The Arecibo Arp 220 Spectral Census I: Discovery of the Pre-Biotic Molecule Methanimine and New Cm-wavelength Transitions of Other Molecules

C. J. Salter, T. Ghosh, B. Catinella\textsuperscript{1}, M. Lebron\textsuperscript{2}, M. S. Lerner, R. Minchin and E. Momjian\textsuperscript{3}

Arecibo Observatory, HC 03 Box 53995, Arecibo, PR 00612
csalter@naic.edu

ABSTRACT

An on-going Arecibo line search between 1.1 and 10 GHz of the prototypical starburst/megamaser galaxy, Arp 220, has revealed a spectrum rich in molecular transitions. These include the “pre-biotic” molecules: methanimine (CH$_2$NH) in emission, three $v_2 = 1$ direct l-type absorption lines of HCN, and an absorption feature from either $^{18}$OH or formic acid (HCOOH). In addition, we report the detection of two, possibly three, transitions of $\lambda$4-cm excited OH not previously detected in Arp 220 which are seen in absorption, and a possible absorption feature from the 6.668-GHz line of methanol. This marks the first distant extragalactic detection of methanimine, a molecule with high relevance to the origins of life. Also, if confirmed, the possible methanol absorption line presented here would represent the first extragalactic detection of methanol at a distance further than 10 Mpc. In addition, the strong, previously undetected, cm-wave HCN $v_2 = 1$ direct l-type lines will aid the study of dense molecular gas and active star-forming regions in this starburst galaxy.

Subject headings: Radio lines: galaxies, galaxies: starburst, galaxies: individual (Arp 220), galaxies: ISM

\textsuperscript{1}also, MPIfA, Karl-Schwarzschild-Straße 1, Postfach 1317, D-85741, Garching, Germany

\textsuperscript{2}also, Department of Physical Sciences, University of Puerto Rico, P.O. Box 23323, San Juan, PR 00931-3323

\textsuperscript{3}also, NRAO, Array Operations Center, P.O. Box O, 1003 Lopezville Road, Socorro, NM 87801-0387
1. Introduction

Star formation in galaxies occurs in two modes – “quiescent”, in which stars form at a relatively modest rate over a long period of time across the entire galaxy disk, and “starburst”, in which gas is turned into stars extremely rapidly, confined to compact regions of the galaxy, often its circumnuclear volume (Kennicutt 1998). Starbursts are often triggered by external dynamical disturbances such as galaxy mergers. The dust heating associated with these intense bursts of star formation within giant molecular clouds can produce hugely increased IR luminosity and conditions favorable for maser emission. The strong 18-cm OH maser emission in some of these galaxies is many orders of magnitude more luminous than its counterparts in our Galaxy (Baan et al. 1982; Baan 1989; Martin et al. 1989; Lonsdale et al. 1998; Darling & Giovanelli 2002). Ultra Luminous Infrared Galaxies (ULIRGs) are thought to be systems where all of these processes are occurring simultaneously (Elston et al. 1985; Lawrence et al. 1986; Sanders & Mirabel 1996). The existence of obscured AGNs in these objects are also considered as an alternative (or additional) energy source where these are fueled by molecular gas falling into the central regions of these merging systems (Surace & Sanders 1999; Scoville et al. 2000; Sanders & Mirabel 1996; Genzel et al. 1998; Lutz et al. 1999; Veilleux et al. 2002, 2006). Molecular gas is thus one of the most important constituents of the ISM and plays a critical role in the evolution of galaxies.

To date, over 140 molecules have been identified in space, mostly in the ISM of our Milky Way Galaxy. Some of these are rather complex, containing as many as 8 to 13 atoms. Interstellar organic molecules are thought to form mostly on the surface of dust grains. Heating events, such as the formation of a protostar, release the icy grain mantles into the gas phase (Nomura & Millar 2004; Caselli et al. 1993). Once released, these molecules may form amino acids by the combination of organic species known as “pre-biotic” molecules (Blagojevic et al. 2003). (We note that non-gas phase reaction pathways for the formation of extraterrestrial amino acids have been described in Elsila et al. 2007). Methanimine is one such molecule (CH$_2$NH; Kirchoff et al. 1973) which can form the simplest amino acid, glycine (NH$_2$CH$_2$COOH), either by (i) first combining with hydrogen cyanide (HCN) to form aminoacetonitrile (NH$_2$CH$_2$CN), with subsequent hydrolysis (Strecker Synthesis; Dickerson 1978), or (ii) directly combining with formic acid (HCOOH) (Godfrey et al. 1973). Methanimine has been previously detected in the interstellar medium of our own Galaxy (Godfrey et al. 1973; Dickens et al. 1997) and tentatively in the nearby galaxy, NGC 253 (Martín et al. 2006), but never beyond the neighborhood of our Galaxy (i.e. beyond ~5 Mpc).

