Timescales for Prebiotic Photochemistry Under Realistic Surface Ultraviolet Conditions

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Abstract

Ultraviolet (UV) light has long been invoked as a source of energy for prebiotic chemical synthesis, but experimental support does not involve sources of UV light that look like the young Sun. Here we experimentally investigate whether the UV flux available on the surface of early Earth, given a favorable atmosphere, can facilitate a variety of prebiotic chemical syntheses. We construct a solar simulator for the UV light of the faint young Sun on the surface of early Earth, called StarLab. We then attempt a series of reactions testing different aspects of a prebiotic chemical scenario involving hydrogen cyanide (HCN), sulfites, and sulfides under the UV light of StarLab, including hypophosphite oxidation by UV light and hydrogen sulfide, photoreduction of HCN with bisulfite, the photoanomerization of α-thiocytidine, the production of a chemical precursor of a potentially prebiotic activating agent (nitroprusside), the photoreduction of thioanhydridouridine and thioanhydroadenosine, and the oxidation of ethanol (EtOH) by photochemically generated hydroxyl radicals. We compare the output of StarLab to the light of the faint young Sun to constrain the timescales over which these reactions would occur on the surface of early Earth. We predict that hypophosphite oxidation, HCN reduction, and photoproduction of nitroprusside would all operate on the surface of early Earth in a matter of days to weeks. The photoanomerization of α-thiocytidine would take months to complete, and the production of oxidation products from hydroxyl radicals would take years. The photoreduction of thioanhydridouridine with hydrogen sulfide did not succeed even after a long period of irradiation, providing a lower limit on the timescale of several years. The photoreduction of thioanhydroadenosine with bisulfite produced 2′-deoxyriboadenosine (dA) on the timescale of days. This suggests the plausibility of the photoproduction of purine deoxyribonucleotides, such as the photoproduction of simple sugars, proceeds more efficiently in the presence of bisulfite. Key Words: Photochemistry—Young Sun—Early Earth—Prebiotic chemistry—Origins of life. Astrobiology 21, 1099–1120.

1. Introduction

The starting points for origins of life research are diverse and researchers do not agree about the best place to begin. Some start with biology and investigate how extant metabolic pathways may have come to fruition, concentrating on bacterial and archaeal life that most closely resembles the last universal common ancestor. Others start with chemistry and find synthetic reactions and pathways that result in greater chemical complexity that synthesize the building blocks of the macromolecules that constitute life, such as DNA, RNA, proteins, and lipids. Yet others start with geology or exogeology, considering the mineral surfaces, volcanic degassing, atmospheric conditions, astrochemistry, and cometary chemistry and delivery of these building blocks.

Here we explore a particular scenario that started with chemical synthesis and then fruitfully underwent refinements using inputs from biochemistry and geochemistry (Sasselov et al., 2020). Starting with hydrogen cyanide (HCN), hydrogen sulfide, cyanamide, cyanoacetylene, phosphates (or reduced phosphorus species, see Ritson et al., 2020) in liquid water and exposing these mixtures to ultraviolet (UV) light and combining them in particular sequences result in high

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and selective yields of simple sugars (Ritson and Sutherland, 2012; Xu et al., 2018), pyrimidine ribonucleotides (Powner et al., 2009), purine deoxyribonucleosides (Xu et al., 2020), other purines (Stairs et al., 2017), several amino acids and phospholipid precursors (Patel et al., 2015), and mechanisms for nonenzymatic activation of these (Mariani et al., 2018b; Bonfio et al., 2019; Liu et al., 2020), or pathways to form oligomers ab initio (Canavelli et al., 2019). Chemical pathways, especially at the start of prebiotic synthesis, must be selective and result in reasonable yields, or else the products have little chance to interact with each other to produce the macromolecules that constitute life (Sutherland, 2015; Rimmer et al., 2018). This scenario therefore provides a promising chemical starting point for origins of life research (Sutherland, 2017).

The chemical starting point for this scenario has intersected multiple times with geological starting points. Hydrogen sulfide is difficult to concentrate in solution from volcanic degassing without also building up concentrations of bisulfite from dissolution of sulfur dioxide (Ranjan et al., 2018, see also Gaillard and Scaillet, 2014). In closed lake environments, Ranjan et al. (2018) found that sulfites could have plausibly achieved 1 M concentrations on early Earth. The photochemistry works with bisulfite as well as hydrogen sulfide (Xu et al., 2018). Bisulfite serves better for this prebiotic photochemical reduction than cyanocuprates (Todd et al., 2018) or hydrogen sulfide (Rimmer et al., 2018), because UV photodetachment of bisulfite electrons has a much larger cross section than the UV photodetachment of electrons from cyanocuprates or hydrogen sulfide.

These intersections do not leave this scenario unchallenged. Nonenzymatic activation of nucleotides, lipids, and amino acids can be accomplished through the intermediacy of imidazolides, which can be derived from methyl isocyanide and imidazoles (Mariani et al., 2018b; Bonfio et al., 2019; Liu et al., 2020). Methyl isocyanide can be formed by aqueous nitropurusside chemistry, and HCN, ferrous iron, and nitrite (NO$_2^-$) react to form nitropusside in water under UV light (Mariani et al., 2018b). Fixed oxidized nitrogen on early Earth would mostly have been in the form of nitrates, whereas nitrates would likely have been less abundant on early Earth (Ranjan et al., 2019). Another problem for this scenario is that, as we show, the <250 nm wavelength end of the near ultraviolet (NUV; 200–300 nm) spectrum can photodetach electrons from HO$^-$, producing hydroxyl radicals that may destroy many of the photochemical products in this scenario.

This chemical scenario is directly connected to starlight, which provides a further intersection between prebiotic chemistry and astronomy. This intersection allows us to predict the planets on which life is most likely to originate via this photochemical scenario (Rimmer et al., 2018). We can use this criterion to delineate an “abiogenesis zone,” outside of which the star is not bright enough in UV light to permit the photochemical scenario. This zone provides criteria informed by prebiotic chemistry for exoplanet candidate selection and ranking in the search for extrasolar biosignatures. In addition, this zone provides a test for the scenario: if life is found outside the abiogenesis zone, then there must at least be some other plausible way for life to originate. This chemical scenario also provides predictions for prebiotic chemistry on the martian surface, relevant for future missions (Sasselov et al., 2020). Starlight and stellar activity affect the chemistry in other ways, by generating atmospheric HCN, relevant for prebiotic chemistry, and N$_2$O, which may affect the climate (Airapetian et al., 2016). The changing gas-phase chemistry will affect the average pH of open bodies of water (Krisiussen-Totton et al., 2018), although local lakes, ponds, and streams will plausibly have a wide range of pH, from highly acidic to alkaline (Cousins et al., 2013; Toner and Catling, 2019; Kadoya et al., 2020). It is also plausible that the pH will depend on the location in the body of water and will change over time (Cousins et al., 2013; Mariani et al., 2018; Rimmer and Shorttle, 2019).

We consider an atmosphere favorable for UV transmission to the surface of early Earth, where photons impinge on a shallow aqueous surface environment with sufficient concentrations of cyanide, bisulfite, phosphate, or precursor reduced phosphorus, and measure the amount of time required to build up the chemical products with high (>50%) steady-state yields. To investigate these timescales, we need to simulate the young Sun’s UV output within the laboratory.

All of the experiments that support, test, or challenge this scenario have been performed by using a discrete light source with most of its UV emission at a single wavelength. Most of the published experiments made use of sixteen low-pressure Hg lamps that emit in the UV almost exclusively at 254 nm. Some recent novel experiments, such as those of Roberts et al. (2018) and Todd et al. (2018, 2019), use a tunable light source and explore multiple wavelengths. None of these light sources is realistic. The Sun, like any star, is a broadband light source with a wavelength range spanning from <0.1 nm to more than 100 μm.

Habitable planets require atmospheres, and plausible abiotic atmospheres will block light <200 nm (Ranjan and Sasselov, 2016), and light at wavelengths greater than 400 nm will not typically contribute to the types of photochemistry invoked in prebiotic synthesis (e.g., see: Sanchez and Orgel, 1970; Ritson and Sutherland, 2012; Xu et al., 2018). This leaves a broad wavelength range for photochemistry, and photochemistry is inherently nonlinear. The chemistry that is produced by a 254 nm light source and a 365 nm light source cannot be added together to determine what chemistry will occur when both are present. Prebiotic chemical scenarios that use starlight need to be investigated by using a Sun-like light source. We provide such an investigation here, and find that in some cases the chemistry does not work as well, in some cases it works better, and in some cases it seems not to work at all.

In this article, we present StarLab, an experimental stellar simulator set up to simulate the light of the Sun as it was roughly 4 billion years ago. In Section 2, we provide the schematic for the experiment and compare its output to the output of young Sun analogues. In Section 3, we discuss a group of experiments that are key for different aspects of UV-driven prebiotic chemistry, for which we use StarLab. We also run these experiments under the more “classic” illumination source of a mercury lamp reactor (the Rayonet RPR-200) to demonstrate the differences in outcome between using a more realistic broadband source such as StarLab and narrow-band sources that emit primarily at wavelengths of 254 or 365 nm (depending on the specific bulb used). We show the results of these experiments. In Section 4, we discuss how these experiments can be directly compared with an early Earth environment to constrain whether the light of the young Sun was sufficient for these reactions on early Earth. We summarize the article in Section 5.
2. Simulating the Young Sun

Here we discuss the design of the UV source we use in all the following prebiotic chemical experiments. In simulating the young Sun, first we discuss the design of the experimental apparatus (Section 2.1) and then discuss the plausible UV spectrum of the young Sun and compare this spectrum with the apparatus (Section 2.2).

