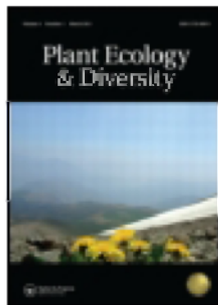


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A simple method for determining limiting nutrients for photosynthetic crusts

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Background: Photosynthetic crust communities are important to the functioning of many desert and early successional ecosystems. Little is known about the factors that limit the growth of these communities, especially during early stages of primary succession or following disturbance.

Aims: Our main goal was to develop a method to study nutrient limitations of crust growth in laboratory microcosms. We used the new method to test the hypothesis that phosphorus limits the growth of crusts in newly deglaciated soils of the high Andes.

Methods: We modified the point–intercept method used in plant ecology to quantify the spread of cyanobacteria, algae and mosses on the soil surface in response to additions of nitrogen and phosphorus.

Results: Fertilization with phosphorus significantly increased the growth rate and final percentage cover, and decreased the lag time for growth of cyanobacterial and algal communities in recently deglaciated soils. By contrast, nitrogen additions had no significant effect on the growth of microbial phototrophs, and all nutrient additions suppressed the growth of early successional mosses.

Conclusions: We propose that the method described here offers a valuable tool for assessing the nature of nutrient limitation of photosynthetic organisms in early successional and desert ecosystems. The information provided by using this approach can increase our understanding of the earliest stages of ecosystem development and may help inform strategies for the reclamation of disturbed arid ecosystems by identifying potential limiting nutrients.

Keywords: Glacial retreat; global warming; limiting nutrients; Peru; photosynthetic crust; primary succession

Introduction

Approximately 40% of the Earth's terrestrial surface is covered by warm or cold deserts (Bowker et al. 2005), both of which often support diminutive photosynthetic communities, variously referred to as cryptogamic crusts and biological soil crusts, among many other names (Belnap 2003; Bowker et al. 2005; Davey and Clarke 1992; Zelikova et al. 2012). These phototrophic communities are especially prevalent in areas that lack vascular plants, where they are able to exploit intermittent pulses of water to photosynthesise and grow (Belnap 2003; Garcia-Pichel and Pringault 2001). Beyond water limitation, little is known about the soil nutrients that might further limit the growth of these communities (Belnap 2003; Bowker et al. 2006); however, there have been correlative studies indicating that micronutrients such as calcium (Ca), manganese (Mn), sulphur (S) and zinc (Zn) may influence the distribution and abundance of some crust-forming organisms (Bowker et al. 2005; Ullmann and Büdel 2001). In addition, some studies indicated that nitrogen (N) additions may adversely affect mosses in desert crusts (Stark et al. 2011) and in montane ecosystems (Pearce et al. 2003), but according to Bowker et al. (2005), no work has directly examined N or phosphorus (P) limitation of crust growth in desert ecosystems. There is also an obvious gap in the literature

concerning experimental evidence for N or P limitation of photosynthetic microbes in soils in an early phase of development; however, based on previous studies of soil heterotrophs in glacial forelands (Göransson et al. 2011, Schmidt et al. 2011b; Yoshitake et al. 2007), it is possible (if not likely) that the lack of available N or P may limit the establishment and growth of biological soil crusts during primary succession.

Most of what is known about nutrient limitation during primary succession is derived from plant ecological studies (Walker and Syers 1976; Chapin et al. 1994; Matthews 1992; Vitousek 2004) that indicated that N most often limits plant productivity during early primary succession (Walker and Syers 1976; Vitousek 2004). Support for this hypothesis comes from the observation that all of the elements (except N) required for plant growth are usually found in geologic substrates on which primary succession occurs (Jenny 1980; Tilman 1988); however, while phosphorus (P) may be present in early successional soils or bedrock, it is often bound in unweathered minerals and is therefore unavailable to organisms (Schlesinger et al. 1998; Walker and Syers 1976). For example, recently deglaciated soils of the high Andes contain relatively high concentrations of P in primary mineral forms ($>600 \mu\text{g g}^{-1}$) but very small stocks of biologically available P that can be accessed

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by heterotrophic microbes ($<1 \mu\text{g g}^{-1}$; Schmidt et al. 2011b). In addition, recent studies of terrestrial microbial communities in early and late successional ecosystems have indicated that P can limit microbial activity and growth (Cleveland et al. 2002; King et al. 2008), but these studies have been confined to the effects of P on heterotrophic processes (e.g. soil respiration). We do not know if phototrophic microbes have access to the same pools of P as heterotrophic microbes (Cleveland and Liptzin 2007; Reiners 1986) or if they are limited by P or N, or both (or neither) early in succession. This shortcoming limits our understanding of the earliest stages of ecosystem succession because cyanobacteria and other microbial phototrophs are among the dominant organisms that colonise recently deglaciated soils (Kaštovská et al. 2005; Nemergut et al. 2007; Schmidt et al. 2011a, Yoshitake et al. 2010).

