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
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## *Gallionella* and *Sulfuricella* populations are dominant during the transition of boreal potential to actual acid sulfate soils

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Acid sulfate soils release metal laden, acidic waters that affect the environment, buildings, and human health. In this study, 16S rRNA gene amplicons, metagenomes, and metatranscriptomes all demonstrated distinct microbial communities and activities in the unoxidized potential acid sulfate soil, the overlying transition zone, and uppermost oxidized actual acid sulfate soil. Assembled genomes and mRNA transcripts also suggested abundant oxidized acid sulfate soil populations that aligned within the Gammaproteobacteria and *Terracidiphilus*. In contrast, potentially acid tolerant or moderately acidophilic iron oxidizing *Gallionella* and sulfur metabolizing *Sulfuricella* dominated the transition zone during catalysis of metal sulfide oxidation to form acid sulfate soil. Finally, anaerobic oxidation of methane coupled to nitrate, sulfate, and ferric reduction were suggested to occur in the reduced parent sediments. In conclusion, despite comparable metal sulfide dissolution processes e.g., biomining, *Gallionella* and *Sulfuricella* dominated the community and activities during conversion of potential to actual acid sulfate soils.

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Natural deposits containing sulfidic sediments<sup>1</sup> are widespread along coastal areas worldwide such as Baltic Sea, Asia, and Australia<sup>2</sup>. These sediments are predominantly stable if they remain anoxic, such as under the sea or the groundwater table<sup>3</sup>. However, coastal areas containing sulfidic sediments have been dredged for purposes such as agriculture and housing. This exposes the sulfides to atmospheric oxygen and initiates microbial-aided chemical reactions<sup>4</sup> that turn potential acid sulfate soil materials (PASS) into actual acid sulfate soil materials (ASS; pH ≤ 4), which are described as the “nastiest soils on earth”<sup>5</sup>. This conversion to ASS releases large quantities of acidity that concomitantly increases the mobility of toxic metal(loid)s such as Al, As, Cd, and Zn that are transported to surrounding waters where they cause severe problems for the environment<sup>6,7</sup>, economy<sup>8</sup>, and human health<sup>5</sup>.

Pyrite (FeS<sub>2</sub>) oxidation is a complex biogeochemical process that involves a series of chemical reactions that are catalyzed by microbial re-oxidation of ferrous iron<sup>9</sup> and metabolism of intermediate sulfur compounds formed during mineral breakdown<sup>10</sup>. Pyrite dissolution is extensively studied in the context of the industrial process of “biomining”<sup>11</sup> and acid mine drainage<sup>12</sup> and the geochemical reactions that occur when PASS are oxidized to ASS have also been elucidated<sup>2,13</sup>. Culture based studies of ASS identify sulfur-oxidizing microbes<sup>14</sup> and acidophiles such as *Acidithiobacillus ferrooxidans* and *Acidithiobacillus thiooxidans*<sup>15</sup>. In addition, 16S rRNA gene amplicon-based studies of Australian coastal PASS are dominated by sequences aligning with Proteobacteria<sup>16</sup> while sub-tropical ASS contain iron-oxidizers such as *Alicyclobacillus tolerans*, *Leptospirillum* sp., *Sideroxydans lithotrophicus*, and *S. paludicola*<sup>17</sup>. Metagenomic analysis also demonstrates a shift in microbial populations when a Chinese sub-tropical PASS was exposed to oxygen<sup>18</sup>. Finally, Australian coastal ASS split into “high organic,” “surface tidal,” and “sulfuric” zones shows a disproportionately higher number of active microbial populations from the “rare biosphere” (defined as the great number of low abundance microorganisms found in an environment) that were involved in iron- and sulfur cycling<sup>19</sup>. Despite the importance of ASS in the release of metals to the environment, the microbial communities and activities in the underlying sediments and during the process of metal sulfide oxidation have been largely neglected.

The Risöfladan experimental field is situated on the Ostrobothnia coast in Western Finland. The Risöfladan parent marine sediments contain approximately 50% FeS<sub>2</sub> and 50% metastable iron sulfides (FeS<sub>n</sub>; *n* = 1.0–1.3) that were drained over 60 years ago and have since developed into a typical clay-type ASS<sup>20,21</sup>. The experimental field is utilized to investigate the mitigation of metal and acid release<sup>22–27</sup> and the remediation of ASS. Microbial communities in the Risöfladan ASS include populations with 16S rRNA gene sequences most similar to acidophiles such as *Acidocella* sp. and *Acidithiobacillus ferrivorans* as well as populations similar to species identified from low temperature and metal contaminated environments<sup>13,26</sup>. Finally, 16S rRNA gene amplicons assigned to sulfur and/or iron cycling populations including *Acidithiobacillus*, *Gallionella*, *Sulfuricurvum*, and *Sulfurimonas* were identified from metal sulfide containing river sediments from the vicinity of Risöfladan when they were exposed to air<sup>28</sup>. However, most previous ASS studies only used 16S rRNA gene sequencing to identify relative abundances of populations during the generation of boreal ASS and the metabolic potential and active processes of the populations during this transition remain uncertain.

In this study, soil samples were collected from a Risöfladan depth profile that were used to investigate the microbial community structure by high throughput amplicon sequencing of the 16S rRNA marker gene; metabolic processes coded within

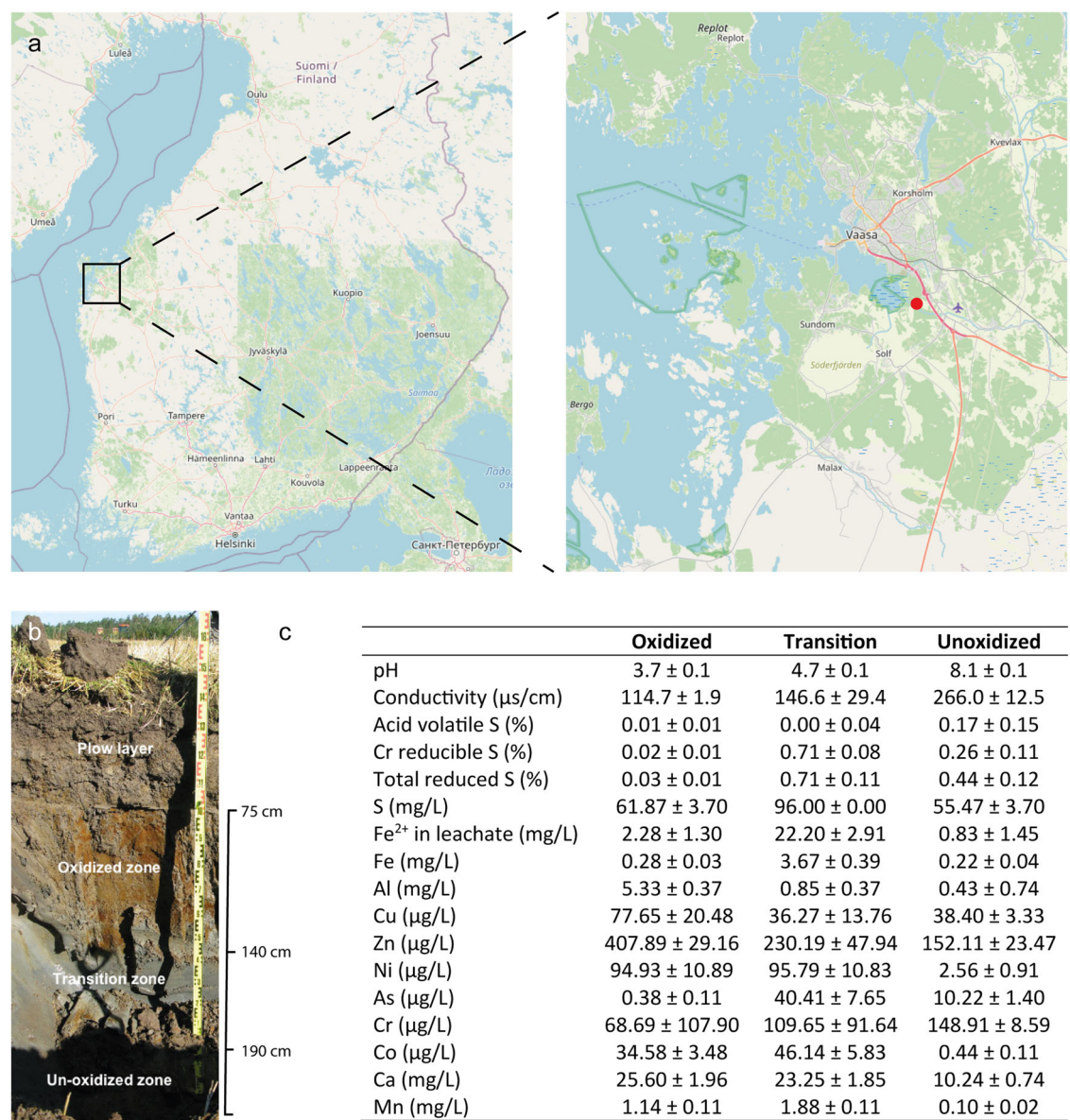
reconstructed genomes by community DNA (metagenomics) sequencing; and community RNA (metatranscriptomics) based activity of the reconstructed genomes that shows the processes being carried in the oxidized, transition, and unoxidized soil zones. The tested hypotheses were that (i) the community structure of microbes is altered during PASS conversion to ASS and (ii) in contrast to extreme acidophiles found in biomining or acid mine drainage environments<sup>29,30</sup>, the conversion of PASS to ASS selects for moderate acidophiles.