2.1. StarLab: the apparatus

The top-level requirements for StabLab are as follows:

1. To provide a broadband illumination that approximates the spectral irradiance profile of the Sun on the surface of the early Earth (preozone layer).
2. To provide a means to deliver that illumination into a 10-mm-sided quartz cuvette sample holder, minimizing loss of photons until it arrives at the sample.
3. To ensure that the sample does not overheat during the course of the experiment—for our purposes we defined this to be $T < 40^\circ C$ over the length of the experiment, which can range from 30 min to several days.
4. To have a means of measuring the spectral irradiance received by the sample and to be able to monitor the change in spectral irradiance after passing through the sample.

The ozone layer formed in Earth’s atmosphere sometime after the Great Oxygenation Event about 2.3 Ga (Luo et al., 2016), while life on Earth is believed to have arisen around 4 billion years ago. The ozone layer blocks radiation in the 200–310 nm wavelength range. To replicate the illumination on the surface of early Earth, around the time that life was first forming, the illumination source must provide UV photons down to 200 nm. This top-level requirement, derived from Requirement (1), determines the choice of lamps available for the experiment and the types of optical materials that can be used.

The lamps available that provide broadband illumination in the 200–400 nm region are as follows:

- Xenon arc lamp, usable wavelength range: 200–2500 nm.
- Deuterium arc lamp, usable wavelength range: approximately 160–400 nm.
- Laser-driven light source, wavelength range 170–2100 nm.

Modifications to this wavelength range are dependent on the exact lamp system configuration. "Ozone-free" versions of most lamps are available, which usually block wavelengths below 260 nm (thus preventing the conversion of $O_2$ to $O_3$, which is hazardous in laboratories that cannot provide sufficient ventilation). Other coatings can be available for the lamps to tune the lower cutoff wavelength and fiber-coupled lamps generally limit the lower wavelength cutoff to $\approx 190$ nm.

For the first version of our StarLab experimental configuration, we chose a 75 W Xenon arc lamp in the PowerArc range manufactured by Horiba. The lamp housing for this model uses an ellipsoidal reflector and so is very efficient at collecting and focusing most of the light emitted by the Xe bulb at a single focus (as a point of comparison, the more traditional vertical lamp housings, at one of their foci, would contain five to six times less flux for the same wattage lamp).

One feature of the spectral irradiance of a Xe lamp that needs consideration is the high-emission region from $\approx 750$ to $\approx 1100$ nm. This high-level emission is not present in the early solar spectrum and additionally brings problems into the experimental setup due to strong heating effects. Without cooling of the sample or a prefiltering of the Xe spectrum, the sample to be illuminated by our simulated starlight would rapidly boil. To meet Requirement (3), we devised an apparatus that can provide both the functions of filtering the unwanted spectral regions and allow for the possibility of cooling the sample if required. The schematic of the StarLab apparatus is shown in Fig. 1, and a picture of the experiment is shown in Fig. 1. Color images are available online.

FIG. 1. Schematic of StarLab, the stellar simulator for the young Sun. The focal distance from the aperture of the Xe lamp to the sample is 150 mm, which contains an optical depth of water in the quartz attenuator (water-filled tube) of 130 mm. The spectrometer fiber can be placed at the sample focus (via the sample loader) to measure the flux received by the sample before sample loading, or at the position shown in the drawing to monitor changes while a sample is being illuminated.
shown in Fig. 2. The setup is based around a piece of customized glassware—the glass in this case is GE 214, a type of fused quartz that has a high transmittance over our required wavelength range. This custom glassware, which we call the quartz attenuator, is filled with water and is designed to hold a sample cuvette at the focus of the Xe arc lamp. The water acts like an optical filter, preferentially absorbing the emissions from the Xe arc lamp in the near infrared and so preventing the sample from overheating. If required, the quartz attenuator can be connected to a circulating water supply so that the temperature of the water is maintained at a given value.

Figure 3 shows the spectral irradiance of StarLab in the wavelength region of interest, as measured at the sample position (through the water-filled quartz attenuator), and Figs. 4 and 5 show the ratio of StarLab to the young Sun, above the atmosphere and at the surface of early Earth, respectively.

![FIG. 2. Photograph of the quartz attenuator tube inside its mounting box. The Xe arc lamp is mounted to the left, outside of the box, the black interface plate can be seen to the left inside the mounting box. The sample is loaded through the circular opening in the top of the tube so that it is positioned at the focus of the Xe lamp. Color images are available online.](image)

For the fourth requirement (see list above), we used an Ocean Optics (now Ocean Insight) Flame UV-VIS (visible) Spectrometer to measure the spectrum of light received by the sample at the lamp focus and to monitor any changes in the spectrum during the course of an experiment (Fig. 1)—that is, from absorption changes due to chemical evolution in the prebiotic sample. The measurement range of the Flame spectrometer is given to be 200–850 nm, however, on usage, we quickly realized that for spectral regions <220 nm, the measurements had extremely low signal-to-noise, which effectively made this region unusable for our purposes. For the set of experiments presented here, we essentially have no flux below 220 nm due to the absorption of those wavelengths by nonpure water. This is a realistic scenario for aqueous chemistry on early Earth—the scenario itself requires several chemical species to be dissolved in the “prebiotic pondwater,” which rapidly attenuates the transmission of the shorter wavelengths of UV light (see Quickenden and Irvin,

![FIG. 3. Spectral irradiance (W/(cm²·nm)), versus wavelength (nm), for StarLab at the location of the sample (blue), compared with 75x the spectral irradiance 1 AU from the young Sun as observed above the atmosphere (black). Color images are available online.](image)

![FIG. 4. Ratio of the spectral irradiance of StarLab over the spectral irradiance 1 AU from the young Sun as observed above the atmosphere, as a function of wavelength (nm).](image)
2.2. Comparing StarLab with the young Sun

Wavelength chemistry may occur, except for at the actual surface of the water, where shorter optical depths, almost negligible amounts of UV light assuming a few mmol of ferrocyanide, even for very shallow column depth of water in StarLab—most of the mineral waters in that sample (similar to the Cambridge tap water used in the StarLab quartz attenuator) show a 0–0.3% throughput at 220 nm. This demonstrates that realistic water sources are very effective blockers of UV light below 220 nm. In addition, we note that for waters containing ferrocyanide we would expect the UV attenuation to be significantly higher (Chakrabarti and Roberts, 2008). So that in a postimpact pond on early Earth, the UV attenuation to be significantly higher (Chakrabarti and Roberts, 2008). So that in a postimpact pond on early Earth, assuming a 0.3 bar atmosphere with 33% N₂, 33% CO, and 33% CO₂ with water at vapor pressure.

Roberts, 2008). So that in a postimpact pond on early Earth, assuming a few mmol of ferrocyanide, even for very shallow optical depths, almost negligible amounts of UV light <220 nm would penetrate.

Using the experimental setup shown in Fig. 1 with tap water in the quartz attenuator, we have thus considered a realistic flux for most of a body of water on early Earth, except for at the actual surface of the water, where shorter wavelength chemistry may occur.

2.2. Comparing StarLab with the young Sun

The Sun is a G2V-type star and is about 4.6 billion years old (Bonanno et al., 2002). It exhibits a typical activity level of stars of similar age and mass (Mittag et al., 2016). Using the Yonsei-Yale evolutionary track (Kim et al., 2002; Yi et al., 2003), its effective temperature has remained nearly constant with age, and its radius has increased with age. This means that in the past its radius was smaller. The Sun <3.5 Ga was 75% the size of the Sun today and this is the reason for the faint young Sun paradox (Sagan and Mullen, 1972). The Sun can be compared with other young solar metallicity stars that fit this evolutionary track, and the spectra of these stars can be measured.

These spectra are a proxy of the spectrum of the Sun at that age. The star κ1-Ceti is a star with solar metallicity and has an age of 300–400 My and matches the Yonsei-Yale evolutionary track for the Sun when it was 300–400 My old, and so, κ1-Ceti is an analog young Sun, and its UV spectrum is used as a proxy of the UV spectrum of the young Sun (Ribas et al., 2005, 2010). Claire et al. (2012) use κ1-Ceti, along with EK Dra, π1-Uma, γ1-Ori, β Com, and β Hyi, to construct an evolutionary model of the Sun’s UV spectrum from 4.5 Ga to the present. Since κ1-Ceti is approximately the age of the Sun 4.0 Ga, and since this seems consistent with the time life approximately originated on Earth, we compare the spectrum of our lamp with the spectrum of κ1-Ceti, which we hereafter call the young Sun or young solar spectrum.

