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Infiltration in Icelandic Andisols: the Role of Vegetation and Soil Frost

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Abstract

Soil frost formation, snow distribution, and winter/spring/summer terminal infiltration rates (TIRs) were quantified in Icelandic Andisols with contrasting vegetation cover types (grassland, spruce and birch woodland, lupine, and sparsely vegetated lava site). TIRs (mm h^{-1} ; determined with double-ring infiltrometers) were generally higher in unfrozen than in frozen soils (102–369 vs. 9–306, respectively in sandy soils; 28–94 vs. 3–72 in finer-textured soils) and differed between land cover types, being consistently highest in birch woodlands. TIR was an inverse function of soil frost depth. Lowest TIRs were associated with deep and dense soil frost, which formed in spruce woodland and grassland communities where snow depth was shallow. Results suggest conditions conducive to erosion by water are most likely to occur during winter warm spells and in spring in vegetation types where snow cover is low or ephemeral. Threefold increases in TIRs occurred one year after livestock grazing was discontinued, suggesting Andisols are hydrologically resilient where vegetation cover is relatively continuous and soil organic carbon content is high.

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Introduction

When ecosystems become dysfunctional, hydrological processes are significantly altered, often resulting in high runoff and loss of soil and nutrients. Hence, criteria related to hydrologic function are often used to evaluate ecosystem stability and resistance to disturbance. The capacity of a site to capture and store precipitation, run-on, and snowmelt, and the stability of this capacity, are key determinants of ecohydrological function (Ludwig et al., 1997; Pellant et al., 2000; Ludwig et al., 2005). Infiltration is an important component of water capture potential.

Vegetation enhances infiltration by increasing surface roughness, by reducing raindrop impact, and by improving soil porosity via root channels, addition of organic matter, and providing habitat for soil biota (e.g., Branson et al., 1981; Thurow, 1991). These influences vary with plant functional groups, with infiltration generally being highest in soils under trees and shrubs, lower in grasslands, and least in sparsely vegetated areas (Thurow et al., 1986). In high latitude ecosystems, vegetation additionally influences infiltration via its effects on radiant energy exchange and snow capture which, in turn, affect soil frost formation (e.g., Tyrtikov, 1976; Jones et al., 2001). Occurrence of soil frost and depth of freezing in relation to vegetative cover has been found to decrease in the rank order: bare field > grassland > evergreen conifer forest > deciduous forest (Pierce et al., 1958; Shanley and Chalmers, 1999).

Soil frost can impede or enhance infiltration. Concrete frost, which consists of many thin ice lenses and small crystals, may be

essentially impermeable to water (Post and Dreibelbis, 1942; Trimble et al., 1958; Dunne and Black, 1971). Soil frost with loose crystals may be porous (Post and Dreibelbis, 1942; Storey, 1955); and may have infiltration rates greater than their unfrozen counterparts (Trimble et al., 1958; Haupt, 1967; Blackburn and Wood, 1990) due to soil structural changes accompanying freezing (e.g., enhanced aggregation; Luo et al., 2003). Concrete frost tends to develop in soils with no or delayed snow cover (Haupt, 1967; Fahey and Lang, 1975) and in open land susceptible to deep frost penetration (Pierce et al., 1958; Stoeckeler and Weitzman, 1960); it is less common in forested areas, particularly deciduous forests (Pierce et al., 1958; Stoeckeler and Weitzman, 1960; Fahey and Lang, 1975). High soil water content promotes formation of dense soil frosts that retard infiltration (Kane and Stein, 1983). Consequently, rainfall or snowmelt during warm spells followed by freezing temperatures enhances concrete frost formation (Proulx and Stein, 1997; Shanley and Chalmers, 1999). An understanding of the ecohydrology of high latitude ecosystems thus requires an understanding of how vegetation and climate interact to affect the type of soil frost that may form.

Soils in the Andisol order are common in volcanically active regions of the world (Soil Survey Staff, 1999). Although Andisols have high infiltration rates (Forsythe, 1975; Shoji et al., 1993), they are highly vulnerable to erosion by water (Nishimura et al., 1993; Poulenard et al., 2001) and wind (Arnalds, 2000; Arnalds et al., 2001), due to the low-density soil aggregates, lack of particle cohesion, and high soil water retention (Wada et al., 1992; Kimble et al., 2000). High water retention also makes high latitude

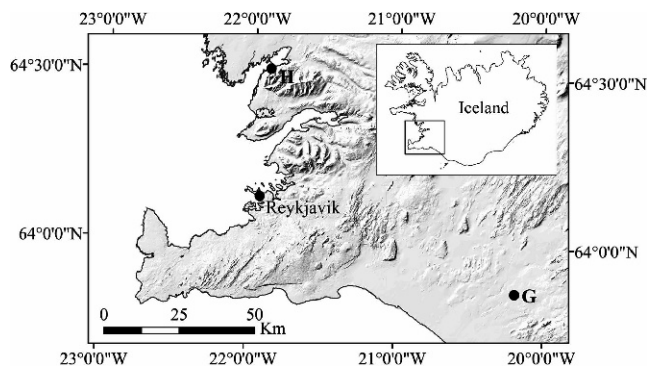


FIGURE 1. Location of the Gunnarsholt (G) and Hafnarskogur (H) study sites in Iceland.

Andisols susceptible to frost heaving and needle ice cryoturbation. Natural or anthropogenic reduction of the plant biomass thermal barrier may thus intensify cryogenic processes that hasten formation of bare patches (Archer and Stokes, 2000) while promoting the formation of impermeable soil frost types.

Andisols are the dominant soil order in Iceland (Arnalds, 2004) where extensive vegetation change, loss of plant cover, and soil erosion have occurred (Aradottir and Arnalds, 2001; Arnalds et al., 2001). Since settlement about 1100 years ago, birch woodland cover is estimated to have declined from 25% (Sigurdsson, 1977) to only 1% (Gudjónsson and Gíslason, 1998); and lands classified as barren desert have increased from 10–15% cover to >35% cover (Arnalds, 2000). Erosion by water in Iceland is significant and a significant fraction of this erosion occurs in conjunction with winter rains and spring snowmelt (Arnalds et al., 2001). An understanding of how vegetation and land cover change affects hydrological processes influencing erosion is central to the development of land management protocols aimed at curtailing land degradation and restoring dysfunctional ecosystems. In this study, we sought to quantify the relationship between vegetation cover, soil frost formation, and infiltration in Icelandic Andisols.

