The $X_{\text{CO}}$ Conversion Factor from Galactic Multiphase ISM Simulations

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Abstract

CO($J=1\rightarrow0$) line emission is a widely used observational tracer of molecular gas, rendering essential the $X_{\text{CO}}$ factor, which is applied to convert CO luminosity to $H_2$ mass. We use numerical simulations to study how $X_{\text{CO}}$ depends on numerical resolution, non-steady-state chemistry, physical environment, and observational beam size. Our study employs 3D magnetohydrodynamics (MHD) simulations of galactic disks with solar neighborhood conditions, where star formation and the three-phase interstellar medium (ISM) are self-consistently regulated by gravity and stellar feedback. Synthetic CO maps are obtained by postprocessing the MHD simulations with chemistry and radiation transfer. We find that CO is only an approximate tracer of $H_2$. On parsec scales, $W_{\text{CO}}$ is more fundamentally a measure of mass-weighted volume density, rather than $H_2$ column density. Nevertheless, $\langle X_{\text{CO}} \rangle = (0.7\text{--}1.0) \times 10^{20} \text{cm}^{-2} \text{K}^{-1} \text{km}^{-1} \text{s}$, which is consistent with observations and insensitive to the evolutionary ISM state or radiation field strength if steady-state chemistry is assumed. Due to non-steady-state chemistry, younger molecular clouds have slightly lower $\langle X_{\text{CO}} \rangle$ and flatter profiles of $X_{\text{CO}}$ versus extinction than older ones. The CO-dark $H_2$ fraction is 26%--79%, anticorrelated with the average extinction. As the observational beam size increases from 1 to 100 pc, $\langle X_{\text{CO}} \rangle$ increases by a factor of $\sim2$. Under solar neighborhood conditions, $\langle X_{\text{CO}} \rangle$ in molecular clouds is converged at a numerical resolution of 2 pc. However, the total CO abundance and luminosity are not converged even at the numerical resolution of 1 pc. Our simulations successfully reproduce the observed variations of $X_{\text{CO}}$ on parsec scales, as well as the dependence of $X_{\text{CO}}$ on extinction and the CO excitation temperature.

Key words: galaxies: ISM – ISM: abundances – ISM: clouds – ISM: molecules – molecular processes

1. Introduction

Molecular clouds are the birthplaces of stars. In addition, molecular gas is the dominant component of the interstellar medium (ISM) in dense and shielded environments. Measuring the properties of molecular clouds is therefore critical to understanding the ISM and star formation in the Milky Way and beyond. However, the most abundant molecule in the ISM, molecular hydrogen $H_2$, is not directly observable in emission at typical ISM temperatures because of its low mass and lack of dipole moment. As a result, the second most abundant molecule, CO, is often used as an observational tracer for $H_2$. The standard technique employs a conversion factor $X_{\text{CO}}$ to relate the observed velocity-integrated intensity of CO($J=1\rightarrow0$) line emission $W_{\text{CO}}$ to the $H_2$ column density $N_{H_2}$,

$$N_{H_2} = X_{\text{CO}} W_{\text{CO}},$$

(1)

Although the CO($J=1\rightarrow0$) line emission is bright and easy to detect with ground-based radio telescopes, it is often very optically thick. Many observational studies have measured $X_{\text{CO}}$ by deriving the $H_2$ mass independently of CO emission, via dust emission or extinction, gamma-ray emission, or the virial theorem (e.g., Solomon et al. 1987; Strong & Mattox 1996; Dame et al. 2001; Lombardi et al. 2006). Surprisingly, the value of $X_{\text{CO}}$ only varies within a factor of $\sim2$ for many molecular clouds in the Milky Way and local disk galaxies. This has motivated the adoption of a constant standard $X_{\text{CO}}$ conversion factor in the literature, $X_{\text{CO},MW} = 2 \times 10^{20} \text{cm}^{-2} \text{K}^{-1} \text{km}^{-1} \text{s}$ (see review by Bolatto et al. 2013).

It is important to note that this standard $X_{\text{CO}}$ is an average value for nearby molecular clouds on scales of tens of parsecs. $X_{\text{CO}}$ is empirically known to vary both on small scales and for molecular clouds in different environments. One of the earliest studies of $X_{\text{CO}}$, by Solomon et al. (1987), suggested that $X_{\text{CO}}$ varies by a factor of a few for molecular clouds in the Milky Way, decreasing with increasing CO luminosity. Recent high-resolution observations have found that $X_{\text{CO}}$ can vary by more than an order of magnitude on parsec scales, although the averages of $X_{\text{CO}}$ over individual molecular clouds are within a factor of $\sim2$ of the standard Milky Way value (Pineda et al. 2008; Ripple et al. 2013; Lee et al. 2015; Imara 2015; Kong et al. 2015). Beyond nearby molecular clouds, $X_{\text{CO}}$ in the Galactic center is a factor of $\sim4$ lower than the mean value in the disk (Blitz et al. 1985; Ackermann et al. 2012), and similar results are found for the central regions in nearby spiral galaxies (Sandstrom et al. 2013). High surface density starburst regions have $X_{\text{CO}}$ significantly below $X_{\text{CO},MW}$ (e.g., Downes & Solomon 1998; Bolatto et al. 2013, and references therein). Observations also indicate that $X_{\text{CO}}$ can be much higher than the standard Milky Way value in low-metallicity galaxies (Israel 1997; Leroy et al. 2011).

Theoretical models and numerical simulations have provided insights into the $X_{\text{CO}}$ conversion factor. Wolfire et al. (1993) constructed spherical cloud models with a photodissociation region (PDR) code, and suggested that $X_{\text{CO}}$ is only weakly dependent on the incident far-ultraviolet (FUV) radiation field strength, and insensitive to the small variations in metallicity up to a reduction of metallicity by a factor of 5 relative to the solar neighborhood. These models rely on simple assumptions about cloud structure and kinematics. To model molecular clouds with more realistic structure, many numerical simulations have been carried out to study 3D turbulent molecular clouds with self-consistent, time-dependent chemistry and radiation transfer.
(e.g., Glover & Mac Low 2011; Shetty et al. 2011a, 2011b; Glover & Clark 2012; Szűcs et al. 2016). Shetty et al. (2011b) and Szűcs et al. (2016) found similar cloud-average \( X_{\text{CO}} \) to the standard observed value (with significant variations on smaller-than-cloud scales). Shetty et al. (2011b) concluded that \( X_{\text{CO}} \) has a weak dependence on gas density, temperature, and velocity, and that the nearly constant \( X_{\text{CO}} \) is the result of the limited range of physical properties found in nearby molecular clouds. However, these simulations consider molecular clouds to be isolated from the large-scale galactic ISM, and their key physical properties such as the average density and velocity dispersion are set artificially based on the initial conditions of the simulations and prescribed turbulent driving.

In recent years, more efforts have been made to investigate \( X_{\text{CO}} \) in global galaxy simulations (Narayanan et al. 2011, 2012; Feldmann et al. 2012; Duarte-Cabral et al. 2015). With resolutions of tens of parsecs, however, global galaxy simulations cannot resolve substructures in molecular clouds, and subgrid models are generally required to estimate the CO conversion factor in molecular clouds. Furthermore, as observations of galactic and extragalactic data, little comparison has been made regarding the variation of \( X_{\text{CO}} \) within molecular clouds on parsec or smaller scales. Furthermore, as observations of galactic and extragalactic molecular gas probe a range of scales, it is important to understand how \( X_{\text{CO}} \) may vary with the effective area of a radio beam.

In this paper, we present a new study of the \( X_{\text{CO}} \) conversion factor in MHD galactic disk simulations with solar neighborhood conditions and 1–4 pc resolutions. The high-density clouds are formed and destroyed self-consistently within the turbulent, multiphase, magnetized ISM by gravity and stellar feedback. In our models, the distribution of H\(_2\) and CO is obtained by postprocessing the MHD simulations with chemistry and radiation transfer. While ideally all dynamics and chemistry would form within the regions of the CNM where the gas is dense and shielded. We model the formation of molecular gas by postprocessing the \( X_{\text{CO}} \) simulations with chemistry and shielding, which is described in detail in Section 2.2.

We adopt the fiducial solar neighborhood model of KO2017. The simulation domain size is \( L_x = L_y = 1024 \text{ pc} \) and \( L_z = 4096 \text{ pc} \). The initial gas surface density \( \Sigma = 13 \ M_\odot \text{ pc}^{-2} \). The simulation reaches a quasi-steady state after \( t \approx 200 \text{ Myr} \). The total mass of the gas in the simulation slowly declines as the gas turns into stars or leaves the simulation domain as galactic winds. In this paper, we focus on the simulation during the time frame \( t = 350–420 \text{ Myr} \) when the surface density of the gas is in the range \( 9 \ M_\odot \text{ pc}^{-2} < \Sigma < 10 \ M_\odot \text{ pc}^{-2} \).

In order to study the effect of numerical resolution on \( X_{\text{CO}} \), we consider the simulation with three different resolutions: \( \Delta x = 4, 2, \) and 1 pc. The 4 pc simulation starts from \( t = 0 \) with the initial condition described in KO2017, and runs until...
$t = 700 \text{ Myr}$. To save computational time, we use an “extraction” method to refine the resolution. We use the output of the 4 pc simulation at time $t = 350 \text{ Myr}$ as the initial condition of the 2 pc simulation, and run that for 70 Myr (until $t = 420 \text{ Myr}$). Similarly, we use the output of the 2 pc simulation at $t = 378 \text{ Myr}$ as the initial condition of the 1 pc simulation, and run that for 4 Myr (until $t = 382 \text{ Myr}$). We also reduce the domain size in the $z$-direction to $L_z = 2240 \text{ pc}$ for the 2 pc simulation and to $L_z = 896 \text{ pc}$ for the 1 pc simulation. Because the scale height $H \sim 100 \text{ pc}$ for the CNM and $H \sim 400 \text{ pc}$ for the WNM, the simulation domain in the $z$-direction is big enough to capture most of the mass in the neutral and molecular ISM.

