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Photoredox cycling during UV irradiation of ferrocyanide ($[\text{Fe}^{\text{II}}(\text{CN})_6]^{4-}$) in the presence of stoichiometric sulfite (SO_3^{2-}) is shown to be an extremely effective way to drive the reductive homologation of hydrogen cyanide (HCN) to simple sugars and precursors of hydroxy acids and amino acids.

Our previous, potentially prebiotic, Kiliani-Fischer-like reductive homologation of hydrogen cyanide (HCN 1) to the simple carbohydrates glycolaldehyde 2 and glyceraldehyde 3, required the use of either HCN 1 itself, or hydrogen sulfide (H_2S) as stoichiometric reductants to effect copper(II) \leftrightarrow copper(I) photoredox cycling (Scheme 1).^{1,2} In this chemistry intended to demonstrate 'proto-metabolism',³ protons delivered by general acids facilitate direct reduction of nitrile groups by photochemically-generated hydrated electrons. The reaction network is initiated by reduction of HCN 1 to methanimine 4 and hydrolysis of the latter to formaldehyde 5. Formation of the cyanohydrin of 5, glycolonitrile 6, is followed by further reduction and hydrolysis to glycolaldehyde 2. Another cycle of reductive homologation, *via* glyceronitrile 7, gives glyceraldehyde 3. Although prebiotically plausible,⁴ these syntheses are either problematic as regards subsequent use of the sugars in RNA synthesis, or invoke distinct and rather specific geochemical scenarios. Thus, using HCN 1 as the stoichiometric reductant, isocyanic acid 8 (formed upon hydrolysis of cyanogen 9) traps 2 and 3 in the form of cyclic adducts (Scheme 1).¹ Using H_2S as the reductant presents difficulties associated with concentrating such a species in water – its low solubility means that it could most readily be concentrated as its conjugate base, hydrosulfide (HS^- , pK_a of H_2S (~ 7))⁵ in alkaline groundwater. Furthermore, the relatively low abundance of copper in Earth's crust would have restricted either chemistry to copper-rich environments, such as those enriched through impact metallogenesis.

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Photochemical reductive homologation of hydrogen cyanide using sulfite and ferrocyanide[†]

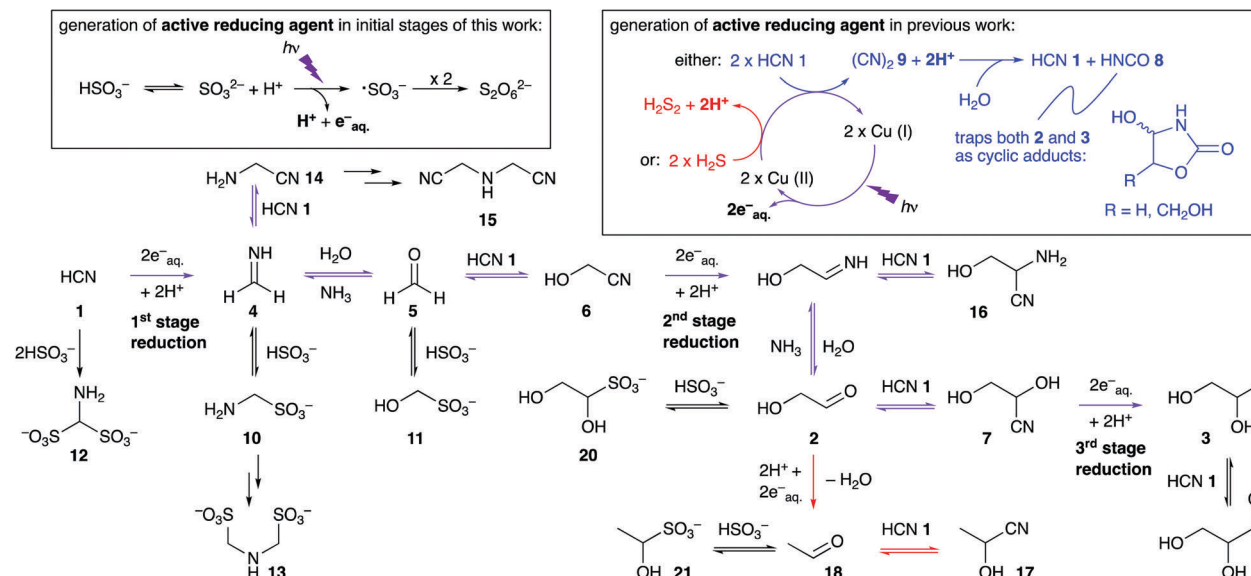
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For reductive homologation of HCN 1 to have been widespread, an alternative to either HCN 1 or H_2S as reductant would have to have been more globally available, and, if a catalyst was also required, it would ideally be based on a much more abundant metal. Here we describe a potentially widespread prebiotic synthesis of simple sugars and amino acid precursors from HCN 1 using sulfite (SO_3^{2-} , deriving from dissolution of atmospheric SO_2) as stoichiometric reductant with ferrocyanide ($[\text{Fe}^{\text{II}}(\text{CN})_6]^{4-}$) promoting the production of hydrated electrons (ESI 1.1†).

