A COMPREHENSIVE SURVEY OF HYDROGEN CHLORIDE IN THE GALAXY

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ABSTRACT

We report new observations of the fundamental J = 1−0 transition of HCl (at 625.918 GHz) toward a sample of 27 galactic star-forming regions, molecular clouds, and evolved stars, carried out using the Caltech Submillimeter Observatory. Fourteen sources in the sample are also observed in the corresponding H(^37)Cl J = 1−0 transition (at 624.978 GHz). We have obtained clear detections in all but four of the targets, often in emission. Absorptions against bright background continuum sources are also seen in nine cases, usually involving a delicate balance between emission and absorption features. From RADEX modeling, we derive gas densities and HCl column densities for sources with HCl emission. HCl is found in a wide range of environments, with gas densities ranging from 10^5 to 10^10 cm^{-3}. The HCl abundance relative to H_2 is in the range of (3–30) x 10^{-10}. Comparing with the chlorine abundance in the solar neighborhood, this corresponds to a chlorine depletion factor of up to ~400, assuming that HCl accounts for one-third of the total chlorine in the gas phase. The [35Cl]/[37Cl] isotopic ratio is rather varied, from unity to ~5, mostly lower than the terrestrial value of 3.1. Such variation is highly localized, and could be generated by the nucleosynthesis in supernovae, which predicts a _3Cl deficiency in most models. The lower ratios seen in W3IRS4 and W3IRS5 likely confine the progenitors of the supernovae to stars with relatively large mass (\gtrsim 25 M_\odot) and high metallicity (Z \sim 0.02).

Key words: ISM: abundances – ISM: molecules – submillimeter: ISM

1. INTRODUCTION

Initially of interest as a potential coolant for molecular clouds, hydrogen chloride (HCl) was first detected in the J = 1−0 transition at 625.918 GHz (De Lucia et al. 1971) toward IRc2 in Orion A, in emission, by Blake et al. (1985), and in absorption against dust continuum in Sgr B2, by Zmuidzinas et al. (1995), both on board the NASA Kuiper Airborne Observatory with a beam width of 2′.3. Ground-based observations at a higher spatial resolution of 12″ were carried out using the 10.4 m telescope of the Caltech Submillimeter Observatory (CSO) by Schilke et al. (1995) and Salez et al. (1996) toward OMC-1 and MonR2. Salez et al. (1996) also detected the H(^37)Cl J = 1−0 transition toward OMC-1. These studies found the HCl abundance to be 10^{-9} to 10^{-10} relative to molecular hydrogen, indicating that chlorine is depleted by a factor of several hundred compared to its solar neighborhood abundance and that HCl is not a significant coolant of molecular clouds. Salez et al. (1996) also reported a [35Cl]/[37Cl] isotopic ratio that is 1.3–2 times higher than the solar neighborhood value, indicating variations of the isotopic ratio in the Galactic interstellar medium.

Chemistry of chlorine and hydrogen chloride in the interstellar medium has been investigated by a number of authors (Jura 1974; Dalgarno et al. 1974; Blake et al. 1986; Schilke et al. 1995; Neufeld & Wolfire 2009), establishing a relatively simple chemical network involving less than a dozen other atomic and molecular species, as well as electrons and cosmic-ray particles, in the production and destruction of HCl. In dense molecular clouds, such a network yields, among other species, predominantly HCl, taking up about one-third of the chlorine in the gas phase (Schilke et al. 1995; Neufeld & Wolfire 2009).

HCl J = 1−0 transition falls on the shoulder of a relatively strong atmospheric water absorption line. Consequently, ground-based observations of HCl have been limited to the rare occasions of superb weather at high sites, such as Mauna Kea, and relatively little is known about HCl in different categories of objects, such as evolved stars, active star-forming regions, molecular clouds with no apparent star formation activity, etc. In particular, the [35Cl]/[37Cl] isotopic ratio is of interest in different categories of objects, as variations of this ratio can be produced through nucleosynthesis in massive stars followed by mixing of the processed material into the surrounding molecular cloud (Salez et al. 1996). The chlorine isotopic ratio could thus offer insights into the evolutionary history of the molecular material.

