

Characterization of Natural and Affected Environments

Organic carbon amendments affect the chemodiversity of soil dissolved organic matter and its associations with soil microbial communities

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1 **Organic carbon amendments affect the chemodiversity of soil dissolved organic**
2 **matter and its associations with soil microbial communities**

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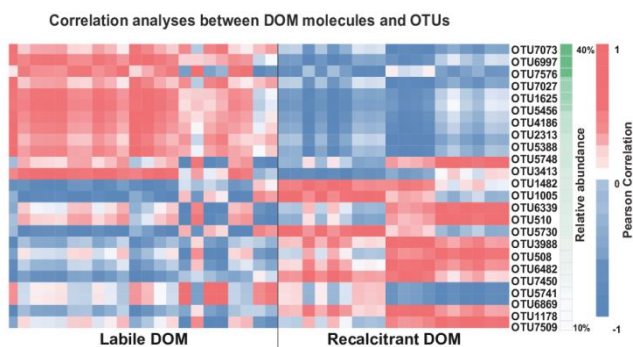
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18

19 **Abstract**

20 The “4 per mil” initiative recognizes the pivotal role of soil in carbon re-
21 sequestration. The need for evidence to substantiate the influence of agricultural
22 practices on chemical nature of soil carbon and microbial biodiversity has become a
23 priority. However, owing to the molecular complexity of soil dissolved organic matter
24 (DOM), specific linkages to microbial biodiversity have eluded researchers. Here, we
25 characterized the chemodiversity of soil DOM, assessed the variation of soil bacterial
26 community composition (BCC) and identified specific linkages between DOM traits
27 and BCC. Sustained organic carbon amendment significantly ($P < 0.05$) increased total
28 organic matter reservoirs, resulted in higher chemodiversity of DOM and emergence of
29 recalcitrant moieties ($H/C < 1.5$). In the meantime, sustained organic carbon
30 amendment shaped the BCC to a more eutrophic state while long-term chemical
31 fertilization directed the BCC towards an oligotrophic state. Meanwhile, higher
32 connectivity and complexity were observed in organic carbon amendment by DOM-
33 BCC network analysis, indicating that soil microbes tended to have more interaction
34 with DOM molecules after organic matter inputs. These results highlight the potential
35 for organic carbon amendments to not only build soil carbon stocks and increase their
36 resilience but also mediate the functional state of soil bacterial communities.

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TOC / Abstract Art

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42 **Introduction**

43 Soil organic matter (SOM) represents the largest pool (1500 ~ 2300 Pg C) of
44 terrestrially organic carbon in the biosphere ¹, this being more than two times the
45 amount of carbon in the atmosphere ². Hosting the largest diversity of organisms on
46 land, soils play a pivotal role in regulating major global biogeochemical cycles (carbon,
47 nutrients and water) ³. As supporters of human food production systems, soils
48 contribute to the economic status of nations ⁴. The role of SOM as an essential resource
49 for heterotrophic life, its directing influence on soil food webs and the subsidiary effects
50 it has on physical and chemical soil attributes (such as, structure, moisture content,
51 nutrient availability, infiltration capacity) affirm SOM status as a key indicator of soil
52 quality ⁵.

53 Despite the importance of the soil carbon reservoir, manifold pressures have
54 resulted in substantial degradation of soil and SOM ^{6,7}. A further antagonism to SOM
55 depletion, of relevance to modern agriculture, is the use of inorganic fertilizers. With
56 literature spanning almost a century, and field measurements, Mulvaney et al. ⁸
57 evidenced chemical-N fertilizers ability to increased mineralization of native soil
58 carbon. This enhancement in microbial utilization of SOC lead to demonstrable
59 depletions of both soil carbon and soil nitrogen stocks. Depletion of SOC under
60 continuous cultivation has led to the declines in crop yields in multiple cropping
61 systems ^{9,10}. The most obvious means to increase soil carbon is to augment new carbon,
62 here numerous candidates (e.g. composts ¹¹, animal sludges ¹², sewage sludges ^{13, 14}),
63 plant residues ¹⁵ and biochars) have potential. Significantly, not only can these
64 amendments assist in rebuilding soil carbon stocks, they also stimulate microbial
65 populations ¹⁶, that in turn can influence the delivery of soil ecosystem services ^{17, 18}.

66 Dissolved organic matter (DOM) is an important constituent of SOM as it provides
67 soluble organic substrates that support and sustain heterotrophic microbial communities
68 ¹⁹⁻²¹. The variation of DOM composition regulates microbial growth and activity, and
69 *vice versa* ^{22, 23}, therefore the association between individual DOM molecules and soil
70 microbes lies at the heart of the DOM cycle ²⁴. Our understanding of how soil microbes
71 and DOM interact and the implications of these interactions for: soil microbial ecology,
72 the carbon cycle, and the delivery of soil ecosystem services, are fundamental to our
73 ability to use soils sustainably.

74 Only recently have tools emerged that can resolve the vast chemodiversity of
75 DOM with analytical precision. Specifically, Fourier Transform Ion Cyclone
76 Resonance Mass Spectrometry (FT-ICR MS) has paved the way for in-depth
77 appreciation organic carbon profiles. Recently, this technique has successfully been
78 used to characterize DOM chemodiversity in different environments, specifically,
79 ocean water ²⁵, lake water ²⁶, soil pore water ²⁷, sediment ²⁸, and the atmospheric
80 particulates ²⁹. Several studies have characterized DOM chemodiversity in forest soils
81 ^{30,31} by FT-ICR MS, and they have shown that soil DOM chemodiversity changed with
82 soil depth and significantly affected by soil pH and nitrate. However, few of these
83 publications have provided any insight into the DOM chemodiversity and relationships
84 between DOM and microbial communities in agricultural soils.

85 In this study we investigated the DOM molecular composition and bacterial
86 community composition in an agricultural field with long-term fertilization. We
87 characterized chemodiversity of soil DOM and analyzed its relationship with soil
88 bacterial community. We hypothesized that the soil DOM chemodiversity significantly
89 associated with soil bacterial community composition under long-term organic carbon
90 amendments.

91 **Materials and methods**

92 **Field experiment.**

93 The soil samples were collected at experimental station (37°20'N, 116°38'E) of
94 the Chinese Academy of Agricultural Sciences (CAAS), located in Dezhou, Shandong
95 Province, China. The annual average temperature in the experiment site was 12.9°C,
96 and the annual average rainfall was 522 mm. A long-term carbon-amendment/fertilizer
97 experiment was conducted on an agricultural field with fluvo-aquic soil. The soil
98 texture was clay loam, and the soil moisture capacity was 23%. The field experiment,
99 from which samples were drawn, comprised randomized block of treatments (N = 3)
100 maintained under a continuous rotation of winter-wheat/summer-maize for a decade.
101 Eight treatments were instated (Figure S1) ³². Detailed information of the application
102 rates of fertilizers was shown in Table S2. The fertilizers of phosphorus and potassium
103 were applied as basal fertilizers. Annually (each June), all plots received the same
104 application of basal fertilizer (superphosphate (600 kg hm⁻²) and potassium sulphate
105 (240 kg hm⁻²)). Control treatments (CK) received basal fertilizer but no further carbon
106 and nitrogen amendment. While additional inorganic fertilizer (0.5N and 1N), in the
107 form of urea, was applied at 65 kg hm⁻² (0.5N) and 130 kg hm⁻² (1N) to provide low
108 and high inorganic fertilizer regimes, respectively. To establish sewage sludge
109 augmented plots, urea was again applied at the lower application rate (65 kg hm⁻²) and
110 sewage sludge applied (dry weight equivalent) at 4.5 t hm⁻² (0.5SS), 9 t hm⁻² (1SS), 18
111 t hm⁻² (2SS) and 36 t hm⁻² (4SS). Finally, plots containing chicken manure were
112 established with urea applied at the lower application rate (65 kg hm⁻²) and chicken
113 manure applied (dry weight equivalent) at 10 t hm⁻² (CM). Thus, the CK and 0.5N
114 treatments provided points of reference to discern SOC and BCC shifts directed by, i)
115 urea fertilizer and ii) carbon amendments. In each plot, the fertilizers were spread over

116 the fields and mixed well with the soil (0 - 15 cm) immediately following the
117 application. Sewage sludge (SS) was collected from Beijing sludge disposal plants, and
118 then underwent composting. The chicken manure (CM) was purchased from a fertilizer
119 company (Hebei Beautiful Day Fertilizer Technology Co., LTD.) in Hebei Province.
120 Soil samples were collected in 2015 (ten months after their annual fertilizer and/or
121 organic carbon amendments were added). Physical and chemical properties of soil,
122 sewage sludge and chicken manure are provided in Table S1. Surface soil (0-15 cm)
123 samples were collected from plots (~ 2 kg per plot). Each soil sample was a mix of ten
124 soil cores per plot. After sampling, soil samples were immediately transported to
125 laboratory on dry ice and stored at -80 °C.