We initially explored the photoreduction chemistry of HCN 1 with bisulfite/sulfite alone using direct analysis by ^{13}C NMR spectroscopy. After 2.5 h of irradiation, the expected first-stage reduction products of HCN 1, namely methanimine 4 and its hydrolysis product formaldehyde 5, were not observed by ^{13}C NMR spectroscopy (ESI). Instead, aminomethanesulfonate 10 and hydroxymethanesulfonate 11, the bisulfite adducts of 4 and 5, respectively, were observed together with aminomethane-disulfonate 12⁶ and iminodimethanesulfonate 13 (Scheme 1). The identities of these products were confirmed by comparing their spectral properties with those of authentic compounds (ESI). After a longer irradiation time (5 h), the first-stage Kiliani-Fischer homologation products, glycolonitrile 6, glycine nitrile 14 and iminodiacetonitrile 15 were observed. Most importantly, the second-stage product, glyceronitrile 7, was also detected in the reaction mixture at this stage. Comparing ^{13}C NMR spectra at different time points revealed that the bisulfite adducts of the first-stage reduction products, aminomethanesulfonate 10 and hydroxymethanesulfonate 11, were gradually converted to the first-stage homologation products, glycolonitrile 6 and glycine nitrile 14 as the bisulfite and sulfite in the mixture were consumed.

Our initial experiments with HCN 1 and bisulfite/sulfite had simulated the delivery of SO_2 from the atmosphere into groundwater containing cyanide salts derived from the prior thermal metamorphosis of sodium or potassium ferrocyanide salts in the dry-state.⁶ Alternatively, bisulfite and formaldehyde 5, produced atmospherically by photoreduction of CO_2 ,⁷ could





Scheme 1 Reductive homologation of HCN **1**. Colour scheme: previously observed chemistry using photoredox cycling of copper(II) \leftrightarrow copper(I) with the stoichiometric reductant being HCN **1** (blue), or H₂S (red); previously observed transformations using either reductant (purple); additionally observed new transformations (black).

have rained-in to cyanide containing groundwater as hydroxymethanesulfonate **11**. We therefore explored the chemistry starting directly from **11** and observed efficient production of glycolonitrile **6** when a solution of **11** was mixed with potassium cyanide.

Starting with an initial ratio of cyanide to bisulfite/sulfite (all in the form of hydroxymethanesulfonate **11**) of 1:1, the ratio of glycolonitrile **6** to hydroxymethanesulfonate **11** in the mixture had reached 4:1 after equilibration (ESI), liberating free bisulfite/sulfite to act as a reductant in subsequent photochemistry. To determine the extent of Kilian-Fischer homologation upon irradiation, we again used ¹³C-labelled reagents with analysis by ¹³C NMR spectroscopy (Fig. 1).

The equilibration reaction between hydroxymethanesulfonate **11** and cyanide with glycolonitrile **6** and bisulfite/sulfite was mimicked by mixing 1 equivalent of ¹³C-labelled formaldehyde **5** with 2 equivalents of disodium sulfite and 2 equivalents of ¹³C-labelled potassium cyanide and adjusting the pH of the solution to **7**. Initially, only hydroxymethanesulfonate **11**, glycolonitrile **6** and excess HCN **1** were observed in the ¹³C NMR spectrum. However, after irradiation for 12 h, nearly all the hydroxymethanesulfonate **11** had been converted into glycolonitrile **6**, glycercenitrile **7**, serine nitrile **16** (convertible to serine by hydrolysis) and iminodiacetonitrile **15**. Interestingly, a small amount of acetaldehyde cyanohydrin **17** (convertible to lactate by hydrolysis) was also produced in the reaction, which could lead to alanine nitrile (convertible to alanine by hydrolysis) if sufficient ammonia was present in the system at a later stage.²

