Methanol masers and millimetre lines: a common origin in protostellar envelopes
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CHAPTER 3

Distribution and excitation of thermal methanol in 6.7 GHz maser bearing star-forming regions
I. The nearby source Cepheus A

Abstract

Candidate high-mass star-forming regions can be identified through the occurrence of 6.7 GHz methanol masers. In these sources the methanol abundance of the gas must be enhanced, because the masers require a considerable methanol path length. The place and time of origin of this enhancement is not well known. Similarly, it is debated in which of the physical components of the high-mass star-forming region the masers are located. The aim of this study is to investigate the distribution and excitation of the methanol gas around Cep A and to describe the physical conditions of the region. In addition the large-scale abundance distribution is determined in order to understand the morphology and kinematics of star-forming regions in which methanol masers occur. The spatial distribution of methanol is studied by mapping the line emission, as well as the column density and excitation temperature, which are estimated using rotation diagrams. For a limited number of positions the parameters are checked with non-LTE models. Furthermore, the distribution of the methanol abundance is derived in comparison with archival dust continuum maps. Methanol is detected over a $0.3 \times 0.15 \text{ pc}$ area centred on the Cep A HW2 source and shows an outflow signature. Most of the gas can be characterised by a moderately warm rotation temperature ($30$–$60 \text{ K}$). At the central position two velocity components are detected with different excitation characteristics, the first related to the large-scale outflow. The second component, uniquely detected at the central location, is probably associated with the maser emission on much smaller scales of $2''$. A detailed analysis reveals that the highest densities and temperatures occur for these inner components. In the inner region the dust and gas are shown to have different physical parameters.

Abundances of methanol in the range $10^{-9} - 10^{-7}$ are inferred, with the abundance peaking at the maser position. The geometry of the large-scale methanol is in accordance with previous determinations of the Cep A geometry, in particular those from methanol masers. The dynamical and chemical time-scales are consistent with a scenario where the methanol originates in a single driving source associated with the HW2 object and the masers in its equatorial region.
3.1 Introduction

While massive stars ($\geq 10 \, M_\odot$) have an enormous impact on the evolution of the Galaxy during their life and death, their birth process is still shrouded in mystery (e.g. Kurtz 2005). To a large extent this is because of the environment in which the high-mass stars form. Most high-mass stars are observed at large distances (compared to low-mass stars), evolve rapidly, and form deeply embedded in their molecular clouds. Additionally, they typically form in clusters. Thus, very high-resolution data are needed to resolve them and the disruptive influence they have on their direct environments (e.g. Martín-Pintado et al. 2005).

Methanol maser sources are exciting targets for the detailed study of high-mass star-formation. Continuum emission of warm dust at sub-millimetre wavelengths has been detected at well over 95% of the observed 6.7 GHz methanol maser sites (Hill et al. 2005), while 25% of the same masers are associated with a detectable ultra-compact (UC) H II region (Walsh et al. 1998). This seems to indicate that the masers probe a range of early phases of massive star-formation and that the methanol maser emission disappears when the UC H II region is created (Walsh et al. 1998). The maser sources require high methanol

Figure 3.1: Sample spectra from the 3 positions identified in Sec. 3.3.2. **Top** spectrum at the "Centre" position, coincident with the HW2 object. **Bottom Left** at the position of "NE Outflow" and **Bottom Right" Envelope". The small symbols at the top indicate the frequencies of methanol transitions in this band.
abundance for their excitation, which points to a recent release of methanol into the gas phase (van der Tak et al. 2000a). We can thus expect to observe the thermal methanol lines in these sources, which will then yield diagnostics for the large-scale kinematics and excitation. Besides being signposts, masers are excellent tracers of the geometry and small-scale dynamics of these regions. Much effort has focused on studying the kinematics, and several claims of circumstellar disks, expanding spherical shells, and jets have been presented (e.g. Pestalozzi et al. 2004, Minier et al. 2002, Bartkiewicz et al. 2005b, 2009). The evidence suggests that masers trace an evolutionary sequence or possibly a mass range in the formation process, or even both (e.g. Breen et al. 2010). However, it is not clear exactly with which of the physical structure(s) of the complex surroundings of young massive stars they are associated. By trying to make a link between the maser distribution and the large-scale methanol kinematics and excitation, one can attempt to reach a better understanding of the physical and chemical state of the gas of the regions in which masers occur.

To carry out such a study, we have observed thermal methanol gas towards a sample of 15 high-mass star-forming regions for which we have very long baseline interferometry (VLBI) data of the 6.7 GHz methanol maser. The observations yield maps of the distribution and physical condition of the methanol gas in relation to the methanol maser. These observations of methanol allow one to probe a reasonably wide range of temperature and density, because of its closely spaced $\Delta K = 0$ ($\Delta J = 1$) transitions with a wide range in excitation energies and critical densities (Leurini et al. 2004). Moreover, we can measure the methanol abundance and velocity field. Clearly, the methanol abundance needs to be significant in these regions ($> 10^{-7}$, Sobolev et al. 1997); evaporation of CH$_3$OH from the dust grains in these regions, at temperatures $\sim$100 K is thought to be responsible for this. This specific enhancement is supposed to be so short-lived that it can be used as a chemical clock of the region (van der Tak et al. 2000a). The enhancement can be used to determine the age of these regions and thus the evolutionary stage of the star-formation process.

In this chapter, we describe the observations and the procedure we used to analyse the data. Because of its proximity and because a wealth of data exists on this source, we will focus on Cepheus A East as a first example. The other sources are presented in Chapter 4.

