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Peatland Acidobacteria with a dissimilatory sulfur metabolism

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19

21 Abstract

22 Sulfur-cycling microorganisms impact organic matter decomposition in wetlands and consequently greenhouse gas emissions 23 from these globally relevant environments. However, their identities and physiological properties are largely unknown. By 24 applying a functional metagenomics approach to an acidic peatland, we recovered draft genomes of seven novel Acidobacteria 25 species with the potential for dissimilatory sulfite (dsrAB, dsrC, dsrD, dsrN, dsrT, dsrMKJOP) or sulfate respiration (sat, 26 aprBA, qmoABC plus dsr genes). Surprisingly, the genomes also encoded DsrL, which so far was only found in sulfur-27 oxidizing microorganisms. Metatranscriptome analysis demonstrated expression of acidobacterial sulfur-metabolism genes in 28 native peat soil and their upregulation in diverse anoxic microcosms. This indicated an active sulfate respiration pathway, 29 which, however, might also operate in reverse for dissimilatory sulfur oxidation or disproportionation as proposed for the 30 sulfur-oxidizing Desulfurivibrio alkaliphilus. Acidobacteria that only harbored genes for sulfite reduction additionally encoded 31 enzymes that liberate sulfite from organosulfonates, which suggested organic sulfur compounds as complementary energy 32 sources. Further metabolic potentials included polysaccharide hydrolysis and sugar utilization, aerobic respiration, several 33 fermentative capabilities, and hydrogen oxidation. Our findings extend both, the known physiological and genetic properties of 34 Acidobacteria and the known taxonomic diversity of microorganisms with a DsrAB-based sulfur metabolism, and highlight

35 new fundamental niches for facultative anaerobic Acidobacteria in wetlands based on exploitation of inorganic and organic

36 sulfur molecules for energy conservation.

37

39 Introduction

40 Specialized microorganisms oxidize, reduce, or disproportionate sulfur compounds of various oxidation states (-II to +VI) to 41 generate energy for cellular activity and growth and thereby drive the global sulfur cycle. The capability for characteristic 42 sulfur redox reactions such as dissimilatory sulfate reduction or sulfide oxidation is not confined to single taxa but distributed 43 across different, often unrelated taxa. The true extent of the taxon-diversity within the different guilds of sulfur 44 microorganisms is unknown (Wasmund et al., 2017). However, ecological studies employing specific sulfur metabolism genes 45 (e.g., dissimilatory adenylyl-sulfate reductase-encoding aprBA, dissimilatory sulfite reductase-encoding dsrAB, or soxB that 46 codes for a part of the thiosulfate-oxidizing Sox enzyme machinery) as phylogenetic and functional markers have repeatedly 47 demonstrated that only a minor fraction of the sulfur metabolism gene diversity in many environments can be linked to 48 recognized taxa (Meyer et al., 2007; Müller et al., 2015; Watanabe et al., 2016). A systematic review of dsrAB diversity has 49 revealed that the reductive bacterial-type enzyme branch of the DsrAB tree contains at least thirteen family-level lineages 50 without any cultivated representatives. This indicates that major taxa of sulfate-/sulfite-reducing microorganisms have not yet 51 been identified (Müller et al., 2015).

52 Wetlands are among those ecosystems that harbor a diverse community of microorganisms with reductive-type DsrAB, most 53 of which cannot be identified because they are distant from taxonomically classified DsrAB sequences (Pester et al., 2012). 54 Sulfur-cycling microorganisms provide significant ecosystem services in natural and anthropogenic wetlands, which are major 55 sources of the climate-warming greenhouse gas methane (Kirschke et al., 2013; Saunois et al., 2016). While inorganic sulfur 56 compounds are often detected only at low concentration (lower µM range), fast sulfur cycling nevertheless ensures that 57 oxidized sulfur compounds such as sulfate are rapidly replenished for anaerobic respiration. The activity of sulfate-reducing 58 microorganisms (SRM) fluctuates with time and space, but at peak times can contribute considerably to the anaerobic 59 mineralization of organic carbon in wetlands (Pester et al., 2012). Simultaneously, SRM prevent methane production by 60 rerouting carbon flow away from methanogenic archaea. Peat microorganisms that are affiliated to known SRM taxa, such as 61 Desulfosporosinus, Desulfomonile, and Syntrophobacter, are typically found in low abundance (Loy et al., 2004; Costello and 62 Schmidt, 2006; Dedysh et al., 2006; Kraigher et al., 2006; Pester et al., 2010; Steger et al., 2011; Tveit et al., 2013; Hausmann 63 et al., 2016). In contrast, some microorganisms that belong to novel, environmental dsrAB lineages can be considerably more 64 abundant in wetlands than species-level dsrAB operational taxonomic units of known taxa (Steger et al., 2011). However, the 65 taxonomic identity of these novel dsrAB-containing microorganisms and their role in sulfur and carbon cycling has yet to be 66 revealed.

To identify these unknown DsrAB-encoding organisms and further investigate their fundamental ecological niches, we recovered thirteen metagenome-assembled genomes (MAGs) encoding DsrAB from a peat soil through a targeted, functional metagenomics approach. We analyzed expression of predicted physiological capabilities of the MAGs by metatranscriptome analyses of anoxic peat soil microcosms that were periodically stimulated by small additions of individual fermentation products with or without supplemented sulfate (Hausmann *et al.*, 2016). Here, we show that facultatively anaerobic members of the diverse *Acidobacteria* community in wetlands employ one or more types of dissimilatory sulfur metabolism.

73

75 Materials and methods

76 Anoxic microcosm experiments, stable isotope probing, and nucleic acids isolation

77 DNA and RNA samples were retrieved from a previous peat soil microcosm experiment (Hausmann et al., 2016). Briefly, soil

- 78 from 10–20 cm depth was sampled from an acidic peatland (Schlöppnerbrunnen II, Germany) in September 2010, and stored at
- 79 4 °C for one week prior to nucleic acids extractions and set-up of soil slurry incubations. Individual soil slurry microcosms
- 80 were incubated anoxically (100% N atmosphere) in the dark at 14 °C, and regularly amended with low amounts (<200 μM)
- 81 of either formate, acetate, propionate, lactate, butyrate, or without any additional carbon sources (six replicates each). In
- 82 addition, half of the microcosms for each substrate were periodically supplemented with low amounts of sulfate (initial
- 83 amendment of 190–387 μM with periodic additions of 79–161 μM final concentrations). DNA and RNA were extracted from
- 84 the native soil and RNA was additionally extracted from the soil slurries after 8 and 36 days of incubation.
- Furthermore, DNA was obtained from the heavy, ¹³C-enriched DNA fractions of a previous DNA-stable isotope probing
- 86 (DNA-SIP) experiment with soil from the same site (Pester et al., 2010). Analogous to the single-substrate incubations, anoxic
- soil slurries were incubated for two months with low-amounts of sulfate and a ¹³C-labelled mixture of formate, acetate,
- 88 propionate, and lactate. DNA was extracted, separated on eight replicated density gradients, and DNA from a total of 16 heavy
- fractions (density $1.715-1.726 \text{ g mL}^{-1}$) was pooled for sequencing.

