## The Early Faint Sun Paradox: Organic Shielding of Ultraviolet-Labile Greenhouse Gases

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Atmospheric mixing ratios of  $\sim 10^{-5 \pm 1}$  for ammonia on the early Earth would have been sufficient, through the resulting greenhouse warming, to counteract the temperature effects of the faint early sun. One argument against such model atmospheres has been the short time scale for ammonia photodissociation by solar ultraviolet light. Here it is shown that ultraviolet absorption by steady-state amounts of high-altitude organic solids produced from methane photolysis may have shielded ammonia sufficiently that ammonia resupply rates were able to maintain surface temperatures above freezing.

 $\mathbf{T}$ he compositions and surface pressures of the early atmospheres of Earth and Mars after the end of the heavy bombardment are unknown. Models for Earth range from reducing atmospheres with surface pressures  $p_s$ ~ 1 bar (1) to  $p_s \sim 10$  bar of CO<sub>2</sub> (2). Atmospheres near a neutral oxidation state are currently favored for two reasons. First, a vast reservoir of  $\mathrm{CO}_2$  is present in Earth's sediments in the form of carbonates ( $p_s \simeq$ 60 bar) and in the atmosphere of Venus ( $p_s$  $\simeq$  90 bar). This carbon reservoir could have been in the early atmosphere partly as CO or CH<sub>4</sub> if some abundant reducing agent were present. However, metallic iron, one possible agent, is thought to have been segregated from the surface early and quickly during core formation (3), suggesting that carbon was present in the atmosphere primarily as CO2. The presence of other reduced gases such as NH3 also depends on the oxidation state of the upper mantle. If early continents were largely absent, the resulting inhibition of carbonate sedimentation may have led to dense CO<sub>2</sub> atmospheres; Kasting (4) estimates that  $p_s(CO_2)$ was 0.1 to 10 bar 4 billion years ago (Ga). However, it is possible that the mantle may be more reducing than once thought (5), and geochemical arguments for early atmospheric composition are not yet conclusive. A second argument for an early atmosphere that was in a neutral oxidation state comes from photochemical kinetics, which suggests that atmospheric NH<sub>3</sub> and CH<sub>4</sub> would be photodissociated by solar ultraviolet (UV) radiation on short time scales (6-9). Here we argue instead that photodissociation of CH<sub>4</sub> may have led to the production of a high-altitude organic aerosol that ab-

sorbed UV rays and allowed reducing gases to exist for much longer times. If this is correct, the second of the two arguments against reducing gases in the early atmosphere is no longer compelling, with implications for the origin of life and the resolution of the early faint sun paradox.

The origin of life requires sources of preexisting organic molecules, which are much more readily generated in reducing rather than in neutral oxidation-state atmospheres (10). With allowance for exogenous sources, reducing atmospheres give organic mixing ratios in the early oceans some three orders of magnitude higher than those for neutral atmospheres (11). Thus, models for Earth's primitive atmosphere that postulate no significant abundance of reduced gases suffer the drawback that the origin of life is more difficult to understand.

### The Early Faint Sun Dilemma

The equilibrium temperature of an airless, rapidly rotating planet is  $T_e \equiv [SR^2(1 - T_e)]$ A)/ $4a^2\varepsilon\sigma$ ]<sup>1/4</sup>, where  $\sigma$  is the Stefan-Boltzmann constant,  $\varepsilon$  the effective surface emissivity, A the wavelength-integrated Bond albedo, R the planet's radius, a the planet's distance from the sun, and S the solar constant at *a*. For Earth today,  $T_e \simeq 255$  K, and the difference  $\Delta T \simeq 33$  K between  $T_{\rm e}$  and the observed mean surface temperature  $T_s$  is due to the mainly H<sub>2</sub>O-CO<sub>2</sub> atmospheric greenhouse effect. Stars like the sun increase their luminosity through time as the hydrogen in the star's core is converted to helium. Estimates of the change in S between 4 Ga and today are  $\approx 0.25S_0$  (12), with an error of about  $\pm 0.1S_0$ , where  $S_0$  is the present-day value. Holding A,  $\varepsilon$ , a, and the CO<sub>2</sub> mixing ratio constant at their current values for Earth, and letting the  $H_2O$ abundance be determined by its equilibrium vapor pressure, Earth's  $T_s < 0^{\circ}$ C about 2 Ga (13-15). Allowance for an ice-albedo feedback implies a fully glaciated planet as recently as 1 Ga (16). These results contradict geological evidence for abundant, globally distributed liquid H<sub>2</sub>O before 3 Ga (13, 14, 17, 18). The discrepancy is enhanced by evidence that before 2.5 Ga, Earth may have been considerably warmer than it is today (19). The implications are even more severe for Mars; yet there is strong evidence for liquid water on Mars 3.5 to 4.0 Ga (20).

