Acetone in Orion BN/KL *

High-resolution maps of a special oxygen-bearing molecule

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ABSTRACT

Aims. As one of the prime targets of interstellar chemistry study, Orion BN/KL clearly shows different molecular distributions between large nitrogen- (e.g., C_2H_5CN) and oxygen-bearing (e.g., HCOOCH₃) molecules. However, acetone (CH₃)₂CO, a special complex Obearing molecule, has been shown to have a very different distribution from other typical Obearing molecules in the BN/KL region. Therefore, it is worth investigating acetone in detail at high angular resolutions.

Methods. We searched for acetone within our IRAM Plateau de Bure Interferometer 3 mm and 1.3 mm data sets. Twenty-two acetone lines were searched within these data sets.

Results. Nine of the acetone lines appear free of contamination. Three main acetone peaks (Ace-1, 2, and 3) are identified in Orion BN/KL. The new acetone source Ace-3 and the extended emission in the north of the hot core region have been found for the first time. An excitation temperature of about 150 K is determined toward Ace-1 and Ace-2, and the acetone column density is estimated to be $2-4\times10^{16}$ cm⁻² with a relative abundance of $1-6\times10^{-8}$ toward these two peaks. Acetone is a few times less abundant toward the hot core and Ace-3 compared with Ace-1 and Ace-2.

Conclusions. We find that the overall distribution of acetone in BN/KL is similar to that of N-bearing molecules, e.g., NH_3 and C_2H_5CN , and very different from those of large O-bearing molecules, e.g., $HCOOCH_3$ and $(CH_3)_2O$. Our findings show the acetone distribution is more extended than in previous studies and does not originate only in those areas where both N-bearing and O-bearing species are present. Moreover, because the N-bearing molecules may be associated with shocked gas in Orion BN/KL, this suggests that the formation and/or destruction of acetone may involve ammonia or large N-bearing molecules in a shocked-gas environment.

Key words. Interstellar medium (ISM), ISM: clouds, ISM: molecules, Astrochemistry

1. Introduction

The Orion Molecular Cloud 1 (OMC-1) is a unique environment, especially in the surroundings of the Orion BN/KL region (Becklin & Neugebauer, 1967; Kleinmann & Low, 1967), and one of the closest massive star-forming regions (413±10 pc, Menten et al., 2007; Hirota et al., 2007; Sandstrom et al., 2007) for the study of interstellar chemistry. In this region, evidence of active star formation is overwhelming from detections of multiple strong infrared emission sources or embedded compact radio sources and strong or moderate molecular outflows. For example, a recent unusual explosive event (see, e.g., Allen & Burton, 1993; Zapata et al., 2009) causes the finger-like shock structures seen in vibrationally excited H₂ and high-velocity (30– 100 km s⁻¹) CO emission. Although several sources in BN/KL have been considered as the origin of the explosive outflow, e.g., the continuum source SMA-1¹ proposed by Beuther & Nissen (2008), this explosion is very likely related to the disintegration of a stellar system, which involves the Orion BN object, Source *I*, and maybe Source *n* (see, e.g., Gómez et al., 2005; Zapata et al., 2009; Goddi et al., 2011a). In addition, another outflow with a lower velocity (about 18 km s⁻¹) was detected through the proper motions of H₂O masers (e.g., Genzel et al., 1981; Gaume et al., 1998) accompanied by SiO masers near the infrared source IRc2 (e.g., Greenhill et al., 1998; Doeleman et al., 1999). Evidence indicates that this low-velocity bipolar outflow along a NE-SW axis is driven by source *I* (see e.g., Plambeck et al., 2009).

As one of the prime targets of interstellar chemistry study, Orion BN/KL clearly shows different molecular distributions between large nitrogen- (e.g., C_2H_3CN and C_2H_5CN) and oxygenbearing (HCOOCH₃ and CH₃OCH₃) molecules (see, e.g., Blake

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¹ SMA-1 is a submillimeter continuum peak detected by Beuther et al. (2004) located in the southwest of source I.

Table 1. Observational	parameters o	of the PdB	I data sets
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Bandwidth	Observation date	Configuration	Flux conversion	RMS noise	$\theta_{\mathrm{HPBW}}{}^{a}$	$\delta \mathbf{v}^b$	$\theta_{\rm syn}^{c}$	
(GHz)			(1 Jy beam^{-1})	(mJy beam ⁻¹)	('')	$({\rm km}~{\rm s}^{-1})$	(''×'')	PA (°)
101.178-101.717	2003-2006	BC	15.8 K	2.9	49.7	1.85	3.79×1.99	22
203.331-203.403	10-11/1999	CD	7.0 K	170.1	24.2	0.92	2.94×1.44	27
223.402-223.941	2003-2007	BC	17.3 K	21.2	22.5	0.84	1.79×0.79	14
225.675-225.747	09-11/2005	D	2.9 K	40.8	22.3	0.42	3.63×2.25	12
225.805-225.942	09-11/2005	D	2.9 K	40.8	22.3	0.42	3.63×2.26	12
228.976-229.113	09-11/2005	D	1.8 K	43.9	21.5	0.42	6.02×2.27	10

^(a) Primary beam size ^(b) Channel separation ^(c) Synthesized beam size with natural weightings

Table 2	2. S	pectral	parameters	of	observed	acetone	transiti	ons
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Frequency	Transition	$E_{\rm up}/k$	$S\mu^2$	$g_{\rm s}$	Comment
(MHz)	(J_{K_a,K_c})	(K)	(D^2)		
101297.450 (0.061)	24 _{13,12} – 24 _{12,13} EE	235.5	105.7	16	Partially blended with an NH ₂ CHO line
101426.664 (0.010)	$9_{1,8} - 8_{2,7}$ AE	26.9	61.9	6	
101426.759 (0.009)	$9_{1,8} - 8_{2,7} \text{ EA}$	26.9	61.9	4	
101427.041 (0.010)	$9_{2,8} - 8_{1,7} \text{ AE}$	26.9	61.9	2	
101427.130 (0.009)	9 _{2,8} – 8 _{1,7} EA	26.9	61.9	4	
101451.059 (0.008)	$9_{1,8} - 8_{2,7} \text{ EE}$	26.8	61.9	16	
101451.446 (0.008)	$9_{2,8} - 8_{1,7} \text{ EE}$	26.8	61.9	16	
101475.332 (0.011)	9 _{1,8} – 8 _{2,7} AA	26.7	61.8	10	Partially blended with an H ₂ CS line
101475.734 (0.011)	9 _{2,8} – 8 _{1,7} AA	26.7	61.8	6	Partially blended with an H ₂ CS line
203336.247 (0.036)	27 _{4,23} – 27 _{3,24} EA	225.8	39.0	4	Blended with an NH ₂ CHO line
	27 _{5,23} – 27 _{4,24} EA	225.8	39.0	4	Blended with an NH ₂ CHO line
203336.291 (0.037)	27 _{4,23} – 27 _{3,24} AE	225.8	13.0	6	Blended with an NH ₂ CHO line
	27 _{5,23} – 27 _{4,24} AE	225.8	117.1	2	Blended with an NH ₂ CHO line
223512.409 (0.012)	16 _{8,9} – 15 _{7,8} EE	103.8	60.9	16	Possibly blended with a $C_2H_5^{13}CN$ line ^{<i>a</i>}
223621.690 (0.016)	16 _{8.9} – 15 _{7.8} AA	103.7	60.9	10	Partially blended with an HCOOCH ₃ line
223684.608 (0.012)	17 _{6,11} – 16 _{7,10} EA	110.8	80.6	4	Blended with the U223686.7 line
223684.610 (0.012)	17 _{6,11} – 16 _{7,10} AE	110.8	80.6	6	Blended with the U223686.7 line
223692.004 (0.012)	17 _{7,11} – 16 _{6,10} EA	110.8	80.6	4	Partially blended with the U223694.8 line
223692.104 (0.012)	17 _{7,11} – 16 _{6,10} AE	110.8	80.6	2	Partially blended with the U223694.8 line
223732.826 (0.032)	$12_{9,4} - 11_{8,4}$ EE	66.1	8.7	16	Blended with the U223733.6 line
223767.585 (0.010)	17 _{6,11} – 16 _{7,10} EE	110.7	80.6	16	Blended with the U223767.6 line
223775.253 (0.010)	$17_{7,11} - 16_{6,10} \text{ EE}$	110.7	80.6	16	Possibly blended with a ${}^{13}C_2H_5CN$ line ^{<i>a</i>}
223850.417 (0.014)	$17_{6,11} - 16_{7,10}$ AA	110.7	80.5	10	Partially blended with an HCOOCH ₃ line
223858.308 (0.145)	17 _{7 11} – 16 _{6 10} AA	110.7	80.5	6	Possibly blended with a ¹³ C ₂ H ₅ CN line ^a
225744.082 (0.011)	$19_{4,15} - 18_{5,14} \text{ EE}$	122.1	117.6	16	•
	$19_{5,15} - 18_{4,14} \text{ EE}$	122.1	117.6	16	
225811.979 (0.046)	$13_{95} - 12_{84} \text{EE}$	75.4	36.9	16	Possibly blended with a ${}^{13}C_2H_5CN$ line
228979.750 (0.023)	$12_{10,2} - 11_{9,2}$ EA	68.7	62.6	4	Blended with the U228979.0 line
229033.736 (0.015)	$22_{121} - 21_{220}$ AE	133.1	173.2	2	
	$22_{2,21} - 21_{1,20}$ AE	133.1	173.2	6	
229033.771 (0.015)	$22_{1,21}^{2,21} - 21_{1,20}^{2,20}$ EA	133.1	173.2	4	
	$22_{2,21} - 21_{2,20}$ EA	133.1	173.2	4	
229041.826 (0.024)	$12_{10,3} - 11_{9,3}$ EE	68.6	61.1	16	
229055.797 (0.011)	$22_{121} - 21_{220} EE$	133.1	26.4	16	
	$22_{1,21} - 21_{1,20}$ EE	133.1	146.8	16	
	$22_{2,21} - 21_{2,20} \text{ EE}$	133.1	142.2	16	
	$22_{2,21} - 21_{1,20}$ EE	133.1	31.0	16	
229058.049 (0.010)	$14_{9.6} - 13_{8.5}$ EE	85.4	35.4	16	
229077.788 (0.018)	$22_{1,21} - 21_{2,20}$ AA	133.0	173.2	6	
	22 _{2,21} – 21 _{1,20} AA	133.0	173.2	10	

Notes. Acetone spectroscopic data were taken from the JPL database based on the compilation of several data sets (Groner et al., 2002; Oldag & Sutter, 1992; Vacherand et al., 1986; White, 1975; Peter & Dreizler, 1965). Upper-state energy E_{up}/k , effective line strength $S\mu^2$, and spin statistical weight g_s are also given. Additionally, only the transitions with an Upper-state energy below 300 K and a frequency uncertainty smaller than 1 MHz are listed here. Transitions with E_{up}/k higher than 300 K are all blended and not listed in this table. ^(a) Possibly blended with ¹³C-substituted ethyl cyanide lines at HC, but their contributions are minor (<10%, see also Fig. 5).

et al., 1987; Liu et al., 2002; Friedel & Snyder, 2008, and the references therein). Since the critical densities are fairly similar for N- and O-bearing species, Friedel & Snyder (2008) have proposed the age effect, the time that the molecular gas has had to evolve from an initial composition after being released from ice mantles, may be the most critical factor in the Orion BN/KL N-O differentiation. In addition, they find that the distribution of acetone (CH₃)₂CO is clearly different from those of typical N-or O-bearing molecules.

