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# Geochemistry of Soils and Streams on Surfaces of Varying Ages in Arctic Alaska

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## Abstract

Climate change may bring about geochemical changes in arctic regions as a result of increasing thaw depth. In order to better understand current watershed geochemistry and mineral weathering and provide a basis for predicting the geochemical effects of active-layer thickness increase, we examined elemental chemistry and  $^{87}\text{Sr}/^{86}\text{Sr}$  of streams and sequential and total digests of soils, permafrost, and soil parent materials from seven glacial deposit surfaces of varying geomorphic ages in arctic Alaska in the vicinity of the Philip Smith Mountains quadrangle (69°N, 150°W). We found overall greater exchangeable K concentrations, exchangeable and acid digestible P and Ca concentrations, acid digestible and total Ca/Na and Ca/Sr, and lower acid digestible  $^{87}\text{Sr}/^{86}\text{Sr}$  in permafrost than in active-layer mineral soil. Of the surfaces with similar parent material, stream and soil data suggest that weathering has progressively depleted calcium carbonate in the active layer with increasing surface age. Our results suggest that increasing thaw depth will lead to increasing carbonate and Ca supply to soils and streams, as well as spatially variable increases in P and K supply. Geochemical differences between active-layer soil and permafrost suggest the possibility of using stream geochemistry to detect changes in active-layer thickness in watersheds.

## Introduction

Chemical weathering of minerals is an integral part of soil formation and is ecologically important as a source of nutrients such as phosphorus, potassium, and calcium (Schlesinger, 1997); silicate weathering is geologically important as a sink for atmospheric carbon in the long-term carbon cycle (Walker et al., 1981). It is important to understand mineral weathering and related soil and stream geochemistry in arctic regions because global climate change may cause changes in both the nature and extent of mineral weathering in these regions. Continuous permafrost in arctic regions restricts mineral weathering to the active (seasonally thawed) layer, and warming climate is likely to cause increased active-layer thickness (Anisimov et al., 1997). Increased active-layer thickness will allow minerals previously contained in permafrost to weather, thereby increasing the total mineral surface area that is exposed to weathering. Additionally, permafrost in some areas may contain very soluble minerals that largely have been removed by weathering in overlying thawed soils. Thawing and weathering of these minerals could cause potentially important changes in stream and soil geochemistry for years to come; the nature of these changes will depend on the spatial and temporal progression of permafrost degradation.

Much progress has been made toward understanding soil chemistry and mineral weathering on the North Slope of Alaska. Soils on the North Slope were classified and physically described in early work by Tedrow et al. (1958) and in more recent studies by Ping et al. (1998) and Munroe and Bockheim (2001). In an investigation of the soil chemistry differences between non-acidic and acidic tundra, Bockheim et al. (1998) analyzed soil exchangeable chemistry and other soil characteristics across the Kuparuk River Basin of the North Slope, an area that overlaps the study area of this paper and includes loess, till, and colluvium deposits of different ages. Their study found a relationship between non-

acidic tundra soils and greater amounts of exchangeable base cations. Munroe and Bockheim (2001) found that among the four most recently deglaciated surfaces in this region, soil profile concentrations of weathering products (clay and silt) increased significantly with age, suggesting that even in areas where cryoturbation is common, surface age is a factor in mineral weathering and soil development.

Although cycling of biogeochemically important elements such as carbon and nitrogen in soils and catchments of the Alaskan Arctic has been the subject of many studies (e.g. Giblin et al., 1991; Nadelhoffer et al., 1991; Hobbie et al., 2002; Judd and Kling, 2002), the geochemistry of other elements including macro- and micronutrients released by mineral weathering is less well known. Stutter and Billet (2003) described the stream and soil chemistry of a Swedish arctic region more recently glaciated than the study area of this paper, but with similar sedimentary bedrock (including sandstone and limestone units), and found the system to be dominated by carbonate dissolution. Kling et al. (1992) investigated the chemistry of lakes and rivers over a wide region of the Alaskan North Slope and interpreted the water chemistry in the region of their study to also be dominated by the products of carbonate dissolution and related to surface age.

Both terrestrial (Shaver and Chapin, 1995) and aquatic (Hobbie et al., 1999) ecosystems in this region are nutrient limited, either by P alone or by N and P. Therefore, enhanced release of mineral nutrients via weathering of newly thawed minerals may have an important ecological impact. Hobbie et al. (1999) measured a twofold increase in P concentrations in a stream passing through glacial deposits disturbed by construction and predicted that weathering of thawed permafrost minerals may increase aquatic P availability, causing bottom-up trophic effects.

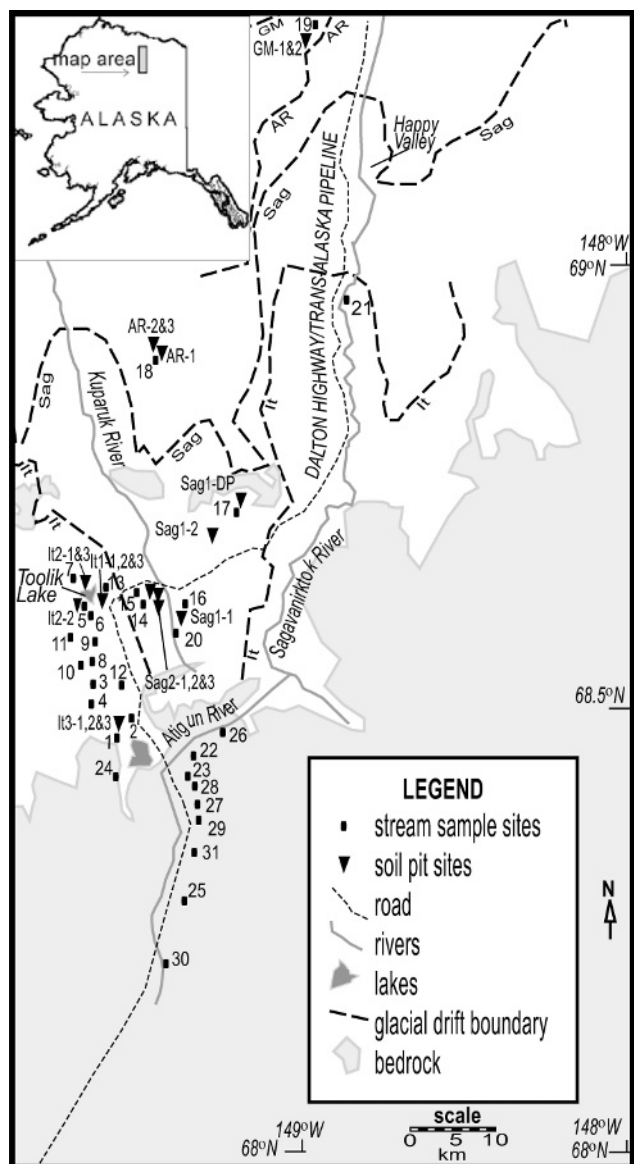
Because of the potential for a unique, permafrost-related geochemical response of arctic systems to climate change, it is

important to understand the current mineral weathering processes and geochemical characteristics of these systems. This study describes the elemental chemistry of soils and streams in the central Alaskan North Slope, examines mineral weathering as a function of surface age, and addresses the geochemical implications of thaw depth increases brought about by climate change. We analyzed stream water and several chemical fractions of soil and bedrock, and used elemental and Sr isotope ratios to identify the mineral sources of weathering products. These results will serve to further our understanding of weathering in arctic regions, as well as provide baseline information for predicting how warming climate may influence the biogeochemistry of ecosystems.

## Study Area

Field research was conducted on the eastern North Slope of arctic Alaska near the foothills of the Brooks Range (Fig. 1), primarily in the Philip Smith Mountains quadrangle (69°N, 150°W). The entire region is underlain by continuous permafrost (Brown and Krieg, 1983). For the years 1990–2000, the mean active-layer thickness at three sites within the study area (Happy Valley, Imnavait Creek, and Toolik Lake) ranged from 33 to 60 cm (Brown et al., 2002); the maximum thaw depth we observed during field work was 80 cm. Mean annual air temperatures on this part of the North Slope range from  $-5.9^{\circ}\text{C}$  in the Brooks Range to  $-12.8^{\circ}\text{C}$  at Prudhoe Bay (Haugen, 1982). The dominant vegetation types include tussock tundra, wet sedge tundra, and riparian willow communities (Giblin et al., 1991). The landscape descends from the steep, rocky foothills of the Brooks Range to gently rolling tundra with poorly developed drainage and many kettle and thermokarst lakes.

North of the foothills, the landscape is developed on till and outwash formed during several glaciations of varying ages, during which glaciers moved northward from the Brooks Range (Hamilton, 1978; Brown and Krieg, 1983; Hamilton, 2003). The oldest and most northerly recognizable glacial deposits resulted from the Gunsight Mountain (GM) glaciation, which is inferred to have occurred in the late Tertiary based on the similarity of erosional features to Pliocene deposits south of the Brooks Range (Hamilton, 1994). The second oldest glacial deposits resulted from the Anaktuvuk River (AR) glaciation, which is assigned an early Pleistocene age by paleomagnetic analysis of overlying sediments (Hamilton, 1986) and correlations with other Alaskan glacial sequences (Hamilton, 1994). The next two younger glaciations are each broken up into separate units consisting of advances and readvances separated by interglacial periods. The Sagavanirktok River glaciation is dated broadly to the middle Pleistocene (780,000–125,000 years B.P.; Hamilton, 2003) based on paleomagnetic data and correlation to glaciations elsewhere in Alaska (Hamilton, 1994), and is separated into two units: Sagavanirktok River 1 (Sag1) and 2 (Sag2). At least five separate phases of the Itkillik glaciation have been identified (Hamilton, 2003), but for the purpose of simplification in this paper we use the three main phases initially described by Hamilton and Porter (1975) and used by Hamilton (1986): Itkillik I (It1), Itkillik II (It2), and the latest Itkillik II re-advance (It3). It1 deposits are beyond the range of radiocarbon dating ( $>40$  ka) but have an inferred age between 50 and 120 ka; radiocarbon dating suggests ages between approximately 11.5 and 24 ka for It2 deposits and 11.4 ka and 12.8 for It3 deposits (Hamilton, 1986, 2003). These surfaces have loess cover ranging from greater than 15 m on the oldest surfaces, through thin but still continuous cover on intermediate-aged surfaces, to



**FIGURE 1.** Map of study area with locations of soil (triangle) and stream (square) sampling sites. Glacial deposit and bedrock boundaries after Hamilton (1978, 2003), Mull and Adams (1985), and Brown and Krieg (1983).

loess-free moraine crests on the youngest surfaces (Hamilton, 1994, 2003).

The bedrock formations in the Brooks Range foothills, from which the glacial till is derived, consist primarily of conglomerates, sandstones, and limestones, with some shale and phyllite (Menzie et al., 1985; Mull and Adams, 1985). Based on the gentle topography of the youngest (It3) moraines, Hamilton (2003) inferred that they contain abundant fine sediments derived from large nearby lake beds and therefore may be compositionally different from other glacial deposits.

The soils developed on these deglaciated surfaces are classified broadly as Gelisols, and more specifically as Turbels and Orthels, depending on the amount of apparent cryoturbation (Munroe and Bockheim, 2001). In most areas, the mineral soil is overlain by a thick (10 or more centimeters) organic horizon. The soils are characterized by medium texture, poor drainage, and cryogenic features such as warped horizons and ice lenses (Ping et al., 1998).

## Methods

### WATER SAMPLING AND ANALYSIS

We sampled small first- or second-order streams in watersheds entirely contained within each of the seven glacial surfaces of different ages in the study area, and within each of the major bedrock types in the foothills north of the Brooks Range (Fig. 1); we also sampled the major rivers (up to fourth order) in the study area. Precipitation samples were collected in acid-washed polyethylene containers. All water samples were filtered through 0.45  $\mu\text{m}$  polypropylene filters into acid-washed polyethylene bottles in the field, acidified with ultra-pure hydrochloric acid, and refrigerated at  $<10^{\circ}\text{C}$  until analysis.

