

Do P Cycling Patterns Differ Between Ice-Free Areas and Glacial Boundaries in the Maritime Antarctic Region?

Authors: Chacón, Noemí, Ascanio, Mildred, Herrera, Rafael, Benzo, Diana, Flores, Saúl, et al.

Source: Arctic, Antarctic, and Alpine Research, 45(2) : 190-200

Published By: Institute of Arctic and Alpine Research (INSTAAR), University of Colorado

URL: https://doi.org/10.1657/1938-4246-45.2.190

BioOne Complete (complete.BioOne.org) is a full-text database of 200 subscribed and open-access titles in the biological, ecological, and environmental sciences published by nonprofit societies, associations, museums, institutions, and presses.

Your use of this PDF, the BioOne Complete website, and all posted and associated content indicates your acceptance of BioOne's Terms of Use, available at www.bioone.org/terms-of-use.

Usage of BioOne Complete content is strictly limited to personal, educational, and non - commercial use. Commercial inquiries or rights and permissions requests should be directed to the individual publisher as copyright holder.

BioOne sees sustainable scholarly publishing as an inherently collaborative enterprise connecting authors, nonprofit publishers, academic institutions, research libraries, and research funders in the common goal of maximizing access to critical research.

Do P Cycling Patterns Differ between Ice-Free Areas and Glacial Boundaries in the Maritime Antarctic Region?

Noemı **Abstract** *´Chaco´n*†§*

§Corresponding author: nutrients in the Antarctic Peninsula. nchacon@ivic.gob.ve

*Mildred Ascanio** Antarctic studies have indicated that during summer, ice-free areas experience greater temperatures than those along the glacial boundaries. This allows for the availability of *Rafael Herrera*‡* liquid water and thus influences the biogeochemical processes of Antarctic terrestrial ecosystems. In this study *Diana Benzo** ecosystems. In this study we explore whether the patterns of soil phosphorus cycling differ **between the glacial boundary and ice-free areas.** To do so, we chose two sites on the **Sau´l Flores** \overline{S} **Sau** \overline{S} $\overline{S$ Filmsula in Antarctica, one at the boundary of the Collins Glacier and another within an ice-free area close to Lake Uruguay. In each location, we determined soil phos-
*Belkis García** phorus distribution, iron and alumin *Belkis Garcı´a** phorus distribution, iron and aluminum fractions, soil mineralogy, alkaline phosphatase *Centro de Ecología, Instituto activity, and fluorescein diacetate hydrolysis. The results showed that soils of the ice-free Venezolano de Investigaciones area had a greater content of phosphorus sorbed on iron/aluminum oxyhydroxides and Científicas, Apdo. 20632 occluded forms. An opposite pattern was obtained for the calcium-bound phosphorus Caracas 1020-A, Venezuela nool Accordingly soil samples from the ice-free area showed the greatest levels of iron/ Caracas 1020-A, Venezuela

†Centro de Oceanología y Estudios

†Centro de Oceanología y Estudios

Antárticos, Instituto Venezolano de aluminum oxyhydroxides and the presence of secondary minerals such as hematite and

Inves Caracas 1020-A, Venezuela fluorescein diacetate hydrolysis were also favored at the ice-free area. The overall results ‡Grupo de Evaluación y Restauración de suggest that the less extreme microclimatic conditions and the presence of liquid water Sistemas Agrícolas y Forestales, in the ice-free area promote the biogeochemical cycling of soil phosphorus. In a context
Universidad de Córdoba, Edf. Leonardo
da Vinci s/n, Campus Universitario de
Rabanales, 14071 Córdoba

DOI: http://dx.doi.org/10.1657/1938-4246-45.2.190

Introduction

A number of studies have concluded that Antarctic soils can sustain communities of heterotrophic organisms in spite of the extremely low temperatures and dry conditions to which they are exposed (i.e., Burkins et al., 2000, 2001; Hopkins et al., 2006a, 2006b; Hopkins et al., 2008). These findings have given rise to several research projects focused on the biotic and/or abiotic factors driving the soil nutrient cycling in the terrestrial ecosystems of this region (i.e. Tscherko et al., 2003; Barrett et al., 2005, 2006; Gajananda, 2007; Hogg et al., 2006; Hopkins et al., 2006a, 2006b; Aislabie et al., 2008; Cannone et al., 2008).

Among the essential nutrients, phosphorus (P) is considered to be the element that most commonly limits net primary productivity in various soil conditions (Vitousek, 1984; Hinsinger, 2001). This is a consequence of its poor mobility due to high reactivity with numerous soil constituents such as iron (Fe) and aluminum (Al) oxyhydroxides, calcium (Ca), organic matter, and clay minerals (Hinsinger, 2001; Chacón and Dezzeo, 2004). In soils of the Antarctic continent there is some controversy regarding P as a limitation. In Antarctic field experiments it has been shown that primary productivity increases mainly by the addition of nutrients such as nitrogen (N) and P (Wasley et al., 2006; Hopkins et al., 2008). However, the elevated levels of available P, and the low C: P ratio of these soils, allowed other authors to suggest that P is not biologically limiting in the terrestrial ecosystems of this region (Bate et al., 2008). Nevertheless, it is well recognized that soils adjacent to Antarctic streams and lakes can be a significant source of available P to these biologically P limiting waters (i.e. Priscu, 1995; Dore and Priscu, 2001; Barrett et al., 2009).

Soil P is generally characterized by various pools that are a function of the state of soil development; whereas the primary inorganic pool is characteristic of young soils, the secondary inorganic- and organic-P pools predominate in more mature soils (Walker and Syers, 1976). Previous studies have shown that soil development is the result of the interplay among parent material, topography, climate, organisms, and time (i.e. Jenny, 1941; Kelly and Yonker, 2005; Bockheim et al., 2005). However, several authors agree that climate (temperature and humidity) plays an important role in soil formation because it controls the rate of physical, chemical, and biological soil processes (Cooper, 1960; Birkeland et al., 1989; Kelly and Yonker, 2005; Egli et al., 2006a; Barrett et al., 2009). In fact, in non-Antarctic cold regions the rate of soil evolution is considered to be distinctly determined by the availability and flow of soil water (Egli et al., 2003; 2006a, 2006b). However, in Antarctic regions, where the liquid water availability is restricted to austral summer (Chen and Blume, 2000; Gooseff et al., 2003), several authors have also recognized that climate is the main driving factor of soil genesis (Ugolini, 1963; Everett, 1976; Matsuoka, 1995; Zhao and Li, 1996; Chen and Blume, 1999; Gooseff et al., 2003; Cannone et al., 2008; Navas et al., 2008; Barrett et al., 2009; Bölter, 2011; López-Martínez et al., 2012).

Taking into account the above findings, in Antarctic terrestrial ecosystems it is expected that P distribution could be affected by climate. In this region only two studies have reported the soil P fractions (Blecker et al., 2006; Bate et al., 2008), and, in both, authors concluded that this element lies mainly in the primary inorganic pool. However, Blecker et al. (2006) indicated that differences in the weathering intensity between sites account for spatial variability in the soil P pools; while Bate et al. (2008) proposed that geologic legacies strongly influence the soil P distribution. It is noteworthy that these studies are restricted to continental sites in Antarctica, while for maritime Antarctica this question remains unclear.

