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Protoplanetary discs: sensitivity of the chemical composition to various model parameters

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ABSTRACT

Protoplanetary discs are challenging objects for astrochemical models due to strong density and temperature gradients and due to the UV photons 2D propagation. In this paper, we have studied the importance of several model parameters on the predicted column densities of observed species. We considered: (1) two-phase (gas and homogeneous grains) or three-phase (gas, surface, and bulk of grains) models, (2) several initial compositions, (3) grain growth and dust settling, and (4) several cosmic-ray ionization rates. Our main result is that dust settling is the most crucial parameter. Including this effect renders the computed column densities sensitive to all the other model parameters, except cosmic-ray ionization rate. In fact, we found almost no effect of this parameter for radii larger than 10 au (the minimum radius studied here) except for N$_2$H$^+$. We also compared all our models with all the column densities observed in the protoplanetary disc around DM Tau and were not able to reproduce all the observations despite the studied parameters. N$_2$H$^+$ seems to be the most sensitive species. Its observation in protoplanetary discs at large radius could indicate enough N$_2$ in the gas phase (inhibited by the three-phase model, but boosted by the settling) and a low electron abundance (favoured by low C and S elemental abundances).

Key words: astrochemistry – Planetary systems – protoplanetary discs – ISM: abundances – ISM: individual objects: DM Tau – ISM: molecules.

1 INTRODUCTION

Modelling the chemistry in protoplanetary discs is a complex task because of the strong gradients in the physical properties that exist (Dutrey et al. 2014). Considering the radial and vertical geometries of these objects, the temperature can go from a few 100 K to a few K (excluding of course the most upper atmosphere of the discs where the temperatures are of a few thousand kelvins) while the density varies between $10^4$ and $10^{12}$ cm$^{-3}$. Both the temperature and density radially decrease from the central star. The temperature also decreases towards the mid-plane of the disc while the density has an inverse behaviour. The chemistry is also driven by the irradiation of UV photons coming from the central star. The consequence of this 2D geometry is a vertical chemical structure. The mid-plane is depleted in gas-phase molecules while the outer most layers of the disc are populated by atoms as photodissociation dominates. In between, there is a layer of material where we find most of the molecules (Dutrey et al. 2014). As a consequence, the chemical composition of the gas and ices presents strong variations in the radial and vertical directions. Because of the competition between gas-grain interactions, bimolecular gas-phase chemistry and photodissociation in the molecular layer, the chemical model predictions are very sensitive to the local conditions. There exists a variety of astrochemical models with different types of approximations for the geometry (2D or 1D + 1D), the chemistry (gas-phase or gas-grain, steady-state or time dependent), the dust properties (single grain size or size distribution), the dynamics (included or not), etc. Henning & Semenov (2013) have done a nice compilation of the different existing models. Because of these differences and the high sensitivity of the disc chemistry, it is very difficult to compare one model result to the other. The philosophy of the work presented here is to explore the importance of some of the chemical model approximations and/or parameters on the predicted column densities. We focused on four different aspects:

1) The two- or three-phase approximation for surface chemistry: In three-phase astrochemical models (gas, surface, and bulk of the ice mantles on top of the grains), we make a difference between the first few (usually two or four) most external layers of ices...
on the grains (considered as the surface) and the rest of the ice below (considered as the bulk). Most of the chemistry occurs at the surface layers while the bulk is protected against evaporation. In the two-phase models (gas and surface of grains), we do not make any difference between the surface and the bulk. The entire ice is equally chemically active and can desorb. The first question we address is then the impact of these two approximations on the chemical model predictions.

2) The initial chemical composition: Protoplanetary discs are the result of a long evolutionary sequence starting from the diffuse medium and ending by the formation of stars and planets. During this sequence, the chemistry is never at steady state. It is very common to simplify the sequence and simply start from an initial chemical composition and use static disc physical properties. The second question we address then is the importance of the initial chemical composition on the disc computed column densities.

3) The third aspect we explored is the grain growth and dust settling.

4) Finally, we tested several values of the cosmic-ray ionization rate based on the idea that this rate may be smaller in protoplanetary discs than the typical value of $10^{-17}$ s$^{-1}$ as claimed by Cleev et al. (2015).

This paper is organized as follows. The disc physical and chemical models are presented in Section 2 while the tested parameters and approximations are described in Section 3. Results are presented in Section 4, compared with observations in the DM Tau protoplanetary disc in Section 5, and compared with other previous models in Section 6. Our conclusions are summarized in the last section.