All water samples were analyzed using a Perkin Elmer Optima 3300 DV Inductively Coupled Plasma Optical Emission Spectrometer (ICP-OES) using five- to eight-point calibration curves for element concentrations (including Ba, Ca, Fe, K, Mg, Na, S, Si, and Sr). Two High-Purity<sup>®</sup> standards (Trace Metals in Drinking Water and River Sediment-B) and one in-house standard were used for quality control. These standards were analyzed to within  $\pm 10\%$  of the known values for Fe and Sr;  $\pm 17\%$  for S, and  $\pm 5\%$  for all other elements.

### SOIL SAMPLING AND ANALYSIS

During late July and August of 2002 and 2003, two to four soil sampling pits on each glacial deposit surface were excavated (Fig. 1), with the exception of the GM surface, on which a single large trench was excavated and sampled on opposite sides. Sites were randomly selected on flat (slope  $<2^{\circ}$ ) moraine crests or shoulders. Vegetation at the pit sites indicated that all of the sites except one were located on moist acidic tundra; one of the It2 pit sites was located on non-acidic tundra. Each pit was excavated through active-layer soil and at least several cm into frozen ground. Soil profiles and visible horizons were measured and described, and homogenized samples were taken from each visible horizon. We took 3–6 horizon samples per pit, usually consisting of layers designated “O” (peat), “A” (an oxidized layer with live roots), “B1” (the upper section of a thick gleyed layer), “B2” (the lower section of the same layer; this division was meant to split the thick “B” horizon approximately in half in order to look for differences with depth), “P1” (the first few centimeters of thickly laminated frozen soil, which may infrequently be part of the active layer), and “P2” (deeper, thinly laminated sediment in permafrost). Several soil profiles contained evidence of cryoturbation such as irregular horizon boundaries or buried peat inclusions. A deeper ( $\sim 3$  m) permafrost sample (labeled Sag1-DP on Fig. 1 and in tables) was obtained from the Sag1 surface from the site of a thermokarst tundra collapse within a few days of that event.

Soil samples were dried at  $35^{\circ}\text{C}$ , and mineral soils were sieved to separate the  $<2$ -mm size fraction, which was used for further analyses. We consider the  $>2$  mm fraction to contribute a negligible amount to overall mineral weathering of the soil due to its low ratio of surface area to volume; the  $>2$  mm fraction appeared to consist mostly of chert pebbles and quartzite cobbles and was on average less than 35% by mass of each sample. Percent organic matter was determined for samples from one representative pit on several surfaces by loss on ignition (LOI) via combustion at  $450^{\circ}\text{C}$  for 24 h, following the method of Heiri et al (2001).

Following the methods of Blum et al. (2002), 0.5 g soil samples were leached and digested sequentially with 5 mL of each of the following ultra-pure solutions: (1) 1M  $\text{NH}_4\text{Cl}$  (pH = 7) at

room temperature for 20 h to obtain the exchangeable fraction; (2) 1M  $\text{HNO}_3$  at room temperature for 20 h to dissolve easily soluble minerals (which in these soils are mostly carbonates but include minor amounts of phosphate and some sulfides) and to leach remaining labile elements bound to organics; and (3) concentrated  $\text{HNO}_3$  at  $150^{\circ}\text{C}$  for 3 h to partially digest less soluble but still potentially weatherable minerals such as biotite, plagioclase, potassium feldspar, sulfides, and some oxides and clays. For convenience, we will refer to the solutions obtained via the second and third steps in the sequential digest procedure as the “cold acid digestible” and “hot acid digestible” fractions, respectively. While we aim to estimate “easily weatherable” and “less weatherable” fractions of the soil with these sequential digests, these digests are operationally defined and thus do not represent clearly defined reservoirs within the soil. Furthermore, while elemental P and S (possibly occurring as  $\text{PO}_4^{3-}$  and  $\text{SO}_4^{2-}$ ) released by  $\text{NH}_4\text{Cl}$  are not necessarily “exchangeable,” they represent the most labile fraction of those elements, and for convenience they are referred to as part of the exchangeable fraction.

To obtain a total digest of mineral soil samples, about 0.1 g of each sample was fused with 1 g of technical grade  $\text{LiBO}_2$  in a graphite crucible at  $1100^{\circ}\text{C}$ , then dissolved in approximately 60 mL of trace metal grade 5%  $\text{HNO}_3$  and filtered through 25- $\mu\text{m}$  pore size cellulose fiber paper. Two procedural blanks ( $\text{LiBO}_2$  with no sample added) and two U.S. Geological Survey Geochemical Reference Standards were also digested.

The soil leachate and digest solutions were analyzed for elemental concentrations by ICP-OES using six- to nine-point calibration curves. Quality control standards included High-Purity<sup>®</sup> ICP-Stock Solution for Si and CRM Soil Solution A and Trace Metals in Drinking Water for other elements. For the sequential digest solutions, these standards were analyzed within  $\pm 7\%$  for Ba, Ca, Fe, K, Na, and Sr;  $\pm 12\%$  for Si, and  $\pm 14\%$  for P and S in the same concentration ranges as the samples. For the total digests, these standards plus an in-house standard were analyzed to within  $\pm 7\%$  for Ba, P, and Si; and  $\pm 12\%$  for Ca, Fe, K, Na, S, Sr in the same concentration ranges as the samples.

### BEDROCK SAMPLING AND ANALYSIS

Samples of each major rock unit on the eastern North Slope (Philip Smith quadrangle) were collected during the summer of 2002. Where possible bedrock samples were collected from outcrops in the same watersheds in which streams were sampled.

Polished thin sections were made of each rock sample, and mineral constituents were identified visually using a petrographic microscope or by an energy dispersive spectrometer (EDS) on a Hitachi S3200N scanning electron microscope (SEM). Sub-samples of each rock sample were pulverized in a tungsten carbide ring mill and then subjected to the same sequential leach and digest procedure described above. After the hot acid digest, the remaining sample was digested with a mixture of concentrated ultra-pure HF,  $\text{HNO}_3$ , and HCl until all sample was in solution. The leachate and digest solution was analyzed by ICP-OES as described above.

### SR ISOTOPIC ANALYSES

Sr was separated from selected water samples and soil and rock leachate and digest solutions by eluting a subsample through Eichrom Sr-Spec resin in a quartz cation exchange column. Fifty to 100 ng of Sr along with 1  $\mu\text{L}$  of  $\text{H}_3\text{PO}_4$  were then loaded onto a tungsten filament with  $\text{Ta}_2\text{O}_5$  powder matrix.  $^{87}\text{Sr}/^{86}\text{Sr}$  was



TABLE 1

Chemistry of local rock units likely incorporated into glacial till deposits. Formation names are defined in the text.

Fm. name	Rock type	Fraction	Ba ( $\mu\text{mol/g}$ )	Ca ( $\mu\text{mol/g}$ )	Fe ( $\mu\text{mol/g}$ )	K ( $\mu\text{mol/g}$ )	Mg ( $\mu\text{mol/g}$ )	Na ( $\mu\text{mol/g}$ )	P ( $\mu\text{mol/g}$ )	S ( $\mu\text{mol/g}$ )	Si ( $\mu\text{mol/g}$ )	Sr ( $\mu\text{mol/g}$ )	Ca/Na (molar)	Ca/Sr (molar)	$^{87}\text{Sr}/^{86}\text{Sr}$
<b>Db</b> s	phyllite/slate	exch.	0.051	25.3	1.22	6.92	13.6	2.64	<0.09	1.77	0.83	0.110	9.6	230	0.727052
<b>Dke</b>	ss/cong.	exch.	0.031	2.94	0.63	9.68	4.12	1.59	<0.09	0.15	0.30	0.020	1.8	147	0.723253
<b>PMI</b>	limestone	exch.	0.009	132	0.28	1.19	2.63	0.40	0.23	<0.08	0.36	0.092	326	1430	0.708986
<b>Pe</b>	shale	exch.	1.35	30.9	1.28	12.7	10.9	0.53	<0.09	0.42	4.79	0.097	58.7	319	0.715667
<b>Kf</b>	sandstone	exch.	2.57	11.9	0.14	20.4	19.4	4.50	<0.09	0.40	0.64	0.052	2.6	227	0.717860
<b>Kn</b>	sandstone	exch.	0.277	35.1	0.47	6.46	12.3	4.35	<0.09	4.13	<0.06	0.066	8.1	533	0.713092
<b>Db</b> s	phyllite/slate	cold dig.	0.016	660	348	4.17	314	0.62	20.4	54.6	60.4	0.831	1060	793	0.725383
<b>Dke</b>	ss/cong.	cold dig.	0.085	12.3	67.5	6.12	6.73	0.47	2.88	<0.08	39.2	0.029	26.3	425	0.719324
<b>PMI</b>	limestone	cold dig.	0.044	5970	1.02	1.01	35.9	0.09	17.1	<0.08	6.66	1.69	64000	3530	0.708058
<b>Pe</b>	shale	cold dig.	2.97	55.4	80.6	8.29	71.3	<0.073	2.01	1.74	44.6	0.103	>758	540	0.714122
<b>Kf</b>	sandstone	cold dig.	2.49	45.7	141	12.5	97.5	3.12	21.2	2.71	97.3	0.096	14.7	477	0.712404
<b>Kn</b>	sandstone	cold dig.	0.415	85.1	127	2.69	62.4	<0.073	15.8	23.7	84.5	0.122	>1160	695	0.713419
<b>Db</b> s	phyllite/slate	hot dig.	<0.001	15.1	518	13.2	252	<0.073	<0.09	19.2	350	0.060	>207	251	0.731134
<b>Dke</b>	ss/cong.	hot dig.	<0.001	<0.04	229	20.1	8.04	<0.073	2.30	0.15	150	0.060	—	<0.59	0.723511
<b>PMI</b>	limestone	hot dig.	0.019	534	2.38	1.19	2.17	<0.073	1.37	0.18	14.0	0.099	>7310	5380	0.713099
<b>Pe</b>	shale	hot dig.	0.097	<0.04	365	51.5	196	<0.073	<0.09	1.29	184	0.022	—	<1.60	0.753962
<b>Kf</b>	sandstone	hot dig.	1.52	<0.04	189	57.1	127	11.1	3.73	2.50	281	0.031	<0.003	<1.11	0.735151
<b>Kn</b>	sandstone	hot dig.	0.610	5.67	334	41.9	210	1.27	0.37	6.44	495	0.057	4.5	100	0.727727
<b>Db</b> s	phyllite/slate	res. dig.	2.66	<0.04	91.3	452	1.32	290	0.28	1.48	1090	0.263	>0.0001	<0.13	0.740233
<b>Dke</b>	ss/cong.	res. dig.	<0.001	<0.04	109	27.8	6.26	<0.073	0.95	<0.08	1560	0.208	—	<0.17	0.714283
<b>PMI</b>	limestone	res. dig.	0.035	1.23	0.17	0.50	<0.0001	<0.073	<0.09	0.47	547	0.007	>16.4	168	0.708846
<b>Pe</b>	shale	res. dig.	1.22	<0.04	259	583	41.2	91.3	0.22	1.25	780	0.412	<0.0003	<0.0850	0.741406
<b>Kf</b>	sandstone	res. dig.	2.30	<0.04	26.9	50.7	29.2	95.1	0.29	1.82	939	0.162	<0.0004	<0.22	0.715210
<b>Kn</b>	sandstone	res. dig.	4.31	5.83	91.9	319	44.2	273	1.65	1.00	926	0.638	0.02	9.1	0.723082

PMI elemental data and cold digestible  $^{87}\text{Sr}/^{86}\text{Sr}$  represent a mean of data from 5 samples of the PMI from different locations and members; other PMI  $^{87}\text{Sr}/^{86}\text{Sr}$  data and data for all other units are from one representative sample. Errors for elemental data and  $^{87}\text{Sr}/^{86}\text{Sr}$  reported in text. Exch. = exchangeable fraction; cold dig. = cold acid digestible fraction; hot dig. = hot acid digestible fraction; res. dig. = residual (HF) digestible fraction.

determined by multiple-collector thermal ionization mass spectrometry (TIMS) on a Finnigan MAT 262. To correct for instrumental mass bias,  $^{86}\text{Sr}/^{88}\text{Sr}$  was normalized to 0.1194. Between 50 and 200 replicate  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios were measured for each sample. Internal precision ( $\pm 2\sigma$ ) calculated from these replicates was generally less than  $\pm 0.000030$ ; exceptions are noted in tables and graphs. The Sr standard NBS-987 was analyzed after every 12 samples with a mean value of  $0.710231 \pm 0.000016$  ( $2\sigma$ ) during the period of sample analyses.

