

## Understanding How Microbiomes Influence The Systems They Inhabit: moving from a correlative to a causal research framework

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## 1 **Abstract**

2 Translating the ever-increasing wealth of information on microbiomes (environment,  
3 host, or built environment) to advance the understanding of system-level processes  
4 is proving to be an exceptional research challenge. One reason for this challenge is  
5 that relationships between characteristics of microbiomes and the system-level  
6 processes they influence are often evaluated in the absence of a robust conceptual  
7 framework and reported without elucidating the underlying causal mechanisms. The  
8 reliance on correlative approaches limits the potential to expand the inference of a  
9 single relationship to additional systems and advance the field. In this perspective  
10 piece we propose that research focused on how microbiomes influence the systems  
11 they inhabit should work within a common framework and target known microbial  
12 processes that contribute to the system-level process of interest. Here we identify  
13 three distinct categories of microbiome characteristics (microbial processes,  
14 microbial community properties, and microbial membership) and propose a  
15 framework to empirically link each of these categories to each other and the broader  
16 system level processes they affect. We posit that it is particularly important to  
17 distinguish microbial community properties that can be predicted from constituent  
18 taxa (community aggregated traits, CATs) from those properties that are currently  
19 unable to be predicted from constituent taxa (emergent properties, EPs). We discuss  
20 how a series of existing methods in microbial ecology can be applied to more  
21 explicitly elucidate properties within each of these categories and connect these  
22 three categories of microbial characteristics with each other. We view this proposed  
23 framework, gleaned from a breadth of research on environmental microbiomes and  
24 ecosystem processes, as a promising pathway with the potential to advance  
25 microbiome science across a broad range of disciplines.

## 26 ***Current Approaches Linking Microbial Characteristics and Ecosystem*** 27 ***Processes***

28 In all ecosystems, virtually all processes are influenced by microorganisms and  
29 many processes are carried out primarily/exclusively by microorganisms. This has  
30 sometimes led to the assumption that a better description of the microbiome  
31 (including its associated transcripts, proteins, and metabolic products) should lead to  
32 a better understanding and predictions of system level processes. However, such  
33 justifications assume that measurable characteristics of the microbiome (e.g. 16S  
34 rRNA gene libraries, metagenomes, enzymatic activities) can inform our ability to  
35 better understand and predict system-level processes. Unfortunately, additional  
36 information about the microbiome does not always provide a clearer understanding  
37 of ecosystem processes beyond what can be predicted by environmental factors  
38 alone<sup>1,2</sup>.

39 Two recent meta-analyses<sup>3,4</sup> suggest that research at the intersection of  
40 ecosystem science and microbial ecology often rely on assumed correlations  
41 between microbiome characteristics and ecosystem processes and less frequently  
42 test to see if those correlations are present. The first, an examination of 415 studies,  
43 found little evidence that protein-encoding genes (sometimes referred to as  
44 “functional genes”) or gene transcripts correlate with associated biogeochemical  
45 processes<sup>3</sup>. Although all studies attempted (or presumed) to link microbial genes or  
46 transcripts with function, only 14% measured both the copy number of genes or  
47 transcripts and the corresponding process (n = 59 studies, comprising 224 individual  
48 effects as most studies had measured multiple gene-function relationships). Of the  
49 224 effects where both characteristics were measured, only 38% exhibited a positive  
50 relationship between molecular and process measurements that many assume to

51 exist. The effect size for the relationship between gene copy number and process  
52 rate had an approximately normal distribution with a mean near zero<sup>3</sup>. This result  
53 was consistent whether functional gene or transcript abundance was used as the  
54 response variable. The second study, compiled a separate dataset of 148 studies  
55 that examined microbial membership and ecosystem processes in response to  
56 experimental manipulations<sup>4</sup>. Whereas 40% of included studies reported  
57 concomitant changes in microbial membership and an ecosystem process, only one  
58 third of those cases reported the relationship between microbial membership and an  
59 ecosystem process. Interestingly, of the 53 studies that posed a hypothesis or  
60 objective about links between microbial membership and ecosystem processes,  
61 more than half (53%) did not test for a statistical structure-function link<sup>4</sup>.

62 Microbiomes are the engines that power system-level processes<sup>5</sup>. However the  
63 meta-analyses described above illustrate that the current approach to study the links  
64 between microbiome characteristics and ecosystem processes are not well  
65 formulated and relationships between microbiome characteristics and system level  
66 processes are rarely tested. When linkages are explicitly tested, significant  
67 correlations between microbiome characteristics and ecosystem processes are  
68 sometimes present, but more frequently not present<sup>3,4</sup>. One reason for the ambiguity  
69 between microbiome characteristics and system level processes is that many studies  
70 are conducted in the absence of a conceptual framework that illustrates how different  
71 measurable microbial characteristics relate to one another and to the system level  
72 process of interest.