2 MODEL DESCRIPTION

The physical model, we assume, for protoplanetary discs is the same as in Wakelam et al. (2016). Briefly, the radial and vertical density profiles are computed assuming that the gas is in hydrostatic equilibrium. The temperature gradients are computed following the prescriptions of Dartois, Dutrey & Guilloteau (2003) and Williams & Best (2014). For our nominal model (without grain growth), the local visual extinction ($A_v$) is derived assuming a conversion factor of $(A_v/N_H) = 6.25 	imes 10^{-22}$ (Wagenblast & Hartquist 1989) with $N_H$ the vertical H column density. The UV flux at a radius $r$ from the star is assumed to be absorbed at the surface of the disc and half of the flux diffuses in the vertical direction. The stellar mass and UV fluxes are similar to those of the DM Tau star and all parameters to compute the 2D physical structure are listed in Table 1 of Wakelam et al. (2016). DM Tau is a T Tauri star of mass 0.53 $M_\odot$ (Simon, Dutrey & Guilloteau 2000), its Keplerian disc has a total mass of 0.03 $M_\odot$, with a surface density of 0.8 g cm$^{-2}$ (Guilloteau et al. 2011) and an outer radius of 800 au (Guilloteau & Hersant 1998). The computed density and temperature gradients are shown in Figs 1 and 2, respectively. Throughout the paper, $r$ refers to the radius from the star and $z$ is the altitude above mid-plane, both are in unit of au. $h$ is the scale height $h = \sqrt{\frac{k_B T_{\text{mid}}}{\mu m_a G M}}$, with $k_B$ the Boltzmann constant, $T_{\text{mid}}$ the mid-plane temperature, $\mu$ the mean molecular weight of the gas, $m_a$ the atomic mass unit, $G$ the gravitational constant, and $M_*$ the mass of the star. We assume a duration of $4 \times 10^6$ yr for the chemical evolution. Although the chemistry in the disc has not reached steady state (even in the mid-plane where the gas-grain exchanges dominates the chemistry), it only slowly evolves in the disc for ages beyond $10^6$ yr for all the conditions (including the cosmic-ray ionization rates) used in this study. For this reason, the results presented here would still stand if the disc were two times younger or older.

The 2D chemical composition in the disc is computed with the gas-grain code Nautilus fully described in Ruaud, Wakelam & Hersant (2016) and using the physical structure described above as input parameters. The chemical composition of the gas and the volatile ice on top of interstellar dust grains is computed as a function of time by solving a set of differential equations. The gas-phase chemical network is derived from kida.uva.2014 (Wakelam et al. 2015a), which contains all relevant and common processes for the gas-phase chemistry (unimolecular reactions, such as photodissociation and ionization by direct UV photons and cosmic-ray induced UV photons, bimolecular reactions such as ion–neutral and neutral–neutral reactions, and electronic recombination). This network has been updated from more recent publications from Wakelam et al. (2015b), Loison et al. (2016), Hickson, Wakelam & Loison (2016), Vidal et al. (2017), and Loison et al. (2017). CO, H$_2$, and N$_2$ self-shielding are computed using tabulated values from Lee et al. (1996), Visser, van Dishoeck & Black (2009), and Li et al. (2013), respectively. In the three-phase version of the code, the surface layer is defined by the two most external layers of molecules while the rest below defines the bulk of the mantle. Species from the
gas-phase can be physisorbed on the grain surfaces upon collision while physisorbed species can desorb through thermal and non-thermal processes. For non-thermal desorption, the model includes the desorption of species due to cosmic-ray induced heating (Hasegawa & Herbst 1993), (direct and indirect) UV photons, and exothermic surface reactions (i.e., chemical desorption). The surface network is originally the one included in the gas-grain model developed in Prof. Herbst’s team with modifications described in Ruad et al. (2015), Wakelam et al. (2015b), Loison et al. (2016), Hickson et al. (2016), Vidal et al. (2017), and Loison et al. (2017). The binding energies have been updated from Wakelam et al. (2017). All the data on the surface reactions and binding energies have been included in the online KIDA database (Wakelam et al. 2012, ApJ 199, article id.21). For photo-desorption, considering the complexity of the mechanism, we followed the advice of Bertin et al. (2013) and used a constant photo-desorption yield of 10^{-4} molecule/photon for all molecules (see also Ruad et al. 2016, for discussions). For the chemical desorption, we are using the model described by Garrod, Wakelam & Herbst (2007) with a parameter of 0.1 (which implies that nearly 1 per cent of the singly produced species desorbs). Both the surface and bulk are chemically active but the binding and diffusion barrier are set differently so that the reactivity in the bulk is much slower: the diffusion-to-binding energy ratio is 0.4 on the surface and 0.8 in the bulk. In addition, in the bulk, the binding energies, smaller than the one of water, are set to the one of water (except for H, H₂, C, N, and O) in agreement with the findings of Ghesquière et al. (2015). Diffusion of species occurs through thermal hopping except for oxygen, which has been shown to diffuse through tunneling effects Minissale et al. (2013). Without this tunneling diffusion and with the new estimate of the high binding energy of O on water surfaces, complex organic molecules would not be produced efficiently on the surfaces. The reaction–diffusion competition as proposed by Chang, Cuppen & Herbst (2007) and Garrod & Pauly (2011) is included in the model. The bulk and the surface phases interact in the sense that the species from the surface are incorporated little by little to the bulk while new species land on the surfaces. Similarly, the bulk species become the surface ones while surface species desorb in the gas phase. All details on how the processes are included in the code and the model parameters are given in Ruad et al. (2016).

3 TESTS

We have tested the effect of several assumptions commonly used for disc chemical modeling. The following models have been run:

(1) Two- or three-phase model: The chemical model is either computed with the two-phase or three-phase approximation. Although the three-phase models are physically more realistic, they are numerically difficult to perform. As a consequence, two-phase models are still commonly used for protoplanetary disc chemical modeling but also for many other types of applications.