## Results

Sampling was performed when the microbes were most active i.e., when the cracks in the ASS were depleted of water and filled with air. Triplicate samples were taken from each soil zone (Fig. 1); namely, the ASS oxidized zone (defined as “OX”; depth 75–140 cm), the transition zone (“TR”; 140–190 cm) that has a steep pH gradient from acidic to near neutral, and a pH neutral unoxidized PASS zone (“UN”; >190 cm).

**Geochemical characteristics of the soil zones.** The oxidized zone was characterized as a typical ASS, with a pH <4 and high redox potential (Fig. 1 and Supplementary Table S1). The pH and conductivity significantly increased with soil depth, respectively (Spearman’s rank correlation; pH-depth  $R^2 = 0.97$  and  $p < 0.001$ , conductivity-depth  $R^2 = 0.84$  and  $p = 0.004$ ). The concentration of total leachable S, most likely as sulfate (SO<sub>4</sub><sup>2−</sup>), was significantly different (S-depth  $R^2 = 0.90$  and  $p = 0.001$ ) in the soil zones with the highest concentrations in the transition zone and the lowest in the unoxidized zone. Acid volatile sulfide (AVS; comprising of mainly metastable iron sulfides) showed significantly highest values in the unoxidized zone (AVS-depth  $R^2 = 0.80$  and  $p = 0.01$ ). Although chromium reducible sulfur (CRS; mainly pyrite) showed highest values in the transition zone, no significant differences were found between the soils (CRS-depth  $R^2 = 0.48$  and  $p = 0.193$ ). Furthermore, no significant differences were found between the soils regarding the total reduced sulfur (TRS; all reduced sulfur species; TRS-depth  $R^2 = 0.47$  and  $p = 0.197$ ) although the TRS in the oxidized zone was lower than in the other two zones. All total leachable Fe was identified as ferrous iron in all soil zones with the highest concentrations in the transition zone, although not found statistically significant (FeII-depth  $R^2 = -0.369$  and  $p = 0.33$ ; Fe-depth  $R^2 = 0.16$  and  $p = 0.68$ ). This was likely due to ferric attack on the iron sulfide generating ferrous iron in the bioleaching process along with ferric precipitation at pH values >1.8<sup>11</sup> that gave the highest total Fe value in the transition zone followed by lower values in the oxidized and unoxidized zones. Seasonal contributions to the Fe value in the transition zone may come from the oxidized zone via percolating water and from the unoxidized zone through capillary movement. The concentrations of leachable Al (Al-depth  $R^2 = 0.74$  and  $p = 0.023$ ), Cu (Cu-depth  $R^2 = 0.69$  and  $p = 0.042$ ), and Zn (Zn-depth  $R^2 = 0.95$  and  $p < 0.001$ ) were significantly higher in the oxidized zone compared to the transition and unoxidized zones. Furthermore, Ca (Ca-depth  $R^2 = -0.85$  and  $p = 0.004$ ) and Ni (Ni-depth  $R^2 = 0.84$  and  $p = 0.004$ ) were significantly higher in both the oxidized and the transition zones compared to the unoxidized zone. Finally, the concentrations of leachable As (As-depth  $R^2 = 0.48$  and  $p = 0.195$ ), Cr (Cr-depth  $R^2 = 0.05$  and  $p = 0.893$ ), Co (Co-depth  $R^2 = -0.47$  and  $p = 0.197$ ), and Mn (Mn-depth  $R^2 = -0.47$  and  $p = 0.197$ ) did not differ significantly between the three soil zones.

**16S rRNA gene-based microbial diversity in the soil zones.** The 16S rRNA gene amplicon sequencing generated an average of

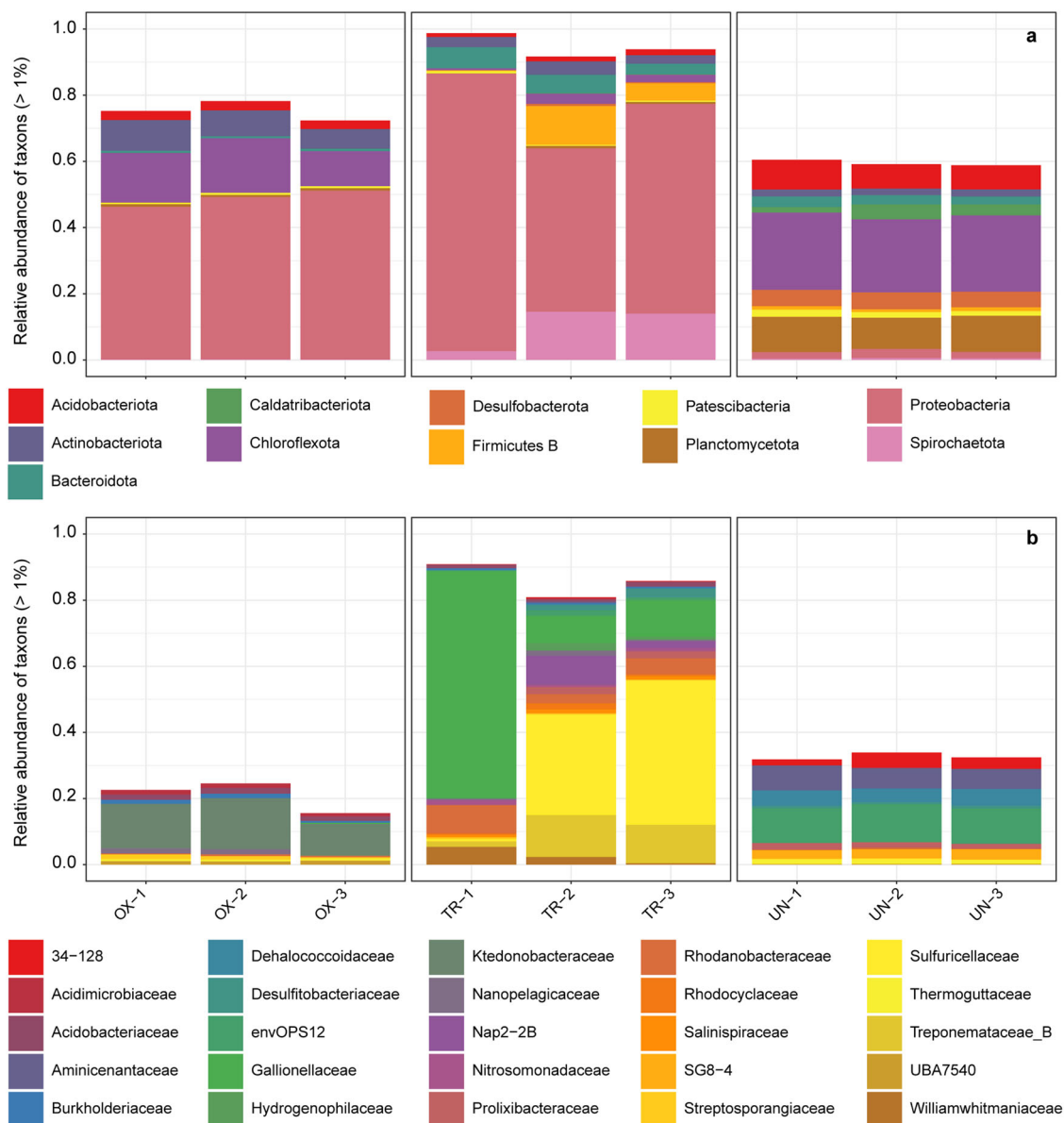


**Fig. 1 Sampling site, depth profile through the soil, and key characteristics.** The figure shows the sampling site near Vaasa, Finland marked with a red dot (a), a photograph of the plough layer at the top, oxidized ASS, transition zone, and unoxidized PASS at the deepest depth (b), and basic geochemical characteristics (c) (data are averages ± SD; n = 3; complete data in Supplementary Table S1). The maps were generated by © OpenStreetMap under an Open Database License.