The differences between acetone and other large O-bearing molecules make acetone an excellent tool for testing different chemical models of complex molecules productions, i.e., in the gas phase by pure cosmic ray induced ion-molecule chemistry, on the surface of grains, or both. The higher temperatures in the vicinity of star-forming regions are certainly an important influence, because these allow some endothermic reactions to proceed. Another influence may be the recent shock event. If so, there would be a dependence on the time at which molecules were released to the gas phase and started to be processed by the gas-phase ion-molecule reactions.

Acetone was first detected in the interstellar medium (ISM) by Combes et al. (1987) and later confirmed by Snyder et al. (2002) toward the hot molecular core Sagittarius B2(N-LHM). They derived an acetone column density of $2 - 9 \times 10^{16}$ cm⁻² with a rotational temperature of 100–270 K in a ~ $13'' \times 7''$ beam toward Sgr B2(N). The first acetone detection toward Orion BN/KL was reported by Friedel et al. (2005) using BIMA, and the follow-up observations at 1 mm with CARMA by Friedel & Snyder (2008) show that the acetone emission is compact in the Orion BN/KL region, and high angular resolution observations have the advantage of detecting weak signals from these complex organic species with compact spatial distributions. The derived acetone column density is $1 - 7 \times 10^{17}$ cm⁻² assuming a rotational temperature of 150–500 K toward the acetone emission peaks in BN/KL.

The formation process of acetone in space is still unclear. Herbst et al. (1990) have shown that the ion-molecule radiative association reaction $CH_3^+ + CH_3CHO \rightarrow (CH_3)_2CHO^+$ proposed by Combes et al. (1987) is very inefficient even at low temperatures, where the rate coefficient for radiative association shows an inverse temperature dependence ($\approx T^{-1}$). Their results suggest that other reactions may in work, e.g., grain surface chemistry, because the radiative association reaction cannot account for the observed acetone abundance in Sgr B2. Therefore, the strategy for determining the most likely formation route of interstellar acetone involves a high angular and velocity comparison of the acetone spectrum with other large Obearing molecules, and an attempt to correlate the distribution of the acetone emission where shocks or heating sources are likely to release molecules from icy grain mantles.

The observations and acetone spectroscopy are presented in §2, and the high-quality images of the acetone emission distribution at < 3" resolution in Orion BN/KL are presented in §3. The spatial and spectral comparisons between acetone and other molecules are discussed in §4. Furthermore, the acetone formation and the cloud structure of BN/KL are discussed in §5. Moreover, we have compared our PdBI data with the science verification (SV) data of the Atacama Large Millimeter Array (ALMA) at 223 GHz in Appendix B. The three-dimensional visualization for different molecules in BN/KL is demonstrated in Appendix C.

2. Observations and spectroscopy

2.1. IRAM observations and line blending

The data used in this study are part of the large 1-3 mm data sets obtained from 1999 to 2007 using the PdBI² toward the Orion BN/KL region (see Favre et al., 2011a, for more observational details). Six data sets (Table 1) were used in this study, and the observations were carried out with five antennas equipped with two SIS receivers. The quasars 0458–020, 0528+134, 0605–085, and 0607-157, and the BL Lac source 0420-014 were observed for the phase and amplitude calibrations. The six units of the correlator allowed us to achieve different bandwidths and spectral resolutions. Using the IRAM 30m single-dish data (J. Cernicharo, priv. comm.), the missing flux for the (CH₃)₂CO lines near 223 GHz is estimated as between 20% and 50%. The large uncertainty in this estimate is due to line confusion and the difficulty of determining spectral baselines in the 30m data. The missing flux in our (CH₃)₂CO line interferometric observations near 101 GHz is estimated to be 25% to 35%.

The PdBI data were reduced with the GILDAS³ package, and the continuum emission was subtracted by selecting linefree channels. Our continuum emission images were presented in Favre et al. (2011a) where the H₂ column densities of selected clumps were also estimated. What we call continuum includes both true continuum emission (from the dust thermal and free-free emission) and a pseudo-continuum made of overlapping faint lines, which cannot be separated from the former true continuum. The data cube was then cleaned channel-bychannel with the Clark algorithm (Clark, 1980) implemented in the GILDAS package.

Possible line blending with acetone was investigated by using the JPL database⁴, the Cologne database for molecular spectroscopy (CDMS⁵), and Splatalogue⁶. It is known that ¹³C-substituted ethyl cyanide (¹³CH₃CH₂CN, CH₃¹³CH₂CN, and $CH_3CH_2^{13}CN$) has intense transitions in mm and submm regimes (Demyk et al., 2007; Fortman et al., 2012), and may blend with acetone lines in Orion BN/KL. We have examined the possible acetone line blending with ¹³C-substituted ethyl cyanide and found: (1) Their spatial distributions are concentrated at the hot core (HC, the strongest mm and submm continuum peak) region judging from the rather isolated $CH_3^{13}CH_2CN 25_{5,20} - 24_{5,19}$ line at 223426.5 MHz (not shown). (2) The synthesized spectrum of ¹³C-substituted ethyl cyanide indicates that their contributions to the acetone intensities are likely to be minor (<10%) in our data (see Fig. 5) toward the main acetone emission peaks. The upper limit of ¹³C-substituted ethyl cyanide column density (beam averaged, 1".8 \times 0".8) at Ace-1 is about 2 \times 10¹⁴ cm⁻² with a rotational temperature of 160 K (see also the synthesized spectra in Fig. 5). Additionally, a ¹³C-substituted ethyl cyanide column density of about $2 - 4 \times 10^{14}$ cm⁻² was estimated toward IRc2 (with a diameter of 7") at 300 K, which is lower than the value $(1.6 \times 10^{15} \text{ cm}^{-2} \text{ with } 300 \text{ K})$ given by Demyk et al. (2007) assuming the same source size toward IRc2. This difference may be because the ¹³C-substituted ethyl cyanide in Orion BN/KL is extended and was largely resolved out in our PdBI observations. Therefore, we conclude that the line blending due to ¹³Csubstituted ethyl cyanide is not severe in our case.

² The IRAM Plateau de Bure Interferometer. IRAM is supported by INSU/CNRS (France), MPG (Germany), and IGN (Spain).

³ http://www.iram.fr/IRAMFR/GILDAS/

⁴ http://spec.jpl.nasa.gov/

⁵ http://www.astro.uni-koeln.de/cdms/

⁶ http://www.splatalogue.net/



Fig. 1. Left: Channel maps of the $(CH_3)_2CO$ emission at 101.43 GHz for a synthesized beam of $3''_{...8} \times 2''_{...01}$, where four $(CH_3)_2CO$ lines, $9_{1,8} - 8_{2,7}$ AE/EA and $9_{2,8} - 8_{1,7}$ AE/EA ($E_{up}/k=26.9$ K), are blended together. Contours run from 9 mJy beam⁻¹ (3 σ) to 114 mJy beam⁻¹ in steps of 15 mJy beam⁻¹, and the dashed contours represent -3 mJy beam⁻¹. The bottom-right panel shows the integrated intensity (from 5 to 11 km s⁻¹) in contours running from 10% to 90% in steps of 10% of the peak intensity (623.1 mJy beam⁻¹ km s⁻¹), and the dashed contours represent -5 mJy beam⁻¹ km s⁻¹ (1 σ). Right: Channel maps of the (CH₃)₂CO emission at 101.45 GHz in a synthesized beam of 3".8 × 2".0, where two $(CH_3)_2$ CO lines, $9_{1.8} - 8_{2.7}$ EE and $9_{2.8} - 8_{1.7}$ EE ($E_{uv}/k=26.8$ K), are blended together. Contours run from 24 mJy beam⁻¹ (8 σ) to 216 mJy beam⁻¹ in steps of 24 mJy beam⁻¹, and the dashed contours represent -3 mJy beam⁻¹. The bottom-right panel shows the integrated intensity (from 5 to 11 km s⁻¹) in contours running from 10% to 90% in steps of 10% of the peak intensity (1.1 Jy beam⁻¹ km s⁻¹), and the dashed contours represent -0.005 mJy beam⁻¹ km s⁻¹ (1 σ). The black square marks the center of explosion according to Zapata et al. (2009). The positions of source BN, HC, infrared sources IRc6/7, source I, methyl formate peaks (MF-1 and MF-3) are marked as triangles. The positions of acetone emission peaks (Ace-1 to Ace-3) are marked as stars.

2.2. $(CH_3)_2CO$ spectroscopy

Acetone is a double methyl-substituted version of the ubiquitous species formaldehyde H₂CO. Owing to its two internal rotators (C_{2V} symmetry), acetone has four torsional substates AA, EE, EA, and AE with slightly different rotational energy levels and different spin statistical weights (see Groner et al., 2002, for more details). The designations AE, EA, EE, and AA refer to the different symmetry states of the molecule. Under normal interstellar conditions, these different symmetry states will not interchange. Because of the different degenerate nature of the torsional substates, the spin weighting has to be taken into account in the column density calculation. The acetone spin weights for the AA : EE : EA : AE symmetry states are 6 : 16 : 4 : 2 for $K_a K_c = ee, oo$ and 10 : 16 : 4 : 6 for $K_a K_c = eo, oe$. The btype asymmetric top selection rules ($\Delta J=0, \pm 1; \Delta K_a=\pm 1, 3, ...;$ $\Delta K_c = \pm 1, 3, ...$) apply to the transitions without severe torsionalrotational interactions (see Groner et al., 2002).

The effective rotational partition function q is approximated by $q = 261.67 T^{1.5}$ for the temperature above 100 K (Groner et al., 2002). The rotational-vibrational partition function used by

Friedel et al. (2005) is $Q_{\rm rv} \approx \sum_{i=0}^{2} e^{-E_i/T} q$, where E_i are the first three vibrational states at 0, 115, and 180 K (Friedel et al., 2005). However, we found that the acetone partition function given in the JPL database starts to deviate from the one used by Friedel et al. (2005) as $T \gtrsim 120$ K (see Fig. A.1). It is likely that the approximation used by Friedel et al. (2005) does not take higher rotational and/or vibrational states into account which start to be essential at higher temperatures. Therefore, whereas the difference is limited for the temperature range (≤ 160 K) derived here for acetone, one should use the more complete acetone partition function given in the JPL database for higher temperatures. The acetone molecular parameters in this paper were taken from the JPL database.

2.3. $(CH_3)_2 CO$ critical density

The critical densities n_c of acetone can be estimated according to

$$n_{\rm c} = \frac{\sum_{l < u} A_{ul}}{\sum_{l \neq u} C_{ul}},\tag{1}$$



Fig. 2. Channel maps of the $(CH_3)_2CO 17_{7,11} - 16_{6,10}$ EE line $(E_{up}/k=110.7 \text{ K})$ at 223775.3 MHz for a synthesized beam of 1''8×0''8. Contours run from 40 mJy beam⁻¹ (2 σ) to 320 mJy beam⁻¹ in steps of 20 mJy beam⁻¹, and the dashed contours represent $-20 \text{ mJy beam}^{-1}$. The bottom-right panel shows the integrated intensity (from 5 to 11 km s⁻¹) in contours running from 20% to 95% in steps of 15% of the peak intensity (151.3 mJy beam⁻¹ km s⁻¹). The black square marks the center of explosion according to Zapata et al. (2009). The positions of source BN, HC, IRc6/7, source *I*, and methyl formate peaks (MF-1 and MF-3) are marked as triangles. The positions of acetone emission peaks (Ace-1 to Ace-3) are marked as stars.

where A_{ul} is the radiative de-excitation rate (Einstein Acoefficient) and C_{ul} the collisional de-excitation rate (Kwok, 2007). Assuming a two-level system, the collisional deexcitation rate can be estimated as (see Friedel & Snyder, 2008; Tielens, 2005; Flower, 2007)

$$C_{ul} = \left(\frac{8kT}{M_{\mu}\pi}\right)^{\frac{1}{2}}\sigma_{ul},\tag{2}$$

where M_{μ} is the reduced mass of acetone and its main collisional partner H₂, and σ_{ul} is the collisional cross section of acetone. Here we assume that the collisional cross section of acetone is approximated by its geometrical cross section (~ 8 × 10⁻¹⁶ cm⁻²). We find $C_{ul} \approx 1.0 \times 10^{-10}$ cm³ s⁻¹ for acetone at 150 K. The critical densities for the detected acetone transitions at 3 and 1.3 mm are estimated to be $4 \times 10^5 - 2 \times 10^8$ cm⁻³. Therefore, the acetone emission generally probes a dense region with a density higher than a few times 10⁵ cm⁻³ at these frequencies, which is similar to HCOOCH₃, C₂H₅CN, and (CH₃)₂O (Favre et al., 2011a; Friedel & Snyder, 2008).

