v_{1,2} = 1,1$ state with E_{upp}/k of 7749 K, which holds the excitation record for all species in our survey. We only detected vibrationally excited lines with Einstein-A coefficients higher than $10^{-2} \,\mathrm{s}^{-1}$, except for the $v_2 = 1 \,J_{K_a,K_c} =$ $1_{1.0}$ – $1_{0.1}$ and $8_{2.7}$ – $7_{3.4}$ lines at 658 and 968 GHz. The 658 GHz line is a maser (Menten & Young 1995), and shows the highest peak flux in our spectral survey. The profile of the $3_{1,2}$ – $3_{0,3}$ line of the $v_{1,2} = 1,1$ state is also suggestive of maser amplification. The profiles and intensities of the other six lines are consistent with thermal emission and only display the central component at $22 \,\mathrm{km \, s^{-1}}$. We did not detect the $v_3 = 1 \, 3_{0.3} - 2_{1.2}$ line, which was observed at the same time as the $v_3 = 16_{3,3}-6_{2,4}$ and should have a similar intensity; for this reason and the low S/N, the detection of this latter line is considered tentative. The remaining non-detected vibrationally excited lines of H₂O do not provide significant constraints on the excitation of these levels. We note that the 658 and 970 GHz maser lines are remarkably smooth and do not show narrow (~1 km s⁻¹ wide) spikes like the strong 22.2 GHz ortho- H_2O v = 0 $6_{1,6}$ – $5_{2,3}$ maser line or strong ²⁸SiO maser lines. (The 620 GHz $v = 0.5_{3,2}$ –4_{4,1}, maser line shows a few narrow features.) In the first detection paper of the 658 GHz line, Menten & Young (1995) argued that the greater *smoothness* of this line's profile compared with that of the 22.2 GHz maser, arise because the 658 GHz maser is saturated while the 22 GHz is not.

The rotational diagrams of vibrationally excited H_2O are shown in Fig. 1. Excluding maser lines, we only detected three excited transitions of ortho- H_2O and four lines of para- H_2O , but they mostly originate from different vibrationally excited states. Therefore we can hardly investigate the rotational temperatures in the vibrationally excited ladders: we only derive an excitation temperature of 3000 K between the two *thermal* $v_2 = 1$ lines of ortho- H_2O . However, we can study the relative excitation between the different vibrationally excited states, for which we



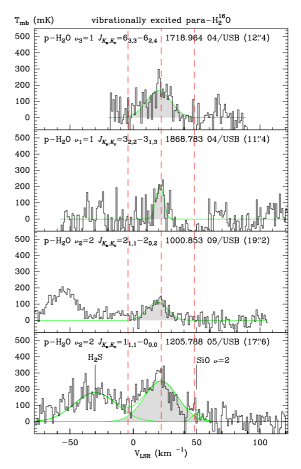


Fig. 6. Same as Fig. 3 for the detected lines of vibrationally excited water. Note the narrow profile of the $v_2 = 1$ $J_{K_a,K_c} = 1_{1,0}-1_{0,1}$ line in the bottom-left panel, indicating its maser origin. This may also be the case of the $v_{1,2} = 1,1$ $J_{K_a,K_c} = 3_{1,2}-3_{0,3}$ line in the top-left panel.

derive excitation temperatures between 1500 and 3000 K. These values are much higher than those derived for the rotational levels of the ground-vibrational state (see before), suggesting that the population of the vibrational excited levels is mostly due to IR pumping, at 6.2 and $2.7\,\mu\text{m}$, similar to the cases of other species such as SiO (Sect. 3.4) and SO (Sect. A.1).

For the rare oxygen isotopic substitutions of water, $H_2^{17}O$ and H₂¹⁸O, we only detected lines from their ground-vibrational levels with $E_{\rm upp}/k$ below 250 K. Several lines with excitation energies of about 500 K are also present in the observed bands, but they happen to blend with other (stronger) lines, and therefore it is difficult to tell whether they are detected. However, we did not detect the 5_{3.2}-4_{4.1} line of any of the two species, whose upper level is at more than 700 K above the ground. Neither did we detect the $v_2 = 1 \ 1_{1,0} - 1_{0,1}$ line of ortho- $\tilde{H}_2^{18}O$. Among the six detected lines (see Fig. 7), we were able to identify the triple profile structure only in the lowest-energy line of para-water $(1_{1,1}-0_{0,0})$ for both H₂¹⁷O and H₂¹⁸O. As was also the case for the main isotope, the line is very asymmetric, but here the absorption due to the blue-HEVW is so deep that the flux is below the continuum level, for which we measured a value of 70-100 mK at these frequencies. These P-Cygni profiles were previously observed by Neufeld et al. (1999) in the main isotopomer in the three cases in which the line shape was resolved in their ISO observations, those of the $J_{K_a,K_c} = 7_{2,5}-6_{1,6}$, $4_{4,1}-3_{1,2}$, and $4_{3,2}-3_{0,3}$ transitions, with $E_{\rm upp}/k$ at 1126, 702, 550 K, respectively. Assuming that the continuum level detected around 1100 GHz has an extent similar to the $H_2^{17}O$ and $H_2^{18}O$ molecules responsible for the P-Cygni profiles, we conclude that the opacity in the J_{K_a,K_c} = $1_{1,1}$ - $0_{0,0}$ line of both species must be $\gtrsim 1$, and that the excitation of the blue-HEVW must be significantly lower than the true dust brightness temperature, which is about $40\,\mathrm{K}$ for a source size of 1''.

The rotational diagrams for the central component do not give a reliable estimate of the excitation temperature. We only have two lines per species at most, so we cannot distinguish between general trends and peculiarities in the excitation of the individual lines. Taking all transitions together, we see that the abundances of $H_2^{18}O$ and $H_2^{17}O$ are very similar and that again an ortho-to-para abundance ratio lower than 3:1 fits the data better. Compared with the main isotopologue, the lines are 10 to 20 times weaker. This is most likely due to the high opacity of the H_2O lines, but note that the opacity of the $H_2^{18}O$ and $H_2^{17}O$ cannot be negligible, as discussed above.

3.3. Other hydrides

3.3.1. Ammonia (NH₃)

We observed four rotational lines of NH₃, two lines of the ortho $(J_K = 1_0^+ - 0_0^- \text{ and } 3_0^+ - 2_0^-)$ and two lines of the para $(J_K = 3_1^- - 2_1^+ \text{ and } 3_2^+ - 2_2^-)$ species. These results are presented in Fig. 8. The results of the $J_K = 1_0^+ - 0_0^-$ have already been published by Menten et al. (2010). The profile of this line is somehow unique in our survey because it is slightly U-shaped. The line is somewhat stronger at positive velocities, suggesting a self-absorption effect or some intrinsic asymmetry in the emitting region, and it displays intensity bumps at 11 and 35 km s⁻¹ in addition to the central component at 22 km s⁻¹ and the HEVWs at -4 and +48 km s⁻¹. This line is very similar to the $J_N = 7_6 - 6_5$ of SO (see Fig. 3 in Tenenbaum et al. 2010b) and resembles

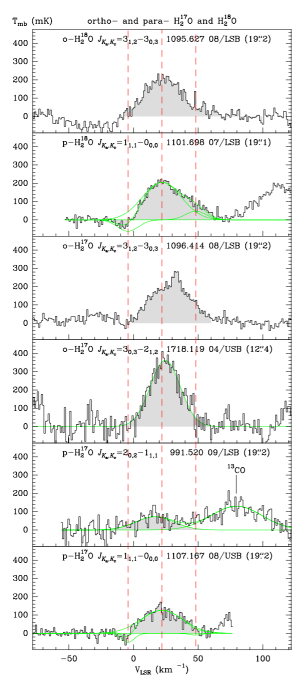


Fig. 7. Same as Fig. 3 for the detected lines of isotopomers of water.

those of $^{12}\text{CO}\ J=2-1$ and J=3-2 (see Kemper et al. 2003; De Beck et al. 2010). Note that the excitation energy of the $J_K=1_0^+$ level of NH₃, 27 K, is in between those of the J=2 and J=3 levels of ^{12}CO (17 and 33 K, respectively), but is significantly lower than that of the upper level of the $J_N=7_6$ of SO (47 K). The total width⁵ of the line is also similar to those of the CO lines, extending from about $-30\,\mathrm{km}\,\mathrm{s}^{-1}$ to $+70\,\mathrm{km}\,\mathrm{s}^{-1}$. In spite of the relatively more complex profile of this line, we also characterized its shape by fitting the three Gaussians; these results are shown in Fig. 8 and Table A.4.