126 **Chemical characterization.**

127 The pH of soil was measured in a solid-to-deionized water ratio of 1:2.5 using a
128 digital pH meter (PHS-3C, Shanghai Lida Instrument Company, China)³³. Total carbon
129 (TC) and total nitrogen (TN) contents were determined by dry combustion in an element
130 analyzer (Vario EL III - Elementar, Germany)³⁴. Briefly, 5 g air dried soil was placed
131 in a centrifuge tube (50 mL), and 25 mL Milli-Q water (18 MΩ) was added (1:5
132 solid:liquid ratio). The tubes were shaken in the shaker (170 rpm) and were centrifuged
133 at $2,800 \times g$ for 10min. The supernatant was filtered (0.45 μm) and kept in the 4 °C
134 until the determination of DOC and DTN concentrations by a TOC analyzer (Liquic
135 TOC - Elementar, Germany)³⁵. Subsamples were extracted with 2 M KCl solution, and
136 the concentration of nitrate and ammonium were determined using a continuous flow
137 analyzer (SAN++, Skalar, Holand)³⁶.

138 **Dissolved organic matter (DOM) analysis.**

139 The solid-phase extraction of dissolved organic matter (SPE-DOM) from soil was
140 performed as described in our previous work³⁷. More specifically, soil DOM was first

141 extracted with Milli-Q water (1:5 w/v) on a reciprocal shaker (170 rpm) for 8 h. Samples
142 were then centrifuged at $2,800 \times g$ for 10 min, and the supernatant were filtered through
143 a $0.45 \mu\text{m}$ mixed cellulose ester membrane. SPE cartridges (Bond Elut PPL, 500 mg, 6
144 mL, Agilent Technologies) were activated by sequentially rinsing with pure methanol
145 (mass spectrometry grade) and 0.01 M HCl ²¹. Acidified DOM samples (pH = 2) were
146 passed over the activated cartridges and then the cartridges were rinsed with acidified
147 Milli-Q water (pH = 2). After the cartridges were completely dried with ultrapure N₂
148 gas, DOM was eluted from the cartridges with methanol (5 mL). and stored at -20 °C
149 ²⁶. Extraction efficiencies were calculated by drying methanol eluate and re-dissolving
150 with ultrapure water. The extraction efficiencies were different among treatments (48
151 $\pm 11.2\%$, $65 \pm 20\%$, $49 \pm 18\%$ and $50 \pm 17.6\%$ on average for CK, N, SS and CM). The
152 final concentration of DOM was about 20 mg/L by methanol dilution. The molecular
153 composition of solid-phase extractable DOM (SPE-DOM) was analyzed using a 9.4 T
154 Bruker apex-ultra FT-ICR MS equipped with an electrospray ionization source (Bruker
155 Apollo II) applied in negative mode ³⁷. SPE-DOM was dissolved in methanol and
156 injected into the electrospray source at $3 \mu\text{L min}^{-1}$ by a syringe pump ^{38, 39}. PPL
157 extraction blanks and solvent blanks were prepared and analyzed to check for possible
158 contamination. Contaminated peaks in these blanks were removed from obtained DOM
159 profiles ^{23,33}. Detected mass peaks with S/N less than 6 were not considered in the
160 following data processing ^{23,33}.

161 **Bacterial community composition (BCC) analysis.**

162 DNA was extracted from soil (0.50 g) using the FastDNA Spin Kit for soil (MP
163 Biomedical, Santa Ana, California, USA) ³⁶. 16S rRNA gene Illumina sequencing was
164 performed on an Illumina Hiseq2000 platform at Novogene (Beijing, China).
165 Community DNA was amplified utilizing amplification primers F515 (5'-

166 GTGCCAGCMGCCGCGG-3') and R907 (5'-CCGTCAATTCMTTTRAGTTT-3')
167 targeting the V4-V5 region of the 16S rRNA³². The protocol described in our previous
168 work, was adopted to process and analyze the obtained sequenced data³². In summary,
169 raw reads were filtered, quantified, and subsequently analyzed using QIIME.
170 Operational taxonomic units (OTUs) were defined at the level of 97% similarity. The
171 alpha diversity index and β -diversity were calculated based on operational taxonomic
172 units (OTUs) table as described in our previous work^{36,40}.

173 **Statistical analysis.**

174 Absolute peak intensities of the FT-ICR MS spectra were normalized to the sum
175 of peak intensities of a given spectrum, and thereafter referred to as *relative* peak
176 intensities in the following statistical analyses²⁵. Compound groups were delineated by
177 the following parameters according to previous studies²¹: elemental ratios⁴¹,
178 aromaticity index (AI)⁴², double bond equivalence (DBE)⁴² and H/C cutoffs²⁶. A one-
179 way analysis of similarities (ANOSIM)²³ was performed (using R version 3.3.3) to
180 determine if different treatments resulted in significantly different DOM molecular
181 composition. The diversity index was calculated with R version 3.3.3 package 'vegan'.
182 Non-metric Multidimensional scaling (NMDS), based on the Bray-Curtis distance, was
183 performed to evaluate the overall pattern of DOM molecules among the treatments³².
184 Correlations among DOM community, BCC and environmental variables were
185 established using a Mantel test and redundancy analysis (RDA). NMDS and RDA were
186 performed using R version 3.3.3 in the 'vegan' package. Linear discriminant analysis
187 (LDA) effect size (LEfSe) was performed (using software sourced at:
188 <http://huttenhower.sph.harvard.edu/lefse/>) with LDA set at > 2.0, to indicate significant
189 difference⁴³ in DOM chemo-markers among treatments.

190 The BCC was expressed in terms of relative abundances of OTUs for each of the

191 phylogenetic resolutions from phylum to species²⁵. Non-metric multidimensional
192 scaling⁴⁴ (NMDS), based on Bray-Curtis distance, was used to compare BCC profiles
193 among treatments. For alpha diversity, the metrics of observed species (i.e. OTUs),
194 Chao 1, and Shannon index were calculated³². RDA analysis was conducted to
195 determine the significant environmental parameters that shaped soil BCC⁴⁵.

196 The specific links between DOM chemodiversity and BCC were revealed by
197 network analysis. Pairwise correlations were calculated using the ‘psych’ package in R
198 version 3.3.3 to determine the relationships between individual DOM molecule and
199 bacterial OTUs using Pearson product-moment correlation ($p < 0.05$)²³. *P*-values were
200 adjusted according to the false discovery rate to correct for multiple correlations⁴⁴.
201 Where correlations revealed a pairwise Pearson’s correlation coefficients $R > 0.6$ they
202 were considered statistically robust³². These values were then taken forward and
203 visualized in a network constructed using Cytoscape software (version 3.6.02)²³ and
204 Gephi 0.9.2^{13, 46}. The top interactions ($r \geq 0.6$ or $r \leq -0.6$) were prioritized to reduce
205 network complexity and thereby allow key linkages to be appreciated. This “funneling”
206 resulting in 24 DOM molecules and 16 OTUs with which to construct the network using
207 Cytoscape⁴⁷. Gephi 0.9.2 was used to generate the separate network plots using the
208 Force Atlas layout to connect DOM molecules and OTUs. Node sizes were correlated
209 to the number of edges they contained, which resulted in larger nodes for OTUs
210 compared to DOM molecules⁴⁸.

211 **Results and Discussion**

212 **Soil chemical properties.**

213 Long-term carbon-amendment or fertilizer application had a significant effect on
214 chemical characteristics of soil (Figure S2). The contents of total carbon (TC) and DOC
215 in sewage sludge (SS) and chicken manure (CM) treatments were higher than those in

216 chemical fertilizer (N) treatments and control (CK) (Figure S2a,b). Compared with
217 0.5N treatment ($69.12 \text{ mg}\cdot\text{kg}^{-1}$), 4SS and CM treatments increased DOC concentrations
218 by 1.93 and 1.63 times, respectively (Figure S2b). High doses of organic carbon
219 amendment (4SS and CM) increased TN and DTN significantly with the DTN increase
220 by 2.4 times in 4SS treatment compared with 0.5N treatments (Figure S2c, d). 4SS and
221 CM significantly decreased the soil C/N ratio ($P < 0.05$) (Figure S2e). NO_3^- -N
222 concentrations showed increasing trends with N application and organic carbon
223 application, and was significantly higher ($168 \text{ mg}\cdot\text{kg}^{-1}$) in the 4SS treatments compared
224 to the control ($22.8 \text{ mg}\cdot\text{kg}^{-1}$) (Figure S2f). NH_4^+ -N concentrations across all treatments
225 were not significantly different (Table S7). Soil pH ranged from 7.55 to 7.99 and no
226 significant differences were observed among treatments (Table S7). Researches have
227 demonstrated that sustained organic amendment could influence soil agroecosystem
228 characters and usually resulted in higher nutrient contents compared with N-containing
229 chemical fertilizer treatments¹¹. Our results are consistent with previous researches that
230 have demonstrated that long-term application of organic fertilizer influenced the
231 organic matter content in soil and improved soil quality^{49, 50}.

232 **An overview of the variation and complexity of DOM composition.**

233 General characteristics of DOM revealed unique molecular composition harbored
234 by different carbon-augmentation and fertilizer application practices (Figure 1, Table
235 1). A total of 6,428 molecular formulae were putatively assigned (average of all samples
236 3,607). Only 23.7% (1,521 molecules) of all molecular formulae were shared in all
237 tested soil samples (Figure S8). Moiety molecular mass covered a range from
238 approximately 132 to 599 Daltons. The DOM composition was comprised of
239 heteroatomic compounds, such as CHO, CHON and CHOS (delineated by elemental
240 formula combinations). CHON were most abundant (50.3% ~ 58.3% of all molecules)

241 followed by CHO (37.0% ~ 44.0%) and CHOS (3.3% ~ 8.7%) (Table 1).

