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Madison T. James

Sophia T. Farrisi

Shreya Shah

Vishal Shah

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**Linking Microbial Community Structure and Ecosystem Functions  
in Acidic Soil from Pennsylvania, USA**

**Madison T. James, Sophia T. Farrisi, Shreya Shah, Vishal Shah\***

West Chester University, West Chester, PA, USA

**RUNNING TITLE:** Soil microbial community structure and function

\*Corresponding Author  
Phone: 610-436-3522  
Email: [vshah@wcupa.edu](mailto:vshah@wcupa.edu)

23 **Abstract:** Microorganisms play a critical role in the structure and functioning of soil ecosystems.  
24 Within acidic soil across the northeastern United States and Canada, we have little understanding  
25 of the microbial diversity present and its relationship to the biochemical cycles. The current study  
26 is aimed at understanding the taxonomical and functional diversities in the acidic soil obtained  
27 from near various types of trees, how the diversities change as a function of depth, and the linkage  
28 between taxonomical and functional diversities. From eight sampling locations, soil samples were  
29 collected from three horizons (depths). The three depths were 0-10 cm (A), 11-25 cm (B), and 26-  
30 40 cm (C). Results indicate that across all the samples analyzed, *Bradyrhizobium* and *Candidatus*  
31 *Solibacter* are the most abundant bacteria in the soil microbiome. The differences in the soil  
32 microbiome across the samples were attributed to the abundance of individual organism's present  
33 in the soil and not to the presence or absence of individual organisms. Subsystem level analysis of  
34 the soil microbiome sequences indicate that there is higher level of abundance of genes attributed  
35 to regulation and cell signaling. A low level of sequences were detected for sulfur metabolism,  
36 potassium metabolism, iron acquisition and metabolism, and phosphorous metabolism. Structure-  
37 functional analysis indicate that *Bradyrhizobium*, *Rhodopseudomonas*, and *Burkholderia* are the  
38 major organisms involved in the nutritional ecosystem functioning within acidic soil. Based on  
39 the results, we propose utilizing a consortium of these organisms as an environmentally friendly  
40 alternative to the use of chemicals to maintain soil fertility and ecosystem functioning.

41

42 **Keywords:** Acidic soil, microbial ecology, shotgun metagenome, microbiome, *Bradyrhizobium*,  
43 *Rhodopseudomonas*, *Burkholderia*.

44

## 45 1.0. INTRODUCTION

46 Soil microorganisms are the largest biodiversity pool on earth, with more than  $10^{30}$  microbial cells,  
47  $10^4 - 10^6$  species, and nearly 1000 Gbp of microbial genome per gram of soil (Vogel et al. 2009;  
48 Mendes and Tsai, 2018). They are the primary factors that affect the soil ecosystem functioning  
49 and play key roles in forming and maintaining a multitude of soil characteristics including  
50 integrity, fertility, ecology, and overall soil function (Shah et al. 2011). Soil microorganisms are  
51 also vital for decomposition, pollutant removal, recycling of essential elements, suppressing plant  
52 diseases found in soil, and promoting growth for vegetation (Garbeva, 2004). Much is known of  
53 the microbial taxa present in soils from across the planet and the impact of perturbation of soil  
54 conditions. A Google Scholar search for the term 'soil microbial diversity' reveals over 1.6 million  
55 hits. Nevertheless, our understanding of how microbial diversity and ecosystem functions are  
56 linked, and how each of the microbial taxa present in the soil are linked to the individual ecological  
57 functions remain limited.

58 Increased use of 16s rDNA metagenomic methodology using pyrosequencing and Illumina  
59 Miseq and Hiseq techniques, has increased our understanding of the taxonomy of soil  
60 microorganisms by orders of magnitude. However, the 16S rDNA sequencing method has  
61 numerous limitations including differentiating closely related species (Hasan et al. 2014), non-  
62 uniform distribution of sequence dissimilarity among taxa, presence of multiple copies of the 16S  
63 rRNA gene (Garrity et al. 2009), failure of target amplification of polymerase chain reaction (PCR)  
64 primers (Venter et al. 2004), and generation of chimeric sequences (Quince et al. 2009). Further,  
65 in majority of these research, the role of individual microorganisms in the soil remains at the level  
66 of hypothesis based on prior literature (examples include our own prior research: Kumar et al.  
67 2011; Collins et al. 2012). Methods such as Biolog, Fungilog, and soil enzyme activity are many

68 times used in studies as indicators of the ecosystem functioning and correlation to the taxonomic  
69 data (Rutgers et al. 2016; Sobek et al. 2003; Nannipieri et al. 2002). While a step forward, these  
70 methods are primarily predictive of soil microbial functional dynamics (Bell et al. 2009).

71 Whole genome shotgun metagenomics provide a better approach for obtaining the  
72 taxonomic and functional aspects of the entire soil microbial genome. This method yields millions  
73 to billions of short reads, providing necessary sequencing depth as needed. It also offers an  
74 opportunity to identify organisms present in the microbiome and the biochemical pathway  
75 information present at the genomic level in each of the identified organisms. In this study, we  
76 employ shotgun metagenomic approach to identify and quantitate bacterial species present in the  
77 acidic soil and elucidate the major ecological functions of major organisms.

78 Acidic soil typically has a pH range of 4.0-4.5, is high in iron and aluminum, and is often  
79 considered nutrient-poor. Across the eastern United States and southeastern Canada, soil is  
80 primarily acidic (Bruulsema 2006). The acidic conditions in the soil of the region is primarily  
81 attributed to the parent materials of the soil and increased precipitation that leaches cations from  
82 the soil. The soil is optimal for the growth of trees like Apple, Beech, Dogwood, Oak, and  
83 Magnolia, and Pears. Literature search indicates that no reports are available studying the structure  
84 – function relationship in the natural acidic soil from the region.

85 The current study focused on understanding the taxonomical and functional diversities in  
86 the acidic soil obtained from near various types of trees, how the diversities change as a function  
87 of depth, and the linkage between taxonomical and functional diversities. We address three  
88 questions in the study: (1) what are the major microorganisms that are common to all soil types

89 and depth? (2) what are the biochemical pathways that can be generalized across all the soil types  
90 and depth? (3) What role do each of these organisms play in the ecosystem functioning in the soil?

