1	Disinfection exhibits systematic impacts on the drinking water microbiome.
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18 ABSTRACT

19 Limiting microbial growth during drinking water distribution is achieved either by maintaining a 20 disinfectant residual or through nutrient limitation without the use of a disinfectant. The impact of these contrasting approaches on the drinking water microbiome is not systematically understood. 21 22 We utilized genome-resolved metagenomics to compare the structure, metabolic traits, and 23 population genomes of drinking water microbiomes across multiple full-scale drinking water 24 systems utilizing these two-distinct microbial growth control strategies. Microbial communities 25 cluster together at the structural- and functional potential-level based on the presence or absence 26 of a disinfectant residual. Disinfectant residual concentrations alone explained 17 and 6.5% of the variance in structure and functional potential of the drinking water microbiome, respectively. 27 despite including samples from multiple drinking water systems with variable source waters and 28 29 source water communities, treatment strategies, and chemical compositions. The drinking water 30 microbiome is structurally and functionally less diverse and less variable across disinfected 31 systems as compared to non-disinfected systems. While bacteria were the most abundant domain, archaea and eukaryota were more abundant in non-disinfected and disinfected systems, 32 33 respectively. Community-level differences in functional potential were driven by enrichment of 34 genes associated with carbon and nitrogen fixation in non-disinfected systems and γ -aminobutyrate 35 metabolism in disinfected systems which may be associated with the recycling of amino acids. Metagenome-assembled genome-level analyses for a subset of phylogenetically related 36 37 microorganisms suggests that disinfection may select for microorganisms capable of using fatty 38 acids, presumably from microbial decay products, via the glyoxylate cycle. Overall, we find that disinfection exhibits systematic and consistent selective pressures on the drinking water 39 microbiome and may select for microorganisms able to utilize microbial decay products 40 41 originating from disinfection inactivated microorganisms.

42

43 INTRODUCTION

44 Drinking water systems harbor diverse and complex microbial communities in bulk water, biofilms on pipe wall, suspended solids, and in loose deposits¹⁻⁵. While treatment processes at the drinking 45 water treatment plants (DWTPs) shape the microbial community that leaves the DWTP⁶⁻⁹, multiple 46 factors can influence the structure and function of the drinking water microbiome in the drinking 47 48 water distribution systems (DWDSs). These factors include, but are not limited to, DWDS size, 49 pipe materials and ages, water age within the DWDS and similar factors within premises plumbing (PP) in buildings and homes¹⁰⁻¹⁴. Managing the microbiological quality of drinking water during 50 51 transport through the DWDS and into the PP is essential for the provision of safe drinking water. 52 Unwanted microbial growth and/or changes in the drinking water microbiome composition during transit through the DWDS and PP are associated with several detrimental outcomes. For instance, 53 this could lead to proliferation of opportunistic pathogens¹⁵⁻¹⁹ and an eukarvotic microbes^{14, 16, 20,} 54

 21 , taste and odor issues²², and impact infrastructure via corrosion damage^{23, 24}.

56 Source-to-tap differences in drinking water systems can range from source water type (e.g., 57 surface, ground, reuse water), process configurations at the DWTP, heterogeneity and condition of the DWDS and PP; yet globally there are two fundamental approaches for managing the 58 drinking water microbiome during transport to the consumer²⁵. The first and most widely used 59 approach involves maintenance of a disinfectant residual (e.g., chlorine) in the DWDS. This is 60 accomplished by ensuring the water leaving the DWTP has a chlorine residual and/or by using 61 booster stations in large complex DWDSs to compensate for disinfectant residual decav²⁶. 62 63 Disinfectant residuals counteract microbial growth through inactivation, thus ensuring stable 64 microbial concentrations during distribution. While disinfectant residuals are effective in managing microbial growth in the DWDS, there are some key issues associated with them. These 65 include aesthetic and corrosion related problems^{25, 27, 28}, but more importantly the formation of 66 harmful disinfection byproducts (DBPs)²⁹⁻³¹, which are also regulated. Further, there is an 67 increasing recognition that the disinfectant residuals may be associated with selection of some 68 opportunistic pathogens^{16, 32} and antibiotic resistance genes (ARGs) in drinking water³³⁻³⁵. 69

The second approach for managing microbial growth in the DWDS, primarily practiced in parts
of western Europe (e.g., Netherlands, Denmark, and Switzerland), involves distribution of drinking
water without any disinfectant residuals³⁶. These systems focus on minimizing nutrient availability

73 in the DWDS to limit microbial growth using high-quality source waters and/or multi-barrier 74 treatment. While some of these drinking water systems may also use chlorine or other chlorine 75 compounds (e.g., chlorine dioxide) at the DWTP, they ensure that chlorine is not detectable prior 76 to distribution. The efficacy of this approach is supported by evidence that incidences of microbial 77 contamination and associated waterborne illnesses are comparable to systems that maintain a disinfectant residual^{25, 37}. This suggest that with appropriate source water quality management, 78 79 treatment, and well maintained infrastructure, drinking water can be safely distributed without disinfectant residuals²⁵. 80

81 Despite reports of comparable biological water quality between systems with and without 82 disinfectant residuals, there are a limited number of studies that have systematically compared the microbial community between these two types of systems. Bautista et al (2016)³⁸ conducted a 83 84 meta-analyses study involving collation, curation, and comparison of 16S rRNA gene amplicon 85 sequencing data from previously published datasets. While this study was confounded my 86 methodological differences between datasets being used, the key conclusions were that 87 presence/absence of disinfectant residuals impact microbial community structure and membership 88 and that systems without disinfectant residuals are more diverse than their disinfected counterparts. Recently, Waak et al (2019)³⁹ compared biofilms between two drinking water systems, one 89 90 chloraminated systems and one without a disinfectant residual. Consistent with previous findings 91 they observed higher cell numbers and higher diversity in the system without disinfectant residual, 92 with higher proportional abundance (proportion of total community) of deleterious microbes (i.e., mycobacteria, nitrifiers, corrosion causing bacteria) in the chloraminated system. Both, Bautista 93 et al (2016)³⁸ and Waak et al (2019)³⁹ utilized gene-targeted assays (i.e., 16S rRNA gene) to probe 94 drinking water microbiome composition and its differences. While gene-targeted assays can 95 provide valuable information on microbial community structure and membership information, 96 they do not provide insight into metabolic differences that may drive the observed differences in 97 98 community structure. Further, gene-targeted assays can be limited by primer-bias and can result in 99 non-detection of microbial community members. Both challenges can be overcome by utilizing 100 metagenomics which can provide insights into structure and functional potential of a microbial 101 communities without being biases against or towards specific community members. This comes 102 with the limitation that differences between samples/systems emerging from low-abundance 103 microbes may not be detected as this may require ultra-deep sequencing.

104 We used metagenome analyses and genome-resolved metagenomics to investigate the potential 105 influence of disinfectant residuals on the drinking water microbiomes by comparing drinking water 106 systems from the United Kingdom (with disinfectant residual) and the Netherlands (without 107 disinfectant residual). The goals of this study were (1) to determine the extent to which disinfectant 108 residual shapes the structure and functional potential of the drinking water microbiome, (2) to 109 determine whether the selective pressures of disinfection are conserved across drinking water 110 systems, and (3) identify metabolic pathways underpinning differences in structure and functional potential of the drinking water microbiome. Addressing these questions across different drinking 111 112 water systems with inherent system-to-system variability (e.g., source water, water chemistry, 113 treatment process, etc.) but one consistent difference - i.e., presence or absence of disinfectant 114 residual - will help highlight disinfection that are conserved and thus generalizable across systems.

115 MATERIALS AND METHODS

116 Sample collection and processing.

Drinking water samples were collected from 12 drinking water systems in Netherlands (n=5) 117 118 between October to December 2013 (Non-disinfected, i.e. ND) and the United Kingdom (n=7) 119 between April to August in 2015 (Disinfected, i.e. D). Samples were collected at two to four 120 locations in each DWDS which resulted in 23 D and 18 ND samples. A total 15 liters of water was 121 filtered through three sterile Sterivex filters with 0.22μ m pore size polyethersulfone membrane (EMD MilliporeTM SVGP01050) using a peristaltic pump (Watson-Marlow 323S/D) to harvest 122 123 microbial cells. Immediately after filtration, the membranes were removed aseptically from the 124 Sterivex cartridge, cut into pieces and then transferred to Lysing Matrix E tubes. The membranes 125 were stored at 4°C for 24 hours or less before being transported to the laboratory and stored at -80°C. Further details of sample treatments and preservation are described in Sevillano-Rivera et 126 al.³⁵, along with detailed description of chemical analyses. Briefly, Orion 5 Star Meter (Thermo 127 128 Fisher Scientific, Waltham, MA) was used to measure temperature, pH, conductivity and dissolved 129 oxygen, total chlorine, and phosphate was also determine on-site using DR 2800 VIS 130 Spectrophotometer (Hach Lange, the UK) and EPA approved HACH kits. Nitrogen species were 131 measured according to standard method, 4500-NH3-F for ammonia, 4500-NO2-B for nitrite, and

4500-NO3-B for nitrate respectively in laboratory⁴⁰, while total organic carbon (TOC) was
 determined using Shimadzu TOC-LCPH Analyzer (Shimadzu, Kyoto, Japan).

