Hydroxide Salts in the Clouds of Venus: Their Effect on the Sulfur Cycle and Cloud Droplet pH

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Abstract

The depletion of SO2 and H2O in and above the clouds of Venus (45–65 km) cannot be explained by known gas-phase chemistry and the observed composition of the atmosphere. We apply a full-atmosphere model of Venus to investigate three potential explanations for the SO2 and H2O depletion: (1) varying the below-cloud water vapor (H2O), (2) varying the below-cloud sulfur dioxide (SO2), and (3) the incorporation of chemical reactions inside the sulfuric acid cloud droplets. We find that increasing the below-cloud H2O to explain the SO2 depletion results in a cloud top that is 20 km too high, above-cloud O2 three orders of magnitude greater than observational upper limits, and no SO above 80 km. The SO2 depletion can be explained by decreasing the below-cloud SO2 to 20 ppm. The depletion of SO2 in the clouds can also be explained by the SO2 dissolving into the clouds, if the droplets contain hydroxide salts. These salts buffer the cloud pH. The amount of salts sufficient to explain the SO2 depletion entails a droplet pH of ~1 at 50 km. Because sulfuric acid is constantly condensing out into the cloud droplets, there must be a continuous and pervasive flux of salts of $\approx 10^{-13}$ mol cm$^{-2}$ s$^{-1}$ driving the cloud droplet chemistry. An atmospheric probe can test both of these explanations by measuring the pH of the cloud droplets and the concentrations of gas-phase SO2 below the clouds.

Unified Astronomy Thesaurus concepts: Venus (1763); Planetary atmospheres (1244); Water vapor (1791); Clouds (1262)

Supporting material: machine-readable tables

1. Introduction

Both sulfur dioxide (SO2) and water vapor (H2O) are known to be depleted in the cloud layer of Venus (see, e.g., Vandeaele et al. 2017a; Biersen & Zhang 2020) and to vary in abundance above the cloud top by an order of magnitude or more both spatially (Jessup et al. 2015; Encrenaz et al. 2019; Marcq et al. 2020) and temporally in years-long cycles (Marcq et al. 2013; Vandeaele et al. 2017b). Both of these species participate in Venus’s atmospheric sulfur cycle (Yung & Demore 1982; Krasnopolsky 1982, 2007, 2010a, 2012; Mills et al. 2007; Zhang et al. 2012; Biersen & Zhang 2020). Their photodestruction in the upper cloud layer (60–70 km) leads to the formation of sulfuric acid (H2SO4), which condenses out and forms the clouds in Venus’s atmosphere (Yung & Demore 1982). The droplets rain out of the clouds at a height of $\approx 48$ km, where they evaporate (Yung & Demore 1982; Krasnopolsky 2007). The H2SO4 then dissociates and replenishes SO2 and H2O in the lower atmosphere (e.g., Krasnopolsky 2007). The behavior of all other known chemically reactive species in Venus’s atmosphere is influenced by this cycle (Krasnopolsky 2007, 2010a, 2012), and many of these species participate in this cycle. The sulfur cycle in the atmosphere of Venus establishes a strong and persistent redox gradient through the atmosphere of Venus. Venus is more reduced above the clouds and more oxidized below the clouds.

Though this cycle is central to the atmospheric chemistry of Venus, it is not fully understood, and no self-consistent full-atmospheric model of Venus yet accounts for this cycle.

There are several models of the lower atmosphere of Venus (0–40 km) that account for the efficient evaporation of H2SO4 and the effect of its dissociation products, SO2 and H2O, on the abundances of carbon monoxide (CO), carbonyl sulfide (OCS), and SO2 (Krasnopolsky 2007, 2013). Other models describe the middle atmosphere of Venus (60–120 km; e.g., Zhang et al. 2012), investigating the chemistry above the clouds (60–80 km) where SO2 is depleted and then reappears between 85 and 105 km (Sandor et al. 2010; Belyaev et al. 2012). Zhang et al. (2010) propose that night-side evaporation of H2SO4 at 85–105 km, followed by rapid displacement to the day side by strong winds and subsequent photodissociation, can explain this behavior. The distribution of upper atmospheric aerosols at the day and night sides (Parkinson et al. 2015) could be applied as constraints for the proposed mechanism of Zhang et al. (2010). Pinto et al. (2021) suggest an alternative hypothesis, involving (SO2) chemistry. There has recently been a model of the upper cloud layer of Venus exploring the SO2 depletion from 1 ppm to ~10 ppb, and the correlation with H2O abundances in the clouds (Shao et al. 2020).

A diagram of the sulfur cycle and its connection to other trace atmospheric species in the atmosphere of Venus is shown in Figure 1.
At least three atmospheric models of Venus also exist, but they either do not predict the observed SO2 depletion (Yung et al. 2009), do not couple the SO2 depletion to the sulfur cycle (Greaves et al. 2020a), or do not consider the H2O and SO2 depletion in concert. The best current full-atmospheric model that accounts for the SO2 depletion, from Bierson & Zhang (2020), reproduces the SO2 reasonably well, though only by fixing the H2O profile and by inhibiting the vertical transport within the clouds, as suggested by Marcq et al. (2018).

To explain the depletion of sulfur dioxide and water in concert, either some unknown chemistry must take place within the cloud layer, or observations of lower atmospheric SO2 and/or H2O must be mistaken. We explore both of these possibilities in this paper.

In Section 2 we show why the sulfur cycle can be explained by decreasing the amount of hydrogen in the lower atmosphere or increasing the amount of hydrogen in the clouds, either by increasing the water vapor in the clouds or by transporting the hydrogen into the clouds in a different form. We propose that hydrogen could be contained either in aerosols that are lifted by winds into the clouds from the surface, delivered exogenously, or contained within the clouds within some unknown chemical species. In Section 3, we discuss our full-atmospheric model for Venus. In Sections 4 and 5 we present our two hypotheses: that the observational constraint on SO2 and/or H2O are wrong and that cloud chemistry may explain the sulfur depletion. We then show the results of our model if the observational constraints on SO2 and H2O are wrong (Sections 6.2 and 6.1) or if we introduce cloud chemistry (Section 6.3). In Section 6.3 we also predict the effect of this source of hydrogen on the cloud chemistry, chiefly on how it would act as a pH buffer in the clouds. We discuss the implications of our results in Section 7, particularly about how rainout and replenishment of hydrogen is needed to sustain the SO2 gradient. We also speculate about possible sources of hydrogen and their delivery into the clouds, and ways of reconciling the changing cloud chemistry with observations. Section 8 contains our conclusions.

2. The Puzzle of Sulfur Depletion

There is good observational evidence that the concentration of SO2, the dominant sulfur-bearing species in Venus’s atmosphere, varies by several orders of magnitude between altitudes of 40–80 km (see Appendix A). The concentration of SO2 is above 100 ppm at 40 km and between 1 and 100 ppb at 80 km, and we call the decrease in SO2 with height the “depletion” of SO2. The standard explanation for this depletion predicts the major presumed constituent of the clouds of Venus: H2SO4. Near the top of the clouds,

\[
\text{H}_2\text{SO}_4 \rightarrow \text{H}_2\text{O} + \text{SO}_2 + \text{O}_2
\]

The H2SO4 condenses out to droplets that then drop down to ∼40 km, where the droplets evaporate. These droplets are predicted to make up the cloud layer. This mechanism ends up reducing the upper atmosphere by replacing CO2 with CO and SO2 with SO. The excess oxygen is bound up in the sulfuric acid that has condensed out of the atmosphere.

An oxygen atom is needed to form SO3, and this O will probably come from either CO2 or SO2 because they are by far the most abundant O-containing molecules. The formation of one molecule of H2SO4 is the destruction of one molecule of H2O and at least one molecule of SO2 (the sum of Reactions (1) and (3)), and at most two molecules of SO2 (the sum of Reactions (2) and (3)). Between one and two molecules of SO2 are lost with every molecule of H2O to make a molecule of H2SO4 this way, so this mechanism predicts that the
below-cloud H₂O and SO₂ concentrations are within a factor of 2 of each other. Therefore, the maximum depletion of SO₂ in the atmosphere by this mechanism is equal to \((\chi + 1)\text{[H₂O]}\), where [H₂O] \((\text{cm}^{-3})\) is the atmospheric mixing ratio of H₂O and \(\chi\) is the fraction of H₂SO₄-bound O that was produced by SO₂ dissociation.

The observational constraints, however, are [SO₂] \(\approx 150\text{ ppm}\) and [H₂O] \(\approx 30\text{ ppm}\). Even if \(\chi = 1\) and all water were converted to H₂SO₄, the SO₂ would only be depleted by \(\sim 20\%\). This is insufficient to explain the several-orders-of-magnitude depletion of SO₂.

Therefore, the SO₂ depletion is a puzzle for which there is no successful solution in the literature consistent with observations. This implies that either the observational constraints on H₂O and/or SO₂ in the lower atmosphere are wrong and their abundances below the clouds are within a factor of 2 of each other, or that some alternative chemistry explains the SO₂ depletion. We are not the first to notice the implications of this puzzle: a missing sulfur reservoir (see, e.g., Yung & Demore 1982; Parkinson et al. 2015; Marqu et al. 2018).

One alternative mechanism to explain the SO₂ depletion is the formation of condensible sulfur allotropes out of SO₂ photodissociation or thermal dissociation products. The remaining sulfur cannot be in the form of SO, because the resulting concentrations of SO would be at least two orders of magnitude greater than indicated by above-cloud observations. This explanation requires photodissociation of both SO and SO₂ near the cloud top that is many orders of magnitude more efficient than predicted by any model, or similarly more efficient thermal dissociation of SO₂, and must explain 80% of the SO₂ depletion. Twenty percent of the SO₂ will be converted into SO₃ and will react with H₂O to form H₂SO₄ condensing out. Another 20% will balance the reducing power of the SO₃ removal. The remaining 60% would have to go through either or both of the total reactions:

\[
2\text{SO}_2 \rightarrow \text{S}_2 + 2\text{O}_2, \quad \text{Thermochemistry};
\]

\[
2\text{SO}_2 + 4h\nu \rightarrow \text{S}_2 + 2\text{O}_2 \quad \text{Photochemistry.}
\]

This would either predict 150 ppm additional O₂ in the upper atmosphere of Venus, exacerbating the O₂ overabundance problem discussed in the Introduction, or would lead to oxidation of CO. However, oxidation of CO would cause it to become depleted, whereas we see that CO increases above the clouds. It is not possible that the excess oxygen would remain in the form of atomic oxygen because atomic oxygen is not chemically stable at above-cloud altitudes. It is possible that the oxygen is stored in some other chemical species that has not yet been identified, but thus far there is no known candidate species at several ppm concentrations needed to contain the excess oxygen. For these reasons we do not consider this explanation further here.

### 3. The Model

For this work we use a photochemistry-diffusion model based on the model of Rimmer & Helling (2016). The model is composed of a solver and a network. The solver, ARGO, solves the time-dependent set of coupled nonlinear differential equations:

\[
d\chi_X/dt = P_X - L_X\chi_X - \partial \Phi_X/\partial z,
\]

where \(\chi_X \,(\text{cm}^{-3})\) is the number density of species X at height z (cm) and time t (s), \(P_X \, (\text{cm}^{-3} \text{ s}^{-1})\) is the rate of production of X at height z and time t, \(L_X \, (\text{s}^{-1})\) is the rate constant for the destruction of species X at height z and time t. The term \(\partial \Phi_X/\partial z \, (\text{cm}^{-3} \text{ s}^{-1})\) describes the divergence of the vertical diffusion flux.

As described by Rimmer & Helling (2016), the chemistry is solved by following the motion of a parcel up and down through a one-dimensional atmosphere described by a grid of set temperature, \(T \,(\text{K})\); pressure, \(p \,(\text{bar})\); and other properties. A parcel starts at the surface with a particular set of initial chemical conditions and then moves through the atmosphere at a velocity determined by the Eddy diffusion coefficient:

\[
t_{\text{chem}} = (\Delta z)^2/2K_z,
\]

where \(\Delta z \,(\text{km})\) is the change in height from one part of the grid to the next, and \(K_z \,(\text{cm}^2 \text{ s}^{-1})\) is the Eddy diffusion coefficient. The volume mixing ratios are recorded for each species at each grid height, constructing chemical profiles for the atmosphere. There is a method to account for molecular diffusion, described by Rimmer & Helling (2016) and corrected by Rimmer & Helling (2019). We do not solve above Venus’ homopause, so molecular diffusion, though included, is not significant, and we do not describe it in detail here. After the parcel makes a single complete trip, a UV radiative transfer model is run on the recorded profiles, completing a single global iteration and recording the actinic flux \(F_\lambda\) (photons cm⁻² s⁻¹), which will be excluded from the units hereafter.