91  
92 **2.0. METHODOLOGY**

93 **2.1. Sample Collection:** Samples were collected initially from eight sampling locations across the  
94 West Chester University Campus in West Chester, PA. No permit was required to obtain samples.  
95 Table 1 describes the analyzed eight locations, the types of vegetation present at each sampling  
96 location, as well as sampling coordinates. The locations were selected based on the vegetation  
97 present and initial sampling of 25 different locations. The final locations were selected based on  
98 the similarity of the vegetation between sampling locations, pH levels, and the quantity/quality of  
99 the DNA isolated. The protocols for the safety of data collection were strictly followed as  
100 recommended by the U.S. Fish and Wildlife Services and the Foundation for Ecological Research  
101 in the Northeast (Batcher, 2005). At each location, soil samples were collected from three horizons:  
102 0–5 cm (Horizon A); 6-15 cm (Horizon B) and 16–30 cm (Horizon C).

103 **2.2. pH Measurement:** 5 g soil samples were mixed with 10 mL d/w and vortexed for 10 minutes.  
104 The solution was allowed to sit for 1 hour and pH measured of the settled solution. All  
105 measurements were done in triplicates.

106 **2.3. DNA Extraction and Shotgun Metagenomics:** DNA extraction from each soil sample was  
107 carried out using the Qiagen DNeasy PowerSoil DNA Isolation Kit, according to the  
108 manufacturer's protocol. DNA concentration in all samples was determined using the Qubit 3  
109 Fluorometer (Invitrogen Technologies). All the samples were diluted to 100ng/μl and used for the  
110 library preparation, using the Nextera Dna Flex Library Preparation kit (Illumina, San Diego, CA),  
111 according to the manufacturer's protocol. Cluster generation and sequencing were performed with

112 the MiSeq Reagent Kit v2 500-cycles Paired-End in a MiSeq instrument at West Chester  
113 University. Samples were sequenced in a batch of 24 samples on a single flow cell. DNA  
114 sequences were annotated with Metagenomics Rapid Annotation (MG-RAST) pipeline version 4.0  
115 for downstream analyses. Taxonomic and functional profiles were generated using the normalized  
116 abundance of sequence matches to the Refseq and Subsystems databases, respectively. All settings  
117 were set at default values prior to analysis. The sequences have been deposited and are available  
118 through the NCBI BioProject Database ID: PRJNA 719140.

#### 119 **2.4. Clustering analysis:**

120 Clustering analysis were performed using Statistica (release 14.0) software. The tree cluster  
121 analysis was performed using Ward's method as the amalgamation rule and the distance measured  
122 as the Euclidean distances. Prior to clustering analysis, data obtained from MG-RAST were log<sub>2</sub>  
123 transformed and DSeq normalized.

## 124 **RESULTS**

125 **3.1. Soil pH:** pH for all the soil samples analyzed in this study were in the acidic range of 4.1 to  
126 6.3 (Table 1). Results show that the type of tree clearly influences the soil pH, with soil around  
127 Douglas Fir being the most acidic soil amongst all the types studied. No significant difference in  
128 pH was observed across the depths, except for soil obtained around the Oak tree (Table 1), where  
129 a stark drop in pH was observed as we go from depth A (pH, 5.7) to depth C (pH, 4.4).

130 **3.2. Sequencing analyses and microbial community diversity:** A total of 22,745,412 raw  
131 sequence reads were generated for the 24 samples using the Illumina Miseq sequencing platform.  
132 96.2% of the sequences passed the Illumina Chasity Filter for a total of 21,880,208 PF reads. MG-  
133 RAST analysis of the submitted reads yielded 8,219,706 total sequences (Table S1). Over 99% of

134 the sequences were annotated, with almost equal distribution of known proteins (4,073,051) and  
135 proteins of unknown function (4,073,868) (Table S1).

136 The rarefaction curves indicate high genetic diversity, with no complete saturation  
137 observed even after almost 8 million sequences (Figure S1). For all the samples, the curve has  
138 slowly begun to flatten, indicating a reasonable number of species have been sampled. The mean  
139 alpha diversity observed was 479, with the range from 417 to 547 species (Table S1).

140 **Taxonomic characterization of soil microbiome:** Taxonomically, all soil samples had bacterial  
141 populations from 50-57 phylum. Bacteria belonging to Proteobacteria and Actinobacteria were  
142 the most predominant bacteria, comprising over 60% of the total microbial community in each of  
143 the samples analyzed (Figure 1).

144 Hierarchical structure analysis was performed on the normalized genus level abundance  
145 data using Ward's linkage method to investigate the link between soil microbiota and plant  
146 type/soil pH and depth. Results indicate that the soil samples analyzed can be divided into six  
147 major clusters, after which the linkage distance separating the sub-clusters is small (Figure 2).  
148 Table 2 describes the members of each cluster and K-means clustering confirms the results. While  
149 overall the samples from individual locations from each of the horizon are clustered together or  
150 are in close clusters, samples from horizon B of location 10 (Pine tree vegetation) and horizon A  
151 of location 12 (Tulip tree) have unique microbiota to form its own cluster. ANOVA analysis  
152 indicates that the mean abundance for all the genera within a cluster are statistically different  
153 between clusters ( $p < 0.05$ ), except for the abundance of seven genera (Table S2). The seven  
154 genera whose abundance are not statistically different between clusters ( $p > 0.05$ ) are  
155 *Nitrosopumilus*, *Carboxydotherrmus*, unclassified genera derived from Deltaproteobacteria,  
156 *Pelotomaculum*, *Oceanicola*, *Thermotoga*, and *Bdellovibrio* (Table S2).



157 Table 3 shows the abundance of the top 30 microbial genera in the representative samples  
158 from each of the clusters and the average abundance of the organisms across all the 24 samples  
159 analyzed. Results show that *Bradyrhizobium* and *Candidatus Solibacter*, both Gram-negative  
160 bacteria, are the most abundant microorganism in the soil samples analyzed. *Streptomyces* and  
161 *Mycobacterium* are the two most abundant Gram-positive bacteria found in the soil samples.  
162 Figure 3 provides an overview of the total microbial community present in the soil sample obtained  
163 from the top horizon near Oak tree (sample 1A). As described earlier, the primary difference in  
164 microbial community in all the samples is the % abundance of individual organisms.

### 165 **Functional characterization of soil microbiome**

166 A heat map illustrating the functional annotation of sequence reads containing predicated proteins  
167 of known functions across all the 24 soil samples is shown in Figure 4. Variation was observed  
168 between samples primarily related to proteins involved in virulence, disease and defense; cell wall  
169 and capsule; membrane transport; DNA metabolism; and respiration. Among the functional  
170 categories identified by MG-RAST, the five most dominant categories based on the relative  
171 abundance of assigned reads were carbohydrates ( $13.3\pm 0.4\%$ ), the clustering-based subsystems  
172 (functional coupling evidence but unknown function;  $12.9\pm 0.2\%$ ), amino acids and derivatives  
173 ( $9.6\pm 0.3\%$ ), miscellaneous ( $6.8\pm 0.2\%$ ), and protein metabolism ( $7.7\pm 0.3\%$ ).