134 DNA extractions.

The total genomic DNA was extracted directly from filter membranes using Maxwell16 DNA 135 136 extraction system (Promega) and LEV DNA kit (AS1290, Promega, Madison, WI, US). The filters 137 with collected biomass in lysing matrix E tubes were incubated with 300μ L of lysing buffer and 30µL of Proteinase K and incubated at 56°C. A total of 500µL of chloroform:isoamvl alcohol 138 (24:1, pH 8.0) was added to the tube, vortexed and this was followed by bead beating for 40 s at 6 139 140 m/s using a FastPrep 24 instrument (MP Biomedicals, Santa Ana, CA, USA), and centrifugation at 14,000g for 10 min. The bead beating and centrifugation steps were repeated twice more with 141 142 transfer of supernatant to clean tube followed by replacement of the aqueous phase with fresh 143 lysing buffer. The aqueous phase was then subject to DNA purification using the Maxwell LEV 144 DNA kit. The extracted DNA was quantified using Qubit HS dsDNA assay with Qubit 2.0 145 Fluorometer (Life Technologies, UK). Negative controls consisting of reagent blanks (no input 146 material) and filter blanks (filter membranes from unused Sterivex filters) were processed 147 identically as the samples for DNA extraction. Genomic DNA extracted from mock community, consisting of 10 organisms, detailed previously³⁵, was spiked into negative controls extracted 148 149 (n=8) from the reagent and filter blanks. These negative controls were also included in following 150 library preparation and high-throughput sequencing (see below).

151 Library preparation and Illumina sequencing.

152 Sequencing libraries were prepared using the Nextera XT DNA Sample Preparation Kit (Illumina Inc.). All DNA extracts (including negative controls) were cleaned up with HighPrep PCR 153 154 magnetic beads (MagBio Inc.) to remove short fragments after library preparation and quantified 155 with qPCR according to Illumina guidelines. All libraries were pooled together in equimolar 156 proportion and pooled library was quantified with Qubit HS dsDNA assay and further concentrated 157 using HighPrep PCR magnetic beads (MagBio Inc). Metagenomic sequencing on prepared libraries were performed on four lanes of Illumina HiSEQ 2500 flow cell (2x250-bp read length, 158 Rapid Run Mode) at University of Liverpool Centre for Genomic Research (Liverpool, UK). 159

160 Metagenomic read based analyses.

161 The FASTQ files were trimmed using Cutadapt v1.2.1 (Martin 2014) with a '-O 3' flag, and Sickle v1.200 (Joshi and Fass 2011) using a threshold of window quality score (≥ 20) and read length 162 after trimming (≥ 10 bp). A further trimming was applied using Trimmomatic v0.35⁴¹ to remove 163 any remaining Illumina Nextera adaptors and trim reads according to quality score with a 4-base 164 165 wide sliding window and a minimum average quality score of 20 and singlet reads were excluded in downstream analyses as well. To estimate metagenome diversity and coverage for each sample, 166 Nonpareil 3.0⁴² was used in kmer mode on the quality filtered reads. Diversity and coverage 167 information for each metagenome was estimated using command 'Nonpareil.set()' in R package 168 'Nonpareil'. MicrobeCensus⁴³ was used on quality trimmed reads to estimate average genome size 169 across samples with flag '-n 100000000' for all samples. To eliminate the potential effects of 170 171 bacteria with small genomes (i.e., Patescibacteria) on average genome size estimations, pre-172 processed reads were mapped against 12 Patescibacteria metagenome assembled genomes (MAGs) from this study (see below) and 1,037 Patescibacteria genomes from GTDB-tk⁴⁴. The 173 reads mapped in proper pair to Patescibacteria were removed using samtools ('-F2' flag). 174 175 MicrobeCensus was used again to estimate average genome size using the same parameters.

176 Metagenome assembly and mapping.

177 Filtered pair-ended reads were then pooled from each drinking water system for co-assembly, 178 which resulted in 12 paired-end FASTQ files for co-assembly, including seven from disinfected 179 (Dis) and five from non-disinfected (NonDis) systems. *De novo* co-assembly was performed using MetaSPAdes v3.10.1⁴⁵ with recommended k-values for 2x250bp reads (21,33,55,77,99,127). All 180 scaffolds shorter than 500bp were discarded and UniVec Core build 10.0 (National Center for 181 182 Biotechnology Information 2016) was used for contamination vector screening and any scaffold 183 with a significant hit to the UniVec database was removed. Reads from each samples were then mapped back to the filtered scaffolds using BWA-MEM v0.7.12 with default settings⁴⁶. 184

To eliminate the scaffolds that may have originated from sample or post-processing contamination, reads from negative controls were first mapped back to mock community genomes using BWA-MEM v0.7.12⁴⁶, and all reads not mapped in proper pair were extracted using samtools v1.3.1 (Li et al. 2009) with '-f2' flag and were considered "contaminant reads". Sample reads (S),

contaminant reads (C) and negative control reads (NC) were mapped back to filtered scaffolds in
each co-assembly. Properly-paired mapped reads were extracted using samtools v1.3.1 with '-f2'
flag from the BAM files. Relative abundance and normalized coverage deviation of each scaffold
was calculated using reads from samples and those identified as contaminant reads in negative
controls:

194 Relative abundance_s =
$$\frac{\text{Scaffold coverage}_s}{\sum_{i=1}^{n} \text{Scaffold coverage}_s}$$

195 Relative abundance_C =
$$\frac{\text{Scaffold coverage}_{C}}{\sum_{i=1}^{n} \text{Scaffold coverage}_{NC}}$$

196 Normalized coverage deviation
$$=$$
 $\frac{\text{Standard deviation of scaffold coverage}}{\text{Average scaffold coverage}}$

197 To distinguish true scaffolds from contamination, relative abundance (RA) and normalized 198 coverage deviation (NCD) estimated using sample reads (S) and contaminant reads (C) was 199 compared for all scaffolds:

200 Scaffold =
$$\begin{cases} \text{True scaffold,} & \text{if} \\ \text{Contaminant scaffold,} & \text{if} \end{cases} \begin{array}{c} RA_{C} = 0 \\ RA_{S} > RA_{C} \text{ and } NCD_{S} < NCD_{C} \\ RA_{S} = 0 \\ RA_{C} > RA_{S} \text{ and } NCD_{C} < NCD_{S} \end{cases}$$

True scaffolds, the scaffolds with higher RA and lower NCD in samples compared to negative controls, were kept for downstream analyses while contaminant scaffolds were excluded from all further analyses.

204 Nucleotide and protein composition analyses.

MASH v1.1.1⁴⁷ was used to estimate the dissimilarity between samples using quality filtered reads (with '-r' and '-m 2' flags) and dissimilarity between drinking water systems using true scaffolds with the sketch size of 100000. Prodigal v2.6.3⁴⁸ was used to identify open reading frames (ORFs) in the true scaffolds and translate ORFs to protein-coding amino acid sequences. Following prediction and translation, HMMER v3.1b2⁴⁹ was used to annotate ORFs against the Pfam database v31.0⁵⁰ with a maximum e-value of 1e – 5 and curated bit score thresholds (the gathering thresholds). Subsequently, MASH distances were calculated between drinking water

metagenomes using predicted ORFs, as well as Pfam annotated proteins with the sketch size of100000 and '-a' flag.

