The Complex Organic Molecules of Embedded Protostars at Perseus

Yao-Lun Yang<sup>1, 2</sup>

<sup>1</sup>RIKEN Cluster for Pioneering Research, Wako-shi, Saitama, 351-0106, Japan <sup>2</sup>Department of Astronomy, University of Virginia, Charlottesvile, VA 22904-4235, USA

## 1. INTRODUCTION

Planet formation may start during the embedded phase of star formation. In the scenario where planets form from the embedded disks, resulting in substructures, the chemistry of embedded disks may play a significant role for the chemical composition of the forming planets. In the recent years, observations discover the emission of carbon-chain molecules and complex organic molecules (COMs) toward the center of several embedded protostars, indicating that embedded protostars have developed a complex chemistry at the disk-forming region. If the forming planets inherit the chemistry of embedded disks, the abundance of complex organic molecules may implicate future developments of organics on the planets.

Heavier or more complex molecules, such as cyclic- $C_3H_2$ , SO, and complex organic molecules (COMs), are in the gas phase at the inner protostellar envelope ( $T \gtrsim 100 \,\mathrm{K}$ ), exclusively tracing the properties of the inner envelope where a disk may be forming (Aikawa 2013; Sakai et al. 2014). The kinematics of a rotating infalling envelope has been analyzed with the observations of heavier or more complex molecules, such as  $CH_3OH$  and  $CH_2DOH$  for  $HH\,212$  (Lee et al. 2017), CS for  $IRAS\,04365+2535$  (Sakai et al. 2016) and L483 (Oya et al. 2017), cyclic- $C_3H_2$  for L1527 (Sakai et al. 2014), OCS for  $IRAS\,16293-2422\,A$  (Oya et al. 2016), and methanol and HCOOH for B335 (Imai et al. 2019).

In the review by Herbst & van Dishoeck (2009), complex molecules are defined as carbon-bearing molecules that contain six atoms or more. Saturated complex molecules are rich in hydrogen atoms, often called complex organic molecules (COMs), while the unsaturated complex molecules are lack of hydrogen atoms, mostly in the form of long carbon-chain molecules. While recent observations show several embedded protostars with rich spectra of complex molecules, the occurrence of complex molecules at embedded protostars and its relation-

Corresponding author: Yao-Lun Yang yaolunyang.astro@gmail.com

ship to the star formation process remain poorly understand. Several protostars are rich in COMs but show little emission of long carbon-chain molecules, such as IRAS 16293–2422 (Jørgensen et al. 2016), NGC 1333 IRAS 4A (Bottinelli et al. 2004), B335 (Imai et al. 2016, 2019), and BHR 71 (Yang et al. 2020 accepted); some protostars are rich in long carbon-chain molecules but not in COMs, such as L1527 (Sakai et al. 2010) and IRAS 15398–3359 (Sakai et al. 2009). While the bimodal chemical appearance hints a bimodal evolutionary path, the chemical evolution at the embedded protostars remain ill-constrained as a few protostars show the emission of both COMs and long carbon-chain molecules at different scales, such as L483 (Oya et al. 2017).

The Perseus ALMA Chemistry Survey (PEACHES) aims to provide the statistics on the occurrence of complex molecules at embedded protostars. This program unbiasedly observes 51 embedded protostars with ALMA around 260 GHz, covering the emission of simple molecules such as CS and  $\rm H^{13}CN$  as well as the emission of complex molecules including CH<sub>3</sub>OH and CH<sub>3</sub>OCHO.

## 2. OBSERVATIONS

### 3. RESULTS

The ALMA image cubes are post-processed to extract 1D spectra for identifying the emission of complex molecules and more in-depth analyses. Because the complex molecules only sublime at the temperature greater than  $\sim\!100\,\mathrm{K}$ , we focus on the spectra toward the continuum sources. Four steps of post-processing reduces the image cubes to 1D spectra, which are summarized below.

• Continuum fitting: We use the CASA task imfit to iteratively fit for continuum sources down to  $5\sigma$  of the residual image within the central 70% of the primary beam size (20"). For Set3-ID09, the fitting uses a threshold of  $4\sigma$  and extends the mask to the entire primary beam as a continuum source is detected toward the edge of the primary beam where the noise is elevated.

 Table 1. PEACHES Sample

Source	Common names	R.A. (J2000)	Decl. (J2000)	$v_{ m lsr}$	Beam	Cont. Size	$T_{ m cont}$	Ref. $(v_{\rm lsr})$
		(hh:mm:ss)	(dd:mm:ss)	$({\rm km\ s^{-1}})$	(")	(")	(K)	
Per-emb 22 B		03:25:22.35	30:45:13.11	4.3	064×039	0."95×0."51	0.92	S19
Per-emb $22 A$		03:25:22.41	30:45:13.26	4.3	$0.64 \times 0.39$	$0.686 \times 0.65$	1.71	S19
L1448 NW	L1448 IRS $3C$	03:25:35.67	30:45:34.16	4.2	$0.64 \times 0.39$	$0.''83 \times 0.''47$	3.15	H18
Per-emb 33 B/C		03:25:36.32	30:45:15.19	5.3	$0.64 \times 0.39$	$0.75 \times 0.48$	5.55	S19
Per-emb 33 A		03:25:36.38	30:45:14.72	5.3	$0.64 \times 0.39$	$0.''73 \times 0.''45$	10.33	S19
L1448 IRS 3A		03:25:36.50	30:45:21.90	4.6	$0.64 \times 0.39$	$0.85 \times 0.59$	3.21	H18
Per-emb 26		03:25:38.88	30:44:05.28	5.4	$0.64 \times 0.39$	$0.''69 \times 0.''45$	8.03	S19
Per-emb 42		03:25:39.14	30:43:57.90	5.8	$0.64 \times 0.39$	064×039	0.66	S19
Per-emb 25	IRAS 03235+3004	03:26:37.51	30:15:27.81	5.5	$0.64 \times 0.39$	$0.69 \times 0.41$	5.27	S18
Per-emb 17	L1455 IRS 1, IRAS 03245+3002	03:27:39.11	30:13:02.96	6.0	$0.64 \times 0.440$	$0.79 \times 0.48$	2.00	S19
Per-emb 20	L1455 IRS 4	03:27:43.28	30:12:28.88	5.3	$0.64 \times 0.440$	$1.29 \times 0.78$	0.14	S19
L1455 IRS 2		03:27:47.69	30:12:04.33	5.1	064×040	0."60×0."38	0.13	H18
Per-emb 35 A	NGC 1333 IRAS 1	03:28:37.10	31:13:30.77	7.4	$0.66 \times 0442$	0".75×0"51	0.93	Y20
Per-emb 35 B	NGC 1333 IRAS 1	03:28:37.22	31:13:31.74	7.3	$0.66 \times 0442$	0".78×0"53	0.75	Y20
Per-emb 27	NGC 1333 IRAS 2A	03:28:55.57	31:14:36.97	6.5	$0.66 \times 0442$	0".93×0".66	5.79	Y20
EDJ2009-172		03:28:56.65	31:18:35.43		$0.66 \times 0442$	0."69×0."44	0.62	
Per-emb 36	NGC 1333 IRAS 2B	03:28:57.37	31:14:15.77	6.9	$0.66 \times 0442$	0".73×0".46	5.56	S19
Per-emb 54	NGC 1333 IRAS 6	03:29:01.55	31:20:20.49	7.9	$0.66 \times 0.42$	0."69×0."40	0.07	S19
SVS 13B	NGC 1333 SVS 13B	03:29:03.08	31:15:51.73	8.5	$0.66 \times 0.42$	0."87×0."68	6.64	S19
SVS 13A2		03:29:03.39	31:16:01.58	8.4	0''66×0''42	0."86×0."53	0.61	S18
Per-emb 44	NGC 1333 SVS 13A	03:29:03.76	31:16:03.70	8.7	0."66×0."42	0."98×0."79	6.84	S19
Per-emb 15		03:29:04.06	31:14:46.23	6.8	0."66×0."42	0."89×0."70	0.17	S19
Per-emb 50	IRAS 03260+3111 A	03:29:07.77	31:21:57.11	9.3	$0.66 \times 0.42$	073×044	4.13	Y20
Per-emb 12 B	NGC 1333 IRAS 4A2	03:29:10.44	31:13:32.08	6.9	0."66×0."42	1."33×0."81	10.04	S19
Per-emb 12 A	NGC 1333 IRAS 4A1	03:29:10.54	31:13:30.93	6.9	$0.66 \times 0.42$	1."11×0."98	21.85	S19
Per-emb 21	NGC 1333 IRAS 7 SM2	03:29:10.67	31:18:20.16	8.6	$0.66 \times 0.42$	074×048	2.05	Y20
Per-emb 18	NGC 1333 IRAS 7 SM1	03:29:11.27	31:18:31.09	8.1	0."66×0."42	0."84×0."73	3.42	S19
Per-emb 13	NGC 1333 IRAS 4B1	03:29:12.02	31:13:07.99	7.1	$0.66 \times 0.42$	1."07×0."83	14.76	S19
IRAS4B'	NGC 1333 IRAS 4B2	03:29:12.85	31:13:06.87	7.1	$0.66 \times 0.42$	0."83×0."74	7.13	S19
Per-emb 14	NGC 1333 IRAS 4C	03:29:13.55	31:13:58.12	7.9	$0.06 \times 0.42$ $0.66 \times 0.42$	$0.03\times0.74$ $0.79\times0.750$	3.05	S19
EDJ2009-235	1100 1000 11010 10	03:29:18.26	31:23:19.73	7.7	$0.65 \times 0.12$ $0.67 \times 0.42$	0."66×0."44	0.26	Y20
EDJ2009-237		03:29:18.74	31:23:25.24		$0.07 \times 0.42$ $0.67 \times 0.42$	$0.00 \times 0.44$ $0.67 \times 0.42$	0.12	
Per-emb 37		03:29:18.97	31:23:14.28	7.5	$0.07 \times 0.42$ $0.67 \times 0.42$	0	0.56	Y20
Per-emb 60		03:29:20.05	31:24:07.35		$0.07 \times 0.42$ $0.67 \times 0.42$	$0.32 \times 0.37$ $0.73 \times 0.747$	0.08	
Per-emb 5	IRAS 03282+3035	03:31:20.94	30:45:30.24	7.3	$0.07 \times 0.42$ $0.45 \times 0.30$	$0.75 \times 0.47$ $0.756 \times 0.741$	15.29	S19
Per-emb 2	IRAS 03292+3039		30:49:47.81		$0.45 \times 0.30$ $0.45 \times 0.30$	$1.35 \times 0.41$ $1.35 \times 0.97$		S19
Per-emb 2 Per-emb 10	B1-d	03:32:17.92	31:06:52.01	7.0	$0.45 \times 0.30$ $0.46 \times 0.30$	$0.49 \times 0.32$	7.41	
Per-emb 40		03:33:16.43		6.4	$0.46 \times 0.30$ $0.46 \times 0.30$	$0.49 \times 0.32$ $0.49 \times 0.32$	1.82	S19
Per-emb 29	B1-a B1-c	03:33:16.67	31:07:54.87	$7.4 \\ 6.1$	$0.46 \times 0.30$ $0.46 \times 0.30$	$0.47 \times 0.32$ $0.56 \times 0.39$	1.44	S19 Y20
B1-b N	В1-С	03:33:17.88	31:09:31.74		$0.46 \times 0.30$ $0.46 \times 0.30$	$0.56 \times 0.39$ $0.56 \times 0.47$	8.41 $7.67$	C16
		03:33:21.21	31:07:43.63	6.6	$0.46 \times 0.30$ $0.46 \times 0.30$			
B1-b S		03:33:21.36 03:43:50.97	31:07:26.34	6.6	$0.46 \times 0.30$ $0.50 \times 0.32$	0."63×0."53 0."61×0."52	14.79	C16
Per-emb 16			32:03:24.12	8.8	$0.50 \times 0.32$ $0.50 \times 0.32$	$0.61 \times 0.52$ $0.56 \times 0.32$	0.35	S19
Per-emb 28	UU 211 MMC	03:43:51.01	32:03:08.02	8.6	$0.50 \times 0.32$ $0.49 \times 0.32$		1.52	S19
Per-emb 1	HH 211 MMS	03:43:56.81	32:00:50.16	9.4		0."68×0."48	4.57	S19
Per-emb 11 B	IC 348 MMS	03:43:56.88	32:03:03.08	9.0	$0.750 \times 0.733$	$0.92 \times 0.69$	0.40	S19
Per-emb 11 A	IC 348 MMS	03:43:57.07	32:03:04.76	9.0	$0.750 \times 0.733$	0."61×0."48	10.47	S19
Per-emb 11 C	IC 348 MMS	03:43:57.70	32:03:09.82	9.0	$0.750 \times 0.733$	1."10×0."86	0.34	S19
Per-emb 55	IRAS 03415+3152	03:44:43.30	32:01:31.22	12.0	0."50×0."32	0."49×0."33	0.32	S19
Per-emb 8		03:44:43.98	32:01:35.19	11.0	0."50×0."32	0."49×0."36	8.51	S19
Per-emb 53	B5 IRS 1	03:47:41.59	32:51:43.62	10.2	$0.751 \times 0.733$	$0.758 \times 0.742$	1.55	Y20

References - C16 = Carney et al. (2016); H18 = Higuchi et al. (2018); S18 = Stephens et al. (2018); S19 = Stephens et al. (2019); Y20 = this study.