This method reproduces results for modern Earth and Jupiter (Rimmer & Helling 2016) and agrees with Eulerian solvers for chemical-quenching heights in hot Jupiter atmospheres (Tsai et al. 2017; Hobbs et al. 2019, 2020).

Some of the values of \(L_X\) (and corresponding production coefficients) are the result of direct photodissociation and photoionization. These are all set to zero for the first global iteration and then are calculated for global iteration \(I - 1\) using the chemical profiles of global iteration \(I - 1\). The photodissociation and photoionization rate constant for species X are

\[
k_\lambda(X) = \frac{1}{2} \int F_\lambda \sigma_\lambda(X) d\lambda,
\]

where \(\lambda \,(\text{Å})\) is the wavelength, \(z \,(\text{km})\) is the atmospheric height, and \(\sigma_\lambda(X) \,(\text{cm}^2)\) are the photochemical cross sections (see Appendix B). The factor of 1/2 is typically included to account for the rotation of the planet. Venus rotates too slowly to include this factor for the same reason and would typically be treated as having a day-side and night-side chemistry. However, the zonal winds of Venus are very strong, horizontal mixing is fast, and for simplicity we consider the atmosphere to be a well-mixed average of day-side and night-side chemistries, which will be true for the long-lifetime and medium-lifetime species. The flux at height \(z\) is given by

\[
F_\lambda(z) = F_\lambda(z_0) e^{-(z + z_0)/\mu_0} + F_{\text{diff}}.
\]

Here \(F_\lambda(z_0) \,(\text{cm}^{-2} \text{s}^{-1} \text{ Å}^{-1})\) is the top-of-atmosphere (TOA) flux as described in Section 3.1, \(F_{\text{diff}} \,(\text{cm}^{-2} \text{s}^{-1} \text{ Å}^{-1})\) is the diffuse flux.

\(8\) The actinic flux is the total (direct and diffusive) spectral irradiance integrated over a unit sphere.
from scattering (see Rimmer & Helling 2016), the cosine of the average zenith angle $\mu_0 = 0.54$ (see Hu et al. 2012), and $\tau$ is the optical depth from molecular absorption calculated from the chemical profiles using the prior global solution as well as the photochemical cross sections (see Appendix B for details). In addition, $\tau_a$ is the additional optical depth due to Venus’s mysterious UV absorber, described in Section 3.1.

Beyond these photodissociation and photoionization rate constants, the coefficients used to construct $P_X$ and $L_X$ are provided by the chemical network, STAND2020, which we introduce here. We start with the sulfur network of Hobbs et al. (2020) and add all relevant reactions involving species without thermochemical data from Greaves et al. (2020a). The network includes 2901 reversible reactions and 537 irreversible reactions involving 480 species composed of H/C/N/O/S/Cl and a handful of other elements, including condensation of H_2O and a host of other species, most of which condense at temperatures far lower than achieved in Venus’s atmosphere; see Appendix B. The network, including added condensation chemistry for sulfuric acid and sulfur allotropes, is described in detail in Appendix B.

In addition, we track the dissolution of SO_2 into the cloud droplets and subsequent liquid-phase chemistry for the cloud chemistry model described in Section 5.

The chemical profiles from the most recent global iteration, $I$, are compared to the profiles from the next most recent global iteration, $I - 1$, in order to determine convergence. Convergence criteria are the same as for Greaves et al. (2020a).

We give the parameters and initial conditions for our model in Section 3.1, including the temperature profile, Eddy diffusion, stellar irradiation, and surface boundary conditions for the chemistry. In Section 3.2, we discuss the consistency of these initial conditions compared to chemical equilibrium at the surface.

### 3.1. Parameters and Initial Conditions

The initial conditions and parameters we use are similar to those used by Greaves et al. (2020a). We use the same fixed temperature profile as Greaves et al. (2020a), which was initially taken from Krasnopolsky (2007, 2012). We use Eddy diffusion profiles from the same sources, though we also explore the effect of using the in-cloud Eddy diffusion coefficients of Bierson & Zhang (2020) in Section 4. The profiles we use are shown in Figure 2.

We use a scaled top-of-atmosphere (TOA) solar spectrum from 1–10000 Å, compiled by Granville-Willet (2017), for our TOA boundary condition. These data were compiled using Matthes et al. (2017) for the 401–1149 Å spectral region and Coddington et al. (2015) for the other wavelengths. The actinic flux is then multiplied by 1.913 to account for the difference in average distances of Earth and Venus from the Sun. The TOA spectrum is provided in Harvard Dataverse: doi:10.7910/DVN/AKUTME.

We also include, in addition to molecular absorption and scattering as described above in Section 3, a parameterization of the mysterious UV absorber present within Venus’s cloud layer. The parameterization originates with Krasnopolsky (2012) and is the same as that used by Greaves et al. (2020a), it takes the form

$$ \frac{d\tau_a}{dz} = 0.056 \text{ km}^{-1} \exp \left\{ \frac{-z - 67 \text{ km}}{3 \text{ km}} - \frac{\lambda - 3600\text{Å}}{1000\text{Å}} \right\}, \quad z > 67 \text{ km}; $$

$$ \frac{d\tau_a}{dz} = 0.056 \text{ km}^{-1} \exp \left\{ -\frac{\lambda - 3600\text{Å}}{1000\text{Å}} \right\}, \quad 58 \text{ km} < z \leq 67 \text{ km}; $$

$$ \frac{d\tau_a}{dz} = 0.0 \text{ km}^{-1}, \quad z \leq 58 \text{ km}, $$
The initial surface mixing ratios we set for Venus’s atmosphere are given in Table 1, defining the bulk atmosphere. For all of the models presented outside of Section 7.4, we do not include PH3. Now that we have defined our model and parameters, we will lay out the different scenarios that we consider for explaining the observed SO2 depletion.

3.2. Equilibrium Surface Composition

Here we discuss the consistency of the chemical boundary conditions (Table 1) compared to chemical equilibrium and the implications these conditions may have on surface mineralogy. We find there are solutions where the gas-phase chemistry is consistent with our chosen boundary conditions and the condensed-phase chemistry is broadly consistent with observed surface mineral compositions. We will explore these conditions and their implications for hypothetical cloud chemistry in Section 7.

In 1985 June, the Vega 2 Lander determined the composition of the Venusian surface rock in the northern region of Aphrodite Terra (Surkov et al. 1986, see also Fegley 2014). The rock was analyzed by X-ray fluorescence employing instrumentation that had been improved based on the experience with the previous Venera 13 and 14 missions (Surkov et al. 1982, 1984). The measured oxide ratios are listed in Table 2. We have used these data, in combination with the observed gas composition at the surface in Table 1, to investigate the question of how far the gas at the bottom of the Venusian atmosphere is in chemical equilibrium and in phase equilibrium with the surface rock.

These investigations were carried out by means of the chemical and phase-equilibrium code GGCHEM (Woitke et al. 2018) taking into account the following 16 elements: H, C, N, O, F, S, Cl, Fe, Mn, Si, Mg, Ca, Al, Na, K, and Ti. No information is available about phosphorous at the Venusian surface, so we have excluded that element from this investigation. GGCHEM finds 442 gas-phase species (atoms, ions, molecules, and molecular ions) and 190 condensed species in its databases for this selection of elements. The thermochemical data for the molecules are based on the NIST-Janaf tables (Chase et al. 1982; Chase 1986, 1998), as fitted by Stock (2008), with some additions for diatomic molecules from Barklem & Collet (2016). Condensed-phase data are taken from the SUPCRTBL database (Zimmer et al. 2016) and from the NIST-Janaf database. Some additional vapor pressure data are taken from Yaws (1999), Weast (1971), Ackerman & Marley (2001), and Zahnle et al. (2016).

We consider a mixture of gas and condensed species at $T = 735$ K and $p = 90$ bars with total (gas+condensed) element abundances $e^0$; see Herbert et al. (2020, their Figure 1). In order to find these element abundances we first convert the solid oxide mass ratios given in Table 2 into element particle ratios. Second, we multiply by an arbitrary factor of 1000 and then add the observed gas-phase element abundances from Table 1. Third, we carefully adjust the total oxygen abundance $e^0_O$ until the gas over the condensates has a SO2 concentration of 150 ppm in the model. The arbitrary factor in preparation step two causes the model to produce mostly condensed phases with only small amounts of gas above it. That factor has little influence on the results as long as it is large. The reason for this behavior is that once GGCHEM has determined the solid composition in the form of active condensates, which have a supersaturation ratio $S = 1$ (all other condensates have $S < 1$), one can add arbitrary amounts of those condensates to $e^0_O$; they will just fall out again without changing the resulting gas composition.

The results of this model are shown in Table 3. The resultant solid composition of the Venusian surface rock is a felsic mixture of enstatite (MgSiO3[s]) and quartz (SiO2[s]). The condensates anorthite (CaAl2Si2O8[s]), albite (NaAlSi3O8[s]), and microcline (KAlSi3O8[s]) are the three main minerals forming feldspar, which is found, e.g., in basaltic rock on Earth. Iron is found to be entirely bound in magnetite (Fe2O3[s]). No carbonates and no phyllosilicates are found to be stable under the assumed conditions, nor any minerals containing chlorine. The only halide found to be stable is magnesium fluoride (MgF2[s]). Therefore, all carbon, nitrogen, hydrogen, and chlorine assumed in the model are present in the gas, which allows us to directly fit the observed gas-phase concentrations of CO2, N23, H2O, and HCl.

Fitting the SO2 concentration is more difficult because sulfur is mostly contained in anhydrite (CaSO4). In the close vicinity of the equilibrium solution outlined in Table 3, we see that additional oxygen is used to form more anhydrite in the model on the expense of gaseous SO2 and anorthite (CaAl2Si2O8[s]) via

$$O + SO_2 + CaAl_2Si_2O_8[s] \leftrightarrow CaSO_4[s] + Al_2O_3[s] + 2SiO_2[s],$$

(12)

which is a potentially important buffer mechanism to understand the SO2 concentration in the lower Venus atmosphere. It
allows us in the model to adjust the total oxygen abundance \( O_2^{0} \) to find the desired \( SO_2 \) concentration (more oxygen means less \( SO_2 \)).

Table 3 shows that it is possible to explain both the observed composition of the near-crust Venustian atmosphere and the solid composition of the surface rock by a simple consistent-phase-equilibrium model. All molecules that are predicted to be abundant in our model (those with percent or ppm concentrations) have observed counterparts. The abundance hierarchy matches between model and observations. Other common molecules like \( CH_4 \) and \( NH_3 \), which have extremely small abundances (<10^{-15}) in our model, are not observed. The Venus atmosphere can hence be classified as a type B atmosphere according to Woitke et al. (2021), along with the atmospheres of Earth and Mars. All predicted molecular concentrations are in reasonable agreement with the observed values, in particular when taking into account the large measurement uncertainties (see Table 4). We note, however, that this does not prove that the Venustian atmosphere is in chemical and phase equilibrium with the surface rock; it only shows that the data can be interpreted that way.

For the water-rich scenario (see Section 4), we can increase the hydrogen abundance to find a model with 200 ppm \( H_2O \), which has little effect on \( CO \) and \( OCS \), but results in slightly increased abundances of the ppb molecules, 149 ppm \( SO_2 \), 12 ppm \( CO \), 9 ppm OCS, 505 ppb HCl, 505 ppb HF, 110 ppb \( S_2 \), 370 ppb \( H_2S \), 110 ppb \( S_2O \), and 18 ppb \( H_2 \), which is arguably still consistent with the observations. For the sulfur-poor scenario, we can increase the oxygen abundance to find a model with 20 ppm \( SO_2 \). In that case, the atmosphere is found to have a purer, more oxidizing character with 30 ppm \( H_2O \), 2 ppm \( CO \), 4 ppb OCS, 505 ppb HCl, 355 ppb HF, and \( S_2 \), \( H_2S \), \( S_2O \), and \( H_2 \) all <1 ppb, which seems inconsistent with the observations.

