

1 RUNNING TITLE: Hyporheic zone hotspot mechanisms

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3 Multi ‘omics comparison reveals metabolome biochemistry, not microbiome composition or
4 gene expression, corresponds to elevated biogeochemical function in the hyporheic zone

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20

21 **Abstract.**

22 Biogeochemical hotspots are pervasive at terrestrial-aquatic interfaces, particularly within
23 groundwater-surface water mixing zones (hyporheic zones), and they are critical to
24 understanding spatiotemporal variation in biogeochemical cycling. Here, we use multi 'omic
25 comparisons of hotspots to low-activity sediments to gain mechanistic insight into hyporheic
26 zone organic matter processing. We hypothesized that microbiome structure and function, as
27 described by metagenomics and metaproteomics, would distinguish hotspots from low-activity
28 sediments through a shift towards carbohydrate-utilizing metabolic pathways and elucidate
29 discrete mechanisms governing organic matter processing in each location. We also expected
30 these differences to be reflected in the metabolome, whereby hotspot carbon (C) pools and
31 metabolite transformations therein would be enriched in sugar-associated compounds. In contrast
32 to expectations, we found pronounced phenotypic plasticity in the hyporheic zone microbiome
33 that was denoted by similar microbiome structure, functional potential, and expression across
34 sediments with dissimilar metabolic rates. Instead, diverse nitrogenous metabolites and
35 biochemical transformations characterized hotspots. Metabolomes also corresponded more
36 strongly to aerobic metabolism than bulk C content only (explaining 67% vs. 42% of variation),
37 and bulk C did not improve statistical models based on metabolome composition alone. These
38 results point to organic nitrogen as a significant regulatory factor influencing hyporheic zone
39 organic matter processing. Based on our findings, we propose incorporating knowledge of
40 metabolic pathways associated with different chemical fractions of C pools into ecosystem
41 models will enhance prediction accuracy.

42 **1. Introduction.**

43 Soils and nearshore sediments contain a vast reservoir of stored carbon. Uncertainty in
44 the fate of these stores is central to constraining future atmospheric CO₂ concentrations (Burd et
45 al., 2016; Luo et al., 2016; Todd-Brown et al., 2013). Advances in molecular technology has
46 given researchers new ability to characterize mechanisms governing carbon (C) bioavailability,
47 and ultimately the conversion of belowground C pools to CO₂. These advances are central to
48 ongoing efforts to improve process-based ecosystem models by incorporating microbial and
49 biochemical complexity in biogeochemical processes (Buchkowski et al., 2017; Luo et al., 2016;
50 Wieder et al., 2013; Wieder et al., 2017). Still, there is little consensus on the roles of diverse
51 environmental and microbial factors in enhancing belowground CO₂ flux predictions (Graham et
52 al., 2016b; Graham et al., 2014; Luo et al., 2016; Rocca et al., 2015; Wieder et al., 2017), and
53 understanding the mechanisms regulating biogeochemistry in natural environments is paramount
54 to conceptualizing the structure and parameterization of models (Luo et al., 2016; Todd-Brown
55 et al., 2013; Wieder et al., 2013).

56 Recent research has vastly improved both molecular methodologies (Aebersold and
57 Mann, 2003; Gabor et al., 2014; Tfaily et al., 2017; Tringe and Rubin, 2005; Viant, 2008; Wang
58 et al., 2009) and biogeochemical mechanisms represented in ecosystem models (Allison et al.,
59 2010; Wieder et al., 2015a; Wieder et al., 2013; Xu et al., 2014). For instance, parallel advances
60 in environmental chemistry and microbiology now allow for detailed characterization of
61 belowground C pools (e.g., fluorescent methods (Gabor et al., 2014), high-resolution mass
62 spectroscopy (Tfaily et al., 2017), and nuclear magnetic resonance (NMR) (Markley et al.,
63 2017)) as well as the structure and function of environmental microbiomes (Aebersold and
64 Mann, 2003; Tringe and Rubin, 2005; Wang et al., 2009). Newly improved process models

65 account for chemical attributes of C (Liski et al., 2005; Wang et al., 2017), microbial biomass
66 and physiology (Wieder et al., 2014; Wieder et al., 2015a), and nutrient limitations (Wieder et
67 al., 2015b) in their representations of biogeochemistry, among other factors (Verheijen et al.,
68 2015; Wang et al., 2017). Despite these advances, cross-system experimental evidence has
69 revealed many inconsistencies in the importance of different biogeochemical and microbial
70 properties explaining respiration rates (Graham et al., 2016b; Graham et al., 2017b; Hartman et
71 al., 2017), suggesting a need for greater spatially-explicit understanding of mechanisms involved
72 in organic matter processing (Graham et al., 2017b; Krause et al., 2017).

73 Terrestrial-aquatic interfaces are recognized active regions of C metabolism, and areas of
74 subsurface groundwater-surface water mixing (hyporheic zones) in particular are critical in
75 determining the fate of organic matter globally (Battin et al., 2009; Boulton et al., 1998; Cole et
76 al., 2007; Marín-Spiotta et al., 2014; Regnier et al., 2013). Hyporheic zones contribute
77 disproportionately to stream and river metabolism (Gomez-Velez et al., 2015; Huizenga et al.,
78 2017; Stegen et al., 2016; Stern et al., 2017), with up to 96% of ecosystem metabolism occurring
79 in this zone in some headwater systems (Naegeli and Uehlinger, 1997). Hotspots and hot
80 moments of enhanced biogeochemical activity at confined locations or time points are common
81 within the hyporheic zone (Boulton et al., 1998; Krause et al., 2017; McClain et al., 2003). These
82 hotspots often correspond to vegetation patterns (Harms and Grimm, 2008; McClain et al., 2003;
83 Schade et al., 2001) and understanding their drivers is critical to accurately representing organic
84 matter decomposition within ecosystem models.

85 Our objective is to determine molecular mechanisms associated with hotspots of aerobic
86 metabolism in the hyporheic zone. Previous work in this system has shown differences in (1)
87 aerobic metabolism across variable mixing of water bodies with distinct C chemistries (Stegen et

88 al., 2016; Stegen et al., 2018) and (2) metabolic processes associated with C oxidation in the
89 presence or absence of vegetation (Graham et al., 2017b). Recent work has also shown that
90 elevated rates of metabolism globally correspond to carbohydrate metabolism in soil
91 metagenomes (Hartman et al., 2017). Based on this research, we hypothesize that microbiome
92 structure and function support elevated carbohydrate metabolism in hyporheic zone hotspots,
93 resulting in higher rates of aerobic metabolism. We expect this dynamic to be reflected in
94 sediment metagenomes, metaproteomes, and metabolomes, whereby hotspot microbiome
95 composition, protein expression, and metabolites are enriched in sugar-associated pathways and
96 compounds, relative to low-activity sediments.

97

98 **2. Materials and Methods.**

99 *2.1. Site Description*

100 This study was conducted along the Columbia River shoreline within the Hanford 300
101 Area (approximately 46° 22' 15.80"N, 119° 16' 31.52"W) in eastern Washington State (Graham
102 et al., 2016a; Graham et al., 2017a; Graham et al., 2017b; Slater et al., 2010; Zachara et al.,
103 2013). The Columbia River experiences shoreline geographic variation in vegetation patterns,
104 substrate geochemistry, microbiome composition, and rates of biogeochemical activity (Arntzen
105 et al., 2006; Graham et al., 2017b; Lin et al., 2012; Peterson and Connelly, 2004; Slater et al.,
106 2010; Stegen et al., 2016; Stegen et al., 2012; Zachara et al., 2013); constituting an ideal system
107 for examining mechanisms associated with biogeochemical hotspots.

108 Liquid N₂-frozen sediment profiles (0-60 cm) were collected along two shoreline
109 transects perpendicular to the Columbia River in March 2015, separated by a distance of ~170m.
110 We collected profiles at three locations in each transect with 5m spacing within a spatial domain

111 of ~175 x 10m. In each transect, the lower bank profile was located at ~0.5m (vertical distance)
112 below the water line and the upper bank profile was located ~0.5m (vertical distance) above the
113 water line (approximately 10m horizontal distance), with the third profile situated at the
114 midpoint. Each profile was sectioned into 10-cm intervals from 0-60cm. To capture a range of
115 biogeochemical activities, one transect had dense vegetation, consisting of a closed canopy of
116 woody perennials *Morus rubra* (Red Mulberry) and *Ulmus rubra* (Slippery Elm), and one
117 transect was characterized by a cobbled armor layer with virtually no vegetation.

