Nanoscale Observations Support the Importance of Chemical Processes in Rock Decay and Rock Coating Development in Cold Climates

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Abstract: Conventional scholarship long held that rock fracturing from physical processes dominates over chemical rock decay processes in cold climates. The paradigm of the supremacy of cold-climate shattering was questioned by Rapp’s discovery (1960) that the flux of dissolved solids leaving a Kärkevagge, Swedish Lapland, watershed exceeded physical denudation processes. Many others since have gone on to document the importance of chemical rock decay in all cold climate landscapes, using a wide variety of analytical approaches. This burgeoning scholarship, however, has only generated a few nanoscale studies. Thus, this paper’s purpose rests in an exploration of the potential for nanoscale research to better understand chemical processes operating on rock surfaces in cold climates. Samples from several Antarctica locations, Greenland, the Tibetan Plateau, and high altitude tropical and mid-latitude mountains all illustrate ubiquitous evidence of chemical decay at the nanoscale, even though the surficial appearance of each landscape is dominated by “bare fresh rock.” With the growing abundance of focused ion beam (FIB) instruments facilitating sample preparation, the hope is that that future rock decay researchers studying cold climates will add nanoscale microscopy to their bag of tools.

Keywords: Antarctica; Anthropocene; arctic; biological weathering; chemical weathering; desert varnish; frost weathering; physical weathering; rock coatings; rock varnish

Forward by Ronald I. Dorn

Dr. David Krinsley passed away 5 November 2017 at the age of 90. His career as a geologist spanned six decades. After completing his Ph.D. at the University of Chicago, then a post-doc at Columbia University, he started the Geology Department at Queens College and eventually became its Provost during the Vietnam era. After an appointment as an Overseas Fellow at Churchill College, University of Cambridge, he took a position as chair of Geology at Arizona State University. Dave’s geological research focused on diagenetic processes, including ongoing mineral decay. I met Dave in 1988 after he stepped down as chair and was immediately challenged about everything I had ever written about what Dave called “desert varnish”—questioning even my switch in terminology to rock varnish. The grilling continued until his passing. Dave retired from ASU in 1991 and moved to the University of Oregon, where we continued our collaboration, often with gifted microscopists at the CAMCOR facility. This manuscript was about a quarter finished when Dave passed away. I have done my best to stay true to our discussions in finishing this paper.
1. Introduction

Joint fractures rest at the core of geomorphic weakness [1–4]. Molnar et al. ([1], p. 12) speculated that the “role of tectonics as a crusher of rock is that in those places where rock has dodged the rock crusher, it may be stronger and less easily removed by erosive agents.” While tectonic deformation and other rock-forming processes generate joints [1,5–11], these initial fractures open the way for physical and chemical rock decay processes. Note that we eschew here the term “weathering,” except in quotations, for reasons explained elsewhere [12] and will use instead the preferred term of rock decay.

There exists little doubt that thermal stress [13–17] and frost cracking (or frost wedging) [18,19] play critical roles in cold climate geomorphic systems. The visual appearance of bare rock in alpine, arctic and Antarctic landscapes certainly encouraged the paradigm of the dominance of “physical weathering” in cold climates championed in physical geology and physical geography textbooks [20–22]. Thus, it was surprising when actual measurements in the mid-twentieth century documented the importance of chemical processes, with Rapp [23] showing that the flux of dissolved solids leaving a Kärkevågge watershed in Sweden exceeded physical denudation processes. Hall [14,24–26], Dixon [27–35], Thorn [36–38], and their collaborators followed in persistently and systematically laying an empirical and theoretical foundation for dismantling the paradigm of the dominance of “physical weathering” in cold climates. Unfortunately, there still exists the assumption by some of “minimal chemical weathering” in cold climates like Antarctica [39]. However, most scholars no longer consider biochemical and chemical rock decay processes as unimportant or even a distant second to physical rock decay, as evidenced by recent research [40–47] and Ph.D. dissertations [48] on cold-climate rock decay not having fight against those prejudiced by the old paradigm.

Nanoscale earth science methods [49–51], including studies of mineral and rock decay processes [52–60], by and large, have not yet reached cold climate rock decay research—with the prominent exception of explorations by Schindler and collaborators of anthropogenic interactions with rock surfaces [61–66]. D.H. Krinsley has been the leading pioneer in the use of nanoscale electron microscopy in the field of rock coating research [67–71], and followed by others [72–74]. Prior to his death, Krinsley had been analyzing nanoscale processes associated with cold-climate rock surface samples from alpine, arctic, and Antarctic settings provided by the second author from different sources, and J.G. Bockheim also sent Krinsley a sample of rock varnish on a desert pavement cobble from Beacon Valley, Antarctica.

Although the purpose of this paper is not to review scholarship on cold-climate rock decay research, there exists an abundance of prior research that laid the fundamental groundwork and prior context [75–88] to understand the importance of nanoscale electron microscopy. Similarly, the research conducted for this paper does not present a systematic investigation of any one cold-climate site with sufficient sampling density to make nuanced conclusions. Instead, the simple purpose of this presentation rests in illustrating the power of nanoscale electron microscopy in the study of chemical processes in cold climate regions. The findings of this paper support fully the thinking of Rapp, Hall, Dixon, Thorn, and others that chemical rock decay processes dominate rock surfaces in cold climates. The hope is that the evidence presented here encourages cold climate rock decay researchers to explore the nanoscale.