242 The application of inorganic fertilizer (1N) increased the content of CHON by
243 6.0%, while the abundance of CHOS maintained at 3.3% with respect to the control
244 (3.3%) (Table 1). It is likely that more N was available to microorganisms due to the N
245 fertilization and more N containing organics were produced through microbial
246 metabolism ⁵¹. Carbon amendment (SS and CM) increased the abundance of CHON
247 and CHOS with the highest abundance of CHON in CM treatments and the highest
248 abundance of CHOS in 4SS treatments (Table 1). The application of sewage sludge
249 increased CHOS abundance by a factor of two to three (Table 1). The increase in S-
250 bearing moieties is most likely related to the delivery of these molecules (and their
251 metabolism post application ⁵² in SS and CM treatments. As sludge(SS) is
252 acknowledged to be rich in sulfur (0.7 – 2.1%) ⁵³.

253 Different heteroatomic classes (i.e. O_x, N_xO_y, O_xS_y) in DOM were existed in all
254 treatments. Compared to the inorganic fertilizer treatments, more types of heteroatomic
255 classes were observed in the treatments with high dose carbon-augmentation (Table 1).
256 Comparisons of DOM features in all treatments were based on H/C and O/C ratios,
257 aromaticity index (AI), double bond equivalence (DBE) and H/C cutoffs. The relative
258 abundances of DOM components were significantly different among treatments (Figure
259 1) (ANOSIM: R = 0.4872, P = 0.001). The relative abundance of each DOM component
260 was significantly different among treatments as well (ANOVA, P < 0.05) (Table S6).
261 Lignin-like DOM compounds were dominant in all soil samples, accounting for 54~63%
262 of all assigned molecules (Figure 1). The proportions of recalcitrant components (H/C
263 < 1.5), such as lignin, condensed aromatics and tannins were higher than the proportions
264 of labile components (H/C ≥ 1.5) in SS and CM treatments. The molecular composition
265 in CM treatment covered much lower H/C and wider O/C ratio (ANOVA, p < 0.05)

266 (Table 1), indicating greater recalcitrance ²⁵.

267 The evidence that carbon amendment practices (i.e. SS and CM) increased the
268 stocks of recalcitrant organic carbon are significant, and of global importance. If
269 rejuvenated soil carbon is labile then gains in soil carbon stocks will be transient (as
270 augmented carbon is mineralized back into CO₂). However, as our results indicate, SS
271 and CM amendment (over a decadal period) resulted in an increase in recalcitrant
272 carbon moieties. This evidence is salient to carbon sequestration and climate change
273 mitigation as it highlights the potential for SS and CM carbon amendments to increase
274 both the size of the soil carbon reservoir and the recalcitrance of this carbon. In relation
275 to the 4‰ Initiative ⁵⁴ our results affirm that soil carbon augmentation, with SS and
276 CM, could make a pragmatic contribution to the 4‰ aspiration.

277 **Characterization of DOM chemodiversity.**

278 The application of inorganic fertilizer and organic fertilizers changed the
279 chemodiversity of DOM molecular composition in soil, with significant differences
280 among treatments being reflected at a molecule level (Figure S3). The Chao 1 diversity
281 significantly increased following the application of SS or CM (ANOVA, $P < 0.0001$, F
282 value = 9.076). High dose application of organic fertilizer significantly increased DOM
283 chemodiversity. CM application increased the Chao 1 diversity of DOM, although the
284 increased level was less than those of high dose sewage sludge treatments. Non-metric
285 multidimensional scaling analysis (NMDs), showed the DOM compositions in organic
286 augmentation treatments to be markedly different from those of control and inorganic
287 fertilizer treatments (stress = 0.05) (Figure 2). Additionally, Spearman rank order
288 correlations revealed significant multicollinearity among Chao 1 index, soil pH, TC,
289 TN, DOC, DTN, C/N ratio and NO₃⁻ (Table S4). When soil chemical variables were
290 fitted to the RDA plot (Figure S4a), DOM molecular composition was found to be

291 significantly correlated with DOC, DTN and NO_3^- ($P = 0.001$). The variations of soil
292 DOM chemodiversity under different fertilization significantly correlated with soil
293 chemical factors, indicating that soil chemical factors that have distinct influence on
294 soil BCC can also affect soil DOM molecular diversity.

295 Using the LEfSe analysis, a total of 651 DOM moieties (referred as chemo-
296 markers) emerged to explain the greatest difference among treatments. By plotting
297 these chemical markers, in a van Krevelen plot according to each moiety's O/C ratio
298 versus H/C ratio ²⁵, observation of major groupings was simplified (Figure 3). Thus, a
299 pronounced boundary, differentiating inorganic fertilizer treatments from organic
300 augmentation treatments, was revealed. DOM molecules in the control were dominated
301 by compounds belonging to the aliphatic category ($\text{H/C} > 1.5$) ²⁵. Similarly, chemo-
302 marker molecules in inorganic fertilizer treatments were noted to be indicative of labile
303 organic compounds ($\text{H/C} > 1.5$) i.e. proteins/amino sugars and carbohydrates ²⁵. This
304 result highlights long-term application of inorganic fertilizer in reducing the abundance
305 of recalcitrant soil carbon moieties. Mulvaney ⁸ reported that the amendment of
306 inorganic fertilizer increased mineralization of native soil carbon and nitrogen.

307 In contrast, chemo-markers in SS and CM treatments were distinct from the
308 chemo-markers in CK and N treatments (Figure 3), these moieties belonged primarily
309 to recalcitrant compounds ($\text{H/C} < 1.5$). These, more recalcitrant compounds noted in
310 the SS and CM treatments, belonged mainly to the category of humic-like compounds
311 i.e. condensed aromatics, phenolic and highly unsaturated compounds and polyphenols
312 ⁵⁵. These results are consistent with previous reports that have indicated sludge
313 amendments to increase humic matter contents in soils ¹⁴. Our results add new insight,
314 in terms of the molecular fingerprint of the chemodiversity of DOM in soils augmented
315 with organic carbon over a decadal period. These results are significant as they

316 highlight the benefits organic carbon augmentation can realize in terms of both building
317 soil carbon stocks and increasing the recalcitrance of these carbon stocks. As
318 highlighted above, these findings support the use of organic carbon amendments with
319 potential to make long-term contributions to achieving 4% aspirations⁵⁴.

320 **Characterization of bacterial community composition (BCC).**

321 After assembling and quality filtering, 28,169 - 172,449 sequences were identified
322 per sample (average of 80,361). These sequences were assigned into 7,872 OTUs at a
323 97% identity level. The most dominant phyla across all samples was the copiotrophic
324 taxa, *Proteobacteria* (27.4% - 33.0%, Figure 4a). Other prevalent phyla across all
325 treatments were, *Actinobacteria*, *Acidobacteria* and *Chloroflexi* (>10%) (Figure 4a).
326 The similar trends in BCC have been reported in other long-term field experiment on
327 fluvo-aquic soils^{11,47}. Various types of inorganic fertilizers and manure were fertilized
328 in fluvo-aquic soil for 24 years. Soil BCC were all dominated by the *Proteobacteria*,
329 *Acidobacteria*, and *Actinobacteria*⁴⁷. Soils amended with organic carbon and inorganic
330 fertilizer had markedly, and significantly, different bacterial communities compared to
331 each other and the control soil (ANOSIM, $R = 0.7666$, $P = 0.001$) (Figure 4). The
332 relative abundance of *Acidobacteria* was lower in 4SS (14.2%) and CM (14.5%)
333 compared to other treatments. The amendment of organic materials (CM & 4SS)
334 increased the relative abundance of *Actinobacteria*, while decreased the relative
335 abundance of *Chloroflexi*, especially at high doses of sludge via significant test (Figure
336 4a). These observations are consistent with previous studies in which organic matter
337 amendment stimulated copiotrophic taxa (i.e. *Proteobacteria* and *Actinobacteria*)
338 growth^{11,49,56}. *Proteobacteria* and *Actinobacteria* taxa have previously been reported
339 to be dominant in soil under long-term organic carbon augmentation^{47,57,58}. In contrast,
340 BCC was directed towards an increase in the oligotrophic taxa, *Acidobacteria* that was

341 present at significantly high frequencies in control, or N treatments ^{11,49}. *Proteobacteria*
342 have been putatively recognized as copiotrophic taxa (taxa that thrives in conditions of
343 elevated C and N availability and exhibits relatively rapid growth rates) ^{34, 56, 58} and
344 favors nutrient-rich conditions and associated with carbon rich regimes. In contrast,
345 *Acidobacteria* is considered to be an oligotrophic taxon that exhibits relatively slow
346 growth rate and ability to metabolize nutrient-poor substrates ⁵⁶. The relative increase
347 (25.5%) in *Acidobacteria* increased reported here is consistent with the BCC shifts
348 under twenty-three years of nitrogen-containing inorganic fertilization applications ¹¹.

