

Understanding the nature of atmospheric acid processing of mineral dusts in supplying bioavailable phosphorus to the oceans

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Acidification of airborne dust particles can dramatically increase the amount of bioavailable phosphorus (P) deposited on the surface ocean. Experiments were conducted to simulate atmospheric processes and determine the dissolution behavior of P compounds in dust and dust precursor soils. Acid dissolution occurs rapidly (seconds to minutes) and is controlled by the amount of H⁺ ions present. For H⁺ < 10⁻⁴ mol/g of dust, 1–10% of the total P is dissolved, largely as a result of dissolution of surface-bound forms. At H⁺ > 10⁻⁴ mol/g of dust, the amount of P (and calcium) released has a direct proportionality to the amount of H⁺ consumed until all inorganic P minerals are exhausted and the final pH remains acidic. Once dissolved, P will stay in solution due to slow precipitation kinetics. Dissolution of apatite-P (Ap-P), the major mineral phase in dust (79–96%), occurs whether calcium carbonate (calcite) is present or not, although the increase in dissolved P is greater if calcite is absent or if the particles are externally mixed. The system was modeled adequately as a simple mixture of Ap-P and calcite. P dissolves readily by acid processes in the atmosphere in contrast to iron, which dissolves more slowly and is subject to reprecipitation at cloud water pH. We show that acidification can increase bioavailable P deposition over large areas of the globe, and may explain much of the previously observed patterns of variability in leachable P in oceanic areas where primary productivity is limited by this nutrient (e.g., Mediterranean).

atmospheric processing | ocean macronutrients | desert dusts

Atmospheric inputs are an important source of externally supplied nutrients to the offshore ocean (1). Although all of the inorganic nitrogen (N) is water-soluble and immediately bioavailable, most phosphorus (P) and iron (Fe) are present as minerals that are not immediately soluble in water, hence not bioavailable (2, 3). Such mineral particles, if deposited to the surface ocean, may pass through the photic zone with no effect on primary productivity, owing to their high settling velocity and low solubility (2).

Atmospheric P can be important as the major external supply to the offshore ocean, particularly in oligotrophic areas of the open ocean (1) and areas that are P-limited, such as the Sargasso Sea (4) and Mediterranean (5). The most important source of atmospheric P is desert dust, which has been estimated to supply 83% (1.15 Tg·a⁻¹) of the total global sources of atmospheric P (6). Of that dust, it is estimated that 10% is leachable P. However, observations suggest that the fraction of leachable P in dust is highly variable (7–100%) (7). Only one global modeling study primitively simulates such variability, considering the reaction of protons with apatite minerals using a kinetic approach (8). The same study indicates that deposition of P from biological particles of terrestrial origin may be as important as leachable P from dust over the ocean in some regions and certain seasons. These recent findings translate to a large predictive uncertainty of

the bioavailable P input to the oceans. Studies show that although atmospheric N and Fe supply are of importance on a global scale, atmospheric P supply plays an important secondary role, especially through colimitation with either N or Fe (9, 10). The varying demands and resilience of different phytoplankton communities can have feedbacks on local limiting nutrients on relatively short time scales (11, 12). Additionally, evidence suggests feedbacks in the surface waters that may enhance the impact of the atmospheric supply of P (1, 13). This complex picture is yet to be replicated in global biogeochemistry models; therefore, it is not currently possible to put a limit on the importance of any atmospherically supplied P.

Previous studies have shown that atmospheric processes can increase Fe bioavailability in dust before being deposited to the ocean. Insoluble Fe, principally in the form of iron oxides, can be solubilized by interaction with acid gases that reduce the pH of atmospheric water to the level where solid phase Fe species can start to dissolve (14), by interaction with organic ligands, or by UV photoreduction (15). By contrast, the principal mineral species of P in aerosols, apatite minerals (e.g., hydroxyapatite [Ca₅OH(PO₄)₃]), are only expected to be solubilized by acid processes in the atmosphere, because they do not undergo photoreduction and

Significance

Mineral dust is the most important external source of phosphorus (P), a key nutrient controlling phytoplankton productivity and carbon uptake, to the offshore ocean. The bioavailable P in dust exhibits considerable and poorly understood variability. Detailed laboratory experiments elucidate and quantify the major processes controlling P dissolution in the atmosphere. Dust exposure to acids is the main driver of P mineral transformations, and a simple direct proportionality is found between the amount of bioavailable P dissolved from the dust and acid exposure. Simulations suggest that dust acidification increases leachable P over large areas of the globe and explains much of its variability in important oceanic areas where primary productivity is limited by this nutrient (e.g., North Central Atlantic, Mediterranean).

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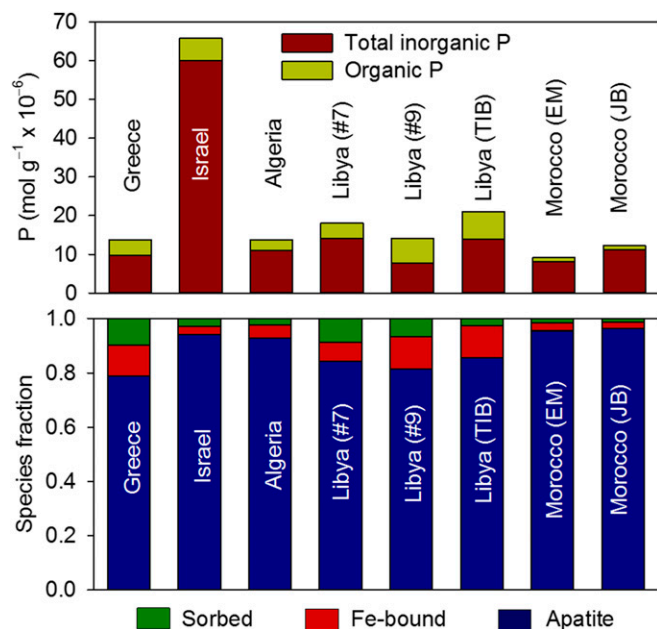


Fig. 1. P speciation in the dust and dust precursor samples, as determined by SEDEX. (*Top*) Concentrations of P present in the different samples. (*Bottom*) Relative fraction of the different inorganic P species.

calcium (Ca) is not strongly complexed by organic ligands. In a study on dust particles sampled in the eastern Mediterranean, Nenes et al. (7) found a correlation between increased leachable inorganic phosphorus (LIP) and increased acid exposure in the aerosol particles. Furthermore, the amounts of P released during acidification were consistent with the thermodynamic limit of solubility. However, no further insight on the mechanism and dissolution kinetics could be obtained.

The principal acid precursor species in the atmosphere are oxides of nitrogen and sulfur (NO_x and SO_x). These precursor species can be the result of natural processes, such as the oxidation of dimethyl sulfide released by phytoplankton in oceanic surface waters (16), volcanic eruptions (17), or lightning. However, at present, the main source of such gases in the northern hemisphere is anthropogenic (18). To a lesser degree, low-molecular-weight carboxylic acids (e.g., formic acid, acetic acid, oxalic acid) are generated in the atmosphere and can contribute acidity, especially in evaporated cloud droplets (18, 19).

Fresh and aged dust particles can be contained within cloud droplets that dissolve acidic gases but with pH levels that do not drop much below 4 (18). Although some cloud droplets condense to form rain, most cloud droplets evaporate to form wet aerosol particles. This process can result in a substantial drop in pH and an increase in ionic strength (IS) of the resultant film of water (20). This cycling between cloud droplets and wet aerosol particles can occur several times [an average of 10 cycles throughout the troposphere (18)] before the aerosol drops to earth by wet or dry deposition. Thus, the chemical conditions within and between clouds are very different, with relatively high pH and low IS in cloud droplets and low pH and high IS in wet aerosols (20–22). This cycling was investigated for its effect on Fe dissolution in the atmosphere by Shi et al. (23), who show that Fe is solubilized in wet aerosols and then reprecipitated as Fe nanoparticles in clouds. By contrast, the impact of pH changes or proton addition to atmospherically processed mineral dusts, and subsequent P dissolution, is virtually unknown.

This study sets out to investigate the nature, magnitude, and controls of atmospheric acid processes on the solubilization of mineral P in dust particles using samples collected during dust storms in Israel and Greece, as well as on dust precursors collected from surface soils in a variety of locations across the Sahara desert (map included in *SI Appendix*, Fig. S1.1). Experiments were

carried out using principally natural dust particles to mimic atmospherically relevant conditions, and thereby represent the amount of P solubilized by atmospheric acid processes. Results are interpreted by modeling the experimental systems using the geochemical PHREEQC (pH-redox-equilibrium calculation) model (24). Calculations using a global 3D atmospheric chemical transport model [Tracer Model 4 of the Environmental Chemical Processes Laboratory (TM4-ECPL)] (25) were used to estimate the potential global importance of these processes.

Results

Properties of Dusts. The highest P concentration was found in the dust sample from Israel (Fig. 1, *Top*). The sequential phosphate extractions (SEDEX) (26) revealed that apatite was the dominant P mineral (Fig. 1, *Bottom*). Total inorganic P [which includes leachable P, Fe-bound P, apatite P (Ap-P), and carbonate P] varied between 7.7 and 60.0 $\mu\text{mol}\cdot\text{g}^{-1}$, and represents 55–92% of the total SEDEX P in the sample. The remainder of the P was made up of organic P (full SEDEX data, mineral composition, and BET (Brunauer, Emmett, and Teller) surface area are provided in *SI Appendix*, section 3.1.2).