214 Taxonomic classification and phylogenetic analyses.

The program 'cmsearch' was implemented in Infernal v1.1.2⁵¹ to search scaffolds against SSU 215 216 rRNA covariance models (CMs) for bacteria, archaea and eukaryota; these are default models used by SSU-ALIGN v0.1⁵² using HMM-only approach and only significant hits were considered. The 217 results were filtered according to length (≥ 100 bp alignment) and e-value (< 1e - 5). SSU rRNA 218 sequences detected in contaminant scaffolds were removed and if more than one SSU gene 219 sequence was found on a single scaffold, only the longest SSU gene sequence was retained. 220 Relative abundance of each SSU gene sequence was calculated for each sampling location as 221 222 follows:

223
$$RPKM_{SSU}^{i} = \frac{\text{Scaffold coverage}^{i}}{\sum_{i=1}^{n} \text{ SSU containing Scaffold coverage per Mb}^{i} \times \text{Scaffold length per kb}^{i}}$$

224 Relative abundance^{*i*}_{SSU} =
$$\frac{\text{RPKM}^{i}_{SSU}}{\sum_{i=1}^{n} \text{RPKM of scaffold containing SSU genei}}$$

SSU rRNA gene sequences were classified using Mothur v1.33.3 (Schloss et al. 2009) with SILVA
 database⁵³ (Release 132) with a minimum confidence threshold of 80%.

227 Annotation and Comparison of functional orthologies and modules between samples

The protein-coding sequences were searched against KOfam, a HMM profile database for KEGG 228 orthology⁵⁴ with predefined score thresholds using KofamScan⁵⁵. Only KEGG orthologies (KO) 229 230 identified on scaffolds with (> 1x) coverage for each sample and those detected more than once 231 across samples within a single drinking water system were retained for further analyses. Average 232 read count for each KO was calculated using scaffold coverage, average length of reads mapped, 233 and total number of reads mapped to each scaffold in a sample using above equations. To assess 234 functions at KEGG module level, BRITE hierarchy file was retrieved from KEGG website, and KO's were categorized into KEGG modules. The abundance of KEGG module in each sample was 235 calculated using the median abundance of the detected KEGG orthologies within each module. 236 237 The completeness of each KEGG module was calculated using 'KO2MODULEclusters2.py'.

238 Metagenome binning and refining.

Anvi'o (versions: v2.2.2, v2.4.0, v4 and v5.1)⁵⁶ was used for metagenome binning and refining. 239 Briefly, CONCOCT⁵⁷ integrated in Anvi'o was used to cluster scaffolds (longer than 2500 bp) into 240 metagenome bins using tetra-nucleotide composition and coverage information across all samples 241 within each metagenomic co-assembly. The 'merge' method of CheckM v1.0.7⁵⁸ was used to 242 identify the bins that that may emerge from the same microbial population, but may have been 243 separated during automated binning process. Following merging of compatible bins, RefineM 244 $v0.0.21^{59}$ was used to automatically refine bins according to genomic properties (i.e., the mean GC 245 content, tetra-nucleotide signature and coverage) and taxonomic classification. The completeness 246 247 and redundancy of each refined bin was estimated using CheckM based on collections of lineage specific single-copy genes resulting in a total of 154 bins with greater than >50% completeness. 248 Among these bins, 130 bins had a redundancy of <10% redundancy, while 24 bins are with >10%249 250 redundancy. Further manual curation of these bins was performed using Anvi'o, resulting in 156 curated metagenome assembled genomes (MAGs). The 156 MAGs were de-replicated using dRep 251 v2.2.2⁶⁰ and MAGS with >10% redundancy were discarded which resulted in 115 dereplicated 252 MAGs with completeness >50% and reduncancy <10%. All raw sequencing data and dereplicated 253 254 MAGs are available on NCBI at BioProject number PRJNA533545.

255 MAG-level analyses

Taxonomy assignment of MAGs was performed using GTDB-Tk v0.1.3⁴⁴ with the flag 256 257 'classify wf'. Genome sizes of MAGs were estimated by multiplying the number of nucleotides 258 in the MAG with the inverse of the CheckM estimated completeness. The MAGs were annotated using the HMM profile database for KEGG orthology with predefined score thresholds using 259 KofamScan⁵⁵. The KO's for each MAG were then categorized into modules based on BRITE 260 hierarchy file retrieved from KEGG⁵⁴, and the completeness of KEGG modules in each genome 261 was calculated using script 'KO2MODULEclusters2.py'. Anvi'o was used to extract a collection 262 of 48 single-copy ribosomal proteins⁶¹ from each MAG using 'anvi-get-sequences-for-hmm-hits' 263 with a maximum number of missing ribosomal proteins of 40. Subsequently, a phylogenetic tree 264 was reconstructed using concatenated alignment of ribosomal proteins sequences using FastTree 265 v2.1.7⁶². Interactive Tree Of Life (iTOL) v4 (Letunic and Bork 2007) was used to visualize the 266 267 phylogenetic tree.

Program 'Union' in EMBOSS v6.6.0.0⁶³ was used to concatenate all scaffolds in each MAG into 268 269 a single sequence. Reads from all samples were cross-mapped to all MAGs using BWA-MEM 270 v0.7.12 with default settings and proportion of each nucleotides in MAG covered by at least 1x coverage was determined using BEDtools⁶⁴. A MAG was considered detected in a sample if $\geq 25\%$ 271 272 of its bases were covered by at least one read from the corresponding sample. This approach was 273 used to determine whether MAGs were detected in all the samples. Further, the MAGs were binned 274 into four categories based on their detection/non-detection within samples. Specifically, MAGs 275 were divided into "D-only" if there were detected in $\geq 20\%$ of the samples from the disinfected 276 systems and not detected in any samples from the non-disinfected systems, "ND-only" if there 277 were detected in $\geq 20\%$ of the samples from the non-disinfected systems and not detected in any 278 samples from the disinfected systems, "both" if there were detected in $\geq 20\%$ of disinfected and 279 non-disinfected systems, while the remaining MAGs were classified in the "other" category. Subsequently, reads from all samples were cross-mapped back to all the MAGs using BBMap 280 v38.24⁶⁵ with a minimum identity of 90%, and 'ambiguous=best' and 'pairedonly=t' flags. After 281 filtering for detection (see above), reads per kilobase of per million reads (RPKM) for each MAG 282 283 and each sample were calculated using equation:

284
$$RPKM = \frac{Number of reads mapped to MAG}{Total number of reads in sample per Million \times MAG length in kbp}$$

285 Statistics

Differences between disinfected and non-disinfected systems for (1) Mash distances distributions, 286 (2) relative abundances were determined using Permutational ANOVA and Pearson's correlations 287 288 distances were estimated in R. BioEnv "sinkr" between pairwise mash in and "vegan"⁶⁶ packages were used to identify (https://github.com/menugget/sinkr) 289 290 environmental parameters (i.e., water chemistry) and their combinations that explain the 291 differences in the structure (i.e., Mash distances between samples estimated using reads) and 292 functional potential (i.e., Bray Curtis distance estimated between samples using KO abundance 293 (i.e., RPKM). BioEnv permutes through 2ⁿ⁻¹ possible combination of selected environmental parameters, 511 combinations in this case, and selects the combinations of scaled environmental 294 295 variables which capture maximum correlation between dissimilarities of community datasets water 296 chemistry and microbial community structure or functional potential. While, BioEnv analyses 297 identified combination of variables that are highly correlated with differences in microbial 298 community structure of functional potential, it does not identify the proportion of variance in 299 microbial community structure of functional potential explained by individual variables or their 300 combination. To this end, we used distance-based redundancy analysis (dbRDA) to perform 301 constrained ordinations on community structure and functional potential to bypass the limitation 302 of usual RDA and CCA, which can only use Euclidean distance measure. Function dbrda() from 303 'vegan' package was used with pairwise Mash distances calculated between samples estimated 304 using reads based Mash distance and Bray-Curtis distances based on KO RPKM to investigate 305 relationships between the environmental variables and community data on both nucleotide 306 composition and KO level. The function varpart() in the vegan package was used to determine the 307 fraction of variation captured parameters identified as significantly associated with read-based Mash and KO relative abundance-based Bray-Curtis distance matrices. DeSeq2 package v1.18.1⁶⁷ 308 309 was used to identify differentially abundant KEGG modules between disinfected and non-310 disinfected systems by only considering KEGG modules with a maximum of one block missing and equal to or greater than 50% complete. The median scaffold-length normalized read count of 311 312 KO's within each module were used in DESeq2 analyses with a maximum adjusted P-value of 313 0.005.