349 Further comparison of the BCC at the class level revealed, *Actinobacteria* as the
350 dominant group (17.4%), closely followed by *Acidobacteria* (16.1%) and
351 *Alphaproteobacteria* (11.8%) (Figure 4b). Another highly abundant class was
352 *Gammaproteobacteria* (6.05%). *Actinobacteria* was the most abundant class in samples
353 of 4SS, while *Acidobacteria* was dominant in 0.5SS, 2SS. In the control (CK), 0.5N,
354 1N, 1SS and CM *Alphaproteobacteria* was the most abundant class. The relative
355 abundance of *Actinobacteria* increased in SS and CM treatments with *Actinobacteria*
356 dominating in 4SS at the class level.

357 The overall pattern of BCC in the NMDS plot (Figure S5) (stress = 0.05) suggested
358 that BCC was altered by organic carbon amendments, and BCC in SS and CM group
359 were markedly different from BCC in 0.5N and 1N group. Alpha-diversity of bacteria
360 was increased in SS and CM treatments (especially at high SS doses) (Table S3). A
361 redundancy analysis elucidated the relationships between the BCC and soil factors in
362 the different fertilization treatments. The RDA ordination plot (Figure S4b) indicated
363 that soil DTN, DOC and NO_3^- were the most important variables that influencing BCC.

364 Previous studies have indicated that organic carbon ⁴⁷ and inorganic fertilizer ⁵⁸
365 application have profound shaping influence on soil microbial community structure,

366 with implications for the cycling of carbon ⁵⁹, the regulation of soil ecosystem services,
367 such as nutrient flows ⁵⁸ and greenhouse gas emission ^{60,61}. The observed shifts in BCC
368 are suggested to be in response to decreasing soil fertility associated with long-term
369 chemical fertilizer application that reduced the nutrient availability and increases
370 nutrient loss. Correspondingly, the low-fertility soils supported oligotrophic
371 ecosystems, whereas high-fertility soils support eutrophic ecosystems ^{11,34}. In contrast,
372 long-term organic carbon-augmentation improved DOM quality, and promoted the
373 growth of copiotrophic taxa. Thus, our results suggested that organic carbon-
374 augmentation supported a bacterial community shift to one indicative of a eutrophic
375 ecosystem.

376 **Linkages between soil bacterial communities, DOM molecular composition and**
377 **soil properties.**

378 The interconnections between chemodiversity of DOM and BCC were explored
379 using co-occurrence network analysis. Results revealed strong and significant
380 associations between DOM molecules and specific taxa (Figure 5). The network pattern
381 indicated taxa of the same phyla or same class had diverse associations with DOM of
382 contrasting chemical characteristics, i.e. having opposite correlations to the same
383 category of DOM compounds or having correlations to the molecules belonging to
384 distinct regions of chemical composition. These results reveal evidence that *Nitrospira*
385 specialized on typical DOM molecules which might be utilized for defining the possible
386 ecological niche.

387 There were 52,319 pairs of correlations, and 7,944 strong correlations ($|R| > 0.9$)
388 between the DOM molecule dataset and the OTUs dataset (Figure S6). The top 100
389 OTUs with the highest relative abundance and the top 100 most abundant DOM
390 molecules were considered for network analysis (Figure 5, Table S5). A total number

391 of 1,105 pairs of correlations were established among these top DOM molecules, OTUs
392 and environmental factors. At $R \geq 0.6$, 135 correlations persisted, of which 40 linked
393 26 DOM molecules and 17 OTUs. All 17 OTUs belonged to the taxa *Acidobacteria* (7),
394 *Proteobacteria* (3), *Actinobacteria* (3), *Nitrospirae* (2), *Chloroflexi* (1) and
395 *Planctomycetes* (1), respectively. Sixteen DOM molecules belonged to recalcitrant
396 compounds ($H/C < 1.5$). *Proteobacteria*, in particular, showed strong positive
397 correlation (red lines) with recalcitrant compounds ($H/C < 1.5$) and negative correlation
398 (black lines) with aliphatics ($H/C > 1.5$) (Figure 5), these chemical traits being
399 consistent with high-dose SS and CM treatments (Figure 3). *Acidobacteria* showed
400 negative correlations with recalcitrant compounds in all treatments (Figure 3). These
401 observations support the hypothesis that infertile soil with low nutrient availability
402 (consistent with CK and 0.5N treatments) generally selected for *Acidobacteria*^{12,34}.

403 A group of ten labile DOM molecules belonged to aliphatic ($1.5 < H/C \leq 2.0$) and
404 carbohydrate ($0.6 < O/C < 1.2$ and $1.5 < H/C < 2.2$). A group of distinct DOM molecules
405 showed strong correlations with more than one OTU (these belonging to seven different
406 phyla) whereas the remaining DOM molecules were correlated with either,
407 *Acidobacteria*, *Nitrospira*, or *Proteobacteria*. Strongly restricted correlations were
408 observed between *Proteobacteria* and aliphatic-like compound, whereas *Acidobacteria*
409 and *Nitrospira* had strong correlations that were almost exclusively with highly
410 unsaturated hydrocarbons, phenolic compounds and lignins. In addition, *Nitrospira*
411 showed strong negative correlations to the recalcitrant compounds, and
412 *Desulfurellaceae* showed strong negative correlations to the CHOS class (especially
413 $C_9H_{18}O_6S_1$, Figure 5). This may suggest that *Nitrospira* and *Desulfurellaceae*
414 specialized on specific DOM categories that are not intensively utilized by other taxa.
415 *Desulfurellaceae* is common sulfate-reducing bacteria in sludge and is capable of

416 consuming CHOS compounds (these noted to be abundant in SS and CM treatments
417 (Table 1)). The co-occurrence in network supports the findings of specialization on
418 specific substrate⁶². The separate network plots showed significant differences between
419 the inorganic fertilizer treatments and organic carbon treatments (Figure 6). In the CK
420 and 0.5N treatment, the network had 350 and 278 edges, and 4SS and CM had 537 and
421 401 edges, respectively. In addition, the more active nodes were detected in the organic
422 carbon-amendments treatments than in the inorganic fertilizer treatments. The node
423 having more than 7 edges was defined as the network hub, which was active in
424 mediating interactions⁵¹. The organic carbon-amendments (especially the high dose
425 treatments) maintain a more complex network structure.

426 Previous research regarding linkages between DOM and BCC has only focused
427 on the effects of the content of soil carbon¹², the appreciation of their relationships with
428 each other is strikingly limited on the broad molecular level due to the complexity in
429 composition of both²⁴. Our results are significant as they reveal, in unprecedented detail,
430 associations between DOM chemodiversity and BCC diversity. Soil bacteria-DOM
431 interaction was demonstrated in strong correlations between specific bacterial taxa and
432 particular DOM molecules, thus, suggesting bacterial specialization on particular
433 substrates. The number of active hubs in the carbon-amendments treatment was more
434 than in the inorganic fertilizer treatments, indicating that a greater diversity of soil
435 bacteria interacted with a greater diversity of DOM molecules in treatments subjected
436 to protracted organic carbon amendment. To the best of our knowledge, this is the first
437 report the co-variation of soil DOM composition and BCC.

438 Taking advantage of technological advances in analytical chemistry, molecular
439 biology and informatics, we explored how the BCC and DOM chemodiversity were
440 altered (in long term, decadal, field experiments) in response to carbon-amendment and

441 application of inorganic fertilizers. This research highlights the manifold associations
442 between the diversity of microbiota and the heterogeneity of soil DOM under long-term
443 organic carbon amendment and inorganic fertilization practices. Our results bring new
444 insight to the negative impacts of protracted inorganic fertilizer application on the DOM
445 resource in soil and the BCC it supports. Our results indicate that protracted organic
446 carbon amendments not only increased soil carbon stocks but also their recalcitrance.
447 In addition, SS and CM amendments shaped BCC to an indicative state of improved
448 soil health and one that has the potential to improve delivery of soil ecosystem service
449 delivery within agroecosystems. These two lines of evidence affirm that soil carbon
450 augmentation, with SS and CM, could make a pragmatic contribution to the 4‰
451 aspiration to (re)build resilient soil carbon stocks while improving soil health and the
452 delivery of beneficial soil ecosystem services. Our results might contribute to the
453 defining of a mechanism to translate current scientific knowledge, regarding soil carbon
454 status, into actionable pathways that might inform new agricultural or land use policy
455 to bring to fruition the 4‰ vision.

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464 **Notes**

465 The authors declare no conflict of interest.

466 **Supporting Information**

467 Details on supportive methods and discussion, additional details of the field layout and
468 experiment (Figure S1, Table S1 and Table S2); soil chemical variables (Figure S2),
469 diversity index of soil DOM and bacterial community (Figure S3 and Table S3),
470 Redundancy analysis (RDA) of DOM molecular composition and bacterial community
471 composition (Figure S4), NMDS analysis of bacterial community composition (Figure
472 S5), Co-occurrence network visualizing the DOM-Bacteria interactions (Figure S6).

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686

Table 1 FT-ICR MS characteristics of DOM composition.