Effect of Protons and Fluid Volume on the Dissolution of Ca and P Minerals.

Fig. 2 shows the combined effects of pH and volume on the release of P from the Israel dust (legends used in this figure are used consistently in all subsequent figures and within *SI Appendix*). Both more acid pH values (at equal water volume) and higher volumes (at equal pH) yield greater P concentrations. The amount of P (Figs. 2 and 3) and Ca (Fig. 4) released from the dust was controlled by the amount of H^+ ions present and not the initial pH. Fig. 3 shows the released dissolved P plotted versus the protons in the experiments (as moles of H^+ per gram of dust). The data showed that below a critical proton concentration of ~ 0.1 mmol/g of dust [$-4 \log \text{mol}(\text{H}^+)/\text{g}$ of dust], the released P was only affected by water volume and not by the proton concentration. Above this proton concentration, an approximately linear increase in P with increasing initial proton concentration was seen, until a plateau was reached [approximately $-2.1 \log \text{mol}(\text{H}^+)/\text{g}$ of dust], where the acid-reactive mineral phosphate pool was exhausted. A similar pattern was seen with Ca (Fig. 4), although the critical proton concentration was slightly lower than seen for P. These figures indicate that above a given proton concentration, both Ca and P mineral phases are subject to dissolution. Fe was only above the detection limits in limited experiments where excess protons allowed the pH to remain low throughout the experiment. Results for the other dust samples yielded similar trends for all analytes (*SI Appendix*, section 3.2).

In all experiments the dust-solution mixtures tended to become buffered to neutral or alkali pH end points after 48 h of reaction.

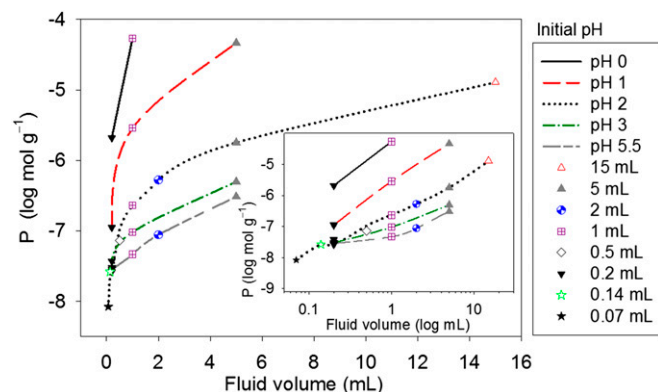


Fig. 2. Dissolved phosphate released from Israel dust in relation to pH and solution volume. Values are presented as moles per gram of dust. Dust masses were ~ 55 mg, and dust-to-volume ratios were 3.5–704.3 $\text{g}\cdot\text{L}^{-1}$. (*Inset*) Same data with the volume axis on a log scale.

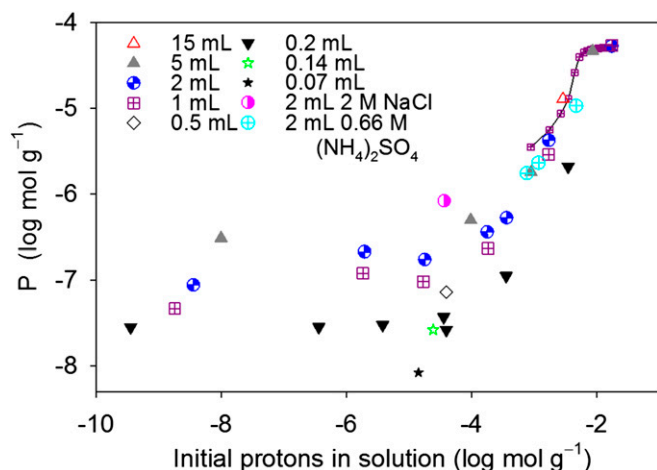


Fig. 3. Dissolved P (moles per gram of dust) released from Israel dust in relation to the absolute concentration of protons that were available for reaction at the start of the experiment. Dust masses were ~ 55 mg, and dust-to-volume ratios were 3.5–704.3 $\text{g}\cdot\text{L}^{-1}$. The crossed boxes connected by a solid line represent a sequential acid addition experiment (*SI Appendix*, section 2.1.3).

The exceptions were experiments where protons remained in excess and all Ca and P minerals were dissolved (*SI Appendix*, section 3.1.1 and Table S3.3). Even using buffered solutions (*SI Appendix*, section 3.1.1), the release profiles for P did not change from those release profiles described above. Similarly, instantaneous acid addition or slow acid diffusion also did not significantly alter the evolution of dissolved P in the experiments (*SI Appendix*, section 3.2.2). When we tested a sequential batch addition of acid, the results (Fig. 3, 1 mL experimental data connected by a solid line; also *SI Appendix*, section 3.2.3) were similar to experiments performed at higher total proton values through a single acid addition, implying consistent Ca and P mineral dissolution as the total proton exposure is increased. Finally, high IS also played a minimal role in driving P release (*SI Appendix*, section 3.2.3).

All experiments with the real dusts described above assumed that the reactive minerals (calcite and apatite) are found in every particle (“internally mixed”) at the ratio that can be calculated from the apatite and Ca concentrations determined in our study (*SI Appendix*, Table S3.1). Although dust particles may be “externally mixed” (particles of a given size have only calcite or apatite but not both), in reality the low-volume experiments brought the acidic fluid into contact with all particles, and thus the system behaved as internally mixed. Calcite (CaCO_3) controlled the H^+ content, and the released P was totally dependent on reactive mineral composition. These results were confirmed experimentally through analog dusts made up of single reactive mineral components (i.e., apatite only) or mixed reactive mineral components (apatite and calcite) that we used as dust proxies (*SI Appendix*, section 3.2.4). The analog experiments demonstrate that if dust particles were externally mixed, then there would have been more P released per H^+ amount compared with our real dust experiments (Figs. 2 and 3) because more protons are available for direct reaction with apatite.

Comparison of Experimental Results with Modeling. The close agreement between our precursor and real dust samples (*SI Appendix*, section 3.2.4) suggests that the interaction between calcite and apatite with acidity is the primary control of P solubility. This relationship is confirmed by thermodynamic calculations for a system containing only calcite, apatite, and solution (Fig. 5 and *SI Appendix*, section 3.3). Above $\sim 10^{-6}$ $\text{mol}\cdot\text{L}^{-1}$ for P and 10^{-4} $\text{mol}\cdot\text{L}^{-1}$ for Ca, there was agreement between measurements and predictions. Below these concentrations, predicted Ca and P deviated somewhat from the measurements, likely owing to desorption of loosely bound P (SEDEX sorbed pool; Fig. 1 and *SI Appendix*, Table S3.1) and Ca.

Discussion

We studied the dissolution of P-containing phases in the surface soils from areas that are known to be sources of Saharan dust (27), as well as two samples of dry deposited Saharan/Arabian dust. There was a relatively constant P speciation in the mineral composition of these dusts (Fig. 1). Given that the principal acid-soluble mineral, Ap-P, was, on average, $89 \pm 7\%$ of the total inorganic P and Fe bound-P was only $7 \pm 4\%$ of the acid-reactive mineral phases, we consider the proton reactions of only Ap-P and CaCO_3 as controlling the atmospheric conversion of mineral P to leachable forms. Other processes that affect the solubilization of Fe, such as photoreduction and organic complexation (12), are likely to have a minor role in increasing P bioavailability because only Fe bound-P will be affected by such processes. There is no evidence of organic P being affected by acidification. In addition to these acid-soluble phases, a small amount ($<7\%$) was in the sorbed inorganic-P pool (extracted by 1 M MgCl_2), which is likely to be solubilized directly into seawater (26).

Although our samples were collected or derived from widespread areas from the Sahara and Arabian deserts, they all had similar relative fractions of Ap-P, Fe-bound P, and adsorbed P. This finding is consistent with a common weathering regime across these deserts, which combines some chemical weathering, little plant growth to convert P minerals into plant biomass in situ, and often reprecipitation of CaCO_3 as caliche. However, we recognize that other regions of the Sahara, with different mineralogy [e.g., the Bodélé (28)], may exert an influence on the bulk properties of some Sahara dust plumes. Our samples had 7×10^{-4} to 3.3×10^{-3} $\text{mol}\cdot\text{g}^{-1}$ of acid-soluble Ca, which is 6–33% by mass of CaCO_3 . Although the P mineral speciation was similar between samples, there was considerable variability in the total P concentration (9.1×10^{-6} to 6.3×10^{-5} $\text{mol}\cdot\text{g}^{-1}$).