118

119 2.2. *Sample Collection*

120 Liquid N₂-frozen sediment profiles were collected as outlined in Moser et al. (2003)
121 using a method modified from Lotspeich and Reed (1980) and Rood and Church (1994). A
122 pointed stainless steel tube (152 cm length, 3.3 cm outside diameter, 2.4 cm inside diameter) was
123 driven into the river bed to a depth of ~60cm. Liquid N₂ was poured down the tube for ~15
124 minutes, until a sufficient quantity of material had frozen to the outside of the tube. The tube and
125 attached material were removed from the riverbed with a chain hoist suspended beneath a tripod.
126 Profiles were placed over an aluminum foil lined cooler containing dry ice. Frozen material was
127 removed with a mallet. The material was then wrapped in the foil and transported on dry ice to
128 storage at -80°C. In the lab, profiles were sectioned into 10cm depth intervals from 0-60 cm (n =
129 6 per profile, except for the unvegetated upper bank profile which was sectioned only from 30-
130 60cm; total n = 33)

131

132 2.3. *Physicochemistry*

133 Details concerning physicochemical assays are provided in the Supporting Information.
134 Briefly, we determined the particle distribution of sediments by separating size fractions via
135 sieving; total nitrogen, sulfur, and carbon content were determined using an Elementar vario EL
136 cube (Elementar Co.Germany); NH_4^+ was extracted with KCl and measured with Hach Kit
137 2604545 (Hach, Loveland, Co); iron content was measured with a ferrozine assay; and all other
138 ion concentrations were measured by inductively coupled plasma mass spectrometry (ICP-MS)
139 on HCl extractions. Aerobic metabolism was determined with a resazurin reduction assay,
140 modified from *Haggerty et al.*(2009). Data are provided in Fig. S1-S3.

141

142 2.4. FT-ICR-MS solvent extraction and data acquisition

143 We leverage state of science chemical extraction protocols combined with Electrospray
144 ionization (ESI) and Fourier transform ion cyclotron resonance (FT-ICR) mass spectrometry
145 (MS) to infer differences in metabolites among our samples. ESI FT-ICR-MS introduces intact
146 organic molecules into the MS without fragmentation and allows for the detection of a wide
147 range of chemical compounds (Tfaily et al., 2015; Tfaily et al., 2017). The use of 12 Tesla (T)
148 FT-ICR-MS offers high mass resolving power ($>1\text{M}$) and mass measurement accuracy (<1 ppm),
149 and while nascent in its application within complex environmental systems, it has emerged as a
150 robust method for determining the chemistry of natural organic compounds (Kim et al., 2003;
151 Koch et al., 2005; Tfaily et al., 2011; Tremblay et al., 2007). Additionally, Tfaily et al. (2015;
152 2017) have optimized metabolite characterization from soils and sediments by sequential
153 extraction with polar and non-polar solvents tailored to the sample set of interest. Tfaily's
154 extraction procedures have been coupled to ESI FT-ICR-MS to distinguish metabolites among
155 ecosystems and soil types (Tfaily et al., 2015; Tfaily et al., 2017) as well as to provide

156 information on the utilization of distinct metabolites among samples within a single environment
157 (Bailey et al., 2017; Graham et al., 2017b; Stegen et al., 2018).

158 Here, we used three solvents with different polarities —water (H₂O), methanol (CH₃OH,
159 ‘MeOH’) and chloroform (CHCl₃)—to sequentially extract a large diversity of organic
160 compounds from samples, according to Tfaily et al. (2015; 2017). Water extractions were
161 performed first, followed by MeOH and then CHCl₃. Previous work has shown that each solvent
162 is selective towards specific types of compounds (Tfaily et al., 2015) and that combining peaks
163 from all extractions provides a more comprehensive description of metabolite composition than
164 any single extraction procedure independently (Tfaily et al., 2017). Water is a polar solvent with
165 a selection bias for carbohydrates with high O/C ratios, amino-sugars, and other labile polar
166 compounds (Tfaily et al., 2015). Conversely, CHCl₃ is selective for non-polar lipids associated
167 with mineral interactions and cellular membranes (Tfaily et al., 2015). Methanol has a polarity in
168 between that of water and CHCl₃ and extracts a mixture of compounds that water and CHCl₃
169 extract (Tfaily et al., 2015). Ultra-high resolution mass spectrometry of the three different
170 extracts from each sample was carried out using a 12 Tesla Bruker Solarix FT-ICR-MS located
171 at the Environmental Molecular Sciences Laboratory (EMSL) in Richland, WA, USA. An
172 expanded description of extraction procedures; instrument calibration and maintenance; and
173 sample injection is presented in the Supplemental Material.

174

175 *2.5. FT-ICR-MS data processing*

176 One hundred forty-four individual scans were averaged for each sample and internally
177 calibrated using an organic matter homologous series separated by 14 Da (–CH₂ groups). The
178 mass measurement accuracy was less than 1 ppm for singly charged ions across a broad *m/z*

179 range (100-1200 m/z). The mass resolution was $\sim 350K$ at 339 m/z . Data Analysis software
180 (BrukerDaltonik version 4.2) was used to convert raw spectra to a list of m/z values applying
181 FTMS peak picker module with a signal-to-noise ratio (S/N) threshold set to 7 and absolute
182 intensity threshold to the default value of 100.

183 For each sample, we combined peaks detected in all three extractions to yield metabolite
184 composition. Peak data were treated as presents/absence data, where a peak was considered to be
185 present in a sample if it was present in at least one of H₂O-, MeOH-, or CHCl₃-extracted
186 samples. Presence/absence data were used because peak intensity differences are reflective of
187 ionization efficiency as well as relative abundance (Kujawinski and Behn, 2006; Minor et al.,
188 2012; Tfaily et al., 2015; Tfaily et al., 2017).

189 Putative chemical formulae were then assigned using in-house software following the
190 Compound Identification Algorithm (CIA), proposed by Kujawinski and Behn (2006), modified
191 by Minor et al. (2012), and previously described in Tfaily et al. (2017). Chemical formulae were
192 assigned based on the following criteria: S/N >7, and mass measurement error <1 ppm, taking
193 into consideration the presence of C, H, O, N, S and P and excluding other elements. To ensure
194 consistent formula assignment, we aligned all sample peak lists for the entire dataset to each
195 other in order to facilitate consistent peak assignments and eliminate possible mass shifts that
196 would impact formula assignment. We implemented the following rules to further ensure
197 consistent formula assignment: (1) we consistently picked the formula with the lowest error and
198 with the lowest number of heteroatoms and (2) the assignment of one phosphorus atom requires
199 the presence of at least four oxygen atoms.

200 The chemical character of thousands of peaks in each sample's ESI FT-ICR-MS
201 spectrum was evaluated on van Krevelen diagrams. Compounds were plotted on the van

202 Krevelen diagram on the basis of their molar H:C ratios (y-axis) and molar O:C ratios (x-axis)
203 (Kim et al., 2003). Van Krevelen diagrams provide a means to visualize and compare the average
204 properties of organic compounds and assign compounds to the major biochemical classes (e.g.,
205 lipid-, protein-, lignin-, carbohydrate-, and condensed aromatic-like).

206

207 *2.6. Identification of putative biochemical transformations using FT-ICR-MS*

208 To identify potential biochemical transformations, we followed the procedure detailed by
209 Breitling *et al.* (2006) and employed by Bailey *et al.* (2017), Graham *et al.* (2017b), and Stegen
210 *et al.* (2018). The mass difference between m/z peaks extracted from each spectrum with S/N>7
211 were compared to commonly observed mass differences associated with biochemical
212 transformations. All possible pairwise mass differences were calculated within each extraction
213 type for each sample, and differences (within 1ppm) were matched to a list of 92 common
214 biochemical transformations (e.g., gain or loss of amino groups or sugars, Table S1). For
215 example, a mass difference of 99.07 corresponds to a gain or loss of the amino acid valine, while
216 a difference of 179.06 corresponds to the gain or loss of a glucose molecule. Pairs of peaks with
217 a mass difference within 1 ppm of our transformation list were considered to be related by the
218 corresponding compound. This approach is feasible with FT-ICR-MS data because the set of
219 peaks in each sample are related by measureable and clearly defined mass differences
220 corresponding to gains and losses of compounds.

