Solving the radiative transfer equation for maser environments

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Abstract. The study of astrophysical maser formation provides a method for probing the chemical composition of the sources they are observed in. In order to understand the pumping mechanisms and physical characteristics of masers, suitable models must contain expressions for each level population from which an inverse function that will cause the mean line intensity to move away from black body form can be derived. Central to this is obtaining numerically consistent solutions to the population density distributions of the maser molecules. The "masers" package is developed in Python and implements the escape probability approximation method. It solves the level population problem for non-LTE statistical equilibrium using molecular data and parameters describing the physical environment. It extends existing radiative transfer software by providing a reasonably fast, stable algorithm that deals with the solution method's inherent sensitivity to oscillations and multiple valid outcomes; allows different maser geometries for calculation; includes the contribution of interacting background radiation fields, as well as other sources of opacity such as line overlap.

1. Introduction

Determining the level population distribution in an arbitrary situation is the central problem in modelling spectral line emission in general and maser radiation in particular. In essence this consists of a two part solution: Solve the rate-equation problem for the populations of the energy levels under conditions that will produce an inversion. Then, calculate the amplification of the maser radiation through the medium sustaining the inversion.

The escape probability approximation method separates population density calculations from the line radiation in the radiative transfer, RT, equation [1]. This is one of the most implemented methods for solving the RT equation and software implementing this method for the study of thermal line emission are readily available. Care should be taken however, since obtaining numeric solutions under conditions causing inversion in maser environments are a little more tricky. In this paper we present easily overlooked numerical failures found during a study investigating physical conditions to investigate population inversion leading to formaldehyde masers.

2. Theory and Calculation

Applying the escape probability to the rate equations, the level populations calculation can be expressed as presented in Equation 2.7.1 from [1].

$$\frac{dn_i}{dt} = -\sum_{j < i} \left\{ A_{ij} \beta_{ij} \left[n_i + W \aleph_{ij} (n_i - n_j) \right] + C_{ij} \left[n_i - n_j \exp\left(\frac{-h\nu_{ij}}{kT}\right) \right] \right\}
+ \sum_{j > i} \frac{g_j}{g_i} \left\{ A_{ji} \beta_{ji} \left[n_j + W \aleph_{ji} (n_j - n_i) \right] + C_{ji} \left[n_j - n_i \exp\left(\frac{-h\nu_{ji}}{kT}\right) \right] \right\}$$
(1)

where $N_i = g_i n_i$, g_i is the statistical weight of level *i*, N_i represents the number density in level *i*, *W* is the dilution factor and \aleph_{ij} the photon occupancy number at transition frequency ν_{ij} .

To obtain the matrix equation, rewrite the parameters of Equation 1, grouping all populating and depopulating levels and substitute the radiative and collisional components

$$R_{ij} = \begin{cases} A_{ji}\beta_{ji}(1+X_{ji}) & i < j \\ A_{ij}\beta_{ij}\left(\frac{g_i}{g_j}\right)X_{ij} & j > i \end{cases}$$
$$R_{ii} = -\sum_{i \neq j} R_{ji}$$
$$\widetilde{C}_{ij} = C_{ji}$$
$$\widetilde{C}_{ii} = -\sum_{i \neq j} C_{ij}$$

Thus obtaining the form:

$$\mathbf{Q}\mathbf{x} = \mathbf{b}$$

where $\mathbf{b} = [0, \dots, 0, 1]^T$, $x_i = \frac{n_i}{n_{mol}}$ the normalised fractional population density with n_{mol} the total population density of the molecule and $\mathbf{Q} = \mathbf{R} + \tilde{\mathbf{C}}$.

The equation can be solved using iterative methods, such as LU-factorization or the singular value decomposition (SVD). The required initial solution is obtained by calculating the level population at local thermal equilibrium (LTE).

3. Masers Python Package

3.1. Implementation

The masers package solves for statistical equilibrium given some parameters to describe the physical environment. The most basic parameters that must be provided are the density of the cloud, the abundance of the maser molecule, various temperatures, and the column density. To calculate the optical depth, the anticipated size and geometry of the cloud must also be provided, as well as any and all diffuse background radiation components that may interact with the line emission.