When refining from a coarser resolution, it takes some time for the turbulence to cascade down to smaller scales and create finer structures. The line-width size relation (e.g., Larson 1981; Solomon et al. 1987; Heyer & Brun 2004; Heyer & Dame 2015),

$$v(l) \sim 0.7 \text{ km s}^{-1} \left(\frac{l}{\text{pc}}\right)^{1/2},$$

(2)
gives the expected timescale for turbulent cascade in the dense ISM:

$$\tau_{\text{turb}}(l) = \frac{l}{v(l)} = 1.4 \text{ Myr} (l/\text{pc})^{1/2}.$$  

(3)

We only use the outputs from the 2 and 1 pc simulations 4–6 Myr after the extraction from coarser resolution, allowing sufficient time for the turbulence to develop at the refined resolution.

The density threshold for creation of sink particles, $\rho_{\text{thr}}$, also depends on the resolution of the simulation. A sink particle is created if the cell is at a local gravitational potential minimum, the flow is converging, and the density of the cell exceeds the Larson–Penston threshold (Larson 1969; Penston 1969) suggested by Gong & Ostriker (2013),

$$\rho_{\text{thr}} \equiv \rho_{\text{LP}}(\Delta x/2) = \frac{8.86}{\pi} \frac{c_s^2}{G\Delta x^2}.$$  

(4)
The typical density threshold at the equilibrium CNM temperature is $n_{\text{thr}} = 2956, 927, \text{ and } 304 \text{ cm}^{-3}$ for resolutions $\Delta x = 1, 2, \text{ and } 4 \text{ pc}$.\(^{5}\)

### 2.2. Postprocessing Chemistry

To model the chemical composition of the gas, we have developed a postprocessing module within the code Athena++ (White et al. 2016). This module reads the output from TIGRESS simulations and performs chemistry calculations assuming that the density and velocity in each grid cell are fixed. We use the simplified chemical network of Gong et al. (2017), which focuses on the hydrogen, carbon, and oxygen chemistry, and gives accurate abundances of H$_2$ and CO. We assume an initial chemical composition of neutral atomic gas, with all hydrogen in the form of H, all carbon in C, all oxygen in O, and all silicon in Si. The initial temperature is the same as the output from MHD simulations. Then we evolve the chemistry, temperature, and radiation field (see below) for time $t_{\text{chem}} = 50 \text{ Myr}$, so that the chemical abundances of the gas reach a steady state. In other words, we do not self-consistently calculate the time-dependent gas dynamics and chemistry, but instead consider the state in which the chemistry and temperature have reached an equilibrium, consistent with radiative heating and ISM structure as determined by the MHD simulations. Because gas cooling is not sensitive to the chemical composition, chemistry has minimal effect on the gas dynamics (Glover & Clark 2012; Gong et al. 2017). However, dust shielding can reduce the gas heating and lower the gas temperature by a factor of $\sim 2$ in shielded regions of the CNM where molecular gas forms.\(^{5}\) In return, gas dynamics can also influence the chemical composition. For example, the timescale for H$_2$ formation can be longer than the turbulent crossing time in the molecular clouds, which may lead to much lower H$_2$ abundance than the equilibrium values (Gong et al. 2017). The temporal dependence of the chemical state and observable CO properties are considered in Section 3.2.

The heating and cooling of the gas are calculated simultaneously with chemistry, with the details described in Gong et al. (2017). We slightly modify the parameter $N(\text{CO})$ for CO cooling in Gong et al. (2017) by setting

$$N(\text{CO}) = \frac{n(\text{CO})}{\max(\langle |dv/dr|\rangle, v_{\text{th}}/v_{\text{esc}})}.$$  

(5)

where $\langle |dv/dr|\rangle$ is the mean (absolute) velocity gradient across the six faces of each grid cell in the simulation, $v_{\text{th}} = \sqrt{2kT/m(\text{CO})}$ the thermal velocity of CO molecules, and $v_{\text{esc}} = 100 \text{ pc}$ the maximum length scale for a photon to escape. Using the maximum of the two terms in the denominator of Equation (5) ensures that there is a minimum probability for the photon to escape when the local velocity gradient is small, given a maximal molecular cloud size of $\lesssim 100 \text{ pc}$. This formalism is consistent with the large velocity gradient (LVG) and escape probability approximation we adopted in carrying out the synthetic observations of CO line emission (Section 2.3).

In order to compute the photoionization and photodissociation rates in the chemistry network, a radiation transfer scheme is needed to calculate the reduction of FUV radiation by dust and molecule shielding. We use the six-ray approximation (Nelson & Langer 1997, 1999; Glover & Mac Low 2007): in each cell, the radiation field is calculated by ray-tracing and averaged over six directions along the Cartesian axes. The incident radiation field is assumed to come from the edge of the computational domain along each ray, and has an initial intensity the same as that in the MHD simulations (the MHD simulations themselves do not include shielding). The main advantage of this approach is the low computational cost. When comparing to ray-tracing along many more different angles, the six-ray approximation gives reasonably accurate results (Safranek-Shrader et al. 2017). Because chemistry and radiation depend on each other, we iterate to solve the chemistry equations and six-ray radiation transfer.

### 2.3. Synthetic Observation of CO Line Emission

To model the CO($J = 1\rightarrow 0$) line emission, we apply the publicly available radiation transfer code RADMC-3D (Dullemond et al. 2012) with chemistry and temperature obtained as described

\(^{5}\) This typical reduction in temperature in high-density regions ($n_H \geq 10 \text{ cm}^{-3}$) is found by comparing the initial temperature output from the MHD simulation to the steady-state temperature from the postprocessing chemistry simulation.
in Section 2.2. We select the midplane region \(|z| < 256 \, \text{pc}\), where almost all molecules are found. H\(_2\) is assumed to be the only collisional partner with CO, and we use a fixed ortho-to-para ratio of 3:1. The synthetic observations are performed along the \(z\)-axis, i.e., the observer is looking at the galactic disk face-on. This avoids cloud blending, because all molecular clouds form near the midplane of the galactic disk.

The CO population levels are calculated by using the LVG and escape probability approximation, which is implemented in RADMC-3D by Shetty et al. (2011a). This approximation allows the population levels to be calculated locally in each cell. The escape probability is

\[
\beta = \frac{1 - e^{-\tau}}{\tau},
\]

and the optical depth \(\tau = \min(\tau_{\text{LVG}}, \tau_{\text{EscProb}})\). The LVG approximation gives

\[
\tau_{\text{LVG}} = \frac{\lambda_0^2}{8\pi} \frac{A_{10} n_{\text{CO}}}{\langle |dv/dr| \rangle} f_1 \left( \frac{f_0}{f_0} - 1 \right),
\]

where \(A_{10}\) is the Einstein A coefficient \(A_{10} = 7.203 \times 10^{-8} \, \text{s}^{-1}\), \(n_{\text{CO}}\) the number density of CO molecules, \(g_0 = 1\) and \(g_1 = 3\) the degeneracies for levels \(J = 0\) and \(J = 1\), \(f_0 = n_0/n_{\text{CO}}\) and \(f_1 = n_1/n_{\text{CO}}\) the fractions of CO molecules in levels \(J = 0\) and \(J = 1\), where \(n_0\) and \(n_1\) are the level populations, and \(\langle |dv/dr| \rangle\) is the same as that in Equation (5). The optical depth from the escape probability approximation is set by a typical length scale \(L_{\text{EscProb}}\), and can be written in the same form as Equation (7) by substituting \(\langle |dv/dr| \rangle\) with \(\sqrt{\pi} \nu_{\text{tot}}/L_{\text{EscProb}}\) (Draine 2011). Here \(\nu_{\text{tot}}\) is the total velocity dispersion (see below). We adopt \(L_{\text{EscProb}} = 100 \, \text{pc}\), consistent with the CO line cooling in Equation (5). In our simulation, the velocity gradient is usually relatively large, and in most cells \(\tau = \tau_{\text{LVG}}\).

Ray-tracing is performed after the CO level populations are obtained. In general, the emission line intensity is determined by radiative transfer (e.g., Draine 2011):

\[
dI = -I_\nu d\tau + S_\nu d\tau,
\]

where \(I_\nu\) is the line intensity at frequency \(\nu\), \(S_\nu\) the source function, and \(\tau_\nu\) the optical depth. \(\tau_\nu\) depends on the line profile, which is set by the velocity dispersion \(\nu_{\text{tot}} = \sqrt{\nu_{\text{mic}}^2 + \nu_{\text{turb}}^2}\). We include a subgrid “microturbulent” velocity dispersion according to the line-width size relation (Equation (2)),

\[
\nu_{\text{turb}} = 0.7 \, \text{km s}^{-1} \left( \frac{\Delta x}{\text{pc}} \right)^{1/2},
\]

where \(\Delta x\) is the resolution of the simulation. We also include a background blackbody radiation field with temperature \(T_{\text{CMB}} = 2.73 \, \text{K}\) from the cosmic microwave background (CMB).

We run RADMC-3D with a passband from \(-20\) to \(20 \, \text{km s}^{-1}\) (wide enough to include all CO emission) and velocity resolution of \(0.5 \, \text{km s}^{-1}\). RADMC-3D produces spectral position–position–velocity (PPV) cubes of the CO\((J = 1-0)\) line. We then interpolate \(I_\nu\) to a finer velocity resolution of \(0.07 \, \text{km s}^{-1}\), and calculate the total CO\((J = 1-0)\) line intensity in each observed pixel, \(W_{\text{CO}}\), by integrating \(I_\nu\) over all velocity channels that have emission above the detection limit, \(T_{\text{det}} = 0.4 \, \text{K}\). This approach matches the typical velocity resolution and sensitivity in observations of nearby molecular clouds (e.g., Ridge et al. 2006; Pineda et al. 2008, 2010; Ripple et al. 2013; Lee et al. 2014, see also Table 1). We define the “CO-bright” region as pixels with \(W_{\text{CO}} > 0.1 \, \text{K km s}^{-1}\), and calculate \(X_{\text{CO}}\) for each pixel in the CO-bright region. The average \(X_{\text{CO}}\), \(\langle X_{\text{CO}} \rangle = \sum N_{\text{H}_2}/\sum W_{\text{CO}}\), is also calculated only within the CO-bright region, similar to the common approach in observations (e.g., Pineda et al. 2008; Ripple et al. 2013). We define the fraction of CO-dark \(H_2\).

### 2.4. The Beam Size in Synthetic Observations

The default beam size in our synthetic observations is the same as the numerical resolution in the MHD simulations. In real observations, the beam size (in physical units) varies depending on the telescope and the distance of the object. The dust extinction or emission map used to derive \(H_2\) column densities typically has coarser resolution than the CO map. To analyze the \(X_{\text{CO}}\) values, the dust map and CO map are smoothed to a common resolution (usually the resolution of the dust map), which we refer to as the “beam size.”

