On the formation of carbonic acid (H$_2$CO$_3$) in solar system ices

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Received 26 April 2007; in final form 29 October 2007
Available online 1 November 2007

Abstract

We investigated the irradiation of H$_2$O–CO$_2$ ice mixtures with energetic electrons in an ultrahigh vacuum chamber. Our laboratory studies confirm that – besides carbon monoxide–carbonic acid (H$_2$CO$_3$) is the dominant reaction product of these irradiation processes. We present the kinetics and dynamics of its formation and elucidate on the temperature-dependence (10–60 K) of the formation processes. Our experimental results indicate that carbonic acid might be present in the solar system ices that contain both water (H$_2$O) and carbon dioxide (CO$_2$) and are exposed by radiation from the solar wind and planetary magnetospheres.

1. Introduction

Unraveling the synthetic routes to form carbonic acid (H$_2$CO$_3$) is of great interest to astrochemists and astrobiologists [1,2]. This acid is a potential reactant to form biologically important molecules like oxalic acid and also has geochemical implications such as the precipitation of carbonates in aqueous solutions [3]. Water and carbon dioxide are the most likely sources of interstellar H$_2$CO$_3$ based on their high cometary [4] and interstellar abundances [5]. Also, both precursors have been found on the surfaces of Europa, Ganymede [6], and Mars [7]. These surfaces are exposed to high energy particles (solar wind, planetary magnetospheres) which can induce a chemical alteration of the pristine environment via non-equilibrium chemistry [8]. Therefore, ices containing water and carbon dioxide are expected to be a residence for carbonic acid. Satellites of Jupiter (Europa, Ganymede and Callisto) are showered with magnetospheric ions [9]. These species can interact with ices and create cascades of secondary electrons in the track of the implanted ions. Due to the absence of a magnetic field, Mars is not bombarded by magnetospheric ions, but its relative proximity to the sun and lack of dense atmosphere exposes its surface to the solar wind.

Moore and Khanna [10] were the first to study binary mixtures of carbon dioxide and water of a ratio of unity. They discovered carbonic acid after irradiating these ices using 0.7 MeV protons. Dellorusso et al. [11] confirmed this finding; these authors also suggested the formation of formaldehyde (H$_2$CO). Subsequent studies of Brucato et al. [12], Gerakines et al. [13], and Wu et al. [14] demonstrated that formaldehyde is not synthesized in such ices. However, previous studies were conducted at ice temperatures between 10 and 20 K; these temperatures do not mimic the majority of solar system ices that harbor these ice mixtures. Would higher temperature systems, such as Triton (38 K) or Europa (>70 K) or Mars (>133 K), still produce carbonic acid? While many studies have been conducted irradiating ices with protons or ultraviolet (UV) photons, little information exists on the effect of high energy electron exposures. Recall that keV electrons are formed in the track of the galactic cosmic ray and solar wind proton implants once they penetrate the ice layers [6]. Therefore, we present studies of water–carbon dioxide ices and irradiate these binary mixtures at 10, 40, and 60 K with energetic (5 keV) electrons. Most important, only a brief speculation of possible mechanisms to form carbonic acid could be found [13]; this falls short because...
the postulated reaction intermediates like radicals and/or ions could not be detected in the experiments. Because neither reaction mechanisms nor rate constants have been derived for the formation of carbonic acid, we found it necessary to conduct an experimental study how carbonic acid is formed in water and carbon dioxide rich ices and how the formation rates depend on the temperature.

2. Experimental

The water–carbon dioxide mixtures were condensed on a silver mirror inside a ultrahigh vacuum (UHV) chamber [15]. A rotatable cold head protrudes from the top of the chamber to steady the silver substrate in the center of the vessel. The temperature of the silver mirror is adjustable from 10 to 340 K by a helium refrigeration system. Connected to the main chamber is an oil-free turbomolecular pump backed by a scroll pump, which allows for pressures to be as low as 7.0 × 10^{-11} Torr. Attached to the chamber is a differentially pumped electron source, which consists of a tungsten filament and electrostatic lenses to accelerate the electrons to 5 keV. The gas mixture was prepared in a separate side chamber connected by a linear transfer mechanism to the main recipient. Twenty mbar of distilled and repeatedly defrosted water was added to 33 mbar of carbon dioxide (BOC Gases, 99.999%) before condensed for 7 min at a pressure of 7.0 × 10^{-3} Torr onto the 10 K silver wafer. Fig. 1a shows an infrared spectrum of the pristine ice mixture taken at 10 K; the infrared absorptions are compiled in Table 1. Utilizing a modified Lambert–Beer law [15], the column densities were calculated to be (1.9 ± 0.1) × 10^{17} molecules cm^{-2} for H2O and (4.6 ± 0.1) × 10^{17} molecules cm^{-2} for CO2. This translates to a carbon dioxide–water ratio of about (2.5 ± 0.5):1. Absorption coefficients for H2O at 3663 cm^{-1} (2.0 × 10^{-16} cm molecule^{-1}) and CO2 at 2341 cm^{-1} (1.0 × 10^{-16} cm molecule^{-1}) were taken from Ref. [16,17], respectively. Based on the densities – CO2: 1.7 g cm^{-3} [18], H2O: 0.93 g cm^{-3} [19] – the equivalent thicknesses of the H2O and CO2 were found to be 62 and 190 nm, respectively.

