A Radical Pathway for Organic Phosphorylation During Schreibersite Corrosion

with Implications for the Origin of Life

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ABSTRACT

Phosphorylated compounds (e.g. DNA, RNA, phospholipids, and many coenzymes) are critical to biochemistry. Thus, their origin is of prime interest to origin of life studies. The corrosion of the meteoritic mineral schreibersite ((Fe,Ni)₃P) may have significantly contributed to the origin of phosphorylated biomolecules. Corrosion of synthetic schreibersite in a variety of solutions was analyzed by nuclear magnetic resonance spectroscopy, mass spectrometry, and electron paramagnetic resonance spectroscopy. These methods suggest a radical reaction pathway for the corrosion of schreibersite to form phosphite radicals (PO₃⁻) in aqueous solution. These radicals can form activated polyphosphates and can phosphorylate organic compounds such as acetate (3% yield). Phosphonates (O₃P-C) are found in the organic P inventory of the carbonaceous meteorite Murchison. While phosphonates are rare in biochemistry, the ubiquity of corroding iron meteorites on the early Earth could have provided an accessible source of organophosphorous for the origin of life allowing the invention of the organophosphates in modern biology as a product of early evolution.
1. INTRODUCTION

Determining the origin of phosphorylated biomolecules is a central goal of origins of life research. Phosphorylated biomolecules are critical components of life important both structurally (e.g. DNA and phospholipids) and energetically (e.g. ATP, UDPG). Contrary to the active role of P in life, P in geochemistry is comparatively inert. Phosphate minerals are sparingly soluble in water and lack reactivity towards most organic molecules. The formation of phosphorylated biomolecules is inhibited by the geochemistry of P on the Earth and many scenarios for the origin of phosphorylated biomolecules have been forced to employ unlikely geochemical settings (e.g., Keefe and Miller, 1995; Peyser and Ferris, 2001; Pasek and Lauretta, 2005). Following the discovery of phosphonic acids in the Murchison carbonaceous chondrite, some researchers have proposed that extraterrestrial material may have provided the initial reactive and soluble phosphorus for life (e.g., de Graaf et al., 1995; Macia et al., 1997).

The meteoritic mineral schreibersite, (Fe,Ni)$_3$P, is a proposed source of reactive P for the origin of phosphorylated biomolecules (Gulick, 1955; Pasek and Lauretta, 2005; Bryant and Kee, 2006). Schreibersite readily corrodes in aqueous solution to form a mixed-valence series of P compounds, including orthophosphate (I, oxidation state +5), pyrophosphate (II, oxidation state +5), phosphite (III, oxidation state +3), hypophosphate (IV, oxidation state +4), and, under some conditions, diphosphite (V, oxidation state +3) (Figure 1). In the presence of acetate/ethanol several organophosphorus compounds are also formed (Pasek and Lauretta, 2005).

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1 Using the nomenclature of Van Wazer (1958) for the salts
A better understanding of the reaction is critical to an evaluation of the prebiotic input of schreibersite to the origin of life, and to the origin of the P-C molecules in the organic inventory of meteorites. Due to the complexity of the corroding schreibersite system, it is difficult to deduce a detailed mechanism. However, a reaction pathway which is consistent with the nuclear magnetic resonance (NMR), mass spectrometry (MS), electron paramagnetic resonance (EPR), and analogy with other reactions can be determined. We propose that the reaction pathway consists of a series of radical reactions culminating in a series of P salts, and that radical chemistry overcomes energy barriers traditionally associated with the phosphorylation of organics.

Two groups of experiments were performed to elucidate a plausible reaction pathway by investigating the process of P speciation from phosphide corrosion. One set of experiments examined the effect of aqueous corrosion of iron phosphide, Fe$_3$P, on a suite of simple compounds (experiments 1-22, Table 1). A second set of experiments (23-32) investigated the stability and reactivity of phosphite, orthophosphate, hypophosphate, and pyrophosphate in the presence or absence of corroding metal; compounds produced during schreibersite corrosion. The stability of hypophosphite (oxidation state +1, Figure 1, VI) was also investigated as this compound is a plausible reaction intermediate.
2. METHODS AND MATERIALS

2.1 Materials and Conditions

Phosphide was introduced as pure Fe₃P powder (-40 mesh, purchased from CERAC, Inc.). High-purity iron powder (99+%, Alfa Aesar) was used in some of the corrosion experiments. Magnetite was from Serro, Minas Gerais, Brazil. The solution volume for all experiments was 25 mL. Aqueous solutions were either pure deionized water, purified in house using a Barnstead NANOpure® Diamond Analytical combined reverse osmosis-deionization system, or were solutions created by adding compounds to the deionized water. Solutions were from salts of Mg- and Ca-chloride (98%, from Alfa Aesar), reagent alcohol (90.5% ethanol, 5% isopropanol, and 4.5% methanol from VWR lab supplies), acetic acid (99.7% from EMD lab supplies), NaHCO₃ (Arm and Hammer, 99+%), glycolaldehyde (MP Biomedicals LLC., 99+%), acetaldehyde (Merck, 99+%), ethylenediaminetetraacetic acid (EDTA was from VWR lab supplies, H₂O₂ (30% in water, VWR lab supplies), pyruvate (Alfa Aesar, 98%), acetonitrile (J.T. Baker, 99.8%), 5-5-dimethyl-1-pyrroline-N-oxide (DMPO from Alexis Biochemicals, 100%, stored at -10°C) or a combination of these components. The pH was between 6.5 – 8 in most experiments with the reported pH determined before and after reaction. The pH of the buffer was adjusted using NaOH (from VWR, 10.0 N stock solution) or NH₄OH (from ADChemco Scientific Inc., 28-30% in water). Half of each solution was isolated from the phosphide and saved for comparison as a blank.
Pure compounds were used as standards for species identification and were prepared at 0.01 M in D$_2$O and NaOH (pH 13). Standards used include H$_3$PO$_3$, H$_3$PO$_4$, Na$_4$P$_2$O$_7$, NaH$_2$PO$_2$, C$_2$H$_5$PO$_3$H$_2$, CH$_3$PO$_3$H$_2$, (CH$_3$)$_3$PO$_4$, Ba(PO$_3$)$_2$ and P$_2$O$_5$ and were acquired from VWR International, and acetylphosphonic acid (H$_2$O$_3$PCH$_2$COOH, 98+%) was acquired from Alfa Aesar. Hypophosphate was synthesized by the method of Yoza and Ohashi (1965) and verified by $^{31}$P NMR. The temperature in all experiments was held at $\sim$20±3°C. The Ar (99.99%) was from University of Arizona Laboratory Stores.