The majority of interstellar molecules (both galactic and extragalactic) have been discovered at millimetric wavelengths, as molecules with small moments of inertia are the most abundant cosmically, with their rotational lines occurring at mm or shorter wavelengths.
However, although less abundant, many complex molecules have spectral lines in the radio regime for $\lambda > 3$ cm, where “line confusion” does not set a limit to their detectability. Many transitions of small polycyclic aromatic hydrocarbons (PAHs), pre-biotic molecules, and even a number of transitions of the simplest amino acid, glycine, fall within the relatively unexplored spectral range between 1 and 10 GHz. In addition, observations in this frequency range are complementary to mm spectral-line surveys which probe colder, lower density gas.

At Arecibo, we are conducting a spectral line census of Arp 220 between 1.1 and 10 GHz, for which the initial observations took place between 31 March and 22 April, 2007. It is planned that the remaining observations will be made in 2008. Here we report the discovery of methanimine emission in this galaxy, plus the detection of three $v_2 = 1$ direct l-type absorption lines of HCN (Thorwirth et al. 2003) from the J=4, 5 and 6 vibrational levels. Also reported is the possible detection of another pre-biotic molecule, formic acid, albeit not unambiguously due to the presence of a nearby $^{18}$OH line. In addition, we present the first detections of two, possibly three, $\lambda4$-cm transitions of the OH radical in absorption, high signal-to-noise ratio detections of the $\lambda6$- and 5-cm transitions of OH, and the possible detection of the 6.7-GHz methanol (CH$_3$OH) transition in absorption.

2. Arp 220

At a distance of $\sim77$ Mpc (redshift, $z=0.018126$), Arp 220 is the nearest Ultra-Luminous Infra-Red Galaxy (ULIRG). Its IR luminosity arises from a powerful, dust-enshrouded starburst, triggered by the merger of two gas-rich galaxies (Sakamoto et al. 1999; Mundell et al. 2001; Sanders & Mirabel 1996). Evidence for this is provided by high resolution optical and radio images revealing a double nucleus with tidal tails and dust lanes. A high supernovae rate has been found from recent high resolution VLBI studies (Lonsdale et al. 2006). Molecules such as the OH radical (Baan et al. 1982; Ghosh et al. 2003), CO (Scoville et al. 1986), formaldehyde (Araya et al. 2004), ammonia (Takano et al. 2005), and mm-transitions of HCN (Aalto et al. 2007) have been detected in Arp 220.

Arp 220 is also known as the prototype OH megamaser galaxy. Its OH $\lambda18$-cm maser emission was first reported by Baan et al. (1982) and was found to have several orders of magnitude higher luminosity than typical Galactic OH masers (hence the name megamaser). High-resolution maps of these OH masers reveal complex structures which could be interpreted either as a dual-component distribution (Diamond et al. 1989; Lonsdale et al. 1998; Rovilos et al. 2003) or, by clumpy unsaturated masers within a single-component medium (Parra et al. 2005; Momjian et al. 2006).
3. Observations

Using several of the complement of receivers on the Arecibo 305-m telescope, we are in the process of making an almost-complete spectral scan of Arp 220 between 1.1 and 10 GHz. To do this, we have employed the WAPP (Wideband Arecibo Pulsar Processor) spectrometer in its recently-commissioned “dual-board” mode. In this mode, eight independent boards, each of 100-MHz bandwidth with 3-level quantization, can be used to cover a spectral band of up to 800 MHz at a single time. Compared to the earlier WAPP capacity, this doubles the total bandwidth covered. The present project serves both for the scientific commissioning of this new option, and as a demonstration of its capabilities.

In practice, we have overlapped the eight boards for each observation such that their centers are separated by 85 MHz. As the final 5-or-so MHz are affected by filter roll-off, this allows high quality data to be acquired for an instantaneous bandwidth of 680 MHz, meaning that those Arecibo receivers which have total bandwidths of 2 GHz can be fully covered via three separate frequency settings. The basic spectral resolution is 24.4 kHz, and both orthogonal polarizations of the celestial signal are being recorded.