Based on studies from maritime Antarctica, Ugolini and Bockheim (2008) provided considerable evidence for climate warming in this region. Some authors have characterized these climate conditions as the warmest and moistest within Antarctica (i.e. Ferron et al., 2004; Ugolini and Bockheim, 2008; Navas et al., 2008). Despite their sensitivity to climate warming, most areas of this region are still covered by glaciers (Lee et al., 2004), while the ice- and snowfree areas are scarce (Navas et al., 2008). There are strong microclimatic differences between glaciers and the ice- and snow-free areas. During the summer months, the relatively greater temperature of the ice- and snow-free areas leads to a deep thawing period that produces a more favorable supply of liquid water (Chen and Blume, 2000; Gooseff et al., 2003). In contrast, in terrestrial areas adjacent to subsurface ice, liquid water is limited to the warmest hours of the day (Motta and Motta, 2004).

Taking into account these findings and the role of water availability in the evolution of the Antarctic soils, we asked whether there were significant differences in the geochemical cycling of soil P between sites near glacial boundaries and those on ice-free areas. Were they to exist, we hypothesized that the concentration of several inorganic P pools defined as P sorbed on amorphous and some crystalline Al and Fe oxyhydroxides and occluded forms should show greater values in the ice-free areas. An opposite trend is expected for the Ca-bound P pool.

In addition to the foregoing hypothesis concerning the chemically defined P fractions, we also hypothesized that biological cycling of P should be favored in the ice-free areas. Support for this assumption comes from the research of Barrett et al. (2006, 2009), Gajananda (2007), Zeglin et al. (2009), and Bölter (2011), who have pointed out that due to the absence of higher plants, soil communities in terrestrial Antarctic ecosystems are largely controlled by abiotic factors. In principle, in those areas with greater liquid water availability due to increasing temperatures, biotic activity would be enhanced. Thus, in order to test these hypotheses, we chose a site along the boundary of the Bellingshausen Ice Dome (Collins Glacier) and another in an ice-free area in the vicinity of Lake Uruguay, Antarctica.

Materials and Methods

STUDY AREA AND SAMPLE COLLECTION

The study site was located in the vicinity of the Artigas Antarctic Scientific Base (Base Científica Antártica Artigas [BCAA]) on the Fildes Peninsula of King George Island, which forms part of the South Shetland Islands in Maritime Antarctica (61°54'-

 $62^{\circ}16'$ S; Fig. 1). The climate is maritime with an annual mean precipitation and temperature of 800 mm and -2 °C (1989–2001; Navas et al., 2008), respectively. During summer the temperatures of the surface soils can reach more than 10° C (Zhao and Li, 1996). Vegetation consists of lichens, mosses, algae, and the grass, *Deschampsia antarctica* (Soave et al., 2000).

The Fildes Peninsula, located at the extreme southwest of King George Island, is the largest ice-free area on the island (Soave et al., 2000; Vogt and Braun, 2004). In this region the last deglaciation occurred at 8–5 ka B.P. (López-Martínez et al., 2012). The peninsula consists of a thick subalkaline volcanic rock stratum with intercalations of volcaniclastic rock that characterize the Fildes Peninsula Group (Almeida et al., 2003). According to Zhao and Li (1996), soil development at this site is controlled by the parent material, landforms, and bioclimatic factors. Soils of the Fildes Peninsula are composed of Histosols, Regosols, Cambisols, and Andisols (Chen and Blume, 1999).

The soil samples were collected from two sites in the vicinity of BCAA: one at the boundary of the Collins Glacier (BG: $62^{\circ}10'49.62''S$, $58^{\circ}54'34.75''W$) and the other in an ice-free area (IFA: 62°11'08.33"S, 58°54'33.02"W) near Lake Uruguay (Fig. 1). We considered only one study site for the ice-free area and another one for the boundary of the glacier because genuine replicates of these sites were not feasible in widely spaced geographic areas. The lack of field replication and consequently the risk of pseudoreplication are almost inevitable in the studies of large-scale systems (i.e. rivers, small watersheds, forest patches) (Hargrove and Pickering, 1992; Oksanen, 2001) such as these. We think this level of pseudo-replication in our study does not undermine our results, though it does mean they should be interpreted with caution, as other authors in similar situations have pointed out.

In each study site, we sampled the surface mineral horizon (0–10 cm) from 15 points selected according to a completely random design (total $n = 30$). The sampling depth was limited to 10 cm due to the presence of stones and rocks. The samples were stored in plastic bags. While those samples destined for the analysis of microbial and enzymatic activity were maintained at $+4^{\circ}$ C, the samples for chemical analysis were air dried at room temperature (Tscherko et al., 2003). A preliminary analysis showed that soil textural class was similar between sites; however, other soil properties such as pH and organic carbon content (OC), showed differences (Benzo et al., unpublished data; samples collected in 2009; Table 1). During the sampling, the field temperatures recorded at lakes adjacent to BG and IFA sites were 2.4 $^{\circ}$ C and 5.3 $^{\circ}$ C, respectively.

P FRACTIONS

Soil P fractions were obtained according to the method of Hedley et al. (1982), which was modified by Tiessen and Moir (1993). This procedure uses a sequence of increasingly strong reagents to remove the labile and more stable P forms (Tiessen and Moir, 1993). Aliquots of 0.50 g of dry soil previously ground and sieved (0.43 mm) were sequentially extracted with anion exchange resin, followed by 0.5 M NaHCO₃, 0.1 M NaOH, 0.5 M HCl, and hot concentrated HCl. Total P was determined in aliquots of each extract by digestion with ammonium persulphate and H_2SO_4 to convert all the organic P forms (Po) to inorganic P forms (Pi).

FIGURE 1. Map of King George Island showing the relative locations of the studied sites. BCAA = Base Científica Antártica Artigas, BG boundary of the glacier, IFA ice-free area. Digitalized image from the Letter 1111: Artigas Antarctic Scientific Base (Service of Oceanography, Hydrography and Meteorology of the Navy of Uruguay, 1991).

Another aliquot of each extract was used to measure Pi after acidification with H_2SO_4 to precipitate organic matter. In each extract, the pH was adjusted, and the P concentration was determined colorimetrically by means of the molybdate-ascorbic acid procedure (Murphy and Riley, 1962). Organic phosphorus was calculated as the difference between total P and Pi. According to Tiessen and Moir (1993), soil P fractions obtained by this procedure have been associated with the following operationally defined pools: (i) P extracted with resin is often considered as the labile P; (ii) bicarbon-

TABLE 1

Some soil properties in the boundary of the glacier (BG) and in the ice-free area (IFA) (D. Benzo et al., unpublished data).

Mean values \pm 1 SE followed by different letters in rows denote significant differences between sites. $OC =$ organic carbon.

ate- and hydroxide-extractable P are distinguished as labile and moderately labile P forms respectively, associated to Fe/Al oxyhydroxides; (iii) diluted HCl P is associated with the primary mineral and/or Ca-bound P; and (iv) hot concentrated HCl P is defined as the highly resistant and unavailable P.

Al AND Fe FRACTIONS

A sequential selective dissolution method was used to estimate the organically bound, non-crystalline and crystalline forms of Fe and Al (Wieder and Lang, 1986). Organically bound Fe and Al were extracted with sodium pyrophosphate (Fe_P, Al_P) (McKeague, 1967). Poor crystalline forms of Fe and Al (Fe $_{\Omega}$, Al $_{\Omega}$) were obtained by extracting the soil with ammonium oxalate ($pH = 4$ in the dark) (McKeague and Day, 1966), and highly structured Fe and Al associated with crystalline Fe oxides were extracted with dithionitecitrate-bicarbonate (Fe_{DCB} , Al_{DCB}) following the method of Mehra and Jackson (1960). Concentrations of both elements in each extract were determined using atomic absorption spectroscopy.

