Fig. 7. Cascade stabilization in diphosphino-substituted carbocations.



corresponding *P*-methyl phosphorus ylide **3** in near-quantitative yield (an ylide is a compound in which a positively charged atom from group 15 or 16 is connected to a carbon atom carrying a negative charge) (Fig. 2). Further investigation of **2'** through analysis of the Laplacian of the electron density (20) and the electron localization function (21) reveals a strong "banana" P–P bond with large p character (the bond ellipticity is 0.45) in the plane of the ring and partial double-bond character (Wiberg bond indices = 1.2) for the P–C bonds (Fig. 5).

The calculations found that the open planar form ($C_{2\nu}$ symmetry) of type **1'a**, analogous to that of amidinium salts, is not a minimum on the potential energy surface. This illustrates the striking difference between phosphorus and nitrogen chemistry (Fig. 6). However, pyramidalization at one of the phosphorus atoms leads to **1'b**, which is not only an energy minimum, but is also lower in energy by 24 kJ/mol than the cyclic form **2'**. The experimental observation of **2** can be explained by the presence of bulky diisopropylamino groups because the steric demand in the linear form is larger than in the cyclic system.

Derivative 2 can be considered as resulting from a "cascade stabilization" of the electrondeficient carbocation center. As in the case of monophosphinocarbenium ions such as A, the first phosphorus atom gives electrons to the carbocationic center and becomes positively charged and, therefore, highly electrophilic (22). The second phosphorus atom then acts as a Lewis base toward the first (Fig. 7).

This type of transformation of a cationic center into an anionic center is unlikely to be unique and should be of important synthetic utility. For example, preliminary calculations predict that the corresponding diphosphinonitrenium ions should exist in the cyclic form, with a negatively charged nitrogen atom. The synthesis of 2, combined with the recent preparation of a tricyclic tetraphosphabenzene valence isomer (23) (calculated in the carbon series to be 933 kJ/mol less stable than benzene), also suggests that the unique electronic properties of heavier main-group elements will allow for the preparation of many new structural moieties that are difficult or impossible to access in the corresponding carbon and nitrogen series.

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 Crystal data for 2 is as follows. Cell constants and an
- 9. Crystal data for 2 is as follows. Cell constants and an orientation matrix for data collection correspond to the monoclinic space group *P*2,/*c*, with the following cell parameters: *a* = 15.031(2) Å, *b* = 10.218(1) Å, *c* = 24.906(3) Å, β = 100.99(1)°, and *V* (cell volume) = 3755.1(8) Å³. There is one molecule, C₂₉H₆₅F₃N₄O₃P₂SSi, per asymmetric unit (number of formula units per cell = 4), giving a formula weight of

696.94 and a calculated density D_c of 1.233 Mg m⁻³. The data of the structure were collected on a STOE Imaging Plate Diffraction System diffractometer at a temperature of 173(2) K with graphite-monochromated Mo K α radiation (wavelength = 0.71073 Å) by using $\boldsymbol{\phi}$ scans. We solved the structure by direct methods, using SHELXS-97 [G. M. Sheldrick, Acta Crystallogr. A46, 467 (1990)]. The linear absorption coefficient, μ, for Mo K α radiation is 0.252 mm⁻¹. The structure was refined with all data on F^2 with a weighting scheme of $w^{-1} = \sigma^2(F_o^2) + (g_1 \cdot P)^2 + (g_2 \cdot P)$ with $P = (F_o^2 + g_1 \cdot P)^2 + (g_2 \cdot P)$ with $P = (F_o^2 + g_1 \cdot P)^2 + (g_1 \cdot P)^2 + (g_2 \cdot P)^2$ 2F 2)/3 using SHELXL-97 [G. M. Sheldrick, Program for Crystal Structure Refinement (Universität Göttingen, Göttingen, 1997)] [g is the weighting factor (g1 0.069100 and $g^2 = 3.406500$), and F_0 and F_c are the observed and calculated structure factors, respectively]. All nonhydrogen atoms were treated anisotropically. The hydrogen atoms were located by difference Fourier maps and refined with a riding model. The final cycle of full-matrix least squares refinement was based on 21,874 measured (5180 unique) reflections and 407 variable parameters and converged with the unweighted agreement factor equal to R1 $[l > 2\sigma(l) =$ 0.0484 (/ is the observed reflection intensity)] and the weighted factor wR2 = 0.1328 for all data. Crystallographic details are available at www.sciencemag.org/ feature/data/1051650.shl.