174 Relative abundance of the predicated proteins annotated at subsystem level 2 for each of  
175 the soil samples is presented in the supplemental table (Table S3). Hierarchical structure analysis  
176 was performed on the normalized values, similar to that performed for taxonomic data. Results  
177 indicate that the soil samples can be divided into 5 clusters, after which the linkage distance  
178 separating the sub-clusters is small (Figure 5). Table 4 describes the members of each cluster and  
179 K-means clustering confirms the results. Similar to taxonomic clustering, samples from horizon

180 B of location 10 (Pine tree vegetation) and horizon A of location 12 (Tulip tree) have unique  
181 composition of functional proteins to form its own cluster.

182 Table 5 shows the abundance of top 30 predicated proteins in the representative samples  
183 from each of the clusters. Results show that unidentified proteins involved in regulation and cell  
184 signaling comprise nearly 1 in 5 proteins predicated from the sequences. Nearly 6% of the  
185 predicated proteins are from the miscellaneous SEED category comprising a diverse set of genes  
186 identified during investigation of plant-prokaryote interactions by a project at the Department of  
187 Energy (DOE), USA (Thureborn et al. 2016). Protein biosynthesis, central carbohydrate  
188 metabolism, and resistance to antibiotics and toxic compounds were the other top predicated  
189 functions of the proteins. Figure 6 provides an overview of the functional hierarchical profiles of  
190 the microbiome from the top horizon soil obtained near Oak tree (sample 1A) analyzed at  
191 subsystem level 3.

### 192 **Linking diversity to function**

193 To identify the key microorganisms playing significant role in the biochemistry of soil,  
194 Refseq and Subsystems analysis were performed together on MGRAST platform. The Subsystem  
195 analysis was performed at level 3 wherever possible. Top 5 genera having the largest quantity of  
196 annotated reads within each of the metabolic class were identified (Table S4). Data indicates that  
197 *Bradyrhizobium*, *Rhodopseudomonas*, and *Burkholderia* are the key bacteria within the soil  
198 microbiota. Both, *Bradyrhizobium* and *Rhodopseudomonas* are top contributors in 24 of the 44  
199 metabolic classes analyzed (Figure 7). *Burkholderia* is a top organism in 16 of the metabolic  
200 classes (Figure 7).

201 From an agricultural perspective, Nitrogen, Phosphorus, Sulphur and Iron metabolic  
202 pathways are significant. In the Nitrogen, Iron and Sulphur pathways, beyond the three genera  
203 identified, *Mycobacteria* also plays a significant role. Organisms from *Anaeromyxobacter* and  
204 *Aromatoleum* genera are key contributors in the nitrosative stress and dissimilatory nitrile  
205 reductase pathways respectively (Table S4). Organisms from *Sorangium* genera have the most  
206 genes coding for Sulphate reduction associated complexes. Similarly, organisms from *Cupriavidus*  
207 and *Pseudomonas* genera are other top bacteria involved in Phosphate pathways (Table S4). In  
208 Iron pathways, *Bacillus*, *Frankia*, and *Pseudomonas* were the top genera involved (Table S4). The  
209 catabolic genes related to the degradation of xenobiotics were also annotated and linked to the  
210 microbial genera. Beyond *Bradyrhizobium*, *Rhodopseudomonas*, and *Burkholderia*, bacteria from  
211 *Pseudomonas* and *Cupriavidus* play a key role in degradation of xenobiotic compounds (Table  
212 S4). Results indicate that for each of the biochemical functions, there is redundancy within the  
213 soil microbiome.

## 214 **DISCUSSION:**

215 We investigated the microbial structural and functional diversity within the top acidic soil  
216 associated with a wide variety of plants. Results indicate that irrespective of the level of acidity  
217 in the soil, most of the microorganisms associated with the soil generally remains the same. The  
218 differences observed between soil samples, could be attributed to the abundance of individual  
219 organism's present in the soil based on the soil chemistry and the vegetation present. The change  
220 in microbial abundance results in change in the abundance of functional genes within the soil  
221 microbiome. Literature is replete with scientific studies showing soil microbiome changes with the  
222 structure of the soil (eg. Fierer and Jackson, 2006; Fierer et al. 2012; Mendes and Tsai, 2018; Shah  
223 et al. 2021). Based on our results, we suggest that one needs to consider whether the type of

224 organisms present in the soil are different or if the abundance of individual organisms is different  
225 before reaching the conclusions related to microbiome difference amongst different soil samples.  
226 Further, current methods of calculating alpha and beta diversity may not capture the true  
227 similarities in the microbiome from different soil types. As further advances are made in the next-  
228 generation sequencing techniques, we believe similarities in the microbiome across soil type could  
229 become more evident.

230 Taxonomically, prior research has shown that Gram-negative organisms are predominant  
231 organisms present in the soil (Shah and Subramaniam, 2018). Results obtained in the current study  
232 supports the prior observation. When one considers similar observations in microbiome studies  
233 conducted in marine environments, and even in human, fish and animals, a theme starts to emerge  
234 - in the microbial communities across the matrices, Gram-negative bacteria are the predominant  
235 organisms.

236  
237 Functionally, high levels of genes attributed to regulation and cell signaling (level 1) appear  
238 to be an identifying indicator for acidic soils. cAMP is a major gene annotated to this category.  
239 Delmont et al. (2012) reported abundance of cAMP related annotation within the soil metagenome.  
240 Considering acidic soil is poor in nutrition and the northeast region of the United States has varying  
241 weather patterns, soil bacteria might be required to deal with constantly fluctuating substrates and  
242 environmental conditions. cAMP is a universal cell energy and metabolism regulator. Higher level  
243 of this and other genes involved in regulation and cell signaling can be attributed to the requirement  
244 of bacteria to adapt to the changing soil chemistry. Surprisingly, we noticed low levels of the  
245 abundance of genes related to nutrient cycling (sulfur metabolism, potassium metabolism, iron  
246 acquisition and metabolism, and phosphorous metabolism). Genes annotated to virulence disease  
247 and defense were significantly prevalent in the soil samples analyzed. The cluster-based

248 subsystems contain diverse functions, such as resistance to antibiotics and toxic compounds, and  
249 pathogenicity islands.