SOIL MINERALOGY

Soil mineralogy was obtained via X-ray diffraction (XRD) using Cu-K α radiation. Four representative soil samples from each

study site were made into pressed powder pellets for XRD analysis. Samples were mounted on glass plates and analyzed from 5° to 90° 20 with a speed of 2° 20 per minute. The software code PDFWIN for windows was used for the evaluation of diffraction patterns.

BIOLOGICAL AND BIOCHEMICAL ANALYSIS

In this study, the activity of alkaline phosphatase (AP) was measured as an index of organic P mineralization (Dick and Tabatabai, 1993), while the fluorescein diacetate (FD) hydrolysis was used as a measure of soil microbial activity (Schnürer and Rosswall, 1982; Green et al., 2006). In alkaline soils such as at the two study sites, it is advisable to measure the activity of AP because it is usually greater than that of the acid phosphatase (Eivazi and Tabatabai, 1977). Briefly, aliquots of fresh soil after incubation at $+37$ -C were used to assess FD hydrolysis and AP activity following the procedures of Schnürer and Rosswall (1982) and Tabatabai and Bremner (1969), respectively.

STATISTICAL ANALYSIS

A one-way analysis of variance (ANOVA) was used to compare soil P fractions, Fe and Al forms, and microbial and biochemical (fluorescein diacetate hydrolysis and alkaline phosphatase) activities. When necessary, data were log-transformed to meet the assumptions of ANOVA (NaOH-Pi, Al_T, FDA). The Tukey Honest Significant Difference (HSD) test was used as a means separation procedure. The Mann Whitney non-parametric test was used when the data did not meet ANOVA assumptions (resin-Pi, NaHCO₃-Pi, NaOH-Po, HCl_{conc}-Po, Al_O, Al_{OCB}, Al_O/_{AlDCB}). Pearson's correlation coefficient (*r*) was used to examine the relationship among parameters (soil P fractions, and microbial and biochemical activities versus P pools). Univariate analyses were performed using STATISTICA for Windows 6.0 (Statistica, 2001).

We used a Principal Component Analysis (PCA) to determine if the concentration of soil inorganic P pools can discriminate between sites influenced by glaciers and the ice-free areas. PCA was performed using the software Multi-Variate Statistical Package (MVSP 3.0 for Windows; Kovach, 1998).

Results

SOIL P FRACTIONS

Concentrations of soil P in the various fractions at both sites are summarized in Table 2. The soil from the BG had less P in the resin-Pi fraction. No difference $(P > 0.05)$ between sites was observed among the mean values of $NaHCO₃-Pi$. None of the sampled sites showed significant differences ($P > 0.05$) in organic P fractions (NaHCO₃-, NaOH-, HCl_{conc}-). The concentration of NaOH-Pi fraction, which is associated with the P strongly chemisorbed on amorphous and some crystalline Al and Fe phosphates (Tiessen et al., 1983), increased significantly in IFA. The soil at IFA also contained greater P in the concentrated HCl-Pi fraction.

TABLE 2

Values of soil phosphorus fractions (mg kg^{-1}) in the boundary of **the glacier (BG) and in the ice-free area (IFA).**

Fraction	BG	IFA
Resin-Pi	27.48 ± 1.32^b	$47.88 \pm 4.06^{\circ}$
$NaHCO3-Pi$	$0.96 \pm 0.19^{\text{a}}$	$3.00 \pm 1.34^{\circ}$
NaOH-Pi	2.96 ± 0.43^b	$24.30 \pm 12.91^{\circ}$
HCl (dil)-Pi	$385.89 \pm 9.21^{\text{a}}$	281.75 ± 16.72^b
HCl(conc)-Pi	26.80 ± 1.97^b	$39.15 \pm 1.68^{\circ}$
$NaHCO3-Po$	$7.95 \pm 1.70^{\circ}$	$8.46 \pm 2.57^{\circ}$
NaOH-Po	$1.98 \pm 0.41^{\circ}$	$9.47 \pm 5.97^{\text{a}}$
$HC1-PO$	$0.68 \pm 0.35^{\circ}$	$2.52 \pm 1.03^{\circ}$

Mean values \pm 1 SE followed by different letters in rows denote significant differences between sites. $Pi = inorganic phosphorus;$ $Po =$ organic phosphorus.

One important difference between sites was that soil of the IFA, which contained greater P concentrations in NaOH-Pi and HCl_{cone} -Pi fractions, had lower P in the diluted HCl-Pi fraction, which is linked to unweathered Ca-bound P phases. At this site the weathering of primary minerals seemed to yield soluble P because a positive correlation ($r = 0.63$, $P < 0.05$; $n = 15$) was obtained between resin-Pi and the diluted HCl-Pi fractions. Both parameters were not correlated at the BG site ($r = -0.03$, $P > 0.05$; $n =$ 15). Soil from the IFA also showed a negative correlation between NaOH-Pi and the diluted HCl-Pi fraction ($r = -0.77$, $P < 0.05$; $n = 15$). This correlation was not observed in the BG soil ($r =$ 0.09, $P > 0.05$; $n = 15$).

Principal Component Analyses (PCA) of the soil inorganic P fractions from the two studied sites is shown in Figure 2. Only the first two axes of ordination are illustrated; these accounted for 65.0% of the variance (Table 3). This analysis clearly showed that, whereas the BG site was associated with soils showing high concentration of the diluted HCl-Pi and low concentrations of resin-, NaHCO₃-, NaOH-, and concentrated HCl-Pi fractions, the IFA findings were opposite.

IRON AND ALUMINUM FRACTIONS

The results of the organically bound, non-crystalline and crystalline forms of Fe and Al are presented in Table 4. In the BG site, the content of total extractable iron (Fe_T = Fe_P + Fe_O + Fe_{DCB}) was well above the total extractable aluminum ($Al_T = Al_P + Al_O$ $+$ Al_{DCB}); however, in the IFA they were in the same order of magnitude.

In both sites, the content of organically bound forms of Fe and Al (Fe_{P}, Al_{P}) was low and their contributions to the content of Fe_T and Al_T were as follows: BG: Fe_P = 1.8%, Al_P = 11.7%; IFA: Fe_P = 3.2%, Al_P = 13.5%. Comparisons of mean values between BG and IFA revealed no significant differences ($P > 0.05$) for Fe_P, whereas Al_P increased sharply in the IFA soil. Poorly crystalline forms of Fe and Al (Fe_O, Al_O) followed a trend similar to that of the Fe $_{\rm P}$ and Al $_{\rm P}$ fractions. The greatest content of crystalline Fe (Fe_{DCB}) was found in the BG soil; however, there was no difference in the content of crystalline Al (AI_{DCB}) between sites. Non-crystallinity ratios (Al_0/Al_{DCB} , Fe_O/Fe_{DCB}), interpreted as the

Component 1

FIGURE 2. Principal Component Analysis for soil P fractions data of the studied sites. IFA ice-free area, BG boundary of the glacier.

larger ratio the lower crystallinity (Beck and Elsenbeer, 1999), were greatest in the IFA soil.

SOIL MINERALOGY COMPOSITION

Soil mineralogical composition was different between studied sites. While in the BG site, XRD analysis showed the presence of labradorite, quartz, and the P-bearing mineral hydroxyapatite, the major mineral phases identified in the IFA were labradorite, quartz, and the secondary minerals hematite and chlorite-vermiculitemontmorillonite (Fig. 3; Table 5). Relative abundance of soil minerals at the study sites, as judged from peak heights of the diffractograms, was as follows: BG: labradorite $>$ quartz $>$ hydroxyapatite; IFA: labradorite $>$ hematite $>$ quartz $>$ chlorite-vermiculite-montmorillonite (Fig. 3).