Early Earth must have had an atmosphere to support liquid water, although this atmosphere may have been more tenuous 4.0 Ga than today (Som et al., 2016; Avice et al., 2017; Rimmer et al., 2019; Lehmer et al., 2020; Payne et al., 2020). The light of the young Sun must traverse this atmosphere to reach the surface and the atmosphere will block some UV light. For this work, we assume a 0.3 bar atmosphere composed of 33% N₂, 33% CO, and 33% CO₂ with water at vapor pressure, the same as the >3.5 Ga atmosphere Rimmer et al. (2019) used to explain the observed micrometeorite oxidation from the work of Tomkins et al. (2016), and consistent with other early Earth model atmospheres invoked to explain the same data (Lehmer et al., 2020; Payne et al., 2020). We solve for the surface UV flux using the model of Rimmer & Helling (2016) and Rimmer and Rugheimer (2019) with the UV absorption and scattering cross sections from PHIDRATES (Huebner et al., 1992; Huebner and Mukherjee, 2015) for most species, with updated cross sections for CO₂ (Ityaksov et al., 2008), HCHO (Chen and Zhu, 2001), H₂O (Ranjani et al., 2020), and CH₄ (Burkholder et al., 2020). Most of these cross sections were obtained from the Mainz database (Keller-Rudek et al., 2013).

3. UV-Driven Prebiotic Chemistry: Methods and Results

Here we discuss the series of experiments we performed to test whether UV light is prebiotic in the context of the prebiotic chemical scenario discussed in Section 1. We answer this question in terms of the timescale of reaction, by which we here mean a photochemical “timescale,” here defined as the time it takes to achieve a 50% yield in a given reaction. This timescale is relevant for prebiotic plausibility of sequences of reactions (see Rimmer et al., 2018, their figure 3). Although StarLab has the same spectral shape as the faint young Sun between 200 and 400 nm, it does not have the same intensity as that expected on the surface of early Earth, and so, the timescale in the laboratory does not directly represent but predicts the timescale on early Earth.

The rate constant at which a pure photochemical reaction takes a reactant to product is called the photoproduction rate, \( k_p \) (s⁻¹), and takes place in a unit volume, depends on the flux of photons of a particular wavelength, \( \lambda \) (nm), entering into a unit sphere surrounding that volume, \( F_\lambda \) \([\text{cm}^2/\text{s} \cdot \text{Å}]\), which is related to the spectral irradiance (which our spectrometer measures) by a factor of \( hc/\lambda \), the individual photon’s energy. It also depends on the probability that a given photon of wavelength \( \lambda \) is absorbed by the reactant, which is represented by the absorption cross section, \( \sigma_{a,\lambda} \) \((\text{cm}^2)\), multiplied by the probability that the product results from the absorption of that photon, called the quantum yield, \( \Phi_p \). We define a reaction cross section as \( \sigma_{\lambda} = \sigma_{a,\lambda} \Phi_p \). The photoproduction rate is then:
\[ k_v = \int \sigma_v F_\lambda \, d\lambda \quad (1) \]

When the photochemical process is known to be linearly dependent on concentration and that there are no other processes that provide significant competition over the same timescale as the irradiation, then the time it takes for a reaction to achieve a yield, \( f \), can be related to the photoproduction rate, \( k_v \) (s\(^{-1}\)), by the following equation:

\[ k_v,\text{exp} = \frac{1}{t_{\text{exp}}} \log\left( \frac{1}{1-f} \right) \quad (2) \]

The advantage of reproducing a plausible UV spectrum for the young Sun is that we can then scale this experimental rate by noting that \( F_{\lambda,\text{exp}} = \alpha(\lambda) F_{\lambda,\text{sun}} \), and so, insofar as \( \alpha \) is constant with respect to \( \lambda \):

\[ k_{v,\text{exp}} = \int \sigma_v F_{\lambda,\text{exp}} \, d\lambda = \alpha \int \sigma_v F_{\lambda,\text{sun}} \, d\lambda = \alpha k_{v,\text{sun}} \quad (3) \]

\[ t_{\text{irr}} = \alpha t_{\text{exp}} \quad (4) \]

We can relate this to Eq. 2 and can then relate experimental time to the time it would take to complete the same reaction on the surface of early Earth, noting that the timescale on early Earth (and for any planet i.e., not tidally locked) will be twice as long to account for the day/night cycle. First, we define an “irradiation time,” \( t_{\text{irr}} \), which is the amount of time a sample would need to be irradiated on early Earth to match the irradiation by StarLab:

\[ t_{\text{Earth}} = \frac{\alpha}{2} t_{\text{exp}}. \quad (6) \]

We return to this concept and explore the wavelength dependence of \( \alpha \) in Section 4. For now, we relate the outcome of a series of experiments exemplifying different parts of the UV-driven prebiotic chemical scenario to this timescale. This timescale can be compared with the degradation timescales of the reactants, or other relevant chemical timescales, to test the plausibility of these reactions on the surface of early Earth.

We begin with a summary of the UV-driven prebiotic chemical scenario (Section 3.1), and then discuss a selection of key photochemical reactions in order. We start with the oxidation of hypophosphite with hydrogen sulfide and UV light (Section 3.2), and then proceed to the photoreduction of HCN by solvated electrons from bisulfite (Section 3.3), then the photoanomerization of \( \alpha \)-thiouridine (Section 3.4), the attempted photoreduction of thioanhydrouridine (Section 3.5), the photoreduction of thioanhydroadenosine (Section 3.6), the photoproduction of nitropyruvate (Section 3.7), and the oxidation of ethanol (EtOH) by photoproduced hydroxyl radicals (Section 3.8).

### 3.1. Overview of the scenario

The series of reactions discussed in this article are best understood in the context of a scenario—a sequence of chemical reactions that takes place in a specific geochemical environment. The scenario we discuss here is shown in Fig. 6.

We begin with Earth after the single moon-sized impactor or multiple smaller impactors provided the material for the late veneer (Genda et al., 2017), which according to Brasser et al. (2020) occurred at \( \sim 4.48 \) Ga. Models of the chemical effect of this bombardment suggest that the impact produces a transient reducing atmosphere that could have persisted for a 100 million years (Zahnle et al., 2020). We focus on one such surface, on a volcanic island.

Another impact, at the tail end of accretion, strikes this surface, generating a plasma shock that results in a considerable amount of HCN, reduced phosphorous species, ammonia (NH3), and, depending on atmospheric redox, some nitric oxide and nitrogen dioxide (NO and NO2). These species subsequently rain out onto an Fe(II)-rich surface, where the HCN forms ferrocyanide salts (Patel et al., 2015, see their figure 2), hypophosphite is among the reduced phosphorous species (Ritson et al., 2020), and the NO and NO2 are oxidized to form nitrates and nitrites (Ranjan et al., 2019), but dominantly nitrates.

Impact craters provide ideal environments for surface hydrothermal vents (Bryant et al., 2013). The pool that intersects with a hydrothermal vent has sulfur dioxide and hydrogen sulfide bubbling through, and if the pH is sufficiently high, \( \equiv 7 \), the sulfur dioxide reacts with the water, forming sulfite and bisulfite salts, and the hydrogen sulfide dissociates, forming sulfides (Ranjan et al., 2018). This pH is plausible for surface hydrothermal systems on volcanic islands, even under CO2-rich atmospheres (Cousins et al., 2013).

As water will self-dissociate producing hydroxide (OH\(^-\)) anions, UV light will impinge on both OH\(^-\) and sulfide (HS\(^-\)), resulting in solvated electrons in addition to OH and HS radicals. The photodetachment of an electron from HS\(^-\) is the first step of the reaction discussed in Section 3.2, and HS radicals can oxidize hypophosphite, resulting in the phosphate required as a chemical and pH buffer for subsequent prebiotic chemistry. The oxidation of hypophosphite with HS radicals is the second step of the reaction discussed in Section 3.2. The photodetachment of an electron from OH\(^-\) is the reaction discussed in Section 3.8 and OH radicals can destroy the products of prebiotic chemistry.

UV light will also impinge on the sulfite salts (SO3\(^2-\)), efficiently producing solvated electrons (Rimmer et al., 2018; Xu et al., 2018). These solvated electrons can add to HCN, which will be reduced and subsequently hydrolyzed. Addition of excess HCN and further reduction can then form glycolaldehyde and glyceraldehyde (Xu et al., 2018). The production of solvated electrons and the subsequent chemistry that results in simple sugars are the reactions discussed in Section 3.3.

