Temperature effects on the first three years of soil ecosystem development on volcanic ash.

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ABSTRACT

Little is known of the earliest stages of soil ecosystem development on volcanic ash, and how this process is affected by temperature. We studied the first three years of soil development in a field-based mesocosm experiment, situated in different climates across Japan. Newly fallen, sterilized volcanic ash from the Sakurajima volcano (Kyushu, Japan) was placed into pots and positioned at six locations with mean annual temperatures ranging from -1.6 °C to 18.6 °C. At 24 months into the experiment, C and N accumulation showed only a weak linear correlation with temperature, but by 36 months there was a clear exponential relationship. This applied only to the top 2 cm of the developing soil, and was not apparent in the lower part of the ash. We suggest that this acceleration in warmer climates relates to a positive feedback involving bryophyte cover, which had become much denser by the third year in the warmer sites. Surprisingly, the abundance of 16S rRNA gene copies of bacteria, fungi, archaea - as well as ammonia oxidizers – did not increase from 12 months to 36 months, and did not show any relationship to temperature, suggesting that input from plants is the major factor in increasing C and N buildup in the soil. Overall it appears that temperature effects on bryophyte cover buildup may be important in controlling the temperature relationship in soil development on volcanic ash.

Keywords: Carbon accumulation, Nitrification, Primary succession, Temperature gradient, Weathering, Volcanic ash
1. Introduction

The process of soil development in primary successional environments has long been a major theme in ecology and geology (Clements, 1916; Huggett, 1998). As well as studies on soil development in debris flows (Turk et al., 2008, 2009), sand dunes (Lichter, 1998), and glacier forelands (Kaye et al., 2003; Mavris et al., 2010), there has been a considerable amount of work on volcanic primary succession – including both lava flows and volcanic ash deposits (Vitousek et al., 1993; Vitousek and Farrington, 1997; Kato et al., 2005). There have been studies on soil development in ash deposits that at the start of the study were already several years old (Fujimura, et al. 2012, 2016), or older (Ohta et al. 2003; Ibekwe et al. 2007; Zeglin et al., 2016), but there has been little work on the very earliest stages of soil ecosystem development. The first three years or so of soil ecosystem development have barely been studied, perhaps due to the difficulties in gaining access to recent volcanic eruption sites.

It is unclear to what extent the major living components of a soil system – bacteria, archaea and fungi, including important functional groups for biogeochemical processes such as ammonia oxidizers – are present in these earliest stages. Studies on volcanic ash deposits that were already several years old have suggested that early stages in volcanic ash soil development are carried out by chemoautotrophs (King, 2003; Fujimura et al., 2012, 2016). For example, Fujimura et al. (2012, 2016) reported the importance of Fe(II)- and H₂-oxidizing chemolithotrophs in a 3.5-6.6 years old ash deposit in Miyake Island, Japan, in initiating the accumulation of organic carbon (C) and then contributing to the development of subsequent microbial communities. Furthermore, Freeman et al. (2009) pointed out the importance of photoautotrophic bacteria (cyanobacteria) in high-elevation barren soils. Only later were larger organisms such as bryophytes seen as having a major role (O'Toole and
It is also unclear what relationship, if any, the process of soil ecosystem development shows to climate - especially temperature. Temperature is seen as a major controlling factor in ecosystem establishment or recovery (Pastor and Post, 1986; Vitousek and Farrington, 1997) as well as pedogenesis (Tsai et al., 2010), but it is unclear how early in the process it becomes important. In this study we were interested in experimentally comparing the effect of temperature on soil ecosystem development from volcanic ash, over the first three years of primary succession. This study was a follow-on from Kerfahi et al. (2017), which discussed the first two years of experimental soil ecosystem development. As we will report here, a pattern was obtained when Year 3 was added to the dataset, with implications for understanding the processes of ecosystem establishment on volcanic ash.

Furthermore, we report here on the inorganic nitrogen (N) dynamics at very early stage of soil development. Nitrogen is not normally present in parent materials such as rock, lava and volcanic ash in certain kinds of metasedimentary and metavolcanic rocks (Holloway et al., 1998). N deposition from the atmosphere is thought to be the primary source in the earliest stages of microbial community development on volcanic ash, while microbial N fixation eventually becomes more important.

Our main working hypothesis here was that there would be a temperature effect on the rate of soil ecosystem development (organic C, organic N, inorganic N, abundance of each of the major microbial groups) would become apparent by Year 3 (36 months) of the experiment. Given the large range of climates studied in this experiment, it was surprising to us that no strong temperature effect on C, N or microbial biomass emerged in the first two years to 24 months (Kerfahi et al., 2017). We anticipated that as the soil ecosystem developed, including development of a bryophyte cover, this effect would eventually show itself more clearly because of the temperature sensitivity of plant growth and metabolism in producing
exudates, dead organic material, etc. that would enrich the soil in C and N. So far, there has been little evaluation of the effect of temperature on the earliest stages of ecosystem development on volcanic substrates, under standardized conditions. While a strong temperature effect would certainly be expected on basic biological and ecological principles, it is important to test such assumptions if ecology is to be rigorously based.

