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Phosphine as a Biosignature Gas in Exoplanet Atmospheres

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Abstract

A long-term goal of exoplanet studies is the identification and detection of biosignature gases. Beyond the most discussed biosignature gas O₂, only a handful of gases have been considered in detail. In this study, we evaluate phosphine (PH₃). On Earth, PH₃ is associated with anaerobic ecosystems, and as such, it is a potential biosignature gas in anoxic exoplanets. We simulate the atmospheres of habitable terrestrial planets with CO₂- and H₂-dominated atmospheres and find that PH₃ can accumulate to detectable concentrations on planets with surface production fluxes of 10¹⁰ to 10¹⁴ cm⁻² s⁻¹ (corresponding to surface concentrations of 10s of ppb to 100s of ppm), depending on atmospheric composition and ultraviolet (UV) irradiation. While high, the surface flux values are comparable to the global terrestrial production rate of methane or CH_4 (10^{11} cm⁻² s⁻¹) and below the maximum local terrestrial PH_3 production rate (10^{14} cm⁻² s⁻¹). As with other gases, PH_3 can more readily accumulate on low-UV planets, for example, planets orbiting quiet M dwarfs or with a photochemically generated UV shield. PH₃ has three strong spectral features such that in any atmosphere scenario one of the three will be unique compared with other dominant spectroscopic molecules. Phosphine's weakness as a biosignature gas is its high reactivity, requiring high outgassing rates for detectability. We calculate that tens of hours of JWST (James Webb Space Telescope) time are required for a potential detection of PH₃. Yet, because PH₃ is spectrally active in the same wavelength regions as other atmospherically important molecules (such as H₂O and CH₄), searches for PH₃ can be carried out at no additional observational cost to searches for other molecular species relevant to characterizing exoplanet habitability. Phosphine is a promising biosignature gas, as it has no known abiotic false positives on terrestrial planets from any source that could generate the high fluxes required for detection. Key Words: Phosphine—Biosignature— Exoplanet—Atmospheric gases—Spectroscopy—Anoxic environments. Astrobiology 20, xxx-xxx.

1. Introduction

IFE MAKES USE of thousands of volatile molecular species I that could contribute toward a biosphere and its associated atmospheric spectrum. Some of these volatiles may accumulate in a planetary atmosphere and be remotely detectable; these are commonly called "biosignature gases."

Theoretical studies of biosignature gases have been recently heavily reviewed elsewhere (Seager et al., 2016; Grenfell, 2018; Kiang et al., 2018; Schwieterman et al., 2018).

Prominent biosignature gases on Earth are those that are both relatively abundant and spectroscopically active (primarily O2, its photochemical by-product O3, and also CH4 and N₂O). Other gases that are not prominent in Earth's atmosphere but might be prominent in exoplanet atmospheres have also been studied, for example, dimethyl sulfide, dimethyl disulfide, and CH₃Cl (Pilcher, 2003; Segura et al., 2005; Domagal-Goldman et al., 2011).

The next generation telescopes will open the era of the study of rocky exoplanet atmospheres. The James Webb Space Telescope (JWST, planned for launch in 2021) is the most capable for transmission spectra studies of a handful of the most suitable rocky planets transiting bright M dwarf stars (Gardner et al., 2006), whereas ESAs Atmospheric Remotesensing Infrared Exoplanet Large-survey (ARIEL, planned for launch in 2028) may be able to detect atmospheric components

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on super-Earths around the smallest M dwarf stars (Pascale *et al.*, 2018). Large ground-based telescopes now under construction, that is, Giant Magellan Telescope, Extremely Large Telescope, and Thirty Meter Telescope (Johns *et al.*, 2012; Tamai and Spyromilio, 2014; Skidmore *et al.*, 2015), can also reach M dwarf star rocky planets by direct imaging, with the right instrumentation.

To the best of our knowledge, phosphine (PH₃) has not yet been evaluated as a biosignature gas. In Earth's atmosphere, PH₃ is a trace gas. It is possible, however, that biospheres on other planets could accumulate significant detectable PH₃ levels. In particular, anoxic biospheres where life would not be heavily dependent on oxygen could produce PH₃ in significantly higher quantities than on modern Earth (Bains *et al.*, 2019b).

Astronomical observations find that phosphine is spectroscopically active and present in stellar atmospheres (namely carbon stars) and in the giant planet atmospheres of Jupiter and Saturn (Bregman *et al.*, 1975; Tarrago *et al.*, 1992; Agúndez *et al.*, 2014). In T dwarfs and giant planets, PH₃ is expected to contain the entirety of the atmospheres' phosphorus in the deep atmosphere layers (Visscher *et al.*, 2006), where it is sufficiently hot for PH₃ formation to be thermodynamically favored.

In both Jupiter and Saturn, phosphine is found on the high observable layers at abundances (4.8 and 15.9 ppm, respectively) several orders of magnitude higher than those predicted by thermodynamic equilibrium (Fletcher *et al.*, 2009). This overabundance of PH₃ occurs because chemical equilibrium timescales are long when compared with convective timescales (Noll and Marley, 1997). PH₃ forms in the hotter deep layers of the atmosphere (temperatures \geq 800 K) and is mixed upward, so that the PH₃ inventory at the cloud-top is replenished. In every astronomical body where phosphine has been detected thus far, other than on Earth, there are regions with high enough temperatures for PH₃ to be the thermodynamically favored phosphorus species.

It has been postulated that elemental phosphorus species originating from the photolysis of phosphine are responsible for the red coloring of Jupiter's red spot and other Jovian chromophores (Prinn and Lewis, 1975), although this hypothesis has not achieved wide community acceptance (Noy *et al.*, 1981; Kim, 1996). For a review of chemical species that are current candidates for the chromophores of Jupiter, see Carlson *et al.* (2016) and references therein.

Phosphine has not been detected in the observable layers of ice giants, such as Uranus and Neptune (Burgdorf *et al.*, 2004; Moreno *et al.*, 2009), despite these planets having sufficiently hot layers to produce PH₃ and strong convection currents, which could transport PH₃ to observable altitudes. Observations put the P/H abundance in Uranus and Neptune at an upper limit of <0.1 solar P/H, which is significantly lower than expected (Teanby *et al.*, 2019).

In this work, we critically assess phosphine as a biosignature gas. We first summarize in what circumstances phosphine is generated by life on Earth (Sections 2.1 and 2.2). We next review the known destruction mechanisms for PH₃ (Section 2.3) and describe our inputs and methods for the assessment of the detectability of PH₃ in a variety of planetary scenarios (Section 3). We then present our results (Section 4): here, we first calculate surface fluxes and associated atmospheric abundances required for the remote spectroscopic detection of PH₃ in transmission and emission spectra (Section 4.1). We then highlight the properties of the PH₃ spectrum that help distinguish it from other molecules (Section 4.2). Next, we present thermodynamic calculations that show that PH₃, in temperate planets, has no substantial false positives as a biosignature gas (Section 4.3). We conclude with a discussion of our results (Section 5).

2. Phosphine Sources and Sinks

On Earth, phosphine is associated with biological production in anaerobic environments and anthropogenic production via a multitude of industrial processes. PH₃ has low mean production rates on Earth, but it is a mobile gas and is found globally, albeit in trace amounts, in the atmosphere. Below, we summarize the known emissions of PH₃ on Earth (Section 2.1), phosphine's association with life (Section 2.2), and its known destruction mechanisms (Section 2.3).

2.1. Phosphine emissions on Earth

Phosphine is a ubiquitous trace component of the atmosphere on modern Earth (Morton and Edwards, 2005). About 10% of the phosphorus in the atmosphere is PH₃; the major phosphorus form is phosphate, mostly as phosphoric acid (Elm *et al.*, 2017). Although PH₃ is found everywhere in Earth's atmosphere, its atmospheric abundance is widely variant, with high concentration regions sometimes having more PH₃ than low concentration areas by a factor of 10,000 (Pasek *et al.*, 2014).

Phosphine has been found worldwide in the lower troposphere of Earth in the ppq to ppb range in daytime, with higher nighttime concentrations than at daytime (due to inhibited ultraviolet [UV]-induced oxidation) (Gassmann, 1994; Gassmann *et al.*, 1996; Glindemann *et al.*, 1996b, 2003; Ji-ang *et al.*, 1999; Han *et al.*, 2000; Zhu *et al.*, 2006a, 2007a, 2007b; Li *et al.*, 2009; Hong *et al.*, 2010a; Zhang *et al.*, 2010). In the high troposphere, PH₃ was found at a peak of 7 ppt during daylight (Glindemann *et al.*, 2003; Han *et al.*, 2011b). This implies that sunlight does not lead to complete destruction of PH₃, unlike previous suggestions (Glindemann *et al.*, 2003; Han *et al.*, 2011b). A sample of locally measured gaseous PH₃ concentrations in a variety of environments on Earth, ranging from ppq to ppb (ng/m³), can be found in Fig. 1.

On Earth, a significant source of phosphine emissions is anthropogenic activity. Because of its broad toxicity to aerobic organisms,* PH₃ is widely used in the agricultural industry as a rodenticide and insecticide (Devai *et al.*, 1988; Bingham, 2001; Glindemann *et al.*, 2005; Perkins *et al.*, 2015; Chen *et al.*, 2017). PH₃ is also used commercially, for example, as a doping agent (Budavari *et al.*, 1996). However, PH₃ emissions linked to biological activity are believed to form the majority of atmospheric PH₃ (Glindemann *et al.*, 2005; Morton and Edwards, 2005). Evidence for the association of PH₃ with anaerobic biology is presented in Section 2.2.

2.2. Biological production of phosphine

All life on Earth relies on phosphorous compounds. The biological phosphorus cycle is heavily, but not exclusively,

^{*}Exposure to phosphine abundances of 400 ppm results in a quick death (Fluck, 1973).

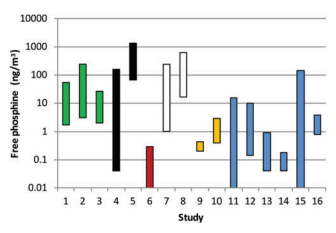


FIG. 1. Measurements of phosphine concentrations in Earth's atmosphere. Study number shown in x axis (references below) and y axis showing the span of locally measured concentration of gaseous PH₃ in units of ng/m³, with maximum values of 600.2 and 1259 ng/m³ (corresponding to concentrations ranging between ppq and ppb). Green bars: marshlands and paddy fields. Black bars: industrial environments. Red bar: Namibia (rural environment). White bars: Arctic and Antarctic environments. Yellow bars: upper troposphere. Blue bars: oceanic samples (coastal and open ocean). References for studies shown: (1) Han et al. (2011a); (2) Han et al. (2000); (3) Niu et al. (2013); (4) Glindemann et al. (1996a); (5) Zhang et al. (2010); (6) Glindemann et al. (1996a); (7) Zhu et al. (2007a, 2007b); (8) Zhang et al. (2010); (9, 10) Glindemann et al. (2003); (11) Li et al. (2009); (12) Zhu et al. (2007a, 2007b); (13) Gassmann et al. (1996); (14) Glindemann et al. (2003); (15) Geng et al. (2005), Han et al. (2011b); (16) Hong et al. (2010a). We do not include measurements of "Matrix-Bound Phosphine" (MBP), material that releases PH₃ when a matrix is treated with high temperatures and strong acid or alkali. Figure adapted from the work of Bains et al. (2019a). Please see Bains et al. (2019a) for more details on MBP and environmental PH3 production. Gaseous PH3 is found in multiple altitudes in Earth's atmosphere above a wide variety of environments, in concentrations ranging from ppq to ppb. MBP, Matrix-Bound Phosphine; PH₃, phosphine. Color images are available online.

reliant on phosphates. Other, less oxidized, phosphorus-containing molecules also play a crucial role in the phosphorus cycle. The exact role of phosphine in the Earth's global phosphorus cycle is not yet fully known. It is, however, likely that, similar to other reduced phosphorus species, PH₃ also has an important role in the global cycling of this essential element. For more details on the role of phosphines in the Earth's phosphorus cycle, please see Appendix E.

Biological phosphine production is associated with microbial activity in environments that are strictly anoxic (lacking oxygen). This finding is in alignment with the fact that the toxicity of PH_3 is intrinsically linked to its interference with O_2 -dependent metabolism (Bains *et al.*, 2019b).

The argument that phosphine is associated with anaerobic life is strengthened by its detection in a wide variety of ecosystems with anoxic niches, including above penguin colonies, rich in bird guano, where it reaches abundances of 300 ppt (Ji-ang *et al.*, 1999; Zhu *et al.*, 2006a, 2007a; Li

et al., 2009; Hong et al., 2010a); animal intestinal tracts, flatus, and feces[†] (Gassmann and Glindemann, 1993; Eismann et al., 1997a; Chughtai and Pridham, 1998; Zhu et al., 2006a, 2006b, 2014); paddy fields[‡] (Han et al., 2011a; Chen et al., 2017); rivers and lakes (Feng et al., 2008; Geng et al., 2010; Hong et al., 2010b; Han et al., 2011b; Ding et al., 2014); wetlands and marshlands (Devai and Delaune, 1995; Eismann et al., 1997b; Glindemann et al., 1996b); and landfills and sludges (Roels and Verstraete, 2004; Ding et al., 2005a, 2005b). Several studies have also reported the production of PH₃ from mixed bacterial cultures in the laboratory (Rutishauser and Bachofen, 1999; Jenkins et al., 2000; Liu et al., 2008); in one case, bacteria turning half of the phosphorus in the culture medium (~180 mg/L) into PH₃ in 56 days (Devai et al., 1988).

Despite a large body of robust circumstantial evidence for the production of phosphine by life, the exact mechanisms for biologically associated production of PH3 are still debated. Recent work postulates that PH₃ production may be associated with the microbial tricarboxylic acid cycle of Enterobacteriaceae (Fan et al., 2020), but the exact metabolic pathway leading to PH₃ production in anaerobic organisms remains unknown. However, we note that the absence of a known enzymatic mechanism is not evidence for the absence of biological production. The synthetic pathways for most of life's natural products are not known, and yet their origin is widely accepted to be biological because of the implausibility of their abiotic synthesis, their obligate association with life, and their chemical similarity to other biological products. For example, a recently published, manually curated, database of natural molecules produced by life on Earth contains $\sim 220,000$ unique molecules of biological origin, produced by thousands of species (Petkowski et al., 2019a), whereas the number of known. experimentally elucidated, metabolic pathways from organisms belonging to all three domains of life is only ~ 2720 (Caspi et al., 2017). Further examples of the complexities in discovering metabolic pathways for molecules associated with biological activity are provided in Appendix E.

There are two proposed explanations for the production of phosphine in anoxic ecosystems [reviewed in Glindemann *et al.* (1998), Roels and Verstraete (2001), Roels and Verstraete (2004), Roels *et al.* (2005), Bains *et al.* (2019a, 2019b)]:

- (1) PH₃ is directly produced by anaerobic bacteria from environmental phosphorus.
- (2) PH₃ is indirectly produced by anaerobic bacteria. Anoxic fermentation of organic matter by anaerobic bacteria results in acid products; these acid products, in turn, could react with inorganic metal phosphides, like those present as trace elements in scrap metal, resulting in phosphine generation.

Of the two proposed explanations for biologically associated production of phosphine, we argue that the direct production as a result of metabolic activity of anaerobic bacteria is the most likely. Our reasoning is based on two lines of evidence:

[†]We note that animal guts are anaerobic, even in small animals. [‡]In agricultural wetlands, both industrial and biological sources of PH₃ are plausible.

(a) PH₃ has been detected in significant amounts in bacterial cultures in controlled laboratory experiments, where no metal phosphides were present, making the indirect acid-dependent production of PH₃ an unlikely scenario (Devai et al., 1988; Glindemann et al., 1996b; Jenkins et al., 2000; Schink and Friedrich, 2000; Ding et al., 2005a, 2005b; Liu et al., 2008).

(b) Several independent studies found that PH₃ was detected in feces from evolutionarily distant animals, inhabiting diverse environments, for example, insects, birds, and mammals (including humans) (Gassmann and Glindemann, 1993; Chughtai and Pridham, 1998; Zhu et al., 2014). It is implausible that there is a significant presence of contaminant metal phosphides in the guts of all the animals, which would be required for an indirect acid-dependent production of PH₃.

We end this introduction to the biological association of phosphine by noting that thermochemical studies on the feasibility of the production of PH₃ in temperate environments have found no plausible thermodynamically favored abiotic pathways, and as such, PH₃ has no substantial false positives for life (see Section 4.3 and Appendix C) (Bains *et al.*, 2019a). Conversely, the production of PH₃ under anoxic conditions by living systems can be thermochemically favorable (Bains et al., 2019a), and biological functions that are accomplished through energy consuming reactions are not uncommon (Bains et al., 2019b). PH₃ could be used by life to perform complex functions that would warrant an energetic investment, such as signaling or a defense mechanism (Bains et al., 2019b). For more information about PH₃ in the context of terrestrial biology and the thermodynamic feasibility of PH₃ production by life, see Bains et al. (2019a, 2019b) and Appendices C-E.

