Gas phase Elemental abundances in Molecular cloudS (GEMS)

II. On the quest for the sulphur reservoir in molecular clouds: the H₂S case

D. Navarro-Almaida¹, R. Le Gal², A. Fuente¹, P. Rivière-Marichalar¹, V. Wakelam³, S. Cazaux ^{4,5}, P. Caselli⁶,
J. C. Laas⁶, T. Alonso-Albi¹, J. C. Loison⁷, M. Gerin⁸, C. Kramer⁹, E. Roueff¹⁰, R. Bachiller¹, B. Commerçon¹¹,
R. Friesen¹², S. García-Burillo¹, J. R. Goicoechea¹³, B. M. Giuliano⁶, I. Jiménez-Serra¹⁴, J. M. Kirk¹⁵, V. Lattanzi⁶,
J. Malinen^{16,17}, N. Marcelino¹³, R. Martín-Domènech², G. M. Muñoz Caro¹⁴, J. Pineda⁶, B. Tercero¹

S. P. Treviño-Morales¹⁸, O. Roncero¹³, A. Hacar¹⁹, M. Tafalla¹, and D. Ward-Thompson¹⁵

(Affiliations can be found after the references)

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ABSTRACT

Context. Sulphur is one of the most abundant elements in the Universe. Surprisingly, sulphuretted molecules are not as abundant as expected in the interstellar medium and the identity of the main sulphur reservoir is still an open question.

Aims. Our goal is to investigate the H_2S chemistry in dark clouds, as this stable molecule is a potential sulphur reservoir.

Methods. Using millimeter observations of CS, SO, H_2S , and their isotopologues, we determine the physical conditions and H_2S abundances along the cores TMC 1-C, TMC 1-CP, and Barnard 1b. The gas-grain model NAUTILUS is used to model the sulphur chemistry and explore the impact of photo-desorption and chemical desorption on the H_2S abundance.

Results. Our modeling shows that chemical desorption is the main source of gas-phase H_2S in dark cores. The measured H_2S abundance can only be fitted if we assume that the chemical desorption rate decreases by more than a factor of 10 when $n_H > 2 \times 10^4$. This change in the desorption rate is consistent with the formation of thick H_2O and CO ice mantles on grain surfaces. The observed SO and H_2S abundances are in good agreement with our predictions adopting an undepleted value of the sulphur abundance. However, the CS abundance is overestimated by a factor of 5–10. Along the three cores, atomic S is predicted to be the main sulphur reservoir.

Conclusions. The gaseous H_2S abundance is well reproduced, assuming undepleted sulphur abundance and chemical desorption as the main source of H_2S . The behavior of the observed H_2S abundance suggests a changing desorption efficiency, which would probe the snowline in these cold cores. Our model, however, highly overestimates the observed gas-phase CS abundance. Given the uncertainty in the sulphur chemistry, we can only conclude that our data are consistent with a cosmic elemental S abundance with an uncertainty of a factor of 10.

Key words. astrochemistry – ISM: abundances – ISM: kinematics and dynamics – ISM: molecules – stars: formation – stars: low-mass

1. Introduction

Sulphur is one of the most abundant elements in the Universe $(S/H \sim 1.35 \times 10^{-5})$, Yamamoto 2017) and plays a crucial role in biological systems on Earth, so it is important to follow its chemical history in space (i.e., toward precursors of the Solar System). Surprisingly, sulphuretted molecules are not as abundant as expected in the interstellar medium (ISM). A few sulphur compounds have been detected in diffuse clouds, demonstrating that sulphur abundance in these low-density regions is close to the cosmic value (Neufeld et al. 2015). This is also the case towards the prototypical photodissociation region (PDR) in the Horsehead Nebula, where the sulfur abundance is found to be very close to the undepleted value observed in the diffuse ISM (Goicoechea et al. 2006), with an estimate of S/H = $(3.5 \pm 1.5) \times 10^{-6}$. A wide variety of S-bearing molecules, including the doubly sulphuretted molecule S₂H, were later detected in the HCO peak in this PDR (Fuente et al. 2017; Rivière-Marichalar et al. 2019). However, sulphur is thought to be depleted inside molecular clouds by

a factor of 1000 compared to its estimated cosmic abundance (Graedel et al. 1982; Agúndez & Wakelam 2013). The depletion of sulphur is observed not only in cold pre-stellar cores, but also in hot cores and corinos (Wakelam et al. 2004). We would expect that most of the sulphur is locked on the icy grain mantles in dense cores, but we should see almost all sulphur back to the gas phase in hot cores and strong shocks. However, even in the well-known Orion-KL hot core, where the icy grain mantles are expected to sublimate releasing the molecules to the gas phase, one needs to assume a sulphur depletion of a factor of ~ 10 to reproduce the observations (Esplugues et al. 2014; Crockett et al. 2014). Because of the high hydrogen abundances and the mobility of hydrogen in the ice matrix, sulphur atoms impinging in interstellar ice mantles are expected to form H₂S preferentially. To date, OCS is the only S-bearing molecule unambiguously detected in ice mantles because of its large band strength in the infrared (Geballe et al. 1985; Palumbo et al. 1995) and, tentatively, SO₂ (Boogert et al. 1997). The detection of solid H₂S (s-H₂S hereafter) is hampered by the strong overlap between the 2558 cm⁻¹ band with the methanol bands at

2530 and 2610 cm⁻¹. Only upper limits of s-H₂S abundance could be derived by Jiménez-Escobar & Muñoz Caro (2011), with values with respect to H₂O ice of 0.7 and 0.13% in W33A and IRAS183160602, respectively.

Sulphur-bearing species have been detected in several comets. Contrary to the interstellar medium (ISM), the majority of cometary detections of sulphur-bearing molecules belongs to H₂S and S₂ (Mumma & Charnley 2011). Towards the bright comet Hale Bopp, a greater diversity has been observed, including CS and SO (Boissier et al. 2007). The brighter comets C/2012 F6 (Lemmon) and C/2014 Q2 (Lovejoy) have also been shown to contain CS (Biver et al. 2016). Currently, some of the unique in-situ data are available from the Rosetta mission on comet 67P/Churyumov-Gerasimenko. With the Rosetta Orbiter Spectrometer for Ion and Neutral Analysis (ROSINA; Balsiger et al. 2007) onboard the orbiter, the coma has been shown to contain H₂S, atomic S, SO₂, SO, OCS, H₂CS, CS₂, S₂, and, tentatively, CS, as the mass spectrometer cannot distinguish the latter from CO₂ gases (Le Roy et al. 2015). In addition, the more complex molecules S₃, S₄, CH₃SH, and C₂H₆S have now been detected (Calmonte et al. 2016). Even with the large variety of S-species detected, H₂S remains as the most abundant, with an ice abundance of about 1.5% relative to H₂O, which is similar to the upper limit measured in the interstellar medium (Jiménez-Escobar & Muñoz Caro 2011).

Gas phase Elemental abundances in Molecular CloudS (GEMS) is an IRAM 30 m Large Program aimed at estimating the depletions of the most abundant elements (C, O, S, and N) in a selected set of prototypical star-forming filaments located in the molecular cloud complexes Taurus, Perseus, and Orion A. The first chemical study using GEMS data concluded that only 10% of the sulphur is in gas phase in the translucent part ($A_v \sim 3-10$ mag, Fuente et al. 2019, hereafter Paper I) of TMC 1. These conclusions were based on millimeter observations of CS, SO and HCS⁺. Here we investigate the chemistry of H₂S, a key piece in the sulphur chemistry, which is still poorly understood.

The formation of H_2S in the gas phase is challenging at the low temperatures prevailing in the interstellar medium. None of the species S, SH, S⁺, and SH⁺ can react exothermically with H₂ in a hydrogen atom abstraction reaction which greatly inhibits the formation of sulphur hydrides (Gerin et al. 2016). Alternatively, H₂S is thought to form on the grain surfaces where sulphur atoms impinging in interstellar ice mantles are hydrogenated, forming s-H₂S and potentially becoming the most important sulphur reservoir in dark cores (Vidal et al. 2017). Recent modeling and experimental studies suggest an active sulphur chemistry within the ices where s-H₂S might be converted in more complex compounds such as OCS or CH₃SH (e.g., Laas & Caselli 2019, El Akel et al., in prep.). Unfortunately, both the surface chemistry reaction rates and the desorption processes are not well constrained, which makes a reliable prediction of both solid and gas phase H₂S abundance difficult. This paper uses a subset of the complete GEMS database to investigate the H₂S abundance in dark cores TMC 1 and Barnard 1b as prototypes of molecular clouds embedded in low-mass and intermediate-mass star forming regions, respectively.

2. Source sample

2.1. TMC 1

The Taurus molecular cloud (TMC), at a distance of 140 pc (Elias 1978), is one of the closest molecular cloud complexes

and is considered an archetypal low-mass star forming region. It has been the target of several cloud evolution and star formation studies (see e.g., Goldsmith et al. 2008), being extensively mapped in CO (Cernicharo & Guelin 1987; Narayanan et al. 2008). The most massive molecular cloud in Taurus is the Heiles cloud 2 (HCL 2, Tóth et al. 2004), which hosts the well-known region TMC 1 (Fig. 1a). As one of the most extensively studied molecular filaments, TMC 1 was included in the IRAM 30 m Large Program GEMS. Fuente et al. (2019) modeled the chemistry of the translucent filament, deriving the gas phase elemental abundances (C/O ~ 0.8-1, S/H ~ $0.4-2.2 \times 10^{-6}$) and constraining the cosmic rays molecular hydrogen ionization rate to ~0.5-1.8 × 10⁻¹⁶ s⁻¹ in this moderate dense $(n_{\rm H_2} < 10^4 \text{ cm}^{-3})$ phase. Here, we will focus on the chemistry of the dense cores TMC 1-CP and TMC 1-C (see Fig. 1a). TMC 1-CP has been extensively observed at millimeter wavelengths and is often adopted as template of C-rich (C/O > 1) dense cores (Fehér et al. 2016; Gratier et al. 2016; Agúndez & Wakelam 2013). Less studied, TMC 1-C was proposed as an accreting starless core with high CO depletion by Schnee et al. (2007, 2010).

2.2. Barnard 1b

Barnard 1 is a young, intermediate-mass star forming cloud, embedded in the western sector of the 30 pc wide molecular cloud complex Perseus (see Fig. 1b). It hosts several dense cores in different evolutionary stages (Hatchell et al. 2005), out of which Barnard 1b is the youngest. The Barnard 1b region was mapped in many molecular tracers, such as CS, NH₃, ¹³CO (Bachiller et al. 1990), N₂H⁺ (Huang & Hirano 2013), H¹³CO⁺ (Hirano et al. 1999), or CH₃OH (Hiramatsu et al. 2010; Öberg et al. 2010). Interferometric observations of Barnard 1b revealed that this compact core hosts two young stellar objects (YSOs), B1b-N and B1b-S (Huang & Hirano 2013; Marcelino et al. 2018), and a third more evolved source, hereafter B1b-Spitzer, with deep absorption features from ices (Jørgensen et al. 2006). The three sources are deeply embedded in the surrounding protostellar envelope, that seems essentially unaffected by the inlaid sources, as shown by the large column density, $N(H_2) \sim 7.6 \times 10^{22}$ cm⁻² (Daniel et al. 2013) and cold kinetic temperature, $T_{\rm K} = 12$ K (Lis et al. 2002). From the chemical point of view, Barnard 1b has a rich chemistry. Indeed, many molecules were observed for the first time in this object, like HCNO (Marcelino et al. 2009) or CH₃O (Cernicharo et al. 2012). Additionally, Barnard 1b shows a high degree of deuterium fractionation and has been associated with first detections of multiple deuterated molecules, such as ND₃ (Lis et al. 2002) or D₂CS (Marcelino et al. 2005), consistent with the expected chemistry in a dense and cold core. Recently, Fuente et al. (2016) proposed that this core is characterized by a low depletion of sulphur, S/H~ 10^{-6} . They proposed that peculiar initial conditions due to the proximity of the bipolar outflows, a rapid collapse of the parent cloud, or the imprint of the two deeply embedded protostellar objects, might be the causes. Within GEMS, we have studied the nine-point cut depicted in Fig. 1b, with visual extinctions (extinctions along the line of sight) ranging from ~ 3 mag to ~ 76 mag.

