SHIELDING BY WATER AND OH IN FUV AND X-RAY IRRADIATED PROTOPLANETARY DISKS

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ABSTRACT

We present an integrated thermal–chemical model for the atmosphere of the inner region of a protoplanetary disk that includes irradiation by both far-ultraviolet (FUV) and X-ray radiation. We focus on how the photodissociation of H$_2$O and OH affects the abundances of these and related species and how it contributes to the heating of the atmosphere. The dust in the atmosphere plays several important roles, primarily as the site of H$_2$ formation and by absorbing the FUV. Large amounts of water can be synthesized within the inner 4 AU of a disk around a typical classical T Tauri star. OH is found primarily at the top of a warm region where the gas temperature is $T_g \approx 650$–1000 K and H$_2$O is found below it, where the temperature is lower, $T_g \approx 250$–650 K. The amounts of H$_2$O and OH and the temperatures of the regions in which they formed are in agreement with recent Spitzer measurements and support the notion of the in situ production of water in the inner regions of protoplanetary disks.

We find that the synthesized water is effective in shielding the disk mid-plane from stellar FUV radiation.

Key words: astrochemistry – protoplanetary disks – X-rays: stars

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1. INTRODUCTION

The gas and dust in protoplanetary disks are both important in the formation of stars and planets (e.g., Dullemond & Monnier 2010; Williams & Cieza 2011). Significant progress has been made in recent years by observing the gas with ground-based facilities and the Spitzer and Herschel space observatories. A variety of molecules have been detected in the inner planet forming regions of these disks, notably water and simple organic molecules (Carr et al. 2004; Carr & Najita 2008, 2011; Pascucci et al. 2009; Pontoppidan et al. 2010a, 2010b; Salyk et al. 2011; Najita et al. 2013). The presence of water is of particular astrophysical and astrobiological significance. Water can be as abundant as CO in protoplanetary disks, making it the third most abundant molecule after H$_2$ and CO. Water and dust can also play important physical and chemical roles, which is a focus of this article.

Numerous models of protoplanetary disk chemistry have been developed for the purpose of establishing and analyzing diagnostic spectral lines of the gas (Markwick et al. 2002; Kamp & Dullemond 2004; Nomura et al. 2005; Kamp et al. 2007, 2009; Agúndez et al. 2008; Gorti & Hollenbach 2008; Woods & Willacy 2009; Woitke et al. 2009a, 2009b; Kamp et al. 2010, 2011; Fogel et al. 2011; Thi et al. 2010; Heinzeller et al. 2011; Walsh et al. 2012; Arens et al. 2011, 2012; Bruderer et al. 2012; Meijerink et al. 2012; Akimkin et al. 2013). These models tend to differ in significant physical details (e.g., how the gas is irradiated and how it is heated), as well as in the chemistry. In addition to our earlier paper (Glassgold et al. 2009), several other models produce significant levels of water in the inner disk (e.g., Woods & Willacy 2009; Thi et al. 2010; Fogel et al. 2011; Heinzeller et al. 2011; Arens et al. 2011), and thus support the idea of in situ formation in the gas phase. If this conclusion can be strengthened, as we attempt to do here, it would diminish the need for an origin of water from the transport of icy particles and bodies (e.g., Ciesla & Cuzzi 2006).

In 2009 Bethell and Bergin called attention to the possibility that the disk mid-plane is shielded from dissociating far-ultraviolet (FUV) radiation by water and OH in the disk atmosphere (Bethell & Bergin 2009, hereafter BB09). They described how including such molecular shielding could account for several features of the water and OH emission observed by Spitzer. Molecular shielding becomes important when grains have settled out of the atmosphere and is no longer the dominant absorber of UV radiation. Shielding by molecules can limit the penetration of the FUV through the disk atmosphere and thus affect the transition from regions dominated by atomic oxygen to regions with oxygen-bearing molecules.

To make these points, BB09 adopted an FUV irradiated isothermal atmosphere and employed a simplified chemistry for OH and H$_2$O. In particular, they assumed hydrogen to be completely in molecular form. One potential difficulty with this assumption is that the formation of H$_2$O and OH requires H$_2$ as a precursor, and the formation of H$_2$ on grains may be limited when the grain surface area is reduced by grain settling. A more complete thermal–chemical model is, therefore, needed to explore how well shielding by H$_2$O and OH works when grain settling is advanced. This topic has not been pursued by earlier thermal–chemical models. For example, the protoplanetary disk model (ProDiMo) of Woitke et al. (2009a; see also Arens et al. 2011; Meijerink et al. 2012), while detailed and inclusive of many thermal and chemical processes, includes molecular shielding by H$_2$ and CO, but not H$_2$O and OH.

In this paper we incorporate UV irradiation and molecular shielding by H$_2$O and OH into our earlier thermal–chemical model of the inner disk atmosphere to explore the role of H$_2$O and OH as UV opacity sources. We carefully treat a number of physical processes related to the abundance of water and OH. Unlike BB09, who prescribed two isothermal layers, we evaluate the temperatures in the transition between these two regions of the atmosphere by considering the chemistry-dependent thermal rate equations. In our previous study of water with a purely X-ray model (Glassgold et al. 2009), we obtained significant levels of water in the inner regions of a typical T Tauri star disk, but much lower levels of OH than were found by Spitzer (NAG11) or in the model of BB09. Therefore, in this paper we
also explore whether a simple UV-irradiated thermal–chemical model can account for the properties of both the H$_2$O and OH emission detected with Spitzer.

In the next section we describe how FUV photodissociation is treated in our model, including how it adds to the heating of the disk atmosphere. One of our goals is to describe our assumptions and the underlying reasons for them in enough detail to allow useful comparison with the results of other investigations. In Section 2, we describe improvements to our model, most importantly the addition of UV photodissociation and its associated heating. We have also added photoelectric and H$_2$ formation heating to our model.