All ices were prepared at 10 K, and were warmed up to a specific temperature (10, 40 or 60 K) prior to an irradiation with 5 keV electrons for 3 h at an electron current of 100 nA. Note the electron beam covers an area of 4030 3947 H2O 4000 3500 3000 2500 2000 1500 1000 500 Wavenumbers (cm^{-1}) H2O.CO2 1384 1384 H2O 1614 1574 H2O 2280 2281 CO2 1638 1626 H2O.CO2 184 13 CO2 750 760 H2O 657654 660654 CO2 660654 663649 H2O.CO2 4030 3947 H2O 3795 3818 H2O.CO2 3702 3707 CO2 3661 3653 H2O 3595 3602 CO2 3385 3332 H2O 3240 3151 H2O 2393 2404 H2O.CO2 2330 2341 CO2 2280 2281 CO2 1638 1626 H2O.CO2 1614 1574 H2O 1384 1384 CO2 750 760 H2O 657654 660654 CO2 660654 663649 H2O.CO2

Fig. 1. (a) Infrared spectrum of the unirradiated H2O/CO2 ice at 10 K. (b) H2O/CO2 infrared spectrum at 10 K after 3 h of irradiation with 100 nA electrons. H2CO3 and CO were the major products. A very small CO3 peak was found at 2044 cm^{-1} but not labeled in the figure. (c) Infrared spectrum of the irradiated sample warmed up to 210 K. This spectrum was collected after 3 h of irradiation at 10 K and warmed to 210 K.

<table>
<thead>
<tr>
<th>Observed wavenumber (cm^{-1})</th>
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<th>Vibration</th>
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<td>3947</td>
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<td>v3</td>
</tr>
<tr>
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<td>H2O·CO2</td>
<td>v1</td>
</tr>
<tr>
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<td>CO2</td>
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<td>H2O</td>
<td>v3</td>
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<tr>
<td>3595</td>
<td>3602</td>
<td>CO2</td>
<td>2v2 + v3</td>
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<td>H2O</td>
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<td>660654</td>
<td>663649</td>
<td>H2O·CO2</td>
<td>v2 *</td>
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</table>

H2O is the literature values and assignments from Ref. [16,22].
CO2 is the literature values and assignments from Ref. [17].
H2O·CO2 is the literature values and assignments taken from Ref. [23].
\* In-plane bend, out-of-plane bend.
3. Results

The first step in the analysis of the data is to examine the initial, pre-irradiated infrared spectrum of the carbon dioxide–water mixture. Based on previous experiments of pure H2O [16], pure CO2 ices [15,17] and H2O/CO2 ice mixtures [22], eleven peaks can be identified as fundamental or combination bands, five belonging to the water and six absorptions to the carbon dioxide (Table 1; Fig. 1a). Ab initio calculations carried out by Danten et al. [23] indicate the remaining peaks belong to a H2O–CO2 complex. These absorptions match up well with the peaks found in our experiments, except for the O–H stretch (ν3), which was off by about 85 cm⁻¹. The peaks at 660 and 654 cm⁻¹ are hidden under the carbon dioxide absorptions, but become visible at 110 K in the warm-up phase after the carbon dioxide has sublimed.

After the sample was irradiated for 180 min, the spectrum clearly shows the presence of carbonic acid (Fig. 1b); six fundamentals were detected and correlate well with the literature data. A list of the wavenumbers and the assignments can be found in Table 2. Two peaks at 690 and 2850 cm⁻¹ are not seen until after the warm-up of the irradiated sample. At 210 K, very little of the water, carbon dioxide, and the water–carbon dioxide complex remained; the carbonic acid spectrum becomes much clearer (Fig. 1c). Among all absorptions identified, only one peak at 1483 cm⁻¹ did not overlap with the absorptions from water and carbon dioxide; therefore, this absorption was used to calculate column densities of the newly formed carbonic acid as discussed below. Of the nine frequencies observed by Gerakines et al. [13], only one could not be confirmed in our experiments. Even after sufficient heating, the COH out-of-plane bending mode at 908 cm⁻¹ could not be witnessed because of lingering water on the silver mirror. Additionally, there is one extra peak to note in the infrared region of irradiated sample (Fig. 1b). The trans-HOCO intermediate could be assigned to a peak at 1810 cm⁻¹ (Fig. 2). Milligan and Jacox [24] identified cis and trans-HOCO radicals at 1797 and 1833 cm⁻¹ respectively. We suggest that the 1810 cm⁻¹ peak is from trans-HOCO, at a slightly lower wavenumber, due to the red-shifting effect of our H2O matrix (unlike Milligan’s 99% CO matrix).