3. Observations

This paper is based on a subset of the GEMS observations carried out with the IRAM 30 m telescope and the 40 m Yebes telescopes. The TMC 1 data used in this paper were already presented in Paper I, where we gave a detailed description of

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Fig. 1. *Left panel*: TMC 1 total visual extinction (the visual extinction along the line of sight) map based on *Herschel* dust emission maps from Jason Kirk (private communication). *Right panel*: visual extinction map of Barnard 1 from the opacity at 850 µm derived by Zari et al. (2016). In both maps the observed positions are in lines of constant declination, or cuts, where the red squares mark those positions in which only IRAM 30 m telescope data is obtained, and green squares mark positions with additional Yebes 40 m data.

the observational procedures used within the GEMS project. The observations towards Barnard 1b were carried out using the same observational strategy as in TMC 1 (see Paper I). We select three lines of constant declination, or cuts, through the three extinction peaks of TMC 1-C, TMC 1-CP, and Barnard 1b. We define the origin of each line as the point of highest extinction in that line. Figures 1a and b show the positions observed across the dense cores TMC 1-C, TMC 1-CP, and Barnard 1b. All the positions were observed in setups 1–4 (see Paper I) using the 30 m telescope. These observations were done using frequency-switching in order to optimize the detection sensitivity. In addition to the 30 m observations, the positions with $A_v < 20$ mag were also observed with 40 m Yebes telescope (setup 0). During the 40 m observations, the observing procedure was position-switching, the OFF-position being $\overrightarrow{RA}(J2000) =$ 03:36:39.571 Dec: 29:59:53.30 for Barnard 1b. This position was checked to be empty of emission in the observed bandwidths before the observations. In the following, line intensities are in main beam temperature, $T_{\rm MB}$, for all observations. Calibration errors are estimated to be $\sim 20\%$ for the IRAM 30 m telescope and ~30% for the Yebes 40 m telescope. In order to constrain the gas physical conditions and the sulphur chemistry, in this paper we use the observed lines of CS and SO, in addition to H₂S (see the list of lines in Table 1, and the corresponding spectra in Figs. A.1, A.2, and B.1).

4. Multi-transition study of CS and SO

A precise knowledge of the gas physical conditions is required to determine accurate molecular abundances. For TMC 1, we adopted the gas kinetic temperatures and molecular hydrogen densities derived in Paper I. These values are summarized in Table A.1, together with the CS column densities also derived in our previous work.

Table 1. Measured transitions.

Species	Transition	Frequency (MHz)	Beam (arcsec)
¹³ CS	$1 \rightarrow 0$	46247.563	42‴
¹³ CS	$2 \rightarrow 1$	92 494.308	27''
¹³ CS	$3 \rightarrow 2$	138 739.335	18″
C ³⁴ S	$1 \rightarrow 0$	48 206.941	42''
C ³⁴ S	$2 \rightarrow 1$	96 412.950	26''
C ³⁴ S	$3 \rightarrow 2$	144 617.101	17″
CS	$1 \rightarrow 0$	48 990.955	42''
CS	$2 \rightarrow 1$	97 980.953	25''
CS	$3 \rightarrow 2$	146 969.029	17"
$H_2^{34}S$	$1_{1,0} \rightarrow 1_{0,1}$	167 910.516	15″
H_2S	$1_{1,0} \rightarrow 1_{0,1}$	168 762.753	15"
SO	$2_2 \rightarrow 1_1$	86 093.96	29''
SO	$2_3 \rightarrow 1_2$	99 299.89	25''
SO	$3_2 \rightarrow 2_1$	109 252.18	23''
SO	$3_4 \rightarrow 2_3$	138 178.66	18″
SO	$4_4 \rightarrow 3_3$	172 181.42	14‴
³⁴ SO	$2_3 \rightarrow 1_2$	97 715.40	25''
³⁴ SO	$3_2 \rightarrow 2_1$	106 743.37	23''
³⁴ SO	$4_4 \rightarrow 3_3$	168 815.11	15''

We estimate the physical conditions towards Barnard 1b from a multi-transition analysis of CS and SO following the same procedure as in Paper I. We fit the intensities of the observed CS, C³⁴S, and ¹³CS J = 1 \rightarrow 0, 2 \rightarrow 1, and 3 \rightarrow 2 lines using the molecular excitation and radiative transfer code RADEX (van der Tak et al. 2007). During the fitting process, the isotopic ratios are fixed to ¹²C/¹³C = 60 and ³²S/³⁴S = 22.5 (Gratier et al. 2016), and we assume a beam filling factor of 1 for all transitions (the emission is more extended than the beam size). Then, we let T_k , n_{H_2} , and N(CS) vary as free parameters. The parameter space $(T_k, n_{H_2}, N(CS))$ is then explored following the Monte Carlo Markov chain (MCMC) methodology with a Bayesian inference approach, as described in Rivière-Marichalar et al. (2019). In particular, we use the *emcee* (Foreman-Mackey et al. 2013) implementation of the Invariant MCMC Ensemble sampler methods by Goodman & Weare (2010). While $n_{\rm H_2}$ and N(CS) are allowed to vary freely, we need to introduce a prior distribution to limit the gas kinetic temperatures to reasonable values in this cold region and, hence, break the temperaturedensity degeneracy that is usual in this kind of calculations. As estimated by Friesen et al. (2017), the gas kinetic temperature in a wide sample of molecular clouds, based on the NH_3 (1,1) and (2,2) inversion lines, is found to be systematically 1 or 2 K lower than that obtained from Hershel maps. This is indeed corroborated in Sect. 5. Thus, we set a Gaussian prior distribution with mean $\mu = T_d$ and $\sigma = 2$ K for the gas kinetic temperature. The collisional rate coefficients for the molecular excitation calculations that involve ¹³CS and C³⁴S isotopologues are those of CS. They include collisions with para and ortho-H₂ as reported by Denis-Alpizar et al. (2018), with a ortho-to-para ratio of 3, and He, taken from Lique et al. (2006). The gas temperature and the density derived for Barnard 1b from the multi-line fitting of CS and its isotopologues are shown in Table B.1.

A similar multi-transition analysis is carried out to derive SO column densities. In this case we have used the collisional coefficients derived by Lique & Spielfiedel (2007). The derived molecular gas densities and SO column densities are shown in Table B.2. In most positions, the densities derived from the SO fitting fully agree with those derived from CS. However, in the vicinity of the visual extinction peak ($A_v > 30$ mag), the molecular hydrogen densities derived from SO data are systematically higher than those derived with CS. Daniel et al. (2013) derived the density structure of the core by fitting the CSO 350 μ m continuum map and the IRAM 1.2 mm image with a power-law radial density profile, and assuming standard values of the dustto-gas mass ratio and dust opacity. The densities derived from the continuum images fitting are consistent with those derived from our SO observations and a factor 5-10 higher than those derived from CS. Therefore, we adopt the densities derived from SO for the inner Barnard 1b core. The lower densities derived from CS are very likely the consequence of a high depletion of this molecule in the densest part of the core.

It is interesting to compare the SO column density obtained in this paper towards Barnard 1b with the S¹⁸O column density derived by Fuente et al. (2016). Comparing both values, we obtain $N(SO)/N(S^{18}O) = 78 \pm 45$, much lower than the isotopic ratio, ¹⁶O/¹⁸O = 550. Recent chemical calculations (Loison et al. 2019) using the gas-grain chemical code Nautilus (Ruaud et al. 2016) in typical conditions for dark molecular clouds, showed that the isotopic fractionation might be important in the case of SO, which yields a ¹⁶O/¹⁸O ratio between 500 and 80. Following their predictions, the $N(SO)/N(S^{18}O)$ ratio is expected to vary with time with changes of more than a factor of 5. The ratio measured in Barnard 1b would be consistent with a chemical age of a few 0.1 Myr.

5. Gas kinetic temperatures from NH₃ data

The low densities found in the translucent phase, with $n_{\rm H}$ of the order of a few 10^3 cm⁻³, might cast some doubts on our assumption that gas and dust are close to thermal equilibrium. To check that this is indeed the case, we have independently

Table 2. Kinetic temperatures, $T_k(CS)$, in Kelvin, of several $A_v < 20$ mag positions in Barnard 1b, compared to the estimated rotation temperature $T_R(NH_3)$ and the corresponding kinetic temperature $T_k(NH_3)$ from the ammonia (1,1) and (2,2) inversion lines.

Offset (",")	$T_{\rm k}({\rm CS})$ (K)	$T_{\rm R}({\rm NH_3})$ (K)	$T_{\rm k}({\rm NH_3})$ (K)
(+80", 0")	13.2 ± 1.8	11.5 ± 1.5	$12.0 \pm 2.1 \\ 12.1 \pm 2.4 \\ 13.4 \pm 2.6$
(+110", 0")	14.4 ± 1.9	11.6 ± 1.8	
(+140", 0")	14.2 ± 1.0	12.7 ± 1.9	

derived the gas kinetic temperature using the NH₃ (1,1) and (2,2) inversion lines as observed with the 40 m Yebes telescope towards the positions with $A_v < 20$ mag in Barnard 1b. To derive the gas kinetic temperatures, first we follow the method described by Bachiller et al. (1987) to derive the rotation temperature $T_{\rm R}(\rm NH_3)$, and then we use the prescription which relates the gas kinetic temperature $T_k(NH_3)$ with $T_R(NH_3)$ found by Swift et al. (2005). Gas kinetic temperatures thus derived from ammonia data (Table 2) fully agree within the error with those derived from the CS multi-transition study (Table A.1). Previous measurements of the gas kinetic temperature (Lis et al. 2002) are in agreement with our values. Thus, we adopt the gas kinetic temperatures derived from CS for our further analysis. It should be noted that we have not observed ammonia lines towards the $A_{\rm v} > 20$ mag positions precluding an independent gas kinetic temperature estimate. However, taking into account the high molecular hydrogen densities measured in these regions, our assumption of gas and dust being close to thermal equilibrium is a plausible approximation. We have not detected the NH₃ lines towards the positions with $A_{\rm v}$ < 10 mag.