73 Microbial characteristics can range in resolution from cellular abundance to the  
74 entire genetic potential of a species-rich community (i.e., a metagenome). Because  
75 all measureable microbial characteristics do not exist within the same plane of detail,

76 a conceptual structure is required to clarify how various microbial characteristics are  
77 related to each other and to the processes they affect. Identifying correlations  
78 between a microbial characteristic and an ecosystem process in the absence a  
79 broader conceptual underpinning creates the potential for correlations to be  
80 mediated by a third (or more) unaccounted variable(s). The absence of an underlying  
81 mechanism also limits the potential for each result to be applied to additional  
82 systems or to expand to broader spatial and temporal scales, for additional testing,  
83 replication, and confirmation.

#### 84 ***Challenges in Linking Microbial Characteristics and Ecosystem Processes***

85 A key challenge in linking microbial information to a system-level process is that  
86 conceptual research frameworks often do not effectively align with the methods  
87 being applied or the data those methods generate. For example, environmental  
88 factors act on the physiology of individual organisms, which alters their competitive  
89 ability, abundance, collective physiology, and ultimately their contribution to an  
90 ecosystem process. However, designing an observational study or experiment from  
91 this framework assumes that environmental and microbial characteristics are  
92 measurable across multiple categories of ecological organization (i.e., individuals,  
93 populations, and communities) at the temporal and spatial scales at which they  
94 influence microorganisms (Figure 1a). In addition, the relationships between  
95 environmental variables and microbial characteristics can be decoupled in both time  
96 and space<sup>4</sup>, and are often non-linear<sup>6</sup>. Recent immigration, phenotypic plasticity,  
97 disequilibrium between the environment and the extant microbiome at the time of  
98 sampling, functional redundancy, and dormancy can all mask the relationship  
99 between measurable microbial characteristics and the processes microorganisms  
100 influence (Figure 1b).<sup>4,7,8,9</sup> As micrometer scale characteristics of microbiomes ( $10^{-6}$

101 m) are scaled to the level of ecosystems (m to km), we assume that our conceptual  
102 understanding is also scalable. However, each of the aforementioned confounding  
103 factors aggregate over multiple orders of magnitude often masking the very  
104 relationships we seek to elucidate (Figure 1b). To formalize how measurable  
105 microbiome characteristics are linked with system-level processes we have  
106 conceptually defined the intersection of microbial and ecosystem ecology and  
107 identified three categories of microbial characteristics that illustrate the relationship  
108 among each category of microbial characteristics and how they may contribute to an  
109 ecosystem process (Figure 2).

### 110 ***Mapping Ecosystem Processes to Microbial Characteristics***

111 Ecosystem processes are defined as a qualitative change in a pool or a flux from  
112 one pool to another (e.g.  $\text{NH}_4^+$  to  $\text{NO}_3^-$ , or dissolved organic matter to  $\text{CO}_2$ ). The first  
113 step to understand how the microbiome influences an ecosystem process, is to  
114 define the ecosystem process of interest and each sub-process that contributes to it  
115 (i.e., the set of constituent reactions that combine to determine net flux). Few, if any  
116 ecosystem processes involve a single metabolic pathway, or are carried out by a  
117 single organism (e.g., a notable exception being the recent discovery of commamox  
118 which can independently carry out nitrification, the conversion of  $\text{NH}_4^+$  to  $\text{NO}_3^-$ )<sup>10,11</sup>.  
119 Rather, ecosystem processes are composites of complementary or antagonistic sub-  
120 processes, carried out by phylogenetically and metabolically diverse  
121 microorganisms<sup>12</sup>. For example, net ecosystem productivity (NEP) is the balance  
122 between antagonistic processes of C-fixation and C-mineralization. Each sub-  
123 process of NEP can be further partitioned into a series of metabolic pathways (e.g.,  
124 chemoautotrophic nitrification and photoautotrophic C-fixation or heterotrophic  
125 fermentation and aerobic respiration). Partitioning each ecosystem process in this

126 hierarchical manner can continue until the sub-processes maps directly to specific  
127 microbial metabolic pathways (e.g., acetoclastic methanogenesis). Subsequently  
128 each of these metabolic pathways can be categorized as either phylogenetically  
129 broad or narrow<sup>13</sup>. Broad processes are phylogenetically common (i.e., widely  
130 distributed among taxa), whereas narrow processes are phylogenetically conserved  
131 (i.e., limited to a specific subset of taxa). For example, denitrification and  
132 photosynthesis are phylogenetically broad processes, while both methanogenesis  
133 and methanotrophy are phylogenetically narrow processes (with at least one notable  
134 exception<sup>14</sup>).

135 The second step is to identify, the controls or constraints on each constituent sub-  
136 process. For example, the kinetics of a single metabolic pathway in a model  
137 organism may help us understand the rate limiting steps of a narrow process, but  
138 insights from model organisms are much less likely to capture the full spectrum of  
139 responses of a broad process where phenotypic variation among phylogenetically  
140 diverse organisms is likely to be much greater<sup>15,16</sup>. Defining the ecosystem process,  
141 its critical sub-processes, and the known phylogenetic distribution of the metabolic  
142 pathways that drive those sub-processes creates an explicit conceptual pathway that  
143 links the ecosystem process to the microorganisms that contribute to it. Once the  
144 ecosystem process has been conceptually partitioned into its component parts and  
145 their primary controls, a concerted approach can be applied to investigate how  
146 characteristics of the microbiome influence the ecosystem process of interest within  
147 the complexity of a natural environment.