In the atmosphere, dust particles cycle between clouds, where they can become cloud condensation nuclei and wet aerosols. In clouds, there is a high water/dust ratio, low IS, and generally high pH, except in the unusual situation of very highly polluted air masses. Wet aerosols, which are generally formed when cloud water evaporates, contain a film of water and can have very low water/dust ratios, high IS, and low pH (23). Our experiments were designed to span these conditions within the practical limitations of laboratory handling. The key parameter in controlling the amount of P (and Ca) liberated from acid processing was the total amount of H^+ ions in the aqueous layer surrounding the dust particle and not the initial pH (Figs. 2–4). For all of our experiments, both CaCO_3 and Ap-P were rapidly dissolved (as measured by Ca^{2+} and orthophosphate liberated) and

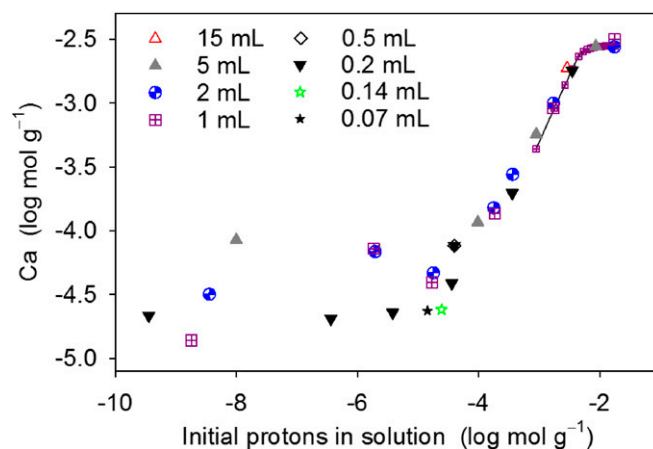


Fig. 4. Dissolved Ca (moles per gram of dust) released from Israel dust in relation to the absolute concentration of protons that were available for reaction at the start of the experiment. Dust masses were ~ 55 mg, and dust-to-volume ratios were 3.5–704.3 $\text{g}\cdot\text{L}^{-1}$.

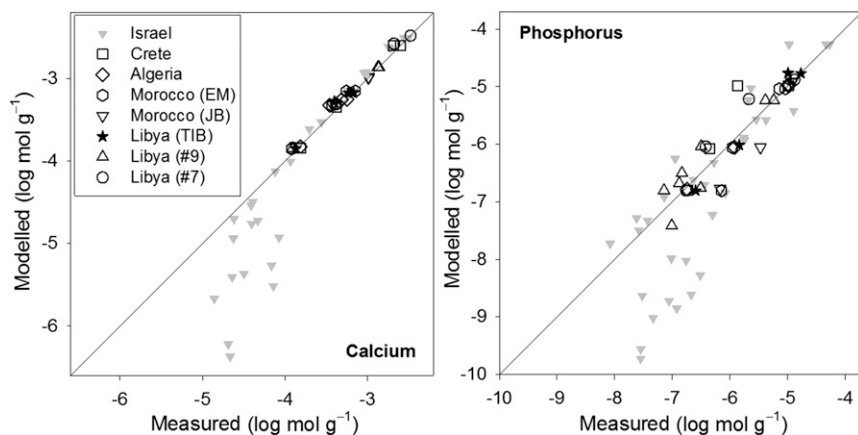


Fig. 5. Comparison of measured P and Ca dissolution from all dusts compared with equilibrium predictions made using PHREEQC.

H^+ ions were consumed on a time scale of seconds to minutes (*SI Appendix, section 3.1.1*). In most cases, the pH was controlled by rapid reaction with $CaCO_3$ to circumneutral values, as shown by the simple relationship between the decrease in H^+ ions being twice the increase in Ca^{2+} cations.

At low H^+ amounts ($<10^{-4}$ mol g^{-1} of dust), P was liberated over a wide range of H^+ ion concentrations. This result was interpreted as being due mainly to simple leaching of dissolved P from the samples into water, independent of the reaction of H^+ with acid-soluble minerals. This observation corresponds to the leachable P previously measured when dust was treated by deionized water or a very-low-strength leach (e.g., ref. 29). The amount of leached P is likely to be independent of whether there is $CaCO_3$ in the dust sample or not.

At proton concentrations above 10^{-4} mol of H^+ per gram of dust, there was a direct proportionality between P (and Ca) released and the amount of H^+ ions consumed (Figs. 2–4). This relatively simple pattern of P release continued until the level where all of the P minerals were exhausted. When the proton-reactive minerals were exhausted, the final solution pH was no longer circumneutral but remained acidic. The P released in experiments with the highest H^+ amounts corresponded approximately to the value of total inorganic P determined by SEDEX extraction. Similar trends were obtained for all of the other dust and dust precursor samples.

The relationship between P, Ca^{2+} , and H^+ ions for all of the test samples was confirmed by PHREEQC equilibrium modeling, in which there were similar increasing trends of P and Ca^{2+} obtained assuming a simple three-component model of calcite, hydroxyapatite, and H^+ ions. The model fit was better for Ca^{2+} (controlled by calcite and H^+ ions) than for P, where there was more scatter. This result is considered reasonable because there is more than one acid-soluble P mineral present in all of the natural samples, and the Ap-P is also likely to be made up from a number of different apatite minerals with different solubility behavior.

Our results also showed that increased IS caused slightly more dissolved phosphate to be liberated into solution. Where the major contributors to increased IS were ammonium and sulfate, the most common ionic species in aerosols (18), it was found that the amount of P released fits on the trend for the HCl-only data if the proton availability from the dissociation of ammonium ions (predicted from a final pH of 8) is allowed for. In our experiments using NaCl (IS = 2), which was used as a surrogate for evaporated sea salt, the liberated P increased by a factor of 4.3. In both cases, even if the IS decreases to that of rainwater, the dissolved P will remain in solution due to very slow precipitation kinetics and will be delivered as bioavailable P to the surface ocean (ref. 30 and references therein).

In the atmosphere, some mineral particles have both $CaCO_3$ and Ap-P on the same particle (internally mixed), whereas other particles have only one or the other mineral (externally mixed). Because acidity in the atmosphere is transferred to dust via the gas phase, the amount of dissolved P will vary between particles, depending on the amount of $CaCO_3$. There were no data available characterizing the degree of external mixing of $CaCO_3$ and Ap-P in

dust samples. However, we expect many of the particles to contain both $CaCO_3$ and Ap-P, because in many sedimentary rocks and desert soil systems, there is a close association between $CaCO_3$ and Ap-P. Both minerals precipitate in marine sediments during diagenesis (31), and such marine sediments are generally the source of Ap-P in desert soils (32). Furthermore, caliche forms in many desert soils, where Ca and bicarbonate produced by in-situ weathering are precipitated (33). However, our results show that in the situation where external mixing does occur, more P would be solubilized because on those particles with only Ap-P, there was no consumption of H^+ by $CaCO_3$ because it was absent from this experiment (*SI Appendix, section 3.2.4*). In reality, although we predict that many dust particles will be internally mixed, there will be a continuum and the balance between these two extremes may not be the same for all dust events.

Comparison with Fe Processes. The fraction of bioavailable Fe supplied to the ocean is also increased by acidic atmospheric processes (34). However, Fe dissolution is much slower than Ap-P (35). This behavior means that for internally mixed particles, essentially all of the H^+ ions will be neutralized by $CaCO_3$ before they react with Fe minerals. However, Fe-bearing particles are often externally mixed as clays (36) or Fe-rich particles (37). Thus, dissolution of Fe and P can happen simultaneously on different particles. Furthermore, Fe dissolution occurs principally in wet aerosols (low pH with high IS) (23). When the aerosol particles are activated into clouds at lower pH, the Fe is likely to reprecipitate as nanoparticles (38), whereas P does not reprecipitate. The only removal of P will be adsorption onto fresh Fe nanoparticles if both processes occur on the same particle/droplet. In that situation, both Fe and sorbed P are likely to be bioavailable.

The Potential Importance to the Global P Cycle and Supply to the Ocean. The main acids in the atmosphere, sulfuric and nitric acids, are generated by the oxidation of sulfur and N gases emitted by biogenic, volcanic, and anthropogenic sources (16–18), the latter of which is dominant at present. It has been noted that there is a higher fraction of LIP in polluted air masses (39, 40). Based on field samples collected in Crete, Nenes et al. (7) provided direct evidence of the increase in LIP in aerosols with increasing aerosol acidity. They suggest that this increased LIP is from polluted air masses from southern Europe bringing acid gases from the north to mix with the Saharan air mass. The increased fraction of LIP in atmospheric aerosols over the Bay of Bengal compared with atmospheric aerosols over the Arabian Sea has also been interpreted as due to the effect of acid processing of aerosols caused by anthropogenic activities (41). The action of the processes identified here may explain why aerosol samples across the North Central Atlantic can have greater fractions of soluble P than dust soil precursors or aerosols collected closer to the dust source (42). Our results demonstrate that atmospheric acid processes are extremely efficient in solubilizing P and provide further support to the finding by Nenes et al. (7) that acids can increase

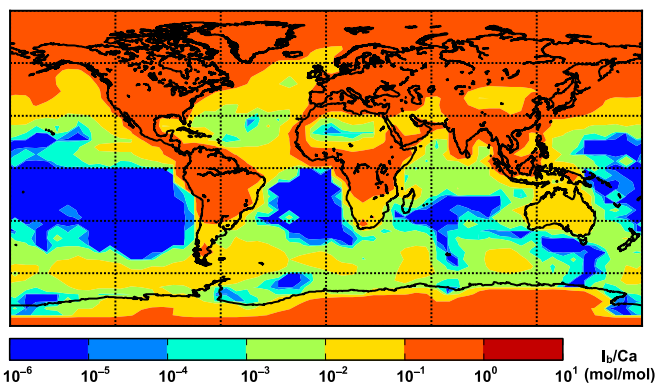


Fig. 6. Annual average ion balance over aerosol Ca. Calculations are performed with the global model framework of Myriokefalitakis et al. (25) using current-day aerosol emissions. Values of the ratio above 10^{-1} indicate regions where considerable solubilization of dust P is expected, where dust aerosol is present.

the delivery of bioavailable P from dust to the oceans. The amount of N fixation is also likely to increase because dust that has been acid-processed will contain increased amounts of both bioavailable P and Fe, which have been shown to limit N fixation (9).