2. Samples and Methods

The samples analyzed in this study derive from alpine, arctic, and Antarctic settings (Table 1). The analysis of rock decay and the rock coatings formed on terrestrial surfaces requires study via electron microscopy (Table 2). Secondary electron (SE) microscopy, back-scattered electron (BSE) microscopy, and energy dispersive X-ray spectroscopy (EDX) are appropriate for micron-scale analyses. Nanoscale analyses requires higher resolution microscopy that often requires thinning samples to permit electrons to pass through them, where resolution varies from ~10 to 50 nm, depending on the material and the techniques used.
Table 1. Sample locations analyzed in this research.

<table>
<thead>
<tr>
<th>Figures</th>
<th>Glacier Location</th>
<th>Location</th>
<th>Rock Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>1,7,13</td>
<td>Greenland outlet glacier</td>
<td>−63.5540, −50.5853</td>
<td>gneiss</td>
</tr>
<tr>
<td>1</td>
<td>Greenland shoreline, near Nuuk</td>
<td>64.2089, −51.6130</td>
<td>gneiss</td>
</tr>
<tr>
<td>2</td>
<td>Sandstone, an older moraine, Beacon Valley, Ross Desert</td>
<td>−77.8220, 160.7148</td>
<td>sandstone</td>
</tr>
<tr>
<td>3</td>
<td>Akesu volcanic field, Tibet [89]</td>
<td>35.7001, 81.5803</td>
<td>andesite</td>
</tr>
<tr>
<td>4</td>
<td>Glacial Polish, Iztaccíhuatl, Mexico</td>
<td>19.1716, −98.6362</td>
<td>andesite</td>
</tr>
<tr>
<td>5,6,9</td>
<td>Desert pavement [90], Beacon Valley, Ross Desert, Antarctica</td>
<td>−77.8321, 160.6293</td>
<td>sandstone</td>
</tr>
<tr>
<td>8</td>
<td>Mt. Van Valkenburg, Clark Mountains, Antarctica</td>
<td>−77.3098, −142.1168</td>
<td>granite</td>
</tr>
<tr>
<td>10</td>
<td>Palisades glacier, acid mine drainage</td>
<td>37.0772, −118.4722</td>
<td>granodiorite</td>
</tr>
<tr>
<td>11</td>
<td>Ngozumpa Glacier, Nepal</td>
<td>28.0069, 86.6898</td>
<td>gneiss</td>
</tr>
<tr>
<td>12</td>
<td>Khumbu Nepal</td>
<td>27.8137, 86.7194</td>
<td>gneiss</td>
</tr>
</tbody>
</table>

Table 2. Microscopy techniques employed in this research.

<table>
<thead>
<tr>
<th>Technique (and Acronym)</th>
<th>Information Obtained</th>
<th>Spatial Resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coupled dual-beam focused ion beam electron microscopy (FIB-EM)</td>
<td>Used to create and image cross sections in situ in TEM analyses. Real time imaging in SEM mode during ion milling.</td>
<td>&gt;~1 nm</td>
</tr>
<tr>
<td>High resolution transmission electron microscopy (HRTEM) and scanning transmission electron microscopy (STEM)</td>
<td>2-D spatial imaging, lattice imaging.</td>
<td>&gt;0.08 nm</td>
</tr>
<tr>
<td>Energy dispersive X-ray analysis (EDX)</td>
<td>Elemental composition, X-ray mapping of elements. SEM provides topographic insight, while average atomic number (Z) revealed through contrast in grayscale BSE image.</td>
<td>&gt;1 nm (HRTEM) &gt;20 nm (SEM)</td>
</tr>
<tr>
<td>Scanning electron microscopy (SEM) with back-scattered electron detector (BSE)</td>
<td>2-D spatial imaging.</td>
<td>&gt;5 nm</td>
</tr>
<tr>
<td>Scanning electron microscopy (SEM) with secondary electrons (SE)</td>
<td>2-D spatial imaging.</td>
<td>&gt;1 nm</td>
</tr>
</tbody>
</table>

High resolution transmission electron microscopy (HRTEM) samples were prepared using a coupled dual-beam focused ion beam electron microscope (FIB-EM) to thin a region until it becomes electron transparent [91]. We also used scanning/transmission electron microscopes (TEM/STEM) where samples consist of polished cross sections, and HRTEMs of thinned lamellae that use much higher electron beam voltages (typically from 80 to 300 kV) than SEM. Compositional analysis employed EDX for higher atomic number elements [92]. The high voltage, along with a thinned sample, allows EDX measurements to be made with spot sizes of <10 nm.

A current leading technique that we would recommend for researchers exploring rock decay in cold climates would be the general category of focused-ion-beam (FIB) instruments. Sample preparation is a critical issue with decayed minerals and rock coating interactions with microorganisms. FIB instruments enable site-specific lamellae via HRTEM, STEM, and EDX (Table 2), where the lamellae can then have a large area of constant thickness. FIB instruments equipped with SEM and micromanipulators enable in situ lift-out approaches where a site of interest can be identified, cut free, and transferred directly to a TEM grid [93]. However, there exists the potential for the sample to resemble a theater curtain. Relatively new approaches to reduce curtaining have been demonstrated recently with xenon plasma focused ion beam systems for cutting large cross sections with high current beams [94].

