PREDICTED ABUNDANCES OF CARBON COMPOUNDS IN VOLCANIC GASES ON IO

LAURA SCHAEFER AND BRUCE FEGLEY, JR.

Planetary Chemistry Laboratory, McDonnell Center for the Space Sciences, Department of Earth and Planetary Sciences, Washington University, Campus Box 1169, St. Louis, MO 63130; bfegley@levee.wustl.edu

Received 2004 June 23; accepted 2004 September 16

ABSTRACT

We use chemical equilibrium calculations to model the speciation of carbon in volcanic gases on Io. The calculations cover wide temperature (500–2000 K), pressure (10⁻⁸ to 10⁻² bars), and composition ranges (bulk O/S atomic ratios ~0 to 3), which overlap the nominal conditions at Pele (1760 K, 0.01 bars, O/S ~ 1.5). Bulk C/S atomic ratios ranging from 10⁻⁸ to 10⁻¹ in volcanic gases are used with a nominal value of 10⁻³ based on upper limits from Voyager for carbon in the Loki plume on Io. Carbon monoxide and CO₂ are the two major carbon gases under all conditions studied. Carbonyl sulfide and CS₂ are orders of magnitude less abundant. Consideration of different loss processes (photolysis, condensation, kinetic reactions in the plume) indicates that photolysis is probably the major loss process for all gases. Both CO and CO₂ should be observable in volcanic plumes and in Io’s atmosphere at abundances of several hundred parts per million by volume for a bulk C/S ratio of 10⁻³.

Subject headings: astrochemistry — planets and satellites: individual (Io)

Online material: color figures

1. INTRODUCTION

Galileo, Hubble Space Telescope (HST), and Earth-based observations show high-temperature silicate volcanism on Io, the most volcanically active object in the solar system (Kargel et al. 2003; Lopes et al. 2001; Spencer & Schneider 1996). Volcanic gases on Io are composed of sulfur and oxygen compounds (SO₂, S, S₂, SO), with smaller amounts of sodium, chlorine, and potassium compounds such as Na, K, and NaCl (Fegley & Zolotov 2000a; Lellouch et al. 2003; McGrath et al. 2000; Spencer et al. 2000; Zolotov & Fegley 1998, 1999, 2001). A comparison of observations with thermodynamic and kinetic calculations shows that volcanic gases on Io, as on Earth, are in chemical equilibrium when they erupt (Zolotov & Fegley 1998, 1999, 2001; Fegley & Zolotov 2000a, 2000b). Terrestrial volcanic gases are mainly H₂O, SO₂, and CO₂, with smaller amounts of H₂S, CO, H₂, OCS, HCl, and HF (e.g., see Table 6.13 of Lodders & Fegley 1998). Io is depleted in hydrogen, and H-bearing gases such as H₂O, H₂S, and H₂ are not expected in Ionian volcanic gases (Fegley & Zolotov 2000a). But what about carbon gases such as CO₂, CO, and OCS?

There are good reasons to suspect that carbon is present on Io, and here we use chemical equilibrium and kinetic calculations to predict carbon chemistry in Ionian volcanic gases. First, Voyager upper limits for C-bearing gases in the Loki plume (Pearl et al. 1979) are fairly high (0.02%–3% relative to SO₂). Second, the Loki plume observed by Voyager was apparently generated by interaction of a lava flow with sulfur-rich surface material (Geissler 2003). The carbon-to-sulfur ratio in this plume may be lower than in volcanic gases evolved at Pele or other hot spots. Third, carbon-to-sulfur atomic ratios are a few percent or higher (Fig. 1) in lunar rocks, basaltic meteorites, and other samples depleted in volatiles relative to solar abundances. It is plausible that carbon remains in the silicate portion of Io and can be volcanically outgassed. Our chemical equilibrium calculations are described in § 2. We discuss constraints on Io’s carbon abundance in the next section (§ 3). This is followed by the results of our chemical equilibrium calculations for the carbon chemistry of Ionian volcanic gases (§ 4). We then use chemical kinetic calculations to model disequilibrium chemistry in the volcanic plume and find that photolysis is the major loss process for carbon-bearing gases (§ 5). Conclusions are given in § 6. Preliminary results of this work are reported by Fegley & Zolotov (2000b).

2. EQUILIBRIUM CALCULATIONS

We computed the chemical equilibrium abundances of gaseous (i.e., atoms, radicals, ions, electrons, and molecules) species in Ionian volcanic gases containing the elements S, O, Na, K, Cl, and C. The abundance and stability of liquids and solids that may condense from the gas were also computed. The calculations were done using a Gibbs energy minimization code of the type described by Van Zeggen & Storey (1970) and the CONDOR mass-balance, mass-action code described by Fegley & Lodders (1994). The two codes give identical results. Computational methods and thermodynamic data are described elsewhere (Fegley & Lodders 1994; Fegley & Zolotov 2000a).