To examine whether acid dissolution of dust P can be globally important, we simulate the acid exposure of dust and examine the extent to which it can occur. To do so, we quantify the ratio between dust Ca and its acid exposure. We quantify the ion balance, $I_b = 2[\text{SO}_4] + [\text{NO}_3] + [\text{Cl}] - 2[\text{Ca}] - [\text{NH}_4] - [\text{Na}] - 2[\text{Mg}] - [\text{K}]$ (where [X] represents the concentration of species X in the aerosol sample in moles per cubic meter of air) over aerosol Ca (Fig. 6; also *SI Appendix, section 3.4.*). Calculations are carried out with the global model framework of Myriokefalitakis et al. (25) using current-day aerosol emissions, and results are shown for the coarse fraction of dust and the model surface layer, which are representative of the dust that deposits to the surface. Values of the ratio above 10^{-1} indicate regions where considerable solubilization of dust P is expected. The simulations clearly indicate that the flux of bioavailable P over considerable regions of the ocean can be substantially increased by acidified dust. This increase is likely to be greater than the 0.59 Tmol of carbon per year that can be calculated from the bioavailable P predicted by the recent global modeling work that accounts for the kinetic Ap-P dissolution process (8).

In many locations, there has been a greater increase in N-containing gases, particularly NO_x and ammonia (NH_3), than in particulate P, which is transported in the atmosphere. The molar ratio is typically greater than 16:1 and often much larger. It has been reported that this supply tends to make surface waters more P-limited (29, 43). Our results suggest that this transformation is somewhat moderated by the relative increase in bioavailable P due to interaction between NO_x and dust particles. As a result, phytoplankton biomass and carbon uptake will increase even in areas of the ocean that are N-limited in the short term because P remains the element that causes a longer term increase in primary productivity (44). The implications of this added carbon export from anthropogenic pollution to ocean ecosystems has the potential to be widespread and considerable, affecting global primary productivity and the carbon cycle.

Methods

Dust and Precursor Dust Sources. Two dust samples and six size-fractionated dust precursor samples were used in this study (Table 1). The majority of the experiments were carried out with a dust sample deposited on a clean, flat surface during a dust storm in Rosh Pina, Israel (collected first in May 2012). A second dust sample was collected between June 1 and June 9, 2013, from a solar panel in Heraklion (Crete, Greece). Based on back-trajectory data from the HYSPLIT model (ready.arl.noaa.gov/HYSPLIT.php), the origins of these dust samples were the deserts of Saudi Arabia/Jordan/Iraq and North Africa, respectively. These two real dusts were used directly as collected, without size

fractioning or other preconditioning. In addition, six dust precursor samples collected from a variety of locations (mainly dry stream or lake bed soils) within the Sahara Desert (Table 1) were used to generate size-fractionated dusts using a dust tower separation and filtration methods (45). We used the $<10\text{-}\mu\text{m}$ fractions for our experiments, similar to previously used dust precursors that have been shown to be analogous to atmospherically sampled dust (45, 46). In this present study, the term “dust” is used to refer to both the dust precursor and the real dust samples unless otherwise stated.

Experimental Procedures.

SEDEX sequential P extraction. P speciation among the different operationally defined P pools was determined on 50–100 mg of each dust sample (Table 1) following the SPEXman SEDEX sequential extraction scheme (47), with the modification that step IIA for Fe-bound P followed the procedure of MacDonald (48). This modification removes the use of citrate, slightly alters the pH to 7.5, and increases the reaction time to 10 h. The absence of citrate means that the resulting solutions do not require any pretreatment, other than dilution, before analysis by the molybdate blue method (discussed below). Five P species were differentiated, namely: leachable or loosely sorbed; Fe-bound; a combined pool containing authigenic apatite, biogenic apatite, and CaCO_3 -bound; detrital apatite plus other inorganic P; and organic P. We defined apatite (Ap-P) as the combined phases extracted as diagenetic and detrital apatite, and CaCO_3 -bound P in the sequential SEDEX extraction scheme. Total inorganic P was defined as the sum of all of the phases except organic P. Methods for mineral composition and surface area analysis are in described in *SI Appendix, section 2.1.*

Phosphorous release experiments. The following experiments were performed on the dust samples using pH-adjusted but unbuffered solutions (HCl, Sigma-Aldrich; $\geq 37\%$ American Chemical Society reagent grade in 18.2 M Ω /cm of Milli-Q water) in an end-over-end stirrer for 48 h. Israel dust (55 ± 3 mg) was used as follows: pH 2 HCl with volumes of 70, 140, 210, and 500 μL and 1, 2, 5, and 15 mL; 0.2 mL of HCl at pH 0, 1, 2, 3, 4, and 5.5; 1 mL of HCl at pH 0, 1, 3, 4, and 5.5; 2 mL of HCl at pH 0.3, 1.3, 2.3, 3.3, 4.3, and 5.5; and 5 mL of HCl at pH 1, 3, and 5.5. Other dusts (30 ± 1 mg) were used at a volume of 1.2 mL at pH 0, 1, 1.8, and 2.4. Several initial experiments were carried out to determine the pH buffering capacities and the proton-dust reaction rates. We showed that almost all protons were consumed within ~ 200 s of starting the reaction and that the phosphate release was equally fast regardless of initial pH (*SI Appendix, sections 2.1.1 and 3.1.1*). Nevertheless, we performed the majority of our experiments over a conservative time period of 48 h, by which time all changes in concentration had been completed. At the end of these experiments, the mixture was passed through a 0.45- μm syringe filter (13-mm Whatman Puradisc polyethersulfone), and dissolved phosphate was determined as described below. In the experiments with solution volumes ≥ 2 mL, the final pH was measured following the 48-h exposures. The effect of high IS was tested by adding either ammonium sulfate or NaCl to reach $IS = 2$ (*SI Appendix, section 2.1.2*). Furthermore, additional experiments were carried out to evaluate the effect of sequential solution addition (*SI Appendix, section 2.1.3*), the effect of acid addition via dialysis (*SI Appendix, section 2.1.4*), and the behavior of simple laboratory-prepared dust analogs (*SI Appendix, section 2.1.5*).

Chemical Analysis of Supernatants. Dissolved inorganic P was analyzed using the molybdate blue reaction (49) after suitable dilution with matrix-matched standards on a segmented flow analyzer. For high concentrations (>50 nmol·L $^{-1}$), this analysis was done on a SEAL Analytical AA3. The relative standard deviation (RSD) was 2.2% ($n = 8$), and the limit of detection (LOD; $3 \times \text{SD}$ of blank) was 12 nmol·L $^{-1}$. Lower concentration samples were analyzed using a 100-cm WPI Liquid

Table 1. Samples used in this study and their source locations

Country of origin	Location	Coordinates
Dusts		
Greece	Crete	35°19'51"N, 25°40'04"E
Israel	Rosh Pina	32°58'12"N, 35°33'32"E
Precursor dusts		
Algeria	Bordj Mokhtar	21°19'30"N, 0°56'46"E
Morocco	El Miyit	30°21'53"N, 5°37'29"W
Morocco	Jebel Brahim	29°56'12"N, 5°37'43"W
Libya	7	32°02'42"N, 22°18'01"E
Libya	9	32°36'47"N, 22°11'42"E
Libya	Tibesti	25°35'N, 16°31'E

Locations are shown in the map included in *SI Appendix, Fig. S1.1*.

Waveguide Capillary Cell coupled to an Ocean Optics USB2000+ spectrophotometer with a precision ($n = 6$ of $60 \text{ nmol}\cdot\text{L}^{-1}$ samples) of 1.6% and a LOD of $2 \text{ nmol}\cdot\text{L}^{-1}$. Dissolved Ca and Fe concentrations in the supernatants were measured using a Thermo Scientific iCAP 7400 Radial inductively coupled plasma-optical emission spectrometer. The Ca and Fe detection limits were $<0.1 \text{ nmol}$ and $500 \text{ nmol}\cdot\text{L}^{-1}$, respectively, with RSDs of 1.5% and 1.8%, respectively [based on eight replicates of a $13 \text{ nmol}\cdot\text{L}^{-1}$ (Ca) or $9.0 \mu\text{mol}\cdot\text{L}^{-1}$ (Fe) standard]. Finally, pH was measured with a Mettler Toledo Seven Excellence meter coupled to an Inlab Expert Pro-ISM pH electrode calibrated with three National Institute of Standards and Technology traceable standard buffers (pH 4, pH 7, and pH 9.2; Mettler Toledo).