369,027 reads per soil zone and were assigned to a total of 4380 different individual gene sequences that are termed “amplicon sequence variants” or ASVs<sup>31</sup>. The rarefaction curves showing the number of individual ASVs versus the sequencing depth (Supplementary Fig. S1) plateaued for all samples indicating sufficient sequencing was performed to identify the majority of the populations. A significantly higher Shannon alpha diversity (Supplementary Fig. S2) was found in the unoxidized zone compared to the oxidized (one-way ANOVA with Tukey’s post hoc test; OX-UN  $p < 0.001$ ) and transition zones (TR-UN  $p < 0.001$ ), while no difference in diversity was found between the oxidized and transition zones (OX-TR  $p = 0.103$ ).

All of the 16S rRNA gene ASVs were classified as Bacteria of which eleven phyla had a total relative abundance >1% (Fig. 2 and Supplementary Fig. S3). The reason why no archaeal 16S rRNA genes were amplified may have been due to primer bias as the amplicons were generated using polymerase chain reaction (PCR) primers designed for bacterial sequences (although they have

been demonstrated to amplify some Archaea<sup>32</sup>) in combination with the low abundance of archaea (described below). A significant portion of the microbial communities in the three soil zones were represented by unassigned Bacteria (21–27, 1–6, and 37–39% relative abundance in the oxidized, transition, and unoxidized zones, respectively). The bacterial community composition and proportions of the identified phyla varied between the soil zones. Proteobacteria (46–51%), Chloroflexota (11–16%), Actinobacteriota (6–9%), and Acidobacteriota (3%) dominated in the oxidized zone. Of these four phyla, unassigned ASVs within the Gammaproteobacteria (43–47%), the Ktedonobacteraceae family (10–15%), and unassigned Actinobacteria (3%) accounted for the majority of the relative abundance. In the transition zone, the Proteobacteria (49–84%) comprised of ASVs aligning to the Gallionellaceae (8–69%), Sulfuricellaceae (1–47%), and Rhodanobacteraceae (3–9%) families. In the unoxidized zone, the community was dominated by Chloroflexota (22–23%), Planctomycetota (9–11%), Acidobacteriota (7–9%), Desulfobacterota



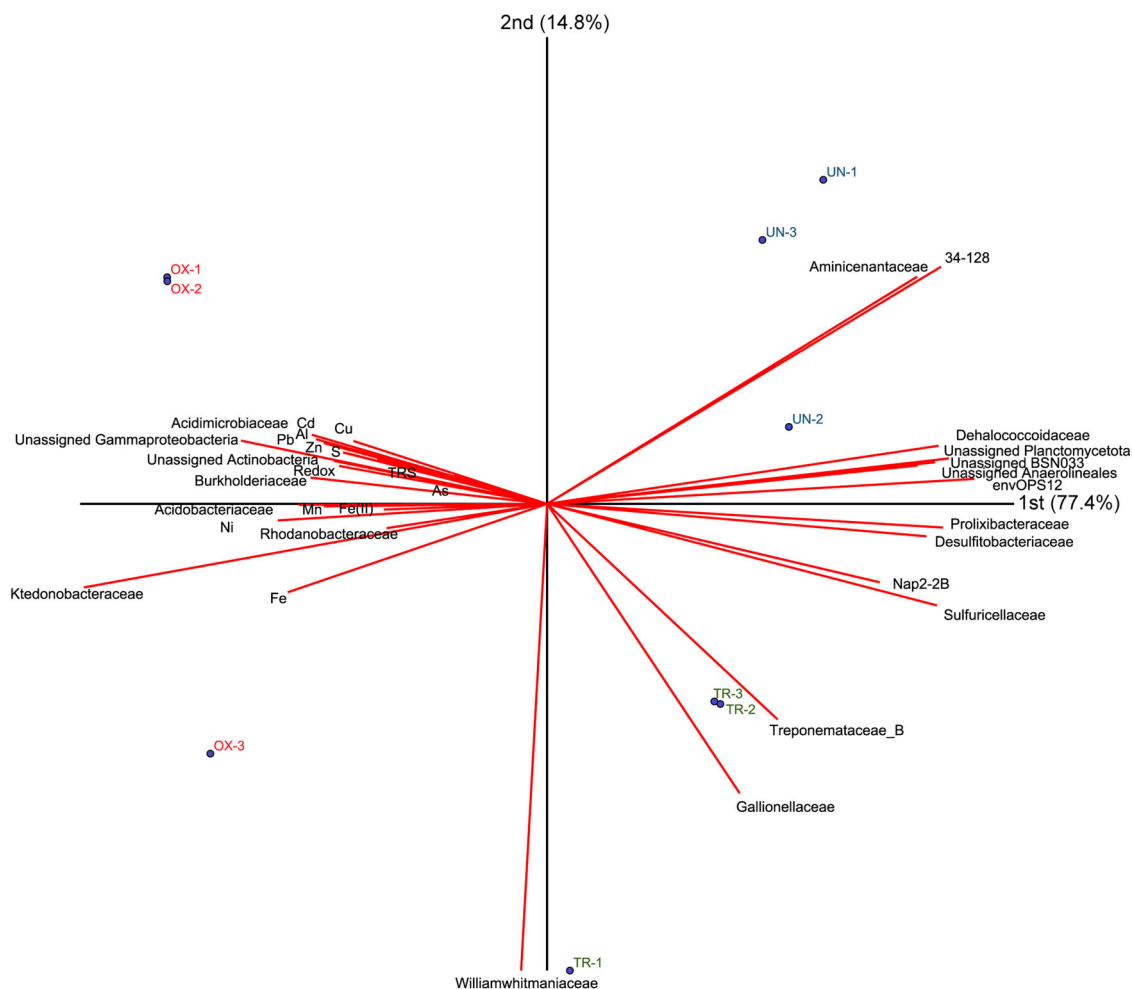
**Fig. 2 Stacked bar relative abundance of 16S rRNA gene amplicon sequencing.** The stacked bars show three replicates for the oxidized (OX), transition (TR), and unoxidized (UN) zones assigned to phyla (a) and family (b). The remaining abundance consists of unassigned ASVs and phyla or families with a relative abundance of <1%.

(5%), and Caldatibacteriota (2–4%). Of these phyla, dominating ASVs belonged to the envOPS12 (10–11%), Dehalococcoidaceae (4–5%), unassigned Planctomycetota (4–5%), Aminicenantaceae (6–7%), unassigned BSN033 (1–2%), and the 34–128 family (2–4%).