2.3. Phosphine chemistry in the atmosphere

Within an atmosphere, phosphine is destroyed by the radicals O, H, and OH in reactions, which are thought to be first order with respect to its reactants and second order overall. PH_3 can also be regenerated by reaction of PH_2 with H and directly photolyzed by UV radiation. These processes are summarized below and discussed in Section 5.2.

Reaction rate constants are expressed via the Arrhenius equation:

$$k = Ae^{-E/RT}$$
.

where k is the reaction rate constant in units of cm³ s⁻¹, A is a constant in units of cm³ s⁻¹, E is the activation energy in units of J mol⁻¹, R is the gas constant in units of J mol⁻¹ K⁻¹, and T is temperature in K.

2.3.1. Destruction by OH radicals. Oxidation with OH radicals is thought to be the main sink for phosphine in Earth's atmosphere via the reaction (Cao *et al.*, 2000; Glindemann *et al.*, 2005; Elm *et al.*, 2017):

$$PH_3 + OH \rightarrow H_2O + PH_2$$

 $PH_2 + [O \text{ species}] \rightarrow \rightarrow \text{ products (including } H_3PO_4).$

For this reaction, $A = 2.71 \times 10^{-11}$ cm³ s⁻¹, E = 1.29 kJ mol⁻¹, corresponding to $k_{\text{PH}_3,\text{OH}} = 2 \times 10^{-11}$ cm³ s⁻¹ at T = 288 K (Fritz *et al.*, 1982). The lifetime of PH₃ due to OH reactions

is calculated to be 28 h at night and 5 h in daytime, with the difference controlled by the concentration of UV-generated OH (Glindemann *et al.*, 2003). The destruction of PH₃ by OH in the atmosphere eventually leads to phosphoric acid, which in turn contributes to the soluble phosphates found in rainwater (Lewis *et al.*, 1985; Elm *et al.*, 2017).

2.3.2. Destruction by O radicals. Phosphine also reacts very rapidly with atomic oxygen (on Earth, generated by photolyzed ozone), with reaction:

$$PH_3 \ + \ O \ \ \longrightarrow \ \ products \ (including \ H_3PO_4).$$

For this reaction, $A = 4.75 \times 10^{-11}$ cm³ s⁻¹, $E \approx 0$, corresponding to $k_{\rm PH_3,O} = 5 \times 10^{-11}$ cm³ s⁻¹ at T = 288 K (*i.e.*, temperature-independent in 208–423 K, Nava and Stief, 1989). Because atomic oxygen is less abundant than OH in Earth's atmosphere and troposphere, PH₃ destruction by OH is still the dominant route despite PH₃ reacting with O at a higher rate. On Earth, OH radicals described above are partially generated from the interaction between O radicals and water vapor, and so the reactions that produce the two radical species are not happening in isolation (Jacob, 1999). In anoxic atmospheres, however, the main source of OH and H radicals is the photolysis of water vapor (Hu *et al.*, 2012).

2.3.3. Destruction by H radicals. Phosphine can be destroyed by the H radical via the reaction:

$$PH_3 + H \rightarrow H_2 + PH_2$$
.

For this reaction, $A = 7.22 \times 10^{-11}$ cm³ s⁻¹, E = 7.37 kJ mol⁻¹, corresponding to $k_{\rm PH_3,H} = 3 \times 10^{-12}$ cm³ s⁻¹ at T = 288 K (Arthur and Cooper, 1997). The reaction of PH₃ with the H radical is most relevant to H₂-rich atmospheres (Seager *et al.*, 2013b).

2.3.4. Recombination from H radicals. Phosphine can be regenerated by the radical recombination reaction:

$$PH_2 \ + \ H \rightarrow PH_3.$$

With rate constant $k_{\rm PH_2,H} = 3.7 \times 10^{-10} \exp(-340 K/T) \, \rm cm^3 \, s^{-1}$, corresponding to $k_{\rm PH_2,H} = 1.1 \times 10^{-10} \, \rm cm^3 \, s^{-1}$ at $T = 288 \, \rm K$ (Kaye and Strobel, 1984). If [PH₂] is high, this reaction can be a major reformation pathway for PH₃ (see Section 5.2).

2.3.5. Destruction through UV radiation. UV radiation is thought to directly photolyze phosphine with unit quantum efficiency upon absorption of irradiation at wavelengths ≤230 nm (Visconti, 1981; Kaye and Strobel, 1984):

$$PH_3 + hv \rightarrow PH_2 + H.$$

This photolysis reaction is not relevant on UV-shielded planets (*e.g.*, modern Earth with its ozone layer) but could be relevant on anoxic planets where UV radiation may penetrate deeper into lower altitudes of the atmosphere.

Overall, phosphine is destroyed by UV irradiation, through both direct photolysis and reactions with UV-generated radical species. The daytime–nighttime PH₃ concentration difference on Earth is large due to the generation of radicals by UV irradiation during the day and their comparative absence at night. However, PH₃ has been detected at concentrations of up to 7 ppt (2.45 ng m⁻³) during daylight in Earth's high troposphere (Glindemann *et al.*, 2003). PH₃ accumulates in the dry upper troposphere on Earth because ozone attenuation of UV and lack of OH-producing H₂O result in low abundances of OH radicals, which slows the PH₃ destruction and its return to the surface in the form of phosphates (Frank and Rippen, 1987; Glindemann *et al.*, 2003).

2.3.6. Solubility and aerosol formation. Phosphine does not easily stick to aerosols and has very low water solubility (Fluck, 1973). PH₃ is therefore a very mobile gas that is less likely to wash out and fall to the surface than other gases, such as hydrogen sulfide, methanethiol, and ammonia (Glindemann *et al.*, 2003).

UV photolysis of phosphine in the presence of hydrocarbons could lead to the formation of complicated alkylphosphines (Guillemin *et al.*, 1995, 1997). Atmospheres that are prone to high concentrations of hydrocarbon radicals, such as H₂-rich atmospheres, could therefore lead to the creation of organophosphine hazes. The plausibility of organophosphine haze formation is discussed further in Section 5.2.

3. Inputs and Methods for the Assessment of Detectability

We have assessed the spectral distinguishability, atmospheric survival, and observational detectability of phosphine in anoxic exoplanets. In this section, we first describe the choice of molecular inputs used for our spectral analyses (Section 3.1). We then provide a brief outline of the photochemical method used to calculate the distribution of molecules throughout the atmosphere (Section 3.2). Finally, we outline the method and detectability criteria for the simulations of observational spectra (Section 3.3).

3.1. Molecular inputs

Molecular spectra can be represented in various forms to best serve as input for spectral representations and atmospheric models. For the comparison of phosphine with other major components of atmospheres (see Section 4.2), we have used cross sections calculated from the most complete spectra available. The PH₃ molecular cross sections come from the recently variationally computed PH₃ line list (Sousa-Silva et al., 2015) and the total internal partition function calculated in the work of Sousa-Silva et al. (2014). For all temperatures under 800 K (which includes all temperate environments), this PH₃ line list represents a complete spectrum containing over 16 billion transitions between 7.5 million energy levels. Even at low temperatures, it is recommended that complete line lists are used for spectral simulations; complete line lists allow for more representative cross sections with improved band shapes when compared with experimental or calculated spectra at room temperature. The carbon dioxide line list is from HI-TEMP (Rothman et al., 2010). All other molecular cross sections are simulated by using complete, theoretically calculated, line lists from the ExoMol database (Yurchenko et al., 2011; Yurchenko and Tennyson, 2014; Sousa-Silva et al., 2015; Tennyson et al., 2016).

For the calculation of the transmission and thermal emission spectra of the model atmospheres (see Section 4.1), molecular opacities for phosphine are adopted from the ExoMol database (Sousa-Silva *et al.*, 2015; Tennyson *et al.*, 2016). For all other molecules, we used the HITEMP and the HITRAN 2016 databases (Rothman *et al.*, 2010; Gordon *et al.*, 2017). Molecular cross sections in the UV region, used to calculate photolysis rates (see Section 3.2), were obtained from the absorption cross sections compendium of Ranjan and Sasselov (2017) and from the work of Chen *et al.* (1991) via the MPI-Mainz Spectral Atlas (Keller-Rudek *et al.*, 2013).

3.2. Photochemical modeling

In this subsection, we provide a brief description of the photochemical model used to calculate the concentration of phosphine as a function of altitude for a range of PH₃ surface fluxes. We also describe the atmospheric and stellar scenarios considered in our photochemical model. We found that it is necessary to use a photochemical model instead of the approximation of fixed radical profiles (Seager *et al.*, 2013b) because of the intense reactivity of PH₃, which can drastically alter the radical profiles of an atmosphere. In particular, at high PH₃ fluxes, the radical concentrations are suppressed due to reactions with PH₃, meaning that PH₃ can build up to much higher concentrations than a fixed radical profile approximation would predict.

3.2.1. Photochemical model. We adapt the photochemical model of Hu et al. (2012) to calculate atmospheric composition for different planetary scenarios. The model is detailed in the work of Hu et al. (2012); in brief, the code calculates the steady-state chemical composition of an exoplanetary atmosphere by solving the one-dimensional chemical transport equation. The model treats up to 800 chemical reactions, photochemical processes (i.e., UV photolysis of molecules), dry and wet deposition, surface emission, thermal escape of H and H₂, and formation and deposition of elemental sulfur and sulfuric acid aerosols. The model is designed to have the flexibility of simulating both oxidized and reduced conditions. UV and visible radiation in the atmosphere is computed by the delta-Eddington two-stream method. The code has been validated by reproducing the atmospheres of modern Earth and Mars. The code and extensive application examples are described in several studies (Hu et al., 2012; Hu et al., 2013; Seager et al., 2013a, 2013b; Hu and Seager, 2014). In calculating convergence, we required that the chemical variation timescale of significant species (>1 cm⁻³) to be at least 10¹⁹ s, that is, longer than the age of the universe.

We added phosphine to the model of Hu *et al.* (2012). We considered surficial production as the only source of PH₃, and rainout, photolysis, and reactions with the main radical species O, H, and OH as the sinks (see Section 2.3). We take the Henry's Law constant for PH₃ from the work of Fu *et al.* (2013) via Sander (2015). For photolysis, we take the PH₃ UV cross sections at 295 K from the work of Chen *et al.* (1991). We follow Kaye and Strobel (1984) in taking the branching ratio of this reaction to be unity and take the quantum yield of PH₃ photolysis to be $q_{\lambda} = 1$ for $\lambda < 230$ nm and $q_{\lambda} = 0$ for $\lambda > 230$ nm. For the reactions with OH,

O, and H, we take the rate constants from the works of Fritz *et al.* (1982), Nava and Stief (1989), and Arthur and Cooper (1997), as detailed in Section 2.3. We are unaware of geochemical constraints on the dry deposition velocity of PH₃; we take this value to be 0 cm s^{-1} , which could lead to an overestimation of PH₃ accumulation rates (see Section 5.2 for a discussion of possible PH₃ deposition). However, our approach neglects the possibility that atmospheric photochemistry may generate PH₃. In particular, we do not consider the recombination reaction, PH₂ + H \rightarrow PH₃. This may lead to underestimating PH₃ accumulation, especially in H₂-dominated atmospheres where H abundances are high.

3.2.2. Planetary scenarios. We model the atmospheres of Earth-sized, Earth-mass planets with two bulk atmospheric compositions: a H₂-dominated atmosphere and a CO₂-dominated atmosphere. We focus on H₂-dominated atmospheres because their low mean molecular masses make them amenable to characterization via transmission spectroscopy (Batalha et al., 2015). We focus on CO₂dominated atmospheres as an oxidizing end-member to complement the reducing H₂-dominated case, and because early Earth is thought to have had a CO₂-rich atmosphere (Kasting, 1993). We only consider anoxic atmospheres because O₂-rich atmospheres are likely to have large quantities of OH radicals, which rapidly destroy phosphine (see Section 2.3). Additionally, the aerobic metabolism of O₂dependent life is likely to be incompatible with widespread PH₃ biological production (Bains et al., 2019b).

Our atmospheres correspond to the H₂- and CO₂-dominated benchmark scenarios of Hu *et al.* (2012), with the key difference that we do not set the rainout rates of H₂, CO, CH₄, C₂H₆, or O₂ to zero, as Hu *et al.* (2012) did to simulate an abiotic planet. In brief, we consider planets with surface pressures of 1 bar, surface temperatures of 288K, and bulk dry atmospheric composition of 10% N₂, 90% H₂/CO₂ for the H₂/CO₂-dominated cases, respectively. The temperature profile is taken to evolve as a dry adiabat until 160 and 175 K for the H₂- and CO₂-dominated cases, respectively, and isothermally thereafter.

The strength of vertical mixing is scaled from that measured in Earth's atmosphere according to the mean molecular mass. The H₂O concentration at the bottom of the atmosphere is set to 0.01, corresponding to 60% humidity. H₂, CO₂, CH₄, SO₂, and H₂S are emitted from the surface at rates corresponding to terrestrial volcanism. The mixing ratio profile of the dominant gases (gases with abundances exceeding 100 ppb) used for the modeling of the H₂-rich atmosphere on a massive super Earth orbiting an active M dwarf is shown in Fig. 2 (see Appendix A for the mixing ratio profiles used to model the remaining atmospheric scenarios). For further details, including the rationale for these parameters, see Hu *et al.* (2012).

Stellar irradiation is a key input for photochemical models. We considered instellation corresponding to the Sun (our "Sun-like" case) (Hu *et al.*, 2012) and from the M dwarf GJ1214 (our "active M dwarf" case) (Seager *et al.*, 2013b). The semimajor axes of the planets for the H₂- and CO₂-dominated cases are taken to be 1.6 AU and 1.3 AU for the Sun-like case, and 0.042 AU and 0.034 AU for the "active M dwarf" case, corresponding to surface temperatures of 288 K at 0 PH₃ emission.

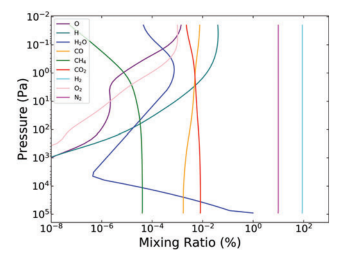


FIG. 2. Mixing ratio profile of a H_2 -rich atmosphere on an Earth-sized planet orbiting an active M dwarf. Vertical axis represents pressure in units of Pa, and the horizontal axis shows the mixing ratio represented as a percentage of the total atmospheric layer. Figure partially adapted from the works of Hu *et al.* (2012) and Seager *et al.* (2013b). Color images are available online.

For a sensitivity test, we also considered a theoretical "quiet" M dwarf simulated by reducing the UV-flux of GJ1214 by three orders of magnitude (corresponding to approximately a factor of 100 less UV radiation than the least active M dwarf known GJ581) (France *et al.*, 2016). See Section 4.1.4 for a discussion of the sensitivity of our results to changes in surface temperature and to low UV irradiation levels.

3.3. Atmospheric spectral simulations

We use the outputs of the photochemical models described above to model observational spectra projected to massive super-Earths with $M_p = 10 \ M_E$ and $R_p = 1.75 \ R_E$. We focus on such large massive planets due to the following observational considerations: (1) such planets are easier to detect via radial velocities and transit observations; (2) large planets have larger thermal emission signatures; and (3) massive planets are more likely to retain H_2 -rich atmospheres, which are much easier to characterize in transmission because of their larger scale heights than other atmosphere types.

3.3.1. Transmission and emission spectral calculations. Transmission and thermal emission spectra were simulated with the program SEAS (Simulated Exoplanet Atmosphere Spectra). The projection from Earth-sized to super-Earths is performed by using equivalent techniques to those in the work of Hu *et al.* (2012); SEAS takes as input a list of molecular mixing ratios as a function of pressure, which, to first order, are invariant to changes in the surface gravity.

The transmission spectrum code calculates the optical depth along the limb path and sums up chords, assuming that the planet atmosphere is homogeneous. The SEAS transmission spectrum code is similar in structure to that described in two studies (Miller-Ricci *et al.*, 2009; Kempton

et al., 2017), with the main difference being that SEAS can accept variable mixing ratio inputs, important for super-Earths whose atmospheres are severely impacted by photochemistry. The temperature–pressure profile, including limits and resolution, is specified by the user. Molecular line lists are taken from HITRAN 2016 and ExoMol databases (Gordon et al., 2017; Tennyson et al., 2016), with cross sections calculated with HAPI (Kochanov et al., 2016) and ExoCross (Yurchenko et al., 2018), respectively. The molecular species are chosen by the user, and all molecules in the HITRAN and ExoMol databases are user-selectable options.