Geochemical Modeling of Experimental Systems. To evaluate the experimental results in terms of predicted equilibrium results, we used the geochemical modeling code PHREEQC (24), with the Lawrence Livermore National Laboratory database. Input conditions were based on the experimental starting solution conditions and assuming only hydroxyapatite and calcite were the reactive minerals present. The relative concentrations of these components were based on the dissolved Ca and P concentrations measured in the experiment for each dust under the most acid conditions.

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Understanding the nature of atmospheric acid processing of mineral dusts in supplying bioavailable phosphorus to the oceans

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Affiliations and footnotes are as in the main manuscript file

Supporting Information Appendix

This Document contains supplementary data as referred to in the main manuscript.

SI.1. INTRODUCTION

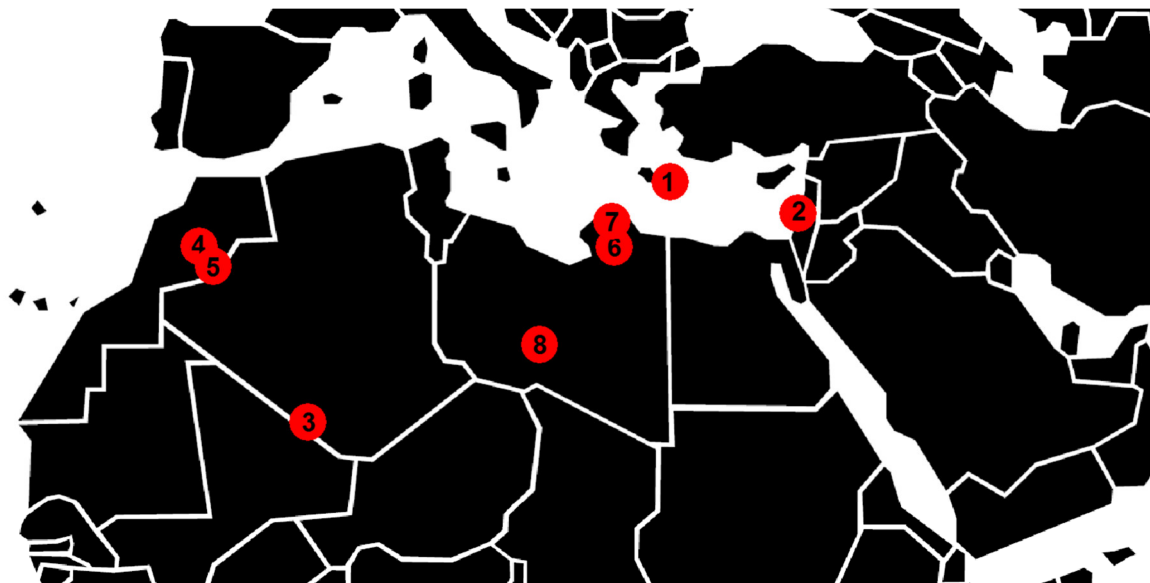


Figure S1.1. Map showing the approximate collection location of the dust/soil samples (see Table 1 in the main manuscript for the precise coordinates). 1, Greece; 2, Israel; 3, Algeria; 4, Morocco (EM); 5, Morocco (JB), 6, Libya (#7); 7, Libya (#9); and 8, Libya (TIB).

SI.2. METHODS

SI.2.1. Experimental procedures

All samples were characterized for their mineralogical composition by X-ray diffraction (XRD; Bruker, D8) for their elemental composition by X-ray fluorescence (XRF; Innovex, X-50) and for their specific surface area using a Micromeritics Gemini VII 2390a instrument after degassing overnight at 75 °C under a flowing nitrogen gas atmosphere.

Due to the limited mass of Israel dust available, some of the supplementary experiments, particularly those involving larger sample masses, were carried out using a sample from Libya (#9). An overview of all supplementary experiments is given in Table S2.1., which includes experimental starting conditions.

Table S2.1. Summary of supplementary experiments and their starting conditions.

Experiment type	Dust(s) used	Starting conditions (T = 18-22°C)
pH tracking (SI.2.1.1.)	Israel (50.1 mg)	pH 4 HCl, 50 mL
Low solution volume to dust mass – high ionic strength (SI.2.1.2.)	Israel (55±3 mg)	0.66 M (NH ₄) ₂ SO ₄ added to pH 1, 2, 3 HCl 2 M NaCl added to pH 3 HCl
P concentration tracking in buffered solution (SI.2.1.1.)	Israel (50.2 mg)	pH 4 potassium phthalate buffer (0.05 M), 200 mL
Sequential solution addition (SI.2.1.3.)	Israel (39.5 mg)	0.7 mL of MQ to wet the dust, then 20 sequential additions of 0.7 mL of pH 1.3 HCl thereafter. Centrifugation and removal of solution was done before each addition.
	Libya (#9) (57.5 mg)	0.5 mL of MQ to wet the dust, then 34 sequential additions of 0.2 mL of 0.001 HCl / 1M MgCl ₂ thereafter. Centrifugation and removal of solution was done before each addition.
	Libya (#9) (51.8 mg)	1.2 mL of pH 2.4 HCl, centrifuged, solution removed, followed by 1.2 mL of 1M NaCl prepared in MQ water
Acid addition to slurry via dialysis (SI.2.1.4.)	Libya (#9) (310±4 mg)	Libya (#9) (49.5 mg)
		1.2 mL of pH 2.4 HCl / 1M NaCl, centrifuged, solution removed, then followed by 1.2 mL of 1M NaCl prepared in MQ water
Low solution volume to mass ratio on dust analogues. Mixture composition mirrors the P and Ca concentration in Israel dust. (SI.2.1.5.)	Mixture of powdered quartz, calcite and hydroxyl-apatite nanopowder (50±2 mg)	24 mL of solution, 9.6×10^{-5} mol of protons added as 1.5 mL of pH 1.2 HCl within a dialysis tube. An additional control experiment added dust directly to a pH 2.4 HCl solution. 1 mL HCl at pH 0, 1, 1.8, 2.4, 3.4

SI.2.1.1. pH and phosphate concentration tracking experiments

In order to assess the rate at which protons were consumed in contact with dust we performed an experiment where pH was tracked at 20 s intervals during reaction in a stirred solution (Israel dust, pH 4 solution, open to the atmosphere). Additionally, we determine the rate of P liberation under buffered pH conditions in a continuously mixed solution (pH 4, 0.05 M potassium phthalate; Sigma-Aldrich, ACS reagent grade). This was achieved by periodic removal and filtration of a small volume of solution at given time intervals.

SI.2.1.2. Experiments testing ionic strength effects

To compare potential differences in high ionic strength (IS) solutions, four experiments were performed. Three solutions were HCl (pH 1, 2 and 3) with ammonium sulphate salts (BDH, Aristar grade) dissolved to give a solution concentration of 0.66 mol L^{-1} (IS = 2, assuming complete dissociation) and one was performed using 2 mol L^{-1} NaCl (Fluka Analytical, TraceSelect) at pH 3 (IS = 2).

SI.2.1.3. Sequential solution addition experiments

To investigate P release over repeated exposure cycles, we performed experiments on two samples where acid addition was repeated sequentially on the same dust sample. In one experiment Israel dust was wetted with 0.7 mL of MQ water in a micro-centrifuge tube. The resulting slurry was placed in an end-over-end shaker for 2 minutes before centrifugation at 13200 rpm for 5 minutes. The supernatant was extracted for analysis after which 0.7 mL of acid (pH 1.3 HCl) was added to the dust, which was then resuspended in the solution. Repeat shaking and centrifugation steps followed, after which the whole cycle was repeated with a further 19 acid additions. Further experiments used Libya dust (#9), 0.5 mL of MQ was used to wet the sample, then 34 sequential additions of 0.2 mL of pH 3 HCl with MgCl_2 (Sigma-Aldrich, ACS reagent grade) to give a concentration of 1M.

Two experiments investigated the effects of high IS, coupled with proton addition. In the first 1.2 mL of pH 2.4 HCl was reacted for ~5 minutes with a known mass of sample (Libya, #9), the slurry was then centrifuged, the solution removed, and this was followed by addition of 1.2 mL of 1M NaCl prepared in MQ water. The second followed the above procedure with the exception that the acid had NaCl added to give a concentration of 1M (IS = 1).

SI.2.1.4. Acid addition to slurry via dialysis

In order to evaluate the potential effects associated with different mineral dissolution rates we performed an experiment where protons were added to the slurry within a dialysis membrane (Spectrum Labs Spectra/Por® Float-A-Lyzer® G2, with a molecular weight cut-off of 100-500 Daltons). This ensures the slurry is not exposed directly to the low pH of the dilute acid, while over the course of the experiment it is still exposed to the same absolute concentration of protons. A control experiment was performed where dust was added directly to an HCl solution that included the same apparatus.