2. Materials and methods

2.1. Source of the volcanic ash

Mt. Sakurajima (31°35′N, 132°39′E, height 1,117 m) is an active volcano located in southern Kyushu Island, Japan. Mt. Sakurajima is situated in the Aira caldera created by catastrophic eruption of around 29,000 years BP (Kobayashi et al., 2013). Large eruptions have periodically occurred since then, interspersed with less active phases. In the last thousand years, three large eruptions occurred in the Bunmei era (1471–1476), An-ei era (1779–1782), and Taisho era (1914–1915) (Biass et al., 2017) – mainly involving lava flowing from upper parts of the volcano and covering parts of the lower slopes. Since 2006, a new phase and different type of volcanic activity started, involving small explosions and ash deposition over the surrounding slopes over the volcano. These have become more active over time (Iguchi et al., 2013), and presently thousands of small eruptions occur annually (Miwa et al., 2013). The volcanic ash of Mt. Sakurajima is characterized as slightly acidic, of relatively low redox potential, and enriched in ions such as Si, Na, Cl, and SO$_4$ (Kawano and Tomita, 2001). Detail of mineral composition of volcanic ash were summarized in Hillman et al. (2012) and Miwa et al. (2013).
2.2. Experimental sites

Six experimental sites were used to position pots of volcanic ash in trays. Six locations across Japan whose mean annual temperature ranged from -2.6 to 18.6°C (Table 1). We used recently deposited ash from Mt. Sakurajima for the experimental pot microcosms, as described below.

i) The Sakurajima site (SJ) is in the warm temperate zone of southern Japan, with a mean annual temperature of 18.6°C (1981-2010) and a mean annual precipitation of 2265.7 mm (1981-2010) according to Kagoshima Meteorological Station, Japan (31°33’N, 130°33’E, 4.0 m a.s.l). The areas sampled were on the lower slopes of the volcano, around 25-50 m above sea level. The site was situated in an open area in the native pine scrub.

ii) The Takakuma site (TK) is in a warm temperate forest site in the surrounding hills near the Sakurajima Volcano, in the Takakuma Experimental Forest of Kagoshima University, southern Kyushu, Japan (31°31’N, 132°46’E, 538 m a.s.l). The site is about 10 km from the crater of Sakurajima with mean annual precipitation is 3410 mm and a mean annual temperature of 14.0°C (1999–2004). The site was situated in an open area by a forest road.

iii) The Kyoto site (KY) is in a warm temperate forest site about 600 km northeast of Sakurajima at Kamigamo Experimental Station, Kyoto University near Kyoto City, Japan (35°04’N, 135°46’E, 140 m a.s.l). Mean annual temperature is 14.6 °C and annual precipitation is 1,538.6 mm (Kamigamo Experimental Station in 1981-2010). The samples were positioned on an open lawn next to the site weather station.

iv, v, vi) Three sites were set at different elevations on Mt. Norikura (summit 3026 m a.s.l.) in central Japan, located 800 km northeast of Sakurajima. The elevations of the three experimental sites on Norikura were 650m, 1450m and 2800m a.s.l. (described here as NK-650m, NK-1450m, and NK-2800m). There were no nearby meteorological stations of
each site for Norikura, but we calculated MAT based on the nearest meteorological stations on the lower slopes and near summit of Mt. Norikura. Assuming a mean lapse rate is 0.6°C/100 m, MAT of NK-650 m 1450 m and 2800 m can be calculated 12.1 °C, 7 °C and −1.6 °C, respectively (Kerfahi et al., 2017).

For comparison of C and N contents of mature developed soils in Japan with the volcanic ash in the experimental pots at weight basis (g kg⁻¹), we used the published data set from 39 sites of well-developed vegetated soils throughout the Japanese archipelago covering 44°20'N to 26°50'N (Urakawa et al. 2015).

2.3. Setting up in-situ incubation experiment

We collected newly fallen volcanic ash on plastic sheeting in an open space (an unused parking lot) near the Sakurajima volcano, over a period of several weeks on March 2012. The freshly accumulated ash was removed and stored every week. The freshly accumulated ash samples were sieved at 2 mm mesh screen to remove large particles before use. The ash was sterilized by heating in portions to 200 °C in a dry oven for 1 h. The ash was handled with sterile gloves.