Here we describe a simple method to determine the effects of added nutrients on photosynthetic microorganisms in early successional soils. Our approach was to microscopically, and non-destructively, monitor the spread of photosynthetic colonies on the soil surface in response to added nutrients in laboratory microcosms. We used theory and adapted methods from plant ecology (Bonham 1989; Jonasson 1988) and soil microbiology (Bowker et al. 2002; Schmidt et al. 2011b) to estimate the percentage cover, growth rates and lag times for development of photosynthetic microorganisms and mosses on the surface of recently deglaciated soils. We used this approach to determine if microbial phototrophs were limited by P or N in recently deglaciated soils in the high Andes of Peru. These early successional soils were just starting to form crust-like communities when collected (Schmidt et al. 2008a, see Materials and methods for details). Based on our previous work showing very low levels of microbe-available P (Schmidt et al. 2011b), and high potential for N fixation at this site (Schmidt et al. 2008a), we hypothesised that P was more limiting than N in this early successional ecosystem.

Materials and methods

Site and organism descriptions

The soils used in this study were collected in the rapidly deglaciating valley of the Sibinacocha Watershed in the Cordillera Vilcanota, Peru ($13^{\circ}46'24''\text{S}$, $71^{\circ}04'17''\text{W}$; Schmidt et al. 2008a, 2008b; Seimon et al. 2007). These early successional soils are extremely oligotrophic and remain mostly devoid of plants (and macroscopic soil crusts) for many years after glacial retreat (Schmidt et al. 2008a). Biological soil crusts slowly form on these soils over a period of ca. 20 to 50 years (Nemergut et al. 2007; Schmidt et al. 2009), but soil darkening is seen in wetter areas within the first four years post-glacial retreat. The soils used in the present study were from sites that had been deglaciated in the previous four years and had not yet formed visible crusts but contained the propagules for crust formation, including a high diversity of cyanobacteria and algae, many of which have not been cultured or described (Nemergut et al. 2007; Schmidt et al.

2008a; Schmidt et al. 2011a). For example, 13 of the 29 cyanobacterial phylotypes in these soils fall into three deeply divergent clades (at the order or higher level) that contain only undescribed cyanobacteria from cold deserts of the Himalayas and Antarctica (supplemental Figure S2 in Schmidt et al. 2011a). The rest of the cyanobacteria in these soils fall into widely divergent groups including the Nostocales, Oscillatoriales and Pseudanabaenales (Schmidt et al. 2011a). The most abundant phylotype is a *Nostoc* sp. (GenBank: GQ306070) related to a free-living strain of *N. sphaeroides* (Lücking et al. 2009). Other abundant phylotypes included close relatives of a Himalayan strain (HQ189014) of *Microcoleus vaginatus* (Schmidt et al. 2011a), a lichen-forming *Nostoc* strain (EF174228) from Chile (Elvebakk et al. 2008), and a *Phormidium*-like isolate (DQ493873) from the high Arctic (Comte et al. 2007). The mosses of these high elevation soils have not been studied previously but they are upright (acrocarpous) and resemble mosses in the Pottiaceae from high volcanoes and Antarctica (Schiavone and Suárez 2009); however, no sporophytes have been observed, nor has molecular work been reported, so positive identification of the mosses is not possible at this time. The crusts that eventually forms along the chronosequence are 'smooth' to 'rugose' to 'rolling' using the classification scheme of Belnap (2003).