SI.2.1.5. Experiments with dust analogues

We tested the ability to replicate experimental results (from Israel dust) in simple analogue systems, and to evaluate the effect of the absence of competition between calcite and apatite for protons within the system. Powdered quartz (surface area of $2.56 \text{ m}^2 \text{ g}^{-1}$, EU JRC Institute for Reference Materials and Measurements, BCR-172) as an inert carrier for the reactive components, was mixed with either only hydroxyapatite nanopowder (Aldrich, <200 nm particle diameter), or this plus calcium carbonate powder (Alfa Aesar ACS grade). Mixture compositions were based upon total Ca and P concentrations as determined from the Israel dust experiments at the lowest pH. The experimental protocol followed that used for the low solution volume to dust mass ratio experiments.

SI.2.2. Global modelling procedures

SI.2.2.1. Global 3-D atmospheric chemistry transport modelling using TM4-ECPL.

The global chemistry – transport model TM4-ECPL (1), simulates the oxidant chemistry accounting for non – methane volatile organics and all major aerosol components, including secondary inorganic (2) and secondary organic aerosols (3). The model is driven by meteorology from the ECMWF (European Centre for Medium – Range Weather Forecasts) Interim re–analysis project (ERA – Interim) meteorology (4) and it uses the anthropogenic and biomass burning emissions from the Representative Concentration Pathway 6.0 (RCP 6.0) emission scenario (5). Biogenic emissions come from the MEGAN – MACC Biogenic Emission Inventory for the year 2008 (6), while soil NO_x and oceanic emissions are taken from the POET inventory database (7). Oceanic emissions of primary organic aerosol, isoprene, terpenes and sea – salt particles are calculated online driven by meteorology (8). Dust emissions fluxes are obtained from the daily AEROCOM inventories (Aerosol Comparison between Observations and Models; (9)), updated to the simulation year of the model. Further details on anthropogenic and natural emissions in the model are provided in (1) and (10), where the atmospheric cycles of Fe and N in TM4-ECPL have been parameterized and evaluated respectively. The current model configuration has a horizontal resolution of 6° in longitude by 4° in latitude and 34 hybrid layers in the vertical, from surface up to 0.1 hPa. The simulation has been performed with meteorology for the year 2008 and a model time-step of 30 min, while a spin-up time of one year (with 2007 meteorology and emissions) has been used.

Aerosol water pH is calculated by the ISORROPIA II thermodynamic model (2) which solves the K⁺–Ca²⁺–Mg²⁺–NH₄⁺–Na⁺–SO₄²⁻–NO₃⁻–Cl⁻–H₂O aerosol system. For mineral dust and sea-salt elements, the model takes into account the following mean percent mass content of particles: Na⁺: 30.6% on sea-salt and 1.7% on dust, Ca²⁺: 1.2% on sea-salt, K⁺: 2.4% on dust and 1.1% on sea-salt and Mg²⁺: 1.5% on dust and 3.7% on sea-salt, Cl⁻: 55% on sea-salt and SO₄²⁻: 7.7% on sea-salt. As far as it concerns the concentrations and the spatial distribution of Ca²⁺ on dust particles, a global soil mineralogy dataset (11) has been applied on dust emissions fields.

SI.3. RESULTS

SI.3.1. Properties of dusts

SI.3.1.1. Initial experiments to investigate reaction rates

Two experiments investigated the rate at which dust minerals reacted in solution. Israel dust added to a pH 4 solution was rapidly neutralised, with 99% of protons consumed within 20 seconds (pH change from 4.0 to 6.2), and 99.9% consumed within 60 s (pH 7.1). A pH of 9 is reached within 200 s (99.999% of protons consumed) before reaching a plateau at pH 9.2. A further experiment, also at pH 4, monitored the P concentration after dust addition to a potassium phthalate (pH buffer) solution (0.05 M). The dissolution of P was rapid, although slower than the pH rise in the unbuffered experiment. After 2 mins over 40% of the dissolvable P was released, rising to over 80% after 1 hr and a maximum was reached after no more than 18 hrs. We used these results to select a conservative reaction time of 48 hours for the majority of our other experiments, except where short time intervals were required to constrain the overall experiment time.

SI.3.1.2. Phosphorus speciation and additional dust properties

The main mineral composition of the dusts as revealed by XRD on a subset of the samples was quartz > feldspar > calcite > kaolinite, confirmed by XRF showing Si as the dominant element (see also ref (35) from the main manuscript). The measured BET surface areas varied from 4.8 to 33.9 m² g⁻¹, with the lowest values recorded for the PM10 dust precursors from Algeria and Morocco (Table S3.1.).

Table S3.1. Results of SEDEX sequential extraction, BET surface area and maximum Ca and P concentrations from experiments in 1 M HCl (0.5 M for Israel dust). Ap-P is the sum of authigenic/biogenic apatite and detrital apatite phases.

Sample	SEDEX phosphorus ($\mu\text{mol g}^{-1}$)						Total SEDEX
	Sorbed	Iron bound	Authigenic / biogenic apatite	Detrital apatite	Total inorganic	Organic	
<i>Dusts</i>							
Greece	0.94	1.09	5.02	2.63	9.69	4.06	13.75
Israel	1.83	1.57	39.19	17.48	60.06	5.69	65.75
<i>Precursor dusts</i>							
Algeria	0.26	0.52	5.57	4.71	11.07	2.57	13.64
Morocco (EM)	0.12	0.24	3.49	4.22	8.08	1.05	9.13
Morocco (JB)	0.14	0.29	4.83	5.97	11.23	0.98	12.22
Libya (#7)	1.22	0.99	9.32	2.51	14.04	3.98	18.02
Libya (#9)	0.50	0.92	4.07	2.20	7.70	6.30	14.00
Libya (TIB)	0.36	1.65	7.70	4.27	13.98	7.00	20.97

Table S3.1. Cont.

Sample	Acid extracted -		BET surface area (m ² g ⁻¹)
	Phosphate (μmol g ⁻¹)	Calcium (mmol g ⁻¹)	
<i>Dusts</i>			
Greece	10.41	2.51	13.4
Israel	53.91	3.16	24.1
<i>Precursor dusts</i>			
Algeria	10.63	0.56	4.8
Morocco (EM)	9.18	0.71	5.0
Morocco (JB)	12.11	1.04	5.5
Libya (#7)	13.20	3.33	31.6
Libya (#9)	5.77	1.38	33.9
Libya (TIB)	17.12	0.68	21.2

SI.3.1.2.1. Elemental and mineralogical characterisation

XRD results showed that the qualitative mineral composition was:

Israel,	quartz > feldspar > calcite > dolomite > kaolinite
Algeria,	quartz > feldspar > calcite > kaolinite > muscovite
Libya (#9),	quartz > calcite > feldspar > kaolinite
Morocco (TIB),	quartz > calcite > kaolinite > muscovite

Chemical composition determined by XRF is shown in Table S3.2.

Table S3.2. Chemical composition of selected dust samples.

Sample	SiO ₂	TiO ₂	Al ₂ O ₃	Fe ₂ O ₃	MnO	MgO	CaO	K ₂ O	P ₂ O ₅	SO ₃	Total
<i>Dust</i>											
Israel	45.22	0.91	7.97	4.51	0.08	1.54	11.80	1.33	0.28	0.18	73.82
<i>Precursor dusts</i>											
Algeria	61.00	0.95	13.34	5.49	0.09	2.85	4.08	2.04	0.17	0.21	90.23
Libya (#9)	53.92	0.81	11.24	4.93	0.08	2.44	6.91	2.14	0.18	0.05	82.70
Morocco (TIB)	54.21	0.91	14.87	7.96	0.11	2.72	3.28	2.04	0.19	0.05	86.34

SI.3.2. Effect of protons and fluid volume on the dissolution of Ca and P minerals

Results for low volume experiments performed on additional dust samples showed a similar trend to the Israel dust data, where there is an approximately linear increase in P before reaching a maximum value (Figure S3.1.).