4. Hypothesis: The Observational Constraints Are Wrong

One possibility to consider is that the below-cloud observational constraints are incorrect. Possibly the below-cloud water vapor is much higher than observations suggest, a possibility considered by Yung & Demore (1982). Or the below-cloud sulfur dioxide is much lower than most observations suggest.

In order to explore the sulfur-poor hypothesis we vary the below-cloud \( SO_2 \) abundance from 80 ppm down to 6 ppm with below-cloud \( H_2O \) kept at the nominal value (30 ppm). To explore the water-rich hypothesis we vary the below-cloud \( H_2O \) abundance from 30 ppm up to 200 ppm with \( SO_2 \) kept at the nominal value (150 ppm). In this case we find that the observed above-cloud \( SO_2 \) depletion is not achieved for any value of below-cloud \( H_2O \) due to \( H_2O \) self-shielding effects. Some support for the water-rich hypothesis is the >100 ppm abundances of water vapor observed in and directly below the clouds of Venus (Mukhin et al. 1982; Surkov et al. 1982; Bell et al. 1991), though these are inconsistent with <100 ppm abundances close to the surface (Bertaux et al. 1996).

To explore the possibility of the water-rich hypothesis further, we test the effect of introducing a trap in the eddy diffusion profile within the cloud layer alongside varying the \( H_2O \) abundance below the clouds. Observational constraints on the eddy diffusivity as a function of altitude in the atmosphere are sparse. Marqu et al. (2018) have suggested that the existence of statically stable layers in the cloud region may inhibit dynamical exchange between the upper and lower regions of the atmosphere, a possibility explored by Bierson & Zhang (2020). In the present work we test this possibility by modifying the nominal eddy diffusion profile taken from Krasnopolsky (2007, 2012) to include a trap of constant lower \( K_{zz} \) across the extent of the cloud layer. The range of values that we explore for \( K_{zz} \) in the trap are 5000, 1000, 500, and 100 cm^2 s^{-1}, shown in Figure 2. Such a \( K_{zz} \) trap, if it exists in the Venus cloud layer, is particularly relevant to the water-rich hypothesis as enhanced water abundance below the clouds would result in lesser thermal heating flux at the cloud base due to \( H_2O \) IR absorption, which in turn increases the convective stability and decreases the eddy diffusivity, shown by Yamamoto (2014). We investigate the results from combining a \( K_{zz} \) trap and enhanced below-cloud water abundance.

5. Hypothesis: Another Source of Hydrogen in the Clouds

The \( SO_2 \) depletion can be explained if there is another source of hydrogen in the clouds. Here we will use NaOH as that source of hydrogen for the sake of convenience when calculating the cloud droplet chemistry. We are not claiming that this is the source of hydrogen. We will discuss possible sources of this excess hydrogen in Section 7.

Some \( SO_2 \) will dissolve into the cloud droplets directly, with a concentration in the droplet, \( c \) (mol/L), linearly proportional to the partial pressure of \( SO_2 \), with the Henry’s Law constant,
Table 4
Observational Constraints of Chemical Species in the Atmosphere of Venus

| Species | \( h_{\text{mix}} \) (km) | \( h_{\text{mix}} \) (km) | Mixing Ratio\(^a\) | Reference | Obs Type |
|---------|-----------------|-----------------|-----------------|-----------|---------|
| \( \text{SO}_2 \) | ... | ... | ppm | ... | ... |
| \( \text{H}_2\text{O} \) | 30.0 | 40.0 | 130. ± 50. | Marcq et al. (2008) | Venus Express |
| \( \text{H}_2\text{S} \) | 30 | 40 | 31 ± 2 | Marcq et al. (2008) | Venus Express |
| \( \text{CO} \) | ... | ... | ppm | ... | ... |
| \( \text{O}_3 \) | ... | ... | ppm | ... | ... |
| \( \text{OCS} \) | ... | ... | ppm | ... | ... |
| \( \text{H}_2\text{SO}_4 \) | 30 | 30 | 14.0 ± 6.0 | Pollack et al. (1993) | Ground |
| \( \text{HCl} \) | 50 | 52 | 3.0 ± 2.0 | Oschlisniok et al. (2012) | Venus Express |
| \( \text{PH}_3 \) | 15 | 25 | 0.41 ± 0.04 | Arney et al. (2014) | Ground |
| \( \text{S}_2 \) | ... | ... | ppm | ... | ... |
| \( \text{S}_4 \) | ... | ... | ppm | ... | ... |
| \( \text{H}_2\text{S} \) | 3.0 | 10.0 | 4. ± 3.9 | Krasnopolsky (2013) | Venera 11 |
| \( \text{H}_2 \) | 10.0 | 19.0 | 6. ± 2.0 | Krasnopolsky (2013) | Venera 11 |
| \( \text{H}_2\text{O} \) | 70.0 | 70.0 | <13.0 | Krasnopolsky (2008) | Ground |
| \( \text{H}_2\text{SO}_4 \) | ... | ... | ppm | ... | ... |
| \( \text{H}_2\text{S} \) | 0.0 | 25.0 | <10. | Donahue et al. (1997) | Pioneer Venus |
| \( \text{H}_2 \) | 22.0 | 64.0 | <10. | Oyama et al. (1980) | Pioneer Venus |
| \( \text{SO}_2 \) | 49.0 | 58.0 | 25. ± 10 | Mukhin et al. (1982) | Venera 13, 14 |

Notes.
\(^a\) Values and errors as found in the cited literature.
\(^b\) If interpreted as mesospheric SO\(_2\), see Section 7.4.
\(^c\) If interpreted as mesospheric PH\(_3\), see Section 7.4.
\(^d\) Based on their wet profile, which fits their data best.

References. Arney et al. (2014); Bell et al. (1991); Bertaux et al. (1996); Bézard et al. (1990); Bézard & de Bergh (2007); Bézard et al. (2011); Chamberlain et al. (2013); Collard et al. (1993); Connes et al. (1967); Connes et al. (1968); Cottini et al. (2012); Cotton et al. (2012); de Bergh et al. (1995); Donahue & Hodges (1992); Donahue et al. (1997); Encenan et al. (2012); Encenan et al. (2015); Encenan et al. (2020); Evans & Ingalls (1969); Fedorova et al. (2008); Fedorova et al. (2015); Fedorova et al. (2016); Fegley (2014); Gelman et al. (1979); Grassi et al. (2014); Greaves et al. (2020a); Greaves et al. (2020b); Hoffman et al. (1980); Iwagami et al. (2008); Krasnopolsky (2008); Krasnopolsky (2010a); Krasnopolsky (2010b); Krasnopolsky (2013); Krasnopolsky (2014); Maiorov et al. (2005); Marcq et al. (2005); Marcq et al. (2006); Marcq et al. (2008); Marcq et al. (2015); Marcq et al. (2018); Meadows & Crisp (1996); Moroz et al. (1979); Moroz et al. (1990); Mukhin et al. (1982); Mukhin et al. (1983); Na et al. (1990); Oschlisniok et al. (2012); Oyama et al. (1979); Oyama et al. (1980); Pollack et al. (1993); Sandor & Canky (2005); Sandor & Canky (2012); Surkov et al. (1982); Surkov et al. (1987); Tsang et al. (2008); Wilson et al. (1981); Young (1972); Zasova et al. (1993).

(This table is available in its entirety in machine-readable form.)

\[ H_{\text{SO}_2}(\text{mol/L bar}), \text{as the constant of proportionality:} \]
\[ e(\text{SO}_2) = H_{\text{SO}_2}p_{\text{SO}_2}. \]  
\[ (13) \]

Here \( p_{\text{SO}_2} \) is the partial pressure of \( \text{SO}_2 \), with the total gas pressure \( p \) (bar) and \( f_{\text{SO}_2} \) as the volume mixing ratio of \( \text{SO}_2 \), equal to \( n_{\text{SO}_2}/(\Sigma n_X) \).

For \( \text{SO}_2 \) dissolved in water, the Henry’s Law constant is \( \approx 10^{-2} \text{ mol/L bar} \) (Burkholder et al. 2020). For \( \text{SO}_2 \) dissolved in sulfuric acid, the constant increases with the sulfuric acid concentration between 0.1 and 1 mol/L bar) (Zhang et al. 1998). The SO\(_2\) then participates in the following reactions (here \( g \) is gas-phase, \( \ell \) is in the droplet). We first consider the dissolution of \( \text{SO}_2 \) and \( \text{H}_2\text{O} \) into the droplets by Henry’s law:

\[ \text{SO}_2(\text{g}) \rightleftharpoons \text{SO}_2(\ell), \]  
\[ \text{H}_2\text{O}(\text{g}) \rightleftharpoons \text{H}_2\text{O}(\ell). \]  
\[ (14) \]
\[ (15) \]

The rate constants for this reaction are balanced such that the concentration of \( \text{SO}_2 \) at any point agrees with Equation (13) when accounting for the droplet volume (see below). The rest
of the reactions are dissociation reactions for which bimolecular rate constants are set to $k_\text{f} = 5 \times 10^{10} \text{ mol}^{-1} \text{ L s}^{-1}$ in order to rapidly achieve equilibrium and avoid any dynamic effects, and the unimolecular rate constant is assigned a value that preserves the equilibrium set by the $pK_a$ or $pK_w$. The rate constant for the reverse reaction is (with units s$^{-1}$)

$$k_r = \frac{\rho k_f}{\mu} 10^{-pK_a},$$

(16)

$$k_r = \frac{\rho k_f}{\mu} 10^{-pK_w},$$

(17)

where $\rho$ (g cm$^{-3}$) is the density of the liquid and $\mu$ (g mol$^{-1}$) is the molar weight of the species. The reactions for the sulfates are

$$\text{H}_2\text{SO}_4(\ell) \rightleftharpoons \text{HSO}_4^-(\ell) + \text{H}^+(\ell), \quad pK_{a1} = -2.8;$$

(18)

$$\text{HSO}_4^-(\ell) \rightleftharpoons \text{H}_2\text{O}(\ell) + \text{H}^+(\ell), \quad pK_{a2} = 1.99;$$

(19)

and the reactions involving sulfuric acid and sulfites:

$$\text{SO}_2(\ell) + \text{H}_2\text{O}(\ell) \rightleftharpoons \text{SO}_3^2-(\ell) + \text{H}^+(\ell), \quad pK_{a1} = 1.857;$$

(20)

$$\text{H}_2\text{SO}_3(\ell) \rightleftharpoons \text{SO}_2^2-(\ell) + \text{H}^+(\ell), \quad pK_{a2} = 7.172.$$  

(21)

The rate constant for $\text{SO}_2$ to react with $\text{H}_2\text{O}$ is $2 \times 10^9 \text{ mol}^{-1} \text{ L s}^{-1}$ (Eigen et al. 1961; Brandt & Van Eldik 1995) and the rate constant for $\text{H}_2\text{SO}_3$ dissociation is $10^8 \text{ s}^{-1}$ (Eigen et al. 1961; Brandt & Van Eldik 1995). These reactions are sufficiently fast to draw the gas-droplet chemistry into equilibrium at all atmospheric heights. It was for this reason that we chose NaOH as our candidate salt. Finally, we consider the self-dissociation of water and the dissociation of sodium hydroxide:

$$\text{H}^+(\ell) + \text{OH}^- (\ell) \rightleftharpoons \text{H}_2\text{O}(\ell), \quad pK_a = 14$$

(22)

$$\text{NaOH}(\ell) = \text{Na}^+(\ell) + \text{OH}^- (\ell), \quad pK_w = 0.2.$$  

(23)

Sodium hydroxide is the example species we will use to buffer the clouds of Venus, freeing up more water in the droplet to react with $\text{SO}_3$, forming sulfurous acid, or with $\text{SO}_2$ to form sulfuric acid. The explanation for these reactions and details about rate constants and equilibrium constants is given in Section 5.