94

Additional DNA was obtained from soils that were sampled from different depths in the years 2004 and 2007 (Steger *et al.*,
2011).

92 **Quantitative PCR and metagenome/-transcriptome sequencing**

93 Abundances of Acidobacteria subdivision 1, 2, and 3 in soil samples from different years and depths were determined by

95 DNA (two libraries), heavy ¹³C-enriched DNA (three libraries), and native soil RNA, and RNA samples from the microcosms

newly-developed 16S rRNA gene-targeted real-time quantitative PCR (qPCR) assays (Supplementary Methods). Native soil

- 96 were sequenced on an Illumina HiSeq2000 system (Supplementary Methods).

97 Binning, phylogeny, and annotation of DsrAB-encoding genomes

98 The differential coverage binning approach by Albertsen et al. (2013) was applied to extract MAGs of interest. The raw 99 FASTQ paired-end reads were imported into the CLC Genomics Workbench 5.5.1 (CLC Bio) and trimmed using a minimum 100 Phred quality score of 20 with no ambiguous nucleotides allowed. TruSeq adapters were removed and a minimum length filter 101 of 50 nt was applied. This resulted in 214, 171, 233, 49, and 294 million reads after quality filtering and trimming for the two 102 native soil and three SIP metagenomes, respectively (84-95% of the raw reads). All reads were co-assembled using CLCs de 103 novo assembly algorithm (kmer size 63, bubble size 50, minimum scaffold size 1000 nt). The reads from all five metagenomes 104 were independently mapped to the assembled scaffolds using CLCs map to reference function (minimum length 0.7, minimum 105 similarity 0.95) to obtained the scaffold coverage. The SIP metagenomes were merged into one mapping. 137, 112, and 376 106 million reads could be mapped to the two native soil metagenomes and the SIP metagenome, respectively (64-66% of quality 107 filtered reads). Gene prediction of the complete assembly was performed using prodigal (Hyatt et al., 2010). In addition to the 108 detection and taxonomic classification of 105 essential marker genes (Albertsen et al., 2013), dsrA and dsrB homologs were 109 identified using TIGRFAM's hidden Markov model (HMM) profiles TIGR02064 and TIGR02066, respectively, with 110 HMMER 3.1 (Eddy, 2011) and the provided trusted cut-offs. Additional dsrAB-containing scaffolds were identified by using 111 tblastn with the published DsrAB database as a query against the assembly (Müller et al., 2015). DsrAB sequences were

- 112 classified by phylogenetic analysis (Supplementary Methods; Müller et al., 2015). Binning and decontamination was finalized
- 113 utilizing the G+C content and tetramer frequencies of the scaffolds, as well as paired-end information, as described and
- 114 recommended in Albertsen et al. (2013). Completeness, contamination, and strain heterogeneity was estimated using CheckM
- 115 1.0.6 (Parks *et al.*, 2015) with lineage-specific marker sets selected at phylum rank (or class rank for *Proteobacteria*). MAGs
- 116 were taxonomically classified by phylogenomic analysis of concatenated marker sequences and calculation of average nucleic
- 117 and amino acid identities (ANI, AAI, Supplementary Methods). MAGs were annotated using the MicroScope annotation
- 118 platform (Vallenet *et al.*, 2017) and eggNOG (Huerta-Cepas *et al.*, 2016). Genes of interest (Supplementary Table S2) were
- 119 manually curated using the full range of tools integrated in MicroScope annotation platform (Supplementary Methods).

120 Genome-centric activity analysis: iRep and metatranscriptomics

121 The index of replication (iRep) was calculated for each MAG with the combined native soil metagenomes. Settings and

- 122 thresholds were applied as recommended (Brown *et al.*, 2016) using bowtie2 (Langmead and Salzberg, 2012) and the iRep
- 123 script with default settings. Quality-filtered metatranscriptome reads were mapped to all genomes using bowtie2 and counted
- 124 with featureCounts (Liao et al., 2014). To determine gene expression changes, we applied the DESeq2 pipeline with
- 125 recommended settings (Love *et al.*, 2014) (Supplementary Methods).

126 Data availability

127 Metagenomic and -transcriptomic data were deposited under the BioProject accession numbers PRJNA412436 and

- 128 PRJNA412438, respectively, and can also be obtained via the JGI's genome portal (JGI Proposal ID 605). MAGs are available
- 129 at MicroScope (https://www.genoscope.cns.fr/agc/microscope/) and were deposited at the European Nucleotide Archive
- 130 (Supplementary Table S1). DsrAB sequences were deposited at NCBI GenBank under the accession numbers MG182080-
- 131 MG182141.
- 132

134 **Results**

135 Functional metagenomics: Recovery of dsrAB-containing genomes from native soil

136 and ¹³C-DNA fraction metagenomes

137 This study was conducted with soil samples from the Schlöppnerbrunnen II peatland in Germany, which is a long-term study 138 site with active sulfur cycling and harbors a large diversity of unknown microorganisms with divergent dsrAB genes (Steger et 139 al., 2011; Pester et al., 2012). We initially generated co-assembled metagenomes from native peat soil DNA (53 Gb) and a 140 pool of DNA extracts from the heavy fractions of a previous DNA-stable isotope probing (DNA-SIP) experiment with soil 141 from the same peat (101 Gb). The heavy fractions, which were obtained from anoxic peat incubations with periodically 142 supplemented sulfate and a mixture of ¹³C-labelled formate, acetate, propionate, and lactate at low concentrations, were 143 enriched in DNA from Desulfosporosinus and also harbored DNA from yet unidentified dsrAB-containing microorganisms 144 (Pester et al., 2010). Based on the metagenome data, the native peat was dominated by Acidobacteria (61%), but also had 145 Actinobacteria, Alphaproteobacteria, and Deltaproteobacteria as abundant (>5%) phyla/classes (Figure 1). Dominance of 146 Acidobacteria, Alpha- and Deltaproteobacteria is typical for peatlands (Dedysh, 2011). Quantitative PCR confirmed that 147 Acidobacteria subdivisions 1, 2, and 3 persistently dominated the Schlöppnerbrunnen II peat microbiota in oxic and anoxic soil 148 layers (Supplementary Methods, Figure 1), as observed in other peatlands (Serkebaeva et al., 2013; Urbanová and Bárta, 2014; 149 Ivanova et al., 2016).