Impacts will also result in cyanamide and cyanoacetylene, either directly (Patel et al., 2015) or from further chemical alteration within a volcanic system (Rimmer and Shorttle, 2019). These and phosphates undergo a series of reactions with simple sugars to produce pyrimidine ribonucleotides,
FIG. 6. A description of the UV-driven prebiotic chemical scenario. (1) An impactor strikes the surface of Earth, generating significant amounts of PH$_3$, HCN, NH$_3$, and NO$_2$. (2) These species rain out and form a crater lake with a variety of organometallic complexes and salts, such as ferrocyanide. (3) Volcanic gas bubbles into the crater lake and UV light impinges on the lake surface, generating OH (R7, Section 3.8) and HS radicals (R1a, Section 3.2). (4, 5) The HS radicals participate in the oxidation of PH$_3$ to phosphates (R1b, Section 3.2) and the UV light generates solvated electrons from bisulfite. The solvated electrons react with the HCN and simple sugars are formed (R2, Section 3.3). These simple sugars undergo a series of reactions with cyanamide and cyanoacetylene to form nucleotide, amino acid, and phospholipid precursors (steps within this series of reactions are discussed in Sections 3.4–3.6, R3–R5). (6) The lake dries, allowing for phosphorylation of the nucleosides and formation of phospholipids. (7) Rain fills the lake with water again, and the phospholipids form into vesicles. Amino acids and precursors to RNA and DNA are also present in the solution. (8, 9) The NO$_2^-$ and nitrate salts react with ferrocyanide to form nitroprusside (R6, Section 3.7), precursors for methyl isocyanide, a nonenzymatic activating agent that leads to the formation of hybrid RNA/DNA oligomers and polypeptides.
purine deoxyribonucleotides, amino acids, and phospholipids (e.g., Patel et al., 2015), the building blocks of RNA, DNA, proteins, and cell membranes. Key reactions along the way to the selective, high-yield synthesis of these species involve UV light, and we examine these in Sections 3.4–3.6.

These building blocks will not react with each other unless activated. In biology, these molecules are activated and then joined together by enzymes. Fortuitously, chemical by-products of the UV-driven synthesis described above react together to activate amino acids, nucleotides, and phospholipids (Bonfo et al., 2019; Liu et al., 2020). This is initiated when the nitrates and nitrates, mentioned above, interact with UV light in the presence of ferrocyanide, producing nitroprusside, via the reaction discussed in Section 3.7.

Nitroprusside undergoes a further series of reactions (Mariani et al., 2018b), resulting in methyl isocyanide, which, in tandem with imidazoles (by-products of reactions involving those in Sections 3.4–3.6), perform the activation and ligation. Although this system has not been tested altogether, the projected result of the scenario would be phospholipid membranes containing a DNA/RNA hybrid system, such as that discussed in the works of Bhowmik and Krishnamurthy (2019) and Xu et al. (2020) (see Section 3.6 for details) with nonenzymatic activating agents, coupled in some way to polypeptides (as discussed broadly in Liu et al., 2020). There is much work to be done to determine the plausible next steps toward the origin of the genetic code and modern metabolism (see Wu and Sutherland, 2019).

We do not test every photochemical reaction in the network, and so, there is ample work for the future. We do test key reactions at each stage of the synthesis and activation of the building blocks of life within this UV-driven prebiotic scenario, which provides an overview of the “landscape” of prebiotic plausibility in terms of the available UV flux. In the same vein, no one need be wedded to this exact scenario, which is provided here mostly to illustrate how the chemistry can be connected. Any scenario that involves these products in an environment accessed by UV light could involve the reactions we now discuss.

3.2. Hypophosphite oxidation with bisulfite

We first consider the photochemical oxidation of hypophosphite ($\text{H}_2\text{PO}_2^-$) to phosphate ($\text{PO}_4^{3-}$) in the presence of hydrogen sulfide.

The availability of phosphorus on early Earth, both in terms of the form the phosphorus would likely take and the amount available for prebiotic synthesis, is unknown. After planetary differentiation, the small amount of phosphorus residing in the lithosphere would have been trapped almost exclusively in the form of apatite (Nash, 1984). The near-insolubility of apatite in water resulted in the so-called phosphate problem, as pointed out many years ago by Gulick (1955), although recently a solution to this problem has been proposed for carbonate-rich environments (Toner and Catling, 2019).

Soluble phosphorus can also result from the aqueous alteration of schreibersite (Tackett et al., 1966), a mineral common in iron/nickel meteorites, but this typically results in reduced forms of phosphorus such as phosphite and hypophosphite (Bryant et al., 2013; Kee et al., 2013). Impacts may plausibly convert large amounts of phosphorus into phosphine (Ritson et al., 2020), which can then be oxidized in the atmosphere—a process that still occurs on Earth today, although in trace amounts, and presumably proceeds via phosphine oxide, hypophosphite, and phosphite, eventually leading to phosphate if oxidation is complete (Glinnemann et al., 2003). As Gulick (1955) also pointed out, phosphite and hypophosphite are more soluble than phosphate in the presence of divalent metal ions (in particular calcium and iron), but would not appear to be as straightforwardly useful for prebiotic chemistry given the enhanced lability of these groups when incorporated into organic molecules and the lack of ability of hypophosphite to form phosphodiester, which would be required for an RNA or DNA analogue.

Hypophosphite can be converted to phosphate by reaction with hydrogen sulfide and UV light (Ritson et al., 2020). This is the reaction we explore here.

Hypophosphorous acid 50% wt (0.055 mmol, 6 μL) was mixed into argon-degassed 10% $\text{D}_2\text{O}/90\% \text{H}_2\text{O}$ (6 mL), and the pH was adjusted to 7.0 by using degassed NaOH. Then NaSH.$\text{H}_2\text{O}$ (0.034 mmol, 1.9 mg) was added to the mixture, and the pH was again slowly adjusted to 7.0 by using degassed HCl. This volume was divided equally into two cuvettes, one irradiated by StarLab and the other by the photoreactor RPR-200. The samples were irradiated for 1 h.

As can be seen from the nuclear magnetic resonance spectrum, shown in Fig. 7, after 1 h of irradiation with StarLab, more than 50%, but not substantially more than 50%, of the hypophosphite has been oxidized to phosphate, which suggests a chemical timescale for this conversion of 1 h, suggesting that this reaction at these concentrations would reach this point after 36–60 h on prebiotic Earth. One-hour irradiation with the RPR-200 results in nearly 100% yield.

3.3. HCN reduction with bisulfite

The next reaction we consider is as follows:

\[ \text{HCN} + \text{SO}_2 \rightarrow \text{HCN} + \text{HCN} \]

(7)

UV light generates solvated electrons from bisulfite, and these solvated electrons can reduce HCN (Xu et al., 2018). This is the first step toward the synthesis of simple sugars and nitriles (Xu et al., 2018), leading to nucleotides, amino acids, and phospholipid precursors (Patel et al., 2015). It is also the key reaction that Rimmer et al. (2018) used to delineate the abiogenesis zone. Aminomethylsulfonate is the adduct of bisulfite with the first reduced product of HCN, and the concentration of aminomethylsulfonate allows us to estimate both the yield and the rate of this reaction.

For this reaction, $^{13}$C-labeled KCN (0.15 mmol, 10.0 mg), $\text{Na}_2\text{SO}_3$ (0.60 mmol, 76.0 mg), and $\text{NaH}_2\text{PO}_4$ (0.61 mmol, 73.0 mg) were mixed in argon-degassed $\text{H}_2\text{O}$ (5 mL, 10% $\text{D}_2\text{O}/90\% \text{H}_2\text{O}$). The pH of the mixture was adjusted to 7.0 with 1 M HCl. The mixture was then divided equally into two 10 mm quartz cuvettes, one of which was irradiated at 254 nm in a photochemical chamber reactor RPR-200, while the other sample was irradiated in StarLab for the same time.
We present the resulting $^1$H and $^{13}$C NMR spectra in Figs. 8 and 9. Comparing the integration of pentaerythritol (added as a reference) and aminomethylsulfonate for the StarLab signals suggests a yield of 10% and a $k_{\text{etp}} = 1.3 \times 10^{-5}$ s$^{-1}$. We can apply this estimate to Eqs. 2 and 3 and apply the measured $\sigma_j = 1.5 \times 10^{-21}$ cm$^2$ (Rimmer et al., 2018) and the StarLab's spectral output for the sample (Figs. 3–5), to test whether the cutoff wavelength for the electron photodetachment reaction really is at 280 nm, or is at a different wavelength. We find that the cutoff wavelength is more likely at 260 nm. If the cutoff wavelength were much higher, the concentration after 1 h should be greater. This cutoff is 20 nm lower than assumed by Rimmer et al. (2018).

3.4. Photoanomerization of $\alpha$-thiocytidine

The third reaction we consider is as follows:

The synthesis of ribonucleotides from 2-aminooxazole and glyceraldehyde followed by reaction with cyanoacetylene and phosphorylation results in $\beta$-ribocytidine, one of the building blocks of RNA (Powner et al., 2007, 2009). However, this synthesis utilized a minor product of the reaction with 2-aminooxazole and glyceraldehyde, arabinose aminooxazoline (AAO, 30% yield), ignoring the major product (isomer), ribose aminooxazoline (RAO, 44% yield) (Sanchez and Orgel, 1970), which selectively crystalizes from the product solution, and allows enantiomeric enrichment during crystallization if the input glyceraldehyde is nonracemic (Anastasi et al., 2006). Reaction of RAO with cyanoacetylene followed by hydrolysis results in $\alpha$-ribocytidine, which is a stereoisomer of the RNA building block $\beta$-ribocytidine. $\alpha$-Ribocytidine can be photochemically converted into $\beta$-ribocytidine, but the conversion is inefficient, due to the competing formation of a by-product, oxazolidinone (Powner et al., 2007).