The observations are being made via a modified version of the Double Position Switching (DPS) technique (Ghosh & Salter 2002). An ON/OFF position-switched observation with 5-min component phases is made on Arp 220, followed by an ON/OFF with 1-min phases on the strong, angularly-nearby, continuum source, J1531+2402, which is used as a band-pass calibrator. In the presence of band-pass ripples or trapped-modes in the orthomode transducer of the feed, this strategy is necessary at all frequencies to produce acceptable spectral baselines. In addition, apparent features observed in the data can often be recognized as astronomical, and not due to the presence of radio frequency interference (RFI), via a comparison of the spectra for Arp 220 and J1531+2402. Data for a total ON-source observing time of about 60 min for Arp 220 will eventually be acquired for each frequency setting.

Data reduction has been performed using the Arecibo IDL analysis package written by Phil Perillat. The individual ON/OFF scans on Arp 220 were processed to yield (ON-OFF)/OFF spectra, and these were bandpass corrected using similar spectra for J1531+2402. Each individual Arp 220 and J1531+2402 scan was inspected for quality, and all RFI present were noted. A number of scans were rejected either due to technical problems or because of excessive RFI. All acceptable scans for a particular frequency setting were then co-added to produce the final spectra. These spectra were smoothed in frequency to a number of resolutions, and the resultant spectra inspected visually to identify the presence of possible emission or absorption lines. Considerable cross-checking was used to ensure that candidate lines were real, and not the effect of RFI or equipment problems.
4. Results & Discussion

A complete cm-wavelength line census for Arp 220, and its full implications for the physical and chemical properties of the interstellar medium of this galaxy, are deferred until completion of the observations. Here we present the detection of methanimine in emission, and of the \( v_2 = 1 \) direct \( l \)-type transitions of HCN, excited-OH transitions, a line that could be either \(^{18}\)OH or HCOOH, and possibly methanol in absorption for this prototypical nearby ULIRG. The spectra are presented in Figs. 1–10, and the parameters derived from them are given in Tables 1 & 2.

Table 1 is ordered as follows: Columns (1–3) present the name of the molecule, the relevant transition and its rest frequency in MHz. Col. (4) is the peak flux density of the line in mJy. Col. (5) is the rms noise of the spectrum in mJy at the velocity resolution of \( \sim 30 \) km s\(^{-1}\). Col. (6) is the radial velocity in km s\(^{-1}\) at the peak of the measured line, derived from the rest frequency of the strongest of the six lines in the multiplet (i.e. that at 5289.813 MHz). Col. (7) is the full-velocity width in km s\(^{-1}\) of the entire emission feature at half of the peak intensity.

For Table 2: Columns (1–3) are the same as those for Table 1. Col. (4) is the maximum optical depth of the absorption line. Col. (5) is the rms noise of the optical-depth spectrum at the velocity resolution of \( \sim 30 \) km s\(^{-1}\). Col. (6) is the radial velocity in km s\(^{-1}\) at the maximum optical depth. Col. (7) is the full width in km s\(^{-1}\) of the line at half of the peak optical depth. Col. (8) is \( \int \tau dV \), the integrated area in km s\(^{-1}\) under the absorption feature.

In the following sections, we discuss the spectra of each molecule separately.

4.1. Methanimine (\( CH_2NH \))

A broad emission feature covering all six \( 1_{10} - 1_{11} \) transitions of the C-band multiplet of methanimine is presented in Fig. 1. In Fig. 1a, the rest frequencies of the transitions in the frame of Arp 220 were derived using a recessional velocity of 5373 km s\(^{-1}\), as found for the western component of the OH megamaser emission of Arp 220, while the heliocentric velocity axis of Fig. 1b is for the transition expected to be the strongest in the methanimine multiplet, that at a rest frequency of 5289.813 MHz (Godfrey et al. 1973). The total velocity width (FWHM) of this emission line is 270 km s\(^{-1}\); we note that a large velocity width is observed for almost all molecular spectra in Arp 220 (e.g. Takano et al. 2005). Using the angular size of \( 0.27'' \times 0.21'' \) measured for the strongest component of the formaldehyde (\( H_2CO \)) emission (Baan & Haschick 1995) as an upper limit for the angular size of the methanimine emission region, we derive a lower limit to the brightness temperature of \( \sim 2800 \) K. Taking into account
the six lines in the multiplet and their relative intensities, we calculate a reduced value of this lower limit to be $\sim 1000$ K for the strongest component. This is similar to the methanimine decomposition temperature of 1300 K (Nguyen et al. 1996). If the solid angle of the emission is smaller than the above, for instance if the emission comes from a number of very compact components, then the brightness temperature could be much higher. We conclude that, as for formaldehyde (Araya et al. 2004), methanimine in Arp 220 is likely to be showing weak maser emission for this transition. We are planning to observe this source at higher spatial resolution in order to determine the brightness temperature more accurately and to discover where exactly in the galaxy the emission is located.