(2) Initial conditions: The disc is usually assumed to form from a cold dense core. The initial chemical composition is usually assumed to be either atomic (for instance Willacy et al. 1998) or molecular (for instance Hersant et al. 2009; Cleeves et al. 2016). Here, we will consider three models. In the first one, all the gas is assumed to be atomic except for H₂ (with atomic abundances listed in table 1 of Vidal et al. 2017, except for sulphur assumed to have the undepleted abundance of 1.5 × 10^{-5}). This is the typical initial condition for a cold core forming from diffuse medium (where hydrogen is usually already converted to H₂). In the two other models, we first compute the composition of a cold shielded core at 10 K, with a H density of 2 × 10^{10} cm^{-3}, and a cosmic-ray ionization rate of 10^{-17} s^{-1} (the grains remain at 0.1μm for this stage). The cold core chemical compositions at 10^3 yr and 10^6 yr are then used as initial compositions for each of the two models as these times represent the chemical ages based on the comparison between chemical models and observations (see for instance Loison et al. 2017; Vidal et al. 2017). This simple assumption of course does not take into account the evolution of the chemistry between the cold core phase and the protoplanetary disc itself (Drozdovskaya et al. 2016).

(3) Dust settling and grain growth: The model is first run assuming a homogeneous distribution of the grains: grains are the same at all altitudes and all radii (grain radius r_d of 0.1μm and dust-to-gas mass ratio of 10^{-2}). Models are also run to simulate the dust settling without and with grain growth. Dust settling depends on the product of the angular velocity and the dust stopping timescale, which scales as the particle size. When this product is larger than 1, particles decouple from the gas. Large grains are assumed to settle by being distributed with a smaller scale height than the gas. The settling ratio, ratio of the dust scale height to the gas scale height, is approximated in the same way as by Boehler et al. (2013). These approximations are based on the numerical study of Fromang & Nelson (2009). The long wavelength dependence of the mm continuum emission of discs suggests that grains have grown to sizes up to several centimetres (e.g. Ricci et al. 2010, and references therein). To evaluate a case of grain growth, we used an effective grain size of 30 μm, which presents the same cross-section as a size distribution with radii between 0.01 μm and 10 cm and an exponent of 3.5 for the number density as a function of size, a value found in the ISM (Mathis, Rumpl & Nordsieck 1977). In our model, dust settling is simulated by changing the dust-to-gas mass ratio (dtogm), in the upper layer of the disc (above a certain altitude z that we will call z_s), to 10^{-2} and putting the extra mass below z_s (the average dtogm is still 10^{-2}). The grain growth is simulated by replacing grains below z_s with larger 30μm grains. Both the dust settling and grain growth changes the number density of grains in the vertical direction. The visual extinction is also changed as the A_v/N_H conversion factor is scaled as follows:

\[(A_v/N_H) = (A_v/N_H)_0 \frac{dtogm}{10^{-2}} \frac{10^{-5}}{r_d(cm)}\]

Since large grains are significantly larger than the optical wavelengths, we assume that the (A_v/N_H) factor scales with the dust size (equation above). Note that in all cases, we do not include a grain size distribution and for each grid of the model we have only one single size (that can change though). Fig. 3 shows the visual extinction obtained in the three types of models described here for z_i = 1h. The computed visual extinction in the three models are the same above z_i. Below this point, in the case of grain growth, A_v remains constant below this layer as the big grains are not contributing to the visual extinction anymore while in the case without grain growth, the visual extinction is increased because of a large amount of small grains.

(4) Cosmic-ray ionization rate: Cleeves, Adams & Bergin (2013), Cleeves, Bergin & Adams (2014), and Cleeves et al. (2015) propose observational and theoretical evidence that the cosmic-ray ionization rate in protoplanetary discs may be smaller than the typical interstellar value due to exclusion by the stellar winds. To test

1http://kida.obs.u-bordeaux.fr/
Figure 3. Computed visual extinction in the case of homogeneous 0.1 μm size grains (upper panel), settling without grain growth on the mid-plane (middle panel), and settling with grain growth on the mid-plane (lower panel). One scale height is approximately where the vertical profiles present a bowing in the upper panel while it is when they present a discontinuity in the two other panels. Xgrain is the abundance of grains with respect to the total proton density.

4 RESULTS

Considering the large number of models we have run and the large number of molecules we can look at, we focus this paper to the molecules observed in the protoplanetary disc around DM Tau. Table 2 lists the observed molecules with their observed column densities at 300 au.

4.1 With homogeneous 0.1 μm grains (models A and B)

Fig. 4 shows the predicted vertical column densities for these species as a function of radius to the central star for models A(1–3) and B(1–3). Here, the molecules are assumed to be optically thin so that the theoretical vertical column densities are multiplied by a factor of 2 to represent the two sides of the disc (on both sides of the mid-plane). For most molecules, all model results with the two-phase mode overlap. The same is found for the model results with the three-phase mode. This result shows that the species column densities, in the ranges of radius considered here, do not depend on the initial conditions in that model. This result is not true for the main ice constituents that present significant differences among two-phase and three-phase models for the inner 100 au. The two- and three-phase models give significant differences for most molecules except for CN, CS, and H2S. The difference in the CO gas-phase column density comes from the fact that in the two-phase model, CO is produced in the molecular layer (between z/h = 2 and 3) by the photodissociation of gas-phase CO2, which is much more abundant with the two-phase model. Fig. 5 shows the computed radial column density of CO2. Between 100 and 200 au in the molecular layer of the disc, we found that the origin of gas-phase CO2 is either chemical desorption during its formation on the surfaces through the reaction s—CO + s—O or photodesorption of surface CO2 (either formed by s—CO + s—O or by s—HCO + s—O; s— means the species physisorbed at the surface of the grains). Both the two desorption mechanisms are more efficient in the two-phase model, where all surface molecules are allowed to desorb equally, explaining the much higher abundance of CO2 in this model. Going outwards from the star, the gas and dust temperature decreases as well as the UV flux. As a consequence, the CO2 production on the grains and further desorption decreases. At 300 au, CO2 is then mostly produced by the gas-phase reaction O + HCO → CO2 + H in both two- and three-phase models. The fact that the CO2 abundance is much smaller in the three-phase model, as compared to the two-phase, at 200 au also explains the smaller SO abundance there. Indeed, SO is mostly formed by the reaction O + HS → H + SO and atomic oxygen is obtained by the photodissociation of CO2. The dramatically lower N2H+ column density with the three-phase model is due to the fact that N2 (precursor of N2H+) cannot evaporate from the grains while it is photodesorbed in the disc mid-plane is the decay of short-lived radionuclides. The upper value is the one commonly used in astrochemical models.