Although the three observed J = 3-2 transitions appear to be blended, it is clear that in these higher excitation lines only

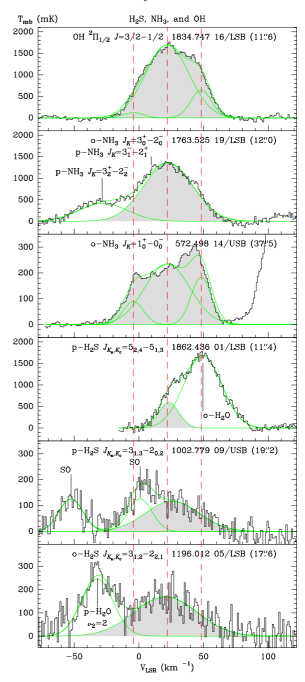


Fig. 8. Same as Fig. 3 for the detected lines of other hydrides: H_2S . NH_3 , and the OH radical.

the central component is present. We separated the contribution from the three lines by simultaneously fitting three Gaussians (only the central component for each transition), assuming that the velocity is the same (see Fig. 8). Our fitting shows that both the $J_K = 3_0^+ - 2_0^-$ and $3_2^+ - 2_2^-$ lines are well detected, showing FWHMs similar to that of the central component of the $J_K = 1_0^+ - 0_0^-$ line (see Table A.4). In contrast, the presence of $J_K = 3_1^+ - 2_1^+$ emission is not required by the fitting; this line should be at least four times weaker than the $3_2^+ - 2_2^-$ line. Using the two lines detected for ortho-NH₃, we derive a rotational temperature of 55 K for the central component, lower than what we found for other species. This excitation value also agrees with the detected intensity of the $3_2^+ - 2_2^+$ transition if we adopt an orthoto-para abundance ratio of 1:1, but is clearly in conflict with the relative weakness of the $3_1^- - 2_1^+$ line, whose upper level lies

⁵ The splitting of the hyperfine structure of these rotational lines of ammonia is very small, just $1.6 \,\mathrm{km}\,\mathrm{s}^{-1}$ in the J=1-0 line and less than $0.5 \,\mathrm{km}\,\mathrm{s}^{-1}$ in the J=3-2 lines.

only 15 K above the ground state of para-NH₃. However, it must be noted that in VY CMa the $3_1^--2_1^+$ line of para-NH₃ blends with the $3_0^+-2_0^-$ one of the ortho species. Since the multiplicity of the K=0 levels is four times that in the K=1 ladder, the $J_K=3_0^+-2_0^-$ line could be significantly more opaque than the $3_1^--2_1^+$ one, and hence the intensity due to the para transition would be practically fully absorbed by the ortho one if this latter is optically thick.

3.3.2. Hydroxyl radical and hydrogen sulphide (OH and H₂S)

We detected the $^2\Pi_{1/2}$ J=3/2-1/2 transition of OH at 1834.747 GHz (see Fig. 8). The line is among the strongest in our survey, only surpassed by the strongest lines of water. The profile⁶ is dominated by the central component and displays a strong asymmetry between the two HEVWs; the intensity of the red-shifted component is nearly six times that of the blue one. The profile is among the widest we have observed in our HIFI survey, extending from -30 to +75 km s⁻¹, together with the low-level lines of water. This OH line, with $E_{\rm upp}/k$ of 270 K, resembles the v=0 32,1-31,2 and 42,2-41,3 lines of H₂O, with $E_{\rm upp}/k$ of 271 and 454 K, respectively. The similarity between the profiles of lines of similar excitation of these two molecules suggests that these two species are also co-located, in agreement with the hypothesis that the main source of OH formation is the photo-dissociation of H₂O.

The velocity range observed in the OH 1834.747 GHz line can be compared with those of the OH masers at 18 cm, as the infrared pumping of these masers via the absorption of stellar IR photons at 34.6 and 53.3 μ m relates these Λ -doubling transitions in the ${}^{2}\Pi_{3/2} J = 3/2$ (ground) state with those connected by the rotational line observed with HIFI. The maser emission at 1612 MHz ($F = 1^+-2^-$) spans the velocity range between -15and +60 km s⁻¹, and comes from a region about 1".8 in diameter (see Benson & Mutel 1979, and references therein), 3×10^{16} cm for a distance of 1.15 kpc. The velocity range of the 1612 MHz maser is only slightly smaller than the 1834.747 GHz thermal line. The extreme velocities are only $10 \, \mathrm{km} \, \mathrm{s}^{-1}$ lower, but the two maxima of the U-shaped 1612 MHz line occur at ~-8 and +50 km s⁻¹, that is, very close to the location of our HEVW components (-4.2 and 48.2 km s⁻¹). These results suggest that the 1612 MHz maser arises from the shells that give rise to our HEVW components, and that these layers have a size of just a few arc-seconds. In addition, the fact that the thermal line detected here is dominated by the central component suggests that the photo-dissociation of water is taking place not only in the outermost layers of the envelope where the OH masers are formed, but also in the accelerated inner regions of the circumstellar shell.

We detected three lines of H₂S: the $J_{K_a,K_c}=3_{1,2}-2_{2,1}$ of ortho-H₂S, and the $3_{1,3}-2_{0,2}$ and $5_{2,4}-5_{1,3}$ of para-H₂S. A total of 20 lines of H₂S lie within the HIFI observed bands, but we only detected intrinsically strong (ΔK_a , $\Delta K_c=\pm 1$) transitions with $E_{\rm upp}/k \lesssim 300$ K. The spectra of the observed lines are shown in Fig. 8. All three lines appear to be blended with lines of SO and H₂O, therefore we determined their intensities by means of multiple-Gaussian fittings. In this case, the fittings are particularly uncertain in view of the resulting line parameters (widths and centroids). Therefore, the results on H₂S need be interpreted with caution. Assuming an ortho-to-para abundance ratio of 3:1,

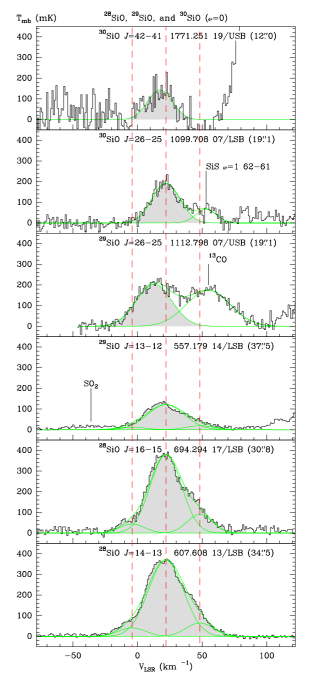


Fig. 9. Same as Fig. 3 for the detected v = 0 lines of ²⁸SiO, ²⁹SiO, and ³⁰SiO

we derive a rotational temperature of 165 K, in agreement with the non-detection of lines with excitation energies $\gtrsim 800$ K.

3.4. Silicon monoxide (SiO)

We detected 15 lines of the three most abundant isotopic substitutions of silicon monoxide, 28 SiO, 29 SiO, and 30 SiO, which have relative abundances of 30:1.5:1 in the solar system. These results are presented in Figs. 9 and 10 and in Table 3. For the main isotopic substitution, 28 SiO, we detected the only two lines from the ground-vibrational state, v = 0, that we observed, the J = 14–13 and 16–15 transitions, with upper levels at excitation energies of 219 and 283 K, respectively. We also detected all the observed lines from the first and second vibrationally excited states, five v = 1 transitions between the J = 13–12

⁶ We note that this transition has three hyperfine components, at -0.6, 0.0, $+1.9 \, \mathrm{km \, s^{-1}} \, \mathrm{w.r.t.}$ the quoted frequency, but this small total separation of $2.5 \, \mathrm{km \, s^{-1}}$ has little influence on the discussion that follows.

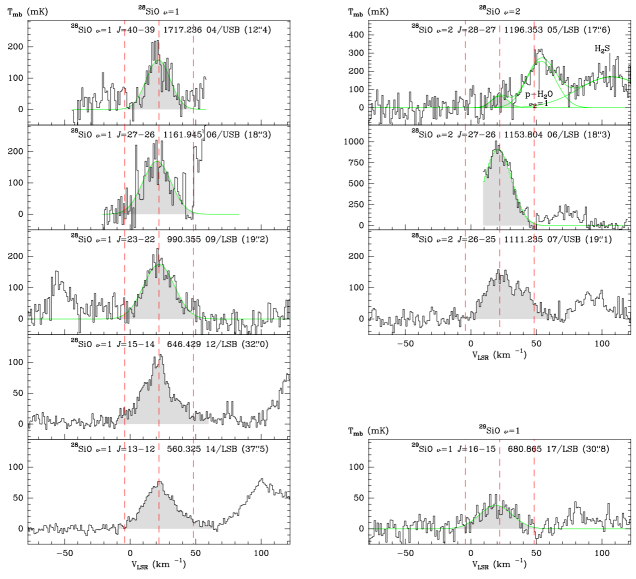


Fig. 10. Same as Fig. 3 for the detected lines of vibrationally excited 28 SiO, and 29 SiO (bottom-right panel). The 28 SiO v=2 J=27–26 line possibly shows some maser amplification.

and the J=40-39, with $E_{\rm upp}/k$ of 1957 and 3462 K, and the contiguous v=2 transitions J=26-25, 27–26, and 28–27, at about 4300 K above the ground level, the detection of the latter one is tentative. We also observed other lines of ²⁸SiO between rotational levels in vibrational states with yet higher excitation, $v \ge 3$, but this yielded no detections.

We detected all the observed rotational lines of 29 SiO and 30 SiO from the ground v=0 vibrational state. These lines range from the J=13-12 of 29 SiO at 187 K excitation energy to the J=42-41 of 30 SiO with $E_{\rm upp}/k$ of 1832 K. We only detected one vibrationally excited line from these two rare isotopic substitutions: the v=1 J=16-15 of 29 SiO. We also observed the v=1 J=18-17 line of 29 SiO, but this transition overlaps the strong $2_{1,1}-2_{0,2}$ line of para-H₂O, therefore we cannot draw conclusions on its detection. We also observed, but failed to detect, the v=1 J=41-40 transition of 30 SiO. No other v=1 lines of 29 SiO or 30 SiO are covered by our observations. We also covered several $v \ge 2$ transitions of the two rare isotopic substitutions of SiO, but these lines remain undetected.