## Results

### BEDROCK COMPOSITION AND GEOCHEMISTRY

Based on the location and extent of rock unit outcrops in the eastern Brooks Range foothills and on cobbles found in soil pits by us and by Hamilton (2003), we identified six units (as mapped by Mull and Adams, 1985) likely to comprise the bulk of the glacial till in the study area: the Devonian Beaucoup phyllite (Db)s, the Devonian Ear Peak sandstone member of the Kanayut conglomerate and sandstone (Dke), the Permian–Mississippian Lisburne limestone (PMI), the Permian Echooka shale (Pe), the Cretaceous Fortress Mountain conglomerate and sandstone (Kf), and the Cretaceous Nanushuk sandstone (Kn). Because the Kn outcrops north of the Sagavanirktok River glacial deposits, it is likely included only in the older and more northerly Gunsight and Anaktuvuk glacial deposits.

Visual and SEM thin section analyses suggest that all three sandstones (Dke, Kf, and Kn) are composed primarily of quartz and chert and contain minor amounts of biotite/vermiculite, muscovite, and oxide and sulfide minerals; the Kf and Kn also

contain minor plagioclase and glauconite. The limestone unit (PMI) appeared chert-rich and argillaceous, which is consistent with the description of this unit by Moore et al. (1994).

The composition of sequential and bulk digests of these six units is reported in Table 1. As expected, the limestone had the lowest  $^{87}\text{Sr}/^{86}\text{Sr}$  values and highest Ca/Sr of the six units; the phyllite (Db)s and the Devonian sandstone (Dke) had the highest exchangeable and cold acid digestible fraction  $^{87}\text{Sr}/^{86}\text{Sr}$  values. The elemental ratios Ca/Na and Ca/Sr are often used as indicators of calcium carbonate contribution because calcium carbonate has high Ca but low Na and Sr concentrations, while the reverse is true for many common silicate minerals. Strontium isotope ratios can be used similarly because most carbonates have low  $^{87}\text{Sr}/^{86}\text{Sr}$  while many silicate minerals have higher  $^{87}\text{Sr}/^{86}\text{Sr}$  values (e.g. Palmer and Edmond, 1992). Because the carbonate and silicate rocks in this area follow these general patterns, we use the elemental concentration ratios Ca/Na and Ca/Sr and the isotope ratio  $^{87}\text{Sr}/^{86}\text{Sr}$  as indicators of the calcium carbonate contribution to soil fractions and stream water solutes.

### SOIL GEOCHEMISTRY

Geochemical characteristics of the soil exchangeable fraction, cold acid digestible fraction, hot acid digestible fraction, and total soil digests are reported in Tables 2–5. The It2 surface has the highest exchangeable Ca concentrations, Ca/Na, and Ca/Sr (Table 2). In the mineral soil of most profiles, there is a trend of increasing Ca concentrations and Ca/Sr and decreasing  $^{87}\text{Sr}/^{86}\text{Sr}$  with depth (Table 2). Statistical tests (ANOVA or Welch ANOVA where the variances are unequal) were performed on concentration

TABLE 2

Mean exchangeable soil fraction (per gram dry soil). Surface and horizon names are defined in the text.

Surface	Horizon	Depth (cm)	Ba ( $\mu\text{mol/g}$ )	Ca ( $\mu\text{mol/g}$ )	Fe ( $\mu\text{mol/g}$ )	K ( $\mu\text{mol/g}$ )	Na ( $\mu\text{mol/g}$ )	P ( $\mu\text{mol/g}$ )	S ( $\mu\text{mol/g}$ )	Si ( $\mu\text{mol/g}$ )	Sr ( $\mu\text{mol/g}$ )	Ca/Na (molar)	Ca/Sr (molar)	$^{87}\text{Sr}/^{86}\text{Sr}^{**}$
It3 ( $n = 4$ )	O	0–7	0.163	92.0	3.31	7.12	0.22	0.58	0.91	2.49	0.087	420	1062	0.713660
	A	7–19	0.296	7.29	0.80	1.42	0.10	0.02	0.07	1.05	0.011	70.4	687	0.720245*
	B1	19–39	0.374	6.55	0.98	1.49	0.12	0.03	0.14	0.60	0.011	55.8	577	0.720483
	B2	39–60	0.518	7.27	0.87	2.61	0.11	0.02	0.19	0.41	0.013	63.6	543	0.719548
	P	62–71	0.779	13.1	3.64	4.62	0.16	0.10	0.47	1.87	0.027	84.1	490	0.716431
It2 ( $n = 3$ )	O	0–16	0.448	174	1.63	1.12	0.34	0.43	2.88	0.80	0.109	509	1596	0.709544
	A	18–29	0.700	52.7	2.31	0.81	0.11	0.12	0.11	1.75	0.041	489	1282	0.710457
	B1	29–46	0.753	57.9	1.99	1.65	0.15	0.08	0.48	1.92	0.040	395	1450	0.709485
	B2	46–62	0.428	83.3	3.95	2.14	0.20	0.15	2.82	3.05	0.060	425	1397	0.709496*
	P	66–73	0.302	88.5	1.85	2.47	0.55	0.10	4.74	1.12	0.071	162	1244	0.709585
It1 ( $n = 3$ )	O***	0–7	0.384	20.5	2.69	7.10	0.20	0.18	1.41	0.44	0.033	105	627	0.721412*
	A	8–23	0.296	4.71	1.03	0.64	0.06	0.02	0.12	0.31	0.007	78.1	657	0.718664
	B1	23–40	0.373	5.48	0.60	0.69	0.06	0.04	0.11	0.22	0.009	93.2	642	0.718510*
	B2	35–49	0.552	6.80	1.50	0.75	0.10	0.05	0.07	0.82	0.010	65.5	701	0.717530
	P1	49–60	0.501	7.38	2.98	0.96	0.10	0.07	0.10	1.60	0.009	75.5	797	0.715944
	P2	62–69	0.489	34.7	0.42	1.76	0.10	0.04	0.14	0.28	0.037	359	951	0.711278
Sag2 ( $n = 3$ )	O	0–8	0.242	32.0	2.72	6.55	0.34	0.68	1.49	0.91	0.048	93.7	670	0.716872
	A	8–25	0.409	4.77	0.58	0.85	0.10	0.05	0.15	0.35	0.009	47.9	541	0.720050
	B1	25–42	0.861	14.2	1.93	1.43	0.23	0.06	0.09	1.52	0.023	61.8	612	0.720851*
	B2	38–55	0.365	3.05	1.52	0.66	0.15	0.04	0.11	1.33	0.006	19.9	504	0.717221
	P1	55–69	0.495	5.39	0.95	1.24	0.15	0.02	0.08	0.49	0.011	35.2	486	0.715970
	P2	71–75	0.861	21.5	1.41	2.43	0.20	0.06	0.14	0.78	0.030	105	709	0.716069*
Sag1 ( $n = 2$ )	O	0–13	0.543	30.6	3.13	3.56	0.29	0.21	1.18	0.65	0.044	106	696	0.717314
	A	13–22	0.611	12.3	1.76	1.22	0.14	0.08	0.34	1.20	0.018	88.0	668	0.717041
	B1	22–40	0.580	5.89	0.45	0.82	0.15	0.04	0.10	0.37	0.009	39.0	635	0.719167
	B2	35–50	0.727	7.17	0.85	1.29	0.16	0.03	0.15	0.54	0.011	44.3	630	0.717241*
	P	50–58	0.680	7.24	2.85	1.62	0.17	0.07	0.20	1.84	0.011	41.5	656	0.716745*
Sag1-DP	P (deep)	200–300	0.340	79.4	0.74	1.74	<0.04	0.10	5.88	1.85	0.102	>1980	780	0.713109
AR ( $n = 3$ )	O	0–11	0.116	68.6	0.78	4.56	0.26	0.87	1.20	0.48	0.070	269	980	0.708721
	A	11–27	0.458	67.1	3.29	1.34	0.21	0.19	0.35	2.10	0.076	322	880	0.708773
	B1	27–50	0.549	73.7	1.38	1.38	0.23	0.09	0.06	1.94	0.080	318	925	0.708416
	B2	45–75	0.616	78.2	1.89	1.50	0.21	0.12	0.00	1.61	0.084	367	931	0.708313*
	P1	70–80	0.691	75.7	1.29	1.56	0.27	0.07	0.08	0.87	0.075	282	1010	0.708253
	P2	85–93	0.688	68.6	4.42	1.55	0.19	0.15	0.01	2.76	0.073	356	942	0.708296
GM ( $n = 2$ )	Oe	0–13	0.185	38.5	1.87	5.93	0.31	0.73	0.86	0.77	0.053	125	731	0.710621*
	B1	13–35	0.755	28.5	1.95	1.10	0.39	0.02	0.24	0.61	0.042	73.1	673	0.711741*
	B2	38–44	0.828	39.0	5.19	1.17	0.42	0.08	0.20	3.16	0.057	92.2	690	0.710258
	IIOa	44–55	0.431	105	3.16	1.04	0.26	0.23	1.21	0.76	0.112	405	937	0.710080
	P1	50–54	0.327	82.8	3.23	1.11	0.19	0.23	0.88	2.17	0.099	425	833	0.709814*
	IIB	57–90	0.278	150	2.28	1.93	0.36	0.16	0.74	1.64	0.145	421	1040	0.709412
	P2	100–110	0.289	85.0	0.59	1.90	0.73	0.05	1.61	0.59	0.155	116	548	0.710615

Errors for elemental data and  $^{87}\text{Sr}/^{86}\text{Sr}$  as reported in text, with the following exceptions:\* Error ( $2\sigma$ ) greater than 0.000030 but less than 0.000075.\*\*  $n = 1$  for all  $^{87}\text{Sr}/^{86}\text{Sr}$  data.\*\*\*  $n = 2$  for It1 O horizon.

data for select elements that were log-transformed to induce normal distribution. Although data sets for individual surfaces are not large enough to test reliably, statistical analyses of data from all surfaces combined show some significant ( $p < 0.05$ ) differences between permafrost and active-layer mineral soil chemistry: exchangeable fraction Ca, K, and P concentrations are significantly greater in the permafrost (all “P” horizons) than in active-layer mineral soil (all “A” and “B” horizons). Despite these overall statistically significant differences, the magnitudes of within-surface differences between permafrost and active-layer mineral soil exchangeable fraction elemental concentrations vary substantially among surfaces (Table 2).

The cold acid digestible fractions of the It2, AR, and GM soil profiles have greater Ca concentrations and Ca/Sr and lower  $^{87}\text{Sr}/^{86}\text{Sr}$  than those of the other surfaces (Table 3). The cold acid digestible fraction of the mineral soil shows a general pattern across all surfaces of increasing Ca concentrations and Ca/Sr and decreasing  $^{87}\text{Sr}/^{86}\text{Sr}$  with depth, and on some surfaces P concentrations also increase with depth. Both Ca and P cold acid digestible fraction concentrations are significantly ( $p \leq 0.05$ ) greater in the permafrost than in the active-layer mineral soil when data from all surfaces are analyzed, and the differences in cold acid digestible fraction K concentrations are nearly significant ( $p = 0.06$ ). These differences between active-layer mineral soil and permafrost vary by

TABLE 3

Mean cold acid digestible soil fraction (per gram dry soil). Surface and horizon names are defined in the text.