Treatment	CHO (%)	CHON (%)	CHOS (%)	Number of Heteroatomic class	C	H	O	MW	DBE	H/C ratio	O/C ratio
CK	44.0a	52.0a	3.33a	49a	15.95ab	17.61d	7.00a	334.66a	8.09a	1.10c	0.44a
0.5N	43.3ab	53.0a	3.33a	52b	15.92ab	16.49bc	7.12a	335.40a	8.63b	1.04b	0.45ab
1N	39.0b	58.0b	3.33a	55c	16.12b	16.33b	7.54b	345.77bc	8.96d	1.01b	0.47c
0.5SS	43.3b	50.3a	6.67b	51b	16.32c	17.73d	7.18a	342.76b	8.39b	1.09c	0.44a
1SS	38.7b	53.7c	7.67c	56c	16.27cd	16.88bc	7.51b	348.41cd	8.82cd	1.04b	0.46c
2SS	37.7b	55.3c	7.00c	56c	16.33cd	16.73bc	7.62b	351.03d	8.97d	1.02ab	0.47c
4SS	37.0b	54.3c	8.67c	56cd	16.46d	17.09d	7.57b	352.60d	8.91cd	1.04b	0.46bc
CM	36.7ab	58.3b	5.33b	57d	15.87a	15.71a	7.66b	345.34cd	9.04d	0.99a	0.48d

687 MW - molecular weight, DBE - double bond equivalent. Significance level: *P < 0.05.

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689

690 **Figure legends**

691 **Figure 1** van Krevelen diagram-derived relative abundance (%) of classification classes
692 of the DOM components (from FT-ICR MS analysis) in: control soil (CK), inorganic
693 fertilizer treatments (0.5N and 1N), sewage sludge treatments (0.5SS, 1SS, 2SS and
694 4SS) and chicken manure treatments (CM).

695 **Figure 2** Distribution patterns of DOM molecules composition. Non-metric
696 Multidimensional scaling (NMDS) analysis of DOM molecules based on Bray-Curtis
697 distance.

698 **Figure 3** Linear discriminant effect size analysis (LEfSe) of DOM chemo-marker
699 molecules that are enriched in different treatments. Number of chemo-marker
700 molecules: 651 in total; 54 in sludge; 293 in manure; 47 in N-chemical fertilizer, and;
701 257 in the control. Molecule compounds with no significant differences are not shown.

702 **Figure 4** (a) Relative abundance of bacteria community composition components at the
703 phylum level. (b) Relative abundances of bacteria community composition at class level.
704 “Others” include low abundance (< 1%) bacteria and the taxonomically unassigned
705 sequences at class level.

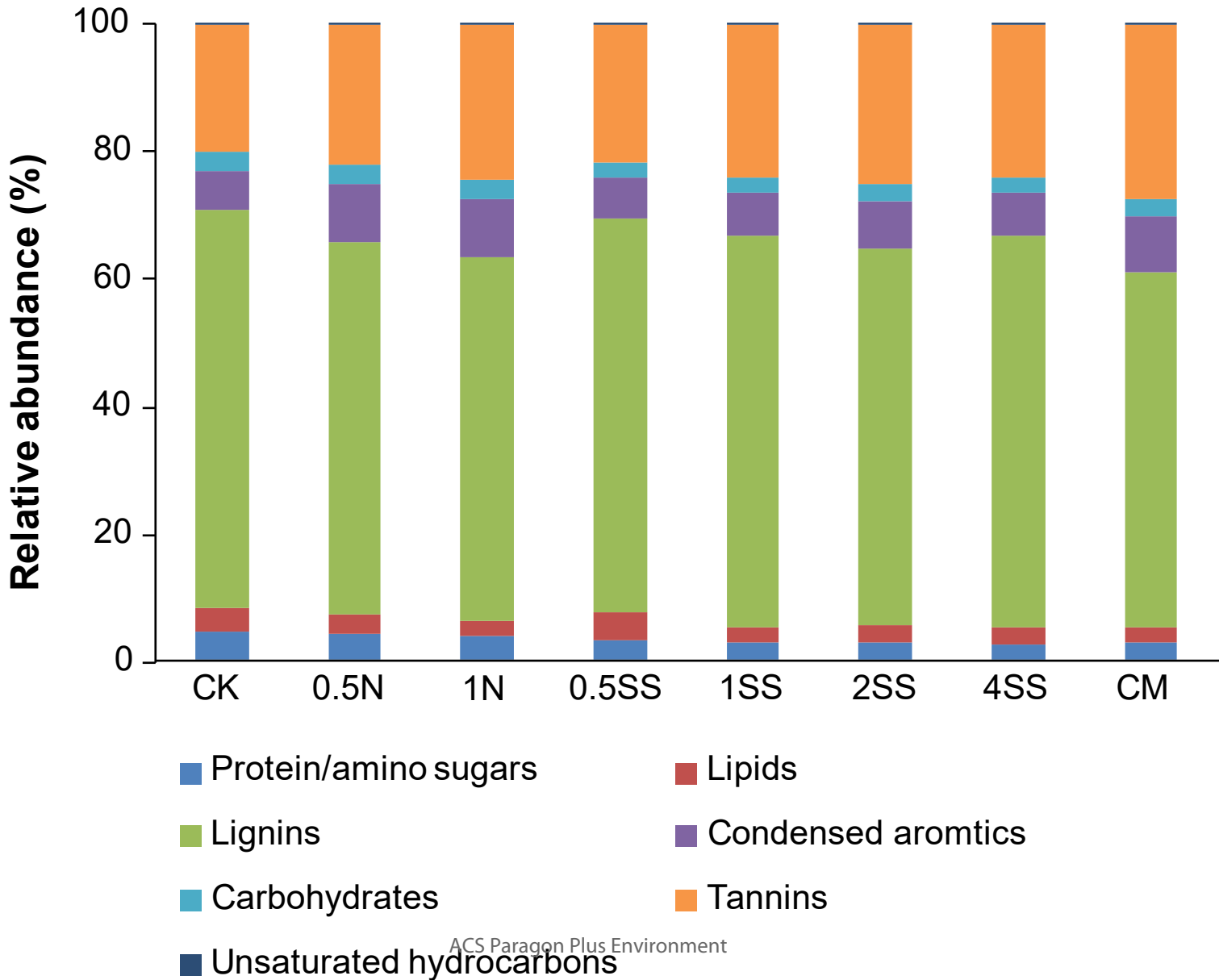
706 **Figure 5** Interaction network analysis of top 100 most abundant bacterial OTUs and
707 top 100 most abundant DOM molecules that were significantly correlated ($P < 0.05$, $|R|$
708 > 0.6). Circles, DOM molecules; Triangles, Bacterial OTUs (green); DOM molecules
709 relative abundances are set proportional to node size. Nodes are colored according to
710 DOM category, e.g. aliphatic compounds (light blue) and recalcitrant compounds (red).
711 Positive correlations are indicated using red lines, negative correlations are indicated
712 using black lines.

713 **Figure 6** Statistically significant and strong co-occurrence relationships between DOM
714 molecules and bacterial OTUs within different treatments. Network plots for (a) CK,
715 (b) 0.5N, (c) 4SS and (d) CM treatments. Nodes represent DOM molecules and OTUs
716 with in significant relationships. The color of each node indicates the OTUs from
717 different phylum and labile (light blue) or recalcitrant (dark blue) DOM molecules (see
718 key). The size of the nodes (circles) is proportioned to the number of the connections
719 (degree).