150 We identified 36 complete or partial dsrAB genes on scaffolds of the co-assembled metagenome and subsequently recovered 151 thirteen MAGs of DsrAB-encoding bacteria by differential coverage binning (Supplementary Table S1, Albertsen et al., 2013). 152 Twenty-eight dsrAB sequences were part of the reductive bacterial-type DsrAB family branch and were closely related to 153 previously recovered sequences from this and other wetlands (Supplementary Figure S1). These dsrAB sequences were 154 affiliated to the known SRM genera Desulfosporosinus (Firmicutes, n=1, one MAG) and Syntrophobacter 155 (Deltaproteobacteria, n=3, two MAGs), the Desulfobacca acetoxidans lineage (n=1), and the uncultured DsrAB family-level 156 lineages 8 (n=19, seven MAGs) and 10 (n=4). Six sequences grouped with the oxidative bacterial-type DsrAB family and were 157 distantly affiliated with Sideroxydans lithotrophicus (Betaproteobacteria, n=5, two MAGs) or Rhodomicrobium vannielii 158 (Alphaproteobacteria, n=1) (Supplementary Figure S2). Interestingly, two of our sequences (n=2, one MAG) and a DsrAB 159 sequence from the candidate phylum Rokubacteria (Hug et al., 2016) formed a completely novel basal lineage outside the four 160 previously recognized DsrAB enzyme families (Supplementary Figure S2) (Müller et al., 2015). The thirteen partial to near 161 complete dsrAB-containing MAGs had moderate to no detectable contamination as assessed by CheckM and manual curation 162 (Supplementary Table S1) (Parks et al., 2015) and derived from Acidobacteria subdivisions 1 and 3 (SbA1-7), 163 Desulfosporosinus (SbF1), Syntrophobacter (SbD1, SbD2), Betaproteobacteria (SbB1, SbB2), and Verrucomicrobia (SbV1), 164 as inferred by phylogenetic analysis of DsrAB sequences (Supplementary Figures S1 and S2) and concatenated sequences of 165 single-copy, phylogenetic marker genes (Supplementary Figure S3). Only the Desulfosporosinus and Syntrophobacter MAGs

- 166 contained rRNA gene sequences.
- 167 Phylogenomic analysis showed that Acidobacteria MAGs SbA1, SbA5, and SbA7 are affiliated with subdivision 1, while
- 168 SbA3, SbA4, and SbA6 are affiliated with subdivision 3 (Supplementary Figure S3). The partial MAG SbA2 lacked the marker
- 169 genes used for phylogenomic treeing, but was unambiguously assigned to Acidobacteria using an extended marker gene set
- 170 (Albertsen et al., 2013) and DsrAB phylogeny. The two near complete (96%) MAGs SbA1 and SbA5 have a size of 5.4 and
- 171 5.3 Mb, respectively. The G+C content of all acidobacterial MAGs ranges from 58% to 63% (Supplementary Table S1). This
- 172 in accordance with genome characteristics of acidobacterial isolates, which have genome sizes of 4.1-10.0 Mb and G+C

- 173 contents of 57-62% (Ward et al., 2009; Rawat et al., 2012,). SbA1 and SbA7 form a monophyletic clade in the Acidobacteria
- 174 subdivision 1 with an AAI (Rodriguez-R and Konstantinidis, 2014) of 63% (Supplementary Figure S3) and DsrAB identity of

175 80% as was calculated with T-Coffee 11 (Notredame et al., 2000) using the unfiltered reference alignment without the

176 intergenic region (Müller et al., 2015). They have 56% AAI to their closest relative, Ca. Koribacter versatilis, which is lower

- 177 than AAIs among members of known acidobacterial genera (60–71%). The third MAG from subdivision 1, SbA5, is affiliated
- 178 with *Terracidiphilus gabretensis* with an AAI of 61%. DsrAB identity of SbA5 to SbA1 and SbA7 is 79%. The three

179 subdivision 3 MAGs form a monophyletic clade with Ca. Solibacter usitatus (Supplementary Figure S3). SbA3, SbA4, and

- 180 SbA6 have AAIs of 59–73% amongst them and 61–62% to *Ca. S. usitatus*. DsrAB identity amongst the three MAGs is 80–
- $181 \quad 94\%$ and 74–79% to the subdivisions 1 MAGs.

182 The DsrAB sequences encoded on all seven MAGs belong to the uncultured DsrAB family-level lineage 8 (Supplementary

183 Figure S1), which so far only consisted of environmental *dsrAB* sequences of unknown taxonomic identity (Müller *et al.*,

184 2015). Based on these MAGs and metatranscriptome analyses of anoxic peat soil microcosms, we describe the putative

185 metabolic capabilities of these novel DsrAB-encoding *Acidobacteria*. Details on the other MAGs will be described elsewhere

- 186 (Hausmann et al., unpublished; Anantharaman et al., unpublished). Functional interpretations of the recovered MAGs are
- 187 made under the premise that the genomes are not closed, and thus it is unknown if genes are absent in these organisms or are
- 188 missing due to incomplete sequencing, assembly, or binning.