2.2 Experimental Setup

Experiments were mostly performed under ambient atmospheric conditions (NMR Air) with some under Ar to remove all dissolved O$_2$ from the water as well as to maintain an inert atmosphere prior to introduction of the phosphide to the water. The corrosion of phosphide under Ar (Pasek and Lauretta, 2005) and N$_2$, (Bryant and Kee, 2006) generates the same products but at lower abundance than under air.

Solutions of 0.01-0.5 M P compound were used for the stability experiments. These solutions were sealed with or without ~0.5-1 g of Fe powder or ~0.5 g of crushed magnetite. Samples were analyzed after one day of constant stirring.

NMR analysis was at pH 13 as the addition of NaOH sharpens NMR peaks and allow for comparison with previous studies. The NaOH precipitated Mg and Ca containing solutions. The carbonate solutions were analyzed at a pH 9.5 since too much volume of NaOH is required to overpower the buffer. The sample was filtered, the
extract concentrated by vacuum and then redissolved in 5 mL of D2O (99.7%, Alfa Aesar).

One experiment replicated the phosphide corrosion in acetic acid under air with NH4OH instead of NaOH. This allowed analysis by ES' ToF-MS (negative Electrospray Time of Flight Mass Spectrometry) to avoid sodium acetate clusters. Prior to analysis, the sample was dried vacuum to remove the majority of NH3 in solution. The remaining material had a slight red coloration.

Corrosion experiments analyzed by Electron Paramagnetic Resonance (EPR) spectrometry were prepared by removing a small quantity of solution and adding the spin trap DMPO (0.2 M final concentration). No NaOH was added to the phosphide sample.

2.3 Analytical Methods

Each solution was analyzed using 31P NMR on a Varian 300 four-nucleus probe FT-NMR (Fourier Transform-NMR) spectrometer at 121.43 MHz and 24.5 °C for 256 to 35000 scans. Spectra were acquired in both H-decoupled and coupled modes.

The bulk dissolved P content was measured using a ThermoFinnigan ELEMENT2 high-resolution Inductively Coupled Plasma Mass Spectrometer (ICP-MS). Analytical details are in Pasek and Lauretta (2005). The ICP-MS data (R^2 = 0.9939) was used to calibrate the NMR to determine P molarity [M] in solution by the Equation 1:

\[
[M] = 0.0075 \times \left( \frac{S}{N} \right)^2 + 0.0007 \times \left( \frac{S}{\sqrt{Scans}} \right) + 0.0001
\]
where $S$ is the signal, $N$ is the noise, and \textit{Scans} is the number of NMR scans taken. This relationship was empirically determined and is accurate to $\pm 10\%$ over the range of $10^{-4}$ to $10^{-2}$ M based on the sample spectra acquired.

The abundance of each P compound was determined by integration of the NMR spectrum. NMR integration is quantitative if the integration is done over a narrow range of frequencies, typically less than 50 ppm. Turning the H-decoupler off during NMR acquisition had no effect on integrations compared with acquisitions with the H-decoupler on, indicating that the Nuclear Overhauser Effect (NOE) did not affect integrations significantly.

Mass spectrometry was performed by infusing a solution into a Waters LCT Premier Time of Flight Mass Spectrometer (TOF-MS) at 100 $\mu$L/min. Soft ionization was performed with ES$^-$ with 4200 V capillary voltage, 300°C desolvation gas temperature, and 10 V cone voltage. The MS was in “W” reflectron mode with a mass resolution of about 10,000 and scanned over 50 to 400 m/z continuously calibrated adenine (-H$^+$) (M/Z = 134.0467 Da) every 10 scans via a dedicated sprayer. Minimal ionization was observed in ES$^+$ mode. A spectrum was acquired every second for 3 minutes after which time the spectra were summed and averaged. The relative abundances of the four species $\text{H}_2\text{PO}_3^-$, $\text{H}_2\text{PO}_4^-$, $\text{H}_3\text{P}_2\text{O}_6^-$, and $\text{H}_3\text{P}_2\text{O}_7^-$ were determined by NMR and used to estimate the MS abundances, differences in ionization efficiency in different compounds make this only an approximation.

EPR analyses were performed using a continuous wave X-band (9-10 GHz) EPR spectrometer, model ESP-300E manufactured by Bruker. Samples were siphoned into
capillary tubes and analyzed with a 1 Gauss modulation and a 20 mW microwave power source. Each spectrum consists of 30 scans from 3290 to 3390 Gauss.

3. RESULTS

In all experiments exposed to air the phosphide powder changed color from gray to a reddish brown. In the experiments isolated from air and flushed under argon, color change was limited to black. Some organic corrosion experiments turned to an iron rust-rich tar and could not be analyzed by NMR. The pH of all systems remained relatively unchanged (pH changes <0.5). Besides external appearance, a number of P species were identified in aqueous solution. The abundances of all P species and total abundance of P in experiments 1-19 is given in Table 2. The inorganic ionic species were identified based on comparison to standards and literature values (Yoza et al. 1994). One organic P compound, acetyl phosphonate, was positively identified by comparison to a prepared standard. Others were identified through literature comparisons.

