NON-EQUILIBRIUM CHEMISTRY OF DYNAMICALLY EVOLVING PRESTELLAR CORES. I. BASIC MAGNETIC AND NON-MAGNETIC MODELS AND PARAMETER STUDIES

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ABSTRACT

We combine dynamical and non-equilibrium chemical modeling of evolving prestellar molecular cloud cores and investigate the evolution of molecular abundances in the contracting core. We model both magnetic cores, with varying degrees of initial magnetic support, and non-magnetic cores, with varying collapse delay times. We explore, through a parameter study, the competing effects of various model parameters in the evolving molecular abundances, including the elemental C/O ratio, the temperature, and the cosmic-ray ionization rate. We find that different models show their largest quantitative differences at the center of the core, whereas the outer layers, which evolve slower, have abundances which are severely degenerate among different dynamical models. There is a large range of possible abundance values for different models at a fixed evolutionary stage (central density), which demonstrates the large potential of chemical differentiation in prestellar cores. However, degeneracies among different models, compounded with uncertainties induced by other model parameters, make it difficult to discriminate among dynamical models. To address these difficulties, we identify abundance ratios between particular molecules, the measurement of which would have maximal potential for discrimination among the different models examined here. In particular, we find that the ratios between NH_3 and CO, NH_2 and CO, and NH_3 and HCO^+ are sensitive to the evolutionary timescale, and that the ratio between HCN and OH is sensitive to the C/O ratio. Finally, we demonstrate that measurements of the central deviation (central depletion or enhancement) of abundances of certain molecules are good indicators of the dynamics of the core.

Key words: ISM: abundances – ISM: clouds – ISM: magnetic fields – ISM: molecules – stars: formation

1. INTRODUCTION

The earliest phases of star formation have been the subject of intense observational and theoretical investigation over the past four decades. Nevertheless, together with the very late stages of star formation (formation of planetary systems), the very early processes of molecular cloud fragmentation into high-density "cores" and the subsequent early contraction of these cores before protostars form in their centers remain the two least constrained and most debated aspects of star formation theory (McKee & Ostriker 2007). The reason for this lack of strong constraints is that it is difficult to extract physical conditions in star-forming regions from observable quantities, such as dust column or molecular line maps; each observable is typically affected by multiple factors which are nontrivial to deconvolve. As a result, theoretical models of cloud fragmentation and core evolution span a large range of possibilities.

A diverse array of mutually interacting microphysical processes influence the evolution of molecular clouds to differing extents: gravity, turbulence, magnetic fields, UV and cosmic-ray ionization, and a complex network of chemical reactions. Without strong observational constraints on the relative importance of these processes, competing star formation theories have arisen based on different assumptions of the physics driving the formation of prestellar cores and their collapse to form protostars.

The path to the appearance of a protostar depends sensitively on the initial conditions and the relative importance of the different physical processes driving the formation and contraction of prestellar molecular cloud cores. The most important distinction among different calculations of prestellar collapse pertains to the nature of any forces opposing gravity during the prestellar collapse. In this context we can identify two broad classes of models: non-magnetic and magnetic.

Non-magnetic collapsing models include early numerical and semi-analytic solutions of the Euler equations of hydrodynamic collapse (e.g., Larson 1969; Penston 1969; Yorke & Krügel 1977; Shu 1977), as well as numerical studies of the gravo-turbulent formation and evolution of bound condensations formed by converging flows in turbulent model clouds (see Mac Low & Klessen 2004 and Scalo & Elmegreen 2004 for recent reviews). The evolution of cores formed by turbulence-driven fragmentation is compatible with models of pure hydromagnetic collapse (e.g., Gong & Ostriker 2009). Turbulence-initiated core formation models can be generally separated into models of rapid global collapse, where the entire molecular cloud is viewed as an evanescent structure (e.g., Hartmann et al. 2001; Heitsch & Hartmann 2008), and models with some initial decaying turbulent support that extends the lifetime of clouds (e.g., Krumholz et al. 2006; Tan et al. 2006). Magnetic models include magnetohydrodynamical (MHD) calculations of the magnetically mediated contraction of prestellar cores, which is typically slower than the pure hydrodynamical collapse (with lower infall velocities and longer evolution timescales, e.g., Fiedler & Mouschovias 1993; Li &

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2 Note, however, that Shu’s singular isothermal sphere models refer to protostellar cores in which a central object exists, since the initial conditions for the inside-out collapse are singular at the center. Thus, this class of models cannot be used to study the early prestellar phase of the core evolution.
Shu 1996; Tassis & Mouschovias 2007), and result in a characteristic hourglass-like configuration of the magnetic field threading the core (see Mouschovias & Ciolek 1999 for a review). For magnetic models the lifetime of the parent molecular cloud is controlled by the initial value of the mass-to-magnetic-flux ratio. If this ratio is smaller than a critical value (the cloud is magnetically subcritical), then the cloud as a whole is long-lived as the magnetic field can support it against gravity. In this case, magnetically supercritical cores form through the process of ambipolar diffusion and collapse to form protostars, while the cloud envelope remains supported by the field. If the mass-to-magnetic-flux ratio of the parent cloud is larger than the critical value, then the cloud as a whole is unstable and short-lived, while the formation and collapse of cores also proceeds on shorter timescales.

The coupling between chemistry and dynamics is an essential step in connecting the dynamics of prestellar cores to observations and has long been recognized as critical in extracting information from molecular line observations. The problem has been pursued for the past decade in a series of groundbreaking studies. Bergin & Langer (1997) identified, through studies of static media, “early” and “late” molecules (molecules which peak in abundance either in the initial or in the later stages of the evolution of a molecular cloud fragment). Aikawa et al. (2001) compared observations of L1544 with coupled chemical/dynamical models. For the dynamics, they used the analytical similarity solution of Larson (1969) and Penston (1969), as well as a “slowed-down” version of it (using an overall slow-down factor) to approximate the slower magnetic collapse and found that their fast models are better fits to L1544. Li et al. (2002) and Shematovich et al. (2003) also modeled L1544, using a non-magnetic model and the Li (1999) magnetic model for the dynamics, which treats the magnetic field in an approximate way, including only magnetic pressure forces and assuming a spherical geometry. They also included grain-surface reactions in their calculations, and they found the magnetic model to be a better fit to the observations. They showed that the magnetic collapse is accelerating, and therefore a uniform slow-down of the non-magnetic collapse is not a good approximation because in magnetic collapse the amount of time spent at different densities changes with respect to the non-magnetic case. Aikawa et al. (2003) updated the Aikawa et al. (2001) calculation to include grain-surface reactions, and they found that moderate retardation factors (factor of three) are not in as strong disagreement with the data as in Aikawa et al. (2001); however, they still found that the faster models are preferred. Li et al. (2002) however noted that their preferred Larson–Penston solution features very large infall velocities that cannot be reconciled with observations of L1544. Lee et al. (2003) modeled the chemistry of various prestellar cores to fit Bonnor–Ebert and Plummer-like radial density profiles using dust continuum and line emission. Lee et al. (2004) looked at the chemical evolution of prestellar cores, aiming to produce line profiles for prestellar and protostellar cores. To model the dynamical evolution of prestellar cores they used a sequence of Bonnor–Ebert profiles, but the assignment of times and velocities to each profile was not treated self-consistently. For the protostellar cores they used the inside-out collapse model of Shu (1977). Flower et al. (2005), using a homologous collapse treatment of prestellar cores, emphasized the effect of grain growth on the chemistry. Aikawa et al. (2005) revisited the prestellar core problem, but instead of using an analytic solution for the dynamics, they used post-processing of a numerical simulation of the collapse of Bonnor–Ebert spheres (one very close to equilibrium and one very unstable), and found significant differences in the system chemistry depending on the initial conditions. This result was confirmed by Keto & Caselli (2008) who coupled one-dimensional dynamical simulations with a simple CO chemistry network and radiative transfer. In a follow-up study, Keto & Caselli (2010), starting from a Bonnor–Ebert sphere of central density $2 \times 10^4 \text{ cm}^{-3}$ in unstable equilibrium, evolving quasi-statically, found that at a density of $2 \times 10^7 \text{ cm}^{-3}$ the abundance of CO is much lower than observations in L1544, and argued for a higher CO desorption rate from grains.

On certain issues a consensus is forming among these studies (see, e.g., Bergin & Tafalla 2007 for a recent review). One such issue is the sensitivity of the chemical state of the contracting core to the duration of the collapse. Characteristic examples are the carbon-bearing molecules, which tend to become depleted on the ice mantles of dust grains and are therefore most prominent in the early stages of evolution of a core, in contrast to nitrogen-bearing molecules, which are not depleted and, as a result, their abundance increases in later stages.

However, core age is not the only contributing factor in determining molecular abundances. Observations have established the existence of chemical differentiation in prestellar cores of otherwise similar properties, which nevertheless show strong variations in the abundances of a number of chemical species. Characteristic examples of chemical differentiation can be seen in observations of prestellar cores L1544, L1498, L1517B, which show CO depletion (Tafalla et al. 2002, 2004), whereas L1521E (a core of similar central density with L1517B) shows no CO depletion (Tafalla & Santiago 2004). This differentiation may be a result of the dynamics of the collapse as suggested by, e.g., Shematovich et al. (2003), and thus reflect the initial conditions in the parent star-forming region.