Data collection and analysis

Soil samples (top 5 cm of soil) were collected in an area that had been uncovered by ice within the four years prior to collection and were within 100 m of the glacial terminus (Nemergut et al. 2007). Soil samples were frozen overnight (-10°C) in the field (Schmidt et al. 2009) and then frozen to -20°C in Lima, Peru before being transported on ice to the University of Colorado, Boulder. Soils were stored at -20°C and thawed for 24 h prior to the beginning of the experiments and were then thoroughly homogenised and sieved (2.36 mm mesh size). Sieving removed only rocks as no established mosses or other macroscopic biological materials were visible in these early successional soils. The water holding capacity of the homogenised soil was determined as described elsewhere (Colores et al. 1996). Equal weight of soil (17.3 g dry weight) was added to all 'microcosms' (three replicates per treatment) in small sterile Petri plates (60 mm diameter \times 15 mm high, Fisher Scientific 8-757-13A) resulting in a 3–4 mm deep layer of soil. N was added (in the initial watering of the soils) to the +N and +N+P treatments as NH_4NO_3 to obtain a final concentration of $75 \mu\text{g N g}^{-1}$ soil. P was added (in the initial watering) to the +P and +N+P treatments as a mix of dibasic and monobasic sodium phosphate to achieve a final concentration of $75 \mu\text{g P g}^{-1}$ soil. The pH of the phosphate solution was adjusted to the pH of the soil (Nemergut et al. 2007) before addition. The concentrations of N and P were chosen based on the work of King et al. (2008). In order to ensure adequate water and prevent water logging, all soils were initially amended to 70% of water-holding capacity and were watered with sterile-deionised water every three days to 70% of water-holding capacity by weight (Scow

et al. 1989). The arrangement of the plates was randomised after each watering in order to control for any variation in light and temperature.

Microcosms were initially incubated in an incubator (Sheldon Manufacturing Inc., Cornelius (OR)) under conditions mimicking the freeze–thaw cycles that these soils experience in the field (Schmidt et al. 2009), with temperatures dropping to 0 °C at night and rising to just above 25 °C during the day. After some growth of the native organisms (no inoculum was added) was evident in the microcosms (nine days), they were incubated at 22–28 °C on a laboratory bench for the remainder of the experiment. Supplemental lighting was provided by a reflective lamp with a 100 W broadspectrum (including UV) bulb (Zoo Med Laboratories, Inc.). Photoperiod was 15 h of light and 9 h of darkness. Data loggers (Hobo Pendant, Onset Computer Corp., Bourne (MA)) were kept with the microcosms at all times to record both temperature and light data.

Growth of photosynthetic colonies (green mats or filaments) on the soil surface was quantified non-destructively using a microscope (with illumination from above) and a point-intercept approach (Bonham 1989; Jonasson 1988). Once experiments were started, every microcosm was examined every three to four days by running multiple transects across each plate and determining the presence or absence of phototrophs in each field of view until at least 50 fields of view (45× magnification) were examined for each replicate at each sampling time. This method is adequate to estimate the percentage cover of colony-forming algae, moss and cyanobacteria, but does not allow for the quantification of unicellular photosynthetic bacteria (which were not examined in this study). Preliminary tests showed that total percentage cover of phototrophs was proportional to the percentage of microscopic fields containing phototrophs as long as at least 50 microscopic views were recorded for each microcosm at each sampling time. This level of sampling is equivalent to randomly examining 35% of the total surface area of each microcosm at each sampling time. A standard curve was generated from separate microcosms ($n = 12$) to those used in the study, and used to convert the percentage of microscopic fields to percentage cover: % cover = 0.937 (positive microscope fields) ($R^2 = 0.96$). This standard curve is only valid up to the point at which percentage cover is equal to 45% of the total surface area, after which the percentage of microscopic fields cannot be used to accurately estimate percent cover because the relationship between the two variables reached an asymptote. Higher values of percentage cover can be estimated if a higher magnification is used because the size of the microscope field more closely approximates a point in space at each higher level of magnification. Higher levels of magnification were not necessary in the present study because the density of phototrophs was low. Percentage cover for soils with higher density of phototrophs can be determined by using a more tedious version of the point–intercept method in which crosshairs on the ocular lens is used as a true point. Fifty to 100 points need to be examined along random transects of

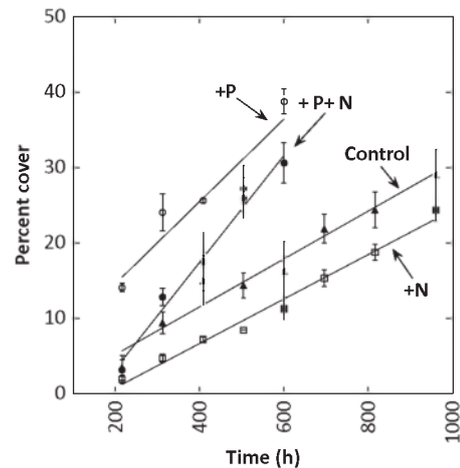


Figure 1. Time course of the growth of microbial phototrophs on the surface of soils from unvegetated areas of the Puca Glacier, Peru forelands.