189 **Dissimilatory sulfur metabolism**

190 Although Acidobacteria are abundant in diverse environments with active sulfur cycling (Serkebaeva et al., 2013; Urbanová 191 and Bárta, 2014; Sánchez-Andrea et al., 2011; Wang et al., 2012), this is the first discovery of members of this phylum with a 192 putative dissimilatory sulfur metabolism. SbA2, SbA3, and SbA7 encode the complete canonical pathway for dissimilatory 193 sulfate reduction, including homologs for sulfate transport (sulP and/or dass, not in SbA7) and activation (sat, ppa, hppA), 194 adenosine 5'-phosphosulfate (APS) reduction (aprBA, qmoABC), and sulfite reduction (dsrAB, dsrC, dsrMKJOP) (Figure 2, 195 Supplementary Table S2a) (Santos et al., 2015). In the AprBA tree, the acidobacterial sequences are part of a large cluster of 196 yet uncultured organisms and Deltaproteobacteria and Firmicutes that respire sulfate, sulfite, or thiosulfate (Supplementary 197 Figure S4) (Watanabe et al., 2016). SbA1, SbA4, SbA5, and SBA6 have an incomplete sulfate reduction gene set but contain 198 all dsr genes for sulfite reduction. Several other dsr genes were present on some of the MAGs. The dsrD and dsrN genes 199 occurred in pairs. The acidobacterial DsrD sequences have the same conserved, hydrophobic residues as Desulfovibrio vulgaris 200 DsrD (Supplementary Figure S5) (Mizuno et al., 2003). Ubiquity of DsrD among SRM suggests an essential function in sulfate 201 reduction, but the physiological role of this small protein is unresolved (Hittel and Voordouw, 2000). DsrN is a homolog of 202 cobyrinate a,c-diamide synthase in cobalamin biosynthesis and may be involved in amidation of the siroheme prosthetic group 203 of DsrAB (Lübbe et al., 2006). DsrV, a homolog of precorrin-2 dehydrogenase, and DsrWa, a homolog of uroporphyrin-III C-204 methyltransferase, may also be involved in siroheme biosynthesis (Holkenbrink et al., 2011). DsrT is required for sulfide 205 oxidation in Chlorobaculum tepidum, but also found in SRM (Holkenbrink et al., 2011). The presence of dsrMK-paralogs 206 (dsrM2, dsrK2) upstream of dsrAB is not uncommon in SRM (Pereira et al., 2011). DsrMK are present in all dsrAB-containing 207 microorganisms and are a transmembrane module involved in reduction of cytoplasmic DsrC-trisulfide in SRM, the final step 208 in sulfate reduction (Santos et al., 2015). DsrC encoded on the MAGs have the two essential cysteine residues at the C-209 terminal end for full functionality (Venceslau et al., 2014). Interestingly, dsrC forms a gene duo with dsrL downstream of 210 dsrAB in all seven MAGs. This is surprising, because dsrL is not found in SRM but in sulfur oxidizers. DsrL is highly 211 expressed and essential for sulfur oxidation by the purple sulfur bacterium Allochromatium vinosum (Lübbe et al., 2006;

- 212 Weissgerber et al., 2014). DsrL is a cytoplasmic iron-sulfur flavoprotein with proposed NAD(P)H:acceptor oxidoreductase 213 activity and was copurified with DsrAB from the soluble fraction of A. vinosum (Dahl et al., 2005). The acidobacterial DsrL 214 sequences are shorter than their homolog in A. vinosum (Supplementary Table S2a), but have the same functional domains 215 (Supplementary Figure S6). Given the possible role of DsrL in sulfur oxidation, we sought to detect additional genes indicative 216 of oxidative sulfur metabolism in the acidobacterial MAGs. However, genes for Sox enzyme machinery (soxABXYZ), 217 thiosulfate dehydrogenase (tsdA), sulfide:quinone reductase (sqr), adenylyl-sulfate reductase membrane anchor subunit 218 (aprM), flavocytochrome c sulfide dehydrogenase (fccAB), sulfur reductase (sreABC), thiosulfate reductase (phsABC), 219 polysulfide reductase (psrABC), membrane-bound sulfite oxidizing enzyme (soeABC), cytoplasmic sulfur trafficking enzymes 220 (tusA, dsrE2, dsrEFH), or DsrQ/DsrU (unknown functions) were absent (Laska et al., 2003; Holkenbrink et al., 2011; Lenk et 221 al., 2012; Wasmund et al., 2017). SbA1, SbA3, SbA4, and SbA6 contain genes that have only low homology to soxCD/sorAB, 222 periplasmic sulfite-oxidizing enzymes (Supplementary Results) and, thus, might have another function (Ghosh and Dam, 223 2009).
- 224 Despite ongoing sulfur cycling, concentrations of inorganic sulfur compounds such as sulfate are low (lower µM range) in the 225 Schlöppnerbrunnen II peatland (Schmalenberger et al., 2007; Küsel et al., 2008; Knorr and Blodau, 2009). Enzymatic release 226 of inorganic sulfur compounds from organic matter might thus represent a significant resource for sulfur-dissimilating 227 microorganisms. Therefore, we specifically searched for genes coding for known organosulfur reactions that yield sulfite 228 (Wasmund et al., 2017). Genes for cysteate sulfo-lyase (cuyA), methanesulfonate monooxygenase (msmABCD), 229 sulfoacetaldehyde acetyltransferase (xsc), and taurine dioxygenase (tauD) were absent. However, suyAB, coding for the (R)-230 sulfolactate sulfo-lyase complex that cleaves (R)-sulfolactate into pyruvate and sulfite (Denger and Cook, 2010), were present 231 in SbA4 and SbA5 (Supplementary Table S2a). Intriguingly, SbA4 and SbA5 only have capability for sulfite reduction. SbA5 232 also encodes the racemase machinery for (S)-sulfolactate to (R)-sulfolactate, (S)-sulfolactate dehydrogenase (slcC) and (R)-233 sulfolactate dehydrogenase (comC); the regulator gene suvR or the putative importer SlcHFG were absent (Denger and Cook, 234 2010). Pyruvate may be used as an energy and carbon source, while sulfite could be used as an electron acceptor for anaerobic
- respiration (Simon and Kroneck, 2013).

236 **Respiration**

237 Cultivated Acidobacteria of subdivisions 1 and 3 are strict aerobes or facultative anaerobes (e.g., Eichorst et al., 2007; 238 Kulichevskaya et al., 2010, 2014; Pankratov and Dedysh, 2010; Dedysh et al., 2012; Pankratov et al., 2012). Accordingly, we 239 found respiratory chains encoded in all acidobacterial MAGs (Figure 3, Supplementary Results), with (near) complete operons 240 for NADH dehydrogenase 1, succinate dehydrogenase (lacking in SbA2), one or both types of quinol-cytochrome-c 241 reductase, low-affinity terminal oxidases, and ATP synthase (lacking in SbA2) (Supplementary Tables S2b-h). High-affinity 242 terminal oxidases, putatively involved in detoxification of oxygen (Ramel et al., 2013; Giuffrè et al., 2014), are limited to four 243 MAGs (Supplementary Table S2g). Genes for dissimilatory nitrogen or iron metabolisms are absent, with the exception of a 244 putative metal reductase in SbA2 of unclear physiological role (Supplementary Results).