6. H₂S abundance

The gas physical conditions derived in previous sections are used to compute the H_2S column densities. First, we estimate the ortho- H_2S column density using RADEX with the ortho- H_2O collisional coefficients calculated by Dubernet et al. (2009), scaled to ortho- H_2S . The H_2S abundance is then calculated adopting an ortho-to-para ratio of 3. The resulting molecular abundances are listed in Tables A.1 and B.1, and shown in Figs. 2a–c.

In these plots, we try to find correlations between the H_2S abundance and physical quantities like visual extinction, gas temperature, and H density. Here, hydrogen nuclei number density is twice the $n_{\rm H_2}$ density obtained with the MCMC approach described earlier. As seen in such plots, the relationship between the H₂S abundance and the physical parameters is mostly monotonic. To assess the degree of correlation, we compute the Kendall's tau coefficient (Kendall 1938) in each case: 1 for perfect correlation, 0 means no correlation, and -1 stands for perfect anti-correlation. Figure 2a shows the estimated H₂S abundances as a function of the visual extinction towards the observed positions. They are loosely anti-correlated (Kendall test results: $\tau_{\rm K}$ = -0.3, *p*-value = 0.06), as the H₂S abundance is, in general, at its minimum towards the visual extinction peak, with values of the order of 10⁻¹⁰, and increases towards lower visual extinctions (Fig. 2a). In TMC 1, the maximum H₂S abundance is measured at $A_v \sim 5$ mag, with values of $\sim 2 \times 10^{-8}$. In Barnard 1b, the H₂S abundance peaks at $A_v \sim 10$ mag with values of $\sim 5 \times 10^{-8}$ and then, decreases again, reaching values of $\sim 2 \times 10^{-8}$ at $A_v \sim 5$ mag. We do not detect H₂S when $A_v < 5$ mag (Fig. 2a).



Fig. 2. *Panel a*: observed H_2S abundance as a function of visual extinction in TMC 1-C (blue), TMC 1-CP (red), and Barnard 1b (green). In a similar fashion, *panel b*: observed H_2S gas-phase abundance against the H number density in TMC 1 and Barnard 1b cores. Crosses represent upper bound values. Finally, *panel c*: the H_2S abundance against the gas kinetic temperature.

The H₂S gas-phase abundance is moderately anti-correlated with the hydrogen nuclei density ($\tau_{\rm K} = -0.4$, *p*-value < 10⁻²). This kind of anti-correlation is expected when the freeze-out of molecules on grain surfaces is determining the molecular abundance. The probability of collisions between gas and grains is proportional to the gas density and the thermal velocity.



Fig. 3. Observed H₂S gas-phase abundance plotted against $n_{gas} \sqrt{T}$ (cm⁻³ K^{1/2}) in log scale, showing the sticking behavior described by Eq. (1). Crosses mark upper bound values, which were not taken into account in this analysis.

Assuming that the sticking efficiency is ~ 1 , that is, the H₂S molecules stick on grains in every collision, we can define a depletion timescale given by the inverse of this probability such as

$$X(H_2S) \propto t_{dep} \propto \frac{1}{n_{gas}\sigma_{gr}v} \propto \frac{1}{n_{gas}\sqrt{T_{kin}}},$$
 (1)

where $X(H_2S)$ is the abundance of H_2S respect to H nuclei, n_{gas} is the density of the gas, $\sigma_{\rm gr}$ is the grain cross-section, and v is the thermal velocity of the species, related to the kinetic temperature T_{kin} . In Fig. 3 we plot the H₂S gas-phase abundance against $n_{\text{gas}} \sqrt{T_{\text{kin}}}$. The H₂S abundance decreases with $n_{\text{gas}} \sqrt{T_{\text{kin}}}$ following a power-law with index $m \approx -1$ in the three studied cores, which confirms this behavior (we do not include upper bound values in this analysis). Interestingly, the abundance of H_2S in Barnard 1b is more than a factor of ~ 3 higher than towards TMC 1 for each density, suggesting a higher formation rate or lower destruction rate in Barnard 1b. In Fig. 2c, we plot the H₂S gas-phase abundance as a function of the kinetic temperature. The H₂S abundance is strongly correlated with the gas kinetic temperature (Kendall test gave $\tau_{\rm K} = 0.6$ and *p*-value $p < 10^{-3}$) as long as $A_v > 5$ mag, which translates into a lower abundance towards the cooler quiescent region TMC 1 compared with slightly warmer Barnard 1b core that is located in an active star forming region. In the following we determine the physical structure and model the chemistry of these two regions.

7. Physical structure: core density profiles

The densities at the observed positions in TMC 1 and Barnard 1b (those marked in Figs. 1a and b) were derived using the procedure described in Sect. 4. To fully characterize the density structure of the cores along the line of sight, we consider the simplest case of spherically symmetric, thermally supported, and gravitationally bound clouds. The solution for such a cloud is the well-known Bonnor-Ebert sphere (BE, hereafter). We assume that the radial density structure of the filament is that of a BE, which has been widely parameterized by the approximate analytical profile (see e.g., Priestley et al. 2018; Tafalla et al. 2002):

$$n_{\rm H}(r) = \frac{n_0}{1 + \left(\frac{r}{r_c}\right)^{\alpha}},\tag{2}$$



Fig. 4. H number density profiles of TMC 1-C and TMC 1-CP (*panel a*) and Barnard 1b (*panel b*) as the result of fitting the data in Tables A.1. and B.1, respectively, to the Eq. (2).



Fig. 5. Effective visual extinction in magnitudes of TMC 1-C and TMC 1-CP (*panel a*) and Barnard 1b (*panel b*) as the result of fitting the data in Tables A.1 and B.1, respectively, to the Eq. (3).

where r is the distance from the origin, that is, either TMC 1-C, TMC 1-CP, or Barnard 1b, n_0 is the hydrogen nuclei number density at the origin r = 0, r_c is the flat radius of the BE, and α is the asymptotic power index. At high distances, density falls as $\propto r^{-\alpha}$. As Figs. 4a and b show, the family of curves defined by Eq. (2) is in good agreement with the observed values (to include the origin in a logarithmic scale, we use the symmetrical logarithmic scale from Matplotlib Python library, which by default uses a linear scale between 0 and 1 and the logarithmic scale for the rest of the data range). The central density, n_0 , ranges between $(5.23-7.55) \times 10^4$ cm⁻³ in TMC 1 and $(3.43-9.11) \times 10^5$ cm⁻³ in Barnard 1b. The value of r_c , the flat radius of the BE sphere, is found to be $(0.28 \pm 0.11) \times 10^4$ au and $(1.11 \pm 0.32) \times 10^4$ au in Barnard 1b and TMC 1, respectively. Finally, the asymptotic power index α ranges from 1.76 in the case of Barnard 1b to 4.29 in TMC 1. In a simple parametrization of a BE collapse, a larger flat radius corresponds to a younger, less evolved BE (Aikawa et al. 2005; Priestley et al. 2018). This is in agreement with our parameterizations, since TMC 1-C and TMC 1-CP, starless cores, have larger flat radii than Barnard 1b, which hosts a first hydrostatic core (FHSC for short) and is, therefore, a more evolved core than TMC 1-C and TMC 1-CP. Similarly, the higher asymptotic power index α of TMC 1 is indeed a feature of younger BEs, according to the parameterizations derived by Priestley et al. (2018).

The visual extinction along the line of sight of the selected positions in TMC 1 and Barnard 1b cores is taken from the visual extinction maps in Figs. 1a and b. A parameter that will be needed in the chemical modeling of the cores (Sect. 9) is the shielding from the external UV field at each point. Assuming spherical symmetry and isotropic UV illumination, each point inside the cloud is shielded from the external UV field by an extinction that is, approximately, half of that measured in the extinction maps. We thus define an effective visual extinction A_{veff} as half of the visual extinction measured in the extinction maps: $A_{\text{veff}}(r) = A_{\text{v}}(r)/2$. In Figs. 5a and b, the effective visual extinction is plotted as a function of the distance towards the origin. The effective visual extinction across TMC 1 is interpolated using a cubic spline passing through the average of the effective extinction at each position (Fig. 5a). In Barnard 1b, we fit the effective visual extinction with the following family of functions:

$$A_{\rm veff}(r) = \frac{A_{0\,\rm eff}}{\sqrt{1 + r/r_{\rm c}}},$$
(3)

where *r* is the distance from the origin, $A_{0 \text{ eff}}$ is the effective visual extinction at the origin, and r_c is the flat radius of the profile, the distance at which effective visual extinction starts falling significantly. This kind of profile is well-suited for BEs with an asymptotic power index $\alpha \sim 2$ (see Dapp & Basu 2009).



Fig. 6. Kinetic temperature in the observed positions of TMC 1-C and TMC 1-CP (*panel a*) and Barnard 1b (*panel b*) and the quadratic spline interpolation of the error-weighted average at each position (solid black line).



Fig. 7. *Panel a*: values of the UV field following the Hocuk et al. (2017) parameterization for TMC 1-C (blue) and TMC 1-CP (red) cuts (dashed lines). We also show the best fit (solid line) for the whole TMC 1 region, with $\chi_{UV}(TMC1) = 5$. Similarly, in *panel b*, the parameterization yields a Draine field of $\chi_{UV}(B1b) \sim 24$ for Barnard 1b. It is worth noticing the raise of dust temperature in the $A_v > 30$ mag region, probably due to the FHSC heating of dust or the presence of a thick ice covering.

The flat radius of Barnard 1b is found to be in agreement with the one previously estimated via BE fitting. Finally, as shown in Figs. 6a and b, we parameterize the gas temperature of the observed positions.

Incident UV field in TMC 1 and Barnard 1b. The incident UV field is a key parameter in the photodesorption of H_2S . The increase of the grain temperature at the cloud borders is only understood as the consequence of dust heating by the interstellar radiation field (ISRF). However, to our knowledge, it is not straightforward to derive the incident UV field from the dust temperature. On the one hand, the exact relation between dust temperature and the ISRF is dependent on the poorly known grain composition and its detailed variation across the cloud. On the other hand, in our part of the Galaxy, the dust heating is dominated by the visible part of the ISRF, and the visible and IR parts of the ISRF do not scale in a simple way with the UV part. Remaining aware of all these problems, in Paper I we obtain a first guess of the local UV flux in TMC 1 using the analytical expression by Hocuk et al. (2017). This expression relates the dust temperature of a region with a given visual extinction and the incident UV field in units of the Draine field (Draine 1978). In this paper, we use the same procedure to estimate the incident UV radiation in Barnard 1b.

Figures 7a and b show the $T_{dust} - A_v$ plots for the three cuts considered in this paper. None of the cuts can be fitted with a single value of the UV field. In TMC 1, the dust temperature of the dense core $(A_v > 7.5 \text{ mag})$ is better fitted with an illuminating UV field such that $\chi_{\rm UV} \sim 6.4$, while the translucent region ($A_v < 7.5$ mag) is better fitted with $\chi_{UV} \sim 3.6$. In Barnard 1b, dust temperatures are best fitted with $\chi_{\rm UV} = 24$ in the region $A_v < 20$ mag. However, this value of the incident UV field underestimates the dust temperature in the region $A_v > 30$ mag. The reason for this underestimation is unclear. A thick layer of ice would allow the dust to be warmer by up to 15% at high visual extinctions (Hocuk et al. 2017). It is interesting to note that Barnard 1b is located in an active star forming region, and hosts the two very young protostellar cores B1b-N and B1b-S. In fact, as seen with ALMA, dust temperatures above 60 K have been detected in scales of 0.6" in the Barnard 1b core. Therefore, we cannot exclude an additional dust heating because of the presence of these young protostars. Due to the large column density $N(H_2) \sim 7.6 \times 10^{22} \text{ cm}^{-2}$ in Barnard 1b, this heating would only affect high extinction areas. Since we are interested in the effects of photo-desorption in the low-extinction layers of the cloud, we will adopt $\chi_{UV} = 5$ in TMC 1 and $\chi_{UV} = 24$ in Barnard 1b in our calculations.