Surface	Horizon	Depth (cm)	Ba ( $\mu\text{mol/g}$ )	Ca ( $\mu\text{mol/g}$ )	Fe ( $\mu\text{mol/g}$ )	K ( $\mu\text{mol/g}$ )	Na ( $\mu\text{mol/g}$ )	P ( $\mu\text{mol/g}$ )	S ( $\mu\text{mol/g}$ )	Si ( $\mu\text{mol/g}$ )	Sr ( $\mu\text{mol/g}$ )	Ca/Na (molar)	Ca/Sr (molar)	$^{87}\text{Sr}/^{86}\text{Sr}^*$
It3 ( $n = 4$ )	O	0–7	0.682	183	57.3	3.97	0.45	5.12	0.58	29.4	0.144	405	1270	0.713346
	A	7–19	0.123	7.77	147	1.41	0.29	2.06	0.68	42.1	0.011	26.7	681	0.713351
	B1	19–39	0.191	8.54	173	1.46	0.28	1.82	0.58	44.7	0.013	30.6	660	0.714043
	B2	39–60	0.189	8.76	157	1.80	0.28	3.05	0.75	44.7	0.013	31.4	672	0.713707
	P	62–71	0.399	12.6	285	2.46	0.44	8.31	1.25	51.7	0.014	28.9	875	0.713848
It2 ( $n = 3$ )	O	0–16	2.17	360	53.8	1.01	0.56	4.70	5.90	11.9	0.218	646	1650	0.710522
	A	18–29	1.09	60.7	119	1.35	0.39	11.3	2.47	59.8	0.070	156	864	0.710246
	B1	29–46	1.34	62.4	122	1.79	0.46	11.2	4.05	68.8	0.072	136	869	0.709875
	B2	46–62	2.17	122	117	1.99	0.41	12.2	7.12	72.2	0.118	297	1040	0.709850
	P	66–73	2.39	257	125	1.97	0.42	9.76	4.01	66.9	0.195	610	1320	0.709610
It1 ( $n = 3$ )	O*	0–7	0.350	10.4	68.1	3.64	0.18	3.73	1.19	18.8	0.017	59.3	627	0.720647
	A	8–23	0.152	6.03	125	0.95	0.10	2.04	0.34	28.5	0.009	62.5	702	0.713331
	B1	23–40	0.180	8.67	121	1.04	0.10	3.23	0.39	32.1	0.012	88.3	729	0.712963
	B2	35–49	0.186	7.23	124	1.14	0.13	3.10	0.19	39.2	0.011	57.6	658	0.712858
	P1	49–60	0.136	6.28	95.7	1.11	0.13	2.41	0.31	31.9	0.009	47.8	685	0.712326
	P2	62–69	0.423	79.8	125	1.40	0.09	5.60	0.64	42.2	0.052	929	1540	0.710896
Sag2 ( $n = 3$ )	O	0–8	0.537	28.1	52.1	2.91	0.44	5.93	1.33	13.2	0.039	64.6	714	0.715214
	A	8–25	0.182	6.84	92.0	1.01	0.13	2.36	0.33	31.1	0.009	50.9	758	0.712909
	B1	25–42	0.414	7.99	138	1.52	0.14	2.71	0.86	45.1	0.011	57.5	703	0.713031
	B2	38–55	0.079	4.04	60.5	0.94	0.18	3.05	0.15	26.9	0.006	22.3	672	0.712395
	P1	55–69	0.128	5.99	65.4	1.25	0.18	2.44	0.38	28.1	0.009	33.1	655	0.712871
	P2	71–75	0.493	12.2	95.4	1.57	0.11	2.18	0.70	37.0	0.015	114	807	0.712010
Sag1 ( $n = 2$ )	O	0–13	0.677	22.6	85.8	3.28	0.40	3.03	1.56	23.2	0.030	57.1	749	0.715044
	A	13–22	0.272	4.0	133	1.29	0.23	1.54	0.66	34.4	0.006	17.0	669	0.713519
	B1	22–40	0.147	2.65	104	1.26	0.26	1.36	0.28	35.9	0.004	10.1	649	0.713252
	B2	35–50	0.186	3.29	115	1.34	0.27	1.84	0.34	39.1	0.005	12.0	602	0.713607
	P	50–58	0.235	6.16	166	1.53	0.31	4.38	0.45	41.4	0.008	20.0	738	0.713144
Sag1-DP	P (deep)	200–300	1.75	244	236	1.46	18.8	21.5	9.08	169	0.279	13.0	874	0.711083
AR ( $n = 3$ )	O	0–11	0.391	166	37.1	4.89	0.40	6.33	1.82	16.3	0.154	418	1080	0.708762
	A	11–27	0.727	82.7	171	1.97	0.73	19.7	1.66	65.2	0.164	113	503	0.709060
	B1	27–50	0.726	183	137	2.37	0.63	20.3	0.90	85.9	0.213	288	857	0.708580
	B2	45–75	0.446	280	70.8	2.16	0.49	13.2	0.01	66.4	0.250	573	1120	0.708235
	P1	70–80	0.777	423	142	2.59	0.80	20.8	0.12	92.8	0.353	530	1200	0.708352
	P2	85–93	0.740	320	94.2	2.45	0.58	19.0	0.01	84.3	0.309	549	1040	0.708425
GM ( $n = 2$ )	Oe	0–13	0.394	54.8	29.4	6.32	0.34	5.52	1.21	7.06	0.075	160	729	0.710627
	B1	13–35	0.310	14.1	162	3.18	1.21	4.66	3.19	62.6	0.029	11.6	490	0.711870
	B2	38–44	0.467	18.3	286	1.86	0.39	3.20	1.72	57.0	0.034	47.3	538	0.711035
	IIOa	44–55	1.10	186	241	3.73	1.24	7.52	5.48	44.0	0.219	150	849	0.710836
	P1	50–54	0.811	101	131	1.96	0.34	6.93	2.43	36.7	0.132	295	762	0.709522
	IIB	57–90	1.05	342	139	4.22	1.34	9.92	3.85	64.0	0.313	255	1090	0.709681
	P2	100–110	1.07	1220	177	3.21	1.37	13.9	2.55	72.4	0.850	891	1440	0.709076

Errors for elemental data and  $^{87}\text{Sr}/^{86}\text{Sr}$  as reported in text; maximum error ( $2\sigma$ ) for any  $^{87}\text{Sr}/^{86}\text{Sr}$  value used in calculating means is 0.000068.\* For  $^{87}\text{Sr}/^{86}\text{Sr}$ ,  $n = 3$  for It3,  $n = 1$  for AR and GM; for other surfaces,  $n$  is as reported.

surface: all surfaces have higher permafrost concentrations of Ca; all surfaces except the GM and It2 have higher permafrost concentrations of K; and cold acid digestible fraction P concentrations are higher in the permafrost on four of the surfaces.

The hot acid digestible fractions of mineral soil horizons on all surfaces contain some Ca, but have lower Ca/Sr and higher  $^{87}\text{Sr}/^{86}\text{Sr}$  than either the exchangeable or cold acid digestible fractions (Table 4). This suggests that the majority of carbonate was dissolved before this step in the sequential digest procedure and that the Ca in this fraction is largely the result of silicate mineral dissolution.

The total digests of the mineral soil (Table 5) show that Ca generally increases with depth on each surface. The It2, AR, and GM surfaces have the highest Ca concentrations. On average,  $\text{SiO}_2$  comprises approximately 80% of the total mineral soil mass.

For a few samples, elemental concentrations from the acid digests sum to slightly more than the total digest concentration. These values are within analytical errors propagated through the sum in all except for the Ca values for the Sag1 deep permafrost sample. This discrepancy is likely the result of sample heterogeneity.

### STREAM GEOCHEMISTRY

In streams on glacial deposit surfaces, dissolved Ca concentrations ranged from  $12 \mu\text{mol L}^{-1}$  on the Sag1 surface to  $594 \mu\text{mol L}^{-1}$  on the It2 surface. Streams draining only limestone bedrock had Ca concentrations up to  $1606 \mu\text{mol L}^{-1}$  (Table 6). Dissolved P concentrations were generally less than  $1 \mu\text{mol L}^{-1}$ , but several streams on the It2 surface had concentrations greater than  $10 \mu\text{mol L}^{-1}$ . Dissolved Si concentrations were less than

TABLE 4

Mean hot acid digestible soil fraction (per gram dry soil). Surface and horizon names are defined in the text.

Surface	Horizon	Depth (cm)	Ba ( $\mu\text{mol/g}$ )	Ca ( $\mu\text{mol/g}$ )	Fe ( $\mu\text{mol/g}$ )	K ( $\mu\text{mol/g}$ )	Na ( $\mu\text{mol/g}$ )	P ( $\mu\text{mol/g}$ )	S ( $\mu\text{mol/g}$ )	Si ( $\mu\text{mol/g}$ )	Sr ( $\mu\text{mol/g}$ )	Ca/Na (molar)	Ca/Sr (molar)	$^{87}\text{Sr}/^{86}\text{Sr}^{***}$
It3 ( $n = 4$ )	O	0–7	0.309	51.2	196	5.42	16.7	13.0	26.2	10.5	0.045	3.07	1150	0.714412
	A	7–19	0.125	2.44	359	8.40	12.9	5.56	3.09	49.3	0.015	0.19	168	0.731134
	B1	19–39	0.145	2.37	409	10.1	13.0	6.74	3.77	44.2	0.014	0.18	173	0.732188
	B2	39–60	0.183	2.17	441	14.3	12.9	8.67	5.75	68.2	0.019	0.17	113	0.733771*
	P	62–71	0.253	4.38	263	9.05	17.0	7.22	10.6	4.12	0.010	0.26	436	0.730624
It2 ( $n = 3$ )	O	0–16	0.949	87.9	95.1	.78	16.7	15.4	101	9.41	0.056	5.27	1570	0.709652*
	A	18–29	0.284	3.91	298	5.22	11.5	6.81	13.6	87.8	0.009	0.34	429	0.720106
	B1	29–46	0.342	4.10	342	7.61	11.5	4.55	14.6	102	0.012	0.36	340	0.720671*
	B2	46–62	0.353	7.20	341	10.5	11.2	7.20	24.5	47.9	0.015	0.64	485	0.725481
	P	66–73	0.324	13.2	395	4.94	8.34	8.20	27.1	50.9	0.019	1.58	710	0.715267
It1 ( $n = 3$ )	O****	0–7	0.296	6.06	317	9.33	17.2	23.2	25.9	15.4	0.012	0.35	526	0.724344
	A	8–23	0.107	1.76	330	5.85	12.1	5.35	2.54	60.4	0.008	0.15	231	0.727602*
	B1	23–40	0.103	1.69	285	6.82	12.5	4.53	2.45	81.9	0.009	0.14	191	0.730939
	B2	35–49	0.191	2.46	399	7.38	18.4	8.26	2.41	3.19	0.010	0.13	254	0.731677
	P1	49–60	0.192	2.70	394	8.64	17.9	7.61	1.98	4.57	0.010	0.15	275	0.732349
	P2	62–69	0.160	2.77	411	15.0	12.0	6.70	8.16	114	0.017	0.23	163	0.733956
Sag2 ( $n = 3$ )	O	0–8	0.238	11.8	156	5.12	17.5	17.5	23.9	8.03	0.016	0.67	752	0.720922*
	A	8–25	0.126	2.14	283	5.86	12.0	5.98	3.10	45.3	0.007	0.18	295	0.722058
	B1	25–42	0.231	2.38	343	9.99	11.9	7.13	6.14	54.2	0.012	0.20	190	0.722017
	B2	38–55	0.187	3.45	216	4.52	18.3	2.95	1.38	4.15	0.009	0.19	394	0.723151
	P1	55–69	0.191	3.08	386	4.25	17.5	7.44	3.01	4.87	0.007	0.18	465	0.721817*
	P2	71–75	0.209	2.10	386	8.62	12.0	5.38	5.55	67.8	0.010	0.17	206	0.723476
Sag1 ( $n = 2$ )	O	0–13	0.363	7.92	176	8.86	17.1	21.5	27.5	2.43	0.019	0.46	409	0.722149**
	A	13–22	0.307	4.20	459	10.1	18.2	12.1	6.59	4.87	0.018	0.23	227	0.723068
	B1	22–40	0.247	4.30	364	7.33	17.8	7.31	1.91	4.88	0.016	0.24	274	0.722645**
	B2	35–50	0.248	3.88	371	7.76	18.4	7.16	1.56	0.94	0.016	0.21	250	0.723827
	P	50–58	0.274	5.15	302	7.92	16.6	5.88	2.30	5.19	0.014	0.31	369	0.725281
Sag1-DP	P (deep)	200–300	0.257	12.7	435	9.86	18.1	4.10	6.02	3.51	0.041	0.70	309	0.726095
AR ( $n = 3$ )	O	0–11	0.348	70.0	136	4.30	16.0	20.9	58.4	19.8	0.069	4.36	1020	0.708900*
	A	11–27	0.394	14.4	385	3.26	8.52	19.7	20.4	9.08	0.036	1.68	397	0.713011**
	B1	27–50	0.450	14.7	535	7.76	17.6	11.3	8.80	5.44	0.027	0.84	538	0.711376
	B2	45–75	0.359	18.9	502	6.69	16.6	9.57	3.99	2.67	0.027	1.14	708	0.713535
	P1	70–80	0.402	20.1	508	8.50	17.8	8.94	4.06	3.25	0.028	1.13	723	0.711658
	P2	85–93	0.386	9.08	491	7.97	17.2	9.73	4.64	1.91	0.022	0.53	407	0.716740*
GM ( $n = 2$ )	Oe	0–13	0.457	48.1	68.9	5.85	15.9	21.7	38.7	16.2	0.062	3.02	781	0.712801**
	B1	13–35	0.297	5.67	234	6.72	17.5	7.95	9.44	5.31	0.019	0.32	300	0.713987*
	B2	38–44	0.360	5.84	460	8.00	17.1	9.02	8.52	5.91	0.021	0.34	283	0.714585*
	IIOa	44–55	0.438	60.6	204	4.96	16.4	15.9	48.5	16.1	0.066	3.68	919	0.713590*
	P1	50–54	0.428	25.2	189	8.38	17.7	11.5	36.1	19.2	0.042	1.43	604	0.713149*
	IIB	57–90	0.463	58.4	209	14.4	18.1	8.30	46.1	12.6	0.066	3.22	890	0.712507**
	P2	100–110	0.340	31.7	211	14.0	18.8	5.23	23.2	4.57	0.046	1.69	684	0.714035**