This early faint sun paradox is only a paradox if we insist on holding A, a,  $\varepsilon$ , and greenhouse warming constant over time. Were A = 0.3 at 4 Ga, as it is today,  $T_e$ would have been 237 K (21); even if A had been 0 at 4 Ga,  $T_{\rm e}$  would still have been only 259 K. In models of an all-ocean early Earth with a 14-hour day, simulated temperatures rise only an additional 5.5 K (22). Likewise, at most a few degrees can be pried out of plausible long-term variations in infrared emissivity. Although some low-mass planets can chaotically migrate in heliocentric distance a over 4 billion years (23), there is no evidence for the steady increases in a for Earth by the  $\sim 10\%$  necessary to have compensated for the increase in S over its history. Another suggestion proposes that the early sun lost  $\sim 10\%$  of its mass, producing a solar flux at Earth before  $\sim 4$ Ga greater than its current value (24); however, there is as yet no clear evidence for such mass loss (25). Convolving the rapid rotation of Earth, an all-ocean planet, extremely low albedo, and a small solar mass loss, it is possible that early Earth reached 0°C but not much warmer temperatures. Such a conglomerate explanation seems strained. We therefore reconsider changes in greenhouse warming.

The Wien peak of Earth's blackbody emission 4 Ga, as today, laid at 8 to 14  $\mu$ m, a window in the combined absorption spectra of H<sub>2</sub>O and CO<sub>2</sub> at pressures below a few bars and temperatures < 300 K. Because small quantities of NH<sub>3</sub> provide considerable opacity around 10 µm, Sagan and Mullen (13, 14) proposed a volume mixing ratio for NH<sub>3</sub> ([NH<sub>3</sub>]) of  $\sim 10^{-5}$ , pressurebroadened in an  $\sim$ 1-bar atmosphere, as the solution to the early faint sun dilemma for Earth and Mars. Subsequent calculations (6, 15) confirm that  $[NH_3] \sim 10^{-5 \pm 1}$ would suffice. In comparison, background abundances in Earth's current O2 atmosphere are  $[NH_3] \sim 10^{-8}$  (26). These NH<sub>3</sub> abundances represent a steady state after

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loss processes, including oxidation, rainout, high-altitude photolysis, and reaction with CO<sub>2</sub>. Globally averaged,  $[\rm NH_3] \sim 10^{-2}$   $[\rm CH_4]$  today.

Although NH<sub>3</sub> is soluble in water and subject to rapid rainout (27), there is, in the absence of photolysis, an equilibrium atmospheric abundance determined by oceanic mineral equilibria. Independently, a lower limit can be derived from the requirement that the deamination of aspartic acid, a key amino acid for the origin of life, be reversed. Bada and Miller (28) thereby found [NH<sub>3</sub>] for early Earth to be  $10^{-7}$  at  $0^{\circ}$ C,  $3 \times 10^{-5}$ at 25°C, and  $3 \times 10^{-4}$  at 50°C, consistent with the  $10^{-5 \pm 1}$  derived from the greenhouse requirement. Other estimates lie in a similar range (29).