[Table 2]

The total yields of dissolved P compounds relative to the amount of phosphide material employed in corrosion range from 0.1% to 10%. The two major factors affecting yield are solution chemistry and atmosphere composition where solution chemistry appears to have a larger effect.
3.1 NMR Experiment Results

Table 3 summarizes the coupling constants, peak locations, and other pertinent NMR data from the experiments detailed below, including detection of organophosphorus compounds which were identified using data from the ES-ToF-MS.

| Table 3 |

For experiments 1-19, analyses of blank solutions by NMR revealed no P contamination. All P observed by NMR in solution originated from the corroded phosphide. Nearly all experiments had four major $^{31}$P-NMR peaks (hypophosphate, orthophosphate, phosphite, pyrophosphate, Table 2). The most diverse P chemistry occurred in organic solution corrosion.

The experiments with acetone and with glycolaldehyde both formed a rust-colored tar which could not be analyzed by conventional NMR. Experiments with ethanol or acetonitrile aqueous solutions produced no new products. Acetate and to a lower molar yield other carbonyls (pyruvate, acetaldehyde and EDTA) generated a diverse suite of soluble products without polymerizing into tar. Experiment 4 corroded Fe$_3$P in a solution of Mg and Ca salts with acetate and ethanol. This experiment produced mainly phosphite in solution. However, a small quantity of a likely organophosphate was detected (~50 ppm) but was absent in a subsequent analysis after standing ~2 months at room temperature. Repetition of this experiment did not generate the peak again. This suggests that this was artifact of the NMR analysis or a relatively unstable or insoluble organophosphate.
The iron phosphide powder was also corroded in a H$_2$O$_2$ solution (experiment 14). The products detected by NMR was surprisingly similar those generated from acetate (Figure 2), the major exception is the presence of minor organophosphorus species in the acetate experiment.

[Figure 2]

3.2 Mass Spectral Experiments

Mass spectral analysis of the CH$_3$COONH$_4$ and Fe$_3$P experiment (#20) indicates an enormous variety of P compounds were produced during this corrosion experiment (Figure 3). Plausible identifications of these compounds are summarized in Table 4.

[Figure 3]

[Table 4]

The mass resolution (< 3ppm) allows the determination of molecular formulas for the detected species. Several mass peaks correspond well with compounds known from the NMR analysis, including phosphite (H$_2$PO$_3$), orthophosphate (H$_2$PO$_4$), hypophosphate (H$_3$P$_2$O$_6$), pyrophosphate (H$_3$P$_2$O$_7$), and diphosphite (H$_3$P$_2$O$_5$).

3.3 Electron Paramagnetic Resonance Spectrometry Analyses

Samples of Fe and Fe$_3$P were corroded in deionized water to determine the nature and abundance of radicals in solution. Figure 4 shows the EPR spectra from DMPO and
water (background), DMPO and Fe in basic solution, and DMPO and Fe$_3$P in neutral water. No peaks are observed in the background spectrum, indicating that the diradical O$_2$ from air is not trapped as a radical species to any significant observable extent. The peaks shown in Figures 4b and 4c confirm the presence of radical species in solution.

The EPR spectrum of DMPO and Fe (Figure 4b) in basic solution corresponds well to literature spectra of the radical DMPOOH (Sankarapandi and Zweier, 1999) which is formed by reaction of DMPO with 'OH radicals. The EPR spectrum of DMPO and Fe$_3$P (Figure 4c) corresponds to a mixture of DMPOOH and DMPOX (oxidized DMPO) radicals in a 5:3 ratio. DMPOX radicals are formed from the interaction of 'PO$_3^{2-}$ radicals with DMPO (Ozawa and Hanaki, 1989).

[Figure 4]

3.4 Organic Phosphorus Compounds Formed

Plausible structures of identified compounds are shown in Figure 5.

[Figure 5]

The masses and total abundances of each organic P compound are listed in Table 4. Chemical shifts of methyl diphosphite, acetyl phosphonate, and hydroxymethyl phosphonate are consistent with published spectra (Satre et al., 1989; Robitaille et al., 1991; Noguchi et al., 2004). Phosphoglycolate may also be present, and this peak may be visible in the acetate/ethanol and Mg$^{2+}$ and Ca$^{2+}$ solution. Organophosphates are more soluble in solution than phosphate minerals (e.g., Boerio-Goates et al. 2001), and may have been present in this solution. Its peak location may be overcome by the much
stronger phosphite peak in Figure 2b. Phosphonoformate has an NMR peak of between -1 to 1 ppm, depending on pH and may be observed in Figure 2b in this area.

Pasek and Lauretta (2005) discuss the possible presence of a cyclic organic P compound. The present data set rules out the NMR data used to argue for the presence of this compound as the Fe₃P and H₂O₂ experiment produced peaks identical to those of the purported organic P compound, despite the lack of any added organics. Cyclic organic P compounds may have been detected by MS, but several of these putative identifications are isomeric with non-cyclic P compounds, hence they have been left off Table 4.

3.5 P Compound Stability Experiments

The motivation for this set of experiments was to understand the stability and reactivity of P species in the presence of metal. Results are presented in Table 5. Hypophosphite (H₂P₂O₇) is a plausible intermediate in oxidation state between Fe₃P and HPO₃²⁻. It does not appear in any of the above corrosion experiments, but it may serve as a short-lived reactive intermediate. We determined the stability of this compound through monitoring this compound over the course of one day. After one day, ~25% of hypophosphite is converted to phosphite under air, independent of material added to the solution (no addition; Fe added; Fe₂O₄ added).

[Table 5]

Phosphite is more stable than hypophosphite over longer periods of time, with no conversion to other P compounds under ambient conditions. Addition of Fe metal to the solution results in a conversion of ~46% of phosphite to orthophosphate in one day.
Addition of magnetite has no effect and the phosphite concentration was unchanged. Experiments with orthophosphate, hypophosphate, and pyrophosphate added to Fe metal did not change significantly with time. No additional P compounds were produced, indicating the kinetic stability of these three compounds is higher than phosphite. These stabilities are consistent with those observed by other investigators (Van Wazer, 1958; Schwartz and Van der Veen, 1973).