As in our prior work we did calculations over a range of temperatures (500–2000 K), pressures (10⁻⁸ to 10⁻² bars), and bulk O/S atomic ratios (~0 to 3). We use the Pele volcanic vent as a nominal case because it is the best-characterized volcanic vent on Io. Galileo and Cassini observations indicate that temperatures of the Pele volcanic vent range from about 1400 to 1760 K (Radebaugh et al. 2004; Lopes et al. 2001). We use a nominal temperature of T = 1760 ± 210 K from Galileo NIMS observations (Lopes et al. 2001). Zolotov & Fegley (2001) determined the vent pressure as 0.01 bars using chemical equilibrium calculations. As we show later, our key conclusions are relatively insensitive to temperature and pressure over a fairly wide range of plausible values.

We used the same relative atomic abundances for S (1.0), O (1.521), Na (0.05), K (0.005), and Cl (0.045) as in our previous modeling (Schaefer & Fegley 2005; Fegley & Zolotov 2000a, 2000b; Zolotov & Fegley 2001; Moses et al. 2002). These elemental abundances are based on spectroscopic observations of the Pele plume, Io’s extended atmosphere, and the Io plasma torus (IPT). Carbon has not been detected on Io, and...
3. CONSTRAINTS ON IO’S CARBON ABUNDANCE

We use three data sets to constrain the abundance of carbon in Io: (1) geochemical analyses of carbon and sulfur in stony meteorites, lunar, and terrestrial samples, (2) spectroscopic upper limits on carbon-bearing gases in the Loki and Pele volcanic plumes on Io, and (3) spectroscopic upper limits on ionized carbon in the IPT. All data are plotted in Figure 1.

The C/S ratios measured in meteorites are from the METBASE meteorite database (Koblitz 2003) and are for observed meteorite falls. The lunar data are for soils, breccias, and basalts (Fegley & Swindle 1993). The terrestrial data are for the upper and lower continental crust, present-day mantle, and different rock types including basalt, kimberlite, granite, shale, and ultramafic rocks (Lodders & Fegley 1998).

Chondrites are undifferentiated stony meteorites that contain metal, silicates, and sulfides. They were not melted on their parent bodies. The C/S ratios of chondrites range from 0.011 to 4.27. Carbonaceous chondrites have the highest C/S ratios and ordinary chondrites the lowest ones. Achondrites are differentiated stony meteorites that have lost essentially all metal and sulfides during melting on their parent bodies. The different types of achondrites for which analytical data are available have C/S ratios ranging from 0.10 to 25.4. The highest C/S ratios are found in ureilites, which are carbon-rich and contain diamonds and organic compounds, whereas the lowest C/S ratios are found in the enstatite achondrites (aubrites), mainly composed of enstatite (MgSiO$_3$). The lunar samples have C/S ratios from 0.05 to 0.6. The highest C/S ratios are for soils, while the lowest are for basalts. Finally, the terrestrial samples have C/S ratios ranging from 0.07 for ultramafic rocks to 21.6 for diamond-bearing kimberlites.

Infrared spectroscopy by Voyager gave upper limits for the abundances of several carbon-bearing gases in the Loki plume (Pearl et al. 1979). Pearl et al. (1979) reported 100 nanobars (nbar) SO$_2$ in the Loki plume and give two different upper limits for other gases corresponding to gas temperatures of 130 and 250 K. Following Zolotov & Fegley (1999), we use the arithmetic mean of the upper limits for each gas. The results expressed as volume (or mole) percentages relative to SO$_2$ are CO$_2$ ($<0.05\%$), OCS ($<0.1\%$), CS$_2$ ($<0.02\%$), and CH$_4$ ($<3\%$).

Lellouch et al. (1992) reanalyzed the Voyager data and derived lower SO$_2$ abundances (5–40 nbar) as a function of the assumed gas temperature. If we combined their lower SO$_2$ abundances with the upper limits (cm-am) reported by Pearl et al. (1979), the CO$_2$/SO$_2$, OCS/SO$_2$, CS$_2$/SO$_2$, and CH$_4$/SO$_2$ ratios would become larger than the values we give above. However, since Lellouch et al. (1992) did not redetermine the upper limits for the carbon-bearing gases, we use the values based on the SO$_2$ abundance of 100 nbar (0.2 cm-am) from Pearl et al. (1979), rather than combine abundances determined by separate analyses. This is a reasonable action because we are looking at relative rather than absolute abundances.