All organisms on Earth are able to metabolize  $NH_4^+$  (30). Few organisms can process  $N_2$  directly, and ubiquitous metabolic facility with  $NH_4^+$  may be a remnant from earlier times when atmospheric  $NH_3$ was readily available (31). These chemical and biological arguments independently point to  $NH_3$  as a minor constituent of the primitive atmosphere.

## NH<sub>3</sub> Photolysis and CO<sub>2</sub> Greenhouses

Photochemical kinetics demonstrates that  $[NH_3] \sim 10^{-5}$  would be photodissociated by the solar UV continuum at wavelengths  $\lambda < 2300$  Å and irreversibly converted into N<sub>2</sub> within decades (6, 8, 9). Likewise,  $[CH_4] \sim 10^{-5}$  would be photolyzed within centuries. These time scales are so short that hypotheses about resupply (13, 14) seem wholly without merit; an NH<sub>3</sub> outgassing rate  $\sim 5 \times 10^{15}$  g year<sup>-1</sup> would be required to keep pace with photolysis, which implies  $\sim 10^3$  bar of N<sub>2</sub> built up over  $\sim 1$  billion years, contradicting current atmospheric and sedimentary rock inventories.

Owen et al. (8) proposed that massive amounts of CO<sub>2</sub> might close the 8- to 14µm window and provide the missing greenhouse effect. Numerous calculations have since shown that a few bars or less of  $CO_2$ might suffice to raise the surface temperature of Earth 4 Ga to several tens of degrees Celsius (15, 32). But Rye et al. (33) argue from the absence of siderite in paleosols that [CO<sub>2</sub>] between 2.75 and 2.2 Ga was  $\leq 10^{-1.4}$  bar, implying that greenhouse gases other than CO2 contributed to greenhouse warming in the late Archean and early Proterozoic eons. For Mars, there is extensive evidence for ancient outflow channels and valley networks (20), some evidence for lakes (34), and even an argument for oceans (35) 3 to 4 Ga. The general opinion has been that this evidence requires a massive early greenhouse effect on Mars (14, 15, 36), but the higher early subsurface thermal gradient eases the greenhouse demands (37). Massive  $CO_2$  greenhouse models face challenges from  $CO_2$  condensation (38) and the difficulty in finding sufficient corresponding martian carbonate sediments (39).

#### Ultraviolet Shielding

Given doubts about massive CO<sub>2</sub> atmospheres for Earth and Mars, we reexamine models of comparatively thin atmospheres in which minor constituents provide the needed infrared opacity. We ask whether, with no ad hoc assumptions, there is a way to shield NH<sub>3</sub> and other UV-labile gases so that demands on the resupply rate are reasonable. Methane can be dissociated only by UV photons of  $\lambda < 1450$  Å (the dissociation is therefore entirely dominated by solar Lyman  $\alpha$  radiation), whereas NH<sub>3</sub> photolysis is driven by UV at wavelengths as long as 2300 Å. This longer wavelength UV, which in total energy flux exceeds that of  $\lambda < 1450$  Å UV by more than two orders of magnitude, penetrates deeper into the atmosphere. The result is that whereas CH<sub>4</sub> photolysis is expected to occur high in the atmosphere (40, 41), photolysis of NH<sub>3</sub> should occur at a lower altitude, with NH<sub>3</sub> photolysis products spatially separated from those of  $CH_4$  (6).

Photochemical modeling indicates that NH<sub>3</sub> photolysis, for [NH<sub>3</sub>] ~  $10^{-5 \pm 1}$ , peaks at altitudes of 25 to 35 km, depending on [NH<sub>3</sub>] and the assumed eddy diffusion coefficient (6). Zahnle (42) has modeled the altitude at which [CH<sub>4</sub>] photolysis and polymerization should occur as a function of [CH<sub>4</sub>], [CO<sub>2</sub>], UV flux, and eddy diffusion profile. For contemporary values of the eddy diffusion profile and [CO<sub>2</sub>], he finds the altitude of CH<sub>4</sub> polymerization ranging from 40 to 75 km (43) as [CH<sub>4</sub>] varies from



Fig. 1. Solar UV flux at Earth (solid line) and the absorption cross section of NH<sub>3</sub> (dashed line) as a function of wavelength. With the exception of the Lyman  $\alpha$  spike at 1216 Å, solar UV flux drops quickly shortward of 2000 Å, whereas NH<sub>3</sub> absorption falls rapidly longward of 2000 Å.

