DENSITY-ENHANCED GAS AND DUST SHELLS IN A NEW CHEMICAL MODEL FOR IRC+10216

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ABSTRACT

A new chemical model is presented for the carbon-rich circumstellar envelope (CSE) of the asymptotic giant branch star IRC+10216. The model includes shells of matter with densities that are enhanced relative to the surrounding circumstellar medium. The chemical model uses an updated reaction network including reactions from the RATE06 database and a more detailed anion chemistry. In particular, new mechanisms are considered for the formation of \( CN^- \), \( C_2N^- \), and \( C_2H^- \), and for the reactions of hydrocarbon anions with atomic nitrogen and with the most abundant cations in the CSE. New reactions involving \( H^- \) are included which result in the production of significant amounts of \( C_2H^- \) and \( CN^- \) in the inner envelope. The calculated radial molecular abundance profiles for the hydrocarbons \( C_2H \), \( C_4H \), and \( C_6H \) and the cyanopolyynes \( HC_3N \) and \( HC_5N \) show narrow peaks which are in better agreement with observations than previous models. Thus, the narrow rings observed in molecular microwave emission surrounding IRC+10216 are interpreted as arising in regions of the envelope where the gas and dust densities are greater than the surrounding circumstellar medium. Our models show that \( CN^- \) and \( C_2H^- \) may be detectable in IRC+10216 despite the very low theorized radiative electron attachment rates of their parent neutral species. We also show that magnesium isocyanide (\( MgNC \)) can be formed in the outer envelope through radiative association involving \( Mg^+ \) and the cyanopolyyne species.

Key words: astrochemistry – circumstellar matter – ISM: molecules – stars: individual (IRC+10216)

Online-only material: color figures

1. INTRODUCTION

The carbon-rich asymptotic giant branch (AGB) star IRC+10216 is among the richest known sources of molecules in the sky. The star is nearing the end of its life and is rapidly losing mass to the interstellar medium. Over 60 species have been detected in the expanding circumstellar envelope (CSE), which extends more than 10,000 AU from the star. The outflow from the central star is carbon-rich (with \([C]/[O] > 1\)), and contains abundant stable molecules such as \( C_2H_2 \), \( HCN \), and \( CH_4 \), which are formed in the hot inner regions near the stellar surface (Lafont et al. 1982). As the density falls with radius, interstellar ultraviolet (UV) photons penetrate the CSE and cause the photolysis of these “parent” molecules. This results in the production of reactive species that subsequently participate in a complex gas-phase chemistry to produce new “daughter” molecules in the outer envelope. Chemical models of this environment (see Millar et al. 2000, for example) are able to reproduce with good accuracy the observed column densities of outer-envelope species such as the carbon chains \( C_nH \) (for \( n = 2, 4, 6, 8 \)). However, it has been suggested that such chemical models are inconsistent with observed \( C_2H \) rotational emission maps (see, for example, Guélin et al. 1999). The observations show \( C_2H \) emission maxima at around the same radius (for different \( n \)), whereas the results of chemical models show spatially separated peak abundances. The modeled molecular radial distributions are also broader than observed. Using optical imaging, Mauron & Huggins (2000) detected multiple dust shells in the envelope of IRC+10216, spaced at regular intervals of \( \sim 5'' \) to \( 20'' \). These shells were interpreted as arising as a result of modulation of the stellar mass-loss rate, perhaps due to the presence of a binary stellar companion. Brown & Millar (2003) included density-enhanced circumstellar dust shells in their chemical model for IRC+10216 in an attempt to make the model more physically realistic and to address the discrepancies between observed molecular emission maps and the results of previous chemical models. The addition of dust shells modifies the radiation field inside the CSE and results in modeled molecular abundance profiles that are in better agreement with the observations.

Figure 1 shows the effect of the Brown & Millar (2003) density-enhanced dust shells on the modeled \( C_2H \), \( C_4H \), and \( C_6H \) radial profiles in IRC+10216. In the presence of dust shells, the molecular abundance peaks move outward and slightly closer together. The \( C_2H \) profile full width at half-maximum narrows from \( 7'' \) to \( 6'' \) and moves from \( 8'' \) to \( 18'' \). According to Guélin et al. (1999), the molecular emission maps from \( C_2H \), \( C_4H \), and \( C_6H \) all peak in a narrow circumstellar ring about \( 2'' \) wide at a radius about \( 15'' \) from the star. It is therefore clear that although the radial distributions of the species in Brown & Millar’s (2003) model represent a better match than previous models, they are still too broad to fully explain the observed maps. In addition, \( C_6H \) peaks at a larger radius than \( C_2H \) and \( C_4H \), which is inconsistent with the observations.

Dinh-V-Trung & Lim (2008) mapped the molecular shells of \( HC_3N \) and \( HC_5N \) in IRC+10216 at unprecedented angular resolution. The shells were found to be clumpy, co-spatial with each other and with a structure closely matching the distribution of dust shells observed by Mauron & Huggins (2000). It was concluded that the circumstellar cyanopolyyne gas density distribution matches that of the dust and therefore that the gas and dust are coupled. Shell-like structures \( \sim 2'' \) wide are present in the maps of Dinh-V-Trung & Lim (2008), which are inconsistent with the \( \sim 10'' \) wide abundance profiles calculated in the model of Brown & Millar (2003).