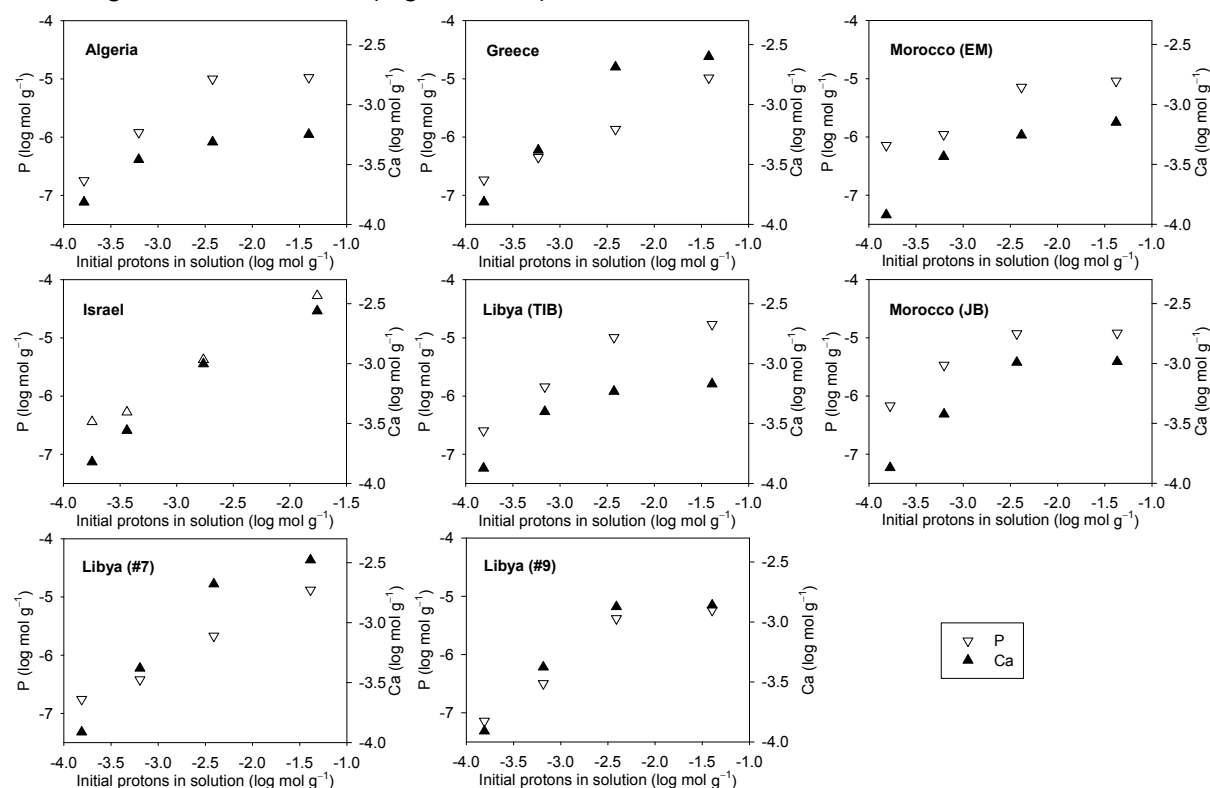


Figure S3.1. Phosphorus and calcium liberated from a range of dusts and soils in relation to the absolute concentration of protons that were available for reaction at the start of the experiment. Values are presented as moles per gram of dust. Experiments were undertaken at a mass to volume ratio of $\sim 25 \text{ g L}^{-1}$.

SI.3.2.1. Endpoint pH values measured in selected experiments

The end state of protons in the low volume experiments was indicated by pH measurements as indicated in Table S3.3. In the majority of experiments protons were completely consumed and pH values in the range 6.8 to 8.0 were reached. Only when the proton concentration exceeded the availability of proton reactive components did the pH remain low. These were the samples where the P released reached a plateau value.

Table S3.3. The pH 48 hrs following initial fluid addition to Israel dust (dust mass $54 \pm 4 \text{ mg}$).

Fluid volume (mL)	Initial pH	Final pH
2	2	7.1
5	1	1.7
5	2	7.6
5	3	8.0
5	5.5	8.0
15	2	6.8

SI.3.2.2. Effect of method of proton addition to dust

We performed an experiment where protons were introduced to the solution/dust slurry via a dialysis membrane. Given the rapid reaction kinetics of acid driven calcite dissolution and the high equilibrium pH of this mineral saturation (pH ~9), it is unlikely that protons diffusing into the slurry will be able to build up to yield low pH values. Figure S3.2. shows the results of this experiment, and a control using the direct addition method. At the end of a 48 hr experiment 74% of the protons had diffused out of the dialysis membrane. The left hand square data point indicates this result, with the right hand square point showing the control.

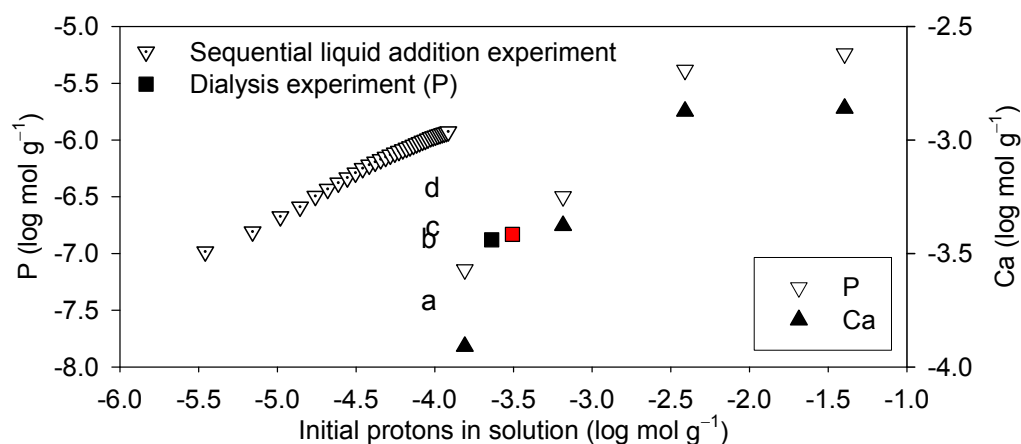


Figure S3.2. Additional experiments conducted on the Libya (#9) dust included with the data from Figure S3.1 (described in Table S2.1). See text for full explanations. Briefly, squares are an experiment where proton addition was via a dialysis membrane, lettered data (a-d) represent the effect of IS, dotted triangles are a sequential solution addition experiment at high IS (discussed in S3.2.3.).

SI.3.2.3. P liberation in solutions of higher ionic strength

Several experiments probed the effect of increasing the ionic strength (IS), in addition to protons added from HCl. Figure 3 in the main manuscript shows results, where in addition to HCl, NaCl or $(\text{NH}_4)_2\text{SO}_4$ salts were added to give an IS of 2 assuming full dissociation (complexation of cations (e.g., Ca^{2+} , NH_4^+) with sulphate and carbonate may reduce the ionic strength below 2, e.g., the IS of the pH 2 $(\text{NH}_4)_2\text{SO}_4$ experiment is 1.34 as calculated by PHREEQC). The presence of NaCl ($I = 2$) increased the liberated P by a factor of 4.3 and the presence of $(\text{NH}_4)_2\text{SO}_4$ increased the liberated P by a factor of 9. The additional P that was liberated may be due to a combination of desorption induced by the increased presence of ions in solution, and changes in solubility of mineral phases in the presence of different solution compositions. For the ammonium sulphate experiments the amount of P released fits on the trend for the HCl-only data if the proton availability from the dissociation of ammonium ions (predicted from a final pH of 8) is considered (data plotted on the x-axis for these experiments includes this predicted proton source).

These effects are observed in experiments on Libya (#9) dust (Figure S3.2.). The sequential experiment used 0.2 mL of pH 3 solution with MgCl_2 to give a 1 M concentration. Results demonstrate that P release is likely to be partly driven by dissolution of the mineral phases into this solution matrix. This matrix effect was further elucidated by data showing that under equal proton concentrations but in the presence of NaCl more P is liberated (S3.2. points 'a' (no NaCl) and 'c' (1 M NaCl)). Further NaCl solution (no HCl) added to both of these dusts

after centrifugation yields further P release (points 'b' and 'd' respectively). These results compare to the SEDEX sorbed phase for this dust of $-6.3 \log \text{ mol g}^{-1}$.

SI.3.2.4. Particle mixing states.

Data in Figure S3.3. describe the two extremes of mixing states in terms of the Ca (as calcite) and P (as apatite) as represented by their concentrations in the Israel dust. The triangles represent the protons required per gram of dust in the absence of calcite. The squares represent a mixed calcite and apatite system and demonstrate that this simple system is analogous to the Israel dust that is shown in the background as grey circles. The data demonstrate that if dust particles were externally mixed then there would be more P released per H^+ amount because more protons are available for direct reaction with apatite.

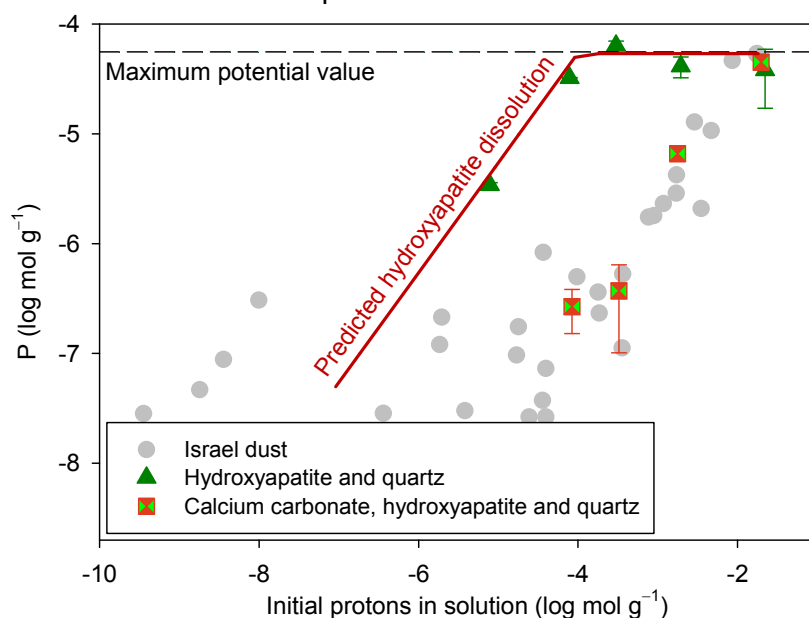


Figure S3.3. Results from analyses of dust proxies. The theoretical dissolution of hydroxyapatite only, in the presence of varying proton concentration is shown as the solid curve. See text for discussion.

