INTRODUCTION

Phosphorus (P) is a crucial bioelement for life on Earth, as it is utilized in the formation of ATP/GTP, genetic materials (DNA/RNA), cell membrane phospholipids, and sugar phosphates formed as intermediate products during sugar metabolism (George et al., 2018). Due to the role P plays in the cycle of life, it is critical to understand P uptake and utilization pathways and their interaction with other key nutrients, such as carbon (C) and nitrogen (N). This is especially true in extreme environments, where organisms are often forced to find novel adaptation strategies for survival and C:N:P ratios may be atypical (Huang et al., 2018; Tian, Chen, Zhang, Melillo, & Hall, 2010). These environments also act as a unique analogue for investigating the potential for life on other planets including Mars, where in the planet's early history active hydrology promoted the release of nutrients making P bioavailable (Komatsu & Ori, 2000;...
Weckwerth & Schidlowsky, 1995). Significantly, Komatsu & Ori (2000) and Weckwerth & Schidlowsky (1995) have suggested that water-mineral interaction on early wet Mars may have allowed for up to 45 times higher phosphate release rates than on early Earth. As a result, the bioavailable phosphate concentration in Martian sediments is about twice that estimated for prebiotic Earth (Adcock, Hausrath, & Forster, 2013), with P<sub>2</sub>O<sub>5</sub> of surface sediments varying between 0.15 and 5.19 wt% as determined by Mars Exploration Rovers (Yen et al., 2006). Since early Mars has all the makings of a planet which could have supported past life, it is critically important to identify biogeochemical signatures that can persist through many millennia of hyperarid conditions such as those found on Mars today.

The Atacama Desert is one of the driest and oldest temperate deserts (Dunai, Lopez, & Juez-Larre, 2005; Hartley, Chong, Houston, & Mather, 2005) that reaches the extreme limit of life (Navarro-Gonzalez et al., 2003). This desert occupies around 105,000 km<sup>2</sup> of northern Chile, and has remained extremely dry for the past 150 Myr (Hartley et al., 2005), with annual precipitation at the hyperarid core less than 2 mm (Houston, 2006) for more than 10 Myr (Sun, Bao, Reich, & Hemming, 2018). Alongside aridity, extremes in UV irradiation are similar to high levels of UV delivered to exposed Martian surfaces, approximately 150 kWh/m<sup>2</sup> UV-A (315–400 nm) and 5 kWh/m<sup>2</sup> UV-B (280–315 nm) (Cordero et al., 2018). UV irradiation causes organic degradation, resulting in an oligotrophic terrain, and the lack of precipitation reduces leaching and runoff, meaning phosphate minerals can be preserved. This location is therefore ideal to investigate both the dynamics of P cycling in relation to other key nutrients (C, N) in hostile environments, and to potentially identify long-lived biomarkers of life in an environment similar to that found on Mars.

The biogeochemical cycling of P is less well-studied than C, N, and S in modern terrestrial soils. This is because P usually remains in the form of orthophosphate (PO<sub>4</sub><sup>3-</sup>, HPO<sub>4</sub><sup>2-</sup>, or H<sub>2</sub>PO<sub>4</sub><sup>-</sup>) and only has one stable isotope, <sup>31</sup>P. These facts render it relatively difficult to track P in biosystems and understand its reaction processes. Recently, however, methods have used phosphate-bound stable oxygen isotopes (hereafter noted as δ<sup>18</sup>O<sub>PO4</sub>) as a tracer for the detection of biological activity (Blake, Chang, & Lepland, 2010; Blake, O’Neill, & Surkov, 2005; Jaisi & Blake, 2014; Stout, Joshi, Kana, & Jaisi, 2014; Tamburini, Pfahler, von Sperber, Frossard, & Bernasconi, 2014), since phosphorus is normally associated with oxygen in both abiotic and organic forms.

Stable oxygen isotope values are reported in parts per thousand (‰) using the standard δ notation:

\[
\delta^{18}O (\%o) = \left( \frac{R_{\text{sample}}}{R_{\text{standard}}} - 1 \right) \times 1000
\]

where R<sub>sample</sub> is the 18O/16O ratio in a given sample and R<sub>standard</sub> is the 18O/16O ratio of Vienna Standard Mean Ocean Water (VSMOW).

We further define an isotopic fractionation factor, α, as follows:

\[
\alpha_{p-r} = \frac{R_{\text{product}}}{R_{\text{reactant}}}
\]

where R<sub>product</sub> and R<sub>reactant</sub> are the 18O/16O ratios in the product and reactant, respectively. The isotopic enrichment factor, ε<sub>p-r</sub>, is then defined as follows:

\[
\epsilon_{p-r} (\%o) = (\alpha_{p-r} - 1) \times 1000
\]

The principle of this method for tracing P cycling using the δ<sup>18</sup>O<sub>PO4</sub> signature relies on the relative stability of the P-O bond under abiotic processing. Abiotic P transformations (including phosphate precipitation, UV photolysis and decay of organophosphorus, and solubilization) do not introduce significant oxygen isotope fractionations in phosphate (Figure 1), resulting in an isotopic signature which is theoretically extremely long-lived (Dahms & Boyer, 1973; Jaisi, Blake, & Kukkadapu, 2010; Kolodny, Luz, & Navon, 1983; Tudge, 1960). However, where biological uptake and processing of P occurs the P-O bond can be broken, enabling isotopic exchange of phosphate O with surrounding O sources (normally soil or cellular water). Hence, deviations in δ<sup>18</sup>O<sub>PO4</sub> from the source mineral signature can potentially be utilized as a biosignature for life (Blake, Alt, & Martini, 2001).

There are three main biological pathways that can cause isotopic fractionations in δ<sup>18</sup>O<sub>PO4</sub> (Blake et al., 2001). These biological processes include the following: (a) microbial P uptake, with ε of about −3‰ (Blake et al., 2005; Gross et al., 2015); (b) phosphoester hydrolisis by various types of phosphoenzymes (e.g., alkaline and acid phosphatase, DNase, RNase, 5′-nucleotidase, and phytases), with ε varying between −30 and +20‰ (Liang & Blake, 2006, 2009; von Sperber, Kries, Tamburini, Bernasconi, & Frossard, 2014); and (c) microbial hydrolysis of diphosphate by inorganic pyrophosphatase,
with $\varepsilon$ as defined by Eq. (4) at equilibration with water and soil temperature (Figure 1) (Chang & Blake, 2015; Granger et al., 2018; Pistocchi et al., 2017).

$$E_{\delta^{18}O_{PO_4}} = -0.18T + 26.3 + \delta^{18}O_{H_2O}$$

(4)

where $E_{\delta^{18}O_{PO_4}}$ is the $^{18}O/^{16}O$ ratio of phosphate at equilibration with water catalyzed by inorganic pyrophosphatase, $T$ is temperature in °C, and $\delta^{18}O_{H_2O}$ is the $\delta^{18}O$ of ambient water.