At a distance of 700 pc (Moscadelli et al. 2009), Cepheus A East is one of the closest high-mass star-forming regions. The region has a luminosity of $2.3 \times 10^4$ L$_\odot$ (Evans et al. 1981) (scaled to the distance of 700 pc), out of which half is thought to come from the well-studied object HW2 (Garay et al. 1996), which is also the site of the 6.7 GHz methanol maser emission. The object shows both large-scale molecular outflows over $\sim$1′ (Gómez et al. 1999) and signs of collapse over a similar extent (Bottinelli & Williams 2004, Sun & Gao 2009). At the core of HW2, a thermal jet extends over $\sim$2″ and proper motion outflow velocities of $\sim$500 kms have been measured in the radio (Hughes & Wouterloot 1984, Curiel et al. 2006). Although a superposition of gas components at different positions cannot be completely ruled out (Brogan et al. 2007, Comito et al. 2007), the evidence is building that there is a disk at the centre of the jet, which is observable in both molecular gas and dust (Patel et al. 2005, Torrelles et al. 2007, Jiménez-Serra et al. 2007), as argued in detail in Jiménez-Serra et al. (2009). VLBI observations of the
6.7 GHz methanol maser suggest that the methanol masers originate in a ring-like structure that extends over ~2” and straddles the waist of HW2, see Chapter 2 (Sugiyama et al. 2008b, Vlemmings et al. 2010, Torstensson et al. 2011).

In Sec. 3.2 we describe the observations and calibration methods. The results are presented in Sec. 3.3 in the form of a number of sample spectra and maps of integrated line strength and velocity. To study the large-scale methanol excitation we construct rotation diagrams and produce maps of the derived physical quantities in Sec. 3.4. We present non-LTE calculations for a number of positions to evaluate the limitations of the diagram analysis in Sec. 3.4.4. Subsequently we discuss estimates of the hydrogen column density before we present the methanol abundance in Sec. 3.4.6. The relation of our findings to the Cep A star-forming process and in particular the occurrence of the methanol maser are discussed in Sec. 3.5.

3.2 Observations and data reduction

The JCMT\(^1\) observations of Cep A East (HW2) \((22^h 56^m 17.9^s +62^\circ 01' 49'' \ (J2000))\) were performed on June 2 and June 17, 2007. We used the array receiver HARP, which has 16 receptors with a spacing of ~30” in a 4×4 grid, resulting in a footprint of ~2’. The receptors are single sideband receivers with a sideband rejection of >10 dB. We used the observing mode HARP5, a type of beam-switching (jiggle-chop) mode where several short scans at different positions of the target source (10 s per jiggle position) are observed before switching to the off-position. As a result, several on-source observations share the same off-source observation, which minimises the overhead while still providing good baselines. The observing mode results in a map of 20×20 pixels covering 2’×2’, thus each pixel is 6”x6”. This ensures proper Nyquist sampling of the 14” telescope beam at 338 GHz.

The ACSIS correlator backend was set up with a 1 GHz (880 kms) bandwidth and 2048 channels centred on the methanol \(7_0 \rightarrow 6_0\) A\(^+\) line at 338.41 GHz. The frequency set-up allowed us to cover 25 CH\(_3\)OH lines (Table 3.1) with a velocity resolution of ~0.43 kms (488 kHz). In total we have an effective (on source) integration time of ~5 min per pixel with a typical system temperature \((T_{\text{sys}})\) of ~280 K. For our analysis we adopted a main-beam efficiency \((\eta_{\text{mb}})\) of 0.6. Regular pointings were made of the normal calibrators, and we estimate to have an absolute pointing accuracy of ~1”.

We used the Starlink package (Gaia and Splat) for an initial inspection of the data. It was found that receptor R06 suffered from bad baselines (large ripples in some scans). For that receptor only these poor scans were removed, after which the scans were combined. The data were then converted into the GILDAS/CLASS format, and the remaining data reduction and analysis was performed in CLASS. A linear baseline was fitted to several spectral regions without any line emission and subtracted from the spectra. Then the data

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\(^1\)The James Clerk Maxwell Telescope is operated by the Joint Astronomy Centre on behalf of the Science and Technology Facilities Council of the United Kingdom, the Netherlands Organisation for Scientific Research, and the National Research Council of Canada.
3 Distribution and excitation of thermal methanol in Cepheus A

**Table 3.1:** CH$_3$OH line data for the observed transitions, adopted from the CDMS (Cologne Database of Molecular Spectroscopy, Müller et al. (2005)). All lines are from the $J = 7 \rightarrow 6$ band, and throughout the chapter a notation like “$-6 E$” refers to the $J = 7_{-6} \rightarrow 6_{-6} E$ transition. Blended lines are indicated by a $^*$. 

<table>
<thead>
<tr>
<th>Frequency (MHz)</th>
<th>$\mu_d S_g$</th>
<th>$E_u$ (K)</th>
<th>Transition</th>
</tr>
</thead>
<tbody>
<tr>
<td>337969.414</td>
<td>5.55</td>
<td>390.1</td>
<td>$-1 A v_7=1$</td>
</tr>
<tr>
<td>338124.502</td>
<td>5.65</td>
<td>78.1</td>
<td>0 E</td>
</tr>
<tr>
<td>338344.628</td>
<td>5.55</td>
<td>70.6</td>
<td>$-1 E$</td>
</tr>
<tr>
<td>338404.580</td>
<td>1.49</td>
<td>243.8</td>
<td>$+6 E$</td>
</tr>
<tr>
<td>338408.681</td>
<td>5.66</td>
<td>65.0</td>
<td>0 A</td>
</tr>
<tr>
<td>338430.933</td>
<td>1.50</td>
<td>253.9</td>
<td>$-6 E$</td>
</tr>
<tr>
<td>338442.344*</td>
<td>1.49</td>
<td>258.7</td>
<td>$+6 A$</td>
</tr>
<tr>
<td>338442.344*</td>
<td>1.49</td>
<td>258.7</td>
<td>$-6 A$</td>
</tr>
<tr>
<td>338456.499</td>
<td>2.76</td>
<td>189.0</td>
<td>$-5 E$</td>
</tr>
<tr>
<td>338475.290</td>
<td>2.76</td>
<td>201.1</td>
<td>$+5 E$</td>
</tr>
<tr>
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<td>2.77</td>
<td>202.9</td>
<td>$+5 A$</td>
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<tr>
<td>338486.337*</td>
<td>2.77</td>
<td>202.9</td>
<td>$-5 A$</td>
</tr>
<tr>
<td>338504.099</td>
<td>3.80</td>
<td>152.9</td>
<td>$-4 E$</td>
</tr>
<tr>
<td>338512.627*</td>
<td>3.81</td>
<td>145.3</td>
<td>$-4 A$</td>
</tr>
<tr>
<td>338512.639*</td>
<td>3.81</td>
<td>145.3</td>
<td>$+4 A$</td>
</tr>
<tr>
<td>338512.856*</td>
<td>5.23</td>
<td>102.7</td>
<td>$-2 A$</td>
</tr>
<tr>
<td>338530.249</td>
<td>3.82</td>
<td>161.0</td>
<td>$+4 E$</td>
</tr>
<tr>
<td>338540.795*</td>
<td>4.60</td>
<td>114.8</td>
<td>$+3 A$</td>
</tr>
<tr>
<td>338543.204*</td>
<td>4.60</td>
<td>114.8</td>
<td>$-3 A$</td>
</tr>
<tr>
<td>338559.928</td>
<td>4.64</td>
<td>127.7</td>
<td>$-3 E$</td>
</tr>
<tr>
<td>338583.195</td>
<td>4.62</td>
<td>112.7</td>
<td>$+3 E$</td>
</tr>
<tr>
<td>338614.999</td>
<td>5.68</td>
<td>86.1</td>
<td>$+1 E$</td>
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<td>338639.939</td>
<td>5.23</td>
<td>102.7</td>
<td>$+2 A$</td>
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<tr>
<td>338721.630</td>
<td>5.14</td>
<td>87.3</td>
<td>$+2 E$</td>
</tr>
<tr>
<td>338722.940</td>
<td>5.20</td>
<td>90.9</td>
<td>$-2 E$</td>
</tr>
</tbody>
</table>
were smoothed in the spectral domain to improve the signal-to-noise ratio, which resulted in a velocity resolution of 0.87 kms and an rms of ∼65 mK.