Ultraviolet spectroscopy from the HST (McGrath et al. 2000) gives upper limits for CS$_2$ in volcanic plumes at Pele ($<0.64\%$) of SO$_2$ and Ra ($<1.33\%$ of SO$_2$). As our calculations will show, CS$_2$ is a very minor C-bearing gas, so these upper limits do not significantly constrain the bulk atomic C/S ratio at Pele or Ra. Earth-based millimeter-wave spectroscopy gave upper limits of less than (0.6–3) $\times 10^{-2}$ cm-am for CO in Io’s global atmosphere (Lellouch et al. 1992). This corresponds to about 30%–150% of SO$_2$ using the typical SO$_2$ abundance of 0.02 cm-am given by Lellouch et al. (1992). Finally, short-wavelength UV observations of the IPT by Feldman et al. (2004) give upper limits of less than 3.7 $\times 10^{-4}$ for the Iogenic C$^{2+}$/S$^{2+}$ atomic ratio and less than 2.5 $\times 10^{-3}$ for the C$^+$/S$^{2+}$ ratio. With the exception of the millimeter-wave upper limit for CO in Io’s global atmosphere, which is clearly too large, the other spectroscopic upper limits for carbon-bearing gases on Io and for ionized carbon in the Io plasma torus are shown together (as the bar labeled “upper limits on Io”) in Figure 1.

The referee noted that if all upper limits are good quality, only the most stringent one is relevant. This would be the upper limit on C/S ions in the IPT. However, there are a number of different processes (e.g., ionization, atmospheric sputtering, and molecular diffusion) that affect the composition of the IPT (Spencer & Schneider 1996), so these values may not be strictly applicable to the bulk atomic C/S ratios in volcanic gases. In addition, as noted above, we have upper limits for C-bearing gases at several different hot spots. There is no reason to assume that the lava at each of these hot spots has the same bulk C/S ratio (see for instance the range of values for terrestrial samples). We therefore compare the range of upper limits to the values for terrestrial, lunar, and meteoritic samples below.

Taken at face value, the upper limits for Io are significantly smaller than the C/S ratios for meteoritic, lunar, and terrestrial samples. This is somewhat surprising because the Moon and the basaltic achondrites (eucrites) are generally depleted in volatiles (e.g., H$_2$O, C, N, S) relative to CI chondritic abundances, and these objects should be good analogs for Io. In particular, Io...
has about the same size, mass, and density as the Moon, and both objects are bone dry.

One could argue that Io lost its carbon by volcanic outgassing, while the Moon, which is geologically quiescent, retained its carbon. However, the giant-impact model of the formation of the Moon predicts that lunar-forming material was heated to very high temperatures of several thousand kelvins (e.g., Canup 2004). Yet the Moon still contains carbon with about 20–30 μg g⁻¹ carbon in basalts and larger concentrations of carbon in other rock types and soils (Fegley & Swindle 1993). Also, the loss of carbon and sulfur via outgassing on Io may be similar because the molecular masses of carbon and sulfur gases that are observed to be outgassed (SO₂, S₂, SO₃) on Io or Earth (CO, CO₂, OCS) are similar to within a factor of 2.3. Thus, it seems difficult to ascribe the apparent lack of carbon on Io to its high-temperature volcanism.

Alternatively, as the referee suggested, Io’s lithosphere and surface may be enriched in sulfur, rather than depleted in carbon. This seems plausible given the evidence for sulfur compounds in Io’s atmosphere, volcanic gases, and surface. This is especially true for volcanic plumes that are formed by interaction of hot lavas with sulfur-rich surface substrate, such as the plume observed by Voyager at Loki (Geissler 2003) and the persistent plume observed at Prometheus (Kieffer et al. 2000; Milazzo et al. 2001). Vaporization of the surface frost will therefore lower the bulk atomic C/S ratios in some volcanic gases. However, plumes at other hot spots such as Pele are directly exsolved from the hot lava (Spencer et al. 1997), and so should be less contaminated by the sulfur-rich lava. At these hot spots, the bulk atomic C/S ratio may be significantly higher.

The upper limits for carbon on Io overlap with the C/S ratios of ordinary chondrites. Geophysical models predict that Io has a Fe/Si ratio similar to that of the L or LL ordinary chondrites (Kuskov & Kronrod 2000, 2001), or to that of the CV3 or CM2 carbonaceous chondrites (Sohl et al. 2002). Cosmochemical models predict that Io accreted from CV3 or CM2 carbonaceous chondrites—like material (Consolmagno 1981; Lewis 1982; Prinn & Fegley 1981), which contains about 0.5%–2.2% carbon and 2.2%–2.7% sulfur by mass. For reference, the L and LL chondrites contain about 0.25%–0.31% carbon and 2.1%–2.2% sulfur by mass. It is plausible that the bulk C/S ratio of Io is similar to that of ordinary, CV3, or CM2 chondrites.

However, some of the carbon and much of the sulfur in chondritic material would go into a metal-sulfide core, such as Io is believed to have (Anderson et al. 1996). Smaller amounts of C and S would be left in the silicate part of Io. Because of the uncertainties involved, we computed chemical equilibria for C/S ratios ranging from 10⁻⁶ to 10⁻¹. This range spans the upper limits for carbon gases derived from observations, as can be seen in Figure 1. We discuss the results of our chemical equilibrium calculations below.