2. THERMAL–CHEMICAL IRRADIATION MODEL

The disk model that we irradiate with both X-rays and FUV is derived from earlier versions with only X-rays (Glassgold et al. 2004, hereafter GNI04; Glassgold et al. 2009, hereafter GMN09; Najita et al. 2011, hereafter NAG11; Ádámkovics et al. 2011, hereafter AGM11). The underlying disk structure is a hydrostatic equilibrium calculation by D’Alessio et al. (1999) for a uniform mixture of dust and gas in Keplerian rotation heated by star light and accretion. GNI04 focused on the gas component separately by including gas cooling and heating (including X-rays) and a chemical model appropriate to the disk atmosphere at small radial distances. In subsequent papers, GMN09 showed how H$_2$O could be formed in situ in inner disk atmospheres if H$_2$ forms on warm grains (Cazaux & Tielens 2004). NAG11 used this model to study the formation of the photodissociation of H$_2$O and OH and then describe results of an improved treatment of X-ray ionization and heating. In this section, we present a detailed discussion of the photodissociation of H$_2$O and OH and then describe several improvements to the thermal part of the model. The present treatment of photodissociation can be extended to other molecules and we will report on these applications in a future publication.

2.1. Photodissociation

The photodissociation rate for a molecule at a particular radial distance $r$ and altitude $z$ above the disk mid-plane is given by,

$$G(r,z) = \int_{\lambda_0}^{\lambda_f} F(\lambda; r,z) \sigma(\lambda) d\lambda,$$

where $F(\lambda; r,z)$ is the spectral distribution of the local FUV photon number flux in cm$^{-2}$ s$^{-1}$ Å$^{-1}$ over the wavelength range from $\lambda_0$ to $\lambda_f$ in Å, and $\sigma(\lambda)$ is the photo-absorption cross section in cm$^2$. There are extensive laboratory measurements of the H$_2$O cross section, and we use the high-resolution results presented in Figure 1 of Mota et al. (2005), displayed here as the solid line in Figure 1. For OH we use the theoretical results from Figure 2 of van Dishoeck & Dalgarno (1984), displayed as the dashed line in Figure 1. We average the cross sections over nine 100 Å bands, and display them as the thick gray lines in Figure 1. We ignore the effects on OH and H$_2$O of FUV radiation below 1100 Å due to strong line self-shielding by abundant CO and H$_2$ of the radiation from 911.7–1108 Å.

We calculate $F(\lambda; r,z)$ from measurements of the FUV luminosity, $L_{FUV}$, of T Tauri stars obtained with the Hubble Space Telescope (HST) by Yang et al. (2012),

$$F(\lambda; r,z) = \frac{L_{FUV}}{4\pi(r^2 + z^2)} \frac{\lambda}{hc} e^{-(\tau_d + \tau_{H_2O} + \tau_{OH})},$$

where $\tau_d$, $\tau_{H_2O}$, and $\tau_{OH}$ are the optical depths due to dust, H$_2$O, and OH, respectively. The measured luminosities imply a value of $F(\lambda; r,z)$ at $\lambda = 1000$ Å and $r = 1$ AU for a classical T Tauri star (CTTS) that is 10 million times larger than the interstellar radiation field (Habing 1968).

Yang et al. (2012) report extinction-corrected FUV luminosities for CTTS between 1250 and 1700 Å excluding Ly$\alpha$. We adopted $L_{FUV} = 0.013 L_\odot = 5 \times 10^{31}$ erg s$^{-1}$, a useful value for comparison with the results of Spitzer spectroscopy, as molecular emission has been reported for sources with both larger and smaller $L_{FUV}$ (Carr & Najita 2011). This value corresponds to a mean luminosity per 100 Å of $L_{band} = 1.1 \times 10^{31}$ erg s$^{-1}$ when taking into account the 450 Å bandpass of the measurements. In the absence of equivalent measurements outside the 1250–1700 Å range, we use the same $L_{band}$ in all nine 100 Å bands from 1100 to 2000 Å for a total FUV luminosity of $0.025 L_\odot$. This value is similar to, but smaller than, the smallest FUV luminosity used by BB09 (0.1 $L_\odot$), who assume that the FUV luminosity includes a large contribution from Ly$\alpha$ radiation.

The Ly$\alpha$ line is estimated to make up 70%–90% of the FUV luminosity of CTTS (Schindhelm et al. 2012). Since we do not include Ly$\alpha$ irradiation in our models, because of its complicated radiative transfer (i.e., H$_1$ scattering), our adopted reference FUV luminosity underestimates the FUV incident on the disk by a factor of 3–10. One way to roughly account for the Ly$\alpha$ luminosity of the source would be to assume that our reference FUV luminosity represents 3–10 times the continuum $L_{FUV}$ values. In that case, our adopted reference FUV luminosity of 0.025 $L_\odot$ would still be relevant to CTTS studied with Spitzer because of the broad $L_{FUV}$ distribution of CTTS (Yang et al. 2012). Consistent with the focus on the photodissociation of H$_2$O and OH, we add two reactions to the chemical network used in AGM11,

$$\text{H}_2\text{O} + h\nu \rightarrow \text{OH} + \text{H} \quad \text{(R1)}$$

and

$$\text{OH} + h\nu \rightarrow \text{O} + \text{H}. \quad \text{(R2)}$$
We ignore the role of molecular extinction in producing excited states of OH and O.

The opacity due to molecular extinction is given by

$$\tau_i = \sigma_i(\lambda) N_{\text{los}}(i),$$  \hspace{1cm} (3)

where $N_{\text{los}}(i)$ is the column density from the star for the molecule $i$ ($i = \text{H}_2\text{O}$ or OH). The calculations of $N_{\text{los}}(i)$ are non-trivial because they depend on the molecular abundances throughout the disk, which themselves depend on column density through Equation (2) and the photochemical reactions R1 and R2. In our previous X-ray models, this complication was unimportant because the X-ray intensity is nearly independent of the chemical composition. Indeed, the physical properties of the disk in X-ray models of a flared disk are largely a function of the vertical column density $N_{\perp}^*$, with $N_{\text{los}}/N_{\perp}^* \approx 10$. For a steady-state calculation in the absence of photodissociation (AGM11; NAG11), the molecular abundances of water and OH in the warm atmosphere are well approximated using $N_{\text{los}}/N_{\perp}^* \approx 30$. We use the same relationship in the calculations reported here, and find that the results are insensitive to changes in this ratio by factors of as much as three, which justifies using this approximation in place of a time-consuming full-disk analysis of the disk chemistry that includes FUV attenuation.