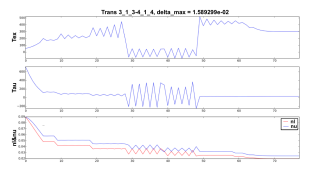
Since the solution of the radiative transfer equation is dependent on the molecular level populations, molecular rate and collision coefficients must also be provided. These coefficients are obtained from the molecular data files in the Leiden LAMBDA database [2], http://home.strw.leidenuniv.nl/~moldata.

During development, we used the RADEX non-LTE excitation and radiative transfer code [3], available from the Leiden website, for comparison and verification; note that both methods solve the rate equations iteratively. Iterative methods are sensitive to oscillation and multiple valid solutions since the solution of the previous iteration is used as the initial estimate for the next iteration and the process is repeated until the input estimate and calculated solution have reached some convergence criteria.

Numerical stability is evaluated by calculating the optical depth and excitation temperature for formaldehyde molecular data [4], gas kinetic temperature $T_k = 300 \text{ K}$, H_2CO fractional abundance $X_{H_2CO} = 5 \times 10^{-7}$, specific column density $\frac{N_{col}}{\Delta v} = 10^6, 10^9, 10^{12} \text{ cm}^{-3}$ s respectively and $T_{BB} = 30 \text{ K}$ as blackbody radiator, private communication Prof van der Walt. The equilibrium solution for total density ranging from $10^3 - 10^{12} \text{ cm}^{-3}$ is calculated using a convergence limit of $|x_n - x_{n-1}|/x_{n-1} < 10^{-3}$, with x_n and x_{n-1} is the level populations calculated during the current and previous evaluation.

Possible convergence criteria includes using population densities or excitation temperature calculations, via the relation $\frac{n_u}{n_l} = \frac{g_u}{g_l} \exp(\frac{-h\nu}{kT_{ex}})$. The effect of choosing either convergence criteria for the parameters above was inspected for specific column density $\frac{N_{col}}{\Delta \nu} = 10^{12} \text{ cm}^{-3}\text{s}$. The masers implementation uses the population densities calculated at each iteration. RADEX evaluates the average difference between the excitation temperature calculated from the previous solution compared to the calculation of the current solution (from source code). Inspection of excitation temperature calculation as convergence criteria showed sensitivity to oscillating solutions, Figure 1, caused by catastrophic cancellation due to the difference of small numbers in the denominator of the excitation temperature calculation, $T_{ex} \propto [\ln(x_lg_u) - \ln(x_ug_l)]^{-1}$. RADEX minimises this by only selecting certain levels to evaluate convergence (from source code).

Excitation temperature values computed at all iterations over total density range are shown in the top graph of Figure 1. The centre and bottom graphs shows the effect carried forward and amplified in the optical depth and level population calculations. The red line in the bottom graph shows the population densities for the lower level of the transition as they were calculated over iterations, while the blue line shows the population densities for the upper transition level. Slight step functions show the total density steps, but the population densities clearly show oscillating behaviour around $n_{H_2} = 10^6 - 10^8 \text{ cm}^{-3}$.



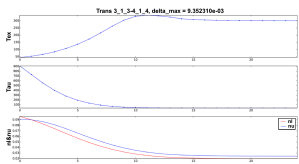


Figure 1. Oscillating behaviour seen in convergence evaluation using excitation temperature evaluation. The convergence results are seen in the 4_{14} – 3_{13} transition showing 80 iterations over total density range $n_{H_2} = 10^6 - 10^8$ cm⁻³ and specific column density $\frac{N_{col}}{\Delta v} = 10^{12}$ cm⁻³s.

Figure 2. A ratio of 9.5:0.5 means that only 5% is given to the new calculation, pulling the iterative solution close to the previous behaviour. This "memory" stabilizes the iterative calculation by ensuring it is not independent, but is influenced by the previous results.

Oscillating behaviour is only pronounced at lower transition levels and the solutions were found to become more stable if the next iteration is given some "memory" of the previous solutions. This was done using a running average calculation, $x_n = c_1 \times x_n + c_2 \times x_{n-1}$ where $c_1 \leq 1$ and $c_2 \leq 1$ are some weighting coefficients with $c_1 + c_2 = 1$. The larger the coefficient for the previous solution x_{n-1} the "longer" the memory of the solution is. For masers this "memory" was found to be fairly large with $x_n = 0.05 \times x_n + 0.95 \times x_{n-1}$ providing solutions under all simulated conditions. It should be noted that a "long memory", $c_2 \rightarrow 1$, requires a more stringent convergence limit. For masers this is found to be $|x_n - x_{n-1}|/x_{n-1} < 10^{-7}$ to obtain solutions shown in Figure 2.