From the comparison of the rotational diagrams for the different vibrational ladders of ²⁸SiO, we can roughly derive the

excitation conditions of these vibrationally excited states. This is a very relevant issue for the pumping of ²⁸SiO masers from low-J v = 1 and 2 states, which are widely observed in O-rich (and some S-type) long-period variables. From this comparison we infer vibrational excitation temperatures of 1500 K between the v = 0 and v = 1 rotational ladders of ²⁸SiO and of 2500 K between the v = 1 and v = 2 states, which are similar to the excitation energy of these v > 0 states (the v = 1 and 2 states are about 1770 K and 3520 K above the ground, respectively). These temperatures are significantly higher than those derived just from the comparison of v = 1 rotational transitions of ²⁸SiO, for which we obtain values between 300 and 550 K, much lower than the excitation energy of the levels, and similar to those derived for the v = 0 states of all SiO species. Radiation only connects levels with $\Delta J = 1$, and therefore the resulting excitation of the v > 0levels can in principle mimic the much lower collisional excitation of the v = 0 states. In contrast, the collisional pumping of v > 0 states would yield similar (high) excitation temperatures (of the order of 1700 K and higher) for both pure rotational transitions in vibrationally excited states and ro-vibrational transitions. Our result supports the radiative-pumping model for the

classical ²⁸SiO masers (J = 1–0 and J = 2–1 in v = 1 and v = 2) against the collisional-pumping model (see Lockett & Elitzur 1992; Bujarrabal 1994, for more details).

The detected v=2 transitions of 28 SiO deserve some attention. The three lines connect the four contiguous levels J=25 to 28. The strength of J=27–26 line is between seven and ten times higher than the other two transitions, suggesting some over-excitation or maser effect in this line. Although the inversion of low-J masers of 28 SiO can be explained with standard radiative models, the presence of masers in higher-J lines typically requires some additional effects, such as line overlapping with ro-vibrational lines of 28 SiO or of some other abundant molecule (such as water). In our case, we identified a potential overlap between the ro-vibrational lines J=27–28 v=2–1 of 28 SiO ($\lambda=8.519845\,\mu{\rm m}$) and $J_{K_a,K_c}=10_{3.8}$ – $11_{4.7}$ $v_2=1$ –0 of ortho-H₂O ($\lambda=8.519933\,\mu{\rm m}$), which are separated only by 3 km s⁻¹.

The line profiles of the v = 0 lines (Fig. 9) display triangular or Gaussian-like shapes, which are characteristic of the envelope layers where the acceleration of the gas is still taking place and the final expansion velocity has not been attained yet. This is not surprising because the high dipole moment of SiO favours the selection of densest gas in the envelope, and SiO is expected to be severely depleted onto grains as the acceleration proceeds along with the dust formation. However, in spite of this, the two detected ²⁸SiO transitions show a velocity range as wide as that covered by CO emission, from about -30 to $+70 \, \mathrm{km \, s^{-1}}$ (about $\pm 50 \, \mathrm{km \, s^{-1}}$ w.r.t. the systemic velocity of the source). Therefore, we can conclude that there are significant amounts of SiO in the gas phase even in the regions where the acceleration of the envelope and the grain growth have both ceased. A three-component Gaussian fit to the SiO lines (see Table A.4) gives FWHMs of 28–32 km s⁻¹ for the central component, that is, about five to ten km s⁻¹ narrower than those of the 12 CO lines with similar excitation energy; only the J =16-15 line of ^{12}CO shows such a narrow central component, but at higher excitation. Considering their total velocity extent and the width for single-component Gaussian fits (Fig. 10), the profiles of the vibrationally excited SiO lines are certainly narrower than those of lines from v = 0 states. ²⁸SiO v = 1 lines extend from -10 to $60 \,\mathrm{km \, s^{-1}}$ at low-J and from 0 to $45 \,\mathrm{km \, s^{-1}}$ at high-J. The Gaussian fits yield FWHMs of about $28 \, \mathrm{km \, s^{-1}}$. This also applies to the the ²⁸SiO v = 2 lines and the v = 1 J = 16-15of ²⁹SiO. These values are very similar to those obtained for the vibrationally excited lines of para-H₂O, suggesting that in the inner regions of the envelope where these vibrationally excited lines arise, the turbulence and/or acceleration are relatively strong.

4. Conclusions

4.1. Sub-mm/FIR spectrum of VY CMa

From our observations we conclude that the sub-mm/FIR spectrum of VY CMa's envelope is very rich in molecular spectral lines, in agreement with previous surveys (see Sect. 1). Considering this high density of lines and the broad intrinsic width of the spectral features in this source, between 50 and $100 \, \mathrm{km \, s^{-1}}$ ($100{-}600 \, \mathrm{MHz}$), we have almost reached the line confusion limit. About 48% of the observed bandwidth is occupied by molecular emission. In some cases, see the results from settings 14 and 08 in Fig. B.1, it has been very difficult to find emission-free areas to use for the residual baseline subtraction. In spite of the wide frequency range covered by our survey and

the large number of detected molecular lines, we have not identified any new species.

We mostly detected diatomic oxides, CO, SiO, OH, SO, AlO, and PO. SiS and PN are the only diatomic species that do not contain oxygen. Most of the polyatomic species are hydrides, H₂O, H₂S, HCN, and the only species with four atoms, NH₃. The only other polyatomic molecule is SO₂, a dioxide. If we include the isotopic substitutions, the spectrum is dominated by lines of ²⁸SiO (ten lines, plus five more from ²⁹SiO and ³⁰SiO), ortho-H₂O (11+3) and para-H₂O (10+3), SO (14), HCN (8+2), CO(3+4), and PO(7). SO_2 is a special case. We may have detected up to 34 lines, but some of them are indistinguishable because they are severely blended. Of the 130 assigned lines, 24 (about 18%) are due to vibrationally excited states. This relatively high prevalence of highly excited lines is most likely due to the strong IR radiation field of the star and the dust envelope. This efficiently pumps molecules from the v = 0 states to vibrationally excited levels, greatly enhancing the emission of these transitions. We found that this effect is particularly strong for H₂O, SiO, SO, and HCN. The measured high intensities of some lines also require the contribution of an additional source of excitation, such as the line overlap of IR lines.

The structure of the lines can be decomposed into several components. The lines are typically dominated by a central component of triangular or Gaussian shape that peaks at the systemic velocity of the source ($22\,\mathrm{km}\,\mathrm{s}^{-1}$ LRS) with FWHM values $\sim 20-45 \,\mathrm{km}\,\mathrm{s}^{-1}$. This emission is sometimes accompanied by two HEVW components, centred at \sim 22 ± 26 km s⁻¹. The width of the HEVWs is narrower, between $11-25\,\mathrm{km\,s^{-1}}$, but suggests a relatively strong (either micro- or most likely largescale) turbulence in the gas. These two HEVW components are almost only present in lines with a relatively low excitation energy. Nevertheless, the extreme velocities displayed by both central and HEVW emission are very similar, reaching values of $\pm 50 \,\mathrm{km \, s^{-1}}$ w.r.t. the systemic velocity of the star. This high nebular expansion rate is similar to that of other super- and hyper-giants in the HIFISTARS sample, except for Betelgeuse, which has a significantly slower wind (see Teyssier et al. 2012).

HIFISTARS observations mostly probe the warm gas in the envelope: only $\rm H_2O$, $\rm NH_3$, and $\rm H_2S$ have lines with upper levels at excitation energies below that of the J=6 of $\rm ^{13}CO$ and $\rm ^{12}CO$ ($E_{\rm upp}/k$ of 111 and 116 K, respectively). In these cases we can see how the intensity of the HEVW components decreases with increasing excitation energy. In other species we mostly see the central component. Their triangular profiles suggest a strong acceleration in the velocity field of the gas: in some species we detected a trend of decreasing line-width with the excitation of the line (see Table A.4).

The clear separation between the HEVWs and the central component suggests that the structures responsible for these two emissions are physically detached. The presence of several mutually detached shells in the envelope around VY CMa agrees with the detailed images of this object (see Sect. 1). These nested structures have also been detected in the envelopes of other massive stars (AFGL 2343 and IRC 10420, see Castro-Carrizo et al. 2007), and may be characteristic of this type of objects. We note that in VYCMa the profiles of optically thin lines, such as ¹³CO J = 6-5 and NH₃ $J_K = 1_0^+ - 0_0^-$, are very similar to other massive sources in the HIFISTARS sample (Teyssier et al. 2012). All of these suggest discontinuous mass loss. Water is detected in the HEVW and central components, and the same applies to OH, meaning that the formation of OH from the dissociation of H₂O does not only occur in the outermost layers of the envelope. The lower excitation and high opacity of water

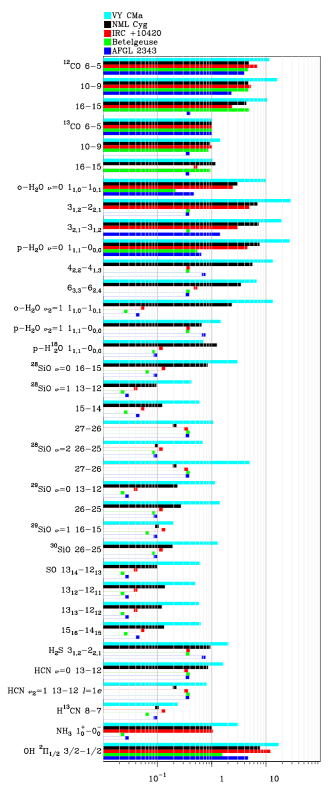


Fig. 11. Comparison of the integrated emission of different lines in HIFISTARS for VY CMa and the other four hyper- and super-giants in the project (see Teyssier et al. 2012). For each object, the integrated intensities of the different lines have been normalized to that of its J=6-5 13 CO transition. Horizontal bars and small squares show, respectively, values and upper limits for detected and undetected lines. See Sect. 4.2 for more details.

in the outer parts of the envelope results in strong self-absorbed profiles in its low-lying lines, which in the case of $H_2^{17}O$ and $H_2^{18}O$ extend below the continuum level.