314 **RESULTS AND DISCUSSION**

315 Water quality parameters across disinfected and non-disinfected DWDS.

316 Sampling was conducted in seven DWDSs with disinfectant residual between April-August of 317 2013 and at five DWDSs without disinfectant residual between October-December 2015. The 318 water chemistry varied between the DWDSs considering they were supplied by different DWTPs, 319 our sampling campaign also captures seasonal differences between locations (Figure 1) (Table S1). 320 Specifically, water temperatures were higher ($\sim 5^{\circ}$ C) for the disinfected samples compared to the 321 non-disinfected samples. While the pH, DO, nitrogen species (i.e., ammonium and nitrate) and TOC measurements were not significantly different between disinfected and non-disinfected 322 323 samples, the measured phosphate and total chlorine concentrations were significantly different 324 (p < 0.05). Specifically, the average total chlorine concentrations in disinfected systems 0.37 mg 325 Cl₂/l (range: 0.1-0.73 mg Cl₂/l) while no disinfectants residuals were measurable in the nondisinfected systems. The average phosphate concentrations were 2.3 mg PO_4^{3-}/l while no 326

phosphate was measurable in non-disinfected samples. Phosphate was higher in the disinfected systems as it is likely to be used for corrosion control⁶⁸. While we were unable to obtain information on source water type (i.e., ground vs surface water) used for production of drinking water supplied to the sampled DWDS, conductivity measurements suggested DWDS in both systems were supplied by a DWTPs drawing from surface and ground water sources (Figure 1).

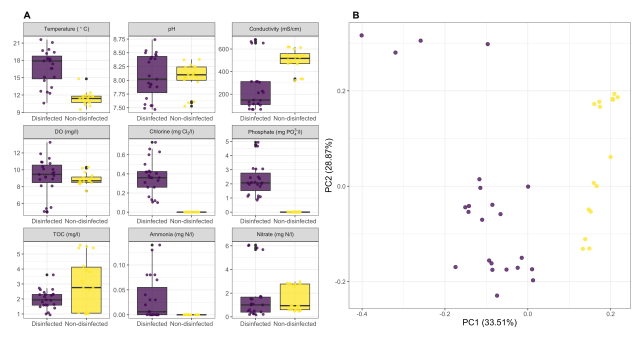


Figure 1: Summary of water chemistry parameters measured for samples collected from disinfected (purple) and non-disinfected systems (yellow). (B) Principle component analyses using Euclidean distances for measured water chemistry parameters indicates distinct clustering of samples from disinfected and non-disinfected systems.

332

333 Summary of metagenomic data set.

334 Metagenomic analyses was used to assess the association between presence/absence of disinfectant 335 residual with the structure and functional potential of the drinking water microbiome. A total of 336 41 drinking water samples were collected from DWDSs with (i.e., chlorine) from the United 337 Kingdom (n=23), while those collected from the Netherlands (n=18) did not have a disinfectant 338 residual. Ouality trimming of raw metagenomic data resulted in the retention of 638 million paired-339 end reads. Co-assembly for each drinking water system was carried out by combining reads from 340 individual sampling location within each drinking water system (Table 1). De novo co-assembly generated 0.04-1.81 million true scaffolds for each sampling location after discarding scaffolds 341 342 shorter than 500bp and contaminant scaffolds (Table 1) with an N50 value ranged from 775 bp to

- 343 3300 bp. The proportion of quality trimmed reads mapping back to true scaffolds ranged from 67%
- to 99% (Table 1) across all samples.

345	Table 1: Sequencing and de novo co-assembly statistics for metagenomes from 12 drinking
346	water systems.

Drinking water system	Paired end Reads (millions)	Scaffolds (>500 bp)	True scaffolds	True scaffold assembly size (Mbp)	% Mapped reads	GC content (%)	N50 (bp)	ORFs per Mbp	Coding density
D1	195.73	555493	546375	615.10	99.02	54.66	1131	1403.68	0.48
D2	46.87	38567	36733	53.03	96.24	55.34	2112	1419.84	0.64
D3	17.40	192457	190882	249.69	91.15	57.82	1531	1498.24	0.63
D4	36.01	123852	122486	204.78	93.03	57.57	3300	1316.54	0.60
D5	36.74	227196	225149	269.12	88.73	59.09	1313	1527.12	0.60
D6	17.39	42209	41459	57.23	95.89	59.16	1641	1504.23	0.65
D8	19.4	77973	76996	108.07	95.38	61.07	1751	1475.71	0.68
ND1	45.52	521371	517773	472.02	83.82	53.75	855	1803.21	0.61
ND2	25.98	363819	361304	316.18	75.03	53.44	802	1807.05	0.56
ND3	48.63	667992	663968	562.73	81.63	52.93	775	1838.06	0.60
ND4	17.78	164328	163361	143.22	66.73	56.48	808	1822.84	0.63
ND5	130.92	1812573	1804048	1834.75	93.74	56.38	1005	1672.04	0.60

347 D=disinfected, ND=non-disinfected, N50=minimum contig length that account for 50% of the

348 bases, ORF=open reading frame.

349 Non-disinfected systems are more diverse than disinfected systems.

350 Non-disinfected systems were significantly (p < 0.0001) more diverse compared to systems that maintained a disinfectant residual (Figure 2A) based on Nonpareil estimated diversity index⁴². This 351 observation is consistent with previous comparisons of bulk water⁶⁹ and biofilm³⁹ samples from 352 353 disinfected and non-disinfected systems. Lower diversity in disinfected systems is likely due to 354 stronger selective pressure of the disinfectant residual as compared to that nutrient limitation in 355 non-disinfected systems. As a result of the higher diversity in non-disinfected systems, the 356 metagenomic sequencing for these samples provided significantly lower coverage of the sampled 357 microbial community (Figure 2B) as compared to systems with a disinfectant residual (p < p358 0.0001).

359 Microbial community membership and structure is different between disinfected and non-360 disinfected systems.

361 We used 2,872 small-subunit (SSU) rRNA genes (2742 genes > 100 bp) identified on the 362 assembled scaffolds to determine community membership and structure across sampling locations 363 (Supplementary file 1, Supplementary table 2). While bacteria were dominant members of the drinking water microbiome in both types of systems (2C, 2D), the relative abundance of archaea 364 and eukaryota were dependent on the presence/absence of disinfectant residual (Figure 2C, 2E). 365 366 Specifically, the relative abundance of eukaryota was higher in disinfected systems as compared to non-disinfected system (2C), while archaea were ubiquitous across non-disinfected samples 367 368 (Figure 2C, E) they were only detected in a single disinfected sample (D2). Non-disinfected 369 systems were taxonomically more diverse, with respect to bacteria and archaea, as compared to 370 disinfected systems. Specifically, a total of 14 bacterial and 6 archaeal phyla were detected in one 371 or more non-disinfected systems that were not detected in any of the disinfected systems. Several 372 of these unique phyla, while not dominant in non-disinfected systems, were detected at relative 373 abundances between 1-5% (e.g., Nitrospirae, Nanoarchaeota).

374 The bacterial community was dominated by Proteobacteria, in particular Alphaproteobacteria and 375 Gammaproteobacteria, in both disinfected and non-disinfected systems with Deltaproteobacteria 376 being much more prevalent and abundant in non-disinfected systems (Figure 2D). Actinobacteria 377 were more abundant than Proteobacteria in two drinking water systems and constituted 44% and 33% of the community in systems D4 and ND1, respectively. Overall, the relative abundance of 378 Proteobacteria was higher in disinfected systems, ranging from 28% to 90%, as compared to non-379 380 disinfected systems, ranging from 30% to 57%. Patescibacteria was the second most abundant 381 phylum across non-disinfected systems, constituting 15% to 29% of the SSU rRNA genes, while 382 they were only detected in one disinfected sample (D2) with a relative abundance of 1%. Within 383 Patescibacteria, Parcubacteria were the most commonly detected phyla followed by Microgenomatia and Gracilibacteria. 384

The observed differences between disinfected and non-disinfected DWDS for bacteria and archaea are largely consistent with a previous meta-analyses of amplicon sequencing data from the 16S rRNA gene⁶⁹. In contrast to bacteria and archaea, results from eukaryotes, which have not been systematically investigated in the drinking water microbiome, were surprising in terms of their

389 higher relative abundance eukaryotic in disinfected as compared to non-disinfected systems. For 390 instance, SSU rRNA genes associated *Nematoda* were detected in nearly every disinfected system, 391 but were not detected in non-disinfected systems. Specifically, SSU rRNA genes from two free-392 living nematode genera, i.e. Araeolaimida and Monhysterida, were detected in five of the eight 393 disinfected systems. Similarly, SSU rRNA genes from the phylum Rotifera were only detected in 394 disinfected systems and were largely associated with the monogonont rotifers within the genera 395 Ploimida. While the relative abundance of scaffolds determined to be of eukaryotic origin was 396 higher in disinfected compared to non-disinfected systems, this does not mean that eukaryotes 397 were proportionally larger part of the drinking water microbiome in disinfected compared to the 398 non-disinfected systems. Genome sizes of picoeukaryotic microbes can be orders of magnitude 399 larger than that of bacteria and archaea and vary significantly between picoeukaryotes themselves. Further, the higher overall diversity and lower sequencing coverage (Figure 1) could also have 400 401 resulted in under sampling of the eukaryotic community in non-disinfected systems.