3. Results

Our data include 22 acetone lines corresponding to 40 transitions (see Table 2) with an upper energy state below 300 K. Nine of these lines appear free of contamination by other molecular emission. Three main acetone emission peaks (Ace-1, 2, and 3) were identified in the lower angular resolution images around 101.4 GHz. The acetone emission toward Ace-3 has been detected for the first time. The acetone transition parameters are summarized in Table 2, and the line measurements toward selected positions are listed in Tables A.1 to A.4. The mean local standard of rest (LSR) velocities and FWHM line widths toward selected positions are listed in Table 3.

The LSR velocities of the selected positions are consistent with those reported by Favre et al. (2011a) for methyl formate and Chandler & Wood (1997) for CS J=1-0. The average FWHM line width of acetone is about 3.5 km s⁻¹, similar to those of HCOOCH₃ and (CH₃)₂O lines (Favre et al., 2011a; Brouillet et al., 2013). It is interesting to note that the acetone line widths of Ace-1 are narrower than those of Ace-2, Ace-3, and HC.



Fig. 3. Acetone population diagrams toward Ace-1 and Ace-2. Only the lines without blending or slightly blended (can be decomposed) are used (black data points) in the fitting. The rest of the data points are shown in gray. The FWHM source size θ_s is estimated to be 2 arcsec for Ace-1 and Ace-2 from the high-resolution acetone images. The beam filling factor $f = \theta_s^2/(\theta_a \theta_b + \theta_s^2)$ is taken into account in the calculation, where $\theta_a \theta_b$ is the synthesized beam size.

3.1. Overall acetone distribution

The acetone emission in BN/KL appears mainly at three peaks (Ace-1, Ace-2, and Ace-3) in our lower resolution images $(3''.8 \times 2''.0, Fig. 1)$, where the acetone emission around Ace-1 seems more extended than those around Ace-2 and Ace-3. The positions of the Ace-1 and Ace-2 peaks were determined at the central velocity channels of our highest resolution images $(1''.8 \times 0''.8)$, and the Ace-3 position was determined on the lower resolution images due to the interferometric filtering effects and lower sensitivities of high-resolution images toward this position. The positions of the three acetone peaks are also consistent with the ALMA-SV data (Fig. B.2). The arc-like structure in the north of HC is revealed for the first time in our highest resolution image (Fig. 2). The reason Friedel & Snyder (2008) did not detect the arc-like extended emission and the Ace-3 peak is likely the different uv plane coverage and lower sensitivities.

Ace-1 corresponds to the strongest methanol and deuterated methanol emission peak (dM-1, Peng et al., 2012a) and the second strongest methyl formate emission peak (MF-2, Favre et al., 2011a). Ace-2 is close to the infrared source IRc7 (see, e.g., Shuping et al., 2004; Gezari et al., 1998) but located about 1" southwest of its emission peak. In addition, Ace-3 is located at the same position of the MF-5 (Favre et al., 2011a) and methanol (including deuterated methanol) peak KL-W (e.g., Peng et al., 2012a), and close to the infrared source IRc6 (see, e.g., Shuping et al., 2004; Gezari et al., 1998). The Ace-3 peak shows weak emission in our highest resolution images near 223 GHz because the extended and weaker acetone emission at Ace-3 is likely to be filtered out by the PdBI. We have also investigated the ALMA-SV data at the same frequency with slightly lower



Fig. 4. Acetone population diagrams toward Ace-3 and HC. Only the lines without blending or slightly blended (can be decomposed) are used (black data points) in the fitting. The rest of the data points are shown in gray. The FWHM source size θ_s is estimated to be 1.7 arcsec for Ace-3 from the ALMA-SV images and 2 arcsec for Ace-2 from our PdBI images. The beam filling factor $f = \theta_s^2/(\theta_a \theta_b + \theta_s^2)$ is taken into account in the calculation, where $\theta_a \theta_b$ is the synthesized beam size.

resolution and found clear detections of acetone toward Ace-3 (see Appendix B for more details). It indicates that the two velocity components seen in the acetone and methyl formate emission are most likely caused by the similar filtering effects. Alternatively, there is a possibility of two large-scale structures associated with Ace-3, which would be too extended to be detected by the PdBI (see also Chandler & Wood, 1997). These large structures may extend to the outer part of the BN/KL region, which may also explain the two velocity components at about 6 and 11 km s⁻¹ of the *Herschel*/HIFI O₂ detection (Goldsmith et al., 2011) toward H₂ Peak 1, about 26" to the northwest of Ace-3.

Acetone shows weak emission in the southern part of Orion BN/KL close to MF-1 and MF-3. However, the emission is extended (e.g., in the low resolution maps Fig. 1) and also suffers from filtering effects in our PdBI data (see Fig. B.5).

3.2. Excitation temperature and column density

We use population diagrams to estimate the column density of $(CH_3)_2CO$ (e.g., Goldsmith & Langer, 1999; Turner, 1991)

$$\ln \frac{N_{\rm up}}{g_{\rm up}} = \ln \frac{3k \int T_{\rm B} dV}{8\pi^3 v S \mu^2 g_{\rm s} f} = \ln \frac{N}{Q} - \frac{E_{\rm up}}{k T_{\rm rot}},\tag{3}$$

where *N* is the total column density, T_{rot} the rotational temperature, *S* the line strength, μ the dipole moment ($\mu_b = 2.93$ D, Peter & Dreizler, 1965), g_s the spin statistical weight, and *f* the beam filling factor. The mean optical depth of the acetone lines is estimated to be 0.02–0.03 toward the selected clumps near 101 GHz and less than 0.06 near 223 GHz with excitation temperatures of 100–150 K, taking the dust continuum emission into account and assuming a 90% continuum flux loss due to filtering. The $(CH_3)_2CO$ beam-averaged column densities and rotational temperatures are estimated from a least-squares fit in the population diagrams. The uncertainties shown in these diagrams reflect the statistical fitting error in data where thermal noise and 10% calibration uncertainty dominate. There is no attempt to estimate an absolute error, and we note that estimating the beam filling factor is uncertain. In addition, the column density uncertainties owing to the missing flux are likely to be less than a factor of two (see Appendix B for the comparison with the ALMA-SV data). The estimated acetone column densities, rotational temperatures, and fractional abundances toward the selected positions are summarized in Table 3.

Ace-1

Toward Ace-1 (Fig. 3), an acetone column density is estimated to be $3.4\pm0.1\times10^{16}$ cm⁻² with an excitation temperature of 159 ± 13 K, assuming a source size of 2". An H₂ column density of $2.4\pm0.2\times10^{24}$ cm⁻² is derived toward the same position by using the 1 mm continuum emission (Favre et al., 2011a), assuming a dust temperature of 150 K and an absorption coefficient $\kappa_{\nu} = 5\times10^{-4}$ cm² g⁻¹. Therefore, the acetone abundance toward Ace-1 is $1.4\pm0.4\times10^{-8}$ and agrees with the result ($0.34-1.84\times10^{-8}$) of Friedel et al. (2005).

Ace-2

We derived an acetone rotational temperature of 145 ± 16 K and a column density of $2.1 \pm 0.1 \times 10^{16}$ cm⁻² toward Ace-2 (Fig. 3), similar to those of Ace-1. From the 1.3 mm continuum emission of Favre et al. (2011a), we derived an H₂ column density of $3.8 \pm 2.0 \times 10^{23}$ cm⁻² toward Ace-2. Owing to its weaker dust continuum emission, the acetone fractional abundance $(5.5 \pm 4.0 \times 10^{-8})$ of Ace-2 seems to be higher than that of Ace-1 but with a larger uncertainty.

Ace-3

The acetone rotational temperature of Ace-3 is estimated to be ~ 100 K, and the acetone column density to be $6.3 \pm 1.9 \times 10^{15}$ cm⁻² (Fig. 4). From the 1.3 mm continuum emission of Favre et al. (2011a), we derived an H₂ column density of $1.3 \pm 0.3 \times 10^{24}$ cm⁻² toward Ace-3 with a dust temperature of 100 K. The acetone abundance toward Ace-3 is $4.8 \pm 1.8 \times 10^{-9}$.

HC

The acetone rotational temperature of HC is estimated to be about 141 K and the column density is estimated to be $1.2 \pm 0.2 \times 10^{16}$ cm⁻² (Fig. 4). From the 1.3 mm continuum emission of Favre et al. (2011a), we have derived an H₂ column density of $1.3 \pm 0.3 \times 10^{24}$ cm⁻² toward HC with a dust temperature of 140 K. The acetone abundance toward HC is $2.1 \pm 0.4 \times 10^{-9}$.

3.3. Synthesized spectra of acetone

The synthesized spectra of acetone (Fig. 5) were produced with Weeds (Maret et al., 2011) in the GILDAS package, assuming that the source sizes equal the PdBI synthesized beams at 101 and 223 GHz. Although the synthesized spectra do not fully reproduce the observed acetone intensities, the uncertainty of



Fig. 6. Comparison of the observed spectra (gray) and the synthesized acetone spectra (green) in LTE near 229 GHz toward Ace-1. The observed spectra are plotted at the original synthesized beam size of 6.0×2.03 . The acetone synthesized spectra were produced assuming a $(CH_3)_2CO$ column density of 1.7×10^{16} cm⁻² and a temperature of 160 K, and the line widths of acetone were fixed at 3.5 km s⁻¹. As in Table 2, only the transitions with $E_{up} < 300$ K are indicated with their frequencies (in MHz), and the others are all blended. In addition, the methyl formate synthesized spectra (blue) are also shown for comparison.

10%–20% is likely contributed by line blending, different beam sizes, the accuracy of the acetone column density, excitation temperature, and possible non-LTE effects.

4. Comparison with other molecules

4.1. Spatial comparison

The detection of two molecules with strong and not blended lines, C₂H₅CN and HCOOCH₃, within the same data sets as acetone allows us to directly compare their distributions with the same observational parameters. In Figure 7, the integrated intensity maps of C₂H₅CN, (CH₃)₂CO, and HCOOCH₃ were overlaid on the same velocity interval. The emission of the large Nbearing molecule C₂H₅CN is mainly located in the northern part of the Orion BN/KL region, i.e., close to HC and IRc7, similar to the highly excited NH₃ (Goddi et al., 2011b). The emission of the typical O-bearing molecule HCOOCH₃ appears mostly in the south, i.e., the Compact Ridge region with a typical V_{LSR} of about 8 km s⁻¹, whereas the acetone emission appears in the regions where the C₂H₅CN emission is also present. It is interesting to note that the HCOOCH₃ emission does not appear in the regions associated with hot (≈ 400 K) and dense ammonia (Goddi et al., 2011b). Since the emissions of C₂H₅CN and highly excited NH₃ (i.e., inversion transitions from $J, K \ge 9, 9$) share a similar distribution, it is likely that C2H5CN traces somewhat higher temperatures than HCOOCH₃ does, and (CH₃)₂CO may be excited in the intermediate temperature range (see also Friedel & Snyder, 2008). For example, the excitation temperature of C₂H₅CN toward the HC region is about 150 K, and is about 100 K for HCOOCH₃ toward the Compact Ridge region (Favre et al., 2011a; Blake et al., 1987). It is probably related to the acetone formation route in the interstellar medium (see §5). The high similarity of the acetone and C₂H₅CN distributions is also seen in the 3D images in Appendix C, where both molecules show a similar arc-like structure that may be associated with the explosive event.