Recently, Xu et al. (2018) optimized the synthesis of canonical pyrimidine ribonucleosides/tides (genetic alphabet C and U), starting with RAO. In this synthesis, $\alpha$-anhydrocytidine which results from RAO, reacts with hydrogen sulfide to produce $\alpha$-thiocytidine, which is efficiently photoanomerized into $\beta$-thiocytidine. $\beta$-Thiocytidine either reacts slowly with water or phosphate to produce $\beta$-ribocytidine (C) and $\beta$-ribouridine (U). The above Eq. 8 is the critical step converting $\alpha$-thiocytidine to $\beta$-thiocytidine.

$\alpha$-Thiocytidine (0.015 mmol, 4.0 mg) and NaSH.xH$_2$O (0.021 mmol, 2 mg, containing >60% of NaSH) were mixed in argon-degassed H$_2$O (5 mL, 10% D$_2$O/90% H$_2$O). The pH of the mixture was adjusted to 7.0 with 1 M HCl. The mixture was then divided into two 10 mm quartz cuvettes, one of which was irradiated at 254 nm in a photochemical chamber reactor RPR-200, while the other sample was irradiated in StarLab for the same time.

As can be seen from the NMR spectra, shown in Fig. 9, the photoanomerization in StarLab after 7 h of irradiation results in a 50% yield of $\beta$-thiocytidine, and a 90% yield after 15 h of irradiation. This corresponds to a timescale of 360–600 h for...
50% yield on the surface of early Earth. If the lifetime of the relevant intermediates is on the order of this timescale, then this pathway will be plausible. The timescale can also be compared with that of the Rayonet, which achieves 50% yield in half the time, but the yield does not improve past 50%. This suggests that the less intense and more realistic broadband source can drive the reaction while suppressing or changing other reactions that destroy the product.

3.5. Attempted photoreduction of thioanhydrouridine

The fourth reaction we consider is as follows:

Many believe that the first informational macromolecule on early Earth was the RNA (e.g., Gilbert, 1986), for reasons involving the capabilities of chemical synthesis (Noorden, 2009), and evidence that certain RNA molecules are capable both of storing information and catalyzing reactions, both in the laboratory (Orgel, 2004) and in modern organisms (Cech, 2000). Finally, it would seem challenging for a system involving eight monomers, four ribonucleotides, and four deoxyribonucleotides to self-assemble into informational molecules that self-replicate with sufficient fidelity (Eigen, 1971), in the absence of macromolecular machines to chaperone, segregate, and facilitate both ribonucleotide and deoxyribonucleotide chemistries. It is worth noting that prebiotic chemists, such as Orgel (1968), were open to the possibility of DNA preceding RNA, or of both arising simultaneously.

Recently, Bhowmik and Krishnamurthy (2019) have postulated and provided some proof of concept that hybrid RNA-DNA systems can segregate themselves into homogeneous RNA and DNA macromolecules. They discovered that certain chimeric templates catalyze the ligation of oligomers, resulting in homogeneous RNA and DNA sequences.

An interesting development in prebiotic nucleoside synthesis is the recent report of the photoreduction of an RNA thionucleoside derivative, thioanhydrouridine, to a DNA thionucleoside, 2¢-deoxy-2-thiouridine (Xu et al., 2019). The discovery provides an explanation for the origin of the building blocks of DNA on early Earth alongside those of RNA, much earlier than usually thought. The reaction was proposed to proceed via photodetachment of an electron from HS\(^-\), which reduces the C2¢–S bond of the thionucleoside, followed by protonation and hydrogen atom transfer. Photodetachment was affected by the RPR-200 with 254 nm lamps, leading to the production of 2¢-deoxy-2-thiouridine in 33% yield after 3 h.

FIG. 8. \(^1\)H NMR spectrum of the photoreduction of HCN in the presence of bisulfite, using pentaerythritol as a standard for comparison, after 1 h. The concentration of product is calculated by integrating under the curve for the aminomethane sulfonate and comparing the value with the integral under the curve of pentaerythritol. (a) Using the RPR-200 reactor. The reduction product is clearly present and the area under the curve exceeds that of the pentaerythritol. (b) Using StarLab. The reduction product is far less than (a), but still visible.
We performed identical experiments in the RPR-200 with 254 nm lamps and in StarLab, to verify the plausibility of this process on early Earth. We found the result reproducible in the RPR-200, and no reaction to occur in StarLab, even with extended radiation times up to 15 h.

To a solution of 2,2'-anhydro-2-thiouridine (0.066 mmol, 16 mg, 1.0 equiv.) in argon-degassed water (2.40 mL, 10% D\textsubscript{2}O/90% H\textsubscript{2}O) was added a solution of NaSH.xH\textsubscript{2}O (60% NaSH; 70 mg, 0.38 mmol, 11 equiv.) dissolved in 2.0 mL degassed water (10% D\textsubscript{2}O/90% H\textsubscript{2}O). The pH of the mixture was adjusted to 7.0 by using degassed 1 M HCl.

The mixture was then divided into two 10 mm quartz cuvettes, each containing \(2.2\) mL with a sample height of 22 mm. One sample was irradiated at 254 nm in a Rayonet photochemical chamber reactor RPR-200 for 3 h. The other sample was irradiated in StarLab for the same time.

After 3 h, the reaction in the Rayonet reactor showed a similar outcome to that reported in the work of Xu et al. (2019), with a mixture of reduction product 2S-dU, starting material, and thiouracil being detected (24%, 64%, and 12%, respectively). The reactions in StarLab showed only recovered starting material.

Similiar experiments were conducted with longer irradiation times in StarLab of 5 and 15 h, with some of the sample being simultaneously submitted to irradiation in the Rayonet reactor for 3 h. In each case, photoreduction proceeded in the Rayonet and did not in the StarLab. Unreacted samples recovered from the StarLab trials were submitted to irradiation in the Rayonet reactor and showed some conversion. Since the photoreduction did not proceed, we cannot make any robust estimates about timescales for plausibility. We can only provide an upper limit of the timescale for this reaction of 1000 h, assuming a detection limit of 1% yield. This implies a lower limit for the timescale of this reaction on the surface of early Earth of several years.

3.6. Photoreduction of thioanhydroadenosine

A related photoreduction to the reaction discussed in the previous section is as follows:

\[
\begin{align*}
\text{Thioanhydroadenosine} & \xrightarrow{\text{hv}} \text{Deoxyadenosine} \\
& \xrightarrow{\text{hv}} 8\text{-Methoxy-deoxyadenosine}
\end{align*}
\]
Thioanhydroadenosine (0.014 mmol, 4.0 mg) and Na$_2$SO$_3$ (0.143 mmol, 18.0 mg) were mixed in argon-degassed H$_2$O (4.0 mL, 10% D$_2$O/90% H$_2$O). The pH of the mixture was adjusted to 7.0 with degassed HCl (1.0 M). The mixture was then divided into two 10 mm quartz cuvettes, one of which was irradiated at 254 nm in a photochemical chamber reactor RPR-200, while the other sample was irradiated in StarLab for the same time.

After 2 h of irradiation in RPR-200, 60% of reduced products (including 2'-deoxyadenosine and 8-mercapto-2'-deoxyadenosine) were observed in crude NMR spectra, while the conversion in StarLab was 40%. After another 2 h of irradiation, the yield of identifiable reduced products had decreased to 40%, possibly due to decomposition by the 185 nm spectral feature produced by the low-pressure Hg lamp. After the same amount of time, the yield of reduced products in StarLab had increased slightly to 43%. The results are shown in Fig. 10.

Although in this case, the kinetics does not simply allow for a simple application of Eq. 6, this equation applies an upper limit to the timescale of 180–300 h for similar chemical starting conditions to result in a yield of ~40% on the surface of early Earth, a much shorter timescale than for the DNA pyrimidines.

3.7. Nitroprusside

We now consider the following reaction:

![Reaction equation]

The synthesis of life’s building blocks is only the very beginning of the origins of life problem (Sutherland, 2017). To construct biological macromolecules from these building blocks, the synthesized molecules need to undergo oligomerization or ligation at the correct positions, typically by being activated with a chemical reagent. In modern cells, this is accomplished with enzymes, but enzymes in modern cells are synthesized by those cells. There is no clear abiotic synthetic pathway for enzymes, and so, it seems that there must be a mechanism for nonenzymatic activation of life’s building blocks.

One promising means of activating ribonucleotides, amino acids, and phospholipids involves methyl isocyanide (Bonfio et al., 2019; Liu et al., 2020), and one prebiotically plausible...
synthesis of methyl isocyanide begins with ferrocyanide and nitrites or nitrates (Mariani et al., 2018b). The immediate reaction these undergo in the presence of UV light is the formation of nitroprusside, and this is the reaction we consider here.

Several possible pathways lead from ferrocyanide ([FeII(CN)₆]⁴⁻) to nitroprusside [FeII(CN)₅NO]²⁻ (Mariani et al., 2018b). Oxidation of ferrocyanide to ferricyanide ([FeIII(CN)₆]³⁻) by NO₂⁻ yields one equivalent of nitric oxide (NO*) and one equivalent of H₂O. Subsequently, photodissociation of one cyanide ligand from ferricyanide gives [FeIII(CN)₅]²⁻, a coordinatively unsaturated intermediate in equilibrium with the aqua complex [FeIII(CN)₅H₂O]²⁻, which then equilibrates with NO* to form nitroprusside ([FeII(CN)₅NO]²⁻). Alternatively, since photolysis of NO₂ provides a hydroxyl radical (OH*) in addition to NO* (Mack and Bolton, 1999), the oxidation of ferro- to ferricyanide may be mediated by the hydroxyl radical. Clearly the production of nitroprusside from ferrocyanide involves a complex multistep mechanism and so analyzing the kinetics is more involved.