We put 200 g portions of the dried ash in poly-vinyl chloride (PVC) columns (67 mm in internal diameter and 150 cm³ in internal volume) with a plastic fine nylon mesh underneath, to avoiding the ash from falling out but allowing water drainage. For estimating leaching of anions and cations from ash columns, we modified the resin core method (Binkley et al., 1986; Shibata et al., 2011; Urakawa et al., 2014). We attached PVC columns (internal diameter is same as ash column) filled with 50 g ion exchange resin (Amberlite MB-1; Organo, Japan) and sealed the base of the pot by nylon mesh.

The combined ash and resin cores were connected tightly using vinyl tape. We put
15 pots with 5 replicates for each year of sampling in each site. Pots were placed together in a plastic basket on corrugated plastic plates to allow drainage. The whole basket was covered with a 1 mm white nylon mesh net to exclude windblown materials, seeds, animals, excessively hard rain and extreme heat in direct sunlight. The 1mm pore size was intended to be large enough to allow microbes and bryophyte propagules to enter. At each site, the tray was located in a flat open area in full sun.

We set sample trays at each site in late June or early July 2012. After one, two and three years (12, 24 and 36 months), we randomly took five replicate pots from each site for sampling. For each sample pot, the surface 2 cm of the ash and remaining ash beneath it were taken separately, for DNA extraction and chemical analysis using a sterilized spatula. We also collected the ion exchange resin from beneath each pot, for chemical analysis of captured ions. The samples were transported to the laboratory at 4°C and any organic debris removed by sterilized spatula and tweezers. Samples were then frozen at -20 °C for further DNA extraction and inorganic N (NH$_4^+$-N and NO$_3^-$-N ) extraction, and remaining ash samples were dried at 105 °C for further chemical analyses. DNA extraction and soil analysis were carried out as detailed below. Details of the experiment were previously reported in Kerfahi et al. (2017).

2.4. Chemical measurements

Extraction and measurement of the NH$_4^+$-N and NO$_3^-$-N concentrations followed Urakawa et al. (2014) with some modifications. Briefly, a 5g subsample of each soil sample was extracted in 50 ml of 2 M KCl and filtered to determine the NH$_4^+$-N and NO$_3^-$-N concentrations. Briefly a 5 g sample of the ion exchange resin was extracted in 50 ml of 1 M KCl twice. Also, NH$_4^+$-N and NO$_3^-$-N concentrations in ash and ion exchange resin extracts
were measured colorimetrically.

Ash samples were dried to a constant weight at 105°C and then weighed. Total C and N concentration of ash samples were determined using an NC analyzer (NC-900; Shimadzu, Kyoto, Japan). We measured pH of ash samples using pH meters (D-51, Horiba, Kyoto, Japan) after extraction from 10 g of dry ash with 25 ml of ion exchange water.

2.5. DNA extraction and Quantitative PCR analysis (qPCR)

DNA was extracted from about 0.3 g volcanic ash samples using the MoBio Powersoil DNA Isolation Kit (MoBio, Carlsbad, CA, USA) according to the manufacturer’s instructions and were stored in a freezer at −20 °C. We quantified the gene abundance of bacterial 16S rRNA, archaeal 16S rRNA, fungal ITS, ammonium-oxidizing bacterial amoA, and ammonium-oxidizing archaeal amoA, real-time quantitative polymerase chain reaction (qPCR) was performed using the Light Cycler 96 (Roche Diagnostics K.K., Mannheim, Germany). Bacterial 16S rRNA, archaeal 16S rRNA, fungal ITS region, ammonium-oxidizing bacterial amoA, and ammonium-oxidizing archaeal amoA were determined using the universal primer sets, 338f (Amann et al., 1990)/518r (Muyzer et al., 1993), 109f (Großkopf et al., 1998)/344r (Raskin et al., 1994), ITS1F_KYO2 /ITS2_KYO2 (Toju et al., 2012), amoA 1F/amoA 2R (Rotthauwe et al., 1997), and CrenamoA 23F/CrenamoA 616R (Tourna et al., 2008), respectively.

The qPCR was performed in 10 µL of final volume containing 5 µL master mix 2x from the Faster Essential DNA Green Master Kit (Roche Diagnostics), 1.5 µL of PCR grade H₂O in the kit, 0.5 µL each of forward and reverse primers (10 µM), and 2.5 µL of template DNA (1/10). The conditions of reaction for bacterial 16S rRNA, archaeal 16S rRNA and fungal ITS genes were as follows: an initial denaturation at 95°C for 10 min, then more than
The conditions of reaction for the bacterial amoA gene were as follows: an initial denaturation at 95°C for 10 min, followed by more than 35 cycles of 95°C for 30 s, 55°C for 30 s, and 72°C for 30 s, followed, with the annealing temperature adjusted to 55°C (Okano et al., 2004). All of the assays were conducted melting curve analysis. The DNA quantity in the standard clone plasmid for each gene was determined using a Qubit 2.0 Fluorometer (Thermo Fisher Scientific, Waltham, MA, USA). Relative quantification of genes was conducted as serial dilution standards and calculated using Light Cycler 96 Software version 1.0 (Roche Diagnostics).