The velocity resolution and sensitivity also vary in observations. We have compiled the observational parameters from the literature of \(X_{\text{CO}}\) observations in the Milky Way and nearby galaxies in Table 1. All the observations listed used \(H_2\) mass estimation from dust extinction or emission. We also list \((X_{\text{CO}})\) obtained by the observations when available.

We investigate the effect of beam size on \(X_{\text{CO}}\) in Section 3.4. The adopted parameters and beam sizes are listed in Table 2, and are designed to match the typical observational parameters listed in Table 1. The synthetic observations with default beam size are based on the original model data (the same as the numerical resolution). To create synthetic maps with larger effective beam, we first smooth out the PPV cubes produced by RADMC-3D to the desired CO map resolution. Then we match the corresponding velocity resolution from the default 0.5 km s\(^{-1}\) in the PPV cubes, by either interpolating to finer or integrating to coarser velocity resolution. We integrate over all velocity channels with emission above the detection limit \(T_{\text{det}}\) and obtain a 2D map of \(W_{\text{CO}}\) at the corresponding CO map resolution. Then both the map for \(A_V(N_{\text{H}_2})\) and the map for \(W_{\text{CO}}\) are smoothed to the common resolution of the beam size, for which \(X_{\text{CO}}\) is calculated. We note that the “beam” is square, not circular. The CO-bright region, for which \((X_{\text{CO}})\) is calculated, is defined as pixels with \(W_{\text{CO}} > 3T_{\text{det}} \Delta \nu\), where \(\Delta \nu\) is the width of the velocity channel.

Note that this is often called “pixel size” in observations. We use “beam size” to distinguish from the “pixel size” determined by the numerical resolution of our simulation and synthetic radiative transfer grid.

We have compared results for our square beam to the result for a circular Gaussian beam, and find that it makes very little difference for \((X_{\text{CO}})\).
3. Results

3.1. Convergence Study: Effects of Numerical Resolution

In this section, we investigate the effect of numerical resolution on both chemistry and $X_{\text{CO}}$. An overview of the models RES-4pc, RES-2pc, and RES-1pc is shown in Figure 1, and the overall properties of the models are listed in Table 4. As the resolution increases, more small structures and dense gas form in the simulations. The locations of molecular clouds are similar in all three models, but the small-scale filamentary structures within the molecular clouds can only be resolved in RES-2pc and RES-1pc. As we shall show (Section 3.1.2), at least 2 pc resolution is needed to accurately determine the average $X_{\text{CO}}$ in molecular clouds for the solar neighborhood conditions of the present simulations.

3.1.1. Molecular Abundances and Dependence of Chemistry on Numerical Resolution

As the numerical resolution increases from 4 to 1 pc, a larger fraction of mass in the simulations is in the dense gas. This is quantified by the increase in $f(100)$ (the fraction of gas with density $n > 100$ cm$^{-3}$) with resolution in Table 4, and the density distributions in Figure 2. The density distributions are similar at low densities where the gas is well resolved. At high densities, the distribution cuts off near the density threshold for creation of sink particles, where the unresolved dense gas is converted into sink particles in the simulations. As resolution increases, the density threshold for creation of sink particles also increases, allowing denser gas to form.

Tables 1, 2, and 3 list observational parameters and model parameters, respectively.

Table 1: Observational Parameters in Selected $X_{\text{CO}}$ Literature

| Reference          | Beam Size | CO Map Resolution | Distance | Object     | Velocity Resolution (km s$^{-1}$) | $T_{\text{tot}}$ (K) | $X_{\text{CO/20}}$ |
|--------------------|-----------|--------------------|----------|------------|-----------------------------------|----------------------|-------------------|
| Ripple et al. (2013) | 0.2 pc    | 0.1 pc             | 420 pc   | Orion      | 0.2                               | 2                    | 1.4               |
| Lee et al. (2015)   | 0.36 pc   | 0.06 pc            | 280 pc   | Perseus    | 0.064                             | 0.8                  | 0.3               |
| Pineda et al. (2008) | 0.4 pc    | 0.06 pc            | 280 pc   | Perseus    | 0.064                             | 0.35                 | 2 ± 1             |
| Leroy et al. (2011) | 60 pc     | 5.8 pc             | 50 kpc   | LMC        | 0.1                               | 0.35                 | 3.0               |
| Leroy et al. (2016) | 60 pc     | 11–60 pc           | 0.05–1.5 Mpc | nearby galaxies$^d$ | 1.6–5 | 0.03–0.2 | ... |
| Smith et al. (2012) | 140 pc    | 90 pc              | 780 kpc  | M31        | 2.6                               | 0.03                 | 1.9 ± 0.4         |
| Sandstrom et al. (2013)$^e$ | 0.6–4 kpc | 0.2–1.2 kpc        | 3.6–21.4 Mpc | spiral galaxies$^f$ | 2.6 | 0.02–0.04 | 1.4–1.8$^g$ |

Notes.

$^a$ Distances of Perseus and Orion molecular clouds are taken from Schlafly et al. (2014).

$^b$ Detection limit for CO($J = 1$–0) line emission. Same as the mean rms noise per velocity channel in observations.

$^c$ Note that the $X_{\text{CO}}$ in Lee et al. (2015) is smaller than that determined by Pineda et al. (2008). Lee et al. (2015) state that the discrepancy mainly results from a different adopted dust-to-gas ratio and the consideration of H1 gas.

$^d$ Antennae, LMC, M31, M33, M51, and M74.

$^e$ Observations used the CO($J = 2$–1) line and assumed a fixed line ratio (2–1)/(1–0) = 0.7.

$^f$ This is the average $X_{\text{CO}}$ in low-inclination galaxies. The dispersion is about 0.3 dex.

Table 2: Parameters for Synthetic Observations

| Beam Size (pc) | CO Map Resolution (pc) | Velocity Resolution (km s$^{-1}$) | $T_{\text{tot}}$ (K) |
|---------------|------------------------|-----------------------------------|----------------------|
| 1             | 1                      | 0.07                              | 0.4                  |
| 2             | 2                      | 0.07                              | 0.4                  |
| 4             | 2                      | 0.07                              | 0.4                  |
| 8             | 2                      | 0.07                              | 0.4                  |
| 16            | 2                      | 0.07                              | 0.4                  |
| 32            | 2                      | 0.07                              | 0.4                  |
| 64            | 4                      | 0.1                               | 0.35                 |
| 128           | 64                     | 2.6                               | 0.03                 |
| 512           | 128                    | 2.6                               | 0.03                 |
| 1024          | 256                    | 2.6                               | 0.03                 |

Table 3: Model Parameters

| ID | Resolution (pc) | $t$ (Myr) | $t_{\text{chem}}$ (Myr) |
|----|-----------------|-----------|--------------------------|
| RES-1pc | 1               | 382       | 50                       |
| RES-2pc | 2               | 382       | 50                       |
| RES-4pc | 4               | 382       | 50                       |
| TCHEM-5Myr | 1           | 382       | 5                       |
| TCHEM-50Myr | 1          | 382       | 50                      |
| T-356Myr | 2               | 356       | 50                       |
| T-416Myr | 2               | 416       | 50                       |

2.5. Model Parameters

We consider three sets of models designed to study different conditions that may affect $X_{\text{CO}}$: the numerical resolution, non-equilibrium chemistry, and variation in the galactic environment (ISM structure and ambient radiation field). The parameters for our models are summarized in Table 3. Model names denote changes in numerical resolution (RES-1pc, etc.), chemical evolution time (TCHEM-5Myr, etc.), and simulation snapshot time (T-356Myr, etc.). Note that $t$ is the time for the MHD simulation, and $t_{\text{chem}}$ is the time for the postprocessing chemistry, as detailed in Section 2.2. RES-1pc and TCHEM-50Myr are two names for the same model, used for clarity in different sections discussing the numerical resolution or evolving chemistry. To do a controlled study, we set the incident radiation field strength $\chi = 1$ (in units of Draine 1978, corresponding to $I_{\text{UV}} = 2.7 \times 10^{-3}$ erg cm$^{-2}$ s$^{-1}$) for all models that intercompare numerical resolution and non-equilibrium chemistry (model IDs starting with RES or TCHEM). In the set of models for studying the variation in galactic environments (model IDs starting with T), $\chi$ is obtained from the star cluster particles as described in KO2017.
Figure 1. The column density of all gas ($N$, first row), molecular gas ($N_{\text{H}_2}$, second row), CO ($N_{\text{CO}}$, third row), and the intensity of the CO($J = 1-0$) line ($W_{\text{CO}}$, last row) in models RES-4pc (left), RES-2pc (middle), and RES-1pc (right). The young (age < 40 Myr) star clusters/sink particles formed in the simulations are shown as filled circles in the first row with $N$. The areas of the circles are proportional to the square root of the cluster masses, ranging from $10^3$ to $10^5 M_\odot$ (see legends in the top left panel), and the color of the circles indicates the cluster age, from 0 (magenta) to 40 Myr (blue).
The change in density distribution with resolution affects the chemical composition of the gas. As the resolution increases from 4 to 1 pc, the total H$_2$ mass stays nearly constant, but the total CO mass increases by a factor of nearly 3 (Table 4). The reason for this is evident in Figure 3: most H$_2$ forms in the density range $n = 10$–100 cm$^{-3}$, which is already well resolved with 4 pc resolution. However, most CO forms at $n \gtrsim 200$ cm$^{-3}$, which is not well resolved with 2 pc, maybe even 1 pc resolution. Using adaptive mesh refinement (AMR) models, Seifried et al. (2017) found that a resolution of $\sim$0.2 pc is needed for the CO abundance to converge.

The chemical composition depends not only on density, which affects the rate of collisional reactions, but also on shielding, which determines the rate of photodissociation by FUV photons. Which factor, density or shielding, is more important in determining the H$_2$ and CO abundances in realistic molecular clouds with complex structures? Figures 4 and 5 plot the probability density distributions (PDFs) of the H$_2$ and CO abundances versus density and shielding in each grid cell. We weight the PDFs by $n_{H_2}$ or $n_{CO}$ so that the color scale is proportional to the H$_2$ or CO mass in each bin. Simple volume-weighted PDFs will show distributions centered at very low density and low molecular abundance, since by volume most gas is atomic.