 $10^{-5}$  to  $6 \times 10^{-4}$  (corresponding to CH<sub>4</sub> fluxes into the atmosphere ranging from  $5 \times 10^{10}$  to  $\sim 4 \times 10^{11}$  cm<sup>-2</sup> s<sup>-1</sup>). In all cases, he takes [H<sub>2</sub>O] at the base of the stratosphere to be  $4 \times 10^{-6}$ ; that is, CH<sub>4</sub> polymerization is not prevented by OH generated from UV dissociation of H<sub>2</sub>O (44). However, a transition from oxidation of CH<sub>4</sub> to polymerization occurs when the local C/O ratio passes through unity, at which point large percentages of CH<sub>4</sub> are polymerized. The CH<sub>4</sub>/CO<sub>2</sub> ratio is therefore important.

Organic solids from atmospheric shocks (as well as interplanetary dust particles) contribute to a high-altitude layer of organic aerosols, depending on atmospheric composition (11). If  $CH_4$  photolysis and other processes produce an organic haze, which in turn provides UV shielding to greater depths (the C<sub>2</sub> hydrocarbons absorb at considerably longer wavelengths than  $CH_4$ ),  $NH_3$  will lie below this screen and be substantially shielded from photolysis. Experimentally,  $N_2$ -CH<sub>4</sub> gas mixtures with [CH<sub>4</sub>] ~ 10<sup>-3</sup> generate upon irradiation complex organic solids with strong UV absorption in the 2000 Å region (45). Destruction of  $NH_3$  by reaction with OH (9) should also be greatly reduced, because photolysis of H<sub>2</sub>O from photons with  $\lambda$  < 2400 Å will be suppressed. Submicrometer aerosols should settle out of the stratosphere on time scales  $t \simeq$ 0.5 to 3 years (46), fairly independent of the atmospheric pressure on early Earth;  $t \simeq 1$ year applies to the present Earth ( $p_s = 1$  bar, gravitational acceleration g = 980 cm s<sup>-2</sup>) and Venus ( $p_s = 90$  bar, g = 890 cm s<sup>-2</sup>).

To calculate the prolongation by shielding of the  $\sim$ 10-year photodissociation time scales of unshielded NH<sub>3</sub>, we must evaluate two photodestruction rates. The photodestruction rate in the absence of any shielding is  $J = \int I_{\lambda} \sigma_{\lambda} d\lambda$ , where  $I_{\lambda}$  is the solar flux at the top of the atmosphere (47) and  $\sigma_{\lambda}$  is the  $NH_3$  absorption cross section (9). If a high-altitude UV shield of optical depth  $\tau_{\lambda}$ is present, this rate becomes J' = $\int I_{\lambda} e^{-\tau_{\lambda} \sec \theta} \sigma_{\lambda} d\lambda$ , where  $\theta$  is the solar zenith angle; we take  $\theta = 45^{\circ}$ , the mean angle. These integrals effectively extend only from 1100 to 2300 Å (Fig. 1), because the solar flux is negligible below 1100 Å and NH<sub>3</sub> absorption is negligible above 2300 Å. The ratio J/J' therefore gives the enhancement of the lifetime of atmospheric NH<sub>3</sub> in the presence of a UV shield of optical depth  $\tau_{\lambda}$ . Because  $\sigma_{\lambda}$  is increasing steeply and the Planck function is decreasing steeply at 2000 Å, these integrals are given approximately by their values near 2000 Å. Before performing the full numerical integrations, we first examine such an approximation.