#### 148 ***Categories of Microbial Characteristics***

149 We propose that attempts to elucidate the microbial contribution to system-level  
150 processes needs to explicitly identify three distinct categories of microbial

151 characteristics: 1) microbial processes, 2) microbial community properties, and 3)  
152 microbial membership (Figure 2). The contribution of the microbiome to ecosystem  
153 processes is exerted through aggregate community properties that are shaped by  
154 both microbial membership and environmental factors. This proposed framework  
155 allows the researcher to clearly identify how different measurements used to  
156 characterize a microbiome interact with each other and identify the potential of each  
157 characteristic to elucidate the microbial contribution to the system level process.  
158 These categories are hierarchically connected (Fig. 2), but they represent distinct  
159 degrees of aggregation that are not simply an additive function of the previous  
160 category. Furthermore, each category is potentially subject to different modes of  
161 regulation, and, each category has different putative linkages to system-level  
162 processes (Fig. 2). All measurable characteristics of microbial communities (e.g.,  
163 abundance of cells, sequence of genes, transcripts, or proteins; enzyme expression  
164 or activity) can be placed within one of the above categories, but most studies rarely  
165 articulate how these measurements differ in their specificity (i.e., the level of  
166 phylogenetic resolution at which they are applied), precision (i.e., the ability of the  
167 method to repeatedly describe the characteristic of interest), or context (i.e., how a  
168 characteristic relates to other characteristics or the ecosystem within which they  
169 were measured). This conceptual structure that orientates each microbial category  
170 within a broader context creates the opportunity to improve the design of  
171 observational and experimental studies in microbiome research.

172 *Microbial Processes* - Microbial processes are the collective metabolism of the  
173 microbiome that contribute to changes in pools and fluxes of elements or compounds  
174 (i.e., Figure 2, Letter K). This is the level of microbial information that can most  
175 readily be incorporated into system-level models because many microbial processes



176 represent the key sub-processes that contribute to a particular ecosystem process  
177 (e.g., methanogenesis + methanotrophy  $\approx$  methane efflux). Commonly measured  
178 microbial processes in ecosystem science include nitrogen fixation, denitrification,  
179 nitrification, phosphorus uptake and immobilization, carbon fixation, and organic  
180 carbon mineralization. The rates of many microbial processes can be approximated  
181 through physiological assays (e.g., biological oxygen demand to estimate microbial  
182 community respiration), and while they do not open the “black box” of the microbial  
183 community, they do directly quantify the microbial contribution (or at least the  
184 potential contribution) to changes in resources moving through the box. Microbial  
185 processes can be distinguished from other microbial characteristics because they  
186 are all rates (i.e., have time in the denominator) and require a bioassay to estimate.

187 Assays used to estimate microbial processes are often logistically challenging,  
188 require manipulations that inevitably deviate from the *in situ* conditions, and often  
189 depend on the environment from which the microbiome was sampled. For example,  
190 the relationship between temperature and microbial processes such as enzyme  
191 activity and phosphorus use efficiency (PUE) vary across latitudinal gradients<sup>17</sup> and  
192 among seasons<sup>18</sup>. Thus, observations of the effect of temperature on either enzyme  
193 activity or PUE depend on where (e.g., at what latitude) and when (e.g., during which  
194 season) they were measured. In the absence of an understanding of the underlying  
195 physiological mechanism (e.g., the physiological change that allows a community to  
196 perform differently at different temperatures), the relationship between and  
197 environmental driver (e.g., temperature) and a microbial process must be measured  
198 through a direct assay at each location and at each time. This limits the inference  
199 possible from relying only on measurements of microbial processes alone.

200 *Microbial Community Properties* - Microbial community properties include a broad  
201 set of microbial characteristics such as community biomass or biomass elemental  
202 ratios (e.g., biomass C:N or C:P ratios) and the majority of phylogenetically  
203 undifferentiated aggregate sequence based measurements (e.g., gene abundance,  
204 metagenomes, transcriptomes). Microbial community properties (Figure 2) represent  
205 an integrated characteristic of the microbiome that has the potential to predict or at  
206 least constrain the estimates of microbial processes. For example, microbial  
207 community biomass C:N (a community property) indicates a microbiome's potential  
208 to mineralize or immobilize N.<sup>19</sup> Community biomass stoichiometry has been shown  
209 to be a useful predictor of nutrient immobilization or mineralization during litter  
210 decomposition<sup>19</sup>, and in soils can predict both respiration and N-mineralization better  
211 than microbial biomass alone<sup>20</sup>. The power of biomass elemental ratios to explain  
212 nutrient cycling has also been shown in freshwater<sup>21</sup> and marine ecosystems<sup>22</sup>  
213 including the seminal paper that demonstrated the similarities between the  
214 stoichiometry of marine algal biomass and that of the dissolved fraction of nutrients  
215 in the ocean<sup>23</sup>.