8. Photodesorption of H₂S: a simple accretion-photodesorption model

To study the role of photodesorption in the formation of gasphase H₂S, we adapted the simplified PDR model proposed by Hollenbach et al. (2009) to the case of H_2S . In this model, the grains are supposed to be covered by an ice layer, from where H₂S is released into the gas phase via photodesorption. Only two processes, freezing onto grain mantles and photodissociation, are considered for gas-phase H₂S destruction. The gas-phase H₂S formation rate is then proportional to the attenuated local UV interstellar flux $1.7\chi_{\rm UV}F_0e^{-0.9A_v} + \Phi_{\rm SP}$, where F_0 is the local interstellar flux, A_v the visual extinction in magnitudes, Φ_{SP} is the flux of secondary photons, and $\chi_{\rm UV}$ is the incident UV field in Draine units. This rate is also proportional to the photodesorption efficiency $Y_{\rm H_2S}$, which is set to 1.2×10^{-3} molecules per incident photon, as calculated by Fuente et al. (2017), the fraction of desorption sites occupied by H_2S ice (f_{s,H_2S}) , and the probability of collision $(n_{\rm gr}\sigma_{\rm gr})$. The destruction rate however must account for dissociation and freezing of H₂S molecules. The probability of a UV photon to dissociate a H₂S molecule is given by the attenuated UV field flux times the density of H₂S, $1.7\chi_{\rm UV}\dot{R}_{\rm H_2S}e^{-0.9A_{\rm v}}n({\rm H_2S})$, where $R_{\rm H_2S}$ is the H₂S photodissociation rate, and $n(H_2S)$ is the H_2S number density. As discussed previously, the freezing probability is given by the probability of collision between grains and H₂S molecules ($n(H_2S)v_{H_2S}n_{or}\sigma_{or}$). In the stationary state, both creation and destruction rates are equal, and therefore:

$$(1.7\chi_{\rm UV}F_0e^{-0.9A_{\rm v}} + \Phi_{\rm SP})Y_{\rm H_2S}f_{s,\rm H_2S}n_{\rm gr}\sigma_{\rm gr}$$

= 1.7 $\chi_{\rm UV}R_{\rm H_2S}e^{-0.9A_{\rm v}}n({\rm H_2S}) + n({\rm H_2S})v_{\rm H_2S}n_{\rm gr}\sigma_{\rm gr}.$ (4)

It should be noted that secondary photons are not considered to contribute to the photodissociation rate, R_{H_2S} , and are assumed to follow the same extinction law as the FUV radiation. Defining $\sigma_{\rm H} \equiv \frac{n_{\rm gr}(A_{\rm v})\sigma_{\rm gr}}{n_{\rm H}(A_{\rm v})}$, and assuming that the ratio $\frac{n_{\rm gr}(A_{\rm v})}{n_{\rm H}(A_{\rm v})}$ is constant for the range of $A_{\rm v}$ considered,

$$X(\mathrm{H}_{2}\mathrm{S}) = \frac{(1.7\chi_{\mathrm{UV}}F_{0}\ e^{-0.9A_{\mathrm{v}}} + \Phi_{\mathrm{SP}})\ Y_{\mathrm{H}_{2}\mathrm{S}}\ f_{s,\mathrm{H}_{2}\mathrm{S}}\ \sigma_{\mathrm{H}}}{1.7\chi_{\mathrm{UV}}R_{\mathrm{H}_{2}\mathrm{S}}\ e^{-0.9A_{\mathrm{v}}} + v_{\mathrm{H}_{2}\mathrm{S}}\ n_{\mathrm{H}}(A_{\mathrm{v}})\ \sigma_{\mathrm{H}}}.$$
(5)

The fractional coverage of the surface by H_2S ice, f_{s,H_2S} , is given by equating the sticking of S atoms to grain surfaces to the photodesorption rate of H_2S

$$(1.7\chi_{\rm UV}F_0e^{-0.9A_{\rm v}}+\Phi_{\rm SP})Y_{\rm H_2S}f_{\rm s,H_2S}n_{\rm gr}\sigma_{\rm gr}=n({\rm S})v_{\rm S}n_{\rm gr}\sigma_{\rm gr},$$

and therefore,

$$f_{\rm s,H_2S} = \frac{n(S)v_{\rm S}}{Y_{\rm H_2S}(1.7\chi_{\rm UV}F_0e^{-0.9A_{\rm v}} + \Phi_{\rm SP})}.$$
(6)

Combining Eqs. (5) and (6), we are able to predict the H₂S abundance for given physical conditions. The abundance relative to water found in comets is of the order of 2% (Bockelée-Morvan et al. 2000), thus we take as saturation value, $f_{s,H_2S} = 0.02$. We obtain the best fitting of the observed H₂S abundances in TMC 1 with $\Phi_{SP} = 2 \times 10^4$ photons cm⁻² s⁻¹ (Fig. 8a). However, we do not reproduce the H₂S abundances in Barnard 1b with the same flux of secondary photons, specially at higher extinction regions (see Fig. 8b). To explain the abundances observed in Barnard 1b we should adopt a higher value, $\Phi_{SP} = 7 \times 10^4$ photons cm⁻² s⁻¹. Secondary photons are generated by the interaction of H₂ with cosmic rays (Prasad & Tarafdar 1983). While the UV photons



Fig. 8. Observed H₂S gas-phase abundance in TMC 1-C, TMC 1-CP, and Barnard 1b (blue, red, and green, respectively), compared to the PDR model prediction (black). The abundances are calculated using expression (5) and assuming $\Phi_{SP} = 2 \times 10^4$ photons cm⁻² s⁻¹ in TMC 1 and $\Phi_{SP} = 7 \times 10^4$ photons cm⁻² s⁻¹ in Barnard 1b. The value of Φ_{SP} has been selected to match the observations.

are absorbed by dust, cosmic rays penetrate deeper into the cloud maintaining a flux of UV photons even in the densest part of starless cores. The flux of UV secondary photons in dense clouds has a typical value of 10^4 photons cm⁻² s⁻¹, with a cosmic ray molecular hydrogen ionization rate of 3.1×10^{-17} s⁻¹, and a factor of 3 uncertainty (Shen et al. 2004). Recent works point to a lower cosmic ionization molecular hydrogen rate in Barnard 1b than in TMC 1 (Fuente et al. 2016, 2019) which is in contradiction with assuming higher Φ_{SP} in Barnard 1b, rendering our assumptions very unlikely. It is true, however, that the uncertainties of these estimates are large, of around a factor of ~5, not allowing us to fully discard this explanation.

As we have seen, this simple model presents several difficulties. It neglects the chemically active nature of the icy grains, as it does not include any chemical reaction on grain surfaces. Furthermore, recent studies have shown that chemical desorption might be an important source of gas-phase H_2S (Oba et al. 2018), which is not included here. Finally, due to the many adhoc parameters introduced, it lacks of any predicting power. It is therefore necessary to consider a more complete chemical modeling of these cores. In the next section we carry out a 1D modeling of the cores using a full gas-grain chemical network to have a deeper insight into the H_2S chemistry.

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Fig. 9. Predicted abundances (solid lines) of the CS (*top row*), H_2S (*middle row*), and SO (*bottom row*) by models with different chemical desorption schemes, together with the observed abundances in TMC 1, at times 0.1-1-10 Myrs. Notice that there is an extinction interval in which the observed H_2S fit to different chemical desorption schemes. This can be interpreted as a change in the grain surface composition. The vertical dashed line corresponds to $A_y = 6$ mag. The interval 2.5–8 mag around this value is shaded in all subplots.

9. Complete modeling of the TMC 1 and Barnard 1b chemistry

NAUTILUS 1.1 (Ruaud et al. 2016) is a numerical model suited to study the chemistry in astrophysical environments. It solves the kinetic equations for the gas-phase and the heterogeneous chemistry at the surface of interstellar dust grains. NAUTILUS is now a three-phase model, in which gas, grain surface and grain mantle phases, and their interactions, are considered. Given a set of physical and chemical parameters, NAUTILUS computes the evolution of chemical abundances. In the following, we adopt the physical properties of TMC 1 and Barnard 1b derived in Sect. 7. Since both TMC 1-C and TMC 1-CP are described with identical physical parameters, we consider a common chemical model for them.

9.1. Chemical network

We have developed an up-to-date sulfur chemical network, based on the KInetic Database for Astrochemistry (KIDA), including recent updates (Fuente et al. 2017; Le Gal et al. 2017, 2019; Vidal et al. 2017). Our chemical network is composed of 1126 species (588 in the gas phase and 538 in solid phase) linked together via 13155 reactions, with 8526 reactions in gas phase and 4629 reactions in solid phase. The gas-phase reactions are composed of (i) bi-molecular reactions, such as radiative associations, ion-neutral and neutral-neutral reactions), (ii) recombinations with electrons, (iii) ionization and dissociation reactions by direct by UV-photons, cosmic rays, and secondary photons (i.e., photons induced by cosmic rays). The solid phase reactions are composed of both surface and bulk iced grain mantle reactions that occurs for most of them through the diffusive Langmuir–Hinshelwood mechanism. The bulk and surface of the grain mantle interact via swapping processes (Garrod 2013; Ruaud et al. 2016). Desorption into the gas phase is only allowed for the surface species, considering both thermal and non-thermal mechanisms. The latter include desorption induced by cosmic-rays (Hasegawa & Herbst 1993), photodesorption, and chemical desorption (Garrod et al. 2007). We further describe this last process in Sect. 9.2. The binding energies considered in this work can be found in Table 2 of Wakelam et al. (2017) and on the KIDA database website¹.

9.2. Chemical desorption: 1D modeling

In Figs. 9 and 10, we show the comparison of our model with the observations of TMC 1 and Barnard 1b, respectively. We have computed the chemical abundances of different sulphuretted molecules in TMC 1 and Barnard 1b using the physical structures derived in Sect. 7. As initial abundances, we adopt undepleted sulphur abundance (see Table 3). Dust and gas temperatures are assumed to be equal. Other relevant parameters are $\chi_{UV} = 5$ and $\zeta_{CR} = 1.15 \times 10^{-16} \text{ s}^{-1}$ for TMC 1

http://kida.obs.u-bordeaux1.fr/



Fig. 10. Predicted abundances (solid lines) of the CS (*top row*), H₂S (*middle row*), and SO (*bottom row*) by models with different chemical desorption schemes, together with the observed abundances in Barnard 1b, at times 0.1-1-10 Myrs. The vertical dashed line represents $A_v = 8$ mag, and the shaded area encloses the range of extinctions 6-19 mag.

Table 3. Initial abundances.