 ${\bf Table~2.~PEACHES~Sample}$ 

PEACHES ID	Source	Common names	R.A. (J2000)	Decl. (J2000)	$v_{ m lsr}$	Beam	Cont. Size	$T_{\mathrm{cont}}$	Ref. $(v_{\rm lsr})$
			(hh:mm:ss)	(dd:mm:ss)	$({\rm km\ s^{-1}})$	(")	(")	(K)	
Set1_ID00	L1448 NW	L1448 IRS 3C	03:25:35.67	30:45:34.16	4.2	0."64×0."39	0."83×0."47	3.15	H18
$Set1\_ID01\_3$	Per-emb 33 A		03:25:36.38	30:45:14.72	5.3	064×039	$0.73 \times 0.45$	10.33	S19
$Set1\_ID01\_4$	Per-emb 33 B/C		03:25:36.32	30:45:15.19	5.3	064×039	$0.75 \times 0.48$	5.55	S19
$Set1_ID01_2$	L1448 IRS 3A		03:25:36.50	30:45:21.90	4.6	0."64×0."39	$0.''85 \times 0.''59$	3.21	H18
$Set1\_ID02$	Per-emb 26		03:25:38.88	30:44:05.28	5.4	0."64×0."39	0."69×0."45	8.03	S19
$Set1_ID02_2$	Per-emb 42		03:25:39.14	30:43:57.90	5.8	0."64×0."39	0."64×0."39	0.66	S19
Set1_ID03	Per-emb 22 A		03:25:22.41	30:45:13.26	4.3	0."64×0."39	0."86×0."65	1.71	S19
$Set1_ID03_2$	Per-emb 22 B		03:25:22.35	30:45:13.11	4.3	0."64×0."39	$0.95 \times 0.51$	0.92	S19
$Set1\_ID05$	Per-emb 25	IRAS 03235+3004	03:26:37.51	30:15:27.81	5.5	0."64×0."39	0."69×0."41	5.27	S18
Set1_ID06	Per-emb 17	L1455 IRS 1, IRAS 03245+3002	03:27:39.11	30:13:02.96	6.0	0''64×0''40	0"79×0"48	2.00	S19
$Set1_ID07$	Per-emb 20	L1455 IRS 4	03:27:43.28	30:12:28.88	5.3	0'.'64×0'.'40	1"29×0"78	0.14	S19
Set1_ID08	L1455 IRS 2		03:27:47.69	30:12:04.33	5.1	0'.'64×0'.'40	0''60×0''38	0.13	H18
Set2_ID00	Per-emb 44	NGC 1333 SVS 13A	03:29:03.76	31:16:03.70	8.7	0''66×0''42	0."98×0."79	6.84	S19
$Set2_ID00_2$	SVS 13A2		03:29:03.39	31:16:01.58	8.4	0''66×0''42	0."86×0."53	0.61	S18
Set2_ID01	Per-emb 12 A	NGC 1333 IRAS 4A1	03:29:10.54	31:13:30.93	6.9	$0.66 \times 0442$	1."11×0."98	21.85	S19
Set2_ID01_2	Per-emb 12 B	NGC 1333 IRAS 4A2	03:29:10.44	31:13:32.08	6.9	0''66×0''42	1."33×0."81	10.04	S19
Set2_ID02	Per-emb 13	NGC 1333 IRAS 4B1	03:29:12.02	31:13:07.99	7.1	0."66×0."42	1."07×0."83	14.76	S19
Set2_ID02_2	IRAS4B'	NGC 1333 IRAS 4B2	03:29:12.85	31:13:06.87	7.1	$0.66 \times 0.42$	0."83×0."74	7.13	S19
Set2_ID03	Per-emb 27	NGC 1333 IRAS 2A	03:28:55.57	31:14:36.97	6.5	$0.66 \times 0.42$	0."93×0."66	5.79	Y20
Set2_ID04	Per-emb 54	NGC 1333 IRAS 6	03:29:01.55	31:20:20.49	7.9	$0.66 \times 0.42$	0. 69×0. 40	0.07	S19
Set2_ID04 Set2_ID05	Per-emb 21	NGC 1333 IRAS 7 SM2	03:29:10.67	31:18:20.16	8.6	$0.00 \times 0.42$ $0.66 \times 0.42$	0. 03×0. 40 0. 74×0. 48	2.05	Y20
Set2_ID06	Per-emb 14	NGC 1333 IRAS 4C	03:29:13.55	31:13:58.12	7.9	$0.00 \times 0.42$ $0.66 \times 0.42$	0. 74×0. 40 0. 79×0. 50	3.05	S19
Set2_ID07	Per-emb 35 A	NGC 1333 IRAS 1	03:28:37.10	31:13:30.77	7.4	$0.00 \times 0.42$ $0.66 \times 0.42$	0. 75×0. 50 0. 75×0. 51	0.93	Y20
Set2_ID07_2	Per-emb 35 B	NGC 1333 IRAS 1	03:28:37.22	31:13:31.74	7.3	$0.00 \times 0.42$ $0.66 \times 0.42$	0. 78×0. 51 0. 78×0. 53	0.75	Y20
Set2_ID07_2 Set2_ID08	SVS 13B	NGC 1333 NVS 13B	03:29:03.08	31:15:51.73	8.5	$0.00 \times 0.42$ $0.66 \times 0.42$	0. 78×0. 53 0. 87×0. 68	6.64	S19
Set2_ID08 Set2_ID09	Per-emb 15	NGC 1939 5 V 5 13D	03:29:04.06	31:14:46.23	6.8	$0.00 \times 0.42$ $0.66 \times 0.42$	0.87×0.08 0.89×0.70	0.04	S19
Set2_ID09	Per-emb 50	IRAS 03260+3111 A	03:29:07.77	31:21:57.11	9.3	$0.00 \times 0.42$ $0.66 \times 0.42$	$0.33 \times 0.70$ $0.73 \times 0.744$	4.13	Y20
Set2_ID11	Per-emb 18	NGC 1333 IRAS 7 SM1	03:29:11.27	31:18:31.09	8.1	$0.00 \times 0.42$ $0.66 \times 0.42$	$0.73 \times 0.44$ $0.84 \times 0.73$	3.42	S19
Set2_ID12 Set2_ID13	Per-emb 37	NGC 1333 II(AS 1 SWI	03:29:18.97	31:23:14.28	7.5	$0.00 \times 0.42$ $0.67 \times 0.42$	$0.84 \times 0.73$ $0.82 \times 0.75$	0.56	Y20
	EDJ2009-235				7.5 7.7	$0.67 \times 0.42$ $0.67 \times 0.42$	$0.82 \times 0.37$ $0.66 \times 0.44$	0.26	Y20
Set2_ID13_2			03:29:18.26	31:23:19.73		$0.67 \times 0.42$ $0.67 \times 0.42$	$0.00 \times 0.44$ $0.67 \times 0.42$		
Set2_ID13_3	EDJ2009-237		03:29:18.74	31:23:25.24		$0.67 \times 0.42$ $0.67 \times 0.42$	$0.67 \times 0.42$ $0.73 \times 0.47$	0.12	
Set2_ID14	Per-emb 60 EDJ2009-172		03:29:20.05	31:24:07.35	• • •			0.08	• • •
Set2_ID15		NGG 1000 ID AG AD	03:28:56.65	31:18:35.43		0'.'66×0'.'42	0."69×0."44	0.62	
Set2_ID16	Per-emb 36	NGC 1333 IRAS 2B	03:28:57.37	31:14:15.77	6.9	$0.66 \times 0.42$	0."73×0."46	5.56	S19
Set3_ID00	B1-b S		03:33:21.36	31:07:26.34	6.6	0."46×0."30	0."63×0."53	14.79	C16
Set3_ID00_2	B1-b N	D1	03:33:21.21	31:07:43.63	6.6	0."46×0."30	0."56×0."47	7.67	C16
Set3_ID01	Per-emb 29	B1-c	03:33:17.88	31:09:31.74	6.1	0."46×0."30	$0.756 \times 0.739$	8.41	Y20
Set3_ID02	Per-emb 10	B1-d	03:33:16.43	31:06:52.01	6.4	0."46×0."30	0."49×0."32	1.82	S19
Set3_ID03	Per-emb 40	B1-a	03:33:16.67	31:07:54.87	7.4	046×030	0."47×0."32	1.44	S19
Set3_ID04	Per-emb 2	IRAS 03292+3039	03:32:17.92	30:49:47.81	7.0	0."45×0."30	1."35×0."97	7.41	S19
Set3_ID05	Per-emb 5	IRAS 03282+3035	03:31:20.94	30:45:30.24	7.3	0."45×0."30	0."56×0."41	15.29	S19
Set3_ID06	Per-emb 1	HH 211 MMS	03:43:56.81	32:00:50.16	9.4	0'.'49×0'.'32	068×048	4.57	S19
Set3_ID07	Per-emb 11 A	IC 348 MMS	03:43:57.07	32:03:04.76	9.0	050×033	0."61×0."48	10.47	S19
Set3_ID07_2	Per-emb 11 B	IC 348 MMS	03:43:56.88	32:03:03.08	9.0	0."50×0."33	0"92×0"69	0.40	S19
Set3_ID07_3	Per-emb 11 C	IC 348 MMS	03:43:57.70	32:03:09.82	9.0	0."50×0."33	1"10×0"86	0.34	S19
Set3_ID08	Per-emb 8		03:44:43.98	32:01:35.19	11.0	0."50×0."32	0."49×0."36	8.51	S19
$Set3_ID08_2$	Per-emb 55	IRAS 03415+3152	03:44:43.30	32:01:31.22	12.0	0."50×0."32	0."49×0."33	0.32	S19
$Set3_ID09$	Per-emb 16		03:43:50.97	32:03:24.12	8.8	0."50×0."32	0."61×0."52	0.35	S19
$\mathbf{Set3\_ID09\_2}$	Per-emb 28		03:43:51.01	32:03:08.02	8.6	$0.''50 \times 0.''32$	$0.''56 \times 0.''32$	1.52	S19
Set3_ID10	Per-emb 53	B5 IRS 1	03:47:41.59	32:51:43.62	10.2	$0.51 \times 0.33$	$0.758 \times 0.742$	1.55	Y20

References - C16 = Carney et al. (2016); H18 = Higuchi et al. (2018); S18 = Stephens et al. (2018); S19 = Stephens et al. (2019); Y20 = this study.

• Extracting spectra: We use the CASA task specflux to extract the mean flux density within the ellipse which has the same major and minor axes as well as the position angle as the fitted continuum sources.