Fig. 5. Comparison of the observed spectra (gray) and the synthesized acetone spectra (green) in LTE at 1.3 and 3 mm toward Ace-1. The observed spectra are plotted at their original synthesized beam sizes of $3''8 \times 2''0$ (101 GHz) and $1''8 \times 0''8$ (223 GHz). The acetone synthesized spectra near 101 GHz (upper panel) were produced by assuming a (CH₃)₂CO column density of 3×10^{16} cm⁻² and a temperature of 160 K, and the acetone synthesized spectra near 223 GHz (lower panel) were produced assuming a (CH₃)₂CO column density of 4×10^{16} cm⁻² with the same temperature. Line widths of acetone synthesized spectra were fixed at 3.5 km s⁻¹. As in Table 2, only the transitions with $E_{up} < 300$ K are indicated with their frequencies (in MHz), and the others are all blended. Synthesized ¹³C-substituted C₂H₅CN spectra are shown in red, assuming a column density of 2×10^{14} cm⁻² and a temperature of 160 K. Their contributions to the acetone lines blending are minor. In addition, the methyl formate synthesized spectra (blue) are also shown for comparison.

Table 3. Acetone line	parameters toward	selected	positions

Position	R.A.	Dec.	$V_{\rm LSR}^{a}$	ΔV^b	$T_{\rm rot}^{\ d}$	$N_{\rm tot}{}^d$	X _{ace} ^d	Note ^c
	$(05^{h}35^{m})$	$(-05^{\circ}22')$	$({\rm km}~{\rm s}^{-1})$	$({\rm km}~{\rm s}^{-1})$	(K)	(cm^{-2})		
Ace-1	14 <u></u> .449	34''.24	7.47 ± 1.07	3.49 ± 1.09	158.7 ± 13.1	$3.4\pm0.1\times10^{16}$	$1.4 \pm 0.4 \times 10^{-8}$	dM-1, MF-2, CS-3
Ace-2	14 <u>°</u> 240	31″55	7.58 ± 1.27	4.46 ± 1.30	145.4 ± 16.2	$2.1\pm0.1\times10^{16}$	$5.5 \pm 4.0 \times 10^{-8}$	CS-5, IRc7
Ace-3	14 <u>°</u> 143	28''06	6.70 ± 1.95	4.20 ± 1.94	~100	$6.3 \pm 1.9 \times 10^{15}$	$4.8 \pm 1.8 \times 10^{-9}$	MF-5, CS-6, IRc6, KL-W
HC	14 <u>°</u> 580	31"00	6.87 ± 1.29	4.92 ± 1.37	141.0 ± 20.9	$1.2 \pm 0.2 \times 10^{16}$	$2.1\pm0.4\times10^{-9}$	CS-1

Notes. Positions are given at the epoch J2000.0. Acetone column densities and fractional abundances were estimated in source sizes of 2" for Ace-1, Ace-2, and HC, and 1".7 for Ace-3. ^(a) Mean LSR velocities measured at peak temperatures. ^(b) Mean FWHM line widths. ^(c) Counterparts as observed in the deuterated methanol (dM, Peng et al., 2012), methyl formate (MF, Favre et al., 2011a), CS J=1-0 (Chandler & Wood, 1997), and infrared (Shuping et al., 2004; Gezari et al., 1998) emission. Note that IRc7 is offset ~ 1" to the southwest from Ace-2, and IRc6 is extended. ^(d) The acetone rotational temperatures T_{rot} , column densities N_{tot} , and fractional abundances ($X_{ace} = [(CH_3)_2 CO]/[H_2]$) were estimated with source sizes of 2 arcsec.

Chandler & Wood (1997) obtained a high-resolution ($\theta_{syn} = 2''.1 \times 1''.7$) CS J=1-0 image with VLA, and they identified six main CS peaks in Orion BN/KL. It is essential to note that all acetone peaks have counterparts in the CS J=1-0 emission. The CS peaks CS-3, 5, and 6 correspond to Ace-1, 2, and 3, respectively, and CS-1 is located at the HC position (see Table 3 for the counterpart summary). In addition, the CS J=1-0 emission is strong in the south of BN/KL (i.e., CS-4) close to MF-1 and 3, where most of the large O-bearing molecules are detected. The overall CS distribution covers the region where large N- and

O-bearing molecules are detected. The sulfur-bearing molecules (e.g., CS, SO, and SO₂) mainly form via warm gas-phase chemistry or shock chemistry (e.g., Charnley, 1997; Pineau des Forêts et al., 1993), and they coincide with both N- and O-bearing molecules in BN/KL (Schreyer et al., 1999; Chandler & Wood, 1997). This suggests that the formation routes via warm gas phase or shock chemistry may lead to the formation of acetone in BN/KL.

Furthermore, Chandler & Wood (1997) noticed that the integrated CS emission does not coincide with any identified



Fig. 7. Spatial comparison of C_2H_5CN , HCOOCH₃, and (CH₃)₂CO emissions with the same resolution (1'' 8×0 ''8) in Orion BN/KL. The C_2H_5CN 25_{3,23} – 24_{3,22} line at 223553.6 MHz is shown in light-red contours. The HCOOCH₃ 11_{4,8} – 10_{3,7} E/A lines at 223465.3 MHz and 223500.5 MHz are combined and shown in light-blue contours. The (CH₃)₂CO 17_{7,11} – 16_{6,10} EE line at 223775.3 MHz is shown in thick olive contours. Contours run from 15% to 95% in steps of 10% of their peak intensities, and dashed contours represent –10% of their peak intensities. The black square marks the center of explosion according to Zapata et al. (2009). The positions of source BN, source *I*, source *n*, HC, SMA-1, and IRc6/7 are marked as triangles. The positions of acetone emission peaks (Ace-1 to Ace-3) and some of the methyl formate emission peaks (MF-1, MF-3, MF-8, and MF-9) are marked as stars. Black crosses mark the three NH₃ column density peaks (also close to the temperature peaks) according to Goddi et al. (2011b).

clumps. This may be because the velocity dispersion in the arclike structure is higher, and the integrated flux is dominated by the line width rather than the peak intensity. Although the acetone integrated intensity shown in Figures 2 and 7 peaks at SMA-1 (Tang et al., 2010; Beuther et al., 2004), it is likely to be caused by increased velocity width, similar to the CS emission peak mentioned above.

4.2. Spectral comparison

In Figure 8, we compare the line profiles of $(CH_3)_2CO$ with C_2H_5CN and $HCOOCH_3$, together with formamide ($HCONH_2$), which is an interesting molecule because it contains one oxy-

gen and one nitrogen atom and was detected in our data set near 223 GHz. It is interesting to note that the acetone spectra toward Ace-1 and Ace-2 show strong blueshifted line wings at about 4 km s⁻¹. The HCONH₂ 11_{1,11}-10_{1,10} spectrum toward Ace-2 also shows a blueshifted line wing. These blue-wing features may result from shocks. On the other hand, the HCOOCH₃ spectra do not have clear line wings.

Toward Ace-3, the $(CH_3)_2CO$ and HCOOCH₃ spectra reveal that at least two velocity components are present. However, a velocity shift in V_{LSR} of the two components is observed, where the velocity difference between the two components is larger in acetone (~ 4 km s⁻¹) than in methyl formate (~ 2 – 3 km s⁻¹). In addition, the ALMA-SV acetone spectrum toward Ace-3 shows



Fig. 8. Comparison of the $(CH_3)_2CO$, HCOOCH₃, HCONH₂, and C_2H_5CN spectra toward Ace-1, Ace-2, Ace-3, and HC with the same resolution (1''8 × 0''.8). Dashed lines indicate the V_{LSR} =8 km s⁻¹, and the dotted lines indicate the (CH₃)₂CO lines at 223767.6 and 223850.4 MHz (17_{6,11} – 16_{7,10} EE/AA) in black and red, respectively. The spectra were shifted in intensity and some of them were multiplied for clarity.

redshifted line wings (see Figure B.4), which is not clear in our PdBI data. The differences observed between the ALMA-SV and PdBI data are most probably due to different uv coverages and sensitivities (see Table B.1). Because of its extended spatial nature, the acetone emission from Ace-3 is less likely to be recorded in our PdBI data.

Similarly, blueshifted wings can be seen in CS J=1-0 toward CS-3 (Ace-1) and CS-5 (Ace-2) (Chandler & Wood, 1997). Additionally, the CS J=1-0 line widths are broader in CS-6 (Ace-3) than in CS-3 and CS-5. The two velocity components seen in CS-6 are also due mainly to the filtering effects similar to acetone and methyl formate, compared with the ALMA-SV data. CS-3 (Ace-1) has a rather narrow line width of 3.3 km s⁻¹, similar to acetone. Therefore, most of the molecular lines in some regions of BN/KL (i.e., HC/CS-1, Ace-2/CS-5, and Ace-3/CS-6) show broader line widths than those in the other regions (i.e., Ace-1/CS-3, MF-3/dM-2/CS-4, and dM-3/MF-1) indicates that some regions may be more directly affected by shocks than others. These shocks could be driven by local outflow phenomena and/or the Orion explosion event.

5. Discussion

5.1. Acetone formation

Some studies have suggested there is formation of carbonyl species, such as H_2CO , CH_3CHO , and $(CH_3)_2CO$, on or within icy grain mantles (e.g., Snyder et al., 2002; Bennett et al., 2005). In the model of Garrod et al. (2008), acetone is said to form by addition of CH_3 to CH_3CO and behaves similarly to $HCOOCH_3$

and (CH₃)₂O. Recent results from Favre et al. (2011b) and Brouillet et al. (2013) show good agreement in the distributions between HCOOCH₃ and (CH₃)₂O in BN/KL. However, acetone clearly has a different distribution from those of HCOOCH₃ and $(CH_3)_2O$, and more similar ones to that of C_2H_5CN . It is not clear whether the acetone binding energy (it evaporates from the ice mantle at about 65 K) used in the model of Garrod et al. (2008) is a critical factor or not. The acetone excitation temperature toward Ace-1 is estimated to be about 150 K, and it agrees with that of HCOOCH₃ (140 ± 14 K, Favre et al., 2011a). Therefore, $(CH_3)_2CO$ and $HCOOCH_3$ seem to trace the same gas toward Ace-1. On the other hand, not detecting HCOOCH₃ toward the Ace-2/IRc7 position indicates that different formation processes from (CH₃)₂CO and HCOOCH₃ are involved. It is possible that some forming pathways of acetone have not been taken into account in the previous modelings. For example, Chen & Woon (2011) carried out a theoretical investigation of possible reactions between carbonyl species and ammonia and show that acetone may react with water and ammonia via nucleophilic addition with low barriers. Although the final products of these reactions on icy grain mantles are not clear, further research is important. Because of its similar distribution to the N-bearing molecules, the formation of acetone may involve ammonia or N-bearing molecules or need similar conditions, e.g., shocks.