4. RESULTS OF EQUILIBRIUM CALCULATIONS

4.1. Chemical Reaction and Eruption Timescales

Our thermodynamic calculations assume that Ionian volcanic gases reach chemical equilibrium inside the volcanic conduit, where the high temperatures and pressures lead to characteristic chemical reaction times that are less than the characteristic eruption times (τchem < τerupt). However, the low temperatures and pressures in the volcanic plume lead to the opposite situation, namely (τchem > τerupt), and chemical equilibrium is not attained. In between these two regions, in the vicinity of the volcanic vent, τchem and τerupt become equal and quenching of the high-temperature chemical equilibria occurs during the supersonic eruptions. Figure 1 of Zolotov & Fegley (1999) illustrates our basic assumption. We earlier showed that chemical equilibrium is attained for sulfur and oxygen chemistry (Zolotov & Fegley 1998) and for alkali and halogen chemistry (Schaefler & Fegley 2005). Here we examine carbon chemistry.

The characteristic eruption time is 150–250 s (Zolotov & Fegley 1998). The characteristic chemical reaction times are defined as

\[ \tau_{\text{chem}} = [i]/(d[i]/dt), \]

where \([i]\) is the number density of species “i” given by its equilibrium mole fraction \((X_i)\) times the total number density, and \((d[i]/dt)\) is its destruction rate computed using the reaction rate constants given below. We considered a number of elementary reactions in the C–O–S system (Table 1), and found that reactions R1–R4 rapidly equilibrate at the nominal temperature (1760 K) and pressure (0.01 bars) of the Pele vent (see Table 2). However, reactions between CO and CO₂ such as
R5–R7 are significantly slower than the characteristic eruption times of 150–250 s (Table 2). Reactions R5–R7 may not equilibrate unless they are catalyzed by either the magma and/or rock in the volcanic conduit, and/or other gas-phase chemistry proposed for Venus (catalytic cycle C19 in Appendix E) by Mills (1998). Alternatively, the well-known reaction R8 could equilibrate CO and CO2 in volcanic vents on Io, but the total hydrogen abundance required is very high (~0.1% of the sulfur abundance).

4.2. Carbon Equilibrium Chemistry in Ionian Volcanic Gases

We described the equilibrium chemistry of S, O, Na, Cl, and K in our earlier papers (Fegley & Zolotov 2000a; Zolotov & Fegley 1999), and we focus on carbon equilibrium chemistry here. All carbon is in the gas, and no pure solid or molten carbon compounds are stable under the conditions we considered. Figure 2 shows the mole fractions of the major carbon gases at our nominal conditions ($T = 1760 \pm 210$ K, $P = 0.01$ bars) as a function of the C/S ratio. Our calculated mole fractions for CO and CO2 are almost identical, with CO2 being slightly larger. Carbonyl sulfide is $\sim 2.5$ orders of magnitude less abundant than CO, and CS2 is $\sim 3.5$–$4$ orders of magnitude smaller than OCS. The abundances of all of the carbon gases increase with increasing C/S ratio, which is not surprising. Equations giving the mole fraction of each carbon gas as a function of bulk atomic C/S ratio in the gas at our nominal temperature and pressure are as follows:

\begin{equation}
\log X_{\text{CO}} = -0.27 + 0.99 \log(C/S),
\end{equation}

\begin{equation}
\log X_{\text{CO2}} = -0.33 + 1.00 \log(C/S),
\end{equation}

\begin{equation}
\log X_{\text{OCS}} = -3.04 + 1.01 \log(C/S),
\end{equation}

\begin{equation}
\log X_{\text{CS2}} = -6.64 + 1.03 \log(C/S).
\end{equation}

At our highest C/S ratio, where carbon is 10% of total sulfur, CO2 has an abundance of $\sim 8%$ relative to SO2. At our lowest C/S ratio, where carbon is 0.01% of total sulfur, CO2 has an abundance of $\sim 0.007%$ relative to SO2. Figure 2 also shows the upper limit for CO2 measured by Voyager for the Loki plume (dotted line). This upper limit is not strictly applicable to the Pele hot spot, which we are using for our nominal temperature and pressure conditions. However, we cannot calculate chemical equilibria at Loki because the bulk composition of volcanic gas and the total pressure are unknown. The only data available for Loki are vent temperatures from Galileo and terrestrial observations, detections of SO2 (Pearl et al. 1979) and SO (DePater et al. 2002), and the upper limits mentioned earlier. We can calculate chemical equilibria for Pele where sufficient data are available, but there are no upper limits (except for CS2) on carbon gases. If we accept the upper limit for CS2 as the actual abundance, the gas at Pele would have a bulk C/S atomic ratio greater than 10,000, which is geochemically implausible. Therefore, until more data become available, we apply the upper limits at Loki to Pele. Figure 2 shows that our calculated CO2 abundance exceeds the upper limit for all C/S ratios greater than $\sim 10^{-3}$. We therefore use C/S = $10^{-3}$ for the nominal carbon abundance in our subsequent calculations.