- Baseline calibration: The continuum has been removed before the imaging process; however, the extracted spectra sometimes still show imperfect baselines. The spectra show significant differences from one to another such as rich emission lines, lack of emission, and broad emission features. Thus, we manually select the frequency ranges for baseline calibration for each spectral window and each field.
- Velocity correction: Finally, the frequency of the extracted spectra are corrected according to the source velocities. We collect the source velocities from the literature as well as from the strong emission lines in our spectra. Table 1 lists the adopted source velocities and the corresponding references.

Figure 1 shows the images of the continuum along with the fitted shapes, while the properties of the continuum sources are listed in Table 1. Our observations detect 50 continuum sources. The continuum emission appears as compact circular or elliptical shape with no sub-structure. Some sources show extended continuum emission resembling the shape of outflow cavities.

Three sources, EDJ2009-237, Per-emb-60, and EDJ2009-172, have no spectral line detected; therefore, we exclude them from spectral extraction as well as the line identification and modeling. These three sources still need to be included for detection number statistics.

## 3.1. Line Identifications and Modeling

Line identification starts with manual identification and verification for a few sources with rich spectra, including Per-emb-12B and B1-bS. We use SPLATALOGUE<sup>1</sup> to identity the molecular species and use XCLASS (Möller et al. 2017) to verify the identification. The XCLASS package is a LTE radiative transfer code that uses the molecular data from the Cologne Database of Molecular Spectroscopy (CDMS; Müller et al. 2001, 2005; Endres et al. 2016) and the Jet Propulsion Laboratory (JPL; Pickett et al. 1998). An identification needs to satisfy the following criteria.

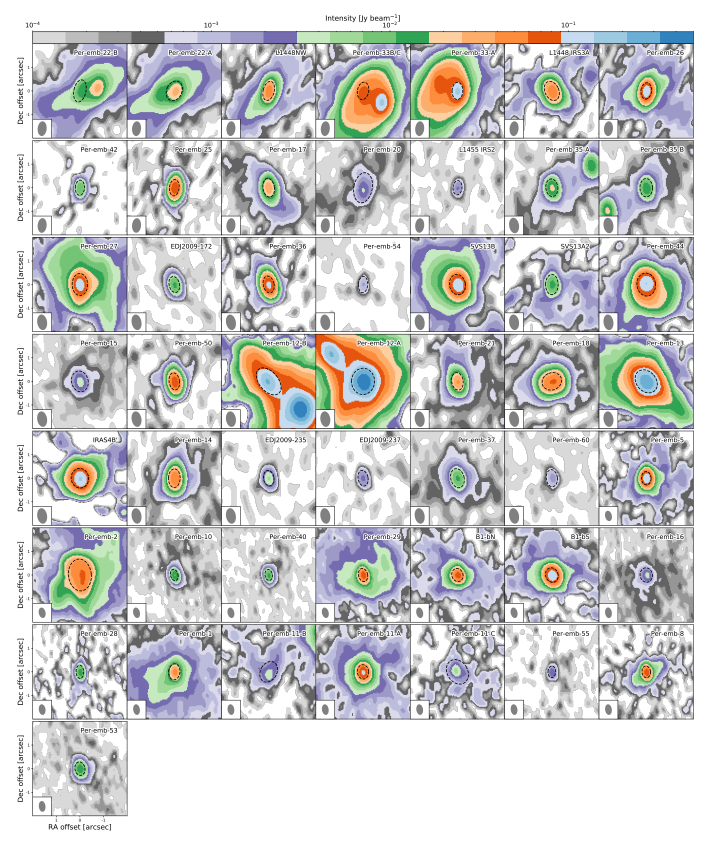
• The spectra agree with the predicted strengths of the model.

- The spectral lines are not all blended with other emission, such as other molecules and the SiO emission tracing the outflows. The emission of a few species, such as HDCO & <sup>13</sup>CH<sub>3</sub>OH, CH<sub>3</sub>CHO & CH<sub>2</sub>DOH, <sup>34</sup>SO & C<sub>2</sub>H<sub>5</sub>OH, and CH<sub>3</sub>OCH<sub>3</sub> & CH<sub>2</sub>DCN, are partially blended (blending occurs at a few lines but other lines remain isolated). The fittings of those species are performed together to verify their identification.
- Identified molecules need to be already found toward young stellar objects as summarized in McGuire (2018).

Table 3 lists the identified species and transitions. Only identifiable transitions are listed. The XCLASS modeling includes all the transitions in our frequency coverage regardless their Einstein-A values and upper energy levels.

Systematic spectral fitting using XCLASS is then applied to all sources using a list of species, compiled from those identifications. The catalogs used in this study are listed in Appendix A The fitting function in XCLASS includes several optimization algorithms that can be used in series to reduce biases. We configure the algorithm chain that starts with the genetic algorithm followed by the Levenberg-Marquardt  $\chi^2$  minimization. The genetic algorithm searches the best-fitting parameters iteratively with generations that evolve like a natural selection, where the better fitting models get less modification over generations. We setup the genetic algorithm to search for the top two best-fitting models after 30 generations. Then, the Levenberg-Marquardt  $\chi^2$  minimization applies to the three best-fitting models for 20 iterations to find the best-fitting models. The genetic algorithm aims to find possible local minimums and the Levenberg-Marquardt minimization further find the best-fitting models in the local minimums. The two best-fitting models found by the genetic algorithm often very similar, suggesting that there is only one minimum. To address the rare cases of two local minimums, we pick the model with the lower  $\chi^2$  values from the two bestfitting models constrained by the Levenberg-Marquardt minimization. We assume the COMs are all concentrated at the center, simplified as a 2D thin circular disk. There are four parameters for the XCLASS modeling, the size of the emitting molecule  $(r_{\text{COM}})$ , the excitation temperature  $(T_{\rm ex})$ , the column density  $(N_{\rm COM})$ , and the line width  $(\Delta \nu)$ . Due to the limited frequency coverage, many species only have a few lines detected, we fix  $r_{\text{COM}}$  as 0.75, similar to our beam size, and optimize the model with five excitation temperatures, 100, 150, 200, 250, and 300 K. We allow the line width varying between  $1.2\,\mathrm{km\,s^{-1}}$  to  $3.5\,\mathrm{km\,s^{-1}}$  for better fitting

http://www.splatalogue.net/



**Figure 1.** The continuum images of all PEACHES protostars. Non-detections toward L1448 IRS 2E and NGC 1333 SVS 3 are not shown. The dashed ellipses illustrate the size of fitted continuum, which is the region for extracting 1D spectra.

quality, and the range of the column density for each molecule is chosen according to the strength of the emission. The range of fitted column densities at different

temperatures indicates the uncertainty of the column densities.

The uncertainty from the fitting.

Table 3. Line Identification

Frequency (MHz)	Transition <sup>a</sup>	log(Einstein-A)	$E_{\rm u}$ (K)	$g_{ m u}$	Ref.			
	Ethynyl (CC	CH)						
$262065.00 \ (0.05)$	$[3, 5/2, 3] \rightarrow [2, 3/2, 2]^{b}$	-4.31	25.16	7	CDMS			
$262067.47 \ (0.05)$	$[3, 5/2, 2] \rightarrow [2, 3/2, 1]^{b}$	-4.35	25.16	5	CDMS			
262078.93 (0.02)	$[3, 5/2, 2] \rightarrow [2, 3/2, 2]^{b}$	-5.22	25.16	5	CDMS			
	Cyclopropenylidene	$(c-C_3H_2)$						
244222.15 (0.01)	$[3, 2, 1] \rightarrow [2, 1, 2]$	-4.23	18.17	21	CDMS			
246557.77 (0.02)	$[16, 10, 7] \rightarrow [16, 9, 8]$	-3.36	397.83	99	CDMS			
$260479.75 \ (0.02)$	$[5, 3, 2] \rightarrow [4, 4, 1]$	-3.79	44.72	33	CDMS			
Methanol (CH <sub>3</sub> OH $v_{ m t}=0$ )								
243915.79 (0.01)	$[5, 1, 4] \rightarrow [4, 1, 3] A$	-4.22	49.66	44	CDMS			
246074.61 (0.02)	$[20, 3, 17] \rightarrow [20, 2, 18] A$	-4.08	537.03	164	CDMS			
246873.30 (0.02)	$[19, 3, 16] \rightarrow [19, 2, 17] A$	-4.08	490.65	156	CDMS			
261805.68 (0.01)	$[2, 1, 1] \rightarrow [1, 0, 1] E$	-4.25	28.01	20	CDMS			
	Methanol ( <sup>13</sup> CH <sub>3</sub> O	$v_{\rm t} = 0$						
246426.12 (0.22)	$[23, 4, 19] \rightarrow [22, 5, 18]$	-4.58	721.02	47	CDMS			
247086.3 (0.5)	$[23, 3, 20] \rightarrow [23, 2, 21] A \rightarrow A +$	-4.07	674.86	47	CDMS			
259036.49 (0.17)	$[17, 3, 15] \rightarrow [17, 2, 16] A + \rightarrow A -$	-4.04	396.48	35	CDMS			
	Methanol (CH <sub>2</sub> DO	$H v_{t} = 0$						
243514.31 (0.01)	$[9, 2, 8] \rightarrow [10, 1, 10] \text{ o}_1$	-5.17	131.85	19	JPL			
246973.11 (0.01)	$[4, 1, 4] \rightarrow [4, 1, 3] e_1$	-4.67	37.69	9	JPL			
260543.63 (0.01)	$[3, 2, 1] \rightarrow [3, 1, 2] o_1$	-4.65	48.34	7	JPL			
	Methanol (CH <sub>3</sub> <sup>18</sup> O)							
246256.60 (0.04)	[11. 2. 10]→[10, 3, 7] A	-4.64	184.27	92	CDMS			
	Sulfur monoxide							
258255.83 (0.01)	$[N, J] = [6, 6] \rightarrow [5, 5]$	-3.67	56.50	13	CDMS			
261843.72 (0.03)	$[N, J] = [7, 6] \rightarrow [6, 5]$	-3.64	47.55	15	CDMS			
	Sulfur monoxide		11100		02.110			
246663.47 (0.1)	$[N, J] = [5, 6] \rightarrow [4, 5]$	-3.74	49.89	11	CDMS			
21000011 (0.1)	Sulfur dioxide		10100		020			
244254.22 (0.01)	$[14, 0, 14] \rightarrow [13, 1, 13]$	-3.79	93.90	29	CDMS			
244204.22 (0.01)	Hydrogen cyanide		33.30		CDMB			
259010.26 (0.01)	$[J, F] = [3, 3] \rightarrow [2, 3]$	-4.07	24.86	7	CDMS			
259011.55 (0.01)	$[J, F] = [3, 2] \rightarrow [2, 1]$	-3.19	24.86	5	CDMS			
259011.80 (0.01) 259011.80 (0.01)	$[J, F] = [3, 2] \rightarrow [2, 1]$ $[J, F] = [3, 3] \rightarrow [2, 2]$	-3.19 $-3.16$	24.86	7	CDMS			
259011.86 (0.01) 259011.86 (0.01)	$[J, F] = [3, 3] \rightarrow [2, 2]$ $[J, F] = [3, 4] \rightarrow [2, 3]$	-3.10 $-3.11$	24.86	9	CDMS			
259011.30 (0.01) 259012.34 (0.01)	$[J, F] = [3, 2] \rightarrow [2, 3]$ $[J, F] = [3, 2] \rightarrow [2, 3]$	-5.46	24.86	5	CDMS			
259012.34 (0.01) 259013.89 (0.01)	$[J, F] = [3, 2] \rightarrow [2, 3]$ $[J, F] = [3, 2] \rightarrow [2, 2]$	-3.40 $-3.92$	24.86	5	CDMS			
239013.89 (0.01)			24.00	J	CDMS			
044095 50 (0.01)	Carbon Monosulf	-3.53	25.07	11	CDMC			
244935.56 (0.01)	$[J]=[5] \rightarrow [4]$		35.27	11	CDMS			
2460246 (0.1)	Formaldehyde (I	*	07.00	0	GDMG.			
246924.6 (0.1)	$[4, 1, 4] \rightarrow [3, 1, 3]$	-3.40	37.60	9	CDMS			
259034.9 (0.1)	$[4, 2, 2] \rightarrow [3, 2, 1]$	-3.44	62.86	9	CDMS			
Methyl formate (CH <sub>3</sub> OCHO)								
245883.2 (0.1)	$[20, 13, 7] \rightarrow [19, 13, 6] \text{ E}$	-3.89	235.98	82	JPL			
245885.2 (0.1)	$[20, 13, 7] \rightarrow [19, 13, 6] \text{ A}$	-3.89	235.98	82	JPL			
245885.2 (0.1)	$[20, 13, 8] \rightarrow [19, 13, 7] \text{ A}$	-3.89	235.98	82	JPL			
245903.7 (0.1)	$[20, 13, 8] \rightarrow [19, 13, 7] E$	-3.89	235.97	82	JPL			
246027.5 (0.1)	$[21, 2, 19] \rightarrow [20, 3, 18] E$	-4.63	139.85	86	JPL			

