Formation of simple organic molecules in inner T Tauri disks

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ABSTRACT

Aims. We present time dependent chemical models for a dense and warm O-rich gas exposed to a strong far ultraviolet field aiming at exploring the formation of simple organic molecules in the inner regions of protoplanetary disks around T Tauri stars.

Methods. An up-to-date chemical network is used to compute the evolution of molecular abundances. Reactions of H2 with small organic radicals such as C2 and C3H, which are not included in current astrochemical databases, overcome their moderate activation energies at warm temperatures and become very important for the gas phase synthesis of C-bearing molecules.

Results. The photodissociation of CO and release of C triggers the formation of simple organic species such as C2H2, HCN, and CH4. In timescales between 1 and 106 years, depending on the density and FUV field, a steady state is reached in the model in which molecules are continuously photodissociated but also formed, mainly through gas phase chemical reactions involving H2.

Conclusions. The application of the model to the upper layers of inner protoplanetary disks predicts large gas phase abundances of C2H2 and HCN. The implied vertical column densities are as large as several 1016 cm−2 in the very inner disk (< 1 AU), in good agreement with the recent infrared observations of warm C2H2 and HCN in the inner regions of IRS 46 and GV Tau disks. We also compare our results with previous chemical models studying the photoprocessing in the outer disk regions, and find that the gas phase chemical composition in the upper layers of the inner terrestrial zone (a few AU) is predicted to be substantially different from that in the upper layers of the outer disk (> 50 AU).

Key words. astrochemistry – stars: circumstellar matter – planetary systems: protoplanetary disks – ISM: molecules

1. Introduction

Protoplanetary disks (PPDs) represent an intermediate stage in the evolution from dark clouds towards planetary systems. A detailed characterization of the physical and chemical conditions in such disks is of great interest as they provide the initial conditions for planet formation (see Najita et al. 2007, Bergin et al. 2007 for recent reviews on the subject). Information about the disk chemical composition has mainly come from radio observations, which are sensitive to the outer cool disk, from ~ 100 AU up to ~900 AU (Dutrey et al. 1997, Thi et al. 2004). Only recently the inner regions of PPDs have been probed by means of infrared observations revealing the presence of warm C2H2 and HCN with large gas phase abundances (Lahuis et al. 2006, Gibb et al. 2007) and of complex organic species such as PAHs (Geers et al. 2006, Habart et al. 2006).

Chemical models have mostly concentrated in the study of the outer disk regions (e.g. Aikawa & Herbst 1999), in part motivated by radio observations of several molecular species. Only a few models have focused on the inner disk (< 10 AU) chemistry. Willacy et al. (1998) studied the chemistry in the disk midplane, shielded from stellar and interstellar UV photons, and found that some organic species could be easily formed on grain surfaces. Markwick et al. (2002) studied the chemistry in the inner 10 AU including stellar and interstellar UV radiation, stellar X-rays and radionuclides decay as sources of ionization, and adsorption/desorption on grains but not mantle chemistry, i.e. formation and destruction of molecules occurs only in the gas phase. They calculated large abundances of organic species such as CH3 but they did not discuss the main chemical routes to form them. Semenov et al. (2004) and Ilgner & Nelson (2006) extensively discussed the ionization degree over a wide range of radii and heights over the midplane. Woods & Willacy (2007) predicted that benzene formation can be efficient in the midplane of the inner disk (< 3 AU), a dense region protected against UV photons where grain surface reactions play an important role in building-up a complex organic chemistry.

In this Paper we investigate gas phase routes to form simple organic molecules, such as acetylene (C2H2), hydrogen cyanide (HCN), and methane (CH4), in a dense and warm oxygen-rich gas exposed to a strong far-UV (FUV) field (hv < 13.6 eV). We apply this model to the inner region (< 10 AU) of a protoplanetary disk around a T Tauri star and compare with the abundances recently derived from observations.