First, we investigated the formation of [FeII(CN)₅NO]²⁻ upon irradiation of a mixture of [FeII(CN)₆]⁴⁻ and NO₂⁻, as described previously (Mariani et al., 2018b). Second, as nitrate (NO₃⁻) would likely have been more abundant than NO₂⁻ on early Earth (Ranjan et al., 2019), and since photolysis of NO₃⁻ also provides NO* and OH* (Mack and Bolton, 1999), we investigated the formation of [FeII(CN)₅NO]²⁻ upon irradiation of mixtures of [FeII(CN)₆]⁴⁻ and NO₃⁻, or both NO₂⁻ and NO₃⁻.

Experiments were performed as follows: Na₄[FeII(CN)₆]·10H₂O (0.1 mmol, 48.4 mg) and NaNO₂ (0.1 mmol, 6.9 mg), NaNO₃ (0.1 mmol, 8.5 mg), or a mixture of NaNO₂ (0.05 mmol, 3.5 mg) and NaNO₃ (0.05 mmol, 4.3 mg) were dissolved in 1.0 mL of an argon-degassed 0.1 M solution of imidazole in water (10% D₂O/90% H₂O). The pH of the mixture was adjusted to 7.0 by using deoxygenated HCl (1.0 M). The mixture was then transferred to a 10 mm quartz cuvette and irradiated in either the RPR-200, with 254 nm lamps or 365 nm lamps, or in StarLab for 1 h. [FeII(CN)₅NO]²⁻ was detected by ¹³C NMR spectroscopy (number of scans: 1000; recycle delay time: 2 s; NOESY mixing time: 0 s; TOCSY mixing time: 0 s; ROESY mixing time: 0 s).

To quantify the amount of [FeII(CN)₅NO]²⁻ formed in each experiment, a standard curve was made by measuring ¹³C NMR spectra (parameters as described above) of different concentrations of Na₂[FeII(CN)₅NO] in an imidazole buffer (0.1 M) at pH 7.0. Each measurement was made three times. A plot of the integral of the [FeII(CN)₅NO]²⁻ signal at δ 134.5 ppm relative to the integral of the imidazole signal at δ 122.0 ppm, versus the concentration of [FeII(CN)₅NO]²⁻, shows a linear correlation. The ¹³C NMR spectra of the reactions described above are shown in Fig. 11 and the

FIG. 11. NMR spectra, namely (a) crude ¹H-NMR spectrum after irradiation in Rayonet (RPR-200) for 2 h; (b) crude ¹H-NMR spectrum after irradiation in StarLab for 2 h; (c) crude ¹H-NMR spectrum after irradiation in Rayonet (RPR-200) for 4 h; (d) crude ¹H-NMR spectrum after irradiation in StarLab for 4 h. As with the photoanomerization reaction (Fig. 10), the yield is better for StarLab, but the reaction is much slower.
linear regression of NMR signal ratios as a function of known quantities of nitroprusside, compared with experimental data, is shown in Fig. 12. We found that irradiation of a mixture of $\text{[Fe}^\text{II}(\text{CN})_6]^{4-}$ and $\text{NO}_2^-$ in an imidazole (0.1 M) buffer at pH 7.0 with light from StarLab for 1 h gave $\text{[Fe}^\text{II}(\text{CN})_5\text{NO}]^{2-}$ in concentrations of 26.9 – 2.7 mM for the RPR-200 (254 nm), 37.6 – 3.8 mM for the RPR-200 (365 nm), and 32.2 – 3.2 mM for StarLab. These concentrations correspond to yields of 27% for the RPR-200 (254 nm), 38% for the RPR-200 (365 nm), and 33% for StarLab.

Irradiation of a mixture of $\text{[Fe}^\text{II}(\text{CN})_6]^{4-}$ and $\text{NO}_3^-$ in an imidazole buffer (0.1 M) at pH 7.0 with light of 254 nm or light from StarLab for 1 h gave $\text{[Fe}^\text{II}(\text{CN})_5\text{NO}]^{2-}$ in concentrations of 8.1 – 0.8 mM for the RPR-200 (254 nm), 1.3 mM for the RPR-200 (365 nm) and 6.7 ± 0.7 mM for StarLab, corresponding to yields of ~10% in the cases of the RPR-200 (254 nm) and StarLab, and ~2% for the RPR-200 (365 nm). Irradiation of this mixture with light of 365 nm provided no detectable $\text{[Fe}^\text{II}(\text{CN})_5\text{NO}]^{2-}$, hence the upper limit. Finally, irradiation of a mixture of $\text{[Fe}^\text{II}(\text{CN})_6]^{4-}$, $\text{NO}_2^-$, and $\text{NO}_3^-$ in an imidazole buffer (0.1 M) at pH 7.0 with light from StarLab for 1 h also gave $\text{[Fe}^\text{II}(\text{CN})_5\text{NO}]^{2-}$ in a concentration of 29.5 ± 2.9 mM, corresponding to a yield of about 30%.

As before, we can use the yields to estimate the timescales of these reactions on the surface of early Earth. For the reaction of ferrocyanide with $\text{NO}_2^-$ or with $\text{NO}_2^-$ and $\text{NO}_3^-$, it would take 36–60 h to achieve a 50% yield of nitroprusside. With only $\text{NO}_3^-$, the timescale is on the order of a month. If ferrocyanide is stable on an anoxic early Earth on these timescales, then nitroprusside is prebiotically plausible in pools of water rich in $\text{NO}_2^-$, $\text{NO}_3^-$, or both.

3.8. Hydroxyl radical production—EtOH

Photochemistry is not only constructive. Photodetachment of electrons from anions can occur with $\text{OH}^-$ anions ever present in liquid water. The product is a solvated electron and a hydroxyl radical, and hydroxyl radicals can be devastating to life’s building blocks and prebiotic intermediate species. The cross section for photodetachment of hydride fortuitously peaks below 200 nm (Zuman and Szafir, 1976) and both $\text{CO}_2$ and $\text{H}_2\text{O}$ vapor effectively shield the surface from these wavelengths (Ranjan and Sasselov, 2016), and so, any planet with a substantial (>0.1 bar) atmosphere will be protected from these wavelengths.

We first found with the Rayonet RPR-200 reactor that photodetachment of an electron from $\text{OH}^-$ may occur above 200 nm. There is then the potential that hydroxyl radicals would be photochemically produced on early Earth but not in

FIG. 12. $^{13}$C NMR of ferrocyanide mixed with NO$_2^-$/NO$_3^-$ and irradiated, as follows. (a) Reaction in StarLab using a mixture of NO$_2^-$ and NO$_3^-$. (b) Reaction in StarLab using NO$_3^-$. (c) Reaction in Rayonet (365 nm) using NO$_3^-$. (d) Reaction in Rayonet (254 nm) using NO$_3^-$. (e) Reaction in StarLab using NO$_2^-$. (f) Reaction in Rayonet (365 nm) using NO$_2^-$. (g) Reaction in Rayonet (254 nm) using NO$_2^-$. The area under the curve for nitroprusside product can be compared with the imidazole (which acts as a standard), and the NMR ratios can be compared with nitroprusside standards, as shown in Fig. 13. $\text{NO}_3^-$, nitrate.
**FIG. 13.** Known concentrations of nitroprusside (mM) as a function of the $^{13}$C NMR peak integral ratio of nitroprusside to imidazole (see Section 3.7 for details). These clearly follow a linear relationship, and a line is fit to the data with 1σ uncertainties shown. The same ratios were calculated for the ferrocyanide irradiation data (Fig. 12), and the concentrations are estimated with 1σ errors. Color images are available online.

**FIG. 14.** Irradiation of ethanol for 2 h by the Rayonet RPR-200 (top), StarLab (middle), and left in the dark (bottom, control). No observable oxidation products after 2 h for either StarLab or the Control. Some oxidation products were visible with the Rayonet. Color images are available online.
some photoreactors typically used in the laboratory, although hydroxyl radicals may be overproduced in other laboratory reactors, such as low-pressure mercury lamps by their 184 nm emission feature. We test this possibility with StarLab. We explore the oxidation of EtOH by hydroxyl radicals in this context.

We mix EtOH (0.18 mmol, 10.5 μL) in argon-degassed water (6 mL, 10% D2O/90% H2O). The solution has a pH of 7.5. The mixture is divided into three cuvettes, one irradiated by StarLab for 2 h, one irradiated by the RPR-200 for 2 h, and a third that is not irradiated. The third cuvette is a control in the event that the argon degassing is not sufficient to prevent some oxidation of EtOH. We dissolved pentaerythritol (25 μmol, 3.4 mg) in argon-degassed H2O (0.5 mL) resulting in a 50 mM solution. After irradiation, we added the pentaerythritol solution (166 μL) to each of the three samples as a standard to estimate the concentrations of the oxidation products.