The thermal emission code integrates a blackbody exponentially attenuated by the optical depth without scattering (Seager, 2010). The code uses the same input temperature–pressure profile and molecular cross sections as described above.

SEAS considers clouds in the emergent spectra for thermal emission by averaging cloudy and cloud-free spectra (resulting in 50% cloud coverage). We omit clouds or hazes for the transmission spectra model; if the atmosphere is cloudy or hazy at high altitudes, the spectral features in transmission will be muted. Consequently, our calculations represent upper bounds on the magnitude of the transmission features with respect to cloud or haze effects. We discuss the impact of clouds in our modeled transmission spectra in Section 5.2.

The SEAS transmission code has been validated by comparing results with the Atmospheric Chemistry Experiment data set (Bernath *et al.*, 2005) for transmission spectrum and with the MODTRAN spectrum (Berk *et al.*, 1998) for thermal emission spectrum. We also compared results related to this phosphine work with transmission spectra generated by the code described in the work of Hu *et al.* (2012).

3.3.2. Detectability metric. We study the spectroscopic detectability of phosphine in H_2 - and CO_2 -rich atmospheres, in transmission and emission observation scenarios. In transmission, we compare with the mean transit depth of the planet radius, and in emission with the blackbody curve.

We consider a 6.5 m space telescope, having a quantum efficiency of 25% observing with a 50% photon noise limit. We consider our 1.75 $R_{\rm Earth}$ -planet to be orbiting (1) a 0.26- $R_{\rm sun}$ M dwarf star at 5 pc with an effective temperature of 3000 K; and (2) a Sun-like star. Stellar flux is the source of the noise, combining in-transit and out-of-transit flux noises. The theoretical transmission spectra are based on JWST and its NIRSpec and MIRI instruments (Bagnasco *et al.*, 2007; Wright *et al.*, 2010). To calculate theoretical thermal emission spectra, we consider a secondary eclipse scenario as observed from JWST with the MIRI instruments (both mid- and low-resolution spectrometers). We binned the data to a resolution of $R \sim 10$ to increase the significance of detection.

We investigate the detectability of phosphine in exoplanet atmospheres by adapting the detection metrics defined by Seager *et al.* (2013b) and Tessenyi *et al.* (2013). The detectability metric is a theoretical metric that uses simulated data.

We first simulate model-independent observational data for all planetary scenarios considered (*e.g.*, using the instrumental constraints of JWST). For an analysis of the transmission spectra models, we then compare the wavelength-dependent transit depth of the planet to the "white-light" transit depth

in each wave band (corresponding to the coverage of each instrument).

Phosphine is considered detectable if we can detect opacity at wavelengths corresponding to PH_3 absorption features with statistically significant confidence. To establish the statistical significance of opacity detection in transmission, we assume a simulated spectrum and then assign binned values for the transit depth. We then calculate the wavelength-dependent one-sigma $(1-\sigma)$ error bar for each binned value (*i.e.*, standard deviation) using only stellar photon noise. The significance of the deviation is calculated with:

$$\frac{|\mu_{\lambda}-\mu_{\lambda}|}{\sqrt{\sigma_{\lambda}^2+\sigma_{\lambda}^2}},$$

where μ_{λ} is the wavelength-dependent transit depth of the simulated atmosphere, μ_{λ} is the mean transit depth of the white-light averaged waveband, and σ is the uncertainty on the measurement. The uncertainties are estimated based on shot noise. We then assess the detectability of phosphine by simulating a model atmosphere with and without PH₃ and comparing the deviation of each atmosphere from their associated mean. This comparison establishes whether a model atmosphere with PH₃ fits the simulated observational data better than one without PH₃.

In thermal emission, we use a similar detectability metric to the transmission analysis described above, with the distinction that we calculate the deviation of our modeled atmosphere spectra from its best-fit blackbody continuum (instead of the white light average used for transmission comparisons). The temperature of the blackbody is set by fitting a blackbody curve to the simulated data.

The integration time is a variable parameter in the SEAS models, but features are only considered detectable if they achieve at least a $3-\sigma$ interval with 200 observation hours or less (considering 100 h in-transit and 100 h out-of-transit).

3.3.3. Scaling to smaller planets. We performed our spectral simulations for a massive super-Earth planet (M_p = 10 M_E and R_p =1.75 R_E). In this section, we consider how the prospects for atmospheric characterization scale to smaller, more Earth-sized worlds.

The amplitude of the atmospheric absorption signal in transmission is characterized by Brown (2001):

$$\frac{\delta A}{A} = \frac{2\pi R_p (kT/\mu g)}{\pi R_*^2} = \frac{2\pi R_p \left(kTR_p^2/\mu G M_p\right)}{\pi R_*^2} \propto \frac{R_p^3}{M_p},$$

where R_p is the planet radius and M_p is the planet mass. This implies that the transmission spectroscopy signal from a 1 R_E , 1 M_E planet should be twice the signal from the 1.75 R_E , 10 M_E planet we consider here, and the phosphine surface fluxes required to produce a detectable atmospheric signal should be half of what we model for our super-Earth scenario. §

 $^{^{\}S}$ We note that it is uncertain whether a small planet can retain a H_2 -dominated atmosphere over geological time in the face of atmospheric escape.

The amplitude of the thermal emission spectral signal is characterized by:

$$F(\lambda) \propto [B(T_{cont}, \lambda) - B(T_{line}, \lambda)]R_p^2$$

where λ is the wavelength, B is the blackbody function, and T_{cont} and T_{atm} are, respectively, the brightness temperature in and out of the spectral line under consideration. The above equation implies that the thermal emission signal from a 1 R_E , 1 M_E planet should be a third of the signal from the 1.75 R_E , 10 M_E planet we consider here, and the PH₃ surface fluxes required to produce a detectable atmospheric signal should be three times larger than what we model for our super-Earth scenario.

We conclude that spectrally characterizing Earth-sized planets is comparable in difficulty to characterizing super Earth planets, but that characterizing the atmospheres of smaller worlds is somewhat easier in transmission and somewhat harder in emission. These differences do not affect our main conclusions.

4. Results

We find that phosphine as a detectable biosignature gas has three encouraging properties: (1) PH_3 can accumulate to detectable levels in an exoplanet atmosphere, provided it has a high production rate at the planet's surface (Section 4.1); (2) PH_3 has unique spectral features, namely strong bands around 2.7–3.6, 4.0–4.8, and 7.8–11.5 μ m, which allow it to be distinguishable from other dominant atmosphere molecules (Section 4.2); and (3) based on the abundances and surface fluxes needed to produce detectable levels of PH_3 , it has no known false positives provided that the planet's surface temperature is below 800 K (Section 4.3). In addition to the above, our results show that, at surface fluxes near the minimum flux necessary to allow for PH_3 detection, a "runaway" effect for PH_3 occurs that drastically alters the atmosphere (Section 5.1).

We present results for the phosphine detection in H_2 - and CO_2 -rich atmospheres, for planets orbiting Sun-like stars and active M dwarf stars.

4.1. Phosphine detection in exoplanet atmospheres

We performed a series of simulations and calculations to explore the prospects for detecting phosphine in an exoplanet atmosphere using transmission and thermal emission spectroscopy. We consider H₂- and CO₂-dominated atmospheres, and stellar irradiation environments corresponding to the modern Sun ("Sun-like") and GJ1214 ("active M dwarf"); see Seager *et al.* (2013b) for details.

In this section, we first present a set of simulated spectra, both in transmission (Section 4.1.1) and in emission (Section 4.1.2), for atmospheric scenarios with and without phosphine. Here, we predict the minimum abundances required for PH₃ to be detectable in each atmosphere considered. We calculate the PH₃ surface production rates needed for PH₃ to achieve the abundances required for detection in Section 4.1.3, using our photochemical model to simulate the equilibrium distribution of atmospheric gases throughout each atmosphere. Finally, we assess the sensitivity of our results to a variable surface temperature and a host star with low levels of radiation (Section 4.1.4).

4.1.1. Amount of phosphine required for detection via transmission spectroscopy. We find that phosphine is detectable in anoxic atmospheres only if it is able to accumulate to the order of ppb to 100s of ppm, for H_2 - and CO_2 -rich atmospheres, respectively. For comparison, PH_3 is present at the ppt to ppb level on modern Earth. We estimate the photochemical plausibility of PH_3 accumulating to such large abundances in Section 4.1.3.

Unfortunately, even with high concentrations of phosphine in the atmosphere, many tens of hours of JWST time are needed to detect it. The atmosphere mixing ratios, the surface production rates required, and the number of observation hours needed for different planet and host star scenarios are provided in Table 1.

Planets with H₂-rich atmospheres orbiting active M dwarfs require the smallest phosphine abundances for its detection (10s to 100s of ppb; Fig. 3), which can be expected due to their lower mean radical concentrations compared with an oxidized atmosphere (Seager *et al.*, 2013b). H₂-rich atmospheres also have transmission spectra that are easier to detect than planets with higher mean molecular-weight atmospheres (*e.g.*, CO₂) because of their larger scale height, that is, a "puffier" atmosphere.

For H_2 -rich atmospheres, only the strongest band of phosphine at 4.0–4.8 μ m can be detectable (Fig. 3). The other PH_3 features are either too weak or contaminated by other atmospheric molecular species.

For CO₂-rich atmospheres (Fig. 4), several spectral bands of phosphine show substantial spectral absorptions when compared with atmospheres without PH₃. Nonetheless, no spectral band of PH₃ can achieve a 3-σ statistical significance even after 200 observation hours when considering a planet orbiting an active M dwarf.

Phosphine is very difficult to detect on planets orbiting Sun-like stars. Planets with CO_2 -dominated atmospheres require longer than 200 observation hours for the detection of PH₃ in transmission spectra, even with the highest surface fluxes considered $(3 \times 10^{13} \text{ cm}^{-2} \text{ s}^{-1})$. The detection of PH₃ can only achieve a 3- σ statistical significance on planets with H₂-rich atmospheres for fluxes of $10^{14} \text{ cm}^{-2} \text{ s}^{-1}$ (Table 1); this flux is comparable to the highest recorded PH₃ flux on Earth (above sewage plants) (Devai *et al.*, 1988) and above the values for the biological production of methane, which on Earth corresponds to $1.2 \times 10^{11} \text{ cm}^{-2} \text{ s}^{-1}$ (Segura *et al.*, 2005; Guzmán-Marmolejo and Segura, 2015).

The results presented above show that it is possible, but difficult, to detect phosphine in anaerobic atmospheres if it is present as a trace gas. However, if PH₃ production rates increase sufficiently, they outpace the ability of stellar NUV photons to destroy PH3, whether via direct photolysis or via generation of radical species. PH3 may then become a significant component of the atmosphere (e.g., 100s to 1000s of ppm), and its detectability increases dramatically. The PH₃ surface fluxes required to reach this runaway phase ($\sim 10^{12}$ cm⁻² s⁻¹) are not significantly higher than those required for detection ($\sim 10^{10-11}$ cm⁻² s⁻¹). For example, with surface production rates only nine times larger than those that produce the atmospheric spectrum shown in Fig. 3, PH₃ reaches the runaway threshold and can be detected with $5-\sigma$ statistical significance after only 3 h of observation (Fig. 5). The plausibility of this runaway effect is discussed further in Sections 4.1.3 and 5.1.

Table 1. Phosphine Mixing Ratios Needed for Detection in Transmission for Different Atmospheric and Stellar Scenarios, with Associated Observation and Surface Flux Requirements

Atmospheric scenario	Required mixing ratio for detection	Minimum observation hours (in-transit+out- of-transit)	Associated confidence interval for phosphine detection (σ)
H ₂ -dominated, Sun-like star	780 ppm	56	3
H ₂ -dominated, active M dwarf (Fig. 3)	220 ppb	91	3
H ₂ -dominated, active M dwarf	220 ppb	200	4.4
H ₂ -dominated, active M dwarf	5 ppb	200	2.5
H ₂ -dominated, active M dwarf	0.28%	3	5
CO ₂ -dominated, Sun-like star	N/A	Not detectable	N/A
CO ₂ -dominated, active M dwarf (Fig. 4)	310 ppm	200	2.7
CO ₂ -dominated, active M dwarf	7.6%	32	3

For planets orbiting an active M dwarf, phosphine requires minimum abundances of 220 ppb and >310 ppm to be detectable with a 3- σ confidence interval on H₂- and CO₂-rich atmospheres, respectively. Values in italics correspond to atmospheric scenarios where PH₃ is at a runaway threshold (see Fig. 5 and Sections 4.1.3 and 5.1). For planets orbiting a Sun-like star, PH₃ must become a major component of the atmosphere for its detection to be possible with less than 200 observation hours or, in the case of H₂-rich atmospheres, with surface production rates above those found anywhere on Earth (*i.e.*, >10¹⁴ cm⁻² s⁻¹). For comparison, biological production of CH₄ on Earth corresponds to 1.2×10^{11} cm⁻² s⁻¹, of which a significant proportion is due to human activity (Houghton, 1995; Segura *et al.*, 2005; Guzmán-Marmolejo and Segura, 2015).

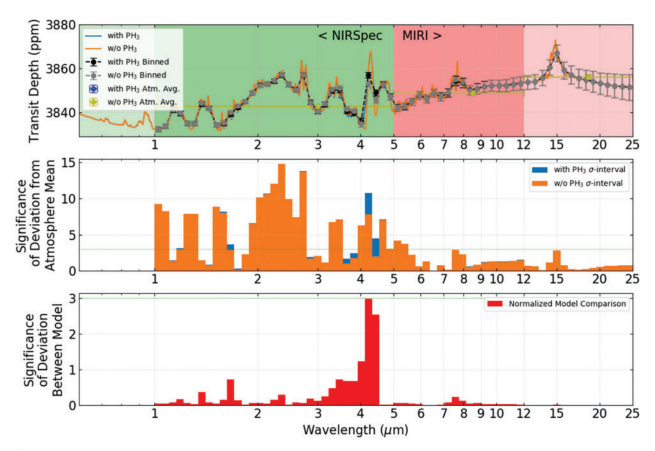


FIG. 3. Theoretical transmission spectra for a H_2 -rich atmosphere on a $10\,M_E$, $1.75\,R_E$ planet with a surface temperature of 288 K orbiting an active M dwarf (1 bar atmosphere composed of 90% H_2 and 10% N_2), after 91 h of observation. *Top panel:* vertical axis shows transit depth of the simulated atmosphere spectra in units of ppm (y axis) and horizontal axis showing wavelength in microns. The orange curve corresponds to the simulated atmosphere spectrum without PH_3 , and the blue curve to an atmosphere spectrum with PH_3 , simulated considering a PH_3 concentration of 220 ppb. Blue error bars correspond to the wavelength-averaged uncertainty within the instrumental waveband; black and gray error bars correspond to the uncertainty of each wavelength bin for atmosphere models with and without PH_3 , respectively. Green and pink shading represent the wavelength coverage of the NIRSpec and MIRI instruments (Bagnasco *et al.*, 2007; Wright *et al.*, 2010). *Middle panel:* vertical axis shows the statistical significance of detection for two different model atmospheres, with PH_3 (blue) and without PH_3 (orange). *Bottom panel:* statistical significance of the detection of PH_3 opacities at each wavelength bin. Vertical axis shows size of the statistical deviation between atmosphere models with and without PH_3 (units of σ-interval). In the middle and bottom panels, the horizontal green line represents the 3-σ statistical significance threshold, and the horizontal axes show the individual wavelength bins (microns). For H_2 -dominated atmospheres, the 4.0–4.8 μm spectral band of PH_3 is the most promising feature for detection in transmission. Color images are available online.