We quantify the shielding by calculating the effective extinction $A_{\nu,\text{eff}}$ for the photoelectric heating (Gong et al. 2017),

$$\chi_{\text{PE}} = \chi \exp(-1.8A_{\nu,\text{eff}}),$$

where $\chi_{\text{PE}}$ is the actual radiation field intensity obtained from the six-ray radiation transfer.

As shown in Figure 4, the H$_2$ abundance has a much tighter correlation with density than with shielding. This is because H$_2$ self-shielding is so efficient that the photodissociation rate of

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**Table 4** Overall Properties of Models for Comparisons in Numerical Resolution and Non-equilibrium Chemistry

| Model ID       | $M_{\text{H}}(M_\odot)^a$ | $M_{\text{H}_2}(M_\odot)$ | $M_{\text{CO}}(M_\odot)$ | $L_{\text{CO}}$ (K km s$^{-1}$ pc$^2$)$^b$ | $\langle X_{\text{CO}} \rangle_0$ | $f_{\text{dark}}$ | $f_{100}$ | $2/(f_{\text{H}_2})$ |
|----------------|-----------------------------|-----------------------------|---------------------------|---------------------------------|--------------------------|-----------------|----------|----------------|
| RES-4pc        | $7.48 \times 10^6$         | $5.76 \times 10^4$         | $4.82 \times 10^4$       | $7.63 \times 10^4$              | 1.45                     | 69%             | 0.4%     | 11%           |
| RES-2pc        | $7.41 \times 10^6$         | $5.55 \times 10^4$         | $5.49 \times 10^4$       | $8.27 \times 10^4$              | 1.07                     | 75%             | 0.9%     | 10%           |
| RES-1pc (TCHEM-50Myr) | $7.41 \times 10^6$         | $6.89 \times 10^4$         | $1.21 \times 10^2$       | $1.22 \times 10^4$              | 1.02                     | 71%             | 2.3%     | 13%           |
| TCHEM-5Myr     | $7.41 \times 10^6$         | $2.46 \times 10^5$         | $8.96 \times 10^4$       | $9.06 \times 10^4$              | 0.56                     | 67%             | 2.3%     | 5%            |

Notes.

$^a$ Total mass $M_{\text{H}} = 1.4m_\odot \int n dV$. The factor 1.4 is from the helium abundance $f_{\text{He}} = 0.1$.

$^b$ Total luminosity of CO($J = 1\rightarrow0$) line.

$^c$ Average $X_{\text{CO}}$ in CO-bright regions. $\langle X_{\text{CO}} \rangle_0 = \langle X_{\text{CO}} \rangle / (10^{20} \text{ cm}^{-2} \text{ K}^{-1} \text{ km}^{-1} \text{s})$.

$^d$ CO-dark H$_2$ gas fraction (see Equation (10)).

$^e$ Fraction of mass with density $n > 100$ cm$^{-3}$.

$^f$ Fraction of hydrogen in H$_2$: $f_{\text{H}_2} = M_{\text{H}_2}/(M_{\text{H}} + M_{\text{H}_2})$.

---

$^{10}$ In Table 4, $M_{\text{H}}$ first decreases slightly when the resolution increases from 4 to 2 pc, then increases again at 1 pc resolution. This nonsecular variation of $M_{\text{H}}$, with resolution is actually a result of temporal variations in the simulations. Because the supernova feedback from the sink/cluster particles is stochastic, simulations with the same initial condition can develop slightly different density structures over time. We compared $M_{\text{H}}$ and $M_{\text{CO}}$ in models RES-4pc and RES-2pc between the time when they have the same initial condition (350 Myr) and the time of comparison in Table 4 (382 Myr). We found that the H$_2$ masses in the two models are similar (up to $\sim$20% variations), but the CO mass increases significantly (up to a factor of $\sim$3) in the RES-2pc model. The H$_2$ and CO mass-weighted density histograms at different times also show very similar features to Figure 3. Therefore, the conclusion from Figure 3 is robust despite the temporal variations.
H₂ is very small in most regions that have a significant amount of H₂. In the absence of photodissociation by FUV radiation, the H₂ abundance is then determined by the balance between H₂ formation on dust grains,

\[ \text{H} + \text{H} + \text{gr} \rightarrow \text{H}_2 + \text{gr}, \]  
(12)

with a rate coefficient \( k_{gr} = 3.0 \times 10^{-17} \text{ cm}^3 \text{s}^{-1} \) (assuming solar neighborhood dust abundance), H₂ formation by H₂⁺,

\[ \text{H}_2^+ + \text{e} \rightarrow \text{H}_2 + \text{H}, \]  
(13)

with a rate coefficient \( k_{13}, \) H₂ destruction by cosmic rays,

\[ \text{CR} + \text{H}_2 \rightarrow \text{H}_2^+ + \text{e}, \]  
(14)

with a rate coefficient \( k_{CR} = 2g_{\text{HI}}(2.3f_{\text{HI}} + 1.5f_{\text{H}}) \), and H₂ destruction by H₂⁺,

\[ \text{H}_2^+ + \text{H}_2 \rightarrow \text{H}_2^+ + \text{H}, \]  
(15)

with a rate coefficient \( k_{15}. \) Reactions (14) and (15) are also the main pathways for H₂⁺ destruction and creation. Equilibrium of H₂⁺ requires

\[ f_{\text{HI}}k_{CR} = f_{\text{HI}}f_{\text{HI}}n_k l_5. \]  
(16)

H₂⁺ is mainly created by Reaction (15), and destroyed by the reaction H₂⁺ + e, which forms H₂ + H (Reaction (13)) or 3H with a branching ratio of 0.35:0.65. Equilibrium of H₂⁺ requires

\[ f_{\text{HI}}f_{\text{HI}}n_k l_5 = \frac{1}{0.35}f_{\text{HI}}f_{\text{HI}}n_k l_5. \]  
(17)
Finally, equilibrium of fully shielded H$_2$ (Equations (12)–(17)) requires

\[
\begin{align*}
    f_{H_2} n k_{gr} + f_{H}^e f_{e} n k_{13} &= f_{H_2}^e k_{CR} + f_{H}^e f_{H_2} n k_{15} \\
    f_{H_2} n k_{gr} + 0.35 f_{H}^e f_{e} n k_{15} &= f_{H_2}^e k_{CR} + f_{H}^e f_{H_2} n k_{15} \\
    f_{H_2} n k_{gr} &= 1.65 f_{H_2}^e k_{CR}.
\end{align*}
\]  

In the above, each $f$ is the abundance of a given species relative to hydrogen nuclei.

Equation (18) can be solved with the conservation of hydrogen nuclei $f_{H_2} = 2 - f_{H}^e$, giving the equilibrium H$_2$ abundance as a function of $n$, plotted as the green dashed line in the left panel of Figure 4. This agrees very well with the upper limit of $f_{H_2}$ in the simulations. The spread of $f_{H_2}$ at a given density is due to the incomplete shielding of FUV radiation in some regions where destruction of H$_2$ from photodissociation brings its abundance below that in completely shielded regions. This can also be seen in the right panel of Figure 4: there is a large spread of $A_{V,eff}$ at a given $f_{H_2}$, and there are many grid cells with $A_{V,eff} < 1$ and significant H$_2$ abundance.

Contrary to the case of H$_2$ abundance, which is determined mostly by density, the CO abundance is determined by both density and shielding, as shown in Figure 5. CO forms mainly in regions with $n \geq 100$ cm$^{-3}$ and $A_V \geq 1$. This agrees very well with the results from 1D slab models in Gong et al. (2017, see their Figures 5 and 6). The main reason why H$_2$ and CO form under different conditions is that the self-shielding of CO and cross-shielding of CO by H$_2$ are much less efficient than the H$_2$ self-shielding. As a result, CO formation is limited by photodissociation, and CO can only form in regions with $A_{V,eff} \geq 1$ where the FUV radiation field is sufficiently shielded by dust. Moreover, CO formation also requires higher densities, because C$^+$ and He$^+$ formed by cosmic rays destroy CO at lower densities. Figures 4 and 5 again show that the H$_2$ mass in our simulations is converged, but the CO mass is not, due to the lack of resolution for gas of very high density (see also Figure 3 and discussion).

Because formation of H$_2$ and formation of CO require different conditions, CO is only a very approximate tracer of H$_2$. Figure 6 shows the distribution of density $n$ versus the effective extinction $A_{V,eff}$ for each grid cell. At a given density, there is a large range of $A_{V,eff}$. We roughly delineate loci where H, H$_2$, and CO form: H$_2$ exists in high-density regions, and $f_{H_2} > 0.5$ corresponds to densities $n \gtrsim 30$ cm$^{-3}$. CO forms in denser and well shielded regions, and $f_{CO} > 10^{-5}$ roughly corresponds to $n \gtrsim 100$ cm$^{-3}$ and $A_{V,eff} \gtrsim 1$. Figure 6 clearly shows that a significant fraction of H$_2$ would not be traced by CO emission (see $f_{dark}$ in Table 4). As $\Delta x$ decreases from 4 to 1 pc, more and more high-density gas is resolved, as also shown in Figure 2. Nevertheless, for all resolutions considered in our models, there is gas in the three different regimes—atomic, CO-bright molecular, and CO-dark molecular.

To validate that we can accurately simulate chemistry in molecular clouds, we compare the CO column densities $N_{CO}$ in our simulations to that in the UV absorption observations of diffuse molecular clouds. Figure 7 shows the comparison between the simulations and observations, as well as the result from the one-sided slab model in Gong et al. (2017). The x-axis of Figure 7 is the extinction from only H$_2$:

\[
A_V(N_{H_2}) = \frac{2 N_{H_2}}{1.87 \times 10^{21} \text{cm}^{-2}}.
\]  

To avoid foreground/background contamination, we compare $N_{CO}$ to $N_{H_2}$ instead of the total column $N$.\footnote{Gong et al. (2017) discussed that the dispersion in observations is much smaller when comparing $N_{CO}$ to $N_{H_2}$ instead of $N$.} Compared to the simulations, the one-sided slab model gives higher CO abundance at $A_V(N_{H_2}) \sim 1$. This is because the six-ray radiation transfer in the 3D simulations considers extinction of FUV radiation from all directions along the Cartesian axes, which is generally lower than the extinction only along the z-axis, $A_V(N_{H_2})$ (that is, $A_{V,eff} \lesssim A_V(N_{H_2})$). At $A_V(N_{H_2}) \ll 1$ or $A_V(N_{H_2}) \gg 1$, the CO abundances in the one-sided slab model and in 3D simulations are more similar, because either the FUV radiation is only weakly shielded at low $A_V(N_{H_2})$ so that the...
photodissociation rate is insensitive to \( A_V(N_{\text{H}_2}) \), or it is already completely shielded at high \( A_V(N_{\text{H}_2}) \) so that the limiting factor for CO formation is no longer photodissociation. The UV absorption observations can only be conducted in diffuse molecular clouds with \( A_V(N_{\text{H}_2}) \lesssim 1 \), and there is a lack of observations at higher extinctions. For the range of \( A_V(N_{\text{H}_2}) \) where the observational data are available, the RES-1pc simulation successfully reproduces the observed range of \( N_{\text{CO}} \). Lower resolution simulations RES-2pc and RES-4pc also show similar average values (magenta lines) and range (not shown in the figure) of \( N_{\text{CO}} \) at \( A_V \lesssim 1 \). At \( A_V > 1 \), models with lower resolutions start to show that the CO mass is not resolved at high densities.