In this article, a new chemical model for IRC+10216 is presented which builds on previous models and includes density-
enhanced shells of gas in addition to the dust shells of Brown & Millar (2003). Thus, the new model assumes kinematical coupling between the gas and dust in the CSE. The aim is to model the chemistry of the narrow shell-like structures observed in the IRC+10216 outflow by Dinh-V-Trung & Lim (2008) and Guélin et al. (1999) to test the idea that coupling between the gas and dust plays an important role in determining the morphology of the molecular distributions.

The molecular anions C$_2$H$^-$, C$_3$H$^-$, C$_4$H$^-$, and C$_2$N$^-$ have been recently detected in the envelope of IRC+10216 (for a review of the astrophysical anion detections reported so far, see Herbst & Osamura 2008 and the newly published detections of Gupta et al. 2009). Thus, the new model assumes that magnesium isocyanide (MgNC) is produced in the outer CSE by gas-phase chemistry is reconsidered upon the anion chemistry in IRC+10216 studied by Millar et al. (2000, 2007). In particular, the C$_2$N$^-$ chemistry is reconsidered using the reaction rates between N atoms and carbon-chain anions published by Eichelleberger et al. (2007). The possibility that magnesium isocyanide (MgNC) is produced in the outer CSE by gas-phase chemistry is also examined.

2. THE CHEMICAL MODEL

2.1. Physics

The new chemical model for IRC+10216 is based on the model of Miller et al. (2000). The underlying density distribution in the CSE is derived using a mass-loss rate of 1.6 $\times$ 10$^{-5}$ $M_\odot$ yr$^{-1}$ (after Men'shchikov et al. 2001), and assuming that the matter expands in a spherically symmetric outflow with a velocity of 14 km s$^{-1}$. The resulting gas number density profile $n(r)$ falls as $1/r^2$, added to which is a series of steplike density enhancements of the form $\beta n(r)$. Based on the dust shell parameters deduced from scattered light observations by Mauron & Huggins (2000), each shell is 2" thick with an intershell spacing of 12", which corresponds to a timescale of approximately 530 years between successive episodes of enhanced mass loss. The parameter $\beta$ is set to 5 for all shells in the present model (see Section 2.3). To convert between physical length units and angular distances on the sky, the distance to IRC+10216 is taken to be 130 pc (after Men'shchikov et al. 2001). The chosen shell parameters are not intended to provide an accurate representation of the dust shell structure observed in IRC+10216 (for which a three-dimensional, time-dependent model of the gas and dust in the envelope would be required), but to permit the study of the general effects on the chemical model of the addition of density-enhanced gas and dust shells. The radial H$_2$ gas density distribution used in the model is shown in Figure 2.

The adopted temperature profile is based on an empirical fit to the gas kinetic temperature profile derived by Crosas & Menten (1997), and takes the form

$$T(r) = \max \left[ \left( \frac{2.81 \times 10^{15}}{r} \right)^{4.7} + \left( \frac{3.34 \times 10^{17}}{r} \right)^{1.05} \right], 10 \right]$$

with a lower limit of 10 K to prevent the temperature becoming unrealistically low in the outer envelope. Parent species (with abundances shown in Table 1) are injected into the model at the inner radius of $r_i = 10^{15}$ cm, where the gas density is $8.6 \times 10^7$ cm$^{-3}$. At this radius, the gas kinetic temperature is 575 K and the interstellar radiation field is attenuated by an effective radial extinction of 47 magnitudes in the V band. Initial abundances are the same as in Millar et al. (2000), with the addition of Mg (see Section 3.3), and with C$_2$H$_2$ and HCN abundances taken from the recent measurements by Fonfría et al. (2008).

The standard interstellar radiation field (Draine 1978) is assumed to impinge on the outside of the CSE from all directions. The extinction is calculated for the underlying $1/r^2$ density distribution of the CSE using the approach of Jura &
Morris (1981). Additional terms are added to the extinction due to the contributions of the density-enhanced shells (for further details of the extinction calculation in the presence of dust shells, see Brown & Millar 2003).

2.2. Chemistry

The chemical reaction network is based on that used by Petrie et al. (2003) and Millar et al. (2007). The reaction rates have been updated to be consistent with those in the (dipole-enhanced) RATE06 database (Woodall et al. 2007). Additional new reactions from RATE06 for those species in the Petrie et al. (2003) model have also been added. The following reactions were deleted from the reaction network in order to increase computational speed, with negligible effect on the chemistry of the species of interest in this study: those with activation energies greater than 300 K; those reactions involving H$_3$O$^+$ as a reagent (except for H$_3$O$^+$ + e$^-$); those with H$_2^+$ as a reagent (except for H$_2^+$ + e$^-$ and H$_2^+$ + H$_2$); and any reactions involving C$^-$, S$^-$, CO$^-$, HOC$^-$, C$_2$H$_5^-$, and C$_2$H$_5$. Carbon-chain species, hydrocarbons, cyanopolyynes and their associated anions and cations are of principal importance for the chemistry of this study. The following species are among those included in the chemical model: carbon chains C$_n$ (n = 1–23), C$_n^+$ (n = 1–23), C$_n^-$ (n = 3–23); C$_{2n-1}$S (n = 1–3); hydrocarbons C$_n$H$^{(n-1)}$ (n = 2–23), C$_n$H$_2^+$ (n = 1–23), C$_n$H$_3$ (n = 1–4), C$_n$H$_4$ (n = 1–3), C$_n$H$_5$ (n = 1–9), C$_n$H$_6^+$ (n = 2–9) and cyanopolyynes C$_{2n-1}$N$^{(n-1)}$ (n = 1–11), H$_2$C$_{2n-1}$N$^-$ (n = 1–12), H$_2$C$_{2n-1}$N$^+$ (n = 1–11) H$_3$C$_{2n-1}$N$^-$ (n = 1–11), H$_3$C$_{2n-1}$N$^+$ (n = 2–5).