4. DISCUSSION

We now summarize the salient points of our experimental results with respect to possible reaction pathways. Any proposed reaction pathway should explain these data: 1) The generation of phosphite, orthophosphate, hypophosphate, and pyrophosphate in nearly all phosphide corrosion experiments, 2) the reactivity towards acetate, and the similarity between the $\text{H}_2\text{O}_2$ and the acetate phosphide corrosion experiments, 3) the complexity of the mass spectra, 4) the presence of radicals in solution, and 5) the relative stability of the P compounds generated.

The solution chemistry is dominated by phosphite, orthophosphate, hypophosphate, and pyrophosphate. All were detected by NMR and MS of solutions in nearly all phosphide corrosion experiments. All experiments have more phosphite and orthophosphate compared to hypophosphate and pyrophosphate, though exact ratios depend on solution chemistry. MS data reveals complex solution chemistry for high mass species, and detected peroxyphosphates in solution. EPR data suggests that at least two radicals are formed in solution during the corrosion of the phosphide. The peak
patterns of these spectra are consistent with the reaction of 'OH with DMPO to form DMPOOH, and with the reaction of a strong oxidant with DMPO to form DMPOX. The strong oxidant is most likely 'PO\textsubscript{3}\textsuperscript{2-}, as DMPOX is not formed during the corrosion of Fe metal. The DMPOOH radical is also formed during the iron metal powder corrosion.

The most complex spectra result from phosphide corrosion experiments in solutions of H\textsubscript{2}O\textsubscript{2} or of sodium acetate. The acetate corrosion solution produced several organic P compounds including acetyl-P compounds and methyl-P compounds, most of which have P-C bonds. A few species have P-O-C bonds similar to those found in modern biochemistry, highlighting the relevance of this study to the origins of life field.

Phosphite, orthophosphate, hypophosphate, and pyrophosphate are all stable in solution in the absence of metal. Phosphite is oxidized to orthophosphate during the corrosion of iron metal under air. No other P compounds were formed during this corrosion, but 'OH radicals were detected in iron corrosion experiments. Hypophosphate and pyrophosphate are both thermodynamically unstable relative to orthophosphate, yet neither compound decays to orthophosphate over the timescales of these experiments.

4.1 Other Work

The synthesis of hypophosphate by Schwartz and Van der Veen (1973), the synthesis organic P compounds by de Graaf et al. (1995), and the isotopic work of Bryant and Kee (2006) are relevant to this work. Schwartz and Van der Veen (1973) describe a reaction producing hypophosphate from phosphite in water and in the presence of UV light:

\[
2\text{HPO}_3^{2-} (aq) \rightarrow \text{P}_2\text{O}_6^{4+} (aq) + \text{H}_2 (g)
\]
UV light breaks the P-H bond and allows the dimerization of phosphite to form hypophosphate via a phosphite radical.

Following the discovery of phosphonic acids in the Murchison meteorite by Cooper et al. (1992), de Graaf et al. (1995, 1997), and Gorrell et al. (2006) proposed that phosphonic acids may form through a series of radical reactions. The de Graaf et al. (1995) reaction pathway synthesized phosphonic acids through reaction of phosphite with simple organics like formaldehyde under UV light. The Gorrell et al. (2006) model hydrolyzed the compound CP to synthesize phosphonic acids, also through a radical reaction pathway. Both proposed origins of meteoritic phosphonic acids rely on radicals to produce phosphonic acids.

Bryant and Kee (2006) corroded schreibersite in the presence of isotopically-labeled water, H$_2^{18}$O under air. The resulting NMR splitting patterns suggested that three of the oxygen atoms on phosphite and orthophosphate originated from water, but that one of the oxygen atoms on orthophosphate may have originated from the air. This suggests a common origin for both phosphite and orthophosphate, with modification occurring as a secondary step. This data agrees with the oxidation of phosphite under air in the presence of iron metal. This transition from H-PO$_3$ to HO-PO$_3$ is likely due to a process other than the formation of the initial PO$_3$ group.

5. PROPOSED REACTION PATHWAY

We propose that the reaction pathway involves the generation, propagation, and termination of phosphate, phosphite, hydrogen, hydroxyl, and acetate radicals when
acetate is added to solution. This reaction pathway provides a pathway to the P salts and to phosphorylated organic compounds.

The EPR data proves the existence of radicals in solution, however, the existence of radicals alone does not imply a radical reaction pathway for the corrosion of schreibersite. The evidence from EPR, NMR, MS, and prior work strongly suggest a radical corrosion pathway when considered together. This is in contrast with an oxidation/reduction model.

Pasek and Lauretta (2005) described a reaction pathway using oxidation/reduction reactions to produce the diversity of P compounds present in solution:

\[
\begin{align*}
\text{Fe}_3\text{P} + 7 \text{H}_2\text{O} & \rightarrow \text{Fe}_3\text{O}_4 + \text{H}_3\text{PO}_3 + 5 \text{H}_2(g) \quad (3) \\
\text{H}_3\text{PO}_3 + \text{H}_2\text{O} & \rightarrow \text{H}_3\text{PO}_4 + \text{H}_2(g) \quad (4) \\
2 \text{H}_3\text{PO}_3 & \rightarrow \text{H}_4\text{P}_2\text{O}_6 + \text{H}_2(g) \quad (5) \\
\text{H}_4\text{P}_2\text{O}_6 + \text{H}_2\text{O} & \rightarrow \text{H}_4\text{P}_2\text{O}_7 + \text{H}_2(g) \quad (6) \\
\text{H}_4\text{P}_2\text{O}_7 + \text{H}_2\text{O} & \rightarrow 2 \text{H}_3\text{PO}_4 \quad (7)
\end{align*}
\]

This proposed reaction pathway was investigated in the series of experiments detailed in the P compound stability experiment section, but did not produce the products predicted. Furthermore, hypophosphate (P2O6\(^4^−\)) and pyrophosphate (P2O7\(^4^−\)) are stable and unreactive for the timescales and reaction conditions studied in the experiments described above. Phosphite and iron metal react to form orthophosphate and Fe\(^{2+}\), but this reaction alone is not an oxidation/reduction reaction since no species is reduced. Additionally, the relative abundances of the various P species are roughly constant even when removed from contact with the powder, which would not be expected if one species turned to another through reaction with water or other P salts. Finally, there is an energetic barrier...
to the formation of hypophosphate ($\Delta G^\circ = +58 \text{ kJ, pH 7, reaction 5}$). The active agent leading to hypophosphate must be more energetic than phosphite.