All organic solids strongly absorb at  $\lambda < 2300$  Å. The optical depth is given by  $\tau_{\lambda} =$ 

 $\phi_{\gamma} t \alpha_{\lambda} / 4 \pi \rho R^2$ , where  $\phi_{\gamma}$  is the production flux (mass per time) of UV-shielding organic solids, *R* is the planet's radius,  $\rho$  is the bulk density of the organic solids,  $\alpha_{\lambda} =$  $4\pi k / \lambda$  is the linear absorption coefficient of the organic solids, and *k* is the imaginary part of the complex refractive index (Table 1). We choose a conservative value of  $k(2000 \text{ Å}) \approx 0.19$ . The higher the proportion of elemental carbon, the higher *k* will be. We adopt  $\rho \approx 1.4$  to 1.6 g cm<sup>-3</sup>, the range for the organic solids produced at low pressures from the radiation chemistry of 10:1 N<sub>2</sub>-CH<sub>4</sub> gas mixtures (48).

Syntheses of organics in CH<sub>4</sub>-N<sub>2</sub> models of Titan's atmosphere imply a production efficiency of  $1.2 \times 10^{-12}$  g erg<sup>-1</sup> at  $\lambda <$ 1550 Å (49). This value is within a factor of 2 of the photodissociation efficiency of CH<sub>4</sub> in pure CH<sub>4</sub> atmospheres at  $\lambda \approx 1295$  and 1470 Å (50). We calculate organic production efficiencies in UV-irradiated CH<sub>4</sub>-NH<sub>3</sub> atmospheres (51) to be a factor of ~5 below those just cited for CH<sub>4</sub>-N<sub>2</sub> atmospheres.

Most of the products of charged-particle irradiation of  $CH_4$ -N<sub>2</sub> atmospheres are in the form of organic heteropolymer (52). A lower limit on the fraction *f* of the irradiation products that are organic solids is taken to be 0.1 (52). We assume a similar range for the products of  $CH_4$ -N<sub>2</sub> photolysis. We estimate the net flux of UV energy below 1500 Å on Earth 4.0 Ga to have been 1 × 10<sup>27</sup> erg year<sup>-1</sup> (11, 53), three times the current value. The production of high-altitude organic aerosols from  $CH_4$  photolysis in the terrestrial atmosphere 4 Ga was then  $\phi_{\gamma} \approx (1.2 \times 10^{15}) f g year^{-1}$ . Other sources of high-altitude organic aerosols (11) amount to  $\approx 10^{13} f g year^{-1}$ .

This calculation assumes the production of organic aerosols to have been UV-limited. The contemporary carbon outgassing rate (mainly as CO<sub>2</sub>) from mid-ocean ridges is  $1 \times 10^{13}$  to  $1 \times 10^{14}$  g year<sup>-1</sup>; estimates (54) of outgassing rates 3 Ga range from  $4 \times 10^{13}$  to  $6 \times 10^{14}$  g year<sup>-1</sup> (or  $\sim 10^{11}$  cm<sup>-2</sup>  $s^{-1}$ ). The outgassing rate 4 Ga must have been greater still but with an unknown  $CH_4$  $CO_2$  ratio. It seems possible, even without exogenous input (55), that the carbon outgassing source would have been able to keep pace with the organic solid production sink, provided  $CH_4/CO_2 \gtrsim 1$ . Integrated over 1 billion years, the organic production would have been of the same order as the total estimated terrestrial inventory of carbon in the mantle and crust (56).