Typically, we measured rotational temperatures between 150 and $500\,\mathrm{K}$ for the central component, even among transitions originating from vibrationally excited levels. These values are usually lower for the HEVW components, between 75 and $200\,\mathrm{K}$.

4.2. VY CMa and other massive stars in HIFISTARS

To compare our HIFISTARS results for VY CMa with those obtained for other hyper- and super-giants in the project, which have been published by Teyssier et al. (2012), we followed a procedure similar to the one used in that paper. These authors compared the peak fluxes of the different lines detected in NML Cyg, IRC +10420, Betelgeuse, and AFGL 2343, correcting for the effects of the distance, size of the envelope, and amount of molecular gas, by normalizing all values to the peak flux of the J = 6-5 ¹²CO line. We included VY CMa in the comparison and increased the number of studied transitions. Here we chose to represent the integrated area values for the lines, because these figures are better correlated with the total amount of molecular gas, and, for non-detections, the corresponding upper limits provide better constraints. For these undetected lines, we took as upper limit the usual value of $3\sigma \times \delta v \times \sqrt{n}$, where δv is the velocity resolution and n is the number of channels typically covered by the lines in the source at zero power. This corresponds to $60 \,\mathrm{km}\,\mathrm{s}^{-1}$ for NML Cyg, IRC +10420, and AFGL 2343, and 30 km s⁻¹ for Betelgeuse. To avoid problems with the possible high opacity of the ¹²CO lines, we used the integrated intensity of the 13 CO J = 6-5 transition as the normalization factor. The results of this comparison are presented in Fig. 11.

As we can see in this figure, VY CMa is the strongest emitter in lines of less abundant molecules (other than ¹²CO and H₂O). This result cannot be due to a relatively weak intensity of the 13 CO J = 6-5 in VY CMa, which might be the case if 13 C were less abundant than in the other sources, since VY CMa is also the strongest emitter in the 13 CO J = 10-9 and H 13 CN J =8–7 lines. Therefore we must conclude that these rare molecular species are more abundant in VYCMa than in the other four sources. VY CMa is followed by NML Cyg, and then by IRC +10420. Betelgeuse and AFGL 2343 are the weakest of all. The contrast between VY CMa and the other sources is stronger for lines with higher E_{upp}/k , such as v > 0 SiO lines and the ortho-H₂O $v_2 = 1$ 1_{1.0}-1_{0.1} maser line, but also for the relatively low excitation lines of SO included in the plot. Interestingly, the only two lines in VY CMa that are surpassed by NML Cyg are 13 CO J = 16-15 and para- H_2^{18} O $1_{1,1}-0_{0,0}$. The relative weakness of the para-H₂¹⁸O in VY CMa, w.r.t. NML Cyg, contrasts with the strength of all other H₂O lines, suggesting a lower abundance of this isotopic substitution in our target. We also note that C¹⁸O remains undetected in our VY CMa spectra.

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to the study of the warm gas and water vapour contents of the molecular envelopes around evolved stars: AGB stars, red super- and hyper-giants; and their descendants: pre-planetary nebulae, planetary nebulae, and yellow hyper-giants. HIFISTARS comprises 366 observations, totalling 11 186 min of Herschel/HIFI telescope time. See http://hifistars.oan.es; and Key_Programmes.shtml and UserProvidedDataProducts.shtml in the Herschel web portal (http://herschel.esac.esa.int/) for additional details. This work has been partially supported by the Spanish MICINN, program CONSOLIDER INGENIO 2010, grant "ASTROMOL" (CSD2009-00038); and by the German Deutsche Forschungsgemeinschaft, DFG project number Os 177/1-1. A portion of this research was performed at the Jet Propulsion Laboratory, California Institute of Technology, under contract with the National Aeronautics and Space Administration. R.S.z. and M.S.ch. acknowledge support from grant N203 581040 of National Science Center. K.J., F.S., and H.O. acknowledge funding from the Swedish National Space Board. J.C. thanks for funding from the Spanish MICINN, grant AYA2009-07304.

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Appendix A: Detailed results for species not discussed in Sect. 3

A.1. Sulphur monoxide (SO)

We detected three rotational J_N – $(J-1)_{(N-1)}$ triplets (N=J, J-1)and J+1) of SO, the J = 13-12, 15–14 (only one component), and 23–22, with $E_{\rm upp}/k$ from about 190 to 580 K. We thus detected all the high line-strength SO transitions within the observed settings, except for one component of the J = 41-40, which lies at one of the edges of the setting 03 observation, and for which the upper limit obtained is not significant. Some other lines of SO of much lower line-strength, with J_N – $(J-1)_N$ or with $\Delta J > 1$, also lie within the observed frequency ranges but yielded non-detections. In general, the upper limits of these lines are not significant either, except for those of the relatively low-lying lines $J_N = 9_9 - 8_9$ and $10_{10} - 9_{10}$, at 568.741 and 609.960 GHz respectively, and with upper-level energies of 106 and 127 K above the ground. Using the information from detections and the relevant upper limits, we built a rotation diagram for SO for the central component (see Tables A.1 and A.4 and Fig. A.1). We derive a rotation temperature of about 200–250 K, similar to that derived for other species from levels of similar excitation energies. The upper limits obtained for the lines at 106 and 127 K excitation energy indicate that the central component of SO cannot have a significant contribution from gas colder than about 60 K.

In addition to these ground-vibrational lines, we detected some SO lines from the first vibrationally excited state v=1, the J_N – $(J-1)_{(N-1)}$ triplets J=26–25 and J=27–26 at 1108 and 1150 GHz, and one component (the other two are blended with a very strong water line) of the J=23–22 triplet at 980.531 GHz. The upper levels of these lines lie between 2178 and 2386 K above the ground. As in the case of SiO, when comparing the rotational diagram of these v=1 transitions with those from the v=0, we see that the rotational temperature is very similar in spite of the much higher excitation energy of the v=1 levels. In contrast, when we compare the intensity of

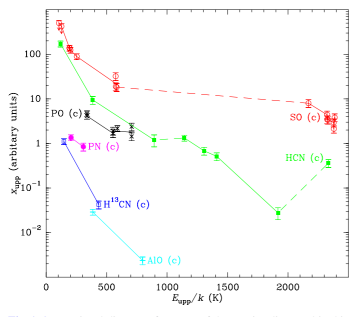


Fig. A.1. Rotational diagrams for some of the species discussed in this appendix: central component (c) only. Upper limits are marked with downward arrows. Solid lines join upper levels in the same vibrational state, while dashed lines join upper levels in different vibrational states.

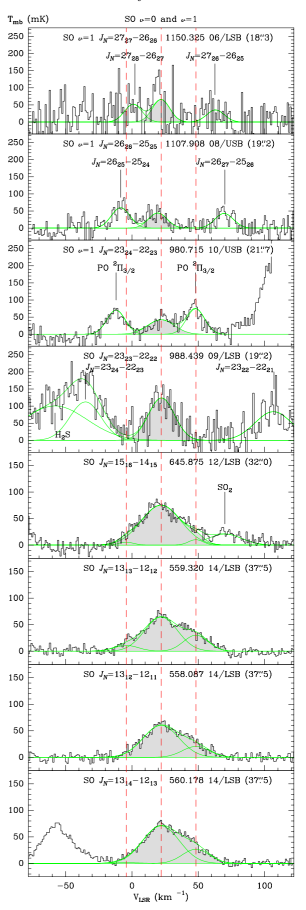


Fig. A.2. Same as Fig. 3 for the detected lines of SO. The v = 0 J = 23-22 and v = 1 J = 26-25 and 27-26 triplets are plotted in one panel each (*fourth*, *second*, and *first from top*).

Table A.1. Spectral-line results for sulphur monoxide, phosphorous monoxide, aluminium monoxide, silicon sulphide, hydrogen cyanide, and phosphorous nitride.