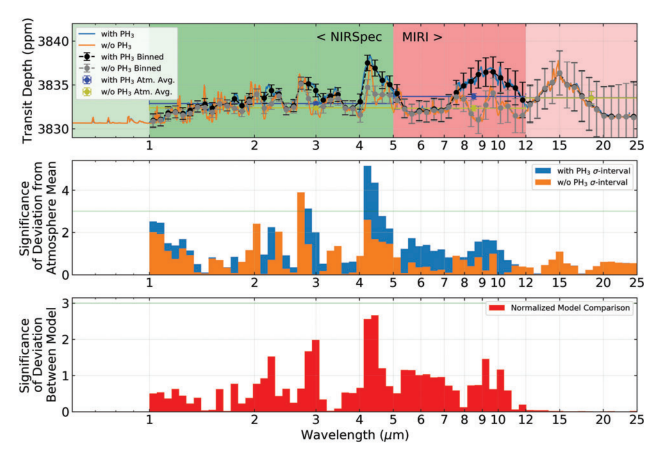


FIG. 4. Theoretical transmission spectra for a CO₂-rich atmosphere on a 10 M_E , 1.75 R_E planet with a surface temperature of 288 K orbiting an active M dwarf (1 bar atmosphere composed of 90% CO₂ and 10% N₂), after 200 h of observation. *Top panel:* vertical axis shows transit depth of the simulated atmosphere spectra in units of ppm (y axis), and horizontal axis showing wavelength in microns. The orange curve corresponds to the simulated atmosphere spectrum without PH₃, and the blue curve to an atmosphere spectrum with PH₃, simulated considering a PH₃ concentration of 310 ppm. Blue error bars correspond to the wavelength-averaged uncertainty within the instrumental waveband; black and gray error bars correspond to the uncertainty of each wavelength bin for atmosphere models with and without PH₃, respectively. Green and pink shading represent the wavelength coverage of the NIRSpec and MIRI instruments (Bagnasco *et al.*, 2007; Wright *et al.*, 2010). *Middle panel:* vertical axis shows the statistical significance of detection for two model atmospheres, with PH₃ (blue) and without PH₃ (orange). *Bottom panel:* statistical significance of the detection of PH₃ opacities at each wavelength bin. Vertical axis shows size of the statistical deviation between atmosphere models with and without PH₃ (units of σ-interval). In the middle and bottom panels, the horizontal green line represents the 3-σ statistical significance threshold, and the horizontal axes show the individual wavelength bins (microns). In CO₂-dominated atmospheres, several spectral features of PH₃ have substantial opacities, but no feature achieves a 3-σ statistical significance when compared with the model atmosphere without PH₃. Color images are available online.

4.1.2. Amount of phosphine required for detection via emission spectroscopy. We now examine the influence of phosphine in the simulated emission spectra of H_2 - and CO_2 -rich planets orbiting Sun-like stars and active M dwarfs.

Our findings on the amount of phosphine required for detection in thermal emission are similar in transmission, that is, PH_3 can only be detected with many tens of hours of observation time (Table 2). We find that, in emission, PH_3 is detectable in anoxic atmospheres only if it accumulates to at least abundances in the order of ppb; for comparison, PH_3 is present at the ppt to ppb level on modern Earth. The photochemical plausibility of PH_3 accumulating to such large abundances is presented in Section 4.1.3.

Detecting an opacity corresponding to an atmospheric spectral feature is much easier than assigning an atmospheric feature to a particular molecule. Our models show that the detection of a CO_2 - or H_2 -rich atmosphere with high statistical significance (>5- σ) is feasible with only a few tens of observation hours. However, the unambiguous attribution of opacity to phosphine requires much longer observation times (Table 2). As an observer, a detection can be considered as an offset to the blackbody curve, but these are only reliable if the blackbody temperature has been accurately estimated. Our detection metric uses a blackbody curve created from a best fit to the simulated observations, which biases toward low temperatures by non-PH $_3$ absorbers. In reality the temperature of the planet may be obtained through other methods, so our results can be considered a conservative estimate for the minimum PH $_3$ abundances required for detection.

We find the most detectable spectral region of phosphine in thermal emission is the broad band at $7.8-11.5\,\mu m$ (Figs. 6 and 7). In emission, planets orbiting an active M

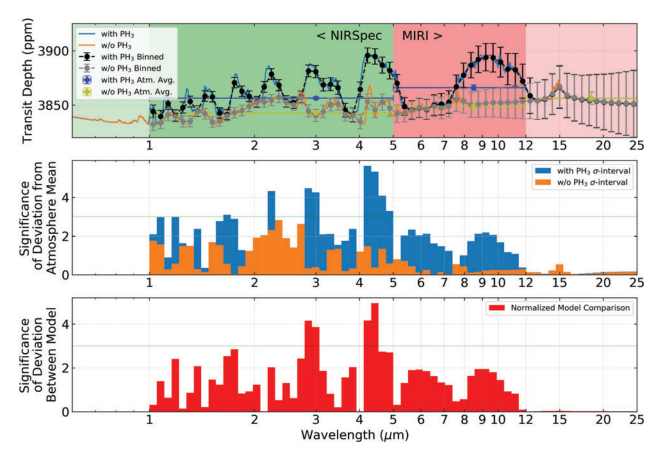


FIG. 5. Theoretical transmission spectra for a H_2 -rich atmosphere on a $10 \, M_E$, $1.75 \, R_E$ planet with a surface temperature of 288 K orbiting an active M dwarf (1 bar atmosphere composed of 90% H_2 and 10% N_2), at the threshold of the phosphine runaway phase. *Top panel:* vertical axis shows transit depth of the simulated atmosphere spectra in units of ppm (y axis), after 3h of observation, and horizontal axis showing wavelength in microns. The orange curve corresponds to an atmosphere spectrum without PH_3 , and the blue curve to an atmosphere spectrum with PH_3 , simulated considering a PH_3 concentration of 0.28%. Blue error bars correspond to the wavelength-averaged uncertainty within the instrumental waveband; black and gray error bars correspond to the uncertainty of each wavelength bin for atmosphere models with and without PH_3 , respectively. Green and pink shading represent the wavelength coverage of the NIRSpec and MIRI instruments (Bagnasco *et al.*, 2007; Wright *et al.*, 2010). *Middle panel:* vertical axis shows the statistical significance of detection for two model atmospheres with PH_3 (blue) and without PH_3 (orange). *Bottom panel:* statistical significance of the detection of PH_3 opacities at each wavelength bin. Vertical axis shows size of the statistical deviation between atmosphere models with and without PH_3 (units of σ -interval). In the middle and bottom panels, the horizontal green line represents the 3- σ statistical significance threshold, and the horizontal axes show the individual wavelength bins (microns). Once PH_3 enters the runaway phase, it can be detected after a few hours of observations, through its two strong features in the 2.7–3.6 and 4– $4.8 \, \mu m$ regions. Color images are available online.

Table 2. Phosphine Mixing Ratios Needed for Detection in Emission for Different Atmospheric and Stellar Scenarios, with Associated Observation and Surface Flux Requirements

Atmospheric scenario	Required mixing ratio for detection	Minimum observation hours (in-transit+out- of-transit)	Associated confidence interval for phosphine detection (σ)
H ₂ -dominated, Sun-like star	N/A	Not detectable	N/A
H ₂ -dominated, active M dwarf (Fig. 6)	220 ppb	131	3
H ₂ -dominated, active M dwarf	4 ppm	52	3
CO ₂ -dominated, Sun-like star	N/A	Not detectable	N/A
CO ₂ -dominated, active M dwarf (Fig. 7)	15 ppm	150	3
CO ₂ -dominated, active M dwarf	310 ppm	48	3

For planets orbiting an active M dwarf, phosphine requires minimum abundances of 220 ppb and 15 ppm to be detectable on H_2 - and CO_2 -rich atmospheres, respectively. For planets orbiting a Sun-like star, no scenario where PH_3 is not a major component of the atmosphere could allow for its detection with less than 200 observation hours.

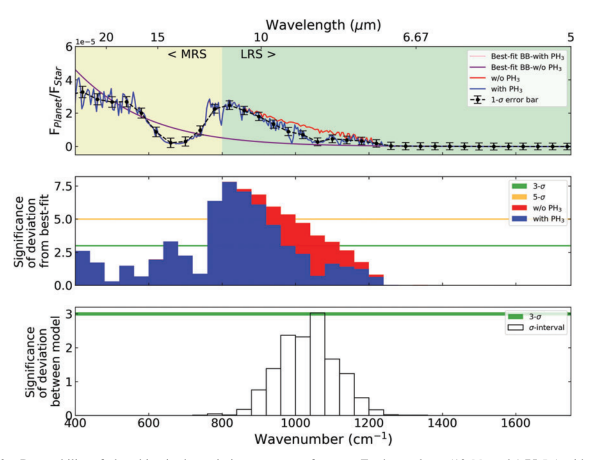


FIG. 6. Detectability of phosphine in the emission spectrum of a super Earth exoplanet (10 M_E and 1.75 R_E) with a H_2 -rich atmosphere orbiting an active M dwarf, after 131 h of observation. Horizontal axes show wavelength in microns (top) and wave numbers in cm⁻¹ (bottom). *Top panel*: vertical axes show the flux ratio between the star and the planet; pink and purple lines represent the blackbody curves fitted to the simulated observational data for atmospheres with and without PH₃, respectively; blue and red curves represent a modeled atmosphere with a PH₃ mixing ratio of 220 ppb and an atmosphere without PH₃, respectively; black error bars represent the 1-σ uncertainty in the observed data; MRS (yellow shading) and LRS (green shading) represent the coverage of the JWST mid- and low-resolution MIRI instruments, respectively. *Middle panel*: statistical significance of the detection of an atmosphere with (blue) and without (red) PH₃ when compared with their best-fit blackbody curves, in units of σ-interval; the horizontal green and orange lines represent the 3-σ and 5-σ statistical significance threshold, respectively. *Bottom panel*: statistical significance of the deviation between an atmospheric model with and without PH₃; the horizontal green line represents the 3-σ statistical significance threshold. The detection of PH₃ achieves a 3-σ confidence interval through the high-frequency wing of its strong broad band at 7.8–11.5 μm. JWST, James Webb Space Telescope. Color images are available online.

dwarf require the smallest PH_3 abundances (100s of ppb to 100s of ppm) to confirm its detection, achieving a 3- σ confidence interval with a minimum of 52 and 48 h of observation, for H_2 - and CO_2 -rich atmospheres, respectively.

Detection of phosphine, in emission, on planets orbiting Sun-like stars is difficult. In these scenarios, the detection of any modeled super-Earth atmosphere cannot achieve a $3-\sigma$ confidence interval even with 200 observation hours.

We note that, at sufficiently high phosphine concentrations, our model shows that the wings of the PH_3 absorption features become opaque (e.g.), the strong broad band at $7.8-11 \, \mu m$) and our emission spectra probe the isothermal stratosphere. Consequently, if PH_3 concentrations are high enough, our models show that it is not possible to detect wavelength-dependent opacities due to PH_3 on the basis of emission data alone. At face value, this observation implies a maximum PH_3 concentration and flux, past which it is impossible to detect PH_3 in emission. However, in reality this effect is an

artifact of our assumption of an isothermal stratosphere. The stratosphere may have temperature variations, which would facilitate the detection of wavelength-dependent opacity variations due to PH₃. A coupled climate-photochemistry model that can provide self-consistent temperature-pressure profiles is required to probe this scenario.

4.1.3. Phosphine surface fluxes required for detection. More critical than atmosphere abundances is the surface flux (*i.e.*, the biological production rate) required for phosphine to accumulate to detectable abundance levels in anoxic atmospheres. This quantity plays a key role in determining the efficacy of PH₃ as a biosignature: if the presence of detectable levels of PH₃ in an atmosphere requires surface fluxes of PH₃ that are higher than those which a biosphere could plausible generate, then it is disfavored as a biosignature gas; if, however, PH₃ accumulates to detectable concentrations at fluxes within the range of

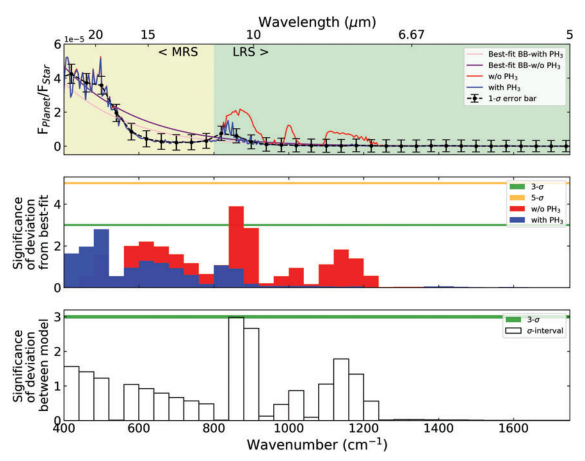


FIG. 7. Detectability of phosphine in the emission spectrum of a super Earth exoplanet (10 M_E and 1.75 R_E) with a CO₂-rich atmosphere orbiting an active M dwarf, after 48 h of observation. Horizontal axes show wavelength in microns (top) and wave numbers in cm⁻¹ (bottom). *Top panel:* vertical axes show the flux ratio between the star and the planet; pink and purple lines represent the blackbody curves fitted to the simulated observational data for an atmosphere with and without PH₃, respectively; blue and red curves represent a modeled atmosphere with a PH₃ mixing ratio of 310 ppm and an atmosphere without PH₃, respectively; black error bars represent the 1-σ uncertainty in the observed data; MRS (yellow shading) and LRS (green shading) represent the coverage of the JWST mid- and low-resolution MIRI instruments, respectively. *Middle panel:* statistical significance of the detection of an atmosphere with (blue) and without (red) PH₃ when compared with their best-fit blackbody curves, in units of σ-interval; the horizontal green and orange lines represent the 3-σ and 5-σ statistical significance threshold, respectively. *Bottom panel:* statistical significance of the deviation between an atmospheric model with and without PH₃; the horizontal green line represents the 3-σ statistical significance threshold. The detection of PH₃ achieves a 3-σ confidence interval through its strong broad band at 7.8–11.5 μm. Color images are available online.

plausible biological productivity, it is favored as a biosignature gas.

As phosphine moves up the atmosphere, its destruction rate and consequent mixing ratio change, due to the varying levels of radical concentrations and radiation at different altitudes. The dominant PH₃ reaction in H₂-dominated atmospheres is PH₃ + H. The dominant reaction in CO₂-dominated atmospheres is PH₃ + O. However, in high-PH₃ atmospheres, H produced from PH₃ photolysis becomes an increasingly important sink for PH₃, even in CO₂-dominated atmospheres. PH₃ is unlikely to dissolve into water and condense into aerosols (as ammonia, hydrogen sulfide, and methanethiol are) (Glindemann *et al.*, 2003), meaning rainout is not expected to be a sink.

We use our photochemical model to estimate the minimum surface production flux, P_{PH3}, for the detectability of phosphine in transmission and emission for a range of planetary

scenarios (Table 3). We find that, for planets orbiting active M dwarfs, PH_3 can build to concentrations detectable by transmission and emission spectroscopy if produced at the surface with rates of the order of 10^{11} cm⁻² s⁻¹. We note that PH_3 requires similar surface flux rates in H_2 - and CO_2 -dominant atmospheres to reach detectable abundance levels, even though those correspond to much lower PH_3 concentration requirements in H_2 -rich atmospheres than in CO_2 -rich atmospheres. We speculate that this occurs because UV penetrates deeper into the more transparent H_2 -rich atmosphere, allowing more radical accumulation and more photolysis at depth [see Fig. 4 of Hu *et al.* (2012)].

The phosphine surface fluxes required to generate the detectable levels of PH₃ are large when compared with global PH₃ emissions on Earth but are comparable to the production rates of other major biosignature gases. For comparison, biological CH₄ and isoprene production on

Table 3. Phosphine Mixing Ratios Needed for Detection in Transmission and Emission for Different Atmospheric and Stellar Scenarios, As Well As Associated Surface Flux Requirements $(P_{PH_3} \text{ [cm}^{-2} \text{ s}^{-1}])$

Atmospheric scenario	Required mixing ratio for detection (in transmission and emission)	$P_{PH_3} \over (cm^{-2} s^{-1})$
H ₂ -rich planet,	780 ppm	1×10^{14}
Sun-like star	(transmission)	10
H ₂ -rich planet, active	5 ppb	1×10^{10}
M dwarf	(transmission)	
H ₂ -rich planet, active	220 ppb	1×10^{11}
M dwarf	(emission)	
H ₂ -rich planet, active	0.28%	9×10^{11}
M dwarf	(transmission)	
(PH ₃ runaway)		
CO ₂ -rich planet, active	310 ppm	3×10^{11}
M dwarf	(transmission)	
CO ₂ -rich planet, active	15 ppm	1×10^{11}
M dwarf	(emission)	
CO ₂ -rich planet, active	7.6%	1×10^{12}
M dwarf	(transmission)	
(PH ₃ runaway)	,	

The values in bold represent surface fluxes and associated atmospheric abundances where phosphine would be able to approach detection but would require longer than 200 hours of observation (which is longer than our allowed limit for detectability). Values in italics correspond to atmospheric scenarios where PH₃ is at a runaway threshold (see Fig. 5 and Sections 4.1.3 and 5.1). For comparison, the maximum recorded surface flux of PH₃ on Earth is $10^{14}~\rm cm^{-2}~s^{-1}$ (above sewage plants, Devai *et al.*, 1988), and the biological production of CH₄ on Earth corresponds to $1.2 \times 10^{11}~\rm cm^{-2}~s^{-1}$ (Houghton 1995; Segura *et al.*, 2005; Guzmán-Marmolejo and Segura, 2015).

Earth are of the order of 10^{11} cm⁻² s⁻¹ (Guenther *et al.*, 2006), where a significant proportion of modern terrestrial CH₄ production is anthropogenic (Houghton, 1995; Segura *et al.*, 2005; Guzmán-Marmolejo and Segura, 2015). As a further comparison, the highest recorded surface flux of PH₃ on Earth is above sewage plants, where PH₃ production reaches 10^{14} cm⁻² s⁻¹ (Devai *et al.*, 1988).