In this paper, we self-consistently follow the chemistry and dynamics of contracting prestellar cores. We do so by adding to the system of equations of Eulerian hydrodynamics, or MHD (as appropriate), one continuity equation with sources and sinks given by the chemical reactions network for each species followed. In this way, the dynamics and non-equilibrium chemistry evolve simultaneously. This is particularly important in the case of MHD models, as abundances of charged species have a feedback effect on the dynamics, since the magnetic field couples only to charged molecules and grains, which in turn determine the evolution of the magnetic field and thus the dynamics of the contraction. In a companion publication (Tassis et al. 2012, hereafter Paper II), we discuss in more detail the interplay between chemistry and dynamics in magnetic models.

Our paper is organized as follows. In Section 2, we discuss our dynamical models (hydrodynamic and MHD) of core evolution. The chemical network we implement is discussed in Section 3. Details on the parameter study we have conducted are given in Section 4. We present our results in Section 5 and we discuss our findings and conclusions in Section 6.

2. DYNAMICAL MODELS OF MOLECULAR CLOUD CORES

2.1. Hydrodynamic Core Models

We model the contraction of a non-magnetic, self-gravitating, isothermal prestellar core in spherical symmetry. The core is assumed to be embedded within a uniform-density isothermal
cloud. We solve the equations of hydrodynamics in spherical symmetry:

\[
\frac{\partial \rho}{\partial t} + \frac{1}{r^2} \frac{\partial (r^2 \rho v_r)}{\partial r} = 0, \tag{1a}
\]

\[
\frac{\partial \rho_j}{\partial t} + \frac{1}{r} \frac{\partial (r \rho_j v_r)}{\partial r} = S_j, \tag{1b}
\]

\[
\frac{\partial (\rho v_r)}{\partial t} + \frac{1}{r^2} \frac{\partial (r^2 \rho v_r^2)}{\partial r} = -C_s^2 \frac{\partial \rho}{\partial r} + \rho g_r, \tag{1c}
\]

\[
g_r = -\frac{4\pi G}{r^2} \int_0^r \rho r^2 dr', \tag{1d}
\]

where \(v_r\) is the radial velocity, \(\rho\) is the total density of the gas, and \(\rho_j\) is the density of each individual chemical species considered (listed in Table 1). We solve a density continuity equation for each of the chemical species in the model (Equation (1b)). In these equations, \(S_j\) are source/sink terms describing mass interchange between the species due to chemical reactions.

The initial state is one of uniform density equal to \(10^3 \text{ cm}^{-3}\) and zero velocity. The radius of the core is 0.4 pc, making it significantly thermally supercritical. At the outer boundary, we require constant pressure and zero velocity. This way, the core experiences no influx of mass.

### 2.2. Magnetohydrodynamic Core Models

We follow Basu & Mouschovias (1994) in modeling the formation and contraction of prestellar cores in self-gravitating, magnetic, rotating, isothermal model clouds in the presence of ambipolar diffusion and magnetic braking. In the presence of an ordered magnetic field the cloud relaxes rapidly along field lines, until thermal-pressure forces balance gravity, before any significant contraction perpendicular to the field lines takes place (Fiedler & Mouschovias 1993; Desch & Mouschovias 2001; Kunz & Mouschovias 2010). Force balance along magnetic field lines is maintained even when the contraction becomes dynamic in the perpendicular direction. Consequently, the thin-disk approximation is adopted for the model cloud cores (Ciolek & Mouschovias 1993). The model cloud is axisymmetric with the rotation axis coinciding with the axis of symmetry, which is taken to be the \(z\)-axis of a cylindrical polar coordinate system \((r, \phi, z)\). The upper and lower surfaces of the disk are located at \(z = \pm Z(r, t)\). The cloud is embedded in an “external” medium of constant thermal pressure \(P_{\text{ext}}\), and density \(\rho_{\text{ext}}\), and is threaded by a magnetic field which, far from the cloud, becomes uniform and equal to a “reference” value \(B_{\text{ref}}\).

The thin-disk evolution equations, obtained by integrating the two-fluid (plasma and neutrals) MHD equations over \(z\), are

\[
\frac{\partial (\rho v_r)}{\partial t} + \frac{1}{r} \frac{\partial (r \rho v_r^2)}{\partial r} = -\frac{\partial \rho L}{\partial r} + \frac{1}{r} \frac{\partial (r \rho L v_r)}{\partial r} = 0, \tag{2a}
\]

\[
\frac{\partial \sigma}{\partial t} + \frac{1}{r} \frac{\partial (r \sigma v_r)}{\partial r} = 0, \tag{2b}
\]

\[
\frac{\partial (\rho v_r)}{\partial t} + \frac{1}{r} \frac{\partial (r \rho v_r^2)}{\partial r} = -C_s^2 \frac{\partial \rho}{\partial r} + \rho g_r, \tag{2c}
\]

\[
g_r = -\frac{4\pi G}{r^2} \int_0^r \rho r^2 dr', \tag{2d}
\]

\[
\frac{\partial (\rho v_r)}{\partial t} + \frac{1}{r} \frac{\partial (r \rho v_r^2)}{\partial r} = -C_s^2 \frac{\partial \rho}{\partial r} + \rho g_r, \tag{2e}
\]

\[
\frac{\partial (\rho v_r)}{\partial t} + \frac{1}{r} \frac{\partial (r \rho v_r^2)}{\partial r} = -C_s^2 \frac{\partial \rho}{\partial r} + \rho g_r, \tag{2f}
\]

\[
\frac{\partial \rho L}{\partial t} + \frac{1}{r} \frac{\partial \rho L v_r}{\partial r} = \frac{1}{2\pi} \int_0^{\pi} \rho d\theta Z(r, \theta), \tag{2g}
\]

\[
\frac{\partial \rho L}{\partial t} + \frac{1}{r} \frac{\partial \rho L v_r}{\partial r} = \frac{1}{2\pi} \int_0^{\pi} \rho d\theta Z(r, \theta), \tag{2h}
\]

\[
B_{\text{ref}}(r, \phi, z) = -\frac{2}{r} \frac{(4\pi \rho_{\text{ext}})^{1/2} \Phi}{\rho_{\text{ext}}} (\Omega - \Omega_0), \tag{2i}
\]

\[
B_{\phi, z}(r) = -\frac{2}{r} \frac{(4\pi \rho_{\text{ext}})^{1/2} \Phi}{\rho_{\text{ext}}} (\Omega - \Omega_0), \tag{2j}
\]

\[
B_{\phi, z}(r) = -\frac{2}{r} \frac{(4\pi \rho_{\text{ext}})^{1/2} \Phi}{\rho_{\text{ext}}} (\Omega - \Omega_0), \tag{2k}
\]

\[
\mathcal{M}(r, r') = \frac{d}{dr} \left[ \int_0^r dk J_0(kr) J_0(kr') \right] = \frac{2}{\pi} \frac{d}{dr} \left[ \frac{1}{r} K_0 \left( \frac{r}{r'} \right) \right], \tag{2l}
\]
from the algebraic solution of the force equation for the ion fluid (we are studying) advected with the velocity of the ions. Grains are overwhelmingly negatively charged in the densities for the gas-phase ions and all the grain mantle species (since dust flux tube balances the external and gravitational pressures. The thermal pressure at the equatorial plane in each magnetic terms are corrections due to the variation of the half-thickness of the cloud, $Z$, with $r$. Equation (2d) is Faraday’s law in the two-fluid approximation with the magnetic flux frozen in the ion fluid. The toroidal component of the magnetic field ($B_{\phi,Z}$) is determined assuming a steady state condition on the transport of angular momentum by torsional Alfvén waves in the external medium and given by Equation (2c). The radial component of the magnetic field at the top surface of the thin disk ($B_{r,Z}$) is given by Equation (2e), obtained by assuming a force-free external medium, whereby the half-thickness of the disk ($Z$) is obtained from Equation (2f) after $\rho_{0}(r,t)$ is calculated from Equation (2e). The $r$-component of the gravitational field (2k) is calculated from Poisson’s equation for a thin disk. $J_{0}$ is the zeroth-order Bessel function of the first kind, $K$ is the complete elliptic integral of the first kind, $r_{c} = \min[r,r']$, and $r_{s} = \max[r,r']$. Equation (2m) is the mass continuity equation for neutral gas-phase chemical species advected with the neutral bulk velocity $v_{n,r}$. Equation (2n) is the mass continuity equation for the gas-phase ions and all the grain mantle species (since dust grains are overwhelmingly negatively charged in the densities we are studying) advected with the velocity of the ions $v_{i,r}$. $\nabla\sigma$ and $\nabla v_{i,r}$ represent the source/sink terms due to chemical reactions. Note that the ions are advected with a different velocity than the neutrals, so two equations are required in the magnetic model to replace Equation (1b) of the non-magnetic model.

Equation (2o) gives the velocity of the ions and is obtained from the algebraic solution of the force equation for the ion fluid where the acceleration terms have been neglected because of the small inertia of these species. The mean (momentum exchange) collision time of a neutral particle with ions in Equation (2o) is

$$\langle \tau_{\text{nl}} \rangle = \frac{1.56 \langle \mu_{i} \rangle m_{p} + 2m_{p}}{\rho_{i} (\sigma w)_{\text{in}}},$$

where $m_{p}$ is the proton mass, $\rho_{i} = \sum_{j+} \rho_{j+}$ is the total ion density, the mean molecular weight of the ions is

$$\langle \mu_{i} \rangle = \sum_{j+} m_{j+} n_{j+} / \sum_{j+} n_{j+},$$

where $\sigma_{n,\text{ref}}(r)$ is the number volume density for each one of the ions considered, and $(\sigma w)_{\text{in}}$ is the average collisional rate between ions and hydrogen molecules. The value of $(\sigma w)_{\text{in}}$ varies slightly with the mass of the ion (McDaniel & Mason 1973), we use the canonical value $(\sigma w)_{\text{in}} = 1.69 \times 10^{-9}$ cm$^{3}$ s$^{-1}$ (Ciolek & Mouschovias 1993).