5.1 Generation of Radicals

Orthophosphate is the stable form of P in water. Phosphorus in schreibersite exists as oxidation state 0. Thus, the driving force during corrosion of schreibersite is oxidation of P from 0 to +5. We propose that the reaction between water and Fe$_3$P produces phosphite radicals, and several equivalents of H$_2$ gas:

$$3\text{Fe}_3\text{P} + 12\text{H}_2\text{O}^+ + 2\text{H}_2\text{O} \rightarrow 5\text{H}^+ + 5\text{OH}^- + 3\text{PO}_3^{2-} + 15\text{H}_2 + 9\text{Fe}^{2+} \quad (8)$$

Both P and Fe are oxidized in this net reaction. The oxidation state of P changes from 0 to nominally +4 and Fe from 0 to +2. Three radical species are formed: H', OH, and 'PO$_3$$^{2-}$. The production of significant quantities of H$_2$ gas occurs as the Fe$_3$P corrodes and is verified by the stoichiometric release of H$_2$ (see Pasek and Lauretta, 2005). The ratio of phosphite to hydroxyl radicals is derived from the EPR spectrum, and depends on the relative efficiencies of the chemical reactions that form DMPOOH and DMPOX. Hydrogen radicals are hypothesized based on the production of H$_2$ and phosphite.

Most phosphide corrosion solutions without Mg$^{2+}$ and Ca$^{2+}$ have at least four P species: orthophosphate, pyrophosphate, hypophosphate, and phosphite. The production of three of these P species is readily explained by radical termination reactions, like the production of phosphite through reaction of a phosphite radical with a hydrogen radical:

$$'\text{PO}_3^{2-} + \text{H}^+ \rightarrow \text{HPO}_3^{2-} \quad (9)$$
All termination reactions from the inorganic radical mixture are summarized in a radical matrix (Table 6). The H₂ evolved during corrosion is partly due to a radical termination of 2H atoms. The production of orthophosphate, hypophosphate, and phosphite is predicted from the initial radical termination products of schreibersite corrosion.

[Table 6]

One of the consequences of this proposed reaction scheme is that orthophosphate and ergo hydroxyl radicals are produced even in anoxic conditions. This is verified by the presence of orthophosphate in all solutions, even those performed under argon. Morton et al. (2003) and Bryant and Kee (2006) reported similar results for corrosion experiments performed in under N₂ atmospheres.

Hydrogen peroxide is a predicted product from radical termination of two hydroxyl radicals. Peroxide forms hydroxyl radicals through reaction with ferrous iron:

$$\text{Fe}^{2+} + \text{HOOH} \rightarrow \text{Fe(OH)}^{2+} + \cdot \text{OH}$$  \hspace{1cm} (10)

This is the Fenton reaction (Walling 1976). In most of the inorganic corrosion experiments, this reaction removes all H₂O₂, generating ferric iron and hydroxyl radicals. The net result is that hydrogen peroxide is removed from the system and thus prevented from affecting P chemistry. However, in solutions with significant chelating molecules like acetate, the Fe²⁺ forms organic complexes allowing H₂O₂ to build up and react with the powder and dissolved P species. Thus the addition of excess H₂O₂ in water to Fe₃P produced species similar to the addition of CH₃COONa (Figure 2). Other solutions with chelating agents (like EDTA, pyruvate, and bicarbonate) also have diverse P speciation products.
Pyrophosphate is a ubiquitous product of all corrosion experiments, but its abundance is the most variable of the four main P species. Pyrophosphate is not predicted as a product resulting from a radical termination step, given the radicals described in reaction 8. Pyrophosphate is probably formed in a second-generation radical termination step. Pyrophosphate is likely formed through the reaction:

\[ \cdot \text{PO}_3^{2-} + \cdot \text{OPO}_3^{2-} \rightarrow \text{P}_2\text{O}_7^{4-} \]  

(11)

This necessitates a source of phosphate (\(\cdot \text{OPO}_3^{2-}\)) radicals. Phosphate radicals could be formed through reaction of orthophosphate and a phosphite radical via a radical propagation reaction:

\[ \cdot \text{OPO}_3^{2-} + \cdot \text{PO}_3^{2-} \rightarrow \cdot \text{OPO}_3^{2-} + \text{HPO}_3^{2-} \]  

(12)

\[ \cdot \text{PO}_3^{2-} + \cdot \text{OPO}_3^{2-} \rightarrow \text{P}_2\text{O}_7^{4-} \]  

(13)

These reactions use the strong oxidizing potential of the phosphite radical and are consistent with the reactivity towards organic compounds. Adding this phosphate radical to the radical matrix (Table 6) provides a few more P species, including peroxyphosphate and peroxypyrophosphate. Both compounds are present in MS data.

The mass spectra shown in Figure 3 and Table 4 demonstrate that there is a huge diversity of P compounds in solution when acetate is added to iron phosphide. Among the diverse chemistry of the P compounds in solution, some are organophosphorus compounds; these are discussed in the next section. The majority are multi-P oxides. How could these compounds form?