2. Chemical model

Physical models of protoplanetary disks suggests a flared-up structure where both the gas density and temperature vary enormously depending on the radius r from the star and height z over the disk equatorial plane (see Dullemond et al. 2007 for a review). The gas density decreases radially outward as a power law of r, and in the vertical direction it decreases as z increases in roughly an exponential way. The gas heating across the disk is dominated by the stellar irradiation of the surface layers at large radii (r ≥ 10 AU), thus the kinetic temperature increases with z, while at small radii (r ~ a few AU) viscous dissipa-
tion may become an important heating mechanism in the midplane regions (D'Alessio et al., 1998, 1999). A flared disk is significantly exposed to the strong UV field from the central star. Therefore, there exists a layer of low AV where the disk material is being photoprocessed (Aikawa & Herbst, 1999; Willacy et al., 2000). Here we focus on the chemistry that takes place in the photon-dominated region (PDR) of the inner disk (r < 10 AU), where typically temperatures are several hundreds of K and gas densities range from 10⁶ to 10¹¹ cm⁻³.

In order to qualitatively understand the chemical routes to form simple organic molecules in the PDR of the inner disk, we have firstly performed a few time dependent chemical models in which chemical abundances evolve under fixed physical conditions representative of this region. We consider an O-rich gas with all the hydrogen initially as H₂, all the carbon as CO, the oxygen in excess as H₂O and all the nitrogen as N₂. We use solar abundances (Asplund et al., 2005) and assume that 50 % of C and 65 % of O are in carbon and silicate grains respectively. Therefore, the initial abundances relative to the total number of H nuclei n_H, where n_H = 2 n(H₂) + n(H), are x(CO) = 1.25 × 10⁻⁴, x(H₂O) = 3.5 × 10⁻³, x(N₂) = 3.0 × 10⁻⁵, and x(He) = 8.5 × 10⁻². We adopt a gas density of n_H = 2 × 10⁸ cm⁻³ and run several models with five different kinetic temperatures: T = 100, 300, 500, 750 and 1000 K.

We first consider a FUV illuminated model with a FUV field strength of Φ = 50,000 (relative to the Draine interstellar radiation field, Draine, 1978), the value at 10 AU according to FUV observations of various T Tauri disks (Bergin et al., 2003, 2004). For this FUV field and gas density, photoprocessing of the gas occurs in a range of visual extinctions AV ~ 0.1 - 5. If AV ≤ 0.1 the gas is very exposed to the UV radiation and molecules are destroyed, and in the case of AV ≥ 5 photoprocessing occurs only marginally. Here we investigate the chemical evolution at a mean extinction value of AV = 2.5. We then consider a separate X-ray illuminated model, in which the gas is solely exposed to X-rays but not FUV photons, aiming at evaluating the influence of these two energy sources on the chemistry. There is evidence for X-ray emission in PPDs (Reijerse & Montmerle, 1999). Its main effect on the chemistry is the ionization of the gas producing high energy photoelectrons (ph-e⁻) which further ionize and dissociate the species. To simulate this effect, we have enhanced the cosmic rays ionization rate by a factor of 1000 over the standard interstellar value, thus we take ζ = 1.2 × 10⁻¹⁴ s⁻¹, which is reasonable for the inner regions of PPDs (Ilee & Glassgold, 1999).

The chemical code used has been described in Cernicharo (2004). The species included in the model are given in Table 1. They were selected on the basis of previous calculations including a larger number of species which allowed us to identify the most important chemical processes forming C-bearing species. The chemical network used was taken from Cernicharo (2004) and Agúndez & Cernicharo (2006), and was checked against the NIST Chemical Kinetics Database (http://kinetics.nist.gov) and the latest version of the UMIST database for astrochemistry (Woodall et al., 2007). We include neutral-neutral and ion-molecule bimolecular reactions, three body processes and thermal dissociations.

Important reactions involved in the built-up of small organic molecules at high temperatures are those of H₂ with radicals such as C₂ and C₂H. These reactions have been studied in the laboratory over a relatively wide temperature range: C₂ + H₂ → C₂H + H between 295 and 493 K (Pitts et al., 1982) and C₂H + H₂ → C₂H₂ + H in the range 178-440 K (Peeters et al., 1996; Opansky & Leon, 1996). They have moderate activation barriers, of about 1400 K, which make them very slow in the cold interstellar medium, although at high temperatures they become rapid enough to control the abundance of C-bearing species. Their rate constant expressions are not included in current astrochemical databases, which are somewhat biased toward low temperature chemistry, but can be found in Table 1 of Cernicharo (2004). The formation of H₂ on grain surfaces is included with a rate constant of 3 × 10⁻¹⁷ cm³ s⁻¹.