The maritime winter climate of Iceland is characterized by freeze-thaw cycles and thin and fluctuating snow cover (Einarsson, 1984). We therefore expected that high frequencies of freeze-thaw events, coupled with the high water holding capacity of Andisols, would cause the formation of impermeable, ice-rich zones. We hypothesized that (a) infiltration rates in such zones would be reduced, hence increasing runoff and soil erosion potential following winter rain and snowmelt runoff, but that (b) the formation of such zones would vary as a function of the type and amount of vegetation cover. As Andisols have intrinsically high infiltration rates, we also hypothesized that (c) type of vegetation cover would not affect infiltration rates in unfrozen soil. To test these hypotheses, we quantified rates of water infiltration and soil frost dynamics in contrasting land cover types during one year.

Study Sites

Studies were conducted at two sites in the lowlands of South and West Iceland. These sites represented contrasting land cover types spanning a range of vegetation degradation and regeneration states. The Gunnarsholt site (G-site; 63°52'N, 20°12'W; ca. 100 m a.s.l.) consisted of revegetated landscapes, whereas the Hafnarskogur site (H-site; 64°30'N, 21°55'W; ca. 30 m a.s.l.) represented degraded landscapes (Fig. 1). Mean January and July temperatures at these sites are -1 to -2 °C and 10 to 11 °C, respectively; mean annual precipitation is 1200 mm (G-site) and 1500 mm (H-

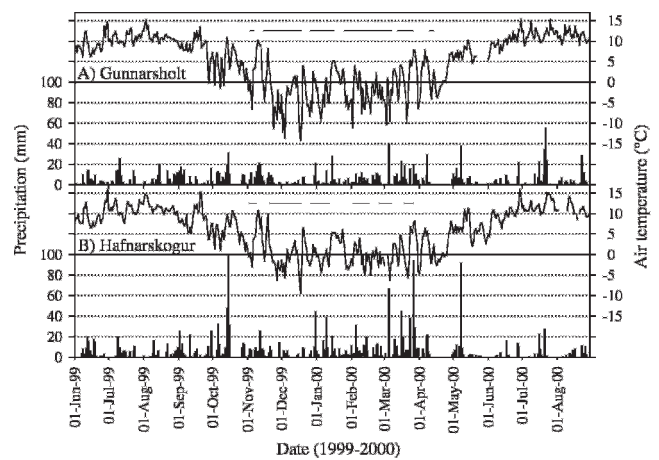


FIGURE 2. Daily precipitation (bars) and air temperature (line), and days with 100% snow cover (horizontal lines) at (A) Gunnarsholt and (B) Hafnarskogur, June 1999 to August 2000 (Veðráttan, 1999, 2000). Gunnarsholt, data from weather station Hella (11 km WSW of site); Hafnarskogur, temperature from Hafnarmelar (5 km SSW of site), precipitation and snow cover from Andakilsárvirkjun (12 km ENE of site).

site). Temperatures are typically less extreme and snow cover shallower at the H-site compared to the G-site, owing to its proximity to the sea (<1 km versus 25 km, respectively). Weather conditions at both sites during the study years 1999 to 2000 are shown in Fig. 2.

The G-site, excluded from livestock grazing since the late 1920s, is comprised of small hills formed by post-glacial basaltic lava flows onto otherwise relatively level terrain. Soils are deep (>1 m) Vitricryands (O. Arnalds, unpublished; Strachan et al., 1998) or Brown Andosols (Icelandic classification scheme; Arnalds, 2004) developed in eolian and tephra materials deposited on the lava field. Sand encroachment covered much of the area with dunes in the late 1800s (Sigurjónsson, 1958). Reclamation efforts in the 1900s stabilized these dunes and created a mosaic of plant communities, five of which were used in this study. These communities occur in two areas (referred to as western and eastern) located about 1 km apart. Birch (*Betula pubescens* Ehrh.) woodland, lupine (*Lupinus nootkatensis* Donn ex Sims) fields, and lava fields sparsely vegetated with *Festuca* grasses and ericaceous dwarf shrubs occur in the western area; and a Sitka spruce (*Picea sitchensis* (Bong.) Carrière) woodland and grassland dominated by *Festuca* and *Agrostis* grasses and *Galium* herbs occur in the eastern area. We refer to these different plant communities as G-birch, G-lupine, G-lava, G-spruce, and G-grassland, respectively. The birch woodland developed from seed planted in 1939 and 1945 (Magnússon and Magnússon, 1989), and consists of birch (mean height 2.2 m) and willows (*Salix phylicifolia* L. and *S. lanata* L.) with a density of stems >30 cm height = 9774 stems ha⁻¹. The first lupines were introduced at the site in 1988 (Sveinn Runólfsson, personal communication). Vegetation on the lava field is self-regenerated. The grassland community developed from *Leymus arenarius* (L.) Hochst. seeds sown around 1930. The Sitka spruce stand (15 × 100 m; 2525 trees ha⁻¹; mean height 5–6 m) was planted in early 1960s (Sveinn Runólfsson, personal communication). Plant nomenclature follows Kristinsson (2005).

The H-site includes hummocky birch woodland (density of stems > 30 cm height = 5344 stems ha⁻¹; mean height = 1.7 m) and *Agrostis-Festuca-Deschampsia* grassland considered to have had some woodland cover in earlier times. These will subsequently

be referred to as H-birch and H-grassland. Soils are deep (>1m) Haplocryands (O. Arnalds, unpublished; Strachan et al., 1998) or Histic Andosols (Icelandic classification scheme; Arnalds, 2004) formed in eolian and tephra materials deposited on an old (about 10,000 yr) sand/gravel shoreline. Sheep grazing was discontinued in the early 1990s, but horses graze the grassland, except for short periods in mid-summer. The H-grassland research plot was fenced in August 1999 to exclude livestock and prevent damage to instrumentation.