250  
251 Results of our study indicate that in the acidic soil, *Bradyrhizobium*, *Rhodopseudomonas*,  
252 and *Burkholderia* are the major organisms in the soil involved in the nutritional ecosystem  
253 functioning. *Bradyrhizobium* and *Candidatus solibacter* are taxonomically the most abundant  
254 organisms in the soil samples analyzed. Collectively, it is evident from taxonomic and functional  
255 analysis of the soil microbiome, bacteria from *Bradyrhizobium* are highly critical to maintaining  
256 soil fertility, irrespective of soil type. Analyzing the microbiota present in 52 soil samples from  
257 different countries, Shah and Subramaniam (2018) found that bacteria from *Bradyrhizobium*  
258 genera were the most abundant organisms in the microbiota. The structure-function linkage results  
259 indicate that the organism is not only responsible for nitrogen fixation and other pathways in N  
260 cycle, but also plays a key role in S and Fe cycles, and degradation pathways of xenobiotic  
261 compounds. *Bradyrhizobium* bacteria are present as symbiotic and non-symbiotic organisms in the  
262 soil, and literature is replete with the importance of the organism in the Nitrogen cycle (Ormeño-  
263 Orrillo and Martínez-Romero, 2019). Many strains of *Bradyrhizobium* are used commercially to  
264 improve crop production (Environmental Protection Agency, n.d.). We suggest that the beneficial  
265 impact of the organism in improving soil fertility could also be attributed to its role in other  
266 biochemical pathways.

267 Acidic soils provide a unique environment for soil microorganisms due to iron, manganese  
268 and aluminum toxicity, low nitrogen, phosphorus, and molybdenum levels, toxic levels of phenolic  
269 acids, and hydrogen ion toxicity (Kidd and Proctor, 2001; Shah et al. 2011). Often to overcome  
270 this issue, nitrogen fertilizers and other chemicals are used to improve soil fertility, but these  
271 methods can cause other environmental issues including increase in nitrous oxide emissions (Xu,

272 2014). As a substitute to the use of chemicals for improving soil fertility and crop production, we  
273 suggest to the scientific community to study the possibility of using consortia of organisms  
274 including *Bradyrhizobium*, *Rhodopseudomonas*, and *Burkholderia*. Considering the importance  
275 and ubiquity of these organisms in the soil, the consortia could be used by farmers across the globe,  
276 irrespective of soil chemistry and geographical location.

277         Next-generation sequencing methods are increasingly used to study how the soil  
278 microbiome responds to changes in environmental conditions or to addition of contaminants in the  
279 soil. We suggest that in addition to analyzing general community-based diversity changes,  
280 scientists should specifically look for changes in the *Bradyrhizobium*, *Rhodopseudomonas*, and  
281 *Burkholderia* population to understand the impact. Our results suggest that changes in abundance  
282 of these organisms may greatly impact the soil fertility.

283         Considering that the soil samples analyzed were from the West Chester, PA region only,  
284 further studies are warranted using acidic soil samples from across the globe to validate the  
285 observations. Nevertheless, the metagenomic data reported here furthers our knowledge on the  
286 acidic soil microbial communities at structural and functional level. There is a large degree of  
287 similarity in the soil microbiome associated with different vegetation and soil pH. Increasing our  
288 attention to similarities in soil microbiome may allow us to further the biotechnological potential  
289 of microbial based products to improve soil fertility in the future.

290

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293 **Conflicts of Interest:** Authors have no conflicts of interest to declare.