720

721

Figure 1



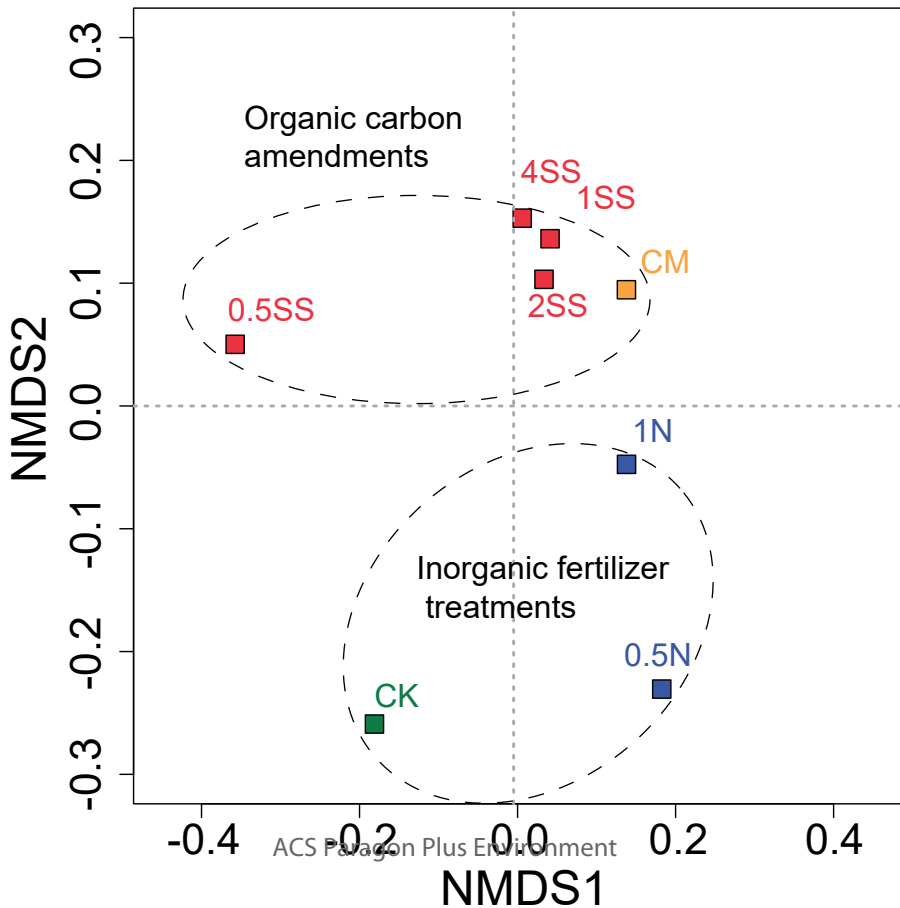


Figure 3

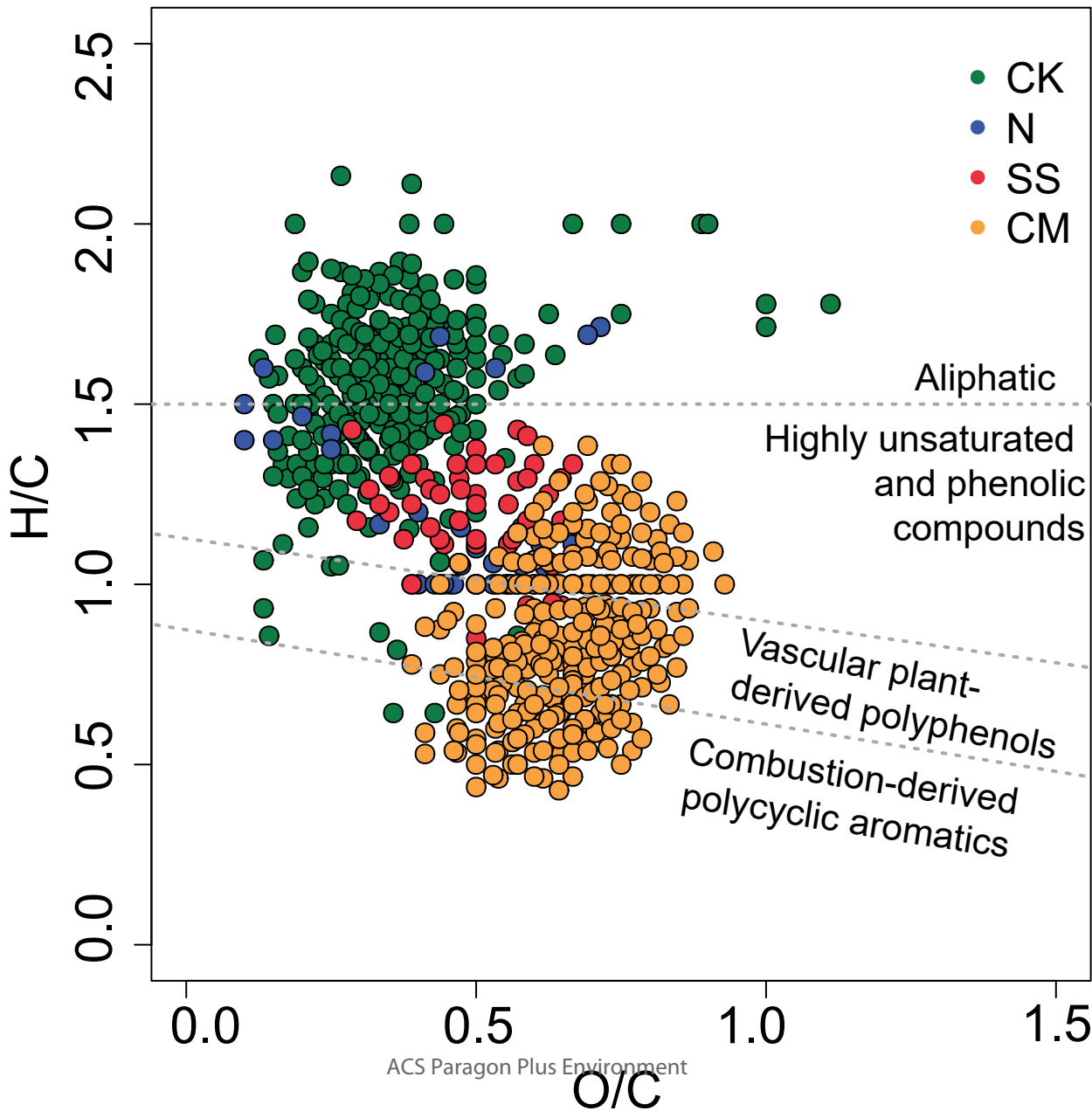
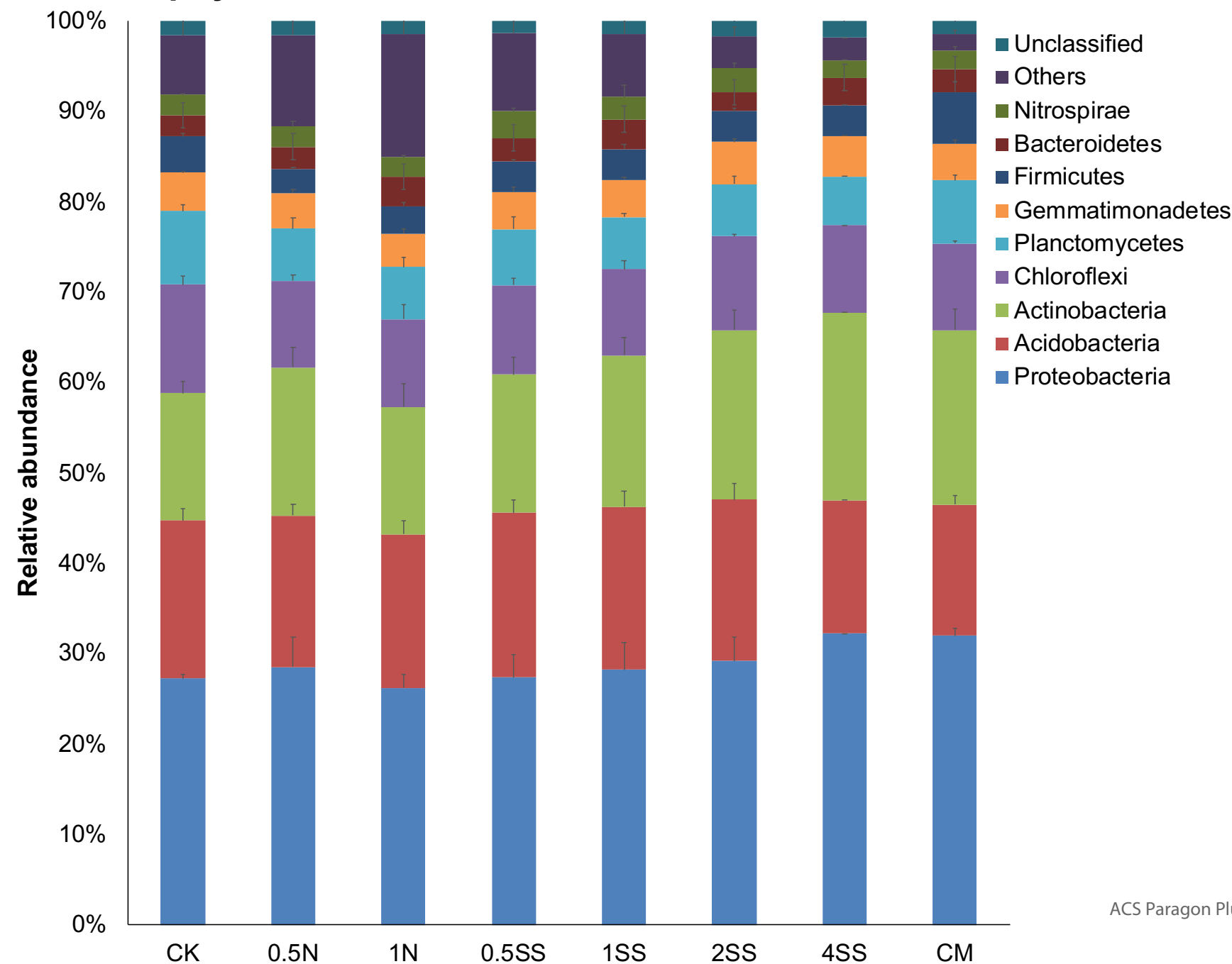
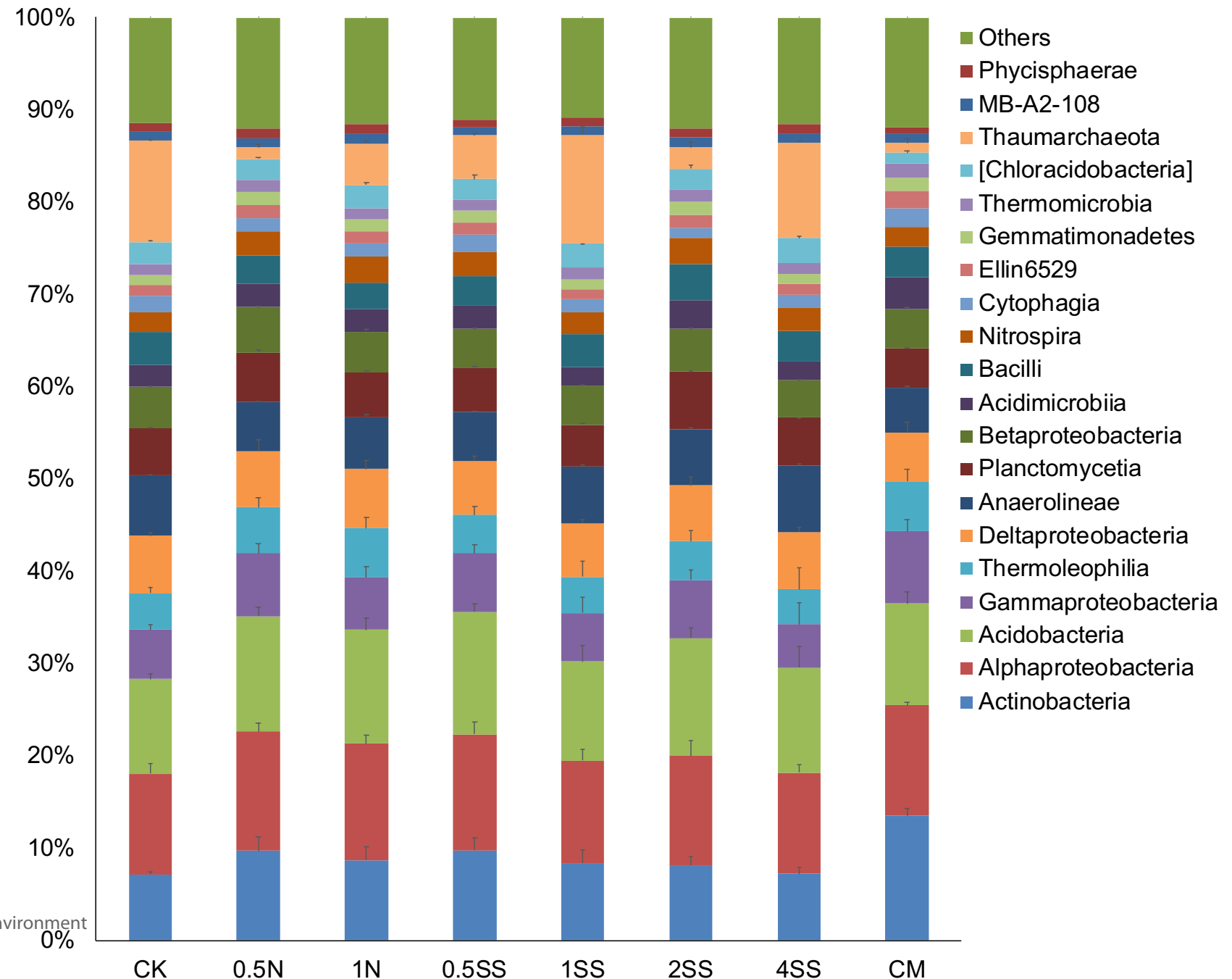