The models we have run are summarized in Table 1. We did not make a full grid of models because of computational capabilities but selected some of the models as examples. For the settled and grain growth models, for example, we have used only the age of 106 yr for the molecular initial conditions because these conditions are the most evolved ones as compared to the atomic one, and so may produce the largest differences. For the cosmic-ray ionization rate, we also limited our tests to the two models presenting the largest differences.

In the effect of decreasing the cosmic-ray ionization rate, we run the model with two values of ζ: 10−17 and 10−19 s−1 (same value in the entire disc). The lower value is the upper limit suggested by Cleeves et al. (2015), which is based on the exclusion of Galactic cosmic rays by TTauri wind, and in that case the dominant source of ionization in the disc mid-plane is the decay of short-lived radionuclides. The upper value is the one commonly used in astrochemical models.
two-phase model. At 200 au, the larger column density of HCN with the two-phase model is due to the fact that HCN is formed efficiently on the surfaces and is desorbed by UV photons. The larger CCH column density in the three-phase model, between 100 and 300 au, is due to a larger abundance peak of this molecule at z/h = 2 (while the molecular abundance peaks only at z/h = 3 in the two-phase model). This difference is due to a larger abundance of s–CH$_4$ (because less carbon is locked in CO and CO$_2$) in the three-phase model at this altitude and these radii. The larger s–CH$_4$ abundance produces the following paths: s–CH$_4$ + s–CCH → s–C$_2$H$_2$, s–C$_2$H$_2$ + s–H → s–C$_1$H$_3$, s–C$_1$H$_3$ + s–H → C$_3$H$_4$, C$_3$H$_4$ + hv → C$_2$H$_2$ + H$_2$, and C$_2$H$_2$ + hv → CCH. This difference is at the origin of the higher (although moderate) CCS column density in the three-phase model as CCS forms from CCH + S.

### 4.2 With grain growth and dust settling (models C–E)

We now assume that grains have settled (dust-to-gas mass ratio is 10$^{-3}$ above z/h = 1) and that the grains in the mid-plane have grown (to 30μm). The species column densities computed with these models are shown in Fig. 6. The main first result is that all species column densities depend on these assumptions. The effect depends on the species and radius but the differences can be orders of magnitude. Considering CO at 100 au, the column densities spread over 1 order of magnitude (between 10$^{17}$ and 10$^{18}$ cm$^{-2}$). Beyond 200 au, the column densities are rather similar for all models and higher by 2 orders of magnitude than in the case of homogeneously distributed 0.1μm grains. To understand what is happening, we also run a model in which we assume dust settling but not grain growth, and two initial conditions (molecular and atomic) (models E1 and E2). In these cases, the CO column density is smaller but highly dependent on the initial conditions. When starting with molecular initial conditions (model E2), we obtain similar column densities as in the case without settling (model B3) while starting with atoms (model E1), we end up with 10 times higher values (but lower than in the case with grain growth – model D1). Fig. 7 shows the vertical CO abundance and cumulative column density computed by the two models with settling and without grain growth (models E1 an E2). Superimposed is the result of the model with settling and grain growth (and molecular initial conditions – model D2). In the models starting with molecular initial conditions, a significant fraction of the carbon is already locked as CO on grains at the beginning of the simulations. This explains the fact that the peak abundance of gas-phase CO around z/h = 2 is much larger while starting from atoms. The fact that the CO column density is much smaller in the

### Table 1. Summary of the different models.

| Model type | Initial conditions | Settling | Grain growth | $\zeta$ (s$^{-1}$) |
|------------|--------------------|----------|--------------|-------------------|
| A1         | Two-phase          | Atomic   | No           | No                | 10$^{-17}$       |
| A2         | Two-phase          | Atomic   | No           | No                | 10$^{-17}$       |
| A3         | Two-phase          | Atomic   | No           | No                | 10$^{-17}$       |
| B1         | Three-phase        | Atomic   | No           | No                | 10$^{-17}$       |
| B2         | Three-phase        | Atomic   | No           | No                | 10$^{-17}$       |
| B3         | Three-phase        | Atomic   | No           | No                | 10$^{-17}$       |
| C1         | Two-phase          | Atomic   | Yes          | Yes               | 10$^{-17}$       |
| C2         | Two-phase          | Atomic   | Yes          | Yes               | 10$^{-17}$       |
| D1         | Three-phase        | Atomic   | Yes          | Yes               | 10$^{-17}$       |
| D2         | Three-phase        | Atomic   | Yes          | Yes               | 10$^{-17}$       |
| E1         | Three-phase        | Atomic   | Yes          | No                | 10$^{-17}$       |
| E2         | Three-phase        | Atomic   | Yes          | No                | 10$^{-17}$       |
| F1         | Two-phase          | Atomic   | No           | No                | 10$^{-19}$       |
| F2         | Three-phase        | Atomic   | Yes          | Yes               | 10$^{-19}$       |