Each magnetic model is initiated at a reference state as in Basu & Mouschovias (1994). The reference state has a column density profile of the form

$$\sigma_{n,\text{ref}}(r) = \frac{\sigma_{n,\text{ref}}}{\left[1 + (r/l_{\text{ref}})^{2}\right]^{3/2}},$$

which implies an almost uniform column density for $r \ll l_{\text{ref}}$, which is where the core forms. The high-$r$ tail has little effect on the calculations and is included for numerical convenience, allowing the calculation of the disk gravitational potential to converge. The profile of the reference state angular velocity is obtained by using a linear dependence of specific angular momentum on mass, corresponding to the angular momentum distribution of a uniformly rotating disk, yielding

$$\Omega_{\text{ref}}(r) = 2\Omega_{c,\text{ref}} \left(\frac{l_{\text{ref}}}{r}\right)^{2} \left[1 - \frac{1}{\sqrt{1 + (r/l_{\text{ref}})^{2}}}ight].$$

The reference state is first allowed to relax to an equilibrium configuration, whereby ambipolar diffusion and chemistry are turned off. It is from this equilibrium state that the subsequent evolution is initiated. For this reason, the initial central density to which each model relaxes is not always the same, but it depends on the initial mass-to-flux ratio (see Figure 1).
The free parameters in the magnetic models are the central density of the initial reference state (which we take to be $10^{3} \text{ cm}^{-3}$); the background angular velocity $\Omega_{b} = 10^{-15} \text{ rad s}^{-1}$, approximately corresponding to the rate of galactic rotation in the solar neighborhood; the reference state angular velocity ($\Omega_{\text{ref}} = 0.48 \text{ km s}^{-1} \text{ pc}^{-1}$); the external pressure $P_{\text{ext}} = 0.1 \pi G \sigma_{\text{ref}}^2 / 2$ corresponding to 10% of the central “gravitational” pressure; the characteristic size of the column density profile $l_{\text{ref}} = 1 \text{ pc}$; and the external density $\rho_{\text{ext}} = 10 \text{ cm}^{-3}$, and the mass-to-magnetic-flux ratio, which is varied (see discussion of parameter study below).

With the exception of the magnetic mass to flux ratio, the dynamical evolution of the collapsing core is not sensitive to any of the free parameters (Basu & Mouschovias 1995).

At the outer boundary of the disk we require continuity of thermal as well as magnetic pressures (see Section 5 of Tassis & Mouschovias 2005). No influx of mass is allowed at the outer boundary, so the total mass of the cloud is conserved.

### 3. CHEMICAL NETWORK

We use a subset of the UMIST chemical reaction network (Woodall et al. 2007), which includes data on 420 species and 13 elements followed in gas-phase reactions. Each chemical species is treated as a separate fluid in our formalism. The source and sink terms on the right-hand side of Equation (1b) and in Equations (2m) and (2n) describe the interactions between chemical species. They are given by

$$S_{j} = \frac{\partial \rho_{j}}{\partial t} = \sum_{k \neq j, l \neq j} \alpha_{klj} \rho_{k} \rho_{l} - \sum_{k \neq j, l \neq j} \beta_{klj} \rho_{l} + \sum_{j \neq k} (\beta_{jk} \rho_{k} - \beta_{kj} \rho_{j}),$$

(6)

where $\alpha_{klj}$ is the rate coefficient for the production of species $j$ due to the interaction of species $k$ and $l$, $\beta_{jk}$ is the rate of production of species $j$ from species $k$ by photoreactions, and $\delta_{ij}$ is the Kronecker delta function.

In our models the source terms are implemented in an operator-split fashion, and the change of abundances due to the source terms is updated after the advection terms. The species we consider are listed in Table 1, and they include 78 gas-phase atomic and molecular species, and 33 grain ice mantle species, which are composed of the elements H, He, C, N, O, and Si as a representative low ionization potential metal. The chemical network follows 1553 reactions.

All the models studied in this work use the isothermal approximation (small deviations from isothermality have been shown to not affect the dynamical evolution of cores significantly, e.g., Keto & Caselli 2010). We adopt as a fiducial temperature $T = 10 \text{ K}$, consistent with line intensity ratios of different molecular transitions showing that starless cores are almost isothermal, with $T \sim 10 \text{ K}$ (Tafalla et al. 1998, 2002, 2004), although there may be small variations of the order of a few kelvin (e.g., Evans et al. 2001). In our parameter studies, we test for the effect of small temperature variations on our chemical abundances.

Since the prestellar cores we model are embedded in a molecular cloud, we assume a visual extinction $A_V = 3 \text{ mag}$ at the outer boundary, so that the results are not much affected by photodissociation. For the inner layers of the core we obtain the visual extinction from the column density to the surface using the empirical conversion relation $N_{H_2} / A_V = 9.4 \times 10^{20} \text{ cm}^{-2} \text{ mag}^{-1}$ (Bohlin et al. 1978). For our deeply embedded cores, no $H_2$ self-shielding is required; CO self-shielding on the other hand is accounted for based on the model of Lee et al. (1996).

We assume a standard MRN distribution of grains (Mathis et al. 1977), assumed to be well mixed with the gas. Since most of the grains are negatively charged at the densities of interest in prestellar cores, we also take into account the electrostatic attraction between ions and grains through an enhanced rate of accretion for the ions (Rawlings et al. 1992). We use a fixed grain abundance ($n_{\text{grain}} / n_{\text{neutral}} = 10^{-12}$) and charge (we take all grains to be singly negatively charged). Grain processes modeled include freezeout, surface reactions, thermal desorption, and cosmic-ray heating. We have used a sticking probability of 0.3.

The discrete nature of grains means that chemistry on their surfaces should properly be modeled by stochastic methods (e.g., Stancheva et al. 2002; Chang et al. 2007). However, those methods are computationally very expensive, especially for a large chemical network and the large number of hydrodynamic and MHD models we run. Instead, we use the rate equations method, with modifications as suggested by Vasyunin et al. (2009), which brings the results of this method into good agreement with Monte Carlo modeling for the range of temperatures we are concerned with. The grain reaction network is taken from Hasegawa & Herbst (1993), as are the rates for desorption by cosmic-ray heating of grains (updated for new binding energies where necessary). Where available, we use binding energies determined in the lab (see Willacy 2007). Other binding energies are taken from Hasegawa & Herbst (1993). The CO$_2$ chemistry on the grains is based on Garrod & Pauly (2011).

The gas-phase reactions are better constrained as they have been studied for a longer time, and there appears to be convergence in the literature among different groups (e.g., Aikawa et al. 2003) with the exception of sulphur-bearing molecules (Wakelam et al. 2006), which, however, are not considered in our model. In our study we aim to assess the effect of the dynamics on chemical abundances, while the chemical network remains fixed.

We use low-metal initial abundances given in Table 2. The calculations are initiated with all elements other than molecular hydrogen in the form of atoms. Of course in nature there is some chemical evolution taking place during the formation of the parent cloud. A first attempt to calculate initial abundances for molecular clouds for one particular cloud formation scenario was presented by Hassel et al. (2010). However, these results are model dependent and will differ for different cloud formation types.

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Table 2: Initial Abundances

| Species | Abundance |
|---------|-----------|
| H       | $1.00 \times 10^{-2}$ |
| H$_2$   | $4.95 \times 10^{-1}$ |
| He      | $1.40 \times 10^{-1}$ |
| C       | $7.30 \times 10^{-5}$ |
| N       | $2.14 \times 10^{-5}$ |
| O       | $1.76 \times 10^{-4}$ |
| Si      | $2.00 \times 10^{-8}$ |

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5 The activation barrier for the CO + O → CO$_2$ reaction on the grain ice was, however, increased from 290 K to 340 K (within the experimental range).

4 An exception is Si, the abundance of which is higher than in the low-metal model.
models. We will return to the issue of initial abundances in a future publication.

4. PARAMETER STUDY

For each class of models (non-magnetic or magnetic), we examine the dependence of the molecular abundances and their evolution on four parameters: the C/O ratio, the cosmic-ray ionization rate, the temperature, and a parameter controlling the time available for chemical evolution. In non-magnetic models, the latter is a time interval prior to collapse during which chemistry evolves but the core is assumed to not evolve dynamically. In magnetic models, it is the initial mass-to-magnetic-flux ratio which controls the amount of magnetic support: a smaller initial mass-to-magnetic-flux ratio implies a larger amount of magnetic support against gravity, and hence a slower dynamical evolution for the core.

Our “reference” non-magnetic model has a C/O ratio of 0.4, a cosmic-ray ionization rate of \( \zeta = 1.3 \times 10^{-17} \, \text{s}^{-1} \), a temperature \( T = 10 \, \text{K} \), and an initial collapse delay time of 1 Myr. Our “reference” magnetic model similarly has a C/O ratio of 0.4, \( \zeta = 1.3 \times 10^{-17} \, \text{s}^{-1} \), \( T = 10 \, \text{K} \), and a central mass-to-flux ratio equal to the critical value for collapse (Mouschovias & Spitzer 1976).