4.2. Hydrogen Cyanide (HCN)

Fig. 2 shows the energy diagram for the $v_2=1$ direct l-type transitions of hydrogen cyanide (HCN) in the J=1 to J=6 vibrational levels. Our detections of the J=2, 4, 5 and 6 HCN transitions are presented in Fig. 3. The J=4, 5 and 6 transitions are seen in absorption against the continuum emission of Arp 220, and we place an upper limit at the 3-$\sigma$ level of 0.0025 for the optical depth of the J=2 line. The first detailed study of this type of HCN transition was carried out for the Galactic proto-planetary nebula, CRL 618, by Thorwirth et al. (2003). These observers detected a number of the higher vibrational levels in this HCN ladder (J=8–14). However we note that the lower energy transitions presented here seem not to have been previously detected in any celestial source. We also note that these lines represent a high excitation energy above the HCN ground state, (e.g. 1067 K for the J=4 line.)

HCN is a well known indicator of high gas density. Recently, Gao & Solomon (2004) demonstrated that a very strong linear correlation exists between the luminosities $L_{\text{IR}}$ and $L_{\text{HCN}}$ for mm-wave transitions of HCN. This extends over a wide range of $L_{\text{IR}}$ from normal galaxies to ULIRGs. A similar $L_{\text{IR}} - L_{\text{CO}}$ correlation has a less linear form, marking out $L_{\text{HCN}}$ as the best tracer of dense molecular gas mass in galaxies, and hence of active star-forming gas. (However, see Papadopoulos et al. 2007, for a different point of view.) The relative line integrals from the J=4 and 6 lines suggest an approximate excitation temperature of $\sim 150$ K. Given the high excitation that these lines represent, it would seem plausible that they are coming from regions of star formation.

Unlike the relatively “smooth” line profiles seen for the excited-OH lines (e.g. Fig. 5), the HCN lines each show evidence for the presence of a number of discrete components. In fact, the central, strongest HCN component becomes increasingly dominant from J=4 to 6. The absence of the J=2 line requires explanation, as the predicted absorption line is
expected to be an order of magnitude stronger than the 3-σ limit we place above. It is highly improbable that the fraction of the continuum emission “covered” by the clouds producing the HCN absorption have decreased by such a large amount between 4488 and 1347 MHz as to render the J=2 line unobservable. Much more likely is that free-free absorption in the foreground ionized screen of this starburst galaxy is greatly attenuating the J=2 line. For this to be the case, an opacity of \( \gtrsim 2.25 \) at 1347 MHz (in the rest frame of Arp 220) would be required. This would imply an optical depth of \( \gtrsim 1.5 \) at 1630 MHz in the galaxy, as found for the spectra of a number of the SNRs near the twin nuclei of Arp 220 by Parra et al. (2007). These authors attributed this high opacity to the combined presence of “a patchy FFA [free-free absorption] ISM with a median opacity of less than 1”, plus absorption in the regions of ionized circumstellar mass-loss envelopes surrounding the SN progenitors. The HCN lines we see are expected to arise from star-forming regions of high gas density, and a detailed study of all possible lines in the \( v_2 = 1 \) direct l-type transitions of the HCN ladder would provide useful evidence concerning the properties of the foreground gas screen to these regions. Clearly, the spectrum of the HCN J=3 line (with a rest frequency of 2693.3 MHz) will be crucial to such a study, and we will be observing this line during our 2008 campaign to complete the present observations. It should be noted that for the implied optical depths at 1347 MHz, even the J=4 line will be reduced through FFA by \( \gtrsim 20\% \), which would reduce the above derived excitation temperature to \( \sim 120 \) K.

### 4.3. Excited Hydroxyl (OH)

The energy levels for the various OH transitions are shown in Fig. 4. In Figs. 5–8 we present the \( \lambda 6\)-, 5- and 4-cm \( \Lambda \)-doublet transitions of the OH radical seen in absorption against the continuum emission of Arp 220. The relevant quantum numbers, rest frequencies and other measured line parameters are presented in Table 2. The \( \lambda 6\)- and 5-cm lines have been previously detected by Henkel et al. (1987) and Henkel et al. (1986) respectively.