216 Microbial community properties can be separated into two categories, emergent  
217 properties (EPs) and community aggregated traits (CATs). Emergent properties have  
218 been used to refer to a variety of phenomena in ecology, however here we use  
219 emergent properties as it has been defined by Salt (1979)<sup>24</sup>: "An emergent property  
220 of an ecological unit is one which is wholly unpredictable from observation of the  
221 components of that unit", which is consistent with its contemporary use in microbial  
222 ecology.<sup>25</sup> For example, the potential importance of emergent properties to influence  
223 ecosystem processes has been demonstrated in a series of experimental flow-  
224 through flumes that mimicked development and metabolism of stream biofilms<sup>26</sup>.

225 Transient storage (i.e., an increase in residence time of the water and its solutes  
226 near the biofilm relative to the flow around it) increased as the microbial biofilm  
227 density increased<sup>26</sup>. Microbial biofilm formation is an EP<sup>27</sup> that affects the important  
228 ecosystem process of hydrological transient storage<sup>26</sup>. Another example of an  
229 emergent property is the relative abundance of a certain traits within a microbiome.  
230 Trait based approaches have a rich history in ecology and have been increasingly  
231 applied to address questions in multiple areas of microbial ecology.<sup>28</sup> For example,  
232 specific functions (i.e. uptake of an individual organic substrate) are associated with  
233 traits which can involve multiple genes, among different taxa, all capable of  
234 performing the function albeit with differences in the underlying physiology  
235 and efficiency. The distribution and expression of these functional gene variants  
236 generates a trait structure among microbiomes, which determines the overall  
237 performance of the microbiome for that given function (i.e. uptake of a given organic  
238 substrate), but which cannot be predicted simply from the presence of the taxa that  
239 carry the genes conferring that trait.<sup>29</sup> While characterization of EPs may improve the  
240 understanding of microbial processes (Figure 2, Letter G) they cannot, in principle,  
241 be estimated or predicted on the basis of the constituent taxa (i.e. membership)  
242 alone (Figure 2, Letter F), and thus must remain as an intermediary between  
243 environmental drivers (Figure 2, Letter C) and microbial processes (Figure 2, Letter  
244 G).

245 Unlike EPs, CATs can potentially be estimated from characteristics of their  
246 constituents and provide a pathway to link microbial community membership to the  
247 community properties that drive microbial processes (Figure 2, Letter E)<sup>30</sup>. For  
248 example, CATs may include commonly measured community properties such as  
249 functional gene abundance as estimated from qPCR (e.g., *pmoA* which encodes a

250 subunit of the enzyme involved in methane oxidation, can be used to estimate  
251 potential for methanotrophy and as a phylogenetic marker for methanotrophs)<sup>31</sup>. A  
252 recent perspective article discussed the role of CATs in microbial ecology and noted  
253 a series of additional CATs (e.g., maximum growth rate, dormancy, osmoregulation)  
254 that could be inferred from metagenomic data of the extant community<sup>30</sup>.

255 Understanding when, and which, community properties that shape microbial  
256 processes can be predicted by membership is a critical research question, and an  
257 important step in understanding how the microbiome contributes to system level  
258 processes. Whether or not a community property is an EP or a CAT is an exciting  
259 area of research and provides an important framework to advance research at the  
260 microbial-ecosystem nexus. New approaches, like studying higher-level interactions  
261 in ecological communities could help understand how a microbiome's constituents  
262 interact to form emergent properties.<sup>32</sup> This is not a trivial task, yet a suite of existing  
263 methods, discussed below, already provides the ability to directly pursue this  
264 challenge.

265 *Microbial Community Membership* - Although the now commonplace analysis of  
266 community membership by sequencing phylogenetic marker genes (e.g., regions of  
267 the 16S, 18S, or ITS genes) or suites of phylogenetically conserved protein  
268 sequences identifies constituent microbial taxa, the direct coupling of microbial  
269 phylogeny to physiology and ecology remains elusive (Figure 2, Letter H).<sup>33,34,35</sup> In  
270 general the paucity of associated physiological data or information on population  
271 phenotypes that accompany phylogenetic sequence data limits the system-level  
272 inference that is possible from analyses of community membership. This constrains  
273 our ability to attribute microbial processes to community membership of even  
274 relatively simple environmental consortia. Whereas it is clear that microbial

275 populations are not randomly distributed in space and time<sup>35</sup>, and that some  
276 microbial traits are conserved at coarse taxonomic scales<sup>28,36,37</sup>, the physiological  
277 mechanisms underlying non-random distributions of microbial taxa across  
278 environmental gradients is often unknown. The limited understanding of the  
279 metabolism of most bacterial phyla limits an explicit understanding between the  
280 organism's abundance and its role in the microbial process that contributes to an  
281 ecosystem process.