Methods

INFILTRATION RATES

Metal double-ring infiltrometers (Bouwer, 1986) were installed in each plant community ($n = 6$) in June/July 1999. The rings were 40 cm high, with 22.5 and 39.0 cm inner and outer diameters, respectively. Rings were driven 30 cm into the soil. Placement of rings was between hummocks, and midway between trees in woodlands. Because one of our objectives was to quantify infiltration rates on frozen soils, and because installation of rings in frozen soils would have been impossible without causing unacceptable levels of disturbance, rings were left in place for the duration of the study. A repeated-measures design was thus used (e.g., van der Kamp et al., 2003). Initial readings were made between 17 July and 5 August 1999. Due to time constraints, no measurements were made on G-grassland. Winter and spring measurements were made during three periods in 2000: 5–11 January, 15–21 April, and 14–22 May. The G-birch, G-lupine, and G-lava communities were inaccessible in January due to snow.

A water head of 4 cm was maintained within rings during infiltration trials, and the water required to maintain the head in the inner ring was recorded at five minute intervals for one hour. Terminal infiltration rates (TIRs; mm h^{-1}) were calculated as the mean of the last three measurements. The temperature of water used approximated that of rainfall during the measurement periods: 0°C in January, $2\text{--}4^{\circ}\text{C}$ in April, and $6\text{--}8^{\circ}\text{C}$ in May and July/August.

SNOW, ICE, AND SOIL FROST

Snow cover (%) and depth (cm) were measured within infiltration rings and along transects in each plant community before infiltration trials in January, but only within rings in April, as snow measured in rings and transects were comparable. Ice cover within rings was estimated visually.

Soil frost depth (cm) and type were quantified during infiltration trials in January, April, and May. Frost depth was measured in small pits near each infiltrometer by visually noting the occurrence of ice crystals. The size and number of ice crystals observed with a $10\times$ hand lens was used to classify frost into one of three categories using criteria in Post and Dreibeilbis (1942) and Stoeckeler and Weitzman (1960): *concrete frost*—a high density of small crystals; *porous concrete frost*—resembled concrete frost, but with lower crystal density; and *porous frost*—fewer and larger ice crystals than the concrete frost types. Soil frost measurements were not possible in G-birch, G-lupine, and G-lava in January because of snow. Shallow standing water in depressions in H-grassland in January limited soil frost measurements ($n = 3$).

GROUND COVER AND SOIL

Ground cover and plant composition of inner infiltration rings was recorded in July/August 2000 using a point-frame (5 cm

pin interval; 15–17 pins per ring). For each pin drop, all contacts with vegetation, litter, stones, and soil were recorded. Above-ground biomass (g m^{-2}) of the ground layer was quantified by clipping vegetation and litter within the inner rings, sorting by life form, and oven-drying at 60°C . Vascular plant species diversity was computed as richness (number of species) and as an abundance-weighted richness (Simpson's Diversity Index; Huston, 1994) in 30 plots per community using a point-frame ($n = 5$ pins per plot).

Soil properties within each inner infiltration ring (to 30 cm depth; by horizon) were determined on samples obtained at the end of the study (August/September 2000). Bulk density (BD) was determined by the core method (Blake and Hartge, 1986). Organic carbon (OC) was quantified by dry combustion (Leco CR-12 carbon analyzer; Nelson and Sommers, 1982). Sand content was determined on field-moist samples. Field texture and water content of field-moist surface soil samples at -0.033 MPa (pressure chambers; Klute, 1986) were determined on samples from the soil pedons described in each plant community.

The sequence and designation of genetic horizons in the soils varied greatly between sites (Table 1). Therefore, BDs in the different communities were not compared on the basis of horizon designations but on their three upper horizons: surface horizons were denoted *horizon 1*; the subtending horizon was denoted *horizon 2*; and the deepest horizons (down to ca. 30 cm depth) were denoted as *horizon 3* (and *4* at H-grassland). The properties of the genetic horizons represented in these groups are summarized in Table 1.

EXPERIMENTAL DESIGN AND DATA ANALYSIS

Logistical constraints forced us to choose between surveying numerous plant communities or studying one or two replicated communities. We opted for the former, as information on the infiltration/frost dynamics of Andisols with different vegetation cover is lacking. Infiltration rings were treated as the unit of replication ($n = 6$). Extrapolation of our results are thus predicated on the assumption that the stands we sampled are broadly representative of the community type, an assumption that should be evaluated in future research. Statistical analyses were performed with SAS v.8.01 (SAS, 2000). Ground cover and depth of soil frost were analyzed using analysis of variance (ANOVA), where communities (G-spruce, H-birch, etc.) were treated as main effects. Between-community comparisons of snow depth measured along transects were ascertained with a Kruskal-Wallis test. Spearman rank correlation coefficients (r_s) were determined for the relationship between January soil frost depth and snow depth within the infiltrometers.

TIR data were analyzed separately for communities in the revegetated and degraded sites. The effect of soil frost depth on TIR was tested with analysis of covariance (ANCOVA) for the H-site April data. Simple linear regression was used to describe the relationship between soil frost depth and TIR at the G-site in April, excluding rings with 100% ice cover. At the H-site, plant community differences in TIR were analyzed separately with ANOVA on a given date. This was also done at the G-site for the January and April TIRs, but plant community effects on TIR in unfrozen soils were assessed with univariate, repeated measures ANOVA, with an error term consisting of the nested variation of the infiltration rings within the plant communities (Littell et al., 1998). Correlation coefficients (r_s) were determined for the relationship between vegetation and soil properties and the July/August 2000 TIRs. When overall tests were significant ($p < 0.05$),

TABLE 1
Topographic setting and selected soil and vegetation characteristics in the contrasting land cover types at the Gunnarsholt (G) and Hafnarskógur (H) study sites in South and West Iceland. See Table 2 for bulk density data.