We also used lower resolution method to examine anthropogenic lead pollution in the surface-most layer of a Greenland rock varnish, where we employed a wavelength dispersive electron microprobe to analyze lead concentrations at the micron scale. Lead is present in natural
manganiferous varnish, but at trace element concentrations typically less than 0.03% PbO [95]. Much higher abundances of lead occur in the surface-most layer of rock varnishes in warm desert settings, and this is attributed to anthropogenic practices adding lead to the atmosphere; the lead has a preference to adsorb onto the manganese and iron hydroxides in rock varnish [96–98]. Where EDX detected the presence of lead in the Greenland ice margin sample, wavelength dispersive spectroscopy (WDS) of a JEOL superprobe electron microanalyzer quantified lead abundance under operating conditions of 20 nA, a take-off angle of 40°, an accelerating voltage of 15 kv, and a 300 s counting time to increase sensitivity to a detection limit of about 0.03% weight PbO.

3. Results

Nanoscale observations of arctic, Antarctic, and alpine cold-climate settings contextualized by micron-scale studies reveal the ubiquitous presence of chemical processes operating at rock surfaces. Even where we observed physical decay processes cracking rocks by iron hydroxides, the iron precipitation was preceded by chemical reactions that first mobilized and then reprecipitated the iron. Nanoscale processes involve a complex interaction of the accretion of rock coatings, the mobilization of rock coating material down into the underlying rock, and the chemical decay of rock minerals. A human footprint was also found in the analyzed samples, the result of direct action such as acid mine drainage and the result of global processes such as lead pollution.

3.1. Iron Cracking as a Physiochemical Process

Beneath the surface of iron films and rock varnishes at the ocean shoreline near Nuuk, Greenland (Figure 1), minerals experience physical cracking due to the precipitation of iron, combined with a suite of other elements. EDX elemental analyses reveal the material to be similar to many iron films, with about 20% iron, 60% clay minerals (Na, Mg, Al, Si peaks), and the remainder with K, Ti, Cu, and Zr (Figure 1e). The brightness of the cracking material seen in BSE (Figure 1b–d) originates from high Z materials of Fe, Cu, and Zr. The iron was remobilized from surface coatings of iron films and rock varnish, as well as Fe-bearing minerals such as magnetite (Figure 1a) and minerals too decomposed to identify with EDX (Figure 1d).

3.2. Quartz Alteration

Using the FIB-EM, HRTEM observations revealed two different processes of quartz alteration in an Antarctic sample from Beacon Valley (Figures 2 and 3). Nanoscale dissolution pores developed within the unaltered quartz (Figure 2b,e). Dissolution pores were more commonly close to internal fractures in the quartz. In addition, a style of alteration developed around the outer boundaries of quartz that resulted in the quartz showing an amorphous texture (Figure 2a,c), where phosphorus external to the quartz (Figure 2e) chemically reacted in such a way as to become a part of this amorphous zone (Figure 2d,f). We do not know the source of the phosphorus, but it could potentially be biologically derived from proximate rock varnish [99].

Quartz alteration had a direct effect on geomorphic interactions on the Tibetan plateau. An andesite lava flow in the Akesu volcanic field contains free quartz. The surface of the lava flow’s free quartz was not a smooth surface created from conchoidal fracturing. Instead the quartz was altered in such a way as to develop a rough surface at the <10 nm scale (Figure 3a). HRTEM observations reveal that the very surface consists of bumpy ridges of unaltered quartz alternating with furrows (Figure 3c) that contain phosphorus intermixed with the quartz in EDX analyses. This roughened texture enabled dust particles to adhere to the quartz surface. The attachment of dust was not a temporary condition. Rather, the surface was rough enough to allow the accretion of iron film on the dust (Figure 3a,b).
Figure 1. Orange iron films dominate near the Nuuk shoreline, while darker rock varnish occurs further away from the coast. All of the BSE images (a–d) are 120 µm wide. (a) The iron derives in part from iron-rich minerals such as magnetite seen in BSE from the Nuuk shoreline, and (b) the dissolution of iron from rock varnish at the Greenland outlet glacier site. Image (b) shows quartz being split apart into silt-size fragments. Images (c) and (d) also show physical splitting of quartz at the Nuuk shoreline. (e) EDX analysis of the cracking material was from the letter E in image (d); the spectra reveal a mixed composition that includes Fe, Si, Al, and Mg, and smaller peaks of Zr, K, Ti, and Cu. The brightness of the line in image (d) highlighted by the arrow originates from the high atomic number elements of iron and copper.
Figure 2. Alteration of quartz from Beacon Valley. (a) Altered quartz is underneath a sequence of an iron film consisting of a granular texture, silica glaze, and Mn-rich rock varnish. Between the fresh quartz and the rock coating is a zone of about 100 nm of alteration. (b) Some of the alteration consists of nanoscale dissolution pores, while (c) another form of alteration is similar to the hydration observed by Pope in temperate settings [59]. While the quartz itself does not contain phosphorus (e), the altered quartz does include phosphorus in EDX analyses (images d and f).
Figure 3. Alteration of free quartz on the surface of a lava flow, Akesu volcanic field, Tibet. (a) A dust particle adheres to the quartz surface that maintains a roughness from nanoscale dissolution. Note the bumpy appearance of the quartz. The dust particle has adhered to the surface long enough to accumulate a titanium-iron-rich film. (b) EDX analysis of the iron-rich film that is composed of mostly Si and Al, but with major amounts of Ti and Fe. (c) The nanoscale surface of the quartz consists of microtopography of dissolution grooves.