245 Hydrogen utilization and production

246 We identified [NiFe] hydrogenases of groups 1, 3, and 4 (Greening et al., 2016) in SbA1-7 (Supplementary Table S2j).

- 247 Membrane-bound group 1 hydrogenases (SbA1, SbA3, SbA5) consume hydrogen from the periplasm as an electron donor to
- generate energy, possibly coupled to sulfate/sulfite reduction. In contrast to other Acidobacteria, no group 1h/5 hydrogenases,
- 249 which are coupled to oxygen respiration, were identified (Greening et al., 2015). Cytoplasmic group 3 hydrogenases (all

- 250 MAGs) are bidirectional and proposed to be involved in energy-generating hydrogen oxidation and/or fermentative hydrogen
- production. Membrane-bound group 4 hydrogenases (SbA1, SbA5, SbA4, SbA6) produce H and are postulated to conserve
- energy by proton translocation by oxidizing substrates like formate (group 4a) or carbon monoxide (via ferredoxin, group 4c)
- 253 (Figure 3).

254 A versatile heterotrophic physiology

255 Acidobacteria are known for their capability to degrade simple and polymeric carbohydrates (Kulichevskaya et al., 2010, 256 2014; Pankratov and Dedysh, 2010; Dedysh et al., 2012; Eichorst et al., 2011; Pankratov et al., 2012; Rawat et al., 2012; 257 Huber et al., 2016), supported by many diverse carbohydrate-active enzymes encoded on their genomes (Ward et al., 2009; 258 Rawat et al., 2012). Accordingly, the MAGs recovered in our study also contain many genes encoding diverse carbohydrate-259 active enzymes (Supplementary Methods, Figure 4). These include glycoside hydrolases (GH, 1.0-4.0% of all genes), 260 polysaccharide lyases (0.07-0.3%), and carbohydrate esterases (0.7-1.4%) that are generally involved in degradation of 261 complex sugars, but also glycosyltransferases (0.9–1.4%) for biosynthesis of carbohydrates. Functional GH families (assigned 262 by EC number) putatively involved in cellulose and hemicellulose degradation were most prevalent (Supplementary Table S4). 263 Specifically, the most often encountered EC numbers encompassed by the different GH families represented cellulose (EC 264 3.2.1.4, e.g., GH5, GH74), xyloglucan (EC 3.2.1.150, EC 3.2.1.151, e.g., GH5, GH74), or xylan (EC 3.2.1.8, EC 3.2.1.37, e.g., 265 GH5) degradation, which is similar to the situation found in other members of Acidobacteria subdivision 1 and 3 (Ward et al., 266 2009; Rawat et al., 2012). Further EC numbers that were often encountered in the various detected GH families were 267 associated with oligosaccharide degradation (EC 3.2.1.21, e.g., GH2) or α -N-acetylgalactosaminidase activity (EC 3.2.1.49, 268 e.g., GH109). Degradation of cellulose and hemicellulose yields glucose and all MAGs encode glycolysis and pentose 269 phosphate pathways (Figure 3, Supplementary Results). α-N-acetylgalactosaminidase releases N-acetylgalactosamine residues 270 from glycoproteins that are commonly found in microbial cell walls and extracellular polysaccharides (Bodé et al., 2013). N-271 acetylgalactosamine can not be directly utilized via glycolysis, however the additionally required enzymes are present (Figure 272 3; Supplementary Results). Under oxic conditions, organic carbon could be completely oxidized to CO via the citric acid 273 cycle (Figure 3). Alternatively, we also identified fermentative pathways. SbA3 encodes the bifunctional aldehyde-alcohol 274 dehydrogenase AdhE that yields ethanol (Figure 3). All MAGs encode additional aldehyde and alcohol dehydrogenases 275 without clear substrate specificity that could also ferment acetyl-CoA to ethanol. SbA7 and SbA5 encode a L-lactate 276 dehydrogenase (Ldh) yielding lactate from pyruvate, while six MAGs encode L-lactate dehydrogenases (LldD, GlcDEF, 277 LutABC) that presumably perform the reverse reaction (Figure 3). Similarly, we identified pathways for acetate and/or 278 propionate production or utilization in all MAGs (Figure 3; Supplementary Results). SbA1 and SbA3 potentially produce H 279 via formate C-acetyltransferase PfID, which cleaves pyruvate into acetyl-CoA and formate. SbA1 encodes for the membrane-280 bound formate hydrogenlyase complex (fdhF, hyf operon) that produces H and might also translocate protons. SbA3 281 harbours an uncharacterized, cytoplasmic, monomeric FDH (*fdhA*) to transform formate to H . SbA1, SbA3, SbA4, and SbA6 282 also encode membrane-bound, periplasmic FDH (fdo operon) that transfers electrons into the membrane quinol pool, as a non-283 fermentative alternative of formate oxidation (Figure 3, Supplementary Table S2i).

284 **DsrAB-encoding** *Acidobacteria* are metabolically active under anoxic conditions

We calculated the index of replication (iRep) based on the native peat soil metagenomes to assess whether DsrAB-encoding *Acidobacteria* were active *in situ* (Brown *et al.*, 2016). SbA1 and SbA5, which were sufficiently complete (\geq 75%) for a reliable estimate, had iRep values of 1.21 and 1.19, respectively. This shows that a fraction of each population was metabolically active, i.e., on average 21% of SbA1 and 19% of SbA5 cells were actively replicating at the time of sampling.

- 289 Concordantly, SbA1–7 were also transcriptionally active in the same native soil samples. 35–46% of the SbA1–7 genes were
- 290 expressed in at least one replicate. SbA1 and SbA5 contributed a considerable fraction (0.4% and 1.8%, respectively,

291 Supplementary Table S1) of the total mRNA reads in the native soil metatranscriptome. These data likely underestimate the

292 metabolic activity of SbA1-7 in situ because freshly sampled soil was stored at 4 °C for one week prior to nucleic acids

extraction.

294 We further analyzed metatranscriptome data from a series of anoxic incubations of the peat soil with or without individual 295 substrates (formate, acetate, propionate, lactate, or butyrate) and with or without supplemental sulfate (Hausmann et al., 2016). 296 While the incubations were not designed to specifically test for the MAG-inferred metabolic properties, they still allowed us to 297 evaluate transcriptional response of the DsrAB-encoding Acidobacteria under various anoxic conditions (Supplementary 298 Methods and Results). All treatments triggered shifts in genome-wide gene expression; more genes were significantly (p < 0.05) 299 upregulated (73–933) than downregulated (14–81) as compared to the native soil. Upregulated genes included sulfur 300 metabolism enzymes, high-affinity terminal oxidases, group 1 and 3 hydrogenases, aldehyde-alcohol dehydrogenase AdhE, 301 glycoside hydrolases, and other carbon metabolism enzymes (Supplementary Table S3, Supplementary Figure S7). 302 Significantly upregulated glycoside hydrolase genes belonged to GH family 2, 3, 5, 9, 10, 18, 20, 23, 26, 28, 29, 30, 33, 35, 36, 303 38, 43, 44, 50, 51, 55, 74, 76, 78, 79, 88, 95, 97, 105, 106, 109, and 129 in MAGs SbA1-6. None of the GH genes were 304 significantly downregulated in the incubations. Noteworthy genes that were significantly downregulated were superoxide 305 dismutases (sodA) in SbA2 and SbA4 (Supplementary Table S3a).