221

222 *2.7. Metagenome sequencing and annotation*

223 To release biomass from sediment particles, thawed samples were suspended in 20mL of
224 chilled PBS/0.1% Na-pyrophosphate solution and vortexed for 1 min. The suspended fraction

225 was decanted to a fresh tube and centrifuged for 15' at 7000 x g at 10°C. DNA was extracted
226 from the resulting pellets using the MoBio PowerSoil kit in plate format (MoBio Laboratories,
227 Inc., Carlsbad, CA) following manufacturer's instructions, with the addition of a 2-hour
228 proteinase-K incubation at 55°C prior to bead-beating to facilitate cell lysis.

229 Purified genomic DNA was submitted to the Joint Genome Institute under JGI/EMSL
230 proposal 1781 for paired-end sequencing on an Illumina HiSeq 2500 sequencer (Table S2).
231 Reads were processed by BBDuk to remove adapters (ktrim=r, minlen=40, minlenfraction=0.6,
232 mink=11, tbo, tpe, k=23, hdist=1, hdist2=1, ftm=5) and trim for quality <12 (maq=8, maxns=1,
233 minlen=40, minlenfraction=0.6, k=27, hdist=1, trimq=12, qtrim=rl) and screened for
234 contaminants against a masked version of human HG19 using BMap (fast local minratio=0.84
235 maxindel=6 tipsearch=4 bw=18 bwr=0.18 usemodulo printunmappedcount idtag minhits=1).
236 (BBDuk and BMap are available at <https://sourceforge.net/projects/bbmap>) Remaining reads
237 were assembled with megahit (Li et al., 2016) using default parameters and k-list=23,43,63,
238 83,103,123. Gene calling and functional and taxonomic annotation was performed by MGAP
239 v4.11.4 (Huntemann et al., 2015). Data sets are available through the JGI Genome Portal
240 (<http://genome.jgi.doe.gov>). Project identifiers are listed in Table S2.

241

242 *2.8. Metaproteomics.*

243

244 Sediment samples were prepared for proteome analysis as per Nicora et al. (in press) and
245 detailed below. Additional details are provided in the Supplemental Material.

246 Proteins were extracted from 30 g of lyophilized sediment using MPLex direct SDS
247 buffer extraction. Sediment was weighed into 50 mL screw-cap self-standing tubes (Next

248 Advance, Averill Park, NY) along with 0.9-2.0 mm stainless steel beads, 0.1 mm zirconia beads
249 and 0.1 mm garnet beads and 8 mL of 60% MeOH in nanopure water. The samples were all bead
250 beat in a 50mL Bullet Blender (Next Advance, Averill Park, NY) at speed 12 for 15 minutes at
251 4°C and transferred into chemical compatible polypropylene 50 mL tubes (Olympus Plastics,
252 Waltham, MA). The dirty tube was rinsed with 2 mL of 60% MeOH and combined with the
253 sample along with 12 mL of ice-cold chloroform and probe sonicated at 60% amplitude for 30
254 seconds on ice, allowed to cool on ice and sonicated again. Samples were incubated for 5 min at-
255 80°C, vortexed for 1 min and centrifuged at 4,500x g for 10 min at 4°C. The upper aqueous
256 phase was collected into a large glass vial, being careful not to touch the protein interphase. The
257 interphase was collected using a large flat spatula into a separate tube and 5mL of ice cold 100%
258 methanol was added to the protein, vortexed and centrifuged at 4,500x g for 5 mins at 4°C to
259 pellet the protein. The supernatant was decanted off and the protein allowed to dry upside down
260 on a Kim-wipe. Meanwhile, the bottom organic phase was collected into a separate large glass
261 vial and 5mL of nanopure water was added to the large soil particulates along with 25mL of cold
262 (-20 °C) chloroform:methanol (2:1, v:v) solution and probe sonicated for 30 seconds as
263 described previously. The sample was allowed to cool at -20°C and centrifuged at 4,500x g for
264 10 min at 4°C. The upper aqueous phase metabolites and bottom organic phase lipids were
265 collected together with the supernatant metabolites and lipids then dried down completely in a
266 vacuum concentrator (Labconco, Kansas City, MO) and stored at -20 °C for analysis. The dirty
267 protein interphase had 20 mL of an SDS-Tris buffer (4% SDS, 100mM DTT in 100 mM Tris-
268 HCl, pH 8.0) added and probe sonicated at 20% amplitude to bring into solution and incubated at
269 95°C for 5 minutes to solubilize the protein and allowed to cool for 20 mins at 4°C. The samples
270 were centrifuged at 4,500x g for 10 mins and the supernatants were decanted into chemical

271 compatible polypropylene 50 mL tubes. The proteins were precipitated by adding up to 20%
272 trichloroacetic acid (TCA), vortexed and placed in a -20°C freezer for 1.5 hrs. The samples were
273 thawed and centrifuged at 4,500x g at 4°C for 10 mins to collect the precipitated protein. The
274 supernatant was gently decanted into waste and 2 mL of ice cold acetone was added to the pellet
275 and re-suspend by vortexing. The sample was placed at -80°C for ~5 mins to ensure the samples
276 were cold and centrifuged for 10 mins at 4,500x g at 4°C. The acetone was removed by gently
277 pouring into waste. The pellets were allowed to dry inverted on a Kim Wipe for ~15 mins. The
278 protein interphases from the supernatants and the protein interphases from the particulates were
279 combined using 100-200ul of SDS-Tris buffer and FASP digested (described in the
280 Supplemental Material).

281 Extractions were analyzed on a Q-Exactive Plus mass spectrometer (Thermo Electron,
282 Waltham, MA) coupled to a Waters NanoAcquity high performance liquid chromatography
283 systems (Waters Corporation, Milford, MA) through 75 um x 70 cm columns packed with
284 Phenomenex Jupiter C-18 derivatized 3 um silica beads (Phenomenex, Torrance, CA). Samples
285 were loaded onto columns with 0.05% formic acid in water and eluted with 0.05% formic acid in
286 Acetonitrile over 100 minutes. Ten data-dependent MS/MS scans (17.5K resolution, centroided)
287 were recorded for each survey MS scan (35K resolution) using normalized collision energy of
288 30, isolation width of 2.00, and rolling exclusion window of +/- 1 Th lasting 30 seconds before
289 previously fragmented signals are eligible for re-analysis.

290 For protein identification, a single reference protein file was developed from the
291 combination of 33 metagenome files ('assembled.faa' and 'product_names' files) available from
292 the Joint Genome Institute (JGI Project IDs and related download links in Supplemental
293 Material). Exact sequence duplicates were removed, and 16 commonly observed contaminants

294 (e.g. tryptic fragments, human keratins, and serum albumin precursors) were included. Final file
295 contained 10,063,272 protein entries from 1,299,102,456 amino acids, 2.04GB in size.

296 The MS/MS spectra from all LC-MS/MS datasets were converted to ASCII text (.dta
297 format) using MSConvert (<http://proteowizard.sourceforge.net/tools/msconvert.html>) which
298 more precisely assigns the charge and parent mass values to an MS/MS spectrum. The data files
299 were then interrogated via target-decoy approach
300 (<http://www.ncbi.nlm.nih.gov/pubmed/20013364>) using MSGFPlus
301 (<http://www.ncbi.nlm.nih.gov/pubmed/25358478>) using a +/- 20 ppm parent mass tolerance,
302 partially tryptic digestion enzyme settings, and a variable posttranslational modification of
303 oxidized Methionine. All MS/MS search results for each dataset were collated into tab separated
304 ASCII text files listing the best scoring identification for each spectrum.

305 Collated search results were further combined into a single result file. These results were
306 imported into a Microsoft SQL Server database. Results were filtered to >1% FDR using an
307 MSGF+ supplied Q-Value that assesses reversed sequence decoy identifications for a given
308 MSGF score across each dataset. Using the protein references as a grouping term, unique
309 peptides belonging to each protein were counted, as were all peptide spectrum matches (PSMs)
310 belonging to all peptides for that protein (i.e. a protein level observation count value). PSM
311 observation counts reported for each sample that was analyzed. Cross-tabulation tables were
312 created to enumerate protein level PSM observations for each sample, allowing low-precision
313 quantitative comparisons to be made. Identified proteins were searched against dbCAN v5 (Yin
314 et al., 2012) using hmmer v3.1 (Finn et al., 2011). CAZy families were assigned based on cutoff
315 suggestions of the maintainers of dbCAN. We removed singletons and proteins not present in at

316 least 25% of samples prior to statistical analysis. Samples with less than 10 total protein counts
317 ($n = 4$) were removed from our dataset to yield 29 samples.