### 3.1.2. Dependence of \( X_{\text{CO}} \) on Numerical Resolution

To understand the relation between physical properties of molecular clouds and CO emission, a helpful reference point is the simple uniform slab model for molecular clouds. In a molecular clouds with \( A_V \approx 1 \), models with \( A_V > 1 \), models with lower resolutions start to show that the CO mass is not resolved at high densities.

In the limit of \( \tau_v \to \infty \), Equations (20) and (21) become

\[
T_A = \frac{T_0}{e^{T_{\text{exc}}/T_0} - 1},
\]

where \( T_0 = 5.5 \, \text{K} \) is \( h\nu_0/k \), with \( \nu_0 = 115.3 \, \text{Hz} \), the frequency of the CO(J = 1–0) line.

Typically, the CO(J = 1–0) line profile (in terms of \( T_A \) and \( \nu \) is not too far from a Gaussian profile, and to first order, the total CO line intensity \( W_{\text{CO}} \) is determined by two parameters: the peak of the line profile \( T_{\text{peak}} \) and the width/velocity dispersion of the line \( \sigma_v \). Under the assumption that the line center is optically thick so that Equation (22) applies, the observed peak antenna temperature, \( T_{\text{peak}} \), would be directly related to the excitation temperature \( T_{\text{line}} \).

\[
T_{\text{line}} \equiv \frac{5.5 \, \text{K}}{\ln(5.5 \, \text{K}/T_{\text{peak}} + 1)}.
\]

We use the notation \( T_{\text{line}} \) for the excitation temperature derived from the line profile (\( T_{\text{peak}} \)) to distinguish it from the true excitation temperature in the molecular clouds \( T_{\text{exc}} \). Although \( T_{\text{line}} \) is \( T_{\text{exc}} \) in a uniform slab cloud as long as the CO line center is optically thick, in real molecular clouds and also in our numerical simulations, the excitation temperature along the line of sight is not constant, and \( T_{\text{line}} \) serves as an estimate of the excitation temperature where most CO emission comes from. For \( T_{\text{peak}} \gtrsim 5.5 \, \text{K} \), Equation (23) gives \( T_{\text{line}} \approx T_{\text{peak}} \).

Another important parameter for the CO line, the velocity dispersion, is calculated using \( \sigma_v \equiv \sqrt{(v^2)_{\text{rms}} - \langle v \rangle^2} \), where \( \langle v \rangle \equiv \int v T_A \, dv / \int T_A \, dv \) is the intensity-weighted average of velocity, and similarly \( (v^2)_{\text{rms}} \equiv \int v^2 T_A \, dv / \int T_A \, dv \).

The relations between \( W_{\text{CO}} \) and \( T_{\text{line}} \) for \( \sigma_v \) are shown in Figure 8. \( T_{\text{line}} \) ranges

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\( ^{12} \) In observations, the intensity is often referred to as the value after background subtraction \( I_{\text{obs}} = I_v - I_0(0) \). Then Equation (20) is often written as \( I_{\text{obs}} = (B_v(T_{\text{exc}}) - I_0(0))(1 - e^{-\tau_v}) \).
from ~2 K (from the CMB background) to ~20 K (from the kinetic temperature of dense gas as discussed below), similar to the range of excitation temperature observed in Perseus and Taurus molecular clouds (Pineda et al. 2008, 2010). The velocity dispersion spans a relatively narrow range \( \sigma_v \approx 1-2 \text{ km s}^{-1} \), and the lower limit for \( \sigma_v \) is set by the subgrid microturbulence velocity in Equation (9). The observations of nearby molecular clouds have higher resolutions of ~0.2-0.4 pc, and therefore a slightly lower but still limited range of velocity dispersions \( \sigma_v \approx 0.8-1.5 \text{ km s}^{-1} \) (Pineda et al. 2010; Kong et al. 2015). \( W_{\text{CO}} \) increases with both \( T_{\text{line}} \) and \( \sigma_v \). For a Gaussian profile with \( T_{\text{peak}} \gtrsim 5.5 \text{ K} \), \( W_{\text{CO}} = \sqrt{2/\pi} T_{\text{peak}} \sigma_v \approx \sqrt{2/\pi} T_{\text{line}} \sigma_v \). Because the variation in \( \sigma_v \) is small, \( W_{\text{CO}} \) correlates very well with \( T_{\text{line}} \) except for regions where \( T_{\text{line}} \) saturates around 20 K. There is no saturation of \( W_{\text{CO}} \), and \( W_{\text{CO}} \) keeps increasing with increasing \( \sigma_v \).

\( W_{\text{CO}} \) is largely determined by the excitation temperature, and the excitation temperature in turn depends on the gas density and temperature. Figure 9 shows the excitation temperature \( T_{\text{exc}} \) and gas temperature \( T_{\text{gas}} \) versus the gas density in each grid cell. \( T_{\text{gas}} \) decreases with increasing density, as the gas cooling becomes more efficient, and heating is also reduced by shielding of the FUV radiation field in dense regions. On the other hand, \( T_{\text{exc}} \) increases with increasing density, because the collisional excitation rate of CO is proportional to density, and because radiative trapping increases in dense regions.

The lower solid magenta line in Figure 9 shows the median \( T_{\text{exc}} \) from model RES-1pc as a function of density. \( T_{\text{exc}} \) only reaches equilibrium with \( T_{\text{gas}} \) at \( n \gtrsim 400 \text{ cm}^{-3} \), implying that the approximation of local thermal equilibrium (LTE) would fail in most regions.

At a given density, \( T_{\text{exc}} \) is higher at lower resolutions for two reasons. First, the velocity gradient \( |dv/dr| \) is smaller at lower resolutions, leading to higher \( T_{\text{gas}} \) and thus lower escape probability \( \beta \) and higher \( T_{\text{exc}} \) at a given density (see Equations (28) and (30) below). Second, at lower resolutions, less high-density gas is resolved, and a larger fraction of the CO gas is found in lower-density gas (see Figure 3(b)). This shifts the distributions of \( T_{\text{exc}} \) and \( T_{\text{gas}} \) in Figure 9 to the left at lower resolution in models RES-2pc and RES-4pc (dashed and dotted magenta lines).

In general, thermalization is expected for densities above a critical value at which collisional de-excitation exceeds spontaneous emission. For CO collisions with H\(_2\), the collisional de-excitation rate is \( n_{\text{H}_2} k_{10} \) for

\[
k_{10} \approx 6 \times 10^{-11} \left( \frac{T_{\text{gas}}}{100 \text{ K}} \right)^{0.2} \text{ cm}^3\text{s}^{-1},
\]

at 10 K \( \lesssim T_{\text{gas}} \lesssim 250 \text{ K} \) (Flower & Launay 1985; Flower 2001; Draine 2011). The spontaneous emission rate is \( \beta A_{10} \), where Equation (6) gives the escape probability \( \beta \), so that

\[
n_{\text{crit}} = \frac{\beta A_{10}}{k_{10}}.
\]

For \( T_{\text{gas}} = 20 \text{ K} \), Equation (24) gives \( A_{10}/k_{10} \approx 2.1 \times 10^3 \text{ cm}^{-3} \).

With increasing density, the optical depth \( \tau_{\text{LVG}} \) increases, leading to decreasing \( \beta \) (Figure 10); at large \( \tau_{\text{LVG}} \), \( \beta \approx 1/\tau_{\text{LVG}} \). For model RES-1pc, we fit the average \( \tau_{\text{LVG}} \) at a given density with a broken power law (magenta line in Figure 10):

\[
\tau_{\text{LVG}} = 2.4 \times 10^{-5} (n/\text{cm}^{-3})^{2.3}, \quad n < 350 \text{ cm}^{-3}
\]

\[
\tau_{\text{LVG}} = 0.21 (n/\text{cm}^{-3})^{0.73}, \quad n \geq 350 \text{ cm}^{-3}.
\]

Combining Equations (25) and (26) yields \( n_{\text{crit}} \sim 300 \text{ cm}^{-3} \). Thus, in regions where \( n \geq 400 \text{ cm}^{-3} \), the CO(J = 1) level is expected to be thermalized, and this is indeed consistent with the median \( T_{\text{exc}} \) for model RES-1pc.\(^{13}\)

13 We note that \( \tau_{\text{LVG}} \) depends on the density and velocity structure, which is resolution-dependent, so the density for thermalization is not expected to be the same for models RES-2pc and RES-4pc as for model RES-1pc. In fact, the velocity gradient \( |dv/dr| \) is smaller at lower resolutions, leading to higher average \( \tau_{\text{LVG}} \), and lower density for thermalization in models RES-2pc and RES-4pc (see Equation (7) and discussions of Figure 9).
The excitation temperature can be estimated as a function of density by Equations (27) and (28) (assuming \( \tau = \tau_{\text{LVG}} \) in Equation (6) and using the average value of \( T_{\text{gas}} \) at a given density). The analytic two-level system approximation for simulation RES-1pc (green line) agrees well with the result from radiation transfer by the RADMC-3D code (lower solid magenta line) at low and high densities, while there are differences within a factor of 2 at intermediate densities \( n \sim 300-1000 \, \text{cm}^{-3} \). This is because the CO rotational levels \( J = 1, 2, \) and 3 have energies of 5.5, 16.6, and 33.2 K, all lower than or comparable to the gas temperature. Indeed, there are significant populations in the \( J \geq 2 \) levels, as expected given that \( T_{\text{gas}} > 5.5 \, \text{K} \). The analytical expression in Equation (28) takes into account only the \( J = 0 \) and \( J = 1 \) levels, and therefore cannot predict the excitation temperature very accurately. At low and high densities the differences are small because the excitation temperature there is determined by the background CMB temperature or the gas temperature as the CO rotational levels approach LTE. Nonetheless, the analytical two-level system approximation agrees with the general trend from the radiation transfer calculations, and gives some insight into the relation between \( T_{\text{exc}} \), \( T_{\text{gas}} \), and \( n \). As a further test, we ran RADMC-3D including only the first \( J = 0 \) and \( J = 1 \) rotational levels of CO, and found that it can indeed reproduce the analytical result of the two-level system model. Figure 23 shows this comparison.