If the excess source of hydrogen is a salt, as is the case with NaOH, $\text{H}^+$ is replaced by $\text{Na}^+$. We calculate the pH where $pH = -\log_{10}(a_{\text{H}^+})$, where $a_{\text{H}^+}$ is the $\text{H}^+$ activity. The concentration of NaOH needed in the droplets to sequester $\text{SO}_2$ is determined by considering the cloud droplet volume as a function of height:

$$V_d(h) = \int \frac{4}{3} \pi r_d^3 \frac{\partial N_d}{\partial r_d} dr_d,$$

(26)

where $r_d$ (μm) is the droplet radius, and the function $r_d(\partial N_d/\partial r_d)$ is the droplet size distribution, which we take from Gao et al. (2014):

$$\frac{\partial N_d}{\partial r_d} = \frac{\partial n_d}{\partial r_d} V_{\text{atm}},$$

(27)

where $n_d$ (cm$^{-3}$) is the droplet number density (across all sizes) and $V_{\text{atm}} = 4\pi R_p^2 \Delta z$, with $R_p = 6052$ km as the radius of Venus and $\Delta z$ (km) is the model height step. We can calculate the amount of NaOH needed to deplete the $\text{SO}_2$ to the observed levels. We do this by dividing the total number of $\text{SO}_2$ molecules by the total droplet volume at height $z$:

$$c(z) = \frac{\rho_{\text{SO}_2}(z)}{N_A k T (z)} \int \frac{4}{3} \pi r_d^3 \left( r_d \frac{\partial n_d(z)}{\partial r_d} \right) dz,$$

(28)

where $N_A = 6.022 \times 10^{23}$ is Avogadro’s number and $k = 1.38065 \times 10^{-16}$ erg/K is Boltzmann’s constant. The above equation is used to prescribe the initial concentration of NaOH (in units of mol cm$^{-3}$) for our solver. The NaOH will rapidly dissociate and the OH$^-$ will then recombine with $\text{H}^+$ to form $\text{H}_2\text{O}$, and this, along with dissolved $\text{H}_2\text{O}$ from the gas phase, will react with $\text{SO}_2$ to form $\text{H}_2\text{SO}_3$. This will rapidly dissociate to form HSO$_3^-$ and H$^+$, and will buffer the solution, meaning that if the $\text{H}^+$ activity is perturbed, the reactions between the anions and $\text{H}^+$ will bring the activity of $\text{H}^+$ back to a given value determined by the $pK_a$. It should be emphasized that Equations (14) and (15) do not describe condensation of either of these species, which generally have partial pressures throughout Venus’s atmosphere that are well below the vapor pressure. Rather, these are the equilibrium concentrations in the droplets due to dissolution, balanced with the partial pressures in the gas.

Prescribing the NaOH this way introduces extra hydrogen into the model in a way not accounted for by mass balance or atmospheric redox balance. A full solution of the atmospheric chemistry coupled with surface chemistry would preserve this balance but that would require us to identify the source of hydrogen. Though we speculate on possible sources of hydrogen, from the surface (delivered via volcanism or winds) or exogenous, in Section 7, we do not know enough about Venus’s atmosphere, surface, or clouds to confidently identify a source and therefore do not self-consistently include this source in our model.

6. Results

The depletion of $\text{SO}_2$ can be explained by the $\text{SO}_2$ dissolving into the clouds, if the cloud pH is higher than previously believed. It can also be explained by varying the below-cloud $\text{SO}_2$ and $\text{H}_2\text{O}$. We discuss the consequences of varying the $\text{SO}_2$ in Section 6.1 and $\text{H}_2\text{O}$ in Section 6.2. The results of incorporating cloud chemistry are presented in Section 6.3.

6.1. Sulfur-poor Venus

$\text{SO}_2$ begins to significantly decrease through and above the clouds when surface $f_{\text{SO}_2} \sim 50$ ppm and achieves best results for below-cloud concentrations of 15 ppm. See Figure 3. The depletion occurs at 70–80 km unless the surface $\text{SO}_2$ is lower, around 15 ppm. Besides being inconsistent with most below-cloud observational constraints on $f_{\text{SO}_2}$, by almost an order of magnitude, the sulfur-poor model predictions agree with observations for all the species we consider reasonably well. The results agree with observations as closely as when we consider cloud droplet chemistry below with one possible exception of $\text{SO}_2$, which is below 1 ppb between 80 and 100 km. This may be brought into better agreement by including a mesospheric source of sulfur acid vapor or by adjusting the below-cloud concentrations of $\text{SO}_2$ because we have found that the above-cloud $\text{SO}_2$ is
very sensitive to the below-cloud SO2 and the Eddy diffusion profile when $f_{SO2} \approx 15$ ppm. The 15 ppm below-cloud SO2 model also predicts H$_2$ concentrations of $\sim 10$ ppm above 70 km.

6.2. Water-rich Venus

For the water-rich case, SO2 does not deplete for any value of below-cloud H$_2$O unless a $K_{zz}$ trap extending to 85 km in altitude is introduced in the eddy diffusion profile. Upon introduction of this trap the SO2 begins to significantly deplete at the top of the atmosphere, and this depletion height then lowers with increasing below-cloud water abundance, achieving best results around surface $f_{H2O} = 200$ ppm. See Figure 4. For in-cloud $K_{zz} = 5000$ cm$^2$ s$^{-1}$, the depletion height is higher than observations suggest (Encrenaz et al. 2019), dropping off at $\gtrsim 75$ km, and the O$_2$ concentration in the upper atmosphere exceeds 100 ppm. The depletion height can be lowered further to $\sim 70$ km by decreasing the $K_{zz}$ value in the trap; however, this exacerbates the overabundance of O$_2$ and causes CO to deviate from smooth monotonic growth with altitude. The CO profile has a downward spike at $\sim 70$ km, consistent with the profile of Pollack et al. (1993), as scaled by Marcq et al. (2005). This is likely due to the change of $K_{zz}$; see Section 7. Neither the SO2 depletion nor O$_2$ abundances agree with observations as well as in the sulfur-poor and cloud chemistry cases. The reason for this is the self-shielding of the excess water, which is not as effectively removed as SO2 within the clouds. The 200 ppm below-cloud H$_2$O model also predicts H$_2$ concentrations of $\sim 100$ ppm above 70 km, inconsistent with observations (see Appendix A).

6.3. Cloud Chemistry

In this scenario, described in Section 5, SO2 depletion through the clouds is accomplished by removing the SO2 in H$_2$SO$_3$ and H$_2$SO$_4$ via droplet chemistry. In our model, the droplet chemistry is driven specifically by sodium hydroxide (NaOH), and the NaOH itself buffers the cloud pH. The NaOH is a proxy used for convenience in modeling and represents other plausible sources of delivered hydrogen, discussed in Section 7. We adjust the amount of NaOH as a function of height to reproduce the observationally constrained SO2 profile (see Figure 5). This initial NaOH is prescribed for each height and is not solved for within the model. The function that reproduces the SO2 depletion, given the estimated droplet volume, was determined by solving the aqueous chemistry and cloud chemistry for different amounts of NaOH,
and the concentration of NaOH and the speciation of sulfuric and sulfurous acid as a function of height that results in the observed SO2 profile are shown in Figure 5. The prediction that the bulk of the clouds is HSO3 only holds if there is no mechanism for oxidizing the sulfur in the clouds. Sulfite aerosols are rapidly oxidized in Earth’s atmosphere (Townsend et al. 2012), and this may be the case for the atmosphere of Venus as well. We show the predicted mass loading in Figure 6. The mass of the lower cloud is largely in the form of sulfate, and this region overlaps with the larger mode 3 aerosols (Knollenberg & Hunten 1980). Profiles are similar for Ca(OH)2, assuming the kinetics are identical between Ca(OH)2 and NaOH. The similarity is due to the large $pK_a$, which results in all Ca(OH)2 becoming fully dissociated, along with the similar mass to basicity potential: the mass of a calcium atom is almost double that of a sodium atom, and each calcium carries two hydrides, which replace two protons.

We can then consider our solution of the droplet chemistry, set out in Equations (14)–(25), which predicts the H$^+$ activity, from which we can calculate the cloud droplet pH. As described in Section 5, the SO2 depletion is set by the capacity of the liquid to hold SO2, which is controlled by the amount of NaOH, and by the total volume of the liquid, which is determined by the cloud droplet size distribution. The predicted droplet pH is plotted as a function of height in Figure 5.

In this model, throughout the clouds the gas-phase SO2 is in equilibrium with the concentration of sulfur in the droplet that is specifically in the form of SO2; i.e., SO2(g) $\times$ SO2(ℓ) and adding or removing gas-phase SO2 results in proportionally changing the droplet SO2 and the balance of the other sulfur species. This equilibrium allows us to write out effective rate constants for SO2 and H2O dissolution into the droplets with rate constants tuned to reproduce the results from solving Equations (14)–(25). For SO2 the effective reaction is

$$\text{SO}_2(g) + \text{H}_2\text{SO}_4(\ell) \rightleftharpoons \text{SO}_2(\ell) + \text{H}_2\text{SO}_4(\ell).$$  \hspace{1cm} (29)$$

The rate constants for the forward reaction, $k_f$ (cm$^3$ s$^{-1}$), and reverse reaction $k_r$ (cm$^3$ s$^{-1}$) are

$$k_f = 10^{-32} \text{ cm}^3 \text{ s}^{-1} e^{9000K/T},$$ \hspace{1cm} (30)

$$k_r = 6.67 \times 10^{-36} \text{ cm}^3 \text{ s}^{-1} e^{9000K/T}.$$ \hspace{1cm} (31)
For H₂O, the effective reaction is

\[ \text{H}_2\text{O}(g) + 2\text{SO}_2(\ell) \rightarrow \text{H}_2\text{SO}_3(\ell) + \text{SO}_2(\ell). \]  

Neither of these reactions accounts for condensation. We determine the effective rate constants for the forward reaction, \( k_f \) (cm³ s⁻¹), and reverse reaction \( k_r \) (cm³ s⁻¹) to be

\[ k_f = 2.53 \times 10^{-36} \text{ cm}^3 \text{s}^{-1} e^{90000 K/T}, \]  
\[ k_r = 8.43 \times 10^{-38} \text{ cm}^3 \text{s}^{-1} e^{90000 K/T}. \]

Finally, an effective reaction needs to be included to encapsulate the release of SO₂ when the droplet rains out and evaporates:

\[ \text{SO}_2(\ell) + \text{H}_2\text{SO}_4(g) \rightarrow \text{SO}_2(g) + \text{H}_2\text{SO}_4(g), \]

with a rate constant of \( 2.2 \times 10^{-4} \text{ cm}^3 \text{s}^{-1} e^{-10000 K/T}. \)

The results of our model agree within an order of magnitude for all species considered and within a factor of 3 for all species except for OCS and the sulfur allotropes. Our model underestimates S₄ and does not predict the steep below-cloud gradient of OCS. In addition, it overpredicts O₃ in the upper atmosphere by a factor of 2–3. Comparison of this model and the best sulfur-poor model, with \( f(\text{SO}_2) = 20 \) ppm and Krasnopolsky’s Eddy diffusion profile (from Section 6.1) is shown in Figure 7.

7. Discussion

Below-cloud SO₂ < 50 ppm is inconsistent with most observations but not all. Vega 1 and 2 observed \( \geq 100 \) ppm concentrations of SO₂ directly below the clouds, though the error bars are large and 50 ppm abundances would be within 2σ of the measurements (Bertaux et al. 1996). Even lower SO₂ was measured within 20 km of the surface (Bertaux et al. 1996), which could indicate rapid surface depletion of SO₂ (Yung & Demore 1982).

The reported observations of below-cloud SO₂ at 100–200 ppm also have large uncertainties, typically on the order of 50 ppm (see, e.g., Marcq et al. 2008), so true values below 50 ppm would amount to discrepancies of 2–3σ. Exploring the hypothesis of low below-cloud SO₂ as an explanation for the above-cloud SO₂ depletion will require both more precise and more frequent observations of the below-cloud SO₂ to see whether it varies and by how much. Such data may only be obtainable with in situ measurements.