### Table 2. Observed vertical column densities (in cm$^{-2}$) in DM Tau at 300 au. E2.1–E2.6 refers to the results of several additional models described in Section 5.1.

| Molecule | Observed value | E2 | E2.1 | E2.2 | E2.3 | E2.4 | E2.5 | E2.6 |
|----------|----------------|----|------|------|------|------|------|------|
| CO       | (1.1 ± 0.3) × 10$^{17}$ | 6.7 × 10$^{10}$ | 3.1 × 10$^{11}$ | 9.7 × 10$^{10}$ | 4.9 × 10$^{12}$ | 5.0 × 10$^{12}$ | 3.3 × 10$^{12}$ | 3.1 × 10$^{10}$ |
| HCO$^+$  | (1.1 ± 0.3) × 10$^{13}$ | 6.7 × 10$^{10}$ | 3.1 × 10$^{11}$ | 9.7 × 10$^{10}$ | 4.9 × 10$^{12}$ | 5.0 × 10$^{12}$ | 3.3 × 10$^{12}$ | 3.1 × 10$^{10}$ |
| N$_2$H$^+$ | (1.1 ± 0.3) × 10$^{11}$ | 6.3 × 10$^{8}$ | 1.2 × 10$^9$ | 6.9 × 10$^{8}$ | 3.5 × 10$^7$ | 4.4 × 10$^7$ | 3.9 × 10$^5$ | 6.2 × 10$^7$ |
| CCH      | (2.8 ± 0.2) × 10$^{13}$ | 4.1 × 10$^{13}$ | 9.9 × 10$^{13}$ | 3.7 × 10$^{13}$ | 4.7 × 10$^{14}$ | 4.7 × 10$^{14}$ | 9.5 × 10$^{14}$ | 3.7 × 10$^{13}$ |
| HCN      | (2.8 ± 0.2) × 10$^{13}$ | 4.1 × 10$^{13}$ | 9.9 × 10$^{13}$ | 3.7 × 10$^{13}$ | 4.7 × 10$^{14}$ | 4.7 × 10$^{14}$ | 9.5 × 10$^{14}$ | 3.7 × 10$^{13}$ |
| CN       | (3.5 ± 0.9) × 10$^{12}$ | 3.1 × 10$^{13}$ | 2.6 × 10$^{14}$ | 1.5 × 10$^{14}$ | 3.9 × 10$^{13}$ | 3.9 × 10$^{13}$ | 1.7 × 10$^{12}$ | 6.5 × 10$^{13}$ |
| CS       | (3.5 ± 0.9) × 10$^{12}$ | 3.1 × 10$^{13}$ | 2.6 × 10$^{14}$ | 1.5 × 10$^{14}$ | 3.9 × 10$^{13}$ | 3.9 × 10$^{13}$ | 1.7 × 10$^{12}$ | 6.5 × 10$^{13}$ |
| SO       | ≤7.5 × 10$^{11}$ | 5.0 × 10$^{12}$ | 5.0 × 10$^{13}$ | 3.1 × 10$^{13}$ | 3.3 × 10$^{12}$ | 3.3 × 10$^{12}$ | 3.5 × 10$^{12}$ | 7.3 × 10$^{13}$ |
| H$_2$S    | ≤1.4 × 10$^{11}$ | 5.3 × 10$^{14}$ | 2.6 × 10$^{15}$ | 7.6 × 10$^{13}$ | 5.1 × 10$^{13}$ | 5.1 × 10$^{13}$ | 4.7 × 10$^{12}$ | 3.7 × 10$^{12}$ |
| CCS      | ≤1.1 × 10$^{12}$ | 2.7 × 10$^{13}$ | 3.5 × 10$^{13}$ | 3.7 × 10$^{13}$ | 3.2 × 10$^{13}$ | 3.2 × 10$^{13}$ | 3.7 × 10$^{12}$ | 2.0 × 10$^{13}$ |
| HC$_3$N  | ≤7.5 × 10$^{11}$ | 1.0 × 10$^{11}$ | 2.5 × 10$^{12}$ | 4.5 × 10$^{11}$ | 1.9 × 10$^{12}$ | 2.0 × 10$^{12}$ | 9.1 × 10$^{10}$ | 6.9 × 10$^{10}$ |

**Note:** References for the observed column densities are the following: Dutrey et al. (2011) for CS, SO and H$_2$S, Chapillon et al. (2012a) for HC$_3$N and CCS, Chapillon et al. (2012a) for CN and HCN, Dutrey et al. (2007) for N$_2$H$^+$, Henning et al. (2010) for CCH, Piétu, Dutrey & Guilloteau (2007) for HCO$^+$ (value indicated with the $b$ exponent) and CO, and Teague et al. (2015) for HCO$^+$ (value indicated with the $b$ exponent). All the observed surface densities have been derived using a proper radiative transfer disc model (DISKFIT; see Piétu et al. 2007; Dutrey et al. 2011, for details).
Figure 4. Column densities of a selection of gas-phase species as a function of radius to the central star for two-phase models A(1–3) and three-phase models B(1–3). The points at 300 au represent the observed column densities.