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history was significantly different from the

preanthropogenic sulfur cycle (4-9). Here we

report sulfur multiple-isotope measurements

(of $\delta^{33}S$, $\delta^{34}S$, and $\delta^{36}S$) of sulfide and sul-

fate minerals from Precambrian sedimentary

and metasedimentary rocks and use them to

document that a profound change occurred in

the sulfur cycle between 2090 and 2450 Ma.

processes produce isotopic fractionations that

depend on the relative mass differences be-

Thermodynamic, kinetic, and biological

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Atmospheric Influence of Earth's Earliest Sulfur Cycle

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Mass-independent isotopic signatures for δ^{33} S, δ^{34} S, and δ^{36} S from sulfide and sulfate in Precambrian rocks indicate that a change occurred in the sulfur cycle between 2090 and 2450 million years ago (Ma). Before 2450 Ma, the cycle was influenced by gas-phase atmospheric reactions. These atmospheric reactions also played a role in determining the oxidation state of sulfur, implying that atmospheric oxygen partial pressures were low and that the roles of oxidative weathering and of microbial oxidation and reduction of sulfur were minimal. Atmospheric fractionation processes should be considered in the use of sulfur isotopes to study the onset and consequences of microbial fractionation processes in Earth's early history.

The present-day sulfur cycle is strongly influenced by anthropogenic emissions, biological processes, and oxidative weathering of continental sulfides (1-3). It has been debated whether the sulfur cycle early in Earth's

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tween different isotopes of sulfur and oxygen. As a result, observed isotope variation can be related by $\delta^{33}S = 0.515 \times \delta^{34}S$, $\delta^{36}S = 1.90 \times \delta^{34}S$, and $\delta^{17}O = 0.52 \times \delta^{18}O$ (10). The quantities $\Delta^{33}S$, $\Delta^{36}S$, and $\Delta^{17}O$ (11) reflect the deviation of measured isotope compositions ($\delta^{33}S$, $\delta^{34}S$, and $\delta^{36}S$, or $\delta^{17}O$ and $\delta^{18}O$) from mass fractionation arrays with origins at $\delta^{33}S_{CDT} = 0$, $\delta^{34}S_{CDT} = 0$, and $\delta^{36}S_{CDT} = 0$ for sulfur, and at $\delta^{17}O_{SMOW} = 0$ and $\delta^{18}O_{SMOW} = 0$ for oxygen (CDT, Canyon Diablo Troilite; SMOW, standard mean ocean water).

In addition, several isotopic fractionation processes are also known to produce massindependent compositions ($\Delta^{33}S \neq 0$, $\Delta^{36}S$ \neq 0, or Δ^{17} O \neq 0) (12–14). These include fractionations that result from hyperfine interactions in solid and liquid phases and an increasing number of gas-phase reactions (13). The hyperfine effect derives from spinorbit coupling in isotopes with odd-mass nuclei and is therefore limited to isotopomers that contain these nuclei (such as ³³S and ¹⁷O). Gas-phase mass-independent fractionations have been documented for a number of sulfur phases in laboratory experiments (SO₂, H_2S , CS_2 , and S_2F_{10}) (12, 15, 16), but the physical chemical origin of the effect is still uncertain. The role of gas-phase mass-independent chemistry in determining the oxygen isotopic compositions of many atmospheric species is unequivocal (13), and these types of reactions may also play an important role in determining the sulfur isotope compositions of atmospheric species (15).

To examine sulfur isotope variablility in early Earth, we extracted sulfur from sulfide and sulfate minerals in a variety of Precambrian samples. Sulfur isotopic analyses fall into two groups defined on the basis of Δ^{33} S values and geological age (Fig. 1; also see Web table 1 at www.sciencemag.org/feature/ data/1052160.shl). Samples younger than 2090 Ma display a range of Δ^{33} S values from -0.11 per mil (‰) to 0.02‰ and are consid-

Fig. 1. Plot of Δ^{33} S values versus sample age. Variable Δ^{33} S values for samples with geological ages greater than the transition interval at 2090 to 2450 Ma are interpreted as an indication of atmospheric influence of the sulfur cycle. Homogenous Δ^{33} S values for samples with geological ages less than 2090 Ma are interpreted as



ered to be consistent with fractionation by mass-dependent processes. Sulfide and sulfate samples older than 2090 Ma but younger than 2450 Ma exhibit a range of Δ^{33} S varying between 0.02 and 0.34‰, and samples older than 2450 Ma exhibit a much larger range of Δ^{33} S, varying between -1.29 and 2.04‰. This variation is consistent with large massindependent compositions. The relation between Δ^{33} S and Δ^{36} S (Fig. 2) and between $\delta^{36}S$ and $\delta^{34}S$ for samples older than 3000 Ma (Fig. 3) rules out the possibility that hyperfine interactions account for the observations. The most likely explanation of observed $\Delta^{33}S$ and $\Delta^{36}S$ values for early Proterozoic and Archean sulfide and sulfate is, therefore, the presence of one or more gasphase, mass-independent chemical reactions in the sulfur cycle.

Photochemical reactions are thought to be important in the early Proterozoic and Archean atmosphere and may be relevant to the sulfur cycle before 2090 Ma. Photochemical reactions have been suggested as the source of nonzero Δ^{33} S values in martian samples (martian meteorites) (15). Our data indicate that a profound change occurred in the sulfur cycle between 2090 and 2450 Ma. This change might represent the onset of a process capable of homogenizing mass-independently fractionated sulfur reservoirs or the suppression of one or more atmospheric reactions that had occurred before this interval.