Lastly, the calculation of optical depth between the 2 implementations is also found to be significant. The masers solver applies the large velocity gradient, LVG, definition for the escape

probability calculation [1]. When results are compared to solutions obtained from RADEX care must be taken to ensure no large negative optical depth values. For thermal radiative transfer calculations the optical depth is always expected to be positive, thus RADEX substitutes the turbulent medium escape probability calculation for $|\tau| \ge 7$ (from source code). This escape probability calculation requires the computation of $\sqrt{\tau}$, which fails for maser regions when inversion causes the optical depth to be negative [3].

Stabilising masers against the highlighted numerical sensitivities provides a robust solver to use during the inspection of maser pumping parameter space. Figure 3 shows the improvement at selected levels known to cause numerical failures in RADEX. The top row shows results from the masers solver, while the bottom row shows results from the RADEX code. The four columns show Tex, log10(Tex), Tau, log10(Tau) for selected transition levels $1_{10}-1_{11}$ and $12_{111}-11_{110}$.

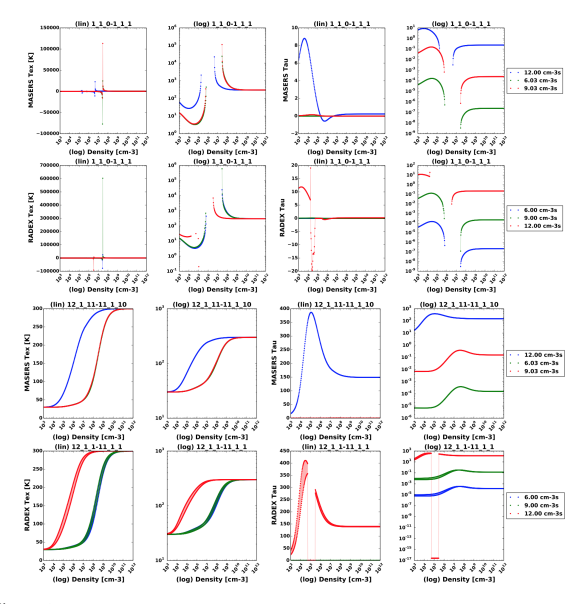


Figure 3. The first and third rows show results from the masers solver with colour scales showing specific molecular column density $\frac{N_{col}}{\Delta v} = 10^6$ (green), 10^9 (red), 10^{12} (blue) for all H_2CO transitions. The second and last rows show results obtained from the **RADEX** solver with specific molecular column density $\frac{N_{col}}{\Delta v} = 10^6$ (blue), 10^9 (green), 10^{12} (red) for all H_2CO transitions.

3.2. Verification

Following methodology suggested by Prof van der Walt, in a private communication, the basic functionality of the masers software could be verified. This methodology describes the inspection of the pumping and inversion of H_2CO masers in star-forming region G37.55+0.20, specifically looking at the optical depth calculation for the 4.8 GHz maser associated with the $1_{10}-1_{11}$ transition as a function of the specific column density.

Assuming fractional molecular density $X_{H_2CO} = 5 \times 10^{-7}$ and electron temperature $T_e = 10^4$ K, Figure 4 shows example calculations of the following three physical environments:

- $n_{H_2} = 4.4 \times 10^5 \,\mathrm{cm}^{-3}$, $T_d = 100 \,\mathrm{K}$, $T_k = 300 \,\mathrm{K}$, $w_d = 0.1$ and no H_{II} region,
- $n_{H_2} = 10^4 \,\mathrm{cm}^{-3}$, $T_k = 20 \,\mathrm{K}$, $w_{H_{II}} = 0.1$, $EM = 6 \times 10^9 \,\mathrm{cm}^{-6} \,\mathrm{pc}$ and no dust,
- $n_{H_2} = 10^4 \,\mathrm{cm}^{-3}, T_d = 100 \,\mathrm{K}, w_d = 0.1, T_k = 300 \,\mathrm{K}, w_{H_{II}} = 0.1, EM = 6 \times 10^9 \,\mathrm{cm}^{-6} \,\mathrm{pc}.$