In the absence of detailed information on the dust in the inner regions of protoplanetary disks, we use the dust model in Appendix A of GNI04, characterized by the dust to gas mass density ratio $\rho_d/\rho_g$, and the geometric mean grain size $a_g$. This model is based on the distribution of Mathis et al. (1977) with minimum and maximum dust sizes, $a_1$, and $a_2$, and $a_g = \sqrt{a_1 a_2}$. The resulting grain surface area per H nucleus is

$$S_d = \frac{n_d(\pi a^2)}{n_H} = \frac{3}{4} \frac{\rho_d/\rho_g}{(\bar{\rho}/1.35m_H)a_g},$$  \hspace{1cm} (4)

where $\bar{\rho} \approx 3 \text{ g cm}^{-3}$ is the mean internal mass density of a grain, $n_d$ is the volumetric density of hydrogen nuclei, $n_d$ is the volumetric density of dust particles, and $1.35 m_H$ is the mass per H nucleus. With these parameters and $\rho_d/\rho_g = 0.01$ and $a_g = 0.707 \text{ m}, S_d \approx 8 \times 10^{-3} \text{ cm}^2$. By choosing $a_g$ larger than the interstellar value of $a_g = 0.035 \text{ m},$ the model takes into account that large grains settle to the mid-plane and a reduced population of small grains remains in the upper atmosphere. The values used here and in past publications, $\rho_d/\rho_g = 0.01$ and $a_g = 0.707 \text{ m}$ or $7.07 \text{ m},$ correspond to reductions in the grain surface area of roughly 20 and 200 relative to the interstellar medium (ISM). We use the temperatures from the dust-gas mixture in D’Alessio et al. (1999) as the dust temperature in all of our calculations.

When we express the optical properties of protoplanetary dust in terms of standard absorption, scattering and extinction coefficients, $Q_{\text{abs}}, Q_{\text{sc}}$ and $Q_{\text{ext}} = Q_{\text{abs}} + Q_{\text{sc}}$, the extinction cross section per H nucleus is $\sigma_{\text{ext}} = \sigma_{\text{ext}} S_d$ and the dust extinction associated with a hydrogen column $N_H$ is

$$\tau_d = \sigma_{\text{ext}} N_H = Q_{\text{ext}} N_H S_d.$$  \hspace{1cm} (5)

The grain surface area per H nucleus also plays a central role in the formation of H$_2$ and in the thermal accommodation of gas and dust, as discussed in the next section.

### 2.2. Heating and Cooling

We revise and extend the treatment of the heating and cooling of protoplanetary disk atmospheres in GNI04, especially the heating mechanisms. We emphasize the heating mechanisms because the underlying physics of these processes is generally less well founded than the cooling. These include the heating associated with FUV radiation and the formation of molecular hydrogen on grains. Figure 2 illustrates the roles of the most important heating and cooling processes for our reference model (Table 1) at a radial distance of 0.95 AU. Dust-gas cooling is the most important coolant, whereas accretion heating dominates the heating. Other heating processes such as photodissociation and grain formation play roles over limited ranges of column density. Water cooling has been included by adapting the work of Neufeld & Kaufman (1993). Other cooling mechanisms in the model (e.g., Oi fine-structure lines and recombination lines) have rates that fall below the plotted range in Figure 2.

#### 2.2.1. Accretion Heating

Viscous accretion heating is the most important heating process in our model. Without it, the surface layers would not be warm enough to take advantage of the warm neutral chemistry that is at the heart of our thermal–chemical model of the inner disk atmosphere. We continue to use the formulation in Equation (12) of GNI04,

$$\Gamma_{\text{acc}} = \frac{9 \alpha_h \rho c^2}{4} \Omega^2,$$ \hspace{1cm} (6)

where $\Omega$ is the angular rotation speed and where, to a good approximation, the local mass density of the gas, $\rho \approx 1 + 4x_{\text{He}}m_H$, includes He but no heavier atoms. The helium abundance, $x_{\text{He}}$, as with all abundances in this paper, is defined relative to total hydrogen. The isothermal sound speed for gas temperature $T_g$ is $c^2 = kT_g/m$, with the mean particle mass given by

$$m = \frac{1 + 4x_{\text{He}}}{1 - x_{\text{He}} + x_{\text{He}} + x_e} m_H.$$ \hspace{1cm} (7)

This quantity varies substantially in the transition region where hydrogen changes from atomic to molecular form in the disk atmosphere. With these definitions, Equation (6) can be written in a numerically useful form, with units of erg cm$^{-3}$ s$^{-1}$, as

$$\Gamma_{\text{acc}} = 6.18 \times 10^{-23} \alpha_h (1 - x_{\text{He}} + x_{\text{He}} + x_e) \times n_H T_g^2 \left( \frac{M}{M_\odot} \right)^{1/2} \left( \frac{\text{AU}}{r} \right)^{3/2}.$$ \hspace{1cm} (8)

Equation (6) is based on the hydrodynamic theory of a viscous accretion disk, where the viscosity $\nu$ is expressed as $\alpha_h \rho c^2/\Omega$. 

![Table 1: Reference Model Parameters](image-url)
protoplanetary disks (Glassgold et al. 2012). We calculate the substantially revised, with the result that the ionization rate according to AGM11. The X-ray luminosity, $L_{\text{X}}$ (Shakura & Sunyaev 1973). GNI04 regarded this as a phenomenological formula with $\alpha_h$ being a parameter that could vary with position and also differ from the more familiar “alpha” parameter that describes the average accretion rate. Some support for this idea comes from simulations of the magnetorotational instability (MRI; e.g., Hirose & Turner (2011)), but a more complete understanding of dissipation in the MRI would help put disk accretion heating on a sounder footing. For the reference model in this paper, we take $\alpha_h = 0.5$.

2.2.2. X-Ray Heating

The heating associated with X-ray irradiation has essentially the same form as in GNI04,

$$\Gamma_X = Q \zeta n_H,$$

where $Q$ is the heating per ion pair produced by X-rays and $\zeta$ is the X-ray ionization rate per H nucleus. Our treatment of the processes described in Section 2.2 of GNI04 has been substantially revised, with the result that $Q \approx 18 \text{ eV}$ in protoplanetary disks (Glassgold et al. 2012). We calculate the ionization rate according to AGM11. The X-ray luminosity, $L_X = 2 \times 10^{30} \text{ erg s}^{-1}$ is the same as that adopted by GNI04 (Section 2.2), who chose the average YSO X-ray luminosity found with Chandra for stellar masses in the 0.8–1.2 $M_\odot$ range for the Orion Nebula Cloud (Garmire et al. 2000). Our choice of $L_X$ is also close to the median value of the pre-main sequence sample studied by Güdel et al. (2010).