2.6. Bryophyte cover

We estimated visually percentage bryophyte cover on the tops of the pots using photographs. For the three Norikura sites at the 12 months and 24 months stage there was no observed bryophyte cover at the time of sampling. We divided each photograph into eight portions radially, and estimated percentage of bryophyte cover for each 12.5% interval. We divided the pot surface into 24 portions if percentage of cover were less than 12.5% or more than 87.5%.

2.7. Statistical analysis

We tested three models (linear, quadratic, and exponential) to describe relationships between MAT and ash C and N accumulation rates. Model selection was carried out based on adjusted $R^2$ and root mean square error (RMSE). Significance level was defined as $< 5\%$.

All statistical analyses were conducted using IBM SPSS Statistics (IBM SPSS 22.0, IBM
3. Results

3.1. Changes in soil organic matter accumulation

Both C and N accumulation rate and content were not significantly correlated with MAT for the 12 month and 24 months stage, except for N at 24 months stage (Fig. 1). However, by the 36 month stage C and N accumulation rate and total C and N increased exponentially with MAT (Fig. 1), in the top 2cm. These patterns were more obvious in the upper 2cm layer, where C and N accumulation rates were much faster than in the lower layer of ash (Fig. S1 and S2).

3.2. Bryophyte cover

Percentage of bryophyte cover was summarized in Table 2. At the 12 month stage, there was no bryophyte cover in almost all of the pots. Only a few very small bryophyte plants (Polytrichum sp.) were observed in SJ and KY sites by this stage. By 24 months, percentage bryophyte cover had increased in the warmest three sites, but still no bryophyte cover was observed in cooler three Norikura sites. An incomplete surface covering of bryophytes (Polytrichum sp. and Plagiochila sp.), small cyanobacterial mats (Nostoc sp.) and small gray foliose lichens (tentatively identified as Parmelia sp.) were observed. By 36 months, percentage bryophyte cover had further increased in warmer sites (Table 2 and Fig. S3). In the two warmest sites, TK and SJ, the percentage bryophyte cover exceeded more than 90 %, while coolest two sites still had no apparent bryophyte cover. Almost complete surface
covering of bryophytes (*Polytrichum* sp. and *Plagiochila* sp.) in SJ and TK, and incomplete but substantial covering of (*Polytrichum* sp.) in KY, and small cover by bryophytes (tentatively identified as *Bryum argentum*) and lichens (*Parmelia* sp.) in NK-650 m were observed. There were significant correlations between percentage bryophyte cover and content of C and N (Fig. 2).

### 3.3. Inorganic N dynamics and pH change

Total pool size of NO$_3^-$-N, NH$_4^+$-N, inorganic N (NO$_3^-$-N + NH$_4^+$-N) did not show clear patterns in relation to MAT and age of ash except for NH$_4^+$-N and inorganic N of 24 months stage (Fig. 3). Pool size of NO$_3^-$-N was far less than NH$_4^+$-N pool, while NH$_4^+$-N were constantly existed even in early soil development (Fig. 3).

Percentage of inorganic N out of total N ranged from 11.6 ± 3.5 to 22.3 ± 6.5 % after 12 months and did not show clear patterns in relation to MAT (Fig. 4). Percentage of inorganic N to total N ranged from 7.1 ± 1.4 to 14.2 ± 2.5 % after 24 months and 4.2 ± 10.9 to 23.1 ± 4.4 % after 36 months stage. There were significant negative correlations between percentage of inorganic N to total N and MAT at the 24 months and 36 months stages.

Leaching of NO$_3^-$-N, NH$_4^+$-N, and inorganic N (NH$_4^+$-N + NO$_3^-$-N) tended to increase with MAT, but decreased again in the warmest site, SJ (Fig. 5). While leaching of NO$_3^-$-N, NH$_4^+$-N, and inorganic N (NO$_3^-$-N + NH$_4^+$-N) did not show any significant relationship with MAT (Fig. 5). The pattern of NO$_3^-$-N leaching was comparable to NH$_4^+$-N leaching despite the pool size of NO$_3^-$-N being far less than the NH$_4^+$-N pool (Fig. 5).