Errors for elemental data and  $^{87}\text{Sr}/^{86}\text{Sr}$  as reported in text, with the following exceptions:\* Error ( $2\sigma$ ) greater than 0.000030 but less than 0.000075.\*\* Error ( $2\sigma$ ) greater than 0.000075 but less than 0.000211.\*\*\*  $n = 1$  for all  $^{87}\text{Sr}/^{86}\text{Sr}$  data.\*\*\*\*  $n = 2$  for It1 O horizon.

1  $\mu\text{mol L}^{-1}$  in two streams that were lake outlets, but other Si concentrations ranged from 10 to 87  $\mu\text{mol L}^{-1}$ .  $^{87}\text{Sr}/^{86}\text{Sr}$  ranged from 0.71023 to 0.71881 in streams draining glacial deposit surfaces and from 0.70847 to 0.71989 in streams draining bedrock surfaces.

## Discussion

### ANALYSIS OF SOIL PARENT MATERIAL VARIABILITY

Before we can evaluate whether geochemical patterns are dependent on varying degrees of weathering on geomorphic

surfaces of different ages, we must ascertain the degree of variability of the composition of the glacial till soil parent among the surfaces. Figure 2 shows a comparison of the  $^{87}\text{Sr}/^{86}\text{Sr}$  of the hot acid digestible fractions of the soil profiles on each surface. Because easily weathered material has already been removed from this fraction by previous steps in the sequential digest procedure, we make the assumption that the chemistry of this fraction reflects minerals that would only be affected by intense weathering, making them unlikely to have been significantly affected yet by weathering in this arctic region. Therefore, the  $^{87}\text{Sr}/^{86}\text{Sr}$  and elemental ratios of this fraction should be very similar for soils with similar parent material, regardless of their age. We used



TABLE 5

Total soil digest chemistry (per gram dry soil). Surface and horizon names are defined in the text.

Surface	Horizon	Depth (cm)	Ba ( $\mu\text{mol/g}$ )	Ca ( $\mu\text{mol/g}$ )	Fe ( $\mu\text{mol/g}$ )	K ( $\mu\text{mol/g}$ )	Na ( $\mu\text{mol/g}$ )	P ( $\mu\text{mol/g}$ )	Si ( $\mu\text{mol/g}$ )	Sr ( $\mu\text{mol/g}$ )	Ca/Na (molar)	Ca/Sr (molar)	Fraction ( $<2$ mm)	LOI (%)
It3 ( $n = 4$ )	O	0–7	—	—	—	—	—	—	—	—	—	—	—	—
	A	7–19	3.01	20.5	547	282	128	12.73	13600	0.583	0.017	35.2	76%	5%
	B1	19–39	3.50	19.9	685	325	129	14.68	14100	0.624	0.018	31.9	76%	7%
	B2	39–60	3.51	19.7	644	327	122	16.77	12800	0.585	0.019	33.7	80%	8%
	P	62–71	4.66	35.1	730	383	128	23.84	13200	0.643	0.023	54.6	63%	—
It2 ( $n = 3$ )	O	0–16	—	—	—	—	—	—	—	—	—	—	—	—
	A	18–29	4.77	124	504	213	151	22.23	13900	0.577	0.023	214	65%	12%
	B1	29–46	5.32	113	565	271	155	19.90	14000	0.591	0.023	192	52%	20%
	B2	46–62	5.73	202	606	265	141	23.04	12400	0.626	0.024	323	59%	8%
	P	66–73	6.67	330	831	335	168	27.77	11800	0.759	0.021	434	82%	7%
It1 ( $n = 3$ )	O*	0–7	—	—	—	—	—	—	—	—	—	—	—	—
	A	8–23	2.65	18.2	541	251	148	12.80	15300	0.508	0.019	35.8	49%	5%
	B1	23–40	2.85	18.7	484	262	118	15.31	15400	0.526	0.019	35.5	57%	8%
	B2	35–49	3.30	20.4	638	286	108	17.46	15100	0.562	0.020	36.3	54%	—
	P1	49–60	3.24	19.9	586	304	112	15.13	17200	0.555	0.020	35.9	53%	—
	P2	62–69	3.45	108	653	305	117	16.98	15100	0.635	0.020	170	71%	10%
Sag2 ( $n = 3$ )	O	0–8	—	—	—	—	—	—	—	—	—	—	—	—
	A	8–25	2.66	19.4	466	197	101	14.25	15600	0.466	0.021	41.6	66%	7%
	B1	25–42	4.37	30.1	705	315	124	20.55	13500	0.517	0.024	58.3	79%	23%
	B2	38–55	2.61	23.8	343	195	115	8.60	16100	0.505	0.019	47.1	69%	—
	P1	55–69	2.55	22.5	514	179	93.5	14.53	15200	0.454	0.021	49.4	61%	—
	P2	71–75	3.82	37.5	743	265	112	17.95	13400	0.432	0.026	86.8	77%	22%
Sag1 ( $n = 2$ )	O	0–13	—	—	—	—	—	—	—	—	—	—	—	—
	A	13–22	3.67	29.3	565	236	137	17.1	14000	0.560	0.023	52.3	69%	15%
	B1	22–40	3.55	26.6	519	229	157	12.36	14000	0.548	0.023	48.5	65%	7%
	B2	35–50	4.08	26.8	586	264	167	13.29	14400	0.576	0.024	46.5	58%	—
	P	50–58	3.77	25.9	525	243	140	13.79	14200	0.570	0.024	45.4	61%	7%
Sag1-DP	P (deep)	200–300	7.13	255	896	675	153	29.46	13400	1.14	0.019	224	88%	—
AR ( $n = 3$ )	O	0–11	—	—	—	—	—	—	—	—	—	—	—	—
	A	11–27	4.13	138	581	200	181	30.1	11900	0.589	0.021	234	46%	—
	B1	27–50	5.34	270	631	267	239	31.6	13500	0.797	0.019	339	40%	—
	B2	45–75	4.58	428	529	255	199	26.89	11900	0.858	0.017	498	30%	—
	P1	70–80	5.45	483	593	268	235	32.41	13000	0.915	0.020	528	41%	—
	P2	85–93	5.25	612	596	277	227	48.97	13700	1.27	0.019	480	37%	—
GM ( $n = 2$ )	Oe	0–13	—	—	—	—	—	—	—	—	—	—	—	—
	B1	13–35	4.47	67.6	475	259	222	20.99	13200	0.685	0.023	98.7	47%	—
	B2	38–44	4.37	75.7	845	209	183	20.33	12600	0.682	0.024	111	76%	—
	IIOa	44–55	3.16	289	379	128	95.5	28.8	5860	0.614	0.032	471	69%	—
	P1	50–54	3.65	219	336	164	126	24.9	8240	0.659	0.028	333	57%	—
	IIB	57–90	4.60	593	396	249	203	25.30	10600	0.982	0.020	604	79%	—
	P2	100–110	4.82	1320	452	283	208	27.27	11400	1.63	0.011	814	57%	—

Errors for elemental data as reported in text. LOI = loss on ignition.

$^{87}\text{Sr}/^{86}\text{Sr}$  rather than elemental ratios as an indicator of differences in parent material because different parent materials of the same type (e.g. different sandstone units) may have very similar elemental ratios due to similar mineralogical compositions, but because of their unique protoliths, they are more likely to have detectable differences in  $^{87}\text{Sr}/^{86}\text{Sr}$ . We examined the  $^{87}\text{Sr}/^{86}\text{Sr}$  of the hot acid digestible fraction with respect to depth, to account for any minor effects of weathering or addition of new material (e.g. eolian loess) to the soil profile.

Overall the composition of the parent material has low variability but showed some grouping among the different surfaces. The GM and AR profiles have significantly lower mean  $^{87}\text{Sr}/^{86}\text{Sr}$  than the other surfaces (Fig. 2;  $p < 0.05$ ). This suggests that the AR and GM soils are derived from different parent material than the other surfaces. Differences in parent material

may be explained by the incorporation of various rock units in different proportions based on different surface expressions of these units at the time of the glacial advances. Additionally, rock units (the Kn and three minor units not analyzed here) that outcrop north of the Sag1 surface may have been incorporated into the GM and AR parent till. It is also possible that long-term eolian loess deposition has contributed to these differences in soil composition. The  $^{87}\text{Sr}/^{86}\text{Sr}$  of the hot acid digestible fraction of the It1, It2, It3, Sag1, and Sag2 soil profiles is similar, suggesting that these surfaces have similar parent material, although the  $^{87}\text{Sr}/^{86}\text{Sr}$  of the upper part of the It3 profile is slightly higher than that of the other surfaces. This is consistent with the suggestion by Hamilton (2003) that the It3 surface was derived from slightly different parent material than the It1 and It2 surfaces, based on surficial geomorphology. For the purposes of further analyses

TABLE 6

Water chemistry for streams on different glacial deposit and bedrock surfaces. Surface names are defined in the text.