Contrary to the findings of Henkel et al. (1987), we find that the 4660, 4751 and 4766 MHz lines have intensity ratios closely in agreement with their expected local thermodynamic equilibrium (LTE) values of 1:2:1.

The \( \lambda 5\)-cm main lines presented here have considerably higher signal-to-noise ratio than the measurements of Henkel et al. (1986) but show similar form. The expected LTE intensity ratios for the 6017, 6031, 6035 and 6049 MHz lines are 1:14:20:1. However, we measure ratios of 1:13:13:<0.4. The two main lines of the \( \lambda 5\)-cm transition (6031 and 6035 MHz) are blended due to their large velocity widths and give the appearance of being “saturated”, as suggested by their similar optical depths. This may be due to their having high optical
depths resulting in saturation of the lines against a continuum component containing of order 10% of the total flux density of Arp 220. However, we note that there is a low-velocity component to (presumably) the 6031-MHz absorption line, seen in Fig. 6 at 6026 MHz, which if also present for the 6035-MHz line would contribute significantly at the frequency of the 6031-MHz line. The satellite lines are seen at much lower signal-to-noise level. However, the greater strength of the 6017-MHz line relative to that at 6049 MHz is interesting. A similar effect was also found by Gardner & Martín-Pintado (1983) for four compact HII regions in our own Galaxy. While they saw the 6017-MHz lines enhanced above the LTE ratio to the main lines, the 6049-MHz line was weaker than predicted, and may even sometimes have appeared in weak emission.

The two $\lambda4$-cm OH lines at 7761 and 7820 MHz (Fig. 7) are detected for the first time in Arp 220, and show similar velocity widths to most other molecular species in the galaxy. The expected ratio for these $2\Pi_{1/2}, J = 3/2$ main lines (7761:7820 MHz) is 1:1.8 in thermal equilibrium, very close to the derived ratio of the peak and integrated brightnesses for the lines of 1:1.89 (see Table 2). The satellite lines in this multiplet are expected to be present only at the 1-$\sigma$ level, and indeed are not detected. Considering the absorption lines in the $2\Pi_{1/2}$ ladder of Fig. 4 at 4750 and 7820 MHz, in thermal equilibrium the ratio of their integrated optical depths would imply an excitation temperature of about 88 K. In a detailed Large Velocity Gradient (LVG) study of their CO observations, Greve et al. (2006) find that in Arp 220, the spectra of low density tracers such as CO (and OH) can only be fitted with a two-phase molecular ISM with the kinetic temperature of one being $>30$ K, while their mm-wave spectra for the high-density tracer molecules, HCN and CS, indicate kinetic temperatures of $\sim50-70$ K.

A very tentative detection of the $2\Pi_{1/2}, J = 5/2, F = 2–2$ line of OH at 8135 MHz is shown in Fig. 8. If confirmed via the remaining data that will be acquired in this project, this would be at least twice as strong as expected for an excitation temperature of 88 K. Further, this line is expected to be only 0.7 times as strong as for the other $2\Pi_{1/2}, J = 5/2$ main line at 8189 MHz. While the 8189 MHz line is also possibly detected, contrary to expectations this would be at an even lower signal-to-noise level than the 8135 MHz transition.

4.4. Formic Acid (HCOOH) or “Heavy” Hydroxyl ($^{18}$OH)

In Fig. 9, we present a high signal-to-noise detection of an absorption line from our L-band spectrum. Despite the presence of nearby RFI caused by Glonass emissions (at $\sim1605$ MHz), the reality of this absorption line has been verified by its presence with similar optical depth in each of 13 individual spectra of Arp 220, but in none of the spectra of the
bandpass calibrator. As demonstrated by the horizontal lines in Fig. 9, there is an ambiguity as to the species responsible for this absorption, which could be either the 1639.5-MHz main line of $^{18}$OH or the 1638.8-MHz line of formic acid (HCOOH). Given the prevalence of OH in Arp 220, it would perhaps not be unreasonable to detect the presence of $^{18}$OH as well. However, since formic acid is relevant to the chemical origin of life, it would be most interesting to resolve this ambiguity.

As is seen in Fig. 9, as well as the strong absorption line detected near 1611 MHz, an apparently weaker absorption is also seen near 1609 MHz, close to where the second, 1637.6-MHz, main line of $^{18}$OH should be found. If this feature were indeed to be real, the ratio of the peak intensity of the higher frequency absorption to this would be 3.1:1. This is higher than the expected ratio of 1.8:1 were the pair to be the main lines of $^{18}$OH (Barrett & Rogers 1964) and in local thermal equilibrium. The frequencies of the peak depths of the two features are separated by an amount that would correspond to a rest frequency separation of about 2.05 MHz were they both to be due to absorption in Arp 220. The laboratory separation of the $^{18}$OH main lines is 1.939 ± 0.003 MHz (Lovas 1986), and hence given the weakness of the feature near 1609 MHz, the agreement is considered to be satisfactory.