TABLE 3

Eigenvalues and cumulative percentage variance of soil phosphorus fractions for the Principal Component Analysis carried out with data of the studied sites.

ALKALINE PHOSPHATASE AND FLUORESCEIN DIACETATE HYDROLYSIS. IMPLICATIONS IN THE GEOCHEMICAL CYCLING OF P

There was a significant difference between sites with respect to enzyme activity and fluorescein diacetate hydrolysis (Table 6). The greatest values for both parameters were detected at the IFA.

TABLE 4

Values of iron and aluminum fractions $(mmol kg⁻¹)$ in the bound**ary of the glacier (BG) and in the ice-free area (IFA).**

Fraction	BG	IFA
Fe _P	$3.01 \pm 0.39^{\circ}$	$3.92 \pm 0.36^{\circ}$
Fe _O	$53.31 \pm 1.95^{\circ}$	$55.45 \pm 2.76^{\circ}$
Fe _{DCB}	$107.77 \pm 3.67^{\text{a}}$	61.13 ± 3.49^b
Fe _T	164 ± 4^a	$121 \pm 5^{\rm b}$
Al_{P}	3.39 ± 1.50^b	21.21 ± 3.46^a
Al _O	$5.45 \pm 2.39^{\rm b}$	$109.70 \pm 9.40^{\circ}$
Al_{DCB}	$20.21 \pm 0.62^{\circ}$	$25.68 \pm 2.80^{\circ}$
Al_T	$29 + 3^b$	$157 \pm 12^{\rm a}$
Fe _O /Fe _{DCB}	0.50 ± 0.02^b	$0.93 \pm 0.05^{\text{a}}$
Al_0/Al_{DCB}	0.30 ± 0.14^b	$4.72 \pm 0.54^{\circ}$

Mean values \pm 1 SE followed by different letters in rows denote significant differences between sites. Fe_O-Al_O , $Fe_{DCB}-Al_{DCB}$, and Fe_P-Al_{-P} = ammonium-oxalate, dithionite-citrate-bicarbonate, and pyrophosphate extractable Fe or Al, respectively; $Fe_T-AI_T = \sum Fe_O$ Al_O, Fe_{DCB}-Al_{DCB}, and Fe_P-Al-_P, respectively.

Downloaded From: https://bioone.org/journals/Arctic,-Antarctic,-and-Alpine-Research on 15 Sep 2024 Terms of Use: https://bioone.org/terms-of-use

FIGURE 3. Typical X-ray diffraction patterns showing the three main peaks of the identified minerals at the studied sites. IFA icefree area, BG = boundary of the glacier, Lab = labradorite, Qz = quartz, Hem = hematite, $HyAP$ = hydroxyapatite, Chl-Vrm-Mnt **chlorite-vermiculite-montmorillonite.**

To investigate the relationships between geochemical and biological cycling of P in each site, correlations between chemically defined P pools and the biological parameters (alkaline phosphatase activity and fluorescein diacetate hydrolysis) were established. At the IFA, there was a positive correlation between the activity of AP and the hydrolysis of FD ($r = 0.95$, $P < 0.05$; $n = 15$). For this site, both biological parameters were positively correlated with the

organic and inorganic P fractions obtained in alkali extracts: NaHCO₃ ($r = 0.96$ for AP and $r = 0.91$ for FD in inorganic pool; $r = 0.90$ for AP and $r = 0.92$ for FD in organic pool; $P \le 0.05$, $n=15$) and NaOH ($r=0.98$ for AP and $r=0.98$ for FD in inorganic pool; $r = 0.91$ for AP and $r = 0.81$ for FD in organic pool; $P < 0.05$, $n = 15$). The relationships between microbial and enzymatic activities and the soil P pools were not evident at the BG site.

TABLE 5

Chemical composition and d-spacing values (A˚) of the three main peaks of identified minerals by XRD analysis in each studied site.

 PM = pure minerals, BG = boundary of the glacier, IFA = ice-free area.

Discussion

CHANGES IN P POOLS, Fe AND Al FRACTIONS, AND SOIL MINERALOGY FROM THE BOUNDARY OF THE GLACIER TO ICE-FREE AREA

It has been widely accepted that in the Antarctic region the extremely low temperatures and the limited availability of liquid water restricts chemical weathering and consequently soil development (i.e. Ugolini, 1963; Chen and Blume, 1999; Gooseff et al., 2003; Bölter, 2011). However, in some locations, during the summer, soil microclimatic conditions allow the presence of liquid water and favor chemical weathering. This is the case of the icefree areas at the Fildes Peninsula, which are exposed to a deep thawing period that lasts between 5 and 6 months each year (Chen and Blume, 1999). During this period, soils of these areas are subjected to greater temperatures and availability of free water than at other sites in Antarctica (Zhao and Li, 1996; Chen and Blume, 1999, 2000; Gooseff et al., 2003). In contrast, in terrestrial areas adjacent to subsurface ice, the liquid water content is limited due

TABLE 6

Alkaline phosphatase (AP) activity and the fluorescein diacetate (FD) hydrolysis of soils in the boundary of the glacier (BG) and in the ice-free area (IFA).

Mean values \pm 1 SE followed by different letters in rows denote significant differences between sites.

to the low air temperature and precipitation (Barrett et al., 2006). During summer, melting water from the glaciated areas may remain in a liquid state only during the warmest hours of the day and refreezes as solar radiation decreases (Motta and Motta, 2004). Taking into account the differences in the microclimatic soil conditions of these two environments, and considering the implications of the water availability in soil chemical processes, we predicted in our first hypothesis that from the boundary of the glacier (BG) to the ice-free area (IFA) P associated with primary minerals should decrease, while P bounded to secondary minerals should increase. To test this, we interpreted our results of soil P fractions along with the contents of secondary minerals (oxyhydroxides of Fe and Al) and soil mineralogy following the theory of Walker and Syers (1976). This theory proposes a conceptual model where availability and forms of soil P vary during its development as a consequence of changes in mineral composition. According to the model, as soil evolution takes place, the weathering of primary minerals (Cabound phosphates) releases P, which is readily chemisorbed into the secondary minerals (clay minerals as well as Fe and Al oxyhydroxides).

Our results advance four lines of evidence that help us support our first hypothesis. First, from BG to IFA, the concentration of primary mineral P (diluted HCl-Pi) declined, while the inorganic P chemisorbed and occluded to Fe/Al oxyhydroxides (NaOH- and concentrated HCl-) increased. This pattern led to an inverse relationship between the diluted HCl-Pi and NaOH-Pi at the IFA, but not at the BG site. Such findings were consistent with those obtained using PCA, which showed that inorganic P pools were distinct between the edge of the glacier and the ice-free area. Second, the greatest levels of Fe/Al oxyhydroxides were found in the IFA where the content of NaOH-Pi was also the greatest. Third, at the BG site, the content of Al_T was sixfold lower than that of the Fe_T, and fivefold lower than the content of Al_T in the soils of the IFA.

In the latter site, the contents of Fe_T and Al_T were similar. The observed trend could be considered a signal of advanced soil development at the IFA because, in the soil environment, high contents of secondary Al minerals are indicative of increased weathering (Hsu, 1977). Fourth, the presence of epigenetic minerals such as hematite and the intergrade chlorite-vermiculite-montmorillonite in the soil of the IFA is strong evidence that chemical alteration of bedrocks seems to be favored in this site. This was not the case at the BG site, where soil mineralogy was characterized by the presence of hydroxyapatite and the absence of chemical weathering products. The soil mineralogy observed at the IFA was in good agreement with the results obtained for other sites in the Antarctic Peninsula (Boyer, 1975; Kuzmann et al., 1998), where chemical weathering is an important pedogenetic process.