Below-cloud H₂O is better constrained, so it is less likely that there is an undetected large source of water vapor beneath the clouds. There are some variations in the measurements, from less than 10 ppm to 60 ppm, but with relatively small error bars. There is some observational support for a large reservoir of water within the clouds, with 700–20,000 ppm concentrations observed in the...
cloud layers by Vega 1, 2 (Surkov et al. 1987) and Venera 13, 14 (Mukhin et al. 1982; Surkov et al. 1982), and 100–200 ppm concentrations observed from the ground (Bell et al. 1991). The high quantities may have been due to incidental sampling of cloud droplets, which are expected to be composed of ∼15–25 wt% water, even without considering the cloud pH buffer hypothesis proposed here. Even if these constraints were not so tight, the model where we increase the below-cloud water vapor predicts that the clouds to extend up to ∼80 km, and the depletion would be more gradual and higher in the atmosphere. Further to this, the water-rich model overproduces O2 above ∼70 km, pushing it beyond the observed upper limit by two orders of magnitude, overproduces H2, and does not allow for SO2 to return to the ∼1 ppb level at 90 km, problems that are not shared by the other best-fitting models.

As we have shown, cloud chemistry is a possible explanation for the depletion of SO2. Aerosols could provide the excess hydrogen capable of depleting gas-phase SO2. This hydrogen could be bound in salts or could be in some other form, such as hydrocarbons. It is important that whatever the source of hydrogen, it is replenished to sustain the SO2 gradient. Otherwise the hydrogen will be consumed, the clouds will be saturated with sulfur, and the gradient of SO2 will disappear. It is also worth mentioning that three phases can participate in this chemistry: the gas phase, the liquid of the droplet (at atmospheric heights where the droplet has not frozen), and the solid aerosol material, either at the core, surrounding the droplet, or suspended within the droplet. This provides a rich and complex multiphase chemistry worth exploring in the lab.

What follows in this section is a discussion of the requirements for and implications of cloud chemistry. In Section 7.1, we discuss possible sources of hydrogen within the clouds. The measured optical constants and spectral features of the clouds of Venus are consistent with cloud droplets composed of a large percentage of sulfuric acid. Any proposed cloud chemistry must either preserve sulfuric acid as the dominant species in the clouds or must propose a species with similar optical properties and spectral features. We discuss the observable implications of our cloud chemistry in Section 7.2. The dissolution of hydrogen halides into the clouds is discussed in Section 7.3. The cloud chemistry also has implications for above-cloud radical concentrations, which affect the lifetime of hypothetical PH3 within the clouds. We discuss the status of PH3 and its lifetime within the clouds in Section 7.4. Finally, in Section 7.5, we briefly discuss the
implications of different cloud chemistry for hypothetical life within the cloud droplets of Venus.

7.1. Possible Sources of Hydrogen in the Clouds

If a buffer explains the sulfur depletion, it is possibly a salt. Salts will dissociate quickly, and some will provide efficient buffers. The salts must get into the clouds in order to buffer them. This can be accomplished either by transport from the surface or exogenous delivery.

**Exogenous Delivery**: Exogenous delivery, meaning delivery of material from the interplanetary medium, is unlikely to provide significant material to buffer the clouds of Venus based on the estimated incoming flux of interplanetary dust. The clouds must be able to retain virtually all of the SO₂ over the timescale of transport through the clouds, requiring a flux of salts, \( \Phi_s \) (mol cm\(^{-2}\) s\(^{-1}\)), of

\[
\Phi_s = \frac{n_{SO_2} H_0}{\tau_{dyn}} = \frac{f_{SO_2} n k T K_{\infty}}{\mu_m m_p N_A g (\Delta h)^2},
\]

(36)
where \( n_{SO_2} \) (cm\(^{-3}\)) is the number density of SO\(_2\) at the height where SO\(_2\) begins to deplete (50 km), \( H_0 \) (km) is the scale height, \( R = 6052 \) km is the radius of Venus, \( \tau_{syn}(s) \) is the dynamic timescale of the atmosphere at 40 km, \( \rho_{SO_2} = 150 \) ppm is the volume mixing ratio of SO\(_2\), \( n = \Sigma n_{i} x_{i} = 2.189 \times 10^{19} \) cm\(^{-3}\) is the gas density at 50 km, the height where the depletion begins, \( k = 1.38065 \times 10^{-16} \) erg/K is Boltzmann’s constant, \( T = 349.7 \) K is the temperature at 40 km, and \( K_N \approx 100 \) cm\(^2\) s\(^{-1}\) is the Eddy diffusion of the droplets within the cloud, a low estimate more favorable for exogenous delivery. \( \mu_{av} = 44 \) is the mean molecular weight of the atmosphere, \( m_p = 1.6726 \times 10^{-24} \) g is the mass of a proton, \( N_A = 6.022 \times 10^{23} \) is Avogadro’s number, \( g = 8.87 \) m s\(^{-2}\) is the surface gravity of Venus, and \( \Delta h = 20 \) km is the thickness of the cloud layer. Applying all these estimates yields

\[
\Phi_s \approx 10^{-13} \text{ mol cm}^{-2} \text{s}^{-1}. \tag{37}
\]

Continuous exogenous delivery of material is insufficient to match this flux even assuming that 100% of the material is in the form of hydrated minerals that will deplete SO\(_2\). If we assume that delivery of exogenous material to Venus is comparable to delivery to Earth, 20–70 kt/yr (Peucker-Ehrenbrink 1996; Greaves et al. 2020a), which translates to \( \sim 10^{-17} \) mol cm\(^{-2}\) s\(^{-1}\), or four orders of magnitude too little to account for the missing hydrogen.

One other possibility is stochastic delivery. If a recent airburst, a large impact that breaks up into the atmosphere, occurred in Venus’s atmosphere, the metals released could permeate the clouds, providing a transient in-cloud source of hydrogen. If this is the case, then the depletion of SO\(_2\) will be temporary, lasting as long as the below-cloud store of these elements persists, on the order of the diffusion timescale or ~1000 years.

**Dust from the Surface:** Transport from the surface also struggles to meet the required flux of hydrated material. Although calculations of dust transport favor a more dusty atmosphere for Venus than Earth (Sagan 1975), Venera measurements suggest that the lower atmosphere is clear, placing upper limits on dust transport to the clouds. It is possible that there is heterogeneous low atmospheric weather, with dust rapidly transported to tens of kilometers above the surface, and that Venera happened to land in a region where the vertical diffusion and winds were insufficient to change the atmospheric opacity. A low-level haze inferred from Venera probe data (Grieger et al. 2004), and consistent with heavy metal frost at higher elevations on Venus’s surface (Schaefer & Fegley 2004), may itself be the suspended dust (Titov et al. 2018), and winds may cause that dust to periodically move into the clouds. The reaction products of the salt with SO\(_2\) in and below the clouds could result in transient high abundances of water, which may then be removed by the leftover oxides. This could reconcile the transient observations of >100 ppm of water vapor in and below the clouds (Mukhin et al. 1982; Surkov et al. 1982; Bell et al. 1991), with the ~25 ppm abundances near the surface (Bertaux et al. 1996).

Explaining the sulfur depletion, with dust containing 5 wt.% salt, requires a dust flux to the clouds of ~16 Gt/year, well within the estimates of surface dust fluxes estimated from analog experiments (Greeley et al. 1984). The composition of the dust and the form of the salts is unknown. Here we will speculate on some possible candidates. Our speculation is based on the chemical and physical stability of the salts:

1. **NaOH:** Sodium hydroxide is the example salt we use for our calculations, but it is an unrealistic candidate salt. It is sufficiently stable to heat, persisting as a liquid up to 1661 K (Haynes 2014). However, it is known to react rapidly with SO\(_2\) to form sodium sulfite and water vapor. Given the concentrations of below-cloud SO\(_2\), NaOH cannot plausibly survive to reach the clouds unless it is injected rapidly, e.g., via a volcanic plume.

2. **Ca(OH)\(_2\):** Calcium hydroxide is stable as a solid at the surface of Venus. Strictly speaking, calcium hydroxide has no melting point. Instead, at 93 bar and ~1000 K, it is expected to decompose into CaO and H\(_2\)O, based on extrapolations of the vapor pressure curve of Halstead & Moore (1957). Ca(OH)\(_2\) will undergo carbonation, and there is ample CO\(_2\), but this reaction is very slow, and Ca(OH)\(_2\) is kinetically stable at temperatures above 723 K (Materic & Smedley 2011). Ca(OH)\(_2\) will also react rapidly with SO\(_2\) but only in the presence of water vapor at concentrations of >3000 ppm (Liu et al. 2010). The kinetic stability versus the dynamic timescale for Ca(OH)\(_2\) aerosols is unknown, but presently Ca(OH)\(_2\) cannot be ruled out as a candidate.

3. **Mg(OH)\(_2\) and Fe(OH)\(_2\):** Neither magnesium hydroxide nor iron hydroxide (either the Fe(II) hydroxide or Fe(III) hydroxides) are stable at Venus’s surface temperature and pressure (Wang et al. 1998; Haynes 2014).

4. **Other Hydroxides:** It may be that more exotic hydroxides, such as Al(OH)\(_3\), could deliver hydrogen to Venus’s cloud layer. The requirements are sufficient concentrations to satisfy the required fluxes and the thermochemical and kinetic stability of the salt in the presence of major atmospheric constituents.

5. **Oxides:** Oxides, either resident surface oxides or oxides produced by the dehydration of hydroxides, may participate in cloud chemistry in unknown ways, sequestering SO\(_2\) directly; for example, [Mg, Fe]O + SO\(_2\) may be converted into [Mg, Fe]SO\(_4\) directly. The subsequent dissociation in sulfuric acid will buffer the cloud droplet pH. There is some indirect evidence of the presence of oxides from the near-surface haze because these oxides can react with hydrochloric acid vapor to form FeCl\(_3\), which has been observed in the clouds and is a candidate for the mysterious UV absorber (Krasnopolsky 2017).

6. **Sulfates:** It is possible hydrogen-bearing sulfates could find their way into the clouds, but no known hydrogen-bearing sulfate is thermally stable at Venus’s surface pressure and temperature. It is possible that they are produced from gas-phase reactions, e.g., the possible production of ammonium sulfate from reaction with NH\(_3\) and SO\(_2\) (Titov 1983). Sulfates that do not contain hydrogen are plausible aerosols. Indeed, we predict they are produced via cloud chemistry. But these aerosols will not deliver hydrogen and cannot directly participate in the depletion of gas-phase SO\(_2\).

The above list is not intended to be exhaustive and does not consider whether these salts are expected at the temperatures and pressures at the surface of Venus, where we would generally expect chemistry to tend toward thermodynamic equilibrium. The results of comparing our surface boundary conditions (Section 3.1) to chemical equilibrium (Section 3.2) predict that no phyllosilicates or hydrated minerals are present at the surface of Venus if the surface and gas are in equilibrium.
The model presented in Section 3.2 can be used to explore the sensitivity of the mineral composition to surface temperature. Figure 8 shows the results of our GGCHEM model at the same constant pressure and total element abundances when varying the surface temperature between 450 K and 850 K. Venus is just about 15 K too warm to have pyrite (FeS$_2$[s]) as a stable condensate on the surface according to this model. For surface temperatures lower than about 720 K, the formation of FeS$_2$[s] would start to remove the SO$_2$ from the atmosphere according to the following complex net reaction:

$$\frac{15}{11} \text{SO}_2 + \frac{1}{11} \text{Fe}_2\text{O}_3[s] + \text{CaAl}_2\text{Si}_2\text{O}_8[s]$$

$$\rightarrow \frac{2}{11} \text{FeS}_2[s] + \text{CaSO}_4[s] + \text{Al}_2\text{O}_3[s] + 2 \text{SiO}_2[s],$$

(38)

which is a variant of Equation (12), where the oxygen on the left side is provided by Fe$_2$O$_3$[s]. At temperatures below about 580 K in this model, the first carbonate magnesite (MgCO$_3$[s]) becomes stable, which could initiate a dramatic change of the atmosphere as the main atmospheric molecule CO$_2$ could deposit at the surface to form MgCO$_3$[s]:

$$\text{CO}_2 + \text{MgSiO}_3[s] \rightarrow \text{MgCO}_3[s] + \text{SiO}_2[s],$$

(39)

leaving an atmosphere that is dominated by N$_2$ with more H$_2$O. Finally, below a temperature of about 520 K, the first phyllosilicate, talc (Mg$_3$Si$_4$O$_{10}$H$_2$[s]), becomes stable, which could partly remove the water from the atmosphere. Only for temperatures below 520 K does our model predict the presence of hydrogen-containing minerals.