306

308 **Discussion**

309 Diverse members of the phylum Acidobacteria are abundant in various ecosystems, particularly in soils and sediments with

- 310 relative abundances typically ranging from 20-40% (Janssen, 2006). Acidobacteria are currently classified in 26 subdivisions
- 311 based on their 16S rRNA phylogeny (Barns et al., 2007). Given their phylogenetic breadth, comparably few isolates and
- 312 genomes are available to explore their metabolic capabilities. Currently isolated species of subdivisions 1, 3, 4, and 6 are
- 313 aerobic chemoorganotrophs that grow optimally at neutral or low pH (Dedysh, 2011; Eichorst *et al.*, 2011; Huber *et al.*, 2014,
- 314 2016). Furthermore, subdivision 4 contains an anoxygenic phototroph (Garcia Costas et al., 2012; Tank and Bryant, 2015),
- 315 subdivisions 8 and 23 contain anaerobes (Liesack *et al.*, 1994; Coates *et al.*, 1999; Losey *et al.*, 2013), subdivisions 1, 3, and
- 316 23 fermenters (Pankratov et al., 2012; Kulichevskaya et al., 2014; Losey et al., 2013; Myers and King, 2016) and subdivision
- 217
- 317 4, 8, 10 and 23 thermophiles (Izumi *et al.*, 2012; Losey *et al.*, 2013; Crowe *et al.*, 2014; Tank and Bryant, 2015).
- 318 Acidobacteria are known as dominant inhabitants of wetlands worldwide, in particular members of subdivision 1, 3, 4, and 8
- 319 (Dedysh, 2011). Strains in the genera Granulicella (Pankratov and Dedysh, 2010), Telmatobacter (Pankratov et al., 2012),
- 320 Bryocella (Dedysh et al., 2012) and Bryobacter (Kulichevskaya et al., 2010) have been isolated from acidic wetlands and are
- 321 presumably active in plant-derived polymer degradation (such as cellulose) (Dedysh, 2011; Pankratov et al., 2011; Schmidt et
- 322 al., 2015; Juottonen et al., 2017), and in nitrogen and iron cycling (Küsel et al., 2008; Kulichevskaya et al., 2014).

323 Here, we provide metagenomic and metatranscriptomic evidence that the newly discovered species represent at least three 324 novel genera in Acidobacteria subdivision 1 and 3 (Supplementary Figure S3) and possess a dissimilatory sulfur metabolism. 325 The seven acidobacterial MAGs from the Schlöppnerbrunnen II peatland encode the complete canonical pathway for 326 dissimilatory sulfite or sulfate reduction. The sulfate reduction pathway, however, could also operate in reverse as proposed for 327 a sulfur-oxidizing deltaproteobacterium (Thorup et al., 2017). The phylogenetic separation into two subdivisions as based on 328 the concatenated marker gene tree is also apparent in the DsrAB phylogeny (Supplementary Figure S1). The acidobacterial 329 DsrAB sequences are distributed among two monophyletic clades within the uncultured family-level lineage 8, which is part of 330 the reductive, bacterial-type DsrAB branch (Müller et al., 2015). The phylogenetic breadth of the acidobacterial DsrAB 331 sequences is representative for the intra-lineage sequence divergence within uncultured DsrAB lineage 8, which suggests that 332 this entire lineage represents yet uncultivated bacteria of the phylum Acidobacteria. Members of this uncultured DsrAB 333 lineage are widespread in freshwater wetlands (Supplementary Figure S1) (Pester et al., 2012). In particular, they represented 334 an abundant fraction of the DsrAB diversity and were permanent autochthonous inhabitants of oxic and anoxic soil layers in

the analyzed Schlöppnerbrunnen II peatland (Steger *et al.*, 2011; Pester *et al.*, 2010).

336 Presence of the complete gene set for canonical dissimilatory sulfate reduction suggests that the pathway is functional, as the 337 genetic capability for sulfate reduction can be rapidly lost by adaptive evolution if unused (Hillesland et al., 2014). Except for 338 a truncated aprB on SbA6, we found no indications of pseudogenes, i.e., unexpected internal stop codons or reading frame 339 shifts, for any of the sulfate/sulfite reduction genes on the acidobacterial MAGs (Müller et al., 2015). In addition, sulfur 340 metabolism genes of each MAG were expressed in the native soil and the anoxic microcosms (Supplementary Table S3a). 341 Many sulfur metabolism genes were even significantly upregulated in the anoxic microcosms, with dsrC and aprBA among the 342 top 10 most expressed genes in SbA7 (Supplementary Table S3a). These findings further support full functionality of the 343 acidobacterial dissimilatory sulfur pathways under anoxic condition.

- 344 Known SRM typically couple sulfate respiration to oxidation of fermentation products such as volatile fatty acids, alcohols, or
- 345 hydrogen (Rabus et al., 2013). While other microorganisms in the Schlöppnerbrunnen II soil, such as Desulfosporosinus,

346 showed sulfate- and substrate-specific responses in our microcosms, hundreds of acidobacterial 16S rRNA phylotypes did not 347 (with the exception of two) (Hausmann et al., 2016). Gene expression patterns of DsrAB-encoding Acidobacteria in the 348 individual anoxic microcosms as analyzed in the present study were ambiguous. Genes for putative oxidation of the 349 supplemented substrates (formate, acetate, propionate, lactate, butyrate) were not specifically upregulated, neither without nor 350 with supplemental sulfate. However, sulfur metabolism genes were upregulated in several incubations as compared to no-351 substrate-controls, suggesting indirect stimulation of a sulfur-based metabolism (Supplementary Results, Supplementary Table 352 S3a). Indirect changes in microbial activity after the addition of fresh organic matter is often observed in soils (priming effects, 353 Blagodatskaya and Kuzyakov, 2008). One explanation for this priming effect is the co-metabolism theory stating that easily 354 available substrates provide the energy for microorganisms to produce extracellular enzymes to make immobile carbon 355 accessible, which is then also available to other microorganisms. The DsrAB-encoding Acidobacteria have a large genetic 356 repertoire to utilize carbohydrates and monomeric sugars (Figure 3). This is in accordance with the carbohydrate utilization 357 potential previously described for subdivision 1 and 3 Acidobacteria (Ward et al., 2009; Rawat et al., 2012). Yet utilization of 358 monomeric sugars is a rare feature of known SRM (Cord-Ruwisch et al., 1986; Stetter, 1988) and utilization of poly- or 359 oligosaccharides by sulfate-reducing bacteria was not yet reported. While the studied Acidobacteria expressed many of their 360 glycoside hydrolase genes in our anoxic peat soil microcosms, further experiments are required to confirm if DsrAB-encoding 361 Acidobacteria couple degradation of carbohydrate polymers or monomers to sulfate reduction.