He	9.00×10^{-2}
Ν	6.20×10^{-5}
0	2.40×10^{-4}
C^+	1.70×10^{-4}
S^+	1.50×10^{-5}
Si ⁺	8.00×10^{-9}
Fe ⁺	3.00×10^{-9}
Na ⁺	2.00×10^{-9}
Mg^+	7.00×10^{-9}
\mathbf{P}^+	2.00×10^{-10}
Cl^+	1.00×10^{-9}
F	6.68×10^{-9}

(Fuente et al. 2019), and $\chi_{\rm UV} = 24$ and $\zeta_{\rm CR} = 6.5 \times 10^{-17}$ for Barnard 1b (Fuente et al. 2016). To compare the output of the chemical model with the observations, we compute a weightaveraged abundance over the line of sight, weighted with the density at each position. Assuming the spherically symmetric density profiles $n_{\rm H}$ given in Figs. 4a and b, the weight-averaged abundance [X]_{ac} of an element X along the line of sight with offset *r* is calculated here as:

$$[\mathbf{X}]_{\rm ac}(r) = \frac{\sum_{i} (l_{i+1} - l_i) (n_{\rm H}(s_i) [\mathbf{X}](s_i) + n_{\rm H}(s_{i+1}) [\mathbf{X}](s_{i+1}))}{\sum_{j} (l_{j+1} - l_j) (n_{\rm H}(s_j) + n_{\rm H}(s_{j+1}))}, \quad (7)$$

where $s_i = \sqrt{r^2 + l_i^2}$, l_i is a discretization of the segment along the line of sight $l_{\text{max}} > \cdots > l_{i+1} > l_i > \cdots > 0$, with $l_{\text{max}} =$ $\sqrt{r^2 + r_{\text{max}}^2}$ and r_{max} being the radius of the density profile $n_{\text{H}}(r)$. In dark clouds, where the temperature of grain particles is below the sublimation temperature of most species, non-thermal desorption processes are needed to maintain significative abundances of molecules in gas phase. This is especially important in the case of H₂S that is thought to be mainly formed on the grain surfaces. Differently from the other processes, chemical desorption links the solid and gas phases without the intervention of any external agents such as photons, electrons, or other energetic particles (Garrod et al. 2007). In other words, it could be efficient in UV-shielded and low-CR environments where photodesorption or sputtering cannot be efficient, becoming more likely the most efficient mechanism in dark cores. This mechanism seems also to be responsible for the abundance of complex molecules such as methanol and formaldehyde in dense cores and PDRs, becoming dominant against phototodesorption (Esplugues et al. 2016, 2019; Le Gal et al. 2017; Vasyunin et al. 2017). New laboratory experiments proved that chemical desorption might be important for H₂S formation on water ice (Oba et al. 2018).

Achieving the correct inclusion of chemical desorption in chemical models is difficult. The efficiency of this process depends not only on the specific molecule involved, but also on the detailed chemical composition of the grain surface on which the reaction occurs, making a correct estimate of its value very difficult to obtain. The chemical desorption process starts from

Table 4. Most important reactions to H₂S in the model.

Visual extinction	TMC	21				
visual extinction	Minissale bare	Minissale ice				
	H ₂ S production					
$A_{\rm v} \sim 5 { m mag}$	$\begin{array}{l} \text{s-H} + \text{s-HS} \rightarrow \text{H}_2\text{S} \ (91.0\%) \\ \text{H}_3\text{CS}^+ + \text{e}^- \rightarrow \text{H}_2\text{S} + \text{CH} \ (8.2\%) \end{array}$	$ s-H+s-HS \rightarrow H_2S \ (90.3\%) \\ H_3CS^+ + e^- \rightarrow H_2S + CH \ (9.2\%) $				
$A_{\rm v} \sim 20 \text{ mag}$	s-H + s-HS → H ₂ S (55.1%) H ₃ S ⁺ + e ⁻ → H + H ₂ S (31.9%)	$s-H + s-HS \rightarrow H_2S (43.6\%)$ $H_3S^+ + e^- \rightarrow H + H_2S (43.3\%)$				
	H ₂ S destr	ruction				
$A_{\rm v} \sim 5 { m mag}$	$\begin{array}{c} H_2S + S^+ \to H_2 + S_2^+ \; (40.1\%) \\ C + H_2S \to HCS + H \; (16.7\%) \end{array}$	$\begin{array}{c} H_2S + S^+ \to H_2 + S_2^+ ~(39.1\%) \\ C + H_2S \to HCS + H~(18.0\%) \end{array}$				
$A_{\rm v} \sim 20 { m mag}$	$ \begin{array}{c} H_2S + H_3^+ \rightarrow H_2 + H_3S^{+(\dagger)} \ (51.7\%) \\ H_2S + S^+ \rightarrow H_2 + S_2^+ \ (11.2\%) \end{array} $	$\begin{array}{c} H_2S + H_3^+ \rightarrow H_2 + H_3S^+ \ (72.9\%) \\ H_2S + H^+ \rightarrow H + H_2S^+ \ (6.4\%) \end{array}$				
Visual extinction	Barnard 1b					
visual extinction	Minissale bare	Minissale ice				
	H ₂ S production					
$A_{\rm v} \sim 5 { m mag}$	$\begin{array}{l} s\text{-}H+s\text{-}HS \rightarrow H_2S~(99.2\%)\\ S+H_2S^+ \rightarrow H_2S+S^+~(0.3\%) \end{array}$	$\begin{array}{l} \text{s-H}+\text{s-HS}\rightarrow\text{H}_2\text{S}\ (85.4\%)\\ \text{s-H}_2\text{S}\rightarrow\text{H}_2\text{S}\ (0.3\%) \end{array}$				
$A_{\rm v} \sim 70 { m mag}$	$\begin{array}{c} H_{3}S^{+}+e^{-}\rightarrow H+H_{2}S~(48.4\%)\\ s\text{-}H+s\text{-}HS\rightarrow H_{2}S~(42.8\%) \end{array}$	$\begin{array}{c} H_{3}S^{+}+e^{-}\rightarrow H+H_{2}S~(47.0\%)\\ s\text{-}H+s\text{-}HS\rightarrow H_{2}S~(46.8\%) \end{array}$				
	H ₂ S destruction					
$A_{\rm v} \sim 5 {\rm mag}$	$\begin{array}{l} H_2S + S^+ \to H_2 + S_2^+ ~(39.2\%) \\ C + H_2S \to HCS + H~(32.1\%) \end{array}$	$\begin{array}{l} H_2S + S^+ \to H_2 + S_2^+ ~(38.0\%) \\ C + H_2S \to HCS + H~(32.4\%) \end{array}$				
$A_{\rm v} \sim 70 { m mag}$	$\begin{array}{c} H_2S + H_3^+ \to H_2 + H_3S^+ \ (73.8\%) \\ H_2S + H^+ \to H + H_2S^+ \ (10.9\%) \end{array}$	$\begin{array}{c} H_2S + H_3^+ \rightarrow H_2 + H_3S^+ \ (69.8\%) \\ H_2S + H^+ \rightarrow H + H_2S^+ \ (20.6\%) \end{array}$				

Notes. (†)This reaction recycles H₂S since the products are reactants in the production reactions.

the energy excess of some reactions. In bare grains, as described in Minissale & Dulieu (2014), its efficiency essentially depends on four parameters: enthalpy of formation, degrees of freedom, binding energy, and mass of newly formed molecules. However, this efficiency becomes lower when dust grains are covered with an icy mantle because part of the energy released in the process is absorbed by the ice matrix. Minissale et al. (2016), based on experimental results, suggested that when taking into account the icy surface, the efficiency of the chemical desorption should be reduced to 10% of the value calculated for a bare grain, when dealing with the water-dominated icy coated mantles of dense regions. We run our chemical model to compare with our observations considering the three chemical desorption scenarios, (i) bare grains following Minissale prescription, (ii) icy coated grains following Minissale prescription, and (iii) ice coated grains assuming Garrod prescription, in which the ratio of the surface-molecule bond-frequency to the frequency at which energy is lost to the grain surface is set to 0.01. The calculations following these three different scenarios are shown in Figs. 9 and 10.

The observed H_2S abundances are in reasonable agreement with those predicted by NAUTILUS at 1 Myr. At this time, the main reactions that lead to the creation and destruction of gas-phase H_2S at low and high extinctions in TMC 1 and Barnard 1b are found in Table 4. In all scenarios, chemical desorption is the main formation mechanism of gas-phase H_2S , and its prevalence diminishes at high extinctions, where density increases and gas-phase reactions become more important to form gas-phase H₂S. Interestingly, the H₂S observations cannot be fitted using the same chemical desorption scheme at every position. In fact, we see that the observational data are in better agreement with the bare grain chemical desorption scheme as described by Minissale et al. (2016) towards the edge of the clouds and to an ice-covered grain chemical desorption scheme towards the more shielded regions. This suggests a change in the chemical composition of grain surfaces, which become covered by a thick ice mantle as the density increases towards the core center, and opens the possibility to use the H₂S abundance to estimate the transition from bare to ice coated grains. Based on our observations, grains become coated with a thick ice mantle at $A_v \sim 8 \text{ mag} (A_{v \text{ eff}} = A_v/2 \sim 4 \text{ mag})$ in TMC 1 and $A_v \sim 12 \text{ mag} (A_v \text{ eff} \sim 6 \text{ mag})$ in Barnard 1b. As a sanity check for our interpretation, we compute a rough estimation of the upper bound of the visual extinction required to photodesorb the ice covering the grains with the expression (Tielens 2010):

$$A_{\rm veff}^{\rm ice} < 4.1 + \ln\left[1.7\chi_{\rm UV}\left(\frac{Y_{\rm pd}}{10^{-2}}\right)\frac{10^4 \ {\rm cm}^{-3}}{n}\right],$$
 (8)

with $\chi_{\rm UV}$ the incident UV field in Draine units, $Y_{\rm pd}$ the photodesorption yield, and *n* the number density. Assuming $Y_{\rm pd} = 10^{-3}$ photon⁻¹ (Hollenbach et al. 2009), we obtain that grains would mainly remain bared for $A_{\rm veff}^{\rm ice} < 6$ mag in TMC 1 and for $A_{\rm veff}^{\rm ice} < 8$ mag in Barnard 1b. These values are in qualitative



Fig. 11. Detail of the predicted H_2S gas-phase abundance in TMC 1 (*left*) and Barnard 1b (*right*) according to the different desorption schemes in a 0-dimensional simulation, and the observed H_2S gas-phase abundances, plotted as a function of density and the time according to a freefall collapse. The shaded band encloses the density interval where the change in desorption scheme occur.

agreement with our results, within a factor of ~2, as grains are expected to remain bare deeper in Barnard 1b than in TMC 1 because of the higher incident UV field. However, this simple expression does not quantitatively reproduce our values of A_{veff}^{ice} . This is not surprising taking into account that our estimate of the local ISRF is uncertain. Furthermore, the above expression is an approximation based on the equilibrium between photo-desorption and freeze-out of water molecules on the grain surfaces. To test the consistency of our result, we will use our chemical model to investigate the ice composition in Sect. 10.