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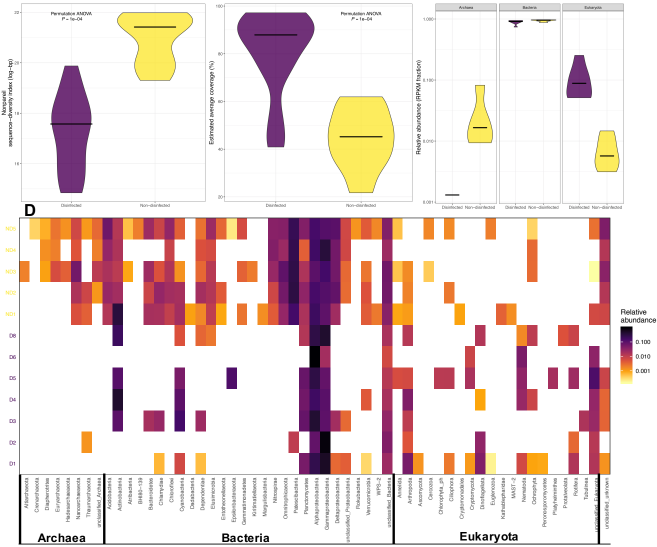


Figure 2: Comparison of (A) diversity and (B) coverage between disinfected and non-disinfected drinking water systems estimated using Nonpareil. (C) Comparison of relative abundance of bacterial, archaeal, and eukaryotic communities in drinking water systems with and without disinfectant residuals. (D) Log10 transformed relative abundance of different phyla (classes for phylum Proteobacteria) across sampling location for the bacteria, archaea, and eukaryota.

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403 Drinking water systems cluster at the nucleotide level based on presence/absence of 404 disinfectant residuals. Samples (for read based analyses) and drinking water systems (for scaffold 405 based analyses) clustered with each other based on the presence/absence of disinfectant residuals 406 (Figure 3A and 3B) based on Mash distance estimates. We further evaluated the significance and 407 explanatory power of measured water chemistry parameters in explaining the observed clustering 408 between disinfected and non-disinfected systems. To do this, we initially performed BioEnv 409 analyses to identify water chemistry parameters and their combinations that were highly correlated 410 with observed Mash distances between samples (Supplementary Table 3). This identified chlorine 411 as being strongly correlated with the Mash distances between samples (R=0.54, p<0.001) while 412 the maximum correlation between water chemistry and Mash distances was observed for a 413 combination of chlorine, phosphate, and TOC (R=0.62, p<0.001). We subsequently utilized 414 dbRDA to independently determine the environmental/water chemistry variables most 415 significantly associated with Mash distances between samples. While chlorine was identified as a 416 significant variable (p < 0.01), dbRDA identified conductivity (p < 0.001) and DO (p < 0.01) as 417 significant variables. Finally, variance partitioning analyses was used to determine the proportion 418 of variance in the Mash distance matrices explained by individual and combination of variables 419 identified as significant by dbRDA (Table S5). This resulted in chlorine, conductivity, and DO 420 individually explaining $\sim 17\%$, 12%, and 1% of the variance in the Mash distance matrix, with 421 $\sim 60\%$ of the variance unexplained by these three variables.

422 We further compared the distribution of Mash distances between drinking water metagenomes 423 within disinfected, within non-disinfected, and between disinfected and non-disinfected systems. 424 Mash distances between drinking water metagenomes from disinfected systems were significantly 425 different (p < 0.0001) and exhibited a lower mean for disinfected as compared to non-disinfected 426 systems. Further, the pairwise Mash-distances between disinfected and non-disinfected systems 427 were significantly different and higher from those estimated within each category (i.e., disinfected 428 or non-disinfected). This was consistent for both read- and scaffold-based analyses (Figure 3D, 429 3E). Finally, the average pairwise Mash distances estimated using reads (i.e., between samples) 430 and scaffolds (i.e., between DWDSs) were highly correlated (Pearson's R = 0.95, P < 0.05) (Figure 431 3C), indicating the *de novo* assembly process did not result in loss of information on factors driving 432 the differences between disinfected and non-disinfected systems.

These analyses provide a few key insights. First, Mash distance-based (both read and scaffold based) clustering of samples occurs depending on presence and absence of disinfectant residual suggests that the microbial communities are more similar within each group (i.e., disinfected and non-disinfected) and dissimilar between the two groups (i.e., disinfected vs non-disinfected). Second, while disinfected and non-disinfected samples cluster distinctly from each other, disinfected systems exhibit lower nucleotide-level heterogeneity as compared to their non439 disinfected systems indicating that the factors governing microbial community in disinfected 440 systems likely impose stronger selective pressures on the microbial community as compared to 441 those in non-disinfected systems. Third, non-disinfected systems exhibit greater diversity not only 442 within a system (Figure 2) but also across systems as compared to disinfected systems. Despite the 443 strong correlation between pairwise Mash distances of reads and scaffolds (Figure 3F), the median 444 Mash distances for pairwise comparison of samples within each type of system (i.e., disinfected 445 and non-disinfected) is higher for the scaffold-based analyses as compared to the read-based 446 analyses. This is likely from the omission of low abundance microorganisms during *de novo* 447 assembly and thus suggests that composition of medium-to-high abundance organisms are likely 448 to be more variable between non-disinfected systems as compared to disinfected systems.

449 Finally, while the water chemistry and environmental parameters between disinfected and non-450 disinfected systems were distinct (Figure 1B), the parameters that most strongly correlated with 451 Mash distances between samples were limited to a combination of chlorine, phosphate, and TOC 452 for BioEnv analyses and chlorine, conductivity, and DO based on dbRDA. Both independent 453 exploratory analyses consistently identified chlorine presence/absence and concentration as one of 454 the key drivers of difference in microbial communities across the samples. Further, variance 455 partition analyses indicated that ~17% of the variance in the Mash distance matrix was driven 456 exclusively by chlorine; this make chlorine the most important parameter measured as part of this 457 study in terms of differentiating between drinking water metagenomes. The significance of 458 phosphate determined by BioEnv analyses is likely because chlorine and phosphate concentrations 459 are inherently associated due to common use of the latter for corrosion control in DWDSs that maintain a chlorine residual⁶⁸. Further, while it is unlikely that DO (identified as significant by 460 461 dbRDA) directly affects microbial community composition (all DO concentrations were near or 462 greater than saturation), it is possible that this may reflect the use of advanced oxidation process 463 (e.g., ozonation) during drinking water treatment. Similarly, conductivity (identified as significant by dbRDA) is unlikely to directly influence the microbial community, but rather this may reflect 464 465 the source water type and treatment processes being used for drinking water production. 466 Specifically, source water derived from ground water sources or from reservoirs under the 467 influence of ground water typically have much higher conductivities than those that rely on surface 468 water supply. Similarly, chemicals used for softening and coagulation/flocculation processes may 469 influence water conductivity. Thus, we speculate that the influence of conductivity may serve as

470 surrogate for a combination of source water and treatment process. These analyses clearly identify

- 471 chlorine as one of the major measured parameters driving the Mash distances between samples,
- followed by conductivity (potential surrogate for source water and treatment process). Further, the
- 473 fact the major proportion of the variance remains unexplained suggests that additional aspects such
- 474 as treatment process configuration, DWDS characteristics, and other water chemistry parameters
- 475 which were not characterized/measured as part of this study also likely play a strong role in
- 476 differentiating between microbial communities in disinfected and non-disinfected drinking water
- 477 systems.