 $Table \ 3 \ continued$ 

7

 ${\bf Table} \ {\bf 3} \ (continued)$ 

Frequency (MHz)	Transition <sup>a</sup>	log(Einstein-A)	<i>E</i> <sub>u</sub> (K)	$g_{ m u}$	Ref.
246038.9 (0.1)	$[21, 2, 19] \rightarrow [20, 3, 18] A$	-4.63	139.85	86	JPL
246054.8 (0.1)	$[20, 12, 8] \rightarrow [19, 12, 7] E$	-3.84	219.43	82	$_{ m JPL}$
246060.8 (0.1)	$[20, 12, 8/9] \rightarrow [19, 12, 7/8] A$	-3.84	219.43	82	$_{ m JPL}$
246076.9(0.1)	$[20, 12, 9] \rightarrow [19, 12, 8] E$	-3.84	219.41	82	$_{ m JPL}$
246285.4 (0.1)	$[20, 11, 9] \rightarrow [19, 11, 8] E$	-3.80	204.21	82	$_{ m JPL}$
246295.1 (0.1)	$[20, 11, 10] \rightarrow [19, 11, 9] A$	-3.80	204.21	82	$_{ m JPL}$
246295.1 (0.1)	$[20, 11, 9] \rightarrow [19, 11, 8] A$	-3.80	204.21	82	$_{ m JPL}$
246308.3 (0.1)	$[20, 11, 10] \rightarrow [19, 11, 9] E$	-3.80	204.20	82	$_{ m JPL}$
246456.1 (0.1)	$[10, 5, 6] \rightarrow [9, 4, 5] E$	-5.52	49.09	42	$_{ m JPL}$
246600.0 (0.1)	$[20, 10, 10] \rightarrow [19, 10, 9] E$	-3.77	190.34	82	$_{ m JPL}$
$246613.4\ (0.1)$	$[20, 10, 11] \rightarrow [19, 10, 10] A$	-3.77	190.34	82	$_{ m JPL}$
246613.4 (0.1)	$[20, 10, 10] \rightarrow [19, 10, 9] A$	-3.77	190.34	82	$_{ m JPL}$
246623.2 (0.1)	$[20, 10, 11] \rightarrow [19, 10, 10] E$	-3.77	190.34	82	$_{ m JPL}$
246660.5 (0.1)	$[10, 5, 6] \rightarrow [9, 4, 5] A$	-4.74	49.08	42	$_{ m JPL}$
246675.4 (0.1)	$[15, 4, 12] \rightarrow [14, 3, 11] E$	-4.93	81.85	62	$_{ m JPL}$
246683.5 (0.1)	$[15, 4, 12] \rightarrow [14, 3, 11] A$	-4.93	81.84	62	$_{ m JPL}$
246752.9(0.1)	$[10, 5, 5] \rightarrow [9, 4, 5] E$	-4.90	49.10	42	$_{ m JPL}$
246891.6 (0.1)	$[19, 4, 15] \rightarrow [18, 4, 14] E$	-3.66	126.22	78	$_{ m JPL}$
246914.7 (0.1)	$[19, 4, 15] \rightarrow [18, 4, 14] A$	-3.66	126.22	78	$_{ m JPL}$
246945.7 (0.1)	$[10, 5, 6] \rightarrow [9, 4, 6] E$	-4.90	49.09	42	$_{ m JPL}$
247040.7 (0.1)	$[20, 9, 11] \rightarrow [19, 9, 10] E$	-3.74	177.83	82	$_{ m JPL}$
247044.1 (0.1)	$[21, 3, 19] \rightarrow [20, 3, 18] E$	-3.66	139.90	86	$_{ m JPL}$
247053.5 (0.1)	$[21, 3, 19] \rightarrow [20, 3, 18] A$	-3.66	139.89	86	JPL
247057.3(0.1)	$[20, 9, 12] \rightarrow [19, 9, 11] A$	-3.74	177.83	82	$_{ m JPL}$
247057.7 (0.1)	$[20, 9, 11] \rightarrow [19, 9, 10] A$	-3.74	177.83	82	JPL
247063.7(0.1)	$[20, 9, 12] \rightarrow [19, 9, 11] E$	-3.74	177.83	82	$_{ m JPL}$
247124.3 (0.1)	$[10, 5, 5] \rightarrow [9, 4, 6] E$	-4.74	49.08	42	$_{ m JPL}$
258275.0 (0.1)	$[21, 13, 8] \rightarrow [20, 13, 7] E$	-3.79	248.37	86	$_{ m JPL}$
258277.4 (0.1)	$[21, 13, 8] \rightarrow [20, 13, 7] A$	-3.79	248.37	86	$_{ m JPL}$
258277.4 (0.1)	$[21, 13, 9] \rightarrow [20, 13, 8] A$	-3.79	248.37	86	$_{ m JPL}$
259341.9 (0.1)	$[24, 0, 24] \rightarrow [23, 1, 23] E$	-4.37	158.23	98	$_{ m JPL}$
259342.0 (0.1)	$[24, 1, 24] \rightarrow [23, 1, 23] E$	-3.58	158.23	98	JPL
259342.1 (0.1)	$[24, 0, 24] \rightarrow [23, 0, 23] E$	-3.58	158.23	98	JPL
259342.3 (0.1)	$[24, 1, 24] \rightarrow [23, 0, 23] E$	-4.37	158.23	98	JPL
259342.7 (0.1)	$[24, 0, 24] \rightarrow [23, 1, 23] A$	-4.37	158.22	98	JPL
259342.9 (0.1)	$[24, 1, 24] \rightarrow [23, 1, 23] A$	-3.58	158.22	98	JPL
259343.0 (0.1)	$[24, 0, 24] \rightarrow [23, 0, 23] A$	-3.58	158.22	98	JPL
259343.2 (0.1)	$[24, 1, 24] \rightarrow [23, 0, 23] A$	-4.37	158.22	98	JPL
261822.3 (0.1)	$[17, 10, 7] \rightarrow [17, 9, 8] \text{ A}$	-4.73	156.63	70	JPL
262088.2 (0.1)	$[16, 10, 6] \rightarrow [16, 9, 7] \text{ A}$	-4.76	146.59	66	JPL
262088.2 (0.1)	$[16, 10, 7] \rightarrow [16, 9, 8] \text{ A}$	-4.76	146.59	66	JPL
` ,	Methyl formate (CH <sub>3</sub>	OCHO v = 1)			
243511.5 (0.1)	$[20, 12, 8] \rightarrow [19, 12, 7] E$	-3.85	407.25	82	JPL
245846.9(0.1)	$[21, 3, 19] \rightarrow [20, 3, 18] E$	-3.66	326.30	86	$_{ m JPL}$
246106.8 (0.1)	$[20, 7, 14] \rightarrow [19, 7, 13] A$	-3.70	343.77	82	$_{ m JPL}$
246184.2 (0.1)	$[20, 8, 13] \rightarrow [19, 8, 12] E$	-3.72	353.27	82	$_{ m JPL}$
246187.0 (0.1)	$[21, 2, 19] \rightarrow [20, 2, 18] \text{ A}$	-3.66	326.62	86	$_{ m JPL}$
246233.6 (0.1)	$[20, 7, 13] \rightarrow [19, 7, 12] \text{ A}$	-3.70	343.79	82	$_{ m JPL}$
246274.9 (0.1)	$[20, 7, 13] \rightarrow [19, 7, 12] E$	-3.70	343.86	82	JPL
246410.95 (0.01)	$[10, 5, 5] \rightarrow [9, 4, 6] \text{ A}$	-4.73	236.70	42	JPL
246422.7 (0.1)	$[22, 1, 21] \rightarrow [21, 2, 20] \text{ A}$	-4.51	330.43	90	JPL
246461.2 (0.1)	$[22, 2, 21] \rightarrow [21, 2, 20] \text{ A}$	-3.65	330.43	90	JPL
( /	. , , , , , , , , , , , , , , , , , , ,	2			
246488.4 (0.1)	$[22, 1, 21] \rightarrow [21, 1, 20] A$	-3.65	330.43	90	$_{ m JPL}$

 $Table \ 3 \ continued$ 

Table 3 (continued)