For non-magnetic models, we examine two additional values of delay: 0 and 10 Myr. For magnetic models, we examine two additional values of the initial mass-to-magnetic-flux ratio: 1.3 times the critical value (a faster-evolving, magnetically supercritical model) and 0.7 of the critical value (a slower, magnetically subcritical model). The three values of the delay and the three values of the mass-to-flux ratio set our dynamical model variations. When these models are combined with reference values for the C/O ratio, \( \zeta \), and \( T \), we will refer to them as the six basic dynamical models.

The carbon-to-oxygen ratio is varied from its reference value by keeping the abundance of C constant and changing that of O (as in Terzieva & Herbst 1998). The two other values of C/O ratio examined are 1 and 1.2. We have varied each of the six basic dynamical models by changing its C/O ratio to one of the values above, and we have thus studied 12 models (6 magnetic and 6 non-magnetic) with the C/O ratio varied from its reference value.

We have also considered different temperatures for each of the six basic dynamical models. We have varied the temperature by a factor of \( \sim 1.5 \) from its reference value of 10 K and examined models with \( T = 7 \, \text{K} \) and \( T = 15 \, \text{K} \), and we have thus studied 12 models (6 magnetic and 6 non-magnetic) with a temperature differing from its reference value.

Finally, to test the effect of the cosmic-ray ionization rate we have studied two additional magnetic and two non-magnetic models, which are, in each case, identical to the corresponding “reference” models, but with different cosmic-ray ionization rates, namely, a factor of four above (\( \zeta = 5.2 \times 10^{-17} \, \text{s}^{-1} \)) and below (\( \zeta = 3.3 \times 10^{-18} \, \text{s}^{-1} \)) its reference value (covering the range of observational estimates, e.g., McCall et al. 2003; Hezareh et al. 2008).

Thus, we have examined a total of 34 models (6 basic dynamical models, 12 with varied C/O ratios, 12 with varied \( T \), and 4 with varied \( \zeta \)). These 34 models are depicted in our figures in Section 5 with different line types, thicknesses, colors, and symbols, as follows.

1. Line type (solid, dashed, dotted) and symbols (X, diamond, cross) depict the evolutionary timescale (“reference,” “slow,” and “fast,” respectively).

2. Color denotes the type of model (magnetic or non-magnetic), with bluish hues corresponding to non-magnetic models, and reddish hues corresponding to magnetic models.

3. The line thickness/symbol size (for red and blue lines and symbols) indicate the values of the cosmic-ray ionization rates: thin red/blue solid lines and small red/blue symbols correspond to the low value of \( \zeta \), while thick red/blue solid lines and large red/blue symbols to the high value of \( \zeta \), with all other parameters fixed at their “reference” values.

4. The effect of different C/O ratio values is shown as a shaded area (orange for magnetic runs, and cyan for non-magnetic runs). The edge of the shaded area in contact with the corresponding (“fast,” “reference,” or “slow,” magnetic or non-magnetic) model line corresponds to \( \text{C/O} = 0.4 \), while the edge of the shaded area furthest away from the line corresponds to \( \text{C/O} = 1.2 \).

5. Finally, the effect of the temperature is shown with the colors brown for magnetic runs and purple for non-magnetic runs, either as a shaded area, or as line thickness/symbol size. In the latter case, the thick brown/purple lines (or the large brown/purple symbols) correspond to \( T = 15 \, \text{K} \) and the thin brown/purple lines (small brown/purple symbols) to \( T = 7 \, \text{K} \).

5. RESULTS

5.1. Dynamics

Figure 1 shows the evolution of the central density of contracting core models with time. Red lines correspond to magnetic runs and blue lines to non-magnetic runs. The different magnetic runs correspond to different values of the mass-to-magnetic-flux ratio: critical for the solid lines/\( \times \) symbols; 0.7 times the critical value (magnetically subcritical, slow-evolving magnetic model) for the dashed lines/diamonds; and 1.3 times the critical value (magnetically supercritical, faster-evolving magnetic model) for the dotted lines/+ symbols. The different non-magnetic runs correspond to different assumed delay times, during which the chemistry operates, before the core is allowed to evolve dynamically (collapse): 1 Myr delay for the solid lines/\( \times \) symbols; 10 Myr delay (slowest-evolving non-magnetic model) for the dashed lines/diamonds; and no delay (fastest-evolving non-magnetic model) for the dotted lines/+ symbols. The symbols mark the times when full radial outputs are obtained and analyzed, and correspond to successive enhancements of the central density by an order of magnitude.

Different dynamical models differ with respect to the two fundamental ingredients affecting the chemical evolution of a core: density, and time spent at each density. Non-magnetic models with substantial time delays (representing the time it takes for some level of turbulent support to dissipate) can take as much time as substantially magnetically subcritical models to reach very high densities. However, once the collapse sets in (support dissipates), the evolution occurs essentially within one free-fall timescale, so that the bulk of the time delay is spent at the initial, low density (\( 10^3 \, \text{cm}^{-3} \)). In contrast, slow magnetic models allow the core to spend substantial time at higher densities, and as a result the chemical state of these models are expected to be different.

The thin and thick red lines correspond to magnetically critical models but have \( \zeta \) a factor of four above and below the “reference” value, and they bracket the normal-thickness red solid line. Because the ambipolar diffusion timescale depends
on the degree of ionization, which in turn depends on the cosmic-ray ionization rate, $\zeta$, affects the evolutionary timescale of the magnetic models: a higher degree of ionization implies increased coupling between magnetic field and neutral fluid, and a resulting slower evolution, as magnetic forces are more effective in opposing gravity (see Paper II).

In addition to the mass-to-flux ratio and the degree of ionization, the evolutionary timescale of the magnetic models is also affected by the C/O ratio. The effect of different values of the C/O ratio can be seen in Figure 1 as the orange shaded area, which denotes a range of C/O values between 0.4 (reference) and 1.2. The reason for this dependence is that the C/O value affects the chemical reaction network and consequently the degree of ionization at each density, resulting in a different evolution rate. In general, higher values of the C/O ratio result in slower evolution, and the effect is more pronounced for models with a low mass-to-flux ratio.

Finally, the evolutionary timescale of the magnetic models also depends on the temperature. The effect of different $T$ values is shown in Figure 1 as the brown shaded area, which denotes a range of $T$ values between 7 K and 15 K. The dependence is stronger for slower models, and it enters through the dependence of the degree of ionization on temperature (see Paper II). The evolutionary timescale increases with increasing temperature, and the difference between 10 K and 15 K is significantly larger than the difference between 7 K and 10 K, although the fractional increase is comparable (about 50% in each case).

Radial profiles of the density, velocity, and (edge-on for the magnetic runs) column density of the magnetic and non-magnetic reference models and for the different time snapshots corresponding to the $\times$ symbols in Figure 1 are given in Figure 2. Again the magnetic model is represented by red lines and the non-magnetic model by blue lines. Each snapshot corresponds to a time when the volume density is an order of magnitude higher than the previous one. Time elapsed per snapshot in magnetic runs: 3.53 Myr, 4.4 Myr, 4.6 Myr, 4.65 Myr; in non-magnetic runs: 1.88 Myr, 2.01 Myr, 2.05 Myr, 2.06 Myr.

Figure 2. Radial profiles of the number volume density $n$ (panel (a)), velocities of the neutrals in units of the isothermal sound speed (panel (b)), and (edge-on for the magnetic runs) column density (panel (c)). Red lines: magnetic runs; blue lines: non-magnetic runs. Each curve corresponds to a snapshot in time when the volume density is an order of magnitude higher than the previous one. Time elapsed per snapshot in magnetic runs: 3.53 Myr, 4.4 Myr, 4.6 Myr, 4.65 Myr; in non-magnetic runs: 1.88 Myr, 2.01 Myr, 2.05 Myr, 2.06 Myr.

5.2. Non-equilibrium Chemistry, Production, and Destruction of Commonly Studied Molecules

The need for the inclusion of non-equilibrium chemistry in our dynamical models is explicitly demonstrated in Figure 3. Here we plot the ratio $P/D$ between the total production and
destruction rates of various molecules, as a function of density, at the center of our reference core models. The left panel corresponds to the magnetic reference model, and the right panel to the non-magnetic reference model. The total production and destruction rates are defined as including all chemical processes leading to the production or destruction of a specific molecule, respectively. Each line corresponds to a different molecule. Note that the y-axis in the two plots is logarithmic and extends over 10 orders of magnitude. A large number of molecules are very much out of equilibrium \( P/D \) is very far from 1, and they remain so throughout each simulation. In the magnetic run, where more time is available for the system to approach chemical equilibrium even at high densities, the majority of molecules eventually move closer to \( P/D = 1 \); in the non-magnetic model the reverse trend is observed, and as the density increases and the evolution timescale decreases, most molecules deviate further away from \( P/D = 1 \). However, even for the magnetic model the spread about \( P/D = 1 \) is substantial, as demonstrated by the inset on the left panel, which zooms in around central densities of \( 10^5-10^6 \) particles per cm\(^3\) and \( P/D \) within 50% of unity displayed on a linear scale. In the inset, the values of \( P/D \) exhibit a spread of more than 20% around unity.