362 It is intriguing to propose that MAGs SbA2, SbA3, and SbA7 derive from acidobacterial SRM as they lack known sulfur 363 oxidation genes, except dsrL (Supplementary Figure S6), and express the complete dissimilatory sulfate reduction pathway 364 (Supplementary Table S2a), including reductive, bacterial-type dsrAB, and dsrD that may be exclusive to SRM (Hittel and 365 Voordouw, 2000; Dahl and Friedrich, 2008; Ghosh and Dam, 2009; Rabus et al., 2015). However, the functions of DsrL and 366 DsrD are yet unresolved, which prevents functional predictions based only on these genes. The proposal of an alternative 367 hypothesis that these novel Acidobacteria reverse the sulfate reduction pathway for dissimilatory sulfur oxidation or sulfur 368 disproportionation, bases on findings with the deltaproteobacterium Desulfurivibrio alkaliphilus (Thorup et al., 2017). D. 369 alkaliphilus also lacks known sulfur oxidation genes (including dsrL), except for sqr, and is proposed to gain energy by 370 coupling sulfide oxidation via a reversed sulfate reduction pathway (with a reductive-type DsrAB) to the dissimilatory 371 reduction of nitrate/nitrite to ammonium. Sulfide oxidation in acidobacterial MAGs SbA2, SbA3, and SbA7 could proceed 372 analogous to the pathway models proposed by Thorup et al. (2017) and Christiane Dahl (Dahl, 2017). Briefly, hydrogen 373 sulfide might react with DsrC either spontaneously (Ijssennagger et al., 2015) or via an unknown sulfur transfer mechanism to 374 form persulfated DsrC. Persulfated DsrC is then oxidized by DsrMKJOP, thereby transferring electrons into the membrane 375 quinone pool, and releasing a DsrC-trisulfide, which is the substrate for DsrAB (Santos et al., 2015; Dahl, 2017). It was 376 hypothesized that electrons released during DsrC-trisulfide oxidation to sulfite and DsrC are transferred to DsrL (Dahl, 2017). 377 Further sulfite oxidation to sulfate would be catalyzed by AprBA-QmoABC and Sat.

The acidobacterial MAGs have the genomic potential to use oxygen as terminal electron acceptor and might thus couple sulfide oxidation to aerobic respiration. Alternative electron acceptors for biological sulfur oxidation in wetlands could include nitrate/nitrite and metals such as Fe(III) (Küsel *et al.*, 2008). However, known genes for dissimilatory nitrate reduction and metal reduction (Weber *et al.*, 2006) were absent from these acidobacterial MAGs. Only SbA2 encodes a putative metal reduction complex that was recently characterized in *Desulfotomaculum reducens* (Otwell *et al.*, 2015). At this time, it is unclear whether DsrAB-encoding *Acidobacteria* are capable of Fe(III) respiration, as seen in *Geothrix fermentans* (Coates *et al.*, 1999) and certain isolates in subdivision 1 (Blöthe *et al.*, 2008; Kulichevskaya *et al.*, 2014).

387 Proposal of the acidobacterial *Candidatus* genera 388 Sulfotelmatobacter, Sulfotelmatomonas, and Sulfopaludibacter

389 Based on combined interpretation of phylogeny (concatenated phylogenetic marker genes, DsrAB), genomic (ANI, AAI) and

390 genetic (DsrAB) distances, and characteristic genomic features of dissimilatory sulfur metabolism (Figure 3), in accordance

391 with Konstantinidis et al. (2017), we classify MAGs SbA1, SbA7, SbA5, SbA3, SbA4, and SbA6 into three new acidobacterial

392 Candidatus genera, including Candidatus species names for the >95% complete MAGs SbA1 and SbA5. In-depth

393 phylogenomic analysis of SbA2 was not possible and therefore it is tentatively assigned to Acidobacteria subdivision 3.

394 Acidobacteria subdivision 1

- Ca. genus Sulfotelmatobacter (Sul.fo.tel.ma.to.bac'ter. L. n. sulfur, sulfur, Sulfur; Gr. n. telma, -tos, swamp, wetland; N.L.
 masc. n. bacter, bacterium; N.L. masc. n. Sulfotelmatobacter, a bacterium from a swamp metabolizing sulfur) with
 Ca. Sulfotelmatobacter kueseliae MAG SbA1 (kue.se'li.ae. N.L. gen. n. kueseliae, of Kuesel, honouring Kirsten
 Küsel, for her work on the geomicrobiology of wetlands) and Ca. Sulfotelmatobacter sp. MAG SbA7.
- *Ca.* Sulfotelmatomonas gaucii MAG SbA5 (Sul.fo.tel.ma.to.mo.nas. L. n. *sulfur*, sulfur; Gr. n. *telma*, *-tos*, swamp,
 wetland; N.L. fem. n. *monas*, a unicellular organism; N.L. fem. n. *Sulfotelmatomonas*, a bacterium from a swamp
 metabolizing sulfur; gau'.ci.i. N.L. gen. n. *gaucii*, of Gauci, in honour of Vincent Gauci, for his pioneering work on
 the interplay of wetland sulfate reduction and global methane emission).

403 Acidobacteria subdivision 3

- Ca. genus Sulfopaludibacter (Sul.fo.pa.lu.di.bac'ter. L. n. sulfur, sulfur; L. n. palus, -udis, L. swamp; N.L. masc. n.
 bacter, bacterium; N.L. masc. n. Sulfopaludibacter, a bacterium from a swamp metabolizing sulfur) with Ca.
 Sulfopaludibacter sp. MAG SbA3, Ca. Sulfopaludibacter sp. MAG SbA4, and Ca. Sulfopaludibacter sp. MAG SbA6.
- 407 Acidobacteria bacterium sp. MAG SbA2.