### 282 ***A Path Forward***

283 We suggest that a challenging but necessary step for microbiome science is to  
284 move away from identifying correlative relationships between characteristics of the  
285 microbiome and system level processes, and towards identifying more causative and  
286 mechanistic relationships. The conceptual diagram (Figure 2) is a road map to  
287 organize and link the diverse suite of measureable microbial characteristics that are  
288 currently available to researchers. Figure 2 does not represent how these  
289 components necessarily interact in the environment; rather it is a map that identifies  
290 potential links between measureable microbial and system-level characteristics that  
291 can help structure our exploration of how microorganisms influence the systems they  
292 inhabit. Ecosystem ecology has traditionally been confined to interactions between  
293 environmental parameters and ecosystem processes (depicted within the horizontal  
294 arrow, Figure 2). Similarly, microbial ecology (depicted within the vertical arrow,  
295 Figure 2) has historically focused on phylogenetically undefined aspects of microbial  
296 communities (e.g., bacterial abundance) and microbial processes (e.g., bacterial  
297 production) or on the physiology of microbial isolates (e.g., sulfate reducing bacteria)  
298 or the collective physiology of highly reduced communities with known membership  
299 (e.g., waste water treatment microbiome). The routine inclusion of sequence-based

300 approaches in studies of environmental microorganisms has lead to an increasingly  
301 detailed description of the world's microbiomes and an increasing interest in how  
302 constituents of those communities interact to influence the system as a whole.

303 The drive to include microbial characteristics into system-level science has led to a  
304 range of approaches for linking characteristics of the microbiome to ecosystem  
305 processes. Direct connections between microbial membership and ecosystem  
306 processes (Figure 2, Letter I), or community properties and ecosystem processes  
307 (Figure 2, Letter J), have proven consistently difficult to establish<sup>3,4</sup>. We propose 1)  
308 identifying which microbial processes are likely to contribute to ecosystem-level  
309 pools and fluxes *a priori* (Figure 2, Letter K), 2) determining which microbial  
310 community properties best describe and predict these microbial processes (Figure 2,  
311 Letter G), and 3) identifying whether the community properties that best describe  
312 each process are a CAT or an EP (Community Properties, Figure 2). If the  
313 community property is a CAT then exploring the link between microbial membership  
314 and community properties may lead to further understanding and perhaps an  
315 enhanced predictive power (Figure 2, Letter E). However, if the community property  
316 is an EP elucidating the microbial membership that contributes to the EP is unlikely  
317 to improve understanding of the drivers of that community property (Figure 2, Letter  
318 F) and understanding how environmental drivers structure the EP will be more  
319 insightful. Formalizing microbiome research into a structured, conceptual framework  
320 will allow the research community to better focus on potential links between  
321 microbiome characteristics and system-level processes that are most likely to be  
322 detected empirically. This approach will also allow researchers working in different  
323 systems to test the same pathways among defined microbiome characteristics and  
324 thus increase the possibility of understanding the casual mechanism (or absence of

325 causality) for observed correlations. Equally as important, we suggest that  
326 attempting to link community membership (Figure 2, Letter I) or community  
327 properties (Figure 2, Letter J) directly to ecosystem processes is by definition  
328 correlative and therefore a less powerful approach to integrating microbiome  
329 characteristics into system-level science. Thus future research endeavors will be  
330 most powerful if they focus on elucidating connections through the complete path of  
331 microbial ecology (Figure 2, blue arrow, Letters E, F, and G) and not direct  
332 connections between microbial membership or community properties and ecosystem  
333 processes (Figure 2, Letter I and J).

### 334 ***Applying and Testing the Proposed Framework***

335 Applying and testing the proposed framework will depend on the ability to more  
336 robustly evaluate each category of microbial characteristics and to directly measure  
337 the arrows that connect each category (Figure 2). Both labeling/sorting approaches  
338 and phenotypic description of isolates provide an opportunity to better understand  
339 how microbial membership contributes to community properties (Figure 2, Letter E or  
340 F). Labeling and cell sorting approaches (e.g., fluorescent in situ hybridization (FISH)  
341 coupled with flow cytometry cell sorting<sup>38</sup>, or immunocapture such as with  
342 bromodeoxyuridine, BrdU)<sup>39</sup> provide powerful tools to constrain the complexity of the  
343 microbiome and directly test hypotheses that link membership to community  
344 properties or microbial processes. Labeling and sorting techniques allow the cells  
345 that can be targeted with a stain or other label to be separated from the broader  
346 community and then assayed for membership or phenotypes such as activity or  
347 biomass composition. For example, a study of an Arctic Ocean bacterial community  
348 labeled the actively growing component of the community using BrdU and then  
349 separated those populations from the rest of the community using an immunocapture

350 technique to better understand the portion of the microbiome that was driving  
351 community dynamics<sup>39</sup>. By simplifying the community, researchers were able to link  
352 membership to secondary production (a microbial process, Figure 2, Letter H) and  
353 begin to better understand which members of the community were contributing to  
354 secondary production.

355 In addition, physiological studies of isolates from a broader distribution of  
356 representative phyla are key to advancing our understanding of how membership  
357 contributes to community properties (Figure 2, Letter E). Finding isolates that are  
358 representative of important community properties has the potential to better  
359 understand phenotypic plasticity and how constituent populations do (CATs) or do  
360 not (EPs) contribute to a community property<sup>15</sup>. For example, work on the marine  
361 bacterioplankton SAR11 has led to an increased understanding of how this  
362 ubiquitous member of the marine microbiome interacts with elemental cycles in the  
363 open ocean<sup>40</sup>. Similarly, a rich body of work on multiple isolates of the comparably  
364 ubiquitous photoautotroph *Prochlorococcus* has advanced our understanding of the  
365 ecology and physiology of one of the most abundant phototrophs on the planet<sup>41</sup>.  
366 Detailed studies of isolates of common environmental OTUs have clearly  
367 demonstrated immense variation within a given OTU (i.e., “microdiversity”) that in  
368 part explains the challenge of linking membership to a community property<sup>16</sup>. For  
369 example, work on *Prochlorococcus* has led to a better understanding of how  
370 ecotypes within a single taxonomic unit (OTU) can lead to specialization in  
371 temperature and substrate affinity<sup>41</sup>. OTUs that form a substantial portion of the  
372 microbiome’s sequence abundance provide potential candidates for further  
373 investigation of possible phenotypic plasticity and or microdiversity<sup>16</sup>. For example, a  
374 single phylotype of the class *Spartobacteria* within the phyla *Verrucomicrobia* was



