

Microbial and Metabolic Succession on Common Building Materials Under High Humidity Conditions

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1 ABSTRACT

2 Despite considerable efforts to characterize the ecology of bacteria and fungi in the built
3 environment (BE), the metabolic mechanisms underpinning their colonization and successional
4 dynamics remain unclear. Here, we applied bacterial/viral particle counting, qPCR, 16S and ITS
5 rRNA amplicon sequencing, and metabolomics to longitudinally characterize the ecological
6 dynamics of four commonly used building materials maintained at high humidity conditions
7 (~94% RH). We varied the natural inoculum provided to each material by placing them in different
8 occupied spaces, and we wet the surface of half of the samples of each material to simulate a
9 flooding event. As expected, different materials showed different bacterial and viral particle
10 abundance, with wet materials having higher growth rates and lower alpha diversity compared to
11 non-wetted materials. Wetting described the majority of the variance in bacterial, fungal and
12 metabolite structure, and material type only influenced bacterial and metabolic diversity, while
13 location of inoculation was only weakly associated with bacterial and fungal beta diversity.
14 Metabolites indicative of microbial activity were identified, as were those that were native to the
15 surface material. Glucose-phosphate was abundant on all materials (except mold-free gypsum) and
16 was correlated with *Enterobacteriaceae*, which could indicate a potential bacterial nutrient source.
17 A compound consistent with scopoletin, a plant metabolite with antimicrobial activity, was
18 significantly negatively correlated with *Bacillus* and positively correlated with *Pseudomonas* and
19 enriched in medium density fiberboard (MDF) materials. In wet samples, the alkaloids nigragillin
20 and fumigaclavine C, both with antimicrobial properties, were significantly positively correlated
21 with the fungal phylum Ascomycota. Nigragillin, was also negatively correlated with *Bacillus* and
22 *Pseudomonas* abundance. Thiabendazole and azoxystrobin (anti-fungal compounds) were highly
23 abundant on mold-resistant gypsum wallboard and likely directly influenced the decreased fungal
24 growth observed on this material. The mold-resistant gypsum material also showed a significant
25 increase in bacterial alpha diversity, and bacterial and viral particle abundance, as well as a
26 decrease in metabolite diversity, likely a result of reduced fungal growth. *Penicillium* taxa were
27 positively correlated with thiabendazole, which suggested the persistence of resistant strains. Also,

28 specific to the wet samples, *Bacillus* abundance was positively correlated with the azoxystrobin,
29 suggesting bi-directional competitive adaptation, and positively correlated with metabolites known
30 to interfere with *Pseudomonas* biofilm formation, which could explain the anti-correlation
31 between these taxa. As expected, high moisture conditions enabled faster growth of inoculating
32 microorganisms, whose composition, chemistry, and competition was shaped by surface material,
33 suggesting that both fungal and bacterial growth need to be considered when determining the
34 impact of dampness in built environments.

35

36 INTRODUCTION

37

38 The microbiology of the built environment comprises bacteria, archaea, fungi, viruses and
39 protists, all of which maintain growth potentials under varying physicochemical regimes. Many
40 recent studies of this ecosystem have applied molecular sequencing techniques to characterize
41 microbial community relationships and dynamics under varying occupant density, building type
42 and location, environmental conditions, and material type (Lax *et al.*, 2017; Adams *et al.*, 2016;
43 Chase *et al.*, 2016; Stephens, 2016; Lax *et al.*, 2014). However, most of these studies have
44 investigated communities sampled from relatively dry materials on which microbes are likely
45 biologically inactive unless they experience liquid water or high relative humidity (RH) (Chase *et al.*,
46 2016). It is widely accepted that fungal growth can occur at RH >75-80% and material decay
47 can occur at RH >95%, depending on material (Viitanen *et al.*, 2010, Johansson *et al.*, 2012).

48 Dampness is a fairly common occurrence in buildings, with approximately half of all
49 homes in the U.S. having experienced dampness or mold (IAQ Report - Prevalence of Building
50 Dampness). Building material dampness occurs for different reasons, including: bulk liquid entry
51 from floods, extreme weather events, and plumbing system problems; rain or snow entry through
52 leaks in building envelopes and roofing systems; and high water vapor content resulting from
53 moisture migration through building materials or condensation of warm humid air on cold surfaces
54 (IAQ Report - Nature and Causes of Building Dampness). Dampness and the presence of visible
55 mold have been consistently associated with adverse human health outcomes, including respiratory
56 and allergic effects (Mendell *et al.*, 2011, Quansah *et al.*, 2012, Fisk *et al.*, 2010, Jaakkola *et al.*,
57 2013). Hypotheses to potentially explain these associations include a combination of exposure to
58 specific microbial agents (Institute of Medicine, 2004), varied gene expression and metabolism
59 (Hegarty *et al.*, 2018), and the release of fungal metabolites including mycotoxins (Miller *et al.*,
60 2014) and microbial volatile organic compounds (mVOCs) (Roze *et al.*, 2013).

61 Although fungal growth on building materials has been studied for decades (Hyvärinen *et al.*,
62 2002, Gravesen *et al.*, 1999, Hoang *et al.*, 2010; Pasanen *et al.*, 1992), only a limited number
63 of studies have used molecular techniques to investigate bacterial and fungal growth, microbial
64 community dynamics, and/or metabolic activity on common buildings materials exposed to liquid
65 water and/or high humidity conditions (Coombs *et al.*, 2017). Therefore, we characterized the
66 bacterial and fungal concentration and diversity, as well as the production of microbial metabolites,
67 on samples of four common building materials incubated at ~94% relative humidity: oriented
68 strand board (OSB), medium density fiberboard (MDF), gypsum wallboard, and mold-resistant
69 gypsum wallboard. We varied the BE source of inoculation and purposely wet half of the samples
70 to assess how indoor microbial sources and the presence of liquid water influence community
71 structure and metabolite profiles of these materials over multiple time points. We used several
72 techniques to quantify microbial growth and microbial community composition and functional
73 metabolism including: bacterial and viral like particle counts, image processing of visible mold

74 growth, qPCR, amplicon sequencing of 16S and ITS rRNA marker genes, and metabolomics.
75 Results from these different methods were integrated via co-occurrence network approaches,
76 which provided insights into microbial community organization and environmental interaction.
77 Improved understanding of how bacterial and fungal metabolism is shaped by environmental
78 properties (e.g., the presence of water, surface material composition) and inoculating source (e.g.,
79 building location, occupancy patterns) could have important implications for architectural design,
80 construction, building management, and occupant health (Rand *et al.*, 2017). Therefore,
81 determining the microbial metabolic dynamics in these high RH environments should be an
82 important research priority.

83

84 RESULTS

85

86 *Experimental Setup*

87

88 Our study used four building materials types: oriented strand board (OSB), medium density
89 fiberboard (MDF), regular gypsum wallboard, and mold-resistant (i.e., mold-free, or ‘MF’)
90 gypsum wallboard. Coupons of 5 cm × 5 cm of each material type were naturally inoculated at
91 two different locations for about 50 days each. After the inoculation period concluded, the time
92 that occupants were coming in close proximity was reported with similar values, a 0.16% and
93 0.18% of total time. The material coupons were sampled for off-line biological and chemical
94 analysis at 7 different sampling time points, referred to here as TP0, TP1, TP2, TP3, TP4, TP5,
95 and TP6. The initial samples (TP0) were taken just after retrieving the samples from the field
96 inoculation and represent non-wetted, naturally inoculated samples previously held at normal
97 residential humidity condition. After inoculation, half of each set of materials from each location
98 were submerged in tap water in separate pans for ~12 hours to simulate the process of wetting of
99 building materials due to a flood or leak. Different sampling strategies were tested and after
100 statistical verification all samples of the same type were combined as technical replicates (**Figure**
101 **S1**). Microbial datasets were later rarefied to same sequencing depth: 1,000 reads for bacteria and
102 10,000 reads for fungi. Unfortunately, rarefaction removed all bacterial samples from MDF
103 materials, which had very low read counts. After rarefying the data, a comparison of the control
104 and non-control samples reflected that control samples looked very similar in bacterial and fungi
105 diversity than non-control samples, (mantel ≥ 0.49 and ≥ 0.43 for location 1 and location 2
106 respectively, all with a $p < 1E-05$), perhaps because air could still transmit through the non-
107 hermetic foil cover and microbes from the interior of the wood (not killed with the sterilization)
108 could have found their way to the surface. Highlighting that the coupon itself could still be an
109 important contributor to microbial diversity for all samples. From these results, the control location
110 was treated indistinctly than the other two locations. For more details see Materials and Methods
111 section.

112

113 *Visible growth, particulate counts and qPCR*

114

115 Visible microbial growth occurred much faster and covered a far greater percentage of the surface
116 area on wet tiles than on non-wetted tiles (**Figure 1A**). OSB and MDF had the greatest coverage
117 and fastest growth: all wet OSB and MDF tiles reached at least 50% visible microbial coverage by
118 day 20, while non-wetted tiles of these types reached $< 25\%$ coverage. No growth was ever visible
119 on the mold-resistant gypsum tiles. Epifluorescence microscopy revealed that counts of bacterial

120 like particles (BLP) and viral-like particles (VLPs) calculated on samples TP0 to TP3 were
121 strongly correlated ($R^2 = 0.65$, $p = 2.8e^{-23}$) (**Figure 1B**), with VLP counts statistically lower than
122 bacterial counts in all samples (ANOVA $\leq 10^{-4}$) and in both wet and non-wet conditions (two
123 sided non-parametric t-test $p \leq 0.035$) **Figure 1C**). This is in keeping with previous research that
124 found very low VLP:bacteria ratios in the indoor environment (Prussin *et al.*, 2015, Gibbons *et al.*,
125 2015). In our dataset, the mean VLP-bacteria ratio was 0.86 ± 0.07 , with a minimum of 0.61 and
126 a maximum of 1.02 across all samples.

127
128 While BLP (only estimated for TP0 to TP3 samples) and bacterial qPCR agree that wetted samples
129 had higher counts than non-wetted samples, cell counts inferred from these two methods
130 drastically differ for different material types and over time. Most notably, the MF-gypsum had the
131 greatest BLP counts but also the lowest 16S rRNA qPCR counts (6-fold or more lower than other
132 materials). Moreover, the BLP cell counts were essentially constant over time, while qPCR counts
133 steadily increased, with TP6 being 4-fold greater than TP1 and 209-fold greater than TP0 counts.
134 To further confirm the differences, we calculated the overall correlation between paired bacterial
135 qPCR and BLP counts and the results were not significant, emphasizing different biases for each
136 method.

137
138 For fungal qPCR we observed MF-gypsum had the lowest abundance, while all other materials
139 had a range of 20 to 118-fold increase over MF-gypsum. Wetted samples revealed a 4-fold increase
140 in qPCR read abundance over non-wetted samples. Also, the qPCR read abundance increased
141 steadily over time, in such a way that TP6 was 11-fold greater than TP1 and 750-fold greater than
142 TP0 counts.

143 *Bacterial, fungal and metabolite diversity*

145 The bacterial and fungal communities in our study tended to decrease in diversity over time, as
146 measured by the Shannon Index (Shannon H'), which incorporates both the richness and evenness
147 of the community. Given that our data was rarified to an even depth before analysis, this decrease
148 in diversity is indicative of the increasing relative abundance of certain community members, and
149 suggests the preferential proliferation of certain taxa in the inoculating community. In our 16S
150 dataset, wet samples experienced faster declines in diversity than non-wetted samples, and were
151 significantly lower in diversity at the end of the study than non-wetted samples (**Figure 2A**),
152 suggesting that certain bacterial taxa grew quickly in the wet environment and became dominant
153 within the community. In our ITS dataset, we also observed a faster decline in diversity in wet
154 samples, although wet samples were significantly more diverse than non-wetted samples by the
155 end of the study (**Figure 2B**). The decrease in fungal diversity in wetted samples was not
156 monotonic, with an initially steep decline and a subsequent increase. This may reflect fast growth
157 by a small number of taxa that quickly dominated the community, followed by the growth of other
158 taxa with slower growth rates. Similar patterns are observed when looking at the diversity changes
159 for each individual material (**Figure S2**) with the exception of a lack of bacterial growth for wet
160 MDF samples and reduced bacterial growth on dry OSB after the study was half way completed.
161 In contrast, we observed no significant changes in the metabolic diversity over time for either wet
162 or non-wetted samples (**Figure 2C**).

Microbial Compositional Changes

163 Across all samples, the diversity of bacteria within the community was significantly correlated to
164 the diversity of fungi (Corr = 0.28, $p=0.0003$) (**Figure 3A**). Interestingly, neither bacterial nor
165 fungal diversity was significantly correlated to the metabolite diversity, perhaps because of a
166 narrower range of observed metabolite diversity compared to the taxonomic datasets. We observed
167 striking changes in the relative abundance of certain bacterial (**Figure 3B**) and fungal (**Figure 3C**)
168 genera over time, which were largely dependent on wetting condition. In the 16S rRNA dataset,
169 *Bacillus* almost immediately came to dominate wet samples, with an average relative abundance
170 as high as 50% after the 2nd time point, even though it represented a negligible part of the
171 community at the start of sampling. *Bacillus* abundance also increased in non-wetted samples,
172 although to a much smaller extent. A similar pattern was observed for the genera *Pseudomonas*
173 and *Erwinia*, which also represented a very small fraction of community diversity at the start of
174 sampling but quickly increased in abundance in wet (but not non-wetted) samples. Interestingly, a
175 very large percentage of reads from early time point samples, both wet and non-wet, were of
176 chloroplast origin. In wet samples, the number of chloroplast reads quickly declined as the
177 bacterial genera proliferated. In non-wetted samples, chloroplast read abundance remained high,
178 and dominated the sequencing effort to such an extent that discarding those reads would have
179 dropped the majority of non-wetted samples below the rarefaction depth. While these likely
180 represent residual DNA signatures from the plant material used to construct each tile material, we
181 have chosen to keep them in the analysis. **Figure S3A** shows how this dynamic slightly vary for
182 each different material type.

183 The majority of reads in the ITS dataset that could be taxonomically assigned to a genus belonged
184 to one of two genera: *Eurotium* and *Penicillium*. *Eurotium* abundance was negligible at the
185 beginning of community succession but quickly flourished in non-wetted samples, becoming the
186 most abundant known genus in those samples by time point 2 (**Figure 3C**). By contrast, *Eurotium*
187 did not become abundant across wet samples. *Penicillium* abundance was, on average, consistently
188 higher in wet samples than in non-wetted samples, and its abundance was significantly anti-
189 correlated to *Eurotium* relative abundance (corr = -0.12, $p = 0.033$). These taxa-specific changes
190 were mirrored by community level differentiation, where wet vs. non-wetted tiles of the same
191 material and inoculating location became significantly more dissimilar (Bray Curtis, Spearman's
192 Correlation, $p<0.01$) in both their bacterial and fungal community structure over time (**Figure 3D**).
193 **Figure S3B** shows how this dynamic slightly vary for each different material type.