408

409

410

Conclusion 412

413 Sulfur cycling exerts important control on organic carbon degradation and greenhouse gas production in wetlands, but

414 knowledge about sulfur microorganisms in these globally important ecosystems is scarce (Pester et al., 2012). Here, we show

415 by genome-centric metagenomics and metatranscriptomics that members of the phylum Acidobacteria have a putative role in

416 peatland sulfur cycling. The genomic repertoire of these novel Acidobacteria species encompassed recognized acidobacterial

417 physiologies, such as a facultative anaerobic metabolism, oxygen respiration, fermentation, carbohydrate degradation, and 418

hydrogen metabolism, but was additionally augmented with a DsrAB-based dissimilatory sulfur metabolism (Figure 5). Based 419

on their genetic repertoire and previous findings on reversibility of the dissimilatory sulfate reduction pathway (Dannenberg et 420

al., 1992; Fuseler and Cypionka, 1995; Fuseler et al., 1996; Thorup et al., 2017), it is intriguing to speculate that the described

421 peatland Acidobacteria could use the same pathway for both sulfate reduction and sulfide oxidation. The described DsrAB-422

carrying Acidobacteria that only encoded the pathway for dissimilatory sulfite reduction had additional genes for sulfite-423 producing enzymes, which suggests that organosulfonates might be their primary substrate for sulfur respiration. Our results

424 not only extend the current understanding of the genetic versatility and distribution of dissimilatory sulfur metabolism among

425 recognized microbial phyla, but also underpin the challenge to unambiguously differentiate between reductive or oxidative

426 sulfur metabolism solely based on (meta-)genome/transcriptome data (Thorup et al., 2017).

427

429 **Conflict of Interest**

430 The authors declare no conflict of interest.

431

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660 Figure legends

661 **Figure 1**

- 662 Microbial community composition in Schlöppnerbrunnen II peatland in samples from different years and soil depths. (a)
- 663 Abundance of phyla and proteobacterial classes in the native soil (relative to all classified reads/amplicons). Taxa less
- abundant than 1% are grouped in grey. Coverage abundance is based on metagenomic reads mapped to classified scaffolds.
- 665 Amplicon abundance is based on rrn operon-copy number-corrected abundance of 16S rRNA gene operational taxonomic
- 666 units (Hausmann et al., 2016). (b) Relative abundance of acidobacterial subdivisions (SD) in the native soil samples as
- 667 determined by 16S rRNA gene qPCR assays. In addition, all subdivisions more abundant than 1% in a 16S rRNA gene
- 668 amplicon dataset are shown (Hausmann *et al.*, 2016).

669 **Figure 2**

- 670 Organization of dissimilatory sulfur metabolism genes on acidobacterial MAGs SbA1-7. Red: sat; orange: aprBA, qmoABC;
- 671 green: dsrABCMKJOPM2K2; blue: dsrD; turquoise: dsrL; violet: dsrNVWa; pink: suyAB, comC, slcC; 1-4 (grey): syntenic
- 672 genes encoding for conserved proteins of unknown function; white: genes of unknown function or not involved in sulfur
- 673 metabolism. In SbA2 all genes are on one scaffold (scaffold 0lkb). Gene fragments at contig borders are indicated by an
- 674 asterisk. *aprB* in SbA6, indicated by two asterisks, is truncated, which indicates a pseudogene or is due to an assembly error.
- 675 Scaffolds are separated by two slashes.

676 **Figure 3**

- 677 Metabolic model as inferred from analysis of acidobacterial MAGs SbA1-7. Sulfur metabolism is highlighted in yellow.
- 678 Enzymes and transporters are shown in blue font. Glycoside hydrolases are shown in pink font (Supplementary Table S2).
- 679 Extracellular compounds are in parentheses. A slash (/) indicates isozymes, i.e., enzymes that perform the same function, but
- 680 are distinctly different or have more than one established name. AcdA+B, MaeB+Pta, MeaB+Mce, Tal+Pgi: bifunctional
- 681 fusion genes/proteins. Otherwise the plus sign (+) indicates protein complexes. TCA: tricarboxylic acid cycle. FDH: formate
- 682 dehydrogenase. Hase: hydrogenase. NDH: NADH dehydrogenase. HCO: haem-copper oxidase. TO: terminal oxidase. KDG:
- 683 2-dehydro-3-deoxy-D-gluconate. KDGP: 2-dehydro-3-deoxy-D-gluconate 6-phosphate. Expression of at least one copy of
- 684 every enzyme and transporter was observed in the incubation samples.

685 Figure 4

- 686 Glycoside hydrolase genes are enriched in acidobacterial genomes/MAGs compared to genomes from other taxa that encode
- 687 DsrA/DsrB. DsrAB-containing MAGs SbA1-7 are shown as solid symbols and numbered accordingly. X-axis shows the total
- 688 number of predicted CDS per genome/MAG.

689 **Figure 5**

690 Putative lifestyles of DsrAB-encoding Acidobacteria.

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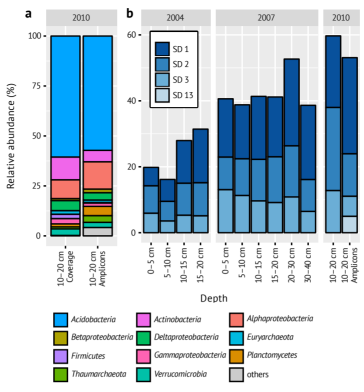


Fig. 2.

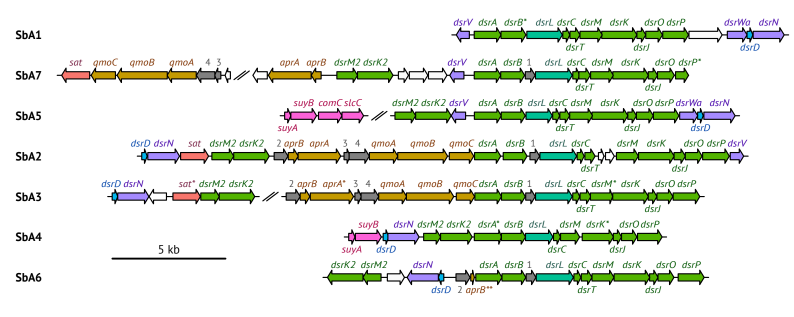


Fig. 3.