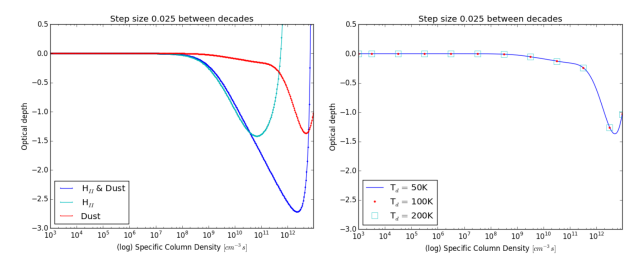


Figure 4. Three examples showing variation of optical depth calculation of the $1_{10}-1_{11}$ transition. Physical parameters were kept constant over the range of specific column density values, showing the maximum optical depth identified.

Figure 5. These results indicates that the inversion is unrelated to the dust temperature, which suggests that radiative pumping by the dust infrared radiation field is not significant for the maser.

Similarly, Figure 5 shows results for optical depth calculation with dust temperatures using $T_d = 50, 100, 200 \text{ K}$, given $n_{H_2} = 4.4 \times 10^5 \text{ cm}^{-3}$, $T_k = 300 \text{ K}$, $w_d = 0.1$ and no H_{II} region.

Numerical results obtained were compared to results obtained by Prof van der Walt using independent numerical code solving the rate equations using Heun's method.

3.3. Pumping

Applying the masers software to investigate the pumping of the H_2CO maser in G37.55+0.20 as described in [5] showed good comparison between results. Both methods showed that collisional excitation with H_2 as well as radiative excitation by the free-free radio continuum radiation from a nearby ultra- or hyper-compact H_{II} region can invert the 1_{10} - 1_{11} (4.8 GHz) transition.

Collisional excitation Figure 6 shows the optical depth of $1_{10}-1_{11}$ transition as a function of H_2CO specific column density over range 10^8-10^{14} cm⁻³ s and $T_k = 140, 180, 220, 260, 300$ K, $n_{H_2} = 4 \times 10^5, 2.6 \times 10^5, 1.6 \times 10^5, 1.3 \times 10^5, 7.5 \times 10^4$ cm⁻³. Visual comparison shows results are duplicate of those shown in Fig 1 presented in [5].

- **Radiative excitation via the dust continuum emission** The dependence of optical depth on the specific column density is shown in Figure 7, for $T_k = 180 \text{ K}$, $n_{H_2} = 1.3 \times 10^5 \text{ cm}^{-3}$, $w_d = 1.0$ and $T_d = 0 \text{ K}$ (no dust), 100, 180 K. Comparison shows results are duplicate of those shown in Fig 3 presented in [5].
- Radiative excitation through the free-free continuum emission from an ultra- or hypercompact H_{II} region is shown in Figure 8 using emission measure of the H_{II} region $EM = 10^{10} \text{ pc cm}^{-6}$, as well as $T_e = 10^4 \text{ K}$, $w_{H_{II}} = 1.0$, $T_k = 140, 180, 220, 260, 300 \text{ K}$, and $n_{H_2} = 4 \times 10^5, 2.6 \times 10^5, 1.6 \times 10^5, 1.3 \times 10^5, 7.5 \times 10^4 \text{ cm}^{-3}$. Comparison shows results are duplicate of those shown in Fig 6 presented in [5].

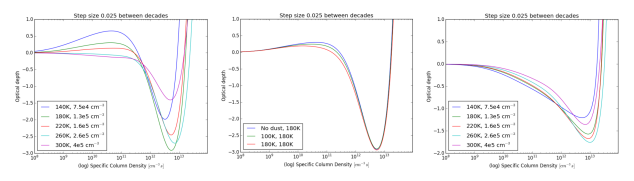


Figure 6. Effect of collisions on the level populations in the absence of radiation fields.

Figure 7. Optical depth as a function of specific column density.

Figure 8. Free-free emission has a significant effect on the inversion.

4. Summary

The paper highlights various reasons why standard matrix inversion methods used in the calculation of thermal line transfer models need to be more robust to ensure good results in the stimulated maser environments used to model maser pumping schemes.

Verification could be achieved through personal communication with Prof van der Walt who has independent software for solving the rate equations using Heun's method.

Validation of the implementation is followed by comparison between the results obtained with the masers package to results published in [5]. Visual comparison of graphs showing results of example parameters inspecting the pumping of the H_2CO maser in G37.55+0.20 shows good agreement.

References

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