We can therefore assess the extent to which abiotic or biotic processes have impacted P by measuring the $\delta^{18}O_{PO_4}$ signature of soils. To investigate deeper into the P cycle, we extract different forms of P using well-developed chemical extractions (Olsen, 1954) and analyze $\delta^{18}O_{PO_4}$ for individual pools of phosphate. By measuring the $\delta^{18}O_{PO_4}$ of each extract, we can identify the extent to which each pool has been subjected to biological utilization. In this study, we focused on four extracted pools of soil phosphate, operationally defined as (a) resin-extractable P, (b) microbial P, (c) NaOH-extractable Al/Fe oxide-bound P and organic P, and (d) HCl-extractable calcium-bound P (Ca-P).

In normal soil environments, resin-extractable P is turned over on timescales from seconds to minutes, oxide-bound P is recycled on weekly to monthly timescales, and Ca-bound P over years to millennia (Helfenstein et al., 2020; Helfenstein et al., 2018). The least bioavailable P pool is therefore normally Ca-bound P. In an alkaline environment like the Atacama Desert (Crits-Christoph et al., 2013; Makhalanyane et al., 2015; Shen, Zerkle, Stueken, & Claire, 2019) or on Mars (pH 7.2–8.2; (Hecht et al., 2009; Smith et al., 2009)), where resin-extractable P (bioavailable P) is high, it would tend to be sorbed by calcite (Reddy, Wetzel, & Kadlec, 2005) and incorporated through precipitation into Ca-bound P mineral phases (Kreuzeder, 2011), making it less bioavailable (Oburger, Jones, & Wenzel, 2011; Urrutia et al., 2014). However, where this Ca-bound P is a major source of P within a biological system it can be reversibly solubilized by the excretion of acids by specialized microbial communities, breaking down Ca-P and once again making it bioavailable. Alternatively, Ca-bound P can dissolve abiotically over an extremely prolonged period under alkaline conditions (Guidry & Mackenzie, 2003) such as those found in Atacama soils. In non-arid soil samples, the concentration of Ca-P is usually low and hard to detect (Amelung et al., 2018). However, the relative pool size increases significantly in desert environments (Angert, Weiner, Mazeh, & Sternberg, 2012; Helfenstein et al., 2018).

Where exchange is rapid, or P cycling long enough, biogenic P becomes the main component of the Ca-P pool; however, this can take thousands of years (Tamburini et al., 2012). Previous studies have shown that in low precipitation environments, the exchange rate between resin-extractable P and Ca-P is slowed and pedogenic P may persist in the Ca-P pool for hundreds of thousands of years (Helfenstein et al., 2018). In the case of the Atacama Desert where precipitation is extremely scarce (<20 mm/year, an order of magnitude lower than the samples studied by Helfenstein et al., 2018), this turnover time may be expected to be further inhibited, or P cycling completely retarded, unless biological communities can access less conventional forms of water to support nutrient cycling.

For example, organisms under hyperarid conditions are able to utilize hydration water from gypsum minerals to supplement the extremely scarce rain water (Palacio, Azorin, Montserrat-Martí, & Ferrio, 2014).

This research aims to investigate the microbial P utilization in the Atacama Desert and to examine the extent to which biological processing can overprint a mineral-dominated pool, even under the harshest of conditions. Previous studies using $\delta^{18}O_{PO_4}$ in Mars-analogous or arid terrestrial environments all focused on relatively water-rich locations. These studies suggested that $\delta^{18}O_{PO_4}$ values are sensitive to levels of enzymatic activity and microbial metabolism wherever liquid water might be present (Blake et al., 2001; Blake et al., 2005; Jaisi & Blake, 2010). To improve interpretation of $\delta^{18}O_{PO_4}$ analyses, we performed soil metagenomic analyses and cell culture experiments: metagenomic analyses can provide information about the existence and relative abundance of P-processing enzymes and pathways to support isotopic findings, and cell culture experiments can indicate how microbes grow in soils of atypical C:N:P ratios and how they respond to bioavailable P. Therefore, we combine this novel $\delta^{18}O_{PO_4}$ proxy for phosphorus cycling with enzyme pathway analyses and cell culture experiments to trace P dynamics in soils spanning an aridity gradient from hyperarid to arid environments.

2 | MATERIALS AND METHODS

2.1 | Study site and soil characterizations

The Atacama Desert is bounded by two high mountain ranges (the Coastal Cordillera and Altiplano/Andes Mountains) (Garreaud, Molina, & Farias, 2010; Garreaud, Vuille, Compagnucci, & Marengo, 2009; Houston & Hartley, 2003; Rech et al., 2010; Veblen, Young, & Orme, 2015), which prevent Pacific and Atlantic Ocean moisture from reaching the hyperarid core. Modern hyperaridity of the Atacama Desert is also a result of subtropical atmospheric subsidence (Takahashi & Battisti, 2007), coupled with a temperature inversion caused by the interaction of the constant strong Pacific anticyclone (Trewarth, 1961) and coastal cold upwelling of the Humboldt ocean current (McKay et al., 2003). For this study, five sites were sampled on a latitudinal gradient from 22°S to 28°S (Figure 2). This sampling strategy was selected to provide an aridity gradient from hyperarid in the north to arid in the south (Figure 3); the sites are María Elena South (MES, 1,493 m altitude) (Azua-Bustos, Caro-Lara, & Vicuna, 2015), Point of No Return Dos (PONR-2, 1,493 m altitude), Yungay (1,007 m altitude), Transition Zone 4 (TZ-4, 1,658 m altitude), and Transition Zone 5 (TZ-5, 588 m altitude), with 0–23 mm average annual precipitation estimated by Shen et al. (2019) shown in Table 1. We note that visual small plant vegetation appeared in TZ-4 and TZ-5, and dried grass roots appeared in their soil samples (Figure 3b). These locations had substantially less visible plant life when visited in 2012, suggesting that the strong rainfall events of 2015–2017 (Azua-Bustos et al., 2018; Uritskiy et al., 2019) might have stimulated a rare “desert bloom” at these locations prior to our sampling (Shen et al., 2019).
Briefly, MES, PONR-2, and TZ-4 are located on mixed sedimentary–volcanic terrane, while Yungay and TZ-5 are located on plutonic terrane (Figure 2). In detail, MES soils are fed by Quaternary and Jurassic sedimentary rocks, Mesozoic volcanics, and Mesozoic–Cenozoic plutonic rocks; PONR-2 soils form on Quaternary sedimentary and Mesozoic volcanic rocks; Yungay soils form from Paleozoic–Mesozoic and Mesozoic–Cenozoic plutons, and Quaternary sedimentary rocks; TZ-4 soils form on a mixture of Cretaceous–Tertiary volcanics, Quaternary sediments, and Paleozoic–Mesozoic plutonic rocks; and TZ-5 soils are affected by

**TABLE 1** Altitude, mean annual temperatures, average precipitation, grain size, and pH from Shen et al. (2019) and C:N:P ratios (this study) of our five study sites. MES, PONR-2, and Yungay are from the hyperarid core; TZ-4 and TZ-5 are from the arid region.