The pH of ash across sites ranged from 4.18 ± 0.09 to 6.10 ± 0.09, 4.66 ± 0.08 to 6.28 ± 0.03, and 4.06 ± 0.93 to 6.47 ± 0.14 after 12, 24, and 36 months stages, respectively.
(Table S1). The pH of ash tended to decreased with MAT however not significantly (Table S1). Changes in pH of the ash did not show any consistent pattern across the sites (Table S1).

3.4. Microbial gene abundance

Bacterial and archaeal 16S rRNA gene abundance and fungal ITS gene abundance did not show clear patterns in relation to the time stage of the experiment except for some exceptions, but in all cases tended to be higher in upper 2cm layer of the pot (Fig. 6).

Bacterial and archaeal amoA gene abundance also did not show any significant clear patterns in relation to time stage, while tending to be higher in upper 2cm (Fig. 7). Abundance of each of these genes increased with MAT, but decreased again in the warmest site SJ for some, such as fungal ITS and the amoA gene of bacteria and archaea (Fig. 6-7).

4. Discussion

4.1. How much soil ecosystem development can occur in 3 years?

In this volcanic ash system, we found a measurable presence of bacteria, archaea and fungi, and archaeal/bacterial ammonia oxidizers, in the first three years of the experiment, and a build up of organic C and N. A previous study (Kerfahi et al., 2017) from the same experiment found that many common soil bacterial groups were already present by 24 months, although there was a high proportion of ‘unclassified’ bacteria – possibility including novel groups peculiar to this environment. However, the overall concentration of C and N in ash samples after 36 months were up to 3.0 gC kg\(^{-1}\) for C and 0.18 gN kg\(^{-1}\) for N, which were still several orders of magnitude less than fully developed forest soils across Japan, which
averaged 92.4, 51.2, 31.9 gC kg\(^{-1}\) in 0-10 cm, 10-30cm and 30-50cm, respectively for C and
from 5.4, 2.9, 1.9 gN kg\(^{-1}\) in 0-10 cm, 10-30cm and 30-50cm, respectively for N (Urakawa et
al. 2015, Fig. S4). Furthermore, bacteria, archaea, and ammonia oxidizer abundance of ash
samples ranged 10\(^6\)-10\(^8\), 10\(^4\)-10\(^6\), and 10\(^2\)-10\(^5\) gene copy per gram soils (Fig. 6, 7) which were
several orders of magnitude less than in developed temperate forest soils reported previously,
i.e. 10\(^7\)-10\(^10\), 10\(^7\)-10\(^9\), and 10\(^5\)-10\(^7\) gene copy per gram soils for bacteria, archaea, and
ammonia oxidizer abundance, respectively (Kemnitz et al., 2007; Isobe et al., 2015, 2018).
While it was not possible to quantify the main source of N entering the system in this
study sites, reported total N deposition across Japan ranged from 0.30 to 1.65 g m\(^{-2}\) y\(^{-1}\)
(Chiwa et al., 2015; Ban et al. 2016), and then either accumulating in the soil, or being
leached and captured in the resin under the pots. In early primary successional systems, the
main source is biological N fixation by cyanolichen (Crews et al., 2001). The fluxes of N in
the first 12 months of the experiment are more likely to reflect the background of
atmospheric deposition, and represent a baseline from which fluxes increase over time during
the experiment. However, the observed fluxes to the resin are complicated by the likely
capture and build up of N in the bryophyte layer (a reservoir which was not included in this
study), and its sequestration in the soil total N pool, preventing N from leaching out of the pot.
By the 36 months stage, some of the warmer climate samples had less N flux to the resin,
possibly reflecting greater N sequestration into growing bryophyte biomass. This is a sign of
the ecosystem becoming more ‘closed’ and efficient during succession (Odum, 1966).
In our sites the amount of NO\(_3^-\)-N leaching was similar to the rate of NH\(_4^+\)-N
leaching from ash pots, which accorded with the pattern observed in total N deposition across
Japan (Chiwa et al., 2015; Ban et al. 2016). Certain part of N deposition may directly reach
out from our experimental pots. As another possible explanation, ammonia oxidizing bacteria
and archaea were always present in pots, suggesting the importance of ammonia oxidizing
activity in early soil ecosystem development and source of $\text{NO}_3^-$-N leaching from pots would
derived from ammonia oxidizers. By contrast, ammonia oxidation and methane oxidation
were found to be negligible at most sites in Hawaian lava flow successional series from about
18 to 300 years old (King, 2003). Soil physical properties of ash deposits could be preferable
environments for ammonia oxidizers compared to those of lava flow, and thus ammonia
oxidizer could be important microbes in early stage of ash soil development.