One of our most interesting findings is the existence of a critical phosphine surface production flux, past which PH₃ accumulation is efficient and the atmosphere transitions to a PH₃-rich state. We term this critical flux the "tipping point." This effect appears analogous to the "CO runaway" effect identified for early Earth (Kasting *et al.*, 1983, 1984, 2014; Zahnle, 1986; Kasting, 2014). Past the tipping point, PH₃ production outpaces the ability of stellar NUV photons to destroy PH₃, whether via direct photolysis or via generation of radical species. In this runaway phase, PH₃ can accumulate to percent levels and pervade the atmosphere. In this case, our models show that PH₃ can be detected with observation times reaching under 10 h (Fig. 5). The plausibility of such a PH₃ runaway effect is discussed in Section 5.1.

4.1.4. Sensitivity analysis to temperature and radiation levels. Our approach prescribes a temperature–pressure profile and considers only two possible stellar scenarios (Sun-like stars and active M dwarfs). We conducted sensi-

tivity analyses to assess the dependence of our results on these assumptions.

4.1.4.1. Sensitivity analysis to temperature. In our study, we assumed surface temperatures of 288 K; in reality, worlds with a broad range of temperatures may be habitable. Temperature may affect phosphine concentrations through varying reaction rates of PH₃ with radicals and through changing the concentration of H₂O in the atmosphere, from which the radical species are largely derived. To test the sensitivity of our results to surface temperature, we calculated PH₃ profiles for CO₂- and H₂-rich atmospheres which have detectable concentrations of PH₃ at 288 K, for new surface temperatures of 273 and 303 K. For simplicity, in calculating the dry adiabatic evolution of the lower atmosphere, we approximated the specific heat capacities at constant pressure of CO₂ and H₂, by their values at 273 K and 303 K (Pierrehumbert, 2010). We adjusted the surface mixing ratio of water vapor to 0.0036 and 0.026, corresponding to the vapor pressures at 273 and 303 K, respectively, for the same 60% humidity assumed at 288 K. Figures 8 and 9 present the results of these sensitivity tests in the case of a H₂-dominated atmosphere orbiting an M dwarf star.

We find phosphine abundances to be weakly sensitive to surface temperature. For both CO_2 - and H_2 -dominated atmospheric scenarios, the total PH_3 column varies by a factor of ≤ 3 relative to the value at 288 K across 273–303 K, with the strongest variation occurring in H_2 -dominated atmospheres. Such variations, while potentially significant for retrievals, do not affect our order-of-magnitude conclusions regarding the detectability of PH_3 .

We attribute this relatively modest variation of phosphine column with temperature to the comparatively small variation of both the PH₃ radical reaction rates and the total water vapor column across this temperature range. Other atmospheric constituents, such as methane, show a much greater sensitivity to lower temperatures than PH₃. We are unsure why this is the case. One possibility is that PH₃ reaction rates are less sensitive to temperature changes than other atmospheric constituents (e.g., from 288 to 303 K, the rate constant for H + CH₄ increases by a factor of 2.3, whereas the rate constant for $H + PH_3$ increases by a factor of 1.16). Another possible explanation for CH₄ having a greater sensitivity to temperature than PH₃ is that CH₄ is primarily removed by OH (and therefore most sensitive to H₂O), whereas PH₃ is primarily removed by O and H (i.e., less sensitive to H₂O).

We crudely considered the potential impact of high phosphine abundances on the temperature profile of a planet. The greenhouse gas potential of PH₃ is not known (Bera *et al.*, 2009), but it is plausible that a significant accumulation of phosphine on an atmosphere would contribute to an increase in the global temperature since PH₃ is a strong infrared absorber.

To first order, the change in surface temperature due to phosphine can be estimated by calculating the surface temperature required to produce enough outgoing radiation to balance the arriving stellar radiation (Pierrehumbert, 2010). We executed this procedure for an atmosphere with and without PH₃. We estimate that, if PH₃ accumulates to the abundances required for its detection (see Section 4.1.3), PH₃ can lead to an increase of surface temperature between 10

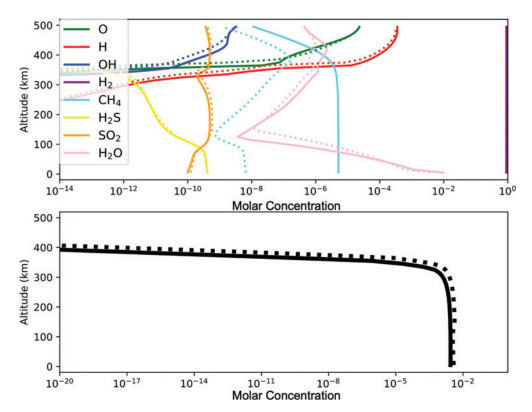


FIG. 8. Distribution of abundances of atmospheric constituents (top panel) and phosphine (bottom panel) throughout the atmosphere of a H₂-rich planet orbiting an active M dwarf. Vertical axes represent altitude in units of km, and horizontal axes represent molar concentration. Solid lines and dotted lines correspond to mixing ratios with surface temperatures of 288 and 273 K, respectively. When comparing low temperatures (273 K) to our standard 288 K models, the PH₃ mixing ratio remains mostly unchanged. Color images are available online.

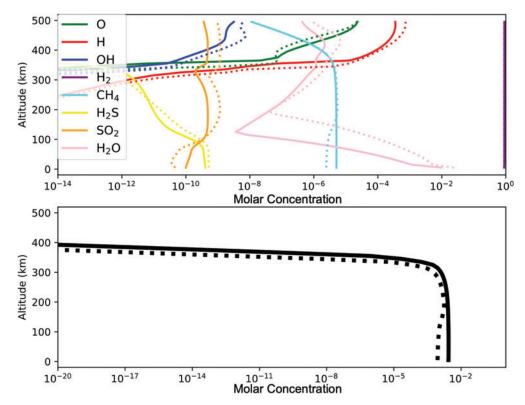


FIG. 9. Distribution of abundances of atmospheric constituents (top panel) and phosphine (bottom panel) throughout the atmosphere of a H₂-rich planet orbiting an active M dwarf. Vertical axes represent altitude in units of km, and horizontal axes represent molar concentration. Solid lines and dotted lines correspond to mixing ratios with surface temperatures of 288 and 303 K, respectively. When comparing high temperatures (303 K) to our standard 288 K models, the PH₃ mixing ratio remains mostly unchanged. Color images are available online.

and 30 K, depending on the atmospheric scenario. Further studies on the greenhouse gas potential of PH₃ are needed to fully explore the impact of its accumulation on the temperature profile of exoplanet atmospheres.

Overall, we conclude that our results are insensitive to variations in surface temperature of ± 15 K.

4.1.4.2. Sensitivity to UV irradiation. UV irradiation limits phosphine concentrations through direct photolysis and radical production. We considered the hypothesis that PH_3 would build to higher concentrations on a planet orbiting a star with low UV output, such as a quiet M dwarf, as considered by Domagal-Goldman et al. (2011). To test this hypothesis, we simulated CO_2 - and H_2 -rich planets orbiting a theoretical "quiet" M dwarf.

We constructed our quiet M dwarf model by reducing the instellation at wavelengths <300 nm of our active M dwarf case (corresponding to GJ1214) by a factor of 1000. This corresponds to ~100 times less UV than GJ581, the quietest M dwarf observed by the MUSCLES survey (France *et al.*, 2016). A truly quiet M dwarf may not exist, as practically all M dwarfs observed to date have at least some chromospheric activity (France *et al.*, 2013, 2016). Our quiet M dwarf case may therefore be considered a theoretical limiting case to study the effect of UV radiation on PH₃ buildup, with the understanding that this limiting case may not exist in reality. We nonetheless note that this limiting case is less extreme than photosphere-only limiting cases considered in past work (Domagal-Goldman *et al.*, 2011; Seager *et al.*, 2013b; Rugheimer *et al.*, 2015).

We find that, for the equivalent surface production rates, phosphine concentrations are two orders of magnitude higher on planets in the quiet M dwarf cases compared with the active M dwarf cases. Low UV emission favors buildup of PH₃ due to lower radical concentrations and photolysis rates. Consequently, as with other proposed biosignature gases, planets orbiting quiet M dwarfs are the best targets for detecting biogenic PH₃ (Segura *et al.*, 2005; Domagal-Goldman *et al.*, 2011; Seager *et al.*, 2013b). We also find that, in planets orbiting a quiet M dwarf, PH₃ is able to enter a runaway phase with two orders of magnitude lower surface fluxes than those required in more active stars (Section 5.1).

Our overall main finding is that, because phosphine is easily destroyed either directly by UV or indirectly by UVmediated creation of H, O, or OH radicals, a UV-poor environment is favorable for the detection of PH₃. This result is consistent with past work (Segura et al., 2005; Domagal-Goldman et al., 2011; Seager et al., 2013b). If there are no sufficiently quiet M dwarf stars, we speculate that a UVpoor environment can be created by a UV shield on the planet itself (Wolf and Toon, 2010). Since PH₃ is readily destroyed in an O₂-rich environment, an ozone UV shield is unsuitable because other oxygen-containing radicals would destroy PH3. However, elemental sulfur aerosols generated on planets with high volcanism and reducing atmospheres may provide such a UV shield (Hu et al., 2013). Additionally, if PH₃ fluxes are high enough, they can overwhelm the supply of destructive UV photons and build up to higher concentrations (a runaway effect). For more context and for a comparison with CH₃Cl, another proposed biosignature gas, see Sections 5.1 and 5.3.

4.2. Phosphine spectral distinguishability

Phosphine's spectral features can be easily distinguished from those of other gases expected to be main components of rocky planet atmospheres. Such gases include water vapor, methane, carbon dioxide, carbon monoxide, ammonia, and hydrogen sulfide (Fig. 10). Ammonia might be present in hydrogen-rich atmospheres (Seager *et al.*, 2013b).

The infrared spectrum of phosphine has three major features: 2.7-3.6, 4-4.8, and 7.8-11 µm, corresponding to polyad (P) numbers 3, 2, and 1, respectively (Sousa-Silva et al., 2013). The 2.7–3.6 μ m region (P=3) is dominated by a hot band and an overtone band, both associated with the symmetric bending mode of PH₃, and six additional combination bands. The 4–4.8 μ m region (P=2) is dominated by both the fundamental symmetric and asymmetric stretching bands of the PH₃ molecule. The P=2 feature is also where several weaker, bending overtone and combination bands occur, combining with the fundamental bands to result in the strongest overall spectral feature for PH3. When compared with water, methane, ammonia, and hydrogen sulfide (but not CO_2), the P=2 feature is uniquely attributable to PH_3 (Fig. 10). In the 7.8–11 μ m region (P=1), the fundamental symmetric and asymmetric bending modes, as well as hot bands, combine to produce a strong and broad absorption feature. The P=1 PH₃ feature overlaps with the ammonia spectrum but is easily distinguishable from the remaining molecules in this comparison (Fig. 10).

For more detailed comparisons focusing on the 2.7-5 and $7.8-11.5 \,\mu m$ regions, see Appendix B.

The strongest band of phosphine, in the 4–4.8 μ m region, is particularly salient when comparing PH₃ with all available spectra of volatile molecules (Fig. 11). However, the second strongest feature of PH₃, the broad band in the 7.8–11 μ m region, is easily obscured by other gases as it absorbs in a heavily populated wavelength region, where many molecules have strong fundamental rovibrational modes.

It is worth noting that, of the 534 molecules for which there are available spectra, only a few dozen have been adequately measured or calculated, and consequently their spectra should be considered preliminary. Furthermore, there are thousands of volatile molecules that could contribute to an atmospheric spectra (Seager *et al.*, 2016) for which there is no available spectral data, so further studies are required to reveal the full extent of the spectral comparison highlighted in Fig. 11 (Sousa-Silva *et al.*, 2018, 2019).

4.3. Phosphine false positives

On Earth, the only significant amounts of phosphine found in the atmosphere are produced anthropogenically or biologically (Sections 2.1 and 2.2). The formation of PH₃ on temperate rocky planets is thermodynamically disfavored, even in high-reducing environments, unlike the abiological production of methane or hydrogen sulfide.

In thermodynamic equilibrium, phosphorus can be conservatively expected to be found in the form of phosphine only at T>800K, and at P>0.1 bar (Visscher *et al.*, 2006), and only in environments rich in H_2 , which is why PH_3 has been detected in Jupiter and Saturn, where these extreme temperatures occur (in the deep layers of the atmosphere). In H_2 -poor environments, much higher temperatures and pressures are required for PH_3 to be thermodynamically

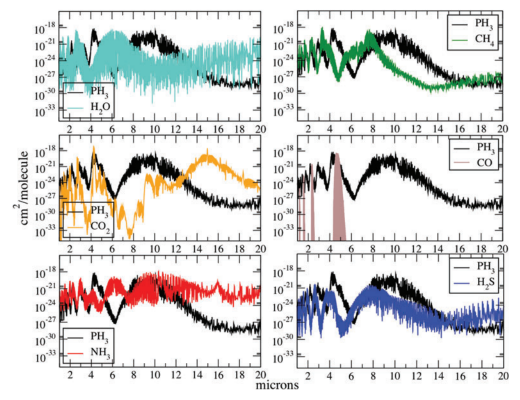


FIG. 10. Comparison of the spectral cross sections of phosphine with other molecular gases at room temperature. Intensity on y axes in a log-scale with units of cm²/molecule, and wavelength represented on the x axes in microns. All cross sections are calculated at zero-pressure (*i.e.*, Doppler-broadened lines only) using the procedure described by Hill *et al.* (2013). PH₃, shown in black, is distinguishable from all compared molecules due to its strong bands in the 2.7–3.6, 4–4.8, and 7.8–11 μ m regions. Color images are available online.

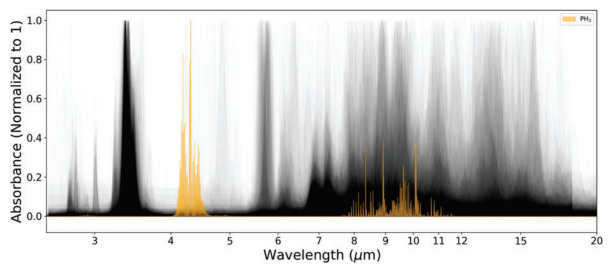


FIG. 11. Comparison of the spectral cross sections of phosphine (orange) with all the available cross sections for molecules that are volatile at room temperature (Lemmon *et al.*, 2010). Intensity on *y* axes in a linear scale representing absorbance (normalized to 1), and wavelength represented on the *x* axes in microns, with the spectral range constrained to 2.5–18.5 μm for fair comparison (many molecules have incomplete spectra beyond this region). Opacity for all molecules is plotted at 1% so that heavily populated regions are highlighted. All cross sections are calculated with SEAS, using molecular inputs from NIST (Linstrom and Mallard, 2001) and ExoMol (Tennyson *et al.*, 2016). The strongest band of PH₃ (4.0–4.8 μm) is easily distinguishable from all other gases, but the broad band at 10 μm can become obscured by other molecules. SEAS, Simulated Exoplanet Atmosphere Spectra. Color images are available online.

favored. We also note that the critical temperature of water is 647~K so there are no surface conditions that favor both PH_3 production and allow for the presence of liquid water. Consequently, in a temperate rocky planet, it is implausible that PH_3 can be produced without biological intervention, so its detection in such an environment is a promising indication of biological activity. We summarize below the potential false-positive scenarios for PH_3 as a biosignature gas and their expected impact on the global concentrations of PH_3 .

Overall, nonbiological phosphine formation is not favored on temperate rocky worlds, and no abiotic pathways can produce PH₃ with the production rates necessary for its detection on habitable exoplanets. We therefore conclude that, in contrast to molecules such as ammonia and methane, a detection of PH₃ on a temperate exoplanet is likely to only be explained by the presence of life.

4.3.1. Formation from phosphite and phosphate. We considered the hypothesis that phosphine could be formed geochemically as a "false positive" by reduction of phosphate or phosphite to PH₃. Phosphate is a dominant form of phosphorus on Earth. Phosphite is much less abundant but was detected in ground water and in mineral deposits (Han et al., 2012, 2013; Yu et al., 2015) where it is likely to be the result of biological activity (Bains et al., 2019a). Phosphite is also postulated to have been much more abundant on early anoxic Earth (Pasek, 2008; Pasek et al., 2013; Herschy et al., 2018). We calculated the Gibbs free energy of formation of PH3 from both phosphate and phosphite under geochemical source conditions at neutral pH, for T = 273 and 413 K and pH₂ = 10^{-6} and 1 bar. In all cases, the formation of PH3 was thermodynamically disfavored; see Appendices C and D for details. We conclude that PH₃ formation from phosphate or phosphite is unlikely in the absence of a biological catalyst; for more details on the thermodynamic plausibility of the reduction of phosphites or phosphate into PH₃, see Bains et al. (2019a).