#### NH<sub>3</sub> Lifetimes, Diffusion Flux, and Resupply

Choosing k(2000 Å) = 0.19, appropriate for kerogen, with  $\rho = 1.6 \text{ g cm}^{-3}$ , t = 1year, and f = 0.5, we find  $\alpha_{\lambda} = 1.2 \times 10^5$  cm<sup>-1</sup> and  $\tau \approx 8.8$ , yielding an attenuation of NH<sub>3</sub> photolysis by a factor exp( $-\tau \sec \theta$ ) = 3.9 × 10<sup>-6</sup>. The lifetime of [NH<sub>3</sub>] ~ 10<sup>-5</sup>, instead of 10 years, would then be 2.5 × 10<sup>6</sup> years. The rate of NH<sub>3</sub> photodestruction would be ~2 × 10<sup>10</sup> g year<sup>-1</sup> in this case or, integrated over 1 billion years, about 2 × 10<sup>19</sup> g, or 0.5% the current atmospheric N<sub>2</sub> abundance. By comparison, the contemporary biogenic value is ~10<sup>13</sup> to 10<sup>14</sup> g NH<sub>3</sub> year<sup>-1</sup> (9, 31, 57). In CH<sub>4</sub>rich reducing atmospheres, NH<sub>3</sub> is also a degradation product of HCN; steady-state production is estimated at ~10<sup>12 ± 1</sup> g NH<sub>3</sub> year<sup>-1</sup> (58).

Models with k(2000 Å) = 0.22, appropriate for N<sub>2</sub>-CH<sub>4</sub> irradiation residues (Titan tholin), with  $\rho = 1.4 \text{ g cm}^{-3}$ , t = 1 year, and f = 0.5 give  $\tau \approx 11.7$ , or a [NH<sub>3</sub>]  $\sim 10^{-5}$  lifetime of  $1.5 \times 10^8$  years. Averaging the values for kerogen and Titan tholin, the lifetime of a [NH<sub>3</sub>]  $\sim 10^{-5}$  primeval atmosphere as a function of f and sedimentation time t for  $k(2000 \text{ Å}) \approx 0.21$  and  $\rho = 1.5 \text{ g cm}^{-3}$  shows that effective shielding of NH<sub>3</sub> by organic solids fails only for low t and low f (Table 2).

The full integrations of J/J' for kerogen optical constants,  $\rho = 1.6 \text{ g cm}^{-3}$ , t = 1 year, and f = 0.5 give  $J/J' = 1.0 \times 10^5$ , or lifetimes for [NH<sub>3</sub>] ~  $10^{-5}$  only a factor of 2.5 less than those found by the 2000 Å approximation (Fig. 2). [The net solar UV flux 4 Ga for

**Table 1.** Values of the imaginary part of the complex refractive index k at 2000 Å for candidate organic solids that could serve as a UV shield.

Material	k <sub>λ</sub>	
Organic residue from irradiation of 6:1 $H_2O:C_2H_6$ ice mixture*	0.17	
Kerogen (67)	0.19	
Organic residue from irradiation of 10:1 N <sub>2</sub> :CH <sub>4</sub> gas mixture (48)	0.22	
Organic residue from irradiation of $H_2O$ -CO-CH <sub>4</sub> -NH <sub>3</sub> ice mixture (68)	0.21 to 0.29	
Organic refractory grains modeling interstellar dust (69)	~0.2	
Organic refractory grains modeling cometary dust (70)	0.28	
Elemental carbon (71)	0.39	
Gas-phase-deposited amorphous C (72)	0.47	
Hydrogenated amorphous C from C electrodes (73)	0.55	
Hydrogenated amorphous C from benzene oxidation (74)	0.50	
Arc-evaporated C (75)	0.65	
CH, heated to 500 K† (76)	0.82	
Soot in hydrocarbon flames:	~1.0	

\*Results are similar to those for  $6:1 \text{ H}_2\text{O:CH}_4$  ice irradiation (78). †At 0.276 µm. ‡Estimated from dispersion relations at 300, 1000, and 1600 K at 0.3 µm; all curves are rising toward shorter wavelengths.  $\lambda < 2300$  Å was about the same as it is today (11, 53).] For Titan tholin optical constants and  $\rho = 1.4~{\rm g}~{\rm cm}^{-3}$ , the result is  $J/J' = 9.6 \times 10^6$ , or a lifetime of  $\sim 10^8$  years. As a consistency check, we note that for many of the organic solids (Table 1),  $\tau(5500~{\rm \AA})/\tau(2000~{\rm \AA}) \sim 0.1$  or less, so that even for  $\tau(2000~{\rm \AA}) \sim 10$ , there is still enough sunlight reaching Earth's surface to drive a significant net greenhouse effect.