194

195 *Environmental Factors Associate with Microbial and Metabolite Diversity*

196 We used ANOSIM to calculate the factors significantly correlated with differences in the microbial
197 communities across our three datasets. Bray-Curtis dissimilarity was calculated for the 16S, ITS,
198 and metabolite datasets, and ANOSIM was used to determine whether distances between samples
199 of the same metadata factor (i.e. wetting condition, inoculating location, and material) were
200 significantly lower than distances between samples of different types (**Figure S4**). In our 16S
201 rRNA dataset, wetting condition, location, and material each had a significant impact on bacterial
202 community structure (all $p < 0.0001$ based on 10^5 randomized permutations), with wetting having
203 the most pronounced effect ($R = 0.418$). In general, non-wetted samples tended to be more similar
204 to each other than wet samples were to each other, which is likely due to the dominance of a single

205 chloroplast OTU. Material had a less pronounced effect ($R=0.247$) and location had the least
206 evident effect on bacterial community structure ($R=0.133$).

207 Interestingly, fungal community structure was not significantly described by variance in material,
208 while location had a relatively weak ($R = 0.129$) though highly significant ($p < 0.0001$) association,
209 suggesting that variations in fungal communities that settle on materials (which have been shown
210 to be driven largely by outdoor fungal communities, e.g., Adams *et al.*, 2013) influence community
211 structure upon experiencing wetting and high RH conditions. Wetting condition was by far the
212 most influential factor influencing fungal community structure ($R = 0.564$, $p < 0.0001$), and in
213 contrast to the bacterial data, wet samples were much more similar to each other than were non-
214 wetted samples. Metabolite diversity within the community was also affected by wetting condition
215 ($R = 0.276$, $p < 0.0001$), with non-wetted samples more similar to each other than wet samples.
216 Material also played a significant role in metabolite diversity ($R = 0.231$, $p < 0.0001$), and mold-
217 free gypsum samples were particularly metabolically similar, likely due to the lack of fungal
218 growth and the underlying chemical composition of the material. Inoculating location had no
219 significant effect on the diversity of metabolites despite having a significant effect on both the
220 bacterial and fungal community membership. We visualized sample similarity using non-metric
221 multi-dimensional scaling (NMDS) ordination based on Bray-Curtis dissimilarity (**Figure 4**). We
222 converted material, location, and wetting condition into binary variables (1 = yes, 0 = no), which
223 were fit onto the ordination, keeping only the significant vectors (R^2 values for each vector and
224 their significance is presented in **Table S1**). Visually, both bacterial and fungal beta diversity was
225 more differentiated by wetting condition due to the significant increase in growth on wetted tiles,
226 while metabolites were visually differentiated by both wetting condition and surface material,
227 likely due to the underlying chemistry of the material and then the subsequent metabolic activity
228 of the microbes when tiles were wetted.

229

230 *Bacterial and fungal network co-occurrence*

231 Using SparCC (Friedman & Alm, 2012), an algorithm developed to quantify correlations on
232 microbial compositional data (data that has been subject to rarefaction), and a correlation threshold
233 >0.4 , uncovered co-occurrence patterns between taxa from each kingdom. In the bacterial network
234 (**Figure S5**) three co-occurrence clusters were identified, the *Bacillus* cluster, *Pseudomonas*
235 cluster, and a cluster comprising chloroplasts and mitochondria. As expected these groups
236 correspond with the most abundant taxa. On all wet materials and on all samples of gypsum (both
237 wet and non-wet), 95% of associations between *Bacillus* and *Pseudomonas* were negative
238 correlations (**Figure S6 and S7**). On non-wetted OSB, MDF and MF-Gypsum there were no
239 negative correlations between *Pseudomonas* and *Bacillus*. Interestingly, there is a dramatic
240 increase in the absolute number of significant co-occurrence relationships between bacterial OTUs
241 in wet (74) versus non-wetted samples (48), which is a 54% increase in the number of edges. In
242 the fungal correlation network, *Penicillium* OTUs co-occurred with many unknown fungal genera,
243 while OTUs corresponding to *Aspergillus* and its subset, *Eurotium*, maintained monophyletic
244 clusters (**Figure S8**). As with the bacterial co-occurrence networks, fungal OTUs associated with
245 wet tiles had negative correlations among each other, although the number was much smaller than
246 for bacteria. Only 7 fungal OTUs were negatively correlated on wet materials, mainly between
247 unknown genera and an abundant *Penicillium* OTU (**Figure S9**). Strikingly, unlike bacteria, the
248 absolute number of significant co-occurrence relationships between fungal OTUs declined in wet

249 (555) versus non-wetted samples (1,133), which was a 104% decrease in the number of edges,
250 suggesting an inverse co-abundance response between bacteria and fungi during growth.

251 To better understand the co-associations between bacteria and fungi, 16S and ITS OTUs were co-
252 correlated in a single network. A random walk-based method uncovered four distinct modules
253 within the network, with a modularity of 0.45 (**Figure 5A**). In general, the taxa present in each
254 sample tended to cluster within an individual network module (median sample association to
255 module = 0.88). We correlated various metadata factors to module membership (**Figure 5B**) and
256 observed that wetting condition had a significant impact on which samples dominated each
257 module: modules 1 and 3 were associated with wet samples, while modules 2 and 4 were associated
258 with non-wetted samples (**Figure 5C**). Location 1 samples dominated module 3, while Location 2
259 samples dominated module 1 (**Figure 5D**). Overall, wetting condition appears to be the most
260 important factor driving community succession, resulting in two different community structures
261 even when the source community is identical. We also visualized the nodes that were assigned to
262 the genera previously discussed in Figure 3. Nodes in the bacterial genera *Bacillus*, *Pseudomonas*,
263 and *Erwinia*, as well as the fungal genus *Penicillium*, were nearly exclusively enriched in the two
264 wet-associated modules (1 and 3), while chloroplast reads and *Eurotium* nodes all clustered within
265 the non-wetted modules (2 and 4; **Figure 5E**).

266 *Metabolite network co-occurrence*

267 A co-occurrence network correlation was calculated for the sample metabolite profiles (**Figure 6**).
268 As these data are not compositional, we built this network using significantly positive Spearman
269 correlations between nodes and included only the 1,000 most abundant metabolites in the dataset.
270 This resulted in a network with 149,316 edges (density = 0.30) when the significance threshold
271 (alpha) was set to 0.001. Using the same module discovery method described above, we uncovered
272 7 distinct modules (modularity = 0.32), excluding 12 metabolites around the periphery of the
273 network that clustered into modules of <5 nodes. Three modules (3, 4, and 7) were significantly
274 correlated with wet samples, while modules 1, 2, 5 and 6 were associated with non-wetted samples.
275 There was almost no correlation between network modules and inoculating locations, further
276 suggesting that while location (and hence the primary inoculating microbes) may influence
277 community taxonomic diversity, it does not appear to strongly affect metabolic diversity during
278 growth. The abundance of metabolites in module 7 were anti-correlated with all other modules,
279 but specifically with module 2 (corr = -0.87, $p < 0.001$). Module 7 is dominated by wet samples at
280 later time points, suggesting that community succession in wet environments may converge to a
281 common metabolic profile, which is wholly distinct from the non-wetted samples in module 2.

282 *Metabolite features can predict sample type*

283
284
285 Random Forest analysis was employed to determine the metabolites associated with various
286 sample types. Models classifying whether a sample was wet had an average accuracy of 98% (error
287 ratio = 25, with expected random error 0.5), and wet-samples were never misclassified as a non-
288 wetted sample in any of the 10 model iterations. Models classifying samples based on material
289 were similarly successful, with an average accuracy of 97% (error ratio = 25, with expected
290 random error 0.75). Metabolomics models were much less successful at predicting the inoculating
291 location, with a mean success of 72% (error ratio = 2.36 with expected random error 0.67). We
292 sought to gain insight into the chemical composition of metabolites that comprise the signatures

293 observed in these models. Feature importance scores were assigned to compounds based on their
294 relative contributions to predicting sample type. For both the wetting condition dependent and
295 material dependent groups, we selected the 100 highest-scoring metabolite features for further
296 examination and chemical identification (**Figure S10**). Wet samples were enriched with 98 of the
297 100 top-scoring metabolites that differentiated wet and non-wetted. None of these compounds
298 were automatically identified by mzCloud, so the metabolites were analyzed via external database
299 searches, and compound classes were designated based on fragmentation spectra. A diverse set of
300 compound families was observed, including compounds likely to be carbohydrates and
301 glycoconjugates, fatty acids, prenol lipids, sterol lipids, polyketides, and glycerolipids, as well as
302 several pyridine derivatives including a form of vitamin B6, indicative of microbial activity and
303 growth and compounds associated with the surface materials.

304
305 Metabolites that were highly enriched in wet vs non-wetted conditions underwent additional
306 manual analysis for confident structural identification. One of these metabolites was identified as
307 Nigragillin ($C_{13}H_{22}N_2O$, accurate mass = 222.1723), which is a fungal alkaloid first identified in
308 *Aspergillus niger* (Isogai *et al.*, 1975). Nigragillin abundance was significantly enriched in wet
309 MDF and OSB samples (505- and 280-fold, respectively) compared to non-wetted samples.
310 However, no significant differences in nigragillin were observed for gypsum or MF-gypsum. In
311 both wet MDF and OSB the nigragillin concentration increases over time (**Figure S11**). Another
312 high-scoring metabolite showed MS/MS fragmentation consistent with Fumigaclavine C
313 ($C_{23}H_{30}N_2O_2$, accurate mass = 366.2291), which is a fungal alkaloid first identified in *Aspergillus*
314 *fumigatus* (Cole *et al.*, 1977). Fumigaclavine C was enriched in wet samples of gypsum, MDF,
315 and OSB (23-, 26-, and 13-fold increase in comparison to non-wetted samples, respectively), with
316 equivalent abundance in mold-free gypsum regardless of wetting. While the concentration of
317 Fumigaclavine C remained flat or increased slowly in most materials, wet gypsum showed a
318 dramatic increase in abundance at TP3 and 4 (**Figure S11**).

319
320 Metabolites that were predictive of material type (OSB, MDF, Gypsum and MF-Gypsum) were
321 also further analyzed to determine how these materials influence the chemical composition of
322 metabolites. Of these metabolites, 80% eluted with a retention time of >7 minutes, indicating a
323 skew toward more hydrophobic compounds. This suggests that hydrophobic compounds are more
324 diverse between the materials and therefore could have greater influence on microbial metabolism
325 than the ubiquitous hydrophilic components. Two of these metabolites were identified by MzCloud
326 search: glucose-phosphate, which was about 10-fold less abundant in MF-gypsum compared to all
327 other materials, and scopoletin, a metabolite produced by the plants with antimicrobial activity
328 Lerat *et al.*, 2009, Gnonlonfin *et al.*, 2012, Nascimento *et al.*, 2013), which was about 60-fold more
329 abundant in MDF samples than in other materials and could be influencing the reduced bacterial
330 growth on this material (**Figure S11**). Thiabendazole and azoxystrobin, known anti-fungal
331 compounds (Clausen & Yang, 2007, Balba, 2007), were highly overrepresented on MF-Gypsum,
332 333 and 595-fold respectively more abundant than the average content for the other three materials,
333 and as such are likely some of the active compounds in MF-Gypsum.

334 *Microbe-metabolite co-occurrences*

335
336
337 The abundances of Nigragillin and Fumigaclavine C were each significantly positively correlated
338 with a fungal OTU annotated to the phylum Ascomycota (corr = 0.66, FDR p = 0.0004), which

339 contains species known to produce such alkaloids. Both Nigragillin and Fumigaclavine C have
340 been reported to display antibacterial activity (Magdy *et al.*, 2017 & Pinheiro *et al.*, 2013).
341 Interestingly, Nigragillin was negatively correlated with the abundance of *Bacillus* and
342 *Pseudomonas* OTUs; this could suggest fungal competition for space and resources (Mille-
343 Lindblom *et al.*, 2006) against bacteria, and in the specific case of MDF, when Nigragillin
344 abundance was greatest no bacterial growth was detected (**Figure S2, Figure S11**). The abundance
345 of glucose-phosphate was significantly correlated to the proportion of a dominant
346 Enterobacteriaceae OTU, a genus which is known to synthesize it (Herter *et al.*, 2006) (corr = 0.72,
347 FDR p = 0.000002). Thiabendazole was positively correlated with *Penicillium* abundance (corr =
348 0.80, FDR p < 10⁻⁹). As thiabendazole is prevalent and persistent in the natural environment, this
349 correlation may indicate the presence of thiabendazole-resistant *Penicillium* strains colonizing the
350 material from the built environment (Holmes & Eckert, 1999).

351
352 Co-occurrence networks were constructed between the bacterial OTUs and metabolites (SparCC
353 correlation of >0.4; **Figure S12**) to explore further specific microbe-metabolite associations and
354 possible mechanistic interactions. Significant correlations were observed between *Bacillus* OTUs
355 and lipids including fatty acids and monoacylglycerophosphocholine compounds, which are likely
356 to indicate cell wall and biofilm formation (Diomande *et al.*, 2015, Dubois-Brissonnet *et al.*, 2016).
357 Interestingly, *Bacillus* OTUs were also positively correlated to other lipid classes including diols
358 and flavonoids, which have all been observed to interfere with *Pseudomonas* biofilm formation
359 (Kong *et al.*, 2014; Jensen *et al.*, 2014). In addition, the abundance of two fatty acids, one diol,
360 and azoxystrobin, were positively correlated with *Bacillus* and negatively correlated with
361 *Pseudomonas*. Conversely, scopoletin was positively correlated with *Pseudomonas* and negatively
362 correlated with *Bacillus* (**Figure S13**). These additional antagonistic compound interactions
363 between *Bacillus* and *Pseudomonas* could represent either competitive interactions between these
364 organisms or different adaptation to the different materials and wetting conditions.

365 **DISCUSSION**

367 As expected, wetted materials had higher bacterial and fungal growth rates and were dominated
368 by a few particular microbes, most notably the bacterial genera *Bacillus*, *Erwinia*, and
369 *Pseudomonas* and the fungal genera *Eurotium* and *Penicillium*. This dominance led to an overall
370 lower alpha diversity compared to non-wetted tiles. Wetting condition and material type described
371 the majority of the variance in bacterial, fungal and metabolite structure. Interestingly, each wetted
372 material showed its own unique microbe-metabolite dynamics.