$\textbf{Table 3} \; (continued)$							
Frequency (MHz)	Transition <sup>a</sup>	log(Einstein-A)	<i>E</i> <sub>u</sub> (K)	$g_{ m u}$	Ref.		
246706.5 (0.1)	[22, 2, 21]→[21, 2, 20] E	-3.65	329.89	90	JPL		
246731.7(0.1)	$[22, 1, 21] \rightarrow [21, 1, 20] E$	-3.65	329.89	90	$_{ m JPL}$		
246985.2(0.1)	$[20, 6, 15] \rightarrow [19, 6, 14] A$	-3.68	335.37	82	$_{ m JPL}$		
259003.9(0.1)	$[21, 7, 14] \rightarrow [20, 7, 13] A$	-3.63	356.22	86	$_{ m JPL}$		
259025.8(0.1)	$[21, 7, 14] \rightarrow [20, 7, 13] E$	-3.63	356.29	86	$_{ m JPL}$		
260479.6 (0.1)	$[44, 9, 36] \rightarrow [44, 8, 37] A$	-4.59	828.74	178	JPL		
	Dimethyl ether (C	H <sub>3</sub> OCH <sub>3</sub> )					
246499.29 (0.01)	$[37, 6, 31] \rightarrow [37, 5, 12] AA$	-4.01	693.72	750	CDMS		
246505.09 (0.01)	$[37, 6, 31] \rightarrow [37, 5, 12] AE$	-4.01	693.72	450	CDMS		
246505.09 (0.01)	$[37, 6, 31] \rightarrow [37, 5, 12] EA$	-4.01	693.72	300	CDMS		
$246697.43 \ (0.01)$	$[27, 4, 23] \rightarrow [26, 5, 21] \text{ AA}$	-4.70	367.61	330	CDMS		
246697.87 (0.01)	$[27, 4, 23] \rightarrow [26, 5, 21] EE$	-4.70	367.61	880	CDMS		
$246698.31 \ (0.01)$	$[27, 4, 23] \rightarrow [26, 5, 21] AE$	-4.70	367.61	110	CDMS		
$246698.31 \ (0.01)$	$[27, 4, 23] \rightarrow [26, 5, 21] EA$	-4.70	367.61	220	CDMS		
$259305.22 \ (0.01)$	$[33, 3, 31] \rightarrow [34, 6, 28] \text{ AA}$	-6.61	563.02	670	CDMS		
259308.39 (0.01)	$[33, 3, 31] \rightarrow [34, 6, 28] \text{ AE}$	-6.61	563.02	402	CDMS		
259308.39 (0.01)	$[33, 3, 31] \rightarrow [34, 6, 28] EA$	-6.61	563.02	268	CDMS		
259309.47 (0.01)	$[17, 5, 12] \rightarrow [17, 4, 13] \text{ AE}$	-4.06	174.54	210	CDMS		
259309.76 (0.01)	$[17, 5, 12] \rightarrow [17, 4, 13] EA$	-4.06	174.54	140	CDMS		
259311.95 (0.01)	$[17, 5, 12] \rightarrow [17, 4, 13] \text{ EE}$	-4.06	174.54	560	CDMS		
259314.28 (0.01)	$[17, 5, 12] \rightarrow [17, 4, 13] \text{ AA}$	-4.06	174.54	350	CDMS		
	Acetone (CH <sub>3</sub> C						
244218.91 (0.01)	$[20, 5, 15] \rightarrow [19, 6, 14] \text{ AE}$	-3.32	139.69	82	$_{ m JPL}$		
244218.91 (0.01)	$[20, 6, 15] \rightarrow [19, 5, 14] \text{ AE}$	-3.32	139.69	250	JPL		
244218.92 (0.01)	$[20, 5, 15] \rightarrow [19, 6, 14] \text{ EA}$	-3.32	139.69	160	JPL		
244218.92 (0.01)	$[20, 6, 15] \rightarrow [19, 5, 14] \text{ EA}$	-3.32	139.69	160	JPL		
245831.34 (0.09)	$[13, 10, 3] \rightarrow [12, 9, 4] \text{ EE}$	-3.80	77.84	432	JPL		
246400.99 (0.05)	$[34, 7, 28] \rightarrow [34, 5, 29] \text{ EE}$	-4.17	364.98	1100	JPL		
246400.99 (0.05)	$[34, 6, 28] \rightarrow [34, 5, 29] \text{ EE}$	-4.03	364.98	1100	JPL		
246400.99 (0.05)	$[34, 7, 28] \rightarrow [34, 6, 29] \text{ EE}$	-4.03	364.98	1100	JPL		
246400.99 (0.05)	$[34, 6, 28] \rightarrow [34, 6, 29] \text{ EE}$	-4.17	364.98	1100	$_{ m JPL}$		
246404.27 (0.01)	$[22, 3, 19] \rightarrow [21, 4, 18] \text{ AE}$	-3.23	$149.62 \\ 149.62$	90 270	$_{ m JPL}$		
246404.27 (0.01)	$[22, 4, 19] \rightarrow [21, 3, 18] \text{ AE}$	-3.23 $-3.23$	149.62	180	$_{ m JPL}$		
246404.29 (0.01) 246404.29 (0.01)	$[22, 3, 19] \rightarrow [21, 4, 18] \text{ EA}$ $[22, 4, 19] \rightarrow [21, 3, 18] \text{ EA}$	-3.23 -3.23	149.62	180	$_{ m JPL}$		
246450.40 (0.01)	$[22, 4, 19] \rightarrow [21, 3, 18] \text{ EA}$ $[22, 4, 19] \rightarrow [21, 3, 18] \text{ EE}$	-3.23 -3.23	149.57	720	$_{ m JPL}$		
246450.40 (0.01)	$[22, 3, 19] \rightarrow [21, 3, 18] \text{ EE}$ $[22, 3, 19] \rightarrow [21, 3, 18] \text{ EE}$	-5.25 -5.09	149.57	720	JPL		
246450.40 (0.01)	$[22, 3, 19] \rightarrow [21, 4, 18] \text{ EE}$	-3.24	149.57	720	JPL		
246450.40 (0.01)	$[22, 4, 19] \rightarrow [21, 4, 18] \text{ EE}$	-4.92	149.57	720	JPL		
246496.17 (0.46)	$[25, 14, 12] \rightarrow [24, 15, 9] \text{ AE}$	-5.01	257.11	100	JPL		
246496.47 (0.02)	$[23, 14, 12] \rightarrow [21, 4, 18] \text{ AA}$	-3.23	149.51	270	JPL		
246496.47 (0.02)	$[22, 4, 19] \rightarrow [21, 3, 18] \text{ AA}$	-3.23	149.51	450	JPL		
246714.12 (0.05)	$[9, 8, 1] \rightarrow [8, 5, 4] \text{ EA}$	-5.84	40.59	76	JPL		
246714.94 (0.05)	$[32, 4, 28] \rightarrow [32, 4, 29] \text{ EA}$	-3.97	305.61	260	JPL		
246714.94 (0.05)	$[32, 5, 28] \rightarrow [32, 3, 29] \text{ EA}$	-3.97	305.61	260	JPL		
246715.04 (0.05)	$[32, 5, 28] \rightarrow [32, 4, 29] \text{ AE}$	-3.97	305.61	390	JPL		
246715.04 (0.05)	$[32, 4, 28] \rightarrow [32, 3, 29] \text{ EA}$	-3.97	305.61	130	JPL		
246719.92 (0.04)	$[33, 6, 28] \rightarrow [33, 4, 29] \text{ EE}$	-5.62	344.85	1100	JPL		
246719.92 (0.04)	$[33, 5, 28] \rightarrow [33, 4, 29] \text{ EE}$	-3.87	344.85	1100	JPL		
246719.92 (0.04)	$[33, 6, 28] \rightarrow [33, 5, 29] \text{ EE}$	-3.87	344.85	1100	JPL		
246719.92 (0.04)	$[33, 5, 28] \rightarrow [33, 5, 29] \text{ EE}$	-5.61	344.85	1100	JPL		
261818.11 (0.01)	$[20, 7, 13] \rightarrow [19, 8, 12] \text{ EA}$	-3.31	151.17	160	JPL		
261818.17 (0.01)	$[20, 7, 13] \rightarrow [19, 8, 12] \text{ AE}$	-3.31	151.17	82	JPL		
261819.09 (0.01)	$[20, 8, 13] \rightarrow [19, 7, 12] \text{ EA}$	-3.31	151.17	160	JPL		
	. , , , , , , , , , , , , , , , , , , ,						

 $Table \ 3 \ continued$ 

Table 3 (continued)

Frequency (MHz)	Transition <sup>a</sup>	log(Einstein-A)	<i>E</i> <sub>u</sub> (K)	$g_{ m u}$	Ref.				
261819.17 (0.01)	$[20, 8, 13] \rightarrow [19, 7, 12] AE$	-3.31	151.17	250	JPL				
Methyl cyanide (CH <sub>3</sub> CN)									
257507.56 (0.01)	$[N, K] = [14, 2] \rightarrow [13, 2]$	-3.00	121.28	58	JPL				
257522.43 (0.01)	$[N, K] = [14, 1] \rightarrow [13, 1]$	-2.99	99.84	58	$_{ m JPL}$				
257527.38 (0.01)	$[N, K] = [14, 0] \rightarrow [13, 0]$	-2.99	92.70	58	JPL				
	Acetaldehyde (CH <sub>3</sub> CHO $v_{\rm t}=0$ )								
246330.73 (0.01)	$[15, 3, 13] \rightarrow [15, 2, 14] A$	-4.29	131.49	62	$_{ m JPL}$				
$260530.40 \ (0.01)$	$[14, 1, 14] \rightarrow [13, 1, 13] E$	-3.20	96.39	58	$_{ m JPL}$				
$260544.02\ (0.01)$	$[14, 1, 14] \rightarrow [13, 1, 13] A$	-3.20	96.32	58	$_{ m JPL}$				
260547.46 (2.07)	$[9, 4, 5] \rightarrow [9, 3, 7] E, v_t = 2$	-6.06	456.38	38	JPL				
	gauche-Ethanol $(g\text{-}\mathrm{C}_2\mathrm{H}_5\mathrm{OH})$								
$246414.76 \ (0.05)$	$[14, 3, 11] \rightarrow [13, 3, 10] v_t = 0 \rightarrow 0$	-3.89	155.72	29	$_{ m JPL}$				
$246524.28 \ (0.01)$	$[13, 2, 12] \rightarrow [12, 1, 12] v_t = 0 \rightarrow 1$	-4.50	136.95	27	$_{ m JPL}$				
$246658.18 \ (0.01)$	$[32, 5, 28] \rightarrow [32, 4, 29] v_t = 0 \rightarrow 0$	-6.33	527.94	65	$_{ m JPL}$				
$246662.98 \ (0.01)$	$[4,2,3]{\to}[3,1,3]\ v_{\rm t}=1{\to}0$	-4.36	74.77	9	$_{ m JPL}$				
$259322.64\ (0.01)$	$[14, 3, 11] \rightarrow [13, 2, 11] v_t = 0 \rightarrow 1$	-4.39	155.72	29	$_{ m JPL}$				
260457.73 (0.01)	[15. 4. 12] $\rightarrow$ [14, 4, 11] $v_{\rm t} = 1 \rightarrow 1$	-3.83	181.10	31	JPL				
trans-Ethanol ( $C_2H_5OH$ )									
$246663.62 \ (0.05)$	$[24, 1, 23] \rightarrow [24, 0, 24]$	-3.73	252.35	49	$_{ m JPL}$				
261815.99 (0.05)	$[28, 3, 26] \rightarrow [28, 2, 27]$	-3.96	350.98	57	JPL				
	Glycolaldehyde ( $cis$ -C	H <sub>2</sub> OHCHO)							
$246773.09 \ (0.02)$	$[30, 2, 28] \rightarrow [30, 1, 29]$	-4.04	252.68	61	CDMS				
$246778.28 \ (0.02)$	$[30, 3, 28] \rightarrow [30, 2, 29]$	-4.04	252.68	61	CDMS				
$262056.78 \ (0.01)$	$[25, 2, 24] \rightarrow [24, 1, 23]$	-3.34	158.25	51	CDMS				
$261795.48 \ (0.01)$	$[25, 11, 14] \rightarrow [25, 10, 15]$	-3.57	254.23	51	CDMS				
261798.96 (0.01)	$[25, 11, 15] \rightarrow [25, 10, 16]$	-3.57	254.23	51	CDMS				
	Methyl cyanide (C	H <sub>2</sub> DCN)							
$259315.51 \ (0.01)$	$[15, 1, 15] \rightarrow [14, 1, 14]$	-2.82	104.97	31	CDMS				
260523.05 (0.01)	$[15, 2, 13] \rightarrow [14, 2, 12]$	-2.82	121.60	31	CDMS				
	Ethyl cyanide ( $\mathrm{CH_3CH_2CN}$ )								
$246268.74 \ (0.01)$	$[27, 2, 25] \rightarrow [26, 2, 24]$	-2.90	169.80	55	CDMS				
$246421.92 \ (0.01)$	$[28, 2, 27] \rightarrow [27, 2, 26]$	-2.90	177.26	57	CDMS				
$246548.70 \ (0.01)$	$[27, 3, 24] \rightarrow [26, 3, 23]$	-2.90	174.06	55	CDMS				
260535.69 (0.05)	$[29, 5, 25] \rightarrow [28, 5, 24]$	-2.84	215.06	59	CDMS				
	Formamide (NH	<sub>2</sub> CHO)							
243521.04 (0.01)	$[12, 1, 12] \rightarrow [11, 1, 11]$	-2.98	79.19	25	CDMS				
	Formic acid ( $t$ -H	COOH)							
262103.48 (0.01)	$[12, 0, 12] \rightarrow [11, 0, 11]$	-3.69	82.77	25	CDMS				

 $<sup>^</sup>a\mathrm{The}$  typical quantum numbers are listed as  $[J,\,K_\mathrm{a},\,K_\mathrm{c}]$  unless specified.

## 4. CONTINUUM OPACITY

## 5. DETECTION STATISTICS

We summarize the fraction of sources with detections of molecules in Figure 2. The detection statistics include COMs, carbon-chain molecules, and the simple organic molecules, such as CS, H<sup>13</sup>CN, SO, <sup>34</sup>SO, and SO<sub>2</sub>. The PEACHES protostars show a great chemical diversity from no molecule detected (B1-bN) to rich spectra of COMs (e.g. Per-emb-12B). Detections of COMs and the number of COMs detected show no obvious correlation

with the bolometric luminosity and bolometric temperature of the protostars, which are conventional evolutionary indicators. Low luminosity sources have fewer COMs detected; however, if COMs mostly come from thermally desorption, the region with  $T>T_{\rm desorption}$  may be smaller for the low luminosity sources, making the emission of COMs fainter and reducing our sensitivity to detect COMs. We also compare the detection statistics with the mass derived from 9 mm observations that resolved the sources as a proxy of the central mass (Tychoniec et al. 2018). The detection statistics show no clear correlation with the central mass; however, the

<sup>&</sup>lt;sup>b</sup> The quantum numbers are [N, J, F]

sources with smaller central mass have fewer detections of COMs, which may due to their low luminosity.