Plant community	Slope inclination and aspect	Horizon, horizon depth (cm) ^a	Soil surface horizon			Soil horizons below surface horizon to 30 cm depth			Vascular plants in ground layer				
			Field texture	Sand (%) ^b	OC (%)	Water content at -0.033 MPa (%)	Sampled horizons ^c	Field texture	Weighted average sand (%) ^d	Weighted average OC (%) ^d	Species richness	Simpson's diversity index	Dominant species
G-spruce	5°, SE	Oi, 3 (5)	—	—	17.4	125	A1, 3Bw1	1	72	1.9	3	0.42	<i>Agrostis capillaris</i>
G-grassland	8°, SE	A1, 5	sil	44	7.3	81	A2, A3	1	70	2.0	10	0.82	<i>Festuca richardsonii</i> , <i>Agrostis capillaris</i> , <i>Galium boreale</i> , <i>G. verum</i> , <i>Equisetum arvense</i>
G-birch	20°, S	A1, 7	1	49	6.5	54	A2, 2C	1, s	77	1.1	17	0.86	<i>Agrostis capillaris</i> , <i>Poa pratensis</i> , <i>Festuca richardsonii</i> , <i>Hieracium spp.</i> , <i>Equisetum arvense</i>
G-lupine	25°, S	A1, 16 (5)	sl	70	2.3	—	A1, A2	sl, 1	54	1.6	—	—	<i>Lupinus nootkatensis</i>
G-lava	25°, S	A1, 16 (5)	sl	68	2.3	72	A1, A2	sl, 1	62	2.4	15	0.85	<i>Calluna vulgaris</i> , <i>Empetrum nigrum</i> , <i>Festuca richardsonii</i> , <i>F. vivipara</i> , <i>Galium verum</i>
H-birch	0°, —	Oi, 8	1	—	19.6	162	A	cl	37	8.0	17	0.77	<i>Deschampsia flexuosa</i> , <i>Agrostis capillaris</i> , <i>Anthoxanthum odoratum</i> , <i>Gymnocarpium dryopteris</i>
H-grassland	3°, NW	Oi, 5	cl	—	25.0	167	A1, A2, Bw1	cl	—	20.2	11	0.71	<i>Agrostis capillaris</i> , <i>Festuca richardsonii</i> , <i>Deschampsia caespitosa</i>

^a Number in brackets show sampling depth of surface horizons if they are not equal to the horizon thickness.

^b Sand content was not determined on soils containing >10% OC as carbon cannot be removed effectively from Andisols (Arnalds et al., 1995).

^c Horizons 2 and 3, respectively, in Table 2.

^d Represents a weighted average of 2 samples (3 for H-grassland).

TABLE 2

Mean (± 1 SE) bulk density (g cm^{-3}) of soil horizons from the surface to 30 cm depth in plant communities at Gunnarsholt (G) and Hafnarskogur (H). Means are n of 6 samples except: horizons 1 and 2 in G-lupine ($n = 5$); and horizon 3 in G-lupine ($n = 4$). See Table 1 for horizon designations for each plant community.

Plant community	Bulk density \pm SE (g cm^{-3})		
	Soil horizon 1	Soil horizon 2	Soil horizon 3
G-spruce	0.25 ± 0.028	0.62 ± 0.037	0.82 ± 0.025
G-grassland	0.45 ± 0.016	0.71 ± 0.018	0.89 ± 0.013
G-birch	0.44 ± 0.018	0.80 ± 0.019	0.92 ± 0.019
G-lupine	0.74 ± 0.045	0.79 ± 0.074	0.74 ± 0.060
G-lava	0.76 ± 0.035	0.77 ± 0.018	0.56 ± 0.050
H-birch	0.24 ± 0.020	0.48 ± 0.025	0.52 ± 0.018
H-grassland ^a	0.25 ± 0.013	0.43 ± 0.007	0.37 ± 0.024

^a BD \pm SE of horizon 4 was $0.33 \pm 0.02 \text{ g cm}^{-3}$.

sequential Bonferroni multiple comparison tests (Rice, 1989) were used to make pair-wise comparisons of means. Data were transformed prior to analysis if assumptions were not met.

Results

GROUND COVER AND SOILS

Soils, vegetation, and topography are summarized in Table 1. OC content was highest in surface (19.6–25.0%) and subsurface (8.0–20.2%) horizons at the H-site and lowest in G-lupine and G-lava soils (1.6–2.4%). At the H-site, soil field texture was finer (loam, clay loam) than at the G-site (sand, sandy loam, loam, silt loam). Water-holding capacity of the H-birch and H-grassland surface soils was high (162% and 167% at -0.033 MPa, respectively) as was that of the G-spruce surface soil (125%); water-holding capacity in other communities was $\leq 81\%$. Surface horizon BD ranged from 0.24 to 0.76 g cm^{-3} and was lowest in the G-spruce, H-birch, and H-grassland, and highest in G-lupine and G-lava (Table 2).

Vascular plant species richness of ground cover was greatest in the G- and H-birch woodlands and G-lava communities and least in G-spruce woodland (Table 1). Bare soil cover was high in the G-lava (31%) community, but $<10\%$ in all others (Fig. 3). Litter dominated the G-spruce surface. Vascular plant ground cover differed between plant communities ($p < 0.001$): cover was lowest in G-spruce, highest and comparable in G-birch, G-grassland, G-lupine, H-birch, and H-grassland, and intermediate in G-lava. Total aboveground biomass ranged from 17 g m^{-2} in G-spruce to 840 g m^{-2} in G-grassland (Fig. 3).