2.2.3. Photodissociation Heating

Photodissociation of H$_2$O and OH produces fast H atoms that can heat the gas, as noted by BB09. The kinetic energy of the heavy products OH and O can also contribute, but this is a relatively small effect ($\sim 5\%$). The photodissociation products can also be left in excited states. At sufficiently high densities, realized in some parts of protoplanetary disks, they may be collisionally de-excited and lead to heating. In the case of H$_2$O, photodissociation leads to highly excited levels of the ground electronic state of OH. The fluorescence of these levels has been detected in protoplanetary disks in the mid-infrared with Spitzer and ascribed to photodissociation (e.g., Najita et al. 2010; Carr & Najita 2011). Calculations of the heating in H$_2$O photodissociation (Crovisier 1989; Rodgers & Charnley 2005) are consistent with roughly equal amounts of direct heating and fluorescence.

We estimate photodissociation heating by first calculating the difference between the photon and dissociation energies,

$$E_{\text{phdiss}} = (h\nu - D_{\text{diss}}),$$

for each molecule and each band. The dissociation energies $D_{\text{diss}}$ are 5.13 eV for H$_2$O and 4.41 eV for OH. After averaging over the FUV bands, $\langle E_{\text{phdiss}}(\text{H}_2\text{O}) \rangle = 3.52 \text{ eV}$ and $\langle E_{\text{phdiss}}(\text{OH}) \rangle = 5.54 \text{ eV}$. These quantities enter the thermal balance through the heating rates per unit volume for molecule $i$ (H$_2$O or OH),

$$\Gamma_{\text{phdiss}}(i) = G(i) n(i) \eta(i) \left\langle E_{\text{phdiss}}(i) \right\rangle,$$

where $\eta(i)$ is the fraction of $E_{\text{phdiss}}$ going into heat. For H$_2$O this fraction is about 0.5 ($\sim 1.8 \text{ eV}$). A similar fraction might well apply to the photodissociation OH because the outcomes suggested by the cross sections calculated by (van Dishoeck & Dalgarno 1984) indicate the production of significant fluorescence (e.g., the 6300 Å lines from the $^1D_2$ level of atomic oxygen). Photodissociation heating of H$_2$O and OH is weak unless one or both of these molecules is near maximum abundance, $\sim 10^{-4}$. For example, photodissociation heating becomes dominant in the narrow transition region from log $N_H = 21.1$ to 21.2 in Figure 2.

2.2.4. Photoelectric Heating

Another component of the heating from FUV radiation arises from the photoelectric effect on grains and polycyclic aromatic hydrocarbons (PAHs). The underlying physics is complicated because it depends on the poorly known properties of the grains, such as the size and charge distributions. In addition
to the grain surface area and the FUV flux, some of the other relevant properties include the absorption coefficient, \( Q_{abs} \); the probability for ejecting an electron, \( Y \); and the energy of the liberated photoelectron, \( E_{phel} \). All of these quantities depend on the incident photon energy. In view of the uncertainties in the physical and chemical properties of the grains and small particles, we adopt average values for most of the quantities that enter into the heating rate per unit volume,

\[
\Gamma_{phel} = Y E_{phel} Q_{abs} S_d n_H F_{FUV}(\lambda), \tag{12}
\]

and use Equation (4) for \( S_d \) and set \( Q_{abs} = 1 \). For the other grain properties we rely on the theory of Weingartner & Draine (2001) developed for the ISM to estimate the first two factors in Equation (12). The effective band for the photoelectric effect extends from \( \sim 8 \) eV to \( 13.6 \) eV (911.7–1550 Å), and the typical threshold (work function) is \( \sim 5.5 \) eV. On this basis, the mean photoelectron energy is \( E_{phel} = 5.3 \) eV. Weingartner & Draine (2001) give the optical properties of single silicate or carbonaceous grains as a function of photon energy and grain size. We estimate the efficiency to be \( Y = 0.05 \) by roughly interpolating between a grain size of 0.03 \( \mu \)m and bulk material in Figure 5 of Weingartner & Draine (2001). The efficiency of grain photoelectric heating will be reduced relative to the ISM with the reduction in the grain surface area per \( H \) nucleus \( S_d \).3

2.2.5. \( H_2 \) Formation Heating

Ever since the suggestion by Spitzer & Cochran (1973) that newly formed \( H_2 \) molecules might carry a significant fraction of the dissociation energy (4.48 eV), the amount of the resulting gas heating due to formation on grains has remained uncertain. The basic issue is how much energy goes into kinetic and internal excitation energy of the \( H_2 \) molecule in addition to gas heating due to formation on grains has remained uncertain. In view of the uncertainties in the physical and chemical properties of the grains and small particles, we adopt average values for most of the quantities that enter into the heating rate per unit volume (with units of \( \text{erg cm}^{-3} \text{s}^{-1} \)) as,

\[
\Gamma_{H_2, \text{form}} = 6.91 \times 10^{-31} T_g^{1/2} n(H) n_H \\
\times \epsilon S \left( \frac{\rho_d/\rho_g}{0.01} \right) \left( \frac{0.05 \mu \text{m}}{a_g} \right) \left( \frac{\Delta E}{eV} \right). \tag{14}
\]

In practice we follow GMN09 and use a revised estimate of curves 2 and 3 in the left panel of Figure 1 of Cazaux & Tielens (2010) for the formation efficiency \( \epsilon \) as a function of \( T_d \). We approximate \( \epsilon = 1 \) for \( T_d < 25 \) K, \( \epsilon = 0.6 \) for \( 25 < T_d < 80 \) K, \( \epsilon = 0.33 \) for \( 80 < T_d < 900 \) K, and \( \epsilon = 0 \) for \( T_d > 900 \) K. We estimate \( S = 0.2 \) for silicate grains from Tielens (1983).