#### 6. EXCITATION TEMPERATURES

6.1.  $CH_3OH$ 

6.2. CH<sub>3</sub> OCHO

#### 7. THE SPECTRA OF CCH

Due to the absorption and irregular line profile, the XCLASS fitting routine often fails to faithfully reproduce the observed CCH spectra.

#### 8. CORRELATIONS OF COMS

The chemical evolution of protostars may leave certain patterns in the abundance of molecules as the dynamical evolution determines the density and temperature structures, regulating chemical reactions. Thus, the abundance of COMs and their correlations provide critical information to constrain the chemical evolution at embedded protostars. The fitted column density of COMs indicates the abundance of COMs around protostars. Typically COMs are locked into the ices on dust grains at outer envelope. Therefore, we take the column density of COMs as a proxy of the abundance of COMs.

As described in Section 3.1, we fit the column density and line width with different excitation temperatures, resulting in a range of column density as its uncertainty. The comparison between CCH and CH<sub>3</sub>OH shows no correlation between these two molecules (Figure 18), similar to the conclusion in Higuchi et al. (2018). The single dish survey by Graninger et al. (2016) shows a correlation between C<sub>4</sub>H, a more complex carbon-chain molecules, and CH<sub>3</sub>OH. Outflow activity can promote the formation of CCH, which is more efficiency at warm temperature. In face, the morphology of CCH often traces the outflow cavities seen from CS. Therefore, the lack of correlation between CCH and CH<sub>3</sub>OH may be affected by outflows.

Figure 20 shows the correlations of several COMs selected from their detection rates. The column density of  $\mathrm{CH_3OH}$  best correlates with that of  $\mathrm{CH_3CN}$ . Belloche et al. (2020) also found the tight correlation between these two molecules. The column densities of  $\mathrm{CH_3OCH_3}$  and  $\mathrm{CH_3OCHO}$  also show a tight correlation. To quantify the goodness of correlation, we calculate the Pearson's correlation coefficient (r), which tests the linearity of two variables. A simple calculation of the Pearson's correlation coefficient would ignore the uncertainties of the column density as well as the upper limits. Thus, we use the bootstrapping method to sample the fitted column densities and upper limits to calculate Pearson's r. For detections, we assume a normal distribution centers on the best-fitted values with the uncer-

tainty as the width of the normal distribution; for non-detections, we assume a normal distribution centers at zero with a width of one third of the upper limit, which is the  $3\sigma$  upper limit. Figure 19 shows the distributions of the Pearson's r for the correlation between CH<sub>3</sub>OH and CH<sub>3</sub>OCHO after 10000 resampling for all derived column densities (detections & non-detections) and the detection-only sample. With the detection-only sample, the mean Pearson's  $r_d$  is 0.92, as expected for a tight correlation, with a Gaussian-like distribution skewed toward lower values. After including the upper limits, the mean Pearson's r decreases to 0.65 with larger uncertainty (the 95% credible interval increases by 76%). If we take  $r \geq 0.5$  and  $r_d \geq 0.9$  as the criteria for having correlations, (to be updated after updating the figure).

## 9. SPATIAL EXTENT OF COMS

## 10. DISCUSSION

10.1. Notes on the 1D Spectra

L1448 IRS 3A-

• The emission of SO and CS shows additional redshifted component, separated by  $\sim 4.6 \,\mathrm{km}\,\mathrm{s}^{-1}$ .

Per-emb-33-A —

- Strong absorption at 246509 MHz.
- HDCO seems to have two components for the transition at 246925 MHz, but the one at 259035 MHz only has one component, which is underestimated by the model.

Per-emb-33-B/C-

• CCH shows three components leading to an inaccurate fit.

Per-emb-42-

- Double-peaked CS line
- Triple-peaked CCH line

Per-emb-26—

- Red-shifted excess appears in the CS, SO, CH<sub>3</sub>OH, H<sup>13</sup>CN, and HDCO lines.
- Broad SO lines peak at slightly blue-shifted velocity ( $\sim 1 \,\mathrm{km\,s^{-1}}$ ).
- The best-fitting model overestimates the CH<sub>3</sub>CH<sub>2</sub>CN, CH<sub>3</sub>CHO, and c-C<sub>3</sub>H<sub>2</sub> lines, possibly due to the contamination of SiO, which should have been excluded for fitting. Tests are running now.
- The secondary CH<sub>3</sub>OH line becomes overestimated at  $T_{\rm ex} \ge 200$  K.
- Unidentified lines at 246525 MHz and 244249 MHz.

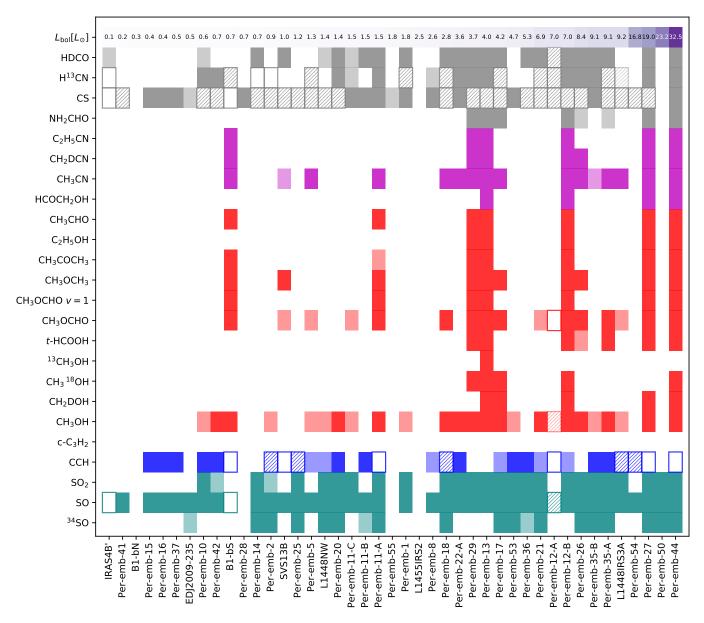


Figure 2. The detection statistics sorted by their bolometric luminosity.

## Per-emb-22-B-

- The SO lines have small red-shifted excess. The best-fitting line strength decreases significantly at  $T_{\rm ex}$  =200 K. Not sure why.
- The SO line profiles are slightly skewed toward blue-shifted velocity, while the CS line shows another brighter peak at blue-shifted velocity.

# Per-emb-22-A —

 $\bullet$  Hints of CH<sub>3</sub>OCHOemission, but not significant enough to warrant a detection.

Per-emb-17—

- Many line profiles exhibit a broad double-peaked profile, separated by  $\sim 5-6\,\mathrm{km\,s^{-1}}$ . Per-emb-17 is a binary system unresolved by our observations. However, the channel maps suggest that the two components are likely to surrounding the southern source, Per-emb-17-B.
- The CH<sub>3</sub>OCHO line at  $\sim$ 259343 MHz may be optically thick.

## Per-emb-20—

 The CCH lines have a narrow double-peaked profile. The CS line shows a similar double-peak profile.

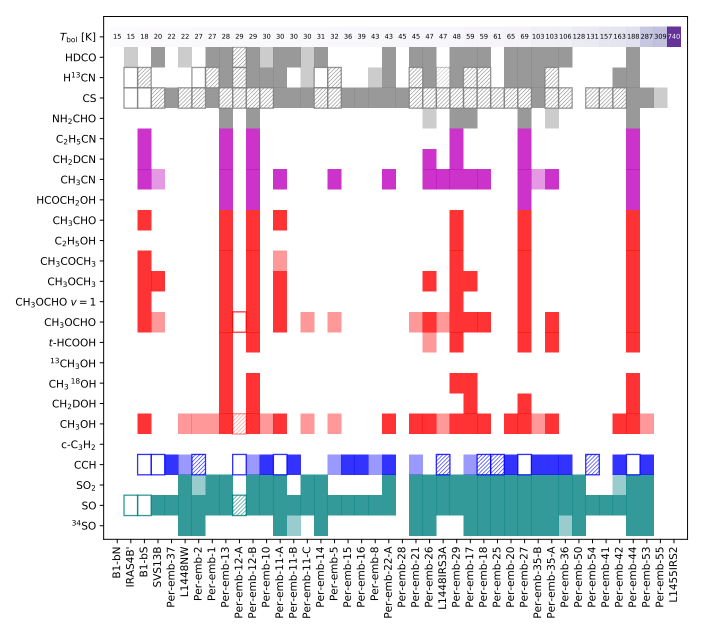


Figure 2 (Cont.). The same figure as Figure 2 but sorted by their bolometric temperature.

• The SO lines have a broad component underneath the typical narrow lines.

#### SVS13 A2-

- Weak indication of the unidentified line at 246525 MHz, which has been detected in other sources.
- The model strength of CCH suddenly decreases at  $T_{\rm ex}$  =200 K by  $\sim$ 0.5 K over a 2 K line.

## Per-emb-44—

 Unidentified lines at 244248 MHz, 246219 MHz, 246254 MHz, 246344 MHz, 246389 MHz, 246434 MHz,

- $246525\,\mathrm{MHz},\,246838\,\mathrm{MHz},\,258268\,\mathrm{MHz},\,258271\,\mathrm{MHz},$  and  $262068\text{-}262070\,\mathrm{MHz}.$
- Higher temperatures (Tex > 100 K) provide better fittings. Probably should adopt the temperature fitted from CH<sub>3</sub>OCHO (previous MCMC fitting suggests a temperature of 263 K).

#### Per-emb-12-B-

 $\bullet$  Unidentified lines at 244248 MHz, 246254 MHz, 246314 MHz, 246322 MHz, 246389 MHz, 246434 MHz, 246525 MHz, 246696 MHz, 246838 MHz, 246873 MHz, 247082 MHz, 258268 MHz, 258271 MHz, and 262068-262070 MHz.

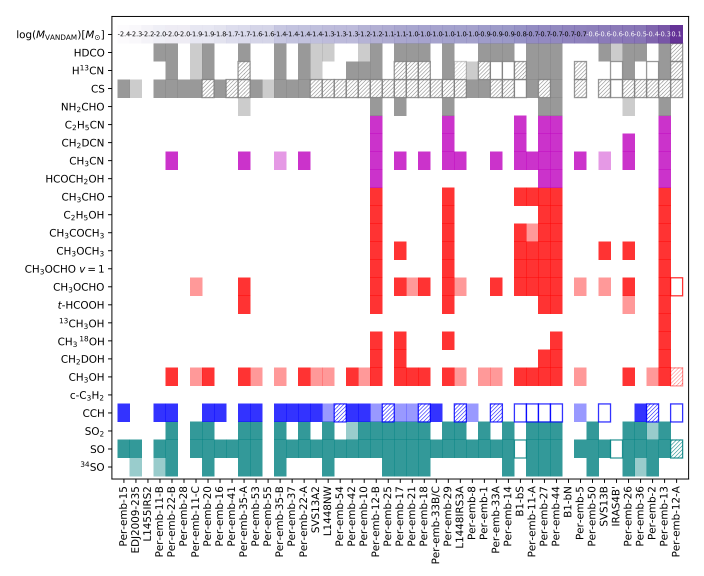


Figure 2 (Cont.). The same figure as Figure 2 but sorted by their mass derived from their 9 mm observations (Tychoniec et al. 2018).

### Per-emb-12-A —

• Strong absorption features detected across the spectra, CCH, SO, H<sup>13</sup>CN, CS, CH<sub>3</sub>OH, HDCO, CH<sub>3</sub>CN, and CH<sub>3</sub>OCHO.

## IRAS4B1—

• Spectra show no emission along with absorption at SO, CS, and CH<sub>3</sub>OH lines.

## Per-emb-13—

- The CH<sub>3</sub>OCHO emission needs  $T_{\rm ex} > 100$  K to have a good fit.
- All three CH<sub>3</sub>OH lines are detected but two of them show clear sign of self-absorption, therefore, not ideal for fitting the excitation temperature.