If the absorption near 1611 MHz is indeed the higher frequency component of the $^{18}$OH main lines, then its radial velocity is closer to that of the peak velocity found for other molecules in Arp 220 than would be the case for a formic acid detection (see Table 2 and the horizontal bars displayed in Fig. 9.) However, we note that even for $^{18}$OH, this velocity of $\sim 5265$ km s$^{-1}$ is significantly lower than that of the normal molecular peak velocity in Arp 220. In addition, its width of about 190 km s$^{-1}$ is narrower than for any other species that we detect. Were this absorbing gas to represent $^{18}$OH, we note that a velocity of 5265 km s$^{-1}$ is “allowed” in Arp 220 from the HI absorption study of Mundell et al. (2001). However, a peak velocity of 5135 km s$^{-1}$, appropriate for the formic acid transition, has little associated HI absorption. Nevertheless, if the absorption were to be due to $^{18}$OH at 5265 km s$^{-1}$ this would imply a remarkably high isotopic ratio of $^{18}$OH:$^{16}$OH for a cloud at this velocity, especially so as this is well away from the peak velocity range of the OH megamaser line as seen at low angular resolution (Baan et al. 1982).

Given the proximity of the Glonass RFI, it is difficult to establish the origins of this absorption feature. However, $^{18}$OH would seem a more likely identification than formic acid. Confirmation of the reality of the second, weaker, absorption component near 1609 MHz would effectively establish this identification. The possibility that the feature represents absorption in our galaxy from the 1612.231 MHz satellite OH line has to be very small given the high galactic latitude of the line-of-sight ($b = 53^\circ.0$), the implied large radial velocity ($\sim +185$ km s$^{-1}$), and large line width ($\sim 190$ km s$^{-1}$).
4.5. Methanol ($\text{CH}_3\text{OH}$)

In Fig. 10, we present a possible detection of the $5_1 - 6_0$ $A^+$ methanol line in absorption. Although this line is apparently detected with a signal-to-noise ratio of almost 6:1, and the entire 100-MHz band in which it is observed is basically RFI-free, the quality of the baseline in this case is rather poor. We await our remaining observations for confirming the reality of this line. However, we can conclude that at the 1 mJy level ($5\sigma$) no maser emission (such as is commonly seen for this line from regions of massive-star formation in our own Galaxy) is seen.

Assuming the reality of the methanol absorption line, an excitation temperature for the transition of $T_{ex} = 20$ K, and an Einstein coefficient of $A = 0.1532 \times 10^{-8}$ s$^{-1}$ (Cragg et al. 1993), in LTE the column density of the lower energy level, $N_{6,0}$ is;

$$N_{6,0} = 1.34 \times 10^{15} \int \tau \delta v \text{ cm}^{-2}$$  \hspace{1cm} (1)

where the integral $\int \tau \delta v$ is expressed in units of km s$^{-1}$.

An approximation for the total column density of methanol can be obtained assuming a Boltzmann distribution for the methanol energy-level population. If the relative abundances of the A and E species of this molecule is taken to be 1:1 based on their having a relatively small ground state energy difference, then the total column density of methanol molecules is;

$$N = N_{6,0} \frac{2}{13} Q(T_{ex}) \frac{E_{6,0}}{k T_{ex}}$$  \hspace{1cm} (2)

where $Q(T_{ex})$ is the partition function, taken to be 39.8 (Cragg, D., quoted in Houghton & Whiteoak 1995), and $E_{6,0}$ is the energy above the ground state of the $6_0A^+$ level. Thus, the presence of a 6668-MHz methanol absorption line in Arp 220 would imply a total column density for this molecule of $N_{\text{CH}_3\text{OH}} \sim 2.5 \times 10^{17}$ cm$^{-2}$.

5. Concluding Remarks

The Arecibo Arp 220 Spectral Census is an on-going project and a detailed analysis will be presented in a separate paper, as will the final spectra from the completed project. We have presented here the discovery of methanimine in this galaxy and the detection of three previously unseen cm-wavelength transitions of HCN. We have also observed, for the first
time in Arp 220, an absorption line that may be either formic acid or $^{18}$OH, two (possibly three) λ4-cm transitions of excited OH, and what may be the first extragalactic detection of methanol.