Besides the absorptions from carbonic acid and trans-HOCO, the electron irradiation of the water–carbon dioxide ices only produced two additional products. These are carbon monoxide (CO) and the cyclic carbon trioxide isomer (CO3). Both molecules were identified via their absorptions at 2140 and 2044 cm⁻¹. These positions agree well with previous band assignments [15,17]. No hydrogen peroxide (HOOH) was seen in the FTIR spectra, but could be assigned to the proper mass to charge ratio, m/z, of 34 as witnessed by the quadrupole mass spectrometer (QMS). In the blank experiment, no ion current was received for m/z = 34, but in all three irradiations, a small amount was observed. The peak for this mass-to-charge formed just after the sublimation of water at 190 K, and was approximately three orders of magnitude lower than H2O. Formaldehyde (H2CO) and formic acid (HCOOH) were not detected by the FTIR. As noted by Wu et al. [14] in their H2O/CO2 experiments, the carbonyl stretch from H2CO (1720 cm⁻¹) could be hidden underneath the broad peak of carbonic acid, we are not able to confirm the formation of H2CO. In our QMS data, HCOOH (m/z = 46) was not found in any of the four experiments.

Fig. 3 presents the decay of H2CO3 absorption features during the post-irradiation warming up in the temperature range 210–256 K. The decay of these absorption features is correlated to the detection of H2CO3 (m/z = 62) mass signal in the temperature range. These results indicate that

<table>
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<th>Literature wavenumber (cm⁻¹)</th>
<th>Vibration</th>
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<td>2626</td>
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<td>1719</td>
<td>C=O stretch</td>
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<tr>
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<td>COH a-sym stretch</td>
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<td>COH in plane bend</td>
</tr>
<tr>
<td>1038</td>
<td>1038</td>
<td>COH sym stretch</td>
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<tr>
<td>812</td>
<td>813</td>
<td>CO2 out of plane bend</td>
</tr>
<tr>
<td>685</td>
<td>690</td>
<td>CO2 in plane bend</td>
</tr>
</tbody>
</table>

* Ref. [11,13].

![Fig. 2. Deconvoluted peaks of the irradiated infrared spectra (1850-1550 cm⁻¹) at 10 K.](image-url)
H$_2$CO$_3$ was generated by irradiation and were released into the gas phase at temperature between 210 and 256 K.

4. Discussion

To investigate the evolution of carbonic acid during the irradiation in time, one must understand the kinetics of the reaction. Fig. 4 shows the column density of H$_2$CO$_3$ as the ice sample is irradiated. The absorption coefficient (6.5 × 10$^{-17}$ cm molecule$^{-1}$) at 1505 cm$^{-1}$ was taken from Geraikines et al. [13]. The best fit includes a two-step mechanism, i.e. A → B → C, where A is the H$_2$O · CO$_2$ complex, B is an intermediate, and C presents the carbonic acid. To calculate the best fits as shown in Fig. 4, (1)–(3) were used. This leads to the rate constants (Table 3) for the first step from A to B, $k_1$, and the second step from B to C, $k_2$. Based on our fits, the changes in rate constants with temperature are not statistically significant. Therefore, at irradiations from 10 to 60 K, neither step is significantly temperature dependent suggesting that non-equilibrium chemistry dictates the chemical processing of the sample. However, the number of H$_2$O · CO$_2$ complexes formed capable of reacting, $[A]_0$, were found to increase with temperature. This likely indicates that as temperature increases, the number of H$_2$O · CO$_2$ complexes with the correct orientation geometry for the reaction to form carbonic acid present in the pristine sample increases.

\[
[A]_t = [A]_0 e^{-k_1 t} 
\]

\[
[B]_t = [A]_0 \left( \frac{k_1}{k_2 - k_1} \right) (e^{-k_1 t} - e^{-k_2 t}) 
\]

\[
[C]_t = [A]_0 \left[ 1 - \left( \frac{k_2}{k_2 - k_1} e^{-k_1 t} \right) \left( \frac{k_1}{k_2 - k_1} e^{-k_2 t} \right) \right] 
\]

Now that the kinetics has been solved, let us consider the likely reaction mechanism(s). We suggest that the reaction

![Fig. 3. Decay of the infrared features of the new species upon warming. The H$_2$O/CO$_2$ ice was irradiated at 10 K with electrons for 3 h, then, was warmed up to these temperatures. The spectra are offset for clarity.](image)