SI.3.3. Comparison of experimental results with modelling

Figure S3.4. shows the Figure 5 data (of the main manuscript) separately for the different Israel dust experiments. Modelling of the ammonium sulphate experiments excluded ammonia oxidation reactions. Additionally we modelled (with PHREEQC) the relative effect of different acid types and ionic strength on P and Ca release for a scenario of Israel dust at a mass to volume ratio of 28 g L^{-1} (Figure S3.5.).

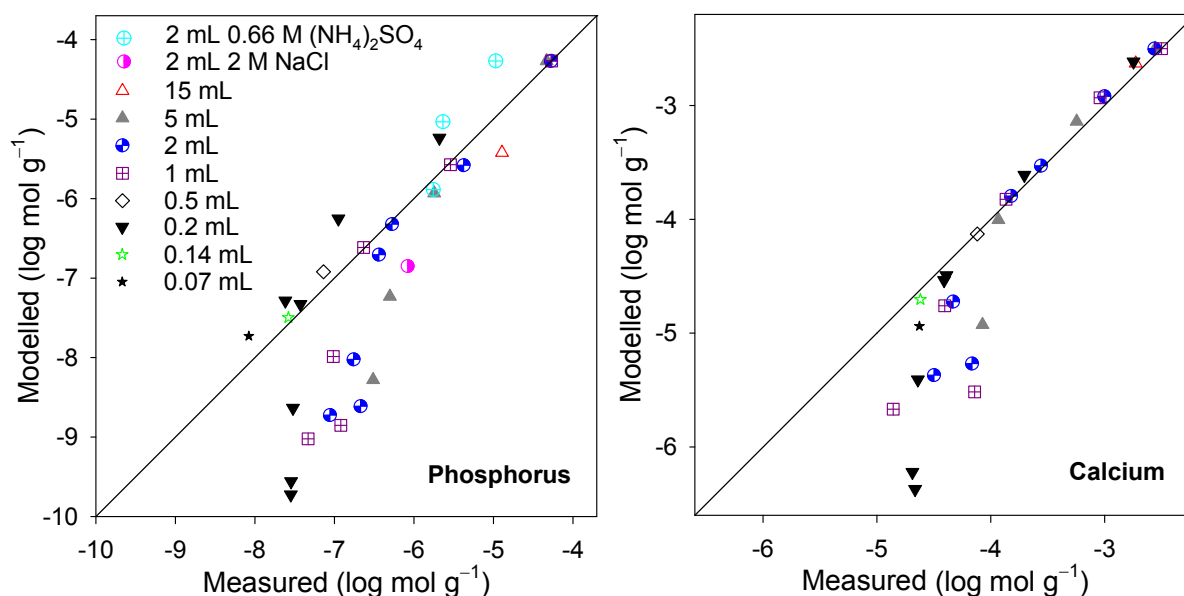


Figure S3.4. Comparison of measured P and Ca dissolution from Israel dust compared to equilibrium predictions made using PHREEQC.

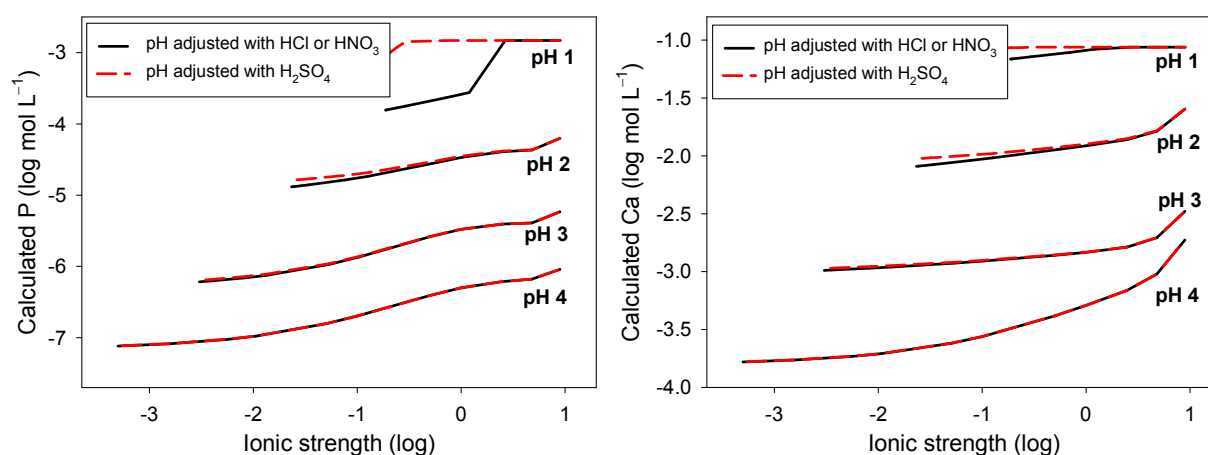


Figure S3.5. Theoretical release of P (left panel) and Ca (right panel) from dust under varying ionic strength and pH for three different acid types.

SI.3.4. Supplementary global modelling data

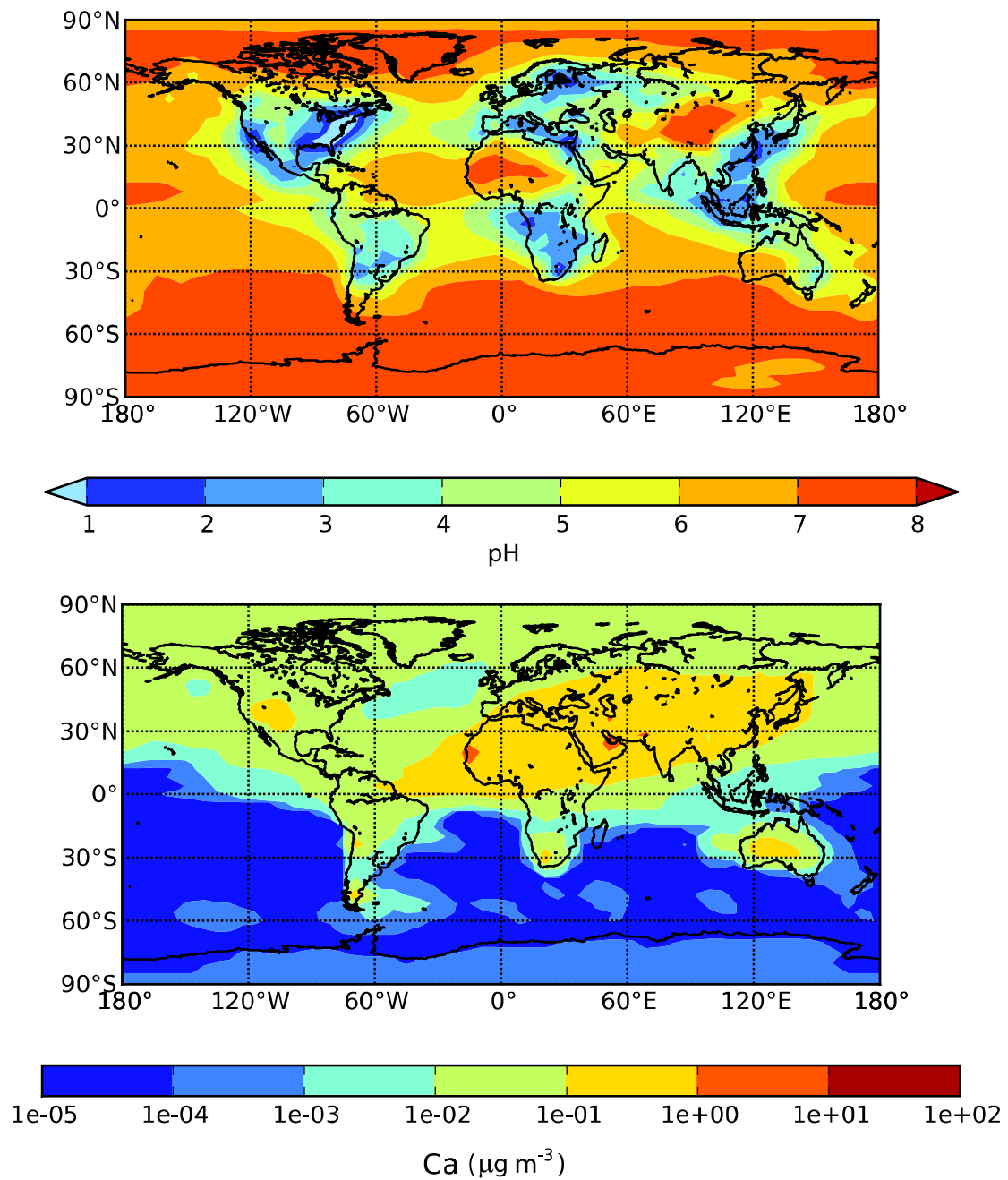


Figure S3.6. Annual average maps of coarse mode aerosol pH (top panel) and Ca concentration (bottom panel). Calculations are carried out with the global model framework of Myriokefalitakis et al. [1] using current day aerosol emissions; results are shown for the model surface layer.

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