Figure 3 illustrates our results for carbon equilibrium chemistry as a function of temperature and total pressure. The calculations for Figure 3a were performed at the nominal temperature of 1760 K as a function of variable pressure. Our nominal pressure is highlighted on the graph. Figure 3a shows that the abundances of CO2 and OCS increase with increasing pressure, whereas the abundance of CO increases with decreasing pressure. At high total pressures, CO2 is the most abundant gas, followed by CO and OCS. The maximum abundance of OCS at our nominal temperature is only $\sim 0.8%$ of CO2 at a pressure of $10^5$ bars. At pressures greater than our nominal pressure, the CO2 abundance exceeds the Voyager upper limit.

\[ \text{Reaction} \quad \text{Species Destroyed} \quad \tau_{\text{chem}} \quad (s) \]

\begin{tabular}{lccc}
R1 & OCS & $3.00 \times 10^0$ &  \\
R2 & OCS & $3.00 \times 10^{-2}$ &  \\
R3 & CO & $7.00 \times 10^0$ &  \\
R4 & CS2 & $8.00 \times 10^{-1}$ &  \\
R5 & CO & $7.94 \times 10^5$ &  \\
R6 & CO & $1.58 \times 10^6$ &  \\
R7 & CO2 & $2.00 \times 10^6$ &  \\
\end{tabular}
for the Loki plume. The CS₂ mole fraction is less than 10⁻⁸, and we do not expect CS₂ to be observable under any conditions on Io. We therefore believe that the upper limits for CS₂ from McGrath et al. (2000) are much higher than the actual abundances.

Figure 3b shows carbon equilibrium chemistry at our constant nominal pressure for a range of temperatures. Our nominal temperature is highlighted on the graph. At temperatures above our nominal temperature, CO is the major gas, followed by CO₂, OCS, and CS₂. The CO abundance decreases with decreasing temperature, whereas that of CO₂ increases with decreasing temperature. At temperatures below our nominal value of 1760 K, the abundance of CO₂ exceeds the upper limit determined by Voyager for the Loki plume. Loki is apparently a lava lake that periodically overturns (Rathbun et al. 2002). It typically has temperatures of 400 K (Geissler 2003), much less than those of Pele, but peak temperatures at Loki exceed 900 K (Rathbun et al. 2002). We can get CO₂ abundances less than the Voyager upper limits if we decrease (1) the pressure, (2) the C/S ratio, or (3) both. Carbonyl sulfide has a fairly constant abundance from 2000 K down to about 600 K, where the abundance drops sharply. The CS₂ mole fraction is less than 10⁻⁸ and is not shown on the graph.

Figure 4 illustrates the effect of the O/S molar ratio on carbon equilibrium chemistry. This ratio is the bulk O/S ratio of the volcanic gas. Pure SO₂ has O/S = 2, pure SO₃ has O/S = 3, and pure sulfur vapor has O/S equal to zero. Oxygen-to-sulfur ratios from 0 to 2 can be thought of as mixtures of SO₂ + S₂, whereas O/S ratios from 2 to 3 can be thought of as mixtures of SO₂ + O₂. To date, there is no evidence for O/S ratios greater than 2 in volcanic gases on Io. The oxygen fugacity (fO₂, essentially the O₂ partial pressure) varies with temperature, total pressure, and the O/S molar ratio of the volcanic gas (Zolotov & Fegley 1999). The range of fO₂ values as a function of the O/S.

![Fig. 3a](image1.png) ![Fig. 3b](image2.png)

**Fig. 3.—**Equilibrium chemistry of carbon in a volcanic gas containing C, S, O, Na, K, and Cl for a C/S ratio of 0.001 (a) at a constant temperature of 1760 K as a function of pressure, and (b) at a constant pressure of 0.01 bars as a function of temperature. The shaded region in (a) shows the nominal Pele pressure plus uncertainties, and the shaded region in (b) shows the nominal Pele temperature, plus uncertainties. [See the electronic edition of the Journal for a color version of this figure.]