Besides the above-mentioned, it is important to note that, according to the theory of Walker and Syers (1976), the chemical alteration of primary minerals provides available P to the soil solution. In this study, this was observed in the IFA soils, since a direct relationship between resin-Pi and the diluted HCl-Pi fraction was obtained. Despite the fact that the alteration of primary minerals increased the available P at the IFA soils, such increase was not reflected in the organic P pool, which was very low and similar between the sites. This result is expected in an environment where the primary productivity is negligible and where, consequently, the content of soil organic matter is low (i.e. Hopkins et al., 2006a, 2006b).

The observed differences in P cycling patterns as well as the formation of epigenetic minerals seem to confirm our first hypothesis. At this point it is important to mention that pedogenetic time was not considered in our discussion, because differences in soil properties could be explained as a function of time when soils are being formed under the same climatic condition (Kelly and Yonker, 2005). However, this is not the case at the studied sites, since, as was mentioned earlier, soils of the ice-free areas are exposed to a less extreme microclimatic soil condition than the terrestrial areas adjacent to subsurface ice.

Despite that soils of the IFA are significantly more developed than soils of the BG site, in this study it is also important to recognize that the alteration of primary minerals in the IFA is still at an early stage since the content of Ca-bound P (281.75 \pm 16.72 mg kg^{-1}) is at the lower limit of the values reported by other authors (Blecker et al., 2006 [\approx 267–349 mg kg⁻¹]; Bate et al., 2008 $[\approx 250 - 980$ mg kg⁻¹]) for soils of the Antarctic continent, where the weathering is more restricted.

Fe AND Al FRACTIONS AND THEIR IMPLICATIONS ON THE SOIL P CYCLING

The non-crystalline forms of Fe and Al were found to be more abundant in the IFA site, where the chemical weathering was more intense (\sum Fe_O-Al_O: BG = 58.76 mmol kg⁻¹, IFA = 165.15 mmol kg^{-1}). As a result, this led to an increase of the non-crystallinity ratios (Al_0/Al_{DCB} , Fe_O/Fe_{DCB}) in the IFA soils. Non-crystalline forms of Fe and Al (Fe_O-Al_O) are considered the main sorbents of phosphate, in relation to crystalline forms, due to their comparatively larger specific surface area (Parfitt, 1989; Ugolini and Dahlgren, 2002).

Whilst the soils of the IFA were enriched with non-crystalline minerals, crystalline forms of Fe were dominant at the BG site. This result seems to be closely related to chemical weathering. According to Ugolini and Dahlgren (2002), when volcanic materials, such as those that make up the Fildes Peninsula Group, are exposed to chemical weathering, the precipitation of non-crystalline minerals is kinetically favored over precipitation of crystalline minerals. In comparison, crystalline minerals are formed at the expense of metastable compounds (Ugolini and Dahlgren, 2002). These authors consider the time of exposure of the weathering processes that are associated with warm and dry soil conditions as the main factor promoting the formation of crystalline minerals. Soils of the BG site are not exposed to warm temperatures, but they are subjected to longer periods of liquid water limitation, which probably favors the formation of crystalline minerals rather than non-crystalline ones.

In addition to poorly crystalline minerals, the organically bound forms of Fe and Al (Fe_P, Al_P) are also active surfaces that control soil P retention capacity (Gerke and Hermann, 1992; Chacón and Dezzeo, 2004; Chacón et al., 2005a, 2008). In this study, the content of these compounds was low as a result of the generally low content of soil organic matter. However, Al_P was more abundant in the soil from the IFA site than that from the BG site. This finding, along with the great content of non-crystalline minerals obtained for this site, allowed us to conclude that, potentially, IFA soils have a greater P retention capacity than soils at the BG. The implications of these mineral surfaces on P dynamics have been well recognized for highly weathered acid soils (i.e. Schwertmann and Taylor, 1977; Parfitt, 1978; Chacón and Dezzeo, 2004; Chacón et al., 2005a, 2008). However, it is important to bear in mind that these mineral surfaces are positively charged over the whole pH range commonly encountered in the soil (Hinsinger, 2001) and also control the P dynamics of alkaline soils (Matar et al., 1992; Bertrand et al., 1999; Hinsinger, 2001). In fact, in alkaline soils of the Antarctic continent, the increases of Al and Fe phosphates in soils where weathering is less restricted have been explained by the presence of non-crystalline and crystalline Fe and Al oxyhydroxides (Blecker et al., 2006).

ALKALINE PHOSPHATASE AND MICROBIAL ACTIVITY AND THEIR RELATIONSHIP WITH THE BIOGEOCHEMICAL CYCLING OF P

On the basis that biological activities in the terrestrial ecosystems of Antarctic region are largely controlled by abiotic factors (soil temperature and availability of liquid water), we formulated our second hypothesis wherein we predicted an increase in microbial and enzymatic activities for the ice-free area. Indeed, AP activity followed this expected trend. However, before we conclude that such behavior is promoted by the less extreme microclimatic conditions in the IFA, we ascertained within the context of our results, other concepts that could explain the observed pattern with this enzyme.

On a global scale, it has been indicated that extracellular phosphatase activities increase with soil organic matter concentration (Sinsabaugh et al., 2008). In this study, the content of soil organic carbon at the IFA was 13-fold lower than the recorded value at the BG site (Benzo et al., unpublished data; samples collected in 2009), even though the AP activity was fivefold greater at the IFA than at the BG site. Comparing our findings with results obtained by Sinsabaugh et al. (2008), it can be concluded that in our study sites there is no indication that soil organic carbon plays an important role in the activity of AP. However, other factors inherent to soil matrix could be masking our results. Several studies have found that the interaction of phosphatase enzymes with non-crystalline Al and Fe oxides and clay minerals decreases enzyme activity when compared to the non-immobilized enzyme activity (Huang et al., 1995; Rao et al., 1996, 2000; Shindo et al., 2002; Chacón et al., 2005b). As was already discussed above, soils of the IFA were enriched with non-crystalline compounds, whereas crystalline forms dominated the soils of the BG site. Hence, from this mineralogical composition we could not explain the observed pattern of this soil enzyme. Generally speaking, most secondary minerals are found in the fine fraction of the soil. According to Benzo et al. (unpublished data; samples collected in 2009), soil from the BG site showed a relatively greater content of fine particles (clay) than that from the IFA. In both sites, the correlation of these data with the values of AP activity obtained in this study was not significant (BG: $r = -0.217$, $P > 0.05$, $n = 15$; IFA: $r = 0.117$, $P > 0.05$, $n = 15$). This trend and the previous findings allow us to conclude that soil matrix is not driving the AP activity in the studied sites.

Once the influence of soil matrix on AP activity was discarded, it could be assumed that the activity was likely enhanced by soil microclimate conditions such as humidity and temperature. For Antarctic soils this assumption is reasonable considering that in this harsh environment microbial activities, like carbon and nitrogen mineralization and enzyme production (i.e. alkaline phosphatase), are favored in those areas where temperature allows greater soil water availability (Barrett et al., 2005; Zeglin et al., 2009).

Considering the findings mentioned above and the fact that FD hydrolysis has been widely used to determine the overall soil microbial activity potential (Nannipieri et al., 2003), it was not surprising that this parameter followed the same behavior detected for AP activity. Unfortunately, to the best of our knowledge, for Antarctic soils there are no studies evaluating the influence of soil microclimate conditions on FD hydrolysis. However, for temperate ecosystems it has been reported that suitable conditions of soil temperature and moisture increased the levels of this biochemical parameter (Sidari et al., 2008; Lillo et al., 2011; Reyes et al., 2011).