Two basic models have been suggested for Earth's early sulfur cycle. The first is that the Archean sulfur cycle did not differ significantly from the preanthropogenic sulfur cycle (7–9) and that the dominant source of oceanic sulfate was oxidative weathering of continental sulfides and weathering of continental sulfates. The second is that oxidative weathering did not play a significant role in the Archean sulfur cycle and that the principal source of oceanic sulfate was photochemical oxidation of volcanogenic sulfur species in the Archean atmosphere (δ). Our mass-



It is also possible that the atmospheric chemistry responsible for producing the observed mass-independent sulfur isotopic compositions before 2090 Ma may have stopped operating as a result of a change in atmospheric composition or actinic flux. Some have suggested that changes in the solar spectrum [ul-







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traviolet (UV) and visible wavelengths] resulting from main sequence brightening (18) could have affected atmospheric chemistry in Earth's earliest atmosphere (19, 20). Although this change may account for our observations, changes in the abundance of absorbing species in the upper atmosphere exert a much stronger influence on lower atmospheric UV and on photochemistry (21). Higher surface UV resulting from a reduced ozone column depth in an atmosphere with low oxygen concentrations (22-24) may have played a role in determining atmospheric sulfur chemistry and in generating isotopically distinct sulfur reservoirs. Photolysis of SO₂, for instance, results in production of SO_3 (25) that converts to H_2SO_4 upon contact



Fig. 3. Three isotope plots of (A) δ^{36} S versus δ^{34} S and (B) δ^{33} S versus δ^{34} S for samples older than 3000 Ma. The array formed on (A) follows the relation δ^{36} S = 2.17 (±0.1) × δ^{34} S [calculated using York (31)] rather than the mass-dependent relation δ^{36} S = 1.90 (±0.01) × δ^{34} S (10). The data in (B) do not follow the tightly constrained mass-dependent relation δ^{33} S = 0.515 (±0.005) × δ^{34} S (10) but scatter on the diagram. Barite and chert-plus-barite samples are plotted as triangles. All other samples are plotted as diamonds. These arrays are inconsistent with biological fractionation processes and are consistent with a sulfur cycle that is strongly influenced by atmospheric chemical reactions and atmospheric oxidation reactions.

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with water. This photochemical oxidation sequence has been shown to produce mass-independently fractionated sulfur (15). Whereas oxygen isotopic signatures of present-day products of atmospheric oxidation of reduced sulfurbearing gases have been shown to possess mass-independent oxygen isotopic compositions that are thought to be the result of oxidation by isotopically anomalous ozone and hydrogen peroxide (26, 27), our oxygen isotope data for sulfate from barites from 3300 to 3400 Ma are mass-dependent ($\Delta^{17}O = 0.01 \pm$ 0.06‰). The lack of a similar mass-independent signature in our Archean barites may be a further indication that the sulfur oxidation pathways were different from that of the current atmosphere, possibly reflecting the exchange of sulfite (a product of SO₂ photooxidation) with water during the formation of sulfate. If this is the case, the buildup of atmospheric oxygen may have helped shut down the atmospheric chemical reactions that were responsible for generating isotopically anomalous sulfur-bearing reservoirs. An alternative explanation of the oxygen isotope data is that the original atmospheric oxygen isotopic signature has been lost because of exchange processes that occurred after sulfate formation.

If δ^{33} S fractionations reflect atmospheric fractionation processes, it is also possible that the same applies to δ^{34} S fractionations. Some workers have divided the sulfur isotope record into a period before 2750 Ma, when the range of δ^{34} S was less than about 10‰, and a period after 2750 Ma, when the range of δ^{34} S was greater than 10% (5, 28). These workers suggest that the change in magnitude for the range of observed δ^{34} S fractionations at 2750 Ma is evidence of the onset of microbial sulfate reduction. Other workers have suggested that the smaller range of $\delta^{34}S$ before 2750 Ma also results from microbial sulfate reduction but at higher temperature conditions, possibly at lower oceanic sulfate concentrations or in closed systems (7–9). On three isotope plots (Fig. 3), our data do not follow the mass-dependent arrays that would be formed by microbial sulfate reduction (for example, $\delta^{33}S\sim 0.515\times \delta^{34}S$ and $\delta^{36}S = 1.90 \times \delta^{34}S$), and they allow that the δ^{34} S record for samples older than 2750 Ma may be entirely atmospheric in origin. Although our data do not rule out the possibility of microbial sulfate reduction, they imply that δ^{34} S values cannot be used alone to argue for the operation of (or examine the consequences of) metabolic processes that fractionated sulfur isotopes before 2750 Ma.

Similar sulfur isotope measurements can be used to resolve atmospheric inputs and to gain new insights into biogeochemical element cycles. The recent observation of massindependently fractionated sulfur in this study and in martian meteorites (15) and of mass-independently fractionated oxygen in terrestrial and martian sulfates (27, 29) opens up possibilities for identifying additional components of the sulfur cycle on Earth and on other bodies in the solar system.

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