However, atmospheres in which NH<sub>3</sub> lifetimes are sufficiently prolonged will violate the assumption that NH<sub>3</sub> photolysis occurs below the organic aerosol shield. For example, for f = 0.5 and 1-year sedimentation times (Table 2), shielding lengthens NH<sub>3</sub> photolysis lifetimes by a factor  $\sim 10^6$ . In this case, the time scale for NH<sub>3</sub> to reach the altitude of CH<sub>4</sub> polymerization would be shorter than the photolysis lifetime, and the NH<sub>3</sub> would no longer be shielded. Then the required NH<sub>3</sub> resupply flux  $\phi$  will be given by its flux, dominated by eddy diffusion (59), to the CH<sub>4</sub> polymerization altitude. Contemporary eddy diffusion constants K vary from  $\sim 10^3$  cm<sup>2</sup> s<sup>-1</sup> just above the tropopause to  ${\sim}10^6~\text{cm}^2~\text{s}^{-1}$  at an altitude of 80 km (60). If we choose  $K = 10^5$  $cm^2 s^{-1}$  for the early atmosphere (6), and

**Table 2.** Lifetimes for atmospheres with  $[\rm NH_3] \sim 10^{-5}$ . The sedimentation times are from Oberbeck *et al.* (46). E indicates a value >4.5 billion years.

Sedi- men- tation time (years)	Lifetime (years) for given fraction of irradiation products in organic solids		
	0.1	0.5	0.9
0.5 1 3	40 200 6 × 10 <sup>4</sup>	$2 \times 10^{4}$ $2 \times 10^{7}$ E	5 × 10 <sup>6</sup> E E



**Fig. 2.** Solar UV flux with no attenuation (heavy line), compared with the flux at Earth's surface in the presence of a high-altitude organic solid formed by reactions of UV light with  $CH_4$ . The dashed line shows the UV flux if the high-altitude haze is assigned kerogen optical constants. The dotted line shows the same, but with optical constants appropriate to Titan tholin.

an atmospheric scale height H = 8.4 km, an upper limit (61) for the NH<sub>3</sub> eddy diffusion flux is  $\phi_e \approx (K/H)[NH_3]n(z) \approx$  $2 \times 10^9 \text{ cm}^{-2} \text{ s}^{-1} \text{ or } \sim 10^{13} \text{ g year}^{-1} \text{ at } z =$ 70 km; here n(z) is the contemporary atmospheric number density at altitude z. {Lower polymerization altitudes would give higher  $\phi_e$  due to larger n(z) [n(z = 40km) ~ 40n(z = 70 km)], although this effect is mitigated by lower values of K at lower n(z). In this case,  $[NH_3] \sim 10^{-5}$  has a lifetime of ~ $[NH_3]M_a/\phi$  ~ 5000 years (where  $M_a = 5 \times 10^{21}$  g is the mass of the contemporary atmosphere), so that the current inventory of N<sub>2</sub> would be generated by NH<sub>3</sub> photolysis in  $\sim 5 \times 10^8$  years. Given resupply, half-billion-year time scales for the duration of  $[\rm NH_3] \sim 10^{-5}$ atmospheres appear possible.

Would CH<sub>4</sub> have been irreversibly converted to organic solids and NH<sub>3</sub> to N<sub>2</sub> and organic solids on the early Earth, or are there mechanisms for recycling? Tholins are stable and require temperatures of 600° to 1000°C for half their mass to be vaporized (62). If plate tectonics were in operation 4 Ga, with subduction of the crust down to hundreds of kilometers depth and a geothermal gradient substantially steeper than that which exists today, tholins may have been reprocessed into  $\mathrm{NH}_3,\,\mathrm{CH}_4,\,\mathrm{and}$  other simple organic gases. Recycling of N<sub>2</sub> into NH<sub>3</sub> may occur by means of electrical discharges and NO (30) and TiO<sub>2</sub> photochemical catalysis in titanium-rich deserts (63). These processes reduce the need for outgassing or exogenous resupply. Because methanogens were among the earliest microorganisms, and life seems to have been widespread by 3.5 and possibly by 3.85 Ga (18), there is no need to seek nonbiological sources of CH<sub>4</sub> (or NH<sub>3</sub>) in that period or later. Ultraviolet optical depths above 5 to 10 on Earth would have played an important role in shielding early organisms from UV damage (64).