| Site  | Altitude (meter above sea level/m a.s.l.) | Temperature °C | Precipitation (mm/year) | Median grain size (μm) | pH | C:N:P     |
|-------|------------------------------------------|----------------|--------------------------|------------------------|----|-----------|
| MES   | 1493                                     | 21             | 0.7                      | 444                    | 8.38 | 128:1:7   |
| PONR-2| 1493                                     | 17             | 1.1                      | 486                    | 8.09 | 323:1:4   |
| Yungay| 1007                                     | 17             | 2.0                      | 464                    | 8.11 | 610:1:4   |
| TZ-4  | 1658                                     | 17             | 15.1                     | 291                    | 8.26 | 15:1:1    |
| TZ-5  | 588                                      | 18             | 22.7                     | 322                    | 9.12 | 216:1:8   |

**FIGURE 2** Geologic units (Kirkham et al. 1995) of five study sites (MES, María Elena South; PONR-2, Point of No Return Dos; TZ, Transition Zone)

**FIGURE 3** Landscapes of (a) hyperarid (MES) and (b) arid (TZ-4) sites of the Atacama Desert
Quaternary and Cretaceous sedimentary rocks, and Paleozoic–Cenozoic plutonic rocks (Schenk et al. 1999; Tapia et al., 2018). Detailed sampling methods are described in Shen et al. (2019). In short, 3 pits were dug to 10–20 cm depth at each site, where ~ 200 g of material was removed for geochemical analyses. From each study site, one of the pits was further sampled for $\delta^{18}$O analyses (~500 g of sediment), DNA extraction (~50 g), and cell spreading experiments (~100 g) using sterile techniques. Soils of an additional TZ-5 pit that was surrounded by dead dried shrub, similar to all TZ-4 pits, were sampled for $\delta^{13}$C analyses. Samples were stored at 4°C to avoid any change in the microbial community structure and minimize microbial activity.

Mean annual precipitation and mean temperature (Table 1) for each sampling site were determined from the nearest meteorological stations between the years of 1951 and 2017, from Explorador Climático, Center for Climate and Resilience Research (available at: http://explorador.cr2.cl/). General soil properties (pH, electrical conductivity, total organic carbon (TOC), total organic nitrogen (TON), nitrate, grain sizes, and major elements in the regolith such as P$_2$O$_5$, SiO$_2$, Al$_2$O$_3$, MgO, and Fe$_2$O$_3$) were reported in Shen et al. (2019). Regolith P is referred to total P (TP). Carbonate from each sampling pit was determined in triplicate using the decarbonation method during the preparation step for TOC and TON measurements.

2.2 Sequential phosphate extraction

Sequential extractions were undertaken on each soil sample to isolate different P pools of potential interest for $\delta^{18}$O analyses and P concentrations (Hedley, Stewart, & Chauhan, 1982). Each bulk extraction started with large volumes of soil (-100 g) in an attempt to ensure enough P was obtained for isotope purification and analysis, following Tamburini et al. (2010). Alongside this bulk method, we extracted a subsample of the soil using the same soil: solution ratios for concentration analysis. The concentrations of phosphate from each of the four extractable pools were quantified on an Aquahem 250 analyzer (Thermo Fisher Scientific) using a molybdenum blue reaction (Murphy & Riley, 1962).

The stages of extraction followed Tamburini et al. (2018); briefly these were:

- Resin-extractable P (resin-P), extracted from approximately 600 g of soil in 3.6 L of 18.2 μM ultrapure water with 3 pre-conditioned 125 mm x 125 mm anion-exchange resin strips (VWR International Ltd). It should be noted that as these soils were extremely dry, there was a possibility that a water-based extraction would promote some rapid microbial response and release of P captured in this extraction. This is an unavoidable laboratory effect, but was mitigated as best possible by shaking at 4°C to limit microbial activity within the extract (Tamburini, Pistocchi, Helfenstein, & Frossard, 2018).
- Microbial P, extracted from 100 g of fresh soil with 12.5 ml of hexanol in 1 L of 18.2 μM ultrapure water with one 125 mm x 125 mm pre-conditioned anion-exchange resin strip. This extraction captured the mixture of inorganic (resin-P as above) and microbial phosphate (Granger et al., 2018; Kuno, Tuchiya, & Ando, 1995; Mclaughlin, Alston, & Martin, 1986). Microbial P was calculated by subtracting resin-P from hexanol-extractable P.
- Oxide-bound and organic P pools were extracted using a NaOH/ethylenediaminetetraacetic acid (EDTA) mix (6.5 mg/ml NaOH and 12.0 mg/ml EDTA), added to hexanol extracted samples.
- HCl-extractable P (Ca-P), 50 g of dried soil was added to 100 ml of 1 M HCl (Tamburini, Bernasconi, Angert, Weiner, & Frossard, 2010).

2.3 $\delta^{18}$OPO$_4$ isotope analyses

As we aimed to identify a long-lived biosignature of life in the Atacama sediments, we focused our $\delta^{18}$O analyses on the most stable P pool, Ca-P. Phosphate in these extracts was converted to Ag$_3$PO$_4$ using a slightly adapted purification protocol described by Tamburini et al (2010). Briefly, phosphate was precipitated as ammonium phospho-molybdate (APM) at a pH of 1 (ensured with the addition of 1 ml of concentrated sulfuric acid) and then dissolved in NH$_4$-citrate solution and re-precipitated as magnesium ammonium phosphate (MAP). The MAP precipitate was further dissolved in HNO$_3$ and shaken with cation resin (AG 50 8X). The phosphate in solution was converted to Ag$_3$PO$_4$ by adding 7 ml of Ag-ammine solution, and the precipitate was oven-dried at 50°C. This method does not use the DAX-8 resin step of Tamburini et al (2010) as this is not necessary unless organic contamination is present in the subsequent Ag$_3$PO$_4$ (Granger et al., 2018).

Analysis of $\delta^{18}$OPO$_4$ was undertaken by weighing approximately 300 μg of Ag$_3$PO$_4$ into a silver capsule, this sample was then converted to carbon monoxide (CO) using a thermal conversion elemental analyzer (TC-EA, Thermo Finnigan) at 1,400°C in the presence of glassy carbon chips. The product CO mixed with a helium carrier gas and was analyzed on a Delta + XL mass spectrometer (Thermo Finnigan) at the British Geological Survey (UK). The $\delta^{18}$OPO$_4$ values were calculated by comparison to a internal Ag$_3$PO$_4$ standard, ALFA-1 (ALFA-1 = $\delta^{18}$O VSMOW value of ± 14.2‰). In the absence of an internal Ag$_3$PO$_4$ reference material, the value for ALFA-1 is derived by comparison to the Ag$_3$PO$_4$ standard “B2207” (Elemental Microanalysis Ltd., England), which has been measured in an inter-laboratory comparison study to have a $\delta^{18}$O value of ± 21.7‰ versus VSMOW. Samples were run in triplicate, with a typical precision σ ≤ 0.4‰. Sample purity was assessed by determining the CO yield compared with the yield of Ag$_3$PO$_4$ standards and rejecting samples where this differed by 10%.