The relation between \( W_{\text{CO}} \) and \( T_{\text{gas}} \), as well as the relation between \( T_{\text{exc}} \) and \( T_{\text{gas}} \), gives rise to the strong correlation between \( W_{\text{CO}} \) and the average (mass-weighted) density \( \langle n \rangle_M \) along the line of sight (Figure 11 left panel). Moreover, we found that in the simulations, \( N_{\text{H}_2} \) increases systematically with \( \langle n \rangle_M \) (see Figure 6). This results in a correlation between \( W_{\text{CO}} \) and \( N_{\text{H}_2} \) (Figure 11 right panel). Although \( X_{\text{CO}} \) is measured in terms of \( W_{\text{CO}} \) and \( N_{\text{H}_2} \), there is a smaller dispersion in the correlation between \( W_{\text{CO}} \) and \( \langle n \rangle_M \). This suggests that the CO emission is more fundamentally a measure of \( \text{H}_2 \) density than of column density.

The effect of numerical resolution on \( W_{\text{CO}} \) is already evident in Figures 9 and 11. As the resolution increases, more high-density gas forms in the simulation, and thus there are more pixels with high \( W_{\text{CO}} \). Numerical resolution also has an effect on \( X_{\text{CO}} \), as shown in Figure 12, the histogram of \( X_{\text{CO},20} = X_{\text{CO}}/(10^{20} \, \text{cm}^{-2} \, \text{K}^{-1} \, \text{km}^{-1} \, \text{s}) \) weighted by \( W_{\text{CO}} \). The
average $X_{\text{CO}}$ in a certain region can be written as

$$\langle X_{\text{CO}} \rangle = \frac{\sum n_{\text{H}_2}}{\sum W_{\text{CO}}} = \frac{\sum n_{\text{H}_2} W_{\text{CO}}}{\sum W_{\text{CO}}} = \frac{\sum W_{100} W_{\text{CO}}}{\sum W_{\text{CO}}}. \quad (31)$$

In other words, $\langle X_{\text{CO}} \rangle$ is simply the $W_{\text{CO}}$-weighted average of $X_{\text{CO}}$ in each pixel. Therefore, the peak of the histogram in Figure 12 roughly indicates the average $X_{\text{CO}}$ in the whole simulation domain. The distributions of $X_{\text{CO}}$ in models RES-1pc and RES-2pc are very similar, with a slightly higher peak in RES-1pc. As a result, $\langle X_{\text{CO}} \rangle$ is almost the same in RES-1pc and RES-2pc (Table 4). The model RES-4pc, however, peaks at larger $X_{\text{CO}}$ than the higher resolution models, and therefore has a higher $\langle X_{\text{CO}} \rangle$. This is because the peak of the $X_{\text{CO}}$ distribution, $X_{\text{CO},20} \approx 0.5$, comes from regions with moderately high density $n \approx 100$ cm$^{-3}$ and CO emission $W_{\text{CO}} \approx 40$ K · km s$^{-1}$, which can only be resolved at a resolution finer than 2 pc (see histograms of $n_{\text{H}_2}$ and $W_{\text{CO}}$ in Figure 11). Therefore, we conclude that a numerical resolution of at least 2 pc is needed in order to resolve the average $X_{\text{CO}}$ in molecular clouds for solar neighborhood conditions.

Finally, we compare the distribution of $W_{\text{CO}}$ versus $A_V(n_{\text{H}_2})$ in model RES-1pc to observations of the Orion A and B molecular clouds by Ripple et al. (2013), as shown in Figure 13. Considering the noise level in the observation, we use a higher threshold of $W_{\text{CO}} > 1$ K km s$^{-1}$ to compare to the CO-bright region in Orion. Because most CO emission comes from regions with $W_{\text{CO}} > 1$ K km s$^{-1}$, $X_{\text{CO}}$ is not sensitive to the $W_{\text{CO}}$ threshold. Our simulation shows a similar distribution of $W_{\text{CO}}$ versus $A_V(n_{\text{H}_2})$ to that in Orion. The dispersion of $W_{\text{CO}}$ at a given $A_V(n_{\text{H}_2})$ is large, even more than an order of magnitude at low $A_V(n_{\text{H}_2})$. However, despite the large dispersion of $W_{\text{CO}}$, the average $X_{\text{CO}}$ (which is inversely proportional to the slope) in different $A_V(n_{\text{H}_2})$ bins is very similar, only varying by a factor of $\sim 2$. Similar features are observed in many Milky Way molecular clouds by Lee et al. (2018), and we discuss this in more detail in Section 3.3.

There are also some differences between the simulation and observations. The average $X_{\text{CO}}$ in RES-1pc is a factor of 1.4 lower than that in Orion. As we shall show based on other analyses and comparisons, the typical $X_{\text{CO}}$ in our simulations is
about a factor of \( \sim 2 \) lower than the standard Milky Way value; we discuss possible reasons for this discrepancy at the end of Section 3.3. We also note that because the observation in Ripple et al. (2013) has a higher spatial resolution of \( \sim 0.2 \) pc, there are more pixels at \( A_V(N_{H_2}) \geq 4 \) in the observation in Figure 13. The simulation has more pixels at \( W_{CO} \gtrsim 60 \text{ K km s}^{-1} \), a result of the slightly higher velocity dispersions (see Figure 8 and discussion). In spite of these differences, the general agreement between the models and observations indicates that the simulations can successfully reproduce the basic physical properties of observed molecular clouds.

### 3.2. Non-equilibrium Chemistry

The realistic ISM is highly dynamical: turbulence constantly creates and disperses molecular clouds, and moves gas to environments with different density, temperature, and radiation field strength. As a result, non-equilibrium chemistry is likely to be important, especially in low-density diffuse gas where the chemical timescales are long compared to the dynamical timescales. This is especially an issue for \( H_2 \). Molecular hydrogen can form in low-density gas because of its effective self-shielding, but its formation timescale, \( \tau_{H_2} \approx 10 \) Myr \((n/100 \text{ cm}^{-3})^{-1}\) (Gong et al. 2017), can be longer than the dynamical timescale (Equation (3)). Because CO formation chemically relies on the existence of \( H_2 \), the CO abundance in molecular clouds can also be far from equilibrium. In this section, we carry out comparisons between models with different \( t_{chem} \) (model IDs start with TCHEM in Table 3) to investigate the effect of non-equilibrium chemistry on \( X_{CO} \).

Both \( H_2 \) abundance and CO abundance increase over \( t_{chem} \), reaching a steady state at \( t_{chem} \approx 50 \) Myr, as shown in Figure 14. Over timescales relevant for clouds of size \( \gtrsim 10 \) pc (Equation (3)), there is a larger increase in the \( H_2 \) abundance than in the CO abundance: from \( t_{chem} = 5 \) to 50 Myr, the \( H_2 \) abundance increases by a factor of \( \sim 3 \), while the CO abundance increases only by \( \sim 30\% \).

The difference in the evolution of \( H_2 \) and CO abundances comes from their different distributions. As shown in Figure 15, both \( H_2 \) and CO abundances are closer to equilibrium at higher densities, because the rate of collisional reactions increases with density. In fact, at a given density in the range \( \sim 40 \)–\( 400 \) \( \text{cm}^{-3} \), at 5 Myr, the abundance of \( H_2 \) is closer to its final value than the abundance of CO. However, in equilibrium most of the \( H_2 \) is in gas at intermediate densities \( n \approx 10 \)–\( 100 \) \( \text{cm}^{-3} \), whereas most CO is in gas at high densities \( n \gtrsim 200 \) \( \text{cm}^{-3} \) (Figure 3). This leads to a shorter timescale for the overall CO abundance to reach equilibrium than for \( H_2 \). Since the CO luminosity also increases much less than the \( H_2 \) mass does, this leads to a lower \( X_{CO} \) value at early \( t_{chem} \) (Table 4).

Non-equilibrium chemistry also has an effect on the distribution of \( W_{CO} \) versus \( A_V(N_{H_2}) \). For model TCHEM-5Myr (Figure 16), the distribution of the pixels is shifted to the left compared to that in TCHEM-50Myr (left panel of Figure 13). This is because \( W_{CO} \) is close to equilibrium, but \( N_{H_2} \) is a factor of \( \sim 2 \) smaller than the equilibrium values, for the same reasons discussed above. Moreover, the distribution of \( W_{CO} \) versus \( A_V(N_{H_2}) \) in TCHEM-5Myr shows some hints of a plateau for \( W_{CO} \) at high \( A_V(N_{H_2}) \), especially in the binned average value of \( W_{CO} \), which is not present in TCHEM-50Myr. This implies that younger clouds may have not only lower \( X_{CO} \) on average, but also different distributions of \( W_{CO} \) versus \( A_V(N_{H_2}) \) compared to older clouds. We discuss this further in Section 3.3. Note that \( N_{H_2} \) includes all \( H_2 \) along the line of sight, both in high-density clumps where CO forms, and in the foreground/background low-density envelopes with only \( H_2 \) and no CO. Because most \( H_2 \) (in equilibrium) lies in these low-density envelopes, the fractions of \( H_2 \) in CO-bright and CO-dark regions increase by similar proportions with increasing \( t_{chem} \), and \( f_{dark} \) stays constant from \( t_{chem} = 5 \) Myr to \( t_{chem} = 50 \) Myr (Table 4).