3.3. Complex Intercalation of Cold Climate Rock Coatings

The Iztaccíhuatl volcano, Central Mexico, has a complex glacial history at high altitudes [100]. The surfaces of the Holocene moraines are commonly covered with silica glaze (Figure 4) much in the same way that the most common rock coating on Mauna Kea moraines is also silica glaze [101], but mixed with other rock coatings. Iztaccíhuatl silica glazes contain a strong aluminum peak (Figure 4d2) but with variable amounts of minor elements of P and Ti. Iztaccíhuatl silica glazes commonly contain internal nodules (e.g., Figure 4a–c) that are dominated by iron but have variable amounts of Mg, P, Ca, and Mn (Figure 4d2).
Figure 4. Silica glaze formed on glacial polish of Iztaccíhuatl, Mexico. Three BSE images (a–c) show silica glaze as the dark gray material. The silica glaze is not homogeneous. Oval zones of iron-rich material occur as discrete entities encapsulated by the silica glaze. EDX spectra in images d1 and d2 were acquired at the d1 (silica glaze) and d2 (iron-rich micronodule) spots indicated in image (b).

Several different types of silica glazes were observed in Antarctic samples from Beacon Valley, Ross Desert (Figures 2a, 5c and 6a). In FIB-EM imagery, some of the silica glaze appears to have incorporated 10-nm-scale iron granules (Figure 2a). Another texture consists of 40-nm-scale spheroids (Silica Glaze 2 in Figure 5c) that have EDX spectra indicating that a combination of aluminum glaze and silica glaze (Figure 5d). A third texture is more homogeneous (Silica Glaze 1 in Figure 5c) and is dominated by Si, with minor amounts of Al and Fe (Figure 5e).
Figure 5. Complex series of rock coatings formed on the underside of desert pavement cobble in Beacon Valley. (a) The cobble underside has a light orange coloration. (b) SE image showing patches of the rock coating formed on quartz. (c) FIB-EM section through a rock coating patch, revealing a sequence of rock coatings on top of the unaltered quartz. There are two silica glaze textures, designated 1 and 2, where Locations D and E correspond with the EDX spectra (d) and (e). There is a transition between Silica Glaze 2 and the Fe-rich granules, identified by G with the corresponding EDX analysis (g). (f) A typical EDX spectra of the uppermost layer with granular textures.
3.4. Mn-Rich Bacteria Casts

We found fossilized evidence of budding bacteria within cold climate rock varnishes. Rock varnish formed on a morainal boulder on the edge of a Greenland outlet glacier containing micron-scale casts of bacteria (Figure 7a). When viewed at the highest resolution with a BSE detector, the forms resolved as bacteria undergoing budding (Figure 7b), where the inside of the cells consists of carbon and oxygen (Figure 7c), while the casts on the outside of the cells had EDX spectra similar to those of warm desert.

Figure 6. Complex series of rock coatings formed on the top side of desert pavement cobble in Beacon Valley. (a) FIB-EM section through a rock coating patch, revealing a sequence of rock coatings on plagioclase and quartz. The base of the rock coating consists of Mn-rich varnish, followed by silica glaze. On top of the different rock coatings and rock material is a coating of Fe granules. (b) SE image of what the surface of the Fe granules look like. A few uncoated areas of sandstone quartz occur (identified by SS), but the rest of the surface is coated with Fe-rich granules that have a lamellate texture. (c) EDS analysis of the Mn-rich varnish acquired at the asterisk next to the letter C in image (a). (d) EDS analysis of the Fe granules acquired at the asterisk next to the letter D in image (a).

Many iron films accrete on rock surfaces in, for example, Antarctic [96] and arctic [31] settings. Samples from Beacon Valley, Ross Desert, reveal a nanoscale texture of iron granules not seen previously (Figures 2a, 5c and 6a). This granular texture can be produced as an artifact of sample preparation, but we are confident that this is not the case. In SE views, the iron granules occur in depressions on the underside of desert pavement cobbles (Figure 5b) and also as more continuous coatings on the top surface of the pavement cobble (Figure 6b). Additionally, these iron granules occur in different samples regardless of how they were prepared. The EDX composition of the iron granules resembles iron films on the underside of warm desert pavement clasts with higher peaks associated with clays (Mg, Al, and Si) compared with the iron hydroxides, along with minor amounts of Na, K, and Ca (Figures 5f and 6d).

Beacon Valley samples in a subaerial setting also contained a layer of Mn-rich varnish that formed the bottom layer of the observed rock coating sequence (Figures 2a and 6a). The Mn-rich varnish formed on top of the quartz and was then covered by silica glaze. Its elemental composition contained much more Mn than is typical for warm arid varnish (Figure 6c). Otherwise, the EDX spectra were similar with strong Al and Si peaks, as well as minor elements including Mg, P, K, Ca, Ti, Fe, and Cu.
Figure 7. Mn-rich casts of budding bacteria observed within rock varnish collected from a millimeter-scale depression on a meter-sized boulder on the edge of a Greenland outlet glacier. (a) BSE image of varnish formed on quartz, where the arrows identify bacteria forms. The white box is a B in the middle identifies the location of image (b) that is a BSE close-up image of bacterial casts, caught in the act of starting to bud. Budding bacteria reproduce by developing buds, and this appeared to be taking place when the bacteria were fossilized by Mn encrustation. (c) EDX analysis of the center of one of the bacteria (identified by the letter C in image b) indicates only an organic composition with carbon and oxygen. (d) EDX analysis of the outer cast of the same bacteria (identified by the letter D in image (b)) has a composition similar to rock varnish found in other environmental settings, with the exception of a stronger sulfur peak than is typical.