Note: Each point is the mean of three replicates (error bars are SEM). The soils treated with P showed the most rapid response, with high levels of soil coverage being attained early in the experiment. Lines are the best fit to the linear portion of each data set (after 600 hours the +P and +P+N treatments levelled off at the maximum percentage cover values shown in Figure 4). A two-way ANOVA with interaction and corrected for repeated measures showed high levels of overall significance $P < 0.0001$ and significant differences between all treatments on most dates (Tukey's Honestly Significant Differences test).

each microcosm to obtain reliable estimates of cover, but this procedure was not necessary in the present study.

A two-way ANOVA with interaction and corrected for repeated measures (Devore 2004) was performed on the growth data (see Figure 1) using R (<http://www.r-project.org/index.html>). Percentage cover was modelled by time, and treatment, with an error term for the treatment replicates. Interaction between time and treatment was included in the model. The repeated measures approach was needed because measurements of a single microcosm over time are not independent of each other. Growth rates were estimated by linear regression of growth versus time for the linear portion of the growth curve in each replicate microcosm ($n = 3$) for each treatment. R^2 values for these fits ranged from 0.77 to 0.98. One-way ANOVA were carried out to compare mean growth rates and other calculated parameters (see Figures 2, 3 and 4) as a function of nutrient treatment.

Results

Overall our results showed that P was more limiting than N to soil phototrophs under the experimental conditions used in the present study. The addition of P allowed the growth of phototrophs to begin sooner, grow faster and reach a higher percentage cover than the treatment just receiving N or the control (see Figure 1). The stimulatory effect of P resulted in significantly higher growth rates (see Figure 2; $P < 0.05$) and significantly reduced lag times (see Figure 3; $P < 0.005$) in the +P and +P+N treatments compared to the control and +N treatment. For example, the growth rate was increased by 71% and the lag time was reduced

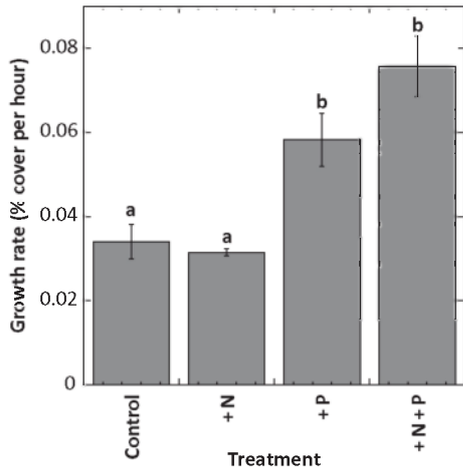


Figure 2. Growth rates (percentage of soil covered per hour) in all treatments and the control.

Note: Each column is the mean of three replicates (error bars are SEM). Columns with different letters were significantly different ($P < 0.05$, one-way ANOVA and Tukey's Honestly Significant Differences test).

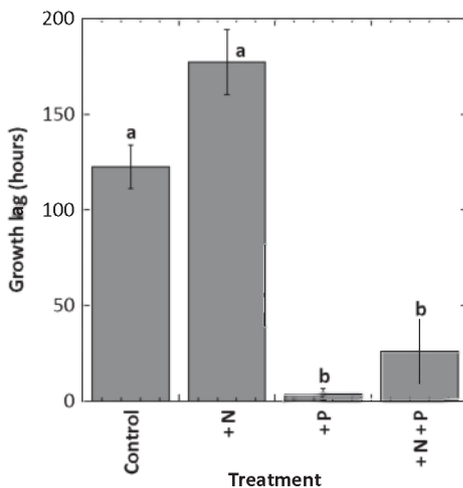


Figure 3. Growth lags (hours before growth started) in all treatments and the control.

Note: Each column is the mean of three replicates (error bars are SEM). Columns with different letters were significantly different ($P < 0.005$, one-way ANOVA and Tukey's Honestly Significant Differences test).

by 97% in the +P treatment compared to the control (see Figures 2 and 3). Likewise, the maximum percentage cover reached during the experiment was significantly higher in microcosms receiving P (see Figure 4; $P < 0.05$), being 31% higher in the +P treatment compared to the control. Our approach also allowed us to monitor the effects of nutrients on larger organisms in the microcosms. Unlike microbial phototrophs, moss growth was inhibited by all nutrient additions compared to the water-only control (see Figure 5).