318

319 *2.9. Statistical analyses.*

320 After removing samples with low protein detection, we retained 29 samples for statistical
321 analyses. To identify hotspots vs. low-activity sediments, we examined the distribution of
322 aerobic metabolism rates in all 29 samples (Fig. S4). The distribution was approximately normal
323 with a minimum of 361.83, a mean of 810.49, and a maximum of 1472.86. We defined low-
324 activity sediments as those falling in the bottom quartile of the distribution (maximum rate of
325 aerobic metabolism: 619.45) and high activity samples as those falling in the top quartile of the
326 distribution (minimum rate of aerobic metabolism: 1033.66), yielding a sample size of 7 for each
327 set of samples. All statistical analyses were conducted using R software ([https://www.r-](https://www.r-project.org/)
328 [project.org/](https://www.r-project.org/)) and graphics were generated with either base packages or ‘ggplot2.’

329 We used two approaches to determine differences in the composition of metabolites,
330 microbial phylogenies, metagenomic functional potential (i.e., all annotated genes), and
331 metaproteomes between hotspots and low-activity sediments. Metabolomes were analyzed as
332 present/absence data, and all other data types were analyzed as relative abundances. First, we
333 examined differences in full compositional profiles across groups using permutational
334 multivariate analysis of variance (PERMANOVA, 999 permutations, ‘vegan’ package) and
335 accounting for potential non-independence between samples within the same core by stratifying
336 this analysis by depth (as per Graham et al., 2017b). Because metabolite data were in
337 presence/absence form, a Sorenson dissimilarity distance was constructed and used to compare

338 metabolite data. All other datasets were analyzed using Bray-Curtis dissimilarity to account for
339 differences in relative abundance.

340 Secondly, we used linear mixed effect models to investigate variation in specific
341 microbial attributes between hotspots and low-activity sediments. Following the convention of
342 Graham et al. (2017b), we included depth as a random effect in these models to account for
343 possible non-independence among sediments. A given microbial attribute was the dependent
344 variable and sediment type was the independent variable. Mixed models were compared to null
345 expectations (i.e., a model including only random effects) with analysis of variance (ANOVA) to
346 determine significance. For significant models ($P < 0.05$), R^2 values for microbial attributes
347 independently and for the entire mixed model (i.e., the effects of the attribute plus depth) were
348 determined with the 'r.squaredGLMM' function in the 'MuMIn' package (Barton, 2009).
349 Because few compounds differed across activity levels, P-values were not adjusted for multiple
350 comparisons in order to maximize the likelihood of detecting differences.

351 We selected specific attributes for analysis as follows: For metabolomic data, we grouped
352 peaks by their Van Krevlen assignment and compared the relative abundance of each Van
353 Krevlen class across activity groups (e.g., the relative abundance of amino sugars in hotspots vs.
354 low-activity sediments). To identify specific microbial phylogenetic and functional groups of
355 interest, we employed the following procedure: (1) we ranked the relative abundance of each
356 assigned phylogeny (class level) or functional annotation (pfam, COG, and KEGG) within each
357 sample; (2) we compiled a master list containing the 25 most abundant phylogenies or functional
358 annotations in every sample; and (3) we determined the relative abundance of every unique
359 phylogeny or functional annotation in the master list within each sample. In this way, we
360 examined every taxon or metabolic pathway that was present in high abundance in at least one

361 sample. For metaproteomic data, we employed a similar pipeline but because the proteomic
362 dataset was smaller, we comprised our master list from any protein in the top 5% of relative
363 abundance in a given sample (equivalent to more than 10 proteins per sample).

364 Because (1) metagenomes did not vary strongly between hotspots and low-activity
365 sediments, (2) proteins are tightly coupled to function conceptually (i.e., they are proximate
366 catalysts of reactions), and (3) metabolites were dramatically different across activity levels; we
367 focused subsequent analyses only on metaproteomic and metabolomics data. We first
368 investigated the extent to which metaproteome structure corresponded to metabolome structure
369 in the full dataset ($n = 29$) by comparing dissimilarity in metabolite composition (Sorenson
370 dissimilarity) to differences in proteomic composition (Bray-Curtis dissimilarity, Mantel test
371 using Spearman correlation and stratifying by depth). We also examined the extent to which each
372 data type corresponded to changes in aerobic metabolism by fitting linear and quadratic mixed
373 effects models across the full dataset. As above, depth was included as a random effect in each
374 model. To reduce multidimensional metaproteomic and metabolomic data into vectors for model
375 construction, we extracted the first two principle components of variation within data type (PCA
376 analysis, 'prcomp'). PC1 and PC2 were predictors in the mixed effects models, and aerobic
377 metabolism was the response variable. Mixed models were compared to null expectations (i.e.,
378 models including only random effects) with ANOVA to determine significance. R^2 values were
379 determined with the 'r.squaredGLMM' function in the 'MuMIn' package. Since both PC1 and
380 PC2 from metabolomes were significantly related to aerobic metabolism, we also constructed a
381 model that included both PC1 and PC2 as predictors. Lastly, to examine additional explanatory
382 power that could be gained by incorporating bulk C as predictors along with metabolite
383 chemistry, we constructed a final model with PC1, PC2, and percent C. This model was

384 compared to the model with PC1 and PC2 only using ANOVA to determine if percent C
385 significantly improved explanatory power of aerobic metabolism.

386 Finally, because metabolomes were most distinguishing between hotspots and low-
387 activity sediments and corresponded to rates of aerobic metabolism, we concluded with set of
388 analyses with metabolomic data only. We deciphered specific metabolites that distinguished
389 hotspots from low-activity sediments using ‘random forest’ machine learning algorithm (Liaw
390 and Wiener, 2002, ‘randomForest’ package). Forests were constructed using metabolite peaks
391 that were assigned chemical formulae, removing singletons for computational limitations. For
392 each forest, 1000 trees were constructed with replacement, and error rates converged near zero
393 (Fig. S5). Peak importance and partial dependence in distinguishing high vs. low activity
394 samples were calculated with the ‘imp’ and ‘PartialPlot’ functions, respectively. Peaks that were
395 assigned a ‘Mean Decrease in Accuracy’ of greater than zero were considered to be
396 distinguishing metabolites. This approach yielded 55 peaks distinguishing low activity samples
397 and 272 peaks distinguishing high activity samples.

398 We also used metabolite transformations to infer biochemical processes associated with
399 aerobic metabolism in hotspots. We identified transformations of interest using the following
400 procedure: (1) we summed the relative abundance of each transformation across all samples (n =
401 29) and (2) we chose the transformations whose abundance was in the top 20% of all
402 transformations (18 out of 92, mean relative abundance: 1.6% to 14.5%). By doing so, we
403 analyzed transformations that occurred most frequently in our dataset, irrespective of their
404 occurrence within individual samples.

405 We then compared the relative abundance of specific transformations between hotspots
406 and low-activity sediments using linear mixed effect models using the same procedure as for

407 abundant microbial phylogenies, metabolic pathways, expressed proteins, and metabolites
408 (described above). Because many abundant transformations in hotspots seemed to contain
409 nitrogen (N) while transformations in low-activity sediments did not, we also examined the role
410 in nitrogenous compounds in biochemical transformations in hotspots. We separated
411 transformations into those containing N and N-free transformations (Table S3 and S4) and
412 determined the relative abundance of transformations in each sample that were either nitrogenous
413 or N-free. We then compared the relative abundance of nitrogenous and N-free compounds
414 between hotspots and low-activity sediments using mixed effects models. Because the
415 distribution of transformations within nitrogenous and N-free categories was highly non-
416 Gaussian, we applied a Box-Cox transformation to each data type ('boxCox' function in 'car'
417 package, Sakia, 1992), and subsequently constructed linear mixed effects models. We further
418 separated amino-acid nitrogenous transformations from non-amino acid nitrogenous
419 transformations (hereafter 'complex N', Table S5 and S6), applied a Box-Cox transformation,
420 and constructed linear mixed effects models with these data.