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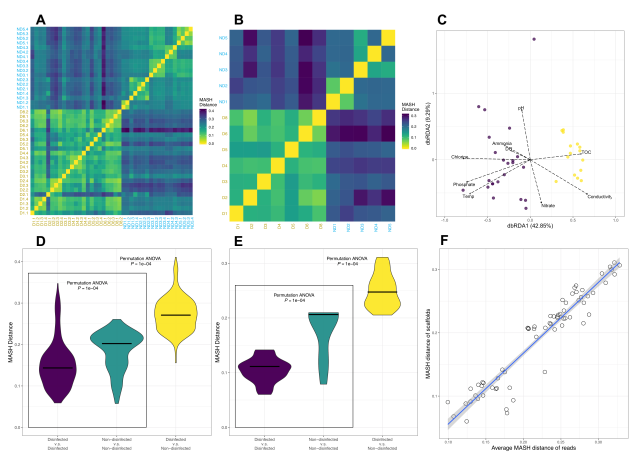


Figure 3: Comparison of nucleotide composition using paired reads each from each sample and true scaffolds in each drinking water system according to Mash distance. (A, B) Heatmaps based on pairwise Mash distances of reads and scaffolds. Heatmaps are colored according to Mash distance; yellow denotes a distance of 0. Labels on x- and y-axis are colored according to disinfection strategies. (C). NMDS clustering of read based Mash distances between samples with vectors representing water chemistry/environmental parameters implemented using dbRDA. (D, E) Violin plots indicating the distribution of pairwise Mash distances of reads and scaffolds. Plots are colored according to the system type for which pairwise comparisons were conducted. Purple denotes comparisons between disinfected samples, yellow denotes comparisons between non-disinfected samples, and green denotes comparisons between disinfected and non-disinfected samples. (F) Correlation between average Mash distances of reads across samples and Mash distances of scaffolds across sampling locations.

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483 Protein coding sequences cluster based on presence/absence of disinfectant residuals. A total 484 of 8 million protein coding sequences were predicted and translated from true scaffolds, of which 485 approximately 17 to 27% were annotated against KEGG database (Table S6). Consistent with the 486 nucleotide-level analyses, samples clustered based on the presence and absence of disinfectant 487 residual (Figure 4A, 4B, 4C) rather than by DWDS. Further, BioEnv analyses identified the 488 combination of chlorine, phosphate, and ammonia as being strongly and significantly correlated 489 (R = 0.392, P < 0.001) with Bray-Curtis distances between samples estimated using abundance (i.e., 490 RPKM) of KOs (Table S7). Similar to nucleotide based analyses, chlorine presence/absence and 491 concentration was the measured parameter more strongly and significantly associated with 492 differences in functional potential between samples at the single parameter level (R=0.382, 493 p<0.001). In contrast to nucleotide based analyses, conductivity and chlorine were the only two variables identified as significantly associated with Bray-Curtis distances between samples 494 495 estimated using relative abundance of KO's in samples using dbRDA (Table S8). Variance 496 partitioning indicated that both conductivity and chlorine individually explained approximately 497 6.5% of the variance in Bray-Curtis distance matrix estimated using KO abundance. A comparison 498 of the pairwise Mash distances within each group (i.e., disinfected, non-disinfected) and between 499 them indicated that the diversity in functional potential was significantly different for both 500 predicted protein coding-sequences and KEGG annotated proteins (p < 0.0001). The median value 501 of Mash distances between the non-disinfected samples was greater than that for disinfected 502 samples (Figure 4D, 4E) and the differences in Mash distances between two groups was larger 503 than the distances within each group. And finally, despite the fact that only 17-27% of predicted 504 proteins were annotated against the KEGG database, the Mash distances between metagenomes 505 estimated using all predicted protein coding sequences and those that were annotated against the 506 KEGG database were highly correlated (Pearson's $R \approx 1.00$, P < 0.05) (Figure 4F), suggesting 507 that focusing on annotated proteins does not result in significant loss of information while 508 performing direct comparisons between samples from disinfected and non-disinfected systems.

509 These analyses based on protein coding sequencing provide several key insights. First, clustering 510 of samples into disinfected and non-disinfected groups is consistent for both community 511 composition (i.e., read-based nucleotide composition analyses) and functional potential, 512 irrespective of the use of all predicted ORF's and KEGG annotated protein sequences. Non-513 disinfected systems are significantly more heterogeneous across systems as compared to their

514 disinfected counterparts. This suggests that selection pressures exerted within disinfected systems 515 are not only evident at community structure/membership (Figure 3), but also evident at the 516 community functional potential level. Further, consistent with microbial community composition, 517 chlorine was also identified as one of the key measured parameters driving differences between 518 samples based on functional potential using both BioEnv and dbRDA analyses. In contrast to TOC 519 which was included in the BioEnv parameter combination for microbial community composition 520 level analyses, ammonia was identified as part of the combination at the functional potential level. 521 While the exact reason behind this difference cannot be ascertained in this study, this may likely 522 be associated with the fact that non-disinfected systems are severely nitrogen limited as compared 523 to disinfected systems, while both systems were likely not carbon limited. Similar to the nucleotide 524 level analyses, both conductivity and chlorine were identified as significantly (p<0.01) associated 525 with differences between samples, with variance partitioning analyses allocating equal amount of 526 variation to both parameters (Table S9). As speculated above, if conductivity is considered a signal 527 for source water and treatment process type, then the impact of these two parameters on the 528 functional potential of microbial community is relatively similar to that of presence/absence of the 529 disinfectant residual. Finally, the residuals from the variance partitioning analyses were noticeably 530 larger (84%) for functional potential analyses as compared to the microbial community 531 composition (60%), suggesting that the impact of unmeasured/uncharacterized factors/parameters 532 on microbial community functional potential was significantly larger than their impact on 533 community composition. While it cannot be ruled out, it is unlikely that the higher fraction of 534 unexplained variation was due to only a proportion of ORFs being annotated; this is because Mash 535 distances estimated using only KEGG annotated ORFs were highly correlated with those estimated 536 using all predicted ORFs using suggesting little to minimal loss of discriminatory power while 537 using only annotated proteins.

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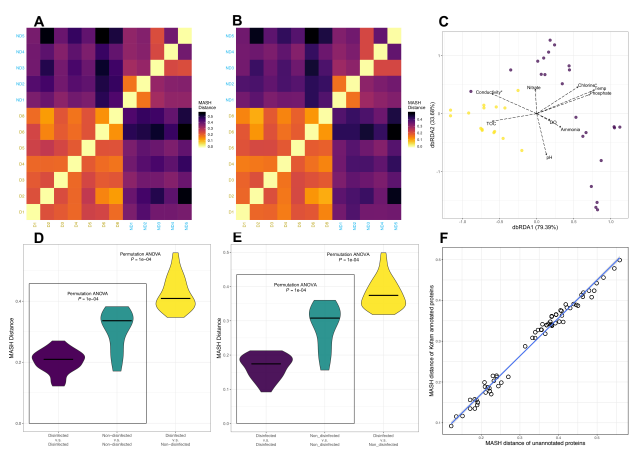


Figure 4: Comparison of functional potential among all and KEGG protein-coding amino acid sequences across sampling locations. This analysis estimates dissimilarity in amino acid composition of samples, similar to the nucleotide composition analyses presented earlier. (A, B) Heatmaps based on pairwise Mash distances of all protein coding sequences and Bray-Curtis distances using KO. Heatmaps are colored according to Mash/Bray-Curtis distance; yellow denotes a distance of 0. Labels on x- and y-axis are colored according to disinfection strategies; dark golden denotes samples with chlorine, while blue denotes samples without disinfectant residuals. C). NMDS clustering of using Bray-Curtis distances using KO abundances between samples with vectors representing water chemistry/environmental parameters implemented using dbRDA. Violin plots indicating the distribution of pairwise (D) Mash distances of all and (E) Bray-Curtis distances KEGG annotated proteins. Crossbars indicate the median value of Mash distances. (F) Correlation between pairwise Mash distances estimated using all and Bray-Curtis distances for KEGG annotated proteins. Crossbars indicate the median value of Mash distances.

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542 Differentially abundant metabolic modules are consistent with microbial growth control

- **strategies.** A total of 7,281 KOs were identified in all samples with 5,922 remaining post-filtering
- based on scaffold coverage (>1x) and frequency of KO detection in each drinking water system
- 545 (detected more than once) (Table S10). The 5,922 KO's were further categorized into 540 KEGG
- 546 modules and upon further filtering to remove KEGG modules with no more than one missing block
- and greater than equal to 50% completion, a total of 208 KEGG modules were retained (Table

548 S11). Of these, a total of 57 KEGG modules exhibited significantly differential abundance between 549 disinfected and non-disinfected samples (p-value < 0.005) (Table S12, S13). Modules associated 550 with ribosomal synthesis, ribonucleotide biosynthesis, and RNA polymerase were ignored from 551 further consideration. Similarly, modules most likely associated with plant metabolism (e.g., Crassulacean acid metabolism) were also ignored. This resulted in 29 and 22 KEGG modules that 552 553 were more abundant in non-disinfected system and disinfected systems, respectively. These 554 included modules associated with energy metabolism (disinfected, i.e. D=2, non-disinfected, i.e., 555 ND=5), carbohydrate and lipid metabolism (D=11, ND=10), nucleotide and amino acid 556 metabolism (D=5, ND=13), and secondary metabolism (D=4, ND=1).