2.2.6. Thermal Accommodation

When gas collides with dust particles, energy is exchanged in a classical kinetic theory process known as thermal accommodation. In principle the rate of energy exchange depends on the nature of the gas and the dust surface, on the gas and dust temperatures, and possibly on the dust particle size. We use the formulation in GN104, rewriting their Equation (6) for the rate of energy exchange between gas and dust as,

\[
\Lambda_{dg} = 4.76 \times 10^{-33} A_H \sqrt{T_g(T_g - T_d)} n_H^2 \\
\times \left( \frac{\rho_d/\rho_g}{0.01} \right) \left( \frac{0.05 \mu \text{m}}{a_g} \right). \tag{15}
\]

Temperatures are in K and densities in \( \text{cm}^{-3} \) so that the units are \( \text{erg cm}^{-3} \text{s}^{-1} \). \( \Lambda_{dg} \) is defined so that it is a cooling function for \( T_g > T_d \), which is usually the case in the present calculations. The strength of the dust-gas cooling is determined by a mean accommodation coefficient \( A_H \), which takes the form, omitting the small contributions of He and heavy atoms,

\[
A_H = x(H)A(H) + \frac{1}{\sqrt{2}} x(H_2)A(H_2). \tag{16}
\]

The treatment given here of dust-gas cooling is based on the widely cited work of Burke & Hollenbach (1983). These authors used a rather simple semi-classical theory to fit experiments on tungsten. They then calculated accommodation coefficients for \( H \) and \( H_2 \) on water ice, silicates, and graphite. We fit Figure 4 of Burke & Hollenbach (1983) for the individual coefficients, and in the present application assume that the dust is pure silicate. It would be very helpful to have laboratory measurements of the accommodation coefficients for astrophysical dust analogs for temperatures relevant to the inner regions of protoplanetary disks, \( T_d \approx 100 \) K and \( T_g \approx 300–1000 \) K.

3. RESULTS

The results presented here for the FUV and X-ray model of the previous section are obtained by integrating the time-dependent heat equation for the gas temperature \( T_g \) and the chemical rate equations for 117 species, using the D’Alessio et al. (1999) flared disk model, which specifies the disk structure in terms of \( n_H \) and \( T_g \). The model is divided into cells located by their radial and vertical coordinates, \( r \) and \( z \). At each radius \( r \), we begin at the “top” of the atmosphere, specified by the vertical column density log \( N_H = 18 \), where we assume atomic
beginning to dominate Lyα cooling with increasing density and to balance accretion heating. Concomitant with the formation of molecules, CO cooling rapidly forces the temperature down close to $\sim 1000$ K. The transition region includes a column of $\sim 10^{19}$ cm$^{-2}$ of H$_2$ in the temperature range $T_g = 1000$–$4000$ K in the inner disk. These conditions are similar to those inferred from HST observations of H$_2$ fluorescence (Herczeg et al. 2004; Schindhelm et al. 2012; France et al. 2011, 2012).

The temperature of the transition region in Figure 3 falls slowly below 1000 K, reaching $T_g = 250$ K for a vertical column log $N_H = 21.9$. Thus the transition region in the X-ray and FUV irradiation model consists of roughly three layers of gas with different temperatures and chemical properties: (1) a warm molecular region that lies below (2) a warmer photodissociation region where H$_2$ and H$_2$O change from partial to full association, and (3) a hot atomic region at the top of the atmosphere. Regions (1) and (2) are distinguished in Figure 3 by the light orange and red shading, respectively. Throughout the disk the temperature of $\sim 650$ K roughly separates the region of warm H$_2$O from warmer OH. At the start of the transition, where the temperature is near 1000 K, OH is as abundant as H$_2$O. By contrast, below the transition region, the temperature drops below 250 K and the abundance of OH is typically 3–4 dex smaller than that of H$_2$O. Once the dust temperature falls below $\sim 125$ K, the freeze out of H$_2$O has to be taken into account. At the radius $r = 0.95$ AU in Figure 3, the freeze out would occur where log $N_H > 22.0$ if the freeze out temperature for H$_2$O is 125 K.

Further insight into the nature of the transition region is provided in Figure 4 where the effects of varying the X-ray and FUV luminosity are illustrated. The abundance curves at $r = 0.95$ AU are displayed for models with neither X-ray nor FUV radiation (dotted curves), X-rays in the absence of FUV (dashed curves), and variations of FUV that range from 0.1 to 10 times the reference FUV luminosity value, $L_{\text{FUV}} = 1.3 \times 10^{-2} L_\odot$ (the solid curves) at fixed X-ray luminosity. The atomic to molecular transition in the inner region of an FUV and X-ray irradiated protoplanetary disk occurs in steps over a layer of finite thickness. This is to be contrasted with abundances calculated in a model with neither FUV nor X-ray irradiation. Indeed, in the absence of X-ray and FUV irradiation, the abundances of H$_2$, H$_2$O, and CO all rise up sharply at log $N_H = 20.9$ and contain essentially all the available hydrogen, oxygen, and carbon. Inclusion of X-rays without FUV shifts the transition down to log $N_H = 21.1$. However, the atomic to molecular transition still occurs in a very thin layer. When both X-rays and FUV are included, the molecular abundances in Figure 4 rise rapidly at log $N_H = 21.1$, close to the pure X-ray case, but not to their maximum values. Beyond this depth, the H$_2$O abundance continues to increase, controlled by photodissociation mediated by the absorption of FUV by dust and then by self-shielding. Maximum abundances of H$_2$ and H$_2$O are achieved at log $N_H = 21.2$ in the reference model. One result of larger FUV is to increase the amount of OH and decrease the amount of H$_2$O in the warm photodissociation region. The gradual atomic to molecular transition is a special characteristic of a protoplanetary disk that is irradiated by both X-rays and FUV.

3.2. The Chemistry of Water and OH

The chemistry is governed by the familiar temperature-sensitive radical reactions used by GNI04 and in subsequent papers (e.g., GMN09 and NAG11). The formation of OH plays
a key role, starting with the endothermic reaction,

\[ \text{O} + \text{H}_2 \rightarrow \text{H} + \text{OH}; \]  \hspace{1cm} (R3)

the rate coefficient recommended by Baulch (2005) is

\[ k_3 = 8.5 \times 10^{-20} T^{2.67} e^{-3163/T} \text{ cm}^3 \text{s}^{-1}. \] (17)

Then water is made by an exothermic reaction with a modest barrier,

\[ \text{OH} + \text{H}_2 \rightarrow \text{H}_2\text{O} + \text{H}; \]  \hspace{1cm} (R4)

its rate coefficient is

\[ k_4 = 8.5 \times 1.70 \times 10^{-16} T^{1.60} e^{-1660/T} \text{ cm}^3 \text{s}^{-1}. \] (18)

These reactions are effective for intermediate temperatures ranging from \( \sim 250-1000 \text{ K} \). Below 250 K, the barrier in

Equation (17) suppresses formation, whereas above 1000 K the barrier to the inverse reaction in R3 can be overcome. For example, in the reference model at the bottom of the hot layer just below \( \log N_{\text{H}} = 21.0 \), the balance of forward and backward reactions in R3 and R4 leads to OH abundances in the range \( 10^{-8}-10^{-4} \), and to much smaller but rapidly growing \( \text{H}_2\text{O} \) abundances. Once the temperature drops to \( \sim 1000 \text{ K} \), the inverse reactions are significantly reduced and the dominant destruction pathway for OH and \( \text{H}_2\text{O} \) becomes FUV photodissociation.