To address these questions, we took advantage of several periods of superb weather on Mauna Kea to conduct a comprehensive survey of HCl in a sample of well-known objects. In the following sections, we first describe the observational details, then present the results and data analysis, followed by discussions and conclusions.

2. OBSERVATIONS

Observations described here were carried out on 2008 September 11, 2010 March 2, and April 7–15 with the 10.4 m CSO telescope. The 690 GHz facility heterodyne receiver was tuned to 625.918756 GHz and 624.977821 GHz, respectively, for observations of HCl J = 1−0 and H(^37)Cl J = 1−0 lines. The lines fall on the shoulder of a prominent atmospheric water absorption line, therefore requiring premium weather conditions. The atmospheric opacity measured at 225 GHz was mostly 0.03–0.04, and the typical double-side-band system temperature was 2500 K. There were marginal periods on 2010 April 7–15 when atmospheric opacity was around 0.05–0.06, resulting in a typical system temperature of 6000 K. Both the FFTS and
4 GHz AOS facility spectrometers were used in 1 GHz mode as backends.

The dish surface optimization system (Leong et al. 2006) was used to correct gravitational deformations of the telescope surface, maintaining a clean 13.5 beam over a wide range of elevations. Temperature calibration using a hot chopper wheel yields $T_A^*$, the antenna temperature corrected for atmosphere and telescope hot spillover. A main beam efficiency of ~50% measured on Jupiter and Uranus is used to convert $T_A^*$ to $T_{mb}$, the main beam brightness temperature. Telescope pointing was checked on Jupiter, Mars, and Uranus, and was found to be within 4″.

Observations were conducted in the beam switching mode at a frequency of 0.523 Hz and with a chop throw of 300″. Typical integration time on each source was 300 s, resulting in an rms of ~0.15 K at a velocity resolution of 0.31 km s$^{-1}$. The source selection was primarily based on its accessibility to reasonable elevations, with a secondary goal of covering a diverse collection of objects. Due to its high critical density ($\sim 10^6$ cm$^{-3}$), the excitation of the HCl $J = 1–0$ transition is most likely subthermal, and radiative excitation, in the presence of continuum emission, is likely to play an important role in the excitation of the transition. The presence of dense gas and/or continuum radiation in an object would likely yield a detection. In the end, 27 sources/positions were observed, among them were active star-forming regions, giant molecular clouds with no apparent star-forming activities, evolved stars, shocked gas in a supernova remnant (SNR), and ultra-compact H II regions. Table 1 lists observational details for all observed sources.

### 3. RESULTS

For sources with matching HCl and H$^3$Cl $J = 1–0$ spectra, we present in Figures 1–3 HCl spectra in the right column and H$^3$Cl spectra in the right column. The antenna temperature scale is kept the same for each source for easy comparison of the line intensities. The velocity scale, $V_{LSR} - V_0$, refers to the main hyperfine component; values of $V_0$ are denoted in each panel. Toward IRC2 in OMC-1, the HCl and H$^3$Cl line profiles are blended heavily with spectral lines from other species, in the case for H$^3$Cl predominantly methanol. It is not easy to untangle the blending and derive useful information from the resulting line profile. Instead we opted to observe two positions south of IRC2 with declination offsets of ~60° and ~100°, respectively. The two positions correspond to local maxima in the HCl map of OMC-1 by Schilke et al. (1995). The (0, ~60°) position also coincides with the CS clump LS1 (Mundy et al. 1986) and the (0, ~100°) position Orion S. Figure 4 shows the HCl $J = 1–0$ spectra for the remaining sources in the sample.