Radical reactions lack specificity, and radical reactions frequently produce unusual minor products. These minor products are formed from propagation / terminations steps like reaction of hypophosphate with a radical inducer (like \(\cdot \text{PO}_3^{2-}\)): 
\[ \text{P}_2\text{O}_6\text{H}_3^- + \text{PO}_3^{2-} \rightarrow \text{P}_2\text{O}_5\text{H}_3^- + \text{PO}_4^{3-} \]  \hspace{1cm} (14)

\[ \text{P}_2\text{O}_5\text{H}_3^- + \text{PO}_3^{2-} + \text{H}^+ \rightarrow \text{H}_4\text{P}_3\text{O}_8^- \]  \hspace{1cm} (15)

This tri-P compound is detected in the MS data. This process can repeat several times to form species like \( \text{H}_5\text{P}_4\text{O}_{10}^- \), \( \text{H}_6\text{P}_5\text{O}_{12}^- \), and so on. These reactions occur less and less frequently as the products become rarer and rarer, consistent with MS abundances.

Figure 6 summarizes the radical reaction pathway for the inorganic components of the solution. Phosphorus species formed from radical terminations in turn react with other radicals from the corrosion, resulting in an increasingly complex suite of P compounds. Each cycle or generation has progressively less total abundance for each compound but is more diverse in terms of molecular arrangement. All compounds will eventually oxidize/hydrolyze to the most stable P compound, orthophosphate.

[Figure 6]

5.2 Proposed Organic Phosphorus Reaction Pathway

Acetate is phosphorylated in solution to form a series of P-C and P-O-C linkages. The total yields for organic P compounds are ~3.6% of the total dissolved P compounds. This yield is consistent with the 0.7% mole fraction of acetate in solution. The phosphorylation products are consistent with the behavior of acetate radicals.

The reaction of acetate with a radical agent forms two distinct radicals: ‘\( \text{CH}_3\text{COO}^- \)’ and \( \text{CH}_3\text{COO}' \). The radical \( \text{CH}_3\text{COO}' \) is unstable and decarboxylates:

\[ \text{CH}_3\text{COO}' \rightarrow \text{CH}_3 + \text{CO}_2 \]  \hspace{1cm} (16)
though CO$_2$ may also form a radical. The ‘CH$_2$COO’ radical is comparatively stable (Wang et al., 2001).

The organic P compounds are broadly grouped into three sets: the P-acetyl compounds, the P-CH$_2$ compounds, and phosphonoformate (Figure 5). The most abundant species are the ones with the lowest molecular weight, consistent with relative abundances of radical species. P-acetyl compounds are formed through radical termination reactions between the inorganic radicals resulting from corrosion described in section 5.1 and the stable acetate radical described above. The P-CH$_2$ compounds may be formed through a decarboxylation reaction of acetyl phosphonate:

\[
\text{O}_3\text{PCH}_2\text{COO}^{3-} + \cdot\text{OH} \rightarrow \text{O}_3\text{PCH}_2\text{COO}^{-2} + \cdot\text{OH} \rightarrow \text{O}_3\text{PCH}_2^{-2} + \text{HCO}_3^{-} \quad (17)
\]

followed by a termination reaction to form a new group of organic P compounds (Figure 7). Alternatively, the CO$_2$ can retain the free radical and react with a phosphite radical to form phosphonoformate:

\[
\text{O}_3\text{PCH}_2\text{COO}^{-2} \rightarrow \text{O}_3\text{PCH}_2^{-} + \cdot\text{CO}_2^{-} \rightarrow \cdot\text{CO}_2^{-} + \cdot\text{PO}_3^{2-} \rightarrow \text{O}_2\text{CPO}_3^{3-} \quad (18)
\]

[Figure 7]

Only acetate, pyruvate, and possibly EDTA and acetaldehyde appear to react with the corroding phosphide powder to produce organic P compounds, whereas ethanol and acetonitrile were not phosphorylated and glycolaldehyde and acetone turned to tar in corroding solutions. Acetate, EDTA, pyruvate, glycolaldehyde, acetone, and acetaldehyde are capable of a keto-enol tautomerization transformation whereas neither ethanol nor acetonitrile tautomerize. Keto-enol tautomerization increases the acidity of protons on $\alpha$ carbons next to carbonyl groups and stabilizes radicals. Acetate has a charged functional group at a pH of 7 which repels other acetate groups, limiting
polymerization. Glycolaldehyde and acetone lack negative charges and hence both polymerized to form tar.

EDTA forms similar radicals, losing an H from the methyl bridge but the concentration of EDTA (0.025 M) used in these experiments was much lower than the concentration of acetate (0.4 M), so organic P compounds were more difficult to detect in those solutions. Acetaldehyde lacks a negative charge, but it is possible that some acetate was formed by oxidation of acetaldehyde, and that it is reaction of P radicals with this acetate that is being detected instead of reaction with acetaldehyde.

5.3 Phosphite Oxidation and Isotopic Evidence

Figure 4b demonstrates that OH radicals are produced during the corrosion of iron metal. These ‘OH radicals are produced through reaction of O₂ gas with the iron metal:

\[
\text{Fe} + \text{O}_2 \rightarrow \text{Fe}^{2+} + \text{O}_2^{2-} \quad (19)
\]

\[
\text{O}_2^{2-} + 2\text{H}^+ \rightarrow \text{H}_2\text{O}_2 \quad (20)
\]

\[
\text{Fe}^{2+} + \text{H}_2\text{O}_2 \rightarrow \text{Fe(OH)}^{2+} + \cdot\text{OH} \quad (10)
\]

An analogous reaction is known for cobalt (Leonard et al., 1998). To the best of our knowledge, this is the first report of ‘OH radical production from the corrosion of iron metal.

The detection of OH radicals is coupled to the phosphite oxidation experiment results (see P compound stability experiments). Phosphite is oxidized by ‘OH radicals produced during the corrosion of iron in this experiment through a radical propagation reaction:
This experiment demonstrates that the conversion of phosphite to orthophosphate occurs through radical reactions.