**Correlation between geochemical factors and microbial populations.** The compositional principle component analysis biplot (Fig. 3; explaining 92.2% of the variation) showed that the biogeochemical environments in the three zones were distinct. The oxidized zone microbial populations were linked with high redox and high concentrations of Al, Cu, Ni, and Zn. In contrast to the analysis of the geochemical data alone (described above) and when ratios between all factors were taken into account; S,  $\text{Fe}^{2+}$ , Mn, As, Cd, and Pb were all grouped with the samples from the oxidized zone. Although the ASVs from the Burkholderiaceae, Acidobacteriaceae, and Acidimicrobiaceae families showed low total relative abundances (1, 1–2, and 1%, respectively) in the 16S

rRNA gene amplicon data, the short links between metals and these ASVs suggested they have a substantial oxidizing role in this zone. The transition zone samples had some of the longest links from the center (e.g., ASVs from the Williamwhitmaniaceae, Gallionellaceae, and Treponemataceae B families), which indicated that their relative abundance were the most dissimilar to the other zones. The negative correlation between ASVs belonging to the Sulfuricellaceae and Nap2-2B families and TRS indicated that these ASVs might be sulfur oxidizers. Finally, the unoxidized zone ASVs also had some of the longest links from the center, i.e., ASVs from the Aminicenantaceae and 34–128 families that were not identified in the other zones.

**Metagenome and metatranscriptome data.** The triplicate metagenomes per soil type had an average mapping percentage of 84% between DNA reads and metagenomic assemblies<sup>31</sup> (Supplementary Table S2) suggesting a good coverage of the three soil communities. Bioinformatic assignment of the community DNA



**Fig. 3 Bi-plot geochemical and 16S rRNA sequencing data.** The compositional principal component bi-plot shows the combined geochemical and 16S rRNA sequencing data from triplicate samples from the oxidized (OX; red), transition (TR; green), and unoxidized (UN; blue) zone.

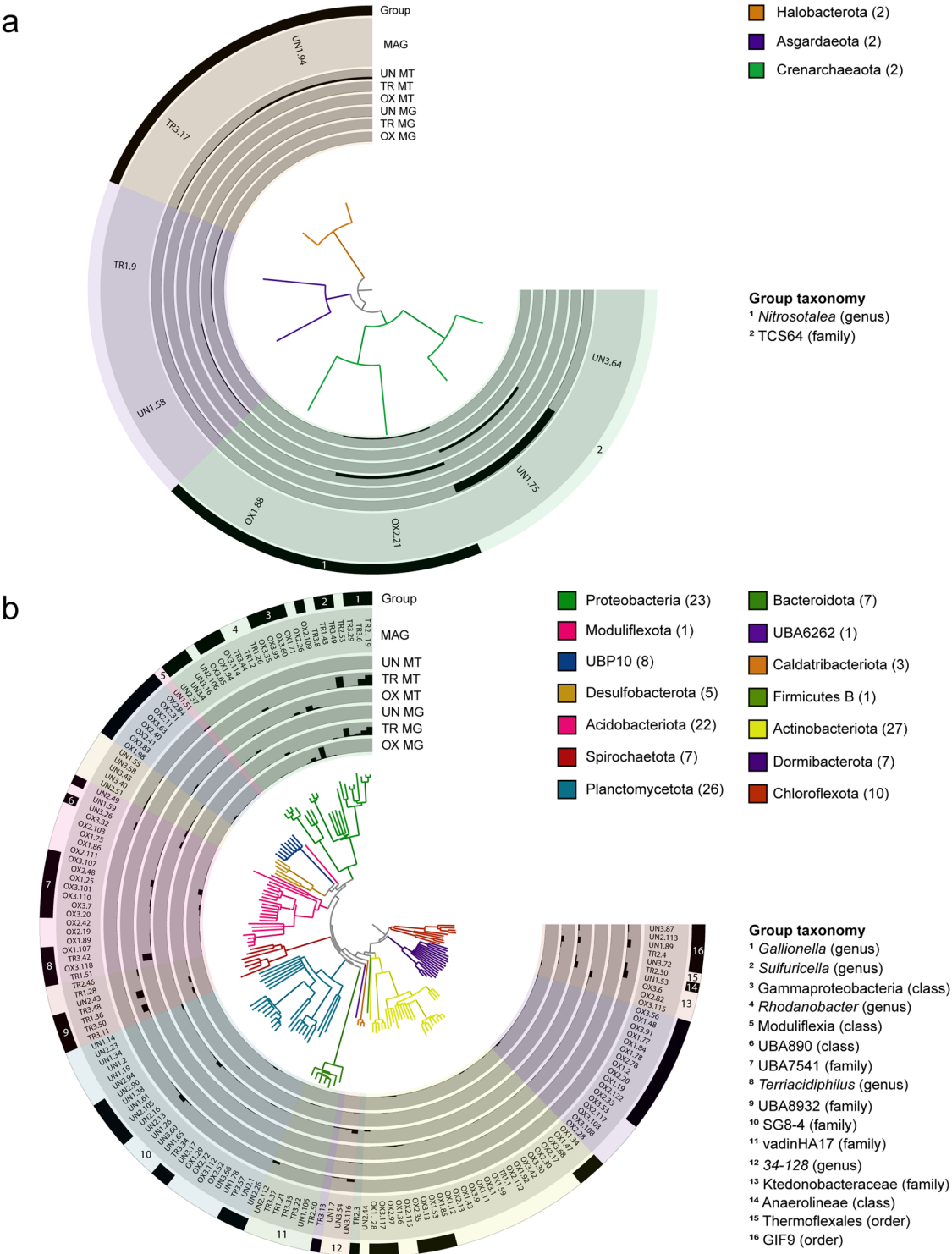
reads (binning) generated 882 metagenome assembled genomes (MAGs) with >75% completeness and <10% contamination that were de-replicated to give 293 unique populations (Supplementary Data S1). Based upon the minimum information for publication<sup>33</sup>, the data included 140 “high-quality draft” MAGs with >90% completeness and <5% contamination. The three metatranscriptomes per soil type had an average read mapping percentage of 85% to the assemblies<sup>31</sup> (Supplementary Table S2). Non-metric multidimensional scaling analysis of the reads and transcripts separated the soils on the first axis while the second axis separated the metaomics data for the transition zone community (Supplementary Fig. S4). This indicated that MAG presence was proportional to RNA transcript activities in the unoxidized PASS and oxidized ASS zones.

In many cases, phylogenetic assignment of the MAGs within the GTDB taxonomy was not achieved beyond, e.g., the class (Supplementary Data S1). For instance, the Planctomycetota MAGs UN2.108, UN2.11, and UN3.86 were not assigned to a class and four unclassified Gammaproteobacteria MAGs (OX1.71, OX3.35, OX3.60, and OX3.95) clustered with the Woessiales and Steroidobacterales orders (Supplementary Fig. S5).

#### MAG based microbial diversity and activity in the soil zones.

All MAGs with a relative abundance or RNA transcript-based activity >1.0% in any one of the three soil zones were categorized into taxonomic groups (Supplementary Data S2 and S3). The phylum with the largest DNA and RNA based relative abundance

in the oxidized zone was Proteobacteria with 34.7% and 21.5%, respectively (Fig. 4 and Supplementary Data S4). Of these abundances and activities, 31.2 and 15.1% were comprised of MAG group Ox\_Gammaproteobacteria compared to <0.1% abundance and activity in the transition and unoxidized zones. Four MAG groups assigned to the Acidobacteriota soil bacteria Acidobacteriota<sup>34</sup> made a further 18.4 and 34.8% of the relative abundance and activity, respectively. OxTr\_Terracidiphilus genus MAGs had a low relative abundance of 1.2% but a high activity of 19.3% of populations typically growing at pH 3 to 6 that matches the acidic conditions in the oxidized ASS<sup>35</sup>. MAGs grouped as Ox\_UBA7541 (13.9 and 10.7%) mediate iron cycling in tropical soils and are suggested to be named *Candidatus Acidoferrales*<sup>36</sup>. Actinobacteriota MAG groups constituted 11.7 and 10.1% of the oxidized zone community and activity, respectively and were dominated by MAGs assigned to the Acidimicrobiales that are typically associated with metal contaminated and acidic environments with the RAAP-2 family being identified in acid mine drainage<sup>37</sup>. In addition, five MAGs in two groups were assigned to the Streptosporangiaceae family found in soils and marine sediments<sup>38</sup> named Ox\_UBA9676 (3.4 and 3.9%) and OxTr\_UBA8262 (2.1 and 2.8%). Groups Ox\_Dormibacteraceae and Ox\_UBA8260 assigned to the Dormibacteria class that has been identified from acid mine drainage<sup>39</sup> accounted for 8.1 and 2.6% of the relative abundance and activity, respectively that was significantly higher than the deeper soil zones (Kruskal–Wallis test with Bonferroni correction; all  $p < 0.05$ ; Supplementary



**Fig. 4** Archaeal and Bacterial domains showing phyla from the MAG groups. Numbers in brackets refer to the number of representative MAGs assigned to each phylum in the GTDB and the group taxonomy define selected abundant MAG groups for Archaeal (a) and Bacterial (b) domains. Bars indicate the relative abundance as percentages of the total reads mapped to the MAGs in the oxidized (OX), transition (TR), and unoxidized (UN) soil zones (metagenome, MG) and protein coding RNA transcripts mapped to the MAGs in the three soil zones (metatranscriptome, MT).