The origin of life is still an open question. The discovery of high abundances of “prebiotic” molecules, such as methanimine, HCN and possibly formic acid in Arp 220 adds a new dimension to the consideration of this problem. It also raises the possibility that other ULIRGs might contain similarly high abundances of such molecules, which could be detectable with the Arecibo 305-m telescope.

We thank Paul Goldsmith (JPL), Sven Thorwirth and Karl Menten (MPIfR) for fruitful discussions. The Arecibo Observatory is a part of the National Astronomy and Ionosphere Center (NAIC) operated by Cornell University under a cooperative agreement with the National Science Foundation (NSF).

Facilities: Arecibo (L-wide, S-high, C, C-high, X)

REFERENCES

Dickerson, R.E. 1978, Sci. Am., 239, 3, 70
Kennicutt, R.C. 1998, ARAA, 36, 189


Fig. 1.— The blended emission line from the six $1_{10} - 1_{11}$ multiplet transitions of methanimine. The upper panel shows the rest frequencies of the transitions in the galaxy for an assumed heliocentric velocity of 5373 km s$^{-1}$ (corresponding to the western nucleus of Arp 220) as indicated by vertical lines. The lower panel shows the emission line as a function of the heliocentric velocity expected for the strongest line in the multiplet. The velocity resolution is $\sim$30 km s$^{-1}$. 
Fig. 2.— Term value diagram for HCN in its $v_2=1$ vibrational state from $J = 1$ to $J = 6$. Only the direct $l$-type transitions with $\Delta J = 0$ are shown. The diagram on the right-hand side shows the $J = 4$ direct $l$-type transition at 4488.4718 MHz in detail. This diagram was drawn using the energy level values obtained from the CDMS database (Müller et al. 2005, 2001)
Fig. 3.— The first astronomical detection of the $v_2 = 1$ direct l-type absorption lines of HCN with vibrational levels $J=4$, 5 and 6. The non-detection of the $J=2$ vibrational level is also included in the figure. The velocity resolution is $\sim 30$ km s$^{-1}$. 
Fig. 4.— The energy diagram for various Λ-doublet transitions of the OH radical.
Fig. 5.— The three $\lambda$6-cm excited OH transitions from $2\Pi_{1/2}$, $J=1/2$, $F=0-1$, 1-1 and 1-0. The $F=1-1$ line is about twice as strong as the other two lines. The velocity resolution is $\sim 30$ km s$^{-1}$. 
Fig. 6.— The four λ5-cm excited OH transitions as a function of rest frequency for an assumed heliocentric velocity of 5325 km s\(^{-1}\) (as appropriate for the λ6-cm OH lines). The frequencies of the four transitions \(2\Pi_{3/2}, J=5/2, F=2-3, 2-2, 3-3, 3-2\) are indicated by vertical lines. The two main lines form a blended absorption line profile. The velocity resolution is \(\sim 30\) km s\(^{-1}\).
Fig. 7.— First detection of the two λ4-cm excited OH main-line transitions, $2\Pi_{1/2}$, $J=3/2$, $F=1-1$ and $2-2$. The velocity resolution is $\sim 30$ km s$^{-1}$. 
Fig. 8.— The possible detection of the λ4-cm excited OH transition $2\Pi_{1/2}$, $J=5/2$, $F=2-2$. The velocity resolution is $\sim30$ km s$^{-1}$. 
Fig. 9.— This plot shows one (or possibly two) absorption line(s) that could be either formic acid or $^{18}$OH. The expected location of the formic acid line and of the two main $^{18}$OH lines are indicated with horizontal bars. The band is affected by interference from the Glonass system at about 1605 MHz, but the authenticity of the deep absorption feature has been verified by its non-appearance in the spectra of the bandpass calibrator. The velocity resolution is $\sim 30$ km s$^{-1}$. 
Fig. 10.— A possible detection of the $5_1-6_0$ $A^+$ methanol line in absorption. However, more data is needed to confirm the detection. The velocity resolution is $\sim 30$ km s$^{-1}$. 
Table 1. Derived parameters for the emission multiplet of methanimine (CH$_2$NH).

<table>
<thead>
<tr>
<th>Molecule</th>
<th>Transition</th>
<th>Rest frequency (MHz)</th>
<th>$S_p$ (mJy)</th>
<th>$\sigma$ (mJy)</th>
<th>$v_p$ (km s$^{-1}$)</th>
<th>FWHM (km s$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methanimine (CH$_2$NH)</td>
<td>$1_{10}-1_{11}$, $\Delta F = 0, \pm 1$</td>
<td>5289.813$^1$</td>
<td>3.5</td>
<td>0.22</td>
<td>5362</td>
<td>270</td>
</tr>
</tbody>
</table>

The subscript $p$ stands for “peak”.