The results above match well with the isotopic data accumulated by Bryant and Kee (2006), who state that the three of the oxygen atoms share the same origin for both phosphite and orthophosphate. The origin of these three oxygen atoms is the water corroding the schreibersite. The results above provide evidence for the change from phosphite to orthophosphate is promulgated by radicals from O₂ corroding iron.

6. CONCLUSION

Phosphide corrodes in water to form three major species- hydroxyl, hydrogen, and phosphite radicals. The termination of these three radicals produces the orthophosphate, phosphite, and hypophosphate species. Secondary radical propagation reactions produce pyrophosphate and many minor P species. Interactions of these radicals with organic radicals in turn forms organophosphorus compounds, with P-C and P-O-C linkages.

The P products are most likely produced through radical termination reactions. They may be removed through oxidation, hydrolysis, or radical propagation reactions. Radical reactions are faster than oxidation or hydrolysis reactions due to the vast difference in energy between the reaction intermediates and products for radical reactions. Thus the suite of P species is the most complex while the phosphide corrodes. After all the phosphide has corroded the P compounds will oxidize to form
orthophosphate. The C-P and the P-P bonds are among the strongest bonds present, and may not be broken except on geologic timescales. The final products predicted from the long timescale of corrosion are orthophosphate with some C-P and P-P compounds. C-P compounds are well known in meteorites (Cooper et al., 1992), but P-P linkages are not. The detection of parts per thousand (with respect to total P) hypophosphate concentrations in meteorites would provide evidence for schreibersite corrosion acting on the parent bodies of meteorites. Unfortunately, hypophosphate is the least soluble of the four major inorganic P compounds, and its detection may be difficult.

6.1 Implications for the Origin of Life

Inorganic phosphates or polyphosphates have long been thought to be the most important source of phosphate for prebiotic synthesis, although reduced forms have occasionally been considered in this context (Schwartz, 1997; Peyser and Ferris, 2001; de Graaf and Schwartz 2005). Only orthophosphates are abundant in terrestrial rocks, principally calcium phosphate. As a result, numerous approaches to prebiotic synthesis using inorganic orthophosphates or polyphosphates have been explored.

Many of the early attempts to phosphorylate nucleosides used condensing agents such as cyanamide or cyanate. Unfortunately, such reactions are inefficient in aqueous solutions because water competes for the activated phosphate intermediate (Lohrmann and Orgel, 1968). More recently it has been shown that AMP can be converted to ADP and ATP by cyanate in the presence of insoluble calcium phosphates (Yamagata, 1999), though the initial phosphorylation to AMP is still unclear.
Despite its importance to biology and intense research, the mechanism by which P was incorporated into biomolecules is still unknown (Schwartz, 1997). With the exception of apatite, none of the inorganic P sources used in successful phosphorylation experiments occur in abundance in natural systems. Polyphosphates are produced through heating in μM quantities (Yamagata et al., 1991), with phosphorylation yields of ~1% (Schwartz and Ponnampemura, 1968) for a total organophosphate concentration of ~10⁻⁸ M, too low for most origin of life scenarios.

Radical termination reactions overcome the energetic difficulties traditionally associated with origin of life studies. The reaction of a P radical with an organic radical or organic compound produces an organic P compound without the need for condensing agents, elevated temperatures, or geologically rare P compounds. Researchers are increasingly turning to radical reactions to produce organic compounds, including organophosphates (e.g., Simakov and Kuzicheva 2005).

Organic P compounds produced during schreibersite corrosion have P-C and P-O-C linkages. Phosphonates have P-C linkages and are ubiquitous in biochemistry, though little is known of their exact biochemical function (Hilderbrand and Henderson, 1983). The formation of the critical P-O-C linkage via simple schreibersite corrosion emphasizes the role that meteorites may have played in the “priming” of the early Earth with organic P compounds. The total yield of P-O-C type compounds is approximately 1% of the total soluble P compounds produced. The thermodynamic instability of schreibersite will result in complete corrosion of this mineral with time, and the total yield of organophosphates from schreibersite will approach 1%. The main organophosphate produced is phosphoglycololate, which is minor metabolic byproduct in modern
metabolism, but the production of this compound indicates that changing the initial conditions or varying the organic chemistry may produce critical P-O-C compounds.

Schreibersite is a ubiquitous mineral in many meteorites, especially the iron meteorites. Iron meteorites are the main component of heavy meteorite falls \(10^3 \text{-} 10^{10}\) kg, see Bland and Artemieva, 2006) and during the late heavy bombardment, could have deposited between \(10^6 \text{-} 10^{10}\) kg of phosphorus as reactive schreibersite across the surface of the Earth per year (Pasek and Lauretta, 2005; in prep.). Using the yields from the acetate experiments, this would correspond to a production rate of \(10^5 \text{-} 10^8\) kg of organophosphate compounds per year through schreibersite corrosion.

This study provides the results of a series of experiments and analyses of synthetic schreibersite corrosion. It demonstrates that a schreibersite most likely corrodes by a radical reaction pathway, and that radicals are capable of overcoming energetic barriers traditionally associated with the prebiotic synthesis of organic P compounds. As such, it provides strong evidence that meteorites may have been critical to the development of life on the Earth.

**Acknowledgements**

We thank S. Benner and G. Cody for helpful discussions. This work was supported by NASA Exobiology, grant NAG5-13470 (DSL), and by a NASA GSRP grant (MAP).
References


Table Captions

Table 1. Experiments reported in this work. NMR is nuclear magnetic resonance spectrometry, ES-TOF-HR-MS is negative electron spray time of flight-high resolution-mass spectrometry, ESR is electron spin resonance spectrometry. More details can be found in Pasek (2006).

Table 2. Results of corrosion experiments. P compounds given in μM. Pi is orthophosphate, P3 is phosphite, P4 is hypophosphate, and PPi is pyrophosphate. “Other” includes a number of varied species with definite identifications listed. Errors are ~1% for ICP-MS, and ~10% for NMR.

Table 3. NMR peak location data, pH of 13.