Discussion

The main goal of this research was to develop a simple method to determine if and what nutrients limit the

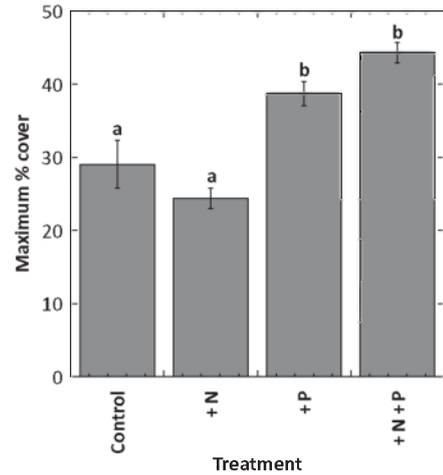


Figure 4. Maximum percentage cover during the experiment.

Note: Each column is the mean of three replicates (error bars are SEM). Columns with different letters were significantly different ($P < 0.05$, one-way ANOVA and Tukey's Honestly Significant Differences test). The highest percentage cover values for the +N+P treatment occurred at time point 960 hours, which was after the linear growth phase (see Figure 1) had stopped.

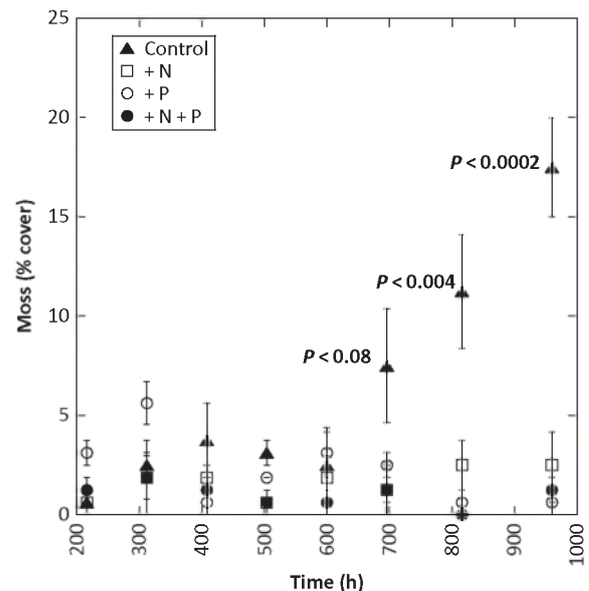


Figure 5. Time course of the growth of moss on the surface of soils from un-vegetated areas of the Puca Glacier forelands (error bars are SEM).

Note: One-way ANOVAs were carried out at each time point; results from Tukey's Honestly Significant Differences tests are shown.

growth of photosynthetic microbes during primary succession in glacier foreland soils. Although we developed the method to examine early successional soils, it could also be applied to any soils that have populations of microbial phototrophs, such as desert crust (e.g. Bowker et al. 2005; Zelikova et al. 2012), or geologically old, unvegetated soils of alpine, Arctic and Antarctic environments (e.g. Freeman et al. 2009; Fell et al. 2006). One advantage of this approach is that it requires minimal equipment, just a dissecting

microscope, a light source, a balance and small Petri plates; therefore experiments similar to those reported here can be carried out at even remote field stations if necessary. Other advantages of this approach are that it is non-destructive and much less cumbersome compared to previous efforts to microscopically monitor the growth of photosynthetic crusts (Bowker et al. 2002).

Despite the simplicity of the approach, the results of this study strongly support our hypothesis that the microbial communities at the study site are limited by the availability of P. Previous work at this site showed potential for P limitation of heterotrophic microbes (King et al. 2008; Schmidt et al. 2011b) but phototrophs were not investigated in those studies, necessitating the development of the approach described here. Our results show that P is limiting to microbial phototrophs in this system; P increased the growth rate, reduced the lag time and increased the maximum percentage cover of phototrophs compared to the treatment receiving just N or the control (see Figures 1, 2, 3 and 4). The obvious linear growth kinetics in the control and +N treatments (see Figure 1) also suggests that P is the primary limiting nutrient in this system. Linear kinetics in microbial systems are indicative of diffusion-limited processes (e.g. Kaplan and Greenberg 1985; Tanzer et al. 1969), which, in this case, is likely to be the diffusion of P through the soil matrix because N addition did not change the linear growth rate, whereas P did (see Figures 1 and 2). To our knowledge the present study is the first to directly show P limitation of phototrophic microbial communities in soil, although P limitation of microbial phototrophs in the oceans has been demonstrated many times (e.g. Martiny et al. 2009; Sanudo-Wilhelmy et al. 2001; Wu et al. 2000).