Rock varnish on a Mt. Van Valkenburg sample has a BSE micron-scale textural appearance (Figure 8a), similar to other observations of microstromatolitic textures in rock varnishes found in warm arid settings [102] and varnishes in humid temperate regions [103]. Upon investigation with higher resolution BSE, the laminated texture resolved into micron-scale bacteria-sized forms with a bright (high Z) atomic number. EDX analyses of these forms showed much higher concentrations of Mn (Figure 8e) than the adjacent varnish material (Figure 8f). A light treatment of the polished surface with HF to remove most of the clay minerals allowed the texture of the bacterial forms to emerge with greater clarity (Figure 8c). These forms appear to be broken segments of what could be the hyphae of budding bacteria. Certainly, the scale of these segments is appropriate for the Mn-rich casts of pieces of budding bacteria hyphae.
The importance of scale in interpreting Antarctic rock varnish can be emphasized through Figure 9 from Beacon Valley. The surface of the sample (Figure 9a) appears dark like any Mn-rich rock varnish. In a BSE image of the polished cross section, the varnish appears laminated and about 20 \( \mu \text{m} \) thick (Figure 9b) with bacteria-size forms on the surface (indicated by the letter C in Figure 9b). At the nanoscale, bacteria resolves into a granular texture (Figure 9c) that consists of equal amounts of Mn and clays (Al and Si) with minor amounts of Mg, P, K, Ca, Ba, Fe, and Cu (Figure 9d).
Figure 9. Antarctic rock varnish from Beacon Valley at vastly different scales. (a) Photograph of a 10 cm cobble from Beacon Valley. (b) BSE image of a pocket of rock varnish from one of the rock-surface depressions seen in image (a). (c) HRTEM image of the cast of bacteria located at the letter C in image (b). The granular nodules with a diameter of about 2 nm derive from ongoing dissolution of bacteria cell walls, where the letter D indicates the location of the EDX analysis (d). Note that the texture to the right of letter D has lineaments that are from clay mineral structures altered by Mn-rich granule precipitation. Also note very strong Mn Kα and Mn Kβ peaks in (d).
3.5. Anthropogenic Geochemical Impacts on Cold-Climate Rock Surfaces

Human alteration of rock surfaces in cold climates occurred in some settings that are obviously anthropogenic, such as acid mine drainage water. Other geochemical changes to cold-climate settings were a result of anthropogenic changes such as removing vegetation that in turn changes the geochemistry of rock surfaces or physically disturbs the rock surfaces associated with yak animal husbandry. Still another observed effect resulted from lead air pollution distant from cold-climate settings. This section explores each of these anthropogenic influences to cold region rock surfaces in turn.

3.5.1. Acid Mine Drainages

An abandoned titanium mine at the margin of the Palisades Glacier [104] is the source of acidified drainage water with measured pH values ranging from 2.4 to 3.5. Where this water flows over rocks, they are coated with an orange iron film. The iron film consists mostly of bacteria-sized forms, about a micron in diameter, all agglomerated together (Figure 10a) along with silica glaze that appears to be filling void space. EDX analyses of the bacteria forms reveal a composition that is more than 50% elements found in clay minerals (Mg, Al, and Si), with the second largest component being iron. Minor elements include K, Ti, Cu, and Zr (Figure 10b).

3.5.2. Coatings Altered by Human Activities in the Khumbu of Nepal

Figure 11 illustrates a common scene in the Khumbu of Nepal, where glacial stone fences made of outwash cobbles are part of yak animal husbandry and other activities. The building of such stone structures modifies rock coatings that originally developed on the stones, for example, placing rock varnish developed in a subaerial environment upside down—burying the varnish in the ground. The result is a clay coating formed over the varnish (Figure 11a). Sherpa combined yak manure with some soil material and put these yak patties on walls to dry as fuel. Upon peeling these fuel pellets off stone walls, some material is left behind—material that then lithifies and results in a clay film mixed with yak manure (Figure 11c). Yaks themselves modify rock coatings by physically stepping on them and, in the case of Figure 11b, detaching the original rock varnish coating from the stone.

Vegetation change in the Khumbu of Nepal alters the biological environment stone surfaces that used to be under a coniferous cover. Removing the acidic coniferous leaf litter increases the pH of rock surfaces. Thus, black fungi that used to thrive under conifers are being replaced by silica glaze. Figure 12a presents one example of silica glaze entombing the desiccated remains of black fungi.