2.4 Metagenomic extraction, sequencing, and enzyme/pathway predictions

To understand the microbial pathways in P dynamics, we performed soil metagenomic analyses. All implementations for gene experiments were either filter-sterilized, autoclaved, or UV-irradiated.
to prevent any external contamination; water was molecular biology grade and nuclease-free. DNA of MES, PONR-2, Yungay, TZ-4, and TZ-5 was extracted with one negative control using the FastDNA® SPIN Kit for Soil (MP Biomedicals) following a modified manufacturer’s protocol: During the cell lysis step, mixtures were incubated at room temperature for 1 hr. DNA extracts were amplified for barcoded Illumina 16s metagenomic sequencing using KAPA HiFi HotStart ReadyMix (KAPA Biosystems) and 16s rRNA primer pair 341F, 5’-CCTACGGGNGGCWGCAG-3’, and 785R, 5’-GACTACHVGGGTATCTAATCC-3’ (Herlemann et al., 2011; Klindworth et al., 2013). Amplicon concentration was quantified using a Qubit™ 3.0 Fluorometer (Invitrogen). The index PCR thermocycling was conducted using Nextera index kit (Illumina®), and only samples with sufficient yields, namely MES, TZ-4, and TZ-5, were passed for further preparation of sequencing. 16S rRNA gene amplicons for Illumina MiSeq System were prepared and sequenced using paired-end 300 bp reads with v3 Chemistry by following the 16s Metagenomic Sequencing Library Preparation protocol (Illumina®) alongside a 20% PhiX control, slightly modified by extending the amplicon PCR from 25 to 29 cycles.

Bacterial sequences were analyzed by the open-source program Quantitative Insights into Microbial Ecology 2 (QIIME 2) (Bolyen et al., 2019). Illumina-sequenced paired reads were combined and demultiplexed into Casava 1.8 paired-end format. Reads of low quality and chimeric sequences were removed with DADA2 (Callahan et al., 2016). Reads were filtered by trimming the first 13 base pairs and of a length less than 300 base pairs. Total sequences were clustered into operational taxonomic units (OTUs) at 99% identity and filtered through chimera checking using VSEARCH (Rognes, Flouri, Nichols, Quince, & Mahé, 2016) and UCHIME (Edgar, Haas, Clemente, Quince, & Knight, 2011). The representative taxonomic identities were aligned at 99% full-length sequence similarity using pre-trained SILVA 132 16s rRNA marker gene reference database (Quast et al., 2013).

Potential functions of these 16S sequences were predicted with Phylogenetic Investigation of Communities by Reconstruction of Unobserved States 2 (PICRUSt2) (Douglas et al., 2019). Briefly, reads were assigned into phylogenetic units with HMMER (Eddy & Wheeler, 2007; Johnson, Eddy, & Wheeler, 2010), EPA-ng (Barbera et al., 2019), and GAPPA (Czech & Stamatakis, 2019). The hidden-state prediction was performed with the R package caster (Louca & Doebeli, 2018). Finally, microbial enzymes and metabolic pathways were identified using the Minimal set of Pathways (MinPath) approach (Ye & Doak, 2009) referencing the MetaCyc pathway database (Caspi et al., 2016; Caspi et al., 2018).

Here, we focused only on data pertaining to biogeochemical P dynamics in the sampled soils, as it was beyond the scope of this study to provide a full metagenomic dataset. Enzymes and metabolic pathways related to microbial phosphate dynamics were selected from all records of predicted pathways. The abundance of phosphate-involved pathway copy numbers was plotted with OriginPro 2019 (OriginLab Corporation).

2.5 Phosphate amendments and cell culture experiments

To explore the microbial growth in these extremely dry soils of atypical C:N:P ratios, we performed cell culture experiments and phosphate amendments. About 10 g of soil from each sampling site was amended with 4.5 ml sterilized 10% monosodium phosphate (NaH2PO4) and 4.5 ml sterilized ultrapure water (as a control) and left for 4 days at room temperature (Shen et al., 2019). Soils were then refrigerated at 4°C. Duplicate amended soils and original soils without amendment were suspended in sterilized ultrapure water by an applicable dilution factor and spread on Luria–Bertani (LB) agar and plate count agar plates, and left at 21°C. Colonies were counted after 20 days of growth (Bagaley, 2006). Colony-forming units (CFUs) were calculated by multiplying the number of colonies formed on agar plates by the correspondent dilution factor, and a factor 1.45 to account for the addition of 4.5 ml solution to 10 g of soil. CFUs on phosphate amendments were further transformed into logarithmic scale, and normalized as the difference relative to water only amendments.

3 RESULTS

3.1 Nutrients, mafic elements, and δ18O PO4

We saw no correlation between TOC and TP at the hyperarid core, and the TOC and TP of arid sites were both higher than hyperarid sites (Figure 5a). The P/N ratio anticorrelated with TOC, with 4 of the 5 sites showing a distinctive relationship. However, site TZ-5 (the wettest site) displayed a different decreasing trend from the other four sites (Figure 5b). P/N significantly correlated ($R^2 = 0.88***$ and $p = .000$) with SiO2/Al2O3 (used to approximate quartz to clay ratios (Broadhurst & Loring, 1970; Olivares et al. 2017; Xiao, Porter, An, Kumai, & Yoshikawa, 1995)) (Figure 5c), which was generally taken as a proxy of sedimentation rates. Additionally, sites with lower levels of total P commonly had larger grain sizes (Table 1).

Mg and Fe were two common mafic elements, the richness of which indicated the abundance of igneous rock sources (Barker, 1978). MgO in the sediments of sampling sites ranged from 0.9 to 1.4% in hyperarid sites, and was higher in arid sites, between 2.2 and 3.3% (Table 2). Fe2O3 (%) had a linear in arid sites, between 5.2 and 6.6% (Table 2), Fe2O3 (%) had a linear relationship with MgO (Figure 5d).

3.2 Distribution of different P pools

Compared to the hyperarid sites (MES, PONR-2, and Yungay, <2 mm/ year), arid sites (TZ-4 and TZ-5, 15–20 mm/year) had a greater pool size of microbial P and NaOH-P (including Al/Fe oxide-bound P and organic P), but a relatively smaller pool size of resin-P and HCl-P (usually calcium-bound P, hereafter noted as Ca-P) (Figure 4). Microbial P
positively correlated with NaOH-P ($R^2 = 0.88^*, p = .022$). In addition, sites with smaller microbial P and NaOH-P pools generally had larger grain sizes (Table 1). Plots of microbial P and total P clustered together within hyperarid and arid sites (Figure 6a). The $\delta^{18}$O values of soil Ca-P from the hyperarid sites (MES, PONR-2, and Yungay) varied between 7.7 and 13.8‰, and from arid sites (TZ-4 and TZ-5) between 19.5 and 25.3‰ (Table 2). The contents of TOC, TON, and microbial P correlated with the $\delta^{18}$O of Ca-P (Figure 6b–d).