It is also interesting to compare the output of our models with the observations of the other two most abundant S-bearing molecules, CS and SO. One first result is that our models overestimate by more than one order of magnitude the CS abundance in TMC 1 and Barnard 1b. The problem of the overestimation of the CS abundance was already pointed out by Vidal et al. (2017) and Laas & Caselli (2019). Vidal et al. (2017) suggested that it could come from wrong branching ratios of the dissociative recombination of HCS⁺ or a potentially new sink for CS that might be the O + CS reaction whose reaction rate has not been measured at low temperatures. The predicted SO abundances are in reasonable agreement with the observed values if we assume the Garrod prescription to describe chemical desorption even at low visual extinctions. The difficult match between chemical model predictions for S-bearing species and observations make it challenging to determine the sulphur elemental abundance with an accuracy better than a factor of 10.

10. Discussion: is H₂S tracing the snow line in dark cores?

As commented above, the H₂S chemical desorption efficiency seems to decay at a visual extinction of $A_v \sim 8$ mag in TMC 1 and $A_v \sim 12$ mag in Barnard 1b. We propose that this jump in the chemical desorption efficiency might be caused by a change in the chemical composition on the surface of grains, in particular the formation of a thick ice mantle. In this section, we use our chemical model to explore the link between the efficiency of the H₂S chemical desorption as traced by our observations and the ice chemical composition in TMC 1 and Barnard 1b.

In NAUTILUS, two ice phases are considered, the mantle and the surface. The mantle is assumed to grow by incorporating material from the surface. The mantle is hence a fossil record of the grain history. To determine the chemical composition of the ice mantle we should reproduce the trajectory of a cell of gas and dust during the core contraction. As a reasonable approximation, we have converted our 1D profile in a cell trajectory as follows. First, the density, temperature and visual extinction profiles shown in Figs. 4a and b, 5a and b and 6a and b are divided in 300 bins that we use to define the cell trajectory. Second, the time elapsed by the material to reach a bin i is equal to the difference between the free-fall time $t_{\rm ff}(\rho) \approx \sqrt{3/2\pi G\rho}$ of the initial bin and the current bin, that is, $t(i) = t_{\rm ff}(\rho_0) - t_{\rm ff}(\rho_i)$, and consequently, the duration of the whole trajectory is equal to the free-fall time assuming the density of the first bin. This time, for the lowest density in our cuts $n_{\rm H} \sim 1 \times 10^3 \text{ cm}^{-3}$ (see Table B.1), is $t_{\rm ff} \sim 10^6$ yr. It is remarkable that this is the time that best fit the observations in Sect. 9, thus suggesting this choice to be a good guess of the collapse time. Third, in each time-step, the chemical abundances of the previous bin are taken as initial abundances for the next step. These trajectories describe the changes in density, temperature and local UV radiation experienced by a given cell once the collapse has started. Finally, we need to establish the chemical composition of the initial molecular cloud. To set the initial abundances of the cloud, we run the chemical model in a pre-phase, using the physical conditions of the first bin and letting chemistry evolve during 1 Myr, the typical time for this stage. The abundances at the beginning of the cloud collapse are then those that result from the pre-phase.

Following this procedure, we predict the chemical composition of the gas and dust as a function of time. In Fig. 11 we show the H_2S abundance as a function of H number density in TMC 1 and Barnard 1b. The density is continuously increasing with time and provides an easier comparison with observations. We recall, however, that in this comparison we are not taking into account the different physical and chemical conditions along the line-of-sight. According to our results in Sect. 9, for the low densities prevailing in the cloud border, we need to assume the chemical desorption prescription for bare grains to account for the high observed H_2S abundances, but at higher densities, our model overestimates the H_2S abundance. In both sources we find



Fig. 12. Predicted abundances of different molecules in gas phase (*top row*), ice surfaces (*middle row*), and ice mantles (*bottom row*) of TMC 1 (*left column*), and Barnard 1b (*right column*), according to the different desorption schemes. They are plotted as a function of the number density, together with the visual extinction that corresponds to such densities (see Figs. 4a and b, 5a and b), and the time according to a freefall collapse.

the same behavior with a break point at densities around $n_{\rm H} = 2 \times 10^4$ cm⁻³. In the case of Barnard 1b, the abundances are well accounted using the Minissale prescription for ice coated grains for $n_{\rm H} > 2 \times 10^4$ cm⁻³. In TMC 1, the observed abundances are 5–10 times lower than our predictions in any of the scenarios. To relate the H₂S abundance with the ice composition, we show the surface and mantle chemical composition as a function of density in TMC 1 and Barnard 1b in Fig. 12. It is worth noting that the density $n_{\rm H} = 2 \times 10^4$ cm⁻³ corresponds to the phase in which basically the whole ice mantle is formed, achieving the highest values for the s-H₂O and s-CO abundances. Moreover, solid H₂S (s-H₂S) becomes the most abundant S-bearing molecule locked in the ices, with an abundance of ~3 × 10⁻⁶ in both targets, one fifth of the cosmic value. Furthermore, according to our

predictions, the main sulphur reservoir is gaseous atomic sulphur all along the cores.

One additional result is the different ice composition predicted in TMC 1 and Barnard 1b. While the predicted abundance of water ice is as high as $\sim 1.3 \times 10^{-4}$ in TMC 1, its abundance is 1.7×10^{-5} in Barnard 1b. In Barnard 1b, the oxygen is found mainly in the form of CO₂ ice, with an abundance of $X(s-CO_2) \sim 7.4 \times 10^{-5}$. Vasyunin et al. (2017) proposed, based on extrapolations from Minissale's experimental results, that, in general, chemical desorption is enhanced if the ice surface is rich in CO or CH₃OH (instead of water). This could explain the higher abundance of H₂S in the high-density regions of Barnard 1b compared to TMC 1. However, this is not in agreement with *Spitzer* data on the ice composition towards the young

Molecule	Predicted ab	oundances (/H)	Predicted a	bundances (/s-H ₂ O)	Abundance in	67P/Churyumov-
Molecule	TMC 1	Barnard 1b	TMC 1	Barnard 1b	comets $(/s-H_2O)^{(1)}$	Gerasimenko ⁽²⁾
СО	4.0×10^{-5}	2.2×10^{-5}	3.0×10^{-1}	1.3	0.002-0.23	_
CO_2	9.0×10^{-6}	7.4×10^{-5}	6.8×10^{-2}	4.2	0.025-0.30	_
H_2S	3.4×10^{-6}	3.3×10^{-6}	2.5×10^{-2}	1.9×10^{-1}	0.0013-0.015	0.0067-0.0175
OCS	3.8×10^{-7}	1.1×10^{-6}	2.9×10^{-3}	6.2×10^{-2}	$(0.3-4) \times 10^{-3}$	0.00017-0.00098
SO	5.5×10^{-9}	6.3×10^{-8}	4.1×10^{-5}	3.6×10^{-3}	$(0.4-3) \times 10^{-3}$	0.00004-0.000014
SO_2	4.0×10^{-11}	1.4×10^{-8}	3.0×10^{-7}	8.0×10^{-4}	2×10^{-3}	0.00011-0.00041
CS	3.7×10^{-8}	7.9×10^{-8}	2.8×10^{-4}	4.6×10^{-3}	$(0.2-2) \times 10^{-3}$	_
H_2CS	4.0×10^{-8}	5.2×10^{-8}	3.0×10^{-4}	3.0×10^{-3}	$(0.9-9) \times 10^{-4}$	_
NS	1.8×10^{-6}	2.7×10^{-6}	1.3×10^{-2}	1.5×10^{-1}	$(0.6 - 1.2) \times 10^{-4}$	_
S_2	1.2×10^{-9}	2.8×10^{-9}	8.7×10^{-6}	1.6×10^{-4}	$(0.1-25) \times 10^{-4}$	0.000004-0.000013
H ₂ O	1.3×10^{-4}	1.7×10^{-5}	1	1	_	_

Table 5. Abundances of selected molecules in ice mantles.

References. ⁽¹⁾Data taken from Bockelée-Morvan & Biver (2017). ⁽²⁾Taken from Calmonte et al. (2016).

Fig. 13. Predicted ice abundances in the mantle of grains in TMC 1 (light green) and Barnard 1b (dark green) of selected molecules and those observed in comets (dark and light purple) from Table 5. Light purple represents the abundance in comets given in Table 5.

stellar object Barnard 1b-S (Boogert et al. 2008), a star in the moderate density neighborhood of Barnard 1b, which do not show any sign of low s-H₂O abundance or s-CO₂ overabundance. This might be caused by several factors such as, for example, that Spitzer observations might show the ice composition of only the envelope of these objects instead of the dense core. It should be noted, however, that the chemistry on grain surfaces is not very well understood yet and, in addition to the surface chemical network, depends on many poorly known parameters such as the diffusivity of the grain surface. Indeed, the ratio s-CO₂/s-H₂O is very sensitive to the diffusivity of the surface and the grain temperature since the efficiency of surface processes involving O, S, CO, and SO is enhanced when surface diffusivities and ice mobilities are increased (Semenov et al. 2018). The same remains true for s-OCS abundance whose abundance is very sensitive to high mobility of heavy atoms. Further observational constraints are required to fine-tune models and give a more accurate description of the ice composition.

Under the assumption that comets are pristine tracers of the outer parts of the protosolar nebula, it is interesting to compare the composition of comets with that predicted for the ice mantles in the high extinction peaks of TMC 1 and Barnard 1b (see Table 5). There are significant variations in the observed abundances of S-bearing species among different comets. To account for this variation, we compare our predictions with a range of values derived from the compilation by Bockelée-Morvan & Biver (2017). In addition, we have added a specific column to compare with the most complete dataset on comet 67P/Churyumov-Gerasimenko (Calmonte et al. 2016). The ice abundances predicted for TMC 1 and Barnard 1b at the end of the simulation are, in general, close to the values observed in comets (see Table 5 and Fig. 13). These values, however, need to be taken with caution. The abundances of the species in comets are usually calculated relative to water. Our calculations show that the abundance of water in solid phase varies of about one order of magnitude from TMC 1 and Barnard 1b, making the comparison with comets very dependent on the abundance of solid water itself.

Sulphur abundance. A large theoretical and observational effort has been done in the last five years to understand the sulphur chemistry that is progressively leading to a new paradigm (Vidal et al. 2017; Fuente et al. 2016, 2019; Le Gal et al. 2019; Laas & Caselli 2019). Gas-grain models provide a new vision of the interstellar medium in which both gaseous and ice species are predicted. Vidal et al. (2017) updated the sulphur chemistry network and used this new chemical network to interpret previous observations towards the prototypical dark core TMC 1-CP. They found that the best fit to the observations was obtained when adopting the cosmic abundance of S as initial condition, and an age of 1 Myr. At this age, more than 70% of the sulphur would remain in gas phase and only 30% would be in ice. Using the same chemical network but with 1D modeling, Vastel et al. (2018) tried to fit the abundances of twenty-one S-bearing species towards the starless core L1544. The authors found that all the species cannot be fitted with the same sulphur abundance, with variations of a factor of 100, being models with S/H = 8.0×10^{-8} at an age of 1–3 Myr, those that best fitted the abundances of all the twenty-one species. A new gasgrain astrochemical network was proposed by Laas & Caselli (2019) to account for the complex sulphur chemistry observed in comets.