We propose that acetaldehyde **18** originates from deoxygenation of glycoaldehyde **2** (Scheme 1) as we had found using H₂S as the stoichiometric reductant in our earlier work.² To quantify the yields of reduced products, a known amount of ¹³C-labelled sodium formate was added to the solution as an

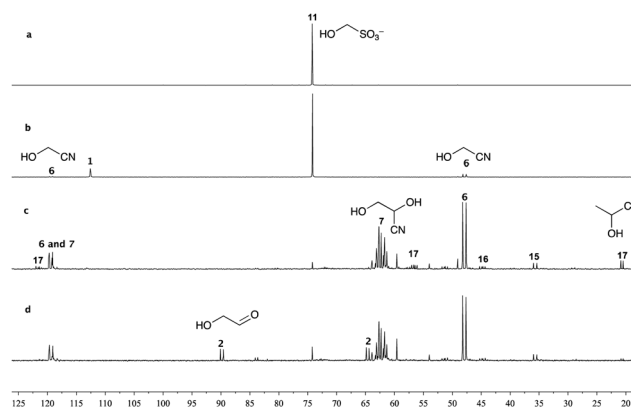


Fig. 1 ¹³C NMR Spectra of the reaction mixtures with 200 mM ¹³C-labelled KCN, 200 mM Na₂SO₃ and 100 mM ¹³C-labelled formaldehyde (in 10% D₂O in H₂O). (a) ¹³C-Labelled **11**; (b) as (a), then mixed with ¹³C-labelled KCN and NaH₂PO₄ at pH 7; (c) the mixture from (b) after irradiation at 254 nm for 12.5 h; (d) the mixture from (c) after sparging with argon for 13 h.

external standard, and the mixture was analyzed by quantitative ¹³C NMR spectroscopy (ESI). Reduced products constituted 34% of the mixture including glycercenitrile **7** (26%), serine nitrile **16** (4%) and acetaldehyde cyanohydrin **17** (4%). Theoretically, reduced products could be obtained in up to 50% yield from a 1:1 mixture of cyanide and bisulfite/sulfite, as the reduction of one nitrile group requires two electrons released from two equivalents of sulfite. After sparging argon through the reaction mixture for 13 h to expel HCN **1** from the solution, free glycolaldehyde **2** could be observed in the ¹³C NMR spectrum (Fig. 1d).

In our previous synthesis using H₂S as the reductant, copper(I) cyanide was found to accelerate the photoreduction



of glycolonitrile **6**, providing **2** in 42% yield after 4 h of irradiation.² In comparison, the new photoreduction with sulfite alone as the reductant, gave reduced products in a lower yield with longer irradiation times (12 h), and this raised concerns about its prebiotic plausibility. We therefore looked for an Earth-abundant compound to accelerate the sulfite reduction chemistry.

It is known that photoionization of ferrocyanide ($[\text{Fe}^{\text{II}}(\text{CN})_6]^{4-}$, ESI 1.2†), effected by UV irradiation at short wavelengths, provides ferricyanide ($[\text{Fe}^{\text{III}}(\text{CN})_6]^{3-}$) and hydrated electrons.⁸ Indeed we had previously attempted using ferrocyanide for reductive homologation chemistry, but it proved inefficient on its own, which we put down to efficient geminate recombination of the electrons and ferricyanide regenerating ferrocyanide. However, in the context of using sulfite as the stoichiometric reductant, ferrocyanide piqued our interest again because it is known that sulfite reduces ferrocyanide to ferrocyanide and, in the process, is converted to sulfate.^{9–11} Thus, depending on the relative rates of several processes, added ferrocyanide might double the reducing capacity of sulfite and accelerate the photochemically-driven reductive homologation of HCN **1**. To investigate whether ferrocyanide might act in this way, a solution of 1 equivalent of ¹³C-labelled KCN and 1 equivalent of Na_2SO_3 in phosphate buffer was divided in two and 10 mol% $\text{K}_4[\text{Fe}(\text{CN})_6]$ was then added to one portion. The two solutions were then irradiated side-by-side for 3 h (Fig. 2). The reaction mixture lacking ferrocyanide gave only the first-stage reduction products **6**, **10**, **11** and a trace of **13**, while the reaction mixture including ferrocyanide furnished mainly the second-stage reduction product glyceronitrile **7** together with a third-stage reduction product **19**, the cyanohydrin of glyceraldehyde **3**. A similar comparison (ESI) was also made of the reactions starting from mixtures of ¹³C-labelled hydroxymethanesulfonate **11** and ¹³C-labelled HCN **1** with and without added ferrocyanide. In the reaction mixture including ferrocyanide, most of the HCN **1** and the hydroxymethanesulfonate **11** had been consumed within 3 h, providing the reduced product glyceronitrile **7** as well as free glycolaldehyde **2** and the cyanohydrin of glyceraldehyde **19**. By comparison, the reaction mixture lacking ferrocyanide showed