375 found to be present in a broad range of soil ecosystems and comprised as much as  
376 31% of all 16S sequences returned from prairie soils<sup>42</sup>, making it an excellent  
377 candidate for targeted isolation and physiological studies. Whereas it is challenging  
378 to isolate and culture many microorganisms from the environment, existing  
379 approaches to isolation have been led to successful isolation of both abundant and  
380 rare members of environmental microbiomes. A recent study isolated members of an  
381 apple orchard soil microbiome where most isolates were from the least abundant  
382 members of the community<sup>43</sup>. Previous studies have had success isolating members  
383 of the pelagic marine microbiome by using filter-sterilized seawater with a dilution to  
384 extinction approach<sup>44</sup>. Thus there is a potential to target both abundant<sup>40,41</sup> and rare<sup>43</sup>  
385 members of diverse microbiomes to learn more about their influence on community  
386 level properties. Studies of environmental isolates are essential in building a broader  
387 understanding of how community membership does or does not contribute to  
388 community properties (H2, Letter E and F).

389 The most commonly measured microbial characteristics can be associated with  
390 one of the three categories of microbial characteristics defined here. However, the  
391 key to moving from a correlative and descriptive approach to a causative and  
392 mechanistic approach comes in measuring the arrows represented by letters in  
393 Figure 2. There is a suite of powerful methods already being employed in microbial  
394 ecology that can actively measure many of the arrows illustrated in Figure 2. These  
395 include: stable isotope probing of mixed communities<sup>45</sup>, single cell methods that can  
396 assay cells in the physiological state they occur in in the environment<sup>46</sup>, and labeling  
397 individual cells with stable isotopes for single cell analyses<sup>46</sup>. Studies that use stable  
398 isotope probing or any form of tracking stable isotopes into a population have been  
399 successful in linking microbial membership to microbial processes (Figure 2, Letter

400 H). For example, a study of sulfate reduction in a Scottish peatland using SIP  
401 revealed that a single species of *Desulfosporosinus* was most likely responsible for  
402 the totality of sulfate reduction within the peatland even though it only comprised  
403 0.006% of the retrieved 16S rRNA gene sequences<sup>47</sup>. In this case, the  
404 *Desulfosporosinus* species represented the only known sulfate reducer within the  
405 community and thus the kinetics of this organism seemingly defined the kinetics of  
406 sulfate reduction for the entire system. Whereas this is a single example of using SIP  
407 to link microbial membership to microbial processes (Figure 2, Letter H), there is a  
408 suite of culture-free techniques (such as Raman microspectroscopy (MS),  
409 NanoSIMS, or X-ray microanalysis, XRMA) that complement sequence-based  
410 microbiome analyses by reporting on the physiological and phenotypic  
411 characteristics of individual cells *in situ*<sup>46,48,49</sup>. For example, Raman MS has the  
412 ability to elucidate the macromolecular composition of uncultured individual cells *in*  
413 *situ*<sup>46</sup>. Incorporation of stable isotopes into a cell's macromolecules can be visualized  
414 as a shift in the Raman spectra. This provides information not only on which cells are  
415 incorporating the substrate but what macromolecular pool those substrates are being  
416 allocated to. Similarly, NanoSims allows for visualization of isotopes within a cell,  
417 and while it cannot be used to identify which macromolecule pool an isotopic label  
418 went into, it can visualize which cells are actively incorporating isotopically labeled  
419 substrates. Both Raman and NanoSims can be coupled with a range of *in situ*  
420 hybridization techniques (e.g., fluorescent *in situ* hybridization, FISH) to identify  
421 which populations are contributing to community properties (Figure 2, Letter E) or  
422 microbial processes (Figure 2, Letter H). For example, the study of a microbial  
423 consortia from the Sippewissett Salt Marsh on the coast of Massachusetts, USA  
424 used a combination of FISH and NanoSIMS to confirm a syntrophic association

425 between a population of autotrophic purple sulfur bacteria and heterotrophic sulfate  
426 reducing bacteria (SRB)<sup>50</sup>. Whereas several pieces of evidence pointed to a  
427 syntrophic association, the authors confirmed the association by first using FISH to  
428 visualize the physical association of each population. Further analysis with  
429 NanoSIMS after incubation with <sup>34</sup>S enriched sulfate and <sup>14</sup>C enriched bicarbonate  
430 confirmed the presence of carbon fixation by the purple photosynthetic bacteria with  
431 sulfide as the electron donor was coupled to the reduction of sulfate by the SRB.  
432 Using existing methods of confirmatory ecophysiology allows for direct  
433 measurements of the arrows connecting membership with microbial processes in a  
434 stable microbial consortia (in this case both carbon fixation and sulfate reduction,  
435 Figure 2, Letter H). These approaches applied in concert with sequence-based  
436 analyses have the potential to empirically link the categories of microbial information  
437 defined here (Figure 2), moving microbiome science from a descriptive and  
438 correlative approach to a mechanistic and causative approach.