In order to properly take into account FUV line self-shielding, we explicitly calculated a grid of H₂ and CO photodissociation rates and C photoionization rates, appropriate for the physical conditions in the inner disk. This calculation was performed with the Meudon PDR code (Le Bourlot et al., 1993). The model has been described in detail elsewhere (Le Petit et al., 2004; Goicoechea & Le Bourlot, 2007). In par-

### Table 1. Species included in the model

| H | C₂H | NH₃ | C⁺ | CH⁺⁺ | HCO⁺ | CN⁺ |
|---|-----|-----|----|------|------|-----|
| H₂ | C₂H₂ | NH₃ | C⁺ | C⁺⁺ | HCO⁺ | CN⁻ |
| He | O | CN | O⁺ | CH⁺⁺ | H₂CO⁺ | HCN⁺ |
| O | OH | HCN | O⁻ | CH⁺⁺ | H₂CO⁺ | HCNH⁺ |
| O₂ | H₂O | HNC | N⁺ | CH⁺⁺ | CO⁺⁺ | NH⁺ |
| N | CO | H₂CN | H₂⁺ | OH⁺ | HCO⁺⁺ | H₂NC⁺ |
| CH | CO₂ | NO | H₁⁺ | OH⁻ | N₂⁺ | CNC⁺ |
| CH₂ | HCO | e⁻ | CH⁺ | H₂O⁺ | NH⁺ | NO⁺ |
| CH₃ | H₂CO | H⁺ | CH₂⁺ | H₂O⁺ | NH⁺⁺ | HNO⁺ |
| CH₄ | N₂ | H⁻ | CH⁺⁺ | O₂⁺ | NH⁺⁺ | HNC⁺ |
| C₂ | NH | He⁺ | CH⁺⁺ | CO⁺⁺ | NH⁺⁺ |

![Fig. 1. Evolution of C₂H₂, HCN and CH₄ abundances for a chemistry driven by FUV photons (left) and by X-rays (right). Different curves correspond to different gas temperatures.](http://www.udfa.net)
Fig. 2. Scheme with the main synthetic routes for the formation of C$_2$H$_2$, HCN and CH$_4$ from C, C$^+$ and N. Reactions with a high activation energy (Ae) are indicated by a thick arrow.

In particular, we computed the depth dependent FUV radiation field for the range of parameters ($\chi$, $n_H$, $T_1$ and initial abundances) considered in the FUV illuminated model. H$_2$, CO and C photorates were then consistently integrated over these fields and used afterwards in the time-dependent chemical calculations. Dust properties are known to influence the resulting rates (e.g. Goicoechea & Le Bourlot 2007). Here we have assumed the standard grain size distribution proposed for the ISM (Mathis et al. 1977). All the other photorates included in the FUV illuminated model have been taken from the UMIST 2006 database. In the X-ray illuminated model we have included reactions induced by X-rays, for which we have adopted the cosmic ray induced reactions rates enhanced by a factor of 1000.

3. Chemical routes to C$_2$H$_2$, HCN and CH$_4$

Left panels in Fig. 1 show the evolution of C$_2$H$_2$, HCN, and CH$_4$ abundances for several gas temperatures in the FUV model. The synthesis of these organic species can be divided into three steps.

(i) Photodissociation of CO and N$_2$ with release of atomic C, C$^+$ (produced by further photoionization of C) and atomic N. The time scale of this step depends only on the FUV field strength, thus on $\chi$ and $n_H$.

(ii) Atom–molecule transition, from the primary species C, C$^+$ and N to simple C-containing molecules. Important reactions which drive this transition are (see Fig. 2) radiative associations such as C + H$_2$ → CH$_2$ + hv and C$^+$ + H$_2$ → CH$_2$ + hv, and rapid neutral-neutral reactions such as C + NO → CN + O, where NO comes from the reaction between N and OH. These reactions are nearly temperature independent, thus their rate depends basically on the gas density: the higher the gas density the faster they proceed. Bimolecular reactions such as C$^+$ + H$_2$ → CH$^+$ + H and C + H$_2$ → CH + H begin to overcome their activation barriers at temperatures above 400 and 700 K respectively. Three body reactions are not competitive, compared to radiative associations, for gas densities below ∼ 10$^{13}$ cm$^{-3}$.