It has been proposed by Remijan et al. (2004) that the formation of acetic acid (CH₃COOH) favors the hot molecular cores with well-mixed N- and O-bearing molecules. This may be another example that the formation of a large O-bearing molecule involves N-bearing molecules or needs similar conditions, although acetic acid has not been detected so far toward BN/KL



Fig. 9. Cartoon presentation of the Orion BN/KL structure.

(Remijan et al., 2003). However, acetone emission is detected toward the positions (i.e., IRc7 and arc-like structure in the north) where there is no strong emission of large O-bearing species like HCOOCH₃ and (CH₃)₂O (Favre et al., 2011a; Brouillet et al., 2013). Furthermore, it is interesting to search for the molecules with distributions similar to acetone in Orion BN/KL in the future. Investigations of molecules related to acetone will be important for understanding the origins of interstellar acetone.

Table 4. Characteristics of different molecules in Orion BN/KL

Species	Example	Distribution			Main	peak ^a			Feature
S-bearing	CS	whole	CS-1	CS-5	CS-6	CS-3	$CS-4^b$	$CS-4^b$	both N- and O-bearing features
N-bearing	NH_3, C_2H_5CN	north	HC	IRc7	KL-W				higher T and broader ΔV
O-bearing	$(CH_3)_2CO$	north	HC	Ace-2	Ace-3	Ace-1			higher T and broader ΔV
	$HCOOCH_3, (CH_3)_2O$	south			MF-5	MF-2	MF-1	MF-3	lower T and narrower ΔV
	CH ₂ DOH, CH ₃ OD	south			KL-W	dM-1	dM-3	dM-2	lower T and narrower ΔV

Notes. ^(a) The strongest peaks of different molecules. Orion KL-W is close to IRc6 and MF-1 is close to IRc4. ^(b) CS-4 is located between MF-1 and MF-3 but closer to MF-3.

5.2. Overall cloud structure of BN/KL

A clearer picture has emerged from the investigations of different molecules' distribution and the recent understanding of the stellar feedbacks inside the Orion BN/KL region. It has been shown that the explosive outflow that happened about 500 years ago may affect the BN/KL region in many ways. For example, Zapata et al. (2011) propose that HC, the strongest mm and submm continuum peak, is probably heated by the explosive outflow owing to its lack of embedded heating sources. In addition, Peng et al. (2012b) suggest that the high velocity bullets from the explosive outflow may be absorbed by the BN/KL cloud and induce the methanol masers seen in the south. The absorbed kinetic energy can then be transferred to thermal energy that affects the chemical processes in the cloud. As expected for the decelerated bullets, the thermal energy (or velocity) deposit is higher close to the explosion center than close to the stop point. This is clearly seen in the north-south differences in excitation temperatures and line widths of many molecules (see Table 4).

Figure 9 shows a cartoon overview of different molecular distributions in BN/KL. As discussed above, because the spectral features of CS are consistent with other molecules toward the same clump (e.g., smaller line widths in the south compared to the north of the cloud), the N-O differentiation seen in BN/KL is very likely caused by physical conditions (i.e., warmer and shocked gas) instead of their chemical properties (e.g., ice mantle composition). On the other hand, the age effect cannot be ruled out since different outflows in the northern region (e.g., the bipolar outflow of source I) may contribute, too, and the reason the northern part is more turbulent may be its more evolved nature. Nevertheless, the bipolar outflow of source I alone cannot explain the blueshifted wings seen in most of the acetone peaks, because the outflow lies along a NW-SE axis (see, e.g., Plambeck et al., 2009) and its orientation tends to be constant when moving from the explosion center to the present position (Bally et al., 2011). In addition, the acetone peaks are located in the direction of the source I southern outflow lobe, which appears to be more redshifted than the northern one, but the line wings associated with Ace-1 and Ace-2 are blueshifted.

6. Conclusions

We have explored the acetone distribution in Orion BN/KL at angular resolutions between $1''.8 \times 0''.8$ and $6''.0 \times 2''.3$ (about 300–2500 AU) from the 3 mm and 1.3 mm Plateau de Bure interferometric data sets. The main findings follow.

1. Our sample consists of 22 lines of acetone with an upper energy state below 300 K, and nine of these lines appear free of contamination by other molecular emission. The FWHM line width of these acetone lines is about 3.5 km s⁻¹, similar to those of HCOOCH₃ and (CH₃)₂O lines.

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- 2. The acetone distribution exhibits three main peaks (Ace-1, 2, and 3) and the extended emission toward the Hot Core and the northern region, which appears to be arc-like. Ace-1 is also the strongest peak of the deuterated methanol emission (dM-1, Peng et al., 2012a) and second strongest peak of HCOOCH₃ (MF-2 Favre et al., 2011a). Ace-2 is located close to the peak of the IR source IRc7 with a separation of about 1". The new weaker source Ace-3, which was not detected by Friedel & Snyder (2008), coincides within 2" with the KL-W/MF-5 position (Peng et al., 2012a; Favre et al., 2011a) and is close to the IR source IRc6.
- 3. The overall distribution of acetone in BN/KL is more extended than the one found by Friedel & Snyder (2008) and is similar to that of the large N-bearing species (i.e., C₂H₅CN). This is likely due to the higher sensitivity and different uv coverage of our PdBI data compared with the CARMA data of Friedel & Snyder (2008).
- 4. Rotational temperatures of 159 ± 14 are determined toward Ace-1 and 145 ± 16 K toward Ace-2. The acetone column density toward these two peaks is about $2-4 \times 10^{16}$ cm⁻² with a relative abundance of $1-6 \times 10^{-8}$, within the range given by Friedel et al. (2005). In addition, Acetone is a few times less abundant toward the hot core and Ace-3 peaks $(2-7 \times 10^{-9})$ compared with Ace-1 and Ace-2.

Current chemical models require a formation of acetone on grain surfaces as for $(CH_3)_2O$ and they cannot explain the difference seen between the two species. Including reactions with N-bearing species either to form or to destroy acetone might be necessary to explain its different distribution (e.g., Chen & Woon, 2011). Additional laboratory and theoretical chemical studies are needed to progress in the understanding of the acetone abundance. High-resolution maps with more sensitive ALMA data will hopefully lead to the discovery of other species with the same spatial distribution, and confirm any association with shocks.

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Appendix A: Complementary figures and tables



Fig. A.1. Comparison of the acetone partition functions taken from the JPL database (black squares) and the approximation (blue curve) used by Friedel et al. (2005). The red curve is a fit to the JPL acetone partition functions.



Fig. A.2. (CH₃)₂CO spectra near 101 GHz toward four positions (three acetone emission peaks and HC) in the Orion BN/KL region. (b)-(c) Spectra correspond to the channel maps shown in Fig. 1 at 101427 MHz and 101451 MHz, respectively. Dashed lines indicate the V_{LSR} of 8 km s⁻¹ and correspond to the (CH₃)₂CO line shown at the top of each panel.

Table A.1.	$(CH_3)_2CO$	measurements	toward	Ace-1
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Frequency	Transition	T _B	V _{LSR}	ΔV	$\int T_{\rm B} dV$	Comment
(MHz)	(J_{k_a,k_c})	(K)	$({\rm km}~{\rm s}^{-1})$	$({\rm km}~{\rm s}^{-1})$	$(\check{K} \text{ km s}^{-1})$	
101297.450	24 _{13,12} - 24 _{12,13} EE	0.85 ± 0.05	7.31±1.85	5.55 ± 1.85	4.97±0.83	Partially blended ¹
101426.664	$9_{1.8} - 8_{2.7} \text{ AE}$					•
101426.759	$9_{1,8} - 8_{2,7} \text{ EA}$	2.04 + 0.05	6 92 1 95	5 14 1 96	10.97 + 0.44	
101427.041	$9_{2,8} - 8_{1,7}$ AE	2.04 ± 0.03	0.82±1.83	3.14 ± 1.80	$10.8/\pm0.44$	
101427.130	9 _{2,8} – 8 _{1,7} EA					
101451.059	$9_{1,8} - 8_{2,7} \text{ EE}$	3 88+0.05	6 80+1 85	5 06+1 85	20 46+0 48	
101451.446	9 _{2,8} – 8 _{1,7} EE	5.88±0.05	0.89±1.85	J.00±1.85	20.40 ± 0.40	
101475.332	$9_{1,8} - 8_{2,7}$ AA	2.25 ± 0.05	6 59+1 85	5 55+1 85	11.27 ± 1.20	Partially blended ²
101475.734	9 _{2,8} – 8 _{1,7} AA	2.25±0.05	0.57±1.05	5.55±1.05	11.27±1.27	I artiany biclided
203336.247	27 _{4,23} – 27 _{3,24} EA					
	27 _{5,23} – 27 _{4,24} EA	14 30+1 19	8 27+0 92	348+094	51 41+4 57	Blended ³
203336.291	$27_{4,23} - 27_{4,24}$ AE	11.30±1.19	0.27±0.92	5.10±0.91	51.11±1.57	Diended
	$ 27_{5,23} - 27_{3,24} \text{ AE}$					4
223512.409	16 _{8,9} – 15 _{7,8} EE	4.37 ± 0.37	7.29 ± 0.84	2.96 ± 0.86	14.20 ± 1.33	Possibly blended ⁴
223621.690	16 _{8,9} – 15 _{7,8} AA	3.56 ± 0.37	7.16 ± 0.84	2.12 ± 0.90	8.56 ± 1.32	Blended ⁵
223684.608	17 _{6,11} – 16 _{7,10} EA	485+037	773+092	203+089	10 32+1 67	Blended ⁶
223684.610	$17_{6,11} - 16_{7,10} \text{ AE}$	1.05±0.57	1.13±0.92	2.03±0.07	10.32±1.07	Diended
223692.004	$17_{7,11} - 16_{6,10}$ EA	4.95 ± 0.37	7.58+0.92	3.00 ± 0.84	16.18+1.69	Partially blended ⁷
223692.104	$ 17_{7,11} - 16_{6,10} \text{ AE}$			010020101	10110_1109	i urtiurij bionaca
223732.826	12 _{9,4} – 11 _{8,4} EE	2.11 ± 0.37	7.18 ± 0.67	4.37 ± 0.81	9.85 ± 2.12	Blended ⁸
223767.585	$17_{6,11} - 16_{7,10} \text{ EE}$	11.61 ± 0.37	7.53 ± 0.84	2.92 ± 0.85	35.27 ± 1.79	Blended ⁹
223775.253	17 _{7,11} – 16 _{6,10} EE	5.88 ± 0.37	7.75 ± 0.84	3.58 ± 0.98	19.99 ± 2.25	Possibly blended ¹⁰
223850.417	17 _{6,11} – 16 _{7,10} AA	3.84 ± 0.37	7.13 ± 0.84	3.30 ± 0.84	14.13 ± 1.32	Slightly blended ¹¹
223858.308	17 _{7,11} – 16 _{6,10} AA	2.48 ± 0.37	7.66 ± 0.84	3.31 ± 0.94	8.35 ± 1.42	Possibly blended ¹²
225744.082	19 _{4,15} – 18 _{5,14} EE	519+043	7 54+0 42	247+042	13 63+1 97	
	19 _{5,15} – 18 _{4,14} EE	5.17±0.15	7.51±0.12	2.17±0.12	15.05±1.97	10
225811.979	13 _{9,5} – 12 _{8,4} EE	1.81 ± 0.34	8.46 ± 0.42	2.99 ± 0.47	5.76 ± 0.64	Possibly blended ¹³
228979.750	$12_{10,2} - 11_{9,2}$ EA					Blended ¹⁴
229033.736	$22_{1,21} - 21_{2,20}$ AE					
	$22_{2,21} - 21_{1,20}$ AE	4.47 ± 0.64	7.38 ± 0.45	2.80 ± 0.41	13.33 ± 1.26	
229033.771	$22_{1,21} - 21_{1,20}$ EA	1.17 ±0.01	7.50±0.15	2.00±0.11	10.00±1.20	
	$ 22_{2,21} - 21_{2,20} \text{ EA}$					
229041.826	$12_{10,3} - 11_{9,3} \text{ EE}$	3.01 ± 0.62	8.46 ± 0.53	3.30 ± 0.41	10.55 ± 1.38	
229055.797	$22_{1,21} - 21_{2,20} \text{ EE}$					
	$22_{1,21} - 21_{1,20} \text{ EE}$	6.92 ± 0.90	7.47 ± 0.44	3.07 ± 0.52	22.58 ± 2.68	
	$22_{2,21} - 21_{2,20} \text{ EE}$					
220059 040	$22_{2,21} - 21_{1,20} \text{ EE}$	2.02 . 0.80	7 27 1 0 50	2.00 ± 0.41	0.21 + 2.05	
229058.049	$14_{9,6} - 13_{8,5} \text{ EE}$	2.92±0.80	/.3/±0.59	3.00 ± 0.41	9.31±2.05	
229077.788	$22_{1,21} - 21_{2,20}$ AA	3.67 ± 0.39	7.06 ± 0.42	4.04 ± 0.45	15.78 ± 0.80	
	222,21 - 211,20 AA					