557 Metabolic modules associated with polyamine biosynthesis, aromatics degradation, terpenoid 558 biosynthesis, and fatty acid metabolism were significantly enriched in disinfected systems. 559 Specifically, metabolic pathways associated with benzene (M00548) and benzoate (M00551) 560 degradation to catechol and methyl catechol were highly enriched in disinfected systems. Further, 561 eukaryota-associated metabolic modules such as terpenoid backbone biosynthesis (M00367) and 562 modules associated with peroxisomal beta-oxidation of very long chain fatty acids (M00861) are 563 likely to be enriched in the disinfected systems due to the higher relative abundance of eukaryota 564 in samples collected from disinfected as compared to non-disinfected systems respectively. 565 Further, modules related to γ -aminobutyrate (GABA) metabolism (M00136, M00027) were 566 enriched in disinfected systems. The GABA shunt pathway converts glutamate to GABA using 567 glutamate decarboxylase (GAD), followed by reversible conversion from α -ketoglutarate to 568 succinate semialdehyde (SSA) through the activity of GABA transaminase (GABA-AT), and 569 finally succinate is formed by succinate semialdehyde dehydrogenase (SSDH) activity. In contrast, 570 the key metabolic modules enriched in non-disinfected systems were associated with carbon 571 fixation and methane metabolism (M00377, M00620, and M00422) and nitrogen fixation 572 (M00175) (Table S13). The differentially abundant carbon fixation modules included the Wood-573 Ljungdahl pathway, Acetyl-CoA pathway, and the incomplete reductive citrate cycle. These 574 pathways can fix carbon dioxide to produce acetyl-CoA which can then be converted to other necessary biosynthetic intermediates of the carbon metabolism^{70, 71}. 575

576 The enrichment of carbon and nitrogen fixation modules in non-disinfected systems is consistent 577 with nutrient limitation as the strategy for microbial growth control in non-disinfected drinking 578 water systems. While the measured total organic carbon concentrations in non-disinfected systems 579 did not indicate carbon limited conditions, DWTP's supplying water to non-disinfected DWDSs typically achieve far superior levels of removal of assimilable organic carbon (AOC)²⁸. Similarly, 580 the nitrogen availability in the form of ammonia was consistently zero for non-disinfected systems 581 582 compared to disinfected systems which has residual ammonia concentrations ranged from 0.01-583 0.15 mg/l of ammonia-nitrogen. In contrast, the enrichment of KEGG modules associated with 584 GABA metabolism in disinfected systems suggests the potential importance of stress protection 585 and utilization of microbial decay products. Previous studies have shown that GABA metabolism 586 is associated with bacterial survival under various types of environmental stresses, including oxidative stress, acidic stress, and osmotic stress ⁷²⁻⁷⁵. Meanwhile, GABA can also play a 587 588 significant role in nitrogen metabolism of bacteria. For instance, putrescine formed due to the 589 breakdown of amino acids potentially from decaying biomass, can be converted to GABA (M00136) and finally metabolized via GABA shunt pathway⁷⁴. The enrichment of GABA 590 metabolism in disinfected systems may thus be associated with greater protection against 591 592 disinfectant stress and by allowing access to decay products from inactivated cells.

593 Average genome size differences between disinfected and non-disinfected system vary 594 between read-based and MAG-based analyses. We further investigated differences in genome 595 sizes between disinfected and non-disinfected systems. Genome sizes can be indicative of the metabolic capacity of microorganisms⁷⁶ and thus provide insights in the whether the 596 597 presence/absence of disinfectants selects for organisms with larger or smaller metabolic 598 repertoire⁷⁷ in comparison to organisms detected in non-disinfected systems. Average genomes 599 size estimates from disinfected systems were significantly larger than those from non-disinfected 600 systems based on MicrobeCensus estimates using entire metagenomic data (Figure 5A); this was 601 consistent even when reads mapping to phyla known to have smaller genomes (e.g., Patescibacteria) were selectively removed from the data set (Figure 5B). This suggests that 602 603 microorganisms in disinfected systems may be metabolically more diverse than their counterparts 604 from non-disinfected systems. Nonetheless, these results were not consistent when compared with 605 estimated genome sizes of MAGs recovered as part of this study. Specifically, we recovered a total 606 of 115 dereplicated MAGs with completeness >50% and redundancy <10% (Table S14). These 607 115 MAGS were binned into four categories based on the detection or non-detection in disinfected 608 samples. Specifically, MAGs were binned in the four groups (i.e., both, D-only, ND-only, and

609 other) based on genome coverage and detection frequency criteria outlined in the materials and 610 methods section (see MAG-level analyses) (Table S15). This resulted in 9, 16, 41, and 49 MAGs 611 were categorized as both, D-only, ND-only, and other (Figure 5C, 5D) (Table S14). In contrast to 612 read-based estimates of average genome size, MAG-based genome size estimates were not 613 significantly different between the three key categories (Both=4.4±0.77Mbp, D-614 only=3.22±0.81Mbp, ND-only=3.48±1.22Mbp) (Figure 5E). Yet, the ND-only category consisted of several smaller genomes (n=17) compared to the D category. The lack of genome size 615 616 differences between disinfected and non-disinfected samples based on MAG-based analyses 617 compared to metagenome-level read-based analyses may be due to the proportion of read-based data represented by the MAGs. Specifically, while 60-90% of the reads from disinfected systems 618 619 mapped to the 115 MAGs with the mapping rate from non-disinfected systems averaging around 620 50% (Figure 5F). Thus, it is likely that the metagenomic assembly and binning process may have resulted in suboptimal recovery of smaller genomes from non-disinfected sample which eliminates 621 622 the signal in genome size differences observed at the metagenome level.

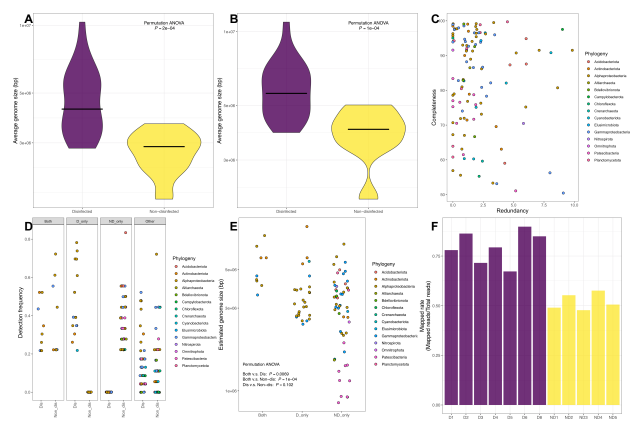


Figure 5: Violin plots indicating the genome size estimated by MicrobeCensus (a) before and (b) after Patescibacteria removal suggest average genome sizes in disinfected systems are larger than those in non-disinfected systems. (C) The 115 MAGs assembled with >50% completeness and <10% redundancy were categorized into (D) four groups based on their detection frequency in disinfected and non-disinfected systems. (E) While the estimated genome sizes of MAGs in D_only, ND_only, and Both categories were not significantly different, the ND_only category consisted of large number of smaller genomes. (F) Barplot indicating the proportion of reads mapped to 115 genomes across samples. Purple and yellow denotes samples from systems with and without a disinfectant residual, respectively.

623 Metabolic capacities differ between metagenome assembled genomes from disinfected and

624 non-disinfected systems. Clustering of MAGs (Figure 6A) based on presence/absence of KEGG

625 metabolic modules was largely driven by phylogenetic placement of MAGs, rather than their 626 classifications into groups based on the detection frequencies in disinfected and non-disinfected 627 systems (Figure 6B). Further, there was insufficient representation of MAGs from D-only/ND-628 only categories across all phylogenetic clusters (e.g., at the species or genus level) to allow for 629 direct comparisons of metabolic potential of closely related MAGs exclusively frequent in 630 disinfected and non-disinfected systems. Nonetheless, there were seven and five high quality (completeness > 90%, redundancy <10%) alphaproteobacterial MAGs that were exclusively 631 632 frequent in disinfected (average detection frequency in disinfected =55%) and non-disinfected systems (average detection frequency in non-disinfected=29%) (Figure 6A). Thus, we focused 633

metabolic module comparisons between these 12 MAGs only. We evaluated differences in metabolic capacity of these MAGs by (1) considering all KEGG modules \geq 75% complete within MAGs to be present in them and (2) all modules present in more than half of the high-quality MAGs within each category to be present within each category (Figure 6C, Table S16). We subsequently confirmed the presence/absence of genes within key metabolic modules using KOlevel annotation for these 12 MAGs (Table S17).