The analysis was performed on a pixel-by-pixel basis by first fitting a Gaussian to the strongest unblended line (7−1 → 6−1 E) in the spectra. The measured velocity and width of this line were then used to place windows around all other features in the spectra for which the emission was integrated. Owing to some overlap between different lines, only lines with a >5σ (∼0.6 K km s⁻¹) detection are included in the subsequent analysis. In the centre area, at the position of HW2, a second velocity component can be distinguished. Where we were able to separate the two velocity components, we performed a manual analysis by fitting individual Gaussians to the lines.

3.3 Results

3.3.1 Methanol lines

In the following we refer to each pixel in the map with its respective offset coordinates (Δα, Δδ), measured in arc seconds, with respect to the centre of the map (J2000 22h56m17.88s +62°01′49.2″). Fig. 3.1 shows the observed spectrum at three positions: (+3″,3″) “Centre”, (+27″,+9″) “NE outflow”, (+21″,-3″) “Envelope”, where the designations can be understood from the subsequent discussion in Sec. 3.3.2. Focusing at the centre spectrum, all but four of the detected lines are originating from methanol (indicated by ticks at the top); the others are identified as SO₂, 34SO₂, and H₂CS. The CH₃OH K=+1 E line is a blend with another SO₂ line, which is also indicated in the figure.

In the centre spectrum, two gas components are observed: the main component with v_lsr = −10.5 km s⁻¹ and a secondary red shifted at −5.2 km s⁻¹, as can be seen in the profile of the −1 E transition (Fig. 3.2). We note that the red-shifted velocity profile is overlapping with the range spanned by the methanol masers, which are observed with velocities between v_lsr = −4.6 and −1.5 km s⁻¹ (Chapter 2). The redshifted component is most readily identified in the higher K transitions. While the first component is brightest in the low K transitions and not detected in the higher K lines, the −5.2 km s⁻¹ component is brighter in the higher K lines and less bright in the lower K lines. For the K=3 transitions, the two components are of similar strength. There is a weak detection of the methanol isotopologue ¹³CH₃OH J=13 → 12, ΔK = +1 in this component, which indicates that optical depth effects may be important for this. Moreover, a careful analysis, which includes averaging over a few pixels, shows that the torsionally excited −1 A v₁=1 line at 337.969 GHz is detected in the “Centre” region in the −5.2 km s⁻¹ velocity component.

Otherwise the observed line profile appears Gaussian in all components and does not show evidence of blue/redshifted wings. However, a modest velocity shift of the line is observed across the source with lower velocities to the NE and higher velocities to the SW (Fig. 3.2).

To begin a consistent analysis of all methanol line emission in our data cube, we extracted the integrated line flux from each line as described above (Sec. 3.2). To illustrate our procedure, we list the integrated line emission of the spectral features for the main
Figure 3.2: Velocity profiles of the -1 E line at three positions. Although the profiles are Gaussian in the outflow, there is a clear shift in the velocity. At the “Centre” position there are clearly two velocity components. The dotted line is taken to be the system velocity, corresponding to the velocity of the dominant component at the “Centre” position.
Table 3.2: Line parameters for the main velocity component measured at the position of HW2 (+3”, +3”)

<table>
<thead>
<tr>
<th>Line $K$, type$^1$</th>
<th>Flux $K$ km s$^{-1}$</th>
<th>Width km s$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 E</td>
<td>2.16</td>
<td>4.21</td>
</tr>
<tr>
<td>−1 E</td>
<td>3.30</td>
<td>4.79</td>
</tr>
<tr>
<td>0 A</td>
<td>3.60</td>
<td>4.79</td>
</tr>
<tr>
<td>−6 E</td>
<td>0.03</td>
<td>1.74</td>
</tr>
<tr>
<td>+6 A</td>
<td>0.29</td>
<td>3.80</td>
</tr>
<tr>
<td>−6 A</td>
<td>0.29</td>
<td>3.80</td>
</tr>
<tr>
<td>−5 E</td>
<td>0.61</td>
<td>4.81</td>
</tr>
<tr>
<td>+5 E</td>
<td>0.31</td>
<td>3.15</td>
</tr>
<tr>
<td>+5 A</td>
<td>0.36</td>
<td>4.18</td>
</tr>
<tr>
<td>−5 A$^2$</td>
<td>0.36</td>
<td>4.18</td>
</tr>
<tr>
<td>−4 E</td>
<td>1.05</td>
<td>4.79</td>
</tr>
<tr>
<td>−2 A$^3$</td>
<td>2.00</td>
<td>4.79</td>
</tr>
<tr>
<td>+4 E</td>
<td>1.12</td>
<td>4.79</td>
</tr>
<tr>
<td>+3 A</td>
<td>1.37</td>
<td>4.79</td>
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<tr>
<td>−3 A$^4$</td>
<td>1.37</td>
<td>4.79</td>
</tr>
<tr>
<td>−3 E</td>
<td>0.93</td>
<td>4.07</td>
</tr>
<tr>
<td>+3 E</td>
<td>0.94</td>
<td>4.97</td>
</tr>
<tr>
<td>+2 A</td>
<td>1.42</td>
<td>5.38</td>
</tr>
<tr>
<td>+2 E</td>
<td>1.55</td>
<td>4.38</td>
</tr>
<tr>
<td>−2 E$^5$</td>
<td>1.55</td>
<td>4.38</td>
</tr>
</tbody>
</table>