#### 3.1. Hyperfine Structure Fitting

The HCl hyperfine structure, resulting from the interaction between the electric field and chlorine nuclear spin, splits the $J = 1–0$ transition into three components, with the outer two separated by ~6.35 and 8.22 km s$^{-1}$, respectively, from the strongest middle component for H$^3$Cl. For H$^3$Cl, the separations are ~5.05 km s$^{-1}$ and 6.45 km s$^{-1}$, respectively. Statistical weights for the upper level of the hyperfine transitions dictate optically thin line strength ratio of 1:3:2. In the observed sample, the three-component hyperfine pattern is clearly seen in many sources, but the components often do not conform to the optically thin ratio, indicating deviations from the optically thin limit or differences in excitation between the three components.

Under the assumption that the hyperfine components share the same excitation and kinematic characteristics, fits to the line profiles can independently provide the optical depth and excitation temperature (Kwan & Scoville 1975). Such a utility is readily available in CLASS, the spectral line data reduction package developed and maintained by IRAM. The hyperfine structure (HFS) fit in CLASS provides four parameters: $\Delta T$, $\tau_{main}$, $V_{LSR}$, w the line width, and $\tau_{main}$ is the total optical depth from all hyperfine components of the line, and $\Delta T_0 = T_{ex} - T_{bg}$ is the difference between the line excitation temperature and background temperature. Table 2 lists the results of such hyperfine structure fits for sources in our sample that show the $J = 1–0$ HCl or H$^3$Cl emission. Cases involving absorption are more complex, as seen in G34.26+0.15 and W43–MM1 in Figure 3, for example. They are often the result of a delicate balance between emission and absorption features. We defer the treatment and discussions for sources with absorption to a separate paper (H. Yoshida et al. 2010, in preparation).

In fitting the hyperfine profiles in $T_{mb}$ scale, one needs to be mindful that the line profile may suffer from beam dilution when a source is smaller than the telescope beam. The uncorrected line temperatures would affect the excitation temperature and optical depth derived from HFS fit of the HCl line profile. Thermal dust emission as seen in submillimeter continuum maps provides a good measure for gas distribution in the source. We compare the size of the 350 μm dust beams to the telescope beam of 13.5′ to determine whether the HCl emission from a given source suffers appreciable beam dilution. Most of our sources in the sample are observed by Dotson et al. (2010) with the SHARP camera at the CSO, and the FWHM of the dust cores ranges from ~30′′ to ~60′′. We therefore do not consider beam dilution a significant source of error for our sample of objects in deriving the excitation temperature and optical depth for the HCl $J = 1–0$ transition. One exception
Figure 1. HCl and H$^{37}$Cl $J = 1-0$ spectra toward W3IRS4, W3IRS5, W3(OH), and two positions in OMC-1 offset from IRc2 in declination by −60′′ and −100′′, respectively. Velocity scale, $V_{\text{LSR}} - V_0$, refers to the main hyperfine component.

may be G5.89−0.39, where the Smithsonian Millimeter Array (SMA) continuum image at 870 μm reveals a dust core ∼6′′–8′′ in size (Tang et al. 2009), implying a factor of ∼3 in beam dilution in the observed line temperature.