421

422 **3. Results.**

423 *3.1. Multi 'Omic Differences across Rates of Aerobic Metabolism*

424 Microbiome structure, functional potential, and protein expression showed limited
425 differences between hotspots vs. low-activity sediments. No significant difference was observed
426 in the composition of microbiome phylogenies ($P = 0.57$, Fig. 1a), metagenomic annotations ($P =$
427 0.69 , Fig. 1b), and metaproteomes ($P = 0.91$, Fig. 1c) between hotspots and low-activity
428 sediments (PERMANOVA). As well, major clades, pathways, and proteins were similar among
429 sediments (Fig. 1).

430 Metabolome composition, however, differed between hotspots and low-activity samples
431 ($P = 0.01$, $R^2 = 0.28$, Fig. 1d, PERMANOVA). When grouped into major compound classes,
432 hotspots contained more amino sugar- and protein-like metabolites ($P < 0.05$, Fig. 1d, mixed
433 effects). Metabolite pool composition was uncorrelated to proteome structure ($P = 0.62$, Fig. S6,
434 Mantel test across all samples).

435 Finally, when considering aerobic metabolism as a continuous variable, metabolome
436 composition was tightly correlated to aerobic metabolism. The first two principle coordinates of
437 metabolite pool composition had significant independent relationships with aerobic metabolism
438 and together explained 67% of variation in aerobic metabolism (mixed effects, P (PC1, quad.) =
439 0.0006 & P (PC2, linear) = 0.01, P (combined, linear) $\ll 0.0001$, Fig. 2a, Fig. S7). Statistical
440 models including both bulk C and metabolite composition were not significantly different from
441 models including metabolite composition alone (R^2 0.73 vs. 0.67, $P = 1$). Metaproteome
442 composition was uncorrelated to aerobic metabolism despite representing the proximal catalysts
443 of metabolic processes (mixed effects, P (PC1) = 0.29 & P (PC2) = 0.53, Fig. 2b).

444

445 *3.2. Distinguishing Metabolite Chemistry and Transformations*

446 Because hotspots were discriminated from low-activity sediments only in the
447 metabolome, we examined metabolite pools to identify compounds and infer processes that
448 distinguished hotspots from low-activity sediments. Hotspots were characterized by high
449 molecular weight ($P < 0.0001$, Fig. 3a), chemically diverse (Fig. 3b) compounds. In particular,
450 nitrogenous classes of compounds (i.e., amino sugars and proteins) were present in metabolites
451 that described hotspots and absent from metabolites characterizing low activity sediments (Fig.
452 3b).

453 Hotspots also exhibited a more even distribution of abundant biochemical
454 transformations, indicating greater diversity of biogeochemical processes in these sediments
455 relative to low-activity sediments (Fig. 4a). These transformations were inferred by comparing
456 differences between peaks in FT-ICR-MS data to a database of known biochemical
457 transformations (see section 2.6). Further, transformations involving nitrogenous compounds (P
458 = 0.03 R^2 = 0.26, Table S3), particularly those involving complex N (P = 0.04 R^2 = 0.20, Fig. 4b,
459 Table S6)), were more prevalent in hotspots.

460

461 **4. Discussion.**

462 Our multi ‘omics investigation of microbiome structure and function in hyporheic zone
463 hotspots highlights the importance of vegetation in driving local-scale (<200m²) aerobic
464 metabolism in the subsurface. Pronounced spatial heterogeneity in metabolism across our
465 relatively small spatial domain (~175 x 10m) is inconsistent with representations of
466 biogeochemical processes within ecosystem models that often aggregate environmental
467 properties across similarly sized domains, and suggests a need for finer scale representation of
468 processes in these models. Our research supports previous observations of relationships between
469 riparian vegetation and hyporheic zone hotspots due to associated carbon and nutrient inputs
470 (Harms and Grimm, 2008; McClain et al., 2003; Schade et al., 2001). For example, Kuglerova et
471 al. (2014) demonstrated that increased vegetation growth corresponded to enhanced
472 biogeochemical cycling in areas of the hyporheic zone experiencing maximal groundwater-
473 surface water mixing. In our system, hotspots were disproportionately distributed within
474 sediments underlying vegetation and exhibited up to 4 times the rate of aerobic metabolism in
475 low-activity sediments despite a small sampling domain (Fig. S4).

476 Contrary to our hypothesis that microbiome composition (metagenomics) and expression
477 (metaproteomics) would be shifted towards carbohydrate metabolism in hotspots, our results
478 support a predominant role for phenotypic plasticity within sediment microbiomes. Phenotypic
479 plasticity, whereby an organism encodes multiple metabolic pathways and adjusts the expression
480 of each depending on environmental conditions, is common in free-living microbes and
481 manifests as different metabolisms occurring in samples with the same genomic composition.
482 Microbiome phylogeny and functional potential did not vary with the level of aerobic respiration
483 in our sediments. While Hartman *et al.* (2017) demonstrated a consistent metagenomic shift from
484 aromatic to carbohydrate metabolism with increased carbon turnover in global soils, we observed
485 no such relationship in our sediments. A key difference between Hartman *et al.* (2017) and the
486 present study is that Hartman *et al.* demonstrated decreased nutrient uptake with increased
487 activity. In the present study, decreased nutrient uptake would correspond to fewer nitrogenous
488 biochemical transformations, whereas we observed increasing activity of biochemical pathways
489 involving nitrogen in hotspots vs. low-activity sediments. The lack of microbiome change in our
490 study may therefore be constrained by N-limitation. For instance, if microbiomes in our
491 sediments are always under strong selection for N-harvesting metabolisms, they would be
492 structured to contain organisms with the ability to metabolize organic N (Craine et al., 2007;
493 Moorhead and Sinsabaugh, 2006), regardless of the magnitude of available N.

494 Interestingly, while one would expect protein expression to correlate with changes in
495 activity, we saw no such effect in our metaproteomic data. The long residence time of enzymes
496 compared to hydrologic mixing may decouple expression from function in dynamic
497 environments such as the hyporheic zone. Hydrologic exchange in the hyporheic zone occurs on
498 relatively short time scales (commonly minutes to days), as well as longer seasonal exchanges,

499 and creates fleeting periods of nutrient availability (Cardenas et al., 2008; Fritz and Arntzen,
500 2007; Sawyer and Cardenas, 2009; Zarnetske et al., 2011). Recent work in soils has shown that
501 protein decay rates vary widely across enzymes and that some enzymes can persist more than 12
502 weeks in soils (Schimel et al., 2017). Assuming a similar magnitude of decay in hyporheic zone
503 sediments would generate a disconnect between protein expression and resource availability that
504 may ultimately disassociate enzyme pool composition from respiration rates, leading to the lack
505 of relationship between metaproteome composition and aerobic metabolism observed here.

506 Our results indicate that improved resolution into sediment metabolomes provides greater
507 insight into sediment function than bulk C content at a given point in time. While there is a long
508 history of constraining biogeochemical rates with standing resource pools, our work indicates
509 that bulk C alone was an insufficient predictor of aerobic respiration (Fig. S8). Metabolite
510 chemistry explained 67% of variation in aerobic metabolism (in comparison to 42% by bulk C
511 pools, Fig. S8), and including bulk C content as a predictor in addition to metabolome
512 composition did not improve the explanatory power of our statistical models. In particular,
513 nitrogenous metabolites were associated with enhanced metabolism. We therefore suggest that
514 metabolome composition can explain aerobic metabolism better than bulk C stocks in some
515 systems and subsequently that new process-based model structures need to account for multiple
516 C pools with various organic N content. For instance, current generation models incorporate
517 mechanisms reflecting that certain C molecules (e.g., glucose) can enter microbial cells by
518 passing directly through their cellular membranes, while larger molecules require secreted
519 extracellular enzymes to break them down prior to microbial uptake (Wang et al., 2013; Wieder
520 et al., 2014). Extending these new frameworks to further parse C chemistry by organic N content
521 and associated metabolic pathways could advance the performance of these models.

522 Microbiomes associated with hyporheic zone hotspots metabolized a broader range of
523 resources compared to microbiomes in low-activity sediments. Enhanced biogeochemical
524 cycling can be achieved through microorganisms using different portions of resource pools
525 (termed ‘resource partitioning’ or ‘niche complementarity’) in which diverse microorganisms use
526 different portions of resource pools (Cardinale, 2011; Loreau and Hector, 2001; Schoener, 1974).
527 Consistent with this mechanism, we show a divergence in active metabolic pathways (inferred
528 from biochemical transformations) in hotspots compared to low-activity sediments. Thus, despite
529 similar microbiome genomic content, hotspots seem to result from the expansion of realized
530 niches of extant microbiomes. In this case, increased biogeochemical function arises through
531 phenotypic plasticity.