$^a$See Table 1 for transition details
$^b$Blend with +5 A
$^c$Blend with ±4 A
$^d$Blend with +3 A
$^e$Blend with +2 E

velocity component at the “Centre” position from Fig. 3.1 in Table 3.2. For lines that are blended we chose a simple strategy. If the strengths ($S_{\phi\mu_2}$) of the lines are similar and they have comparable upper energy levels, we split the measured flux between the two lines. If, however, one line has a significantly higher line strength and lower upper energy level, we attribute the entire emission to the strong line. For the blend between the $K=-2$ A and $K=±4$ A, where the lines have comparable line strengths and energy levels, we assume the total measured flux as an upper limit to all lines.

### 3.3.2 Spatial distribution of methanol

Fig. 3.3 shows the spatial distribution of the flux, the central velocity, and the width of the methanol $7_{-1} \rightarrow 6_{-1}$ E line, which is the brightest unblended feature in the spectra. The
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Figure 3.3: From top to bottom (a-c) the integrated flux, the velocity field, and the line width of the methanol $^{7-1} \rightarrow {6-1}$ E-type transition, the strongest unblended line in the observed spectra. In all maps the cross marks the “Centre” position, the diamond “NE Outflow” and the circle indicates the position where the “Envelope” spectra are evaluated.

methanol maser emission arises just to the NE of the centre of the map (+0.7′′, +0.4′′) and is indicated with a cross in all figures. The white area in the maps represents regions where the line flux is less than our $5\sigma$ detection limit of 0.6 K km s$^{-1}$.

The line emission is constrained to an area that is measured from the points that are farthest apart from each other in the intensity map (Fig. 3.3a), with an extent of ~85′′ by 43′′ (~0.29 pc by ~0.15 pc), and is orientated in the NE-SW direction. The velocity map (Fig. 3.3b) shows that the methanol gas is entrained in an outflow with the blueshifted emission to the NE and the redshifted emission to the SW. Measured between the highest and lowest velocities in the map, the outflow has a modest velocity gradient of ~3.1 kms$^{-1}$ over ~0.20 pc. The integrated line flux peaks in the centre of the map at ~5.2 K km s$^{-1}$, with a bright extension to the SW over ~13′′ (~0.044 pc), which is associated with the red-shifted emission. Both the velocity field and line width maps (Figs. 3.3b and c) show a maximum in the centre of the map to the N of HW2. This is because of a blending
of the two velocity components that we are unable to decompose. A second maximum in the integrated line flux is observed ∼28″ (∼0.095 pc) to the NE of the centre, which is associated with the blueshifted part of the emission and a secondary peak in the line width map. Throughout the chapter we will refer to this position as the “NE outflow”. The quiescent methanol emission that is not associated with the outflows and does not show any peak in the integrated line flux map we will refer to as the “Envelope”.

3.4 Analysis

3.4.1 Rotation diagrams

From the above spectral cuts in three positions and the brightness distribution of one of the lines, it is clear that the observations trace a complex region in which the dynamic and excitation characteristics change, at least on the scales represented by the beam size of the telescope. To make a representation of the large-scale distribution and excitation of methanol, we created rotation diagrams (Boltzmann plots) at every pixel to be able to study the characteristics of the excitation distribution. Inherent to the method are a number of assumptions, namely that the gas at each velocity and spatial resolution element can be described by a single excitation temperature ($T_{\text{rot}} \equiv T_{\text{ex}}$), that all the lines are optically thin, i.e. $\tau << 1$, and that the size of the emitting region is the same for all lines. Although not all these assumptions may hold for all positions, the strength of the method is that it shows general trends on the relevant scales of our observations. In Sec. 3.4.4 we will test the validity of these assumptions.

For each pixel at which we measured at least three methanol transitions with $S/N > 5$, we plotted the upper energy level $E_u$ [K] of the transitions on the x-axis versus the logarithm of the column density in the upper energy state divided by the statistical weight of that upper level $\log(N_u/g_u)$ on the y-axis. The weighted column density ($N_u/g_u$) can be calculated with Eq. 3.1 (Helmich et al. 1994) and the appropriate coefficients in Table 3.1.

\[
\frac{N_u}{g_u} = \frac{N_M}{Q(T_{\text{rot}})} e^{-E_u/T_{\text{rot}}} \approx \frac{1.67 \cdot 10^{14}}{\nu \mu^2 S_g} \int T_{\text{MB}} dv
\]

By fitting a straight line through these data points we determine the rotation temperature $T_{\text{rot}}$ [K] and column density $N_M$ [cm$^{-2}$] of the methanol gas. The rotation temperature is the negative inverse of the slope of the fitted line and the total column density is where the extrapolated line crosses the y-axis.