The HFS fits are sensitive to the signal-to-noise ratio (S/N) in the observed line profiles. $\Delta T$ and hence $\tau_{\text{main}}$ are not well constrained for relatively noisy line profiles. For sources with both HCl and H$^{37}$Cl detections, one would expect the values of $\Delta T$ derived from HFS fits of the two transitions to be very close. In reality, they tend to agree within error bars, albeit rather large error bars in some cases, resulting in questionable estimates of the optical depths. We attempted to derive alternative estimates of the optical depths for both HCl and H$^{37}$Cl lines by imposing a fixed value of $(\Delta T)^*$, a weighted average of the values obtained...
for the two lines. Using \( (\Delta T)^* \) and the \( T_{\text{mb}} \) from Gaussian fits of the three components in each profile, we derive \( \tau (\text{HCl})^* \) and \( \tau (\text{H}^{37}\text{Cl})^* \) as listed in Table 2. The error of \( T_{\text{mb}}/(\Delta T) \) amplified by \( e^\tau \) translates into the error of \( \tau^* \) for each component, resulting in relatively large errors for \( \tau^* \) derived from this approach when individual components are optically thick. We nonetheless consider \( \tau^* \) a more reliable measure of the optical depth than that obtained directly from HFS fits. For sources that have only HCl observations, \( (\Delta T)^* \) and \( \tau (\text{HCl})^* \) in Table 2 are from direct HFS fits. For G5.89–0.39, we also present fits corrected for the likely case of beam dilution. Compared to the uncorrected case, the differences are mostly in the line excitation temperatures, with optical depth remain largely unchanged.

IRC+10216 shows a rather distinctive line profile. It is likely the result of convolution of the hyperfine structure with a shell profile often seen in circumstellar envelopes. As an
approximation, we chose to fit two Gaussian profiles for IRC+10216 at local standard of rest (LSR) velocities of $-17.8$ km s$^{-1}$ and $-27.4$ km s$^{-1}$, respectively, to simulate the shell profile in the hyperfine fits. As shown by the red curve in the appropriate panel in Figure 2, this approach adequately fits the line profile, yielding a solution that indicates that the HCl emission originated from a circumstellar shell with an expansion velocity of $4.8$ km s$^{-1}$, compared to an observed terminal velocity of $14.5$ km s$^{-1}$ (see, for example, Schöier & Olofsson 2001). The $[\text{H}^{35}\text{Cl}]/[\text{H}^{37}\text{Cl}]$ isotopic ratio in the shell is likely in excess of 4 from the optical depth shown in Table 2.

CIT6 is widely regarded as similar to IRC+10216 with comparable chemical and physical characteristics, but a lower dust-to-molecular mass ratio (Zhang et al. 2009). HCl is no exception in that regard. It is clearly detected but the line is too weak to allow for a good HFS fit. Nevertheless, this observation serves as an additional proof that HCl as a species exists in detectable quantities in circumstellar shells of evolved stars. IC443G1 is a dense core in a ring of shocked material associated with SNR IC443 (van Dishoeck et al. 1993). There are tantalizing signs of HCl detection, but the line is again too weak to derive any useful parameters.

G5.89−0.39 is another interesting case with a rather distinctive line profile. Both HCl and $^{37}\text{Cl}$ show a pronounced red wing, indicating the presence of higher velocity components. A three-component fit, as shown by the red curve in the top left panel of Figure 3, represents an excellent match to the observed line profile. As listed in Table 2, the three components are at LSR velocities of 9.3, 17.9, and 38.3 km s$^{-1}$, with FWHM line widths of 4.9, 13.5, and 5.4 km s$^{-1}$, respectively. From the interferometric study of Hunter et al. (2008), the broad ($\sim13$ km s$^{-1}$) line emission is seen in $^{34}\text{S}$, SiO, CH$_3$OH, HC$_3$N, $^{31}$O, and H$^{13}$CO$^+$ toward the dust emission source SMA-N, in the north–northeast extension of the molecular ridge that bounds the northern and western edges of the UCHII. The 9.3 km s$^{-1}$ feature likely originates from the northwest and southwest.