![Fig. 4](image3.png)

**Fig. 4.—**Equilibrium chemistry of carbon in the same volcanic gas as a function of the O/S ratio at otherwise constant conditions (T = 1760 K, P = 0.01 bars, C/S = 0.001). [See the electronic edition of the Journal for a color version of this figure.]
ratio at constant $T$ (1760 K), $P$ (0.01 bars), and C/S ratio ($10^{-2}$) is shown along the top of Figure 4. Carbon dioxide becomes more abundant, while CO, OCS, and CS$_2$ are oxidized by O$_2$ and become less abundant as the $f$ O$_2$ increases, e.g.,

$$2\text{CO} + \text{O}_2 = 2\text{CO}_2,$$  \(7\)

$$2\text{OCS} + 3\text{O}_2 = 2\text{CO}_2 + 2\text{SO}_2,$$  \(8\)

$$\text{CS}_2 + \text{O}_2 = 2\text{CO}_2 + 2\text{SO}_2.$$  \(9\)

Figure 4 shows that CO and CO$_2$ are about equally abundant at the $f$ O$_2$ for Pele volcanic gases. As we discussed previously, Loki is believed to be more oxidizing than Pele (Zolotov & Fegley 1999). Thus if all other variables were the same, the CO$_2$/CO ratio at Loki would be larger than that at Pele. The lower temperature at Loki would also tend to increase the CO$_2$/CO ratio as shown in Figure 3b. However, neither the pressure nor bulk O/S ratio are known, so their effects on the CO$_2$/CO ratio at Loki cannot be quantified at this time.

5. DISCUSSION

5.1. Evidence for the Presence of Carbon on Io

Garrard et al. (1996) describe observations of Jupiter’s magnetosphere by the Galileo heavy ion counter (HIC). They detected highly ionized solar C, which had variable abundances depending on energy. The abundances outside of Io’s orbit (C/O = 0.026 ± 0.007 at 16–17 MeV nucleon$^{-1}$) were significantly larger than the abundances inside of Io’s orbit (C/O = 0.007 ± 0.003 at 16–17 MeV). Their measured S/O ratios were ~1.2 ± 0.2 over all energy ranges, both inside and outside of Io’s orbit. This gives C/S ratios of ~0.022 outside of Io’s orbit and ~0.0058 inside of Io’s orbit. The upper limits for the logenic C$^{2+}$/S$^{2+}$ and C$^+/S^{2+}$ ratios in the IPT are much smaller than these values and are less than 3.7 × 10$^{-4}$ and 2.5 × 10$^{-3}$, respectively (Feldman et al. 2004). The two sets of observations likely give different ratios because of a sampling bias: the HIC was observing highly ionized solar wind particles, whereas the observations of Feldman and colleagues are sampling ionogenic, low-energy particles. However, the lower C/S ratio inside of Io’s orbit observed by the HIC indicates that Io absorbs carbon atoms at these energies. Therefore, it appears that Io has some limited amount of carbon absorbed from the solar wind. If this carbon sticks to the surface, then it may join the cycle of burial and recycling that has been proposed to take place (Keszthelyi et al. 2004). It would be very interesting to determine how much of Io is absorbing; however, such a calculation is outside of the scope of this paper.

5.2. Photochemical Lifetimes of Carbon Gases

We now consider how carbon-bearing gases are destroyed by photochemistry in the volcanic plume. Table 3 lists the photochemical lifetimes of CO, CO$_2$, OCS, and CS$_2$ at Io. We calculated the photochemical lifetimes (at zero optical depth) from the $J_1$ values at 1 AU given in Appendix 1 of Levine (1985), scaled to the distance of Io from the Sun, where

$$J_{\text{chem}} = \frac{1}{J_1}.$$  \(10\)

Table 3 shows that the lifetimes of all carbon gases are long compared to characteristic timescales of ~17 minutes for gas flow over 100 km on Io’s surface and ~20 minutes for a ballistic lifetime within a large plume (see §6 of Moses et al. 2002). The photochemical lifetimes of CO and CO$_2$ are also long compared to an Ionian day (~42 hr).

5.3. Condensation of Carbon Gases on Io’s Surface

Another possible loss mechanism for carbon gases is condensation onto Io’s surface. To determine whether any of the carbon gases may condense on Io, we looked at the vapor pressures of these ices. It is unlikely that CS$_2$ is abundant enough to condense, and CO is too volatile to condense at Io’s surface temperatures. The vapor pressure data for CO$_2$ and OCS from Honig & Hook (1960) at the surface temperatures of Io (90–120 K; Rathbun et al. 2004) give vapor pressure ranges of 7.58–45,586 nbar for CO$_2$ and 63–68,433 nbar for OCS. The atmosphere of Io, which is primarily SO$_2$, has a total pressure of 1–10 nbar. The atmospheric partial pressure of a gas $P_i$(atm) has to equal its vapor pressure over the solid ice $P_i$(ice) for the frozen gas to be stable:

$$P_i(\text{atm}) = P_i(\text{ice}).$$  \(11\)

Otherwise the ice is unstable and all the gas resides in the atmosphere. Therefore, OCS ice is unstable on the surface of Io because the atmospheric partial pressure of OCS is less than the vapor pressure over OCS ice. Carbon dioxide ice could possibly be condensed in colder areas on Io’s surface ($T < 86$ K; Rathbun et al. 2004). Sandford et al. (1991) suggested that a band at 4705.2 cm$^{-1}$ was caused by clusters of CO$_2$ molecules, which could be present on the surface in SO$_2$ or in the atmosphere as aerosol particles. However, they found that the entire surface of Io would have to be dominated by tetramer clusters of CO$_2$ in order to match both the width and location of the band. Schmitt et al. (1994) instead assigned the same band to an SO$_2$ overtone. We conclude that condensation is unlikely to be a significant loss mechanism for carbon-bearing gases (except possibly CO$_2$) on Io.