(iii) Processing of simple molecules, CH$_2$, CH$_2^+$ and CN, into more complex species, represented by C$_2$H$_2$, HCN and CH$_4$ in our model. As Fig. 2 shows this processing occurs through bimolecular reactions involving H$_2$, C and N. Reactions of neutral species with H$_2$ have energy barriers higher than 1000 K, which introduces a strong temperature dependence. This is very marked for CH$_4$ because its formation involves the reactions with the highest energy barriers, making it abundant only at temperatures above ∼ 700 K. The synthesis of organic molecules in the O-rich gas requires that atomic carbon, produced by the dissociation of CO, incorporates into C-bearing species faster than reverting to CO. This is achieved by different mechanisms at low and high temperatures. Below ∼ 400 K atomic oxygen is not converted into OH (the reaction O + H$_2$ → OH + H has an activation energy of ∼ 5000 K) and the main CO-forming reaction, OH + C → CO + H, is inhibited. Thus atomic carbon can react with other species instead of reverting to CO. The main obstacle to form C-bearing species at low temperature is the energy barrier of several reactions. For example, C$_2$H$_2$ and HCN reach low abundances at 100 K because the reactions of C$_2$H and CN with H$_2$ have activation barriers. Above ∼ 400 K atomic oxygen is efficiently converted into OH, which may react with C to form CO but reacts faster with H$_2$ to form water. Thus, most of the oxygen forms H$_2$O, and CO does not reach its maximum abundance allowing atomic carbon to form C-bearing molecules. At high temperatures most of the reactions shown in Fig. 2 overcome their energy barriers producing a rich C-based chemistry.

The chemistry driven by X-rays (see right panels in Fig. 1) is rather similar to that initiated by FUV photons except that the dissociation of CO and N$_2$, step i, is carried out by collisions with photoelectrons: CO + ph$-e^-$ → C + O and N$_2$ + ph$-e^-$ → N + N and by reactions with He*: CO + He* → C$^+$ + O + He and N$_2$ + He* → N + N$^+$ + He. For our chosen FUV and X-ray field strengths, these reactions are about 100 times slower than photodissociations, which makes the chemical time scale to be increased by roughly the same factor, i.e. molecules form later (see Fig. 1). On the other hand, the lower dissociating strength of X-rays, compared to that of the FUV field, results in a higher steady state abundance of C-bearing molecules. This occurs because these steady state abundances depend on the balance between the rates at which molecules are formed (through chemical reactions) and are destroyed (by either FUV photons or X-rays). The destruction rates are lower in the X-ray model, compared to the FUV model, and thus chemical reactions have
more time to form molecules until steady state is reached. As a consequence, in the X-ray model C$_2$H$_2$ and HCN reach high abundances with little dependence on the temperature, once this is above a threshold value of about 300 K (necessary for some important reactions to overcome their energy barriers). Thus, the main differences in the FUV and X-ray models arise because of the different dissociating strengths. In fact, if the ionization rate is enhanced by a factor of 100 in the X-ray model, then the evolution of the molecular abundances and their steady state values become much closer to those of the FUV model.

The relative intensity between the FUV and X-ray fields will establish whether the chemistry is dominated by FUV photons (PDR) or X-rays (XDR). X-rays will drive the chemistry in those regions of protoplanetary disks with a column density toward the star $\gg 10^{22}$ cm$^{-2}$, which are shielded against FUV photons ($A_V > 5$) but not against X-rays (Maloney et al., 1996). In regions with column density values $\gg 10^{23}$ cm$^{-2}$ X-rays are severely attenuated and do not affect to the chemistry.

In summary, according to the models the formation of acetylene, hydrogen cyanide and methane occurs in time scales between a few years and a few thousands of years, and is clearly favored at high temperatures. They reach a steady state in which they are destroyed by photodissociation (or X-ray induced reactions) and reactions with H but are continuously re-formed through reactions involving H$_2$. Thus, the relative rates between re-formation and destruction will determine the steady state abundance. A high abundance of H$_2$, maintained due to its continuous formation on grain surfaces, is essential for accelerating the process

$$A + H_2 \rightarrow AH + H$$

where A is an atom or simple radical. In this way organic species AH can reach a high steady state abundance. In fact, if the formation of molecular hydrogen on grain surfaces is suppressed, then H$_2$ is effectively dissociated (due to its continuous participation in reactions of the above type) in about $10^3$-$10^4$ years in the FUV model and $10^6$ years in the X-ray model, a time at which most of the molecules are also destroyed, thus becoming transient species.