### 3.3 | Enzymes and pathways in microbial phosphate metabolisms

The overall structure of the relative abundance of phosphate pathways in MES, TZ-4, and TZ-5 was very similar (Figure 7). The least abundant pathways were sucrose degradation IV, glucose and glucose-1-phosphate degradation, Entner–Doudoroff pathway III (sugar degradation), UDP-2,3-diacetamido-2,3-dideoxy-α-D-mannuronate biosynthesis, CMP-legionaminate biosynthesis I (sugar-nucleotide synthesis), pyrimidine deoxyribonucleotides de novo biosynthesis IV, pyrimidine deoxyribonucleotides biosynthesis from CTP (nucleotide synthesis), and phospholipases. Comparatively, the microbiome of MES was more abundant in GDP-mannose biosynthesis, GDP-mannose-derived O-antigen building blocks biosynthesis.
(sugar-nucleotide synthesis), pyrimidine deoxyribonucleotides de novo biosynthesis III (nucleotide synthesis), ketogluconate metabolism (sugar acid degradation), pyridoxal 5'-phosphate biosynthesis and salvage (vitamin synthesis), and tRNA processing. The TZ-4 microbiome was abundant in all pathways but ketogluconate metabolism (sugar acid degradation), pyridoxal 5'-phosphate biosynthesis and salvage, thiamin diphosphate biosynthesis II (vitamin synthesis), S-methyl-5-thio-α-D-ribose 1-phosphate degradation (nucleotide degradation), pyrimidine deoxyribonucleotides de novo biosynthesis II, and purine nucleotides de novo biosynthesis II (nucleotide synthesis). The TZ-5 microbiome was abundant in all pathways but tRNA processing, GDP-mannose biosynthesis (sugar-nucleotide synthesis), GDP-mannose-derived O-antigen building blocks biosynthesis, and pyrimidine deoxyribonucleotides de novo biosynthesis III (nucleotide synthesis).

The copy number of inorganic pyrophosphatase genes increased from ~50,000 in MES to ~70,000 in the two arid sites (TZ-4 and TZ-5). Alkaline phosphatase genes increased from ~14,000 in MES to ~30,000 in arid sites. Both DNase and RNase genes increased from ~160,000 in MES to ~250,000 in arid sites. The abundance of acid phosphatase and 5'-nucleotidase genes were ~4,000 and ~40,000 in these three sites, respectively. Phytase genes increased from ~300 in MES to ~2,000 in arid sites (Table 3).

### 3.4 Microbial growth with phosphate amendments

Cultivable heterotrophic micro-organisms from the hyperarid sites and the drier arid site TZ-4 universally did not prefer growing on excessive bioavailable inorganic phosphate. However, the cultivable microbial communities of site TZ-5 did seem to display a preference for growth on phosphate amended plates (Tables 4 and 5).
DISCUSSION

4.1 Phosphorus distribution and microbial P cycling in Atacama sediments

In hyperarid environments such as the core of the Atacama Desert, P loss induced by leaching is negligible; hence, phosphate can be preserved to high levels in sediments. Release of inorganic P in desert minerals can be stimulated via chemical weathering (Sheldon, 1982) and by various microbial solubilization mechanisms, usually including acidification by 2-ketogluconic acid and chelation of the cations bound to P (Ameen et al., 2019; Osorio & Habte, 2013; Scervino et al., 2010; Sharma, Sayyed, Trivedi, & Gobi, 2013). During long-term aridity, phosphate can be released by microbial death and plant decay, for example, after an episodic rainfall-led desert bloom event (Fabre, Gauquelin, Vilasante, Ortega, & Puig, 2006; Orlando, Alfaro, Bravo, Guevara, & Caru, 2010; Vidiella, Armesto, & Gutierrez, 1999). Although Ca-P can dissolve abiotically, this process is negligible when pH is more than 8 (Guidry & Mackenzie, 2003) (Table 1).

At these high levels of phosphorus (with C:P of 67:1 on average) in spite of a large portion of Ca-P, micro-organisms in Atacama soils might be less phosphorus-limited than in normal terrestrial soils, where C:P is typically 186:1 on average (Cleveland & Liptzin, 2007; Reed & Wood, 2016). We also see that the addition of inorganic P to cell culture experiments makes no positive impact on the colony-forming units (CFU) for all of the sites with rainfall below ~15 mm/year (MES, PONR-2, Yungay, TZ-4; Table 5). However at TZ-5, where annual precipitation levels can be 20 mm or more, addition of phosphate to the agar plates enhances growth. Therefore, it appears that P can once again become a limiting nutrient at higher precipitation levels, even though the C:N:P ratio (258:1:5) still suggests a remarkably greater P reserve than N compared to normal terrestrial soils (186:13:1) (Cleveland & Liptzin, 2007; Reed & Wood, 2016). When amended with excess inorganic phosphate, growth of microbial colonies from TZ-5 increases significantly, by about 2 orders of magnitude (Table 5). However, the phosphate salts used in this amendment are directly available for biochemical use, quite different than the major pool of P at TZ-5.
FIGURE 7  Predicted pathways related to biological phosphate metabolisms. Degradation pathways are shown in boxes.
which is in the form of Ca-P (Figure 4). This trend indicates that biological activity at the least arid site is limited by resin-P even if P reserves in the form of Ca-P are present.

Various P sources can be accessed by microorganisms via immobilization, mineralization, and biological solubilization (Gyaneshwar, Kumar, & Parekh, 1998; Illmer & Schinner, 1992; Nahas, 1996) (Figure 1). It appears that these processes must occur to some extent for our sites, as at higher TP levels, we observe that microbial P pool size is also greater (Figure 6a). This trend is also consistent with the transition from hyperarid to arid sites recorded by the relationship between TOC and TP (Figure 5a). These correlations indicate that when rainfall exceeds ~10 mm/year, some TP can be desorbed by biological solubilization and made bioavailable by native microorganisms in situ. Accordingly, the microbial communities could be utilizing mineral P at TZ-5, but at a much slower pace than the resin-P which was added in the cell culture experiments, hence the hundred-fold increase in colony development (Table 5). This reasoning highlights the possibility for resin-P limitation even where mineral P may be more plentiful than other key nutrients such as nitrogen, and at a site where mineral P breakdown appears to be dominated by biological solubilization.

In all other sites, excessive phosphate amendment displays a detrimental effect on microbial growth (Table 5), contrary to the general eutrophication response seen in most natural environments (Golterman, 1973; Lucassen, Smolders, Lamers, & Roelofs, 2005) (Table 5). Usually, the species that actively react to phosphate eutrophication are members of aquatic phytoplankton (Chislock, Doster, Zitomer, & Wilson, 2013; Schindler & Vallentyne, 2008; Vangeel, Mur, Ralskajasiewiczowa, & Goslar, 1994), which are different from extremophiles in desert ecosystems. The unusually high soil P/N ratios are a result of the low N availability (Shen et al., 2019) as limited by water discharge and weathering (McIsaac, David, Gertner, & Goolsby, 2001; Schindler & Vallentyne, 2008; Smolders, Lucassen, Bobbink, Roelofs, & Lamers, 2010; Vitousek, Porder, Houlton, & Chadwick, 2010; Walker & Syers, 1976). When P/N ratios are more than 1 (Table 1), TOC is negatively associated with P/N ratios (Figure 5b). This result suggests that high ambient P/N ratios can be unfavorable to microbial communities in an extremely dry environment. In effect, N sources can conversely promote microbial utilization of P, and nitrate is undoubtedly the major N source in the Atacama Desert (Michalski, Bohlke, & Thiemens, 2004). Only sufficient nitrate can enhance the availability of phosphate by oxidizing ferrous iron, which inhibits iron–phosphate interaction (Smolders et al., 2010). Thus at low N/P (namely high P/N) ratios, the nitrate-driven promotion of microbial P utilization is reduced, which is further unfavorable to microbial communities.