5.4. Chemical Destruction of Carbon Gases

Photochemical modeling of Io’s atmosphere and volcanic plumes (Moses et al. 2002 and references therein) indicates that the most abundant reactive species are O, S, SO, and O$_2$. We looked at a number of reactions of these species with CO, CO$_2$, OCS, and CS$_2$ to determine which, if any, chemical reactions could destroy these carbon gases (see Table 1). We calculated chemical lifetimes for CO, CO$_2$, OCS, and CS$_2$ assuming surface conditions similar to those used in Moses et al. (2002) of 110 K, $P_T = 10$ nbar, and a total number density of 10$^{11.82}$ cm$^{-3}$. All of these reactions give chemical lifetimes (Table 4) much longer than the photochemical lifetimes (Table 3). The fastest of these reactions is R4, which gives a chemical lifetime for
CS$_2$ of $\sim 146$ days. This is very large compared with the photochemical lifetime of $\sim 58$ minutes for CS$_2$. We conclude that photolysis is probably the primary loss mechanism for CO$_2$, CO, OCS, and CS$_2$ in volcanic plumes and Io’s atmosphere.

6. CONCLUSIONS

Chemical equilibrium calculations predict that CO and CO$_2$ are the two most important carbon compounds in volcanic gases on Io. Carbonyl sulfide and CS$_2$ are much less abundant and will be difficult, if not impossible, to observe. The upper limits reviewed in Figure 1 and our calculations support an average upper limit for the bulk atomic C/S ratio of less than $10^{-3}$ for volcanic gases on Io. Using this bulk C/S ratio we predict CO/SO$_2 < 6 \times 10^{-4}$ and CO$_2$/SO$_2 < 8 \times 10^{-4}$ at Pele. Our considerations of photolysis, condensation, and disequilibrium reactions indicate that photolysis is probably the major loss process for CO, CO$_2$, OCS, and CS$_2$. The atmospheric partial pressures of these gases are too low for their pure ices to be stable on Io’s surface; however, carbon dioxide may condense in cold traps on the surface. Their long photochemical lifetimes indicate that CO and CO$_2$ may be present in Io’s atmosphere at all times. However, CO will most likely be observable during high-temperature (>1500 K) volcanic eruptions. Carbon dioxide will be most abundant in low-temperature/high-pressure eruptions. It may be possible to observe OCS in a volcanic plume during an active eruption with a sufficiently high eruption pressure. A detection of CS$_2$ gas would indicate a lava with a very high C/S ratio. We particularly recommend searching for CO$_2$, especially near active volcanic vents such as Pele, where its abundance should be greater. We also recommend searching for CO during high-temperature eruptions, when it should be more abundant.

We thank K. Lodders and M. McGrath for helpful comments and J. Spencer for a constructive review and helpful comments. This work was supported by the NASA Planetary Atmospheres Program.