Figure 4

a BCC/phylum



b BCC/class



ACS Paragon Plus Environment

Figure 5

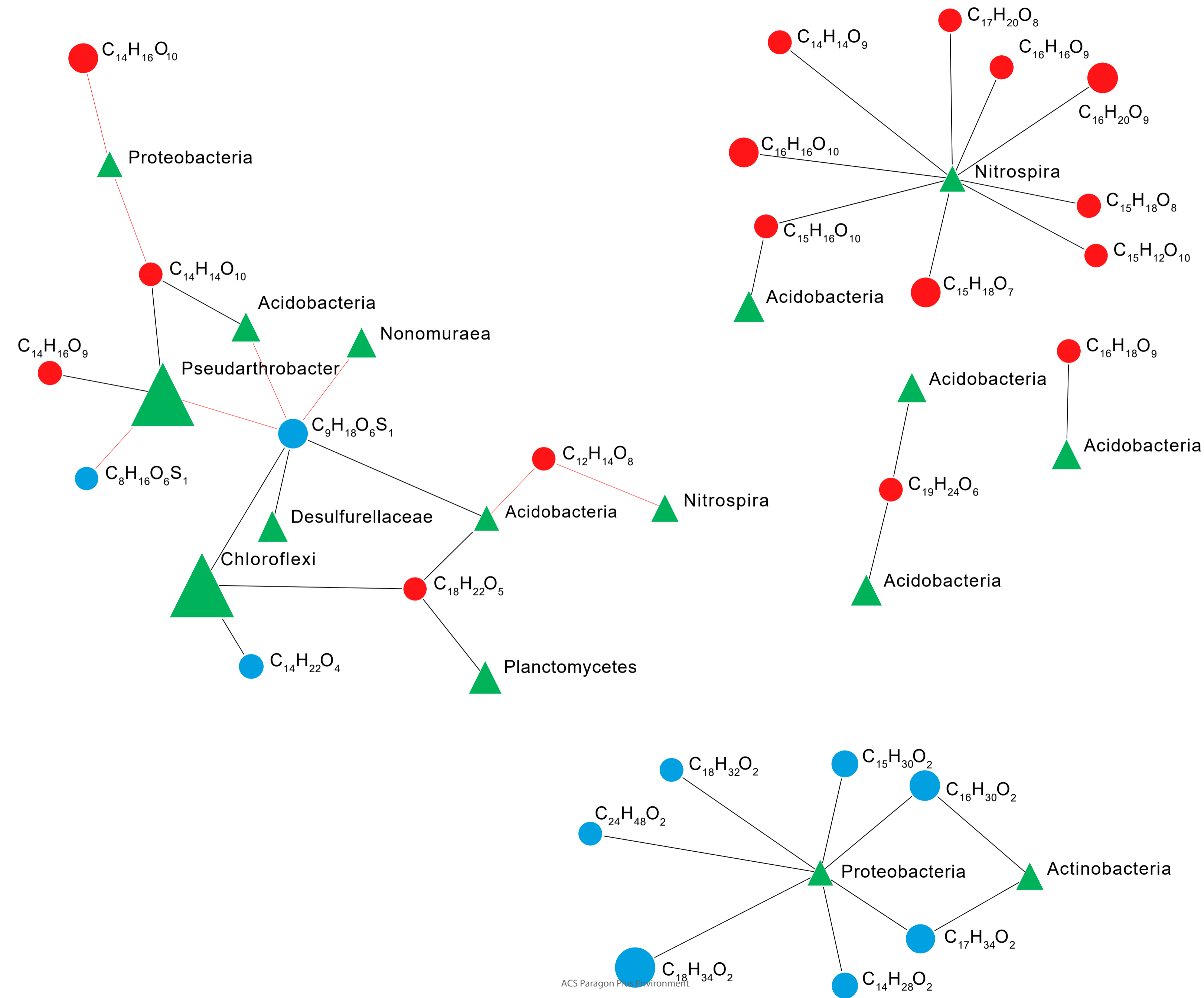
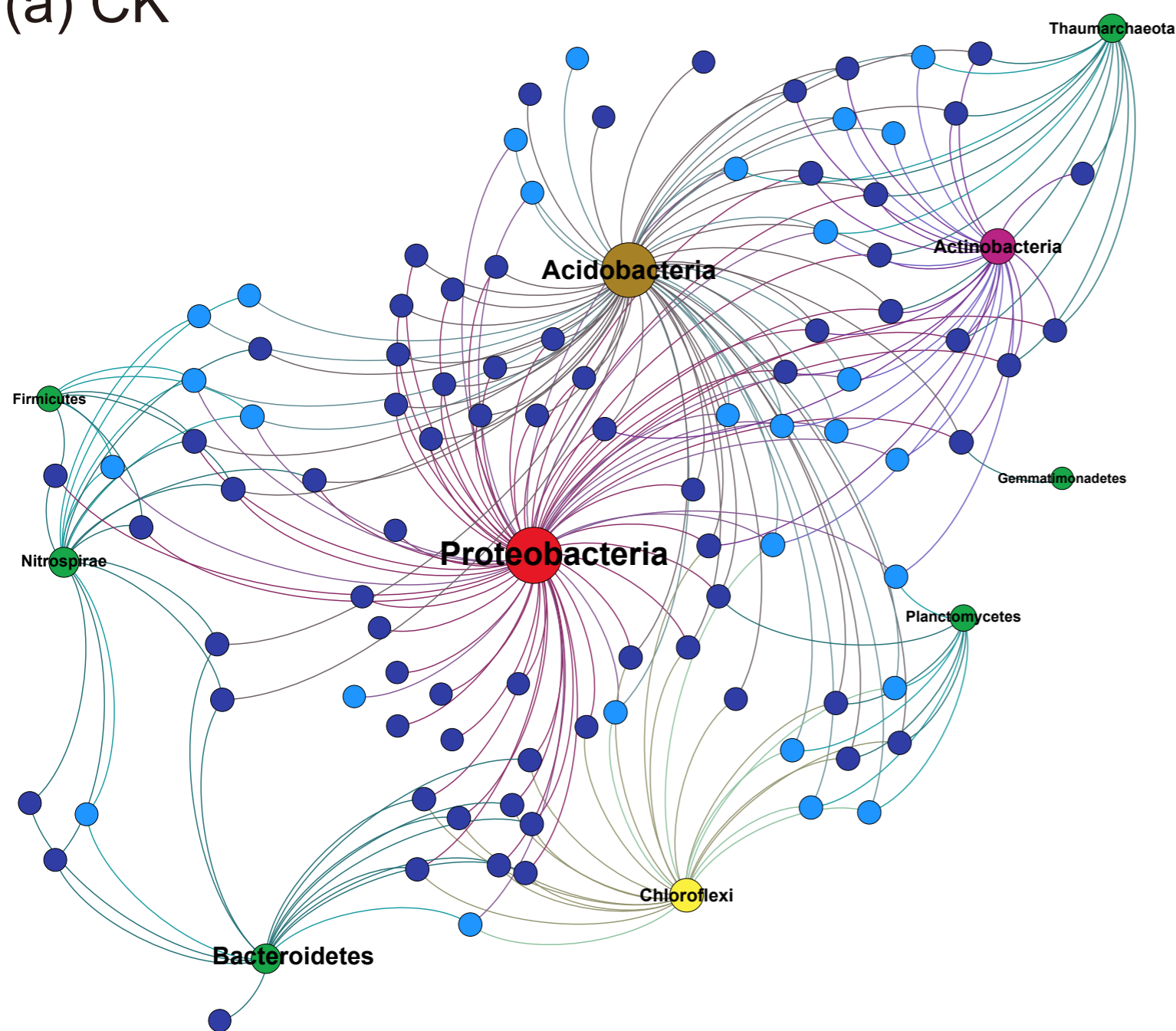