Along the hyperarid to arid gradient, the proportion of resin-P from Atacama soils decreases (Figure 4) as it is presumably immobilized by the more abundant microbial communities in arid sites (Figure 1), although the concentrations of resin-P from Atacama soils are approaching the detection limit for our method (Crain, McLaren, Brunner, & Darrouzet-Nardi, 2018; Lester, Satomi, & Ponce, 2007). Similarly, the relative sizes of microbial P and NaOH-P pools increase in arid sites (Figure 4). Besides the impact of rainfall on microbial P and NaOH-P (as containing organic P) pools, sedimentation potentially plays an important role in diluting these P pools, since the higher microbial P and NaOH-P levels are associated with smaller grain size (Table 1) and higher SiO<sub>2</sub>/Al<sub>2</sub>O<sub>3</sub> (quartz to clay ratios) (Figure 5c) in our study sites. Usually, larger grain sizes cause a decline in the sorption rates of phosphate in organic form (Meng, Yao, & Yu, 2014). Similarly, we recently postulated that bulk N concentrations, largely supplied as nitrate by atmospheric deposition (Ewing et al., 2007; Michalski et al., 2004), could be diluted by higher sedimentation rates and thus more limited to microorganisms (Shen et al., 2019). High sedimentation rates thus promote higher P/N ratios and a small relative pool size of microbial P and NaOH-P, and thus are hostile to these extremophilic micro-organisms.

### TABLE 3 Numbers of predicted genes for enzymes related to P dynamics

| Enzyme abundance       | MES   | TZ-4  | TZ-5  |
|------------------------|-------|-------|-------|
| Inorganic pyrophatase  | 45955 | 76683 | 67800 |
| Alkaline phosphatase   | 14003 | 34644 | 29382 |
| Acid phosphatase       | 3371  | 4440  | 2734  |
| DNase                  | 156246| 287174| 248682|
| RNase                  | 172222| 225415| 232564|
| 5’-nucleotidase        | 33595 | 51364 | 40615 |
| Phytase                | 305   | 3105  | 1343  |

### TABLE 4 Colony-forming units (CFUs) on LB agar and plate count agar plates without amendments, amended with water only, and amended with 10% monosodium phosphate

| Type of culture plate | Amendment | MES   | PONR-2          | Yungay | TZ-4   | TZ-5   |
|-----------------------|-----------|-------|-----------------|--------|--------|--------|
| LB agar               | None      | 17    | 28              | 33     | 2.39 x 10<sup>5</sup> | 2.95 x 10<sup>5</sup> |
|                       | Water     | 7     | 7.98 x 10<sup>3</sup> | 3.71 x 10<sup>5</sup> | 1.23 x 10<sup>5</sup> | 1.17 x 10<sup>3</sup> |
|                       | Phosphate | 0     | 0               | 904    | 4.92 x 10<sup>3</sup> | 3.68 x 10<sup>3</sup> |
| Plate count agar      | None      | 167   | 190             | 5.17 x 10<sup>7</sup> | 2.90 x 10<sup>4</sup> | 6.00 x 10<sup>2</sup> |
|                       | Water     | 1.36 x 10<sup>3</sup> | 8.56 x 10<sup>4</sup> | 6.96 x 10<sup>4</sup> | 1.33 x 10<sup>7</sup> | 1.25 x 10<sup>5</sup> |
|                       | Phosphate | 58    | 75              | 4.84 x 10<sup>5</sup> | 6.38 x 10<sup>6</sup> | 5.80 x 10<sup>6</sup> |
**TABLE 5** The relative microbial growth rate on agar plates with excessive phosphate amendments, illustrating the change in order of magnitude of CFUs with phosphate amendments (all variables are scaled by logarithmic transformation relative to water only amendments). Bold values are those with the absolute change value in the order of magnitude greater than 1

| Site     | LB agar plates | Plate count agar plates |
|----------|----------------|-------------------------|
| MES      | -0.86          | -1.37                   |
| PONR-2   | -3.90          | -3.06                   |
| Yungay   | -0.61          | -1.16                   |
| TZ-4     | -0.40          | -0.32                   |
| TZ-5     | 2.50           | 1.67                    |

**4.2 Phosphoenzymes and phosphate pathways**

Several phosphoenzymes can drive exchange reactions between the stable oxygen isotopes of phosphate and surrounding water. The most studied phosphoenzymes are inorganic pyrophosphatase, alkaline and acid phosphatases, DNase, RNase, 5'-nucleotidase, and phytases, which are all found to be primary phosphoenzymes in Atacama microbiomes (Table 3). These enzymes are broadly involved in P compound degradation pathways.

Pathway prediction analyses show that the majority of phosphate pathways are biosynthesis of microbial P, including pentose phosphate pathways, glycolysis, biosynthesis of sugar-nucleotides, purine and pyrimidine nucleotides, phospholipids, cell walls, vitamins, secondary phosphatic metabolites, and polysaccharides. A relatively small proportion of pathways (13 out of 74) were identified to be degradation pathways or hydrolysis of P compounds (Figure 7), which might be attributed to the dormancy of indigenous microbial communities that were only occasionally activated by moisture elevations (Schulze-Makuch et al., 2018). The most abundant degradation pathways (6 of 13) participate in glycolysis and nucleotide degradation. The less abundant degradation pathways are subclasses of sugar degradation, phospholipid degradation, and methylphosphonate degradation (Figure 7).

Among these pathways, inorganic pyrophosphatase plays a role in methylphosphonate degradation and any hydrolysis of pyrophosphate into two phosphate molecules; it is also a highly conserved phosphoenzyme that ubiquitously exists across all organisms (Knowles, 1980) (Figure 1). DNase and RNase break the phosphodiester bonds of corresponding nucleic acids, and alkaline phosphatase is responsible in completing DNA and RNA degradation by hydrolyzing nucleotides into nucleosides and single phosphate molecules (Liang & Blake, 2006, 2009) (Figure 1). Alkaline phosphatase is also involved in glycolysis II (from fructose 6-phosphate) (Cho, Seo, Kim, Jung, & Park, 2012). Pathways associated with 5'-nucleotidase and acid phosphatase are found to be involved in nucleotide degradation (Passariello et al., 2006). Pathways of phytases, such as phytate degradation I, were not identified in our sequence data, and the abundance of phytases is significantly lower than other phosphoenzymes. These data suggest that phytases are not the dominant drivers of stable oxygen isotope exchange between soil phosphate and the surrounding water.

**4.3 Phosphate δ¹⁸O**

The dominant pool of P was Ca-bound (HCl-extractable) phosphate at all of the sites sampled (Figure 4). This pool either represents (a) the primary mineral P preserved in bedrock, or (b) P formed during early diagenesis as a result of the high pH of the soils, which promotes conversion of resin-P to Ca-P minerals (Oburger et al., 2011; Urrutia et al., 2014). To identify any biosignatures in soil δ¹⁸OPO₄, it is important to first differentiate the stable oxygen isotope fingerprints of biogenic phosphoenzyme-driven water-phosphate equilibration from lithogenic P (Blake et al., 2005; Gross & Angert, 2015) (Figure 1).