Case without grain growth (similar in fact to the case with settling presented in Section 4.1) is caused by the increase of the number of grains in the mid-plane changing the depletion of CO. Indeed, the grain growth increases the cross-section of individual grains hence increasing the probability of collision between gas-phase chemical species and an individual grain. Keeping the total mass constant, grain growth decreases the total number density of grains and so the probability of collision with a population of grains. The abundance (with respect to H) of 30μm grains is only $6.8 \times 10^{-20}$ whereas it is $1.8 \times 10^{-12}$ in the case of 0.1μm grains.

The N$_2$H$^+$ predicted column densities peak inside 80 au and then is smaller than $10^8$ cm$^{-2}$ outwards whatever the model. In the case of settling, the very low N$_2$H$^+$ column density is due to an efficient destruction by electronic recombination in the mid-plane of the disc. Contrary to the case without settling, the UV penetration is more efficient and the electrons are efficiently produced by the reaction S + hv $\rightarrow$ S$^+$ + e$^-$. This result depends on the elemental abundance of sulphur and in our case, we obtain such a strong effect because we have assumed a cosmic value. The electron fractional abundance is about $10^{-6}$ in the case of settling while it is around $10^{-10}$ in the case without settling at 300 au in the mid-plane. The presence of N$_2$H$^+$ at 300 au in the DM Tau disc may then be an indication of a low electron abundance either due to a low UV penetration or a smaller sulphur abundance.
It is quite difficult to compare the column densities computed with and without settling as the settled models produce very different results. For H$_2$S for instance, the effect depends on the radius. Inside 200 au, settling (with and without grain growth) produces larger H$_2$S gas-phase column densities. The reason for this is that the depletion of molecules on the grain surfaces in the molecular layer of the disc is smaller in the case of settling because there are less grains. At radius larger than 200 au, the settling models with grain growth produce smaller column densities because H$_2$S is produced on the grains and desorbed by exothermic surface reactions. The smaller number of grains in the grain growth models results in a smaller production of this species. The case with settling without grain growth produces column densities similar to the cases without settling. Although the number density of grains is smaller above z/h = 1, the number density below is unchanged. The region of the disc below this altitude then contributes mostly to the H$_2$S column density.

4.3 Varying the cosmic-ray ionization rate

Only the column density of N$_2$H$^+$ is changed significantly when the cosmic-ray ionization rate is decreased (models F1 and F2 in Table 1). Decreasing its value by 2 orders of magnitude decreases the N$_2$H$^+$ column density at 300 au by more than 1 order of magnitude. The main reason is that the H$_2^+$ column density is also decreased as N$_2$H$^+$ is formed from the reaction N$_2$ + H$_2^+$ → N$_2$H$^+$ + H$_2$.

5 COMPARING WITH OBSERVATIONS

5.1 Observed and modelled column densities

On the model predicted column densities (Figs 4 and 6), we have reported the observed values at 300 au listed in Table 2. Despite the strong dispersion of model results, some of the observed column densities cannot be reproduced by any of our models at 300 au. HCO$^+$ is always underproduced while CN, HCN, and all S-bearing molecules are overproduced. In the case of N$_2$H$^+$, only models without any settling can produce significant amounts of this molecule at 300 au. Using the settled model (i.e. assuming that settling is occurring in DM Tau protoplanetary disc at 300 au), we have done several other tests to see if we can come closer to the observations. Let’s assume that our most realistic model is the one with settling (occurring at z/h = 1) but without grain growth, starting from an initial cloud with an age of 10$^6$ yr, using the three-phase model (i.e. model E2). For the dust characteristics, one would expect grain growth to have occurred in the disc however, our models show that we need small grains in the mid-plane to deplete CO, otherwise, the CO column density is much higher than the observed one. A more realistic model would require the use of a grain size distribution with both large and small grains. This is the subject of a forthcoming paper (Gavino et al., in preparation). The column densities obtained at 300 au with this model are listed in Table 2 along with the observations. We can see the clear underestimation of HCO$^+$ and N$_2$H$^+$, while CN, CS, H$_2$S, and CCS are overestimated by more than a factor of 10. We have run a number of additional models (starting from model E2) to test some other parameters. These models are summarized in Table 3 and their results are described below. Starting from model E2, we changed the altitude at which the settling occurs: E2.1 (at z/h = 2) and E2.2 (at z/h = 0.5). Since not all neutral abundances peak at the same altitude, the effect is not the same for all species. CS column density for instance is increased in E2.2 while H$_2$S is decreased. The two ions are increased in E2.1 but are still underestimated by 2 orders of magnitude. One way of decreasing the species gas-phase column densities would be to start with a more depleted gas-phase composition. This can be obtained by increasing the density of the parent cloud. We tested a 10 times more dense initial cloud and the resulting column densities are listed as E2.3 in Table 2. The H$_2$S and SO column densities are approximately decreased by a factor of 10 whereas the ones of CS and CCS are not significantly changed. CN and HCN are not sensitive while HC$_3$N is increased, becoming higher than the observed upper limit. The HCO$^+$ column density is strongly increased (becoming close to the observed value) but the value for N$_2$H$^+$ is dramatically decreased. Another way of decreasing the N- and S-bearing species column densities is to decrease the elemental abundances of N and S in the simulations. We tested depletion factors of 10 (E2.4) and 100 (E2.5). If the predicted S-bearing column densities at 300 au are almost linear to the elemental abundance of sulphur, this is not the case for CN and HCN. Decreasing the initial elemental abundances by a factor of 10 have an effect similar to the previous case (E2.3 – 10 times denser initial cloud). Decreasing them by two orders of magnitude goes in the same direction and in this case, even CS and CCS are decreased. Only the ions HCO$^+$ and N$_2$H$^+$ are not changed between models E2.4 and E2.5. In models E2.3–E2.5, the CO column densities are larger by a factor of 10 as compared to the observed value. This is due to the increase in the HCO$^+$ abundance, which produces extra CO upon dissociative recombination. We did one last model (E2.6) similar to E2.3 but with grain growth. In that case, CO is strongly overestimated (more than in the other cases as we have no small grains to depleted gas-phase CO), as are the ions. Only H$_2$S is getting closer to the observations as there are less grains to produce it.