Whether or not this means that the gas at the bottom of the Venusian atmosphere is in fact in chemical equilibrium and whether the element abundances in the gas are regulated by outgassing/deposition via the contact with the hot rock at the surface are yet unsolved questions. Disequilibrium processes might supply phyllosilicates or hydrated salts. For example, geological processes such as volcanism may resurface Venus’s crust with hydrated components (see also discussion below).

Our phase-equilibrium model suggests that hydrogenated rock and salts are not stable on the surface of Venus and will sublimate or react with the atmospheric gases to form other chemicals. However, the salts may be stable enough to be swept up to greater altitudes and cooler temperatures, before they react away and equilibrium is restored.

More research in the lab and in situ observations of the clouds of Venus will be needed to determine if salts are present and, if so, what their chemistry is.

**Volcanic Delivery:** A variant of the dust delivery mechanism to achieve a hydrogen flux to the clouds is volcanism. The residence time of active volcanism on Venus has long been speculated upon, motivated both by the transient SO$_2$ detected at 40 mbar by Pioneer Venus (Table 4; Esposito 1984) and the recognition that SO$_2$ may react with surface minerals and require continual replenishment (Fegley & Prinn 1989).

Volcanism could deliver material to Venus’s clouds in three ways: 1) as solid material deposited at the surface, which is subsequently lofted by winds; 2) as an explosive eruption introducing material into the below-cloud atmospheric, where vertical mixing slowly raises it to the cloud layer; and 3) as a large explosive eruption injecting material directly into the cloud layer.

Scenario (1) is effectively the “dust from the surface” mechanism described above, albeit by explicitly considering volcanism a source of juvenile OH-bearing material is introduced, which could help overcome the short surface lifetime of some OH-bearing phases. The second and third possibilities take advantage of the dynamics of volcanism to shorten the distance, and thereby potentially increase the flux, of mineral sources of OH to the clouds by reducing the time the material spends at high temperature near Venus’s surface. Volcanism may also help loft material above the sluggish surface winds to an altitude where winds more readily carry dust higher (e.g., Linkin et al. 1986; Zasova et al. 2007; Peralta et al. 2017).

Modeling work by Glaze (1999) and Airey et al. (2015) suggests that it is possible for volcanic eruption columns on Venus to reach the cloud base; however, it requires particular circumstances that may or may not be frequently met: in particular, a high elevation...
vent and a magma containing several wt% water. Whether such water-rich magmas exist on Venus is unknown; taking Earth as an analog, magmas with >3 wt% water occur only where subduction is introducing surface water back into the mantle, a tectonic mode that cannot have prevailed on Venus for hundreds of millions of years, if ever. Voluminous water-rich explosive volcanism is also problematic given the tight constraints on the below-cloud H$_2$O mixing ratio of <0.6 ppm. At best, therefore, these constraints would imply a highly stochastic delivery of volcanic material directly into the clouds.

The best case for a volcanic contribution to mineral buffering of Venus’s cloud chemistry is therefore by enhancing background dust levels in the below-cloud atmosphere. Fegley & Prinn (1989) estimate that a volcanic flux of ~1 km$^3$ yr$^{-1}$ is required to sustain atmospheric SO$_2$ at the levels observed. This translates to a mass flux of ~3 $\times$ $10^{12}$ kg yr$^{-1}$ of magma, or ~6 $\times$ $10^{11}$ mol yr$^{-1}$ of hydrated phases assuming terrestrial levels of water in Venus’s magmas. As an upper bound, this volcanic flux could provide a potential 4 $\times$ $10^{12}$ mol cm$^{-2}$ s$^{-1}$ of hydrated phases to Venus’s clouds if the entire mass were mobilized as dust in the atmosphere. Being below the ~$10^{-13}$ mol cm$^{-2}$ s$^{-1}$ of salt delivery estimated above, rates of volcanism either need to be (at least) an order of magnitude larger than assumed here, or the magmas correspondingly more volatile rich, for volcanism to be contributing to chemical buffering of Venus’s clouds by water delivery. We note that although seeming unlikely, given all the current uncertainties on the composition and dynamics of Venus’s interior, this possibility cannot be entirely ruled out.

7.2. Reconciling the Cloud Chemistry with Cloud Observations

There is considerable evidence that the clouds of Venus are mostly sulfuric acid or something very like sulfuric acid. The classic paper by Young (1973) identifies most of the lines of evidence. The refractive index of the clouds obtained from infrared polarimetry, constrained to within 1.425 and 1.455 at the time, is best explained by droplets of ~75% sulfuric acid. In addition, the bottom of the cloud layer matches the condensation temperature of sulfuric acid, and specific spectral features between 8–13 μm are very similar to spectra of condensed sulfuric acid. Sulfuric acid is also expected at concentrations of ~75% based on models of H$_2$SO$_4$ and H$_2$O condensation (Krasnopolsky 2015). Subsequent studies have further refined the estimated concentrations of sulfuric acid in the clouds. Barstow et al. (2012) perform a retrieval on VIRTIS data of the atmosphere from Venus Express and find that the 2.2:1.74 μm radiance ratio is sensitive only to the imaginary index of refraction, and therefore the sulfuric acid concentrations, and that most of the retrieved sulfuric acid concentrations in the lower clouds are between 85–96 wt%, or between 16–18 mol L$^{-1}$. Arney et al. (2014) used the same 2.2:1.74 μm radiance ratio and found that the sulfuric acid concentrations vary by time and latitude between 73–87 wt% in the upper clouds, or between 14–16 mol L$^{-1}$ sulfuric acid concentration, lower than Barstow et al. (2012), suggesting that the concentration of sulfuric acid changes as a function of height. Titov et al. (2018) provide a comprehensive review of the research into Venus’s clouds. None of the UV absorption due to our predicted profiles of gas-phase species, for the cloud model or any other model, explains the mysterious UV absorber, at λ > 200 nm. However, we do not consider the absorption properties of the aerosol particles themselves. It would be useful to determine the UV optical constants of these aerosols, or of sulfuric acid droplets themselves, to see whether a feasible candidate for the mysterious UV absorber is already in our midst.

Recent retrievals rely on infrared bands that constrain the index of refraction. If our proposed cloud chemistry is accurate and if all droplets have the same chemistry, hydrated sulfites and sulfates compose a large fraction of Venus’s clouds. Sulfate achieves an index of refraction of >1.44 at <20 wt% H$_2$O (Cotterell et al. 2017), sufficient to explain the observed index of refraction. In addition, the S–H and S–O bonds are all similar, and so similar spectral features are expected. We predict a significant fraction of the clouds is made up of sulfites, in line with terrestrial SO$_2$ + H$_2$O chemistry (Terraglio & Manganelli 1967). The refractive index of sulfites is not as well known, though it has been measured for cyclic sulfites to be ~1.5 (Pritchard & Lauterbur 1961), which is consistent with Venus cloud observations, particularly those that favor values higher than can be achieved by pure sulfuric acid (Markiewicz et al. 2018; Petrova 2018) but may be achievable by sulfites and sulfates.

One other possibility is that the droplets of Venus do not all have the same chemistry. The salt content of the aerosols could be heterogeneous, either because they derive from different surface, volcanic, or delivered materials, or because some cloud nuclei are produced photochemically, as sulfur allotropes and sulfuric acid. The observed bimodal distribution of cloud droplet sizes (Wilquet et al. 2009, 2012) is consistent with this hypothesis. It may be that the smaller, more numerous cloud droplets lack salts, while less numerous droplets have salts. The salts will afford those droplets a far greater capacity for SO$_2$, and this may explain their larger size. If this is the case, we would predict the small mode droplets have a pH < 0 and the large mode droplets a pH > 1.

7.3. The Effect of Cloud Chemistry on Gas-phase Halogens

Hydrogen halides have been observed on Venus, namely HCl and HF (see Appendix A). Attempts have been made to observe HBr, but thus far only upper limits have been established (Krasnopolsky & Belyaev 2016). These hydrogen halides are likely to dissolve into the droplets, especially if the pH is higher than previously thought, and this may affect both the droplet chemistry and the atmospheric profiles of these halides. We investigate this possibility, considering only HCl, the sole hydrogen halide included in our model.

First, we have to determine Henry’s Law constant for HCl dissolved into water/sulfuric acid mixtures. Williams & Golden (1993) find that the effective Henry’s Law constant of HCl in a solution of 50 wt% H$_2$SO$_4$ is

\[ H^* \approx 10^5 \text{ M/bar} \]  

at room temperature (which we will use as a proxy for the cloud layer). We use the relation (where $H^*$ (mol/(L bar)) is the effective Henry’s law constant and $K_a$ is the acid dissociation constant):

\[ H^* = H \left( 1 + \frac{K_a}{a_{H^+}} \right) \]

(41)

to solve for the hydrogen activity, $a_{H^+}$. The concentration of 50 wt% H$_2$SO$_4$ is approximately 2 mol L$^{-1}$, which will completely dissociate, and result in a pH of roughly −0.5, or $a_{H^+} \approx 2$. For
HCl, \( K_a \approx 2 \times 10^6 \). This means that \( H \approx 0.1 \) for HCl in sulfuric acid.

We can now predict the depletion of HCl for the sulfuric acid case, where \( \text{pH} \sim -3 \). The equation for the capacity, \( \kappa \), of the cloud droplets for a particular species, assuming that the dissociation products are not removed and so a simple equilibrium is achieved, is:

\[
\kappa = \frac{H^+(\text{HCl})N_a}{n(\text{HCl})} \int_{4\pi r^2} \left( r^2 \frac{dn_n}{dr} \right) \text{d}r
\]

(42)

\[
H \left( 1 + \frac{K_a}{a^-} \right) kTN\int_{4\pi r^2} \left( r^2 \frac{dn_n}{dr} \right) \text{d}r
\]

(43)

Though it is true that if there is any Na\(^+\) or other cation, it will form a salt with Cl\(^-\), this will act in solution as though entirely dissociated, and so effectively the amount of Cl\(^-\) and H\(^+\) due to HCl remains unchanged. If \( \kappa < 1 \), then the capacity of the droplets is not great enough to appreciably affect the composition of the gas-phase species. In the pure sulfuric acid case, \( \kappa \approx 1.6 \times 10^{-5} \), and so very little HCl will be in the droplets, compared to the amount in the gas phase.

At higher pH, the situation changes, but still, at \( \text{pH} \sim 1 \), \( \kappa \approx 0.16 \), and so there should not be significant depletion. The value of \( \kappa \) in Equation (43) can be set to 1 and \( a^- \) solved for to predict the pH at which HCl should be significantly depleted, \( \text{pH} \approx 1 \). Though the profiles of gas-phase hydrogen halides should not be affected at lower pH’s, the concentrations of halogens in the droplets could be significant, and it would be informative to measure these concentrations in situ.

7.4. The Presence and Lifetime of Phosphine

A broad time-variable feature has been observed at 267 GHz by both ALMA and JCMT (Greaves et al. 2020a, 2020b). The existence and source of the feature has been disputed (Snellen et al. 2020; Thompson 2021), though see the reply by Greaves et al. (2020b). This feature has been attributed to phosphine (Greaves et al. 2020a, 2020b), consistent with possible in situ detection of phosphine discovered during reanalysis of Venus Pioneer data (Mogul et al. 2021). The updated ALMA data indicate that the 267 GHz feature is now consistent either with 1.5–7 ppb phosphine (PH\(_3\)) or \( \sim 50 \) ppb mesospheric SO\(_2\) (Greaves et al. 2020b; Lincowski et al. 2021), though this amount of SO\(_2\) is not consistent with the nondetection of the 267.5 GHz SO\(_2\) feature, from the same ALMA data, providing a 10 ppb upper limit (Greaves et al. 2020b). Careful modeling of different PH\(_3\) profiles by Lincowski et al. (2021) demonstrates that the proposed in-cloud profile for PH\(_3\) from Greaves et al. (2020a) is insufficient to explain the observed feature. This leaves open the possibility that the signal is due to \( \sim 50 \) ppb mesospheric SO\(_2\), with an anomalous velocity shift, or mesospheric PH\(_3\) in quantities difficult to reconcile with the PH\(_3\) lifetime, unless there is an unknown mesospheric source of PH\(_3\). Further observations will be needed to determine which, if either PH\(_3\) or SO\(_2\), or if some other unidentified molecule, is the source of the 267 GHz feature.