373 Gypsum and MF gypsum were mostly colonized by *Bacillus*, with gypsum being a less selective
374 environment, which allowed for several bacterial species to thrive on the same coupon
375 simultaneously, each of them with high relative abundance and apparently sharing both the
376 physical space and resources. In contrast, MF gypsum prevented most fungal growth and allowed
377 *Bacillus* to dominate with little competition. MDF selected for fungal growth primarily, which
378 allowed for the rapid accumulation of the antibacterial chemical, nigragillin, which is known to be
379 made by the *Aspergillus* fungi. On OSB material, nigragillin and fumigaclavine C, a second
380 fungal-synthesized antibacterial metabolite, may play important roles in microbial growth
381 dynamics. Nigragillin, Fumigaclavine C, and *Aspergillus* relative abundance each gradually
382 increases over time, whereas the abundance of *Pseudomonas* declines after the antibacterial

383 metabolites reach peak abundance, suggesting a dose dependent response (**Figure S11**). These
384 observations bolster our hypothesis that production of antibacterial metabolites by *Aspergillus* may
385 inhibit the proliferation of surrounding bacteria. Also, there is a human health risk associated with
386 the proliferation of the *Aspergillus* fungi in the BE. While the most common species identified in
387 our data was *Aspergillus penicillioides*, a common indoor fungus in damp buildings with known
388 associations to allergies and asthma (Edwards *et al.*, 2012, Hay *et al.*, 1992), other *Aspergillus*
389 species are known to be able to produce mycotoxins (including aflatoxins), molecules that have
390 been associated with cancer and immunosuppression on humans (Roze *et al.*, 2013).

391 *MOLD-RESISTANT GYPSUM*

392
393 Traditional wood-based building materials contain natural polymers such as cellulose and lignin
394 that are susceptible to degradation by fungal colonization (Gravesen *et al.*, 1999, Pasanen *et al.*,
395 1992). With some fungi having been shown to produce mycotoxins including aflatoxins that could
396 affect human health (Roze *et al.*, 2013, Rand *et al.*, 2017), building materials such as mold-resistant
397 gypsum have been developed, which contain antifungal compounds intended to discourage fungal
398 growth. We were particularly interested to examine the microbial communities on these surfaces
399 and as expected, found that fungal growth was diminished on MF-gypsum compared to other
400 materials. However, it appeared that the scarcity of fungal colonies made way for bacterial species
401 to flourish with less competition; on non-wetted materials we observed MF-gypsum bacterial
402 particle counts greater than on the other three materials, and on wetted materials while the MF-
403 gypsum bacterial counts were second to MDF, the abundance level between non-wetted and wetted
404 tiles, unlike MDF, remained minimally changed. This raises the potential that pathogenic bacteria
405 colonization could occur on MF-gypsum and if wetted could grow and lead to negative health
406 outcomes. In terms of metabolite production, thiabendazole and azoxystrobin were some of the
407 anti-fungal compounds found in high abundance and overall a similar subset of compounds
408 accounted for most of the metabolite abundance on this material, indicating lower metabolic
409 diversity when the colonization is dominated by bacterial growth. We also detected a correlation
410 between thiabendazole and *Penicillium*, which suggested the persistence of thiabendazole-resistant
411 fungal strains.

412

413 *DIVERSITY AND INTERACTION BETWEEN MICROBES AND ENVIRONMENT*

414

415 Additionally, certain lipid metabolites (indicative of biofilm formation) showed significant
416 positive correlation with both *Bacillus* and *Pseudomonas* OTUs, and these lipids were negatively
417 correlated with the abundance of chloroplast OTUs, indicating that when bacteria and metabolites
418 indicative of biofilm formation are detected in greater abundance, we see a proportional reduction
419 in plant-associated signal. Similar to the lipids, metabolites annotated to organic molecules and
420 vitamins were also negatively correlated with the chloroplast OTUs, which suggests that bacterial
421 growth, indicated by increased proportion of 16S, cellular counts and associated metabolites, tends
422 to swamp out the background material-chloroplast signal. We hypothesize that this may be because
423 these molecules are being produced by bacteria colonizing and forming biofilms on the woody
424 material. When the relative abundance of the bacteria increases, it reduces our ability to detect
425 chloroplast sequences (based on a given sequencing depth); as such this negative correlation is
426 likely due to the increased abundance of the microbes that mediate the production of these

427 metabolites, reducing the detection frequency of specific chloroplast OTUs, and not due to some
428 mechanistic relationship between the wood and these molecules

429

430 *MICROBIAL-METABOLITE INTERACTIONS.*

431

432 Pseudomonads and Bacillus are often the main contributors to biofilm formation on material
433 surfaces in the built environment (Ronan *et al.*, 2013; Powers *et al.*, 2015; Raaijmakers *et al.*,
434 2010). Biofilms are complex extracellular matrices formed by bacteria through the excretion of
435 lipopeptide biosurfactants, to provide attachment to a surface to support colocalization with a
436 nutrient source and protection from dehydration and chemical activity. Some of these lipopeptide
437 biosurfactants produced by *Pseudomonas* and *Bacillus* species have been shown to have lytic or
438 growth-inhibitory activity against many microorganisms such as bacteria, viruses, mycoplasmas,
439 and fungi (Raaijmakers *et al.*, 2010). Powers *et al.* demonstrated that *Pseudomonas protegens*
440 produces antibiotics that inhibit biofilm formation and sporulation in *Bacillus subtilis*. They also
441 found that *Pseudomonas putida* secretes an unknown inhibitory compound that prevented biofilm-
442 associated gene expression. In our study we demonstrate a number of compounds known to have
443 potential biofilm inhibitory qualities that also co-correlate with either *Pseudomonas* or *Bacillus*
444 abundance, suggesting potential competitive activity between these organisms. While
445 *Pseudomonas*–*Bacillus* interactions have been shown to be competitive, interspecies interactions
446 within the genus *Bacillus* are also important in the formation of biofilms, lipids like hydroxy fatty
447 acids and mono-acyl-glycerophosphocholines could be building blocks or residual products of the
448 biofilm creation (Shank *et al.*, 2011, Diomande *et al.*, 2015, Dubois-Brissonnet *et al.*, 2016).

449

450

451 **CONCLUSION**

452

453 The simultaneous collection of environmental, metabolomic and microbial profiles reveals
454 insights into the chemical signals that may govern BE microbial communities under high humidity
455 conditions, as well as providing evidence that these the membership compete for space and
456 resources. Here we show that wetting condition can profoundly alter both fungal and bacterial
457 community succession, and that the taxa which dominate samples after wetting or exposure to high
458 humidity are not abundant in non-wetted materials and have little relation to the skin-associated
459 taxa which dominate samples of indoor environments. After wetting, the microbial community
460 undergoes a successional trajectory that can result in similar metabolic diversity even when
461 taxonomic diversity remains variable. We further show that while material choice significantly
462 influences bacterial diversity, the same is not true of fungal diversity. In summary, BE microbial
463 ecology once seen as a wasteland (Gibbons, 2016) could rather be seen as a desert environment
464 mostly formed with smaller assemblages that can rapidly become an active ecologically dynamic
465 community if water, in liquid or vapor form, is added. When a material experience high moisture
466 conditions, both fungal and bacterial growth rapidly accelerate and the metabolites associated with
467 their adaptation to different surface materials and competition for resources demonstrate ready
468 made eco-evolutionary adaptation to this sporadic availability of a crucial resource; this
469 phenomenon is very similar to what has been observed in real desert soil microbiomes (Neilson *et*
470 *al.*, 2017), as well as in very different ecosystems, such as sediments exposed to oil pollution
471 (Handley *et al.*, 2017).

472 **Methods**

473 *Test materials*

474 Four building materials were used in this study: oriented strand board (OSB), medium density
475 fiberboard (MDF), regular gypsum wallboard, and mold-resistant (i.e., mold-free, or ‘MF’)
476 gypsum wallboard. All samples were purchased new from a home improvement store in Chicago,
477 IL. The building materials were cut into 5 cm × 5 cm coupons for testing. The material coupons
478 were sterilized by UV irradiation for 20 minutes followed by surface cleaning with a 70% ethanol
479 solution.

480 *Inoculation*

481 The building material coupons were naturally inoculated by placing them uncovered on the floor
482 inside two residences for a period of approximately 50 days each. The goal was to allow for natural
483 settling of microbes from each indoor environment onto the material surfaces. Another set of test
484 coupons was covered with aluminum foil and kept in the laboratory for the same duration to be
485 used as a control group. Each set of test coupons included 44 coupons for each type of building
486 material (i.e., 176 coupons in total) to allow for multiple subsequent sampling strategies. One set
487 of test coupons was placed inside a 6th floor apartment unit with two adult occupants and a medium
488 sized dog located in downtown Chicago, IL (Location 1). The other set of materials was placed
489 inside a 2-story single-family residence without any pets near the main campus of Illinois Institute
490 of Technology, approximately 8 km south of the downtown residence (Location 2). During the
491 inoculation periods, built environment metadata (Ramos & Stephens 2014) were collected in each
492 residence, including temperature (T) and relative humidity (RH) using Onset HOBO U12 data
493 loggers and occupant presence within ~1 m range of the samples using Onset UX90 data loggers.
494 Coupons at a third location (the Built Environment Research Laboratory at the of Illinois Institute
495 of Technology) were covered with aluminum foil to minimize natural inoculation, serving as a
496 control group.

497 *Wetting and incubation*

498 After inoculation, half of each set of materials (i.e., 22 coupons each) from each location, as well
499 as 22 coupons from the control group, were submerged in sterilized tap water in separate pans for
500 ~12 hours to simulate the process of wetting of building materials due to a flood or leak. The other
501 half of each set of materials (i.e., the other 22 coupons each) from each location and the other 22
502 coupons from the control group were not submerged in water. Next, to encourage fungal growth
503 on all of the building materials, all of the material coupons were placed in trays (each tray
504 contained all 22 coupons of one type of material from one location or control group) and were
505 incubated at room temperature ($24 \pm 2.7^\circ\text{C}$) inside a static airtight chamber ($0.9 \text{ m} \times 1.2 \text{ m} \times 0.4$
506 m). Salt solutions (potassium nitrate) were used to maintain high RH near ~94% for the duration
507 of the experiment. Temperature and RH in the chamber were also recorded using Onset HOBO
508 U12 data loggers.

509 *Sampling procedures*

510 The material coupons were sampled for off-line biological and chemical analysis at 7 different

511 sampling time points, referred to here as: TP0, TP1, TP2, TP3, TP4, TP5, and TP6. The initial
512 samples (TP0) were taken just after retrieving the samples from the field inoculation and represent
513 non-wetted, naturally inoculated samples previously held at normal residential humidity
514 conditions. The remaining sampling time points occurred every ~5 days. At each time point, a new
515 coupon of each material from each condition that had never been swabbed before was swabbed,
516 while duplicates of previously un-swabbed samples were also swabbed periodically (at TP0, TP2,
517 TP4, and TP6) for comparison. Two samples ('TP0' and 'TP0 duplicated') were also swabbed at
518 every time point to investigate whether repeatedly swabbing the surfaces impacted the results.
519 Duplicates of both previously swabbed and previously un-swabbed samples were also included to
520 investigate whether or not natural inoculation and subsequent growth was evenly distributed across
521 multiple coupons. **Figure S1** illustrates the experimental setup and **Figure S14** shows coupons'
522 photographs at TP5 and TP6 for each one of the three locations. Details of swabbing procedure at
523 each time point are described below.

524 First, sampling reagents were prepared as follows. Phosphate-buffered saline (PBS) was used for
525 microbial samples that were to be analyzed for DNA and formaldehyde was used for microbial
526 samples that were to be analyzed by microscopy. For PBS, 500 μ l 1X PBS was added to 1.7 ml
527 microtubes for each sample to be collected. For formaldehyde, 100 μ l 4% paraformaldehyde was
528 added to 1.7 ml microtubes for each sample. Microcentrifuge tubes were filled with ethanol
529 solution (200 μ l 50% EtOH) to preserve samples for surface chemistry/metabolomics analysis.

530 For subsequent DNA sequencing and analysis, the entire surface of the test coupons was swabbed
531 using two BD Screw Cap SWUBETM Polyester swabs for approximately 20 sec. The same
532 researcher swabbed every time to keep the swabbing process consistent. One of the double swabs
533 was placed into the tube with PBS and frozen for shipping for subsequent sequencing. The tip of
534 the other of the double swabs was placed into microtubes and the swab tips were vortexed for 10
535 seconds. 100 μ L of sample buffer was removed added to the tube containing 100 μ L 4%
536 paraformaldehyde for fixation. These fixed samples were stored in a refrigerator at 4°C and then
537 sent to the San Diego State University team for running numerical counts of cells and virus
538 particles using microscopy.

539 For surface and microbial chemistry analysis (i.e., metabolomics), another test coupon was
540 swabbed using a cotton-tipped applicator that is dipped in ethanol (Petras *et al.*, 2016). The end of
541 the swabs were cut directly into pre-prepared collection tubes, stored at 4°C for 2-3 hours, and then
542 stored at -20°C overnight. Swabs were then removed with clean forceps the next morning, then re-
543 sealed into the microcentrifuge tubes and sent to the Northwestern University team on ice at -20°C
544 or lower. Overhead photos of each tray of coupons were also taken at each time point for image
545 analysis using ImageJ to calculate the percentage of visible microbial growth coverage (Hoang et
546 al. 2010).

547 *Viral-like particle and bacterial microscopy counts*

548 Epifluorescence microscopy was used to ensure that all samples contained bacteria and virus-like
549 particles and to estimate their abundance. 100 μ L of the paraformaldehyde-fixed samples were
550 resuspended into 5 mL of sterile 0.02 μ m filtered water. Each suspended sample was then filtered
551 onto a 0.02 μ m Whatman Anodisc filter membrane (Thurber *et al.*, 2009). The filters were stained
552 with 1X SYBR Gold and incubated for 10 minutes in the dark. Each filter was washed and mounted

553 onto slides to be observed. Visualization was performed using a QImaging Retiga EXi Fast Cooled
554 Mono 12-bit microscope and Image-Pro Plus software was used to collect digital images and
555 estimate VLP and bacterial abundances.