A point of comparison for this model is provided by Saturn's moon Titan. The UV and radiation processing of Titan's N<sub>2</sub>-CH<sub>4</sub> atmosphere produces an aerosol with average particle radii ~0.25  $\mu$ m and an optical depth of about 7 between 2000 and 3000 Å (65), about the same as the optical depths calculated here for CH<sub>4</sub> photolysis in the early terrestrial atmosphere.

There are clearly uncertainties in many of the input parameters in our calculations. Nevertheless, were the atmosphere of the early Earth reducing, our results suggest that it would have been self-shielding against UV photodissociation. Were this atmosphere instead rich in N<sub>2</sub> with minor CO<sub>2</sub> and CH<sub>4</sub> components, self-shielding may also have been possible, but only if a CH<sub>4</sub>/ CO<sub>2</sub> ratio  $\geq$ 1 could have been maintained. We wish only to suggest that a self-consistent solution to the early faint sun dilemma (66), without invoking massive  $CO_2$  atmospheres, appears to be viable.

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# *Trichodesmium*, a Globally Significant Marine Cyanobacterium

77.

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Planktonic marine cyanobacteria of the genus *Trichodesmium* occur throughout the oligotrophic tropical and subtropical oceans. Their unusual adaptations, from the molecular to the macroscopic level, contribute to their ecological success and biogeochemical importance. *Trichodesmium* fixes nitrogen gas (N<sub>2</sub>) under fully aerobic conditions while photosynthetically evolving oxygen. Its temporal pattern of N<sub>2</sub> fixation results from an endogenous daily cycle that confines N<sub>2</sub> fixation to daylight hours. *Trichodesmium* colonies provide a unique pelagic habitat that supports a complex assemblage of consortial organisms. These colonies often represent a large fraction of the plant biomass in tropical, oligotrophic waters and contribute substantially to primary production. N<sub>2</sub> fixation by *Trichodesmium* is likely a major input to the marine and global nitrogen cycle.

**T**richodesmium, a colonial marine cyanobacterium (1) (Fig. 1), has intrigued naturalists, biologists, and mariners for well over a century (2). These cyanobacteria have been reported throughout the tropical and subtropical Atlantic, Pacific, and Indian oceans, as well as the Caribbean and South China seas (Fig. 2) (3, 4). Modern interest in *Trichodesmium* dates to the early 1960s with the recognition that the biological productivity of large expanses of the ocean is often limited by the availability of nitrogen (5) and the observation that *Trichodesmium* is diazotrophic (that is, an N<sub>2</sub> fixer). The current focus in assessing the global role of the upper ocean in assimilating atmospheric  $CO_2$  has elevated the importance of quantifying marine  $N_2$  fixation.

Although major advances in understanding the biology of Trichodesmium have recently occurred on diverse fronts, several important questions remain largely unresolved: (i) Where does Trichodesmium fit in the broader scheme of cyanobacterial phylogeny? (ii) How does Trichodesmium sustain simultaneous photosynthetic  $O_2$  evolution with nitrogenase activity, and why does it fix N<sub>2</sub> only during daylight periods? (iii) What physiological, morphological, and behavioral adaptations contribute to Trichodesmium's ecological success in the oligotrophic marine environment? (iv) What environmental and ecological factors control production and N<sub>2</sub> fixation in Trichodesmium in situ, and to what extent does it contribute to productivity, nutrient cycling, and trophodynamics in tropical and subtropical seas? (v) What is the over-

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