The primary bedrock of our sampling sites is generally composed of mixed mafic igneous and sedimentary rocks (Figure 5d): Our five study sites are all largely characterized by Mesozoic–Cenozoic sedimentary rocks and igneous rocks ranging in age from the Paleozoic to Cenozoic eras (Farmer, 2003) (Figure 2). As the bedrock compositions remain relatively constant throughout the hyperarid to arid profile, the primary Ca-P endmember should be stable throughout the sites and should reliably reflect the bedrock δ¹⁸OPO₄.

δ¹⁸OPO₄ under magmatic conditions is dominantly constrained by equilibrium isotope exchange at high temperature, while δ¹⁸OPO₄ of clastic sedimentary rocks is much more variable and can reflect decreasing ocean temperatures since the Cretaceous (Shemesh, Kolodny, & Luz, 1988). Normally, igneous rocks contain lower δ¹⁸OPO₄ values than sedimentary rocks, with igneous δ¹⁸OPO₄ varying between 4 and 11‰ (Angert et al., 2012; Blake et al., 2010; Jaisi & Blake, 2010; Markel, Kolodny, Luz, & Nishri, 1994; Mizota, Domon, & Yoshida, 1992). δ¹⁸OPO₄ of sedimentary rocks drops significantly from modern (23‰) to Late Cretaceous (19%), with the lowest value (12.8‰) recorded from the Permian. These trends suggest that ocean temperatures were higher at about 100 Myr ago, and post-depositional ¹⁸O exchange took place between phosphate and the ambient water, as mainly directed by the activity of phosphoenzymes, usually inorganic pyrophosphatase (Shemesh, Kolodny, & Luz, 1983; Shemesh et al., 1988; Tamburini et al., 2014). Accordingly, the δ¹⁸OPO₄ values are always greater in post-Cretaceous sedimentary rocks than in igneous rocks, and can be further enriched by biological activity (Figure 1).

The most hyperarid sites we sampled (with precipitation < 2 mm/year; MES, PONR-2, and Yungay) all have low δ¹⁸OPO₄ values in the Ca-P (7.9, 13.6, and 11.2‰, respectively). Due to the minimal extent of microbial and other biological activity at these sites, this range in isotope values is taken to reflect a bedrock endmember δ¹⁸OPO₄ signature for the wider region. Within the hyperarid core of the Atacama Desert, all three sites have similar proportions of magnesium (1.1 ± 0.1%) and iron (3.4 ± 0.4%) (Table 2), and hence a similar source contribution from igneous rocks. The sedimentary rock source to MES soils is derived from both Jurassic and Quaternary
periods, which is partially older than the solely Quaternary sedimentary rock source to the PONR-2 and Yungay sites. As a result, the $\delta^{18}$O$_{PO_4}$ value of MES soils is the smallest (Table 2). This parent bedrock isotopic endmember is thought to have remained stable for millennia due to the hyperarid nature of the environment and the minimal isotope fractionation associated with abiotic cycling of P (Figure 1) (Azua-Bustos et al., 2015).

Arid sites (TZ-4 and TZ-5), on the other hand, have a higher abundance of magnesium and iron contents than hyperarid sites (Table 2 and Figure 5d), indicating a larger proportion of igneous source rocks. However, the $\delta^{18}$O$_{PO_4}$ of Ca-P plateaus at higher values (22.6 ± 0.7‰) at these two more humid sites. This distinct shift in $\delta^{18}$O$_{PO_4}$ coincides with high TOC, TON, and microbial P (Figure 6b-d), indicating a major change in the source of Ca-P for sites with rainfall in excess of 15 mm/year. This trend suggests that the primary mineral $\delta^{18}$O$_{PO_4}$ signature has been overwritten, controlled by microbial phosphoenzyme hydrolysis and incorporating the $\delta^{18}$O signature of water.

Only hydrolysis by inorganic pyrophosphatase results in a positive $\delta^{18}$O$_{PO_4}$ (Figure 1). The results of enzyme and pathway prediction analyses imply that the hydrolysis by inorganic pyrophosphatase is the most likely reaction to cycle P in these environments, or that this process will rapidly overprint other P utilization strategies, in agreement with previous studies (Blake, 1998; Blake et al., 2005; Gross & Angert, 2015; Jaisi, Kukkadapu, Stout, Varga, & Blake, 2011; Tamburini et al., 2014).

In our sampling regions of the Atacama Desert, available moisture for inorganic pyrophosphatase hydrolysis is minimal. Some possible water sources for life might be sporadic rainfall and infrequent inland fog. For example, previous studies found that biological activity increases in Atacama soils after infrequent rainfall events (Schulze-Makuch et al., 2018; Vidiella et al., 1999) and that Atacama microbial communities and shrubs are capable of utilizing atmospheric water sources (Azua-Bustos et al., 2011; Mooney, Gulmon, Ehleringer, & Rundel, 1980; Stomeo et al., 2013). Groundwater might also act as a minor source of water for life. Importantly, hydration water of hydrated minerals such as the large gypsum reservoir in the Atacama Desert (Dong, Rech, Jiang, Sun, & Buck, 2007; Farias et al., 2014; Rech, Quade, & Hart, 2003) is potentially the main water source of endolithic micro-organisms and plants during the dry period (Palacio et al., 2014).

The inorganic pyrophosphatase-driven biogenic $\delta^{18}$O$_{PO_4}$ signature can be calculated by Eq. (4) using source water $\delta^{18}$O$_{H_2O}$. Due to the long-term strong effects of evaporation, water sources in the Atacama Desert are heavier in $\delta^{18}$O than in most other settings on Earth (Cappa, Hendricks, DePaolo, & Cohen, 2003; Dansgaard, 1964; Surma, Assonov, Herwartz, Voigt, & Staubwasser, 2018). $\delta^{18}$O$_{H_2O}$ of rainwater generally increases as the altitude gets lower, such that between 2,200 and 3,000 m above sea level (a.s.l.), rainwater $\delta^{18}$O$_{H_2O}$ varies between −12 and −2‰ (R Aravena, Peña, Grilli, Suzuki, & Mordecki, 1989; R. Aravena et al., 1999; Fritz, Suzuki, Silva, & Salati, 1981). During March 2015, the $\delta^{18}$O$_{H_2O}$ of rainwater from an extreme precipitation event in the Atacama Desert varied between −9 and −1‰ at 100 to 2,500 m a.s.l. (Jordan et al., 2019), consistent with previous data at this altitude range (Boschetti, Cifuentes, Jacumin, & Selmo, 2019; Schotterer et al. 1996). At about 500 to 1,600 m a.s.l., altitudes spanning our sampling locations (Table 1), the $\delta^{18}$O$_{H_2O}$ value of rainwater is expected to be closer to −2‰ than −10‰. In addition, local fog data for the Atacama Desert show that the $\delta^{18}$O$_{H_2O}$ of fog varies between −3 and −1‰ (R. Aravena, Suzuki, & Pollastri, 1989; Strauch, Oyarzun, Fiebig-Wittmaack, Gonzalez, & Weise, 2006). The range of Atacama groundwater $\delta^{18}$O$_{H_2O}$ is about −11 to + 3‰ (Alpers & Whittemore, 1990; R. Aravena et al., 1989; Herrera et al., 2018; Scheilings, Moya, Struck, Lictveout, & Troger, 2018), but groundwater is less likely to be a significant source to Atacama biological communities. Moreover, the $\delta^{18}$O$_{H_2O}$ of gypsum hydration water has a fractionation factor $\alpha^{18}$O$_{gypsum-water}$ of 1.0035 ± 0.0002 with surrounding water sources (Gazquez, Evans, & Hodell, 2017), so the hydration water $\delta^{18}$O$_{H_2O}$ of Atacama gypsum is anticipated to be slightly higher than the other water oxygen isotope ratios discussed above.