Finally, it is also important to analyze the positive relationship between biological parameters (AP activity and FD hydrolysis) and both organic and inorganic P pools obtained in alkaline extractants of soils of the IFA. This striking finding suggests that in IFA soils the presence of liquid water allowed the co-occurrence of advanced chemical weathering and a greater biological activity. The conjunction of both factors seems to have a strong influence over the P pools at the IFA.

Conclusions

The changes in the inorganic P pools coupled with the increase in secondary minerals at the IFA soils allowed us to conclude that soils from the IFA showed a greater chemical weathering intensity than soils from the BG. This finding was consistent with the pedogenic theory developed by Walker and Syers (1976). However, the prediction, which states that as soil evolution proceeds organic

P pools increase, could not be verified in this study due to the lack of variation between sites in this P pool. This discrepancy is expected in the harsh environments of Antarctica where reduced input of liquid water is responsible for negligible primary productivity. The biochemical transformation of soil P was also favored in the ice-free area. It is likely that the less restrictive microclimatic conditions, to which the soils of the ice-free areas are exposed during the summer, have a considerable effect on the biogeochemical cycling of soil phosphorus. Taken together, our findings demonstrated that, between the boundary of the glacier and the ice-free area, there are differences in the patterns of soil P cycling. In a scenario of climate warming over the Antarctic Peninsula, this information may contribute to the understanding of how the retreat of the glaciers and consequently the advance of the ice-free areas could be altering the biogeochemical cycling of essential nutrients in this region.

Acknowledgments

This research was supported by the Venezuelan Antarctic Program of the Ministerio del Poder Popular para Ciencia, Tecnología e Innovación (MCTI). Chacón thanks Maximiliano Bezada and Daniel Palacios, who helped in the fieldwork. We thank Cristina Lull for her critical comments on an early version of this manuscript. We are very grateful to three anonymous reviewers for their careful revision and valuable and critical commentaries about the original version of the manuscript. We also wish to acknowledge Werner Wilbert and Alessia Bastianoni for revising the English version of the manuscript.