The 1H NMR results show no detectable irradiation products of EtOH, such as acetate, acetaldehyde, and 1,1-ethanediol for either the control cuvette or the StarLab-irradiated cuvette. The RPR-200-irradiated cuvette shows some photooxidation products, likely due to the 184 nm feature (Fig. 13).

We repeated the experiment, but with EtOH (0.36 mmol, 21 μL), and we irradiated the mixture with StarLab for 24 h (Figs. 14 and 15). Some oxidation products were observed. By integrating under the spectral curves for the oxidation products and for pentaerythritol, we can estimate the concentration of the oxidation products.

Taking the total integral of the oxidation products to be 1.88, when accounting for the 1H degeneracies and duplicate peaks, and the total integral of pentaerythritol is 0.71, after accounting for the 1H degeneracies for pentaerythritol (degeneracy of 8). The concentration of the oxidation products is then as follows:

$$c_{\text{ox}} = \frac{c_0 I_{\text{ox}}}{I_0}$$

where $c_{\text{ox}}$ is the concentration of the oxidation products, $I_{\text{ox}}$ is the integral of the lines for the oxidation species, adjusting for degeneracies and duplicate peaks, $c_0$ is the concentration of the pentaerythritol, and $I_0$ the integral of the pentaerythritol feature, accounting for degeneracies.

We find that $c_0 = 0.002 \text{ M}$, $I_0 = 0.71$, and $I_{\text{ox}} = 1.88$, so $c_{\text{ox}} = 0.005 \text{ M}$. Since the starting concentration of EtOH was $0.06 \text{ M}$, this is a yield of 8%. We apply this yield to Eq. 6 to estimate the time it would take to achieve a yield of 50% of 290 h with StarLab, or more than a year under the light of the young Sun.

4. Discussion

Using the results from the beginning of Section 3, we can draw out a relationship between the StarLab spectral irradiance and the UV actinic flux on the surface of early Earth. That relationship is wavelength dependent, because the spectrum of the young Sun is not identical to the StarLab spectrum, even considered at low resolution. Also, the atmosphere and the local aqueous chemistry can affect the UV

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**FIG. 15.** Irradiation of ethanol by StarLab for 24 h. Now some products are detected and their line integrals can be compared with the line integral for pentaerythritol to estimate the concentrations of the oxidation products. Color images are available online.
environment. We made the assumption that the atmosphere is composed of 0.3 bar N₂/CO/CO₂ with H₂O at vapor pressure, which is mostly transparent to UV light above 200 nm.

We also assume an aqueous environment, as may be found in a mineral-rich postimpact crater pond with ferrocyanide at concentrations of >0.1 mM, which at shallow depths of a few mm if rich in ferrocyanide blocks UV light below 220 nm (Higashi and Ozaki, 2004; Chakrabarti and Roberts, 2008). This environment is approximated with the use of mineral water (Cambridge tap water) within the quartz attenuator into which the experimental samples are placed. To be clear, the tap water only attenuates the UV light and is not in contact with the samples themselves.

We show the relationship between the experimental timescales and timescales for reactions on the surface of early Earth as a function of wavelength in Fig. 16. Our results agree reasonably well with those of Ranjan and Sasselov (2016).

Of course, the atmosphere of early Earth when life arose may have been different than we assume. It may have been more reducing, and if so, there would likely have been thick photochemical hazes that can obscure the UV light between 200 and 300 nm (Arney et al., 2016). If life’s origins were coincident with the tail end of accretion, hazes generated by accretionary impact may obscure the UV light (Rimmer et al., 2020). NH₃ would also attenuate UV light below 230 nm (Cheng et al., 2006). The results in this article provide timescales under optimal conditions for surface prebiotic photochemistry.

The surface UV flux predicted by Rimmer et al. (2018) does not match the surface fluxes predicted by Ranjan et al. (2017), and this is due to our assignment of a < 1 single-scattering albedo when calculating Rayleigh scattering. The result has been to effectively include a small particulate haze and overattenuate the short-wavelength NUV compared with a clear sky model. Using the surface UV flux predicted by Rimmer et al. (2018) with single-scattering albedo set to unity, to better approximate clear skies, we find any reaction that achieves a yield of 50% after a 12-h exposure to StarLab meets the abiogenesis zone requirements of Rimmer et al. (2018) for the young Sun. Future work will investigate the implications of these experimental results for abiogenesis zones around other stars. The timescale related to the abiogenesis zone of Rimmer et al. (2018) is shown in Fig. 16.

This abiogenesis zone affords a “ballpark” estimate for degradation timescales, based on a chemist’s experience with these reagents in the laboratory, many of the “stable” intermediates tend to have lifetimes comparable with HCN and bisulfite, although lifetimes for these species depend on temperature and pH and can range over many orders of magnitude (see, e.g., Miyakawa et al., 2002).

To genuinely test whether these reactions would work on early Earth or around another planet, the degradation chemistry needs to be explored over a range of conditions, and applied to the analysis, to determine whether, in the context of a particular scenario, a given reaction is likely to occur with reasonable yield on early Earth, on the basis of the available UV light. Nevertheless, constraints on plausibility can be made based on the known stability of the reactants in the laboratory. Reactants that cannot survive more than a few days must achieve high yields in a matter of a few hours under StarLab, or the degradation will win out, and the path is not prebiotically plausible.

We explore the prebiotic plausibility of six different reactions (Sections 3.2–3.8), and summarize our results in Table 1. We find that most of the reactions, the photooxidation of hypophosphite, the photochemical reaction of HCN and bisulfite, the photoanomerization of thiocytidine, and the conversion of ferrocyanide to nitroprusside, all appear plausible from the standpoint of UV light, although the timescale for photoanomerization of thiocytidine (approximately 360–600 h on early Earth) suggests that the yield for this reaction may be suboptimal, and that further exploration into alternative mechanisms for producing β-ribocytidine is warranted.

In particular, we found that nitroprusside is plausible in water with NO₃⁻, even if no NO₂⁻ is present, and that the photodestruction of nitroprusside by light at short wavelengths is outcompeted by the photochemical production of nitroprusside, likely owing to the spectral shape of StarLab and of the young Sun, producing much more light at longer wavelengths (i.e., >254 nm). This effect will be further enhanced if short wavelength light is attenuated by the water, as it was the case for <220 nm with StarLab. This attenuation may have other consequences for the rest of the network, and opens up the possibility for chemical temporal segregation within the same body of water, where one region of water experiences a different chemical environment. If there is inefficient mixing, or if there are ways to segregate different parts of a pool (pores or vesicles), chemical heterogeneity and relevant chemical gradients will be generated within a pond simply by UV attenuation. Further investigation into this possibility is needed.

We finally turn to the photoreduction of thioanhydrouridine with hydrogen sulfide, photoreduction of thioanhydroadenosine with bisulfite, and the production of hydroxyl radicals from water, quantified via the subsequent oxidation of EtOH. The photoreduction of thioanhydrouridine and production of hydroxyl radicals from water are not plausible. In the case of oxidation of EtOH, this implies that the hydroxyl production by the young Sun on early Earth is very slow, and should not interfere with the prebiotic chemistry.
The failure to photoreduce thioanhydridouridine allowed us to place a lower limit on the timescale for this photochemistry of several years. Even this lower limit may be optimistic, however, considering that it assumes the reaction would occur given longer irradiation times. This may be implausible because the presence of light at other wavelengths may have opened up alternative interfering pathways, or may have altogether disrupted the mechanism allowed under the Rayonet RPR-200.

Finally, the related photoreduction of thioanhydroadenosine, a similar nucleoside derivative, was successful when using bisulfite as the source of solvated electrons. This reaction worked rapidly in StarLab, achieving the final ∼40% yield in 2 h. This suggests that bisulfite is a more plausible source of solvated electrons than hydrogen sulfide, and that more than one step in the prebiotic scenario we consider requires the presence of bisulfite in order for the UV light to be prebiotic.

Future investigations of the prebiotic plausibility of this scenario should focus on the contingency of various reactions in sequence, with a demonstration that these reactions can run in sequence, either without the aid of the experimentalist or with that aid explained by a plausible physical process.

The intervention required to complete the chemistry can be used to estimate the probability that this chemistry could have taken place on early Earth, early Mars, or on other rocky planets outside our solar system. In addition, careful investigation into chemical mechanisms, such as the work done when using pump-probe techniques from the works of Roberts et al. (2018) and Todd et al. (2019), will be key for understanding why some reactions behave differently under a broadband source than under a single wavelength source. This understanding will also be essential for discovering alternative pathways once certain pathways are falsified in terms of the available UV light.

### 4.1. Other effects of stellar irradiation and activity

Stellar irradiation and activity have other effects on planetary atmospheres and the surface environment (for a review of stellar activity through time, see Güdel 2007). Although UV light at wavelengths <200 nm cannot reach the surface of a planet with a >0.1 bar CO₂/N₂ atmosphere, this light drives atmospheric escape (discussed in Gronoff et al., 2020), and can erode atmospheres and can transform them. If the atmosphere is fully eroded, the planet will not be habitable.