Due to the current interest in molecular anions in IRC+10216 (e.g., Millar et al. 2007; Remijan et al. 2007; Cordiner et al. 2008; Thaddeus et al. 2008), the anion chemistry has been extended to include C$_n$H$^-$ down to $n = 2$, utilizing the radiative electron attachment rates from Herbst & Osamura (2008). The following additional CN$^-$ and C$_2$H$^-$ formation reactions have been included:

\[
\text{HCN} + \text{H}^- \rightarrow \text{CN}^- + \text{H}_2 \quad (2)
\]

\[
\text{C}_2\text{H}_2 + \text{H}^- \rightarrow \text{C}_2\text{H}^- + \text{H}_2. \quad (3)
\]

The rate coefficient used for Equation (2) (3.8 $\times$ 10$^{-9}$ cm$^3$ s$^{-1}$) is an estimate taken from Prasad & Huntress (1980). For Equation (3), the rate coefficient (4.42 $\times$ 10$^{-9}$ cm$^3$ s$^{-1}$) was measured experimentally by Mackay et al. (1977). These proton-transfer reactions are likely to be important in the model due to the large HCN and C$_2$H$_2$ abundances in the stellar outflow. H$^-$ is produced in the model mainly by the cosmic-ray (CR) dissociation of H$_2$ (H$_2 + \text{CR} \rightarrow \text{H}^+ + \text{H}^-$). This reaction is slow (with a rate coefficient of 3.9 $\times$ 10$^{-21}$ cm$^3$ s$^{-1}$; Prasad & Huntress 1980), but provides the main source of H$^-$ in the inner envelope. Interior to the second density-enhanced shell ($r \lesssim 2 \times 10^{10}$ cm), the modeled H$^-$ abundance is about $10^{-8}$ cm$^{-3}$. Other reactions similar to Figure 3 were studied by Mackay et al. (1977), who found that many different molecular anions could receive a proton from C$_2$H$_2$ at rapid rates ($\sim$10$^{-9}$ cm$^3$ s$^{-1}$), resulting in the production of C$_2$H$^-$. However, it is presently unknown whether reactions occur between C$_2$H$_2$ and the carbon chain anions C$_n^-$ and C$_n$H$^-$. In light of an observational upper limit for N(C$_2$H$^-$) in IRC+10216, Cordiner et al. (2008) deduced that these reactions probably do not proceed rapidly.

Dominant anion destruction mechanisms are by reaction with H and C$^+$ and by photodetachment. Photodetachment rates were calculated according to Equation (2) of Millar et al. (2007). As a result of its large electron detachment energy, the CN$^-$ photodetachment rate thus calculated is $\sim$100 times less than the value in the RATE06 database.

Carbon chain anions C$_n^-$ (n = 2–7), and C$_n$H$^-$ (n = 2, 4, 6) have been shown to react with atomic nitrogen and result in the formation of products that include the nitrile anions C$_n$N$^-$ (n = 1, 3, 5) (Eichelberger et al. 2007). We have included these reactions in the model as part of our
investigation into the possible mechanisms for the formation of CN$^-$ and the recently discovered C$_2$N$^-$ (Thaddeus et al. 2008). Branching ratios were calculated from Eichelerberger et al.’s (2007) original experimental data by V. M. Bierbaum (2008, private communication). The branching ratio for the associative electron detachment (AED) product channel could not be derived from the experimental data, so we have assumed, arbitrarily, a ratio of 0.5. This assumption constitutes potentially the most significant source of error in these reaction rates.

Mutual neutralization reactions between anions and cations were shown by Lepp & Dalgarno (1988) to have important effects on interstellar chemistry. Thus, reactions of the kind

$$X^+ + Y^- \rightarrow X + Y$$

(4)

have been included for the 20 most abundant cations ($X^+$), and for all anions ($Y^-$), in the model, with a rate coefficient of $7.5 \times 10^{-8}(T/300)^{-0.5}$ cm$^3$ s$^{-1}$ (see Harada & Herbst 2008, for example).

The final chemical network contains 426 gas-phase species coupled by 5539 reactions.

2.3. Evolution of the Shells

Initially, the rate equations are solved as a function of radius (in the same way as Millar et al. 2000), starting from $r_1 = 10^{15}$ cm and moving out to $r_f = 3 \times 10^{18}$ cm where photodissociation destroys all the molecules (apart from H$_2$). Then, in a separate routine the chemical abundances inside the density-enhanced shells are calculated as a function of radius. The chemical rate equations of a density-enhanced packet lying halfway between a shell’s inner and outer radius (at a radius $r_p$ with density ($\beta = 1)n(r_p)$) are solved, starting from $r_1$ and moving out to $2 \times 10^{17}$ cm where the density is sufficiently low that the shell no longer makes any significant contribution to the total amount of matter in the model. The motion of the density-enhanced shells in the outflow are followed so that they move outward over time, synchronized with the outward radial motion of the dense packet. The radiation field is recalculated for the dense packet at each time step, taking into account the new positions of all the shells. The density factor ($\beta$), the intershell spacing and the shell thickness are identical for every shell so that the chemical abundances calculated for the dense packet at a given radius represent the abundances in a density-enhanced shell centered at that radius.