References Cited

- Aislabie, J. M., Jordan, S., and Barker, G. M., 2008: Relation between soil classification and bacterial diversity in soils of the Ross Sea region, Antarctica. *Geoderma*, 144: 9–20.
- Almeida Delia, Del, P. M., Machado, A., Hansen, M. A. F., Chemale, F., Jr., Fensterseifer, H. C., Petry, K., and De Lima, L., 2003: An igneous event at the Fildes Peninsula (King George Island) and around Fort Point (Greenwich Island), South Shetland Islands, Antarctica. *Revista Brasileira de Geocieˆncias*, 33: 339–348.
- Barrett, J. E., Virginia, R. A., Parsons, A. N., and Wall, D. H., 2005: Potential soil organic matter turnover in Taylor Valley, Antarctica. *Arctic, Antarctic, and Alpine Research*, 37: 108–117.
- Barrett, J. E., Virginia, R. A., Wall, D. H., Cary, S. C., Adams, B. J., Hacker, A. L., and Aislabie, J. M., 2006: Co-variation in soil biodiversity and biogeochemistry in northern and southern Victoria Land, Antarctica. *Antarctic Science*, 18: 535–548.
- Barrett, J. E., Gooseff, M. N., and Takacs-Vesbach, C., 2009: Spatial variation in soil active-layer geochemistry across hydrologic margins in polar desert ecosystems. *Hydrology and Earth System Science*, 13: 2358–2358.
- Bate, D. B., Barrett, J. E., Poage, M. A., and Virginia, R. A., 2008: Soil phosphorus cycling in an Antarctic polar desert. *Geoderma*, 144: $21 - 31$.
- Beck, M. A., and Elsenbeer, H., 1999: Biogeochemical cycles of soil phosphorus in southern alpine Spodosols. *Geoderma*, 91: 249–260.
- Bertrand, I., Hinsinger, P., Jaillard, B., and Arvieu, J. C., 1999: Dynamics of phosphorus in the rhizosphere of maize and rape grown on synthetic, phosphated calcite and goethite. *Plant and Soil*, 211: 111–119.
- Birkeland, P. W., Burke, R. M., and Benedict, J. B., 1989: Pedogenic gradients for iron and aluminum accumulation and phosphorus depletion in arctic and alpine soils as a function of time and climate. *Quaternary Research*, 32: 193–204.
- Blecker, S. W., Ippolito, J. A., Barrett, J. E., Wall, D. H., Virginia, R. A., and Norvell, K. L., 2006: Phosphorus fractions in soils of Taylor Valley, Antarctica. *Soil Science Society of America Journal*, 70: 806–815.
- Bockheim, J. G., Gennadiyev, A. N., Hammer, R. D., and Tandarich, J. P., 2005: Historical development of key concepts in pedology. *Geoderma*, 124: 23–36.
- Bölter, M., 2011: Soil development and soil biology on King George Island, maritime Antarctic. *Polish Polar Research*, 32: 105–116.
- Boyer, S. J., 1975: Chemical weathering of rocks on the Lassiter Coast, Antarctic Peninsula, Antarctica. *New Zealand Journal of Geology and Geophysics*, 18: 623–628.
- Burkins, M. B., Virginia, R. A., Chamberlain, C. P., and. Wall, D. H., 2000: Origin and distribution of soil organic matter in Taylor Valley, Antarctica. *Ecology*, 81: 2377–2391.
- Burkins, M. B., Virginia, R. A., and Wall, D. H., 2001: Organic carbon cycling in Taylor Valley, Antarctica: quantifying soil reservoirs and soil respiration. *Global Change Biology*, 7: 113–125.
- Cannone, N., Wagner, D., Hubberten, H. W., and Guglielmin, M., 2008: Biotic and abiotic factors influencing soil properties across a latitudinal gradient in Victoria Land, Antarctica. *Geoderma*, 144: 50–65.
- Chacón, N., and Dezzeo, N., 2004: Phosphorus fractions and sorption processes in soil samples taken in a forest-savanna sequence of the Gran Sabana in southern Venezuela. *Biology and Fertility of Soils*, $40:14-19$.
- Chacón, N., Dezzeo, N., Muñoz, B., and Rodríguez, J. M.; 2005a: Implications of soil organic carbon and the biogeochemistry of iron and aluminum on soil phosphorus distribution in flooded forests of the lower Orinoco River, Venezuela. *Biogeochemistry*, 73: 555–566.
- Chacón, N., Dezzeo, N., and Flores, S., 2005b: Effect of particle-size distribution, soil organic carbon content and organo-mineral aluminium complexes on acid phosphatases of seasonally flooded forests soils. *Biology and Fertility of Soils*, 41: 69–72.
- Chacón, N., Dezzeo, N., Rangel, M., and Flores, S., 2008: Seasonal changes in soil phosphorus dynamics and root mass along a flooded tropical forest gradient in the lower Orinoco River, Venezuela. *Biogeochemistry*, 87: 157–168.
- Chen, B. J., and Blume, H. P., 1999: Study on the dynamics of soil moisture in an ice-free area of the Fildes Peninsula, King George Island, the maritime Antarctica. *Polarforschung*, 66: 11–18.
- Chen, J., and Blume, H. P., 2000: Element enrichment and migration within some soils on Fildes Peninsula (King George Island), the maritime Antarctica. *Journal of Plant Nutrition and Soil Science*, 163: 291–297.
- Cooper, A. W., 1960: An example of the role of microclimate in soil genesis. *Soil Science*, 90: 109–120.
- Dick, W. A., and Tabatabai, M. A., 1993: Significance and potential uses of soil enzymes. *In* Blaine, M. F. (ed.), *Soil Microbial Ecology, Applications in Agriculture and Environmental Management.* New York: Marcel Decker, 95–127.
- Dore, J. E., and Priscu, J. C., 2001: Phytoplankton phosphorus deficiency and alkaline phosphatase activity in the McMurdo dry valley lakes, Antarctica. *Limnology and Oceanography*, 46: 1331–1346.
- Egli, M., Mirabella, A., Sartori, G., and Fitze, P., 2003: Weathering rates as a function of climate: results from a climosequence of the Val Genova (Trentino, Italian Alps). *Geoderma*, 111: 99–121.
- Egli, M., Wernli, M., Kneisel, Ch., and Haeberli, W., 2006a: Melting glaciers and soil development in the proglacial area Morteratsch (Swiss Alps): I. Soil Type chronosequence. *Arctic, Antarctic, and Alpine Research*, 38: 499–509.
- Egli, M., Mirabella, A., Sartori, G., Zanelli, R., and Bischof, S., 2006b: Effect of north and south exposure on weathering rates and clay mineral formation in alpine soils. *Catena*, 67: 155–174.
- Eivazi, F., and Tabatabai, M. A., 1977: Phosphatases in soils. *Soil Biology and Biochemistry*, 9: 167–172.
- Everett, K. R., 1976: A survey of the soils in the region of the South Shetland Islands and adjacent part of the Antarctic Peninsula. Columbus: Ohio State University, Institute of Polar Studies, Report No. 58.
- Ferron, F. A., Simöes, C., Aquino, F. E., and Setzer, A. W., 2004: Air temperature time series for King George Island, Antarctica. *Pesquisa Anta´rtica Brasileira*, 4: 155–169.
- Gajananda, K., 2007: Soil organic carbon and microbial activity: East Antarctica. *European Journal of Soil Science*, 58: 704–713.
- Gerke, J., and Hermann, R., 1992: Adsorption of orthophosphate to humic-Fe-complexes and to amorphous Fe-oxide. Zeitschrift für *Pflanzenerna¨hr Bodenkunde*, 155: 233–236.
- Gooseff, M. N., Barrett, J. E., Doran, P. T., Fountain, A. G., Lyons, W. B., Parsons, A. N., Porazinska, D. L., Virginia, R. A., and Wall, D. H., 2003: Snow-patch influence on soil biogeochemical processes and invertebrate distribution in the McMurdo Dry Valleys, Antarctica. *Arctic, Antarctic, and Alpine Research*, 35: 91–99.
- Green, V. S., Stott, D. E., and Diack, M., 2006: Assay for fluorescein diacetate hydrolytic activity: optimization for soil samples. *Soil Biology and Biochemistry*, 38: 693–701.
- Hargrove, W. W., and Pickering, J., 1992: Pseudoreplication: a *sine qua non* for regional ecology. *Landscape Ecology*, 6: 251–258.
- Hedley, M. J., Stewart, J. W. B., and Chauhan, B. S., 1982: Changes in inorganic and organic soil phosphorus fractions induced by cultivation practices and by laboratory incubations. *Soil Science Society of America Journal*, 41: 970–976.
- Hinsinger, P., 2001: Bioavailability of soil inorganic P in the rhizosphere as affected by root-induced chemical changes: a review. *Plant and Soil*, 237: 173–195.
- Hogg, I. D., Cary, S. C., Convey, P., Newsham, K. K., O'Donnell, A. G., Adams, B. J., Aislabie, J., Frati, F., Stevens, M. I., and Wall, D. H., 2006: Biotic interactions in Antarctic terrestrial ecosystems: Are they a factor?. *Soil Biology and Biochemistry*, 38: 3035–3040.
- Hopkins, D. W., Sparrow, A. D., Elberling, B., Gregorich, E. G., Novis, P. M., Greenfield, L. G., and Tilston, E. L., 2006a: Carbon, nitrogen and temperature controls on microbial activity in soils from an Antarctic dry valley. *Soil Biology and Biochemistry*, 38: 3130–3140.
- Hopkins, D. W., Sparrow, A. D., Novis, P. M., Gregorich, E. G., Elberling, B., and Greenfield, L. G., 2006b: Controls on the distribution of productivity and organic resources in Antarctic Dry Valley soils. *Proceedings of the Royal Society*, B273: 2687–2695.
- Hopkins, D. W., Sparrow, A. D., Shillam, L. L., English, L. C., Dennis, P. G., Novis, P., Elberlin, G. B., Gregorich, E. G., and Greenfield, L. G., 2008: Enzymatic activities and microbial communities in an Antarctica dry valley soil: response to C and N supplementation. *Soil Biology and Biochemistry*, 40: 2130–2136.
- Hsu, H., 1977: Aluminum oxides and oxyhydroxides. *In* Dixon, J. B., and Weed, S. B. (eds.), *Minerals in Soil Environments*. Madison, Wisconsin: Soil Science Society of America, 99–143.
- Huang, Q., Shindo, H., and Goh, T. B., 1995: Adsorption, activities and kinetics of acid phosphatase as influenced by montmorillonite with different interlayer minerals. *Soil Science*, 159: 271–278.
- Jenny, H., 1941: *Factors of Soil Formation*. New York: McGraw-Hill. Kelly, E. F., and Yonker, C. M., 2005: Factors of soil formation/time.
- *In* Hillel, D., Rosenzweig, C., Powlson, D., Scot, K., Singer, M., and Sparks, D. (eds.), *Encyclopedia of Soils in the Environment*. New York: Columbia University Press, 536–539.
- Kovach, W. L., 1998: MVSP—A multivariate statistical package for Windows, version 3.0. Pentraeth, U.K.: Kovach Computing.
- Kuzmann, E., Schuch, L. A., Garg, V. K., De Souza, P. A., Jr., Guimarães, E. M., De Oliveira, A. C., and Vértes, A., 1998: Maritime Antarctica soils studied by Mössbauer spectroscopy and other methods. *Brazilian Journal of Physics*, 28: 434–443.
- Lee, Y. I., Lim, H. S., and Yoon, H. I., 2004: Geochemistry of soils of King George Island, South Shetland Islands, West Antarctica: implications for pedogenesis in cold polar regions. *Geochimica et Cosmochimica Acta*, 68: 4319–4333.
- Lillo, A., Ramírez, H., Reyes, F., Ojeda, N., and Alvear, M., 2011: Actividad biológica del suelo de bosque templado en un transecto altitudinal, parque nacional Conguillío (38°S), Chile. *Bosque*, 32: 46–56.
- López-Martínez, J., Serrano, E., Schmid, T., Mink, S., and Linés, C., 2012: Periglacial processes and landforms in the South Shetland

Islands (Northern Antarctic Peninsula region). *Geomorphology*, 155- 156: 62–79.