Another aspect of vegetation change immediately adjacent to the Ngozumpa glacier involves the removal of moss. Small melt water streams next to the till used to support extensive areas of moss. However, removal of the moss for fuel changed the chemistry of the small streams from acidic to near neutral. pH measurements next to remnant areas of moss had pH values ranging from 3 to 4, but pH measurements where moss had been removed were between 6.5 and 7.5. The change in geochemistry resulted in a change in rock coating development (Figure 12b). Iron films first formed as a result of the acidic water, including the presence of casts of iron bacterial. However, the change in pH resulted in the accumulation of silica glaze over the iron films.
Figure 10. Iron film rock coating that developed on quartz in the drainage leading out of a titanium mine near the Palisades Glacier, Sierra Nevada, California. (a) BSE image where the brighter material has a granular texture where granules are 1 micron in scale or the size of bacteria. The darker material is silica glaze. (b) EDX analysis of a typical bacteria-sized granule.
Figure 11. Rock coating alteration as a result of yak animal husbandry adjacent to the Ngozumpa glacier, Nepal. (a) BSE image of a glacial outwash cobble, coated with rock varnish, placed in the ground as a result of building a stone wall. A clay coating formed over the varnish. (b) A glacial moraine small boulder was kicked by a yak, resulting in spalling of the rock surface. Placement of epoxy over the edge of the spall preserved the spatial relationship, preserving the ability to see in this BSE image a rock varnish coating that was physically detached from the rock. (c) Yak manure combined with some soil material dried on stone walls; removal leaves behind a lithified residue of a new artificial rock coating of a clay film are mixed with manure, as seen in this HRTEM image.
Whereas many geoscientists only see the visual scale evidence of angular fractures that rest at the core of geosciences and of coating rock surfaces. Observations presented here that consistently reveal a persistent story of chemical processes decaying geomorphic weakness [1–4], the hope is that they stop and think about the nanoscale and micron-scale chemical processes at the nanoscale that, we hope, opens the eyes of cold-climate researchers who in cold climate geomorphic settings. However, Figures 1–13 also reveal new details of rock-surface rock-surface processes. The results clearly confirm findings by Rapp [23], Hall [14,24–26], Dixon [27–35], 4. Discussion

He darkest (least electron transparent) materials in Figure 13c.

3.5.3. Lead Contamination of Rock Surface Coatings

A boulder on the side of a Greenland outlet glacier (Figure 13a) hosts rock varnish accumulations in millimeter-scale rock-surface depressions (Figure 13b). WDS electron microprobe analyses in a transect from the base of the varnish to the surface reveal lead contamination in the upper micron of the rock varnish. The WDS analyses in percent PbO had a detection limit of 0.03%. For most of the varnish, a typical condition of <0.03% PbO existed, but the surface-most micron had a lead spike reaching 0.19%. FIB-EM HRTEM imagery of the location of the microprobe measurement (Figure 13c) reveals that the lead is not evenly dispersed. Instead, EDX analyses revealed identifiable PbO peaks at he darkest (least electron transparent) materials in Figure 13c.

3.5.3. Lead Contamination of Rock Surface Coatings

4. Discussion

Figure 12. Alteration of rock coatings as a result of vegetation change in the Khumbu region of Nepal. (a) Secondary electron view of the surface of a rock that used to be underneath a conifer forest. This view shows the desiccated remains of black fungi that is surrounded by silica glaze—where the silica glaze cannot form naturally under the decaying pine needles. However, with deforestation, the less acidic pH allows silica glaze to accumulate. (b) BSE image of a cross section of rock coatings on a cobble in a stream bed next to the Ngozumpa glacier. At first, iron films and casts of iron bacteria coated rock surfaces exposed to acidic waters of moss-rich areas. Removal of the moss changed the water to a less acidic pH and this led to the deposition of silica glaze over the iron film.

3.5.3. Lead Contamination of Rock Surface Coatings

A boulder on the side of a Greenland outlet glacier (Figure 13a) hosts rock varnish accumulations in millimeter-scale rock-surface depressions (Figure 13b). WDS electron microprobe analyses in a transect from the base of the varnish to the surface reveal lead contamination in the upper micron of the rock varnish. The WDS analyses in percent PbO had a detection limit of 0.03%. For most of the varnish, a typical condition of <0.03% PbO existed, but the surface-most micron had a lead spike reaching 0.19%. FIB-EM HRTEM imagery of the location of the microprobe measurement (Figure 13c) reveals that the lead is not evenly dispersed. Instead, EDX analyses revealed identifiable PbO peaks at he darkest (least electron transparent) materials in Figure 13c.

4. Discussion

This discussion section links results presented in Figures 1–13 to prior scholarship on cold climate rock-surface processes. The results clearly confirm findings by Rapp [23], Hall [14,24–26], Dixon [27–35], Thorn [36–38], their collaborators, and others [40–47,105–107] on the importance of chemical processes in cold climate geomorphic settings. However, Figures 1–13 also reveal new details of rock-surface chemical processes at the nanoscale that, we hope, opens the eyes of cold-climate researchers who only perceive “minimal chemical weathering” [39] when they see bare rock (e.g., Figures 1, 11 and 13a). Whereas many geoscientists only see the visual scale evidence of angular fractures that rest at the core of geomorphic weakness [1–4], the hope is that they stop and think about the nanoscale and micron-scale observations presented here that consistently reveal a persistent story of chemical processes decaying and of coating rock surfaces.
were also observed in andesite weathering rinds of a recent Iztacc
with chemical processes. In the case of the Greenland observations (Figure 1), the source of the iron
(Figure 2c). The zone of quartz alteration rests underneath rock coatings (Figure 2a). Amorphous zones
liquid water [108], could influence these quartz alterations.

However, amorphous zones tend to be disorganized along zones of structural weakness. Whereas
Pope found that unweathered crystalline quartz is characterized by a regular lattice arrangement.
were also observed in andesite weathering rinds of a recent Iztacc

4.1. Nanoscale Evidence of Chemical Processes Decaying Rock Surfaces

Iron films form extensive rock coatings in Kärkevagge, Swedish Lapland [31]. In addition to
accreting on rock surfaces, the iron dissolves and mobilizes into mineral and rock fractures in the
underlying rock [31,32]. The precipitation of the iron acts as a wedging agent, breaking apart minerals
into silt and clay-sized particles (Figure 1). Thus, the physical rock decay process of iron cracking starts
with chemical processes. In the case of the Greenland observations (Figure 1), the source of the iron
derives from the chemical decay of iron minerals such as magnetite and the dissolution of rock varnish.