### 3.3. Variations in Galactic Environments

Galactic environment fundamentally impacts the molecular content of the ISM. Supernova feedback creates and destroys molecular clouds; shocks and turbulence shape molecular
and and and and and

\[50 \text{ Myrchem}\]

\[\text{TCHEM} - 5 \text{ Myr}\]

\[\text{H}2\text{ abundance} \sim \text{CO abundance}

\text{at } t_{\text{chem}} = 50 \text{ Myr.}

\[\text{TCHEM-5Myr}\]

\text{CO is well resolved with a resolution of 2 pc in these}

\text{models. The}

\[X_{\text{CO}}\]

\text{is much larger in their}

\text{galactic environments.}

\[X_{\text{CO}}\]

\[X_{\text{CO}}\]

\[\approx 0.6\] with

\[A_V\text{CO}\]

\text{higher}

\[R^2 < 0.4 \text{ in linear regression between } X_{\text{CO}} \text{ and } M_{\text{H}}\text{, the}

\text{radiation field strength } \chi\text{, or the average extinction from H}_2\text{ in CO-bright regions} (A_V)\text{CO}\text{. Remy et al. (2017) measured } X_{\text{CO}}\text{ in individual Milky Way molecular clouds using } \gamma}\text{-ray observations, and they also found no strong correlation between } X_{\text{CO}}\text{ and } M_{\text{H}}\text{ or } (A_V)\text{CO}.\text{14}

Remy et al. (2017) found a slight anticorrelation of \(X_{\text{CO}}\) and \(W_{\text{CO}}\): \(X_{\text{CO}} \sim -0.051 \times W_{\text{CO}}\). We similarly found a slight anticorrelation (Figure 18 left panel), with \(X_{\text{CO}}/W_{\text{CO}} = -0.011 \pm 0.005 (W_{\text{CO}}/\text{K km s}^{-1}) + 1.0 \pm 0.09\), where the uncertainties represent the 90\% confidence intervals for the fitted slope and intercept. The slope of the linear fit is very shallow, and \(X_{\text{CO}}\) is not sensitive to changes in \(W_{\text{CO}}\). We note, however, that Remy et al. (2017) focus on the nearby low-mass molecular clouds with much lower values of \(W_{\text{CO}}\) \(\approx 2-10\) K km s\(^{-1}\) than \((W_{\text{CO}}) \approx 10-20\) K km s\(^{-1}\) in the giant molecular clouds in our simulations, and therefore their work may not be directly comparable to our results.

Large-scale galaxy simulations by Narayanan et al. (2012) found a similar trend that \(X_{\text{CO}}\) decreases with increasing \(W_{\text{CO}}\); although the range of \(W_{\text{CO}}\) is much larger in their simulations because they consider a wide range of galactic environments. Narayanan et al. (2012) found that the \(X_{\text{CO}}\)-(\(W_{\text{CO}}\)) relation is caused by the increase in gas temperature and velocity dispersion at high \((W_{\text{CO}})\), which leads to a faster increase in \(W_{\text{CO}}\) than \(N_{\text{H}_2}\), resulting in the decrease in \(X_{\text{CO}}\). Similarly, we found that the snapshots in our simulations with higher \((W_{\text{CO}})\) also have larger velocity dispersions, although the gas temperature is roughly constant in the CO-forming regions in our models (see discussion of Figure 8 in Section 3.1.2). Interestingly, this is also consistent with the fact that the molecular clouds in the Galactic center have larger velocity dispersions and lower \(X_{\text{CO}}\) than the solar neighborhood clouds. We plan to carry out numerical simulations with Galactic-center-like environments in the future to study the variation of \(X_{\text{CO}}\) in detail.

Unlike \(X_{\text{CO}}\), the fraction of CO-dark \(H_2\), \(f_{\text{dark}}\), does show significant variations and a strong correlation \((R^2 = 0.6)\) with \((A_V)\text{CO}\) (Figure 18 right panel). Linear regression gives

\[f_{\text{dark}} = 0.56\]

\text{sample. Excluding the Perseus cloud, we found no strong correlation}

\[R^2 < 0.6.\] However, this relation is largely driven by one outlier, the Perseus cloud, which has much lower \(X_{\text{CO}}\) and higher \((A_V)\text{CO}\) than the rest of the sample. Excluding the Perseus cloud, we found no strong correlation \((R^2 \approx 0.3)\) between \(X_{\text{CO}}\) and \((A_V)\text{CO}\) for the rest of their sample.\text{14}
\( f_{\text{dark}} = -0.31 \pm 0.14 \langle A_V \rangle_{\text{CO}} + 1.0 \pm 0.2 \), where the uncertainties represent the 90% confidence intervals for the fitted slope and intercept. \( f_{\text{dark}} \) increases with decreasing \( \langle A_V \rangle_{\text{CO}} \). In other words, there is more CO-dark \( \text{H}_2 \) in more diffuse molecular clouds, which is not surprising because CO forms in denser gas than \( \text{H}_2 \). The same trend was identified in the simplified spherical molecular cloud model by Wolfire et al. (2010).\(^{15}\) We note that Wolfire et al. (2010) use a slightly different definition of CO-dark \( \text{H}_2 \), and we use Equation (37) to translate their definition to ours. We have also performed an experiment by running the T-381Myr model and varying only the radiation field strength, and found that \( f_{\text{dark}} \) stays constant over \( \chi = 0.4–3.5 \), confirming the result from Wolfire et al. (2010) that \( f_{\text{dark}} \) is not sensitive to \( \chi \).

Another comparison of \( X_{\text{CO}} \) with observations is shown in Figure 19, where the \( X_{\text{CO}} \) in each pixel is plotted against \( T_{\text{line}} \). Comparing to the California cloud observed by Kong et al. (2015), our simulations show a similar slope for the relation between \( X_{\text{CO}} \) and \( T_{\text{line}} \) at \( T_{\text{line}} > 6 \text{ K} \) (the observational data are not available at lower \( T_{\text{line}} \)). However, the value of \( X_{\text{CO}} \) at a given \( T_{\text{line}} \) is about a factor of \( \sim 4 \) lower than the observations. One reason for this discrepancy may be that Kong et al. (2015)

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\(^{15}\) The result from Wolfire et al. (2010) shown in Figure 18 is taken from their model with metallicity \( Z' = 1.9 \) and incident radiation field \( \chi = 10 \). Wolfire et al. (2010) found that \( f_{\text{dark}} \) is not sensitive to \( Z' \) or \( \chi \) in their studies.
observed the CO($J = 2–1$) line and assumed a fixed line ratio of $W_{\text{CO}}(J = 2–1)/W_{\text{CO}}(J = 1–0) = 0.7$, and this ratio is very uncertain. As discussed below in more detail, generally different observations and also our simulations show similar trends for the variations in $X_{\text{CO}}$, but the absolute value of $X_{\text{CO}}$ can differ by a factor of a few.

Using all of the simulation models, a summary of $X_{\text{CO}}$ as a function of $A_V(N_{\text{H}_2})$ and comparison with observations is shown in Figure 20. Because of the large uncertainties in observations of $X_{\text{CO}}$ at low $A_V(N_{\text{H}_2})$, we plot only the data at $A_V(N_{\text{H}_2}) > 1$. Both in our simulations and in the observations, there is a factor of $\sim 2$ variation in $X_{\text{CO}}$ over $A_V(N_{\text{H}_2}) = 1–12$. Simulations with $t_{\text{chem}} = 50$ Myr (RES-1pc, RES-2pc, T-356Myr–T-416Myr) show a decrease in $X_{\text{CO}}$ at $A_V(N_{\text{H}_2}) \lesssim 3$, regardless of the resolution and variations in galactic environments. Similar trends can be seen in the observations of Orion molecular clouds by Lee et al. (2018) and Ripple et al. (2013).

In contrast, the TCHEM-5Myr model shows a flatter profile at $A_V(N_{\text{H}_2}) \lesssim 3$ and a slight increase in $X_{\text{CO}}$ at $A_V(N_{\text{H}_2}) > 3$. Interestingly, the California cloud observed by Lee et al. (2018) also shows a similar trend. Compared to Orion, the California cloud has similar mass and distance, but a star formation rate that is an order of magnitude lower, and therefore it is believed to be much younger (Lada et al. 2009). This has the interesting implication that the profile of $X_{\text{CO}}$ as a function of $A_V(N_{\text{H}_2})$ may be used as an indicator of the age of molecular clouds.

Although the trend for the correlation between $X_{\text{CO}}$ and $A_V(N_{\text{H}_2})$ is similar in our simulations and observations, there is a discrepancy in the absolute value of $X_{\text{CO}}$. This may be due to systematic errors in either observations or simulations. One major uncertainty in observations of $X_{\text{CO}}$ comes from the assumptions in deriving $N_{\text{H}_2}$. Estimations of $N_{\text{H}_2}$ based on $\gamma$-ray emission systematically give a factor of 2 lower $X_{\text{CO}}$ than dust-based methods, consistent with the value of $X_{\text{CO}}$ in this paper (Bolatto et al. 2013; Remy et al. 2017, see also Figure 20). Even within the dust-based methods, the estimate of $X_{\text{CO}}$ in Orion A based on dust emission is a factor of $\sim 2$ in observations of Lee et al. (2018) compared to that in Ripple et al. (2013) based on dust extinction. As another example, the $X_{\text{CO}}$ in Perseus measured by Lee et al. (2015) (dust emission) is a factor of $\sim 7$ lower than that in Pineda et al. (2008) (dust extinction).

Several possible factors can contribute to the systematics in dust-based observations: different assumptions of the dust-to-gas ratio, uncertainties in foreground/background subtraction, and different resolutions/beam sizes (although the resolution effect is relatively mild, as noted by Lee et al. 2015 and discussed in Section 3.4). Lee et al. (2015) discussed in detail for the case of the Perseus molecular cloud that all these factors can indeed lead to a different estimate of $X_{\text{CO}}$. Sample differences in observations may also play a role. Most observations of $X_{\text{CO}}$ are for nearby low-mass star-forming regions, while most molecular clouds in the Milky Way and our simulations are forming or close to high-mass stars. The feedback from high-mass stars may lead to slightly higher velocity dispersions and lower $X_{\text{CO}}$. The nearby high-mass star-forming molecular cloud is Orion, and it does have a lower value of $X_{\text{CO}}$ than the Milky Way average (Figures 13 and 20).

For the numerical simulations, the main uncertainties lie in the assumptions of equilibrium chemistry and the subgrid model of microturbulence in calculating the CO emission. As a further test, we produced synthetic observations of model RES-2pc with half of the fiducial microturbulence velocity and no subgrid microturbulence (only thermal line-broadening on the grid scale), and found that the values of $X_{\text{CO}}$ increase by factors of 1.4 and 1.8. Therefore, the uncertainty in subgrid microturbulence may account for part but not all of the discrepancies in $X_{\text{CO}}$ between our simulations and observations. Future AMR simulations with higher numerical resolution and non-equilibrium chemistry will be able to provide more insight into these issues.