- Matar, A., Torrent, J., and Ryan, J., 1992: Soil and fertilizer phosphorus and crop responses in the dryland Mediterranean zone. *Advances in Soil Sciences*, 18: 81–146.
- Matsuoka, N., 1995: Rock weathering processes and landform development in the Sor Rondane Mountains, Antarctica. *Geomorphology*, 12: 323–339.
- McKeague, J. A., and Day, J. H., 1966: Dithionite and oxalate extractable Fe and Al as aids differentiating various classes of soils. *Canadian Journal of Soil Science*, 46: 13–22.
- McKeague, J. A., 1967: An evaluation of 0.1 M pyrophosphate and pyrophosphate-dithionate in comparison with oxalate as extractants of the accumulation products in Podzols and some other soils. *Canadian Journal of Soil Science*, 47: 95–99.
- Mehra, O. P., and Jackson, M. L., 1960: Iron oxide removal from soils and clays by a dithionate-citrate system buffered with sodium carbonate. *Clays and Clay Minerals*, 7: 317–327.
- Motta, L., and Motta, M., 2004: Distribution and pattern of shallow melting at the local glaciers of Terra Nova Bay (Antarctica) coast. *Annals of Glaciology*, 39: 483–489.
- Murphy, J., and Riley, J. P., 1962: A modified single solution method for the determination of phosphate in natural waters. *Analytica Chimica Acta*, 27: 31–36.
- Nannipieri, P., Ascher, J., Ceccherini, M. T., Landi, L., Pietramellara, G., and Renella, G., 2003: Microbial diversity and soil functions. *European Journal of Soil Science*, 54: 655–670.
- Navas, A., López-Martínez, J., Casas, J., Machín, J., Durán, J. J., Serrano, E., Cuchi, J. A., and Mink, S., 2008: Soil characteristics on varying lithological substrates in the South Shetland Islands, maritime Antarctica. *Geoderma*, 144: 123–139.
- Oksanen, L., 2001: Logic of experiments in ecology: is pseudoreplication a pseudoissue? *Oikos*, 94: 27–38.
- Parfitt, R. L., 1978: Anion adsorption by soils and soil materials. *Advances in Agronomy*, 30: 1–50.
- Parfitt, R. L., 1989: Phosphate reactions with natural allophone, ferrihydrite and goethite. *Journal of Soil Science*, 40: 359–369.
- Priscu, J. C., 1995: Phytoplankton nutrient deficiency in lakes of the McMurdo Dry Valleys, Antarctica. *Freshwater Biology*, 34: 215–227.
- Rao, M. A., Gianfreda, L., Palmiero, F. M., and Violante, A., 1996: Interaction of acid phosphatase with clays, organic molecules and organo-mineral complexes. *Soil Science*, 16: 1751–1760.
- Rao, M. A., Violante, A., and Gianfreda, L., 2000: Interaction of acid phosphatase with clays, organic molecules and organo-mineral complexes: kinetics and stability. *Soil Biology and Biochemistry*, 32: 1007–1014.
- Reyes, F., Lillo, A., Ojeda, N., Reyes, M., and Alvear, M., 2011: Efecto de la exposición y la toposecuencia sobre actividades biológicas del suelo en bosque relicto del centro-sur de Chile. *Bosque*, 32: 255–265.
- Schnürer, J., and Rosswall, T., 1982: Fluorescein diacetate hydrolysis as measure of total microbial activity in soil and litter. *Applied Environmental Microbiology*, 43: 1256–1261.
- Schwertmann, U., and Taylor, R. M., 1977: Iron oxides. *In* Dixon, J. B., and Weed, S. B. (eds.), *Minerals in Soil Environments*. Madison, Wisconsin: Soil Science Society of America, 145–180.
- Shindo, H., Watanabe, D., Onaga, T., Urakawa, M., Nakahara, O., and Huang, Q., 2002: Adsorption, activity, and kinetics of acid phosphatase as influenced by selected oxides and clay minerals. *Soil Science and Plant Nutrition*, 48: 763–767.
- Sidari, M., Ronzello, G., Vecchio, G., and Muscolo, A., 2008: Influence

of slope aspects on soil chemical and biochemical properties in a *Pinus laricio* forest ecosystem of Aspromonte (southern Italy). *European Journal of Soil Biology*, 44: 364–372.

- Sinsabaugh, R. L., Lauber, Ch. L., Weintraub, M. N., Ahmed, B., Allison, S. D., Crenshaw, C. H., Contosta, A. R., Cusack, D., Frey, S., Gallo, M. E., Gartner, T. B., Hobbie, S. E., Holland, K., Keeler, B. L., Powers, J. S., Stursova, M., Takacs-Vesbach, C., Waldrop, M. P., Wallenstein, M. D., Zak, D. R., and Zeglin, L. H., 2008: Stoichiometry of soil enzyme activity at global scale. *Ecology Letters*, 11: 1252–1264.
- Soave, G. E., Coria, N. R., Montalti, D., and Curtosi, A., 2000: Breeding flying birds in the region of the Fildes Peninsula, King George Island, South Shetland Islands, Antarctica, 1995/96. *Marine Ornithology*, 28: 37–40.
- Statistica 2001. Statistica for Windows. Tulsa, Oklahoma: StatSoft, Inc.
- Tabatabai, M. A., and Bremner, J. M., 1969. Use of p-nitrophenyl phosphate for assay of soil phosphatase activity. *Soil Biology and Biochemistry*, 1: 301–307.
- Tiessen, H., and Moir, J. O., 1993: Characterization of available P by sequential extraction. *In* Carter, M. R. (ed.), *Soil Sampling and Methods of Analysis*. Canadian Society of Soil Science, special publication, 75–86.
- Tiessen, H., Stewart, J. W. B., and Moir, J. O., 1983: Changes in organic and Pi composition of two grassland soils and their particle size fractions during 60–90 years of cultivation. *Journal of Soil Science*, 34: 815–823.
- Tscherko, D., Bölter, M., Beyer, L., Chen, J., Elster, J., Kandeler, E., Kuhn, D., and Blume, H. P., 2003: Biomass and enzyme activity of two soil transects at King George Island, maritime Antarctica. *Arctic, Antarctic, and Alpine Research*, 35: 34–47.
- Ugolini, F. C., 1963: Soil investigations in the Lower Wright Valley, Antarctica. Permafrost International Conference. National Academy of Sciences, National Research Council Publication 1287: 55–61.
- Ugolini, F. C., and Bockheim, J. G., 2008:. Antarctic soils and soil formation in a changing environment: a review. *Geoderma*, 144: $1 - 8$.
- Ugolini, F. C., and Dahlgren, R. A., 2002: Soil development in volcanic ash. *Global Environmental Research*, 6: 69–81.
- Vitousek, P. M., 1984: Literfall, nutrient cycling, and nutrient limitation in tropical forest. *Ecology*, 65: 285–298.
- Vogt, S., and Braun, M., 2004: Influence of glaciers and snow cover on terrestrial and marine ecosystems as revealed by remotely-sensed data. *Pesquisa Anta´rtica Brasileira*, 4: 105–118.
- Walker, T. W., and Syers, J. K., 1976: The fate of phosphorus during pedogenesis. *Geoderma*, 15: 1–19.
- Wasley, J., Robinson, S. A., Lovelock, C. E., and Popp, M., 2006: Climate change manipulations show Antarctic flora is more strongly affected by elevated nutrients than water. *Global Change Biology*, 12: 1800–1812.
- Wieder, R. K., and Lang, G. E., 1986: Fe, Al, Mn and S chemistry of *Sphagnum* peat in four peatlands with different metal and sulfur input. *Water, Air and Soil Pollution*, 29: 309–320.
- Zeglin, L. H., Sinsabaugh, R. L., Barrett, J. E., Gooseff, M. N., and Takacs-Vesbach, C., 2009: Landscape distribution of microbial activity in the McMurdo Dry Valleys: linked biotic processes, hydrology and geochemistry in a cold desert ecosystem. *Ecosystems*, 12: 562–573.
- Zhao, Y., and Li, T., 1996: The pedogenic groups and diagnostic characteristics in the Fildes Peninsula of King George Island, Antarctica. *Antarctic Research*, 7: 70–78.

MS accepted January 2013