5.2 N$_2$H$^+$ and HCO$^+$

In our models, we have found that N$_2$H$^+$ was highly sensitive to some of the model assumptions. The use of the three-phase model prevents the evaporation of N$_2$, precursor of N$_2$H$^+$, in the gas-phase. In the models with grain settling, this effect is weakened as there are less grains to deplete gas-phase N$_2$. Dust settling however also increases the electron donor abundances (atomic carbon and sulphur) in the molecular layer of the disc. There are then more electrons to destroy N$_2$H$^+$. We have looked at the effect of many other parameters, such as elemental carbon and sulphur depletion (up to 2 orders of magnitude), initial cloud density (larger values to get more depletion of the electron donors), smaller UV irradiation.
The points at 300 au represents the observed column density.

Field (to produce less electrons), higher or smaller altitude transitions for the dust settling, more efficient N\textsubscript{2} photodesorption. All these models produce an underestimation of the N\textsubscript{2}H\textsuperscript{+} column density because the electron abundance between $z/h = 1–2$ is too large. The most promising models are the ones with C and S depletion, i.e. with a smaller abundance of electron donors. These models however underestimate the CO column density. All our models underestimate the HCO\textsuperscript{+} column density at 300 au, except for the models in which the elemental abundance of sulphur is decreased (E2.3, E2.4, and E2.5). The peak of column density for this ion is obtained at $z/h = 1$. At this altitude, this ion is mostly produced by the reaction CO + H\textsubscript{2} → HCO\textsuperscript{+} + H\textsubscript{2} while it is destroyed by electronic recombination. Lowering the electron donor abundance, i.e. sulphur, increases significantly the HCO\textsuperscript{+} column density. Both observed N\textsubscript{2}H\textsuperscript{+} and HCO\textsuperscript{+} large column densities at 300 au are an indication of low electronic fractionation at this radius of the observed discs and at the altitude these two species peak (i.e. between $z/h = 1$ and 1.5). At these altitudes, the main cation is C\textsuperscript{+} (it is H\textsubscript{3}\textsuperscript{+} below), orders of magnitude more abundant than N\textsubscript{2}H\textsuperscript{+} and HCO\textsuperscript{+}. The location of the C/C\textsuperscript{+} vertical transition could then be key for N\textsubscript{2}H\textsuperscript{+} and HCO\textsuperscript{+}. The very low N\textsubscript{2}H\textsuperscript{+} column density obtained with the three-phase models is due to a very efficient depletion of N\textsubscript{2} from the gas, an effect much less efficient in the two-phase model. The representation of prefectly spherical...
grains in these models, even for the grains that undergo growth, is very likely a crude approximation. Kataoka et al. (2013) proposed a fluffy growth mechanism for the dust in protoplanetary discs. Such fluffy grains would produce inhomogeneities in the disc dust distribution and very likely more UV penetration towards the mid-plane of the disc. From a chemical point of view, fluffy grains would have a larger cross-section for collisions with gas-phase species as compared to large spherical grains. In addition, the surface layer would be larger, i.e. offering more possibility for N₂ desorption. The models presented in this paper do not include any X-ray chemistry, which should affect the abundance of these two ions (Glassgold et al. 1997; Aikawa & Herbst 1999; Teague et al. 2015; Cleeves et al. 2015).

5.3 The CCH rings

The CCH molecule has been observed by Bergin et al. (2016) in the DM Tau protoplanetary disc. The emission of this molecule was found to form two rings: one peak around 50 au and one around 350 au. Some of our models with settling provide a bimodal profile with a peak between 50 and 120 au and a weaker one after 200 au (see Fig. 4). The exact location of these peaks depends on the model and the maximum difference (still in column density) between the peaks values and the hole in between is less than a factor of 10. The models without settling do not give these bimodal distributions. In these cases, the CCH column density increases towards the outside until 13–160 au, remains approximately flat until 250 au and then increases again. All models, with and without settling predict a drop of CCH column density inside the 50 central au, simply due to photodissociation. It is interesting to see that the models, which produces nice bimodal radial column densities are the ones obtained with the three-phases model, dust settling (with or without grain growth) and with initial atomic conditions. The same models starting from molecular conditions do not give the second peak. Using a gas-phase model, Bergin et al. (2016) argued that the CCH production can only be efficient with a gas-phase C/O ratio larger than 1. Such elemental ratio would be explained by a depletion of oxygen on the grains before the formation of the disc and driven to the mid-plane of the disc. In other words, this oxygen would not participate to the chemistry in the region where they observe CCH. Such model would be close to our settled model in which we start with molecular initial conditions. In our case we have used a standard C/O ratio of 0.6 and we obtain a CCH column density larger than 10¹⁴ cm⁻² for radii larger than 100 au in all our settled models. We did not consider radial drift of the grains but as CCH abundance peaks above \( z/h = 2 \), this should not affect the column density of this molecule. A more detailed work, including radiative transfer calculations would be required to conclude on this matter.