Figure 3. HCl and $^{37}\text{Cl}$ $J = 1−0$ spectra toward G5.89−0.39, W49N, W51FIR2, and DR21(OH). Velocity scale, $V_{\text{LSR}} - V_0$, refers to the main hyperfine component. For sources with detectable continuum, baseline is not removed to show the continuum level.
Figure 4. HCl spectra toward rest of the sources in the sample.
sections of the molecular ridge. While the origin of the 38.3 km s\(^{-1}\) feature is not immediately clear, velocity features as highly redshifted as 78 km s\(^{-1}\) and blueshifted as \(-61\) km s\(^{-1}\), mostly from various masers, were quoted by these authors in establishing a north–south bipolar outflow. In deriving the alternate optical depth of these velocity components in Table 2, we opted to use the \(\Delta T\) from the HCl HFS fits for the 17.9 km s\(^{-1}\) and 38.3 km s\(^{-1}\) components due to their low S/N ratio in the H\(^{37}\)Cl line profile. From the derived \(\tau^*\) in Table 2, the [HCl]/[H\(^{37}\)Cl] isotopic ratio in the source is most likely around 2, though the broad emission feature may have a higher ratio on the order of 3–4.

Sgr A\(^*\), the dynamical center of the Galaxy, yields no HCl detection to an rms level of 0.1 K at a resolution of 0.31 km s\(^{-1}\). Similarly, G1.6–0.025, a giant molecular cloud in the Galactic center region with no apparent sign of star formation, also yields a non-detection. Sgr B2(M) and Sgr B2(N), on the other hand, show broad absorption at both HCl and H\(^{37}\)Cl \(J = 1\rightarrow 0\) transitions.

From Table 2, it is clear that the [H\(^{35}\)Cl]/[H\(^{37}\)Cl] isotopic ratio, based on that of the optical depth, is rather varied, from 4 and over in Mon R2, IRC+10216, and DR21(OH), to around 3 in OMC-1, W3(OH), and IRAS16293, down to around 2 in G5.89–0.39, and to nearly reaching parity in W3IRS4 and W3IRS5. In comparison, the terrestrial [\(^{35}\)Cl]/[\(^{37}\)Cl] abundance ratio is \(\sim 3.1\).

3.2. RADEX Simulations

In addition to the optical depth obtained from hyperfine structure fitting, we can derive other physical parameters from the line profiles by running large velocity gradient (LVG) simulations to fit the three HCl hyperfine components. RADEX, a computer program developed and maintained by van der Tak et al. (2007) for fast non-LTE analysis of interstellar line spectra, was used in the LVG mode to find the best combination of gas density and HCl column density that could reproduce the observed line temperatures under given gas temperature.

The molecular data file for HCl under RADEX includes the hyperfine levels of the lowest eight rotational levels, up to 839 K above ground level, with rates for radiative and collisional transitions among them. The collisional rates between HCl and H\(_2\) are scaled from those of HCl and helium (Neufeld & Green 1994). The average Galactic background radiation field in RADEX simulations was run in off-line mode to cover a large parameter space, and relative \(\chi^2\) was sought between the observed line temperatures and the model results relative to the observed line temperature. Under the assumption that the observed HCl emission suffers no significant beam dilution, this process yields for a given gas temperature the most likely combination of gas density and HCl column density to reproduce the observed line strength of the hyperfine components. Figure 5 shows a RADEX simulation run for OMC-1 (0, −60\(^\circ\)) with a gas temperature of 45 K. Contours are the relative \(\chi^2\) starting from...
between $F_0$ and $F_{\pm 1}$. The solid smooth line and two dotted lines mark the observed optical depth of the HCl line and its quoted errors for HCl optical depth in Table 2 exceed 30%, we generally use 30% to set the error bar for the optical depth. The cross section between the $\chi^2$ contours and the optical depth curves clearly confines gas density and HCl column density in an often tight range.

Table 3 lists for each source the adopted gas temperature and $H_2$ column density, the gas density and HCl column density derived from RADEX simulations, as well as the HCl abundance. Since there is no molecular data file for $H^{37}Cl$ to use under RADEX, we treat it the same as HCl and fit the $H^{37}Cl$ abundance. Since there is no molecular data file for $H^{37}Cl$ to use under RADEX, we treat it the same as HCl and fit the $H^{37}Cl$ abundance. Since there is no molecular data file for $H^{37}Cl$ to use under RADEX, we treat it the same as HCl and fit the $H^{37}Cl$ abundance. Since there is no molecular data file for $H^{37}Cl$ to use under RADEX, we treat it the same as HCl and fit the $H^{37}Cl$ abundance.