Greaves et al. (2020a) and Bains et al. (2020) estimated the lifetime and required flux of 20 ppb PH\(_3\). The amount inferred from the ALMA observations has decreased to 1 ppb (Greaves et al. 2020b), and we can apply our new model results to estimate that a flux of \( 10^7 \) cm\(^{-2}\) s\(^{-1}\) PH\(_3\) is required to explain PH\(_3\) at these abundances, assuming the 267 GHz feature probed the clouds (this also includes effective dry deposition of PH\(_3\) set to fix the abundance of PH\(_3\) at the surface equal to zero, and dry deposition of PH\(_3\) fixed to \( 10^{-3} \) cm s\(^{-1}\)). We also find that PH\(_3\) is efficiently destroyed above 60 km, and the steep gradient in its profile is consistent with the nondetection above 61 km (Encrenaz et al. 2020). The in-cloud PH\(_3\) profile cannot explain the observed feature (Lincowski et al. 2021), but this profile is expected if there were a source of PH\(_3\) in the clouds.

7.5. Implications for Hypothetical Life in the Clouds of Venus

The implications of this cloud chemistry on hypothetical Venusian life depend on how and in what form the sulfur is depleted. If surface, volcanic, or exogenous delivery is responsible, this would explain the provision of various alkaline salts and/or other metals, essential for life as we know it. The higher pH is within the range where known acidophiles can thrive (Messerli et al. 2005). However, although acidophilic and halophilic extremophiles exist on Earth, there are, as far as we know, no known extremophiles that are both acidophilic and halophilic enough to thrive in these conditions (Belilla et al. 2019).

If, on the other hand, the pH is being buffered within the clouds by the organisms themselves, either by use of phosphate or by burning hydrocarbons, sacrificing themselves so that others may live, then the available water will be produced and scavenged by means of the reaction. The remaining question is whether the biomass is sufficient to explain the sulfur depletion, and there is no reliable estimate of the biomass in the clouds of Venus. Early estimates based on tentative detections of phosphine have been made (Lingam & Loeb 2020), but there is significant work left to better constrain the possibility that life is making use of the in-cloud SO\(_2\).
a nebulizer is incorporated onto a mass spectrometer to separate gas and cloud particles and analyze the aerosol chemistry directly (Baines et al. 2021). This would be the most straightforward way to test this hypothesis, and other hypotheses, e.g., Krasnopolsky (2017), that involve cloud chemistry.

Probes to the surface will be relevant for constraining the surface mineralogy and determining whether Venus’ surface composition is in chemical equilibrium. These observations can be combined with a climate history of Venus, based on observations and models, to discover more about Venus’ atmospheric evolution. Specifically, Figure 8 can also be used to speculate on the possible cause for the origin of the thick Venusian atmosphere that we observe today. If Venus once was a cooler planet with a thinner N2-dominated atmosphere, just like Earth, but for some reason it warmed up to temperatures above 580 K, possibly due to large amounts of CO2 released during global resurfacing (Strom et al. 1994), all carbonates in the surface rock would decompose and liberate even more CO2 into the atmosphere. Not only would this increase the greenhouse effect, but it would also make the Venusian atmosphere thicker. Both effects would have increased the surface temperature, leading to a runaway buildup of the thick CO2 Venusian atmosphere that we find today. Observations of surface minerals, possible with VERITAS (Smrekar et al. 2020) and EnVision (de Oliveira et al. 2018), would allow us to test the predictions of this and other hypotheses for the present state of Venus’ atmosphere and climate, in particular, whether the surface and atmosphere of Venus are at thermochemical equilibrium.

To prepare for these missions, experiments are needed in Venus-analog environments to predict the chemistry that takes place, especially the largely unexplored chemistry that may take place within high concentrations of sulfuric acid and the surface chemistry that may take place on efflorescent sulfates. In the meantime, a detailed cloud model of the form published by Gao et al. (2014), but that incorporates this chemistry, would be of value in order to see if any remote predictions would distinguish between sulfuric acid/water droplets and sulfate/sulfite/water droplets. It may be possible to falsify the cloud chemistry hypothesis based on a combination of cloud formation and radiative transfer models and observations. It is unlikely the puzzles addressed in our paper are likely to be resolved without returning to the clouds of Venus. If other missions to other planets are any indication, what we find will be entirely unexpected.

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Software: ARGO (Rimmer & Helling 2016), GGCHEM (Woitke et al. 2018).

Appendix A
Observational Constraints on the Atmospheric Composition of Venus

The species we included in our network robustly and that are also observationally constrained in Venus’s atmosphere are carbon dioxide (CO2), molecular nitrogen (N2), sulfur dioxide (SO2), water vapor (H2O), carbon monoxide (CO), molecular oxygen (O2), carbonyl sulfide (OCS), sulfuric acid vapor (H2SO4), hydrogen chloride (HCl), sulfur monoxide (SO), trisulfur (S3), tetrasulfur (S4), hydrogen sulfide (H2S), and molecular hydrogen (H2). Phosphine (PH3) is also included in our network but not in a robust way. PH3 may have been observed in the atmosphere of Venus (see Section 7.4 for details). There are also many remote observations that constrain cloud properties such as average particle size and indirectly infer that the clouds are made of droplets of high-concentration sulfuric acid (See Section 7.2). To this date no definitive in situ measurements of the cloud droplet chemistry have been made, and so virtually nothing is directly known about the cloud droplet chemistry. Observations have been made by a variety of instruments on the ground, by orbital probes, and by in situ probes. We compiled these data ourselves from a variety of sources, and a more complete compilation has been made by Johnson & de Oliveira (2019), which includes several reactive species not incorporated into our network, such as HF, as well as unreactive species. Our compilation is given in Table 4.

Appendix B
The Chemical Network: Stand2020

We constructed our network starting with Rimmer & Rugheimer (2019), adjusting several reactions relevant for Earth’s atmosphere to improve agreement between our model predictions and observations of formaldehyde and HCN in modern Earth’s atmosphere. We then include the sulfur network developed by Hobbs et al. (2020) and supplement it with reactions from Krasnopolsky (2007) and Zhang et al. (2012). We made extensive use of the NIST database (Manion et al. 2015), the KIDA database (Wakelam et al. 2012), the MPI-Mainz UV/VIS database (Keller-Rudek et al. 2013), and PhiDRates (Huebner & Mukherjee 2015), keeping with the philosophy, quoted from Rimmer & Helling (2016):

1. If there exists only one published rate constant for a given reaction, we use that value.
2. Reject all rate constants that become unrealistically large at extreme temperature.
3. Choose rate constants that agree with each other over the range of validity.
4. If the most recent published rate constant disagrees with (3), and the authors give convincing arguments for why the previous rates were mistaken, we use the most recently published rate.

The network is given in Tables 5, 6, and 7.
### Table 5
Reactions Added and Updated for STAND2020

| Reaction | $\alpha$ | Units | $\beta$ | $E_r$ (K) | Reference |
|----------|----------|-------|---------|----------|-----------|
| $C + C + M \rightarrow C_2 + M$ | $5.46 \times 10^{-31}$ | cm$^3$ s$^{-1}$ | $-1.6$ | $0.0$ | Slack (1976) |
| $C + C + M \rightarrow C_2 + e^-$ | $2.0 \times 10^{-11}$ | cm$^3$ s$^{-1}$ | $0.0$ | $0.0$ | Martinotti et al. (1968) |
| $C + O + M \rightarrow CO + M$ | $2.00 \times 10^{-34}$ | cm$^3$ s$^{-1}$ | $0.0$ | $0.0$ | Fairbairn (1969) |
| $C + O + M \rightarrow CO + M$ | $4.82 \times 10^{-15}$ | cm$^3$ s$^{-1}$ | $-1.0$ | $0.0$ | Est. |
| $H + N + M \rightarrow HN + M$ | $5.02 \times 10^{-32}$ | cm$^3$ s$^{-1}$ | $0.0$ | $0.0$ | Brown (1973) |
| $H + N + M \rightarrow HN + M$ | $1.21 \times 10^{-12}$ | cm$^3$ s$^{-1}$ | $-1.0$ | $0.0$ | Est. |
| $N + N + M \rightarrow N_2 + M$ | $1.25 \times 10^{-32}$ | cm$^3$ s$^{-1}$ | $0.0$ | $0.0$ | Knipovich et al. (1988) |
| $N + N + M \rightarrow N_2 + M$ | $5.00 \times 10^{-16}$ | cm$^3$ s$^{-1}$ | $0.0$ | $0.0$ | Takahashi & Miyazaki (1977) |
| $N + O + M \rightarrow NO + M$ | $3.26 \times 10^{-33}$ | cm$^3$ s$^{-1}$ | $0.0$ | $0.0$ | Campbell & Gray (1973) |
| $N + O + M \rightarrow NO + M$ | $7.87 \times 10^{-14}$ | cm$^3$ s$^{-1}$ | $-1.0$ | $0.0$ | Est. |
| $O + O + M \rightarrow O_2 + M$ | $1.67 \times 10^{-33}$ | cm$^3$ s$^{-1}$ | $-1.0$ | $0.0$ | Javoy et al. (2003) |
| $O + O + M \rightarrow O_2 + M$ | $1.21 \times 10^{-11}$ | cm$^3$ s$^{-1}$ | $-2.0$ | $0.0$ | Est. |
| $Cl^+ + e^- \rightarrow Cl + h\nu$ | $1.13 \times 10^{-10}$ | cm$^3$ s$^{-1}$ | $-0.7$ | $0.0$ | Barth et al. (2021)$^b$ |
| $C_2^+ + e^- \rightarrow C_2 + C$ | $3.00 \times 10^{-7}$ | cm$^3$ s$^{-1}$ | $-0.5$ | $0.0$ | Barth et al. (2021)$^b$ |

**Notes.**

- $^a$ This reaction is not reversed.
- $^b$ The rate constant for this reaction was miswritten in Krasnopolsky (2007); Zhang et al. (2012).