Fig. 3.4 shows the resulting rotation diagrams of the “Envelope” (the lower temperature quiescent gas), the “NE outflow”, and of the first velocity component at the “Centre” position of HW2. For the two positions in the outflow and envelope a single gas component provides a good fit to the data points with a rotation temperature of 30 K to 35 K and
Figure 3.4: Rotation diagrams for three specific components and a population diagram for the fourth component identified in Sec. 3.3.2. Triangles indicate upper limits and error bars the 1σ standard deviation in the flux determination. The diagrams displayed are a: rotation diagram for the main, −10.5 km s$^{-1}$ velocity component at the “Centre” position (+3′′,+3′′). b: population diagram for the redshifted, −5.2 km s$^{-1}$ velocity component at the ”Centre” position (+3′′,+3′′); for this model asterisks indicate a best fit that includes optical depth effects and beam dilution (see Sec. 3.4.2), c: rotation diagram for the “NE Outflow” (+27′′,+9′′), d: rotation diagram for “Envelope” (+21′′,-3′′).
a column density of $10^{16}$ cm$^{-2}$. However, in the centre region things are more complex. The velocity component at $-10.5$ km s$^{-1}$ can be fitted by a straight line, which yields a rotation temperature of 224 K and a column density of 2$\times$10$^{15}$ cm$^{-2}$. For the $-5.2$ km s$^{-1}$ velocity component, the points lie nearly horizontal in the diagram and we were only able to put a lower limit of $>300$ K on the rotation temperature with the simple fit.

3.4.2 Population diagram modelling

Clearly the basic assumptions going into the rotation diagram analysis are violated for the “Centre” position, and a successful explanation must accommodate both an apparently negative rotation temperature and the relative weakness of the low $K$ lines for the redshifted component. Non-LTE conditions, optical depth and beam-dilution effects can all play a role here. To investigate the effect of the line optical depth on the above results, we adopted, as a first step, the method of Goldsmith & Langer (1999), which includes the source size ($\theta_s$) as a free parameter and allows for optical depth effects. We performed a $\chi^2$ analysis for $10 < T_{\text{rot}} < 500$ K, $10^{14} < N_M < 10^{18}$ cm$^{-2}$, and $1'' \leq \theta_s \leq 14''$, i.e. a beam dilution factor between 200 and 1. For the redshifted component, we find the best fit to be for a rotation temperature of 125 K, a column density of $4 \times 10^{17}$ cm$^{-2}$, and $\theta_s = 1''$, which is much smaller than the beam. The merit of this model can be seen in panel Fig. 3.4b. In this specific case the maximum optical depth encountered was $\tau = 6.8$. Because this seems to be the component with the highest column density, it is not surprising that $^{13}$CH$_3$OH is detected in this component. Clearly, the highly excited $K$ lines lead the model towards a high column density, but at the same time, the only way for the model to accommodate the lower $K$ lines by increasing the optical depth is to decrease the source size. It should be noted, however, that this analysis assumes a single excitation temperature. We will discuss non-LTE modelling in Sec. 3.4.4.

3.4.3 Spatial distribution of the excitation

Maps of the derived rotation temperatures and column densities are presented in Figs. 3.5 and 3.6. In the maps we excluded the results of the $-5.2$ km s$^{-1}$ velocity component seen at the position of HW2. The methanol rotation temperature ranges from $\sim20$ K to $\sim100$ K over most of the map, but shows a pronounced peak of $\geq200$ K close to the HW2 position. We find column densities between $10^{15}$ cm$^{-2}$ and $3 \times 10^{16}$ cm$^{-2}$. Overall, the methanol column density distribution is quite smooth and shows a minimum towards the centre position. In Sec. 3.4.4 we will investigate whether this minimum is an artefact of the rotation diagram method. Additionally, some care must be taken when analysing the results in the centre region. Only at the position of HW2 are we able to separate the two velocity components of the gas. In the neighbouring pixels the integrated line fluxes are therefore overestimated due to blending.
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Figure 3.5: Map of derived rotation temperatures. Symbols are as in Fig. 3.3.

Figure 3.6: Map of derived column densities. Symbols are as in Fig. 3.3.
3.4.4 Non-LTE analysis

Method

To check the validity of the assumptions of the rotation diagram analysis, we ran a set of non-LTE models. By comparing the resulting synthetic spectra with the observed spectra, these models also allow us to investigate the physical conditions (i.e., temperature and density) at distinct positions in the methanol-emitting region. We used the RADEX\(^2\) package, which uses statistical equilibrium for the population calculations and an escape-probability method to calculate the optical depth (van der Tak et al. 2007). The main assumptions are that the medium is uniform and that the optical depth is not too high (\(\lesssim 100\)). The most accurate collisional rates are available for the first 100 levels of methanol (Pottage et al. 2004, Schöier et al. 2005), and we therefore restrict this part of the analysis to the 24 relevant lines from Table 3.1.

Initial runs of the non-LTE model indicated that the methanol excitation is more strongly governed by the density than by the temperature. Using a uniform sphere geometry, we considered kinetic temperatures between 30 and 300 K and \(\text{H}_2\) densities between \(10^4\) and \(10^7\) cm\(^{-3}\). Even at the lowest temperature all lines, except the \(K=\pm 5\) and \(K=\pm 6\) transitions, are excited in the high-density case. On the other hand, at low-density and high-temperature, only the lowest three \(K\) levels (\(K=0,\pm 1,\pm 2\)) are populated. A combination of high temperature and high density is required to excite all lines, which renders this methanol band useful as a density and temperature tracer (Leurini et al. 2004).

Starting with model values comparable to the results from Sec. 3.4.1, it is possible to model the observed line strengths of the large-scale emission with the simple assumption that the emitting region is filling the beam. For these parameters all lines are optically thin (\(\tau < 0.3\)), which validates the assumption made in the rotation diagram analysis for the large-scale emission.

In the above calculations the background-radiation field was represented by the cosmic microwave background with \(T_{\text{bg}} = 2.73\) K. Models with background-radiation temperatures in the range 3 – 300 K were run, but this has no clear impact on the lines considered here. Mostly the contrast of the lines with respect to the background was reduced without changing their relative intensities. This result can be understood from the fact that our models consider only the torsional-vibrational ground state of methanol (Leurini et al. 2007). These models are inconclusive with regard to the influence of the radiation field, which is expected to be important in the maser region.

Non-LTE results

Then we used the non-LTE calculations to fit the spectra for the four regions identified in Sec. 3.3.2 (the velocity components at “Centre”, the “NE outflow”, and the quiescent gas in the “Envelope”). In a search for minima of the reduced \(\chi^2\) we used a model grid that includes column densities from \(10^{13}\) to \(10^{19}\) cm\(^{-2}\) at every half dec, densities from \(10^4\)

\(^2\)http://www.strw.leidenuniv.nl/~moldata/radex.html
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Table 3.3: Results of the non-LTE models.