The water abundance increases with increasing depth and density until its maximum value is reached just after \( \log N_{\text{H}} = 21.2 \). At this point, consumption of H by the above reactions is reduced, \( \text{H}_2 \) reaches its maximum abundance, and OH decreases as almost all oxygen is incorporated into \( \text{H}_2\text{O} \). In the layer where the \( \text{H}_2\text{O} \) abundance increases from \( 10^{-6} \) to \( 2 \times 10^{-4} \), FUV photodissociation is the dominant, although not the only, destruction mechanism.

The increase in \( \text{H}_2\text{O} \) is moderated by the absorption of FUV, by dust and by \( \text{H}_2\text{O} \) itself, which is the self-shielding process discussed in BB09. In the FUV dissociation region of the reference model in Figure 3, \( \tau_{\text{FUV}} > \tau_{\text{H}_2\text{O}} + \tau_{\text{OH}} \) midway to where the water abundance peaks (\( \log N_{\text{H}} \approx 21.1 \)), at which point the total FUV–optical depth is large. Once \( \text{H}_2\text{O} \) reaches its maximum abundance, destruction by reactions with X-ray generated molecular ions becomes more important than photodissociation, although one of the major ions, \( \text{H}_3^+ \), is partially recycled to \( \text{H}_2\text{O} \) by dissociative recombination. Going deeper down into the atmosphere, where \( \log N_{\text{H}} > 21.8 \) at \( r = 0.95 \text{ AU} \), the temperature continues to drop and OH is no longer rapidly converted into \( \text{H}_2\text{O} \) according to the rate coefficient in Equation (18), and instead fast exothermic reactions with atomic oxygen lead to high abundances of \( \text{O}_2 \).

The abundance of OH is also reduced, so that most of the oxygen is in the form of O and \( \text{O}_2 \). The transition from \( \text{H}_2\text{O} \) to \( \text{O}_2 \) and O occurs for all radii larger than 0.5 AU.

### 3.3. The Distribution of Water and OH

Figure 5 shows how the abundances of \( \text{H}_2\text{O} \) and OH in the disk atmosphere change with radius and grain surface area, with the reference case (\( \mu_g = 0.707 \mu m \)) on the left and the case where the grain surface area is reduced by another factor of 10 (\( \mu_g = 7.07 \mu m \)) on the right. Results are shown only for small radii, \( r \approx 2 \text{ AU} \), where freeze out does not occur until below the regions of maximum gaseous abundance of \( \text{H}_2\text{O} \) and OH. The photodissociation rates for \( \text{H}_2\text{O} \) are plotted as dotted curves. The unshielded rate at \( r = 0.95 \text{ AU} \) is \( G = 10^{-2} \text{s}^{-1} \), which is \( 10^7 \) times larger than the value for water in the ISM. The effects of self-shielding are evidenced by the sudden drop in \( G \) near where the water abundance peaks. These curves for \( G \) apply over the entire range of vertical columns in Figure 5, even where water is subject to freeze out. In such regions, water vapor may still be produced by the warm chemistry in reactions R3 and R4 because the progenitor species (\( \text{H}_2 \), O and OH) do not freeze out until reaching much lower temperatures. In order for water to freeze out, adsorption onto grain surfaces has to successfully compete with photodissociation.

In the reference case on the left side of Figure 5, the thickness of the warm (shaded orange) region decreases with increasing radius, while the peak abundances remain high. The thickness of the warm region mainly effects \( \text{H}_2\text{O} \) because its maximum abundances are achieved at greater depths than for OH. Beyond \( \sim 0.5 \text{ AU} \), the \( \text{H}_2\text{O} \) abundance always decreases when the temperature falls below \( \sim 300 \text{ K} \). The variation of the

![Figure 4. Vertical profiles of molecular abundances for H2 (black), H2O (blue), OH (green), and CO (magenta), and gas temperatures (red) in the disk atmosphere at \( r = 0.95 \text{ AU} \). Dotted curves are abundances without irradiation. Dashed curves are abundances under X-ray irradiation alone. Solid lines correspond to \( L_{\text{XUV}} = 0.0013, 0.013, \) and 0.13 \( L_{\odot} \), from faint to dark, respectively. \( L_X = 5.2 \times 10^{-2} L_{\odot} \) in each model with X-rays. For H2 and H2O the 0.0013 \( L_{\odot} \) case essentially overlaps X-ray irradiation alone. The red shaded region is where \( T_g \) is in the range 250–650 K.](image)
Figure 5. Vertical distributions of abundances for H$_2$ (black), H$_2$O (blue), and OH (green) for the reference model (left) and the reduced grain area case (right) at several radii (i.e., with $L_X = 5.2 \times 10^{-4} L_\odot$ and $L_{FUV} = 1.3 \times 10^{-2} L_\odot$). $T_g$ is plotted (red) with units along the right axis. The red shaded region is where $T_g$ is in the range 650–1000 K, and the orange shaded region is where $T_g$ is in the range 250–650 K. The line of sight FUV optical depth of unity is indicated by the vertical (dashed black) lines. Curves for the dissociation rate of water, $G$, are shown (dotted black).

(A color version of this figure is available in the online journal.)
H$_2$O and OH abundances with radius shown in Figure 5 is another manifestation of the role of temperature in the warm radical chemistry.