532 Hotspots contained diverse metabolites, were distinguished from low-activity sediments
533 by protein- and amino sugar- like metabolites, and exhibited higher instances of biochemical
534 transformations involving organic N. It has been widely demonstrated that N availability can
535 limit microbial activity (Treseder, 2008), resulting in slower decomposition and ecosystem
536 organic matter processing (Averill and Waring, 2017; Weintraub and Schimel, 2003; Zhang et
537 al., 2008); and previous work in this system has suggested a coupling of C and N cycles (Stegen
538 et al., 2018). An excess of inorganic N relative to other N sources has also been demonstrated to
539 shift microbial metabolism by suppressing oxidative enzyme synthesis (Edwards et al., 2011;
540 Jian et al., 2016) and by changing the ratios of synthesized enzymes that degrade recalcitrant vs.
541 labile organic matter (Moorhead and Sinsabaugh, 2006). Similarly our analysis of metabolite
542 transformations supports different metabolic strategies among hotspot vs. low-activity
543 microbiomes as well as a role for organic N in particular as a driver of aerobic metabolism in
544 hotspots.

545 While C:N ratios do not reflect obvious N limitation in our system (means: 6.55 (hotspot)
546 and 6.19 (low-activity)), these ratios can misrepresent chemical conditions experienced by
547 microbiomes due to N absorption to soil matrices and by aggregation processes that physically
548 separate N from microorganisms (Lützow et al., 2006; Sollins et al., 1996). For instance,
549 Darrouzet-Nardi and Weintraub (2014) demonstrated that bulk methods of C and N
550 determination such as those used here may overestimate N availability by 5-fold. As such,
551 organic N may play a critical role in our system despite ratios of C:N in bulk sediments.

552 Moorhead and Sinsabaugh (2006) proposed a concept known as ‘microbial nitrogen
553 mining’ in which microorganisms in environments with limited availability of labile N
554 metabolize C to access organic N (Craine et al., 2007; Moorhead and Sinsabaugh, 2006).
555 Microbial N mining should result in the preferential decomposition of organic N, and our results
556 indicate that this process may contribute to high respiration rates in hyporheic zone hotspots.
557 Consistent with N mining, we observed biochemical pathways involving nitrogenous
558 compounds, and complex N in particular within hotspots, supporting previous observations that
559 organic N increases the decomposition of organic matter (reviewed in Averill and Waring, 2017;
560 Hu et al., 2001; Zhang et al., 2008). The metabolism of complex nitrogenous organic compounds
561 specifically suggests that the breakdown of more chemically-complex organic material may be
562 stimulated to facilitate microbial access to nitrogen and that more available labile C in vegetated
563 sediments provides additional energy necessary to metabolize more chemically-complex
564 nitrogenous molecules.

565 Based on our results, we propose a new conceptualization of hyporheic zone organic
566 matter transformations in which greater resolution into C metabolites and associated metabolic
567 pathways—beyond bulk C stocks and/or microbiome structure—is critical for predictive

568 hydrobiogeochemical models (Fig. 5). Phenotypic plasticity suggests (1) an ability for
569 microbiomes to rapidly acclimate to environmental conditions, regardless of their underlying
570 genomic content, and (2) that there may be limited contributions of metagenomic information to
571 functional predictions in dynamic systems. Because compositional shifts in microbiomes are
572 often on the order of days to months or longer (Balser and Firestone, 2005; Graham et al., 2016a;
573 Poretsky et al., 2014; Waldrop and Firestone, 2006), the ability of microorganisms to adjust
574 metabolic function to suit prevailing conditions without associated species turnover denotes that
575 the timescale of microbiome response is not limited to these longer timescales and instead that
576 microbial responses in some systems may be more tightly constrained by RNA transcription and
577 translation (~minutes or less, Moran et al., 2013).

578 Further, we suggest that broader metabolic activity in biogeochemical hotspots stimulates
579 the production and consumption of diverse metabolites which alter the composition of future
580 resource pools, and therefore that understanding subsequent feedbacks between metabolite
581 consumption and production is vital for predictive biogeochemical models. For example, as
582 metabolism proceeds through time, future metabolic rates are dependent on the resources (i.e.,
583 metabolites) generated by these pathways. Metabolism of an expanded suite of resources through
584 broader realized niche space within biogeochemical hotspots may therefore fundamentally
585 change the subsequent metabolite and resource profiles (Fig. 5). As hotspots and hot moments
586 occur through time and space, resource pools progressively change in ways that are dependent on
587 microbial transformations of prior resource pools, and knowledge about the specific metabolic
588 pathways that generate propagating changes as well as their rates and limitations is imperative
589 for improving biogeochemical predictions. Our results also demonstrate organic N in particular
590 may be involved these alternative pathways, and thus, that nutrient limitations are important

591 considerations in constraining rates within predictive models, as has been recently explored by
592 Wieder et al (2015b). We further extended this conceptualization to propose that organic N
593 specifically should be considered as a possible regulatory factor in process-based ecosystem
594 models.

595 The fate of organic carbon stored in soils and sediments, particularly at terrestrial-aquatic
596 interfaces (Battin et al., 2009; Boulton et al., 1998; Cole et al., 2007; Marín-Spiotta et al., 2014;
597 Regnier et al., 2013), is responsible for large amounts of uncertainty in predictions of future
598 global biogeochemistry (Luo et al., 2016; Todd-Brown et al., 2013). Here, we provide insight
599 into the molecular mechanisms generating hotspots of elevated aerobic metabolism in the
600 hyporheic zone. While microbiome structure and protein expression did not vary across levels of
601 aerobic metabolism, we found that metabolite chemistry and diverse, N-related biochemical
602 pathways were associated with hyporheic zone hotspots. We posit that microbiome phenotypic
603 plasticity can enable microbiomes with similar structure to expand their realized niche in
604 response to changing environments and rapidly increase function in favorable environments.
605 Because of fleeting environmental changes within hyporheic zones, we hypothesize that
606 hyporheic zone microbiomes may exhibit more phenotypic plasticity than more stable
607 ecosystems, meriting further investigation as we explore spatiotemporal dynamics across
608 multiple systems (Fodelianakis et al., 2017). As we search for consensus on the importance of
609 various environmental factors in predicting future biogeochemical rates (Graham et al., 2016b;
610 Graham et al., 2014; Luo et al., 2016; Wieder et al., 2017), a spatially-explicit understanding of
611 the mechanisms regulating organic matter transformations in areas with elevated biogeochemical
612 rates is essential for improving the conceptualization and parameterization of ecosystem models.
613 We propose that understanding and constraining the rates of specific metabolic pathways that

614 utilize discrete portions of resource pools is a critical consideration when advancing carbon cycle
615 complexity in these models.