The steady state abundances of C$_2$H$_2$, HCN and CH$_4$ are very sensitive to the kinetic temperature (especially in the FUV model), and may vary from $<10^{-10}$ at 100 K up to several $10^{-5}$ at 1000 K. The chemistry of these simple organic molecules also depends, although to a lesser extent, on other parameters of the model. The FUV/X-ray field and gas density affect the chemical time scale (a decrease in any of these parameters will make the chemistry proceed slower) and also to the steady state abundance (higher $n_{H_2}/\chi$ or $n_{H}/\chi$ ratios favor the formation of molecules compared to the destruction by FUV photons or X-rays and thus favor a higher steady state abundance). The C/O ratio is also an important parameter which affects the wealth of C-bearing molecules. Obviously, values close to 1 favor a rich C-based chemistry while low values make it more difficult. We nevertheless find that only adopting C/O ratios substantially lower than the solar value (0.54, Asplund et al., 2005), then the abundances of C-bearing species turn to decrease appreciably. The choice of the initial abundances is also an issue when modelling the chemistry in protoplanetary disks and several options have been used in the literature (see e.g. Willacy et al., 1998, Semenov et al., 2004). If we assume an initial atomic composition, instead of a molecular one, we find that the chemical timescale is noticeably reduced (the step i has been already carried out) although the steady state abundance is essentially the same. The non-dependence of steady state abundances on the initial composition is a consequence of the strong photoprocessing of the material, and indicates that molecular abundances in the PDR of inner disks are not affected by the chemical history of the gas. The actual situation in PPDs may be different if mixing motions bring material from outer cold regions and affect the chemical balance in the PDR.

At this point it would be interesting to discuss our results by comparing with some other PDR models appeared in the literature. A detailed study of the chemistry in PDRs has been carried out by Sternberg & Dalgarno (1995). They modelled a dense ($n_H = 10^6$ cm$^{-3}$) and highly irradiated ($\chi = 2 \times 10^5$) plane parallel cloud. They considered a reduced set of chemical species, including S- and Si-bearing molecules, of low chemical complexity (e.g., they did not included species with more than one carbon atom). The calculated steady state abundances for C-bearing molecules such as HCN and CH$_2$ are lower than $10^{-9}$ in the warm region of $A_V = 0.2$ ($T_k = 3000$-$5000$ K), although radicals such as CH have abundances as high as $10^{-6}$. The most remarkable difference between their model and ours lies in the $n_{H_2}/\chi$ ratio. The much lower value of their model (5 compared to the value of 4000 adopted by us) favors photoprocesses compared to chemical reactions and results in low abundances of closed-shell organic molecules. A more recent PDR model by Teissier et al. (2004) and Fety et al. (2005), focuses on the study of long carbon chains in PDRs such as the Horsehead Nebula. They consider a model with a gas density $n_H = 2 \times 10^4$ cm$^{-3}$ and a FUV field strength $\chi = 60$, thus the $n_{H_2}/\chi$ ratio is 300, much closer to the value adopted by us. Their model nevertheless predicts C-chains such as C$_2$H and C$_2$H$_2$ to have abundances $\lesssim 10^{-8}$. Besides a different chemical network used, another important difference between their models and ours resides in the kinetic temperature. In their model the abundance of hydrocarbon radicals peaks in the region of $A_V = 1$-$2$, where $T_k$ is lower than 100 K. At these relatively low temperatures hydrocarbons do not reach high abundances since their formation is inhibited by various reactions which have moderate activation barriers.