The panels on the right side of Figure 5 lack the red shaded regions, where $T \approx 650$–1000 K, illustrating that the amounts of warmer OH and H$_2$O are significantly reduced by decreasing the grain surface area. When the grain area is reduced, the gas is not thermalized as efficiently and the atmosphere remains hot to larger $N_d$, causing the molecular transition to occur deeper, where densities are larger and the dust is colder. At $r = 2$ AU, near the top of the transition, the small grain area model has larger warm abundances of OH than H$_2$O. This is a direct effect of the role of dust in providing less attenuation of the FUV, as indicated by the plots of the photodissociation rates. The dashed vertical lines in Figure 5 indicate the location where the FUV dust optical depth, from Equation (5), is unity. Increasing the dust grain size reduces the dust surface area $S_d$ according to Equation (4) and lowers the FUV opacity. The value $t_d = 1$ occurs near log $N_{HI} = 20.8$ at $r = 2$ AU in the reference model, before water self-shielding becomes important. In the case of the reduced grain surface area at $r = 2$ AU, the FUV penetrates much deeper into the disk, beyond the transition region, and the large photodissociation rates throughout reduce the abundance of H$_2$O in the warm region, producing OH, and fully self-shielding at log $N_{HI} = 21.8$.

In order to compare with observations, we first calculate the vertical column densities of warm H$_2$O and OH as a function of radius and then integrate them to obtain the cumulative number of molecules within a given radius. The vertical column densities are given in Table 2 for the two temperature regimes, 250–650 K and 650–1000 K, which correspond roughly to where H$_2$O and OH, respectively, are most abundant. Results from both the reference grain area ($a_g = 0.707 \mu$m) and the reduced grain area ($a_g = 7.07 \mu$m) cases are tabulated. To be conservative, freeze out of H$_2$O onto grains has been taken into account by excluding gaseous water from regions where the dust temperature is less than 125 K.

The freeze out temperatures for oxygen-bearing species O, O$_2$, OH, and CO are all much lower and these species stay in the gas phase in the inner disk. Since OH is most abundant in warm regions where H$_2$O remains in the gas phase, freeze out in cold regions does not significantly impact our calculation of OH columns. Where H$_2$O freezes out, the back-reaction in R4 is slow, hydrogen is entirely molecular, and the OH chemistry is dominated by reactions with molecular ions and O, rather than the photodissociation of H$_2$O. In the reference case for H$_2$O, the largest columns occur within 0.5 AU for the temperature range 250–650 K. The relatively slow decrease for larger radii in the reference case is compensated by the increase of annular area with radius. The total numbers of warm H$_2$O and OH enclosed within a given radius are shown in Figure 6. The case of reduced grain area is distinguished from the reference case by the use of dashed lines. The total amount of warm H$_2$O levels off beyond ~4 AU due to freeze out. The total number of warm water molecules in the reference case is ~$4 \times 10^{35}$ or $1.22 \times 10^{23}$ g, which is about 9% of the mass of Earth’s oceans.

### 4. Discussion

The atmosphere of a protoplanetary disk that includes both FUV and X-ray irradiation consists of an upper hot region with $T_g \approx 5000$ K, a cool lower region with temperatures $T_g < 250$ K, and a warm layer with temperatures 250 $< T_g < 1000$ K in between. The transition from atoms to molecules occurs mainly in the warm layer, where large amounts of water are produced. For our reference model, gaseous water would be present in the region where the grain temperature is above the freeze out temperature (~125K; i.e., within a radial distance of 4 AU). However, processes that we did not consider here, such as photodesorption and gas-grain chemistry, could well maintain a gaseous water reservoir over a larger range of radii.

We confirm several of the ideas presented by BB09; for example, molecular shielding by H$_2$O and OH is important in

![Table 2: Column Densities of H$_2$O and OH](https://example.com/table2)

![Figure 6. Number of warm molecules within a radius, r, with $T_g$ in the range 250–1000 K, and where $T_g > 125$ K (i.e., above the freeze out temperature) are shown in black for both the reference model ($a_g = 0.707 \mu$m, solid lines), and for reduced grain area ($a_g = 7.07 \mu$m, dashed lines). Particularly warm molecules with $T_g$ in the range 450–1000 K, are shown in red. (A color version of this figure is available in the online journal.)](https://example.com/figure6)
accounting for the large amounts of warm \( H_2O \) and OH that are found in \textit{Spitzer} observations of FUV-irradiated CTTS disks. In addition, we also find that when significant grain settling has occurred, UV absorption by \( H_2O \) and OH plays an important role in the thermal and chemical properties of the disk atmosphere by shielding the disk mid-plane from UV radiation.

In confirming these ideas, we have made several important improvements in the implementation of molecular shielding. One example is the treatment of molecular hydrogen, which plays a key role in the formation of OH and \( H_2O \). BB09 assumed that all hydrogen was in \( H_2 \) in the atmosphere without considering the chemistry of \( H_2 \), especially the role of grains. They assumed that molecules dominated the FUV opacity due to dust depletion, but did not examine whether depletion was consistent with the abundance of \( H_2 \). We find that molecular shielding is effective even in the presence of significant grain settling (i.e., at the level of grain settling seen in T Tauri stars).

When we examined the model results in Section 3 to see how FUV radiation changes the pure X-ray model, we found that it is the X-rays that still play a role at the start of the transition. However, the rise up to maximum molecular abundances is achieved more gradually under the influence of FUV photodissociation of \( H_2O \) and OH. Once the FUV radiation has been largely extinguished, the X-rays again play a significant role by reshuffling oxygen between the molecular ion \( H_3O^+ \) and O, OH, and \( H_2O \). In this way the abundances accounting for the large amounts of warm \( H_2O \) and OH that are found in different places in the disk atmosphere: the OH is higher and hotter than the \( H_2O \). Looking down into the disk atmosphere, a layer of the OH appears first, and below it are layers of warm and cool \( H_2O \). A similar distinction in the OH and \( H_2O \) temperatures may be present in \textit{Spitzer} observations of CTTS disks. Carr & Najita (2011) analyzed \textit{Spitzer} observations of the disks of six CTTSs in Taurus-Aurigae, and obtained temperatures of \( \sim 600 \) K for \( H_2O \). Working with \textit{Spitzer} observations of a larger and different sample, Salyk et al. (2011) found \( H_2O \) temperatures clustering around \( \sim 450 \) K and significantly higher OH temperatures, with an average value of 1050 K for 15 detections.

While similar amounts of \( H_2O \) are produced in the large and small grain area models, greater differences are seen in the OH that is produced in the two models. As shown in Figure 6 (lower panel), the large grain area model (solid line) produces much more OH than then small grain area model (dashed line). The enclosed number of warm OH molecules in the large grain area model is similar to that observed by \textit{Spitzer} (\( \sim 10^{11} \)). In either model, the OH arises from a larger range of radii than the \( H_2O \). Taken literally, the OH and \( H_2O \) line profiles would differ, with the OH profiles more centrally peaked than the \( H_2O \).