The effect of non-equilibrium chemistry becomes obvious when we examine the dependence of central molecular abundances on central density for different dynamical models. If the chemistry was in equilibrium, then the central volume density would uniquely determine the central molecular abundances. This, however, is not the case, as we can see in Figure 4, where we examine the central abundances, production, and destruction rates due to various reactions for one carbon-bearing (CO) and one nitrogen-bearing (NH\(_3\)) molecule. The left column corresponds to the non-magnetic reference model (1 Myr delay), and the right column corresponds to the magnetic reference model (critical mass-to-flux ratio). The top row shows the evolution with central density (bottom axis) and with time (top axis) of the central abundances of CO (blue lines) and NH\(_3\) (purple lines). Solid lines correspond to gas-phase abundances and dashed lines correspond to abundances on grain ice mantles. The middle and bottom rows show the most important production (black lines) and destruction (red lines) reactions for the two molecules. The importance of each reaction is quantified by the percentage by which the specific reaction contributes to the total production/destruction rate of each molecule (y-axis).

Comparing the top left and right panels, we can immediately see that the central abundances of the two molecules at the same central density are very different for different dynamical models, a result of the chemistry being out of equilibrium. Because each model spends different amounts of time at each density (see Figure 1), non-equilibrium chemistry results in significantly different central abundances. Note that since the magnetic model evolves slower, the gas-phase abundances of both molecules in the magnetic core model decrease more steeply with density, as the molecules freeze out onto the grain mantles; correspondingly, there is a faster increase in the ice mantle abundance of each molecule.

It is evident from the middle and bottom panels that any specific reaction reaches its maximum importance at different densities for different dynamical models. Freezeout onto grain mantles (solid red line), which becomes increasingly important at higher densities, is a prominent example. Furthermore, for each molecule different reactions dominate the production and destruction channels for different dynamical models.

5.3. Evolution of Central Abundances

Figures 5 and 6 show the evolution with central density of the central abundances of selected molecules in the gas phase. Figure 5 shows the effect of the dynamics, C/O ratio, and \( \zeta \) on the gas-phase abundances, whereas Figure 6 shows the effect of the temperature.

In Figure 5 reddish hues correspond to magnetic models and bluish hues correspond to non-magnetic models. Different line types correspond to different dynamical models, with dashed lines representing the “slowest” models (subcritical magnetic model and 10 Myr delay non-magnetic model), solid lines representing the “intermediate” models (critical magnetic model and 1 Myr delay non-magnetic model), and dotted lines representing the “fast” models (supercritical magnetic model and no-delay non-magnetic model). The shaded bands (cyan for non-magnetic models, orange for magnetic models) show the effect of the C/O ratio on the abundances (the edge of the band adjacent to a model line corresponds to C/O = 0.4 and the edge furthest away from the line to C/O = 1.2). The line thickness is representative of the value of the cosmic-ray ionization rate, with normal-thickness lines corresponding to \( \zeta = 1.3 \times 10^{-17}\text{ s}^{-1} \).
Figure 4. Top panels: central abundances for two commonly observed molecules (CO and NH$_3$), for the non-magnetic (left column) and magnetic (right column) reference models. Solid lines: gas-phase abundance; dashed lines: grain ice mantle abundance. Middle and bottom panels: percentage contribution of most important reactions to the total production (black lines) and destruction (red lines) rates for the two molecules.

A general feature of the central gas-phase abundance evolution curves of Figure 5 is their dependence on the dynamical evolution model. Magnetic models, which evolve more slowly between different central densities and spend more time at each value of the density, have an abundance peak at low densities and generally exhibit a faster drop of the gas-phase abundance with increasing central density, due to efficient freezeout onto grain ice mantles. As a result, the abundance evolution curves appear convex. By contrast, non-magnetic models evolve faster, chemistry has less time in which to operate at each density, the abundances peak and drop at higher densities, and the non-magnetic abundance evolution curves are generally concave. The quantitative explanation of this behavior can be given by comparing the dependence of the freezeout timescale and the collapse timescale on density for different models. The steep drop in molecular abundances (especially in carbon-bearing molecules) occurs at the density where the freezeout timescale becomes shorter than the collapse timescale (Walmsley 1991). In magnetic models this happens at much lower densities ($\sim 10^4$ cm$^{-3}$) than in non-magnetic models ($\sim 10^6$ cm$^{-3}$).

The results of Figure 5 make it obvious that different collapse delay times at the initial phases of non-magnetic core evolution are not a good proxy for the effect of magnetic fields. Although the slowest non-magnetic model curves (dashed curves, 10 Myr delay) are generally closer to the magnetic model curves, the difference is still significant even in the qualitative behavior of the abundances (concave versus convex). The reason for this is that allowing more time for chemical evolution at $\sim 10^3$ cm$^{-3}$ does not have the same effect, from a chemistry perspective, as spending a similar amount of time at higher densities, which is what happens in magnetic models.

The behavior of certain molecules, such as H$_2$CO, H$_3^+$, NH, and CH$_3$, is quite different for the non-magnetic models. Whereas the slowest non-magnetic and all the magnetic models have a qualitative similar behavior that resembles that of other molecules (the abundance builds up with increasing density, peaks at $\sim 10^3$ cm$^{-3}$, and then decreases at even higher density), the faster non-magnetic models exhibit a local minimum at these densities, instead of a peak.

The molecules with the greatest sensitivity to the C/O ratio are, unsurprisingly, the oxygen-bearing molecules, the abundance of which is generally lower for higher C/O ratios. Other molecules sensitive to the C/O ratio include CH, CH$_3$, and NH.
Figure 5. Central abundances for representative molecules as a function of central density for different dynamical models, different C/O ratios, and different cosmic-ray ionization rates. Reddish hues correspond to magnetic models (dashed: magnetically subcritical; solid: magnetically critical; dotted: magnetically supercritical), and bluish hues to non-magnetic models (dashed: 10 Myr delay; solid: 1 Myr delay; dotted: no delay). The thin and thick solid lines correspond to cosmic-ray ionization rate $\zeta$, a factor of four above and below the “reference” value. The orange/cyan shaded areas show the effect of a varying C/O ratio in magnetic/non-magnetic models and correspond to a range of C/O values between 0.4 (reference value) and 1.2.

As expected, H$_3^+$ responds directly to the change in the cosmic-ray ionization rate, as it is a direct ionization product (see Paper II for more details). Molecules that are primarily produced through reactions involving direct ionization products, such as H$_3^+$ and He$^+$ (e.g., CH) or free electrons (e.g., NH$_2$, NH, and CH$_3$), are similarly affected, even in the non-magnetic models. The magnetic models have an additional response to $\zeta$ through the modification of their evolutionary timescale (see Figure 1). For this reason, the change in molecular abundances, especially in the case of the largest-$\zeta$ magnetic model (which is also by far the slowest-evolving model), can be dramatic and feature very significant depletion in relatively low densities. The biggest difference is seen in molecules which, for moderate evolutionary timescales, show relatively modest depletion (e.g., N-bearing molecules).

Figure 6 shows the effect of the temperature on the evolution with density of the central abundances of a few representative molecules. The blue and red lines correspond to the reference
non-magnetic and magnetic models (the same ones shown with the normal-thickness blue and red lines in Figure 5). High-temperature ($T = 15$ K) models are shown with the thick brown and purple lines, for magnetic and non-magnetic models, respectively, while low-temperature ($T = 7$ K) models are shown with the thin brown and purple lines; all other parameters are kept at their reference values.

There are some significant differences between the molecular abundances in the $15$ K models compared to those in the $10$ K models, especially for nitrogen-bearing compounds. This is a result of two effects: (1) The reaction

$$\text{N}^+ + \text{H}_2 \rightarrow \text{NH}^+ + \text{H}$$

(7)

has a small activation barrier of $85$ K. At $15$ K the temperature is high enough for this barrier to be overcome and hence this reaction is much more efficient than at $10$ K (see also Wakelam et al. 2010). This reaction is the start of the formation of nitrogen-bearing molecules, and an increase in its efficiency results in higher abundances of these molecules. (2) At $15$ K the grains are warm enough for the desorption of N$_2$ ice to proceed more rapidly than at $10$ K (the rate of desorption is approximately $10^{10}$ times greater at $15$ K compared to $10$ K).

The way in which these processes act to produce the model results depends on the conditions in the models. As can be seen in Figure 6, increasing the temperature to $15$ K results in an increase in the abundance of nitrogen-bearing molecules for the non-magnetic models (bold purple curves). The opposite effect is seen for the magnetic models (bold brown curves). The difference between the two types of models is due to the timescales over which they evolve. The non-magnetic models in Figure 6 evolve quickest and reach their final density within $2$ Myr, whereas the magnetic models take between $5.3$ and $5.5$ Myr (Figure 1). Considering the non-magnetic models first, there is an increase in gaseous N$_2$ caused by its faster desorption from the grains (see Figure 8). This means that the N$_2$ abundance in the gas is higher at earlier times. The increase in N$_2$ increases the destruction rate of H$_3^+$ as they react, forming N$_2$H$^+$, and results in the sharp decrease in H$_3^+$ abundance seen in Figure 6. Most of the N$_2$H$^+$ will be recycled back into N$_2$ by recombination with electrons, but a small amount ($\sim 5\%$; Molek et al. 2009) will form NH. This can go on to form other gaseous molecules, or it can freeze out onto the grains, where it is quickly hydrogenated to form ammonia ice. Ammonia has a much higher binding energy than N$_2$ and once formed remains on the grains at these temperatures. The effect of the enhanced rate for reaction (7) can also be seen in the increased abundance of NH in the gas phase in the $15$ K model.