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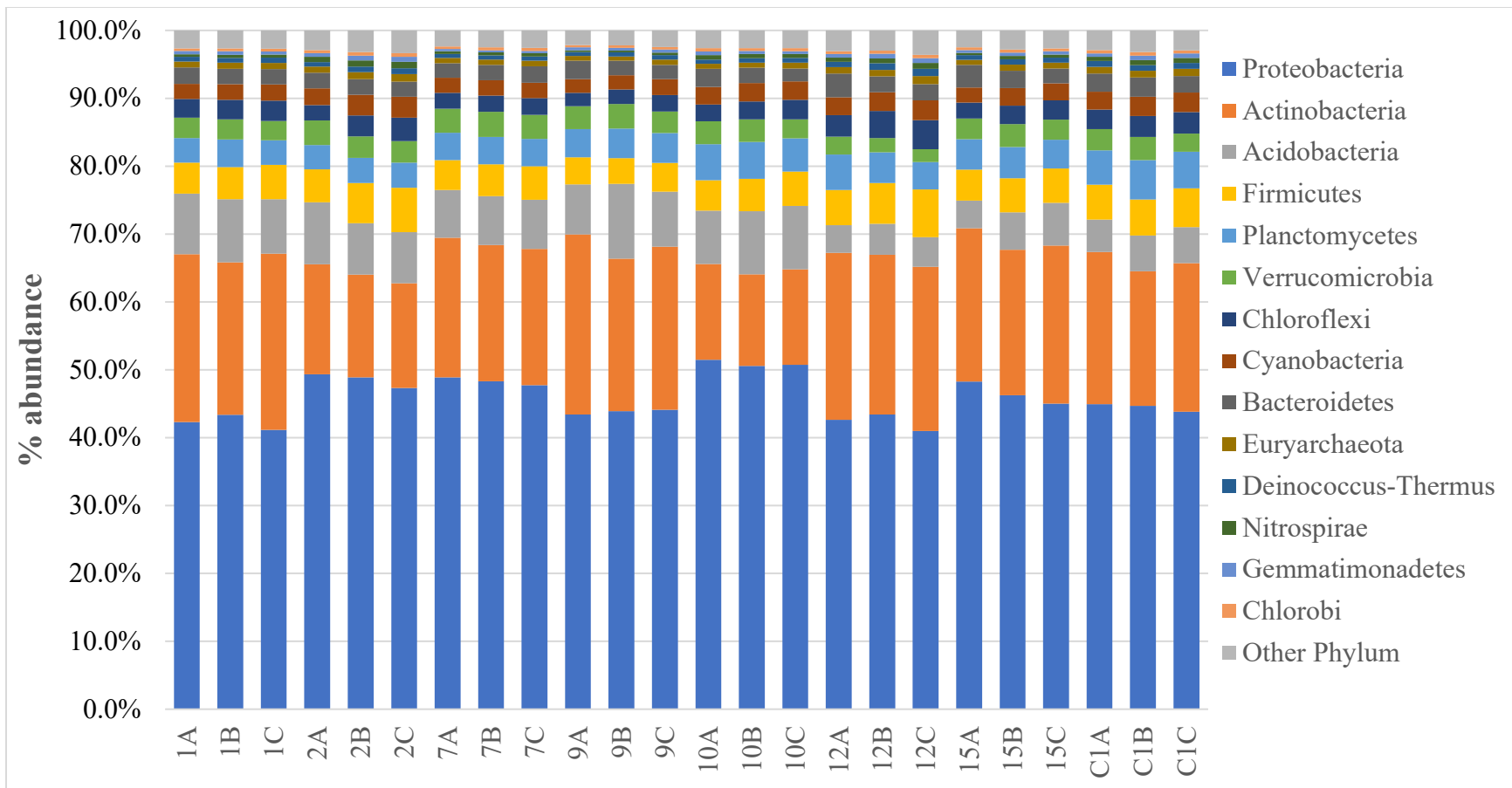
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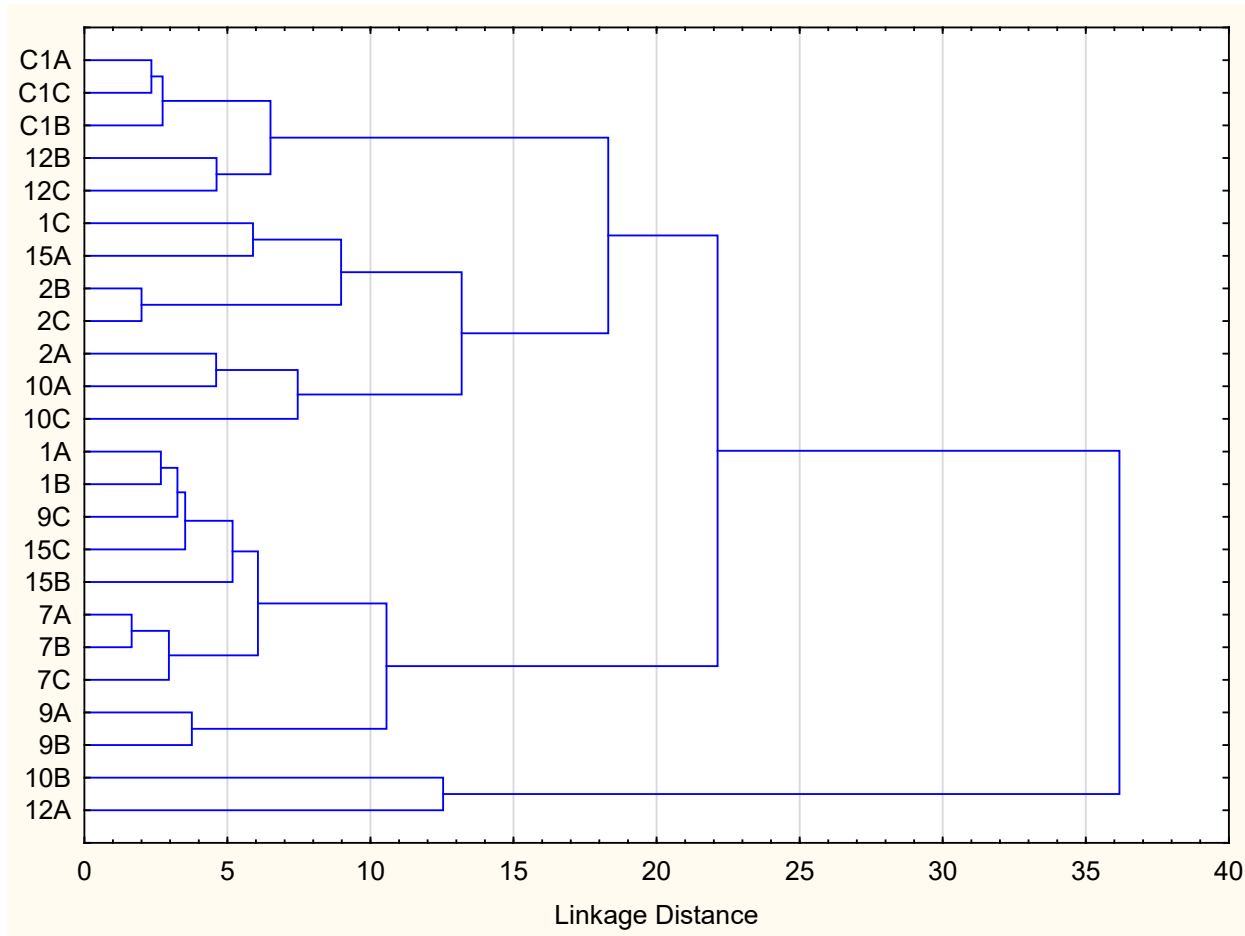
354 Vogel, T. M., Simonet, P., Jansson, J. K., Hirsch, P. R., Tiedje, J. M., Van Elsas, J. D., ... &  
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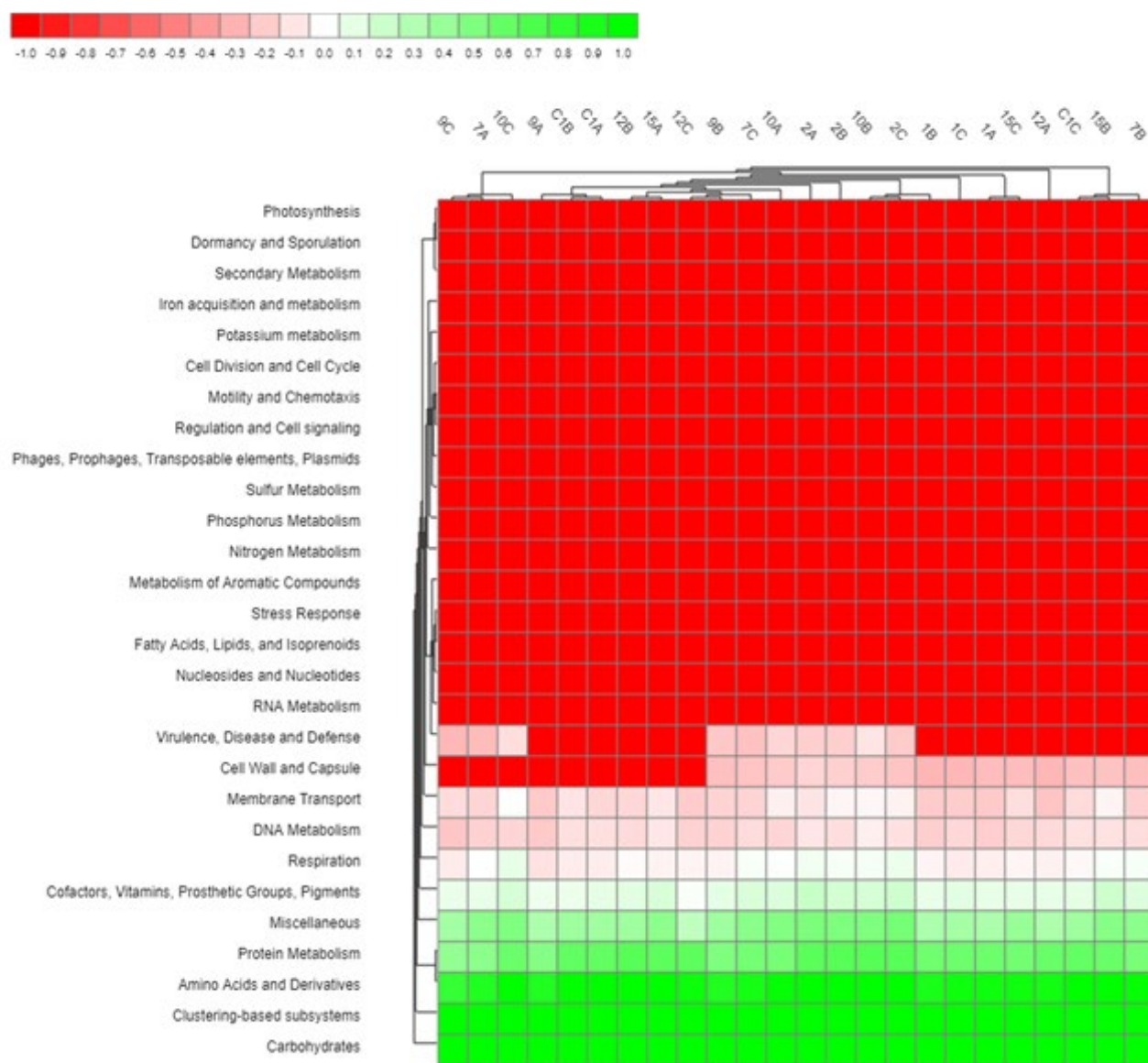