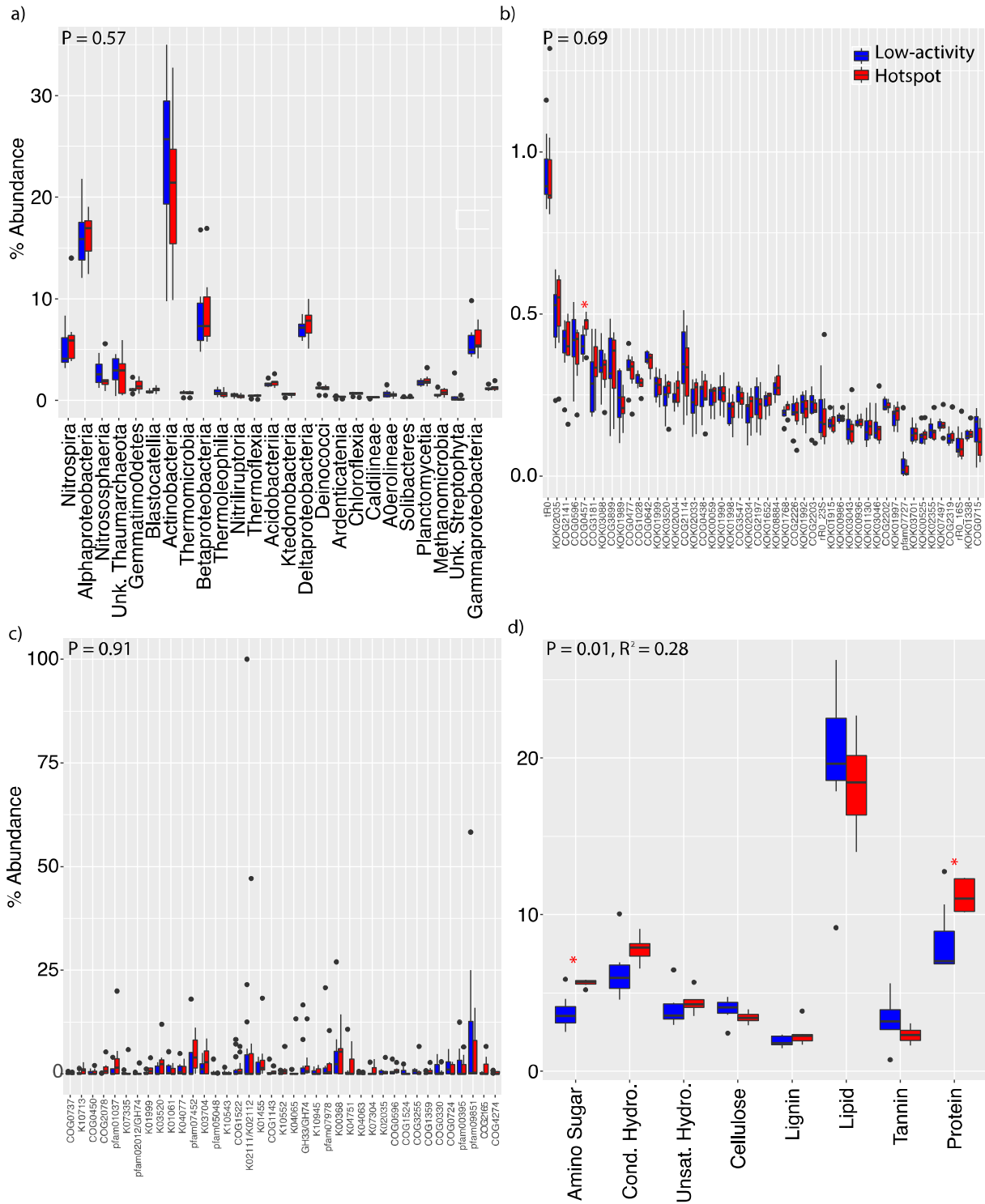
616

617 **Acknowledgements.**

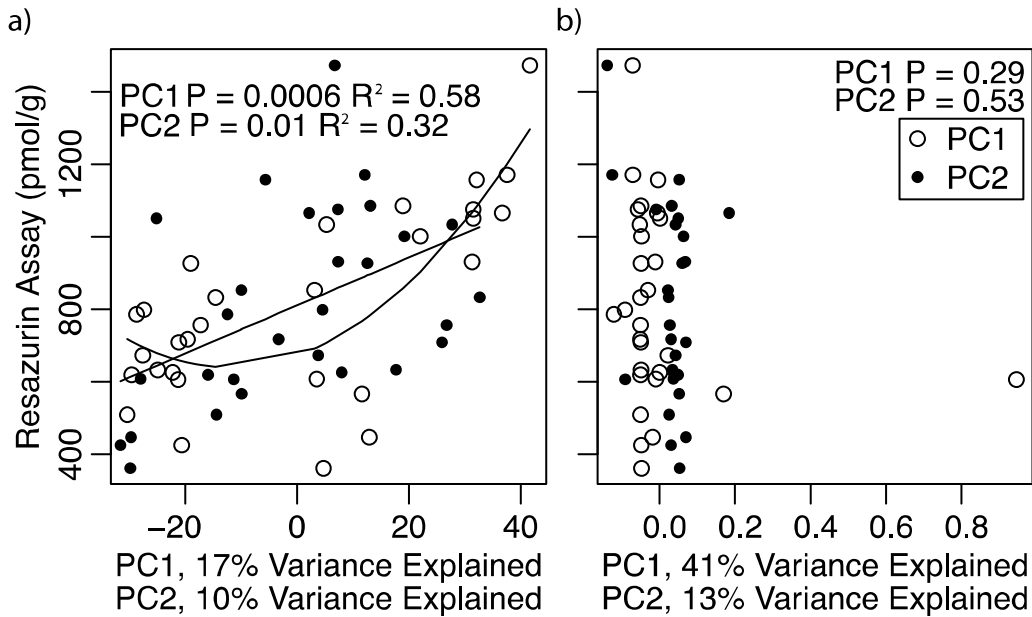
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625

626 **Figures.**



628 **Fig. 1. Multi ‘Omic Differences in Hotspots vs. Low-activity Sediments.** (a-d) show the most
629 abundant classifications only for visual simplicity, but P-values in the upper left-hand corner of
630 each panel are derived from all data. P-values are derived from PERMANOVA with
631 stratification by depth. R^2 values are provided for significant P-values. (a) shows abundant
632 microorganisms (grouped at the class level), (b) shows abundant metagenomics annotations, (c)
633 shows abundant metaproteomic identifications, and (d) shows metabolomics data grouped by
634 compound class. Asterisks denote significant differences between hotspots and low-activity
635 sediments, via mixed models with depth as a random effect ($P < 0.05$).
636
637



638

639 **Fig. 2. Relationships of Metabolomes and Metaproteomes with Aerobic Metabolism.** The

640 first two principle components were extracted from metabolomics and metaproteome data.

641 Correlations between each component and aerobic metabolism were determined using linear and

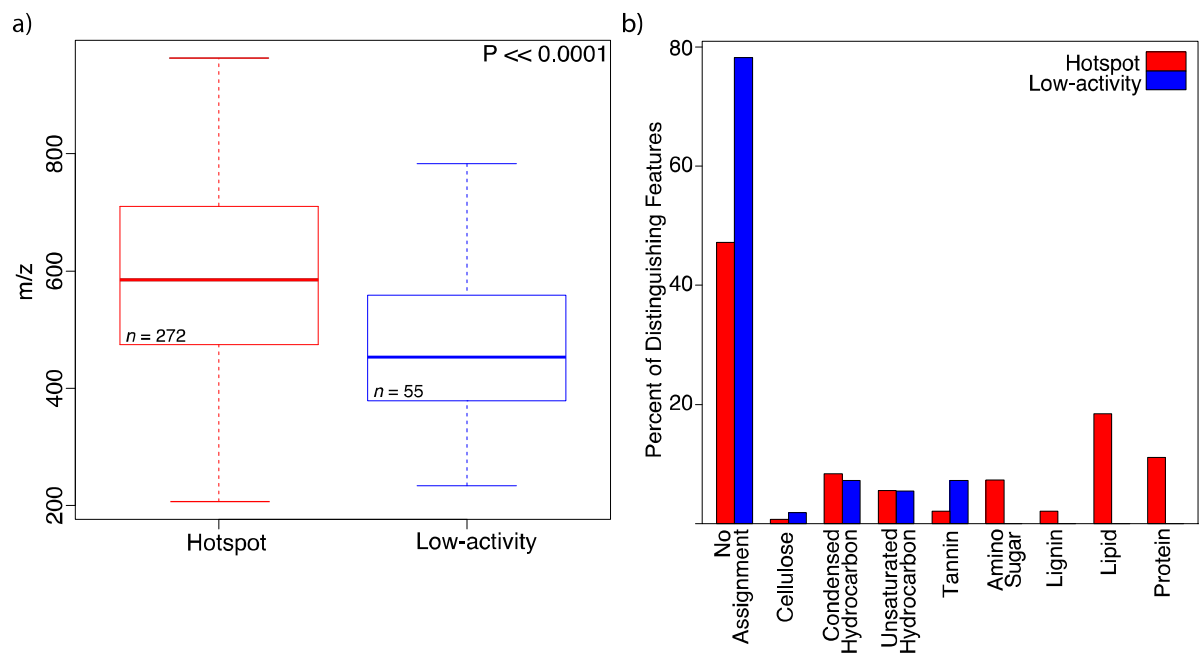
642 quadratic mixed models that included depth as a random effects. Regression lines are shown for

643 the best model; only significant models are shown. (a) Metabolomic data were strongly

644 correlated with aerobic metabolism, while (b) metaproteomic data were uncorrelated.