- Unidentified lines at 244248 MHz, 246254 MHz, 246331 MHz, 246344 MHz, 246434 MHz, 246525 MHz, 246838 MHz, 246974 MHz, 247086 MHz, 257268 MHz, 257271 MHz, 259323 MHz, 259331 MHz, 262098 MHz, and 262109 MHz.
- The best-fitting models have two different widths for the CH<sub>3</sub>CHO lines.
- The best-fitting model for <sup>13</sup>CH<sub>3</sub>OH lines overestimates the line width.

#### Per-emb-27—

 All three CH<sub>3</sub>OH lines are detected, but none of the temperature produce a good fit to all three lines, suggesting that some lines are optically thick. The transitions at 243916 MHz and

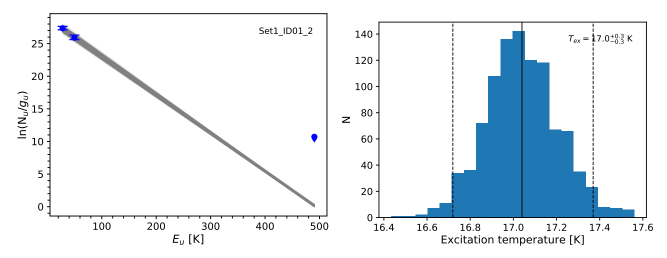


Figure 3. The methanol rotational diagram for L1448 IRS3A and the fitted excitation temperature distribution using the bootstraping method.

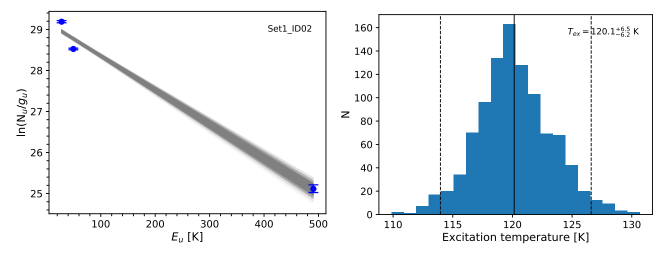


Figure 4. The methanol rotational diagram for Per-emb 26 and the fitted excitation temperature distribution using the bootstraping method.

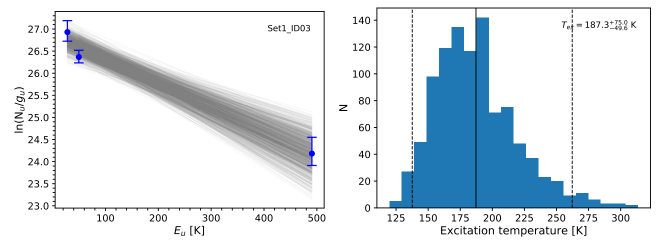


Figure 5. The methanol rotational diagram for Per-emb 22A and the fitted excitation temperature distribution using the bootstraping method.

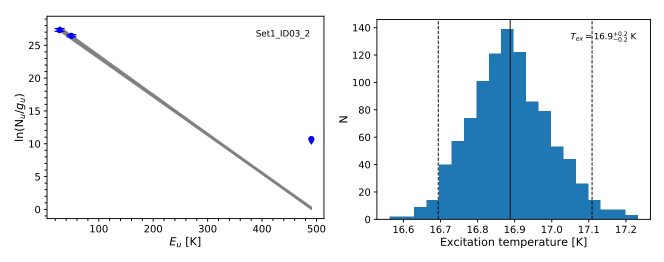


Figure 6. The methanol rotational diagram for Per-emb 22B and the fitted excitation temperature distribution using the bootstraping method.

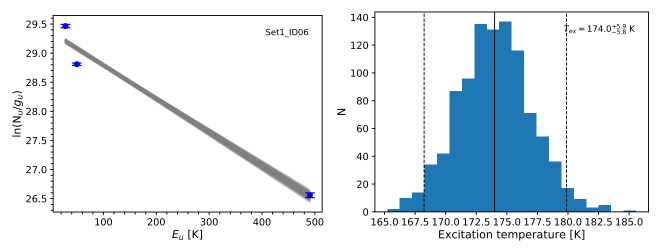


Figure 7. The methanol rotational diagram for Per-emb 17 and the fitted excitation temperature distribution using the bootstraping method.

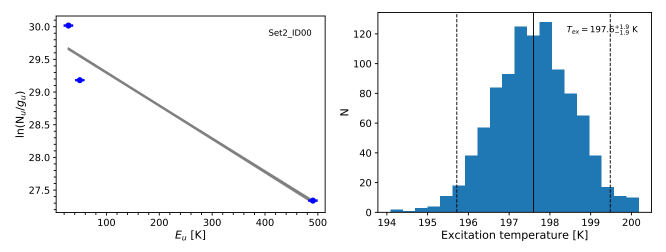


Figure 8. The methanol rotational diagram for Per-emb 44 and the fitted excitation temperature distribution using the bootstraping method.

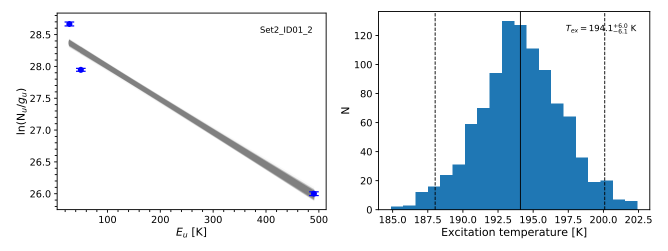


Figure 9. The methanol rotational diagram for Per-emb 12B and the fitted excitation temperature distribution using the bootstraping method.

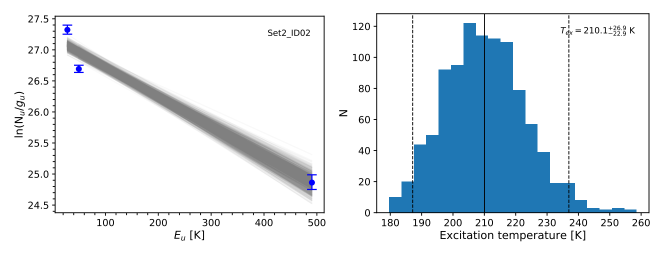


Figure 10. The methanol rotational diagram for Per-emb 13 and the fitted excitation temperature distribution using the bootstraping method.

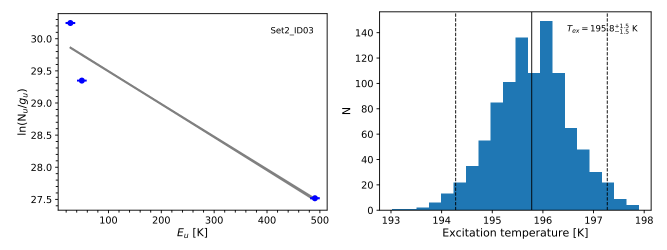


Figure 11. The methanol rotational diagram for Per-emb 27 and the fitted excitation temperature distribution using the bootstraping method.

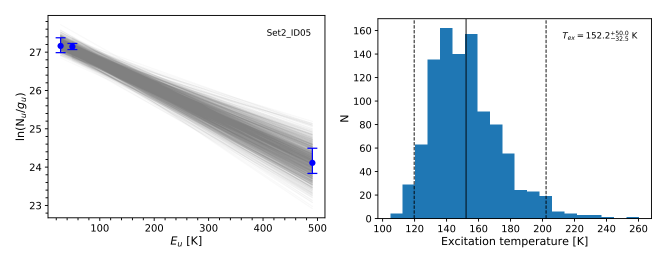


Figure 12. The methanol rotational diagram for Per-emb 21 and the fitted excitation temperature distribution using the bootstraping method.

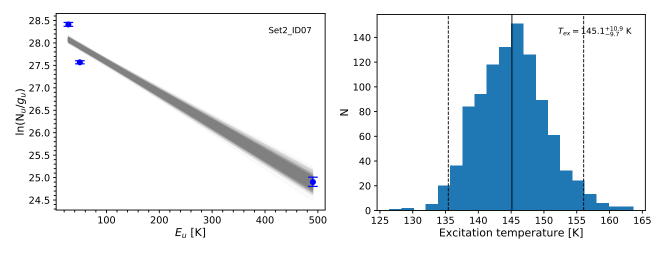


Figure 13. The methanol rotational diagram for Per-emb 35A and the fitted excitation temperature distribution using the bootstraping method.

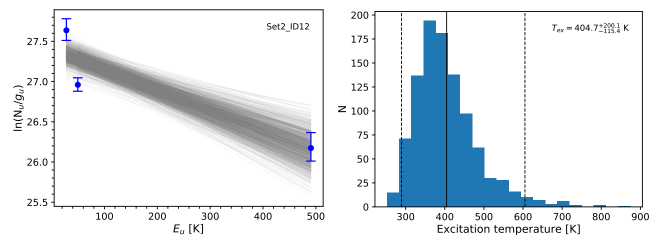


Figure 14. The methanol rotational diagram for Per-emb 18 and the fitted excitation temperature distribution using the bootstraping method.

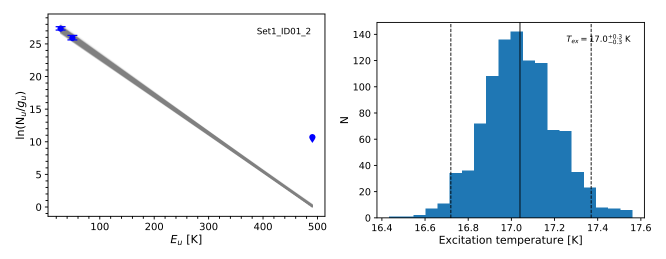


Figure 15. The methanol rotational diagram for L1448 IRS3A and the fitted excitation temperature distribution using the bootstraping method.

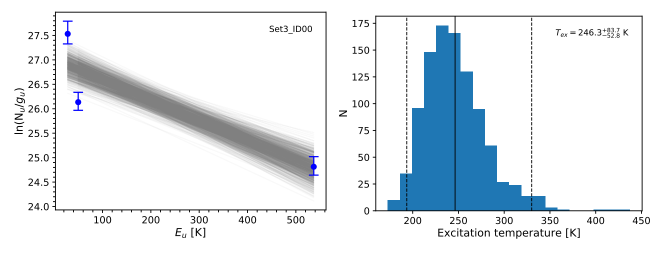


Figure 16. The methanol rotational diagram for B1-bS and the fitted excitation temperature distribution using the bootstraping method.

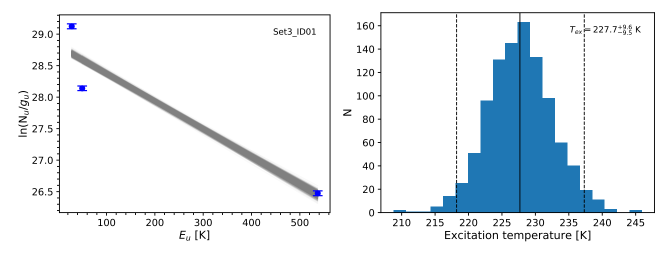


Figure 17. The methanol rotational diagram for Per-emb 29 and the fitted excitation temperature distribution using the bootstraping method.

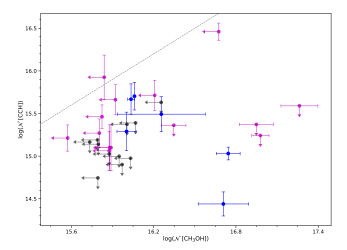


Figure 18. Correlation of the column densities of CCH and  $\mathrm{CH_3OH}$  fitted from the PEACHES protostars. The sources where both molecules are detected are shown in black; the sources where only one molecule is detected are shown in magenta; finally, the sources where both molecules are not detected are shown in black for the corresponding upper limits.

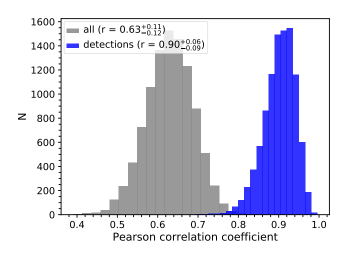


Figure 19. Distributions of Pearson's correlation coefficient from 10000 resamples drawn from detections + non-detections and only detections. The legend indicates the mean values of Pearson's r along with the range of the 95% credible interval as the associated uncertainties.