556 *Metabolomics analysis*

557 Samples were analyzed by High-Performance Liquid Chromatography and High-Resolution Mass
558 Spectrometry and Tandem Mass Spectrometry (HPLC-MS/MS). Specifically, the system
559 consisted of a Thermo Q-Exactive in line with an electrospray source and an Agilent 1200 series
560 HPLC stack including a binary pump, degasser, and autosampler, outfitted with a column (Waters
561 XBridge BEH Shield RP18, 100x2.1 mm, 5 μ m particle size with matching guard). The mobile
562 phase A was H₂O with 0.1% Formic Acid; B was Acetonitrile with 0.1% Formic Acid. The
563 gradient was as follows: 0-0.5 min, 98% A; 5 min, 80% A; 10-10.5 min, 5% A; 10.6-15 min, 98%
564 A, with a flow rate of 400 μ L/min. The capillary of the ESI source was set to 275 $^{\circ}$ C, with sheath
565 gas at 40 arbitrary units and the spray voltage at 4.0 kV. In positive polarity mode, MS1 data was
566 collected at a resolution of 35,000. The precursor ions were subsequently fragmented using the
567 higher energy collisional dissociation (HCD) cell set to 30% normalized collision energy in MS2
568 at a resolution power of 17,500. Data were processed with Compound Discoverer 2.0 (Thermo
569 Fisher) with MS/MS metabolite identifications made by comparing experimental MS/MS spectra
570 with library spectra from MZCloud (lower cutoff score of 90% match).

571
572 For the metabolites that were selected for more in-depth characterization, classification of structure
573 or substructure was performed by searching databases such as the Dictionary of Natural Products,
574 the LIPID MAPS Structure Database, and GNPS (Global Natural Products Social Molecular
575 Networking). Predicted structures resulting from a matched intact mass (≤ 10 ppm error) were
576 subsequently validated through manual analysis of fragmentation mass spectra.

577
578 Metabolite differential abundances (fold calculations) were calculated from Compound Discoverer
579 median peak areas for each compound including all three sampled locations.

580 581 *DNA extraction and sequencing*

582 To perform DNA extraction, the Qiagen DNeasy Powersoil HTP kit was used with a modified
583 protocol optimized for low-biomass samples. Swab tips were inserted into each well of the bead
584 plate, and then cut off using a sterilized wire cutter. The manufacturer's protocol was then
585 followed, with the following modifications: before cell lysis, the bead plates (containing beads,
586 bead solution, swabs, and the C1 solution) were heated for 20 minutes at 60 $^{\circ}$ C in a water bath.
587 Additionally, the protocol steps using solutions C2 and C3 were combined into a single step, by
588 adding 150 μ l each of C2 and C3 together to the lysed sample in the 1 ml plate.

589 The DNA obtained from the DNA extraction was used for both high-throughput 16S/ITS
590 sequencing, and qPCR. The 16S sequencing targeted the V4 region of the bacterial 16S rRNA
591 gene, using the primer pairs 515F/806R. The ITS sequencing targeted the highly variable fungal
592 internal transcribed spacer region located between the 5.8S and 18S rRNA genes. Both primer sets
593 used the same reaction mix and thermocycler instructions: Reaction mix: 9.5 μ L of molecular
594 biology grade H₂O, 12.5 μ L of Accustart II PCR Toughmix, 1 μ L each of forward and reverse
595 primers at 5 μ M, and 1 μ L of sample DNA for a total reaction volume of 25 μ L.

596 To make both the 16S and ITS amplicons, the following PCR program was used: Initial denaturing
597 step at 94°C for 3 minutes, followed by 35 cycle of: 94°C for 45 seconds, 50°C for 60 seconds,
598 and 72°C for 90 seconds, followed by a final extension step of 72°C for 10 minutes. The resulting
599 amplicons were quantified using the Picogreen dsDNA binding fluorescent dye on a Tecan Infinity
600 M200 Pro plate reader and pooled to 70 ng DNA per sample using the Eppendorf epMotion 5075
601 liquid handling robot. Primers and PCR reagents were removed using Agencourt AMPure beads,
602 and then the clean amplicon pool was sequenced at Argonne National Laboratory’s Environmental
603 Sample Preparation and Sequencing Facility, following the Earth Microbiome Protocol (Caporaso
604 *et al.*, 2011). Sequencing was performed on an Illumina Miseq using V3 chemistry, generating
605 2x150nt reads.

606 qPCR was performed using a Roche LightCycler 480 II. The 515F/806R primer pair was used
607 again for amplification, using a mix of 10 µL Light Cycler 480 SYBR Green I Master mix, 6 µL
608 of molecular biology grade H₂O, 1 µL of 515F primer (10 µM), 1 µL of 806R primer (10 µM),
609 and 2 µL of template DNA for a total of 20 µL per reaction. The following thermocycler conditions
610 were used: (1) 95°C for 5 minutes, (2) 95°C for 10 seconds, (3) 45°C for 45 seconds, (4) Measure
611 fluorescence, with steps 2 through 4 repeated 50 times. To determine the copy number of the 16S
612 gene (and therefore the number of organisms per swab), a standard curve was generated using a
613 serial dilution of a plasmid containing the *E. coli* 16S rRNA gene.

614 615 *Treatment of Technical Replicates*

616 We used Mantel test to determine whether the bacterial communities on replicate tiles (tiles on the
617 same tray sampled at the same time) significantly resembled each other and preserved patterns of
618 beta-diversity. We began by calculating the Bray-Curtis dissimilarity between each pair of samples
619 taken from the same tile type using the *beta_diversity.py* function from the software QIIME 1.9.1
620 (Caporaso *et al.*, 2010), producing dissimilarity matrices for each sampling type. Then the Mantel
621 test and false discovery rate adjustment was performed using the *mantel* and *p.adjust* functions in
622 the Vegan and stats R packages. For all comparisons between the sampling types the mantel
623 statistic (which measures the stress in the fit of the two matrices) was significantly high (mantel
624 ≥ 0.67 for fungi and ≥ 0.5 bacteria (all $p < 1E-05$) (**Table S2 (Fungi) and S3 (Bacteria)**). Based
625 on the highly significant resemblance between tile types, we treated all samples of the same type
626 as technical replicates, meaning that all combinations of material, location, wetting condition, and
627 time point had either 2 (time point 0), 3 (time points 1, 3, and 5), or 4 (time points 2, 4, and 6)
628 replicates.

629 *Rarefaction and statistical analyses*

630 After sequencing and sample merging, bacterial and Fungi OTU tables were rarefied to 1,000 and
631 10,000 reads respectively for statistical analyses. Rarefaction and statistical analyses were
632 performed using R.

633 *Random forest analyses*

634 Random forest models were implemented using the “randomForest” R package. Samples from
635 timepoint 0 were removed from the dataset. Models were built with 1000 trees and 10-fold cross

636 validation. For each of the 10 models for each metadata criterion, a randomly drawn 70% of
637 samples (100 samples) were used for model training and the remaining 30% (44 samples) were
638 used for validation.

639 *NMDS*

640 We visualized sample similarity using non-metric multi-dimensional scaling (NMDS) ordination
641 based on Bray-Curtis dissimilarity. Metadata vectors were fit onto the ordination using the *envfit*
642 command in the Vegan R package. We converted material, location, and wetting condition into
643 dummy variables (1 = yes, 0 = no) and, in the case of the bacterial and fungal datasets; also fit
644 vectors of relative abundance for the common genera described in **Figure 3**. We assessed
645 significance of each of the vectors using 10^5 permutations, and removed non-significant vectors
646 from the figure. The R^2 values for each vector and their significance is presented in **Table S1**

647 *Co-occurrence networks*

648 Traditional correlation networks are unsuited to genomic survey data as these data are relative,
649 rather than absolute, measures of community composition. Since the relative abundances of all
650 taxa within each sample must sum to 1, the fractions are not independent and will often exhibit
651 negative correlations to each other regardless of the true correlation in absolute abundance. To
652 avoid these compositional effects, we generated our networks using SparCC (Friedman & Alm,
653 2012), a correlation metric based on log-ratio transformed data that is specifically suited to
654 compositional genomic surveys. Pseudo-p values for each correlation were generated through
655 comparison from 100 to 1,000 bootstraps of the permuted OTU table.

656 For the same kingdom and microbe-metabolite networks only samples where either bacteria or
657 fungi and metabolites were found in detectable levels after rarefaction were used (N=83, N= 91
658 respectively). Additionally, only bacterial OTUs with >9 reads, fungi OTUs with >99 reads, and
659 metabolites with > 5'000,000 abundance in the rarified dataset were used for a total number of 630
660 bacterial OTUs, 352 fungi OTUs and 426 metabolites. Figures were generated using CAVNet R
661 package (Cardona, 2017) and only displayed the higher correlation threshold (positive or negative)
662 greater than 0.4.

663 For the network encompassing both bacteria and fungi, the OTUs reads threshold remained the
664 same but only samples with both 16S and ITS data were kept (N = 153) producing a new subset
665 of bacterial and fungi OTUs, 590 and 581 OTUs respectively. This dataset produced a co-
666 occurrence network with 1,171 nodes. Only positive correlations with a pseudo-p < 0.05 were
667 included, resulting in a network with 33,509 edges (density = 0.052). The network was ordinated
668 using the Fruchterman-Reingold Algorithm (edge-weighted, force-directed) in the *igraph* R
669 package, with node size based on the log read count of each OTU across all samples (with ITS
670 counts first divided by 10 to equalize rarefaction depth between datasets). We used the Walktrap
671 method (Pons & Latapy, 2005) to uncover dense subgraphs (modules) within the network, which
672 may correspond to distinct community structures. We chose Walktrap (which is based on random
673 walks within the network) as our method of community inference due to its computational
674 tractability and its accuracy at uncovering subgraphs regardless of network size (Yang *et al.*, 2016).
675 We used random walks of four steps, which resulted in four distinct modules with a network
676 modularity of 0.45

677

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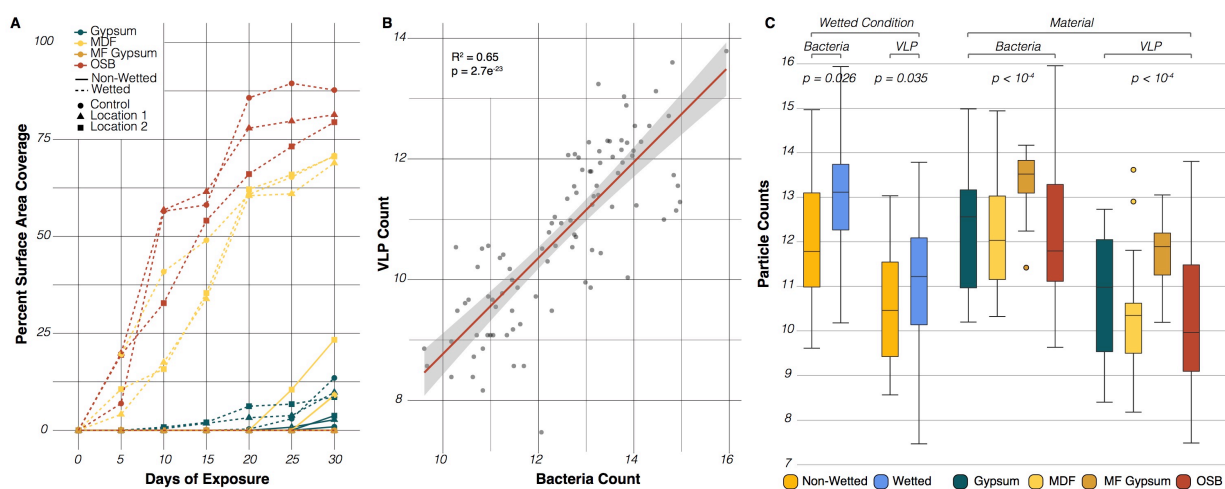
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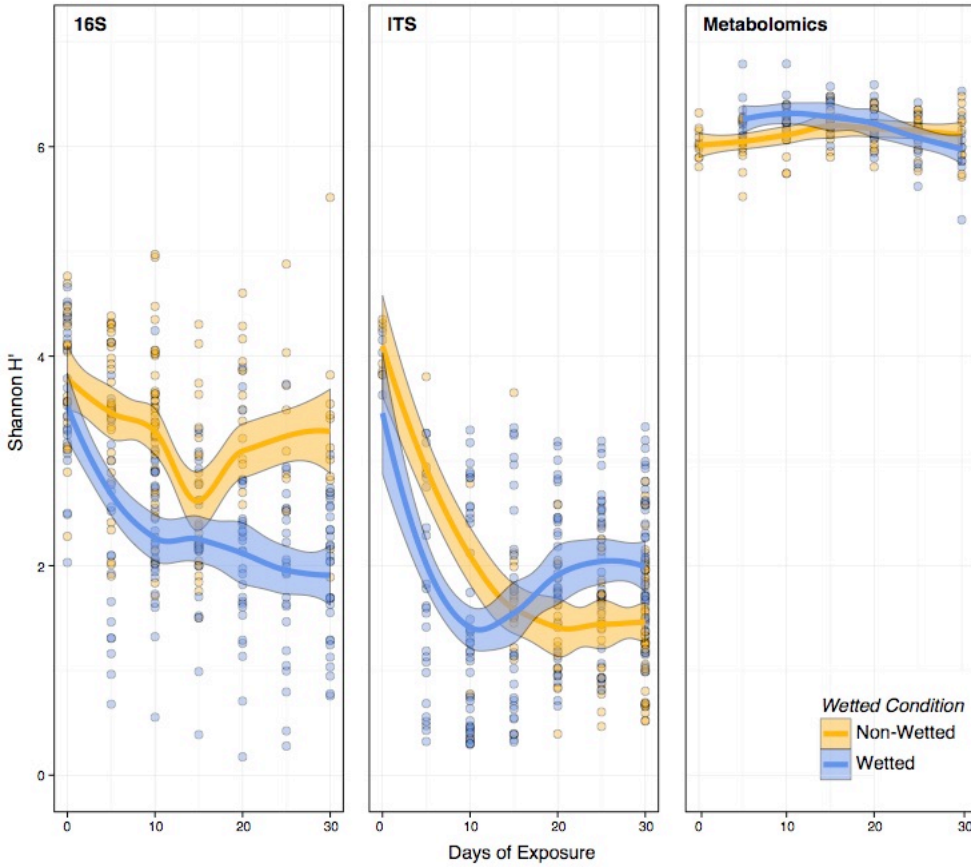
852 FIGURES

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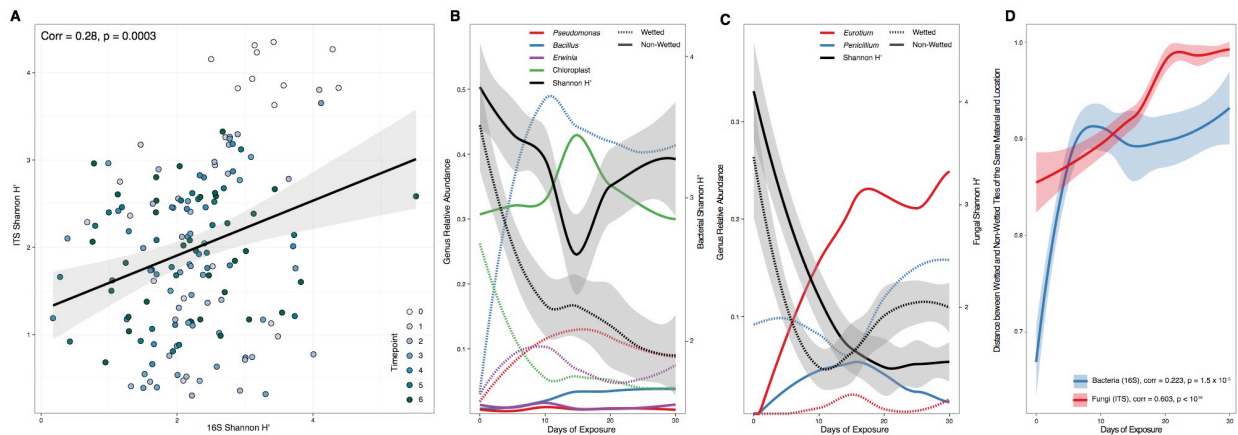


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855 **Figure 1: Microbial growth rates vary across sample types.** (A) Percent of surface area covered
856 by visible microbial growth through time (n=168, 4 materials, 3 locations, 2 wetting conditions, 7
857 time points). Color indicates tile material, point shape indicates inoculating location, and line type
858 indicates whether the tile was wet before incubation. (B) Correlation in the counts of bacteria-like
859 (BLP) and viral-like particles (VLP) across all tiles (n=96 samples, 4 materials, 3 locations, 2
860 wetting conditions, 4 time points). (C) Boxplots of BLP and VLP counts by wetting condition and
861 by material (n=96 samples).