**References.**

Amano et al. (1983); Andersson et al. (2003); Anglada et al. (2017); Arthur & Cooper (1997); Ashmore & Burnett (1962); Atkinson et al. (1989); Atkinson et al. (2004); Barth et al. (2021); Baulch et al. (1981); Baulch et al. (1992); Baulch et al. (1994); Baulch et al. (2005); Becker et al. (1991); Benson (1989); Blitz et al. (2006); Bogdanchikov et al. (2004); Bozzelli & Dean (1989); Bozzelli & Dean (1995); Brown (1973); Brudnik et al. (2009); Bulatov et al. (1990); Burkholder et al. (2020); Campbell & Gray (1973); Cardelino et al. (2003); Caridade et al. (2005); Carl et al. (2005); Choi & Lin (2005); Chung et al. (1975); Clyne & Watson (1974); Clyne & Townsend (1975); Clyne et al. (1984); Cobos et al. (1985); Cohen & Westberg (1991); Cook et al. (1981); Cooper & Hershberger (1992); Craven & Murrell (1987); DeMore et al. (1985); Dorton et al. (1991); Du et al. (2008); Duran et al. (1989); Ebbing & Kaufman (1983); Eiteneer et al. (1998); Ennis & Birks (1988); Estepianian et al. (2001); Fairbairn (1969); Faravelli et al. (2000); Fernandes et al. (2006); Fritz et al. (1982); Fujii et al. (1985); Fulle et al. (1998); Fulle et al. (1998); Gao et al. (2015); Garland (1998); Golden et al. (2003); Gordon et al. (1971); Goumri et al. (1995); Goumri et al. (1999); Greaves et al. (2010); Harada et al. (2010); Harding et al. (2007); Herndon et al. (1999); Hickson & Keyser (2005); Hindiyari et al. (2007); Hobbs et al. (2020); Hong et al. (2012); Huyhn & Violi (2008); Hwang et al. (2010); Iyer et al. (1983); Jacobs et al. (1989); Jasper et al. (2007); Jasper et al. (2009); Javoy et al. (2003); Jodkowski et al. (1995); Jourdain et al. (1979); Ju & Wang (2010); Karkach & Osherov (1999); Keyser (1980); Klippenstein & Kim (1993); Klippenstein et al. (2009); Knipovich et al. (1988); Krasnopol'sky & Pollack (1994); Krasnopol'sky (2007); Kosarapov et al. (2004); Kurbanov & Mamedov (1995); Kurtén et al. (2010); Laidler & Wojciechowski (1961); Lamb et al. (1984); Li et al. (2004); Lilienfeld & Richardson (1977); Lissianski et al. (1995); Lovejoy et al. (1987); Lovejoy et al. (1996); Lu et al. (2003); Markwalder et al. (1993); Martínez et al. (2000); Martinotti et al. (1968); Miller et al. (2003); Mills (1998); Mills et al. (2007); Monks et al. (1993); Montoya et al. (2005); Moses et al. (2002); Mousavi-Pour et al. (2003); Mousavi-Pour et al. (2009); Murakami et al. (2003); Natarajan et al. (1997); Nava & Stief (1989); Nicholais et al. (1979); Nickolaisen et al. (1994); Pugsberg et al. (1979); Robertson & Smith (2006); Roose et al. (1978); Rowe et al. (1998); Schnitler et al. (1996); Seinfeld & Pandis (2016); Sellevag et al. (2009); Sendt et al. (2002); Sendt & Haynes (2005); Senosiain et al. (2006); Sherar et al. (1994); Shum & Benson (1985); Singleton & Cvetanović (1988); Slack (1976); Stoeckel et al. (1985); Stone & Rowley (2005); Straus et al. (1968); Streif et al. (1976); Takahashi & Miyazaki (1977); Tiznit et al. (2010); Tobey et al. (1984); Tsang & Hampson (1986); Tsang (1987); Tsang & Herron (1991); Tsang & Walker (1992); Tsang (1992); Tsuboi & Hashimoto (1981); Tsuboi et al. (1981); Tsuchiya et al. (1994); Tsuchiya et al. (1996); Tsuchiya et al. (1997); Turanay et al. (2012); Vandeputte et al. (2010); Vidal et al. (2017); Wang & Howard (1990); Wang & Hou (2005); Watson et al. (1976); Wei & Timmons (1975); Wine et al. (1988); Woiki & Roth (1995); Wooldridge et al. (1996); Xu & Sun (1998); Xu & Sun (1999); Xu & Lin (2009); Yasunaga et al. (2008); Yoshimura et al. (1992); Yung & Demore (1982); Zanchet et al. (2018); Zhang & Bauer (1997); Zhang et al. (2012).
Table 6
Photochemistry Reactions in STAND2020

| Reactants | Database | References |
|-----------|----------|------------|
| C | PHIDRATES | Verner & Yakovlev (1995); Verner et al. (1996) |
| C\(^1\)D | PHIDRATES | Barfield et al. (1972); Henry (1970) |
| C\(^3\)S | PHIDRATES | Barfield et al. (1972); Henry (1970) |
| H | PHIDRATES | Stobbe (1930); Sauter (1931); Berthe & Salpeter (1957) |
| H\(^2\) | PHIDRATES | Verner & Yakovlev (1995); Verner et al. (1996) |
| NO | PHIDRATES | Verner & Yakovlev (1995); Verner et al. (1996) |
| O | PHIDRATES | Verner & Yakovlev (1995); Verner et al. (1996) |
| O\(^3\)D | PHIDRATES | Barfield et al. (1972); Henry (1970) |
| O\(^5\)S | PHIDRATES | Barfield et al. (1972); Henry (1970) |
| H\(^2\) | PHIDRATES | Gelman (1962); Broad & Reinhardt (1976) |
| C\(^2\) | PHIDRATES | Barfield et al. (1972); Pouilly et al. (1983); Paladil et al. (1985) |
| CH | PHIDRATES | Walker & Kelly (1972); Barsuhn & Nesbet (1978); van Dishoek (1987) |
| CN | PHIDRATES | Henry & McLelroy (1968); Masaoka & Samson (1981); Cairns & Samson (1965) |
| CO | PHIDRATES | Cook et al. (1965); Kronebusch & Berkowitz (1976) |
| N\(^2\) | PHIDRATES | Samson & Cairns (1965); Cook & Metzger (1964) |
| NO | PHIDRATES | Huffman (1969); Samson & Cairns (1964); Cook & Metzger (1964) |
| N\(^2\)O | PHIDRATES | Huffman et al. (1963); Lofthus & Krupenie (1977); Hubert & Herzberg (1979) |
| H\(^2\)O | PHIDRATES | Barfield et al. (1972); Lavender et al. (1984, 1987); Cairns & Samson (1965) |
| H\(^2\)S | PHIDRATES | Ackerman et al. (1970); Herman & Mentall (1982); Huffman (1969) |
| HO | PHIDRATES | Lee et al. (1973); Watanabe et al. (1967) |
| HO\(^2\) | PHIDRATES | Marmo (1953); Kronebusch & Berkowitz (1976) |
| HO\(^2\)C | PHIDRATES | Barfield et al. (1972); Brion et al. (1979); Samson & Cairns (1965) |
| HCO | PHIDRATES | Cook & Metzger (1964); Matsumaga & Watanabe (1967); Watanabe (1958) |
| HCN | PHIDRATES | Lee et al. (1977); Kronebusch & Berkowitz (1976); Bogumil et al. (2003) |
| HCN | PHIDRATES | Burkholler et al. (2020) |
| HO\(^2\) | PHIDRATES | van Dishoeck & Dalgarno (1984); Barfield et al. (1972); Nee & Lee (1984) |
| CO\(^2\) | PHIDRATES | Henry & McLelroy (1968); Cairns & Samson (1965); Nakata et al. (1965) |
| CO\(^2\) | PHIDRATES | Lawrence (1972a, 1972b); Kronebusch & Berkowitz (1976) |
| H\(^2\)O\(^2\) | PHIDRATES | Iyaksoy et al. (2008); Archer et al. (2013); Burkholler et al. (2020) |
| H\(^2\)O\(^2\)C | PHIDRATES | Phillips et al. (1977); Katayama et al. (1973) |
| H\(^2\)O\(^2\)O | PHIDRATES | Watanabe & Junsa (1964); Watanabe & Velikoff (1953); McNesby et al. (1962) |
| HNO | PHIDRATES | Slanger & Black (1982); Kronebusch & Berkowitz (1976); Ranjan et al. (2020) |
| H\(^3\)C \(^1\)O | PHIDRATES | West & Berry (1974); Nuth & Glicker (1982) |
| H\(^3\)C \(^1\)O \(^2\) | PHIDRATES | See references for H\(^2\)O \(^2\) |
| H\(^3\)C \(^1\)O \(^3\) | PHIDRATES | Okabe (1978); Burkholler et al. (2020) |
| H\(^3\)C \(^1\)O \(^4\) | PHIDRATES | Okabe et al. (1971); Moule & Fso (1971) |
| H\(^3\)C \(^1\)O \(^5\) | PHIDRATES | Barfield et al. (1972); Watanabe & Junsa (1964); Goodvee & Stein (1931) |
| S\(^1\) | PHIDRATES | Mills (1998); Krasnopolsky (2007) |
| S\(^2\) | PHIDRATES | Mills (1998); Krasnopolsky (2007) |
| Cl\(^2\)S | PHIDRATES | Mills (1998) |
| SO | PHIDRATES | Barfield et al. (1972); Phillips (1981) |
| SO\(^2\) | PHIDRATES | Barfield et al. (1972); Wu & Judge (1981); Golomb et al. (1962) |
| SO\(^3\) | PHIDRATES | Hinz et al. (2003); Burkholder & McKeen (1997) |
| OCS | PHIDRATES | Barfield et al. (1972); Carnovale et al. (1982); Lee & Chiang (1982) |
| H\(^2\)SO | PHIDRATES | Matsuanna & Watanabe (1967); Molina et al. (1981); Black et al. (1975) |
| ClO | PHIDRATES | Lane & Kjargestad (2008); Farahani et al. (2019) |
| ClO | PHIDRATES | Burkholler et al. (2020) |
| ClO | PHIDRATES | Mills (1998) |
| ClO | PHIDRATES | Mills (1998) |
| ClO | PHIDRATES | Mills (1998) |
| ClO | PHIDRATES | Mills (1998) |
| ClO | PHIDRATES | Barfield et al. (1972); Carnovale et al. (1982) |
| ClO | PHIDRATES | Barfield et al. (1972); Vaghjiani (1993); Steer & Knight (1968) |
| Note. | | a HO has both theoretical and experimental estimates, and one can switch between them. Experimental values are used for all results presented here. |
Table 7
Modified Antoine Equations for Vapor Pressure of Condensable Species

| Species | Equation | References |
|---------|----------|------------|
| CH₄     | \[ \log_{10} p = 4.4251 - \frac{453.92}{T} - \frac{4055.6}{T^2} \times 10^5 + \frac{1.1535 \times 10^6}{T^3} \] | Moses et al. (1992) |
| H₂O     | \[ \log_{10} p = 6.8543 - \frac{3537.0}{T} - \frac{33100.0}{T^4} \times 10^5 + \frac{1.752 \times 10^6}{T^4} + \frac{2.995 \times 10^{10}}{T^5} \] | Rimmer et al. (2020) |
| H₂N     | \[ \log_{10} p = 15.96 - \frac{901.6}{T} \] | Est. |
| HCN     | \[ \log_{10} p = 20.055 - \frac{4522.2}{T^3} \] | Sinozaki et al. (1926) |
| N₂O     | \[ \log_{10} p = 16.22 - \frac{2971.0}{T} \] | Ambrose (1956); Ambrose & Townsend (1964) |
| C₂H₂    | \[ \log_{10} p = 6.2224 - \frac{1644.1}{T} \] | Michels & Wassenaar (1950) |
| C₂H₄    | \[ \log_{10} p = 6.8489 - \frac{1085.0}{T - 2.555} \] | Loomis & Walters (1927) |
| C₂H₆    | \[ \log_{10} p = 7.1349 - \frac{1240.3}{T} \] | Est. |
| C₃H₄    | \[ \log_{10} p = 4.9008 - \frac{1441.4}{T} \] | Est. |
| C₃H₆    | \[ \log_{10} p = 16.413 - \frac{3333.3}{T} \] | Aston et al. (1937) |
| HC₃N    | \[ \log_{10} p = 20.72 - \frac{5087.5}{T} \] | Dannhauser & Flueckinger (1963) |
| NCCN    | \[ \log_{10} p = 16.381 - \frac{2031.0}{T} \] | Stull (1947) |
| C₄H₁₀   | \[ \log_{10} p = 5.5709 - \frac{1461.2}{T} \] | Est. |
| H₂SO₃   | \[ \log_{10} p = 4.4753 - \frac{3229.0}{T} - \frac{3.1723 \times 10^6}{T^2} \times 10^5 + \frac{4.0832 \times 10^8}{T^3} - \frac{2.0321 \times 10^{10}}{T^4} \times 10^6 \] | Greaves et al. (2020a) |
| H₂SO₄   | \[ \log_{10} p = 4.4753 - \frac{3229.0}{T} - \frac{3.1723 \times 10^6}{T^2} \times 10^5 + \frac{4.0832 \times 10^8}{T^3} - \frac{2.0321 \times 10^{10}}{T^4} \times 10^6 \] | Greaves et al. (2020a) |
| S₂      | \[ \log_{10} p = 7.024 - \frac{6091.2}{T} \] | Lyons (2008) |
| S₃      | \[ \log_{10} p = 6.3428 - \frac{6202.2}{T} \] | Lyons (2008) |
| S₄      | \[ \log_{10} p = 6.0028 - \frac{6047.5}{T} \] | Lyons (2008) |
| S₅      | \[ \log_{10} p = 5.1609 - \frac{6741.8}{T} \] | Lyons (2008) |
| S₆      | \[ \log_{10} p = 4.8039 - \frac{2814.1}{T} \] | Lyons (2008) |
| S₇      | \[ \log_{10} p = 5.2127 - \frac{4113.6}{T} \] | Lyons (2008) |
| S₈      | \[ \log_{10} p = 4.1879 - \frac{3269.1}{T} \] | Lyons (2008) |

Notes. \( T \) is in units of K, \( p \) in units of bar.

* The isomer will matter significantly for vapor pressures. This is a way to arbitrarily constrain the abundances of large hydrocarbons in cold reducing atmospheres.

b Estimated to be the same as H₂SO₄ in the atmosphere of Venus.