Notes. Acetone spectroscopic data are taken from the JPL database. Upper-state energy E_{up}/k , effective line strength $S\mu^2$, and spin statistical weight g_s are also given. ⁽¹⁾ Partially blended with the NH₂CHO 18_{2,16} – 18_{2,17} line at 101293.3 MHz. ⁽²⁾ Partially blended with the H₂CS 3_{1,3} – 2_{1,2} line at 101477.8 MHz. ⁽³⁾ Blended with the NH₂CHO 10_{1,10} – 9_{1,9} line at 203335.8 MHz. ⁽⁴⁾ Possibly blended with the CH₃CH₂¹³CN 25_{4,22} – 24_{4,21} line at 223512.8 MHz. ⁽⁵⁾ Partially blended with the HCOOCH₃ 37_{8,30} – 37_{7,31} E line at 223624.5 MHz. ⁽⁶⁾ Blended with the U223686.7 line. ⁽⁷⁾ Partially blended with the U223694.8 line. ⁽⁸⁾ The emission peaks at about 7 km s⁻¹, which is likely be blended with the U223733.6 line. ⁽⁹⁾ Blended with the U223767.6 line. ⁽¹⁰⁾ Possibly blended with the ¹³CH₃CH₂CN 26_{2,25} – 25_{2,24} line at 223773.5 MHz. ⁽¹¹⁾ Slightly blended with the HCOOCH₃ 35_{7,29} – 35_{6,30} A line at 223854.2 MHz. ⁽¹²⁾ Possibly blended with the ¹³CH₃CH₂CN 27_{2,26} – 27_{0,27} line at 225810.7 MHz. ⁽¹⁴⁾ Blended with the U228979.0 Line.

Table A.2.	$(CH_3)_2CO$	measurements	toward	Ace-2
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Frequency	Transition	T _B	V _{LSR}	ΔV	$\int T_{\rm B} dV$	Comment
(MHz)	(J_{k_a,k_c})	(K)	$({\rm km}~{\rm s}^{-1})$	$({\rm km}~{\rm s}^{-1})$	$(\check{K} km s^{-1})$	
101297.450	24 _{13,12} - 24 _{12,13} EE	0.49 ± 0.05	7.31±1.85	5.55±1.85	3.28±0.50	Partially blended ¹
101426.664	$9_{1,8} - 8_{2,7} \text{ AE}$					•
101426.759	$9_{1,8} - 8_{2,7} \text{ EA}$	1 26 1 0 05	6 92 1 95	6 20 + 1 86	9 59 10 40	
101427.041	$9_{2,8} - 8_{1,7}$ AE	1.20 ± 0.03	0.82±1.83	0.29±1.60	0.30±0.40	
101427.130	9 _{2,8} – 8 _{1,7} EA					
101451.059	9 _{1,8} – 8 _{2,7} EE	251 ± 0.05	6 80+1 85	6 56+1 85	17.44 ± 0.40	
101451.446	$9_{2,8} - 8_{1,7} \text{ EE}$	2.51±0.05	0.09±1.05	0.30±1.85	17.44±0.49	
101475.332	9 _{1,8} – 8 _{2,7} AA	1 69+0 05	6 59+1 85	5 55+1 85	8 78+1 49	Partially blended ²
101475.734	$9_{2,8} - 8_{1,7}$ AA	1.07±0.05	0.57±1.05	5.55±1.05	0.70±1.49	I artially blended
203336.247	$27_{4,23} - 27_{3,24}$ EA					
	27 _{5,23} – 27 _{4,24} EA	1.78 ± 1.19	8.27 ± 0.92	4.93 ± 0.94	9.35 ± 5.74	Blended ³
203336.291	$27_{4,23} - 27_{4,24}$ AE	11/021119	0127 2017 2		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Dienaeu
	$ 27_{5,23} - 27_{3,24} \text{ AE}$					~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~
223512.409	16 _{8,9} – 15 _{7,8} EE	< 2.5	6.50 ± 0.84		< 16.0	Possibly blended ⁴
223621.690	16 _{8,9} – 15 _{7,8} AA	1.88 ± 0.37	7.16 ± 0.84	2.36 ± 0.86	4.91 ± 0.86	Partially blended ³
223684.608	$17_{6,11} - 16_{7,10}$ EA	< 3.1			< 14.4	Blended ⁶
223684.610	$17_{6,11} - 16_{7,10} \text{ AE}$					Divideo
223692.004	$17_{7,11} - 16_{6,10} \text{ EA}$	1.99 ± 0.37	8.42 ± 0.84		< 6.4	Partially Blended ⁷
223692.104	$ 1/_{7,11} - 16_{6,10} \text{ AE}$.0.4			1.0	DI 1.18
223732.826	$12_{9,4} - 11_{8,4} \text{ EE}$	< 0.4			< 1.2	Blended ^o
223767.585	$1/_{6,11} - 16_{7,10} EE$	3.50±0.37	8.36±0.84	5.10 ± 0.91	18.94 ± 2.14	Blended ²
223775.253	$17_{7,11} - 16_{6,10} \text{ EE}$	4.34±0.37	7.75±0.84	4.54±0.89	21.8/±1.98	Possibly blended ¹⁰
223850.417	$17_{6,11} - 16_{7,10}$ AA	1.58 ± 0.37	7.13±0.84	4.39 ± 0.94	7.85±1.63	Blended ¹¹
223858.308	$17_{7,11} - 16_{6,10}$ AA	1.58 ± 0.37	7.66 ± 0.84	3.67 ± 0.87	6.06 ± 1.39	Possibly blended ¹²
225744.082	$19_{4,15} - 18_{5,14} \text{ EE}$	2.04 ± 0.46	7.35 ± 0.43	3.71 ± 0.48	8.08 ± 2.99	
005011.070	$19_{5,15} - 18_{4,14} EE$	0.00.000	0.12.0.45	2.00 . 0.41	0 (0 . 0 . 50	D 11 11 1 113
225811.979	$13_{9,5} - 12_{8,4} \text{ EE}$	0.82 ± 0.29	9.13 ± 0.45	3.00 ± 0.41	2.63 ± 0.59	Possibly blended ¹³
228979.750	$12_{10,2} - 11_{9,2} \text{ EA}$					Blended ¹⁴
229033.736	$22_{1,21} - 21_{2,20}$ AE					
220022 771	$22_{2,21} - 21_{1,20}$ AE	< 1.2			< 3.6	
229055.771	$22_{1,21} - 21_{1,20} EA$					
220041 826	$ 22_{2,21} - 21_{2,20} EA$	< 0.0			- 27	
229041.820	$12_{10,3} - 11_{9,3}$ EE	< 0.9	•••	•••	< 2.7	
229033.191	$22_{1,21} = 21_{2,20}$ EE 221 21 = 211 20 FF					
	$22_{1,21}$ $21_{1,20}$ EE $22_{2,21} - 21_{2,20}$ FE	2.70 ± 0.47	8.01 ± 0.45	3.00 ± 0.41	8.62 ± 0.78	
	$22_{2,21} - 21_{2,20} EE$					
229058.049	$14_{0.6} - 13_{8.5} EE$	2.66+0.46	7.72 ± 0.45	3.00 ± 0.41	8.49+0.75	
229077.788	$ 22_{121} - 21_{220} AA$	1.50 0.16	7.5220.13	5.00 20.11	0.00.00	
	$22_{2,21} - 21_{1,20}$ AA	1.58±0.16	/.50±0.42	5.18 ± 0.48	8.69±0.65	

Notes. Acetone spectroscopic data are taken from the JPL database. Upper-state energy E_{up}/k , effective line strength $S\mu^2$, and spin statistical weight g_s are also given. ⁽¹⁾ Partially blended with the NH₂CHO 18_{2,16} – 18_{2,17} line at 101293.3 MHz. ⁽²⁾ Partially blended with the H₂CS 3_{1,3} – 2_{1,2} line at 101477.8 MHz. ⁽³⁾ Blended with the NH₂CHO 10_{1,10} – 9_{1,9} line at 203335.8 MHz. ⁽⁴⁾ Possibly blended with the CH₃CH₂¹³CN 25_{4,22} – 24_{4,21} line at 223512.8 MHz. ⁽⁵⁾ Partially blended with the HCOOCH₃ 37_{8,30} – 37_{7,31} E line at 223624.5 MHz. ⁽⁶⁾ Blended with the strong U223680.0 line. ⁽⁷⁾ Partially blended with the U223694.8 line. ⁽⁸⁾ Blended with the U223733.6 line. ⁽⁹⁾ Blended with the U223767.6 line. ⁽¹⁰⁾ Possibly blended with the ¹³CH₃CH₂CN 26_{2,25} – 25_{2,24} line at 223773.5 MHz. ⁽¹¹⁾ Blended with the HCOOCH₃ 35_{7,29} – 35_{6,30} A line at 223854.2 MHz. ⁽¹²⁾ Possibly blended with the ¹³CH₃CH₂CN 14_{2,12} – 13_{1,13} line at 223859.3 MHz, similar to the 223775.3 MHz (CH₃)₂CO line. ⁽¹³⁾ Possibly blended with the CH₃¹³CH₂CN 27_{2,26} – 27_{0,27} line at 225810.7 MHz. ⁽¹⁴⁾ Blended with the U228979.0 Line.