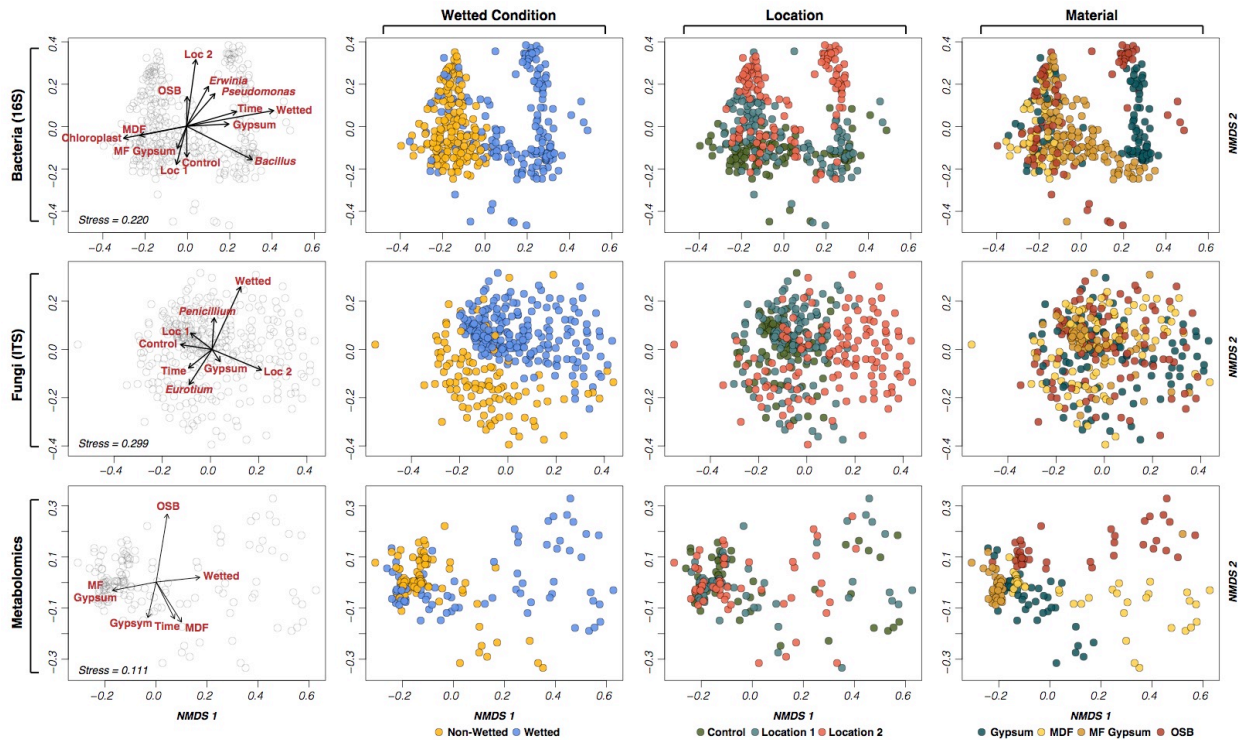


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863 **Figure 2: Change in the Shannon Index of samples over time.** Points represent individual
864 samples and the trend lines are a smoothed moving-average of the mean and shaded regions
865 indicate the standard error (n=338, 330 and 144 samples for 16S, ITS and Metabolomics,
866 respectively)

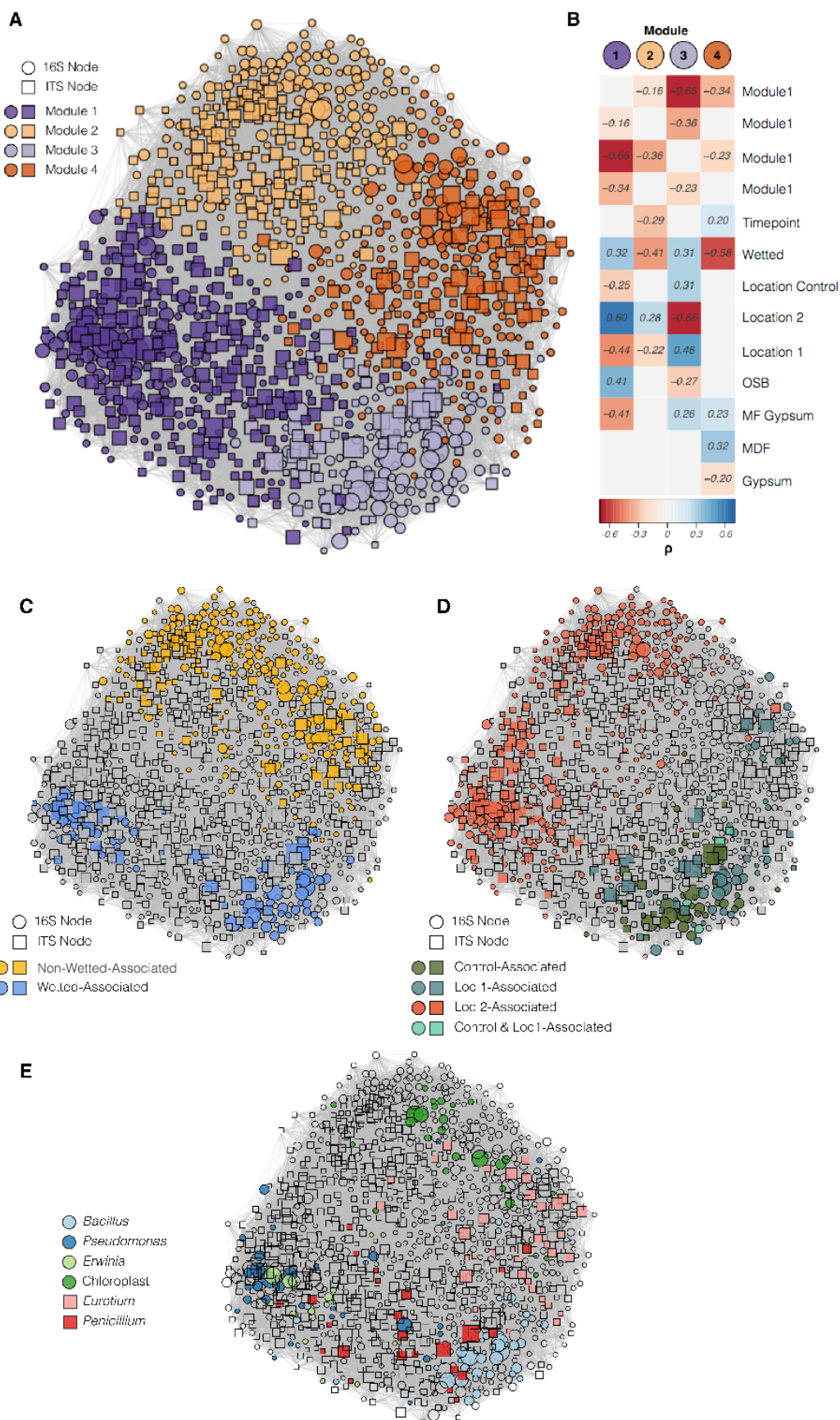


867 **Figure 3: Overview of community succession.** (A) Fungal diversity and bacterial diversity are
868 significantly correlated across communities (n=153 samples). Points represent individual
869 samples, colored by the time point at which the sample was taken. (B) Changes in the relative abundance
870 of selected bacterial genera over the course of succession (n=338 samples). Lines represent a
871 smoothed moving average of the mean. Genus is indicated by color and wetting condition is

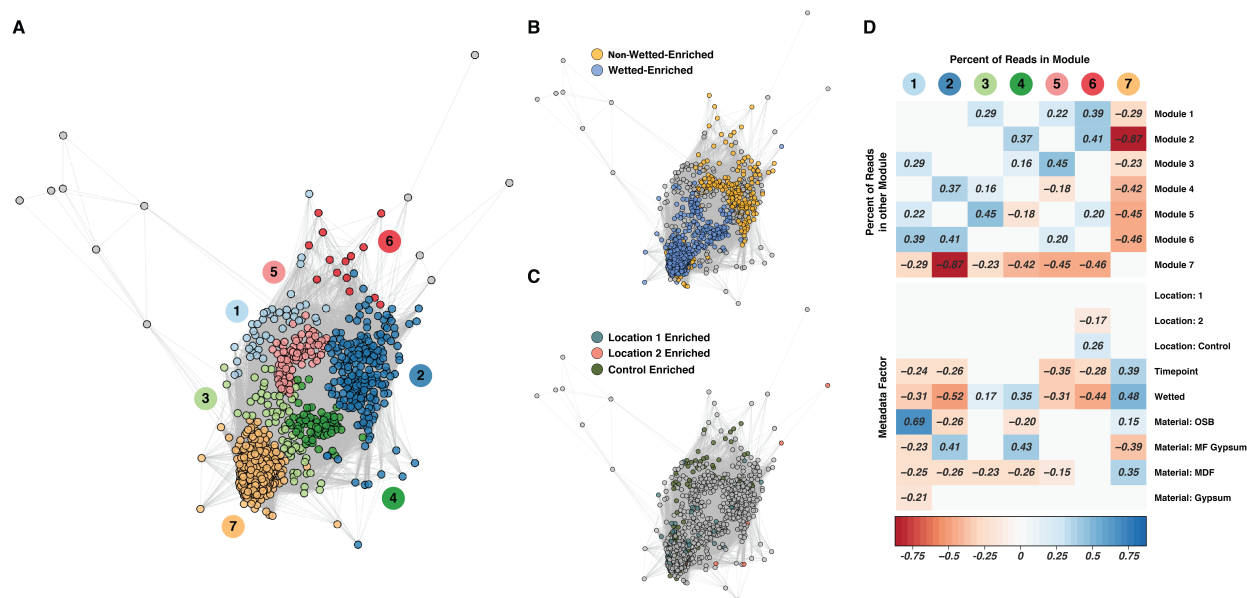
872 indicted by line style. Average community diversity (Shannon H' at OTU level, as in Figure 3) is
 873 indicated by black lines with standard error indicated by the gray shaded region. Genus abundance
 874 is indicated on the left y-axis and Shannon H' is indicated on the right y-axis. (C) Changes in the
 875 relative abundance of selected fungal genera over the course of succession (n=330 samples).
 876 Formatting is as in (B). (D) Wet vs. non-wetted replicates of tiles of the same material and
 877 inoculating location become increasingly dissimilar over the course of community succession
 878 (n=338, 330 samples for 16S and ITS, respectively). The y-axis is the Bray-Curtis distance
 879 between replicates. Spearman correlation between community dissimilarity and time is indicated
 880 in the legend error



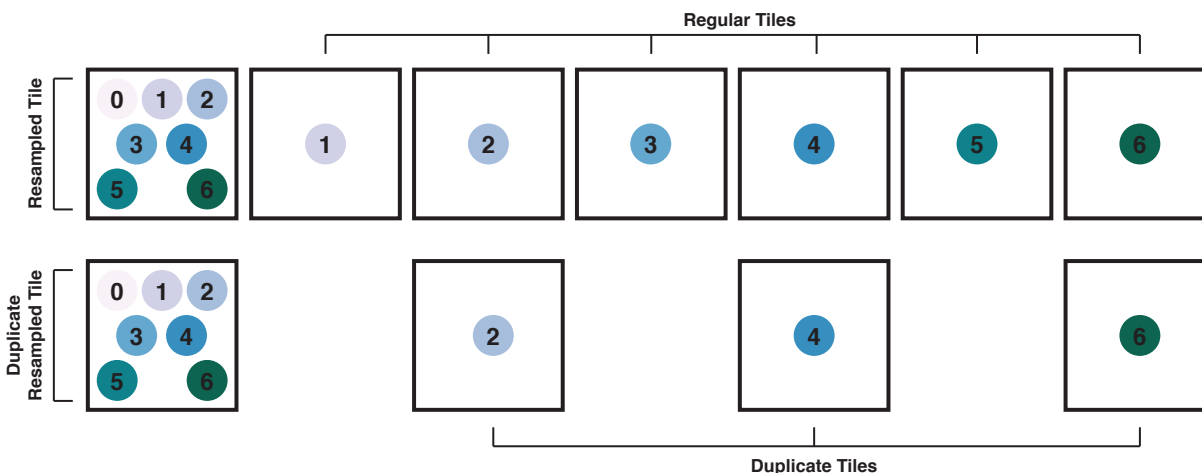
881 **Figure 4: NMDS plots illustrate clustering of sample diversity by sample type.** Each row
 882 comprises four identical NMDS plots (n=338, 330 and 144 samples for 16S, ITS and
 883 Metabolomics, respectively). The leftmost plot illustrates the ordination's association with
 884 environmental variables and the remaining plots color sample points by various metadata factors.
 885 The stress on the NMDS plot is indicated in the rightmost plot in each row error