439 These culture free approaches also create the potential to begin to determine  
440 which community properties are EPs, and which are CATs. For example, microbial  
441 community biomass stoichiometry (e.g., biomass C:N or C:P) cannot currently be  
442 predicted (or even constrained) from a list of it's constituent taxa (Figure 2, Letter F).  
443 However, microbial biomass stoichiometry is a community property with power to  
444 predict the microbial contribution to nutrient cycling (Figure 2, Letter G).<sup>19-22</sup>  
445 Independently, the biomass stoichiometry of microbial isolates grown on the same  
446 media has been shown to differ among different taxa suggesting a relationship  
447 between an organisms' identity and the elemental composition of its biomass.<sup>51,52</sup>  
448 However, there is an abundance of evidence that suggests that the biomass  
449 stoichiometry of many isolates is a function of the stoichiometry of the media they

450 were cultured on<sup>52</sup>. Electron dispersal spectroscopy (EDS) has the power to  
451 measure the C:N:P of individual bacterial cells growing *in situ* (i.e. not in culture)<sup>49</sup>.  
452 The potential to couple EDS analysis with a phylogenetic label presents the  
453 opportunity to assay mixed microbial communities and assess the link between  
454 phylogenetic identity and biomass stoichiometry under natural conditions<sup>53</sup>. Thus,  
455 community biomass stoichiometry can potentially be deconstructed into the biomass  
456 stoichiometry of its constituent taxa<sup>53</sup>. This approach would provide a direct link  
457 between community membership and a community property (e.g., biomass C:N,  
458 Figure 2, Letter E), that influences an important microbial process (i.e. nutrient  
459 recycling).

#### 460 ***Designing microbiome research***

461 It is critical that we recognize the influence of the taxonomic and functional  
462 composition of the microbiome is exerted through multiple pathways, some that are  
463 direct and can be readily identified, some that are indirect, and mediated by complex  
464 interactions at the community level. We must also recognize that the influence of  
465 microbiomes will vary depending on the system-level process in question, because  
466 analysis of microbial characteristics may simply not improve the environmentally-  
467 based prediction of certain processes (Figure 2, Letter A), whereas other system-  
468 level processes may indeed benefit from the inclusion of microbial characteristics  
469 including membership. For the latter, the challenge then is to determine which  
470 microbial category is the most relevant predictor of the system level process of  
471 interest: microbial processes, community properties, or microbial membership.  
472 Establishing the links between these microbial dimensions (Figure 2, Letters E, F,  
473 and G) further contributes to our understanding of the mechanistic underpinnings

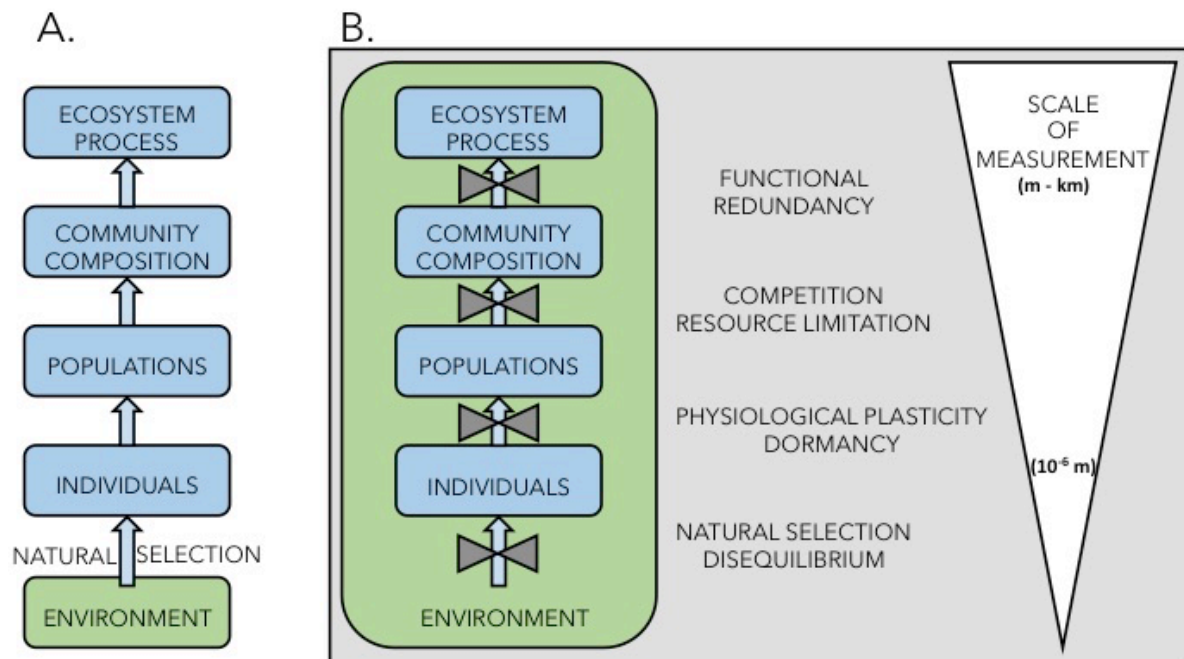
474 that affect system level processes and thus will have greater explanatory power in a  
475 broader range of systems.