6 COMPARISONS WITH OTHER DISC MODELS

We will restrain the comparison to gas-grain models for protoplanetary discs such as DM Tau and studying the effect of grain growth and/or settling. Fogel et al. (2011) studied the importance of dust settling (without grain growth) with a two-phase gas-grain model. Although their physical and chemical model is quite different, we can compare their computed column densities listed in their table 2 (second lines without Ly \( \alpha \) radiation, at 250 au) with our models A and C. Looking at tendencies, we obtain similar trends for CO, HCN, CN, and CCH: more column density with settling. For \( \text{N}_2\text{H}^+ \), we also find a decreased column density with settling. In our case, the decreasing HCO\(^+\) column density is less obvious. We obtain column densities of the same order (for the species shown in the two papers), except for \( \text{N}_2\text{H}^+ \), which seems to be much less abundant in our settled case while the other nitrogen bearing species are more abundant. Using a similar chemical model, Vasyunin et al. (2011) studied the effect of both the dust settling and growth, varying in the radial and vertical directions. If their settled model (GS) produces approximately the same gas-to-dust mass ratio at 100 au as ours, their grain growth is much less important. At this radius, the maximum size increase of the dust by a factor of a few above 0.1 \( \mu \)m, i.e. much smaller than our grain growth model. In that condition, they find that the effect of dust evolution is limited at large distances from the star. Since we do not consider the radial dependency of the dust properties, we do not see this effect. The CO column density in their disc with evolved dust (i.e. with settling and grain growth) is much smaller than in all our models with settling (with or without grain growth) while the CO column density in the case with dust evolution is similar to ours in models A, except at 10 au. Secondly, \( \text{N}_2\text{H}^+ \) and CN do not appear to be sensitive to the dust evolution in their case whereas they are strongly sensitive in ours. Comparing their column densities at 100 au with ours for their A5 model (i.e. without settling and grain growth) with our model A1, we obtain similar CO, much higher \( \text{N}_2\text{H}^+ \), CN, HCN, and CCH while much lower \( \text{H}_2\text{S} \) and \( \text{HC}_3\text{N} \). Following this work, Akimkin et al. (2013) used a similar model but improved the treatment.
of the UV penetration more consistent with the evolving dust properties. Despite the fact that the values of the computed column densities are different from Vasyunin et al. (2011), they find similar trends.

7 CONCLUSIONS

In this paper, we have made several chemical models of the protoplanetary disc around DM Tau in order to understand the effect of several model assumptions and try to identify general trends while comparing these models to observed column densities. For this, we used a physical structure mostly based on observed parameters and assuming hydrostatic equilibrium. This 2D structure is then used as input to the Nautilus gas-grain model. We investigated the effect of the following model approximations: two- or three-phase model (two-phase model means that all species on the grains have the same behaviour whereas three-phase means that the top two monolayers are chemically different from the rest of the mantle), the initial conditions (atomic, molecular with different ages), cosmic ray ionization rate, and grain growth and dust settling. The results of a large number of models were compared with observed column densities at 300 au, published in the literature. Our main results are the following:

(1) Without dust settling, the two- and three-phase models differ but the initial conditions have no influence.
(2) With dust settling, molecular abundances become very sensitive to other model assumptions.
(3) Both grain growth and settling reduce the molecular depletion (and also enhances UV penetration). This boosts the CO gas-phase abundance at radii larger than 100 au.
(4) Only N$_2$H$^+$ is affected by the cosmic-ray ionization rate.
(5) N$_2$H$^+$ is the most sensitive species. Its production requires enough N$_2$ in the gas-phase (inhibited by the three-phase model, but boosted by the settling) and a low electron abundance (favoured by low C and S elemental abundances).
(6) Two key aspects of protoplanetary disc chemistry, yet to explore, are the inhomogeneities within these discs and the fluffiness of the grains.

The main conclusion of this work is that the computed chemical composition of protoplanetary discs is very sensitive to both the intrinsic chemical parameters of the models and the physical structure of the disc. In the case of the physical structure, there are still large uncertainties and the current observations are still limited leading to approximations (e.g. observed inhomogeneities, rings and spirals which are not yet properly incorporated in physical models). However, with future observations, utilizing the high spatial resolution of ALMA, these uncertainties will decrease. On the chemical model itself, there are uncertainties in the chemical data but also on some of the processes (not necessarily studied here such as the chemical desorption; Wakelam et al. 2017). As a consequence, it seems very difficult to accurately derive chemical parameters from observed surface densities in discs, without being very cautious. Protoplanetary discs intrinsically exhibit complex structures (observed large density and temperature gradients, small scale/unresolved dust heterogeneities, etc.) and therefore, they are not the best objects where chemical processes can be checked and studied. Studying poorly known or new chemical processes in simpler objects with 1D structure, before incorporating them in disc chemical models, is more efficient and may provide more quantitative results.

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