 $261806\,\mathrm{MHz}$  are  ${\sim}30$  K, while the transition at  $246873\,\mathrm{MHz}$  is about 24 K. They seems to be optically thick. In comparison, the continuum brightness temperature is only 5.8 K.

 $\bullet$  Unidentified lines at 244232 MHz, 244248 MHz, 246207 MHz, 246254 MHz, 246388 MHz, 246435 MHz, 246525 MHz, 246538 MHz, 246838 MHz, 246973 MHz, 247084 MHz, and 259330 MHz.

• The CH<sub>3</sub>OH line at 243916 MHz and the SO lines become optically thick at 100 K.

#### Per-emb-54—

- The SO lines appears red-shifted by  $\sim 2 \,\mathrm{km \, s^{-1}}$ .
- The emission of CS and CCH shows a blue-shifted peak along with absorption slightly red-shifted compared to the source velocity.

#### Per-emb-21—

• Emission of  $CH_3OH$  is detected. However, the broad widht and noisy spectra lead to a bad fit. The best-fitting model has the maximum line width allowed,  $3.5 \, \mathrm{km \, s^{-1}}$ .

## Per-emb-14-

• The best-fitting model underestimates the HDCO lines, possibly due to the simultaneously fitted <sup>13</sup>CH<sub>3</sub>OH lines.

#### Per-emb-35-B—

- The best-fitting model underestimates the SO lines at  $T_{\rm ex} = 100$  K.
- The CCH lines show a double-peaked line profile.
- The CH<sub>3</sub>OH line at 243915 MHzhas an S/N of 1.2, but hints the existence of CH<sub>3</sub>OH.

#### Per-emb-35-A —

- The goodness of fitting for the  $CH_3OH$  lines is a strong function of temperature, suggesting that the  $CH_3OH$  lines can indicate the  $T_{ex}$ .
- The CH<sub>3</sub>OCHO line at 259342 MHz has an S/N of 1.8, but hint the existence of CH<sub>3</sub>OCHO.

#### SVS13B—

• The fitted width of the HDCO lines is overestimated.

#### Per-emb-15—

• All lines have only the blue-shifted emission, making them blue-asymmetric.

### Per-emb-50—

• The SO and SO<sub>2</sub> lines have a very broad component  $(\Delta \nu = 6 \,\mathrm{km \, s^{-1}})$ , skewing toward the blue-shifted velocity.

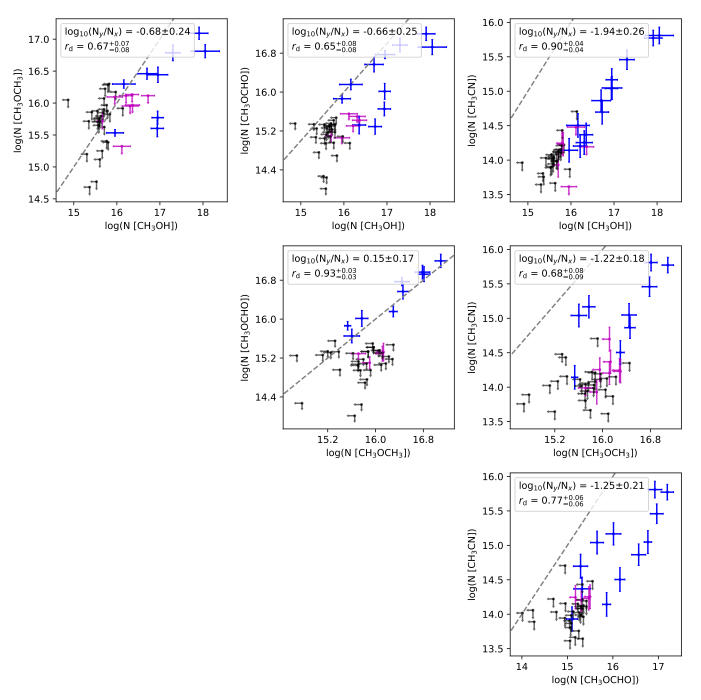


Figure 20. Corner plot of the correlations of the column densities between CH<sub>3</sub>OH, CH<sub>3</sub>CN, CH<sub>3</sub>OCHO, and CH<sub>3</sub>OCH<sub>3</sub>. The color code follows that in Figure 18. The dashed line indicates equality. The legends indicate the Pearson's r for the detection-only sample  $(r_d)$  and the logarithmic ratio of the two molecules  $(N_y/N_x)$ .

Per-emb-18—

- Many transitions of  $\mathrm{CH_3OCHO}$  are tentatively detected; however, none of them has  $\mathrm{S/N} > 3$ . Currently categorized as non-detection.
- $\bullet$  The HDCO lines are broader than the maximum allowed line width,  $3.5\,\rm km\,s^{-1},$  making the fitting inaccurate.

Per-emb-37—

• The fitting of SO lines has a strong variation as a function of temperatures.

Per-emb-36—

• The SO lines are underestimated by the bestfitting model. Perhaps it can be fixed by changing

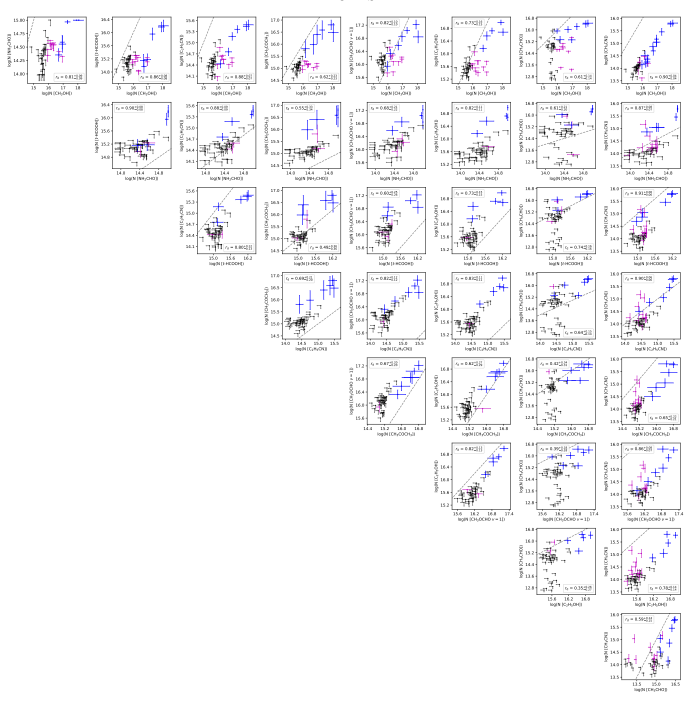


Figure 21. Corner plot of the correlations of the column densities between  $CH_3OH$ ,  $CH_3CN$ ,  $CH_3CHO$ ,  $C_2H_5OH$ ,  $CH_3OCHO v = 1$ ,  $CH_3COCH_3$ ,  $CH_3CH_2CN$ , t-HCOOH, and  $NH_2CHO$ . The legends are similar to Figure 20

the Variation parameter. To be tested. Test run on laptop shows no issue of having a good fit at 100 K for SO with Variatopn  $= 10^{-2}$ .

- The CCH lines only show at the blue-shifted velocities.
- The fitting of  $CH_3OCH_3$  is limited by the minimum line width of  $1.2 \,\mathrm{km \, s^{-1}}$ .

• Higher temperatures produce worse fittings to the

CH<sub>3</sub>OCHO lines. Previous MCMC fitting of the

CH<sub>3</sub>OCHO lines suggests a temperature of 58 K.

- Unidentified lines at 246027 MHz, 246099 MHz, 246143 MHz, 246192 MHz, 246525 MHz, 246674 MHz, and 2467320 MHz.
- The CH<sub>3</sub>OCHO lines around 258278 MHz and the H<sup>13</sup>CN lines have a few dips within the line profile, suggesting absorption or just noisy spectra.

#### Per-emb-29—

• Only two CH<sub>3</sub>OH lines are covered. Both lines have a strength of  $\sim 10$  K, suggesting optically thick.

#### Per-emb-5—

- The CCH fitting is not robust across different temperatures (only 150 K and 300 K fit).
  - 10.2. Comparison to the CALYPSO Survey
- 10.3. Complex Chemistry throughout Star Formation

10.4. 1D Spectra

Y.-L. Yang acknowledges the supports the JSPS Post-doctoral Fellowship from Japan Society for the Promotion of Science. This paper makes use of the following ALMA data: ADS/JAO.ALMA#2016.0.00391.S. ALMA is a partnership of ESO (representing its member states), NSF (USA) and NINS (Japan), together with NRC (Canada), MOST and ASIAA (Taiwan), and KASI (Republic of Korea), in cooperation with the Republic of Chile. The Joint ALMA Observatory is operated by ESO, AUI/NRAO and NAOJ. The National Radio Astronomy Observatory is a facility of the National Science Foundation operated under cooperative agreement by Associated Universities, Inc.

Facilities: ALMA

Software: astropy, XCLASS, spectral-cube, CASA

#### **APPENDIX**

#### A. CATALOGS FOR MOLECULAR DATA

### REFERENCES

Aikawa, Y. 2013, Chemical Reviews, 113, 8961

Belloche, A., Maury, A. J., Maret, S., et al. 2020, A&A, 635, A198

Bottinelli, S., Ceccarelli, C., Lefloch, B., et al. 2004, ApJ, 615, 354

Carney, M. T., Yıldız, U. A., Mottram, J. C., et al. 2016, A&A, 586, A44

Endres, C. P., Schlemmer, S., Schilke, P., Stutzki, J., & Müller, H. S. P. 2016, Journal of Molecular Spectroscopy, 327, 95

Graninger, D. M., Wilkins, O. H., & Öberg, K. I. 2016, ApJ, 833, 125

Herbst, E., & van Dishoeck, E. F. 2009, ARA&A, 47, 427Higuchi, A. E., Sakai, N., Watanabe, Y., et al. 2018, ApJS, 236, 52

Imai, M., Oya, Y., Sakai, N., et al. 2019, ApJL, 873, L21
Imai, M., Sakai, N., Oya, Y., et al. 2016, ApJL, 830, L37
Jørgensen, J. K., van der Wiel, M. H. D., Coutens, A., et al. 2016, A&A, 595, A117

Lee, C.-F., Li, Z.-Y., Ho, P. T. P., et al. 2017, ApJ, 843, 27 McGuire, B. A. 2018, ApJS, 239, 17

Möller, T., Endres, C., & Schilke, P. 2017, A&A, 598, A7

Müller, H. S. P., Schlöder, F., Stutzki, J., & Winnewisser, G. 2005, Journal of Molecular Structure, 742, 215

Müller, H. S. P., Thorwirth, S., Roth, D. A., & Winnewisser, G. 2001, A&A, 370, L49

Oya, Y., Sakai, N., López-Sepulcre, A., et al. 2016, ApJ, 824, 88

Oya, Y., Sakai, N., Watanabe, Y., et al. 2017, ApJ, 837, 174
Pickett, H. M., Poynter, R. L., Cohen, E. A., et al. 1998, JQSRT, 60, 883

Sakai, N., Sakai, T., Hirota, T., Burton, M., & Yamamoto, S. 2009, ApJ, 697, 769

Sakai, N., Sakai, T., Hirota, T., & Yamamoto, S. 2010, ApJ, 722, 1633

Sakai, N., Sakai, T., Hirota, T., et al. 2014, Nature, 507, 78
 Sakai, N., Oya, Y., López-Sepulcre, A., et al. 2016, ApJL,
 820, L34

Stephens, I. W., Dunham, M. M., Myers, P. C., et al. 2018, ApJS, 237, 22

Stephens, I. W., Bourke, T. L., Dunham, M. M., et al. 2019, ApJS, 245, 21

Tychoniec, L., Tobin, J. J., Karska, A., et al. 2018, ApJS, 238, 19