In this paper, we attempt to carry out a comprehensive study of the sulphur chemistry in TMC 1 and Barnard 1b by modeling the CS, SO, and H₂S observations in three cold cores using a fully updated gas-grain chemical network. Our chemical model is unable to fit the observations of the three species with the same parameters. By assuming a cosmic abundance of sulphur, we have a reasonable agreement between H₂S and SO observations with model predictions. However, we overestimate the CS abundance by more than one order of magnitude. Adopting a different approach, Fuente et al. (2019) obtained S/H ~ $(0.4-2.2) \times 10^{-6}$, one order of magnitude lower than the cosmic sulphur abundance, in the translucent part of the TMC 1 filament. Given the uncertainty in the sulphur chemistry, we can conclude that the value of S/H that best fits our data \sim a few 10^{-6} to $1.5\times10^{-5}.$

11. Summary and conclusions

We performed a cloud depth dependent observational and modeling study, determining the physical structure and chemistry of three prototypical dense cores, two of them located in Taurus (TMC 1-C, TMC 1-CP) and the third one in Perseus (Barnard 1b).

- We used the dust temperature to carry out a rough estimate of the incident UV field which is $\chi = 3-10$ Draine field in TMC 1 and $\chi = 24$ in Barnard 1b. Moreover, we modeled the physical conditions of TMC 1- C, TMC 1-CP and Barnard 1b assuming a BE sphere. TMC 1- C and TMC 1- CP are well-fitted with the same parameters while Barnard 1b requires a higher central density consistent with a more evolved collapse.
- Using the full gas-grain model NAUTILUS and the physical structure derived from our observations, we investigated the chemistry of H₂S in these cores. Chemical desorption reveals as the most efficient release path for H₂S in cold cores. Our results show that the abundance of H₂S is wellfitted assuming high values of chemical desorption (bare grains) for densities $n_{\rm H} < 2 \times 10^4$ cm⁻³. For higher densities, our model overestimates the H₂S abundance suggesting that

chemical desorption becomes less efficient. We propose that this critical density is related with a change in the chemical composition of the surface of the grains.

- To further investigate the relationship between the H_2S and grains properties, we examined the chemical composition of the icy mantles along the cores, as predicted by our model. Interestingly, the decrease of the H₂S chemical desorption occurs when the abundance of s-H₂O and s-CO achieves the maximum value in both molecular clouds.
- One additional result is that our model predicts different ice compositions in TMC 1 and Barnard 1b. While the abundance of s-H₂O is as high as $\sim 10^{-4}$ in TMC 1, its abundance is $\sim 10^{-5}$ in Barnard 1b. The abundances of s-H₂S are however very similar in the two clouds, about one fifth of the sulphur cosmic abundance.
- In addition to H_2S , we compared the abundances of CS and SO with model predictions. Our chemical model is unable to fit the observations of the three species simultaneously. Given the uncertainty in the sulphur chemistry, we can only conclude that the value of S/H that best fits our data is the cosmic value within a factor of 10.

This paper presents an exhaustive study of the sulphur chemistry in TMC 1 and Barnard 1b by modeling the CS, SO, and H₂S observations. Our chemical model is unable to fit the observations of the three species at a time, but it does manage to fit, with reasonable accuracy, our H₂S and SO observations. More theoretical and experimental work needs to be done in this area, especially in improving the constraints on the chemistry of CS. Given the uncertainty in the sulphur chemistry, we can only conclude that the value of S/H that best fit our data is the cosmic value within a factor of 10.

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- Observatorio Astronómico Nacional (OAN), Alfonso XII, 3, 28014 Madrid, Spain
- e-mail: dnavarro@oan.es ² Harvard-Smithsonian Center for Astrophysics, 60 Garden St., Cambridge, MA 02138, USA
- ³ Laboratoire d'Astrophysique de Bordeaux, Univ. Bordeaux, CNRS, B18N, allée Geoffroy Saint-Hilaire, 33615 Pessac, France
- ⁴ Faculty of Aerospace Engineering, Delft University of Technology, Delft, The Netherlands
- ⁵ University of Leiden, PO Box 9513, 2300 RA Leiden, The Netherlands
- Centre for Astrochemical Studies, Max-Planck-Institute for Extraterrestrial Physics, Giessenbachstrasse 1, 85748 Garching, Germany
- Institut des Sciences Moléculaires (ISM), CNRS, Univ. Bordeaux, 351 cours de la Libération, 33400, Talence, France
- Observatoire de Paris, PSL Research University, CNRS, École Normale Supérieure, Sorbonne Universités, UPMC Univ. Paris 06, 75005 Paris, France
- Instituto Radioastronomía Milimétrica (IRAM), Av. Divina Pastora 7, Nucleo Central, 18012 Granada, Spain
- Sorbonne Université, Observatoire de Paris, Université PSL, CNRS, LERMA, 92190 Meudon, France
- ¹¹ École Normale Supérieure de Lyon, CRAL, UMR CNRS 5574, Université Lyon I, 46 allée d'Italie, 69364 Lyon Cedex 07, France
- 12 National Radio Astronomy Observatory, 520 Edgemont Rd., Charlottesville VA 22901, USA
- Instituto de Física Fundamental (CSIC), Calle Serrano 123, 28006 Madrid, Spain
- ¹⁴ Centro de Astrobiología (CSIC-INTA), Ctra. de Ajalvir, km 4, Torrejón de Ardoz, 28850 Madrid, Spain
- Jeremiah Horrocks Institute, University of Central Lancashire, Preston PR1 2HE, UK
- 16 Department of Physics, University of Helsinki, PO Box 64, 00014 Helsinki, Finland
- Institute of Physics I, University of Cologne, Cologne, Germany
- ¹⁸ Chalmers University of Technology, Department of Space, Earth and Environment, 412 93 Gothenburg, Sweden
- Leiden Observatory, Leiden University, PO Box 9513, 2300, Leiden, The Netherlands

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Appendix A: Physical conditions and chemical abundances in TMC 1

A.1. Table

Table A.1. TMC 1 physical conditions and chemical abundances.

Source name	$T_{\rm dust}$ (K)	A_v (mag)	$T_{\rm gas}~({\rm K})$	$n_{\rm H} ({\rm cm}^{-3})$	$N(^{13}\text{CS}) (\text{cm}^{-2})^{(a)}$	$N(^{13}\text{CS})/N(\text{H}_{\text{T}})^{(b)}$	$N(\text{ortho} - \text{H}_2\text{S})(\text{cm}^{-2})$	$N(H_2S)/N(H_T)$
TMC 1-CP+0	11.92	18.20	9.7 ± 0.8	$(3.0 \pm 0.8) \times 10^4$	$(3.9 \pm 1.0) \times 10^{12}$	$(1.1 \pm 0.3) \times 10^{-10}$	$(3.1 \pm 0.9) \times 10^{13}$	$(1.1 \pm 0.3) \times 10^{-9}$
TMC 1-CP+30	12.00	16.71	10.2 ± 0.2	$(4.6 \pm 0.4) \times 10^4$	$(1.6 \pm 0.2) \times 10^{12}$	$(4.7 \pm 0.6) \times 10^{-11}$	$(5.1 \pm 1.4) \times 10^{13}$	$(2.0 \pm 0.6) \times 10^{-9}$
TMC 1-CP+60	12.24	13.74	11.3 ± 2.2	$(7.6 \pm 3.8) \times 10^4$	$(1.2 \pm 0.4) \times 10^{11}$	$(4.3 \pm 1.4) \times 10^{-11}$	$(1.2 \pm 0.6) \times 10^{13}$	$(5.7 \pm 0.3) \times 10^{-10}$
TMC 1-CP+120	13.16	7.27	12.5 ± 1.3	$(3.0 \pm 0.8) \times 10^3$	$(2.8\pm 1.1)\times 10^{12}$	$(1.9 \pm 0.8) \times 10^{-10}$	$(5.5 \pm 1.6) \times 10^{13}$	$(5.1 \pm 1.4) \times 10^{-9}$
TMC 1-CP+180	13.86	4.77	16.0 ± 2.6	$(5.4 \pm 1.6) \times 10^3$	$(5.4 \pm 1.5) \times 10^{12}$	$(5.7 \pm 1.6) \times 10^{-11}$	$(4.3 \pm 1.3) \times 10^{13}$	$(5.9 \pm 1.8) \times 10^{-9}$
TMC 1-CP+240	14.39	3.25	14.7 ± 1.1	$(3.2 \pm 2.0) \times 10^3$	$(5.2 \pm 3.2) \times 10^{11}$	$(8.0 \pm 5.0) \times 10^{-12}$	$(2.1 \pm 1.3) \times 10^{13}$	$(4.2 \pm 2.1) \times 10^{-9} (c)$
TMC 1-C+0	11.26	19.85	8.5 ± 2.0	$(9.2 \pm 6.8) \times 10^4$	$(1.1\pm 0.5)\times 10^{12}$	$(2.8\pm 1.2)\times 10^{-11}$	$(2.0 \pm 1.5) \times 10^{13}$	$(6.0 \pm 4.9) \times 10^{-10}$
TMC 1-C+30	11.32	18.47	10.3 ± 2.0	$(8.8 \pm 4.6) \times 10^4$	$(7.0 \pm 3.9) \times 10^{11}$	$(1.9 \pm 1.1) \times 10^{-11}$	$(2.5 \pm 1.3) \times 10^{13}$	$(8.6 \pm 5.0) \times 10^{-10}$
TMC 1-C+60	11.67	13.34	11.6 ± 2.2	$(2.4 \pm 1.0) \times 10^4$	$(9.2 \pm 2.3) \times 10^{11}$	$(3.5 \pm 0.9) \times 10^{-11}$	$(5.4 \pm 2.0) \times 10^{13}$	$(2.6 \pm 1.0) \times 10^{-9}$
TMC 1-C+120	13.13	4.79	11.1 ± 1.9	$(1.1 \pm 0.5) \times 10^4$	$(6.2 \pm 1.8) \times 10^{11}$	$(6.5 \pm 1.9) \times 10^{-11}$	$(1.0 \pm 0.5) \times 10^{13}$	$(1.4 \pm 0.6) \times 10^{-8}$
TMC 1-C+180	14.08	2.20	13.5 ± 1.1	$(1.1 \pm 2.8) \times 10^4$	$(1.8\pm 0.9)\times 10^{11}$	$(4.1\pm2.1)\times10^{-11}$	$(6.5 \pm 3.3) \times 10^{12} \ ^{(c)}$	$(1.9 \pm 1.0) \times 10^{-9} (c)$
TMC 1-C+240	14.53	1.63	13.5 ± 2.7	$(5.2\pm1.8)\times10^3$	$(1.6 \pm 1.0) \times 10^{11}$	$(4.9 \pm 3.0) \times 10^{-11}$	$(1.5\pm1.0)\times10^{13~(c)}$	$(6.0 \pm 4.2) \times 10^{-9} (c)$

Notes. ^(a)When ¹³CS or C³⁴S isotopologues are not detected, ¹³CS column densities are determined from that of C³⁴S or CS, applying the ratios CS/¹³CS \approx 60 and C³⁴S/¹³CS \approx 8/3. ^(b) $N(H_T)$ stands for the total hydrogen column density: $N(H_T) = N(H) + 2 N(H_2)$. ^(c)Upper bound values.