4. Abundances in the inner region of a T Tauri disk

The physical conditions adopted in the chemical models discussed in the previous section are assumed to be representative of those prevailing in the PDR of inner T Tauri disks. Nevertheless, real conditions at different points ($r$, $z$) in such disks span over a wide range around our adopted values. In order to check whether photochemistry can account for the formation of simple organic molecules with abundances comparable to those observed, we have adopted the disk physical structure, gas density and temperature at each point, from a steady state flared accretion disk model provided by P. D’Alessio (D’Alessio et al., 1998, 1999). The disk model has a mass accretion rate $\dot{M} = 10^{-8}$ $M_\odot$ yr$^{-1}$, grain properties similar to those of interstellar dust (maximum dust grain size is 0.25 $\mu$m), and a central star with $M_*= 0.7 M_\odot$, $T_*= 4000$ K and $R_*= 2.6 R_\odot$. The chemical model was run for a grid of $18 \times 16$ ($r \times z$) points covering the upper layers of the very inner disk $r = 0.1$-$3$ AU. The modelled region covers the CO/C/C$^+$ transition, from an inner height $z_{in}$ (where the gas is well shielded against stellar and interstellar light and all the carbon is as CO) up to the disk surface defined by the height $z_{out}$ (where the gas is completely exposed to the FUV field and all the carbon is as C$^+$, see Fig. 3).

We consider that the gas is affected by stellar and interstellar FUV radiation and by cosmic rays. For simplicity we neglect the effect of stellar X-rays, although in the case of strong X-ray emitter T Tauri stars the abundances of certain
molecules, which are sensitive to the ionization rate such as HCO$^+$ or HNC, may be substantially affected. The influence of an enhanced ionization rate on the molecular abundances has been discussed by Aikawa & Herbst (1999) and Markwick et al. (2002). Concerning the formation of simple organics such as C$_2$H$_2$, HCN and CH$_4$, an enhanced ionization rate does not greatly change the results from those obtained when only a FUV radiation field is considered.

We treat the FUV field considering that each point ($r$, $z$) of the disk is exposed to two separate radiation fields, stellar and interstellar. The FUV flux emanating from the star is diluted geometrically and attenuated by the column density of material in the line of sight toward the point ($r$, $z$). The interstellar radiation is assumed to penetrate into the disk in the vertical direction. For simplicity the radiative transfer of stellar and interstellar FUV photons is treated separately, in a plane parallel 1+1D approach. The computed depth-dependent H$_2$, CO and C photorates are then used in the grid of time-dependent chemical models. Although a coupling between the attenuation of the FUV field and the molecular abundances exists, a self-consistent 2D solution of the radiative transfer problem, together with the chemical evolution, is beyond the scope of this study.

The spectral shape of the FUV field emitted by a T Tauri star significantly differs from that of the interstellar radiation field (ISRF). Therefore, photodissociation/ionization rates can be drastically different in the presence of a T Tauri radiation field compared to an enhanced ISRF. This problem has been addressed by van Dishoeck et al. (2006), who calculated photodissociation/ionization rates of various species for a 4000 K blackbody radiation field, typical of T Tauri stars, and compared with the values obtained under the ISRF. They found that, for the same integrated flux between 912 and 2050 Å, the photorates of species such as H$_2$, CO or N$_2$ decrease by several orders of magnitude when the stellar radiation field, instead of the ISRF, is used.

Given the impact of the true stellar radiation field on the photochemistry, we have used the Meudon PDR code to generate a new grid of depth-dependent H$_2$ and CO photodissociation rates and of C photoionization rates. Instead of adopting a given ISRF enhancement, the FUV field is generated by a T Tauri star simulated by a blackbody at 4000 K and a stellar radius of 2.6 $R_*$.

The intensity in the Ly$\alpha$ line, at 1216 Å, can be very important in T Tauri stars (Bergin et al., 2003), and can have a large impact on the photorates of some species, such as OH and CH$_4$, although it does not affect to species such as H$_2$, CO and N$_2$, because their main photoband absorption bands lie short of 1216 Å. For simplicity, here we do not consider enhanced emission at the Ly$\alpha$ line. The resulting radiation field at the disk surface is then determined by the geometrical dilution from the star. We obtain stellar photorates for H$_2$, CO and C that are about 5 orders of magnitude smaller than the ISRF photorates (for the same integrated flux in the FUV), in agreement with the values reported by van Dishoeck et al. (2006). Stellar photorates for species other than H$_2$, CO and C are taken from van Dishoeck et al. (2006) when available, or are assumed to be equal to the ISRF photorates otherwise. The visual extinction of stellar and interstellar light at a given point is assumed to be proportional to the column density of hydrogen nuclei N$_{H}$ in the direction toward the star and outward in the vertical direction respectively. We use the standard relation $A_V = N_H$ (cm$^{-2}$) / 1.87 $\times$ 10$^{21}$ $A_{0.1}$ (Bohlin et al., 1978), adequate for the ISM grain size distribution. In the model region, the stellar FUV field has a strength which is higher than the ISRF by orders of magnitude ($A_V > 10^{-4}$). However, since the stellar photorates are for many species lower than the ISRF photorates by orders of magnitude, the photoprocessing of the material in the PDR layers is to a great extent dominated by the interstellar, rather than stellar, FUV radiation field.