Species and	Rotational	$E_{\rm upp}/k$	Rest freq.	Setting &	rms [‡]	Peak	Area	Veloc. range	
elec./vibr. state	quantum nums.	(K)	(GHz)	sideband	(mK)	(mK)	$(K \text{ km s}^{-1})$	LSR (km s ⁻¹)	Comments
$\frac{\text{SO } v = 0}{\text{SO } v = 0}$	$J_N = 9_9 - 8_9$	106	568.741	14 USB	6.6 (1.05)	≤20	≲0.13	Lore (kins)	Comments
30 0 = 0	$J_N = 99-89$ $J_N = 10_{10}-9_{10}$	127	609.960	13 LSB	5.4 (0.99)	≈20 ≲16	≈0.13 ≲0.10		
	$J_N = 13_{14} - 12_{13}$	193	560.178	14 LSB	6.6 (1.08)	82	3.08	[-13;+75]	
	$J_N = 13_{12} - 12_{13}$ $J_N = 13_{12} - 12_{11}$	194	558.087	14 LSB	6.6 (1.08)	69	2.54	$[-15, \pm 75]$ $[-15, \pm 75]$	
	$J_N = 13_{12} - 12_{11}$ $J_N = 13_{13} - 12_{12}$	201	559.320	14 LSB	6.6 (1.08)	77	2.92	[-15;+75]	
	$J_N = 15_{13} \cdot 12_{12}$ $J_N = 15_{16} - 14_{15}$	253	645.875	12 LSB	7.8 (0.93)	82	3.15	g-fitted	
	$J_N = 23_{24} - 22_{23}$	575	988.616	09 LSB	33.1 (1.06)	156	5.21	g-fitted	
	$J_N = 23_{22} - 22_{21}$	576	988.166	09 LSB	33.1 (1.06)	96	2.82	g-fitted	
	$J_N = 23_{23} - 22_{22}$	583	988.439	09 LSB	33.1 (1.06)	129	2.84	g-fitted	
v = 1	$J_N = 23_{24} - 22_{23}$	2169	980.715	10 USB	19.8 (1.07)	58	1.26	g-fitted	
0 1	$J_N = 26_{27} - 25_{26}$	2169	1108.012	08 USB	23.1 (1.08)	60	1.09	g-fitted	
	$J_N = 26_{25} - 25_{24}$	2324	1107.723	08 USB	23.1 (1.08)	44	0.79	g-fitted	
	$J_N = 26_{26} - 25_{25}$	2331	1107.908	08 USB	23.1 (1.08)	45	0.81	g-fitted	
	$J_N = 27_{28} - 26_{27}$	2378	1150.409	06 LSB	59.5 (1.04)	52	0.86	g-fitted	
	$J_N = 27_{26} - 26_{25}$	2379	1150.165	06 LSB	59.5 (1.04)	52	0.86	g-fitted	
	$J_N = 27_{27} - 26_{26}$	2386	1150.325	06 LSB	59.5 (1.04)	65	1.08	g-fitted	
	0 IV = 1 27 = 0 20				(2101)			8	
PO $^{2}\Pi_{3/2} v = 0$	$J = 35/2 - 33/2 \ l = e$	338	763.121	11 USB	13.5 (0.98)	36	1.04	g-fitted	
- 5/2	$J = 35/2 - 33/2 \ l = f$	338	763.310	11 USB	13.5 (0.98)	27	0.92	g-fitted	
	$J = 45/2 - 43/2 \ l = e$	553	980.644	10 USB	19.8 (1.07)	78	1.06	g-fitted	
	J = 45/2 - 43/2 $l = f$	553	980.834	10 USB	19.8 (1.07)	68	1.21	g-fitted	
$^{2}\Pi_{1/2} v = 0$	J = 31/2 - 29/2	589	680.730	17 LSB	13.4 (1.10)	15	0.62	g-fitted	
$^{2}\Pi_{3/2} v = 0$	$J = 51/2 - 49/2 \ l = e$	707	1110.970	08 USB	23.1 (1.08)	71	2.38	g-fitted	
3/2	$J = 51/2 - 49/2 \ l = f$	707	1111.160	07 USB	22.0 (1.08)	75	1.44	g-fitted	
	, ,							C	
AlO $v = 0$	N = 20-19	386	764.603	11 USB	13.5 (0.98)	73	2.63	g-fitted	
	N = 29-28	798	1106.980	08 USB	23.1 (1.08)	84	1.00	g-fitted	tent. detec.
SiS v = 0	J = 61-60	1168	1102.029	07 LSB	22.0 (1.09)	106	2.59	g-fitted	
v = 1	J = 55-54	2404	989.685	09 LSB	33.1 (1.06)	≲99	≲0.77		
	J = 61-60	2707	1096.635	08 LSB	23.1 (1.10)	44	1.19	g-fitted	tent. assig.
	J = 62-61	2760	1114.431	07 USB	22.0 (1.08)	63	1.61	g-fitted	tent. assig.
	J = 64-63	2870	1149.994	06 LSB	59.5 (1.04)	≲179	≲1.38		
		440	<	40.7700	- 4 (O O -)				
HCN v = 0	J = 7-6	119	620.304	13 USB	5.4 (0.97)	261	12.1	g-fitted	
	J = 13-12	386	1151.452	06 LSB	59.5 (1.04)	266	8.11	g-fitted	
	J = 20-19	893	1769.877	19 USB	77.8 (1.02)	145	2.85	g-fitted	tent. detec.
$v_2 = 1$	J = 7 - 6 l = 1e	1143	620.225	13 USB	5.4 (0.97)	23	0.85	g-fitted	
	$J = 11-10 \ l = 1f$	1306	979.285	10 USB	19.8 (1.07)	64	0.90	g-fitted	
	$J = 13-12 \ l = 1e$	1411	1151.297	06 LSB	59.5 (1.04)	119	4.14	g-fitted	
	$J = 20-19 \ l = 1e$	1917	1769.603	19 USB	77.8 (1.02)	65	1.22	g-fitted	
$v_2 = 2$	$J = 11-10 \ l = 2e$	2334	979.554	10 USB	19.8 (1.07)	31	0.45	g-fitted	
			<00 	4= 7700	40.4.4.00	•	4.40		
$H^{13}CN v = 0$	J = 8-7	147	690.552	17 USB	13.4 (1.08)	38	1.19	g-fitted	
	J = 14-13	435	1207.853	05 USB	88.8 (0.99)	120			
DM 0	N 12 12	207	(10.500	10 1 00	5 4 (0 00°	22	0.44	C 1	
PN v = 0	N = 13-12	205	610.588	13 LSB	5.4 (0.99)	22	0.44	g-fitted	
	N = 16-15	307	751.311	11 LSB	13.5 (1.00)	39	0.64	g-fitted	
v = 1	N = 12-11	2078	559.665	14 LSB	6.6 (1.08)	≲20 <170	≤0.16		
	N = 25-24	2631	1164.391	06 USB	59.5 (1.03)	≲179	≲1.39		

Notes. (‡) For the spectral resolution in km s⁻¹ given in parentheses.

similar rotational lines in the two vibrationally excited states, we derive vibrational excitation temperatures of about 2000 K. This result points out that the excitation of the vibrationally excited states is mostly due to radiative pumping for SO as well.

The profiles of the well-detected lines are triangular or Gaussian-like, see Fig. A.2. The FWHM values obtained for the central component show a clear decreasing trend with increasing excitation energy of the levels, ranging from $35 \,\mathrm{km}\,\mathrm{s}^{-1}$ for the v=0 J=13-12 lines to $20\,\mathrm{km}\,\mathrm{s}^{-1}$ for the v=0 J=23-22

and all the v = 1 detected lines (though in these latter cases the S/N is poor and the results of the fittings are less accurate).

A.2. Sulphur dioxide (SO₂)

We tentatively detected $33~SO_2$ lines in our spectra, see Table A.2. Almost all these lines are only slightly stronger that the detection limit, therefore their parameters are rather uncertain. However, we verified that our identifications are compatible

Table A.2. Spectral-line results for sulphur dioxide.

Species and	Rotational	$E_{\rm upp}/k$	Rest freq.	Setting &	rms [‡]	Peak	Area	Comments
elec./vibr. state	quantum nums.	(K)	(GHz)	sideband	(mK)	(mK)	(K km s ⁻¹)	
$SO_2 v = 0$	$J_{K_{\rm a},K_{\rm c}} = 10_{6,4} - 9_{5,5}$	139	753.060	11 LSB	13.6 (1.00)	34	1.50	
	$J_{K_{\rm a},K_{\rm c}} = 16_{5,11} - 15_{4,12}$	186	765.321	11 USB	13.5 (0.98)	24	1.05	
	$J_{K_{\rm a},K_{\rm c}} = 8_{8,0} - 8_{7,1}$	190	763.982	11 USB	13.5 (0.98)		(a)	
	$J_{K_{\rm a},K_{\rm c}} = 9_{8,2} - 9_{7,3}$	198	763.992	11 USB	13.5 (0.98)		(a)	
	$J_{K_{\rm a},K_{\rm c}} = 10_{8,2} - 10_{7,3}$	207	763.999	11 USB	13.5 (0.98)		(a)	
	$J_{K_{\rm a},K_{\rm c}} = 16_{6,10} - 16_{5,11}$	213	560.614	14 LSB	6.6 (1.08)	11	0.79	
	$J_{K_a,K_c} = 11_{8,4} - 11_{7,5}$	217	764.000	11 USB	13.5 (0.98)		(a)	
	$J_{K_{\rm a},K_{\rm c}} = 12_{8,4} - 12_{7,5}$	229	763.993	11 USB	13.5 (0.98)		(a)	
	$J_{K_a,K_c} = 13_{8,6} - 13_{7,7}$	240	763.976	11 USB	13.5 (0.98)		(a)	
	$J_{K_a,K_c} = 16_{7,9} - 15_{6,10}$	245	969.224	10 LSB	19.8 (1.08)	30	1.26	
	$J_{K_a,K_c} = 18_{6,12} - 18_{5,13}$	246	559.882	14 LSB	6.6 (1.08)	8	0.31	
	$J_{K_a,K_c} = 14_{8,6} - 14_{7,7}$	253	763.948	11 USB	13.5 (0.98)		(a)	
	$J_{K_a,K_c} = 17_{7,11} - 16_{6,10}$	261	988.302	09 LSB	33.1 (1.06)	23	045	
	$J_{K_a,K_c} = 19_{6,14} - 19_{5,15}$	263	559.500	14 LSB	6.6 (1.08)	8	0.18	
	$J_{K_a,K_c} = 15_{8,8} - 15_{7,9}$	267	763.906	11 USB	13.5 (0.98)		(a)	
	$J_{K_a,K_c} = 20_{6,14} - 20_{5,15}$	281	558.813	14 LSB	6.6 (1.08)	8	0.18	
	$J_{K_a,K_c} = 16_{8,8} - 16_{7,9}$	282	763.847	11 USB	13.5 (0.98)		(a)	
	$J_{K_a,K_c} = 17_{8,10} - 17_{7,11}$	297	763.768	11 USB	13.5 (0.98)		(a)	
	$J_{K_a,K_c} = 21_{6,16} - 21_{5,17}$	301	558.391	14 LSB	6.6 (1.08)	9	0.30	
	$J_{K_a,K_c} = 20_{7,13} - 20_{6,14}$	313	661.668	12 USB	7.8 (0.91)	13	0.38	
	$J_{K_a,K_c} = 18_{8,10} - 18_{7,11}$	314	763.668	11 USB	13.5 (0.98)		(a)	
	$J_{K_a,K_c} = 22_{6,16} - 22_{5,17}$	321	557.283	14 LSB	6.6 (1.08)	16	0.52	
	$J_{K_a,K_c} = 22_{7,15} - 22_{6,16}$	352	660.918	12 USB	7.8 (0.91)	21	0.30	
	$J_{K_a,K_c} = 23_{7,17} - 22_{6,16}$	374	1102.115	07 LSB	22.0 (1.09)	25	1.05	
	$J_{K_a,K_c} = 23_{7,17} - 23_{6,18}$	374	660.473	12 USB	7.8 (0.91)	12	0.15	
	$J_{K_a,K_c} = 28_{4,24} - 27_{3,25}$	416	968.697	10 LSB	19.8 (1.08)	12	0.37	
	$J_{K_a,K_c} = 25_{7,19} - 25_{6,20}$	419	659.338	12 USB	7.8 (0.91)	6	0.18	
	$J_{K_a,K_c} = 26_{7,19} - 26_{6,20}$	443	658.542	12 USB	7.8 (0.91)	10	0.40	
	$J_{K_a,K_c} = 32_{0,32} - 31_{1,31}$	459	571.553	14 USB	6.6 (1.05)	10	0.38	
	$J_{K_a,K_c} = 32_{2,30} - 31_{3,29}$	504	571.533	14 USB	6.6 (1.05)	13	0.50	
	$J_{K_a,K_c} = 34_{1,33} - 33_{2,32}$	543	622.827	13 USB	5.4 (0.97)	12	0.32	
	$J_{K_a,K_c} = 37_{1,37} - 36_{0,36}$	609	659.421	12 USB	7.8 (0.91)	16	0.95	
	$J_{K_a,K_c} = 36_{2,34} - 35_{3,33}$	630	661.511	12 USB	7.8 (0.91)	14	0.44	
$v_2 = 1$	$J_{K_a,K_c} = 56_{2,34} - 55_{3,33}$ $J_{K_a,K_c} = 6_{5,1} - 6_{2,4}$	843	1096.851	08 LSB	23.1 (1.10)	93	1.77	maser?