**Figure 1.** Phyla comparison of soil microbiomes collected from eight sampling locations at three different horizons (A, 0-10 cm; B, 10-20 cm; C, 20-30 cm).

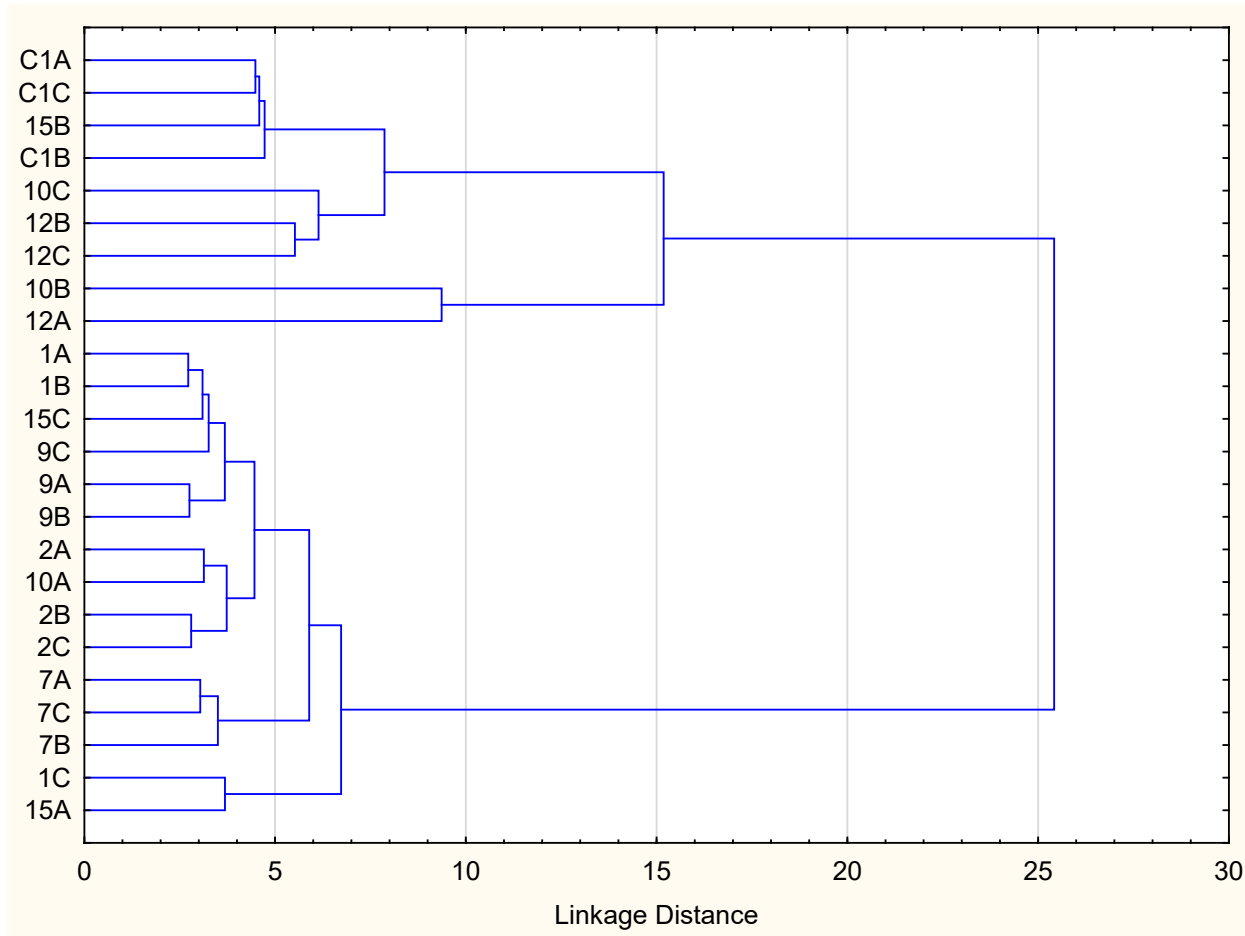


**Figure 2.** Hierarchical structure analysis used to identify the number of major clusters and the cluster members based on the normalized genus level abundance data.