<table>
<thead>
<tr>
<th>Position</th>
<th>Δα</th>
<th>Δδ</th>
<th>log(N$_{\text{M}}$)$_T$</th>
<th>T$_{\text{kin}}$</th>
<th>log(n$_{\text{H}_2}$)$_T$</th>
<th>T$_{\text{rad}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Centre, main</td>
<td>+3</td>
<td>+3</td>
<td>16.0</td>
<td>90</td>
<td>4.0</td>
<td>30</td>
</tr>
<tr>
<td>Centre, red-shifted&lt;sup&gt;b&lt;/sup&gt;</td>
<td>+3</td>
<td>+3</td>
<td>17.0</td>
<td>130</td>
<td>7.0</td>
<td>100</td>
</tr>
<tr>
<td>Outflow (NE)</td>
<td>+27</td>
<td>+9</td>
<td>15.0</td>
<td>90</td>
<td>5.0</td>
<td>30</td>
</tr>
<tr>
<td>Envelope</td>
<td>+21</td>
<td>-3</td>
<td>15.0</td>
<td>90</td>
<td>5.0</td>
<td>30</td>
</tr>
</tbody>
</table>

<sup>a</sup> Offset from centre of map RA 22$^h$56$^m$17.88$^s$, DEC +62$^°$01′49.2″.

<sup>b</sup> This component was modeled with a beam dilution factor of 50.

up to $10^9$ cm$^{-3}$, kinetic temperatures between 30 and 300 K, and radiation temperatures in the range 3 – 400 K. Based on our findings in Sec. 3.4.1 we adopted a beam dilution factor of 50 for the redshifted emission in the “Centre”, corresponding to the 2″ extent of the region where the masers appear. It is found that the models are able to reproduce the lines at all four positions to within 5-10% accuracy. In Fig. 3.7 we plot the observed integrated flux and the synthetic integrated flux calculated with RADEX versus $7K \rightarrow 6K$ number. Also indicated is our 3σ detection limit.

Like before, this analysis is complicated somewhat by the blending of lines by methanol and in the case of the $K=+1$ E by blending with SO$_2$. For the $K=+1$ E line we did not include it in the analysis of the centre position components because the SO$_2$ contributes significantly to it. For the “NE outflow” and for the “Envelope” the SO$_2$ does not appear to contribute significantly, because the other SO$_2$ line is not detected, and we included it in the analysis. At the position of HW2 the methanol gas is highly excited, the blending of methanol lines becomes more severe and confusion prevents one from detecting the higher ($K\geq5$) lines. Also, at this position the $K=\pm2$ lines show a substantial (≈ 50%) deviation from the RADEX model. The reason for this discrepancy is yet unclear, but could be another indication that we do not treat radiative processes completely.

The results of our non-LTE analysis are summarised in Table 3.3. The coolest and most diffuse gas is found in the envelope and outflow, where one should take into account that the granularity of the model grid probably prevents the distinction of small temperature and density differences. In comparison, the gas associated with the rotation temperature peak is much denser and also warm. We find the highest density and temperature in the “Centre” HW2 position where the maser emission occurs. However, given the large beam dilution and the limited set of available levels for collisional and radiative excitation, these numbers should be treated cautiously.

The non-LTE analysis confirms that there are no steep column density gradients across the source. But although generally the values agree with our rotation diagram analysis, they differ in details. In particular, the non-LTE analysis finds local maxima of the column density in the “Centre” components. Although the rotation diagram maps show some shortcomings for the inner components, they are useful for outlining the qualitative large-
3.4 Analysis

Figure 3.7: Plots of the integrated intensity for different CH$_3$OH $7K \rightarrow 6K$ transitions and the best $\chi^2$ model values. In the panels the observed flux (open symbols) is plotted, as well as the non-LTE model values (plusses and crosses). Also indicated is the 3$\sigma$ detection limit. Circles and plusses (blue) are used for E-type lines and squares and crosses (red) for A-type. Labels on the top identify blended lines, for which the combined flux and model value are displayed. The $K=+1$ line which is blended with an SO$_2$ was not included in the analysis for the centre region components (top panels). The parameters of the fits are listed in Table 3.3. $a$: main, $-10.5$ km s$^{-1}$ velocity component at the “Centre” position with beam dilution factor of 1, $b$: red-shifted, $-5.2$ km s$^{-1}$ velocity component at “Centre” with a beam dilution factor of 50, $c$: the “NE Outflow” and $d$: “Envelope”.

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![Image](image.png)

**Figure 3.8:** H$_2$ column densities derived from the SCUBA dust continuum for a constant dust temperature of 30 K.

scale properties and for constraining the distribution of the excitation; running pixel-based non-LTE models is beyond the scope of this chapter.

### 3.4.5 H$_2$ column density

To estimate the methanol abundance we require a measure of the H$_2$ column density to combine with the estimates of the methanol column density. Therefore we used the SCUBA map of Bottinelli & Williams (2004) and derived the H$_2$ column distribution using the formula of Henning et al. (2000) (eq. 3.2), where $S_ν$ is the flux in units of Jy beam$^{-1}$, $κ_d(λ)$ the dust opacity at the observed wavelength, $Ω_{mb}$ the solid angle of the main beam, $B_ν(T)$ the Planck function (Black body), $m_H$ the mass of the hydrogen atom, and $R$ is the gas-to-dust ratio. In the calculation we used: $κ_d(875μm) = 1.4$ and $R=150$ (Henning et al. 2000).

$$N(H_2) = \frac{S_ν}{κ_d(λ)Ω_{mb}B_ν(T)2m_H}R$$

(3.2)

The calculations were run for two cases. In the first case, we assumed a constant dust temperature of $T_d= 30$ K to derive the H$_2$ column density and the methanol abundance, $X_M$. In the second case we assumed the dust temperature $T_d$ to be equal to the rotation temperature that we derived from the rotation diagram analysis in Sec. 3.4.1 and used that as input to the black-body function.