SNOW, ICE AND SOIL FROST

The winter 1999–2000 was characterized by more persistent and greater snow cover than in average years (Vedráttan, 1999, 2000). Data from the weather station nearest each study site provide a crude index of snow cover persistence and continuity (Fig. 2). Transect measurements on our sites documented continuous snow cover in early January in all plant communities, except H-grassland where cover was 86%. However, mean snow depth differed between communities ($p < 0.001$), being deepest in the G-birch, G-lupine, and G-lava communities (72–114 cm), and shallowest (≤ 11 cm) in G-spruce and H-grassland (Fig. 4). By April, snow had disappeared in all but the G-birch community (Fig. 4), where it persisted until the end of April. The snow cover data from weather station Hella (Fig 2A), and unpublished soil temperature data in the G-birch, G-spruce, and G-lava communities suggest that snow in the other G-communities disappeared by the end of March.

G-grassland and H-birch infiltration rings were ice-free in January 2000. Five of six H-grassland rings had ice cover (100% cover in four) as did three of six G-spruce rings (but none had 100% cover). Ice was only observed in two G-spruce rings in April (100% cover in each) and only one G-birch ring (25% cover). The observed occurrence of ice cover in rings was qualitatively representative of the patchiness of ice distribution in the broader communities (personal observation).

Soil frost depth in January ranged from a mean of 0.7 cm in H-birch to 16.7 cm in G-spruce. In April the mean frost depth ranged from 0.6 cm in G-birch to 19.7 cm in G-spruce (Fig. 4). G-

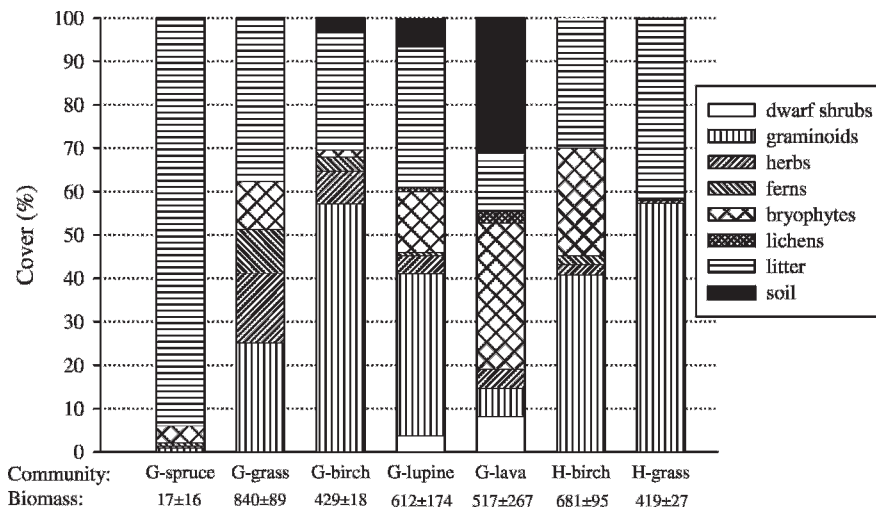


FIGURE 3. Relative ground cover (%) and total aboveground biomass (mean ± 1 SE; g m^{-2}) within inner infiltration rings in plant communities at Gunnarsholt (G) and Hafnarskogur (H). Data are means of 6 rings per plant community ($n = 5$ for G-lupine) in July/August 2000. The biomass of litter (duff layer) in the spruce woodland not included.

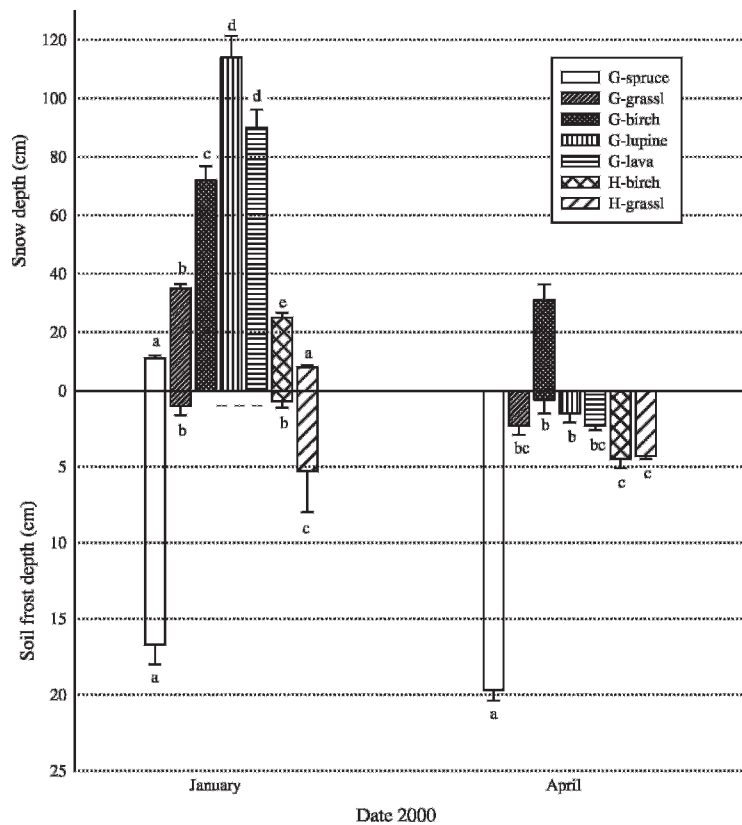


FIGURE 4. Upper panel: mean snow depth (± 1 SE; cm) measured along transects ($n = 13\text{--}49$) in January, and within inner infiltration rings in April. Lower panel: mean soil frost depth (± 1 SE; cm) near infiltration rings in plant communities at Gunnarsholt (G) and Hafnarskogur (H) in January and April 2000. Means followed by different letters within each date were significantly different ($p < 0.05$). Means of soil frost are $n = 6$ samples except: in January $n = 3$ for H-grassland; in April $n = 5$ for G-birch. Concrete frost characterized G-spruce soil; porous concrete frost characterized H-grassland soils; and porous frost characterized soils in all other communities. Deep snow prevented soil frost data collection in January for G-birch, G-lupine and G-lava.

spruce was the only community where soil frost was observed in May (data not shown). By this time, surface soils had thawed but a frost lens beginning at 8.4 ± 2.7 cm depth and extending to 15.0 ± 2.5 cm was still evident. Soil frost depth differed between communities both in January ($p < 0.001$) and April ($p < 0.001$) (Fig. 4). It is noteworthy that at the H-site, although soil frost in January had penetrated to greater depths in the grassland compared to the deciduous birch woodland (5.3 cm and 0.7 cm, respectively), depths were comparable in April (4.3 and 4.5 cm) owing to increases in woodland frost depths. Increased frost depths from January to April were also observed in the evergreen G-spruce woodland, although soil frost there was already deep in January. In contrast, soil frost depths in the grassland communities were comparable in January and April. January soil frost depth and snow depth within infiltrometers were negatively correlated both at the G-site ($r_s = -0.90$, $p < 0.001$; includes the spruce woodland and grassland) and the H-site ($r_s = -0.51$, $p < 0.16$).