Notes. (‡) For the spectral resolution in km s⁻¹ given in parentheses. (a) Lines $J_{K_a,K_c} = J_{8,J-8} - J_{7,J-7}$ for even values of J between 8 and 18, and $J_{K_a,K_c} = J_{8,J-7} - J_{7,J-6}$ for odd values of J between 9 and 17, appear to be severely blended. They were fitted as a group, resulting in a integrated area of 1.20 K km s⁻¹ and peak values between 10 and 20 mK.

Table A.3. Unassigned spectral features.

Rest freq. (GHz) LSB/USB	Setting name	rms [‡] (mK)	Peak (mK)	Area (K km s ⁻¹)	Possible	assignations and comments
559.675 (25)/569.555 (25)	14	6.6 (1.08)	22	0.40	29 SiS $J = 32-31$	(569.528 GHz USB)
607.055 (25)/622.774 (25)	13	5.4 (0.99)	16	0.77	29 SiS $J = 35-34$	(622.760 GHz USB); blended with SO ₂
751.664 (25)/766.264 (25)	11	13.6 (1.00)	60	1.40	33 SO $J_N = 18_{18} - 17_{17}$	(766.263 GHz USB)

Notes. $^{(\ddagger)}$ For the spectral resolution in km s⁻¹ given in parentheses.

with an unresolved, optically thin emission with a rotational excitation of about 250–300 K. Under these assumptions we checked that only the lines listed in the table are visible in the spectra. Other lines may also be present, but they happen to overlap much stronger lines of other species and therefore cannot be analysed. The detected transitions have high line-strengths and $E_{\rm upp}/k \lesssim 630$ K.

In addition to these v = 0 lines, we identified another spectral feature that we tentatively assign to SO₂. For a systemic velocity of $22 \,\mathrm{km \, s^{-1}}$ and using a single-component Gaussian fitting, we derived rest frequencies of 1,096.860 GHz (if from the LSB) and 1,109.215 GHz (if from the USB) for the spectral feature that appears to the left of the $v = 0.3_{1.2} - 3_{0.3}$ line of ortho-H₂O in

setting 08 (Fig. B.1). The spectral feature is clear (Fig. A.3), with a $T_{\rm mb}$ value of about 93 mK. This line is also tentatively detected in the spectra of other O-rich sources in HIFISTARS, such as o Cet, W Hya, and R Dor, and it is clearly absent from the C-rich envelope around IRC+10216, and in the O-rich envelopes around IK Tau, IRC+10011, and R Cas (HIFISTARS team, priv. comm.). These observations also suggest that the spectral feature arises from the LSB. We searched for lines in a 50 MHz (13.5 km s⁻¹) interval around the two possible frequencies in the Splatalogue catalogue (http://www.splatalogue.net/) and in the private line catalogues of some of the members of the HIFISTARS team. According to this, the most likely candidate for our detection is the $v_2 = 1$ $J_{K_8,K_6} = 65,1-62,4$ of SO₂

Table A.4. Triple-Gaussian fitting results for CO, H₂O, NH₃, SiO, SO, and SO₂.

						777.7	
Species and	Rotational	$E_{ m upp}^{\dagger}$	Central co Peak	mponent FWHM	HE\ Blue peak	/W compon FWHM	ents Red Peak
elec./vibr. state	quantum nums.	(K)	$T_{\rm mb}$ (mK)	$(km s^{-1})$	$T_{\rm mb}$ (mK)	(km s^{-1})	$T_{\rm mb}$ (mK)
$\frac{12}{\text{CO } v = 0}$	J = 6-5	116	1091	38	292	19	366
	J = 10-9	304	1499	37	366	22	452
	J = 16-15	752	1296	31	205	17	264
13 CO $v = 0$	1 (5	111	92	40	40	10	20
v = 0	J = 6-5 $J = 9-8$	111 238	82 131	49 47	48	12	38
	J = 10-9	291	173	39			
	J = 16-15	719	257	19			
ortho- $H_2O v = 0$	$J_{K_{\rm a},K_{\rm c}} = 1_{1,0} - 1_{0,1}$	27	1143	43	-253	17	92
$01110-11_200 = 0$	$J_{K_{\rm a},K_{\rm c}} = 1_{1,0} - 1_{0,1}$ $J_{K_{\rm a},K_{\rm c}} = 3_{0,3} - 2_{1,2}$	163	4664	43	-255 -1514	17	538
	$J_{K_a,K_c} = 3_{1,2} - 3_{0,3}$	215	1809	45	-268	21	232
	$J_{K_a,K_c} = 3_{1,2} - 2_{2,1}$	215	2919	46	-425	13	310
	$J_{K_a,K_c} = 3_{2,1} - 3_{1,2}$	271	2009	41	-25	25	364
	$J_{K_{\rm a},K_{\rm c}} = 5_{3,2} - 5_{2,3}$	698	1802	31			
$v_2 = 1$	$J_{K_a,K_c} = 2_{1,2} - 1_{0,1}$	2379	1007	32			
	$J_{K_{a},K_{c}} = 8_{2,7} - 8_{3,4}$	3556	46	16			
para- $H_2O v = 0$	$J_{K_{\rm a},K_{\rm c}} = 1_{1,1} - 0_{0,0}$	53	3140	43	-804	18	373
	$J_{K_a,K_c} = 2_{0,2} - 1_{1,1}$	101	3317	40	-469	12	287
	$J_{K_a,K_c} = 2_{1,1} - 2_{0,2}$	137	1570	45	-101	23	267
	$J_{K_a,K_c} = 4_{2,2} - 4_{1,3}$ $J_{K_a,K_c} = 6_{3,3} - 6_{2,4}$	454 952	1183 1011	50 31		16	167
$v_2 = 1$	$J_{K_a,K_c} = 0_{3,3} - 0_{2,4}$ $J_{K_a,K_c} = 1_{1,1} - 0_{0,0}$	2352	244	28			
$v_2 = 1$ $v_2 = 2$	$J_{K_a,K_c} = 1_{1,1} - 3_{0,0}$ $J_{K_a,K_c} = 2_{1,1} - 2_{0,2}$	4684	114	27			
$v_3 = 1$	$J_{K_a,K_c} = 6_{3,3} - 6_{2,4}$	6342	169	27			
ortho- H_2^{18} O $v = 0$	$J_{K_a,K_c} = 3_{1,2} - 3_{0,3}$	215	209	34			
para- $H_2^{18}O v = 0$	$J_{K_0,K_0} = 1_{1.1} - 0_{0.0}$	53	205	35	-62	19	47
	4, 0						
$ortho-H_2^{17}O\ v=0$	$J_{K_a,K_c} = 3_{0,3} - 2_{1,2}$	162 215	350 232	26 35			
	$J_{K_{\rm a},K_{\rm c}} = 3_{1,2} - 3_{0,3}$	213	232	33			
para- $H_{2}^{17}O v = 0$	$J_{K_a,K_c} = 1_{1,1} - 0_{0,0}$	53	126	38	-49	11	-9
• Z	$J_{K_{\rm a},K_{\rm c}} = 2_{0,2} - 1_{1,1}$	101	72	35			
ortho-NH ₃ $v = 0$	I -1+ 0-	27	235	41	89	19	183
ortho-infig $v = 0$	$J_K = 1_0^+ - 0_0^-$ $J_K = 3_0^+ - 2_0^-$	170	233 1441	41	89	19	165
	$3K - 3_0 - 2_0$	170	1111	15			
para-NH ₃ $v = 0$	$J_K = 3^+_2 - 2^2$	127	450	44			
OH $^{2}\Pi_{1/2} v = 0$	J = 3/2 - 1/2	270	1695	40	122	19	641
28 SiO $v = 0$	J = 14-13	219	369	32	41	25	65
510 0	J = 16-15	283	378	28	43	23	90
29.6.0	. 10 10	107	110	4.4	12	25	20
29 SiO $v = 0$	J = 13-12 J = 26-25	187 722	118 203	41 32	12	25	20
	J - 20-23	122	203	34			
30 SiO $v = 0$	J = 26-25	713	193	23			
	J = 42-41	1832	153	22			
SO $v = 0$	$J_N = 13_{14} - 12_{13}$	193	70	32	2	24	26
30 v = 0	$J_N = 13_{14} - 12_{13}$ $J_N = 13_{12} - 12_{11}$	193	60	31	0	24	20
	$J_N = 13_{12} \cdot 12_{11}$ $J_N = 13_{13} - 12_{12}$	201	64	29	10	21	30
	$J_N = 15_{16} - 14_{15}$	253	75	35	5	13	10
$SO_2 v = 0$	$J_{K_{\rm a},K_{\rm c}} = 16_{5,11} - 15_{4,12}$	186	16	29	-4	14	19
$SO_2 v = 0$							