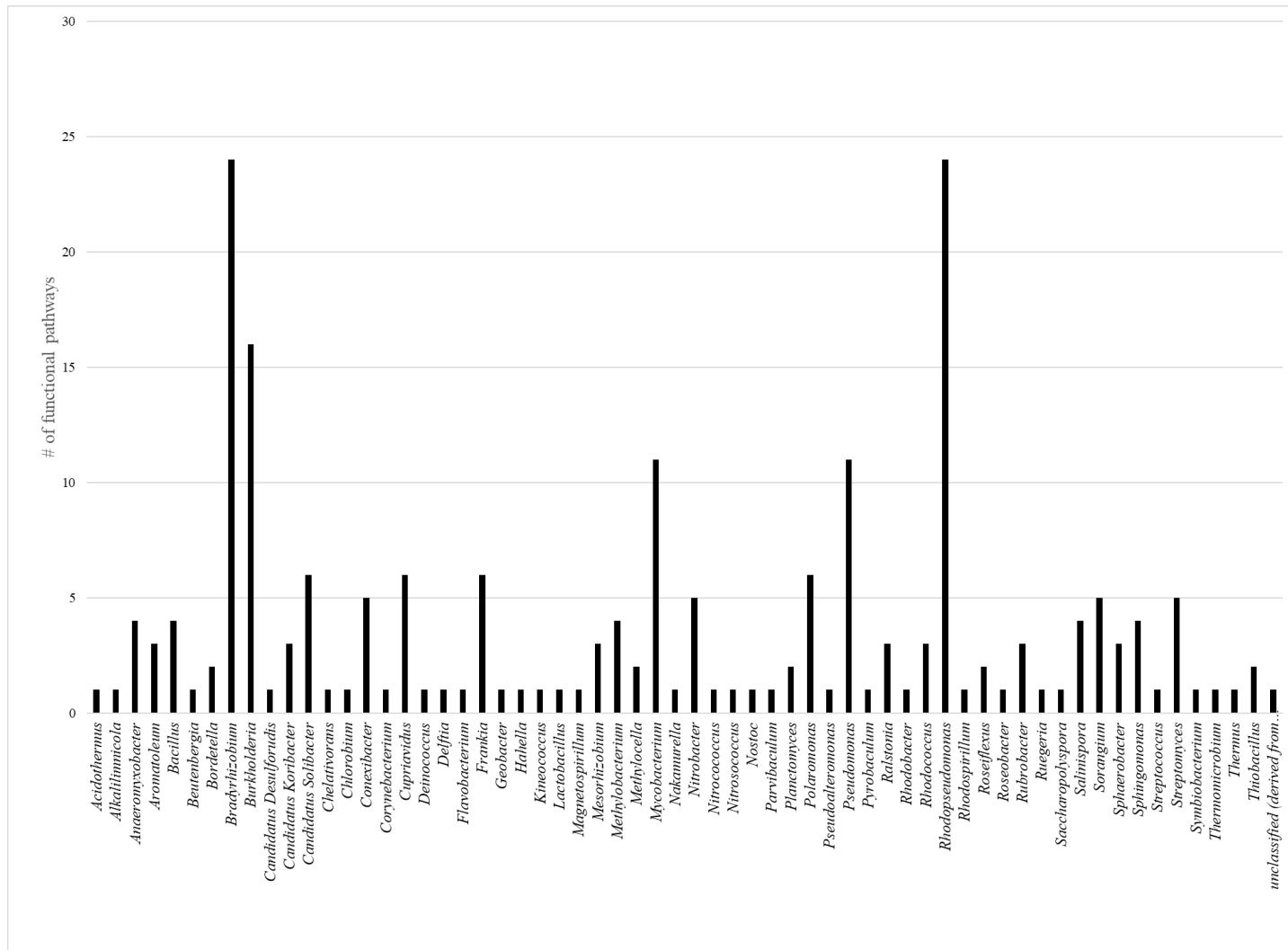


**Figure 4.** Heat map showing the differential abundance of functional categories (subsystem Level 1) between different soil samples.



**Figure 5.** Hierarchical structure analysis used to identify the number of major clusters and the cluster members based on the normalized predicated proteins annotated at subsystem level 2.





**Figure 7.** Total number of functional pathways top microbial genera are involved in within the acidic soil samples analyzed.



Site No.	Plant	Coordinates	Top	Middle	Bottom
			(A)	(B)	(C)
1	Oak	39°57'10.4"N 75°35'54.9"W	5.7 ± 0.3	5 ± 0.2	4.4 ± 0.2
2	Douglas Fir	39°56'59.1"N 75°35'38.9"W	4.1 ± 0.06	4.3 ± 0.06	4.4 ± 0.1
7	Pine - 1	39°55'57.1"N 75°36'09.8"W	5.4 ± 0.1	5.4 ± 0.03	5.4 ± 0.04
9	Pine - 2	39°56'55.9"N 75°36'06.6"W	6.2 ± 0.2	5.9 ± 0.1	5.7 ± 0.1
10	Pine - 3	39°56'22.7"N 75°35'36.5"W	5.8 ± 0.04	5.9 ± 0.1	5.9 ± 0.09
12	Tulip Tree	39°57'09.0"N 75°35'58.2"W	5.8 ± 0.1	6.1 ± 0.1	6 ± 0.6
15	Willow Oak	39°57'00.4"N 75°36'01.3"W	6.2 ± 0.1	6.3 ± 0.05	6.2 ± 0.1
C1	Grass	39°57'04.8"N 75°35'58.7"W	5.7 ± 0.2	6.1 ± 0.08	6.2 ± 0.03

**Table 1.** Vegetation type, coordinates, and pH for the locations from where the soil samples were obtained along with the pH values at each of the three sampling depths for each location. (0–5 cm, Horizon A; 6-15 cm, Horizon B; 16–30 cm, Horizon C)

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Cluster 1	10B
Cluster 2	C1A, C1B, C1C, 12B, 12C
Cluster 3	12A
Cluster 4	2A, 10A, 10C
Cluster 5	1A, 1B, 7A, 7B, 7C, 9A, 9B, 9C, 15B, 15C
Cluster 6	1C, 2B, 2C, 15A

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**Table 2.** Clustering of the soil samples based on the hierarchical structure analysis of taxonomic data at genera level. Soil samples were collected from eight sampling locations at three different horizons (A, 0-10 cm; B, 10-20 cm; C, 20-30 cm).