Phosphite can disproportionate to phosphine at $T \gtrsim 320$ K and acidic pH (Bains et al., 2019a), raising the possibility that "black smoker" hydrothermal systems ($T \le 678$ K, pH=2-3) (Martin et al., 2008) might generate phosphine (Appendix D). Such systems do not dominate volcanic emission on Earth, leading us to propose they would be a negligible contributor on Earth-analog worlds. However, if a world had global, hot, acidic oceans (e.g., due to very high pCO₂), then the theoretical possibility of abiotic phosphine production exists, although likely only in the presence of high H₂ concentrations, very low pH and within a very hot temperature band (Appendix D) (Bains et al., 2019a). Given that these oceans would be unlikely to have pH values below 4 (carbonic acid has a pH of 3.6) and PH₃ formation is only favored at pHs closer to 2, we consider this scenario possible but implausible.

4.3.2. Lightning. We also considered the possibility of phosphine production by lightning. Lightning discharges even in highly reducing atmospheres produce only negligible amounts of reduced phosphorus species, including PH₃, and are very unlikely to provide high flux sources of PH₃ globally. A few studies have examined the production of reduced phosphorus species from phosphate as a result of simulated lightning discharges in laboratory conditions

(Glindemann *et al.*, 1999., 2004); only a very small fraction of the phosphorus was reduced to PH₃ through this process, even in highly reduced atmospheric conditions. Similarly, a mineral fulgurite—a glass resulting from lightning strikes—was also proposed as a potential source of PH₃ given that it could, in principle, contain reduced phosphorus species (Pasek and Block, 2009). However, these sources are rare and localized; they would have minimal impact on a global scale. We are not aware of kinetically favored reactions that would promote the conversion of the thermodynamically favored phosphate to PH₃.

4.3.3. Volcanism. Phosphine is not known to be produced by volcanoes on Earth.

Calculations on the production of phosphine through volcanism on a simulated anoxic early Earth showed that only trace amounts of PH₃ can be created through this avenue; the predicted maximum production rate is 102 tons per year (Holland, 1984), which corresponds to $\sim 10^4~\rm cm^{-2}~s^{-1}$. We note that the estimation of the maximum production of PH₃ through the volcanic processes reported by Holland (1984) is made under the assumption of a highly reduced planet, which provides favorable conditions for PH₃ volcanic production. The volcanic production of PH₃ in other planetary scenarios is even more unlikely. We estimate that the maximum production of PH₃ by volcanoes in any planetary scenario, even H₂-rich atmospheres, is at least seven orders of magnitude lower than the surface fluxes required for detection (see Section 4.1.3).

4.3.4. Exogenous delivery. Finally, we considered the possibility of exogenous meteoritic delivery as a source of reduced phosphorus species that could lead to the abiotic production of phosphine. Reduced phosphorus species can be found in the meteoritic mineral schreibersite (Pech et al., 2011). Schreibersite is (Fe,Ni)₃P, which is present in iron/ nickel meteorites (Geist et al., 2005); it is not present in stony or carbonaceous bodies. The current accretion rate of meteoritic material to Earth is of the order of 20-70 kilotons per year (Peucker-Ehrenbrink, 1996). We calculated the maximum PH₃ production from these sources as follows: considering that $\sim 6\%$ of the meteoritic material is iron/ nickel (Emiliani, 1992) and such meteorites contain an average of 0.25% phosphorus by weight (Geist et al., 2005), and working under the conservative assumption that the totality of the phosphorus content could be hydrolyzed to PH₃, these meteors would deliver a maximum of ~ 10 tons of PH₃ to Earth every year.** Therefore, the contribution from meteoritic sources to the global average PH₃ production rates is still negligible. The above calculations are also in agreement with previous estimations of the phosphine production through meteoritic delivery, which were also found to be negligible (Holland, 1984).

5. Discussion

We find that phosphine is a promising marker for life if detected on a rocky exoplanet. On Earth, PH₃ is naturally

^{**}For comparison, 10.2 million tons per year of methane are produced from ruminants alone (Moss *et al.* 2000).

associated exclusively with anaerobic life and is not expected to have any significant false positives for life on temperate exoplanets. Our models find that, if produced at sufficiently high surface fluxes, PH₃ can accumulate in planetary atmospheres to detectable abundances. Here, we discuss the photochemical impact of high abundances of PH₃ in an exoplanet atmosphere (Section 5.1). We then describe the known limitations of our calculations (Section 5.2) and expand on alternative methods of detecting PH₃ in exoplanet atmospheres (Section 5.3). We summarize our findings in section 5.4.

5.1. Phosphine "tipping point" and its impact on the atmosphere

Our models show that, with global phosphine surface fluxes comparable to those found locally in anoxic ecosystems on Earth, PH₃ can have a significant impact on planets with anoxic atmospheres. We calculate that PH₃ becomes detectable on anoxic planets where it is emitted at the surface with fluxes greater than $10^{11}~\rm cm^{-2}~\rm s^{-1}$. Consequently, PH₃ is accessible to remote detection only if it is a substantial product of the biosphere, emitted in quantities similar to CH₄ and isoprene on Earth (Guenther *et al.*, 2006). We note that the maximum recorded surface flux of PH₃ on Earth is $10^{14}~\rm cm^{-2}~\rm s^{-1}$ (Devai *et al.*, 1988). PH₃ may be emitted by life at these detectable surface fluxes as a product of primary

metabolism, such as CH₄, on warm acidic worlds, or as a secondary metabolite, such as isoprene (Bains *et al.*, 2019a, 2019b).

A surprising result of our models was that, once the phosphine surface flux reaches a *tipping point* (*e.g.*, $>9 \times 10^{11}$ cm⁻² s⁻¹ for planets orbiting active M dwarfs), PH₃ enters a runaway phase and begins to drastically change the atmosphere (Fig. 5). This phase appears analogous to the CO runaway described for early Earth and reviewed in the work of Kasting *et al.* (2014). In this runaway phase, PH₃ production outpaces the ability of stellar NUV photons to destroy PH₃ (whether via direct photolysis or via generation of radical species), and modest increases in PH₃ flux lead to dramatic increases in PH₃ accumulation in an atmosphere.

With fluxes of 10¹² cm⁻² s⁻¹ (10 times higher than the minimum required for detection), phosphine can approach percent concentrations and would be detectable with just less than 10 h of observation (Table 1 and Fig. 12). In this runaway phase, PH₃ can affect the concentrations of other atmospheric constituents; for example, in CO₂-dominated atmospheres, the H₂ concentration increases dramatically in PH₃ runaway, presumably from H₂ generated by PH₃ destruction. This raises the possibility that enhanced H₂ concentrations may be used to confirm PH₃ detections. In summary, PH₃ is readily detectable if it is produced at rates about an order of magnitude higher than methane or isoprene on Earth.

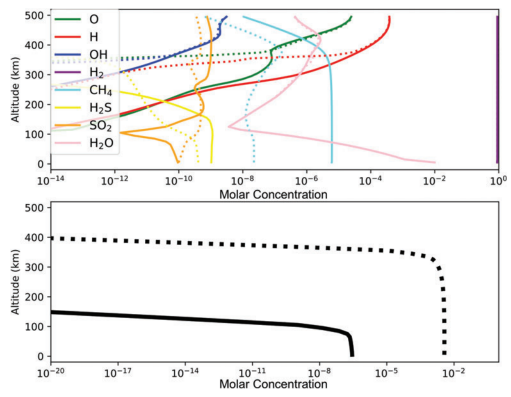


FIG. 12. Distribution of abundances of major atmospheric constituents (top panel) and phosphine (bottom panel) throughout the atmosphere of a H_2 -rich planet orbiting an active M dwarf. Solid and dashed lines show molecular abundances immediately below and above the PH_3 runaway phase, respectively. The x axis shows abundance concentrations, and the y axis shows altitude in units of km. The scavenging effect of PH_3 leads to a decrease in O and OH and, to some extent, H radicals in the atmosphere. Consequently, both PH_3 and other trace gases (e.g., H_2S) are able to accumulate to larger abundances once PH_3 enters the runaway phase. Color images are available online.

Phosphine can affect the spectrum of a rocky planet atmosphere, even at concentrations somewhat below detectability, by driving down radical concentrations due to its intense reactivity with these molecules, effectively becoming a scavenger in the atmosphere. This affects the concentrations of other atmospheric constituents, such as methane. Domagal-Goldman et al. (2011) described a comparable effect in models of organosulfur volatiles, which detectably altered the CH₄ and C₂H₆ abundances despite being themselves undetectable. Domagal-Goldman et al. (2011) reported that elevated C₂H₆/ CH₄ ratios could be diagnostic of high organosulfur flux, and hence a biosignature. Similarly, it may be possible to use the indirect effects of high PH₃ flux to infer the presence of PH₃ even if it is not directly detectable; more detailed measurements of the reaction kinematics of PH₃ and its by-products, as well as more sophisticated atmospheric modeling are required to explore this possibility.

5.2. Model limitations and assumptions

We have assessed the potential of phosphine as a biosignature gas using a set of sophisticated photochemical and radiative transfer models. Nonetheless, given the complexity of simulating both atmospheric composition and subsequent spectral observations, many approximations and assumptions were made. Below is a brief discussion of the major limitations of the work performed here.

5.2.1. Clouds. We assumed cloudless skies for the simulation of exoplanet transmission spectra. To estimate how clouds might affect our results, we re-ran our models for phosphine abundances in the detectable range considering cloud decks at various altitudes, with coverage ranging from 10% to 100%. As expected, the detectability of PH₃ is reduced with the introduction of cloud coverage; for example, for planets with a H₂-dominated atmosphere orbiting an active M dwarf, and with PH₃ surface fluxes of 10¹¹ cm⁻² s⁻¹, the detectability of PH₃ reduced by up to a factor of 10 (from a cloudless model to a model with full cloud coverage at 100 Pa). However, once PH₃ reaches the tipping point (see Section 5.1), it becomes sufficiently abundant in the upper troposphere to be mostly unaffected by the presence of clouds.

5.2.2. Reaction networks and haze formation. In this work, we have focused on the reactivity of phosphine with the dominant radical species O, H, and OH. The photochemistry of PH₃ with radicals originating from other more exotic atmospheric species, although likely to be rare, is insufficiently studied. For example, photochemistry of PH₃ and hydrocarbons, through UV radiation, could lead to the formation of complicated alkyl-phosphines (Guillemin et al., 1995, 1997), and in consequence increase the probability of hazes. Inclusion of these reactions would increase PH₃ destruction rates and hence increase the required surface fluxes for detection; however, since these species are not expected to be dominant radicals, the effect of their inclusion would be minor and should not affect our results. In contrast to the formation of hydrocarbon and sulfur-based hazes that have previously been thoroughly addressed (Domagal-Goldman et al., 2011; Arney et al., 2017), there is very little work on phosphorus-based hazes. The formation of such hazes is possible in theory (Guillemin *et al.*, 1995, 1997; Pasek *et al.*, 2011), and early laboratory experiments implied the possibility of formation of such organophosphine hazes in planetary atmospheres, but further studies are needed to properly address the plausibility and impact of organophosphine haze formation and its associated potential as a PH₃ sink. Our photochemical model will continue to update whenever we are able to expand our reaction networks.

We have neglected the formation of organic hazes in this work. There is evidence that such hydrocarbon hazes occurred on Earth, due to transient high levels of methane (Izon et al., 2017; Zahnle et al., 2019). Organic hazes are predicted to form at $[CH_4]/[CO_2] > 0.12$ for M dwarf stars, and $[CH_4]/[CO_2] \ge 0.2$ for Sun-like stars (Arney et al., 2016, 2017). A methanogenic biosphere producing high CH_4 fluxes is required to generate such high ratios; in our work, we have not considered such a biosphere. The net effect of hazes would be to facilitate phosphine buildup through attenuation of photolytic UV. However, these same hydrocarbon hazes would also cloak some of the PH_3 spectral features, although primarily not in the wavelength bands where PH_3 is a strong absorber.

We did not include in our models the recombination of phosphine via $PH_2 + H \rightarrow PH_3$. The rate constant for this reaction is 1.1×10^{-10} cm³ s⁻¹ at 288 K (Kaye and Strobel, 1984). The rate constant for H attack on PH_3 is 3.3×10^{-12} cm³ s⁻¹ at 288 K (Arthur and Cooper, 1997). Consequently, if $[PH_2]/[PH_3] \geq (3.3 \times 10^{-12}$ cm³ s⁻¹)/ $(1.1 \times 10^{-10}$ cm³ s⁻¹) = 0.03, then reactions with H can reform PH_3 as fast as it is destroyed by H-attack, and substantially lower the PH_3 surface fluxes that are required for PH_3 to accumulate in the atmosphere. Detailed photochemical modeling is required to constrain whether such high $[PH_2]/[PH_3]$ is plausible, or whether other sinks will suppress $[PH_2]$. We note that, in models of Saturn's atmosphere, $[PH_2]/[PH_3] << 0.03$ (Kaye and Strobel, 1984); if the atmospheres of terrestrial H_2 -dominated exoplanets behave similarly, this recombination mechanism will not be able to significantly replenish PH_3 .

Finally, we note that our results reflect the prediction of atmospheric models that anoxic atmospheres should have much higher mean radical concentrations than modern Earth (Hu *et al.*, 2012). Due to their reactivity, one might expect concentrations of radicals to be suppressed even in anoxic atmospheres, as OH is on Earth. While our reaction networks include all known relevant atmospheric chemistry, it is possible that there are chemical reactions, which are relevant to anoxic temperate terrestrial planets, but which have not been considered in the context of the Solar System and hence are not included in the reaction compendia we use (*e.g.*, the NIST database (Linstrom and Mallard, 2001) and the JPL compendium (Sander *et al.*, 2011).

Due to the incompleteness of our reaction network, it is likely that, when phosphine is destroyed, additional radicals are created that are not considered in our models. While PH₃ is a trace gas, this omission should have negligible consequences, as the calculation of the infinite series due to radicals tends to converge quickly. However, when PH₃ approaches the runaway phase and becomes the dominant radical sink the atmosphere, our scenarios become a low-radical regimen. In reality, the radical production would not stop, and the transition to a runaway scenario may be slower

than we predict. If radical concentrations were overestimated in our model, then PH₃ can build to detectable levels with lower surface fluxes. If, at high concentrations of PH₃, the intermediate radical production from its destruction has been underestimated, then PH₃ can only enter a runaway phase with higher surface fluxes than those calculated here. Detailed studies of the chemical reaction networks of anoxic planets are required to explore these possibilities.

5.2.3. Prescribed temperature-pressure profiles. Our photochemical model uses prescribed temperature-pressure profiles (Appendix A) that are isothermal above the stratosphere. In reality, this is an oversimplification; one of the consequences of assuming that there is no temperature inversion at high altitudes is the underestimation of the detectability of phosphine in our simulated emission spectra (Section 4.1.2). We also did not couple the potential heating effect from PH₃ dissociation in our photochemical model, so we performed a sensitivity analysis to small changes in temperature (±15 K) and found that our main conclusions remain unchanged (Section 4.1.4). However, if PH₃ fluxes exceed the "tipping point" and PH3 enters a runaway phase, it may be possible for temperatures to increase beyond the maximum 303 K we consider here. In the most extreme scenario, a PH₃ runaway might trigger a runaway greenhouse state, potentially rendering the planet uninhabitable. A coupled climate-photochemistry model is required to thoroughly investigate this scenario.

5.2.4. Phosphine sinks. Our photochemistry model assumes a dry deposition velocity of zero for phosphine, that is, no consumption of PH₃ by surficial geochemistry or biology. Apart from its efficient oxidation by atmospheric components, there are no other known significant PH₃ sinks on Earth. It is possible, however, that other planets may have PH₃ deposition pathways that we cannot account for. For example, given the opportunistic nature of biology, and the fact that, at least on Earth, phosphorus is a growthlimiting nutrient, life might use any excess atmospheric PH₃ as source of phosphorus. It is plausible that anaerobic life on other planets will not only just produce PH3 but also reabsorb it from the atmosphere; in these scenarios, biology would slow down the PH₃ accumulation in the atmosphere leading to a dampening, and possible avoidance, of a runaway PH₃ effect (Section 5.1). In extreme cases, for example, if biological production of PH₃ can equal its reabsorption, life's recycling of biogenic PH3 might entirely prevent its accumulation in the atmosphere of an exoplanet.

To estimate the effect of a potential sink for phosphine, we tested the variability of our results to a non-zero deposition velocity for PH₃. Deposition velocity tests show that, for a comparable deposition rate to CO and O_2 (10^{-4} cm⁻¹) (Harman *et al.*, 2015), concentrations of PH₃ varied by a factor of <2, which is not enough to affect detectability. We note that this is a conservative estimate given that the only plausible PH₃ sinks are biological.