Pope used HRTEM to study quartz decay in dryland settings, discovering that quartz changes to
an amorphous condition with crystalline defects [59]—perhaps undergoing a form of quartz hydration.

A sample from Beacon Valley reveals what Pope observed previously [59]: amorphous quartz
(Figure 2c). The zone of quartz alteration rests underneath rock coatings (Figure 2a). Amorphous zones
also occur in quartz from the Khumbu region of Nepal near the Ngozumpa glacier (Figure 11c), where
the amorphous zones appear to be in pockets tens of nanometers across, found underneath yak manure.
In addition to quartz hydration, dissolution generates nanoscale pores (Figure 2b). Dissolution pores
were also observed in andesite weathering rinds of a recent Iztaccihuatl moraine as evidenced by
intra-mineral porosity (Figure 4a–c).

What is surprising and could not have been possible for Pope [59] to analyze given the
instrumentation in 1995 is that quartz hydration also imports elements not native to quartz (Figure 2e)
such as phosphorus (Figure 2d,f). It is possible that water vapor interactions, even in the absence of
liquid water [108], could influence these quartz alterations.
Quartz alteration, whether from hydration or another process, has the effect of making surfaces rougher. Mechanical fracturing of quartz produces conchoidal fracturing with surfaces too smooth for particle adherence. In contrast, roughened surfaces facilitate adherence of aeolian particles to rock surfaces through van der Waals forces [109,110]. The altered quartz, mixing with undecayed quartz, creates sufficient surface roughness (Figure 3c) to promote the adherence of dust particles onto exposed free quartz in the cold, dry Akesu volcanic field, Tibetan plateau (Figure 3a).

Quartz is not the only mineral undergoing chemical decay in the studied samples. Biotite feathering in the Khumbu of Nepal occurs in concert with rock varnish remobilization (Figure 11a). As the biotite splits from hydration and iron oxidation, mobilized varnish moves down into this “weathering rind” and plugs up most of the voids generated by biotite splitting. Prior research in cold climates previously found abundant evidence for the chemical decay of a variety of minerals [78,111–116].

4.2. Nanoscale Observations of Rock Coatings

A variety of rock coatings formed on analyzed the cold climate samples, notably different types of silica glaze, iron films consisting of Fe-rich granules, iron films composed of casts of bacteria, rock varnish, and notably fossilized Mn-rich budding bacteria.

Silica glaze has long been noted in Antarctic samples [74,106,117], and others have previously observed intercalation of different rock coatings, namely silica glaze and sulfate crust in Victoria Land, Antarctica [74], and rock varnish and silica glaze on the Tibetan Plateau [89]. Similarly, the different rock coatings observed here have been noted in Sweden [31], Greenland [118], Iceland [119], and Antarctica [120]. Thus, observing different types of silica glaze, different types of iron film and rock varnish is not new in cold climate research.

Sulfate-rich precipitates are frequently noted in a variety of Antarctic contexts from weathering rinds to rock coatings [74,79,107], in the Swedish Lapland [121], and it has been suggested that sulfate plays a role in barium’s behavior in rock varnish [122], sometimes being enriched and correlated with Mn and sometimes not being present in anything other than very minor amounts [123]. Thus, the presence of a strong S-peak in the bacterial sheath observed in Greenland (Figure 7d) should not be surprising.

The nanoscale observations of rock coatings presented here, however, did uncover aspects not previously noted in scholarship. Beacon Valley iron films maintain a granular texture (Figures 2a, 5c and 6a) that is not an artifact of sample preparation. The 20–70 nm diameter of the granules is much too great to be bacterial casts. However, Langworthy et al. [120] discovered that silica glaze formation on the Akesu volcanic field of the Tibetan Plateau, and perhaps elsewhere, starts as soluble Al–Si complexes [Al(OSi(OH))_3^{2+}] released from the weathering of phyllosilicate minerals. These Al–Si complexes are then stabilized through very gentle wetting such as frost sublimation or the movement of capillary water on rock surfaces. The iron-rich granules that are dominated by silica (Figure 5f) could potentially form in a similar way, given that they have a similar diameter. If gentle wetting events are responsible, then these wetting events could explain the granules’ decrease in size with depth in one sample (Figure 2a), as the granules were dissolved and then reprecipitated.

The Mn-rich casts observed in association with fossilized budding bacteria display a fibrous texture of fine spines and long bundles of hair-like manganese oxides very similar to what has been seen on the cell wall of natural microbial communities [124], culturing studies [125], and fossilized budding bacteria in varnishes non-cold-climate settings [126]. These fibrous forms could be the result of different processes, such as Mn(II) absorption [127], bacterial oxidation [126], bacterial use of Mn (IV) as extracellular respiratory electron acceptors [128], or some combination of processes.