4. DISCUSSION

4.1. Where to Look for HCl Emission?

From Table 3 it is clear that HCl emission is associated with warm dense gas with a density on the order of $10^8$ cm$^{-3}$. Given a critical density of a few $\times 10^8$ cm$^{-3}$, molecular excitation is mostly sub-thermal, dominated by radiative processes rather than collisions with $H_2$. Sources with strong continuum and/or dust emission would likely yield strong HCl line emission.

The derived gas densities for the two OMC-1 cores appear to be somewhat higher than $\sim 5 \times 10^6$ established by the multi-transitional study of $HC_3N$ (Bergin et al. 1996). This may...
indicate that the gas temperatures adopted for these cores for the RADEX simulations should have been higher. Another factor may be the average Galactic background radiation field used in RADEX for radiative excitation. As HCl transitions are shortward of ~480 μm in wavelength, radiation from warm dust would contribute significantly to the molecular excitation. Including local warm dust in the radiation field for RADEX simulation would produce a more precise picture of the gas where HCl emission originates.

The non-detection toward Sgr A* likely reflects a genuine lack of HCl molecules in the circumstellar disk surrounding Sgr A*. Latvakoski et al. (1999) show a warm dust ring ~80″ in size, as well as the “minispiral” within the ring. The dust ring coincides with a molecular ring with sharp inner edges seen in HCN (Güsten et al. 1987). With our 13′′ telescope beam, we are clearly probing the molecular cavity around Sgr A*, yielding an upper limit of 8 × 10^{-11} for HCl abundance relative to H_2 toward Sgr A*, markedley lower than those listed in Table 3.

The non-detection in G1.6−0.025 is intriguing. The source is a giant molecular cloud located in the Central Molecular Zone (CMZ) of the Galaxy, a region characterized by elevated gas temperature (typically ~70 K), higher gas density (>10^3 cm^{-3}), and highly supersonic internal velocity dispersion (~15–50 km s^{-1}; Morris & Serabyn 1996). Dense cloud cores are seen in various molecular tracers (Gardner & Whiteoak 1981; Gardner & Boes 1987; Whiteoak & Peng 1989; Kuiper et al. 1993; Menten et al. 2009), yet there is little sign of apparent star-forming activity in the cloud. For the extended 50 km s^{-1} cloud of G1.6−0.025, we derive a C^{18}O column density of 2 × 10^{12} from the C^{18}O J = 2-1 line spectrum taken at the CSO, assuming LTE at a gas temperature of 30 K (Menten et al. 2009). Again with a C^{18}O abundance of 1.7 × 10^{-7}, the cloud has an H_2 column density of 1 × 10^{22} cm^{-2}. Similar to Sgr A*, we obtain an upper limit of 7 × 10^{12} for HCl column density by modeling an HCl line of 0.1 K with 28 km s^{-1} in line width, yielding an HCl abundance of 7 × 10^{-10} relative to H_2. It would look as though the HCl non-detection in the 50 km s^{-1} cloud of G1.6−0.025 is the result of a combination of relatively low H_2 column density and HCl abundance.

It would be interesting to see if other clouds in CMZ have similarly low HCl abundance seen in Sgr A* and the 50 km s^{-1} cloud of G1.6−0.025. A prime target would be M-0.02-0.07, the so-called 20 km s^{-1} cloud ~2″ northeast of Sgr A*. A common trait among the three clouds is the likely interaction with an SNR (Menten et al. 2009). It is not clear though how such interaction would lead to relatively low production of HCl.