Notes. (†) Since the ortho and para spin isomer variants of both water and ammonia behave as different chemical species, for all ortho transition E_{upp}/k values are relative to the corresponding lowest ortho energy level, while for the para transitions they are given w.r.t. the corresponding lowest para energy level.

at $1096.851\,\mathrm{GHz}$. This line is only at $10\,\mathrm{MHz}$ ($2.7\,\mathrm{km\,s^{-1}}$) from the LSB value derived from the fit. Indeed, this is the only catalogued line in these frequency ranges from a species previously detected in VY CMa. Nevertheless, the identification of the line is tentative because the 65,1-62,4 transition has a very low linestrength, it is a $\Delta K = 3$ transition with $\log_{10}(A_{ul}) = -6.85$, but the detected spectral feature is relatively intense. In addition, it is a vibrationally excited line, its upper level being placed at more than 800 K above the ground. If plotted in a rotational diagram for SO₂ lines, we observe that the detected intensity is five orders of magnitude higher than expected for LTE conditions. We can only explain this result if we assume that our line shows a strong maser effect. The FWHM of the line is 14 km s⁻¹, a value similar to that of other lines arising from the inner envelope, such as those of NaCl and other vibrationally excited lines of SO₂ (Ziurys et al. 2007; Tenenbaum & Ziurys 2010; Tenenbaum et al. 2010a). The line is slightly blue-shifted w.r.t. the systemic velocity (as is also the case for o Cet), which could be explained if the maser is due to an IR line overlap.

A.3. Other oxides: phosphorous monoxide and aluminium monoxide (PO and AIO)

We detected several lines of PO, both from the ${}^{2}\Pi_{1/2}$ and ${}^{2}\Pi_{3/2}$ electronic states. The ${}^{2}\Pi_{3/2}$ transitions always come in *paired* features, corresponding to the l = e and l = f Λ -doubling components, while the ${}^{2}\Pi_{1/2}$ always show *single features* because we cannot separate the Λ -doubling splitting in this case⁷. We clearly detected the ${}^{2}\Pi_{3/2}$ J = 35/2-33/2, 45/2-43/2, and 51/2-49/2pairs, with upper state energies at 338, 553, and 707 K above the ground, respectively. Another two ${}^{2}\Pi_{3/2}$ transitions, the J=79/2-77/2 and 83/2-81/2 with E_{upp}/k of 1700 and 2100 K, were also observed, but since they are blended with other stronger lines, we cannot conclusively determine their presence. For the $^{2}\Pi_{1/2}$ transitions, we only detected the J = 31/2-29/2 at 589 K above the ground. The J = 53/2-51/2 and 55/2-53/2 lines, at $E_{\rm upp}/k \sim 1100\,{\rm K}$, are also within the band, but again they are blended with much stronger emission from other species. In all cases the lines only show the central main component, see Fig. A.4. The rotational diagram suggests an excitation temperature of 400 K (Fig. A.1).

We also detected two lines of AlO: the N=20–19 and 29–28 transitions with $E_{\rm upp}/k$ of 386 and 798 K. The N=45–44 and 49–48 transitions were also observed, but we did not detect them, the upper limits not being significant. The N=29–28 line is detected at the edge of the band in the setting 08 observation, and should be considered as tentative (see Fig. A.3). Because of this, the estimate of the excitation temperature from the rotational diagram is very uncertain; we derive a value of 160 K, a rather low figure compared with the 230 K found by Tenenbaum & Ziurys (2009), but still compatible in view of the uncertainties. The N=20–19 line is clearly detected, the profile just consisting in the central component.

A.4. Silicon sulphide (SiS)

We detected only one v=0 rotational line of SiS, the J=61–60 at 1102.029 GHz. The J=104–103 at 1861.245 GHz is observed in setting 01, overlapping the vibrationally excited line of water at 1868.783 GHz, but it is very unlikely that we could have

⁷ We only consider the strongest hyperfine components, for J = j - (j-1) the F = (j+1/2) - (j-1/2) and F = (j-1/2) - (j-3/2) components, which are not resolved in VY CMa for both $^2\Pi_{1/2}$ and $^2\Pi_{3/2}$ electronic states.

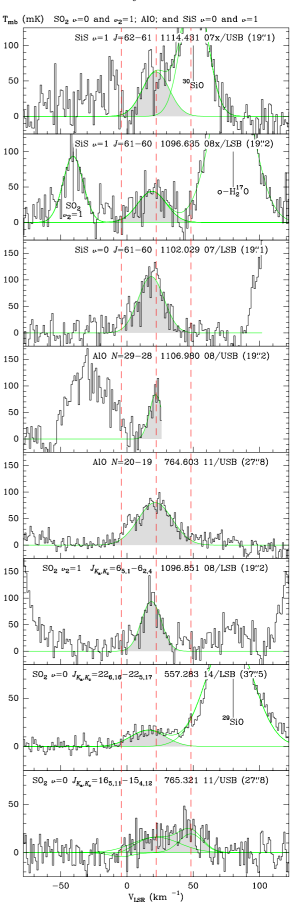


Fig. A.3. Same as Fig. 3 for the detected lines of AlO, and SiS, and some lines of SO₂. The assignation of the $v_2 = 1$ $J_{K_a,K_c} = 6_{5,1}-6_{2,4}$ line of SO₂ is tentative.

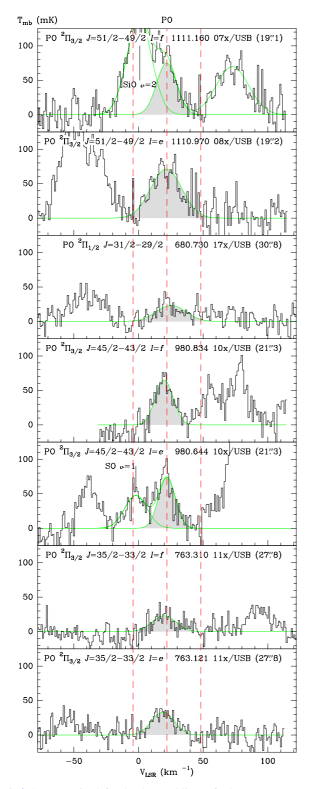


Fig. A.4. Same as Fig. 3 for the detected lines of PO.

detected the SiS line since it has an upper state energy of 4723 K above the ground. No other v=0 lines of SiS were observed, but several other rotational transitions of vibrationally excited states were covered by our survey. We did not detect any lines from states v=2 and higher, but we tentatively detected two lines from the v=1 levels: the J=61–60 at 1096.635 GHz, and the J=62–61 at 1114.431 GHz (this line has been also detected in OH 26.5+0.6, see Justanont et al. 2012). In addition, we obtained significant upper limits for the v=1 J=55–54

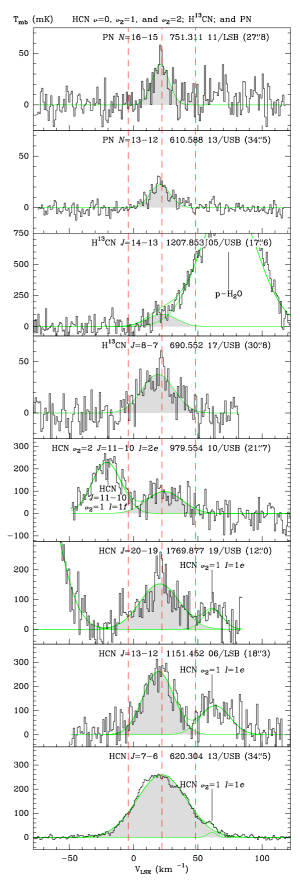


Fig. A.5. Same as Fig. 3 for the detected lines of HCN, H¹³CN, and PN.

and J = 64–63 transitions of SiS. The values obtained for the v = 1 J = 62–61 transition may be affected by the blending of

this line with the v=0 J=26-25 line of 30 SiO entering from the other side-band of the receiver, but they are very similar to the adjacent J=61-60 line, which presents a clear detection. These two lines show just about half the strength of the v=0 J=61-60, even though they have an excitation energy of 2700 K, that is, 1600 K above the v=0 line. This would suggest a relatively high excitation temperature between the v=0 and v=1 ladders due to the IR pumping of the v=1 levels. However, the non-detection of the v=1 J=55-54 and J=64-63 transitions suggests that the strength of the two detected v=1 lines may also be enhanced by an IR line overlap.