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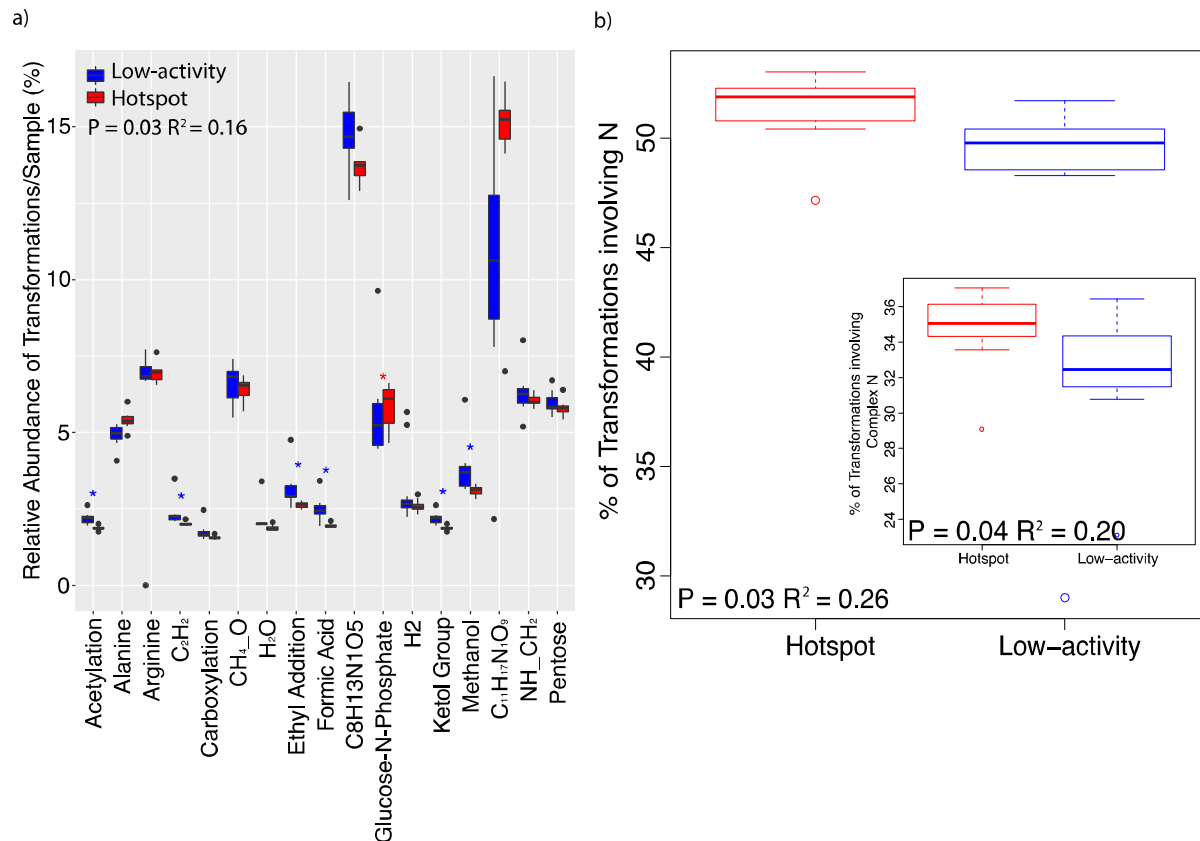
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647
648 **Fig. 3. Metabolites Distinguishing Hotspots from Low-Activity Sediments.** (a) The molecular
649 weight of distinguishing metabolites was significantly higher in hotspots than low-activity
650 sediments. A boxplot of molecular weights are shown, with the center line indicating the mean
651 and the hinges of each box representing the values at the 25th and 75th percentiles. The whiskers
652 represent the range of the data, calculated using default settings in R. P-value is derived from a
653 one-sided Mann-Whitney U test. (b) shows the distribution of distinguishing metabolites in
654 hotspots vs. low-activity sediments. Compounds are grouped by molecular class and displayed as
655 a percent of distinguishing metabolites in each class.

656

657



658

659 **Fig. 4. Differences in Biochemical Transformations between Hotspots and Low-activity**

660 **Sediments.** (a) shows the most abundant transformations only for visual simplicity, but the P-

661 value in the upper left-hand corner is derived from all data. Asterisks denote significant

662 differences between hotspots and low-activity sediments, via mixed models with depth as a

663 random effect ($P < 0.05$). (b) shows a boxplot of the percent of transformations involving N

664 across hotspots and low-activity sediments. The inset shows the same analysis using only

665 transformations involving 'complex N'. Lists of the specific transformations included in each

666 analysis are shown in Tables S3-S6. For each boxplot, with the center line indicating the mean

667 and the hinges of each box representing the values at the 25th and 75th percentiles. The whiskers

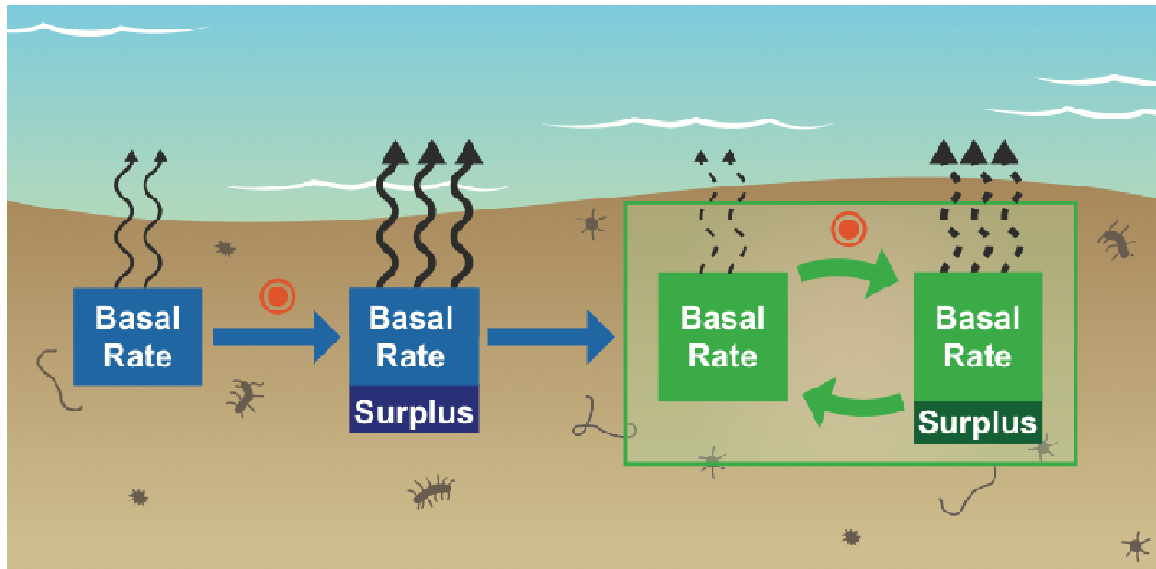
668 represent the range of the data, calculated using default settings in R. P-values and R² are

669 calculated using mixed models with depth as a random effect. Due to non-normality in the

670 distribution of transformations, a Box-Cox transformation was applied prior to model

671 construction.

672



673

674 **Fig. 5. Conceptualization of a Role for Metabolic Pathways in CO₂ Flux Predictions.** We

675 propose that greater resolution into C metabolites and associated metabolic pathways—beyond

676 sediment bulk C and/or microbiome structure—is vital for predictive hydrobiogeochemical

677 models. Broader metabolic activity in biogeochemical hotspots stimulates the production and

678 consumption of diverse metabolites, in particular nitrogenous compounds. Therefore, there is a

679 need to incorporate subsequent feedbacks between metabolite consumption and production into

680 predictive biogeochemical models. Orange bull's eyes represent the occurrence of a hotspot or

681 hot moment. Basal rate is the baseline aerobic metabolism of sediments. Surplus metabolism

682 indicates the increase in aerobic metabolism associated with hotspots and includes stimulated

683 organic N respiration. Arrows denote CO₂ flux, where the number and thickness of arrow

684 denotes the relative size of the flux. Blue boxes indicate the initial condition of sediments and the

685 first occurrence of a hotspot or hot moment. Green boxes show an ongoing feedback loop in

686 which the basal rate of a microbiome is continuously influenced by metabolite production and

687 consumption.

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