The magnetic models are indicated by red/brown lines in Figure 6, where the bold brown line denotes the $15$ K model. The increased temperature here has exactly the opposite effect on nitrogen molecular abundances compared to the non-magnetic models. In this case the abundance of N$_2$H$^+$ falls at $15$ K, as do the abundances of most other nitrogen-bearing species. The abundance of H$_3^+$ is roughly the same for all temperature models. The reason for this variance is the differing dynamical timescales of the models considered. The magnetic models remain at a relatively low density for $> 2$ Myr, before the onset of rapid collapse. During this prolonged period of time the chemistry evolves quite differently depending on the temperature. This is illustrated in Figure 7, which shows the evolution of the N$_2$H$^+$ abundance with time for two static models with a density of $2 \times 10^4$ cm$^{-3}$ and with temperatures of $10$ K (black) and $15$ K (red). At $15$ K the nitrogen chemistry evolves faster, with X(N$_2$H$^+$) reaching a peak value of $\gtrsim 10^{-13}$ cm$^{-3}$ at $1$ Myr, compared
Water ice abundance (Boogert et al. 2011; ¨Oberg et al. 2011); in the magnetic models, rapid collapse does not begin until after ∼4 Myr. From Figure 7 we can see that the $N_2H^+$ abundance is falling rapidly at this time and is more than an order of magnitude higher at 10 K than at 15 K. At 15 K much more nitrogen has been incorporated into NH$_3$ ices by this stage in magnetic models (see Figure 8) and less is available in the gas phase. Hence, the early evolution of the collapse is quite different and results in the different abundances seen in Figure 6.

Figure 8 shows the behavior of central abundances of various molecules in grain ice mantles as a function of central density. Colors and lines are as in Figure 5; however, here the effect of temperature is also overplotted with a brown shaded area for the magnetic models, and a purple shaded area for the non-magnetic models. In all models the composition of ice mantles is dominated by H$_2$O, in accordance with observations of absorption features in the spectra of background stars viewed through dense clouds taken in IR (e.g., Whittet 2003 and references therein). In observed dark quiescent regions (where cores are most likely to be prestellar and the effect of the protostellar UV field is minimal), CO and CO$_2$ are also observed to be important constituents of ice mantles: the contributions relative to the abundance of water ice is at the level of 10%–40% for CO (Chiar et al. 1995; Gibb et al. 2004; Chiar et al. 2011; ¨Oberg et al. 2011) and 20%–30% for CO$_2$ (Whittet et al. 2009; Gibb et al. 2004; ¨Oberg et al. 2011). Our models are in general agreement with these observations. There are, however, a few exceptions. The CO ice abundance in the 10 Myr delay time non-magnetic model shows a great sensitivity to temperature, with the 15 K model abundance being much lower (more than three orders of magnitude) than the rest of the models and observations. Similarly, the CO$_2$ abundance is significantly decreased, compared to observations for the 7 K versions of the 10 Myr delay and 1 Myr delay non-magnetic models and in comparison to all of the magnetic models. In observed clouds CH$_3$OH is also found at the few percent level relative to the water ice abundance (Boogert et al. 2011; ¨Oberg et al. 2011); in our models the CH$_3$OH abundance is of the order of $\lesssim$1%.

Nitrogen-bearing ices are dominated in our models by NH$_3$ ($\lesssim$10% of the water ice abundance), whereas CN ice is very underabundant, consistent with observations (Gibb et al. 2004; Chiar et al. 2011). NH$_3$ ice is shown to be a good chronometer, with its abundance almost monotonically increasing with the evolutionary timescale. HCN is systematically more abundant than HNC. N$_2$ ice is very sensitive to temperature as discussed above, and its abundance decreases with increasing temperature.

Molecular oxygen ice abundance has a sensitive dependence on the elemental C/O ratio (cyan and yellow bands for magnetic and non-magnetic models, respectively), and its abundance decreases as oxygen elemental abundance decreases, as expected. It is this dependence that dominates the range in possible values for the O$_2$ ice abundance.

The evolutionary trends with central density followed in our models by various mantle ice abundances are generally similar: the abundance builds up fast as a result of efficient molecule freezeout onto grains and high enough ($A_V > 3$) extinction, and saturates at higher values of the central density.

5.4. Radial Abundance Profiles

Figure 9 shows radial profiles of gas-phase abundances of representative molecules, taken when the core has a central density of $10^4$ cm$^{-3}$ (upper row) and $10^6$ cm$^{-3}$ (lower row). Colors and lines are the same as in Figure 8. At high densities (lower row), both magnetic and non-magnetic models have experienced significant central depletion of all molecules (although generally to different extent). This is consistent with the qualitative picture of Figure 5: at high densities the molecular abundances in all models have decreased significantly. At lower densities, however (upper row), magnetic models exhibit a qualitatively different behavior from non-magnetic models, at least for certain molecules. The difference is most striking in the case of CO, which for non-magnetic models is centrally peaked at a central density of $10^6$ cm$^{-3}$. By contrast, at the same central density magnetic models have already suffered significant depletion at the center. In the case of NH$_3$ and CN, however, the radial profiles are quite similar (centrally peaked or flat) in both magnetic and non-magnetic models, even at low densities. As we will see in Section 5.6, this effect can be used to discriminate between non-magnetic and magnetic models.

An important point to note is that the plots in Figure 9 are in a logarithmic scale, and, as a result, most of the core mass is located in the larger scales depicted in the plots. For this reason, even in the cases where some or all models have undergone significant central depletion, when a core is not resolved the observed abundances sample mostly the outer layers, where little if any depletion has occurred. This results in degeneracies among models with otherwise significant differences in central depletion, when a core is unresolved. This point is prominently demonstrated in Section 5.5.

5.5. Total Abundances—Unresolved Cores

Figure 10 shows the total gas-phase abundances of selected molecules in the entire core (i.e., abundances mass averaged over the entire extent of the “unresolved” core), as a function of evolutionary stage, quantified by the core central density ($x$-axis). More advanced evolutionary stages are to the right side of the horizontal axis. The effect of C/O ratio is shown with the shaded orange and cyan regions (for magnetic and non-magnetic models, respectively), whereas the effect of temperature is...
Figure 8. Same as Figure 5, for grain mantle ices. Reddish hues correspond to magnetic models (dashed: magnetically subcritical; solid: magnetically critical; dotted: magnetically supercritical), and bluish hues to non-magnetic models (dashed: 10 Myr delay; solid: 1 Myr delay; dotted: no delay). The thin and thick solid lines correspond to cosmic-ray ionization rate $\zeta$ a factor of four above and below the “reference” value. The orange/cyan shaded areas show the effect of varying C/O ratio for magnetic/non-magnetic models and correspond to a range of C/O values between 0.4 (reference value) and 1.2. The brown/purple shaded areas show the effect of varying temperature for magnetic/non-magnetic models and correspond to a range of $T$ values between 7 K and 15 K (reference value is 10 K).

A striking feature of the total gas-phase abundances is that there is little evolution with central density. The cause of this effect is the dominant contribution of the outer layers of the core (the lower-density regions) to the total core mass and their weight in determining the total core abundances as would be observed in unresolved cores. Since the abundances in the outer layers of the cores do not evolve much, the total core abundances also remain relatively stable with evolutionary stage.

Because of the power-law abundance profiles, this feature is generally preserved even if one averages only over radii with densities above the critical density for excitation of a specific molecular line instead of over the entire core. Of course, the fact that a specific line is observed already places a lower limit on the central density of the core, equal to $n_{\text{crit}}$ for the specific line. However, little evolution of the abundance with time is seen beyond that limit, unless $n_{\text{crit}}$ is very high ($\gtrsim 10^6$ cm$^{-3}$). This can be seen in Figure 11, where we have plotted abundances of selected molecules averaged over regions with densities higher than the critical density $n_{\text{crit}}$ for the following molecular lines: CO($J = 1\rightarrow 0$), 115 GHz, $n \geq n_{\text{crit}} = 2.18 \times 10^5$ cm$^{-3}$; HCO$^+$(J = 1$\rightarrow 0$) 89.2 GHz, $n \geq n_{\text{crit}} = 1.63 \times 10^5$ cm$^{-3}$; NH$_3$(J = 11), 23 GHz, $n \geq 10^7$ cm$^{-3}$ (although the critical density for this transition is much lower ($\sim 2 \times 10^3$ cm$^{-3}$), this line requires much higher densities before the line is thermalized; Evans 1989); N$_2$H$^+$(J = 1$\rightarrow 0$), 93.17 GHz, $n \geq n_{\text{crit}} = 2.25 \times 10^5$ cm$^{-3}$; HCN(J$F = 12\rightarrow 01$), 88.631 GHz, $n \geq n_{\text{crit}} = 2.18 \times 10^6$ cm$^{-3}$; and CN(N = 1$\rightarrow 0$), 113.17 GHz, $n \geq n_{\text{crit}} = 1.3 \times 10^6$ cm$^{-3}$. These critical densities are for $T = 10$ K. All molecular data are taken from the Leiden Atomic and Molecular Database (Schöier et al. 2005).5

The large range of possible abundance values for different models at a fixed evolutionary stage (central density) demonstrates the large potential of chemical differentiation in prestellar cores. Even cores that appear identical in, e.g., their total column

5 http://www.strw.leidenuniv.nl/~moldata/
density profiles, can exhibit large variations in their molecular abundances as a result of a diversity in other local physical parameters (temperature, ionization rate, initial elemental abundance ratios, and dynamics).