2.4. Model Variations

The effects on the model results of variations in the mass-loss rate, radiation field strength, gas-to-dust ratio, shell thickness, intershell spacing, density-enhancement factor and stellar distance have been analyzed. The results presented in this study are for the model that uses the parameters that we believe best match the observational constraints. Modification of these parameters, particularly those that affect the radiation field strength, can significantly alter the radial abundance profiles calculated by the model. However, under such circumstances the main conclusions of this study remain the same. The location of the 15$^{\circ}$ density-enhanced shell has been fixed in order to best reproduce the observational data, which may be considered a contrivance of the model. However, in Figure 8 of Mauron & Huggins (2000), this can be identified as the radius at which the first distinct dust shells occur.

To permit comparison between this model and previous models published in the literature (which do not include density-enhanced gas and dust shells), we have also run the model without any density-enhanced shells (i.e., $\beta = 0$) and also with the density-enhanced dust shells of Brown & Millar (2003) to produce the data shown in Figure 1.

2.5. Molecular Excitation

To facilitate comparison of the model results with observed maps of molecular microwave emission line flux, the rotational excitation of some of the molecules of interest has been calculated as a function of radius using a modified version of the mmline computer code (described by Justtanont et al. 1994). The central stellar radius was taken to be $3.1 \times 10^{13}$ cm and the temperature 2650 K (derived from Men’shchikov et al. 2001). The dust opacity was taken from Figure 6 of Men’shchikov et al. (2001). Due to the lack of published collisional excitation rates and vibrational transition strengths, only a rough estimate of the rotational excitation is possible for most molecules. For C$_2$H and C$_2$H$^+$, the HCO$^+$ rates from Flower (1999) are used, and for C$_6$H, C$_4$H$^+$, C$_4$H, and C$_6$H$^+$ the HCN rates from Green & Chapman (1978) are used. For C$_2$H and C$_6$H$^+$, the rates have been extrapolated up to $J = 31$. The collisional rates used are for closed electronic-shell species so the spectroscopic structure of the corresponding (closed electronic-shell) anions has been used for the calculations of the excitation of the open-shell C$_2$H, C$_6$H, and C$_6$H radicals. This approximation is reasonable because the structure of these hydrocarbon anions and neutrals are very similar. Only the $^2\Pi_{1/2}$ states have been considered for the neutral hydrocarbons; the population of the $^2\Pi_{3/2}$ states is not expected to significantly affect the relative populations of the states of interest here. Rotational Einstein A coefficients were calculated using dipole moments from Woon (1995) and Blanksby et al. (2001) for the neutrals and the anions, respectively. Infrared (IR) pumping of rotational levels has been calculated through consideration of the radiative excitation of a single vibrational state $\sim 10$ m$\mu$m above the ground state (e.g., Bieging et al. 1984). C$_2$H has been calculated to have a strong vibrational transition ($A = 0.6$ s$^{-1}$) at 12.5 m$\mu$m (Tarroni & Carter 2004), which we assume to also occur in C$_2$H$^+$. The vibrational spectra of C$_2$H, C$_6$H and their associated anions are less well known. For these species, a transition has been assumed to occur at 12.5 m$\mu$m with $A = 1$ s$^{-1}$. IR pumping has a significant effect on the molecular excitation, but changes in the vibrational transition wavelengths and Einstein A coefficients by up to an order of magnitude do not significantly affect the results of the present study. To assess the impact of errors in the collisional excitation rates on the calculated molecular emission profiles, the rates were varied by an order of magnitude either way. The overall features of the emission profiles remained the same.

3. RESULTS

3.1. Molecular Radial Abundance and Intensity Distributions

The upper panel of Figure 3 shows the calculated radial abundance profiles for the hydrocarbons C$_n$H ($n = 2, 4, 6$). Comparison with Figure 1 shows the impact of the density-enhanced shells on these species. A prominent effect of the shells is to raise of the abundances relative to the surrounding CSE due to the increased gas density. The photon-induced hydrocarbon chemistry is suppressed very near to the star by the dust shell at a radius of $r = 1''$ which provides additional shielding of C$_2$H$_2$ from photodissociation. The abundances all reach a peak in the shell at $r = 15''$. However, this is not where C$_6$H reaches its
greatest abundance, which occurs in the third density-enhanced shell at $r = 29''$. As shown in the middle panel of Figure 4, HC$_3$N and HC$_5$N both reach their greatest abundances in the 15'' shell. CR-induced chemistry also results in the synthesis of a significant amount of HC$_3$N in the innermost ($r = 1''$) shell (through the reaction HCN + CRPHOT $\rightarrow$ CN + H, followed by CN + C$_2$H$_2$ $\rightarrow$ HC$_3$N + H). The anions C$_4$H$^-$, C$_6$H$^-$, and C$_8$H$^-$ (shown in Figure 5) do not reach their maximum abundances in the 15'' shell; C$_4$H$^-$ peaks at a similar radius to the model with no shells whereas C$_6$H$^-$ and C$_8$H$^-$ peak in the 29'' shell.