Table S3). Finally, one Archaeal MAG group, Ox\_Nitrososphaerales (3.4 and 8.0%) contained two MAGs with one further aligning with the autotrophic (i.e., carbon dioxide fixing) ammonia oxidizing *Candidatus* Nitrosotalea genus of phylum Thermoproteota<sup>40</sup>. Proteobacteria also dominated the transition zone community with Gammaproteobacteria MAG groups constituting 54.4 and

59.3% of the relative abundance and activity, respectively (Fig. 4 and Supplementary Data S3). Proteobacteria groups included the Tr\_Gallionella genus (29.1 and 33.2%) that are characterized as iron-oxidizers<sup>41</sup> and the Tr\_Sulfuricella genus (17.0 and 20.8%) that are autotrophic sulfur-oxidizers<sup>42</sup>. Both these MAG groups had significantly higher relative abundance and activity in the transition zone compared to both the oxidized and unoxidized

zones (all  $p < 0.05$ ). MAG group Tr\_Rhodanobacter (4.9 and 4.2%) was significantly more abundant and active compared to the unoxidized zone ( $p < 0.05$ ) with members of the genus identified as acid-tolerant denitrifiers<sup>43</sup>. The sulfur oxidizing *Thiobacillus* genus<sup>44</sup> MAG group Tr\_Thiobacillus (1.8 and 0.2%) was also only present in the transition zone. Finally, two MAG groups assigned to the Spirochaetota phylum were split into the mostly free living anaerobes in the Spirochaetales order<sup>45</sup>.

Planctomycetota with 18.1 and 13.6% relative abundance and activity was most abundant in the unoxidized zone (Fig. 4 and Supplementary Data S3) that was further divided into MAG groups attributed to three Phycisphaerae, one Brocadia, and one Planctomycetes class. The partitioning of these MAG groups differed in the other two zones with UnTr\_Phycisphaerae constituting a minor portion of the transition zone but was all but absent in the oxidized zone (all  $p < 0.05$  except for OxUn\_Thermoguttaceae). Characterized Phycisphaerae are aquatic bacteria associated with marine algae<sup>46</sup> while Brocadia have been detected in marine sediments<sup>47</sup> and therefore, may have been present from before the PASS materials were drained. The second most abundant and active phylum group was attributed to the Chloroflexota group UnTr\_GIF9 from the Dehalococcoidia that constituted 7.5 and 9.2% of the relative abundance and RNA transcript activity, respectively that was equally abundant in the unoxidized and transition zones, but significantly more active in the unoxidized zone ( $p < 0.05$ ). Characterized Dehalococcoidia are obligate anaerobes identified in e.g., soils and aquatic environments<sup>48,49</sup>. Two additional Chloroflexota MAG groups were UnTr\_Anaerolineae and Un\_Thermoflexales that were significantly more abundant and active ( $p < 0.05$ ) in the unoxidized zone compared to oxidized zone. The next most abundant bacterial phylum was the Caldatribacteriota (group UnTr\_34-128) with 10.9 and 7.9% of the total relative abundance and activity in the unoxidized zone, respectively that had significantly higher relative abundance ( $p < 0.05$ ) compared to the oxidized zone and was most similar to candidate bacteria 34–128 that lacks a detailed description<sup>50</sup>. Further abundant and active populations included UnTr\_Desulfobacteria (5.7 and 6.1%) that had significantly higher abundance and activity compared to the oxidized zone (all  $p < 0.05$ ). MAG groups that aligned with the Archaea included Un\_TCS64 from the Thermoproteota class Bathyarchaea (6.7 and 21.7%) that have been identified in e.g., marine environments, soils, and sediments where they may have an ecological role in symbiosis with Methanomicrobia<sup>51</sup>. A further unoxidized zone archaeal group was Un\_ANME-1ex4572 (0.6 and 7.1%) that aligned with the Halobacterota class Syntropharchaea that are characterized as anaerobic methane producers<sup>52</sup>. A third archaeal MAG group was UnTr\_Heimdallarchaea (2.3 and 0.5%) that aligned with the Asgardarchaeota class Heimdallarchaea that partitioned in the soil types with the highest abundance and activity in the unoxidized zone compared to the oxidized zone (all  $p < 0.05$ ).

**mRNA transcript based metabolic processes in the three soil zones.** Key genes were investigated in the RNA transcripts (Figs. 5–7, Supplementary Data S4, and Supplementary Fig. S6) to create a model of the major metabolic processes in the three zones (Fig. 8). While some MAG groups were identified in more than one zone, key groups were predominantly found in a single zone suggesting a clear distinction between the active populations. For instance, the average number of RNA transcripts ( $n = 3 \pm \text{SD}$ ) were considerably higher in the transition versus unoxidized zone encoding ferrous iron ( $7177 \pm 2198$  and  $680 \pm 25$  TPMs) and sulfur compounds ( $1244 \pm 302$  and  $1 \pm 1$  TPMs) oxidation that likely catalyzed the conversion of PASS to actual ASS. In contrast,

transcripts for methanogenesis (i.e., methane production) were predominantly identified in the unoxidized compared to the transition zone ( $4575 \pm 533$  and  $582 \pm 106$  TPMs). Furthermore, transcripts for both ferrous oxidation and ferric reduction genes suggested separate populations in aerobic and anaerobic soil niches or alternatively, multiple populations with different metabolic strategies were included within some MAG groups.

The oxidized zone MAG group (Fig. 5) with the highest mRNA-based activity was OxTr\_Terracidiphilus that was likely involved in organic matter decomposition<sup>35</sup> potentially coupled to the reduction of sulfate (*dsvABC*) as electron acceptor. The group also showed transcripts for ferrous iron oxidation (*cyc1/2* and *foxABYZ*) found in acidophile species that preferentially grow at low pH<sup>53</sup>; adaptation to low pH<sup>54</sup> (e.g., *kdpABCDE* and *shc*); metals resistance<sup>55</sup> including Cu, Co-Zn-Cd, and As (*copA*, *czcABC*, and *acr3* and *arsBC*); and *sodA*, *bcp*, and *dnaX* for oxidative stress<sup>56</sup> likely due to the ferrous ion. The Ox\_Gamma-proteobacteria group RNA transcripts suggested involvement in iron cycling<sup>53</sup> via ferrous iron oxidation genes typically found in acidophiles and species that grow at neutral pH (i.e., neutrophiles) along with iron reduction (e.g., *dmkB*, *fmnB*, *cymA*, *mtrABC*, and *ndh2*) as the terminal electron acceptor. This MAG group also had RNA transcripts annotated for acidic metal containing environments such as low pH stress<sup>54</sup> and metals resistance<sup>55</sup>. The Acidobacteriae Ox\_UBA7541 transcripts also suggested its involvement in iron cycling with low transcript numbers for acidophilic and neutrophilic ferrous oxidation and with ferric iron reductases as a terminal electron acceptor<sup>36,53</sup> along with transcripts typical for ASS such as pH homeostasis and metals resistance. Finally, the Archaea MAG group Ox\_Nitrososphaerales from the Thermoproteota phylum showed transcript based activity in iron oxidation<sup>53</sup> and dissimilatory iron reduction<sup>53</sup>.