The results of this study have broader implications for understanding primary succession, at least at high elevation sites, such as those studied here. In our previous work, we speculated that the rate at which plants establish at this site was likely to be constrained by the extreme abiotic environmental conditions at this high elevation (>5000 m a.s.l.) site; however, our findings of very low levels of bioavailable P (Schmidt et al. 2011b) and the limitation of microbial primary production by P (this study) provide strong evidence that P limitation may be an important factor limiting plant colonisation of these areas. Indeed, plant colonisation has occurred at even higher elevations in the same region, but mostly in areas where there is active animal migration and hence faecal deposition (authors' pers. obs., 2011). Another indication of P limitation of this area is the unexpectedly high prevalence of arbuscular mycorrhizal (AM) fungi and root endophytes in plants that are slowly invading this watershed (Schmidt et al. 2008b). AM fungi increase P uptake by plants, but AM propagules are poorly dispersed and are therefore often absent in early successional soils (Miller 1979; Schmidt and Scow 1986). This lack of AM inoculum in newly deglaciated soils could explain the slow rate of plant succession at this site; however, more work is needed to confirm this hypothesis, as well as to explore how common P limitation is for phototrophs in other early successional soils.

Another intriguing finding of the present study is that unlike microbial phototrophs, mosses were inhibited by all of the nutrient addition treatments (see Figure 5). This finding could indicate that the unidentified moss species present in these soils are extremely well adapted to low nutrient conditions and are therefore inhibited by high nutrient availabilities, as has been suggested for lichens from oligotrophic systems (Welch et al. 2006) and for mosses in various ecosystems (Li and Glime 1990; Stark et al. 2011). Mosses in tundra ecosystems have also been reported to respond negatively to additions of both N and P (Chapin and Shaver 1985; Nemergut et al. 2008), but mechanisms for this inhibition have not been determined. The fact that mosses are inhibited by nutrient additions makes it unclear if the present method is useful for determining what, if any, nutrients are limiting to their growth. Future work with mosses should include using lower concentrations of nutrient or different chemical species to ensure that inhibition by particular ions is not masking any potential growth stimulation that could occur from nutrient additions.

Finally, this work highlights the importance of carrying out basic, manipulative experiments in order to gain a preliminary understanding of the roles of microbial communities in a broader ecological context. Our finding of P limitation in microcosms of early successional soils is just a first step towards understanding what limits microbial growth in the field and some of our future effort will be directed at studying this phenomenon in the field at the remote Andean sites where these soils originated.