Table 4. Mass spectral peaks of NH4OOCCH3 and Fe3P corrosion solution. Abundance is compound abundance in μM calculated from NMR results, Mass is the peak location, and ID is the proposed identification (a “?” implies uncertainty). “NMR” under “Note” indicates identification in the NMR spectrum, “H2” implies either H2 loss or gain from a main peak and may correspond to a radical rearrangement (e.g., Iwahashi et al., 1992), “Org” identifies an organic-P compound. Named compounds are organic-P compounds. Differences between the peak location and the calculated mass for the ID are all less than 100 ppm. Nitrogen has been excluded from peak identifications in table 14. NH4+ is a relatively inert species acting mostly to preserve charge balance. Despite this, N species may be present at low total concentrations.
Table 5. Results in terms of percent of total NMR area for P compound stability experiments, Pi is orthophosphate, P3 is phosphite, P4 is hypophosphate, PPi is pyrophosphate, and P1 is hypophosphite.

Table 6. Radical termination matrix. Non-radical species are formed through the termination of two radical species. The three radicals H, 'OH, and 'PO$_3^{2-}$ can terminate in nine different ways, three of which are redundant. Additional compounds produced if the phosphate radical, 'OPO$_3^{2-}$, is added to the radical matrix are shown in the final column and row of this table. All predicted P compounds are observed in the MS data.
Table 3.

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*Also in H₂O₂ experimental corrosion.
Table 4.

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Table 6.

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Figure 1. Inorganic phosphorus compounds discussed in text. Bryant and Kee (2006) identified I-IV, and produced VI under UV light.

Figure 2. A comparison of NMR spectra. a) Fe$_3$P with H$_2$O$_2$ (0.6 M) and b) Fe$_3$P with sodium acetate ± ethanol (0.4 M). The peak heights of both spectra have been increased substantially to allow for comparison between solutions. Major peaks are otherwise identical to those of figure 4. Note the organic P compounds present between 15-20 ppm in b).

Figure 3. Mass spectrum of NH$_4$OOCCH$_3$ and Fe$_3$P corrosion solution. The tallest peaks are labeled. Note that concentration is proportional to peak height and total mass and that the abundances in table 5 were determined by standardizing the peak heights to known NMR concentrations.

Figure 4. EPR spectra of solutions.

Figure 5. Structures of organic P compounds produced.

Figure 6. Proposed inorganic reaction pathway summary.

Figure 7. Proposed organic P reaction pathway summary.
Figure 1.

I. Orthophosphate

\[ \text{H-P-O-H} \]

II. Pyrophosphate

\[ \text{H-P-O-P-O-H} \]

III. Phosphite

\[ \text{H-P-H} \]

IV. Hypophosphate

\[ \text{H-P-O-P-O-H} \]

V. Diphosphate

\[ \text{H-P-O-P-O-H} \]

VI. Hypophosphite

\[ \text{H-P-H} \]
Figure 2.

a) \( H_2O_2 \)

\[ P_2O_6^{4-} \quad PO_4^{3-} \quad HPO_3^{2-} \quad P_2O_7^{4-} \]

b) \( NaOAc \)

\[ \text{org-P} \quad P_2O_5^{4-} \quad PO_4^{3-} \quad HPO_3^{2-} \quad P_2O_7^{4-} \]

\[ \downarrow \]
Figure 3.
Figure 4.

a) Water and DMPO

![Graph showing EPR amplitude vs magnetic field (Gauss) for water and DMPO.]

b) Basic solution (pH 13), Fe metal, and DMPO

![Graph showing EPR amplitude vs magnetic field (Gauss) for basic solution, Fe metal, and DMPO.]

Magnetic Field (Gauss)
Figure 4 cont’d.
c) Water, phosphide powder, and DMPO
Figure 5.

I. Acetyl-P compounds

\[
\begin{align*}
\text{Acetyl phosphonate} & : \quad \text{HO-}P\text{O-}C_2\text{O}_3 \text{H}_3 \\
\text{Phosphoglycolate} & : \quad \text{HO-}P\text{O-}C_2\text{O}_3 \text{H}_3 \\
\text{Hypophosphatoacetate} & : \quad \text{HO-}P\text{O-}C_2\text{O}_3 \text{H}_3 \\
\text{Tri-P-acetate} & : \quad \text{HO-}P\text{O-}C_2\text{O}_3 \text{H}_3 \\
\text{Quad-P-acetate} & : \quad \text{HO-}P\text{O-}C_2\text{O}_3 \text{H}_3 \\
\text{Penta-P-acetate} & : \quad \text{HO-}P\text{O-}C_2\text{O}_3 \text{H}_3 \\
\end{align*}
\]

II. Methyl-P Compounds

\[
\begin{align*}
\text{Hydroxymethyl phosphonate} & : \quad \text{HO-}P\text{O-}C\text{H}_2 \\
\text{Methyl diphosphite} & : \quad \text{HO-}P\text{O-}C\text{H}_2 \\
\text{Phosphonomethylphosphate} & : \quad \text{HO-}P\text{O-}C\text{H}_2 \\
\end{align*}
\]

III. Phosphonoformate:
Figure 6.

Increasing abundance

Oxidation / Hydrolysis

Non-radical products

- HPO$_3^-$, P$_2$O$_5^+$, HPO$_4^{2-}$
- H$_2$O$_2$, H$_2$O, H$_2$

Radicals generated

- H$_3^+$, OH$^-$, PO$_3^{2-}$

Initiation

- $\text{Fe}_3\text{P}$ + H$_2$O

Termination

- Propagation

P$_2$O$_5^+$, H$_2$P$_2$O$_6^2-$
- HP$_2$O$_4^-$, P$_2$O$_8^-$
- HPO$_4^{2-}$, P$_2$O$_4^-$, ~30 others

P$_3$O$_9$$^{4-}$, P$_4$O$_{10}^{6-}$
- ~3,200 others

Et cetera

Many
Figure 7.