The transition zone MAG group (Fig. 6) with the highest mRNA-based activity was Tr\_Gallionella that was suggested to mediate ferrous iron oxidation via *cyc1/cyc2*, *foxAB*, *foxYZ*, *mtaA*, and *mtrB*. In addition, this group had transcripts for thiosulfate oxidation involving the SOX complex (*soxAX*) and anaerobic sulfate reduction for energy conservation. Once again, either this suggested separate populations within the MAG group or oxic versus anoxic niches mediating sulfur cycling. In addition, transcripts were assigned for pH homeostasis; metals resistance; and oxidative stress. The Spirochaetota phylum Tr\_Treponematales MAG group RNA transcripts were assigned to iron oxidation and methanogenesis from acetate (*acs*, *pta*, and *ackA*) for energy generation; sulfate reduction and denitrification (i.e., conversion of nitrate to  $\text{N}_2$  gas; *norB*) for energy conservation; and adaptation to the low pH, high concentration of metals, and oxidative stress. The Tr\_Sulfuricella RNA transcripts confirmed the geochemical suggestion (Fig. 3) that the populations were involved in sulfur oxidation (*fccAB* for sulfide oxidation and *SoxAX*) coupled to nitrate reduction/denitrification (*narGIY*, *norB*, *napA*, and *nosZ*) or ferric iron reduction (*dmkAB*, *eetB*, *fmnB*, *mtrB*, and *cymA*). Finally, Tr\_Rhodanobacter had transcripts for thiosulfate oxidation (*tsdA*) that was likely coupled to nitrate reduction and denitrification as terminal electron acceptors<sup>57</sup> and to a lesser extent ferric iron reduction.

The unoxidized zone MAG group (Fig. 7) with the highest mRNA-based activity was Un\_TCS64 of the Thermoproteota class Bathyarchaea with transcripts encoding genes involved in organic carbon oxidation potentially coupled to dissimilatory iron reduction, *mttB* for methanogenesis<sup>51</sup>, and metals resistance. Group Phycisphaerae Un\_SG8-4 was suggested to grow via organic carbon oxidation coupled to ferric reduction along with transcripts coding for metal resistance. Archaeal phylum Halobacterota MAG group Un\_ANME-1ex4572<sup>52</sup> showed RNA



**Fig. 5 Bacterial domains showing MAG groups from the oxidized soil zone.** Bars indicate protein coding RNA transcripts mapped to the MAGs in the soil zone metatranscriptome. Maxima for each band was set to: 881 (dissimilatory iron reduction), 29 (dissimilatory nitrate reduction), 1929 (dissimilatory sulfur reduction), 5277 (Fe oxidation), 5612 (low pH), 1654 (metals resistance), 2481 (methanogenesis), 6249 (oxidative stress), and 1259 (sulfur oxidation). Numbers in brackets refer to the number of representative MAGs assigned to each phylum in the GTDB and the group taxonomy define selected RNA transcript based active MAG groups.

based activity of genes involved in anaerobic oxidation of methane via reversed methanogenesis suggested to be initiated by the *mcrA* encoded methyl-coenzyme M reductase<sup>58</sup>. Finally, MAG group UnTr\_Desulfobacteria showed transcripts for genes involved in nitrate reduction (*narHI*), iron reduction, and sulfur reduction (*psrA*, *dsvABC*, *trrB*, and *phsA*) as terminal electron acceptors

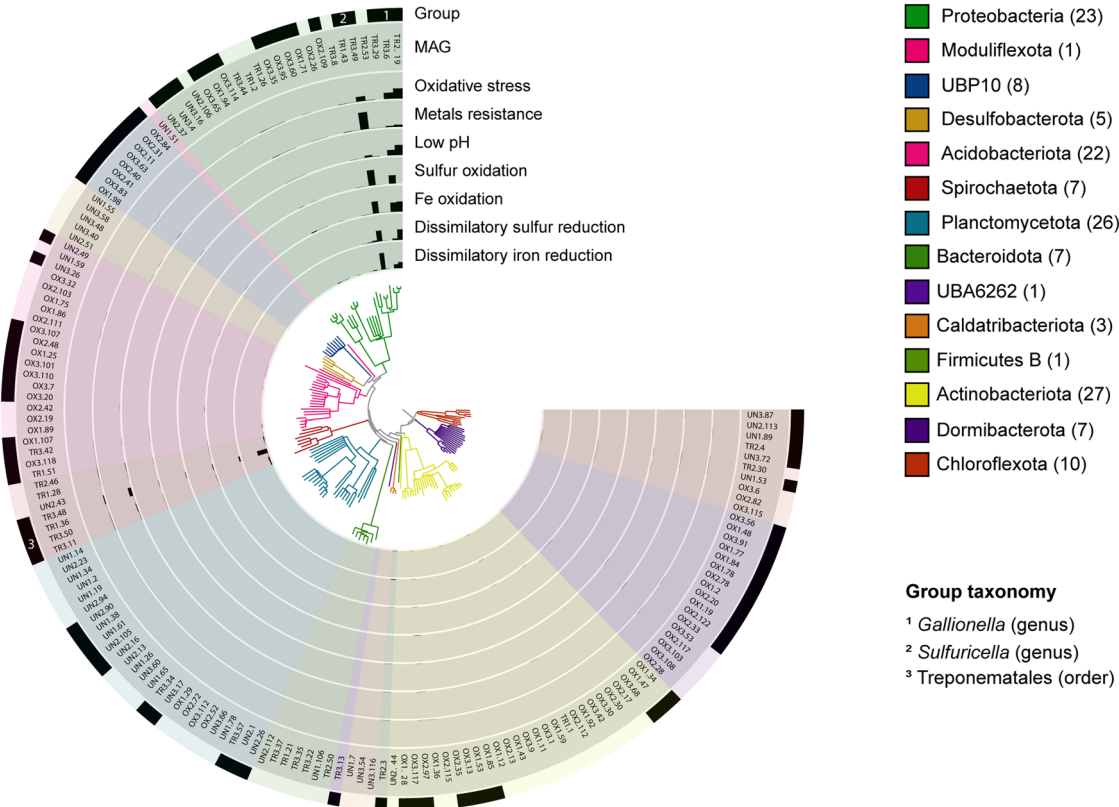
## Discussion

The low pH in the oxidized and transition zones coupled with the highest concentrations of ferrous iron (although not significant) and leachable sulfur (most likely as sulfate) in the transition zone supported that metal sulfide oxidation occurred<sup>20</sup>. The results from the sulfur speciation also supported that pyrite can be preserved in the transition zone while the environment in the oxidized zone favors pyrite oxidation and thus metal mobilization<sup>20</sup>. The data also demonstrated that metals and metalloids were freed from the metal sulfides in the parent sediments and were leachable from the soil during the spring thaw such that they could cause environmental damage such as fish kills<sup>59</sup>.

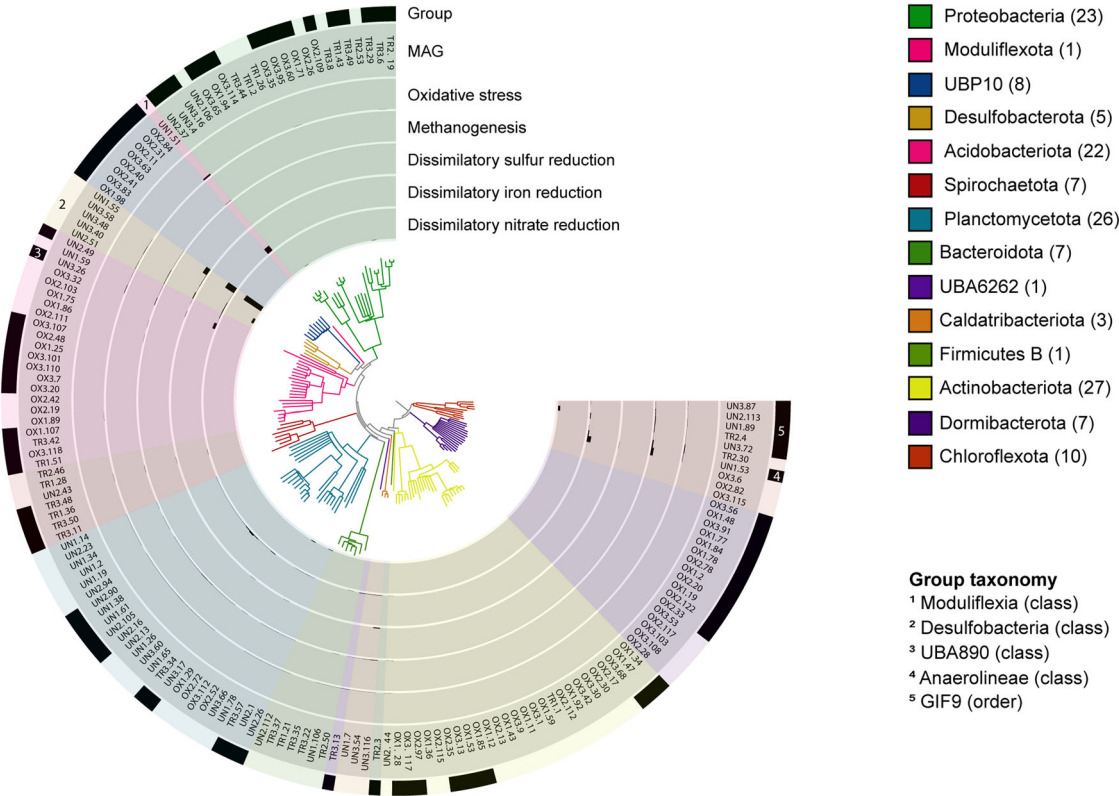
In general, the populations identified as dominating the three soil zones are present in PASS and actual ASS boreal environments suggesting the data can be extrapolated to Finnish and Swedish soils. For instance, the oxidized zone *Terracidiphilus* from the Acetobacteraceae, Ox\_UBA7541 from the Acidobacteriae, and Ox\_Nitrososphaerales have been identified in Swedish<sup>60</sup> and Finnish<sup>61</sup> ASS. The key transition zone genus *Gallionella* are detected in several other boreal ASS environments in Sweden<sup>60</sup> and Finland<sup>13,24,28</sup> along with Australian coastal ASS drains<sup>17</sup> while