### 3.4. Dependence of $X_{\text{CO}}$ on the Observational Beam Size

Observations of molecular clouds often have different physical beam sizes/resolutions, which depend on the telescope as well as the distance of the object. In order to investigate the effect of observational resolution on $X_{\text{CO}}$, we smooth the synthetic observations to different beam sizes as described in Section 2.4.
X\textsubscript{CO} increases by a factor of \sim 2 as the beam size increases from \sim 1 pc to \sim 100 pc, as shown in Figure 21. This is a result of the CO-dark H\textsubscript{2}. The total CO emission remains the same as the beam size increases, because the detection limits for different beam sizes (Table 2) are generally sensitive enough to detect most of the CO emission. This is not surprising because the sensitivity in observations is designed to serve the purpose of accurately measuring the CO emission. However, the CO emission is smoothed out spatially as the beam size increases, resulting in a larger area of CO-bright regions. Although the total mass of H\textsubscript{2} remains the same, because X\textsubscript{CO} is calculated only within CO-bright regions, a larger area of CO-bright regions leads to a larger fraction of H\textsubscript{2} mass accounted for, and therefore an increase in X\textsubscript{CO}. This is clearly illustrated in Figure 22, showing the correlation between f\textsubscript{dark} and \langle X\textsubscript{CO} \rangle. From beam sizes of \sim 100 pc to \sim 1 kpc, some simulations show a continued increase in X\textsubscript{CO} (e.g., T-401Myr), but some simulations with more diffuse molecular clouds (e.g., T-381Myr) start to have part or all of their CO emission falling below the detection limits, leading to a non-detection of W\textsubscript{CO} or reduction of X\textsubscript{CO}. This suggests that some diffuse molecular clouds may not be detected with a beam size coarser than \sim 100 pc in extragalactic observations.

In Figure 21, we plot the observations of X\textsubscript{CO} in Milky Way molecular clouds and nearby galaxies (Table 1). Because of the large uncertainties in the observations (as discussed above, and also seen directly in the different X\textsubscript{CO} from two Perseus observations) and dispersions of X\textsubscript{CO} in different molecular clouds, we cannot identify any obvious trend for the variation of X\textsubscript{CO} with beam size. The
general range of $X_{\text{CO}}$ in the simulations is similar to the observations.

4. Summary

In this paper, we theoretically model the $X_{\text{CO}}$ conversion factor by postprocessing MHD galactic-disk ISM simulations with chemistry and radiation transfer to produce synthetic observations of molecular clouds. We conduct detailed analyses of the dependence of molecular abundances and observed line strengths on ISM conditions, and also consider numerical and observational effects on calculated and measured $X_{\text{CO}}$. Our main findings are as follows.

1. CO is only a very approximate tracer of $H_2$. In our simulations, most $H_2$ forms at intermediate densities $n \approx 10$–100 cm$^{-3}$, but most CO forms at higher densities $n \gtrsim 200$ cm$^{-3}$ (Figure 3). The $H_2$ abundance is determined mostly by density, but the CO abundance by dust shielding (Figures 4, 5). With a 2 pc numerical resolution, $H_2$ abundance is converged, but CO abundance is not. Although there is considerable scatter, the mean relation between the CO and $H_2$ column densities in the simulations is in agreement with observations of UV absorption spectra (Figure 7).

2. For CO emission, the high optical depth of the line further complicates the observable relation to $H_2$. On parsec scales, $W_{\text{CO}}$ is largely determined by the mean excitation temperature of CO (Figure 8), which is in turn determined by the mean gas density. Thus, $W_{\text{CO}}$ most directly probes the mean gas density along the line of sight. However, for the turbulent clouds in our simulations, the mass-weighted mean volume density along a line of sight tends to be correlated with column density. This leads to a correlation between $W_{\text{CO}}$ and $N_{H_2}$ (Figure 11).

3. A numerical resolution of at least 2 pc is needed in order to resolve the average $X_{\text{CO}}$ in molecular clouds for solar neighborhood conditions (Figure 12). In our simulations with environmental conditions similar to the solar neighborhood, we found $(X_{\text{CO}}) = (0.7$–$1.0) \times 10^{20}$ cm$^{-2}$ K$^{-1}$ km$^{-1}$ s, about a factor of 2 lower than the estimate from dust-based observations, and consistent with the $X_{\text{CO}}$ from $\gamma$-ray observations. The value of $(X_{\text{CO}})$ is not sensitive to the variations in molecular cloud mass, extinction, or the strength of the FUV radiation field (Table 5).

4. We found the CO-dark $H_2$ fraction $f_{\text{dark}} = 26$%–79%, which has an anticorrelation with the average extinction of molecular clouds (Figure 18 right panel).

5. The chemical timescale for $H_2$ abundance to reach equilibrium is longer than that for CO abundance (Figure 14), because of differences in characteristic densities. As a result, younger molecular clouds are expected to have lower $(X_{\text{CO}})$ values and flatter profiles of $X_{\text{CO}}$ versus extinction than older molecular clouds (Figures 16, 20).

6. As the observational beam size increases from $\sim 1$ to $\sim 100$ pc, $(X_{\text{CO}})$ increases by a factor of $\sim 2$, due to the decrease in the CO-dark $H_2$ fraction (Figures 21, 22).

7. Our numerical simulations successfully reproduce the observed variations of $W_{\text{CO}}$ on parsec scales, as well as the trends for the dependence of $X_{\text{CO}}$ on extinction and the CO excitation temperature. However, the value of $X_{\text{CO}}$ in our simulations is systematically lower by a factor of $\sim 2$ compared to dust-based observations (Figures 13, 19, 20).

The overall agreement between our numerical simulations and observations of Milky Way molecular clouds gives us confidence that similar simulations can be used to probe the $X_{\text{CO}}$ conversion factors in different environments, such as the Galactic center, low-metallicity dwarfs, and extreme star-forming systems (ultraluminous infrared galaxies and high-redshift galaxies). In a follow-up study, we will investigate the properties of individual molecular clouds in our simulations. In the future, we also plan to integrate full non-equilibrium chemistry with the MHD simulations.

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Appendix A
Definitions of CO-dark \( H_2 \)

In this paper, we define the CO-dark \( H_2 \) as the molecular gas without CO emission along a given line of sight. Wolfire et al. (2010) use a slightly different definition in their spherical cloud model, and they refer to the CO-dark \( H_2 \) as the molecular gas outside the optical depth \( \tau_{CO} = 1 \) surface. Their definition of CO-dark \( H_2 \) includes the \( H_2 \) along the line of sight in the projected CO-bright areas on the plane of the sky, as long as it is outside the \( \tau_{CO} = 1 \) surface (see their Figure 1). In other words, the definition of Wolfire et al. (2010) is in 3D physical space while our definition is in 2D observational space.

To compare the result from Wolfire et al. (2010) to our simulations, we need to translate their definition of CO-dark \( H_2 \) fraction, denoted by \( f_{DG} \) (their Equation (1)) to our definition denoted by \( f_{dark} \) (Equation (10) in this paper). Below we derive the relation between \( f_{DG} \) and \( f_{dark} \). We refer the reader to Figure 1 in Wolfire et al. (2010) for a useful illustration for this derivation.

From Equation (10), \( f_{dark} \) can be written as

\[
f_{dark} = \frac{M_{H_2} - M_{br}}{M_{H_2}} = 1 - \frac{M_{br}}{M_{H_2}} \tag{32}\]

where \( M_{H_2} \) is the total \( H_2 \) mass (same as \( M_{H_2,tot} \) in Equation (10)) and \( M_{br} \) is the mass in CO-bright areas on the projected sky. From Figure 1 in Wolfire et al. (2010), \( M_{br} = M_{CO} + M_{DG} \), where \( M_{CO} \) is the mass with \( r < R_{CO} \), and \( R_{CO} \) is the radius of the cloud where \( \tau_{CO} = 1 \). \( M_{DG} \) is the mass that lies within \( R_{CO} \) in the 2D projected sky, but outside \( R_{CO} \) in the 3D cloud. Compared to the definition in Wolfire et al. (2010),

\[
f_{DG} = 1 - \frac{M_{CO}}{M_{H_2}} \tag{33}\]

\( M_{DG} \) is the part of the cloud that Wolfire et al. (2010) considered to be CO-dark, but we do not.

Wolfire et al. (2010) assume that the cloud has a density profile \( n(r) = n_0(r_0/r) \), where \( n_0 \) and \( r_0 \) are constants. This gives

\[
M_{H_2} = \int_0^{R_{H_2}} 4\pi m_H n r^2 dr = 2\pi n_0 m_H r_0 R_{H_2}^2, \tag{34}\]

and similarly,

\[
M_{CO} = 2\pi n_0 m_H r_0 R_{CO}^2, \tag{35}\]

where \( m_H \) is the mass of the hydrogen atom. \( M_{DG} \) can be estimated from \( M_{DG} \approx 2\pi R_{CO} \Sigma_{DG} \), where \( \Sigma_{DG} = m_H \int_{R_{CO}}^{R_{H_2}} n dr = n_0 m_H r_0 \ln(R_{H_2}/R_{CO}) \). Therefore,

\[
M_{br} = M_{CO} + M_{DG} \approx 2\pi n_0 m_H r_0 R_{CO}^2 \left[ 1 + \ln \left( \frac{R_{H_2}}{R_{CO}} \right) \right]. \tag{36}\]

Figure 23. Comparison of the excitation temperature from the RADMC-3D radiation transfer code (scatter points with median values in density bins shown as the magenta line) to that from the analytical two-level system model (green line) for the simulation RES-1pc. Only the \( J = 0 \) and \( J = 1 \) rotational levels of CO are included in the calculations using RADMC-3D.

Appendix B
Test of the RADMC-3D Code

Figure 23 shows a test for the RADMC-3D radiation transfer code. The level populations of CO are solved with only the first two rotational levels instead of the default 41 levels. The analytical model uses Equations (27) and (28) to compute \( T_{exc} \) versus \( n \). Note that because \( T_{LVG} \) depends on level populations (see Equation (7)), the average values of \( T_{LVG} \) in this case are slightly larger than that given by Equation (26).

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