### 3.4.6 Methanol abundance

The results of the constant temperature hydrogen column density and methanol abundance calculations are presented in Figs. 3.8 – 3.9. The resulting hydrogen column density
3.4 Analysis

Figure 3.9: Methanol abundance derived from the methanol column densities obtained in the rotation diagram analysis and the \( \text{H}_2 \) column densities obtained for the constant dust temperature 30 K.

Fig. 3.8 is the SCUBA map (by Bottinelli & Williams (2004)) scaled by a constant factor to convert it from Jy beam\(^{-1} \) to \( N(\text{H}_2) \) in cm\(^{-2} \). The result is a very smooth map with \( \text{H}_2 \) column densities between \( \sim 10^{21} \text{ cm}^{-2} \) and \( \sim 10^{24} \text{ cm}^{-2} \). The general morphology of the dust emission is similar to that of the methanol gas, though notably the dust emission does not peak at the position of HW2, but closer to the SE at position (-9\",-4\")).

Similarly, the methanol abundance map derived with a constant dust temperature, Fig. 3.9, shows the same structure as the column density map, since there are no large variations in the SCUBA dust map. We derive a methanol abundance of \( X_M \sim 10^{-8.5} \) for the central region. As noted before, the distribution is relatively flat and probably not very accurate for the centre position.

In the second case, when we used the methanol rotation temperature as a measure of the dust temperature, we find the resulting \( \text{H}_2 \) column density to be anti-correlated to the rotation temperature, because the high rotation temperatures derived for the centre region then result in a lower \( \text{H}_2 \) column. In this way we would derive a \( \text{H}_2 \) column of \( \sim 10^{22} \text{ cm}^{-2} \) (much lower than inferred earlier \( \sim 10^{24} \text{ cm}^{-2} \); Martín-Pintado et al. 2005). Although we already encountered the limited validity of the rotation diagram for the “Centre” region in Sec. 3.4.4, clearly there must be a mismatch between the gas and dust temperature on the scales of the JCMT maps. Both the rotation diagram and the non-LTE calculations indicate higher temperatures towards the centre, for which there is little evidence in the dust images, which even peak at an offset of \( \sim 10'' \) from the rotation temperature maximum. Besides the possibility that the rotation temperature is not a good estimate of the kinetic temperature of the gas, it seems plausible that the kinetic temperature of the gas is not equal to the dust temperature at these locations. Therefore, we adopt a constant dust temperature of 30 K to estimate the \( \text{H}_2 \) column density and CH\(_3\)OH abundance, realising that this method will overestimate \( N(\text{H}_2) \) at the centre of the map.
3 Distribution and excitation of thermal methanol in Cepheus A

3.5 Discussion

3.5.1 Outflow morphology

We have presented maps of the large-scale distribution of the thermal methanol associated with Cep A East (Figs. 3.3, 3.5, and 3.6). The emission is constrained to a region of size $0.29 \times 0.15$ pc ($85'' \times 43''$), roughly centred on the location of the radio continuum source HW2, the most massive YSO in the region. Moreover, the velocity field shows that the methanol gas is entrained in a large-scale bipolar outflow. The position angle of this outflow appears to be consistent with the HW2 geometry and earlier results (Gómez et al. 1999).

While the highest flux of the $K=-1$ E line Fig. 3.3a is associated with the receding part of the outflow (towards the SW), which is coincident with the SCUBA peak, the most energetic gas appears at the opposite side of HW2. This is evident already in the velocity width distribution of the brightest line, which has a line width of $>4$ kms, to the NE of HW2, Fig. 3.3c. It becomes clearer when the temperatures are derived from a rotation diagram analysis where the rotation temperature peaks to the NE of HW2 at $\sim 200$ K (Fig. 3.5).

Several authors (Patel et al. 2005, Jiménez-Serra et al. 2007, Torrelles et al. 2007) have inferred obscuring gas in a flattened circumstellar structure of $\sim 1''$ (700 AU) surrounding HW2. The gas shows signs of rotation and is perpendicular to the high-velocity outflow ($\sim 500$ kms) seen in the radio continuum (Curiel et al. 2006). The molecular outflow that we observe in methanol is blueshifted to the NE, which implies that this side of the outflow is pointing towards the observer, Fig. 3.3b. In Chapter 2 we propose a model in which the methanol maser emission arises in a ring-structure in the equatorial plane of HW2. In contrast to the circumstellar molecular material, the masers do not show signs of rotation; rather the observations seem to indicate an infall signature. It is argued that the masers occur in the shock interface between the circumstellar disk and inflow regulated by a large-scale magnetic field (Vlemmings et al. 2010). We derive an inclination of 67.5° and a position angle of 9.3° for the ring in Chapter 2. The masering gas occurs on size scales of $\sim 2''$ and is very likely clumpy on smaller scales $\sim 0.1''$ (Minier et al. 2002). At the resolution of the observations presented in this chapter the masering gas clearly cannot be resolved, nevertheless the geometry that is observed on smaller scales suggests that to the NE of HW2 we have a line of sight into the outflow that probes highly excited gas close to the protostar, which explains why we see such a high rotation temperature in this region. Evidence for this wide-angle molecular outflow has recently been presented by Torrelles et al. (2010).

3.5.2 Methanol distribution

The highest methanol temperatures appear to occur close to the HW2 central source, but we demonstrated in Sec. 3.4.6 that the dust emission is not showing the same distribution. It seems plausible that the highest methanol rotation temperatures occur in a region
where the dust is at least partly destroyed, possibly at the onset of the outflow where the evaporation from the grains occurs that produces the methanol gas in the first place. The considerations on the dust distribution force us to adopt a constant dust temperature in our abundance analysis. On small radii this may have two effects: firstly, the gas-to-dust ratio may be higher because of grain destruction, which increases $N(\text{H}_2)$ and lowers the abundance. Secondly, the dust temperature may be higher than 30 K, which decreases $N(\text{H}_2)$ and increases the abundance. Increasing the gas-to-dust ratio from 150 to 200/250 decreases the abundance with a factor of 0.75/0.60 respectively. Increasing the dust temperature to 60/100/300 K increases the resulting abundance by a factor of 2.3/4.1/13.0 respectively. High optical depth of the dust will lower the derived abundance. These arguments show that at the peak rotation temperature, the methanol abundance is quite sensitive to increased dust temperatures, but is probably correct to within an order of magnitude, without taking beam-dilution effects into account.