*Sulfuricella* are present in Finnish ASS<sup>26</sup>. In addition, *Rhodanobacter* are present in dredged river sediment containing sulfidic materials deposited on land nearby the Risöfladan field<sup>28</sup> along with bioremediation experiments treating the Risöfladan ASS<sup>24,61</sup>. While fewer studies have been carried out on Scandinavian unoxidized PASS, the Desulfobacterota class has been identified in Swedish ASS environments<sup>60</sup>. Finally, despite the distance and contrasting conditions, the unoxidized zone selected for a similar microbial community as an Australian coastal ASS where 16S rRNA gene sequences were identified aligning with sulfate reducing Desulfobacterales, methanogenic Halobacteria, metal reducing Phycisphaerae, organohalide-respiring Dehalococcoidetes, and strictly anaerobic Anaerolineae<sup>16</sup>.

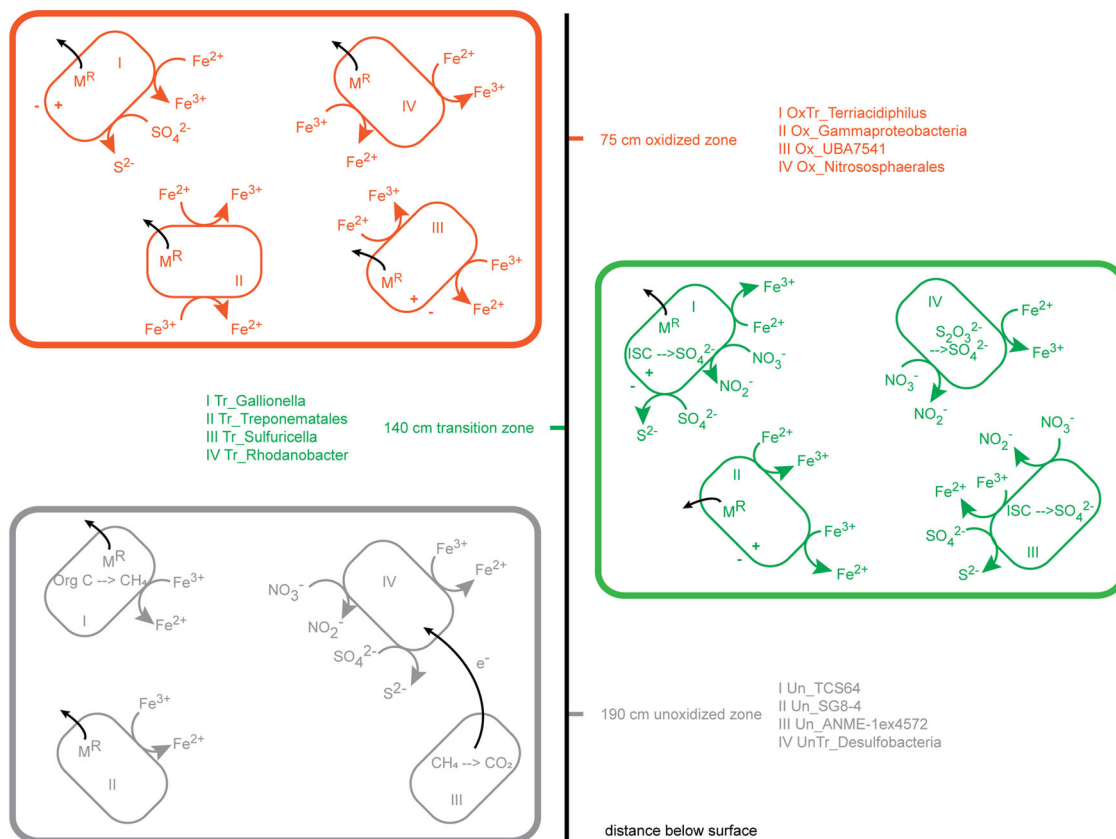
The dominant ferrous iron and sulfur oxidizing microbes in the transition zone were from the *Gallionella* and *Sulfuricella* genera. In contrast to the original description of *Gallionella* species as solely iron oxidizers<sup>62</sup>, but in agreement with results from a low pH acid mine drainage site<sup>63</sup>, the Tr\_ *Gallionella* group was also suggested to oxidize reduced sulfur compounds. In published *Gallionella* genomes, ferrous oxidation is attributed to MtoAB<sup>64</sup>, while the Tr\_ *Gallionella* group in this study was also suggested to code for *cyc1/cyc2* that are involved in ferrous oxidation by the *Acidithiobacillus* genus<sup>65</sup> and *fox* genes from e.g., the acidophile *Sulfuracidifex metallicus*<sup>66</sup>. *Sulfuricella* species oxidize reduced sulfur compounds at near neutral pH conditions in soils<sup>67</sup> and freshwater lakes<sup>42</sup> with the MAG group transcripts also supporting anaerobic sulfate and nitrate reduction for energy conservation<sup>63</sup>. The high relative abundance and activity of *Gallionella* and *Sulfuricella* species in the gradual transition of PASS to ASS in natural environments was in contrast to rapid PASS oxidation in an aerated bioreactor wherein extreme



**Fig. 6 Bacterial domains showing MAG groups from the transition soil zone.** Bars indicate protein-coding RNA transcripts mapped to the MAGs in the soil zone metatranscriptome. RNA transcript maxima for each band and numbers in brackets are as described for Fig. 5.



**Fig. 7 Bacterial domains showing MAG groups from the unoxidized soil zone.** Bars indicate protein-coding RNA transcripts mapped to the MAGs in the soil zone metatranscriptome. RNA transcript maxima for each band and numbers in brackets are as described for Fig. 5.