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Frequency	Transition	T _B	$V_{\rm LSR}$	ΔV	$\int T_{\rm B} dV$	Comment
(MHz)	(J_{k_a,k_c})	(K)	$({\rm km}~{\rm s}^{-1})$	$({\rm km}~{\rm s}^{-1})$	$(\check{K} km s^{-1})$	
101297.450	$24_{13,12} - 24_{12,13}$ EE	0.20 ± 0.05	7.31±1.85	5.55 ± 1.85	1.43 ± 0.63	
101426.664	$9_{1.8} - 8_{2.7} \text{ AE}$					
101426.759	$9_{1,8} - 8_{2,7} \text{ EA}$	0.72+0.05	6 97 1 95	7 10 1 96	5 78 10 16	
101427.041	$9_{2,8} - 8_{1,7}$ AE	0.72 ± 0.03	0.82±1.83	/.10±1.80	3.78 ± 0.40	
101427.130	$9_{2,8} - 8_{1,7} \text{ EA}$					
101451.059	$9_{1,8} - 8_{2,7} \text{ EE}$	1 47+0.05	6 80+1 85	5 04+1 85	0 70±0 38	
101451.446	$9_{2,8} - 8_{1,7} \text{ EE}$	1.47 ± 0.03	0.09±1.05	J.94±1.6J	9.79±0.38	
101475.332	9 _{1,8} – 8 _{2,7} AA	1.72 ± 0.05	6 50+1 85	5 55+1 85	0.63±0.48	Dortiolly blanded
101475.734	9 _{2,8} – 8 _{1,7} AA	1.72±0.05	0.39±1.65	J.JJ±1.6J	9.03±0.48	r artially bielided
203336.247	27 _{4,23} – 27 _{3,24} EA					
	27 _{5,23} – 27 _{4,24} EA	<12			~? ?	
203336.291	27 _{4,23} – 27 _{4,24} AE	N1.2	•••	•••	\2.2	
	27 _{5,23} – 27 _{3,24} AE					
223512.409	16 _{8,9} – 15 _{7,8} EE	< 0.4			< 1.2	
223621.690	16 _{8,9} – 15 _{7,8} AA	< 0.4			< 1.2	
223684.608	17 _{6,11} – 16 _{7,10} EA	< 0.4			< 1.2	blended ²
223684.610	17 _{6,11} – 16 _{7,10} AE	< 0.1	•••		< 1. <u>2</u>	olended
223692.004	17 _{7,11} – 16 _{6,10} EA	< 0.4			< 1.2	
223692.104	17 _{7,11} – 16 _{6,10} AE	< 0.1	•••	•••	< 1. <u>–</u>	2
223732.826	12 _{9,4} – 11 _{8,4} EE	< 0.5			< 1.3	Possibly blended ³
223767.585	17 _{6,11} – 16 _{7,10} EE	1.37 ± 0.35	5.02 ± 0.84	1.73 ± 0.86	2.66 ± 0.62	first component
		1.19 ± 0.35	10.00 ± 0.84	1.65 ± 0.86	2.23 ± 0.59	second component
223775.253	17 _{7,11} – 16 _{6,10} EE	1.29 ± 0.35	6.07 ± 0.84	1.51 ± 0.86	2.08 ± 0.55	first component
		0.83 ± 0.35	10.26 ± 0.84	2.86 ± 0.93	2.25 ± 1.00	second component ⁴
223850.417	17 _{6,11} – 16 _{7,10} AA	< 0.4			< 1.2	
223858.308	17 _{7,11} – 16 _{6,10} AA	< 0.4	•••		< 1.2	
225744.082	19 _{4,15} – 18 _{5,14} EE	< 1.2			< 3.6	
	19 _{5,15} – 18 _{4,14} EE					
225811.979	$13_{9,5} - 12_{8,4} \text{ EE}$	< 0.2	•••		< 0.6	5
228979.750	$12_{10,2} - 11_{9,2}$ EA	•••	•••		•••	Blended ⁵
229033.736	$22_{1,21} - 21_{2,20}$ AE					
	$22_{2,21} - 21_{1,20}$ AE	1.04 ± 0.33	7.18 ± 0.45	3.20 ± 0.41	3.54 ± 0.53	
229033.771	$22_{1,21} - 21_{1,20} \text{ EA}$					
000041.000	$22_{2,21} - 21_{2,20} \text{ EA}$	0.5				
229041.826	$12_{10,3} - 11_{9,3} EE$	< 0.5			< 1.5	
229055.797	$22_{1,21} - 21_{2,20} \text{ EE}$					
	$22_{1,21} - 21_{1,20} \text{ EE}$	< 1.3			< 3.9	
	$22_{2,21} - 21_{2,20} \text{ EE}$					
220059.040	$22_{2,21} - 21_{1,20} \text{ EE}$. 0.0				
229058.049	$14_{9,6} - 13_{8,5} \text{ EE}$	< 0.9	•••		< 2.7	
229077.788	$22_{1,21} - 21_{2,20}$ AA	1.24 ± 0.39	7.75 ± 0.56	3.00 ± 0.41	3.94 ± 0.59	
	$\perp 222 21 - 211 20 \text{ AA}$					

Table A.3. (CH₃)₂CO measurements toward Ace-3

Notes. Acetone spectroscopic data are taken from the JPL database. Upper-state energy E_{up}/k , effective line strength $S\mu^2$, and spin statistical weight g_s are also given. The two velocity components in Ace-3 may be due to an incomplete uv coverage so were measured separately. ⁽¹⁾ Partially blended with the H₂CS $3_{1,3} - 2_{1,2}$ line at 101477.8 MHz. ⁽²⁾ Blended with the U223686.7 line. ⁽³⁾ Possibly blended with the U223733.6 line. ⁽⁴⁾ Possible blended with the ¹³CH₃CH₂CN $26_{2,25} - 25_{2,24}$ line at 223773.5 MHz. ⁽⁵⁾ Blended with the U228979.0 Line.

Table A.4.	(CH ₃) ₂ CO	measurements	toward	HC
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Frequency	Transition	T _B	V _{LSR}	ΔV	$\int T_{\rm B} dV$	Comment
(MHz)	(J_{k_a,k_c})	(K)	$({\rm km}~{\rm s}^{-1})$	$({\rm km}~{\rm s}^{-1})$	$(\check{K} \text{ km s}^{-1})$	
101297.450	$24_{13,12} - 24_{12,13}$ EE	0.42 ± 0.05	7.31±1.85	5.55 ± 1.85	1.43±0.63	Partially blended ¹
101426.664	$9_{1,8} - 8_{2,7} \text{ AE}$					·
101426.759	9 _{1,8} – 8 _{2,7} EA	0.78 + 0.05	6 92 1 95	6 21 + 1 97	2 62 1 0 52	
101427.041	$9_{2,8} - 8_{1,7}$ AE	0.78 ± 0.03	0.82±1.83	0.21±1.87	2.02 ± 0.33	
101427.130	9 _{2,8} – 8 _{1,7} EA					
101451.059	$9_{1,8} - 8_{2,7} \text{ EE}$	176+0.05	6 90 1 95	6 10 1 1 05	11.02 + 0.42	
101451.446	9 _{2,8} – 8 _{1,7} EE	1.70±0.05	0.09±1.05	0.10±1.05	11.95±0.42	
101475.332	$9_{1,8} - 8_{2,7}$ AA	1.15 ± 0.05	6 59+1 85	5 55+1 85	5 07+0 86	Partially blended ²
101475.734	9 _{2,8} – 8 _{1,7} AA	1.15±0.05	0.37±1.03	5.55±1.05	5.07±0.00	I artially biclided
203336.247	27 _{4,23} – 27 _{3,24} EA					
	27 _{5,23} – 27 _{4,24} EA	670 ± 119	735+092	498 ± 094	44 73+6 04	Blended ³
203336.291	27 _{4,23} – 27 _{4,24} AE	0.70±1.17	1.55±0.72	4.70±0.74	++. <i>15</i> ±0.0+	Dicilaca
	$27_{5,23} - 27_{3,24}$ AE					
223512.409	16 _{8,9} – 15 _{7,8} EE	2.03 ± 0.35	4.77 ± 0.84		< 12.7	Possibly blended ⁴
223621.690	16 _{8,9} – 15 _{7,8} AA	2.37 ± 0.37	6.32 ± 0.84	4.45 ± 0.92	10.53 ± 1.70	
223684.608	$17_{6,11} - 16_{7,10} \text{ EA}$	< 5.4			< 16.2	blended ⁶
223684.610	$17_{6,11} - 16_{7,10} \text{ AE}$		•••		10.2	biended
223692.004	$17_{7,11} - 16_{6,10}$ EA	2.07 ± 0.37	5.07 ± 1.85		< 14.7	
223692.104	$17_{7,11} - 16_{6,10} \text{ AE}$					7
223732.826	$12_{9,4} - 11_{8,4} \text{ EE}$	< 1.1			< 3.3	blended'
223767.585	$17_{6,11} - 16_{7,10} \text{ EE}$	2.77 ± 0.37	5.02 ± 0.84	5.38 ± 0.98	15.43 ± 2.18	
223775.253	$17_{7,11} - 16_{6,10} \text{ EE}$	2.60 ± 0.37	6.91±0.84	7.31 ± 1.04	18.56 ± 2.77	Possibly blended ^o
223850.417	$17_{6,11} - 16_{7,10}$ AA	0.82 ± 0.37	6.30 ± 0.84	6.40 ± 1.08	5.18 ± 2.19	
223858.308	$17_{7,11} - 16_{6,10}$ AA	0.81 ± 0.37	5.99 ± 0.84	4.95 ± 0.97	4.04 ± 1.70	Possibly blended ⁹
225744.082	$19_{4,15} - 18_{5,14} \text{ EE}$	< 2.0			< 10.0	
	$19_{5,15} - 18_{4,14} \text{ EE}$					
225811.979	$13_{9,5} - 12_{8,4} \text{ EE}$	< 0.4			< 1.2	Possibly blended ¹⁰
228979.750	$12_{10,2} - 11_{9,2} \text{ EA}$					Blended ¹¹
229033.736	$22_{1,21} - 21_{2,20}$ AE					
	$22_{2,21} - 21_{1,20}$ AE	2.72 ± 0.63	7.74 ± 0.41	3.20 ± 0.41	9.27 ± 1.38	
229033.771	$22_{1,21} - 21_{1,20} \text{ EA}$					
220041.026	$ 22_{2,21} - 21_{2,20} \text{ EA} $	1.00.0.00	0.10.0.00	2 2 0 0 1 1	(75 1 20	
229041.826	$12_{10,3} - 11_{9,3} EE$	1.98 ± 0.63	8.13±0.69	3.20 ± 0.41	6.75±1.38	
229055.797	$22_{1,21} - 21_{2,20} \text{ EE}$					
	$22_{1,21} - 21_{1,20} \text{ EE}$	4.50 ± 0.52	7.52 ± 0.43	3.00 ± 0.41	14.37 ± 0.93	
	$22_{2,21} - 21_{2,20} \text{ EE}$					
220058 040	$22_{2,21} - 21_{1,20} EE$	2 02 10 52	7 25 10 46	2.00 ± 0.41	0.67 + 0.02	
229038.049	149,6 - 138,5 EE	3.03 ± 0.32	1.23±0.40	3.00 ± 0.41	9.07±0.92	
229077.788	$22_{1,21} - 21_{2,20}$ AA	2.19 ± 0.47	6.90 ± 0.42	4.38 ± 0.45	10.21 ± 1.14	
	222,21 - 211,20 AA					

Notes. Acetone spectroscopic data are taken from the JPL database. Upper-state energy E_{up}/k , effective line strength $S\mu^2$, and spin statistical weight g_s are also given. ⁽¹⁾ Partially blended with the NH₂CHO 18_{2,16} – 18_{2,17} line at 101293.3 MHz. ⁽²⁾ Partially blended with the H₂CS 3_{1,3} – 2_{1,2} line at 101477.8 MHz. ⁽³⁾ Blended with the NH₂CHO 10_{1,10} – 9_{1,9} line at 203335.8 MHz. ⁽⁴⁾ Possibly blended with the CH₃CH₂¹³CN 25_{4,22} – 24_{4,21} line at 223512.8 MHz. ⁽⁵⁾ Partially blended with the HCOOCH₃ 37_{8,30} – 37_{7,31} E line at 223624.5 MHz. ⁽⁶⁾ Blended with the strong U223680.0 line. ⁽⁷⁾ Blended with the U223733.6 line. ⁽⁸⁾ Possibly blended with the ¹³CH₃CH₂CN 26_{2,25} – 25_{2,24} line at 223773.5 MHz. ⁽⁹⁾ Possibly blended with the ¹³CH₃CH₂CN 14_{2,12} – 13_{1,13} line at 223859.3 MHz, similar to 223775.3 MHz (CH₃)₂CO line. ⁽¹⁰⁾ Possibly blended with the CH₃¹³CH₂CN 27_{2,26} – 27_{0,27} line at 225810.7 MHz. ⁽¹¹⁾ Blended with the U228979.0 Line.