REFERENCES

Anderson, J. D., J. S. Jeffers, P., Lifshitz, A., & Yadava, B. P. 1971, Int. Symp. Combustion Processes, 13, 417
Canup, R. M. 2004, Icarus, 168, 433
Consolmagno, G. J. 1981, Icarus, 47, 36
DeMore, W. B., et al. 1997, JPL Publ. 97-4, 1-266
De Pater, I., Roe, H., Graham, J. R., Strobel, D. F., & Bernath, P. 2002, Icarus, 156, 296
Fegley, B., Jr., & Lodders, K. 1994, Icarus, 110, 117
Fegley, B., Jr., & Swindle, T. D. 1993, in Resources of Near-Earth Space, ed. J. Lewis, M. S. Matthews, & M. L. Guerrieri (Tucson: Univ. Arizona Press), 367
Fegley, B., Jr., & Zolotov, M. Yu. 2000a, Icarus, 148, 193
—. 2000b, Meteoritics Planet. Sci., 35, A52
Feldman, P. D., Strobel, D. F., Moos, H. W., & Weaver, H. A. 2004, ApJ, 601, 583
Garrard, T. L., Stone, E. C., & Murphy, N. 1996, Science, 274, 393
Geissler, P. E. 2003, Annu. Rev. Earth Planet. Sci., 31, 175
Honig, R. E., & Hook, H. O. 1960, RCA Rev., 21, 360
Kargel, J. S., et al. 2003, EOS, 84, 313
Keszthelyi, L., et al. 2004, Icarus, 169, 271
Kieffer, S. W., Lopes-Gautier, R., McEwen, A., Smythe, W., Keszthelyi, L., & Carlson, R. 2000, Science, 288, 1204
Klemm, R. B., & Davies, D. D. 1974, J. Phys. Chem., 78, 1137
Koblitz, J. 2003, METBASE Ver. 6.0 for Windows
Kuskov, O. L., & Kronrod, V. A. 2000, Planet. Space Sci., 48, 717
—. 2001, Icarus, 151, 204
Lellouch, E., Belton, M., De Pater, I., Paubert, G., Gulkis, S., & Encrenaz, T. 1992, Icarus, 98, 271
Lellouch, E., Paubert, G., Moses, J. I., Schneider, N. M., & Strobel, D. F. 2003, Nature, 421, 45
Levine, J. S. 1985, The Photochemistry of Atmospheres: Earth, the Other Planets, and Comets (London: Academic Press)
Lewis, J. S. 1982, Icarus, 50, 103
Lodders, K., & Fegley, B., Jr. 1998, The Planetary Scientist’s Companion (New York: Oxford Univ. Press)
Lopes, R. C. M., et al. 2001, J. Geophys. Res., 106, 33053
McGrath, M. A., Belton, M. J. S., Spencer, J. R., & Sartoretti, P. 2000, Icarus, 146, 476
Milazzo, M. P., Keszthelyi, L. P., & McEwen, A. S. 2001, J. Geophys. Res., 106, 33121
Mills, F. P. 1998, Ph.D. thesis, Caltech
Moses, J. I., Zolotov, M. Yu., & Fegley, B., Jr. 2002, Icarus, 156, 107
Pearl, J., et al. 1979, Nature, 280, 755
Phleger, R. H., & Yung, Y. L. 1984, Int. J. Chem. Kinetics, 16, 1531
Prinn, R. G., & Fegley, B. Jr. 1981, ApJ, 294, 308
Radebaugh, J. S., McEwen, A. S., Milazzo, M. P., Keszthelyi, L. P., Davies, A. G., Turtle, E. P., & Dawson, D. D. 2004, Icarus, 169, 65
Rathbun, J. A., Spencer, J. R., Davies, A. G., Howell, R. W., & Wilson, L. 2002, Geophys. Res. Lett., 29, 1443, doi: 10.1029/2002GL014747
Sandford, S. A., Salama, F., & Allamandola, L. J. 1991, Icarus, 91, 125
Schafer, L., & Fegley, B., Jr. 2005, Icarus, in press
Schmitt, B., De Bergh, C., Lellouch, E., Maillard, J.-P., Barbe, A., & Douet, S. 1994, Icarus, 111, 79
Sohl, F., Spohn, T., Breuer, D., & Nagel, K. 2002, Icarus, 157, 104
Spencer, J. R., Jessup, K. L., McGrath, M. A., Ballester, G. E., & Yelle, R. 2000, Science, 288, 1208
Spencer, J. R., Sartoretti, P., Ballester, G. E., McEwen, A. S., Clarke, J. T., & McGrath, M. A. 1997, Geophys. Res. Lett., 24, 2471
Spencer, J. R., & Schneider, N. M. 1996, Annu. Rev. Earth Planet. Sci., 24, 125
Tsang, W., & Hamspson, R. F. 1986, J. Phys. Chem. Ref. Data, 15, 1087
Van Zeggern, F., & Storey, S. H. 1970, The Computation of Chemical Equilibria (Cambridge: Cambridge Univ. Press)
Woodbridge, M. S., Hanson, R. K., & Bowman, C. T. 1994, Int. J. Chem. Kinetics, 26, 361
Yung, Y. L., & DeMore, W. B. 1999, Photochemistry of Planetary Atmospheres (New York: Oxford Univ. Press)
Zolotov, M. Yu., & Fegley, B., Jr. 1998, Icarus, 132, 431
—. 2001, Lunar Planet. Sci. Conf., 32, 1474

TABLE 4

| Reaction | Species Destroyed | $t_{chem}$ (s) |
|----------|------------------|----------------|
| R1       | OCS              | $2.51 \times 10^3$ |
| R2       | OCS              | $9.33 \times 10^3$ |
| R3       | CO               | $3.98 \times 10^5$ |
| R4       | CS$_2$           | $1.26 \times 10^7$ |
| R5       | CO               | $1.00 \times 10^7$ |
| R6       | CO               | $7.94 \times 10^9$ |
| R7       | CO$_2$           | $1.00 \times 10^9$ |
| R8       | CO               | $1.58 \times 10^2$ |