640 The metabolic module associated with the glyoxylate cycle (M00012) was present in 86% of the 641 MAGs in the D-only category while being only partially complete in most of the ND-only MAGs. Specifically, isocitrate lyase (aceA: K01637) and malate synthase (aceB: K01638), two key genes 642 643 involved in the glyoxylate cycle, were present in 40% and 100% of the MAGs from D-only, respectively and both genes were absent in all ND-only MAGs included in this analysis. The 644 glyoxylate shunt is associated with use of non-carbohydrate carbon sources (i.e., via 645 gluconeogensis), such as break down products from lipids, fatty acids etc^{78} . The likely benefit of 646 647 the glyoxylate shunt and associated use of lipids and fatty acids as carbon source is further 648 supported by the fact that KEGG module associated with propanoyl-coA metabolism (M00741) 649 was complete in 6/7 as compared to 2/5 MAGs from the D-only and ND-only categories. This 650 metabolic module is associated with the conversion of propanoyl-coA, a toxic byproduct of fatty 651 and amino acid degradation, to succinyl-coA. High biomass turnover rates, due to disinfectant induced microbial inactivation, may result in resource pools enriched in microbial decay products 652 thus allowing a significant advantage for microorganisms capable of necrotrophic growth⁷⁹ aided 653 654 by the glyoxylate cycle. Thus, it is feasible that the ability to utilize microbial decay products may 655 provide a distinct advantage to microorganisms inhabiting disinfected DWDSs.

656 The glyoxylate shunt may provide additional benefits for microorganisms subject to disinfectant stress via enhanced fitness to oxidative stress⁷⁸ and enhanced persistence when challenged with 657 other chemical stressors (e.g., antibiotics)⁸⁰. In contrast to module level analyses at the 658 659 metagenome level where carbon fixation capacity was significantly more abundant in non-660 disinfected as compared to disinfected systems, the alphaproteobacterial MAGs from D-only 661 systems harbored the capacity for carbon fixation via the Calvin-Benson-Bassham cycle (M00165, M00166, M00167) while this capacity was mostly absent from MAGs in the ND-only category. 662 663 Nonetheless, these MAG-based analyses are limited in phylogenetic scope and does not weigh the

664 importance of MAGs to their respective systems based on their relative abundance. Hence, we suggest that metagenome-level analyses should take precedence over findings at the MAG level 665 666 when they conflict. While the glyoxylate shunt was not identified as a significantly enriched in the 667 disinfected systems at the metagenome level analyses, the GABA shunt (metagenome level 668 analyses) and glyoxylate shunt (MAG level analyses) may both be involved in use of non-669 carbohydrate carbon sources suggesting that re-use of microbial decay products may indeed be a 670 key bacterial trait that allows for persistence in disinfected drinking water systems. Further lending 671 support to this is that that propanoyl-coA metabolism was identified as significantly enriched in 672 disinfected systems compared to non-disinfected systems using both metagenome-level and MAG-673 level analyses. Interestingly, only one metabolic module was identified as being more than twice 674 as prevalent in alphaproteobacterial MAGs from ND-only systems compared to those from D-only 675 systems (i.e., M00156: cbb3-type Cytochrome C oxidase). The greater metabolic capacity of 676 alphaproteobacterial D-only MAGs compared to ND-only MAGs was also confirmed at the KO-677 level by evaluating the presence/absence of KO's in the D-only and ND-only category MAGs. 678 Specifically, while only 8 KOs were twice or more as prevalent in ND-only MAGs compared to 679 D-only MAGs, the total KOs that were twice or more as prevalent in D-only MAGs was 109. This 680 supports the conclusion that metabolic repertoire of alphaproteobacterial D-only MAGs is 681 significantly larger than that of ND-only MAGs. Notable among the genes that were twice as 682 frequent in D-only MAGs compared ND-only MAGs included those involve in SOS-response 683 mediated mutagenesis involving trans-lesion synthesis (i.e., *imuA*: K14160, *imuB*: K14161, and dnaE2: K14162)⁸¹, glyoxylate reductase (gyaR: K00015) which may be likely involved in 684 regulating glyoxylate concentrations, and vitamin B12 transporter (*btuB*: K16092). SOS response 685 is typically activated in response to significant cellular accumulation of damaged DNA ⁸² and *imuA* 686 and *imuB* co-expression with *dnaE2* has been shown to be responsive to UV damage 81 . Thus, the 687 688 higher prevalence of SOS response related genes in D only MAGs may be associated with the DNA damage caused by disinfectants. Further, the ability to synthesize vitamin B12, an essential 689 690 co-factor, is limited to certain bacteria and archaea and thus the ability to uptake vitamin B12 from the environment is essential for growth⁸³. The higher abundance of vitamin B12 transporters is 691 692 consistent with metagenome level observations that the microbial community in disinfected 693 systems rely more on scavenging from the environment as compared to non-disinfected systems.

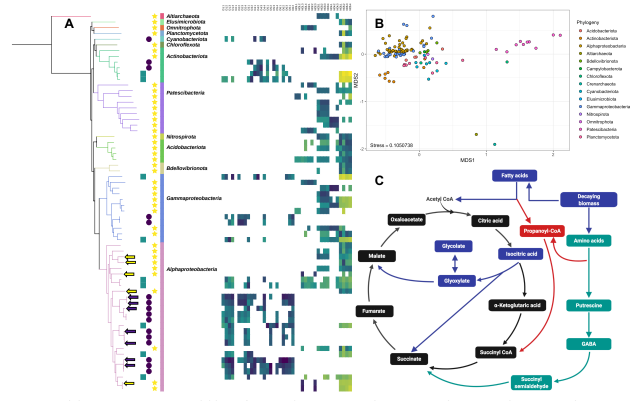


Figure 6: (A) Phylogenomic tree of 66 MAGs classified as D-only (purple circles), ND-only (yellow stars), and both (teal squares) constructed using 48 ribosomal proteins along and their relative abundance (RPKM) in the samples collected from disinfected and non-disinfected systems. RPKM's for MAGs are only reported for samples where 25% of the nucleotides in a MAG were covered by at least one read. (B) Clustering of all MAGs based on their clustering metabolic potential (i.e., completeness of KEGG modules) was primarily drive by phylogeny. (C) The metabolic modules identified as differentially abundant in disinfected systems using metagenome level analyses (Table S12) are shown using teal arrows and squares and those more prevalent in high quality alphaproteobacterial MAGs from D-only (purple arrows - Figure 6A) compared to those from ND-only category (yellow arrows - Figure 6A) are shown using blue arrows and boxes, while red arrows and boxes indicates modules identified as more prevalent in D-only systems using both metagenome and MAG level analyses.

695

696 CONCLUSIONS.

697 To our knowledge, this is the first study to provide metagenomic insights into differences in 698 structure and functional potential of drinking water microbiomes across full-scale drinking water 699 systems that rely on disinfection (i.e., disinfected) or nutrient limitation (i.e., non-disinfected) to 700 manage microbial growth. Understanding the microbial implications of these two microbial 701 growth control strategies is essential to not only develop a better understanding of ecological and 702 metabolic traits guiding community level processes in these system, but is also critical for 703 providing a community-level context to the microbiological safety in either type of drinking water 704 system. In this study, we show that disinfection exhibits consistent, systematic, and significant 705 association with drinking water microbiome at the membership, structure, and functional potential 706 at the metagenome and MAG levels, irrespective of the drinking water system under consideration 707 (e.g., source water type, treatment process, etc.). In doing so, we also identify key metabolic traits 708 associated with carbon and nitrogen metabolism that are over represented in bacteria in disinfected 709 systems compared to non-disinfected systems. This suggests that the influence and efficacy of 710 disinfection on the drinking water microbiome may not simply be associated with differential 711 disinfection resistance⁸⁴, but may also expand to other metabolic traits that include the use of 712 carbon and nitrogen sources made available via microbial inactivation and its regulation. It is 713 important to note that while the impact of disinfection on microbial community structure and 714 functional potential is clear, the metabolic traits identified in this study provide a hypothesis to 715 support future experimental work that will be required to validate the findings of this study.

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