All three structural types of soil frost (concrete, porous concrete, and porous) were observed during January and April. Concrete frost characterized all soil samples in G-spruce. Porous concrete frost characterized all samples in H-grassland, whereas porous frost characterized soils in G- and H-birch, G-grassland, G-lupine, and G-lava. The frost types did not appear to change over time within a community. Qualitative observations indicated ice crystals in G-spruce and H-grassland organic soil horizons were larger than those occurring in the mineral horizons below. Needle ice was observed among the porous frost of G-lava soils (three of six April samples).

INFILTRATION RATES

Terminal infiltration rates (TIRs) were typically achieved within 10 to 20 minutes. Winter TIRs ranged from 3 to 306 mm

h^{-1} . Summer TIRs ranged from 28 to 369 mm h^{-1} . As a rule, rates were lowest in the H-grassland community and highest in the G-birch woodland (Fig. 5).

TIRs within the plant communities varied with time, the greatest differences occurring between the summer and winter dates (Fig. 5). TIRs in H- and G-birch woodlands, and the G-lupine and G-lava field communities varied between dates by a factor of 1.1 to 1.3. G-grassland TIRs varied by a factor of 2 (January vs. May 2000), and G-spruce woodland TIRs varied by a factor of 25 (April vs. July/August 2000). The greatest date difference was observed in H-grassland where the January and July/August 2000 TIRs varied by a factor of 37, and summer measurements varied by a factor of 3. Peak TIR occurred in summer for all communities except G-grassland, where peak TIR occurred in January.

Winter Rates

Infiltration was totally inhibited in rings where surface ice cover was 100%. There was no indication of soil thawing during the 60-minute trials. In January, at the H-site, birch woodland had greater TIR than grassland ($p < 0.001$); however, TIRs in G-grassland were much greater than those in H-grassland and far exceeded those of G-spruce woodland ($p < 0.001$) (Fig. 5). At the H-site in April, TIR was most strongly correlated with BD of the surface soil horizon ($r_s = -0.81$, $p < 0.01$). Correlations between TIR and soil frost depth were not significant ($r_s = -0.30$, $p = 0.35$) and accounting for soil frost depth did not improve prediction of TIR (ANCOVA; $p = 0.24$, $R^2 = 0.40$). TIRs in the two H-site communities differed at $p = 0.067$ with no interaction between plant community type and soil frost depth ($p = 0.85$). At the G-site, April TIRs in the five plant communities were most strongly correlated to soil frost depth ($r_s = -0.75$, $p < 0.001$). Exclusion of the spruce woodland, which exerted a strong

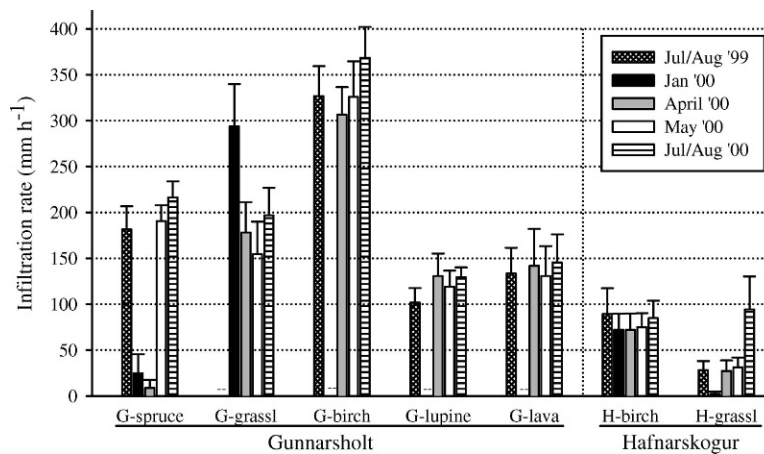


FIGURE 5. Mean (± 1 SE) terminal infiltration rates (mm h^{-1}) in plant communities at Gunnarsholt (G) and Hafnarskogur (H) on five dates. Means are from six double-ring infiltrometers except: G-birch and G-lupine ($n = 5$); G-lava in July/August 1999 ($n = 5$); and G-birch in April 2000 ($n = 4$).

influence on the data spread by virtue of its substantially greater soil frost depth (Fig. 4), reduced the Spearman r to -0.59 ($p < 0.01$). Using all G-site data points, the relationship between soil frost depth and TIR was best described by a non-linear regression function ($p < 0.001$, $R^2 = 0.61$; Fig. 6), whereas, when excluding G-spruce, a linear relationship explained most variability in the data ($p < 0.001$, $R^2 = 0.45$; Fig. 6). TIRs in the five communities differed ($p < 0.001$) in April in the rank order G-spruce < G-lupine \approx G-lava \approx G-grassland < G-birch woodland.

Summer Rates

Summer TIRs differed ($p < 0.001$) between G-site plant communities in the rank order spruce woodland \approx lupine \approx lava field < birch woodland. At the H-site, TIRs in the birch woodland were higher than those in the grassland in July/August 1999 ($p = 0.04$) and May 2000 ($p = 0.04$), but were comparable in July/August 2000 ($p = 0.82$) (Fig. 5). Variables strongly related to July/August 2000 TIR in the G-site communities were surface soil BD ($r_s = -0.59$, $p < 0.01$) and organic cover ($r_s = 0.50$, $p < 0.01$). At the H-site, BD of the surface soil horizon was the only variable significantly correlated with TIR in July/August 2000 ($r_s = -0.85$, $p < 0.001$).