Our model does not treat the Ly\( \alpha \) line, which can be a significant fraction of the incident FUV. For example, Ly\( \alpha \) accounts for \( \sim 80\% \) of the observed total FUV for TW Hya (BB09; Yang et al. 2012). The central portion of the observed line is absorbed out, and it appears as two broad features separated by several \( \AA \). Following Bethell & Bergin (2011), the hot upper atmosphere of the disk is a region of diffuse Ly\( \alpha \) due to multiple scattering of the line by atomic H. In our reference model discussed in Section 3, atomic H is the dominant form of hydrogen almost down to vertical columns of log \( N_H = 21.2 \). Thus the diffuse Ly\( \alpha \) extends throughout the upper part of the warm atmosphere to the level where OH has begun to decrease from its maximum abundance and \( H_2O \) has achieved its maximum.

Bethell & Bergin (2011) used a Monte Carlo code to calculate the transfer of Ly\( \alpha \) for a D’Alessio disk model that includes a simplified chemistry for the \( H_2 \) molecule and three levels of dust depletion. If the dust surface area is reduced by the value 20, as in our reference model, it can be deduced from their Figure 10 that the initial ratio of 6 for the Ly\( \alpha \) to the FUV continuum number flux is reduced to \( \sim 1/3 \) near log \( N_H = 21 \). On this basis, the Ly\( \alpha \) line would increase the dissociation rate of water beyond that calculated in this paper by \( \sim 1/3 \). The increased photodissociation could be approximated by increasing \( L_{FUV} \) in our model, with the results illustrated graphically in Figure 4.

The role of the absorption of Ly\( \alpha \) by \( H_2O \) ignored by Bethell & Bergin (2011), is also important. Using the cross section for Ly\( \alpha \) absorption by \( H_2O \) \( (1.6 \times 10^{-17} \text{ cm}^2) \), Mota et al. (2005), the vertical optical depth through log \( N_H = 21.0 \) is \( \sim 3.2 \), ignoring the fact that the Ly\( \alpha \) photons are not all traveling downward. This reduces the effect of the Ly\( \alpha \) line by a further factor of 25 beyond the effects of dust included by Bethell & Bergin (2011). On this basis we do not expect the total amount of H\( \alpha \) estimated in this paper to be much affected by the omission of Ly\( \alpha \). The details of the transition region may change, as may the properties of the OH radical, which is located in a region with more diffuse Ly\( \alpha \) radiation. Thus it would be of interest to include the transfer of Ly\( \alpha \) in our model of a thermally and chemically inhomogeneous protoplanetary disk atmosphere.
It is interesting to compare our relatively simple model with the very detailed model, ProDiMo. Presented by Woitke et al. (2009a), this model was extended and applied in several subsequent papers, especially Meijerink et al. (2012), which includes X-ray as well as FUV irradiation. ProDiMo includes a very large number of thermal processes. For the purposes of producing the warm temperatures in the inner disk, ProDiMo relies on the photoionization of PAHs and collisional de-excitation of excited Fe II levels. Our main heating mechanism stems from mechanical heating that can be traced to the dissipation produced by the accretion process, most likely the MRI. There are uncertainties associated with these processes (e.g., the PAH abundance in disks and the magnitude of mechanical heating). In both models, dust-gas cooling, discussed in Section 2.2.6, is important.

Another difference between ProDiMo and our model is their use of a relatively large inner radius of 0.5 AU; essentially all the warm water predicted by Woitke et al. arises in the very inner rim of the disk rather than over the inner several AU as in our model. Low H2O abundances ($X_{H_2O} \lesssim 10^{-7}$) are found beyond the inner rim, in contrast with the much larger abundances in our model ($X_{H_2O} \approx 10^{-4}$). These differences imply that ProDiMo would have difficulty accounting for the large amounts of H2O that have been observed in inner CTTS disks if it adopted a smaller radius that is more typical of CTTS. A likely reason for this major difference between the results from ProDiMo and our model is that we have included UV shielding by OH and H2O which both absorb radiation over a wide range of FUV wavelengths, whereas ProDiMo includes UV shielding by CO and H2, which shield over a more limited range of wavelengths.

5. CONCLUSIONS

We have developed an integrated thermal–chemical model for protoplanetary disk atmospheres that includes irradiation by FUV and X-rays, grain settling, and a detailed treatment of the physical processes that affect water and other molecules. Our objective has been to identify some of the key processes and how they operate. We emphasized that several of the processes are uncertain, especially those relating to grains, and we have focused on a reference model that represents a typical CTTS disk.

Our results support the idea that H2O in the inner regions of protoplanetary disks can be formed in situ. This thesis has been discussed in several of the earlier modeling papers cited in Section 1. Especially noteworthy are those that obtain sufficient warm columns of H2O to be consistent with the amounts deduced from Spitzer observations, for example, Glassgold et al. (2009), Woods & Willacy (2009), Heinzelner et al. (2011), and Najita et al. (2011). In the present work, both the physical and astrophysical details of how adequate amounts of H2O can be formed are analyzed in detail. Among the new results presented here is the key role of dust surface area in regulating processes that directly affect the H2O abundance (i.e., H2 formation, dust-gas thermal accommodation, and the extinction of FUV radiation). All of these processes are sensitive to the size distribution of the grains in the upper atmosphere of the disk. We find that, in our updated model, large amounts of H2O and OH are synthesized in the observable warm regions of a typical protoplanetary disk atmosphere, even in the presence of FUV and X-ray radiation and substantial grain settling.

The H2O and OH are located primarily in distinct locations, with the warm H2O layer located below a warmer layer of OH. Both the temperatures and total numbers of molecules are consistent with Spitzer observations of CTTS, with the OH in the atmosphere extending over a larger range of radii than the H2O. In our model, freeze out on dust grains restricts the gaseous water reservoir in the warm atmosphere to radii < 4 AU.

UV absorption by H2O and OH is important in accounting for the large amounts of water that are observed, confirming the ideas discussed by BB09. FUV radiation affects the width and sharpness of the atomic to molecular transition, whereas X-rays and grain settling affect the depth at which the transition starts. X-rays also generate ions that destroy water below the photodissociation region. Many of these conclusions can be tested with suitable observations, as has already been done in some cases. Together with the discussion in previous sections, they present a richer picture of an atomic to molecular in protoplanetary disk atmospheres.

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