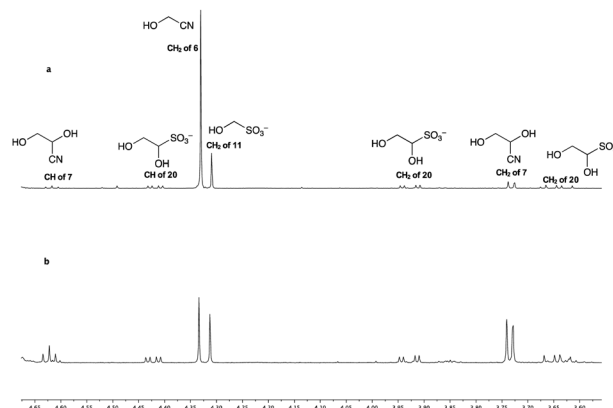


Fig. 3 ¹H NMR Spectra of the reaction mixtures with 25 mM of **11**, 25 mM KCN and 100 mM NaH_2PO_4 in $\text{D}_2\text{O}/\text{H}_2\text{O}$ (10% D_2O in H_2O) after irradiation for 1 h. (a) The reaction with no $\text{K}_4[\text{Fe}(\text{CN})_6]$; (b) the reaction with 10 mol% $\text{K}_4[\text{Fe}(\text{CN})_6]$.

considerably less efficient reduction in the same period of time (ESI). In order to quantify the effect of ferrocyanide on the photoreduction, the reaction of hydroxymethanesulfonate **11** with KCN was repeated with unlabelled **11**. Hydroxymethanesulfonate **11** was mixed with 1 equivalent of KCN in phosphate buffer and the resulting mixture was again divided into two parts, into one of which was added 10 mol% $\text{K}_4[\text{Fe}(\text{CN})_6]$. Reactions were monitored periodically by ¹H NMR spectroscopy and yields of products were calculated by relative integration of their proton resonance signals (Fig. 3).

Comparing the reactions after only 1 h of irradiation, the catalyzed, or promoted reaction was found to have proceeded rapidly, affording 68% yield of total reduced products (glyceronitrile **7** in 40% yield and glycolaldehyde sulfite adduct **20** in 28% yield), while the control reaction gave only 20% yield of reduced products, eventually increasing to 25% after 3 h.

In the control reaction starting from HCN **1** and sulfite, third-stage reduction products such as glyceraldehyde **3** and its cyanohydrin **19** could barely be detected in the photoreduction mixtures. To investigate the effect of ferrocyanide on the later stages of the overall synthetic scheme, we simply mixed glycolaldehyde **2** with 1 equivalent of KCN and 1 equivalent of Na_2SO_3 in phosphate buffer. In the dark, the ratio of the sulfite adduct **20** to cyanohydrin **7** was 2.4:1. As before, the mixture was divided into two parts, into one of which was added 10 mol% $\text{K}_4[\text{Fe}(\text{CN})_6]$. After 1 h of irradiation, the photoreduction reaction including ferrocyanide afforded 30% of **19**, the cyanohydrin of glyceraldehyde and the ratio of **20** to **7** had changed to 0.6:1 (ESI). The sulfite in the mixture was efficiently consumed (as reductant) and the equilibrium was in favor of the formation of cyanohydribs **7** and **19**. In comparison, the control reaction afforded no detectable **19** after 1 h of irradiation, but afforded 3% of deoxygenated products (acetaldehyde sulfite adduct **21** and acetaldehyde cyanohydrin **17**) deriving from glycolaldehyde **2**. After 3 h of irradiation, 10% of **19** and 8% of acetaldehyde derivatives were observed. Based on our experimental findings and results from the literature, reaction