5.2.5. Phosphorus availability. We considered whether the availability of phosphorus in the crust of a planet could be a limiting factor for the accumulation of phosphine in the atmosphere. An estimate of the total phosphorus within Earth's crust shows that, if all the phosphorus was to be

converted to PH₃, it would produce approximately twice as many PH₃ molecules as the total number of all molecules belonging to all gases present in the atmosphere of the modern Earth (Yaroshevsky, 2006). We conclude that, in principle, the total mass of phosphorus in a planetary crust does not limit the development of a high-PH₃ atmosphere.

5.3. Alternative detection methods beyond JWST

For the phosphine detectability calculations in this work, we have considered observations from a JWST-like telescope, with a 6.5 m diameter telescope mirror operating within 50% of the shot noise limit and a quantum efficiency between 20% and 25%. The integration time is assumed to be under 200h for all atmospheric scenarios. For comparison, the cryogenic lifetime of JWST is 5 years, which is equivalent to an integration time of 100h for a planet orbiting the habitable zone of an M dwarf star. The spectral resolution of JWST is R = 100 at 1 - 5 μ m and R = 160 at 5 - 12 μ m, which is more than necessary for distinguishing between PH₃ and other dominant gases in the atmosphere (see Section 4.2).

We considered the possibility of detecting phosphine in anoxic atmospheres using alternative telescopes to JWST. Missions such as TPF-I (Lawson *et al.*, 2008), Darwin (Fridlund, 2000), Origins Space Telescope (OST) (Battersby *et al.*, 2018), Habitable Exoplanet Imaging Mission (HabEx) (Gaudi *et al.*, 2018), Large UV Optical Infrared Surveyor (LUVOIR) (Roberge, 2019), and the 30-m class of ground telescopes (Johns *et al.*, 2012; Tamai and Spyromilio, 2014; Skidmore *et al.*, 2015) could also be able to characterize atmospheres of temperate planets in wavelength regions where PH₃ is spectrally active.

TPF-I was intended to be a nulling interferometer with four 4 m diameter telescopes flying in formation with a baseline range of 40–100 m and operating at 6.5–18 µm with a spectral resolution of 25-50. Darwin was planned as set of 3–4 m diameter telescopes flying in a nulling interferometer configuration. For both instruments, only the nearest (~ 4 pc) M dwarf star habitable zones would be accessible. Winters et al. (2019) estimated that there are only 22 M dwarf stars candidates that could have planets within their habitable zone suitable for atmospheric characterization. TPF-I and Darwin are currently canceled, but similar telescopes [e.g., the proposed Large Interferometer For Exoplanets, or LIFE (Quanz et al., 2019)] may one day be commissioned that perform in a similar interferometer formation and could have the capability to identify phosphine on temperate exoplanets orbiting M dwarfs.

The OST (Battersby *et al.*, 2018) has a large wavelength coverage $(2.8-20\,\mu\text{m})$ and will be able to provide atmospheric spectra for planets orbiting K and M dwarf stars, through transmission and secondary eclipse observations. The OST will have the sensitivity and coverage to detect many spectral signatures of potential biosignature gases, including phosphine.

The HabEx (Gaudi *et al.*, 2018) and the LUVOIR (Roberge, 2019) are planned as powerful telescopes that could launch in the coming decades and are focused on the detection of potential biosignature gases. Both cover a wavelength region where phosphine is spectrally active (UV to near-infrared), although not a particularly strong absorber (1.28–1.79 µm) (Sousa-Silva *et al.*, 2014). We have not

modeled atmospheres with PH₃ in an HabEx/LUVOIR scenario, but these telescopes' high-contrast spectroscopy may allow for some of the high-frequency combination and hot bands of PH₃ to be detected.

Future 30-m ground-based telescopes will be limited by Earth's observing windows, but those capable of M- and N-band spectroscopy (4.5–5.0 and 7.5–14.5 μ m, respectively) would be favorable for the detection of phosphine, given its strong feature at 4–4.8 μ m and its broad band centered at 10 μ m (see Section 4.2).

5.4. Phosphine as a biosignature gas

An ideal biosignature gas lacks abiotic false positives, has uniquely identifiable spectral features, and is unreactive enough to build up to detectable concentrations in exoplanet atmospheres. Phosphine fulfills the first two criteria: PH₃ is only known to be associated with life and geochemical false positives for PH₃ generation are highly unlikely (Bains et al., 2017, 2019a); PH₃ possesses three strong features in the 2.7–3.6, 4–4.8, and 7.8–11 μ m regions that are distinguishable from common outgassed species that may be present in terrestrial exoplanet atmospheres, such as CO₂, H₂O, CO, CH₄, NH₃, and H₂S.

The greatest challenge to the detectability of phosphine at low surface fluxes is its reactivity to radicals and its vulnerability to UV photolysis. We found that, in the most tractable observational scenario (planet orbiting an active M dwarf), PH₃ must be emitted at a rate of 10¹¹ cm⁻² s⁻¹ to build to levels detectable by transmission or thermal emission spectroscopy (e.g., using JWST). The required PH₃ production rates for detection are two orders of magnitude lower for planets orbiting a hypothetical "quiet" M dwarf with extremely low levels of chromospheric activity; this latter scenario is likely unrealistic and corresponds to an extreme lower limit on the required PH3 flux for its detection. An alternative path for a UV-poor surface environment would be a planet with a UV shield, or possibly intense hazes, although hazes might also inhibit a PH3 detection. Finally, our models suggest that, at high but plausible surface fluxes $(10^{12}-10^{14}~\text{cm}^{-2}~\text{s}^{-1})$, depending on the planetary scenario), PH3 is able to exhaust the supply of M dwarf NUV photons and enters a "runaway" phase. If such a runaway effect occurs, PH₃ becomes easily detectable, protects other trace gases from destruction by radicals, and rapidly changes the overall composition of the planetary atmosphere.

We compare phosphine with another proposed biosignature gas: methyl chloride or CH_3Cl (Segura *et al.*, 2005). The surface flux required for CH_3Cl buildup to detectable levels is 5×10^7 to 3×10^{11} cm⁻² s⁻¹ under the same atmospheric scenarios we consider here, or up to four orders of magnitude less than PH_3 . This difference is rooted in the lower reactivity of CH_3Cl with radicals; its reaction rates with O and H are four to six orders of magnitude lower than PH_3 (Seager *et al.*, 2013b). We conclude that it is more difficult for PH_3 to build to detectable concentrations than CH_3Cl because of its higher reactivity with radicals. As a further comparison, the modern methane surface flux is 535 $Tg/year = 1.2 \times 10^{11}$ cm⁻² s⁻¹, meaning that for PH_3 to build to detectable levels in planets orbiting active stars, it must be emitted at a rate comparable to that of methane on Earth

(Houghton, 1995; Segura et al., 2005; Guzmán-Marmolejo and Segura, 2015).

For a final comparison, we consider the highest concentrated fluxes of phosphine on Earth. On Earth, detections of PH₃ in biogas from sewage plants can reach very high levels, for example, 10^{14} cm⁻² s⁻¹ (Devai *et al.*, 1988). It is therefore plausible that a complex anaerobic biosphere, not dissimilar from those found in sludges on Earth, could achieve global PH₃ production levels comparable to the production rates found locally in isolated environments on Earth. In such anoxic environments, PH₃ would be detectable in transmission and emission for planets orbiting M dwarfs (Table 1). In other words, we can imagine, for example, a planet like Earth in the early Carboniferous period (\sim 318 Mya), but without O₂ in the atmosphere: a wet anoxic "tropical paradise" from pole to pole. Such a planet could potentially produce tremendous amounts of PH₃ by its rich anaerobic biosphere.

Phosphine's relevance as a biosignature gas depends on its production by life, on its possible geochemical false positives, and on its detectability. Prospects for the buildup of PH₃ to detectable levels are uncertain with near-future telescopes. However, the wavelength regions where PH₃ is spectrally active are similar to those of other atmospherically important molecules, for example, H₂O and CH₄. Consequently, searches for PH₃ can be carried out at no additional observational cost to searches for other molecular species relevant to characterizing exoplanet habitability.

Ultimately, if detected on a temperate planet, phosphine is an extremely promising biosignature gas, since its lack of high-flux false positives would be a strong reason to hypothesize production by life.

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Abbreviations Used

ARIEL = Atmospheric Remote-sensing Infrared Exoplanet Large-survey

HabEx = Habitable Exoplanet Imaging Mission

JWST = James Webb Space Telescope

LUVOIR = Large UV Optical Infrared Surveyor

MBP = Matrix-Bound Phosphine

NUV = Near UV

OST = Origins Space Telescope

 $PH_3 = phosphine \\$

SEAS = Simulated Exoplanet Atmosphere Spectra

UV = ultraviolet

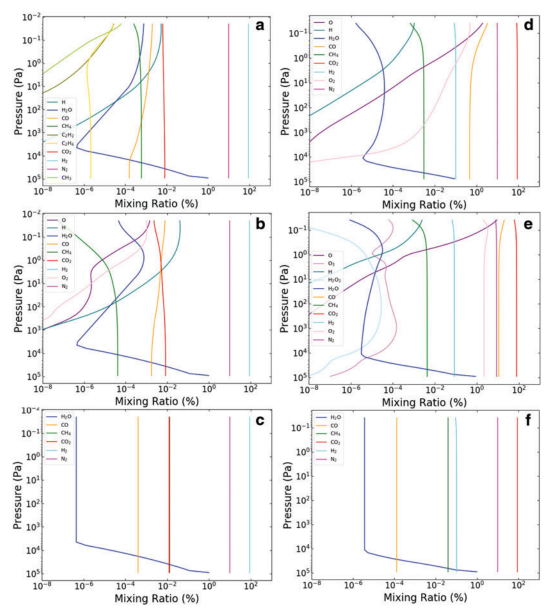
Appendix

Appendix A: Atmospheric Mixing Ratios and Temperature–Pressure Profiles

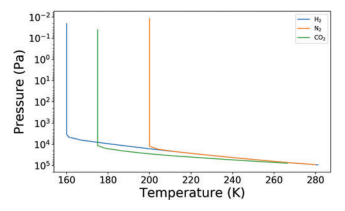
Appendix Figure A1 shows the molecular mixing ratio profiles used to simulate atmospheric spectra for the planets with CO₂- and H₂-rich atmospheres orbiting active M dwarfs, quiet M dwarfs, and Sun-like stars. To establish the

effect of adding phosphine to the atmosphere, we ran the photochemical models with the starting mixing ratios from Appendix Fig. A1 considering surface fluxes of phosphine (PH₃) ranging from 10^3 cm⁻² s⁻¹ to 10^{14} cm⁻² s⁻¹.

Appendix Figure A2 shows the temperature–pressure profiles used to simulate the transmission and emission spectra for the planets with H_2 - and CO_2 -rich atmospheres.



APPENDIX FIG. A1. Mixing ratio profiles of simulated Earth-sized planets with the following parameters: (a) H₂-dominated atmosphere on a planet orbiting a Sun-like star; (b) H₂-rich atmosphere on a planet orbiting an active M dwarf; (c) H₂-rich atmosphere on a planet orbiting a Quiet M dwarf; (d) CO₂-rich atmosphere on a planet orbiting a Sun-like star; (e) CO₂-rich atmosphere on a planet orbiting an active M dwarf; (f) CO₂-rich atmosphere on a planet orbiting a quiet M dwarf. Vertical axis represents pressure in units of Pa, and horizontal axis shows the mixing ratio represented as a percentage of the total atmosphere. Figure partially adapted from the works of Hu *et al.* (2012) and Seager *et al.* (2013).



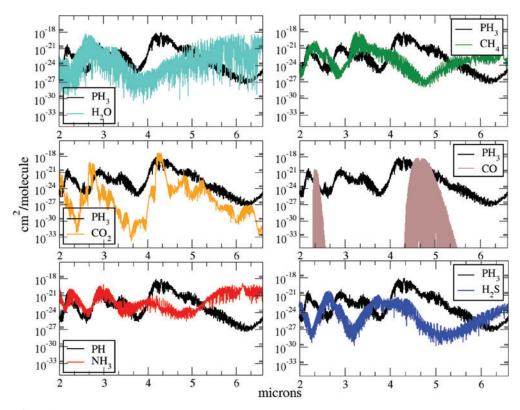
APPENDIX FIG. A2. Temperature–pressure profiles of the simulated massive super-Earths ($M_p = 10 \ M_E$ and $R_p = 1.75 \ R_E$), with H₂- and CO₂-rich atmospheres (N₂-rich atmosphere shown for comparison as an intermediate reducing scenario). Vertical axis represents pressure in units of Pa, and horizontal axis shows the temperature in units of K. Figure adapted from the work of Hu *et al.* (2012).

Appendix B: Detailed Spectral Comparison of Phosphine with Other Atmospheric Components

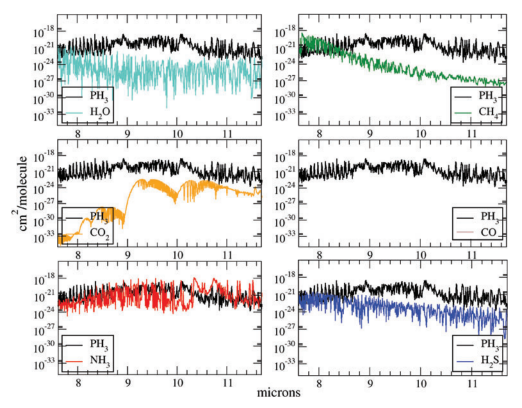
The three strongest spectral features of phosphine occur at 2.7-3.6, 4.0-4.8, and $7.8-11.5\,\mu m$. Appendix Figures B1 and B2 (2–6.5 and $7.5-11.8\,\mu m$, respectively) show a detailed comparison between the room temperature cross sections of PH₃ and six common atmospheric molecules, namely H₂O, CH₄, CO₂, CO, NH₃, and H₂S.

Appendix C: Thermodynamics of Abiotic Phosphine Synthesis at Circumneutral pH

We conduct a thermodynamic analysis to determine whether the formation of phosphine from phosphate and phosphite is thermodynamically favored at circumneutral pH. We calculate the free energy of formation of PH₃ in solution (and solid, in reduction of calcium phosphate) under geochemical source conditions and elemental composition similar to



APPENDIX FIG. B1. Comparison of the spectral cross sections of phosphine with other gases at room temperature, in the 2–6.5 μ m region. Intensity on y-axes with units of cm²/molecule and wavelength represented on the x-axes in microns. All cross sections are calculated at zero-pressure (i.e., Doppler-broadened lines only) using the procedure described by Hill et al. (2013). PH₃, shown in black, is distinguishable in this region from all compared molecules.



APPENDIX FIG. B2. Comparison of the spectral cross sections of phosphine with other gases at room temperature, in the 7.6–11.7 μm region. Intensity on *y*-axes with units of cm²/molecule and wavelength represented on the *x*-axes in microns. All cross sections are calculated at zero-pressure (*i.e.*, Doppler-broadened lines only) using the procedure described by Hill *et al.* (2013). PH₃, shown in black, is distinguishable in this region from all compared molecules except ammonia. Note that CO is not known to absorb in this region.

terrestrial planets for two temperature conditions (0°C and 140°C, or 273 and 413 K) and two extreme H_2 abundance levels (Appendix Table C1) on the basis of established thermodynamic values (Barner and Scheurman, 1978; Conrad *et al.*, 1986; Amend and Shock, 2001; Linstrom and Mallard, 2001; Fu *et al.*, 2013). The free energy of reaction of formation of PH_3 is given by:

$$\Delta G = \Delta G^{\circ} + R.T.ln(Q),$$

where ΔG^{o} is the standard free energy of formation, R is the gas constant, T is the absolute temperature, and Q is the reaction quotient.

The resulting calculated free energy of formation of phosphine can be used as a proxy to estimate the likelihood of PH₃ formation as a geochemical false positive (Bains *et al.*, 2017). The more positive the free energy of formation of PH₃ for a given reaction pathway, the less likely it is to be produced through geochemical processes. We note that the estimation of likelihood of geochemical false-positive scenarios for PH₃ is a part of the larger effort to estimate the possibilities for false positives for all biosignature gases (Bains *et al.*, 2017).

The calculated energy of formation of phosphine is strongly positive for any of the proposed PH₃ formation reaction pathways, given the geochemical concentrations of gases, suggesting that if PH₃ is detected in an atmosphere of an exoplanet, its source is not likely to be geochemical,

increasing the probability for biological production (Appendix Table C1).