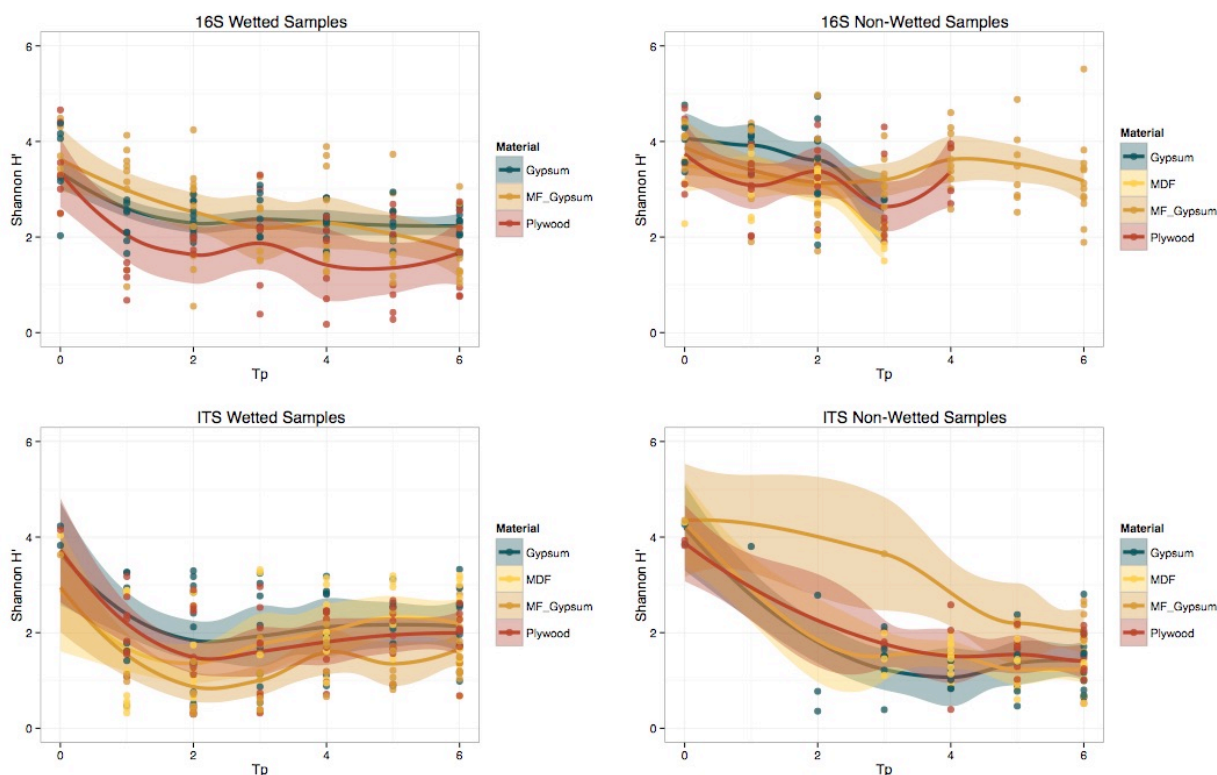
886 **Figure 5: Network of SparCC OTU correlations.** (A) Edge-weighted, spring-embedded
 887 network ordination, with nodes colored by module membership. Node shape represents node type
 888 (16S or ITS) and node size is based on the log-transformed abundance of each node (n=153 with
 889 both 16S and ITS, respectively). (B) Correlations between metadata factors (treated as dummy
 890 variables where true = 1 and false = 0) and the percent of reads in network modules. Non-
 891 significant correlations are not shown. (C) Taxa enriched in wet or non-wetted samples, as
 892 determined through a two-sided non-parametric t-test with 10^5 permutations. (D) Taxa enriched in
 893 samples originating from an individual inoculating location, with statistical methods as in (A). (E)
 894 Taxonomy of nodes in the genera included in Figure 4.



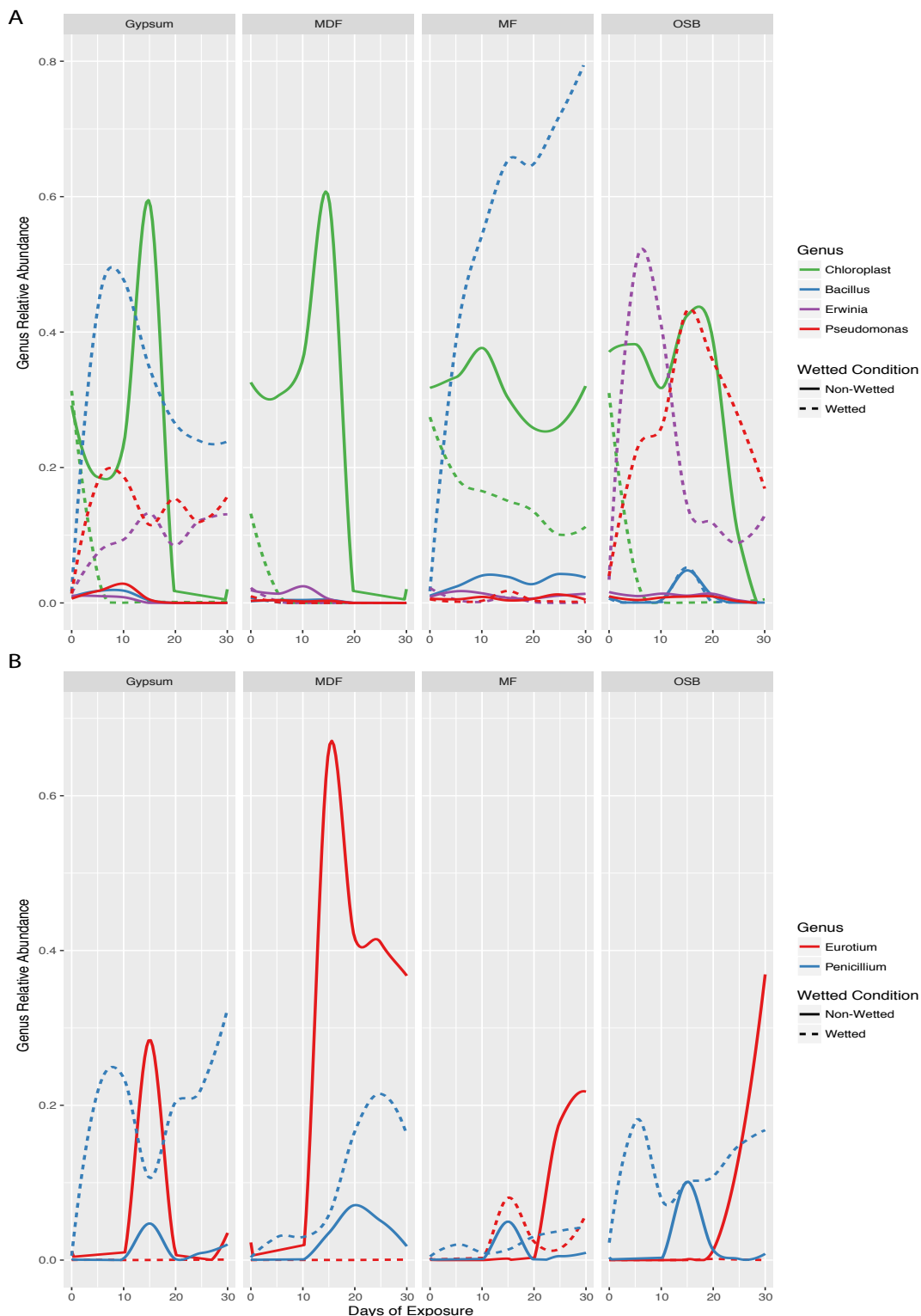
895 **Figure 6: Metabolite co-occurrence network.** (A) Network of significantly positive spearman
 896 correlations between metabolites, with network module indicated by color (n=144 samples). (B)
 897 Metabolites enriched in wet or non-wetted samples, as determined through a two-sided non-
 898 parametric t-test with 10^5 permutations. (C) Metabolites enriched in samples originating from an
 899 individual inoculating location, with statistical methods as in (B). (D) Correlations between
 900 metadata factors (treated as dummy variables where true = 1 and false = 0) and the percent of
 901 metabolites in network modules. Non-significant correlations are not shown.



902 **Figure S1: Experimental setup of project.** Illustration of the experimental setup and coupon
 903 sampling procedures.



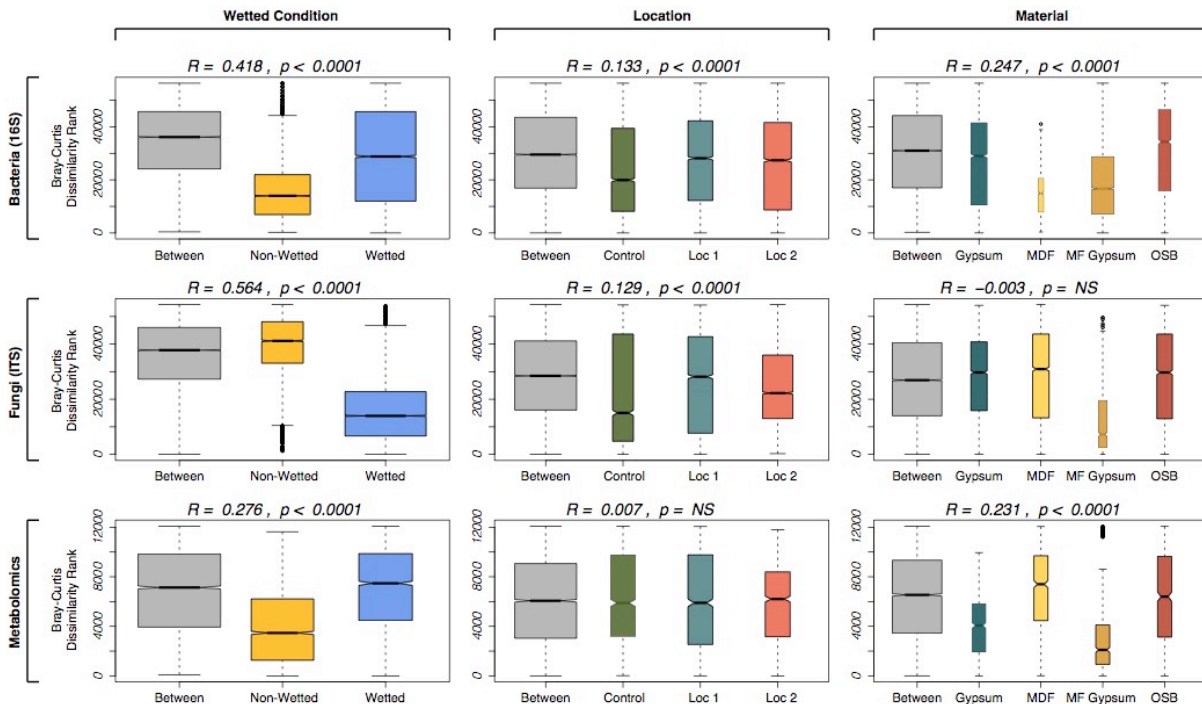
904 **Figure S2: Change in the Shannon Index by material over time.** Points represent individual
 905 samples (n=338, 330 samples for 16S and ITS, respectively) and the trend lines are a smoothed
 906 moving-average of the mean and shaded regions indicate the standard error
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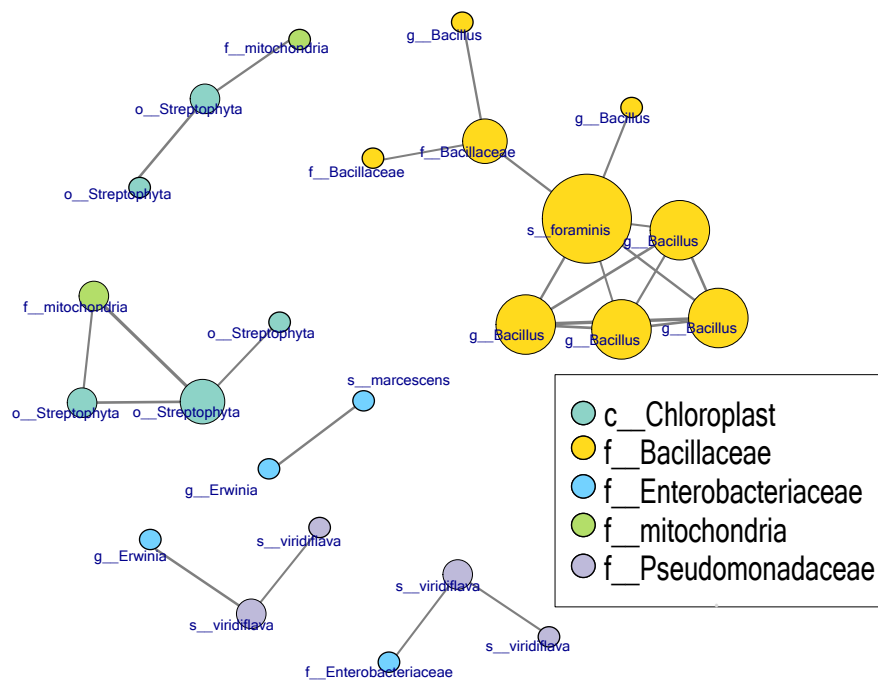
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Figure S3: Microbial succession by material over time Changes in the relative abundance of selected microbial genera for each material over the course of succession. (n=338, 330 samples for 16S and ITS, respectively). (A) Lines represent a smoothed moving average of the mean. Genus