476 The framework presented here provides one approach to formalize inquiry across  
477 microbiome science and encourages empirical linkages between the presence of  
478 organisms in a system and the processes that characterize that system. Whereas we  
479 draw examples from environmental microbiomes and the ecosystems they inhabit,  
480 this structured approach has the potential to benefit the analysis of microbiomes  
481 associated with other systems such as host organisms and those of the built  
482 environment. As important as establishing causal links among microbial  
483 membership, community properties, microbial processes, and ecosystem processes,  
484 is determining when these links are unlikely to be present. Research that  
485 indiscriminately seeks to identify correlations, which does not recognize the  
486 hierarchy of effects, and that places all metrics on an equal plane are susceptible to  
487 confirmation bias and will continue to yield conflicting and ambiguous results that not  
488 only fail to provide new insight into ecosystem processes, but also blurs the  
489 connections that do exist. We suggest that rather than looking for linkages among  
490 microbiome membership and system-level processes in every study, research efforts  
491 would benefit from strategically targeting the linkages and processes for which an *a*  
492 *priori* understanding of microbial physiology should allow us to improve our  
493 understanding of the ecosystem process.

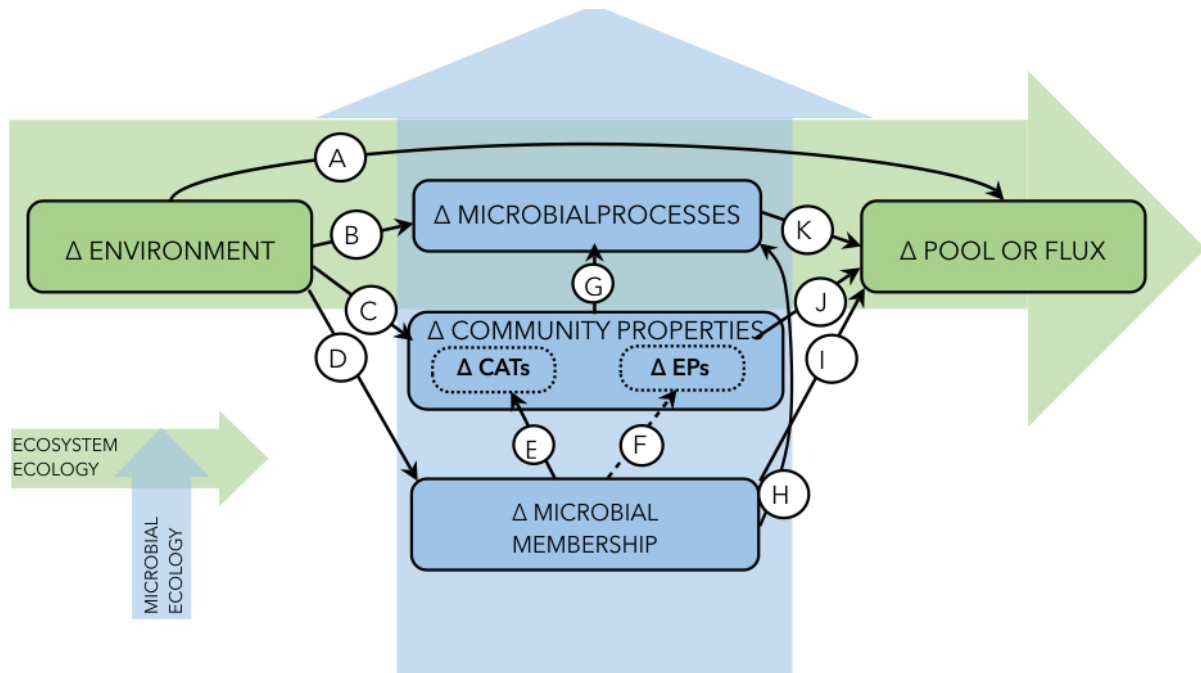
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## Figures



**Figure 1** Diagram of microbial-ecosystem linkages A) how linkages are commonly conceptualized across levels of ecological organization and B) the series of ecological phenomena that create challenges when attempting to link metrics from one level of ecological organization to the other.



**Figure 2** Shown is a conceptual map of the intersection between microbial (vertical) and ecosystem (horizontal) ecology with each of the three categories of microbial characteristics (microbial processes, community properties, and microbial membership) as defined in the text. We argue for an increased focus on studies that elucidate pathways E, F, and G. In addition, we note that pathways I and J are less likely to effectively incorporate microbiome characteristics into system-level science. The delta symbol in each category indicates an emphasis on how changes within a category may lead to a change in a connected category. The dotted arrow for letter F denotes that many emergent properties cannot currently be linked to membership and is an important area for active research.



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