However, the large variation in molecular abundances among different models is neither monotonic nor systematic, and the abundances resulting from different parameters are not well separated. Instead, we often see a nonlinear dependence on certain parameters (a characteristic example is the ionization rate), as well as largely overlapping ranges, resulting in strong model degeneracies. We conclude that discrimination between models is not straightforward, but instead requires the use of a variety of different observations and constraints, in combination with a statistical study accounting for as many different molecular abundances as possible. In the next sections we will discuss certain quantities, such as depletion measures and abundance ratios, which also hold promise as tools for the discrimination among dynamical models and physical parameters.

We caution the reader that the abundances in Figure 10 are not to be compared directly with abundances derived from observations beyond an uncertainty of a factor of at least several. There are various reasons for possible discrepancies. First of all, the mass-averaged abundances presented here are not appropriate when the core is at least partly resolved; in this case, a more detailed study is needed, starting with the radial profile at a specific evolutionary snapshot for each model, and properly accounting for the beam size and the physical scale it corresponds to at the distance of the core. Second, chemical reaction rates have appreciable uncertainties (Wakelam et al. 2010). We have not studied the effect of such uncertainties here, as our purpose was to investigate the effect of varying the core dynamics and other physical parameters on molecular abundances. Finally, obtaining a molecular abundance from observations involves several approximations as well as assuming a value for the H$_2$ column density (which is not always well constrained). Nevertheless, we have verified, for the following cores, that molecular abundances obtained from observations are generally within our model ranges, or only deviate from it within uncertainties: L1498 (Tafalla et al. 2004, 2006); L1517B (Tafalla et al. 2004, 2006; Crapsi et al. 2005); L1544 (Caselli et al. 1999; Jørgensen et al. 2004); L134N (Wakelam et al. 2006); TMC-1(CP) (Smith et al. 2004); L1689B (Jørgensen et al. 2004). For an improved comparison with observations, the appropriate methodology would be to start from our dynamical/chemical models, perform radiative transfer calculations, and directly compare predicted line intensities with observed ones. We plan to return to such a comparison in a future study.

5.6. Depletion Measures

The qualitatively different behavior between the radial profiles of magnetic and non-magnetic models seen in Figure 9 can be used to discriminate between models. We define the central deviation $\Delta$ (central depletion or enhancement) of the molecular abundance as the ratio between the integrated abundance through the central 30% of the radial extent of the core over the integrated abundance over the entire core.

We have confirmed that the results are similar if this fraction is changed to 10% or 20%.

\[ \frac{n_{\text{H}_2}}{n_{\text{H}^+}} \]

\[ \frac{n_{\text{CO}}}{n_{\text{H}}^{10}} \]

\[ \frac{n_{\text{CH}_3}}{n_{\text{H}}^{12}} \]

\[ \frac{n_{\text{CNO}}}{n_{\text{H}}^{14}} \]

Figure 9. Radial profiles of gas-phase abundances of three commonly observed molecules, taken when the core has reached a central density of 10$^4$ cm$^{-3}$ (upper row) and 10$^6$ cm$^{-3}$ (lower row). Reddish hues correspond to magnetic models (dashed: magnetically subcritical; solid: magnetically critical; dotted: magnetically supercritical), and bluish hues to non-magnetic models (dashed: 10 Myr delay; solid: 1 Myr delay; dotted: no delay). The thin and thick solid lines correspond to cosmic-ray ionization rate $\zeta$ a factor of four above and below the “reference” value. The orange/cyan shaded areas show the effect of a varying C/O ratio for magnetic/non-magnetic runs, and correspond to a range of C/O values between 0.4 (reference value) and 1.2. The brown/purple shaded areas show the effect of varying temperature for magnetic/non-magnetic models and correspond to a range of $T$ values between 7 K and 15 K (reference value is 10 K).
Tassis et al.

Figure 10. Abundances mass averaged over the entire core for representative molecules at four different stages in the core’s evolution (quantified by the core central density) for different dynamical models, different C/O ratios, and different cosmic-ray ionization rates. Reddish hues correspond to magnetic models (dashed: magnetically subcritical; solid: magnetically critical; dotted: magnetically supercritical), and bluish hues to non-magnetic models (dashed: 10 Myr delay; solid: 1 Myr delay; dotted: no delay). The thin and thick solid lines correspond to cosmic-ray ionization rate \(\zeta\) a factor of four above and below the “reference” value. The orange/cyan shaded areas show the effect of a varying C/O ratio for magnetic/non-magnetic models, and correspond to a range of C/O values between 0.4 (reference value) and 1.2. The brown/purple lines show the effect of varying temperature for magnetic/non-magnetic models. Thin brown/purple lines: \(T = 7\) K; thick brown/purple lines: \(T = 15\) K (reference value is 10 K).

Despite the significant overlap between magnetic and non-magnetic models both in the radial profiles and the total abundances discussed in Figures 9 and 10, respectively, for many molecules the magnetic and non-magnetic models not only are well separated, but also show little sensitivity to model parameters such as the evolutionary timescale, the temperature, the C/O ratio, and the cosmic-ray ionization rate. Molecules which show such good separation between magnetic and non-magnetic models include CO, CO\(_2\), HCO\(^+\), NO, OH, H\(_2\), and...
HC$_3$N. By contrast, H$_3^+$ and CN show a particularly degenerate behavior between different models.

For non-magnetic models, CO-bearing molecules, in particular CO, CO$_2$ and HCO$^+$, tend to show no depletion ($\Delta < 1$) at early stages of the collapse and only become depleted when the central density reaches $10^5$–$10^6$ cm$^{-3}$. By contrast, for magnetic models, the same molecules are already centrally depleted by central densities of $\sim 10^4$ cm$^{-3}$. NH$_3$, on the other hand, shows no sign of central depletion at central densities of $\sim 10^4$ cm$^{-3}$ for both magnetic and non-magnetic models. It only becomes depleted at later stages of the collapse ($10^6$–$10^7$ cm$^{-3}$ for non-magnetic models and $10^5$–$10^6$ cm$^{-3}$ for magnetic models). N$_2$H$^+$ shows a similar behavior; however, its behavior is complicated by its sensitivity to temperature in the case of non-magnetic models (see also discussion in Section 5.3). Note also that the high cosmic-ray ionization rate magnetic model is always depleted of N-bearing molecules as a result of the very long evolutionary timescale (see also discussion of Figure 5). This difference in behavior between CO and NH$_3$ is consistent with observations (e.g., Caselli et al. 1999; Bacmann et al. 2003; Crapsi et al. 2007).

5.7. Abundance Ratios of Molecules with Discriminatory Potential

Ratios between abundances of different molecules have long been considered to be promising tools to overcome model uncertainties and degeneracies. However, most abundance ratios exhibit large variations and no consistent trend with any single model parameter when other parameters are varied. We have identified several gas-phase abundance ratios (averaged over the entire core) that are most affected by a single model parameter and show little variation as a result of the rest and thus show promising discriminatory potential. We plot these in Figures 13 and 14.

Figure 13 shows the values of the abundance ratio between HCN and OH, mass averaged over the entire core, for our various models. This ratio can serve as an indicator of the C/O ratio. Different models are plotted with different symbols as discussed in the caption. The C/O ratio, shown in the vertical axis, exhibits a trend (increases with increasing value of each abundance ratio). Similarly, Figure 14 shows three abundance ratios that trace the evolutionary timescale of a core. These ratios are $X_{\text{NH}_3}/X_{\text{CO}}$, $X_{\text{NH}_2}/X_{\text{CO}}$, and $X_{\text{NH}_3}/X_{\text{HCO}^+}$. All the ratio values are taken at the time when the central density in each model has reached $10^6$ cm$^{-3}$. However, the averaged abundance over the entire core shows little time evolution, as shown in Figure 10.

Each abundance ratio examined in these two figures shows an appreciable scatter at a fixed value of the parameter that it can help constrain (C/O ratio or evolutionary timescale, respectively). However, a general trend is obvious, and there are significant parts of the C/O or timescale parameter space that can be excluded with measurements of each ratio.

6. DISCUSSION AND CONCLUSIONS

We have combined dynamical and non-equilibrium chemical modeling of both magnetic and non-magnetic contracting molecular cloud cores, in order to predict molecular abundances as a function of core evolutionary stage and to identify observables with high promise to discriminate among core models.

To this end we have undertaken an extensive parameter study, following a total of 34 different models, and varying:
Figure 12. Central deviation of molecular abundance with respect to the average abundance over the entire core ($\Delta$; see the text) as a function of central density. The dotted line corresponds to $\Delta = 1$ (no central enhancement or depletion). Reddish hues correspond to magnetic models (dashed: magnetically subcritical; solid: magnetically critical; dotted: magnetically supercritical), and bluish hues to non-magnetic models (dashed: 10 Myr delay; solid: 1 Myr delay; dotted: no delay). The thin and thick solid lines correspond to cosmic-ray ionization rate $\zeta$ a factor of four above and below the “reference” value. The shaded areas show the effect of a varying C/O ratio, and correspond to a range of C/O values between 0.4 (reference value) and 1.2. The brown/purple shaded areas show the effect of varying temperature for magnetic/non-magnetic models and correspond to a range of $T$ values between 7 K and 15 K (reference value is 10 K).