A.2. TMC 1-C spectra

Fig. A.1. Single dish spectra of ¹³CS $1 \rightarrow 0$, ¹³CS $2 \rightarrow 1$, ¹³CS $3 \rightarrow 2$, C³⁴S $1 \rightarrow 0$, C³⁴S $2 \rightarrow 1$, C³⁴S $3 \rightarrow 2$, CS $1 \rightarrow 0$, CS $2 \rightarrow 1$, C³⁴S $3 \rightarrow 2$ transitions towards TMC 1-C positions with offsets (+0", 0"), (+30", 0"), (+60", 0"), (+120", 0"), (+180", 0"), (+240", 0"). The systemic velocity is $v_{lsr} = 6.5 \text{ km s}^{-1}$.

Fig. A.2. Single dish spectra of ¹³CS 1 \rightarrow 0, ¹³CS 2 \rightarrow 1, ¹³CS 3 \rightarrow 2, C³⁴S 1 \rightarrow 0, C³⁴S 2 \rightarrow 1, C³⁴S 3 \rightarrow 2, CS 1 \rightarrow 0, CS 2 \rightarrow 1, C³⁴S 3 \rightarrow 2 transitions towards TMC 1-CP positions with offsets (+0", 0"), (+30", 0"), (+60", 0"), (+120", 0"), (+180", 0"), (+240", 0"). The systemic velocity is $v_{lsr} = 6.5 \text{ km s}^{-1}$.

Appendix B: Physical conditions and chemical abundances in Barnard 1b

B.1. Tables

Table B.1. Barnard 1b physical conditions.

Source name	$T_{\rm dust}$ (K)	$A_{\rm v}$ (mag)	$T_{\rm gas}~({\rm K})$	$n_{\rm H} ({\rm cm}^{-3})$	$N(^{13}CS) (cm^{-2})^{(a)}$	$N(^{13}\text{CS})/N(\text{H}_{\text{T}})$	$N(\text{ortho} - \text{H}_2\text{S})(\text{cm}^{-2})$	$N(H_2S)/N(H_T)$
B1B-cal-0_0	11.90	76.00	9.8 ± 1.4	$(6.3 \pm 3.0) \times 10^5$	$(2.0 \pm 1.6) \times 10^{12}$	$(2.6 \pm 2.1) \times 10^{-11}$	$(5.2 \pm 2.5) \times 10^{13}$	$(4.5 \pm 2.1) \times 10^{-10}$
B1B-cal-10_0	11.72	59.80	10.1 ± 1.5	$(4.2 \pm 2.8) \times 10^5$	$(1.3 \pm 0.9) \times 10^{12}$	$(2.2 \pm 1.5) \times 10^{-11}$	$(1.8 \pm 1.2) \times 10^{14} {}^{(b)}$	$(1.9 \pm 1.2) \times 10^{-9} {}^{(b)}$
B1B-cal-20_0	11.72	45.80	10.9 ± 1.7	$(1.6 \pm 0.9) \times 10^5$	$(1.9 \pm 1.2) \times 10^{12}$	$(4.0\pm 2.6)\times 10^{-11}$	$(3.2 \pm 2.1) \times 10^{14} {}^{(b)}$	$(4.5 \pm 2.9) \times 10^{-9} {}^{(b)}$
B1B-cal-30_0	11.54	38.70	11.9 ± 1.0	$(1.0 \pm 0.4) \times 10^5$	$(1.1 \pm 0.5) \times 10^{12}$	$(2.8 \pm 1.2) \times 10^{-11}$	$(3.1 \pm 1.4) \times 10^{14}$	$(5.2 \pm 2.3) \times 10^{-9}$
B1B-cal-40_0	11.54	28.39	12.1 ± 1.1	$(1.0 \pm 0.4) \times 10^5$	$(9.5 \pm 4.4) \times 10^{11}$	$(3.3 \pm 1.6) \times 10^{-11}$	$(2.0 \pm 0.9) \times 10^{13}$	$(4.5 \pm 2.0) \times 10^{-9}$
B1B-cal-50_0	12.39	20.00	13.2 ± 1.3	$(4.7 \pm 2.2) \times 10^4$	$(1.7 \pm 0.8) \times 10^{11}$	$(8.7 \pm 4.2) \times 10^{-11}$	$(1.9 \pm 0.9) \times 10^{13}$	$(6.0 \pm 2.9) \times 10^{-9}$
B1B-cal-60_0	12.67	20.00	12.3 ± 0.9	$(3.1 \pm 1.6) \times 10^4$	$(2.2\pm 1.1)\times 10^{12}$	$(1.1\pm0.5)\times10^{-10}$	$(3.4 \pm 1.7) \times 10^{14}$	$(1.1 \pm 0.6) \times 10^{-8}$
B1B-cal-80_0	13.24	17.05	13.2 ± 1.8	$(5.5 \pm 2.4) \times 10^4$	$(1.0\pm 0.3)\times 10^{12}$	$(6.1\pm1.8)\times10^{-11}$	$(1.0 \pm 0.4) \times 10^{14}$	$(3.9 \pm 1.7) \times 10^{-9}$
B1B-cal-110_0	13.98	14.46	14.4 ± 1.9	$(5.2 \pm 2.1) \times 10^4$	$(9.8 \pm 2.7) \times 10^{11}$	$(6.8 \pm 1.9) \times 10^{-11}$	$(8.2 \pm 3.3) \times 10^{13}$	$(3.7 \pm 1.5) \times 10^{-9}$
B1B-cal-140_0	14.53	11.87	14.2 ± 1.0	$(9.5 \pm 2.5) \times 10^3$	$(5.4\pm 2.7)\times 10^{12}$	$(4.5 \pm 2.3) \times 10^{-10}$	$(8.3 \pm 2.2) \times 10^{14}$	$(4.5 \pm 1.2) \times 10^{-8}$
B1B-cal-180_0	16.21	8.57	15.3 ± 1.2	$(3.6 \pm 1.7) \times 10^3$	$(6.2 \pm 3.3) \times 10^{12}$	$(7.3 \pm 3.9) \times 10^{-10}$	$(4.2 \pm 2.0) \times 10^{14}$	$(3.2 \pm 1.5) \times 10^{-8}$
B1B-cal-240_0	16.70	6.16	16.4 ± 1.0	$(3.8 \pm 1.8) \times 10^3$	$(1.6\pm 0.8)\times 10^{12}$	$(2.6\pm1.4)\times10^{-10}$	$(1.5 \pm 0.7) \times 10^{14}$	$(1.6 \pm 0.8) \times 10^{-8}$
B1B-cal-500_0	18.23	3.44	18.0 ± 5.4	$(9.6\pm2.2)\times10^2$	$(5.5\pm 3.0) \times 10^{11}$	$(1.6 \pm 0.9) \times 10^{-10}$	$(1.8\pm0.4)\times10^{13(c)}$	$(3.4\pm0.8)\times10^{-9(c)}$

Notes. ¹³CS and H₂S abundances. ^(a)When ¹³CS or C³⁴S isotopologues are not detected, ¹³CS column densities are determined from that of C³⁴S or CS, applying the isotopic ratios CS/¹³CS \approx 60 and C³⁴S/¹³CS \approx 8/3. ^(b)Column densities are obtained from that of the isotopologue H₂³⁴S, using H₂S/H₂³⁴S \approx 22.5. ^(c)Upper bound values.

Table B.2. Barnard 1b physical conditions and SO abundances.

Source name	$T_{\rm dust}$ (K)	$A_{\rm v}$ (mag)	T _{gas} (K)	$n_{\rm H} ({\rm cm}^{-3})$	$N(SO) (cm^{-2})$	$N(SO)/N(H_T)$
B1B-cal-0_0	11.90	76.00	9.8 ± 1.4	$(6.3 \pm 3.0) \times 10^5$	$(2.4 \pm 0.6) \times 10^{14}$	$(1.5 \pm 0.4) \times 10^{-9}$
B1B-cal-10_0	11.72	59.80	10.1 ± 1.5	$(4.2 \pm 2.8) \times 10^5$	$(2.5 \pm 0.9) \times 10^{14}$	$(2.1 \pm 0.8) \times 10^{-9}$
B1B-cal-20_0	11.72	45.80	10.9 ± 1.7	$(1.6 \pm 0.9) \times 10^5$	$(2.7 \pm 1.0) \times 10^{14}$	$(2.9 \pm 1.1) \times 10^{-9}$
B1B-cal-30_0	11.54	38.70	11.9 ± 1.0	$(1.0 \pm 0.4) \times 10^5$	$(2.4 \pm 1.6) \times 10^{14}$	$(3.2 \pm 2.1) \times 10^{-9}$
B1B-cal-40_0	11.54	28.39	12.1 ± 1.0	$(1.0 \pm 0.4) \times 10^5$	$(9.9 \pm 4.5) \times 10^{13}$	$(1.7 \pm 0.8) \times 10^{-9}$
B1B-cal-50_0	12.39	20.00	13.2 ± 1.0	$(4.7 \pm 2.2) \times 10^4$	$(5.2 \pm 1.7) \times 10^{13}$	$(1.3 \pm 0.4) \times 10^{-9}$
B1B-cal-60_0	12.67	20.00	12.3 ± 1.0	$(3.1 \pm 1.6) \times 10^4$	$(4.8 \pm 1.6) \times 10^{13}$	$(1.2 \pm 0.4) \times 10^{-9}$
B1B-cal-80_0	13.24	17.05	13.2 ± 1.8	$(5.5 \pm 2.4) \times 10^4$	$(2.2 \pm 0.8) \times 10^{13}$	$(6.5 \pm 2.3) \times 10^{-10}$
B1B-cal-110_0	13.98	14.46	14.4 ± 1.9	$(5.2 \pm 2.1) \times 10^4$	$(3.4 \pm 1.9) \times 10^{13}$	$(1.2 \pm 0.7) \times 10^{-9}$
B1B-cal-140_0	14.53	11.87	14.2 ± 1.0	$(9.5 \pm 2.5) \times 10^3$	$(4.2 \pm 2.2) \times 10^{13}$	$(1.8 \pm 0.9) \times 10^{-9}$
B1B-cal-180_0	16.21	8.57	15.3 ± 1.2	$(3.6 \pm 1.7) \times 10^3$	$(4.9 \pm 2.4) \times 10^{13}$	$(2.9 \pm 1.4) \times 10^{-9}$
B1B-cal-240_0	16.70	6.16	16.4 ± 1.0	$(3.8 \pm 1.8) \times 10^3$	$(4.9 \pm 2.6) \times 10^{13}$	$(4.0 \pm 2.1) \times 10^{-9}$
B1B-cal-500_0	18.23	3.44	18.0 ± 5.4	$(9.6 \pm 2.2) \times 10^2$	$(1.5 \pm 0.3) \times 10^{12} \ ^{(a)}$	$(2.2 \pm 0.5) \times 10^{-10}$ (a)

Notes. ^(a)Upper bound values.

Fig. B.1. Single dish spectra of ¹³CS $1 \rightarrow 0$, ¹³CS $2 \rightarrow 1$, ¹³CS $3 \rightarrow 2$, C³⁴S $1 \rightarrow 0$, C³⁴S $2 \rightarrow 1$, C³⁴S $3 \rightarrow 2$, CS $1 \rightarrow 0$, CS $2 \rightarrow 1$, C³⁴S $3 \rightarrow 2$ transitions towards positions with offsets (+0", 0"), (+30", 0"), (+60", 0"), (+120", 0"), (+180", 0"), (+240", 0") in the Barnard 1b filament. The systemic velocity is $v_{lsr} = 6.5 \text{ km s}^{-1}$.