A.5. Hydrogen cyanide (HCN)

We detected all the observed rotational transitions of HCN from the ground vibrational state: the J=7-6, 13-12, and 20-19 lines, with $E_{\rm upp}/k$ of 119, 387, and 893 K. We also detected, although tentatively in some cases, all the HCN rotational lines within our bands from the vibrationally excited states $v_2=1$ and $v_2=2$: the J=7-6, 13-12, and 20-19 l=1e transitions of the $v_2=1$ state, and the $v_2=1$ J=11-10 l=1f and $v_2=2$ J=11-10 l=2e lines. Two other lines from the $v_1=1$ and $v_3=1$ states were also observed, but were not detected.

The profiles of all these lines show only the central component. Including the data for vibrationally excited transitions, we found a noisy trend when plotting the FWHMs of the lines against the excitation energy of the upper levels. Lines with relatively low excitation energy, for instance, the v=0 J=7-6, show FWHM values higher than $40\,\mathrm{km\,s^{-1}}$, while values lower than $20\,\mathrm{km\,s^{-1}}$ are found for transitions with $\geq E_{\mathrm{upp}}/k$ of 1500 K. This result suggests that the vibrationally excited lines of HCN arise from the inner regions of the envelope where the final expansion velocity has not been attained yet.

The rotational diagram for HCN shows excitation temperatures between 90 and 250 K for the v=0 lines, and of about 200 K for the $v_2=1$ lines (Fig. A.1). If we compare similar rotational transitions in different vibrational states, we obtain excitation temperatures of 210–350 K between the v=0 and $v_2=1$ ladders (and of 1500 K between the $v_2=1$ and $v_2=2$ ladders, but this a very tentative result). This would suggest, although less clearly than in the cases of H_2O , SiO, and SO, that the excitation of the $v_2=1$ and 2 levels in HCN is also dominated by the IR radiation pumping.

We also systematically searched for lines of $H^{13}CN$ in our spectra, both from the ground and the $v_2 = 1$ states. We detected the v = 0 J = 8-7 and (tentatively) the J = 14-13 lines, with $E_{\rm upp}/k$ of 145 and 435, respectively. No vibrationally excited lines were detected in our data, which include the frequencies of the $v_2 = 1$ J = 7-6 and J = 8-7 transitions. Because of the severe blending of the v = 0 J = 14-13 line (it is at the edge of the strong $4_{2,2}-4_{1,3}$ line of para- H_2O), the results from the rotational diagram of $H^{13}CN$ can only be considered as tentative. We derive

an excitation temperature of about 90 K, very similar to the value for HCN between similar transitions, suggesting that the lines from both species are optically thin.

A.6. Phosphorous nitride (PN)

In addition to NH₃ and HCN, the other N-bearing molecule identified in VY CMa in our HIFI spectra is PN. We detected the only two (ground-vibrational state) rotational lines covered by our observations, the N=13–12 and 16–15 multiplets (six unresolved components within 4 MHz), with upper level energies of 205 and 307 K above the ground. The lines are relatively narrow, with FWHPs of less than $20 \,\mathrm{km \, s^{-1}}$, suggesting that the emission originates in the inner shells of the envelope. The excitation temperature derived from the rotational diagram is about 220 K. Two v=1 multiplets, the N=12–11 and 25–24 at 559.7 and 1164.4 GHz, were also observed but were not detected, the limits are irrelevant.

A.7. Unassigned spectral features

There are three spectral features in our spectra that we were unable to unambiguously assign to any molecular line. They are listed in Table A.3 and are marked "U" in green in Fig. B.1. For the two in settings 13 and 14, 29 SiS is a good candidate. For the other U-line in setting 11, the most likely candidate is 33 SO ($J_N = 18_{18}-17_{17}$ at 766.263 GHz). In other HIFISTARS observations, the U-features in settings 13 and 14 are also detected in C-rich envelopes, whereas that in setting 11 seems to be present in other O-rich sources, which would support our conjectures.

Appendix B: Full Herschel/HIFI WBS spectra

Here we present the full-bandwidth results of our Herschel/HIFI WBS observations of VY CMa. In Fig. B.1 we show the results for LO frequencies of 1102.9 GHz and lower: these are, from bottom to top, settings 14, 13, 12, 17, 11, 10, 09, and 08. In Fig. B.2 we show the results for LO frequencies of 1106.9 GHz and higher: these are, from bottom to top, settings 07, 06, 05, 04, 03, 19, 16, and 01. The $T_{\rm mb}$ observed with HIFI/WBS is plotted vs. both LSB (lower x-axis) and USB (upper x-axis) restfrequency scales (in GHz), assuming an LSR systemic velocity of 22 km s⁻¹ for the source. The location of the detected lines from the LSB/USB is indicated by the red and blue arrows and labels. The point of the arrow marks the appropriate axis in each case. Unassigned features are labelled "U" in green, and arrows point to the two possible rest frequencies. We note that setting 16 was observed twice using a slightly different LSR velocity for the source; since we cannot average these two spectra simultaneously aligning the two (side-band) frequency scales, in Fig. B.2 we only show the result for OBSID 1342196570.



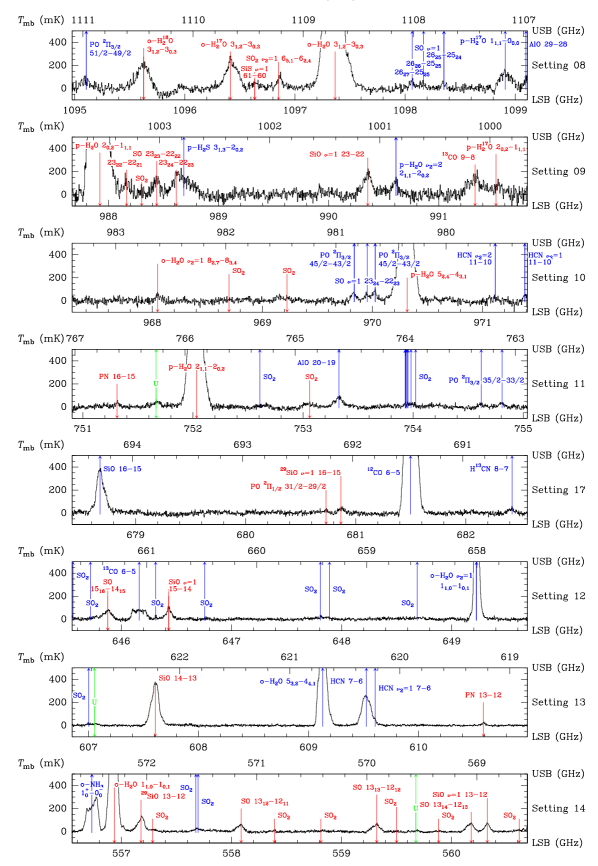


Fig. B.1. Full WBS bandwidth HIFI results for VY CMa for LO frequencies of 1102.9 GHz and lower. Quantum numbers for SO₂ transitions are not given for clarity.

J. Alcolea et al.: HIFISTARS Herschel/HIFI observations of VY Canis Majoris $T_{\rm mb}$ (mK) 1869 1868 USB (GHz) 2000 p-H₂O v₁=1 3_{2,2}-3_{1,3} 1500 1000 Setting 01 500 0 LSB (GHz) 1861 1862 1841 — USB (GHz) 1842 $T_{\rm mb}$ (mK) 1843 2000 OH $^{2}\Pi_{1/2}$ 3/2-1/2 ¹²CO 16-15 1500 1000 Setting 16 500 0 والمساري والمساوي LSB (GHz) 1834 1835 1836 $T_{\rm mb} \ ({\rm mK})1772$ 1771 1770 USB (GHz) 2000 HCN v₂=1 20-19 $0-NH_3 3_0^+-2_0^$ $p-NH_3 \ 3_1^--2_1^+$ 1500 $-NH_3 3_2^+ - 2_2^-$ 1000 Setting 19 ³⁰SiO 42-41 HCN 20-19 500 0 LSB (GHz) 1763 1764 $T_{\mathbf{mb}}$ (mK) 1762 1761 USB (GHz) 2000 ¹³CO 16-15 p-H₂O 6_{3,3}-6_{2,4} o-H₂O v₂=1 2_{1,2}-1_{0,1} 1500 1000 Setting 03 500 0 LSB (GHz) 1753 1754 1755 $T_{\rm mb}$ (mK) 1718 1717 USB (GHz) 2000 $o-H_2^{17}O \ 3_{0,3}-2_{1,2}$ 1500 $p-H_20$ $v_3=1$ $6_{3,3}-6_{2,4}$ 1000 Setting 04 500 LSB (GHz) 1710 1711 $T_{\rm mb}$ (mK) 1209 1208 1206 1205 1207 USB (GHz) p-H₂0 4_{2,2}-4_{1,3} $o-H_2S$ $3_{1,2}-2_{2,1}$ $-H_2O \nu_2 = 1 \ 1_{1,1} - 0_{0,0}$ SiO v=2 27-17 Setting 05 LSB (GHz) 1193 1194 1195 1196 1197 1165 1164 1163 1162 USB (GHz) ¹²CO 10 -H₂O 3_{2,1} SiO v=2 27-26 Si0 v=1 27-26 Setting 06 HCN vo=1 13-12 O Entricumental of many popular representative the residence of the party representative to the residence of the party residence of the reside LSB (GHz) 1151 1152 1153

1101 Fig. B.2. Full WBS bandwidth HIFI results for VY CMa for LO frequencies of 1106.9 GHz and higher. Quantum numbers for SO₂ transitions are not given for clarity.

1113

26-25 13CO 10-9

1114

1100

1112

1102

p-H₂¹⁸0 1_{1,1}-0_{0,0} SiS 61-

1111

1103

Si0 v = 2

USB (GHz)

Setting 07

LSB (GHz)

PO ²Π_{3/2} 51/2-49/2

1000

500

 $T_{\rm mb}~({
m mK})$

1000

500

 $T_{\rm mb}$ (mK) 1115

400

200

1099