Discussion

The results indicate that the formation of ice-rich zones in Andisols reduces infiltration rates. The formation of these ice-rich zones appeared to vary with the type of soils and vegetation cover,

but not with the amount of plant ground cover, as ice-rich zones with low TIRs occurred in both sparsely (G-spruce) and well-vegetated (H-grassland) Andisols. Contrary to expectations, type of vegetation cover also had a significant influence on TIRs of unfrozen soils.

The TIR data presented here were generated using ring infiltrometers left in place for a full annual cycle. Infiltration rates measured with ring infiltrometers do not represent rates occurring under natural rainfall (Branson et al., 1981); therefore, the rates we present are an assessment of the relative differences between plant communities and soils. There are some considerations when using this technique in high latitude environments. Metal rings with their high thermal conductivity may influence soil frost formation. However, similarities in frost type and depth in soils adjacent to and away from rings suggest this was not a problem in our study (data not shown). To avoid ring-induced ice formation, we recommend drilling small holes in the rings just above the ground surface, such that water can freely exit downslope. During measurements, the holes are plugged. Insertion of rings to 30 cm depth was sufficient to prevent ejection by frost heaving.

INFILTRATION RATES IN SUMMER

In a review of infiltration rates in soils derived from volcanic ash in Central America, Forsythe (1975) reported 2-hour rates of $70\text{--}210 \text{ mm h}^{-1}$. Our results for unfrozen Icelandic Andisols extend this range substantially in both directions ($28\text{--}369 \text{ mm h}^{-1}$). TIRs in Icelandic Andisols were generally higher in the sandier soils at the G-site ($102\text{--}369 \text{ mm h}^{-1}$), compared to in the

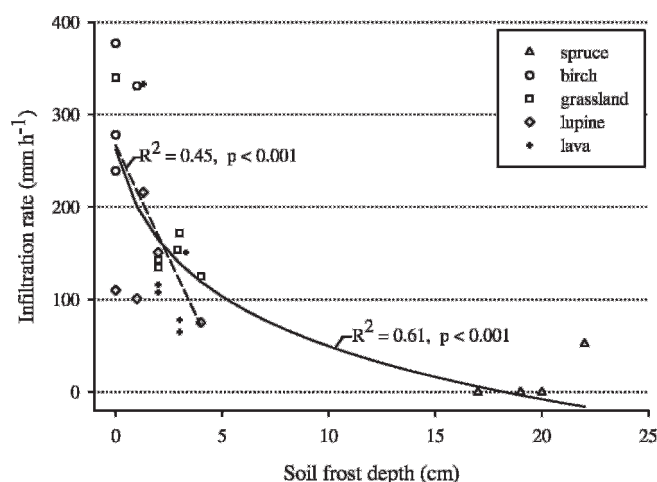


FIGURE 6. Relationship between terminal infiltration rate (TIR; mm h^{-1}) and soil frost depth (cm) in April 2000, in plant communities at Gunnarsholt. Non-linear fit (all data [TIR = $262.11 - 204.17 \times \log(\text{frost depth} + 1)$]) and linear fit (excluding G-spruce [TIR = $266.63 - 48.69 \times (\text{frost depth})$]) are presented.

finer textured soils at the H-site (28–94 mm h⁻¹) (Table 1, Fig. 5). Land cover influences superimposed against this backdrop of texture effects contributed to the broad range of observed TIRs. For example, TIR in G-birch woodland was about 4× that in the H-birch woodland and TIR in G-grassland was 2–6× that of H-grassland.

Plant life-form effects on TIR were substantial (Fig. 5) and these have significant implications for management and restoration of Iceland's young, erosion-prone soils. TIRs were typically highest in woodlands dominated by native birch. The high TIRs in birch woodlands may be attributed to the dominance of shrub growth forms, which typically augment infiltration by enhancing soil structure (Thurow, 1991) and macropore formation (Beven and Germann, 1982). Lower TIRs in the G-spruce compared to G-birch woodland may reflect the sparser plant ground cover in the spruce woodland (Fig. 3). Birch stands are also known to support greater earthworm biomass than spruce stands in Iceland (Gudleifsson, 2007). A greater abundance of other soil biota (Saetre, 1998) which alter soil structure to promote infiltration may also foster higher rates of infiltration in birch communities.

Variability in summer TIRs within plant communities was relatively small, with one exception. In the H-grassland, TIR was three times higher in July/August 2000 compared to July/August 1999 and May 2000 (Fig. 5). Horses were grazing the site in July/August 1999, whereas in July/August 2000 the area had been excluded from grazing for one year and cover and biomass of vegetation were noticeably higher (qualitative observations). We hypothesize that the grazing exclusion led to increased plant root production which acted in combination with winter freeze-thaw events to counteract soil compaction caused by hoof action. The threefold greater TIR, observed one year after grazing was discontinued, suggests that Andisols are hydrologically resilient where vegetation cover is continuous and soil OC content high. Low TIR in H-grassland compared to H-birch communities in summer 1999 may thus reflect grazing rather than plant life-form effects. In the absence of grazing, H-grassland sites may have TIRs comparable to those of birch woodlands in unfrozen soils.

SOIL FROST DYNAMICS AND INFILTRATION RATES IN WINTER

TIRs in frozen versus unfrozen soils were generally comparable within the G- and H-birch woodlands, G-lupine, and G-lava field communities (Fig. 5). These sites all had continuous, relatively deep (≥25 cm) snow cover in January (Fig. 4). By contrast, TIRs in frozen and unfrozen soils associated with the G-spruce woodland and G- and H-grasslands varied dramatically. Moreover, the direction of change in TIR varied from summer to winter, with summer rates being highest in G-spruce woodland and the H-grassland, and winter (January) TIR being highest in the G-grassland. These summer vs. winter differences, or the lack thereof, appear to be related to the distinctive soil frost conditions occurring in each plant community.