In the same vein, IC443 Clump G1 may represent another case where chemistry plays an important role in the detection of HCl. Clump G1 is a dense core in a ring of shocked material associated with SNR IC443 (van Dishoeck et al. 1993). It has a relatively rich chemistry compared to the pre-shock gas, but lacks many of the more complicated molecular species, such as methanol. These authors argued that such deficiency could be the result of the disruption of the chemical network brought on by the infusion of atomic hydrogen to the shocked gas by repeated shocks, as atomic hydrogen rapidly destroys many of the intermediate molecules in the reaction network. To derive an estimate of HCl column density from the low signal-to-noise detection, we use RADEX simulation to reproduce an HCl line of 0.2 K with an FWHM of 10 km s^{-1} (van Dishoeck et al. 1993). The resulting HCl column density of 3 × 10^{12} cm^{-2}, when compared to an H_2 column density of 1 × 10^{22} (van Dishoeck et al. 1993), leads to an HCl abundance of 3 × 10^{-10}. Another, the weak HCl line emission from IC443 Clump G1 may be the result of relatively low HCl abundance and H_2 column density. More observations of similar cores are needed to confirm that shocked gas may have a somewhat lower HCl abundance compared to other similarly dense gas in star-forming regions, PDRs, and ultra-compact HII regions.

Ultra-compact HII regions with reasonably high emission measures have proven to be a treasure trove for detecting HCl absorption (see Figure 4). Such cases typically involve a delicate balance between an emission feature and an absorption feature. Our forthcoming paper (H. Yoshida et al. 2010, in preparation) will discuss these cases.

4.2. The Case of IRC+10216

In Figure 2, we have shown that the HCl line profile for IRC+10216 could be adequately fit by two Gaussians centered at −17.8 km s^{-1} and −27.4 km s^{-1}, respectively, indicating that the HCl emission is associated with a circumstellar shell with an expansion velocity of 4.8 km s^{-1}. The HCl emission is likely from a density-enhanced shell ~15″ from the central star as seen in H^{13}CNO (Schöier et al. 2007) and SiO (Schöier et al. 2006), and modeled by Cordiner & Millar (2009). Radiation from the central star, as well as from local dust, contributes significantly to the molecular excitation. We therefore consider the HCl abundance and [^{35}Cl/^{37}Cl] abundance ratio for IRC+10216 as listed in Table 3 highly uncertain. More sophisticated modeling efforts such as Schöier & Olofsson (2001) and Wyrowski et al. (2006) are required to correctly interpret the results for IRC+10216. Cernicharo et al. (2010b) recently reported an HCl abundance of 5 × 10^{-8} from their Herschel SPIRE and PACS observations of IRC+10216. It may be interesting to note that we derive an upper limit of 8 × 10^{-8} for the HCl abundance in CIT6 by treating the low signal-to-noise detection as a 0.2 K line with 20 km s^{-1} in line width. Similar to the procedure used in Sgr A* and G1.6−0.025, the resulting HCl column density is 5 × 10^{12} cm^{-2}, with an adopted gas temperature of 40 K and an H_2 column density of 6.3 × 10^{19} cm^{-2} (Zhang et al. 2009).

4.3. ^{35}Cl/^{37}Cl Abundance Ratio

As listed in Table 3, the ^{35}Cl/^{37}Cl abundance ratio varies from around unity to around 3 and over. Cernicharo et al. (2000) reported an X[^{35}Cl/^{37}Cl] ratio of 3.1 ± 0.6 from line intensity ratios of NaCl, KCl, and AlCl in IRC+10216. Recent Herschel HIFI observations show an X[^{35}Cl/^{37}Cl] ratio of 2.7 in NGC6334I (Lis et al. 2010) and 2.1 ± 0.5 in W3A (Cernicharo et al. 2010b). These are comparable to the terrestrial value of ~3.1. It is interesting to note that the ^{35}Cl/^{37}Cl abundance ratio is in good agreement with the integration line intensity ratio of the HCl and H^{35}Cl line, as predicted by the modeling efforts of Cernicharo et al. (2010a) for their Herschel HIFI observations of W3A.
One may argue that the low isotopic ratio observed in W3IRS4 might be the result of observing an unresolved, very optically thick source. DR21(OH), which is at a comparable distance (∼2.5 kpc versus ∼2.2 kpc for W3IRS4), with a comparable dust core size at 350 μm (Dotson et al. 2010) and HCl opacity, argues against this scenario.