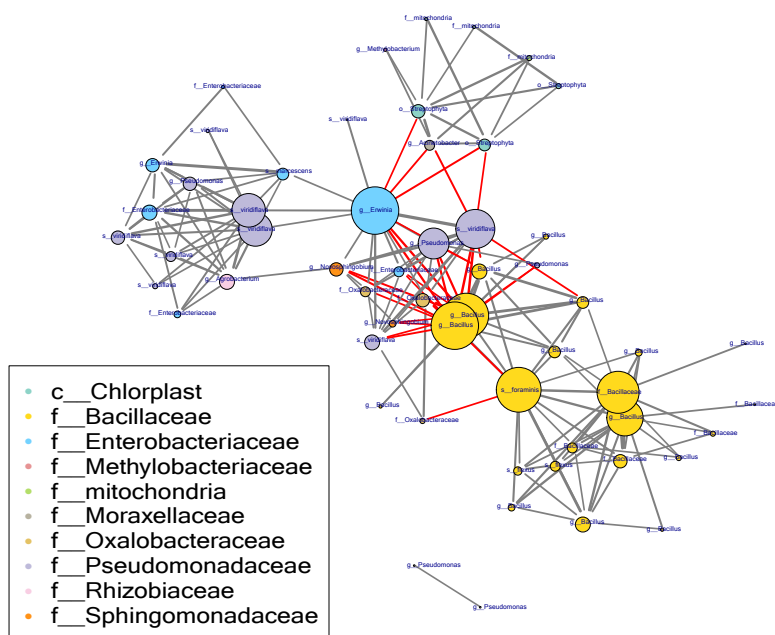
912 is indicated by color and wetting condition is indicated by line style. **(B)** Changes in the relative
913 abundance of selected fungal genera over the course of succession. Formatting is as in **(A)**
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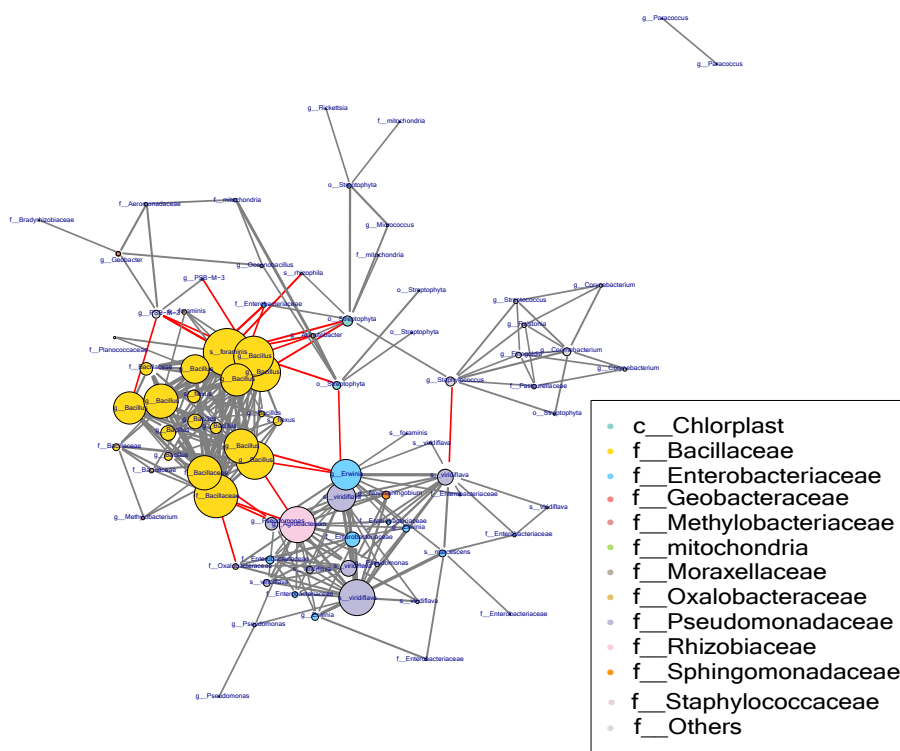
916 **Figure S4: ANOSIM quantifies the influence of metadata factors on the dissimilarity**
917 **between samples.** Columns represent different metadata factors and rows represent the three
918 datasets in this study (n=338, 330 samples for 16S and ITS, respectively). Boxplots depict the
919 range of ranked Bray-Curtis dissimilarities within and between factors (lower rank = lower
920 dissimilarity). Boxplot width indicates the number of samples represented by the boxplot
921



922 **Figure S5: Bacteria-Bacteria co-occurrence network.** Co-occurrence network (from n=83
923 bacteria samples) shows highly correlated bacteria form monophyletic clusters for samples
924 containing more abundant taxa
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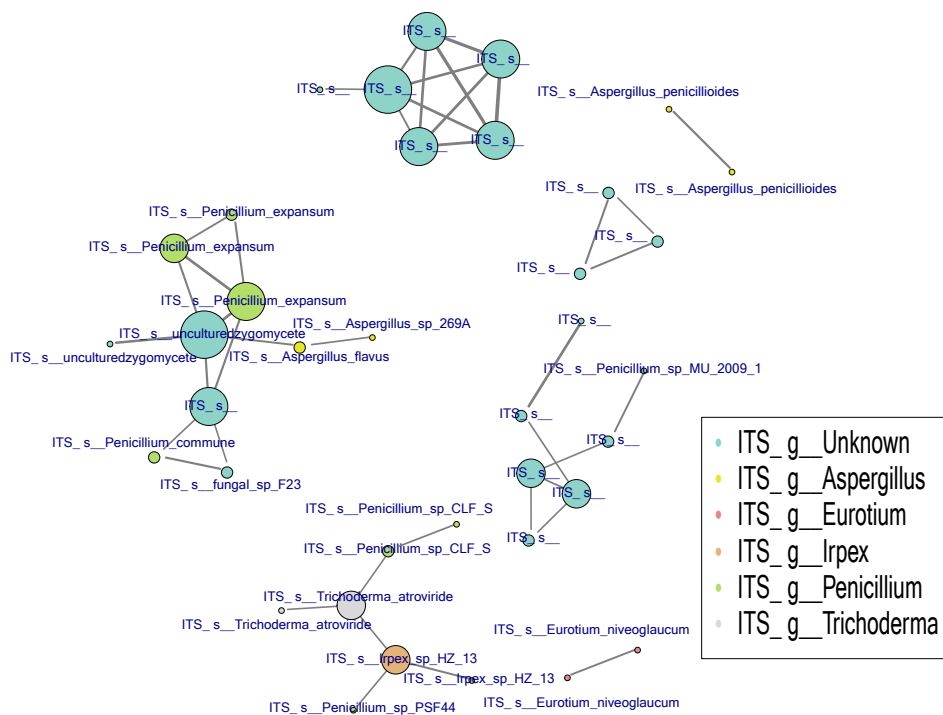


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 927 **Figure S6: Bacteria-Bacteria co-occurrence network on wet samples.** Co-occurrence network
 928 (from n=39 wet samples) shows how *Pseudomonas* and *Bacillus* are anticorrelated on wet samples
 929 for samples containing more abundant taxa
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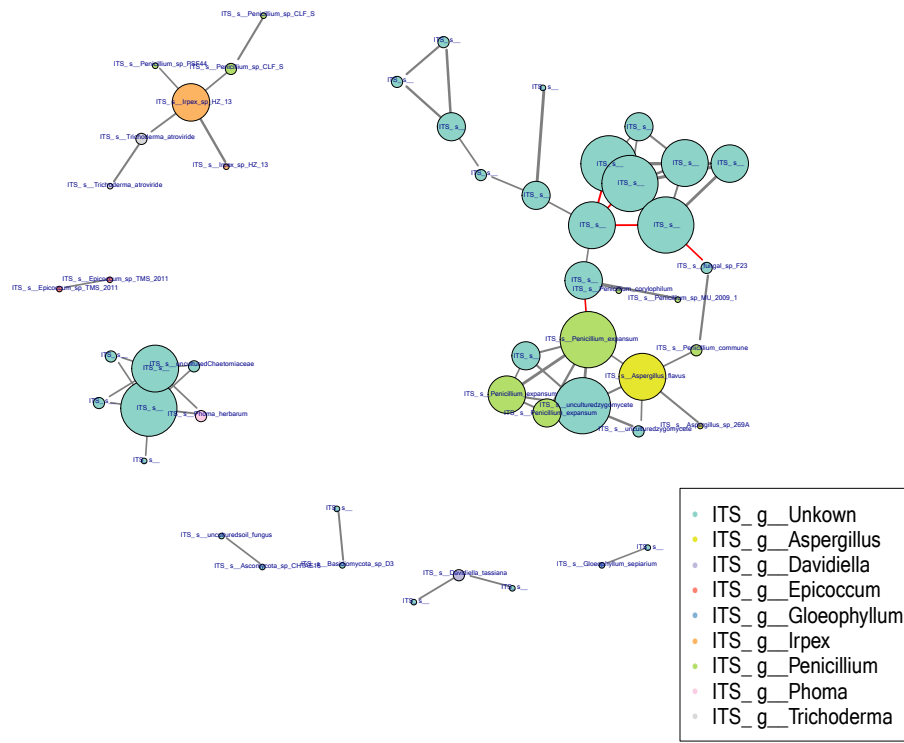
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Figure S7: Bacteria-Bacteria co-occurrence network on gypsum samples. Co-occurrence network (from n=24 gypsum samples) shows how *Pseudomonas* and *Bacillus* are anticorrelated on all gypsum samples



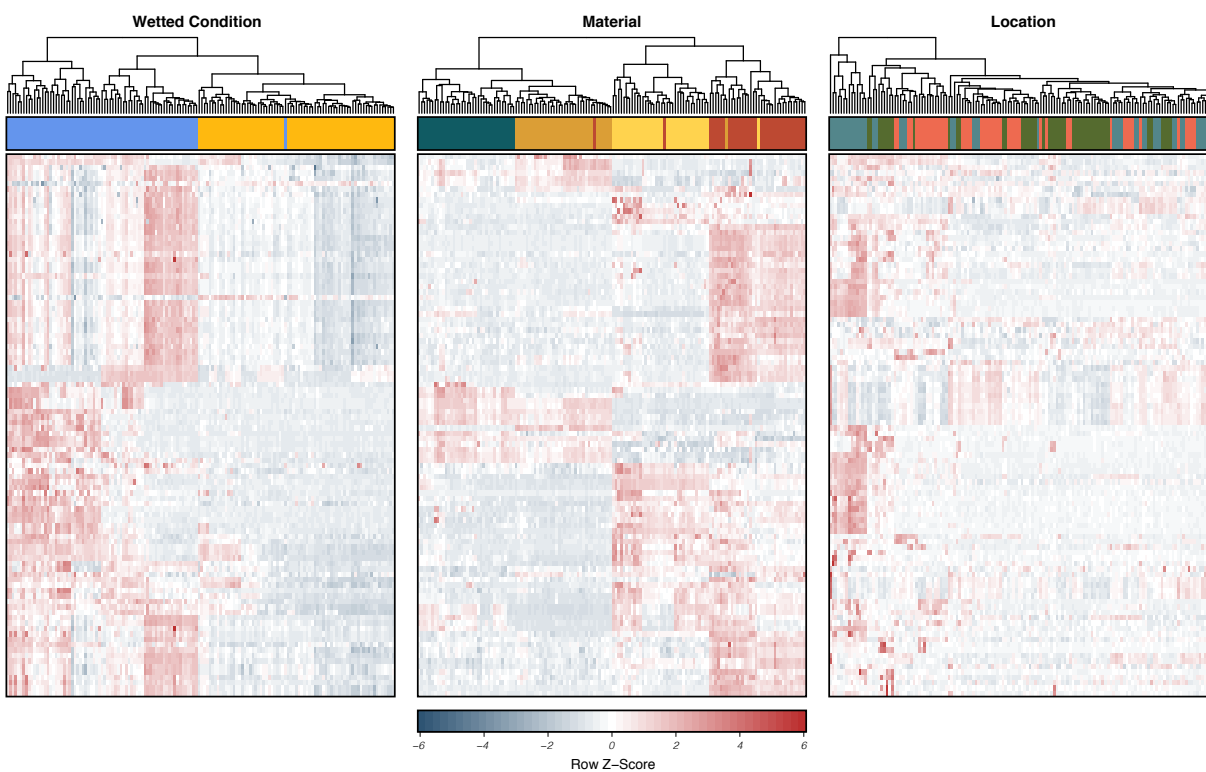
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Figure S8: Fungi-Fungi co-occurrence network. Co-occurrence network (from n=91 fungi samples) shows highly correlated fungi forms mostly monophyletic clusters



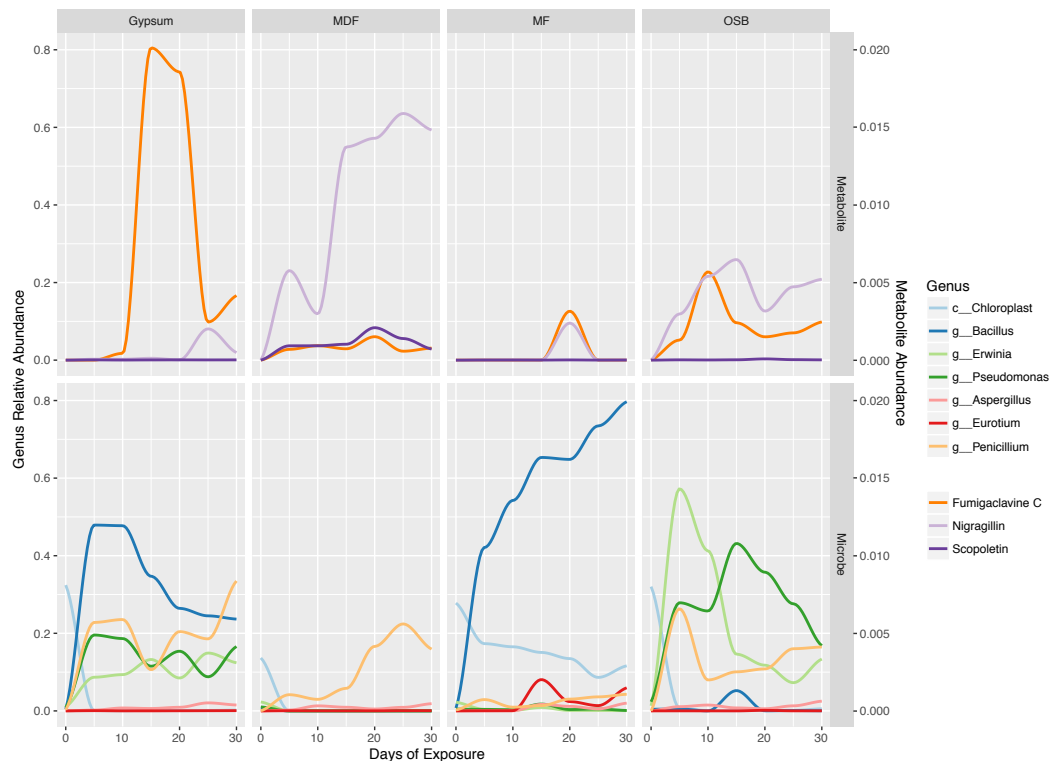
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Figure S9: Fungi-Fungi co-occurrence network on wet samples. Co-occurrence network (from n=58 wet samples) shows how certain Fungi OTUs are anticorrelated on wet samples



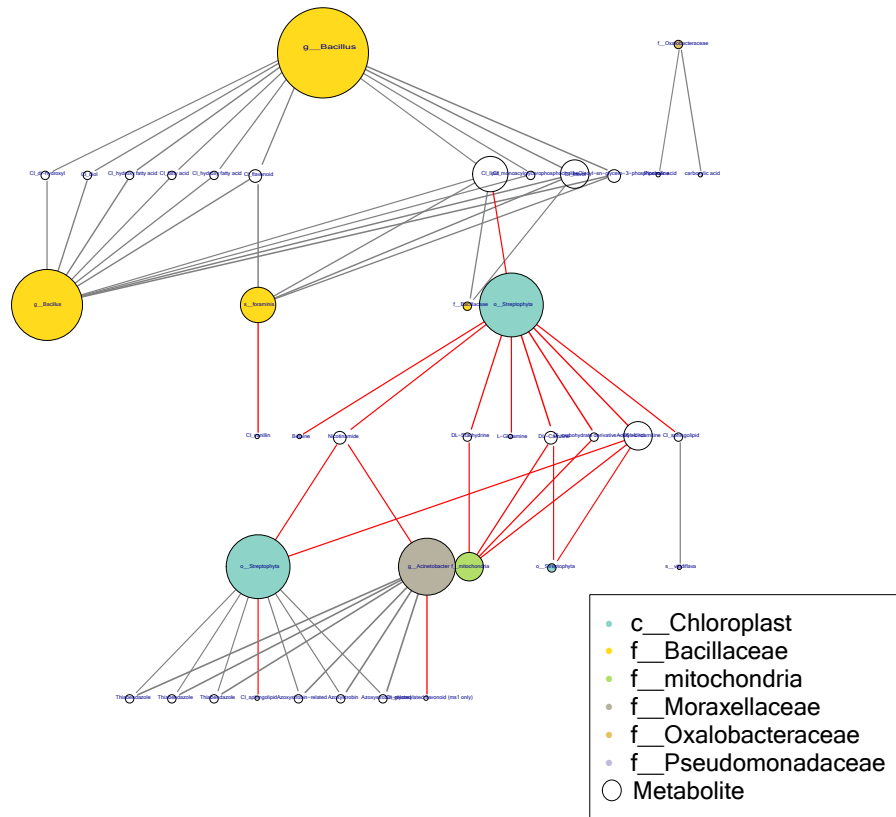
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Figure S10: Random forest metabolite selection heatmap. Random forest learning was used to select the metabolites that most distinctly identify each environmental condition, wetted or non-wetted, wood material type and inoculation location (n=144 samples and 3187 metabolites), the 100 highest-scoring metabolite features for each condition where selected for further examination.