The results of enzyme and pathway predictions imply that the hydrolysis by inorganic pyrophosphatase dominates the biogeochemical P dynamics in Atacama soils (Figure 1). Therefore, if we assume that isotopic equilibrium has been reached at the arid sites (TZ-4 and TZ-5), we can re-arrange the microbial P turnover equation by inorganic pyrophosphatase (Eq. (4)) to establish the primary moisture source for the biological communities surviving and actively cycling P in this environment. Using the average temperature recorded from the nearest meteorological stations (Table 1) and supposed $\delta^{18}$O$_{PO_4}$ equilibrium values for each site (Table 2), we calculate the theoretical $\delta^{18}$O$_{H_2O}$ of source water to be $-3.31 \pm 0.34%$ at TZ-4, $+2.04 \pm 0.16%$ at TZ-5, and $-1.04 \pm 0.17%$ at the TZ-5 pit adjacent to shrubs. All values fall within the higher range expected for potential moisture sources, confirming the highly evaporative nature of the moisture available for biological communities (Hsieh, Chadwick, Kelly, & Savin, 1998).

At site TZ-5, we see differences between theoretical $\delta^{18}$O$_{H_2O}$ from the two samples collected. Considering the relationship between source water $\delta^{18}$O$_{H_2O}$ and biogenic soil $\delta^{18}$O$_{PO_4}$ discussed above, some explanations for this heterogeneity might be that (a) the equilibrium of water–phosphate interaction mainly driven by soil microbial inorganic pyrophosphatase is not fully reached, and (b) the water sources for life are distinct from each other even on a small spatial scale. Further investigation is required to ascertain the driving force behind these small differences in calculated source water $\delta^{18}$O values at site TZ-5.

However, observed differences of ~5% between the calculated isotopic signature of source water at TZ-4 and TZ-5 point to a stronger driving force. One clear difference between the sites is their altitudes, where TZ-5 is at almost 1,000 m lower elevation than TZ-4 (Table 1). When considering this difference, theoretical water isotope values follow the expected trend of decreasing rainwater
\( \delta^{18} \text{O}_{\text{H}_{2}\text{O}} \) with altitude (R. Aravena et al., 1999; Jordan et al., 2019). Again this finding requires further investigation, but if correct, this suggests that rainfall does indeed dominate water resources at these arid sites, even under hostile conditions of < 20 mm/yr rainfall (Orlando et al., 2010; Schulze-Makuch et al., 2018). How exactly this rainfall is accessed by the biological community is still uncertain. It can be assumed that a proportion is available directly after a rain event but that there must also be a critical slow release water reservoir, or regular additional water (e.g., fog and groundwater), which reflects rainfall isotope values and is utilized between the infrequent rainfall events. We suggest that gypsum hydration water is the most likely reserve of bioavailable moisture between rainfall events in these environments (Palacio et al., 2014). The fractionation of \( \delta^{18} \text{O} \) during uptake into gypsum is minimal but slightly positive, potentially explaining the theoretical positive isotope value of + 2.04‰ at TZ-5 which is higher than all theoretical rainfall and fog sources outlined above. The uptake of water into the gypsum mineral phase should also preserve the altitude-driven difference between rainfall at TZ-4 and TZ-5 as suggested by our theoretical \( \delta^{18} \text{O}_{\text{H}_{2}\text{O}} \) calculations.

On current Mars, diurnal cycles promote the formation of a thin layer of liquid water within the subsurface (Altheide, Chevrier, Nicholson, & Denson, 2009; Chevrier, Hanley, & Altheide, 2009; Cull et al., 2010; Kereszturi & Rivera-Valentin, 2012; Martinez & Renno, 2013; Meslin et al., 2013). This process means that small volumes of water are available to nourish indigenous Martian "microbes," possibly allowing them to slowly thrive and cycle key nutrients (Stevenson et al., 2015). Additionally, during portions of its planetary history Mars was far wetter than today (Craddock & Howard, 2002). On an earlier, wetter Mars, moisture might have in principle supported significant biologically catalyzed oxygen atom exchange and departures from the igneous \( \delta^{18} \text{O} \) baseline (Figure 1). The arid sites characterized by biogenic \( \delta^{18} \text{O}_{\text{PO4}} \) signature in this study have experienced similar climatic conditions to the middle-late Hesperian \( \delta^{18} \text{O}_{\text{PO4}} \) values move away from this lithogenic end-member. These results suggest that as biological activity increases, the \( \delta^{18} \text{O}_{\text{PO4}} \) of mineral P gradually shifts toward values closer to the equilibrium function of \( \delta^{18} \text{O}_{\text{H}_{2}\text{O}} \) providing a crucial biosignature if the \( \delta^{18} \text{O}_{\text{H}_{2}\text{O}} \) can be approximated. Critically, we have shown that microbial activity is capable of replacing authigenic P with biogenic P over millions of years, even under relatively inhospitable conditions similar to the middle-late Hesperian Mars. These \( \delta^{18} \text{O}_{\text{PO4}} \) values further suggest that microbial ecosystems living in such extreme environments can presumably utilize water from a variety of sources in addition to rainfall, including hydrated minerals.

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**CONFLICT OF INTEREST**

We have no conflict of interest to declare.

**AUTHOR CONTRIBUTION**

J.S., A.Z., and M.C. sampled the study sites. J.S. performed soil characterization, DNA extraction, cell culture, and all statistical analyses. J.S. and A.S. performed sequential phosphate extraction. A.S. performed stable oxygen isotope measurements in phosphate. All authors wrote and approved the final manuscript. Jianxun Shen administered the project, supervised the study, and reviewed and edited the manuscript. Andrew Smith designed methodology, performed stable and visualized data, and wrote the original draft. Mark Claire acquired funding, supervised the study, and reviewed and edited the manuscript. Andrew Smith designed methodology, performed validation, and reviewed and edited the manuscript. Aubrey Zerkle administered the project, supervised the study, and reviewed and edited the manuscript.

**DATA AVAILABILITY STATEMENT**

Sequencing data for enzyme and pathway analyses in this study can be found in the National Center for Biotechnology Information (NCBI) under the Sequence Read Archive (SRA) accession numbers of SRX7370993, SRX7370998, and SRX7370989.

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