Fig. A.3. (CH₃)₂CO spectra near 223 GHz toward four positions (three acetone emission peaks and HC) in the Orion BN/KL region. Dashed lines indicate the V_{LSR} of 8 km s⁻¹ and correspond to the (CH₃)₂CO line shown at the top of each panel. (c) The dotted line indicates the (CH₃)₂CO 17_{6,11} – 16_{7,10} EA/AE lines at 223684.6 MHz. (d) The dotted line indicates the (CH₃)₂CO 17_{6,11} – 16_{7,10} EE line at 223767.6 MHz. (e) The dotted line indicates the (CH₃)₂CO 17_{6,11} – 16_{7,10} EI line at 223767.6 MHz. (e) The dotted line indicates the (CH₃)₂CO 17_{6,11} – 16_{7,10} EI line at 223767.6 MHz. (e) The dotted line indicates the (CH₃)₂CO 17_{6,11} – 16_{7,10} EI line at 223767.6 MHz. (e) The dotted line indicates the (CH₃)₂CO 17_{6,11} – 16_{7,10} EI line at 223767.6 MHz. (e) The dotted line indicates the (CH₃)₂CO 17_{6,11} – 16_{7,10} EI line at 223767.6 MHz. (f) The dotted line indicates the (CH₃)₂CO 17_{6,11} – 16_{7,10} EI line at 223767.6 MHz. (f) The dotted line indicates the (CH₃)₂CO 17_{6,11} – 16_{7,10} EI line at 223767.6 MHz. (f) The dotted line indicates the (CH₃)₂CO 17_{6,11} – 16_{7,10} EI line at 223767.6 MHz. (f) The dotted line indicates the (CH₃)₂CO 17_{6,11} – 16_{7,10} A line at 223850.4 MHz.

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Fig. A.4. $(CH_3)_2CO$ spectra near 225 and 229 GHz toward four positions (three acetone emission peaks and HC) in the Orion BN/KL region. Dashed lines indicate the V_{LSR} of 8 km s⁻¹ and correspond to the $(CH_3)_2CO$ line shown at the top of each panel. (c) The dotted line indicates the $(CH_3)_2CO$ 22_{1,21} – 21_{2,20} AE/EA and 22_{2,21} – 21_{1,20} AE/EA lines at 229033.8 MHz. (d) The dotted line indicates the $(CH_3)_2CO$ 14_{9,6} – 13_{8,5} EE line at 229058.0 MHz.

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Appendix B: Comparison with the ALMA science verification data

in Figure B.2, and the spectrum comparison around 223 GHz toward different positions is shown in Figures B.3-B.5. The missing flux of the ALMA-SV data is estimated to be 10–20% compared with the IRAM 30m single-dish data (J. Cernicharo, priv. comm.) around 223 GHz, less than our PdBI data (20-50% missing flux).

Spatial comparison

Figure B.1 shows the uv coverages of the PdBI and ALMA-SV data. They are comparable, but the ALMA-SV data show a denser coverage for shorter spacings. The PdBI data have longer baselines, which result in a higher angular resolution. However, due to the few uv tracks in the -50 to 50 m domain, the PdBI observations are likely to filter out larger structures in BN/KL. This is clearly seen in Figure B.2 in which most of the emission toward Ace-3 and in the southern part of BN/KL close to MF-1 and MF-3 is filtered out in the PdBI data. Likewise, it is important to note that the acetone emission associated with the arc-like structure in the north of HC is not resolved out by the PdBI.

Basic observational parameters

Spectral comparison

Figure B.3 shows the ALMA-SV and PdBI spectra toward Ace-1 and 2. The differences between the two data sets are small for the HCOOCH₃ and $(CH_3)_2CO$ weak lines, but greater for the C_2H_5CN and SO_2 strong or broad lines. It is because these lines tend to be spatially extended and are easily resolved out by the PdBI. Filtering effects are more obvious toward Ace-3 and MF-3 (Figs. B.4 and B.5) where most of the line emission, including SO_2 , C_2H_5CN , and $(CH_3)_2CO$, is filtered out.

Summary

The calibrated ALMA-SV data were retrieved from the ESO archive. The observations were carried out in 2012 January with 16 12-m antennas (see e.g., Zapata et al., 2012; Niederhofer et al., 2012, for more observational details). The primary beam at 223 GHz is about 28". Some observational parameters of the ALMA-SV and our PdBI data are listed in Table B.1. The ALMA-SV data were analyzed with the GILDAS package, and the acetone emission images were cleaned with the same Clark algorithm as for the PdBI data to minimize the effects contributed by different cleaning algorithms. The ALMA and PdBI channel maps of the acetone $17_{7,11} - 16_{6,10}$ EE line are shown

Our PdBI spatial distribution and individual spectra are very consistent with the new ALMA-SV data toward Orion BN/KL. However, due to different uv coverages and the smaller PdBI primary beam, some extended emission is filtered out in our data, especially toward Ace-3 and in the southern part (close to MF-3) of BN/KL. It is not clear why the arc-like acetone emission close to HC has not been resolved out. One possible reason is that the acetone emission in the southern part is weaker and suffers more easily from filtering effects, as observed toward the Ace-3 region. The fully functional ALMA in the near future with a denser uv coverage and the short spacings of the Atacama Compact Array will surely help solve these issues and greatly improve the molecular line images.

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Table B.1. Comparison of the ALMA-S'	/ and PdBI observational	parameters at 223.7 GHz
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Data	Antenna number	Integration time ^a	Flux conversion	RMS noise	$\theta_{\mathrm{HPBW}}{}^{b}$	δv^c	$\theta_{\rm syn}^{d}$	
		(min)	(1 Jy beam^{-1})	(mJy beam ⁻¹)	('')	$({\rm km}~{\rm s}^{-1})$	(''×'')	PA (°)
PdBI	5	~ 150	17.3 K	21.2	22.5	0.84	1.79×0.79	14
ALMA-SV	12	~ 100	9.2 K	9.6	28.2	0.65	1.92×1.39	166

Notes. ^(a) On-source integration time ^(b) Primary beam size ^(c) Channel separation ^(d) Synthesized beam size



Fig. B.1. Comparison of the PdBI and ALMA-SV uv coverages.



Fig. B.2. Comparison of the PdBI (black) and ALMA-SV (green) channel maps of the acetone $17_{7,11} - 16_{6,10}$ EE line ($E_{up}/k=110.7$ K) at 223775.3 MHz. Contours run from 0.6 to 8.1 K in steps of 0.9 K, and the dashed contours represent -0.6 K. The bottom-right panel shows the integrated intensity (from 5 to 11 km s⁻¹) in contours running from 15% to 95% in steps of 10% of their peak temperatures, and the dashed contours represent -10% of their peak temperatures. The primary beams of the ALMA-SV and PdBI data are indicated. The black square marks the center of explosion according to Zapata et al. (2009). The positions of source BN, HC, IRc6/7, and source *I* are marked as triangles. The positions of acetone emission peaks (Ace-1 to Ace-3) are marked as stars.

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Fig. B.3. Comparison of the PdBI and ALMA-SV spectra near 223 GHz toward Ace-1 and Ace-2.



Fig. B.4. Comparison of the PdBI and ALMA-SV spectra near 223 GHz toward Ace-3 and HC.

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Fig. B.5. Comparison of the PdBI and ALMA-SV spectra near 223 GHz toward MF-1 and MF-3. MF-3, a more extended region, suffers strong filtering effects on strong broader lines, such as SO_2 and C_2H_5CN in the PdBI data similar to Ace-3. Some strong lines from other molecules appear at the similar frequencies to acetone in MF-1 and MF-3.

Appendix C: Three-dimensional view of molecular structures in BN/KL

The three-dimensional visualization enables us to investigate the distributions and possible structures of different molecules in great detail. Therefore, we have prepared the three-dimensional (R.A., Dec., and $V_{\rm LSR}$) images of different molecules in Orion BN/KL using the 3D Slicer program⁷ and the complemented procedures provided by the Astronomical Medicine Project ⁸. Figures C.1-C.4 show the different viewing angles of the C₂H₅CN, (CH₃)₂CO, and HCOOCH₃ structures in BN/KL. In addition, the superposed images of these molecules were also plotted. In addition, a complementary 3D movie will be available on the A&A website.

C_2H_5CN

The C₂H₅CN line has a larger line width compared with those of (CH₃)₂CO and HCOOCH₃ and is clearly seen in Figures C.3 and C.4. The arc-like structure close to HC is clearer in the 3D images (e.g., Figs. C.1 and C.2). In addition, the slow outflow associated with IRc7 in the NH₃ emission proposed by Goddi et al. (2011b) may actually be the two different clumps in the SE and NW of IRc7 (see Figs. C.3 and C.4) because of the multicomponent structure in the blueshifted clump revealed in the image. This complicated blueshifted clump was detected in the methyl cyanide emission (Wang et al., 2010) toward southeast of IRc7. These components may be due to the explosive outflow. However, we cannot rule out the possibility of a multi-outflow that comes from IRc7.

$(CH_3)_2CO$

The arc-like structure is clearly seen in the acetone 3D images. It is interesting to note that the filtered-out emission toward Ace-3 is mostly centered at 8 km s⁻¹, which creates an artificial shell-like structure in the 3D images.

HCOOCH₃

Part of the arc-like structure seen in C_2H_5CN and $(CH_3)_2CO$ is also seen in HCOOCH₃ where the HCOOCH₃ emission extends from MF-3 to Ace-1 and even to the northwest of HC (Figs. C.1 and C.2). In addition, some structures close to MF-1 are seen and may be related to outflows/shocks.

⁷ http://www.slicer.org

⁸ http://am.iic.harvard.edu



Fig. C.1. Three-dimensional view (R.A., Dec., and V_{LSR}) of C₂H₅CN (red), (CH₃)₂CO (green), and HCOOCH₃ (blue) distributions in Orion BN/KL. The C₂H₅CN 25_{3,23} – 24_{3,22} line at 223553.6 MHz is shown in red surfaces, the (CH₃)₂CO 17_{7,11} – 16_{6,10} EE line at 223775.3 MHz in green, and the HCOOCH₃ 11_{4,8} – 10_{3,7} A lines at 223550.5 MHz in blue. The isothermal surfaces are plotted from 15% to 95% in steps of 20% of their peak intensities.



Fig. C.2. Three-dimensional view (R.A., Dec., and V_{LSR}) of C₂H₅CN (red), (CH₃)₂CO (green), and HCOOCH₃ (blue) distributions in Orion BN/KL with a different viewing angle from Fig. C.1.



Fig. C.3. Three-dimensional view (R.A., Dec., and V_{LSR}) of C₂H₅CN (red), (CH₃)₂CO (green), and HCOOCH₃ (blue) distributions in Orion BN/KL with a different viewing angle from Fig. C.1.



Fig. C.4. Three-dimensional view (R.A., Dec., and V_{LSR}) of C₂H₅CN (red), (CH₃)₂CO (green), and HCOOCH₃ (blue) distributions in Orion BN/KL with a different viewing angle from Fig. C.1.