Stellar activity can also result in the abiogenic buildup of molecular oxygen (Tian et al., 2014), which would interfere with several prebiotic chemical reactions, including the ones discussed here, which generally require anoxic conditions, and any scenario that is inhibited or disrupted by molecular oxygen, for example, chemistry that makes use of cysteine-containing peptides (Bonfio et al., 2017). Energetic photons are not all bad. They can lead to the formation of HCN, although in relatively small quantities (Zahnle, 1986; Tian et al., 2011; Rimmer and Rugheimer, 2019), and of formaldehyde (Pinto et al., 1980), precursors for the UV-driven prebiotic scenario we investigate here.

Energetic particles can penetrate deeper into the atmosphere and may cause more severe problems for atmospheric evolution. Energetic particles, produced from coronal mass ejections that accompany flares, are more relevant for surface chemistry, since their chemical impact lies within the stratosphere and troposphere, rather than being relegated to the thermosphere and mesosphere (Airapetian et al., 2016). These solar energetic particles generate greenhouse gases, which may affect the surface temperature of early Earth and generate substantial amounts of HCN (Airapetian et al., 2016).

Solar energetic particles can themselves drive some prebiotic chemistry in the gas phase (Kobayashi et al., 1998). The light of flares may be relevant for the >200 nm UV environment on the surface of rocky exoplanets (see Section 4.2), but are not relevant for early Earth. We can see this by comparing the energy deposited by the light of the Sun with the energy deposited by a high-energy flare that the active cool star AD Leo produced (Hawley and Pettersen, 1991). This flare has energy 10× the energy of the Carrington event (Cliver and Dietrich, 2013), and is thought to be at the upper limit of flare energies for the Sun (Cliver and Dietrich, 2013). As can be seen from the work of Segura et al. (2010), the intensity of this flare only achieves the quiescent emission of the young Sun.

### Table 1. Timescales for Key Prebiotic Photochemical Reactions on the Surface of Early Earth, Accounting for the Light of the Young Sun

| Reaction                        | Time irradiated in the laboratory (h) | Yield in the laboratory | Irradiation timescale | Early Earth timescale |
|---------------------------------|--------------------------------------|------------------------|-----------------------|----------------------|
| Oxidation of hypophosphite      | 1                                    | 50%                    | 18–30 h               | 36–60 h              |
| HCN+bisulfite                   | 1                                    | 10%                    | 90–150 h              | 180–300 h            |
| Photoanomerization of thiothymidine | 7                                    | 50%                    | 180–300 h             | 360–600 h            |
| Photoreduction of thiothyminodeoxyribonucleosine | 15                        | <1%                    | >5 Years              | >10 Years            |
| Nitroprusside synthesis         | 1                                    | 30%                    | 18–30 h with nitrite, | 36–60 h with nitrite,|
|                                 |                                       | 10% with nitrate,      | ~720 h without nitrate| ~1440 h without nitrate|
| Oxidation of ethanol            | 24                                   | 8%                     | >1 Year               | >2 Years             |

*The time the sample must be irradiated on the surface of early Earth to obtain a yield of 50%.

bThe time the sample must be present on early Earth to obtain a yield of 50%, accounting for the day/night cycle.

HCN = hydrogen cyanide.
4.2. Implication for exoplanets

One great advantage of this scenario is that it depends on starlight, and starlight is the property of exoplanet systems that is best known. As discussed above, the measurement of the rates of these reactions under realistic conditions in the laboratory, and the comparison of these rates with degradation rates for the reactants, allows us to investigate whether these reactions are likely to take place on the surface of planets around other stars. To investigate this chemistry in the context of exoplanets, changes to the StarLab configuration, as presented in this article, will be implemented.

To investigate the chemistry that may be enabled by including the 200–220 nm UV wavelengths in our current broad wavelength range, we use ultrapure water in the quartz attenuator. As already discussed above, most plausible aqueous environments on early Earth would not include these shorter wavelengths, but to investigate chemistry that could happen on the actual surface layer, this band of the UV would be available, and these microenvironments should also be explored in the future.

Looking beyond the environment on early Earth, we also plan to simulate the stellar flux environment as may be experienced on exoplanet surfaces around other stellar types to experimentally constrain the abiogenesis zone based not on one reaction, but based on the chemistry as a whole, using a realistic light source that accurately represents the diverse spectra of these stars.

In addition, we look into the effects of catalysts on the UV-driven chemistry, for example, the catalytic role of ferrocyanide in the cyanide reduction by UV-irradiation of bisulfite. This is especially interesting for M dwarf stars—due to their smaller size and mass, Earth-like, rocky planets are more easily detected compared with Earth-like planets around solar-like stars.

A number of Earth-like exoplanets around M dwarfs have already been discovered, such as the TRAPPIST-1 system (Gillon et al., 2017) and planets similar to these will be good candidates for atmospheric characterization with the James-Webb telescope. It is therefore interesting to investigate if UV-driven origin chemistry on such planets is plausible—on these very red M stars sufficient UV flux only comes from flares. In a future article we will present a reconfiguration of StarLab to simulate the flux environment (including periodic flares), as may be experienced on planets around M stars, and determine if any of the chemical pathways presented here are possible in such a nonconstant UV flux environment, and to provide an update to the abiogenesis zone based on stellar activity.

5. Conclusions

We constructed a stellar simulator that accurately reproduces a multiple of the broadband actinic flux of the young Sun on the surface of early Earth. We then performed a series of seven experiments with StarLab exploring the various stages of a prebiotic chemical scenario with UV-driven aqueous chemistry starting with cyanide, bisulfite, and sulfides, with a focus on the role of the UV light. We first considered the availability of phosphate, an essential feedstock molecule for subsequent prebiotic synthesis, by exploring the photooxidation of hypophosphite with hydrogen sulfide. Then we explored the first steps of the synthesis, which commences with HCN and bisulfite. Next, we turned to a key later synthetic step toward cytidine and uridine, the photoanomerization of theophytidine.

Then we attempted the photoreduction of thioanhydrouridine, a key step in the prebiotic synthesis of deoxynucleotides. This photoreduction failed in StarLab, allowing us to place a lower limit on the timescale for this reaction on early Earth, assuming longer exposures would result in the expected product. In Section 4, we present some reasons why this is unlikely to be the case, and that it is more likely new destructive chemical mechanisms have been initiated due to the broadband nature of the light and the nonlinear nature of the chemistry. We also demonstrated the successful photoreduction of thioanhydroadenosine in the presence of bisulfite, indicating that bisulfite is a necessary condition for the sufficiency of solar UV light for prebiotic photochemistry.

Next, we considered the production of nitroprusside, the first step in the prebiotically plausible production of methyl isocyanide. Methyl isocyanide can be used as a component in the nonenzymatic activation of nucleotides, amino acids, and phospholipid precursors. Finally, we considered destructive hydroxyl radical production by exploring the oxidation of EtOH by UV-generated hydroxyl radicals in water.

We discovered that the photooxidation of hypophosphite, the reaction of HCN with bisulfite, and the production of nitroprusside are all prebiotically plausible, and the photoanomerization of theophytidine is at the edge of plausibility. We additionally discovered that nitroprusside can be produced from ferrocyanide and NO3− alone, and that the rate of this conversion suggests that nitroprusside is prebiotically plausible in both NO3−-rich and NO3−-rich water, so long as the water also has access to ferrocyanide and UV light.

We found that two reactions, photoreduction of thioanhydrouridine and oxidation of EtOH, are not prebiotically plausible. In the former case, there are alternative pathways to RNA pyrimidines and DNA purines that do succeed, although with the use of bisulfite instead of hydrogen sulfide as the source of free electrons. In the latter case, this is a positive outcome for the prebiotic chemistry, because the results of the EtOH oxidation experiment suggest that timescales for producing hydroxyl radicals in water on the surface of early Earth are likely far longer than the lifetimes of even stable intermediates. However, if certain building blocks need to survive for more than a matter of decades, then the production of hydroxyl radicals may become significant.

One advantage of exploring prebiotic plausibility in terms of starlight is that starlight is what exoplanet scientists observe first and know best. Starlight is also one of the best constrained global parameters for early Earth, although even here uncertainty remains (Spalding et al., 2018). Constraining prebiotic plausibility by using starlight will apply to all the local environments on early Earth, and the constraints can be made with a higher level of confidence than those for other parameters such as chemical composition or local temperature or pH.

These constraints can also be transformed into conditions for further delineating the abiogenesis zone for the Sun, and for other stars, including for ultracool stars. This is especially relevant given our present knowledge about the UV environments of planetary systems around ultracool host stars such as TRAPPIST-1 (Ducrot et al., 2020) and systems discovered by TESS (Günther et al., 2020), and future discoveries around these systems, as well as around young exoplanet systems (Bottrill et al., 2020; Rimmer et al., 2020). If biosignatures are discovered on a statistically
significant subset of these planets, the distribution of that subset may provide the only real way to test prebiotic chemical scenarios in the future.

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Abbreviations Used
EtOH = ethanol
HCN = hydrogen cyanide
NH₃ = ammonia
NO = nitric oxide
NO₂ = nitrogen dioxide
NO₂⁻ = nitrite
NO₃⁻ = nitrate
OH⁻ = hydroxide
PH₃ = phosphine
RAO = ribose aminooxazoline
UV = ultraviolet

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