These nanoscale observations of cold climate rock varnish are consistent with the polygenetic model of rock varnish formation where diagenesis breaks apart these cell encrustations into nanoscale granules [68] that are then remobilized and reprecipitated amongst the mixed-layer clays (e.g., Figure 9) in a process first explained by Potter ([129], pp. 174–175):
“Deposition of the manganese and iron oxides within the clay matrix might then cement the clay layer...the hexagonal arrangement of the oxygen in either the tetrahedral or octahedral layers of the clay minerals could form a suitable template for crystallization of the layered structures of birnessite.” [129]

McKeown and Post ([130], p. 712) emphasized that ongoing disequilibria continue even after the varnish formed:

“[e]ven if analysis methods are improved, the situation will remain complicated by the flexibility and great variety of Mn oxide structures. The common elements of these structures enable them to easily intergrow with and transform with one another.” [130]

Thus, while budding bacteria originally enhance the Mn–Fe (Figures 7 and 8), it is the geochemical dissolution at the nanoscale and subsequent reprecipitation of oxyhydroxides in clays (Figure 9) that results in varnish formation.

The left side of Figure 9c shows the outer portion of the bacterial cast, while the right side of Figure 9c started out as clay minerals with a typical linear structure. The granular nanoscale texture seen in this FIB-EM view is typical of how bacterial casts slowly dissolve and are mobilized away from the bacterial casts. The layering typical of clay minerals in HRTEM views of varnish [67] is being broken apart by the dissolution and then the reprecipitation of the Mn-rich granules into the clays—to the point where the layered texture is mostly lost (Figure 9c). These nanoscale movements do not create instability in the varnish laminations seen at the micron scale that are stable for tens of millennia [131–133]. Scale matters and these snanoscale movements would be analogous to cars shuffling in a parking structure composed of micron-scale laminations.

An issue that sometimes arises with regard to rock varnish (or “desert varnish”) is whether this rock coating is the same in warm deserts and cold climates. The clear and simple answer is yes. The phenomenon is the same, and different environments of formation show a similar variability outlined in detail elsewhere [96,120]. A good part of the confusion with regard to “varnish” in Antarctica is that several researchers who published early papers on the topic misnamed the rock decay features under investigation [74,134,135].

Several prior studies lack the direct observation of bacteria and other microorganisms in weathering rinds and rock coatings in Antarctica [75,77,79,134]. Thus, the natural question is why we have observed bacteria (Figures 7–10) where others have not. A number of possibilities emerge, including improvements in sample preparation techniques and microscopy such as FIB-EM (Table 2) that allow better preservation of delicate features as well as real spatial variability in the presence of preserved microorganisms. One issue pertaining to our observations of bacteria involves potential sample contamination; we think that the chance of contamination is truly remote to non-existent because we only examined material entombed naturally in rock coatings. None of the observations reported here pertain to surfaces that could have been contaminated in sample handling and transport.

4.3. Anthropogenic Geochemical Impacts on Cold-Climate Rock Surfaces

In contrast to the granular iron films of Beacon Valley, those observed in high alpine acid-mine drainage (Figure 10a) consist of bacteria-sized forms that agglomerated together. These forms appear similar to those observed in natural microbial communities [124] and resemble iron secreted on the cell walls of bacteria living in acid waters [136]. The high concentrations of titanium and zirconium could be explained by the ability of the iron to scavenge heavy metals [137].

Acidification of temperate rock surfaces results in the dissolution of rock varnish as exemplified by acid fog in Southern California [138] and industrial emissions in Western Australia [139]. In contrast, the reduction of acidity in the Khumbu region of Nepal from deforestation of acid-producing plants had the opposite effect on rock coatings. Fungi that had been active in mineral dissolution became encased in silica glaze (Figure 12a), and iron film that formed in an acidic drainage of a sphagnum bog became encased in silica glaze after the sphagnum was removed for fuel (Figure 12b).
The first study of lead in relation to rock varnish occurred with respect to the Earthen figure (or geolyph) called the Bouse Fisherman. This is the figure of a human holding a spear with a quartz tips ([96], p. 139):

“Lead accumulates in rock varnishes and dust films on desert surfaces. Electron microprobe profiles reveal that lead is a contaminant in the uppermost surfaces of rock varnishes, but these concentrations drop to background levels below the very surface of natural rock coatings that have formed since lead additives were introduced into gasoline in 1922.” (Dorn [96], p. 139)

This observation now has extensive replication [97,98,140–145]. Twentieth-century industrial activities spread lead and other elemental pollutant around the globe, even in areas distant from major lead-pollution sources [146–148]. Thus, the spike in lead on the surface of varnish next to a Greenland outlet glacier (Figure 13) should not be surprising; what is new is that the lead appears to be concentrated in nanoscale granules (Figure 13c). These TEM observations appear similar to the bacterial accumulation of lead in temperate settings [149]. The concentration of lead could also be a product of the affinity of lead for biogenic manganese oxides [150] that have a granular appearance (Figure 9c).

5. Conclusions

Every sample from alpine, arctic, and Antarctic cold climate sites studied here reveals the dominance of nanoscale chemical rock-surface processes in cold climates. Many studied samples also display evidence for the importance of microorganisms in the formation of rock varnish as well as iron film rock coatings. We hope that some of the findings here could assist in interpreting surficial processes on Mars so as to avoid the early history of ignoring chemical rock decay in terrestrial cold climates. While the visual evidence of “fresh rock” may fool some geoscientists some of the time, the ubiquitous evidence of chemical rock decay and chemical processes influencing rock coatings leads to the inevitable conclusion that it is “chemical weathering” that truly dominates rock decay in cold climates.

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