Appendix D: Production of Phosphine from Phosphite at Acidic pH

It has been suggested that abiotic disproportionation of phosphites could be a source of environmental phosphine (Roels and Verstraete, 2001). Indeed, one common laboratory method of obtaining PH₃ is by heating phosphite, which disproportionates to phosphine and phosphate (Gokhale *et al.*, 2007). This process is only favored at high temperature ($T > 50^{\circ}\text{C}$ or T > 323K) or acid conditions (pH $\sim < 1.3$, where H₃PO₃ and/or H₂PO₃ dominate) (Appendix Fig. D1). We also note that the production of phosphite *via* the reduction of phosphate with H₂ on Earth-analog habitable worlds is highly endergonic and is also unlikely (Appendix Table D1). For a detailed analysis of thermodynamic limitations of PH₃ and phosphite production, see Bains *et al.* (2019a).

Such acidic and high-temperature conditions are not characteristic of Earth-analog habitable worlds, although they are available at "black smoker" vents (Martin *et al.*, 2008), and might be more common on a hot world with more acid oceans, for example, a world with high pCO₂ orbiting close to its parent star. We conclude that disproportionation of phosphite is unlikely to generate significant fluxes of phosphines on Earth-analog habitable worlds, but that worlds which are known to have hot extremely acid

APPENDIX TABLE C1. REACTION PATHWAYS AND THE ENERGY OF FORMATION OF PHOSPHINE FROM PLAUSIBLE GEOCHEMICAL VOLATILE CONCENTRATIONS

(1) Oxidized atmosphere (low H_2 levels)

 ΔG (kJ mol⁻¹) at plausible geochemical gas concentrations $H_2 = 10^{-6} \text{ bar}$ $PH_3 = 10^{-6} \text{ bar}$ $CH_4 = 10^{-6} \text{ bar}$ $CO_2 = 0.01 \text{ bar}$ $Phosphate = 10^{-5} \text{ M}$ $Phosphite = 10^{-6} \text{ M}$ pH = 7

Possible geochemical PH_3 formation reaction	$\Delta G^{o} (kJ \ mol^{-1})$	$Temp. = 0^{\circ}C$	$Temp. = 140^{\circ}C$
$\frac{1}{\text{HPO}_4^{2-}_{(aq)} + 4\text{H}_{2(aq)}} \rightarrow \text{PH}_{3(aq)} + 2\text{H}_2\text{O}_{(1)} + 2\text{OH}_{(aq)}^{-}$	225.16	410.14	538.49
$HPO_{4_{-}(aq)}^{2^{-}} + CH_{4(aq)} \rightarrow PH_{3(aq)} + CO_{2(aq)} + 2OH_{(aq)}^{-}$	415.32	503.97	558.16
$HPO_3^{2-(aq)} + 3H_{2(aq)} \rightarrow PH_{3(aq)} + H_2O_{(1)} + 2OH_{(aq)}^{-(aq)}$	130.79	303.69	355.06
$\begin{array}{l} HPO_{4}^{2-}(_{aq}) + 4H_{2(aq)} \rightarrow PH_{3(aq)} + 2H_{2}O_{(1)} + 2OH^{-}_{(aq)} \\ HPO_{4}^{2-}(_{aq}) + CH_{4(aq)} \rightarrow PH_{3(aq)} + CO_{2(aq)} + 2OH^{-}_{(aq)} \\ HPO_{3}^{2-}(_{aq}) + 3H_{2(aq)} \rightarrow PH_{3(aq)} + H_{2}O_{(1)} + 2OH^{-}_{(aq)} \\ HPO_{3}^{2-}(_{aq)} + \frac{1}{2}H_{2}O_{(1)} + \frac{3}{4}CH_{4(aq)} \rightarrow PH_{3(aq)} \\ + \frac{3}{4}CO_{2(aq)} + 2OH^{-}_{(aq)} \end{array}$	358.80	451.88	459.38
$^{1/2}$ Ca ₃ (PO ₄) _{2(s)} + 4H _{2(g)} \rightarrow PH _{3(g)} + 1½Ca(OH) _{2(s)} + H ₂ O _(g)	373.32	465.08	526.75

(2) Reduced atmosphere (high H_2 levels)

 ΔG (kJ mol⁻¹) at plausible geochemical gas concentrations $H_2 = 1$ bar $PH_3 = 10^{-6}$ bar $CH_4 = 10^{-1}$ bar $CO_2 = 0.01$ bar $Phosphate = 10^{-6}$ M $Phosphite = 10^{-5}$ M pH = 7

Possible geochemical PH_3 formation reaction	$\Delta G^{o} (kJ \ mol^{-1})$	$Temp. = 0^{\circ}C$	$Temp. = 140^{\circ}C$
$\begin{array}{c} \text{HPO4}_{2^{-}(\text{aq})}^{2^{-}} + 4H_{2(\text{aq})} \rightarrow \text{PH}_{3(\text{aq})} + 2H_{2}O_{(\text{l})} + 2OH^{-}_{(\text{aq})} \\ \text{HPO4}_{2^{-}(\text{aq})}^{2^{-}} + \text{CH}_{4(\text{aq})} \rightarrow \text{PH}_{3(\text{aq})} + \text{CO}_{2(\text{aq})} + 2OH^{-}_{(\text{aq})} \\ \text{HPO3}_{2^{-}(\text{aq})}^{2^{-}} + 3H_{2(\text{aq})} \rightarrow \text{PH}_{3(\text{aq})} + H_{2}O_{(\text{l})} + 2OH^{-}_{(\text{aq})} \\ \text{HPO3}_{2^{-}(\text{aq})}^{2^{-}} + \text{CH}_{4(\text{aq})} \rightarrow \text{PH}_{3(\text{aq})} + \text{CO}_{(\text{aq})} + 2OH^{-}_{(\text{aq})} \\ \text{$\frac{1}{2}$ Ca}_{3}(\text{PO}_{4})_{2(\text{s})} + 4H_{2(\text{g})} \rightarrow \text{PH}_{3(\text{g})} + 1\frac{1}{2}\text{Ca}(\text{OH})_{2(\text{s})} + H_{2}O_{(\text{g})} \\ \end{array}$	225.16	289.95	356.65
	415.32	483.07	526.54
	130.79	204.40	204.84
	358.80	427.06	421.83
	373.32	339.65	337.00

Free energy of formation of phosphine (ΔG) was calculated under geochemical concentration of gases for two different temperature scenarios and two H₂ levels (1) oxidized atmosphere (low H₂ levels) and (2) reduced atmosphere (high H₂ levels). Under all the tested terrestrial planet conditions, the free energy of formation of PH₃ (ΔG) is positive, making the geochemical formation of PH₃ from phosphate and phosphite an unlikely scenario at circumneutral pH.

oceans ($T > \sim 50$ °C/323 K, pH < ~ 1.3) may be vulnerable to this false-positive scenario.

Appendix E: Phosphine in the Context of Terrestrial Biology

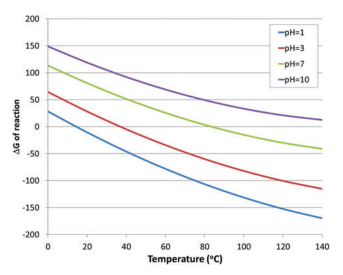
We briefly address a few of the common concerns in the association of phosphine with terrestrial biology: the toxicity of PH₃; the absence of a known metabolic biosynthetic pathway and enzymatic mechanism for the production of PH₃; some context on the ecology of phosphines; and a discussion of the phosphorus cycle.

Phosphine Toxicity

On modern Earth, phosphine is a rare toxic gas. It is highly toxic to aerobically metabolizing organisms (Bond and Monro, 1967), which is reflected by its wide use as

fumigant [e.g., reviewed in Perkins et al. (2015)]. The toxicity of PH₃ appears to be strictly dependent on aerobic metabolism [e.g., reviewed in Valmas et al. (2008)]. The detailed analysis of the toxic effects of PH₃ on aerobic organisms is beyond the scope of this article and is published elsewhere (Bains et al., 2019b). However, if PH₃ chemistry is indeed selectively incompatible with O₂-dependent metabolism of aerobic organisms, as literature suggests, it opens an intriguing possibility that phosphines may safely be produced as secondary metabolites to much greater extent by obligatorily anaerobic life, which does not rely on O₂ metabolism (Bains et al., 2019b).

We note that it is not known why phosphine is so much more toxic to oxygenic organisms than anoxic ones, given that both rely on heme iron atoms that interact with PH₃ (Bains *et al.*, 2019b). We postulate that the highly positive electrode potential of the oxygen-handling enzymes may



APPENDIX FIG. D1. Phosphine production by disproportionation of phosphite is favored under very low pH or high temperature conditions. Y axis: ΔG° (Gibbs free energy of reaction under standard conditions), in units of kJ mol⁻¹. X axis: temperature (°C). The free energy of reaction for four reactions are shown: disproportionation at very low pH, where phosphite is present as unionized H_3PO_3 form; and disproportionation of ionized species present at higher pHs (see Bains *et al.*, 2019a for more details on calculation method).

make them more susceptible targets for PH_3 complexation and hence inactivation.

Biosynthetic Pathways of Phosphine and Other Volatiles

The identification of natural products from anaerobic organisms and elucidation of their biosynthetic pathways is notoriously difficult. Culturing of anaerobic organisms is much more laborious task than aerobic ones, and identification or purification of anaerobic metabolites is much more complex as well. It requires specialized experimental setup, as often isolated molecules or their biosynthetic precursors get destroyed in our oxygen-rich atmosphere before they can be properly studied. It is therefore not surprising that the exact molecular mechanism of biological phosphine formation in anaerobic environment has eluded discovery for such a long time. Even identification of biosynthetic pathways of natural molecules from aerobic organisms that can be easily cultured in the laboratory conditions can take many decades. For example, the biosynthetic pathway of

Appendix Table D1. ΔG (kJ mol⁻¹) for Forming Phosphite by Reduction of Phosphate with H₂: $\text{HPO}_4^{2^-} + \text{H}_2 \rightarrow \text{HPO}_3^{2^-} + \text{H}_2\text{O}$

Temperature		ΔG (kJ mo Γ^{I}); reducing conditions (as above in Appendix Table C1)
0°C/273 K	38.52	17.62
140°C/413 K	95.36	63.73

nucleocidin, an unusual fluorine-containing natural antibiotic produced by a bacterium *Streptomyces calvus*, was only recently elucidated, after ~ 60 years of intensive studies (Petkowski *et al.*, 2018).

The economically important production of volatile molecules by fruiting bodies of truffles (Tuber sp.), responsible for the unique aroma of the fungus, is another good illustration of the complexity of the biological production of volatile and nonvolatile natural metabolites. Despite many decades of studies, and an influx of funding from a multimillion-dollar food industry, the biosynthetic pathways responsible for synthesis of truffle volatiles are not fully understood (Zambonelli et al., 2016). The biological production of truffle volatiles, for example, thiophene derivatives, is further complicated by the fact that the biosynthetic pathways for molecules responsible for the unique aroma of truffles are likely shared between multiple species of symbiotic microorganisms (bacteria and yeast) living within the fruiting body of the fungus. The metabolic processes of the symbionts and the fungus host collectively allow for the formation of the final volatile product (e.g., thiophene derivatives) (Zambonelli et al., 2016). Each of the symbionts, and the host fungus, likely contain only part of the full biosynthetic pathway required for the formation of the final volatile product, and each single species is only responsible for the production of few biosynthetic intermediates for the formation of thiophene derivatives (Zambonelli et al., 2016).

Such ecological complexity of biosynthetic pathways is a common occurrence. It is quite possible that phosphine production requires a similarly complex ecological scenario, where the full biosynthetic pathway for PH₃ synthesis is shared between multiple species of anaerobic microorganisms and only leads to phosphine formation under very specific environmental conditions (Bains et al., 2019a). Such complex ecological scenarios further complicate the elucidation of the direct mechanism of biosynthesis. Recent work postulated that PH₃ production may be associated with the microbial tricarboxylic acid cycle of Enterobacter (Fan et al., 2020). As it is in the case of the complex ecology of biosynthesis of truffle volatiles, the lack of understanding of the details of their biological production does not mean that phosphine or any other natural molecule is not a product of the metabolism of the living organisms.

Phosphines in Earth's Ecology

All life on Earth relies on phosphorous-containing compounds in its metabolism. The great majority of biochemicals used by life on Earth are pentavalent phosphorus-containing molecules, predominantly phosphates (Petkowski et al., 2019b). At first glance, trivalent phosphorus compounds appear to be almost completely absent from biology, yet strong circumstantial evidence seems to suggest that anaerobic life on Earth explores chemistry of phosphines beyond simple phosphine production. One isolated study (Davies, 2008) reports identification of phospholane volatile in biological samples. Phospholane is a volatile, trivalent phosphoruscontaining, five-membered ring saturated hydrocarbon. It is the only molecule reported so far to be isolated from biological samples, which contains a bond between trivalent phosphorus and carbon atoms. Phospholane, like PH₃, is a volatile trivalent phosphorus molecule that was identified in European

badger fresh scat samples (Davies, 2008). It is interesting to note that, so far, all trivalent phosphorus-containing molecules that were isolated from biological samples (*i.e.*, PH_3 and phospholane) appear to be produced in strictly anoxic environments and appear to be exclusively associated with anaerobic O_2 -free dwelling microorganisms, which seems to be in agreement with previous observations that phosphines tend to be poisonous to aerobic life, but nontoxic in anoxic environments.

Both phosphine and phospholane were detected in anaerobic environments, which are generally difficult to study. If natural production of trivalent phosphorus-containing natural chemicals is strictly dependent on anoxic environments, it is likely that more such molecules are going to be discovered in the future. Almost all natural compounds reported to be produced by life on Earth were detected or isolated from organisms living in oxygenic environments, as shown by the repositories of natural molecules produced by life (Bains et al., 2019b; Petkowski et al., 2019a); thus, the small number of known natural molecules containing trivalent phosphorus might be the result of sample bias toward O₂-rich environment. For a detailed study on the chemistry of PH₃ in the context of the terrestrial biology, see Bains et al. (2019b).

The Phosphorus Cycle on Earth

It is generally assumed that life on Earth only consumes phosphorus in its most oxidized form of phosphate and that phosphates are the only form of phosphorus that is useful for life's cellular metabolism. This assumption is incorrect. Organisms can also use phosphorus in many other forms beyond phosphates, such as organic phosphorus esters, polyphosphates, phosphonates, and even reduced phosphorus species such as phosphites. Phosphite and hypophosphite biochemistry is well known (Casida Jr. 1960; Metcalf and Wolfe, 1998; Stone and White, 2012; Pasek et al., 2014; Figueroa and Coates, 2017), and some microorganisms living in strictly anoxic environments, such as anaerobic sludges in wastewater treatment plants, can fulfill all their energy requirements by oxidizing phosphite to phosphate (a process called dissimilatory phosphite oxidation) (Figueroa and Coates, 2017; Figueroa et al., 2018). Studies in recent years established that a complex redox phosphorus cycle (beyond the simple cycling of phosphates) has equally important bioenergetic and ecological consequences as the more recognized microbial nitrogen, carbon, or sulfur cycles (Karl, 2014).

Marine organisms can reduce a significant fraction (up to 15%) of the total phosphates taken up from the surrounding environment (Van Mooy et al., 2015). The reduced phosphorus species (e.g., in the form of the soluble phosphite) are then rapidly released to the surrounding seawater (Van Mooy et al., 2015). The authors argue that the total amount of phosphorus cycled through such complex redox reactions could exceed other oceanic sources of phosphorus (e.g., direct continental and atmospheric sources) (Van Mooy et al., 2015). It is unknown why marine planktonic organisms engage in such an energetically costly endeavor. Production and rapid release of large quantities of reduced phosphorus species is unexpected, especially in the open oceanic waters where phosphorus is a limiting nutrient. The production of reduced phosphorus species is, of course, dependent on the abundance of phosphorus but appears to be independent of inorganic or organic phosphorus concentrations and independent of the overall phosphate turnover time (Van Mooy et al., 2015). The synthesis and rapid export of soluble reduced phosphorus species could stem from the general cooperative behavior of planktonic organisms in the open ocean. One of the more studied producers of reduced phosphorus species is the cyanobacteria Trichodesmium. The colonies of Trichodesmium are also inhabited by many species of epibiotic and symbiotic bacteria that appear to cooperate in the cycling of phosphorus (Hmelo et al., 2012; Van Mooy et al., 2012). Phosphonates and reduced phosphorus compounds (i.e., phosphites) may be part of a barter economy where phosphorus is exchanged for other nutrients between Trichodesmium and other species living in the cooperative colonies. Such examples of biological importance of rare redox phosphorus species clearly show that reduced phosphorus species play an important albeit underappreciated role in the global biogeochemical cycle of phosphorus.

The exact place and role of phosphine in the global phosphorus cycle is not yet fully known. It is, however, likely that, similar to other reduced phosphorus species, PH₃ also has an important role in the global cycling of this essential element.

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