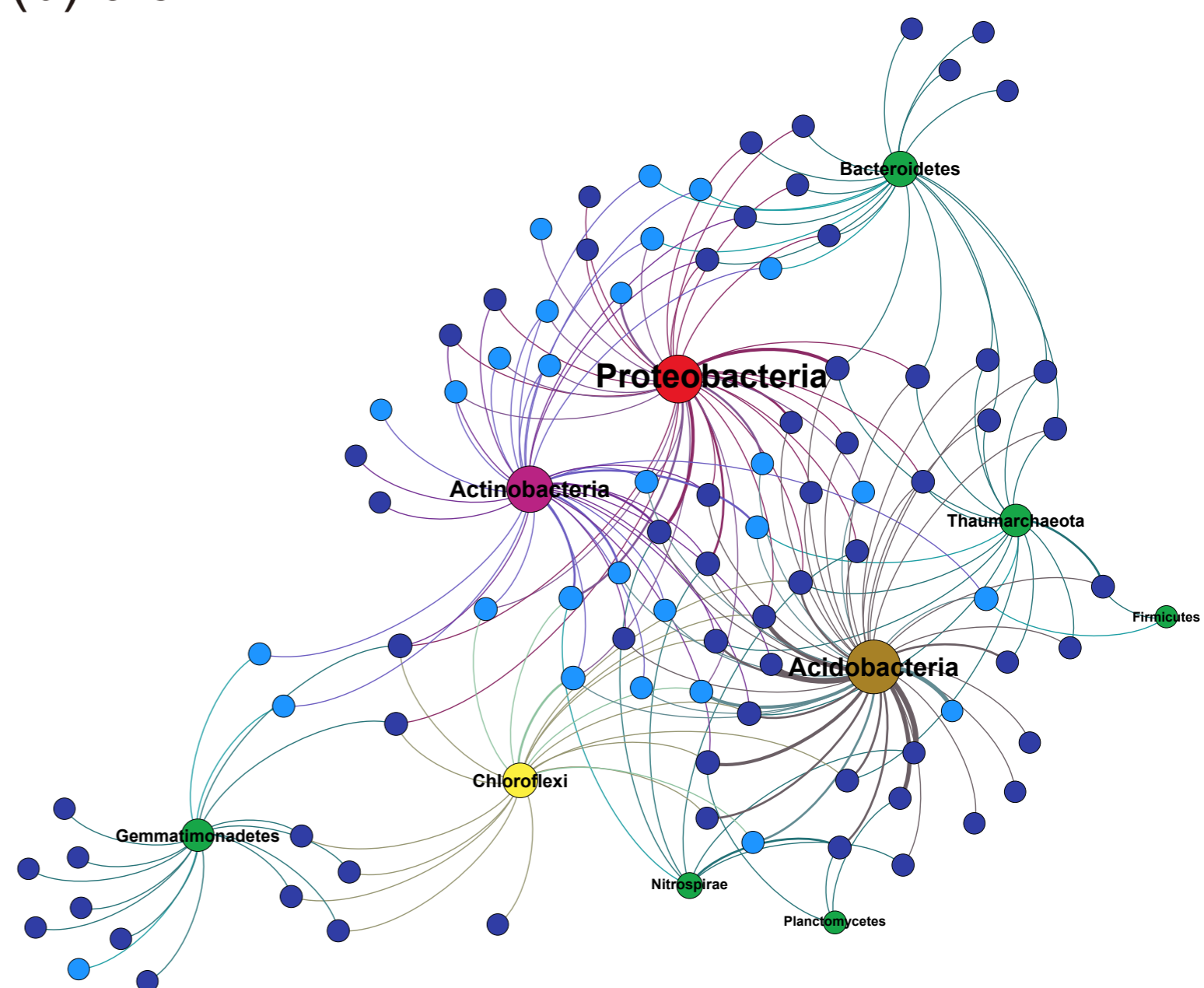
Figure 6



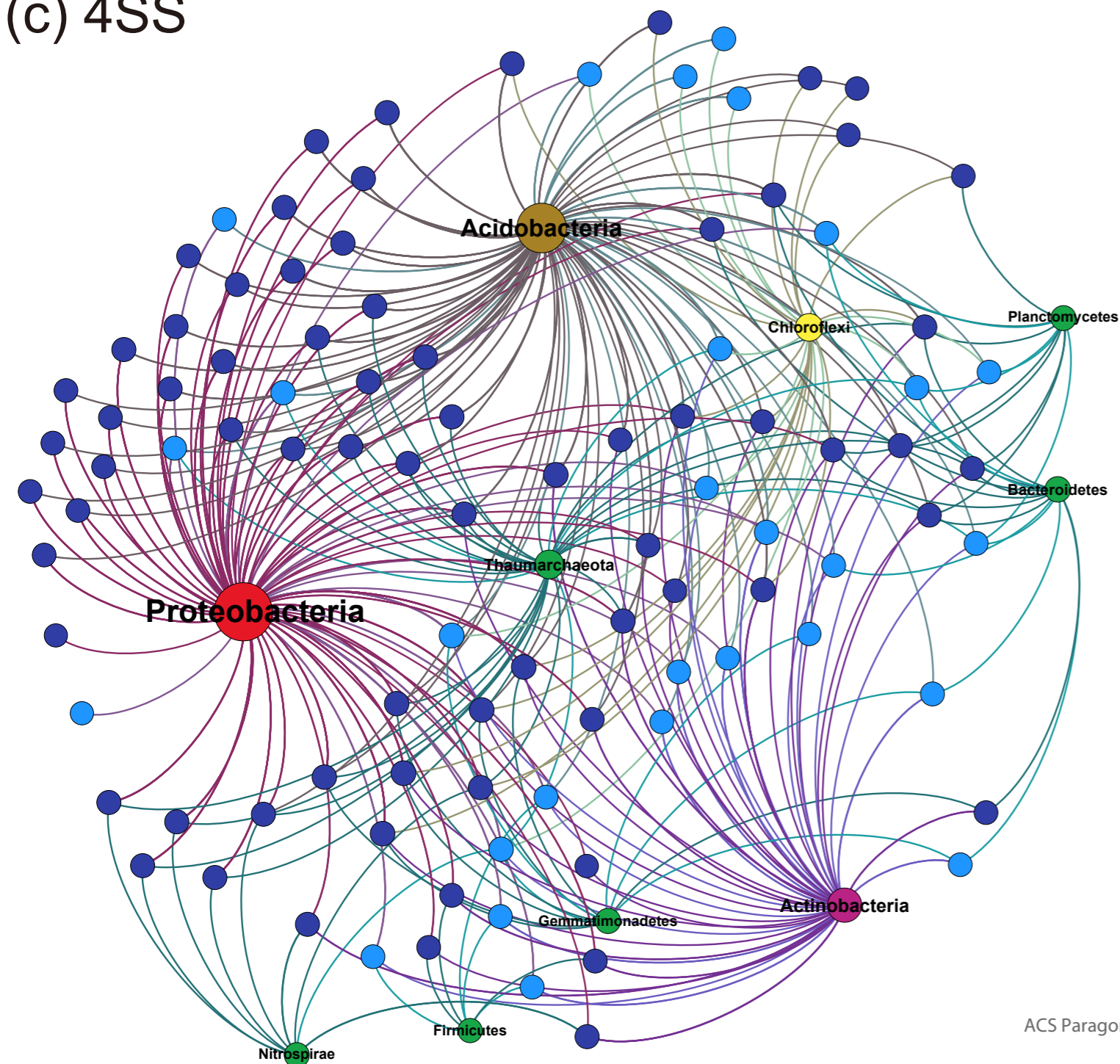
(a) CK



(b) 0.5N



(c) 4SS



(d) CM