Surface	Stream name	Date sampled	Ba (μmol/L)	Ca (μmol/L)	Fe (μmol/L)	K (μmol/L)	Mg (μmol/L)	Na (μmol/L)	S (μmol/L)	Si (μmol/L)	Sr (μmol/L)	Ca/Na (molar)	Ca/Sr (molar)	<sup>87</sup> Sr/ <sup>86</sup> Sr
It3	(1) It 3 stream	7/29/02	0.08	551.3	3.88	2.3	104.9	36	20.8	67.6	0.740	15	745	0.710391*
	(2) Island Lake outlet	8/2/02	0.13	522.0	<0.08	10.3	183.5	77	20.7	85.9	0.736	6.8	709	0.710234
It2 (outwash)	(3) I minus outlet	6/27/02	0.10	219.6	1.03	8.3	61.5	16	20.8	78.2	0.760	14	289	0.713132
	(4) I minus inlet	6/27/02	0.07	136.3	2.26	4.8	37.9	13	20.8	78.8	0.766	11	178	0.712971
	(5) TW stream (lower)	7/21/02	0.10	269.2	0.96	1.0	57.6	7	20.8	77.9	0.761	40	354	0.713314
	(6) Milky Way (upper)	8/10/02	0.12	255.4	1.56	2.3	62.7	27	18.4	40.3	0.205	9.4	1244	0.712085*
It2 (till)	(7) N1 inlet	7/11/02	0.09	543.5	0.64	1.2	117.5	13	20.8	80.3	0.752	42	722	0.710600*
	(8) I1 outlet	8/7/02	0.03	123.1	1.34	6.4	36.5	14	8.1	0.9	0.102	8.8	1204	0.715188**
	(9) I4 outlet	8/7/02	0.05	104.9	2.65	5.0	34.7	13	8.5	<0.4	0.096	7.8	1093	0.715593
	(10) I2 outlet	8/7/02	0.07	129.7	4.61	6.7	50.5	14	6.8	10.4	0.131	9.4	988	0.716473*
It1	(11) I6 HW inlet	8/8/02	0.08	594.3	9.20	1.0	97.9	20	6.4	34.6	0.404	30	1471	0.711281
	(12) I8 Headwater	6/26/02	0.06	168.1	2.32	2.8	52.7	13	20.8	78.8	0.764	13	220	0.713464*
	(13) It 1 stream	7/15/02	0.25	440.6	<0.08	2.5	73.4	19	20.8	76.2	0.746	23	590	0.710995
Sag2	(14) E5 inlet S	7/30/02	0.07	44.2	6.37	0.7	20.2	11	3.1	37.2	0.046	4.0	958	0.716328
	(15) E5 inlet W	8/13/02	0.09	19.7	2.02	0.3	12.2	13	8.5	43.6	0.025	1.6	796	0.718806
Sag1	(16) Imnavait (weir)	8/1/2002	0.08	28.1	13.55	0.6	14.2	4	1.2	20.2	0.036	7	777	0.716905
	(17) Toolik R. trib.	8/16/02	0.06	11.7	8.36	0.5	10.9	4	2.6	29.0	0.018	3	654	0.717341
AR	(18) Anaktuvuk stream	8/15/03	0.08	78.2	4.55	0.6	18.3	12	2.3	47.1	0.079	6.7	990	0.712417
GM	(19) Gunsight stream	8/16/03	0.07	165.8	5.62	<0.3	44.2	9	3.5	58.7	0.148	20	1120	0.711356
Multiple surfaces	(20) Kuparuk R.	8/7/02	0.28	199.1	0.37	6.6	71.2	78	20.8	65.3	0.745	2.6	267	0.707878**
	(21) Sagavanirktok R.	7/31/02	0.22	924.3	<0.08	8.9	282.5	80	20.7	76.8	0.635	12	1456	0.710453
Carbonate bedrock	(22) Atigun R. trib.	7/29/02	0.26	1559.1	<0.08	13.3	287.4	97	20.6	80.2	0.678	16	2301	0.710550*
	(23) Kayak Creek	8/7/02	0.13	1605.7	<0.08	7.5	166.3	20	20.8	80.9	0.569	79	2821	0.708469
	(24) Aufeis stream	7/31/02	0.35	1226.5	<0.08	5.4	272.0	21	20.8	71.0	0.707	58	1736	0.708802
Mixed bedrock	(25) Trevor Creek	8/7/02	0.30	706.7	<0.08	10.5	264.2	94	20.7	83.6	0.704	7.5	1003	0.714855*
	(26) Atigun R.	7/29/02	0.18	691.1	<0.08	12.2	324.6	122	20.7	80.8	0.697	5.67	991	0.714458
	(27) Bad Idea stream	8/18/03	0.18	866.5	<0.08	14.8	167.5	169	397.0	34.7	1.495	5.14	580	0.711382
	(28) Holden Creek	8/18/03	0.09	569.4	0.14	8.6	102.4	88	122.5	21.2	1.212	6.4	470	0.709772
	(29) Roche Moutonee Cr.	8/18/03	0.12	372.6	0.16	9.9	110.1	56	83.0	24.0	0.465	6.7	802	0.713843
Silicate bedrock	(30) Atigun R. headwater	8/7/02	0.07	570.4	<0.08	12.2	395.8	271	20.7	77.5	0.686	2.11	832	0.719655*
	(31) Underwillow stream	8/18/03	0.27	254.5	0.30	22.7	194.9	296	318.6	31.6	0.502	.86	507	0.719893
Precipitation	field station precip. 1	8/5/02	<0.01	102.2	<0.08	83.5	4.4	218	20.8	92.3	0.770	0.47	133	0.709485**
	field station precip. 2	8/6/02	0.03	84.9	<0.08	19.0	2.1	57	20.8	92.0	0.772	1.5	110	0.708781*

Errors for elemental data and <sup>87</sup>Sr/<sup>86</sup>Sr as reported in text, with the following exceptions:

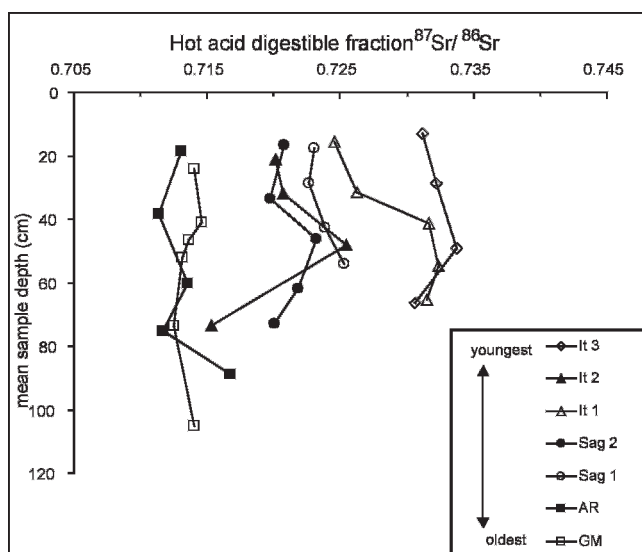
\* Error (2σ) greater than 0.000030 but less than 0.000051.

\*\* error (2σ) greater than 0.000051 but less than 0.000165.

with respect to surface age, we will consider only the It1, It2, Sag1, and Sag2 to be derived from similar parent material.

Besides these overall differences among the profiles from each surface, each individual profile shows a pattern of increasing hot acid digestible fraction <sup>87</sup>Sr/<sup>86</sup>Sr with increasing depth within the active layer (down to approximately 50 cm; Fig. 2). The most likely explanation for this is eolian deposition of a weathering-resistant mineral, with lower <sup>87</sup>Sr/<sup>86</sup>Sr than the soil parent material, which is mixed down through the active layer by cryoturbation or water movement. Such dust could be derived from the Dke sandstone or resistant minerals in the PMI (Table 1), or from sediments outside the study region. This dust may also

have a carbonate component that contributes to the exchangeable and cold acid digestible fractions. However, the effects of this input as shown by increasing <sup>87</sup>Sr/<sup>86</sup>Sr with depth seem to be of similar magnitude in the hot acid digestible fraction, so we can assume that age comparisons between these surfaces based on the other fractions are not substantially affected. An alternate explanation for the pattern of increasing <sup>87</sup>Sr/<sup>86</sup>Sr with depth in the active layer could be removal by weathering of a resistant mineral with relatively high <sup>87</sup>Sr/<sup>86</sup>Sr (such as biotite) in the upper part of the active layer, but this seems unlikely due to the low weathering intensity of this environment as evidenced by the relatively slow carbonate weathering described in the next section.



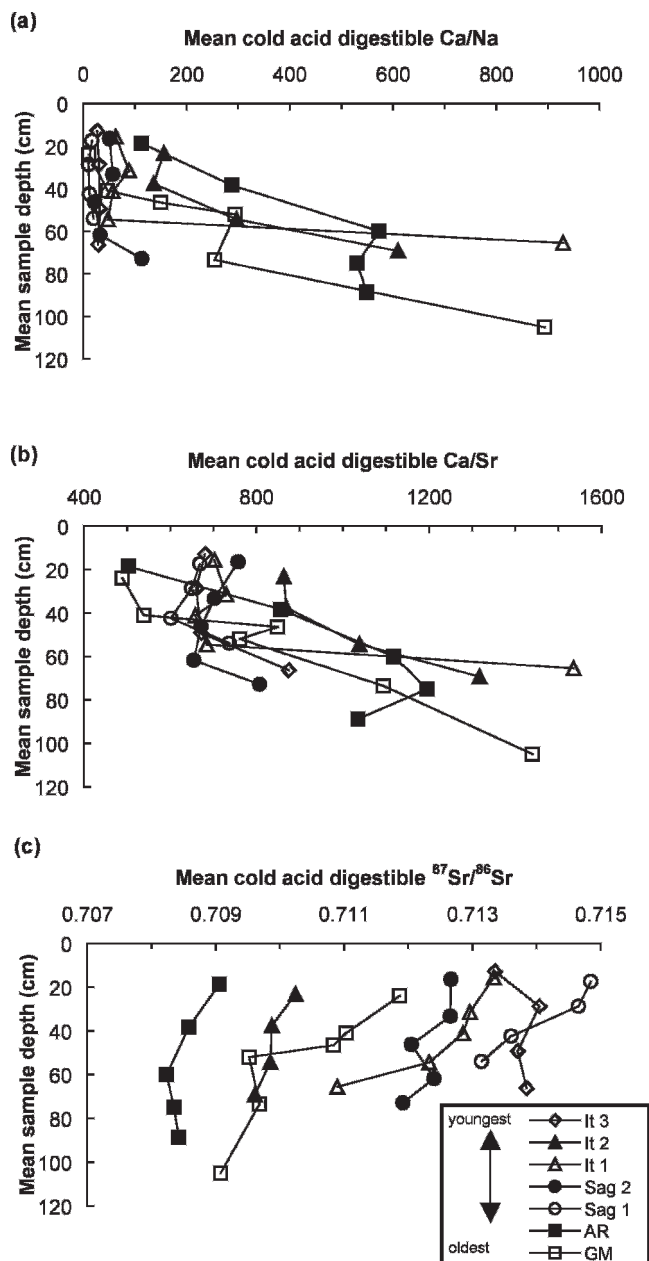
**FIGURE 2.** Hot acid digestible fraction  $^{87}\text{Sr}/^{86}\text{Sr}$  of soil horizon samples vs. mean horizon depth.  $^{87}\text{Sr}/^{86}\text{Sr}$  analytical errors are reported in Table 4.

#### CARBONATE WEATHERING AND ITS RELATIONSHIP TO SURFACE AGE

Calcite depletion via weathering in the active layer is indicated on every surface, by decreasing  $^{87}\text{Sr}/^{86}\text{Sr}$  and increasing Ca/Na and Ca/Sr with depth (Fig. 3), and by greater Ca concentrations in permafrost than in active layer mineral soil as described in the results section. These geochemical trends suggest increasing carbonate mineral concentration in the soil with depth, as frequency of thaw and hence weathering intensity decreases. These results may explain the findings of Everett et al. (1989), who monitored the seasonal geochemistry of the Imnavait stream on the Sag1 surface and found that streamwater Ca concentrations were greatest in the late summer, corresponding to deep seasonal thaw.