Clearly, variations in HCl isotopic ratio are highly localized. Salez et al. (1996) suggested one likely origin for such variations through explosive nucleosynthesis, whereby enrichment of one isotopolog over the other, as synthesized in massive stars, is mixed into the surrounding molecular cloud through explosive processes such as supernova explosions.

Models for nucleosynthesis in Type Ia supernovae (SNe Ia; Travaglio et al. 2004 and references therein) and Type II supernovae (SNe II; Nomoto et al. 1997, 2006; Kobayashi et al. 2006) indeed show a general deficiency of 37Cl in the processed material in most cases. In SNe Ia, the 35Cl/37Cl abundance ratios vary from 3.5 to 5.4 among models of different ignition conditions in the central white dwarf before the thermonuclear runaway (Travaglio et al. 2004). In SNe II, the elemental yields of the nucleosynthesis depend on the mass and metallicity of the progenitor as well as the explosion energy. Kobayashi et al. (2006) show that the 35Cl/37Cl abundance ratios produced in SNe with high explosion energy (≥10E51 with E51 ∼ 1051 erg, so-called supernovae) are typically in the range of 1.1–2.8 for varying progenitor mass and metallicity, while the ratios vary in a larger range (to as high as 18) in normal SNe (with explosion energy ∼ E51). It is interesting to note that in normal SNe II with relatively large progenitor mass (≥25 M_☉) and metallicity (Z ∼ 0.02), the 35Cl/37Cl abundance ratios are consistently below unity to as low as ∼0.5 (Kobayashi et al. 2006). Given enough mixing with the surrounding molecular clouds, these ratios could support the observed range of the 35Cl/37Cl abundance ratio.

W3IRS4 and possibly W3IRS5 present interesting exceptions of relative 37Cl deficiency predicted in most nucleosynthesis models of SNe. This likely confines the progenitors of the SNe to be stars of relatively high mass (≥25 M_☉) and high metallicity (Z ∼ 0.02).

Kawabata et al. (2010) and Perets et al. (2010) recently reported the discovery of a new category of subluminous supernovae. Although the origin for such faint supernovae is still controversial, these authors reported the unusual composition in the supernova ejecta to be rich with helium and calcium. It would be interesting to examine the nucleosynthesis yields in this class of supernovae in regard to the 35Cl/37Cl abundance ratios.

5. CONCLUSION

We conducted a comprehensive search of HCl J = 1–0 line in the Galaxy. Of the 27 sources/positions observed, 14 show emission, 9 show absorption, 2 showed marginal detection in emission, and 2 are non-detections. Fourteen of the sources/positions were also observed in the H3Cl J = 1–0 transition. RADEX simulations show that HCl emission is mostly associated with warm dense gas, of order 100 cm−3. HCl abundance is fairly uniform, in the range 10−10 to a few ×10−9, in general agreement with previous studies of OMC-1 by Schilke et al. (1995) and Salez et al. (1996). However, 35Cl/37Cl abundance ratios are found to vary considerably from around unity in W3IRS4 and W3IRS5 to ~5 in DR21(OH). They are in good agreement to the integrated intensity ratios of the HCl and H37Cl line as suggested by Cernicharo et al. (2010a).

The variations of the 35Cl/37Cl abundance ratios are found to be highly localized. They could be supported by the varying yields of nucleosynthesis of supernovae from different progenitors. The low 35Cl/37Cl abundance ratios seen in W3IRS4 and W3IRS5 likely confine the progenitors of the supernovae to stars of relatively high mass (≥25 M_☉) and high metallicity (Z ∼ 0.02).

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