The proportion of inorganic N relative to total N decreased with increasing MAT, as
well as with time, an indication of the increasing importance of organic N in the developing
soil ecosystem. This organic N would be important source for utilization by microbes as well
as external source such as deposition. Photosynthetic microbes and N fixing microbes may
also be important in the early stages of soil development (Crews et al. 2001). Indeed, Kerfahi
et al. (2017) reported that the relative abundance of Cyanobacteria, Chloroflexi and
Firmicutes, which include photosynthetic and N fixing microbes, were significantly higher in
the same volcanic ash at the 24 months stage than in forest soils. While the energy costs for N
fixing microbes to fix N are far higher than to assimilate N from ammonium, N fixation is
affected by C and N availability of environments (Bottomley and Myrold 2007, Reed et al.
2011). In early stages of soil development, N fixation might be limited more by C availability
than N availability. Further potential studies to investigate this may include metagenome and
metatranscriptome analyses.

It is likely that microbes in the soil play an important role in the transformation of
the original silicate minerals in the ash to clays and other minerals, releasing metal ions (e.g.
$\text{Na}$, $\text{Mg}$, $\text{K}$, and $\text{Ca}$) that can be utilized by the developing plant cover (Kawano and Tomita
2001). The importance of microbes in this process was shown in an earlier laboratory
mesocosm study of Sakurajima volcanic ash, where a sterile system was compared to a
non-sterile one with soil microbes present (Bennett et al. 2001; Kawano and Tomita 2001).
It is unclear what role the development of bryophyte cover itself plays in the
development of the soil ecosystem, but those of our treatments with abundant bryophyte
cover showed evidence of accelerated buildup of N and C (Fig. 2). The role of bryophytes in
soil development in primary succession has been considered previously (O'Toole and Synnott,
1971; Brown and Bates, 1990). While bryophytes do not have a root system as such, they
have hair-like structures (rhizoids) that penetrate the soil, and their turnover supply C and N
to the soil, and dead parts of the photosynthetic thallus are appressed against the soil surface
or fall to it. It is clear that most of the enrichment in C and N is in the top 2 cm layer of our
experimental pots, which has a more rapid accumulation of these two elements – likely due to
the proximity to the bryophyte sources of photosynthetically fixed material. Bryophytes can
survive by taking up N as well as minerals from deposition in rain and by dust, but mineral
availability from substrate is thought to be important to establishment (O'Tool &
Synnott, 1971; Brown and Bates, 1990; Bates and Farmer, 1990). It would be interesting to
investigate whether potentially mutualistic fungi were present in the top later of the
developing soil — although there is no experimental evidence for the transfer of
photosynthates and nutrients between AM fungi and bryophytes (Davey and Currah, 2006).
Plant establishment is an important factor accelerating organic matter accumulation by
litterfall, dead root tissues and root exudate, and thus affecting soil microbes (Zak, et al.,
2003; Wardle, et al., 2004). It is possible that once a complete bryophyte cover is established,
the moist surface conditions and possibly the exudates from the plants, encourage biotic and
abiotic weathering that provides further mineral nutrients for the bryophyte community. This
then is a positive feedback dependent on development of a bryophyte cover. However, the pH
of the ash is apparently not connected to the bryophyte cover as the decrease in pH was
already seen after one year when the bryophyte cover was still very low. Apparently
precipitation, MAT and possibly N deposition are the main factors influencing the pH at
these early stages of soil development.

The overall picture from this study is that within three years starting from sterile volcanic ash, in temperate and warm temperate climates at least, a soil ecosystem can build up with a surface layer of plant primary producers (bryophytes), significant accumulation of soil C and N, with ammonia oxidation occurring. In the warm temperate sites, the system was beginning to show signs of becoming closed, with reduced leaching of N out of the system.

Bryophytes are known to be a strong sink of atmospherically deposited N (Binkley and Graham, 1981, Weber and Van Cleve, 1984). Nevertheless, in all the sites the amount of C and N after three years is still much less than in mature forest soils, such as would be found under forest in Japan (Fig. S4).

It is important to bear in mind that the mesocosm system we used might either accelerate or decelerate the speed of soil ecosystem development on volcanic ash. The gauze covered pots we used might tend to avoid extremes of temperature, and retain humidity. We found that generally, in all sites, the air temperature within the gauze-covered trays next to the pots was about 1 °C warmer than the ambient in sunny weather – though no different in other weather conditions (Kerfahi et al. 2017). This raised temperature may tend to accelerate soil development – although it should affect all sites to a similar extent. On the other hand, it is possible that with the gauze covering excluding windblown seeds, the lack of vascular plants rooting into the ash might have impeded the process of ecosystem development. It has been suggested that vascular plant roots and their associated biota play an important role in chemical weathering (Cochran and Berner, 1996; Landeweert et al., 2001).