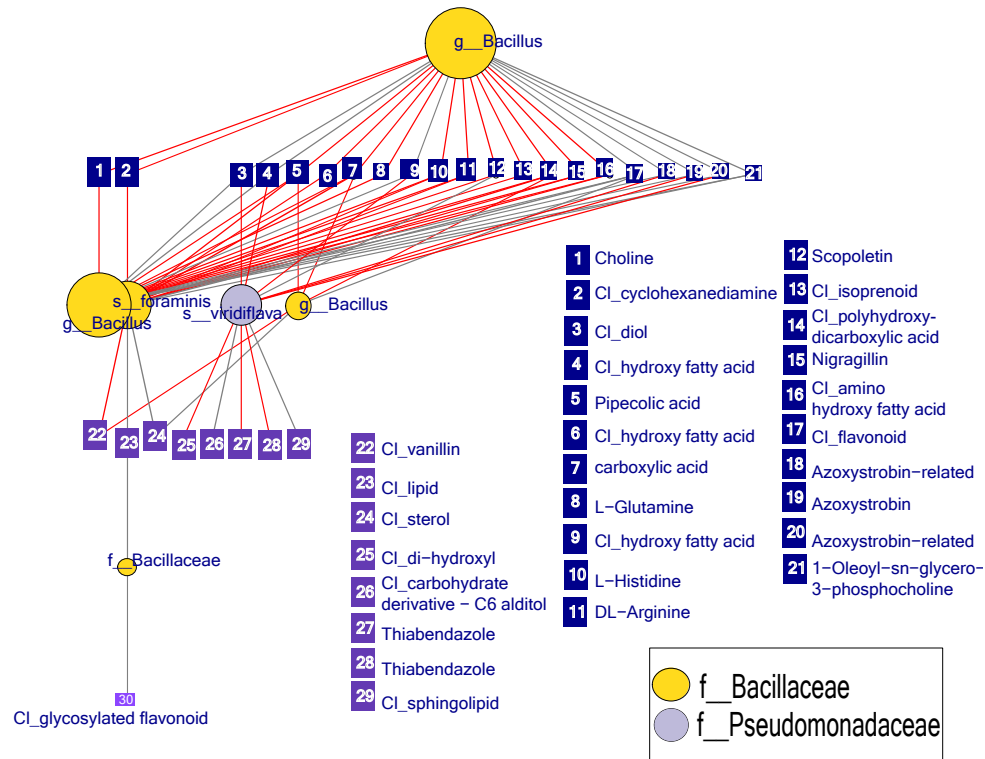
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Figure S11: Metabolite and microbial succession on wet samples by material over time. Changes in the relative abundance of selected bacterial genera for each material over the course of succession (n=39, 58 and 72 wet samples for bacteria, fungi and metabolites, respectively). Lines represent a smoothed moving average of the mean. Genus and metabolites are indicated by different colors

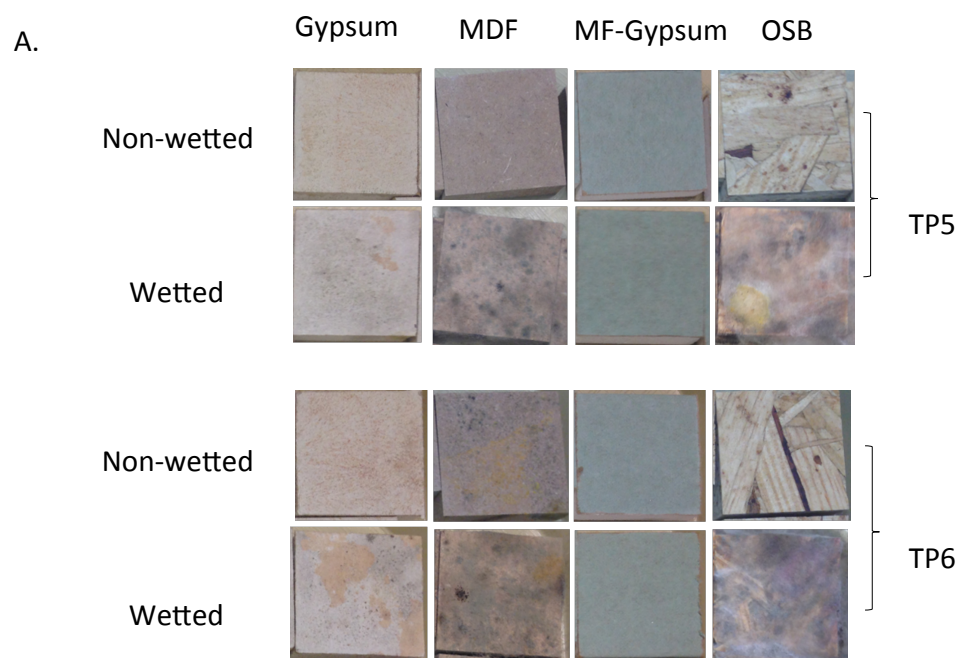


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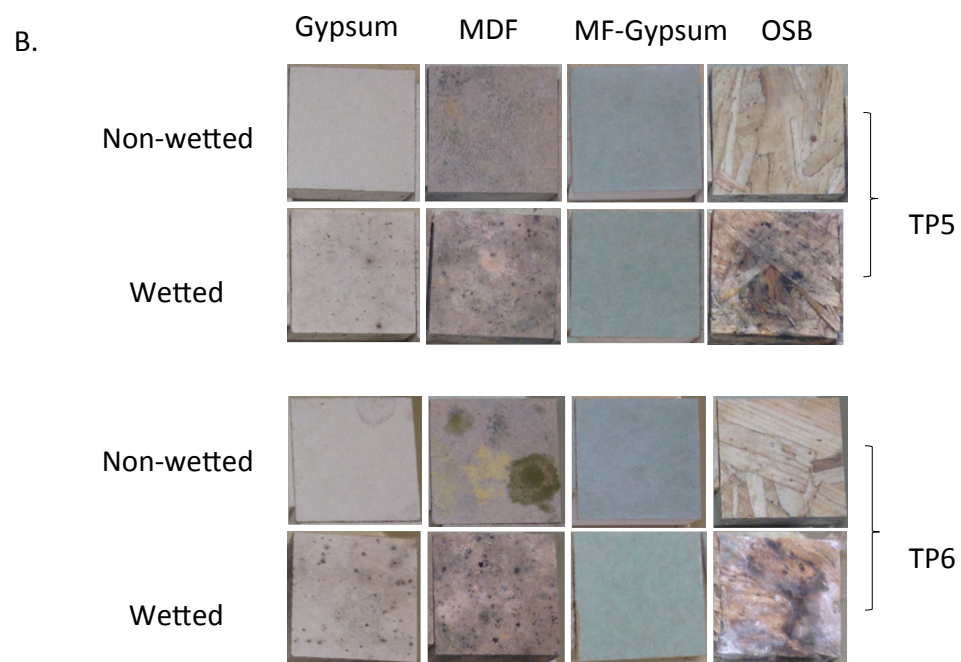
Figure S12: Bacteria-Metabolite co-occurrence network. Bacteria and Metabolite paired co-occurrences suggesting biochemical exchanges (from n=83, 144 samples, respectively). Lipid and hydroxyl compounds are strongly connected to *Bacillaceae* groups. Some specific lipids correlate positively with bacteria and negatively with wood material (plants). Vitamins and small carbon compounds negatively correlate with the wood material (plants).



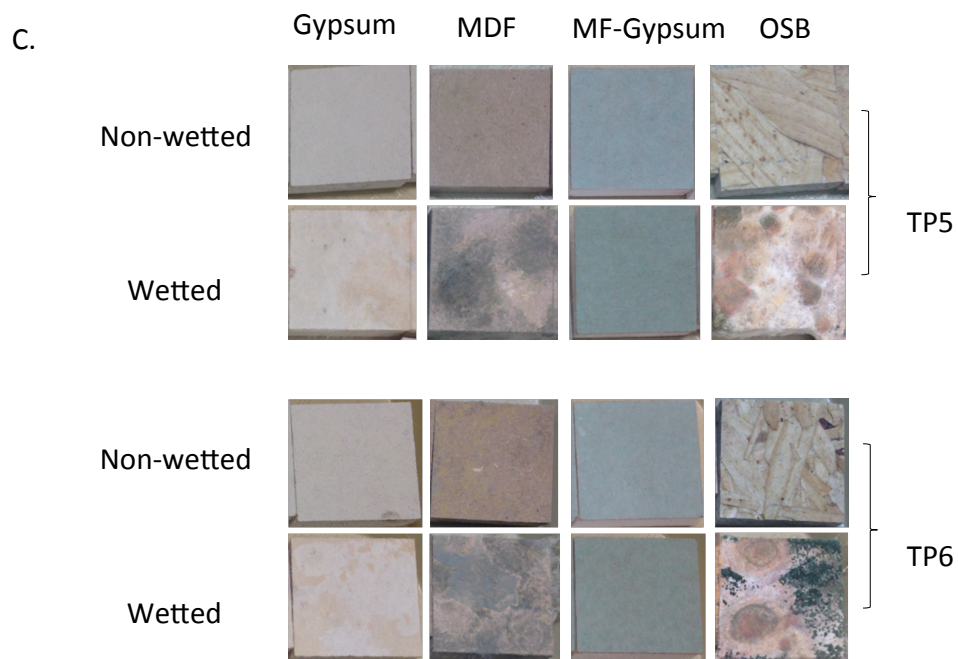
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 967 **Figure S13: Bacteria-Metabolite co-occurrence network for Bacillus and Pseudomonas**
 968 **interactions only.** Bacteria and Metabolite paired co-occurrences suggesting biochemical
 969 exchanges (from n=83, 144 samples, respectively). *Nigragillin* is negatively correlated with both
 970 *Pseudomonas* and *Bacillus*. *Azoxystrobin* correlates negatively with *Pseudomonas*, but positively
 971 with *Bacillus*. Hydroxyl compounds correlates negatively with *Pseudomonas* but positively with
 972 *Bacillus*



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976 **Figure S14: Photographs of wood coupons from different materials and wetting conditions**
977 **at TP5 and TP6.** Bacterial and Fungal growth on coupons surface photographs for (A) location
978 1 (B) location 2 and (C) control location.
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	<u>Bacteria (16S)</u>		<u>Fungi (ITS)</u>		<u>Metabolomics</u>	
	R^2	p	R^2	p	R^2	p
Wetted	0.374	< 0.0001	0.450	< 0.0001	0.204	< 0.0001
Timepoint	0.130	< 0.0001	0.094	< 0.0001	0.178	< 0.0001
Control	0.074	< 0.0001	0.107	< 0.0001	0.001	0.983
Location 1	0.045	0.0006	0.076	0.0006	0.009	0.502
Location 2	0.214	< 0.0001	0.303	< 0.0001	0.007	0.603
Gypsum	0.085	< 0.0001	0.022	0.0271	0.141	< 0.0001
MDF	0.108	< 0.0001	0.009	0.2399	0.238	< 0.0001
MF Gypsum	0.027	0.0095	0.008	0.2720	0.209	< 0.0001
OSB	0.041	0.0015	0.002	0.762	0.503	< 0.0001
<i>Bacillus</i>	0.259	< 0.0001	NA	NA	NA	NA
<i>Pseudomonas</i>	0.088	< 0.0001	NA	NA	NA	NA
<i>Erwinia</i>	0.098	< 0.0001	NA	NA	NA	NA
Chloroplast	0.196	< 0.0001	NA	NA	NA	NA
<i>Penicillium</i>	NA	NA	0.106	< 0.0001	NA	NA
<i>Eurotium</i>	NA	NA	0.185	< 0.0001	NA	NA

981 **Table S1: ANOSIM results calculate the factors significantly correlated with differences in the**
microbial communities across our three datasets, Bacteria, fungi, Metabolomics

	REG			RE			RE.DUP			DUP		
	mantel	significance	n	mantel	significance	n	mantel	significance	n	mantel	significance	n
REG	<NA>			0.67	1E-05	74	0.71	1E-05	79	0.81	1E-05	44
RE	0.67	1E-05	74	<NA>			0.85	1E-05	77	0.75	1E-05	40
RE.DUP	0.71	1E-05	79	0.85	1E-05	77	<NA>			0.79	1E-05	39
DUP	0.81	1E-05	44	0.75	1E-05	40	0.79	1E-05	39	<NA>		

Table S2: Mantel test results calculate the correlation among fungi samples across different sampling strategies.

	REG			RE			RE.DUP			DUP		
	mantel	significance	n	mantel	significance	n	mantel	significance	n	mantel	significance	n
REG	<NA>			0.62	1E-05	75	0.56	1E-05	75	0.60	1E-05	35
RE	0.62	1E-05	75	<NA>			0.72	1E-05	80	0.53	1E-05	36
RE.DUP	0.56	1E-05	75	0.72	1E-05	80	<NA>			0.50	1E-05	35
DUP	0.63	1E-05	56	0.53	1E-05	36	0.50	1E-05	35	<NA>		

Table S3: Mantel test results calculate the correlation among bacteria samples across different sampling strategies.