Molecular abundances tend to peak within the density-enhanced shells because the increased density raises the abundances of chemical reagents which drives the chemistry at a faster rate. It is not always the case, however, that this raises the abundances of daughter species, as can be seen in Figure 3.
where the \( \text{C}_2\text{H} \) abundance is reduced in the \( r = 29'' \) shell due to the increased densities of atoms and ions (including C, C\(^+\), and N), that it reacts with.

A comparison of the abundance profiles in the models with and without density-enhanced shells shows that, in general, the increased shielding of the CSE from interstellar UV by the dust shells (which inhibits the photochemistry) causes photodissociation of parent species to be less efficient in the inner regions and causes the daughter abundances to rise more slowly with radius, moving the profile maxima outward.

The calculated emission intensities for \( \text{C}_2\text{H}, \text{C}_4\text{H}, \text{C}_6\text{H}, \text{C}_8\text{H}^-, \) and \( \text{HC}_3\text{N} \) (shown in the lower panels of Figures 3–5, respectively) are greatest within the \( r = 15'' \) density-enhanced shell. \( \text{C}_3\text{H}^- \) reaches maximum intensity near \( r = 8'' \). The differences between the abundance profiles and the emission profiles for \( \text{C}_2\text{H} \) and \( \text{C}_8\text{H}^- \) are particularly notable. Because the 3 mm emission from these species originates from a high rotational level (around \( J = 30 \)), the strength of the emission is highly dependent on the rate of collisional excitation, and therefore the density. Thus, inside the density-enhanced shells the lower \( J \) levels tend to become depopulated in favor of the higher levels.

For \( \text{C}_2\text{H}, \text{C}_4\text{H}, \text{C}_6\text{H}, \) and their corresponding anions, two-dimensional emission maps have been constructed from the radial emission profiles by integration along (pencil-beam) lines of sight through the CSE. This calculation assumes spherical symmetry of the CSE and therefore cannot show any of the azimuthal structure in observed emission maps. Nevertheless, these maps (shown in Figure 6) provide a useful means for comparing the main features in the modeled and observed molecular emission.

\( \text{C}_2\text{H} \) and \( \text{C}_4\text{H} \) show a strong, narrow emission ring centered at \( 15'' \), in excellent agreement with observed maps of Güellin et al. (1999). In addition to the observed \( 15'' \) ring, the modeled \( \text{C}_4\text{H} \) emission map shows a thick, strong ring at around \( r = 8'' \). The patterns for \( \text{C}_2\text{H}^- \) and \( \text{C}_8\text{H}^- \) are similar to their parent neutrals whereas \( \text{C}_2\text{H}^- \) is markedly different: it has a centrally peaked emission map because it is produced predominantly by the reaction of \( \text{H}^- \) with \( \text{C}_2\text{H}_2 \)—the abundances of which are greatest in the (well shielded) inner CSE—rather than by electron attachment to its parent neutral.

3.2. New Anion Results

Column densities for all anions included in the chemical model are given in Table 2.

Cordiner et al. (2008) identified that the reaction of \( \text{H}^- \) with \( \text{HCN} \) (Figure 2) could dominate the synthesis of \( \text{CN}^- \) in IRC+10216. However, the present study shows that reactions between atomic nitrogen and the carbon chain anions may provide a greater source of \( \text{CN}^- \). Upon inclusion of the reactions \( \text{N} + \text{C}_n\text{H}^- \) and \( \text{N} + \text{C}_m\text{H}^- \) into the chemical network (see Section 2.2), the \( \text{CN}^- \) column density is raised from \( 1.3 \times 10^{10} \) to \( 1.3 \times 10^{12} \) cm\(^{-2} \). Due to the fact that \( \text{C}_7^- \) is the most abundant anion involved in these reactions, the dominant formation reaction for \( \text{CN}^- \) is \( \text{N} + \text{C}_7^- \rightarrow \text{CN}^- + \text{C}_6 \). These results are dependent on the values of the branching ratios assumed for the AED reactions. Even if the AED branching ratio is as large as 0.9 (instead of the assumed value of 0.5), the \( \text{CN}^- \) column density is calculated to be \( 2.7 \times 10^{11} \) cm\(^{-2} \) and \( \text{N} + \text{C}_7^- \) is still the dominant reaction for \( \text{CN}^- \) production.

\( \text{C}_3\text{N}^- \) was detected for the first time in IRC+10216 by Thaddeus et al. (2008), who report a column density of \( 1.6 \times 10^{12} \) cm\(^{-2} \). If \( \text{C}_3\text{N}^- \) is assumed to be formed only by radiative electron attachment to \( \text{C}_3\text{N} \), the modeled column density is \( 1.1 \times 10^{10} \) cm\(^{-2} \), with a corresponding anion-to-neutral ratio of 0.02%. This would imply that the electron attachment rate used (calculated by Petrie & Herbst 1997) is too small. However, upon inclusion of reactions between nitrogen atoms and anions, the modeled \( \text{C}_3\text{N}^- \) column density rises to \( 9.0 \times 10^{11} \) cm\(^{-2} \) (which corresponds to an anion-to-neutral ratio of 1.3%), in reasonable agreement with the observed value. The dominant reactions involved in the production of \( \text{C}_3\text{N}^- \) in our present model are \( \text{N} + \text{C}_7^- \) \((n = 5, 6, 7) \). If these reactions do indeed dominate the \( \text{C}_3\text{N}^- \) synthesis then the rate of radiative electron attachment to \( \text{C}_3\text{N} \) calculated by Petrie & Herbst (1997) may still be accurate.