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References

- Belnap J. 2003. The world at your feet: desert biological soil crusts. *Frontiers in Ecology and the Environment* 1:181–189.
- Bonham CD. 1989. *Measurement for terrestrial vegetation*. New York (NY): Chapman and Hall.
- Bowker MA, Belnap J, Davidson DW, Goldstein H. 2006. Correlates of biological soil crust distribution across a continuum of spatial scales: support for a hierarchical conceptual model. *Journal of Applied Ecology* 43:152–163.
- Bowker MA, Belnap J, Davidson DW, Phillips SL. 2005. Evidence for micronutrient limitation of biological soil crusts: Importance to arid-lands restoration. *Ecological Applications* 15:1941–195.
- Bowker MA, Reed SC, Belnap J, Phillips SL. 2002. Temporal variation in community composition, pigmentation, and Fv/Fm of desert cyanobacterial soil crusts. *Microbial Ecology* 43:13–25.
- Chapin FS, Shaver GR. 1985. Individualistic growth response of tundra plant species to environmental manipulations in the field. *Ecology* 66:564–576.
- Chapin FS, Walker LR, Fastie CL, Sharman LC. 1994. Mechanisms of primary succession following deglaciation at Glacier Bay, Alaska. *Ecological Monographs* 64:149–175.
- Cleveland CC, Liptzin D. 2007. C:N:P stoichiometry in soil: is there a “Redfield ratio” for the microbial biomass? *Biogeochemistry* 85:235–252.
- Cleveland CC, Townsend AR, Schmidt SK. 2002. Phosphorus limitation of microbial processes in moist tropical forests: evidence from short-term laboratory incubations and field studies. *Ecosystems* 5:680–691.
- Colores GM, Schmidt SK, Fisk MC. 1996. Estimating the biomass of microbial functional groups using rates of growth-related soil respiration. *Soil Biology and Biochemistry* 28:1569–1577.
- Comte K, Sabacka M, Carre-Mlouka A, Elster J, Komarek J. 2007. Relationships between the Arctic and the Antarctic cyanobacteria; three *Phormidium*-like strains evaluated by a polyphasic approach. *FEMS Microbiology Ecology* 59:366–376.
- Davey MC, Clarke KJ. 1992. Fine structure of a terrestrial cyanobacterial mat from Antarctica. *Journal of Phycology* 28:199–202.
- Devore JL. 2004. *Probability and statistics for engineering and the sciences*. 6th ed. Belmont (CA): Thomson, Brooks/Cole.
- Elvebakk A, Papaefthimiou D, Robertsen EH, Liaimer A. 2008. Phylogenetic patterns among *Nostoc* cyanobionts within bi- and tripartite lichens of the genus *Pannaria*. *Journal of Phycology* 44:1049–1059.
- Fell JW, Scorzetti G, Connell L, Cary SC. 2006. Biodiversity of micro-eukaryotes in Antarctic Dry Valley soils with <5% soil moisture. *Soil Biology and Biochemistry* 38:3107–3119.
- Freeman KR, Pescador MY, Reed SC, Costello EK, Robeson MS, Schmidt SK. 2009. Soil CO₂ flux and photoautotrophic community composition in high-elevation, ‘barren’ soils. *Environmental Microbiology* 11:674–686.
- Garcia-Pichel F, Pringault O. 2001. Cyanobacteria track water in desert soils. *Nature* 413:380–381.
- Göransson H, Venterink HO, Bååth E. 2011. Soil bacterial growth and nutrient limitation along a chronosequence from a glacier forefield. *Soil Biology and Biochemistry* 43:1333–1340.
- Jenny H. 1980. *Soil genesis with ecological perspectives*. New York (NY): Springer Verlag.
- Jonasson S. 1988. Evaluation of the point intercept method for the estimation of plant biomass. *Oikos*, 52:101–106.
- Kaplan HB, Greenberg EP. 1985. Diffusion of autoinducer is involved in regulation of the *Vibrio fischeri* luminescence system. *Journal of Bacteriology* 163:1210–1214.
- Kaštovská K, Elster J, Stibal M, Santrůčková H. 2005. Microbial assemblages in soil microbial succession after glacial retreat in Svalbard (High Arctic). *Microbial Ecology* 50:396–407.
- King AJ, Meyer AF, Schmidt SK. 2008. High levels of microbial biomass and activity in unvegetated tropical and temperate alpine soils. *Soil Biology Biochemistry* 40:2605–2610.
- Lücking R, Lawrey JD, Sikaroodi M, Gillevet PM, Chaves JL, Sipman HJM, Bungatz F. 2009. Do lichens domesticate photobionts like farmers domesticate crops? Evidence from a previously unrecognized lineage of filamentous cyanobacteria. *American Journal of Botany* 96:1409–1418.
- Martiny AC, Huang Y, Li W. 2009. Occurrence of phosphate acquisition genes in *Prochlorococcus* cells from different ocean regions. *Environmental Microbiology* 11:1340–1347.
- Matthews JA. 1992. *The ecology of recently deglaciated terrain: a geoecological approach to glacier forelands and primary succession*. Cambridge (UK): Cambridge University Press.
- Miller RM. 1979. Some occurrences of vesicular–arbuscular mycorrhiza in natural and disturbed ecosystems of the Red Desert. *Canadian Journal of Botany* 57:619–623.
- Nemergut DR, Anderson SP, Cleveland CC, Martin AP, Miller AE, Seimon A and Schmidt SK. 2007. Microbial community succession in unvegetated, recently deglaciated soils. *Microbial Ecology* 53:110–122.
- Nemergut DR, Townsend AR, Sattin SR, Fierer N, Neff JC, Bowman WD, Schadt CW, Weintraub M, Schmidt SK. 2008. The effects of chronic nitrogen fertilization on alpine tundra soil microbial communities: implications for carbon and nitrogen cycling. *Environmental Microbiology* 10:3093–3105.
- Pearce ISK, Woodin SJ, van der Wal R. 2003. Physiological and growth responses of the montane bryophyte *Racomitrium lanuginosum* to atmospheric nitrogen deposition. *New Phytologist* 160:145–155.
- Reiners WA. 1986. Complementary models for ecosystems. *American Naturalist* 127:59–73.
- Sanudo-Wilhelmy SA, Kustka AB, Gobler CJ, Hutchins DA, Yang M., Lwiza K, Burns J, Capone DG, Raven JA, Carpenter EJ. 2001. Phosphorus limitation of nitrogen fixation by *Trichodesmium* in the central Atlantic Ocean. *Nature* 411:66–69.
- Schiavone MM, Suárez GM. 2009. *Globulinella halloyi* (Pottiaceae), a new species from Argentina. *The Bryologist* 112:584–588.
- Schlesinger WH, Brijnzel LA, Bush MB, Klein EM, Mace KA, Raikes JA, Whittaker RJ. 1998. The biogeochemistry of phosphorus after the first century of soil development in Rakata Island, Krakatau, Indonesia. *Biogeochemistry* 40:37–55.
- Schmidt SK, Lynch RC, King AJ, Karki D, Robeson MS, Nagy L, Williams MW, Mitter MS, Freeman KR. 2011a. Phylogeography of microbial phototrophs in the dry valleys of the high Himalayas and Antarctica. *Proceedings of the Royal Society B* 278:702–708.
- Schmidt SK, Nemergut DR, Miller AE, Freeman KR, King AJ, Seimon A. 2009. Microbial activity and diversity during extreme freeze–thaw cycles in periglacial soils, 5400 m elevation, Cordillera Vilcanota, Perú. *Extremophiles* 13:807–816.
- Schmidt SK, Nemergut DR, Sowell P, Reed SC, Cleveland CC. 2011b. Estimating phosphorus availability for microbial growth in an emerging landscape. *Geoderma* 163:135–140.
- Schmidt SK, Reed SC, Nemergut DR, Grandy AS, Cleveland CC, Costello EK, Weintraub MN, Hill AW, Meyer AF, Martin AP, et al. 2008a. The earliest stages of ecosystem succession in high-elevation (5000 meters above sea level), recently deglaciated soils. *Proceedings of the Royal Society B* 275:2793–2802.