1. the core evolutionary timescale, controlled by the collapse delay time for the case of non-magnetic models, or by the initial mass-to-magnetic-flux ratio, i.e., the amount of magnetic support against gravity, for magnetic models;
2. the elemental C/O ratio in the core;
3. the cosmic-ray ionization rate, which affects the chemistry in both magnetic and non-magnetic models, but also the evolutionary timescale in magnetic models, by affecting the extent of coupling between the magnetic field and the neutral fluid; and
4. the temperature.

Our main results can be summarized as follows. We have explicitly demonstrated that many reactions in our chemical network are out of equilibrium during the dynamical evolution of the core, necessitating the inclusion of non-equilibrium chemistry and the coupled modeling of chemistry and dynamics to properly follow the molecular abundances. For different dynamical models the molecular abundances at the center of the core are not the same, even for identical values of the central number density; rather, the amount of time
Figure 13. Gas-phase abundance ratio between HCN and OH mass averaged over the entire core, which has good potential for discriminating between C/O ratio values. Red/blue symbols correspond to magnetic/non-magnetic models with $T = 10$ K. Diamonds: “slow” models; X: “reference” models; crosses: “fast” models. The size of the red/blue symbols represents the value of $\xi$ (larger symbols correspond to larger $\xi$). Brown/purple symbols correspond to magnetic/non-magnetic models with $T = 7$ K (small symbols) and $T = 15$ K (large symbols), and a “reference” value for $\xi$. The C/O ratio of each model is shown on the vertical axis, while the value of the abundance ratio is shown on the horizontal axis. Points correspond to a central H$_2$ density of $10^6$ cm$^{-3}$; however, the core-average abundance ratios evolve little with density.

that a model core has spent at each density is what determines the actual molecular abundances at each evolutionary stage. Therefore, it is not possible to properly account for magnetic effects in core chemistry by using a collapse retardation factor in non-magnetic models. Even if the overall final age of the core is the same, the fact that the non-magnetic core model has spent very different fractions of this age at various values of the central density compared with a magnetic core model results in significantly different molecular abundances.

Abundances in magnetic models peak at much lower densities and decrease much faster at higher densities. The evolution of grain mantle ice abundances is qualitatively similar in magnetic and non-magnetic models, with a fast initial increase due to freezeout, and an abundance saturation at higher densities and later times. The value of that saturation, however, is different in different models. We find that the abundance of NH$_3$ ice scales roughly with the evolutionary timescale of the model, varying by a factor of four for our model range, which corresponds to an evolutionary timescale range of about 10 Myr. NH$_3$ ice is thus a potentially important chronometer.

Radial profiles of gas-phase abundances reveal that the most significant difference between magnetic and non-magnetic models is at the central parts of the core, whereas most of the core extent tends to evolve much less and is much more similar among different models. However, as only a small fraction of the mass is concentrated in the central layers of the core, the central differences are smeared out in unresolved cores. As a result, the evolution of the total, mass-averaged molecular abundances with central density in any particular model is much milder, if at all present. The situation is similar with the total abundances of grain mantle ices.

Differences in the C/O ratio tend to affect mostly the abundance of the oxygen-bearing molecules, whereas the cosmic-ray ionization rate $\xi$ has its largest effect on the nitrogen-bearing molecules. For magnetic models the direct effect of the cosmic-ray ionization rate on chemistry is compounded by its indirect effect on core evolution timescale, and as a result magnetic models are much more sensitive to changes in $\xi$.

The many effects of different model parameters on molecular abundances, sometimes competing, sometimes compounding, result in severe degeneracies among different models. For this reason we have identified molecular abundance ratios which may help to constrain, even in unresolved cores, specific model parameters: the evolutionary timescale (ratios between NH$_3$ and CO, NH$_2$ and CO, and NH$_3$ and HCO$^+$) and the C/O ratio (ratio between HCN and OH).

In addition, we have demonstrated that the central deviation $\Delta$ (central depletion or enhancement) of molecules can be another useful tool in model discrimination. Whereas values of $\Delta$ for many molecules are degenerate among different models, there are several molecules for which there is a clear separation between magnetic and non-magnetic models. For these molecules the value of $\Delta$ even shows very little sensitivity to other model parameters. Measurements of the central depletion of such molecules therefore have an excellent

Figure 14. Gas-phase abundance ratios mass averaged over the entire core, with high potential for discrimination among evolutionary timescales for the core. Diamonds: “slow” models; X: “reference” models; crosses: “fast” models. Red/blue symbols correspond to magnetic/non-magnetic models with $T = 10$ K and C/O = 0.4. The size of the red/blue symbols represents the value of $\xi$ (larger symbols correspond to larger $\xi$). Orange/cyan points correspond to magnetic/non-magnetic models with $T = 10$ K, C/O = 1.2, and a “reference” value for $\xi$. Brown/purple symbols correspond to magnetic/non-magnetic models with $T = 7$ K (small symbols) and $T = 15$ K (large symbols), C/O = 0.4, and a “reference” value for $\xi$. The time it takes the core to reach a central density of $10^6$ cm$^{-3}$ is shown on the vertical axis, while the value of the abundance ratio is shown on the horizontal axis. Points correspond to a central H$_2$ density of $10^6$ cm$^{-3}$; however, the core-average abundance ratios evolve little with density.
Figure 15. Fractional change of total elemental abundance of Si ($\chi_{Si}$) with respect to the initial one ($\chi_{Si0}$) vs. central density in collapsing core models. Pure hydrodynamical model cores are shown with blue symbols and lines, while MHD core models are shown with red symbols and lines as in Figure 1.

potential for constraining dynamical models of core evolution. For this reason it is very important to place adequate emphasis into understanding differential depletion and grain chemistry effects as much as possible, and thus gain confidence in the chemistry part of these calculations, which still features significant uncertainties (e.g., Wakelam et al. 2010).

The dynamical models we have used in this study have low dimensionality: the non-magnetic models are spherically symmetric (one-dimensional), whereas the magnetic ones are axially symmetric, and integrated over the $z$-direction (1.5 dimensions). We note that as a result of the distinct geometries, the magnetic model does not converge to the non-magnetic model in the limit of zero magnetic field. A zero magnetic field disk model cannot be followed with the current numerical scheme, as the thin-disk approximation breaks down in the absence of a magnetic field. Although higher-dimensional calculations are desirable (e.g., van Weeren et al. 2009), it is currently not possible to perform a parameter study as extensive as the one undertaken here using higher-dimensional models: a single model in this study (including non-equilibrium chemistry) requires more than 800 CPU hours for the magnetic runs and more than 200 CPU hours for the non-magnetic runs. Nevertheless, we plan to consider true two-dimensional calculations for select cases presented here, to assess the effect of low dimensionality on our calculations.

Because the model cores are not multi-dimensional, multiplicity or fragmentation is by definition neither accounted for nor allowed. This, however, is not a severe disadvantage in our case, as multiplicity at these early stages of collapse and for the low densities considered here (up to $10^7$ cm$^{-3}$) is not observed (Schnee et al. 2010).

Finally, our work has not accounted for turbulent mixing; the flows in our model are laminar. Any turbulent mixing would result in making the radial profiles shallower (Xie et al. 1995). However, dense molecular cloud cores are generally not observed to have significant amounts of turbulence since their line widths are approximately sonic (Myers 1983; Barranco & Goodman 1998; Goodman et al. 1998; Kirk et al. 2007).

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APPENDIX

CODE TESTS

As a test of the dynamical part of our code, we have demonstrated that for the (more complex) magnetic model our code returns the same results as Basu & Mouschovias (1994) when identical initial conditions are used.

To confirm that the chemistry implementation in our code behaves as expected, we have checked that the elemental abundances are conserved in our simulations. Figure 15 shows the fractional changes in the abundance of the most underabundant element Si (abundance of $10^{-8}$, see Table 2) as a fraction of central volume density, for magnetic (red lines) and non-magnetic (blue lines) of different mass-to-magnetic-flux ratios and initial delays, respectively. The fractional change always remains $<10^{-3}$ for all simulations. We thus confirm that the codes conserve elemental abundances better than 1 part in 1000 of the total abundance even for the most underabundant elements.

Figure 16. Fractional change of the CO abundance as a function of mass at different times of the collapse, normalized to the initial time $t_0$. Left panel: non-magnetic reference model; right panel: magnetic reference model. The inset shows the initial abundance profile.
Furthermore, we have performed a test on the advection of chemical species. Retaining identical initial and boundary conditions as in the main runs in both magnetic and non-magnetic models, we allowed the chemistry to evolve for 1 million years before any dynamical evolution was allowed to take place. In this way, an abundance profile for the various molecules was developed. During the next step of the test, the dynamics were turned on, while the chemical reactions were turned off, so that molecules were advected, but their abundances could not be altered due to sources or sinks. Advection should not be changing the distribution of abundance with mass, and the abundances within a single mass element should be conserved. The only expected change, as the contraction of the core proceeds, is the distribution of mass elements with scale. The results are presented in Figure 16. The left panel corresponds to the non-magnetic reference model and the right panel to the magnetic reference model. The plots show the relative error (normalized to the initial time) of the abundance of CO for different mass elements for different snapshots in time after the dynamics are turned on. The CO abundance profile is shown in the insets. We conclude that no spurious numerical alteration in molecule abundances is induced because of advection, as expected for a fully conservative advection scheme such as the one implemented here (Morton et al. 1994).

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