Cyanopolyynes \( \text{C}_n\text{N} \) (for \( n > 4 \)), by analogy with the structurally similar linear hydrocarbons \( \text{C}_n\text{H} \) \((n > 5) \) studied by Herbst & Osamura (2008), have been assumed to undergo rapid radiative stabilization upon attachment of a free electron so that the rate of radiative electron attachment used in the model is \( 1.25 \times 10^{-7}(T/300)^{-0.5} \) cm\(^{-3} \) s\(^{-1} \). Consequently, \( \text{C}_5\text{N}^- \) and \( \text{C}_7\text{N}^- \) have large modeled column densities of \( 1.7 \times 10^{13} \) and

![Figure 6. Emission maps for hydrocarbons and their anions, calculated from the modeled radial intensity profiles assuming spherical symmetry of the CSE. The grayscale is set to one at the respective emission maxima of each plot. Spatial units are arcseconds from the central star.](image-url)
anion-to-neutral column density ratios for C4H, C6H, and C8H to determine whether Equation 3 is indeed the dominant production route. Direct comparison of (unresolved) radii, which would cause the molecule to exist in a higher state of rotational excitation. Dense shells in the inner envelope have a significant impact on the C2H− radial abundance profile, as shown in Figure 7. C2H− is concentrated in the region of the envelope around the r = 1″ shell. The presence of this shell raises the C2H− column density from 2.3 × 10^{12} to 5.5 × 10^{12} cm^{-2}. This anion is unusual in the model because its rate of formation by radiative electron attachment is very slow; it is produced predominantly in the inner envelope as a result of Equation 3 (H− + C2H2 → C2H− + H2). This may have important observational consequences because the kinetic temperature of the gas is higher at such radii, which would cause the molecule to exist in a higher state of rotational excitation. Direct comparison of (unresolved) single-dish microwave observations of C2H− and C2H may be made more difficult by the difference in the distributions and therefore the telescope beam-filling factors of these two species. Detailed microwave observations of the spatial distributions of C2H− are clearly required in order to confirm this result and determine whether Equation 3 is indeed the dominant production mechanism for this species.

In the new model for the CSE of IRC+10216, the respective anion-to-neutral column density ratios for C4H, C6H, and C8H are calculated to be 1.4%, 7.4%, and 4.5%. The corresponding observational ratios are 0.024%, 6.3% (Cernicharo et al. 2007), and 26% (Remijan et al. 2007). As highlighted above for C2H, obtaining accurate observational ratios is hindered by the possibility that the anions and neutrals have different spatial distributions within the CSE. Nevertheless, it is clear that our model still severely overestimates the amount of C4H− compared with C4H (see also Herbst & Osamura 2008). It may be the case that the C4H radiative electron attachment rate calculated by Herbst is too large. Alternatively, C4H− may be destroyed more rapidly than our model currently predicts, for example, by reaction with HCN or C2H2 (e.g., Equations (2) and (3)). In that case, such reactions would have to be much more rapid for C3H− than for C2H− and C4H−, which could conceivably result from the different electron binding energies of these species (E. Herbst 2008, private communication). Further laboratory and/or theoretical studies are required in order to determine whether the organic anions considered here react with HCN, C2H2 or other possible proton-donating molecules.

Anion-to-neutral ratios of the Cn and CnH species are typically slightly greater in the present model compared to the no-shells (NS) model. This is largely attributable to the increased electron abundances inside the dominant 15″ shell which leads to increased rates of electron attachment. Figure 8 shows the electron abundances as a function of radius. The density-enhanced shells cause increased shielding of the gas from photoionization which results in electron abundances which are a factor ~2 lower between r = 20″ and 100″. Inside the shells at these radii, contrary to the 15″ shell, the electron densities are up to an order of magnitude lower than the surrounding CSE. Beyond ~100″ where the density becomes very low, the shells begin to have a negligible effect on the photoionization rate as shown for atomic carbon in Figure 8. Accordingly, the electron abundances of the models with and without shells converge at this radius.

### 3.3. MgNC

A number of metal-containing molecules have been detected in IRC+10216. While some of these, for example NaCl, KCl, and AlCl, are expected to be abundant in the local thermodynamic equilibrium region and have spatial distributions that...
peak on the star, other species, particularly the cyanides and isocyanides such as MgCN, MgNC, and AlNC, have shell-like distributions on scales very similar to the cyanopolyynes (Ziurys et al. 2002). Two possibilities exist to account for the presence of these species in the outer envelope: either they form in the gas phase from metals that have not been incorporated into dust in the inner envelope or they have been released from grains by erosion processes in the outer envelope.