When considering only the surfaces with similar parent material (It1, It2, Sag1, and Sag2), some patterns related to surface age become apparent. The mean  $^{87}\text{Sr}/^{86}\text{Sr}$  values of the active-layer cold acid digestible fraction are lowest on the youngest surface (It2) and highest on the oldest (Sag1), with overlapping values for the two other surfaces (Fig. 3c). The decrease in mean cold acid digestible fraction  $^{87}\text{Sr}/^{86}\text{Sr}$  between the permafrost and the "A" horizon is least on the youngest surface. The youngest surface also has the highest exchangeable, acid digestible, and total Ca concentrations and Ca/Sr (Tables 2, 3, and 5). These data suggest that weathering has progressively depleted the calcite in the active layer over time on these surfaces. Furthermore, mean cold acid digestible fraction  $^{87}\text{Sr}/^{86}\text{Sr}$  values for shallow permafrost ("P" samples from within soil pits, excluding the Sag1 deep permafrost sample) increase with surface age among the four surfaces derived from similar parent material (Fig. 3c, Table 3). This trend suggests that the currently frozen soil has been episodically thawed during past intervals of warmer climate and has undergone some weathering or was mixed with weathered material via cryoturbation during these intervals.

Both stream and soil geochemistry suggest that differences in carbonate weathering drive not only depth-related geochemical differences within individual soil profiles, but also age-related differences in whole watershed geochemistry. Simple chemical mixing lines drawn between end-members formed by the cold acid digestible fractions of the PMI limestone (high Ca/Sr, low



**FIGURE 3.** Cold acid digestible (a) Ca/Na, (b) Ca/Sr, and (c)  $^{87}\text{Sr}/^{86}\text{Sr}$  of soil horizon samples vs. mean sample depth. Elemental ratios are molar. Analytical errors are reported in the text.

$^{87}\text{Sr}/^{86}\text{Sr}$ ), Dke sandstone (low Ca/Sr, high  $^{87}\text{Sr}/^{86}\text{Sr}$ ), and Kf sandstone (low Ca/Sr, low  $^{87}\text{Sr}/^{86}\text{Sr}$ ) represent the expected composition of waters containing easily weathered solutes derived from these units based on varying proportional contributions from each unit by mass (Fig. 4). These mixing lines define a range containing 7 out of 15 streams draining watersheds developed on different glacial till surfaces. Streams draining PMI-based watersheds have a lower mean Ca/Sr and very slightly higher mean  $^{87}\text{Sr}/^{86}\text{Sr}$  than the cold acid digestible fraction of PMI samples (Fig. 4, hatched line end-members). This may be explained by a greater influence of precipitation on these watersheds, or a fraction of the PMI which is more soluble under natural conditions and differs chemically from the fraction digestible with stronger acid in the laboratory, or both. The slightly higher Ca/Sr and  $^{87}\text{Sr}/^{86}\text{Sr}$  of streams draining Dke sandstone and undifferentiated Kanayut conglomerate (MDk) bedrock may be explained

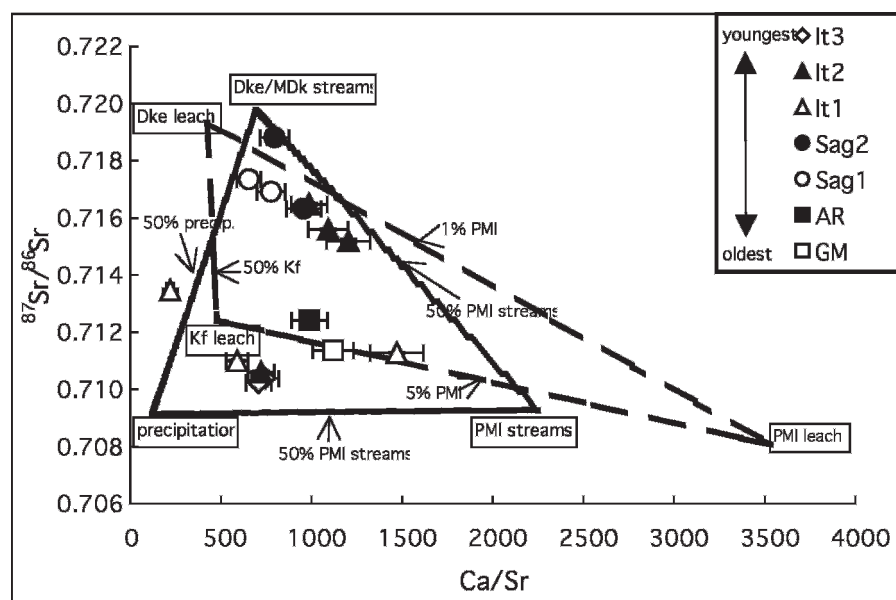


FIGURE 4.  $^{87}\text{Sr}/^{86}\text{Sr}$  vs.  $\text{Ca}/\text{Sr}$  of stream water samples from glacial deposit surfaces. Dashed lines connecting the cold acid digestible fraction values of PMI, Dke, and Kf rock units (see Table 1) represent simple chemical mixing lines for these end-members. Hatched lines connecting the mean values for precipitation and streams draining PMI and Dke/MDk units represent simple chemical mixing lines for these end-members.  $\text{Ca}/\text{Sr}$  is a molar ratio. Error bars represent propagated  $\text{Ca}/\text{Sr}$  analytical errors;  $^{87}\text{Sr}/^{86}\text{Sr}$  errors are smaller than the symbol size and are reported in Table 6. Rock unit abbreviations are explained in the text.

by contributions from other members of the MDk, as well as the slightly higher  $^{87}\text{Sr}/^{86}\text{Sr}$  of the exchangeable fraction of the Dke.

Simple chemical mixing lines drawn between the mean  $\text{Ca}/\text{Sr}$  and  $^{87}\text{Sr}/^{86}\text{Sr}$  of precipitation samples, streams draining PMI limestone bedrock, and streams draining Dke/MDk bedrock (Fig. 4, hatched lines) represent the expected composition of waters containing solutes derived from a combination of inputs based on the solutes measured in unit-specific streams and precipitation. All but one of the 15 glacial till surface streams sampled are contained within these mixing lines (Fig. 4). The reasons that these end-members predict water composition more accurately than the range defined by the cold acid digestible fraction of rock units are likely that (1) the effects of precipitation and other members of the MDk are incorporated, and (2) they are based on weathering in natural settings, rather than on an operationally defined laboratory cold acid digest.

Within the ranges defined by both sets of mixing lines, most streams on younger (It3, It2, and It1) surfaces have  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios similar to those of streams draining limestone bedrock and the cold acid digestible fraction of limestone, but lake outlet streams on the It2 surface have higher  $^{87}\text{Sr}/^{86}\text{Sr}$ . The  $\text{Ca}/\text{Sr}$  ratios of these streams vary widely, perhaps reflecting varying effects of precipitation in different watersheds or at the different times when the streams were sampled. The effects of precipitation on the  $^{87}\text{Sr}/^{86}\text{Sr}$  of these streams would be nearly undetectable because the difference between the  $^{87}\text{Sr}/^{86}\text{Sr}$  of precipitation and streams draining PMI watersheds is less than 0.0002. Streams on older (Sag2 and Sag1) surfaces have  $\text{Ca}/\text{Sr}$  and  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios similar to those of streams draining Dke/MDk bedrock and the cold acid digestible fraction of Dke sandstone, suggesting much less carbonate influence on these surfaces than on younger surfaces. Streams on very old (AR and GM) surfaces have  $\text{Ca}/\text{Sr}$  and  $^{87}\text{Sr}/^{86}\text{Sr}$  which fall very close to the Kf-PMI mixing line, suggesting that the weathering products dissolved in these streams are primarily derived from these two end-members, or perhaps an end-member with similar chemistry such as the Kn (Table 1). The chemistry of the GM and AR streams suggests little input of the Dke/MDk end-member, consistent with the differences in parent material between these and younger surfaces discussed above.

The effects of weathering on shallow mineral soils of different ages was examined by plotting cold acid digestible  $\text{Ca}/\text{Sr}$  vs.

$^{87}\text{Sr}/^{86}\text{Sr}$  for soil “A” horizon samples from all glacial deposit surfaces along with the mixing lines from the cold acid digestible fraction rock unit end-members described above (Fig. 5).  $\text{Ca}/\text{Sr}$  and  $^{87}\text{Sr}/^{86}\text{Sr}$  for the GM, AR, and It2 surfaces fall outside and below the range defined by the mixing lines, suggesting that their chemistry has been influenced by input from precipitation, or low- $^{87}\text{Sr}/^{86}\text{Sr}$  loess deposition, or both. Data from the It3, It1, and Sag2 surfaces overlap, consistent with the overlapping  $^{87}\text{Sr}/^{86}\text{Sr}$  values for It1 and Sag2 soils in Figure 3, but despite the clear differences in the stream water  $\text{Ca}/\text{Sr}$  vs.  $^{87}\text{Sr}/^{86}\text{Sr}$  for these surfaces (Fig. 4). This may be because, as discussed above, the cold acid digestible fraction represents an estimate of potentially “easily weatherable” minerals, but does not necessarily reflect what is actually being weathered. Therefore, the It3, It1, and Sag2 surfaces could have similar reservoirs of “easily weatherable” minerals as estimated by the digest, yet release into streams very different weathering products that represent the most soluble part of the “easily weatherable” fraction. Cryoturbation may also play a role in keeping the cold acid digestible fraction similar in soils close in age (like the It1 and Sag2) via the upward mixing of less-weathered minerals from deeper, less frequently thawed layers.

Despite the overlapping values of data from the It3, It1, and Sag2 surfaces, samples from the youngest (It2) of the four surfaces with similar parent material generally have the highest  $\text{Ca}/\text{Sr}$  and lowest  $^{87}\text{Sr}/^{86}\text{Sr}$ , while samples from the oldest (Sag1) of these four surfaces have the lowest  $\text{Ca}/\text{Sr}$  and highest  $^{87}\text{Sr}/^{86}\text{Sr}$ , plotting near the Dke digest end-member. These data suggest that carbonate weathering causes detectable geochemical differences to develop in the cold acid digestible fraction of shallow mineral soil over hundreds of thousands of years. Additional research will be necessary to explore the relationship between the time scale at which the effects of weathering are apparent and the rate of cryoturbation.

These results show that carbonate content decreases in soils and carbonate weathering products decrease in streams with increasing surface age, but the carbonate weathering regime is still generally dominant even on older surfaces. This interpretation is consistent with previous studies of aquatic and soil chemistry in this region (Kling et al., 1992) and similar arctic regions (Stutter and Billet, 2003). The decrease in carbonate weathering over time (driven by a depletion of carbonate minerals via weathering) that

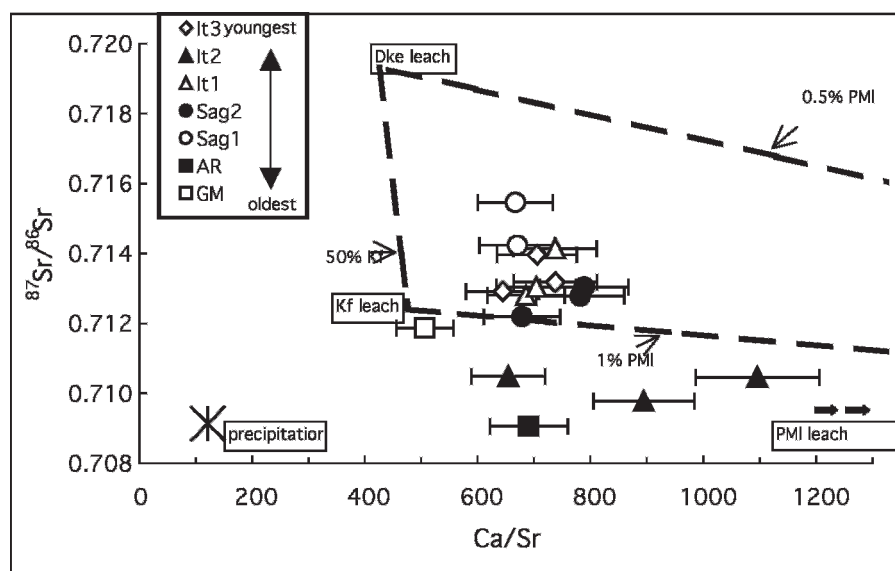


FIGURE 5. Cold acid digestible fraction  $^{87}\text{Sr}/^{86}\text{Sr}$  vs.  $\text{Ca}/\text{Sr}$  of soil “A” horizon samples from glacial deposit surfaces. Dashed lines connecting the cold acid digestible fraction values of PMI, Dke, and Kf rock units (see Table 1) represent simple chemical mixing lines for these end-members, as in Figure 4.  $\text{Ca}/\text{Sr}$  is a molar ratio. Error bars represent propagated  $\text{Ca}/\text{Sr}$  analytical errors;  $^{87}\text{Sr}/^{86}\text{Sr}$  errors are smaller than the symbol size and are reported in Table 3. Rock unit abbreviations are explained in the text.

we interpret from these data is similar to changes that take place in a much shorter time (thousands rather than hundreds of thousands of years) in tropical and temperate climates (e.g. Lichter, 1998). Despite the extended timeframe, this change in the carbonate content of soils with time supports the suggestion of Munroe and Bockheim (2001) that the traditional soil chronosequence model relating age to soil properties is applicable even in cryoturbated soils.