**Fig. 8 Model of microbial mediated processes and populations in the oxidized (OX), transition (TR), and unoxidized (UN) soil zones.** Key MAG groups and selected metabolisms are represented (black arrows denote transport processes).  $+/-$  across the cytoplasmic membrane shows the low pH adaptation of an internal positive membrane potential; MR refers to metal resistance by efflux of the metals from the cell; ISC inorganic sulfur compound oxidation; and  $e^-$  for extracellular electron transfer. Assignment of metabolic characteristics was aided using metagenome analysis by DRAM<sup>115</sup> while the figure was adapted from Wu et al.<sup>116</sup>.

acidophiles including *Acidithiobacillus* and *Acidocella* were also identified<sup>13</sup>. In addition, rapid oxidation of the dredged river sediment mentioned above was dominated by 16S rRNA gene sequences most similar to the extremely acidophilic genus *Acidithiobacillus*<sup>28</sup>. The transition zone ASVs, assembled genomes, and mRNA transcripts also identified Tr\_Rhodanobacter as involved in thiosulfate oxidation. Many of the transition zone MAG groups contained RNA transcripts coding for pH homeostasis<sup>54</sup>. Mechanisms to maintain a near neutral cytoplasmic pH are divided into the “first line of defense” primarily identified in acidophiles (rather than neutrophiles) that stop the influx of protons into the cell followed by the “second line of defense” that removes protons once they have entered<sup>68</sup>. For instance, the Tr\_Gallionella and Tr\_Treponematales groups had mRNA transcripts assigned to the “first line of defense” potassium importer genes *kdpABCDE* suggested to form an internal positive membrane potential to hinder proton influx in acidophiles<sup>69</sup>. Tr\_Gallionella also encoded “second line of defense” genes such as arginine decarboxylase in the arginine-dependent acid resistance (ADAR) system<sup>68</sup> plus a major histone-like protein HU that is suggested to regulate the ADAR system in *Escherichia coli*<sup>70</sup>. However, the lack of RNA transcripts for these genes suggested the Tr\_Gallionella group was not acid stressed in the transition zone. In contrast, the oxidized zone Ox\_UBA7541 group RNA transcripts not only coded for the potassium-transporting ATPase system but also *shc* squalene-hopene cyclase for “first line of defense” cell membrane associated pH homeostasis system<sup>71</sup>. Therefore, the MAG groups in the oxidized and

transition zones were predominantly acid tolerant or moderate acidophiles with some populations potentially being adapted to lower pH values or exhibiting transcripts relate to pH stress. Finally, several oxidized and transition zone MAG groups were suggested to be adapted to metal containing environments with transcripts coding for arsenic, cadmium, cobalt–zinc–cadmium, and mercury resistance<sup>55</sup>.

Several poorly characterized populations including the archaeal MAG groups Un\_ANME-1ex4572 and Un\_TCS64 dominated the unoxidized zone but were not identified in the 16S rRNA gene amplicons (potentially due to the PCR bias mentioned above). The unoxidized zone MAG groups were suggested to grow via anaerobic energy conservation including organic carbon oxidation by Un\_TCS64 plus Un\_SG8-4 populations that may have been retained in the sediments from their marine origins<sup>46,72</sup> and UnTr\_Anaerolineae. Unoxidized zone populations potentially also carried out sulfate reduction by e.g., UnTr\_Desulfobacteria that contained transcripts coding for genes involved in nitrate, iron, and sulfur reduction. Once the inorganic electron acceptors for energy conservation had been consumed in the unoxidized zone, then a transition may have occurred whereby the deeper community altered to favor methanogenic populations such as Un\_TCS64<sup>58</sup>. The community within the unoxidized zone may also have mediated anaerobic oxidation of methane (AOM)<sup>73</sup> by the methanotrophic (i.e., methane consuming) Archaeal Un\_ANME-1ex4572 group from the Halobacterota. However, AOM only occurs during symbiosis between a methanotrophic population in association with, e.g., a sulfate reducing population

such as UnTr\_Desulfobacteria. In support of AOM occurring in the unoxidized zone, MAG groups UnTr\_GIF9 and UnTr\_34-128 that aligned with the Dehalococcoidia and Caldatribacteriota, respectively strongly correlate with AOM rates in Baltic Sea sediments<sup>74</sup>.

In conclusion, the three soil zones had highly distinct populations and activities supporting hypothesis one that the microbial communities altered during PASS conversion to ASS. The transition zone *Gallionella* population contained RNA transcripts for a typical acidophile first line of defense potassium uptake system suggesting at least some of the transition zone populations supported the hypothesis two of moderate acidophiles mediating PASS conversion to ASS. In contrast, anaerobic metabolic processes dominated the unoxidized zone with methanogenic methane being consumed during AOM.

## Methods

**Study site and soil sampling.** Soil was sampled in mid-August 2017 from typical farmland acid sulfate soil in the area at the Risöfladan experimental field in Vaasa, Finland (63° 02' 50.22"N, 21° 42' 41.85"E; Fig. 1). The oxidized (OX), transition (TR), and unoxidized (UN) soil zones were exposed using a mechanical digger, surfaces that had come into contact with the scoop aseptically scraped away, and soil from the three zones collected (see Supplementary Methods for details). Samples were analyzed for pH, oxidation–reduction potential, conductivity, ferrous iron, multi-elements, and sulfur speciation as described in the Supplementary Methods. Three replicates from the OX, TR, and UN zones were used in all analyses.

**RNA/DNA preparation and sequencing.** RNA and DNA were extracted, processed, quality controlled, and sequencing libraries prepared as described in the Supplementary Methods. The RNA concentrations after DNase treatment and rRNA depletion as well as the total amount of RNA and DNA are in Supplementary Table S4 and detailed in Högfors-Rönholm et al.<sup>31</sup>. All metagenome and metatranscriptome sequencing was performed by the Joint Genome Institute in Walnut Creek, CA, USA on the Illumina NovaSeq platform generating 2 × 151 bp sequences. The metagenomes and metatranscriptomes were assembled, binned, mapped, and processed as described in the Supplementary Methods.

**16S rRNA gene amplicon sequencing and analysis.** Two-step 16S rRNA gene PCR amplification (V3–V4 region) using primers from Hugerth et al.<sup>75</sup> and sequencing at the SciLifeLab, Sweden (2 × 300 bp) were carried out as previously described<sup>24,31</sup>. ASVs were derived using the DADA2 pipeline<sup>76</sup> and annotated against the GTDB bacterial small subunit collection (release 86)<sup>77</sup>. Additional downstream analysis is described in the Supplementary Methods.

**Reporting summary.** Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

## Data availability

The 16S rRNA gene amplicons are available from the NCBI BioProject ID PRJNA524144<sup>78</sup>. The raw sequencing data are available from the NCBI Sequence Read Archive under accession numbers SRP18508216<sup>79</sup>, SRP18508417<sup>80</sup>, SRP18508618<sup>81</sup>, SRP18508719<sup>82</sup>, SRP18508820<sup>83</sup>, SRP18508921<sup>84</sup>, SRP18509222<sup>85</sup>, SRP18509123<sup>86</sup> and SRP18522324<sup>87</sup> for metagenomes and SRP18522225<sup>88</sup>, SRP18509426<sup>89</sup>, SRP18509327<sup>90</sup>, SRP18522528<sup>91</sup>, SRP18522429<sup>92</sup>, SRP18522630<sup>93</sup>, SRP18522731<sup>94</sup>, SRP18522832<sup>95</sup> and SRP18522933<sup>96</sup> for metatranscriptomes as detailed in Högfors-Rönholm et al.<sup>31</sup> and summarized in Supplementary Table S5. The final assembled contigs are available from the Integrated Microbial Genomes and Microbiomes (IMG/M) portal under ID numbers 3300031671<sup>97</sup>, 3300031672<sup>98</sup>, 3300031670<sup>99</sup>, 3300031669<sup>100</sup>, 3300031578<sup>101</sup>, 3300031673<sup>102</sup>, 3300031566<sup>103</sup>, 3300031565<sup>104</sup> and 3300031539<sup>105</sup> for metagenomes and 3300030718<sup>106</sup>, 3300030716<sup>107</sup>, 3300030712<sup>108</sup>, 3300030713<sup>109</sup>, 3300030710<sup>110</sup>, 3300030711<sup>111</sup>, 3300030714<sup>112</sup>, 3300030719<sup>113</sup> and 3300030717<sup>114</sup> for metatranscriptomes, and summarized in Supplementary Table S5.

## Code availability

All software versions and scripts needed to reproduce the results are specified in the Materials and Methods section and in the Supplementary Methods.

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## Author contributions

E.H.-R. and M.D. conceived the study. E.H.-R. and S.E. took the samples. E.H.-R., S.C., M.L.-F., and P.Ö. prepared nucleic acids and analyzed samples. D.L., D.B., S.C., and T.L. analyzed the data. E.H.-R. and M.D. wrote the manuscript. All authors commented upon and approved the final version.

## Competing interests

The authors declare no competing interests.

## Additional information

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