In order to test the hypothesis that metal isocyanides are formed by gas-phase chemistry, we have investigated the synthesis of MgNC via radiative association reactions between Mg$^+$ and the cyanopolyynes (Dunbar & Petrie 2002):

$$\text{Mg}^+ + \text{HC}_n\text{N} \rightarrow \text{HC}_n\text{NMg}^+ + h\nu$$

(5)

(with $n = 3, 5, 7, 9$), followed by dissociative recombination reactions of the form

$$\text{HC}_n\text{NMg}^+ + e^- \rightarrow \text{MgNC} + \text{C}_{n-1}\text{H}.$$  

(6)

Figure 9 shows the radial abundance (cm$^{-3}$) of MgNC in both the presence and absence of shells. In these calculations, we adopt an initial Mg/H$_2$ abundance ratio of 10$^{-5}$—the derived abundances and column densities are directly proportional to this ratio. The effect of the density-enhanced shells is, once again, to move the peak of the distribution outward and to concentrate the abundance to the peaks of the gas density, in this case with almost equal peaks in the 15$''$ and 29$''$ shells. The total MgNC column density is calculated to be 5.7 \times 10^{13} \text{ cm}^{-2}, compared to observed values of (0.93–5) \times 10^{13} \text{ cm}^{-2} (Guélin et al. 1995; Higbeberger & Ziurys 2003). The dominant formation reaction is with HC$_2$N which, despite its smaller abundance, has the largest radiative association rate coefficient (Figure 5), 6.59 \times 10^{-5} (T/300)^{-0.47} \text{ cm}^3 \text{ s}^{-1} (Dunbar & Petrie 2002).

It was suggested by Dunbar & Petrie (2002) that the inclusion of Equation 5 into chemical models for IRC+10216 might significantly reduce the abundances of the species HC$_n$N and help to reconcile the discrepancy between modeled and observed HC$_n$N column densities. In the present model, we find that the reaction of Mg$^+$ with HC$_3$N, HC$_7$N, and HC$_9$N results in only \sim 1\% reduction in their calculated column densities, which does not significantly improve the match with observations. However, the depletion of these species is dependent (in a roughly linear fashion) on the initial Mg abundance employed.

4. DISCUSSION

Compared to previous chemical models of IRC+10216 (e.g., Nejad & Millar 1987; Cherchneff et al. 1993; Millar et al. 2000; Brown & Millar 2003; Agúndez et al. 2008), the model presented here is unique in the inclusion of density-enhanced shells of gas and dust. The shells included in our model have physical parameters similar to the dust shells observed by Mauron & Huggins (2000). As expected, the modeled column densities differ from those calculated in the model by Millar et al. (2000, referred to hereafter as MHB). Table 3 gives the column densities for these two models for 33 species for which observational column densities have been published. To highlight the effect of the density-enhanced shells on the chemistry and to permit comparison with models without density-enhanced shells, the column densities from the present model with no density-enhanced shells (referred to as NS), are also given. The column densities calculated by the three different models are generally in good agreement with observations, especially given the complex morphology of the source.

Notable differences between the MHB and NS models include a substantial reduction in the HC$_3$N/HC$_2$N column density ratio in the new model due to the enhanced photodissociation rate used for HC$_3$N (from the RATE06 database). The C$_3$N abundance is reduced as well because it is produced predominantly from HC$_5$N (via an alternative photodissociation channel). The carbon chains C$_6$H, C$_7$H, and C$_9$H, the cyanopolyynes HC$_4$N, HC$_7$N, and HC$_9$N and also CN are significantly more abundant in the present models than MHB. This is primarily due to the increased initial abundances of the parent species C$_2$H$_2$ and HCN (Fonfria et al. 2008) to which their chemistry is closely coupled (see Millar & Herbst 1994). C$_6$H and C$_8$H are also more abundant as a result of the addition of C$_6$H$^+$ to the chemical network which undergoes associative detachment with H to form C$_6$H$_2$. C$_6$H$_2$ reacts with C to produce C$_7$H or with C$_2$H to produce C$_8$H$_2$, which is photodissociated to C$_8$H.

Among the 33 chemical species listed in Table 3, 12 show calculated column densities that differ from observations by more than a factor of 10. In the present model, the moderate-to-large-sized hydrocarbons and cyanopolyynes C$_4$H$_2$, C$_5$N, C$_6$H, C$_8$H, and HC$_9$N might be considered to have column...
It has been brought to our attention by the referee that in a recent study of narrow submillimeter emission lines from IRC+10216 by Patel et al. (2008), a lower limit to the abundance of CS in the inner envelope of $9.3 \times 10^{-6}$ was derived. This is significantly greater than our adopted abundance of $4.0 \times 10^{-6}$. Thus, we have investigated the impact on our model of raising the initial CS abundance to $1.0 \times 10^{-5}$. We find that the only species to be appreciably affected are those containing sulfur and those whose chemistries are closely related to the sulfur-containing molecules. The column densities of HCS, OCS, H$_2$CS, and $C_n$S (for $n = 1-5$) scale approximately linearly with the initial CS abundance. The $C_3$ and $C_3H$ column densities are raised by about 50%, mainly as a result of the photodissociation of CS to produce $C_3 +$ CS. The OH column density increases by a similar factor due to the hydrogen-transfer reaction of HCS with O. No other species in Table 3 have column densities that are significantly affected by the increase in the initial CS abundance.