The gas densities are 10$^8$-10$^{11}$ cm$^{-3}$ in the sampled region and the gas kinetic temperature, assumed equal to the dust temperature in the disk physical model, ranges from 100 to 1000 K. According to Kamp & Dullemond (2004), the gas temperature significantly exceeds the dust temperature in the very upper layers with a visual extinction of stellar radiation $A_V < 0.1$. In this superheated surface layer our model most probably underestimates the gas temperature. The implications however for the calculated abundances of simple organic species are little since this is a region where CO has fully dissociated and most of the carbon is as C$^\cdot$. We do not consider adsorption/desorption on dust grains since at the high temperatures prevailing in the studied layers ($T > 100$ K) all the molecules are supposed to have been evaporated from grain mantles and to be in the gas phase.

The abundance distributions at time 10$^6$ yr, when steady state has been reached, for C$_2$H$_2$, HCN, and CH$_4$ are plotted in Fig. 3. A thin shell of the disk contains simple organic species with abundances ranging from 10$^{-8}$ (in the outer part, at 2-3 AU) up to 10$^{-4}$ (in the inner 1 AU where gas is hot). The amount of material contained in such a thin layer is however remarkably large due to the large gas densities prevailing there. Vertical column densities within the disk PDR can be calculated by integrating the gas molecular densities in the $z$ direction (see Fig. 4). The column densities of H$_2$ and CO are also shown to visualize the molecular abundances relative to these two species.

Labouis et al. (2006) observed toward IRS 46 hot (> 350 K) C$_2$H$_2$, HCN and CO$_2$ with column densities of 3, 5 and 10 $\times$ 10$^{16}$.
The presence of HNC is strongly related to that of HCN, since it is formed through proton transfer to the latter species and subsequent dissociative recombinations. The column density of NH$_3$ has a radial profile similar to that of CH$_4$, but with a value about one order of magnitude lower. Also it is interesting to note that the abundance ratio of a closed-shell molecule and its related radical (e.g. H$_2$O/OH and HCN/CN) is quite large (about 100-1000), in contrast with the values close to 1 predicted in the outer disk, $r > 50$ AU, (e.g. Willacy et al. 2000). This is a consequence of the large gas densities prevailing in the inner disk which, in spite of the strong FUV field, tend to favor the formation of closed-shell species through reaction (1).

5. Conclusions

In summary, we have shown that a strong FUV/X-ray field may efficiently drive a rich C-based chemistry in a dense O-rich gas with kinetic temperatures of several hundreds of degrees Kelvin. The application of our PDR chemical model to the inner regions of a T Tauri disk yields gas phase abundances for simple organic molecules such as C$_2$H$_2$ and HCN that range from $5 \times 10^{-5}$ to $10^{-8}$, depending on the radius. This translates into vertical column densities as large as $10^{17}$ cm$^{-2}$ in the very inner disk (< 1 AU) down to $10^{15}$ at 3 AU. The model thus explains the large column densities of C$_2$H$_2$ and HCN (several $10^{16}$ cm$^{-2}$) observed in the inner regions (a few AU) of IRS 46 and GV Tau disks (Lahuis et al. 2006; Gibb et al. 2007). The huge variations with radius of the gas density and temperature within the PDR layer of protoplanetary disks results in a quite different gas phase chemistry, i.e. different molecular abundances and ratios between them, in the inner terrestrial zone compared to the outer regions at several hundreds of AU. Our results await confirmation from further observations of different molecular species probed in the inner regions of protoplanetary disks.

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