- Schmidt SK, Scow KM. 1986. Mycorrhizal fungi on the Galápagos Islands. *Biotropica* 18:236–240.
- Schmidt, SK, Sobieniak-Wiseman LC, Kageyama SA, Halloy SRP, Schadt CW. 2008b. Mycorrhizal and dark-septate fungi in plant roots above 4270 meters elevation in the Andes and Rocky Mountains. *Arctic, Antarctic and Alpine Research* 40:576–583.
- Scow KM, Schmidt SK, Alexander M. 1989. Kinetics of biodegradation of mixtures of substrates in soil. *Soil Biology and Biochemistry* 21:703–708.
- Seimon TA, Seimon A, Daszak P, Halloy SRP, Schloegel L, Aguilar C, Sowell P, Hyatt A, Konecky B, Simmons J. 2007. Upward range extension of Andean anurans and chytridiomycosis to extreme elevations in response to deglaciation. *Global Change Biology* 13: 288–299.
- Stark LR, Brinda, JC, McLetchie DN. 2011. Effects of increased summer precipitation and N deposition on Mojave Desert populations of the biological crust moss *Syntrichia caninervis*. *Journal of Arid Environments* 75: 457–463.
- Tanzer JM, Wood WI, Krichevsky MI. 1969. Linear growth kinetics of plaque-forming streptococci in the presence of sucrose. *Journal of General Microbiology* 58: 125–133
- Tilman D. 1988. *Plant strategies and the dynamics and structure of plant communities*. Princeton (NJ): Princeton University Press.
- Ullmann I, Büdel B. 2001. Ecological determinants of species composition of biological soil crusts on a landscape scale. In: Belnap J, Lange OL, editors. *Biological soil crusts: structure, function, and management*. Berlin (Germany): Springer Verlag. P. 203–213.
- Vitousek PM. 2004. *Nutrient cycling and limitation: Hawaii as a model system*. Princeton (NJ): Princeton University Press.
- Walker TW, Syers JK. 1976. The fate of phosphorus during pedogenesis. *Geoderma* 15:1–19.
- Welch AR, Gillman MP, John EA. 2006. Effects of nutrient application on growth rate and competitive ability of three foliose lichen species. *The Lichenologist* 38:177–186.
- Wu J, Sunda W, Boyle EA, Karl DM. 2000. Phosphate depletion in the Western North Atlantic Ocean. *Science* 289:759–762.
- Yoshitake S, Uchida M, Koizumi H, Kanda H, Nakatsubo T. 2010. Production of biological soil crusts in the early stage of primary succession on a high Arctic glacier foreland. *New Phytologist* 186:451–460.
- Yoshitake S, Uchida M, Koizumi H, Nakatsubo T. 2007. Carbon and nitrogen limitation of soil microbial respiration in high Arctic successional glacier foreland near Ny-Ålesund, Svalbard. *Polar Research* 26:22–30.
- Zelikova TJ, Housman DC, Grote EE, Neher DA, Belnap J. 2012. Warming and increased precipitation frequency on the Colorado Plateau: implications for biological soil crusts and soil processes. *Plant and Soil* 355:265–282.