4.2. How does temperature affect the rate of soil ecosystem development?

In the first two years (at 12 months and 24 months) of the experiment, there were
only a weakly significant or non-significant effects of temperature on the soil attributes we measured (Fig. 1). However, at the 3 years stage (36 months) a clear effect of temperature emerged, at least in terms of C and N (Fig. 1c, f), and only in the top 2 cm layer of the pots. It is noticeable that the acceleration of C and N accumulation in the soils of the warmest sites is associated with an increase in bryophyte cover. This change further emphasizes the potential importance of bryophytes in the development of this ecosystem – acting as a source of C and N to the developing soil.

The relationship to bryophyte cover may be tentatively seen as evidence for a positive feedback effect between soil ecosystem development and bryophyte vegetation cover, with accelerated plant growth being an important driver at higher temperature. It appears that in the warmer three sites in our series, soil ecosystem development is held back until bryophyte cover develops – at which point there is a rapid increase in C and N, presumably due to dead material arriving in the upper soil from the bryophyte plants. The accumulation of C and N itself can be expected to promote bryophyte growth, by providing a more moisture retentive and nutrient rich soil, and accelerating chemical weathering of the ash - although there was no direct measurement of chemical weathering indicators in this study.

4.3. Microbial abundance did not increase with age or temperature.

We had anticipated that the copy number of microbial genes – an indicator of microbial population density - would increase greatly during the 3 years experiment. Surprisingly, there was no significant time trend in any of the indicators (16S rRNA genes of bacteria, archaea and fungi, and archaeal and bacterial amoA genes). Nor was there any tendency for the warmer climate stations to have greater copy numbers, even though C and N accumulation in the soil was clearly greater in the warmer sites by 36 months (Fig 1c, f).

The microbial abundance may reflect an active population which processes new
input of organic material to the soil from bryophytes and other primary producers such as cyanobacteria. However, despite the throughput of material – which leads to the organic C and N building up in the ash pot soils – microbial abundance does not appear to accumulate. In mature forest soils, by contrast, microbial abundance is much greater (Fig. S5), so on the longer timescale there is clearly some overall relation to age of the soil.

5. Conclusion

Our study provides a rare first glimpse and the very earliest stages of soil ecosystem development on volcanic ash. Following through volcanic ash mesocosms for their first three years reveals a temperature- and time-dependent component, with accelerated ecosystem development once a complete bryophyte cover is established. In warmer climates, this cover establishes early, possibly initiating a feedback between soil development and bryophyte coverage.

It would be interesting to follow up this study with further analysis of the same soils – measurement of conversion of silicates to clays as an indicator of weathering, Ca mineral fluxes, and sequencing of bacterial communities and metagenomes for Year 3. A longer term study, following the ash soil over 20 years or more of ecosystem development, would also be interesting to the understanding of ecosystem development in volcanic landscapes.

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cool-temperate deciduous forest. Soil Biol. Biochem. 124, 90-100.


development of volcanic ash soils along a climosequence in Northern Taiwan. Geoderma 156, 48-59.


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<th>Site Name</th>
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<th>MAP (mm)</th>
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Table 2 Bryophyte cover (%) of ash pots after 1-year, 2, and 3 years of experiment. Mean ± SD (n = 5).

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<th>3-year</th>
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<tr>
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<td>SJ</td>
<td>0.0 (0.0)</td>
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Figure captions

**Fig. 1**  C and N accumulation rates of ash pots for (a, d) 1-year, (b, e) 2, and (c, f) 3 years of experiment in relation to MAT. We tested three models (linear, quadratic, and exponential) to describe relationships with MAT. Model selection was carried out based on adjusted $R^2$ and RMSE. Dotted lines indicate best fitted lines. (c) $y = 0.105x^2 - 0.381x + 2.673$, $R^2 = 0.975$, RMSE = 1.19, (e) $y = 0.959e^{0.031x}$, $R^2 = 0.664$, RMSE = 0.186, (f) $y = 0.180e^{0.135x}$, $R^2 = 0.789$, RMSE = 0.486.

**Fig. 2**  (a) C and (b) N contents of ash in relation to percentage of bryophyte cover. (a) $y = 0.339x + 10.267$, $R^2 = 0.821$, RMSE = 4.680, (b) $y = 0.0258x + 1.119$, $R^2 = 0.777$, RMSE = 0.409.

**Fig. 3**  Pool size of (a) NO$_3$-N, (b) NH$_4^+$-N, and (c) inorganic N (NO$_3$-N + NH$_4^+$-N) of ash pots in relation to MAT. We tested three models (linear, quadratic, and exponential) to describe relationships with MAT. Model selection was carried out based on adjusted $R^2$ and RMSE. (b) $y = -0.002x + 0.144$, $R^2 = 0.824$, RMSE = 0.0058 for 2014, (c) $y = -0.002x + 0.150$, $R^2 = 0.805$, RMSE = 0.0242 for 2014.