<b>Genus</b>	<b>C1A</b>	<b>1A</b>	<b>2A</b>	<b>10B</b>	<b>12A</b>	<b>15A</b>	<b>Average</b>
<i>Bradyrhizobium</i>	2.7	4.6	5.0	5.1	2.7	4.7	4.4
<i>Candidatus Solibacter</i>	2.9	4.1	4.4	4.5	2.3	2.4	3.6
<i>Streptomyces</i>	3.6	4.2	2.4	2.2	4.1	3.6	3.4
<i>Mycobacterium</i>	3.0	2.8	3.6	3.0	2.7	3.1	2.8
<i>Rhodopseudomonas</i>	1.9	2.9	3.3	3.2	1.9	3.1	2.8
<i>Burkholderia</i>	1.9	2.1	2.2	2.6	1.7	2.1	2.2
<i>Candidatus Koribacter</i>	1.1	2.7	2.7	2.7	1.0	0.9	1.9
<i>Conexibacter</i>	2.0	2.5	1.1	0.7	2.3	2.0	1.7
<i>Frankia</i>	1.7	2.3	1.3	1.2	1.9	1.8	1.8
<i>Methylobacterium</i>	1.4	1.5	2.0	1.8	1.3	1.7	1.7
<i>Nitrobacter</i>	1.0	1.4	1.7	1.7	0.9	1.5	1.4
<i>Anaeromyxobacter</i>	1.5	1.0	1.1	1.3	1.4	1.1	1.3
<i>Geobacter</i>	1.3	1.0	1.2	1.3	1.2	1.2	1.2
<i>Planctomyces</i>	1.2	0.9	1.0	1.4	1.3	1.1	1.1
<i>Mesorhizobium</i>	1.1	0.9	1.0	1.0	1.2	1.2	1.0
<i>Acidobacterium</i>	0.6	1.6	1.5	1.6	0.6	0.6	1.1
<i>Gemmata</i>	1.0	0.9	0.8	1.4	1.1	1.0	1.1
<i>Rhodococcus</i>	1.1	1.1	0.8	0.7	1.1	1.0	1.0
<i>Pseudomonas</i>	1.0	0.9	0.9	1.1	0.9	1.0	0.9
<i>Chthoniobacter</i>	0.9	1.0	1.2	1.0	0.7	1.0	1.0
<i>Rhizobium</i>	0.9	0.8	1.0	1.0	0.9	1.1	1.0

<i>Sinorhizobium</i>	1.0	0.7	1.0	0.9	0.9	0.9	0.9
Unclassified (derived from Verrucomicrobia subdivision 3)	0.9	0.8	1.1	1.1	0.8	0.8	0.9
<i>Sorangium</i>	1.1	0.6	0.7	0.9	1.0	1.0	0.8
<i>Cupriavidus</i>	0.9	0.7	0.9	1.0	0.8	0.9	0.8
<i>Rhodopirellula</i>	1.0	0.7	0.7	0.9	1.0	0.9	0.8
<i>Pirellula</i>	1.0	0.6	0.6	0.9	1.1	0.8	0.8
<i>Myxococcus</i>	1.0	0.7	0.7	0.8	0.9	0.8	0.8
<i>Bacillus</i>	0.7	0.6	0.9	0.8	0.9	0.8	0.8
<i>Roseiflexus</i>	0.9	0.7	0.6	0.7	0.9	0.7	0.8

**Table 3.** Relative Abundance of the top 30 bacterial genera in the representative soil samples from each cluster and average values across all the soil samples analyzed.

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Cluster 1	12A
Cluster 2	C1A, C1B, 10C, 12B, 12C, 15B
Cluster 3	10B
Cluster 4	1C, 7A, 7B, 7C, 15A, 15C
Cluster 5	C1C, 1A, 1B, 2A, 2B, 2C, 9A, 9B, 9C, 10A

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**Table 4.** Clustering of the soil samples based on the functional annotation of the sequence reads by MG-RAST at subsystem level 2.

Soil samples were collected from eight sampling locations at three different horizons (A, 0-10 cm; B, 10-20 cm; C, 20-30 cm).

<b>Level 1</b>	<b>Level 2</b>	<b>C1A</b>	<b>1A</b>	<b>2A</b>	<b>10B</b>	<b>12A</b>	<b>15A</b>
Regulation and Cell signaling	NULL	21.4	21.6	21.7	21.6	21.9	21.8
Miscellaneous	Plant-Prokaryote DOE project	6.0	5.9	6.2	6.2	5.9	6.1
Protein Metabolism	Protein biosynthesis	4.7	4.6	4.5	4.3	4.8	4.6
Carbohydrates	Central carbohydrate metabolism	4.2	4.2	4.1	4.1	4.0	4.1
Virulence, Disease and Defense	Resistance to antibiotics and toxic compounds	2.8	3.0	3.3	3.7	2.9	2.8
DNA Metabolism	DNA repair	2.7	2.4	2.5	2.7	2.5	2.6
Amino Acids and Derivatives	Lysine, threonine, methionine, and cysteine	2.5	2.5	2.5	2.5	2.6	2.6
Respiration	Electron donating reactions	2.3	2.2	2.2	2.1	2.4	2.1
RNA Metabolism	RNA processing and modification	2.2	2.1	2.1	2.1	2.3	2.2
Cofactors, Vitamins, Prosthetic Groups, Pigments	Folate and pterines	2.0	2.0	1.9	1.8	2.0	2.0
Amino Acids and Derivatives	Branched-chain amino acids	1.9	2.0	1.9	1.8	1.9	1.9
Membrane Transport	ABC transporters	1.6	1.6	1.6	1.8	1.5	1.7
Amino Acids and Derivatives	Arginine; urea cycle, polyamines	1.8	1.5	1.5	1.6	1.6	1.6
Nucleosides and Nucleotides	Purines	1.7	1.6	1.6	1.6	1.6	1.6
Carbohydrates	Monosaccharides	1.6	1.6	1.4	1.4	1.8	1.6
Carbohydrates	Di- and oligosaccharides	1.4	1.7	1.4	1.3	1.5	1.4
Carbohydrates	One-carbon Metabolism	1.5	1.4	1.4	1.4	1.5	1.3
Protein Metabolism	Protein degradation	1.4	1.4	1.3	1.5	1.4	1.3

Fatty Acids, Lipids, and Isoprenoids	Fatty acids	1.2	1.3	1.4	1.3	1.3	1.3
Amino Acids and Derivatives	Aromatic amino acids and derivatives	1.3	1.3	1.3	1.3	1.3	1.3
Cell Wall and Capsule	Capsular and extracellular polysacchrides	1.1	1.2	1.2	1.1	1.3	1.1
Stress Response	Oxidative stress	1.1	1.0	1.1	1.2	1.1	1.2
DNA Metabolism	DNA replication	1.1	1.1	1.1	1.1	1.1	1.2
Carbohydrates	Fermentation	0.9	1.0	1.1	1.0	1.1	1.1
RNA Metabolism	Transcription	1.0	1.0	0.9	0.8	1.0	0.9
Amino Acids and Derivatives	Glutamine, glutamate, aspartate, asparagine; ammonia assimilation	1.0	0.9	0.9	1.0	0.9	1.0
Respiration	Electron accepting reactions	0.8	1.0	0.9	0.9	0.9	0.9
Cell Wall and Capsule	Gram-Negative cell wall components	0.8	1.0	1.0	1.0	0.8	0.8
Phages, Prophages, Transposable elements, Plasmids	Phages, Prophages	0.8	0.8	0.9	0.8	0.7	0.8
Nucleosides and Nucleotides	Pyrimidines	0.9	0.9	0.8	0.7	0.8	0.8

**Table 5.** Relative Abundance of the top 30 functional genes (level 2) bacterial genera in the representative soil samples from each cluster.