The derived abundances of $\sim 10^{-8.5}$ are higher than expected from gas-phase chemistry, but lower than seen in the solid state (Gibb et al. 2004). High optical depth of the lines is ruled out by the excitation calculations, except in the central region. However, it is possible that there the local methanol abundance is still higher. For example, the methanol maser occurs on much smaller scales than those we are probing in these JCMT observations. The total extent of the CH$_3$OH maser is $\sim<2''$. Moreover, we found that the same small-scale enhancement can explain the peculiar excitation of the “Centre” region. This strengthens the idea that the highly excited methanol gas emission is coming from the same region. Assuming that the dust and the corresponding H$_2$ is smoothly distributed, this component can have a higher column density because of the optical depth and beam-dilution, resulting in methanol abundance estimate of $\sim 10^{-7}$ locally. This value is similar to previous values by other authors for maser regions (Menten et al. 1986, 1988) and still higher than expected from gas-phase chemistry and lower than seen in the solid state. Possibly the methanol evaporated from grain mantles some time ago ($\sim 1000$ yr) and is now destroyed by chemical reactions and/or photo-dissociated.

As noted earlier, all the methanol gas seem to be entrained in an outflow, spanning 0.29 pc. The outflow is oriented close to the plane of the sky, assuming an inclination of 67.5° (Chapter 2) we find an outflow velocity of $\sim 5.8$ km s$^{-1}$. The result implies a dynamical timescale of $\sim 2.4 \times 10^4$ yrs. The dynamical timescale agrees well with the chemical timescale (a few $10^4$ years) for which such a methanol enhancement is believed to be possible (van der Tak et al. 2000a). This supports the argument of a common driving source for all the methanol gas in the region. Alternatively, the methanol may be released in the wake/outflow cavity of the jets described by Cunningham et al. (2009): the uv-radiation produced in the shock interface may shine back into the cavity and photo-desorb the methanol, which causes a local enhancement in the methanol column density.

### 3.5.3 Physical conditions

The rotation diagram analysis shows fairly warm gas entrained in a large-scale outflow. Although the non-LTE analysis demonstrates that the methanol band used in this study is
3 Distribution and excitation of thermal methanol in Cepheus A

quite sensitive to collisional excitation even in the low temperature regime, it qualitatively confirms the large-scale presence of warm methanol gas. Over large areas of the source the total methanol column density appears to be fairly constant at \( \sim 10^{15} \text{ cm}^{-2} \), assuming a uniform distribution with unity beam-filling factor.

The fidelity of the rotation diagrams is breaking down particularly in the centre area, where higher excitation lines appear. Especially the red-shifted component requires non-LTE excitation and/or beam-dilution effects to be explained. At these locations a complete analysis requires high-resolution data and non-LTE modelling that includes excitation by infrared radiation. That more methanol levels should be taken into account is demonstrated by the detection of the torsionally excited \(-1 \text{ A } v_1=1\) line at 337.969 GHz in the redshifted component (Leurini et al. 2007). Radiative pumping is also expected from models of methanol maser excitation (Cragg et al. 2005, Sobolev & Deguchi 1994, Sobolev et al. 1997)

3.5.4 Origin of maser emission

Although a wider area around Cep A has been searched for maser emission (Chapter 2), only one place in this source that is so abundant in methanol does show maser emission. We find the maser position to contain warmer gas than the other positions, to have probably the highest column density, and certainly the density appears to have a maximum here at \( \sim 10^7 \text{ cm}^{-3} \). Moreover, this position does stand out as the position where a second kinematic component at \(-5.2 \text{ km s}^{-1}\) is found. Special properties for this component are the detection of the torsionally excited line, and the requirement to invoke a large beam-dilution to explain the ratios of the higher excited lines. This dilution factor matches quite well with the size of the area of < 2" over which masers are detected in Chapter 2 (Sugiyama et al. 2008b, Torstensson et al. 2011). Combining these facts, we identify this component with the origin of the maser action. We note that the densities we derive for this component are still below \(10^9 \text{ cm}^{-3}\), where the maser lines would be quenched according to model calculations (Cragg et al. 2005). The models have methanol temperature ranges and methanol column densities that are consistent with our estimates, but it is hard to test the dust temperature, which must be >100 K for the maser excitation. We note that many of the values estimated here are inferred indirectly and the relevant processes may occur on even smaller scales than reflected by the beam-dilution factor.

The central abundance of \(X_M \sim 10^{-7}\) is not in excess to that found in low-mass star-forming regions, but the methanol column density \(N_M \sim 10^{17} \text{ cm}^{-2}\) clearly is (Maret et al. 2005). This could point to the condition that the high-density, high-temperature region in low-mass star-forming regions is too small to produce a path length long enough, with sufficient methanol column density, to produce maser emission. Jørgensen et al. (2002) show the high-density, high-temperature region for a low-mass protostar to be \(\sim 10-30 \text{ AU}\). In contrast, for a high-mass protostar, the region extends over \(\sim 1000 \text{ AU}\) (Doty et al. 2002), which possibly facilitates the necessary amplification. Alternatively, Pandian et al. (2008) suggest that in low-mass stars the critical infrared pumping is only available in the inner region, where the density is so high that the maser will be quenched. Both effects
favour maser emission in the environments of high-mass protostars and explain that the masers occur on large distances from the central object, as found in Chapter 2.

To further constrain the physical conditions at the position of the methanol maser and discern radiation and density effects, high-J and $v_r = 1$ line observations are required, for example with the CHAMP+ instrument on the APEX telescope. To probe the excitation of methanol gas on small scales (comparable to that of the maser emitting region), high-resolution interferometry observations are needed, with instruments such as the SMA and the IRAM interferometers, and in the longer term, for lower declination sources, ALMA. The interpretation of these observations requires models that include the excitation of CH$_3$OH by infrared radiation.

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