#### SILICATE WEATHERING

The presence of dissolved Si in low but detectable levels in streams on almost every geomorphic surface in our study area suggests that some silicate weathering is occurring; the range of dissolved streamwater Si concentrations in this study is similar to those reported for streams in similar arctic and subarctic systems with sedimentary or mixed bedrock influences (Huh et al., 1998; Millot et al., 2003; Stutter and Billet, 2003; Humborg et al., 2004). The inference that silicate minerals are weathering is consistent with the observed lower concentrations of K in the cold acid digestible fraction of active-layer mineral soil compared to permafrost on six out of the seven surfaces examined in this study. At least some of this K is likely derived from the most soluble silicate minerals contained in parent material sandstones, which have from 2 to 10 times the amount of cold acid digestible fraction K than the limestone unit (Table 1).

#### NUTRIENT RELEASE VIA MINERAL WEATHERING

The ecological influence of weathering in this region is reflected in the exchangeable fraction of soils. Exchangeable  $^{87}\text{Sr}/^{86}\text{Sr}$  and cold acid digestible fraction  $^{87}\text{Sr}/^{86}\text{Sr}$  for each soil horizon are well correlated ( $R^2 = 0.74$ ) across all surfaces in this study. This suggests that locally derived mineral weathering products are the primary source of exchangeable Sr, and therefore probably also the source of ecologically important elements like Ca, P, and K to the watersheds on these surfaces.

We did not perform soil extractions specifically designed to determine the reservoirs of P in these soils. However, the ratio of Ca to P in the cold acid digestible fraction of active-layer mineral soil samples on all surfaces is consistently greater than or equal to 1.67 (Fig. 6), which is the  $\text{Ca}/\text{P}$  of apatite [ $\text{Ca}_5(\text{PO}_4)_3(\text{F}, \text{Cl}, \text{OH})$ ], the major P-bearing mineral in rocks.  $\text{Ca}/\text{P}$  is lowest and nearest to

1.67 in Sag1 soils, which have the least carbonate content as determined by  $\text{Ca}/\text{Sr}$  and  $^{87}\text{Sr}/^{86}\text{Sr}$ . Only 3 out of 52 active-layer mineral soil samples with measurable cold acid digestible fraction P have a  $\text{Ca}/\text{P}$  ratio less than 1.67 (lower by at most only 26%), suggesting that apatite is the primary source of P in this fraction. High Fe concentrations in the hot acid digestible fraction suggest the possibility that some of the P in this fraction is bound to Fe hydroxide species. However, P and Fe concentrations in this fraction are poorly correlated, so it is likely that P in this fraction is bound in other (e.g. organic) forms as well.

#### GEOCHEMICAL IMPLICATIONS OF INCREASING THAW DEPTH

Our results suggest several possible geochemical consequences of increased active-layer thickness brought about by warming climate. The most conspicuous is increasing carbonate and Ca supply to soil and stream water with increasing active-layer thickness. This change would be ubiquitous across the study area,

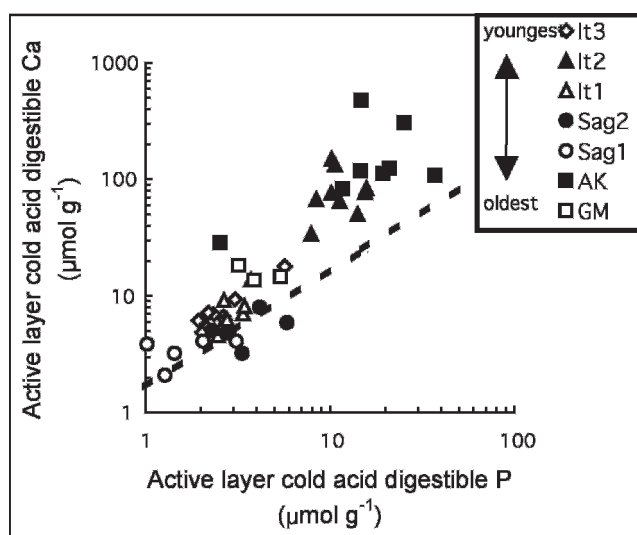


FIGURE 6. Cold acid digestible fraction  $\text{Ca}$  ( $\mu\text{mol g}^{-1}$ ) vs. cold acid digestible fraction  $\text{P}$  ( $\mu\text{mol g}^{-1}$ ) in active-layer mineral soil samples from different glacial surfaces. The dashed line represents the  $\text{Ca}/\text{P}$  of apatite (1.67). Analytical errors are reported in the text.



as all profiles exhibit geochemical indicators of increasing calcite content with depth, and analysis of data from all surfaces shows that cold acid digestible fraction Ca concentrations are significantly higher in the permafrost than in the active layer. The increased carbonate in soils brought about by calcite dissolution may lead to increased pH of acidic tundra, and potentially related changes in vegetation (Bockheim et al., 1998), but these changes would happen concurrently with and could be mitigated by changes in organic matter decomposition and organic acid production that may accompany warmer conditions and deeper thaw.

In addition to increasing carbonate dissolution and general base cation release, our measurements suggest that increased active-layer thickness may increase mineral weathering release of two important nutrients, P and K. On all surfaces there were significantly greater concentrations of exchangeable K in the permafrost than in the active-layer mineral soil. Both exchangeable and cold acid digestible fraction P concentrations were significantly greater in the permafrost than in the active-layer mineral soil overall. The mean increase in P between these layers was 28% for the exchangeable fraction and 61% for the cold acid digestible fraction. The deep permafrost sample from the Sag1 surface was enriched in cold acid digestible fraction P compared to active-layer soil (including "O" horizons) or shallow permafrost on any surface. Despite these mean differences in K and P between permafrost and active-layer mineral soil, the surface-specific differences between active-layer mineral soil and permafrost concentrations of these elements are variable. For example, the It2 surface has active-layer mineral soil concentrations of cold acid digestible fraction P that are actually higher than shallow permafrost concentrations, whereas the It3 and Sag1 surfaces have shallow permafrost concentrations of cold acid digestible fraction P that are more than double the concentrations found in active-layer mineral soil.

These data support the predictions of Hobbie et al. (1999) that thawing of permafrost could increase P availability to ecosystems and provide more information regarding possible mechanisms. Because the permafrost contains small but significantly greater concentrations of exchangeable P than the active layer, increases in available P may occur concurrently with permafrost thawing. Subsequent to initial thaw depth increases, a larger increase in newly available P may occur gradually as previously frozen minerals weather. Based on the differences in exchangeable and cold acid digestible P concentration profiles between surfaces, it is reasonable to expect that different geographic areas or watersheds may undergo different degrees of increased P availability. Some areas, such as those with already high cold acid digestible fraction P in the active layer (It2 and AR) or those where past weathering of currently frozen soil has depleted P in the first few centimeters of permafrost (Sag2), may change little in P availability until large increases (tens of centimeters) in thaw depth occur. Thaw-depth driven increases in K availability are also likely to be spatially variable, based on differences in the exchangeable and cold acid digestible fraction K depth profiles among surfaces.

Sulfur concentrations in soil show no statistically significant increase from the active-layer mineral soil to the permafrost across all surfaces for any fraction, but several individual surfaces have higher S concentrations in the permafrost than in the active-layer mineral soil in either the exchangeable (It3, It2, GM), cold acid digestible (It3, It1), or hot acid digestible (It3, It2, It1) fractions (Tables 2, 3, and 4). The Sag1 deep permafrost sample has exchangeable and cold acid digestible fraction S concentrations 100% and 25% greater, respectively, than any active-layer sample from any surface, including O horizon samples. Our analyses do not

distinguish whether the S in these soils is inorganically or organically bound, but both forms are likely because the soils are organic rich and the sandstone units of the parent material contain accessory sulfide minerals. Sulfur may be an important control on Hg methylation via sulfate-reducing bacteria (Compeau and Bartha, 1985; Gilmour et al., 1992); methylation of Hg in high arctic wetlands is an important step in Hg bioaccumulation in arctic food chains, although soils there appear to have low levels of sulfate-reducing bacteria (Loseto et al., 2004). Further study of the forms and concentrations of S in the permafrost may shed light on how increased thaw depth will influence  $\text{SO}_4$  concentrations in arctic surface waters, potentially affecting Hg methylation pathways.

#### USE OF STREAM GEOCHEMISTRY AS A MONITOR OF THAW DEPTH

The geochemical profiles of soils on the central North Slope described in this study may provide a useful tool for qualitatively detecting watershed-scale changes in thaw depth. We hypothesize that in watersheds which have sufficient changes in soil geochemistry with depth, such as the  $^{87}\text{Sr}/^{86}\text{Sr}$  or Ca/Sr trends described in this study (Fig. 3), stream geochemistry may reflect increases in thaw depth. Specifically, we predict that as thaw depth increases in this study area, more carbonate weathering products will be dissolved in late summer streamwater and will be detectable as increases in streamwater elemental ratios such as Ca/Sr and decreases in streamwater  $^{87}\text{Sr}/^{86}\text{Sr}$  over periods of many years. On some surfaces with very pronounced geochemical differences between active-layer mineral soil and permafrost, such as the It1 surface, regular analysis of stream geochemistry may signal watershed-scale permafrost degradation before it is detectable by traditional physical (e.g., steel probe) methods of measuring thaw depth. Even in permafrost regions with non-carbonate-dominated soil parent material, less intensely weathered soil at depth may provide a geochemical signature as soils thaw more deeply.

### Conclusions

Analysis of stream and soil geochemistry on arctic Alaskan glacial till deposits of varying ages indicates that carbonate weathering is the major controlling factor of geochemical trends related to soil depth and surface age. Carbonate minerals have been depleted by weathering in the active layer on every geomorphic surface, and on the four surfaces with similar parent material, stream and soil geochemistry suggests progressive carbonate depletion with time over hundreds of thousands of years. Dissolved Si concentrations in streams are similar to those in other arctic watersheds, suggesting that silicate weathering is also occurring on these surfaces, but at a much slower rate than carbonate weathering.

Compared to active-layer mineral soils, permafrost has overall greater exchangeable K concentrations, exchangeable and acid digestible P and Ca concentrations, and carbonate content. Some glacial deposit surfaces also have higher S concentrations in permafrost than in active-layer soils. The magnitude of the geochemical differences between permafrost and active-layer soils varies among different surfaces. The results of this study suggest that increasing thaw depth brought about by climate change will lead to overall (albeit spatially variable) increases in carbonate, Ca, P, K, and S supply to soils and streams. This increase may occur concurrently with increased maximum thaw as the exchangeable fraction becomes available, and then progressively as soluble minerals begin to weather. The nature of these

geochemical changes will depend on the rate of thaw; changes are likely to be gradual overall but could be locally abrupt in areas of rapid thermokarst formation. Based on geochemical differences between active-layer soil and permafrost, stream geochemistry may be a useful qualitative monitor of watershed-scale changes in thaw depth.

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