![Fig. 4. Temporal evolution of carbonic acid with time at 10 K (top), 40 K (middle), and 60 K (bottom). Experimental data points have been fit via (1)–(3).](image)

<table>
<thead>
<tr>
<th>$T$ (K)</th>
<th>$k_1$ (s$^{-1}$)</th>
<th>$k_2$ (s$^{-1}$)</th>
<th>$[A]_0$ (molecules cm$^{-2}$)</th>
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<td>10</td>
<td>(0.8 ± 0.4) × 10$^{-4}$</td>
<td>(1.0 ± 0.4) × 10$^{-4}$</td>
<td>(2.1 ± 0.3) × 10$^{15}$</td>
</tr>
<tr>
<td>40</td>
<td>(1.5 ± 0.5) × 10$^{-4}$</td>
<td>(0.6 ± 0.4) × 10$^{-3}$</td>
<td>(2.5 ± 0.2) × 10$^{15}$</td>
</tr>
<tr>
<td>60</td>
<td>(1.1 ± 0.5) × 10$^{-4}$</td>
<td>(1.8 ± 0.6) × 10$^{-3}$</td>
<td>(2.9 ± 0.2) × 10$^{15}$</td>
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commences with an electron-induced O–H bond rupture within the water molecule. Once the O–H bond is broken by the electron irradiation, a hydrogen atom (H) and a hydroxyl radical (OH) are formed. Each open shell species has a potential to react with a neighboring CO2 molecule. Based on these possibilities, we examined two reactions: H+CO2 (4) and OH+CO2 (5). Both reactions are exothermic by the same order of magnitude [25,26]. Conservation of angular momentum and different masses of the hydrogen atom (1 amu) and of the hydroxyl radical (17 amu), the kinetic energy of the hydrogen atom will be significantly greater than that of the hydroxyl radical. The detection of the HOCO radical, but the failed observation of the HO- COO species suggests that the energetic hydrogen atom can add to the carbon dioxide molecule via Eq. (4) even at 10 K:

\[
\begin{align*}
\text{H} + \text{CO}_2 & \rightarrow \text{HOCO} \quad \Delta_r G = -11.7 \text{ kJ mol}^{-1} \\
\text{OH} + \text{CO}_2 & \rightarrow \text{HOCOO} \quad \Delta_r G = -26.4 \text{ kJ mol}^{-1}
\end{align*}
\]

Fig. 5 displays our proposed reaction mechanism. The H2O·CO2 complex undergoes an electron-induced O–H bond rupture, resulting in a hydrogen atom adding to the carbon–oxygen double bond of the CO2 molecule. Note that at 10 K, this reaction cannot proceed without irradiation since excess energy is needed to overcome bond ruptures and the total endoergicity of the reaction of 31.4 kJ mol\(^{-1}\) to form carbonic acid. Therefore, in the interstellar medium, thermal energy alone will not be sufficient to overcome the large barrier (629.3 kJ mol\(^{-1}\)). This potential energy surface requires non-equilibrium chemistry (such as that generated by ionizing radiation). This suprathermal hydrogen now has excess kinetic energy required to overcome the barrier to addition to the carbon dioxide to form cis-HOCO (endoergic by 91.3 kJ mol\(^{-1}\)), which has been proposed before by a number of researchers [25,27–29]. It can isomerize (37.6 kJ mol\(^{-1}\) barrier) to trans-HOCO, and then recombines with the hydroxyl radical to form carbonic acid (exoergic by 471.9 kJ mol\(^{-1}\)).

H2O and CO2 ices are among the most prevalent ices found in the interstellar medium [5]. They are also known to co-exist on the surface of Mars [7], Triton [30], Jovian satellites [6] and on comets [4]. These ices are exposed to irradiation from cosmic ray particles, solar wind and/or planetary magnetospheres. Based on our experimental data, carbonic acid would exist in these objects. Mars in particular is of great significance. Without a dense atmosphere, Mars lacks the power to deflect penetrating solar wind, cosmic rays or any type of high energy particles. In the history of Mars, there might be a high concentration of carbonic acid. H2CO3 is a water soluble acid and if available in sufficient concentrations, could potentially dissolve metal ores and catalyze chemical reactions. The presence of carbonic acid on Mars may also lead to the existence of limestone (CaCO3), magnesite (MgCO3), dolomite (CaMg(CO3)2), and siderite (FeCO3) [3]. These chemicals have been tentatively detected on the Martian surface [31].

**Acknowledgements**

This work was supported by the National Aeronautics and Space Administration through the NASA Astrobiology Institute under Co-operative Agreement No. NNA04CC08A issued through the Office of Space Science. We would also like to thank Ed Kawamura (University of Hawaii at Manoa, Department of Chemistry) for his electrical work as well as Andrew Taylor for his assistance.
References