Deep soil frost was associated with shallow snow depth (Fig. 4), as expected (e.g. Tyrtikov, 1976; Hardy et al., 2001). Shallow snow cover (≤11 cm) was observed in the open H-grassland community where snow was directly exposed to wind and solar radiation, but also in the G-spruce woodland where the dense nature of the spruce canopy seemed to cause high interception loss allowing little snow to reach the forest floor (Shanley and Chalmers, 1999; Jones et al., 2001). Infiltration-retarding concrete and porous concrete soil frost formed in these communities, whereas soils in the other communities developed

porous frost. H-grassland and G-spruce had very high surface soil water retention (>100%; Table 1); but despite similar water retention in the H-birch woodland, infiltration-retarding soil frost did not form there. Deeper snow cover in the H-birch woodland and/or lower soil water content at the time of freezing might be the explanation, but the greater snow cover can likely be attributed to woodland vegetation structure (Tyrtikov, 1976; Jones et al., 2001). Topography (slope inclination; Table 1) appears to account for the difference in snow cover in the western vs. eastern portions of the G-site. Our observations reveal the importance of studying the influence of soil water content and snow cover at the time of freezing on the type and depth of frost formation in Andisols.

Deep concrete soil frost greatly reduced winter TIRs in the spruce woodland. Concrete frost has been found to hinder infiltration completely at frost depths from ca. 8 to 10 cm, measured at small scales (Post and Dreibeilbis, 1942; Megahan and Satterlund, 1962), and to reduce infiltration significantly at spatial scales ranging from points to small catchments (e.g. Dunne and Black, 1971; Kane and Stein, 1983; Blackburn and Wood, 1990; Shanley and Chalmers, 1999). In the spruce woodland, water infiltrated in only two (January) and one (April) of the six infiltrometers. Although the deep matrix of concrete soil frost was largely impermeable, root channels and other macropores that remain free of ice (Trimble et al., 1958; Megahan and Satterlund, 1962; Dunne and Black, 1971) may have allowed modest infiltration to occur in those rings. The negative exponential relationship between soil frost depth and TIR in the five G-site plant communities in April (Fig. 6) supports the existence of such water conducting soil macropores. The logarithmic curve is consistent with what has been described for temperate forests (Rybakova, 1990), where water normally infiltrates through coarse root channels and voids created by soil organisms and freeze-thaw activity.

Porous concrete soil frost and patchy ice cover, common between hummocks, characterized the H-grassland in January when TIR was reduced compared to other times of the year. Even rings with partial or no ice cover had TIRs near zero, which is consistent with field observations of standing water at this site. However, by April, TIR had returned to pre-winter conditions (Fig. 5) and was not influenced by soil frost, despite the fact that soil frost conditions were comparable to those in January (Fig. 4). These seemingly contradictory results may be an artifact of biased sampling of the soil in January, as frost depth could only be determined where water had not ponded. The limited number of measurements made at this time ($n = 3$) may therefore have underestimated average frost depth. Reports on the influence of porous concrete frost on infiltration are few and inconsistent. Stoeckeler and Weitzman (1960) measured reduced rates in soils characterized by porous concrete frost compared to unfrozen soil, whereas Haupt (1967) measured reductions in infiltration on burned or sparsely vegetated plots, but not on plots where plant and litter cover was high. Our results imply that porous concrete frost can reduce infiltration in well-vegetated Andisols (H-grassland), but that depth of frost penetration may govern the influence on TIRs.

Plant communities characterized by soils with porous frost had comparable TIRs in frozen versus unfrozen soils (Figs. 4 and 5) with one exception. G-grassland TIRs in January were elevated relative to those in summer, which may be attributable to increases in the number and/or size of pore spaces associated with frost action. Despite similar soil frost conditions in January and April, G-grassland April TIRs were lower than those recorded in January. This may be due to increased soil ice content following

freeze-thaw cycles during snowmelt periods in January and March (Fig. 2), as snowmelt infiltration followed by soil freezing would increase soil ice content (Johnsson and Lundin, 1991).

Conclusions

The high summer TIRs observed in this study imply that runoff is likely minimal on vegetated, level, unfrozen Andisols, especially since most precipitation comes in relatively small pulses. During winter months, infiltration-retarding soil frost occurred in herbaceous land cover type where exposure to wind and solar radiation limit snow accumulation. Deep, infiltration-retarding, concrete-type soil frost also formed in spruce woodland, a relatively new Icelandic ecosystem type (Pétursson, 1999). Concrete soil frost types were not observed in birch woodlands, and their high snow capture potential make them less prone to such frost formation. Even so, the extensive snow cover in winter 1999–2000 likely resulted in less extensive soil frost than might be expected in average years. Results from winters characterized by frequent thaw periods, and more shallow and ephemeral snow cover may thus prove different. Nonetheless, in this study the soils of birch woodlands were characterized by porous frost and exhibited high TIRs whether frozen or unfrozen. We thus expect that hydrological processes and soil frost dynamics may have changed significantly with the extensive loss of the birch woodlands in Iceland, leading to conditions that would be more conducive to water erosion during winter warm spells and periods of rainfall and snow melt in winter and spring. Open, sparsely vegetated areas have increased substantially in areal extent, because of land degradation. The tendency of such sites to develop deep, infiltration-retarding soil frost in winter suggests that these landscapes would be a likely source of runoff much of the year. Frequent river flooding during winter thaw events (Rist, 1990) supports this hypothesis. We also saw evidence for reductions in TIRs associated with reduced vegetation cover caused by livestock grazing, a widespread, traditional land use of the Icelandic rangelands. Effects of sheep grazing in high altitude Andisolic grasslands has been shown to create conditions for eolian erosion, runoff, and gully erosion (Podwojewski et al., 2002). Extensive harvest of the native birch woodlands in concert with continuous livestock grazing by settlers in Iceland (Aradóttir and Arnalds, 2001) may therefore have combined to effect local changes in hydrologic function that promote soil erosion.

Acknowledgments

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