The inclusion of density-enhanced shells in the model has a profound impact on the calculated radial abundance distributions, both through the extinction effect of the shells which modifies the photochemical reaction rates, and through the effects of their increased densities which accelerate the binary chemical reactions. However, the impact of the shells on the total column densities is generally rather small; even though the chemical abundances may vary by over an order of magnitude inside the shells, the fact that the shells are narrow compared to the radial extent of the CSE means that they contribute toward only a modest fraction of the total column densities. Species whose chemistries are closely linked to the radiation field strength are most affected by the shells, for example $C_3H_4$ and CH$_2$CHCN (whose dominant destruction channels are by photodissociation), are shielded from dissociating radiation out to a larger radius by the increased extinction. The enhanced shielding in the envelope also reduces the ionization rate, resulting in up to an order of magnitude reduction in the electron and $C^+$ column densities at certain radii.

A primary motivation of this study is to examine whether the structure observed by Dinh-V-Trung & Lim (2008) in HC$_3$N and HC$_3$N maps of IRC+10216 is consistent with their suggestion that the gas and dust are coupled, sharing a similar density distribution. The inclusion of density-enhanced shells in our model results in peaks in the radial number density distributions of molecules within the shells, which shows that density enhancements in the molecular gas are able to produce small-scale structure in the molecular distributions similar to those observed.

When molecular excitation is taken into account, the emission intensity profiles for $C_2H$, $C_2H$, $C_3H$, and $C_3N$ all peak inside the dense shell at 15″. This finding is consistent with the 3 mm maps of these species made by Guélin et al. (1999) and Lucas & Guélin (1999) and shows that the presence of dense shells can cause the modeled molecular abundances to peak around the same radius and in a narrow region corresponding to the radius and thickness of the density enhancement. The simulated emission maps (Figure 6) show a similar picture for $C_2H$ and $C_3N$. However, the inner broad intensity maximum of the $C_2H$
radial profile results in a ∼5′ wide, intense emission ring inside of the density-enhanced 15″ ring, which is not seen in the Guélin et al. (1999) map. Possible explanations for this discrepancy might be that the excitation of C$_2$H favors its observation in the narrow shell (contrary to the crude excitation analysis we performed), or that the interferometer used in the observations failed to detect the broader structure of the inner ring.

In the model, the gas and dust share identically shaped density profiles which requires that the gas and dust must be dynamically coupled. This is contrary to the theory (see Truong-Bach et al. 1991; Mauron & Huggins 2000) that radiation pressure accelerates the circumstellar dust to a radial drift velocity ∼2 km s$^{-1}$ faster than the gas. Possible coupling mechanisms may include thermal and turbulent motion of the gas and dust. Future models may need to include a detailed analysis of the coupling between the gas and dust shells.

The shells used in the model represent a gross simplification of the dust shell structure observed in IRC+10216 by Mauron & Huggins (2000). However, the results presented here—that the maximum abundances of species containing carbon chains match the peaks in the circumstellar dust distribution—should be generally applicable to more complex representations of the density structure of the CSE. The density-dependent nature of molecular excitation means that detailed excitation calculations are required to determine the relationship between observed emission intensity and molecular abundance distributions. In our model, the temperature distribution of the gas is assumed to be continuous, with the assumption that the density-enhancements are in thermal equilibrium with the surrounding medium. More detailed observations of the gas and dust will be required to determine if this assumption is realistic. In addition, accurate collisional and vibrational excitation rates will be required in order to more accurately calculate the rotational excitation of the molecules considered.

5. CONCLUSION

Density-enhanced shells of gas and dust have a significant impact on the calculated radial distributions of molecules in the expanding envelope of IRC+10216. Based on the suggestion by Dinh-V-Trung & Lim (2008) that the gas and dust are coupled, we included in a new model for the CSE a set of density enhancements with parameters based on the dust shell observations by Mauron & Huggins (2000). Photochemistry is delayed to greater radii due to increased shielding by the dust shells. Density enhancements in the gas result in molecular abundances that peak inside the narrow shells, showing that the clumpy structure observed in HC$_3$N and HC$_5$N may be the result of density enhancements in the molecular gas. The calculated emission intensity profiles for C$_2$H, C$_3$H, and HC$_3$N all peak within a narrow band about 15″ from the central star, which is consistent with detailed emission maps of these species. The emission profile for C$_3$H$_2$ has a broad maximum around 8″ that is not present in observed maps, which may be indicative of a need for further improvements in the model.

The ethynyl anion C$_2$H$^-$ and the nitrile anions C$_{2n}$−1N$^-$ (for n = 1, 2, 3, 4) have been calculated to reach observable abundances in the CSE. For the smaller nitrile anions for which radiative electron attachment is slow (i.e., CN$^-$ and C$_2$N$^-$), this result is uncertain due to the unknown product branching ratios in the AED channels of the reactions N + C$_2$H and N + C$_{2n}$H$^-$.

Further laboratory measurements (including up to at least m = 10) will be required in order to confirm the importance of these reactions in anion astrochemistry. C$_2$H$^-$ is predicted to be produced in abundance in the inner CSE (at much smaller radii than the other anions in the model), as a product of the reaction of H$^-$ with C$_2$H$_2$. Other possible proton transfer reactions of this kind (i.e., XH + Y$^-$ → X$^-$ + YH$^-$, for the anions listed in Table 2) also need to be considered for possible inclusion in future anion chemical networks.

Our chemical models produce MgNC with a peak abundance in the outer envelope. Dependent on the initial Mg abundance used, the observed MgNC column density matches observation, which shows that gas-phase chemistry is a viable route to the formation of this species.

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