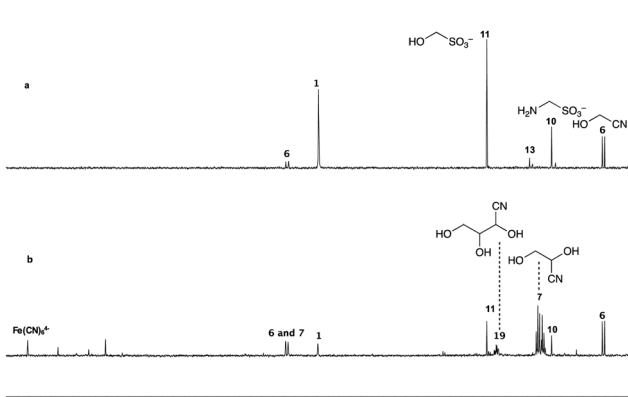
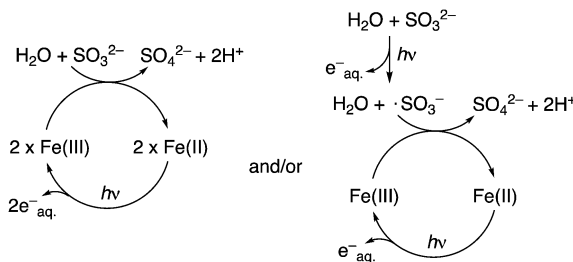


Fig. 2 ¹³C NMR Spectra of the reaction mixtures with 25 mM ¹³C-labelled KCN, 25 mM Na_2SO_3 and 100 mM NaH_2PO_4 in $\text{D}_2\text{O}/\text{H}_2\text{O}$ (10% D_2O in H_2O) after irradiation for 3 h. (a) The reaction with no $\text{K}_4[\text{Fe}(\text{CN})_6]$; (b) the reaction with 10 mol% $\text{K}_4[\text{Fe}(\text{CN})_6]$.





Scheme 2 Proposed mechanisms for the photoredox cycling of iron(III) ⇌ iron(II) in the presence of Na_2SO_3 .

mechanisms involving cyanoferrate photoredox cycling are proposed here (Scheme 2 and ESI 1.3†).

In conclusion, through sulfite (SO_3^{2-}) and catalyzed, or promoted by ferrocyanide ($[\text{Fe}^{\text{II}}(\text{CN})_6]^{4-}$), SO_2 can act as a more efficient and globally available reductant than H_2S in the photochemically-driven homologation of HCN 1 to (precursors of) biomolecules. Considering the ready availability of ferrous iron (Fe^{II}) on early Earth, the ease with which atmospheric SO_2 may be concentrated into groundwater, and the numerous mechanisms for supply of HCN, the sulfite-mediated, ferrocyanide-accelerated photoreduction of cyanide offers a synthesis of sugars and precursors of hydroxy acids and amino acids compatible with a globally plausible geochemical scenario.

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Conflicts of interest

There are no conflicts to declare.

Notes and references

- 1 D. Ritson and J. D. Sutherland, *Nat. Chem.*, 2012, **4**, 895–899.
- 2 D. J. Ritson and J. D. Sutherland, *Angew. Chem., Int. Ed.*, 2013, **52**, 5845–5847.
- 3 K. Ruiz-Mirazo, C. Briones and A. de la Escosura, *Chem. Rev.*, 2014, **114**, 285–366.
- 4 Z. R. Todd, A. C. Fahrenbach, C. J. Magnani, S. Ranjan, A. Björkboom, J. W. Szostak and D. D. Sasselov, *Chem. Commun.*, 2018, **54**, 1121–1124.
- 5 G. M. Marion, J. S. Kargel, J. K. Crowley and D. C. Catling, *Icarus*, 2013, **225**, 342–351.
- 6 R. A. B. Bannard and J. H. Ross, *Can. J. Chem.*, 1954, **32**, 49–50.
- 7 J. P. Pinto, G. R. Gladstone and Y. L. Yung, *Science*, 1980, **210**, 183–185.
- 8 M. Shirom and G. Stein, *J. Chem. Phys.*, 1971, **55**, 3372–3378.
- 9 R. S. Murray, *Chem. Commun.*, 1968, 824–825.
- 10 J. M. Lancaster and R. S. Murray, *J. Chem. Soc. A*, 1971, 2755–2758.
- 11 A. D. James and R. S. Murray, *Inorg. Nucl. Chem. Lett.*, 1976, **12**, 739–742.