$^1$ The strongest of the six lines of this multiplet has this frequency in laboratory measurements.

Table 2. Derived parameters from the molecular absorption lines.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>Transition</th>
<th>Rest frequency (MHz)</th>
<th>$\tau_p$</th>
<th>$\sigma$</th>
<th>$v_p$ (km s$^{-1}$)</th>
<th>FWHM (km s$^{-1}$)</th>
<th>$\int \tau dV$ (km s$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrogen cyanide (HCN)</td>
<td>$v_2 = 1, \Delta J = 0, J = 2$</td>
<td>1346.7650</td>
<td>&lt; 0.0025</td>
<td>0.00085</td>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>$J = 4$</td>
<td>4488.4718</td>
<td>0.016</td>
<td>0.0009</td>
<td>5407</td>
<td>363</td>
<td>5.22</td>
</tr>
<tr>
<td></td>
<td>$J = 5$</td>
<td>6731.9098</td>
<td>0.0205</td>
<td>0.001</td>
<td>5404</td>
<td>330</td>
<td>5.02</td>
</tr>
<tr>
<td></td>
<td>$J = 6$</td>
<td>9423.3338</td>
<td>0.0286</td>
<td>0.002</td>
<td>5398</td>
<td>202</td>
<td>6.14</td>
</tr>
<tr>
<td>Hydroxyl radical (OH)</td>
<td>$2\Pi_{1/2}, J = 1/2, F = 0-1$</td>
<td>4660.242</td>
<td>0.0254</td>
<td>0.00082</td>
<td>5425</td>
<td>300</td>
<td>7.99</td>
</tr>
<tr>
<td></td>
<td>$F = 1-1$</td>
<td>4750.656</td>
<td>0.0549</td>
<td>0.00080</td>
<td>5429</td>
<td>287</td>
<td>15.91</td>
</tr>
<tr>
<td></td>
<td>$F = 1-0$</td>
<td>4765.562</td>
<td>0.0295</td>
<td>0.00080</td>
<td>5447</td>
<td>281</td>
<td>8.41</td>
</tr>
<tr>
<td></td>
<td>$2\Pi_{3/2}, J = 3/2, F = 2-3$</td>
<td>6016.746$^2$</td>
<td>0.0086</td>
<td>0.0016</td>
<td>5313</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>$F = 2-2$</td>
<td>6030.747$^3$</td>
<td>0.11</td>
<td>0.0016</td>
<td>5425</td>
<td>480</td>
<td>56.20</td>
</tr>
<tr>
<td></td>
<td>$F = 3-3$</td>
<td>6035.092$^3$</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>Hydroxyl radical (OH)</td>
<td>$2\Pi_{1/2}, J = 3/2, F = 1-1$</td>
<td>7761.747</td>
<td>0.0087</td>
<td>0.0018</td>
<td>5359</td>
<td>296</td>
<td>2.62</td>
</tr>
<tr>
<td></td>
<td>$F = 2-2$</td>
<td>7820.125</td>
<td>0.0162</td>
<td>0.00153</td>
<td>5432</td>
<td>321</td>
<td>5.04</td>
</tr>
<tr>
<td>Hydroxyl radical (OH)</td>
<td>$2\Pi_{1/2}, J = 5/2, F = 2-2$</td>
<td>8135.870$^2$</td>
<td>&lt; 0.01</td>
<td>0.0021</td>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
</tbody>
</table>
| Hydroxyl radical (OH) | $2\Pi_{1/2}, J = 3/2, F = 2-2$ | 1639.503 | 0.019 | 0.00137 | 5265 | $\sim$192 | $-$
| 18OH or Formic acid (HCOOH) | 1(1,0)–1(1,1) | 1638.805 | ... | ... | ... | ... | ... |
| Methanol (CH$_3$OH) | $5_1-6_0 A^+$ | 6668.5192 | 0.0077 | 0.0013 | 5384 | 360 | 2.53 |

The subscript $p$ stands for “peak”.

$^1$ Since the two main lines $F = 2-2$ and 3-3 are blended, values presented here are for the two lines combined together.

$^2$ For these low signal-to-noise detections some of the parameters were not estimated.

$^3$ Since the identification of this absorption feature is ambiguous, an optical-depth integral was not estimated for it.