**Fig. 4**  Proportion of inorganic N to total N in ash pots in relation to MAT. Solid line, dotted line and broken line indicate the significant relationships between proportion of inorganic N to total N and MAT for 2013 ($y = 0.0091x + 18.618$, $R^2 = 0.0003$, RMSE = 3.729), 2014 ($y = -0.3655x + 13.294$, $R^2 = 0.9716$, RMSE = 0.408) and 2015 ($y = -1.0599x + 22.866$, $R^2 = 0.8551$, RMSE = 2.847), respectively.
Fig. 5 Leaching of (a-c) NO$_3^-$-N, (d-f) NH$_4^+$-N, and (g-i) inorganic N from ash pots in relation to MAT for 1-year, 2, and 3 years of experiment. We tested three models (linear, quadratic, and exponential) to describe relationships with MAT. Model selection was carried out based on adjusted $R^2$ and RMSE. There were no significant relationships with MAT.

Fig. 6 Gene abundance of (a-c) bacterial 16S rRNA, (d-f) archaeal 16S rRNA, and (g-i) fungal ITS for 1-year, 2, and 3 years of experiment in relation to MAT. We tested three models (linear, quadratic, and exponential) to describe relationships with MAT. Model selection was carried out based on adjusted $R^2$ and RMSE. Closed circle and open triangle indicate lower and upper layer of ash pots, respectively. (c) $y = -0.012x^2 + 0.192x + 6.838$, $R^2 = 0.874$, RMSE = 0.156 for Lower layer, (f) $y = -0.015x^2 + 0.261x + 4.285$, $R^2 = 0.959$, RMSE = 0.766 for Lower layer.

Fig. 7 Gene abundance of (a-c) bacterial amoA and (d-f) Archaeal amoA for 1-year, 2, and 3 years of experiment in relation to MAT. We tested three models (linear, quadratic, and exponential) to describe relationships with MAT. Model selection was carried out based on adjusted $R^2$ and RMSE. There were no significant relationships with MAT. Closed circle and open triangle indicate lower and upper layer of ash pots, respectively.
• The first three years of C and N accumulation were investigated in a field mesocosm
• By 24 months, C and N accumulation did not show a clear correlation with temperature
• By 36 months, C and N accumulation correlated exponentially with temperature
• The abundance of soil microbes did not increase with time and temperature
• The faster C and N acceleration in warmer sites may relate to bryophyte cover
Figure

Fig. 1 Tateno et al.

Accumulation rate (g m$^{-2}$ year$^{-1}$) vs. MAT(°C)

(a) C, 1-year
(b) C, 2-year
(c) C, 3-year
(d) N, 1-year
(e) N, 2-year
(f) N, 3-year
Fig. 2 Tateno et al.

(a) C contents (g m$^{-2}$) vs. Percentage of Bryophyte cover (%)

(b) N contents (g m$^{-2}$) vs. Percentage of Bryophyte cover (%)

- C contents: 0 to 60 g m$^{-2}$
- N contents: 0 to 4 g m$^{-2}$
- Percentage of Bryophyte cover: 0 to 100%
Fig. 3 Tateno et al.

(a) 

\[ \text{NO}_3^- \text{-N pool (g m}^{-2}\text{)} \]

(b) 

\[ \text{NH}_4^+ \text{-N pool (g m}^{-2}\text{)} \]

(c) 

\[ \text{Inorganic N pool (g m}^{-2}\text{)} \]

\( \text{MAT}(\degree\text{C}) \)
Fig. 5 Tateno et al.

(a) 1-year

(b) 2-year

(c) 3-year

(d) 1-year

(e) 2-year

(f) 3-year

(g) 1-year

(h) 2-year

(i) 3-year

NO₃⁻-N leaching (g m⁻² period⁻¹)

NH₄⁺-N leaching (g m⁻² period⁻¹)

Inorganic N leaching (g m⁻² period⁻¹)

MAT(℃)
Fig. 6 Tateno et al.

Log number of Bacterial 16S rRNA (g⁻¹)

(a) 1-year

(b) 2-year

(c) 3-year

Log number of Archaeal 16S rRNA (g⁻¹)

(d) 1-year

(e) 2-year

(f) 3-year

Log number of fungal ITS (g⁻¹)

(g) 1-year

(h) 2-year

(i) 3-year

MAT(°C)
Fig. 7 Tateno et al.

Log number of Bacterial AmoA (g⁻¹)
(a) 1-year  (b) 2-year  (c) 3-year

MAT(°C)

Log number of Archaeal AmoA (g⁻¹)
(d) 1-year  (e) 2-year  (f) 3-year

MAT(°C)